A large scale high-throughput screen identifies chemical inhibitors of phosphatidylinositol 4-kinase type II alpha

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Abstract The minor phospholipid, phosphatidylinositol 4-phosphate (PI4P), is emerging as a key regulator of lipid transfer in ER-membrane contact sites. Four different phosphatidylinositol 4-kinase (PI4K) enzymes generate PI4P in different membrane compartments supporting distinct cellular processes, many of which are crucial for the maintenance of cellular integrity but also hijacked by intracellular pathogens. While type III PI4Ks have been targeted by small molecular inhibitors, thus helping decipher their importance in cellular physiology, no inhibitors are available for the type II PI4Ks, which hinders investigations into their cellular functions. Here, we describe the identification of small molecular inhibitors of PI4K type II alpha (PI4K2A) by implementing a large scale small molecule high-throughput screening. A novel assay was developed that allows testing of selected inhibitors against PI4K2A in intact cells using a bioluminescence resonance energy transfer approach adapted to plate readers. The compounds disclosed here will pave the way to the optimization of PI4K2A inhibitors that can be used in cellular and animal studies to better understand the role of this enzyme in both normal and pathological states.

Phosphoinositides represent a small fraction of all phospholipids, but they regulate a whole range of cellular processes (1). These regulatory lipids are formed by sequential phosphorylation of phosphatidylinositol (PI) on its inositol ring at any of three positions (positions 3, 4, and 5). The different combination of these phosphorylations gives rise to the seven known PI species. There are specific enzymes for phosphorylation of each position and in most cases multiple enzymes catalyze the same reaction. A similar multiplicity of phosphatases exists, setting up an elaborate network of PI metabolism (1). Many of these enzymes have been linked to deregulation ofPIP levels and human diseases. For example, numerous studies have investigated the role of PI3Ks and their lipid product, PI(3,4,5)P3, in cancer and immune regulations promoting the development of PI3K inhibitors currently in clinical trials (2). PI 4-kinases (PI4Ks) phosphorylate the 4-position on the ring and have long been viewed only as enzymes that produce an intermediate for the synthesis of PI(4,5)P2, a plasma membrane (PM) lipid of great significance (1, 3). However, research in the last 10 years revealed that the four distinct PI4Ks make PI 4-phosphate (PI4P) in various intracellular compartments, such as the Golgi and endosomes, where the lipid regulates trafficking of various cargos (3). There is now a better understanding of the cellular functions and importance of type III PI4Ks: PI4KB plays a role in the organization of the Golgi and trafficking through this compartment.

Supplementary key words phosphoinositide • endosome • vesicular traffic

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Abbreviations: BRET, bioluminescence resonance energy transfer; HCV, hepatitis C virus; HTS, high-throughput screening; PI, phosphatidylinositol; PI4K, phosphatidylinositol 4-kinase; PI4K2A, phosphatidylinositol 4-kinase type II alpha; PM, plasma membrane; PI4P, phosphatidylinositol 4-phosphate; qHTS, quantitative high-throughput screening.

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The relevance of PI4Ks has largely increased lately after the discovery that hepatitis C virus (HCV) replication requires the host PI4KA enzyme, whereas several enteropathogenic viruses utilize PI4KB for their replication (7). Based on these discoveries, several pharmaceutical companies are developing inhibitors that target the type III PI4Ks. These efforts have been greatly aided by the fact that type III PI4Ks are structural relatives of PI3Ks, and isoform-specific PI3K inhibitors have already been pursued for treatment of cancer and various inflammatory conditions (8). Interestingly, several drugs that have been in the pipeline to inhibit HCV (9) or enteroviruses (10) without solid knowledge of their mechanism of action turned out to inhibit the type III PI4Ks. It is also notable that various HCV strains vary in their PI4K needs; some use exclusively PI4KA, whereas some others use PI4KB and some enteroviruses already show resistance to drugs that we now know inhibit type III PI4Ks of the host cell (10, 11). These observations suggest that viruses can adapt to utilizing different PI4Ks to generate their replication platform, and mutations will allow them to switch enzymes under selection pressure. Because type II PI4Ks belong to a different family of enzymes with no close relatives within the kinase network, they are not targeted by any of the type III PI4K or PI3K inhibitors (12). Because of their localization and functional properties, these enzymes assume some overlapping functions with the type III PI4Ks, especially PI4KB, and will be available for viruses that need PI3P when type III enzymes are inhibited. Modulators of the type II enzymes would provide a better understanding of their physiological functions and their involvement in human disease, and it might trigger an interest in them as targets for development of therapeutics.

Independently, type II PI4Ks, but not the type III enzymes, were found to be needed for Wnt signaling (13–15). It is not clear how the enzyme contributes to the Wnt signaling cascade. Some studies suggested that PI(4,5)P2 at the PM is important for Wnt signaling (13–15), but our research showed no effect of type II PI4K knockdown on PM PI(4,5)P2 levels (12). However, in recent studies, we identified LR10 as an interacting protein of PI4KA (16). LR10 is a transmembrane protein that is a negative regulator of Wnt signaling, inhibiting signaling from the LR6-Fz receptor complex (17). LR10 cycles between the PM and recycling compartments (our unpublished observation). The LR6-Fz complex also undergoes endocytosis, and it is speculated that it signals from endocytic compartments (18). Given the role of PI4KA in the endocytic network, its role in Wnt signaling could be related to the endocytic processing of the Wnt signaling complex. This raises the possibility that, regardless of its mode of action, targeting PI4KA [and PI4KB, as the two enzymes are almost indistinguishable catalytically (19)] could be an effective way of inhibiting Wnt signaling. Wnt signaling is critical in early development, but, as many other developmentally important signaling cascades, it reactivates and drives certain types of cancer cells (20). In particular, together with K-Ras mutations, Wnt signaling was found to be a determinant for lineage selection in pancreatic ductal adenocarcinoma (21, 22). Wnt signaling was also found to be critical in the bone lesions developing in multiple myeloma (23, 24). Targeting Wnt signaling, therefore, is a priority for several pharmaceutical companies (20, 25) (see also http://pharmastrategyblog.com/2010/09/wnt-signaling-and-cancer.html/).

Other cellular processes that have been linked to type II PI4Ks include Akt activation and cancer cell growth (26), an effect that can also be related to the signaling features of Akt in endosomes (27–29) and the role of PI4KA in endocytic trafficking. Our recent studies suggest that PI4KA has an important role in the delivery of the GBA enzyme (an enzyme whose defective lysosomal activity is the cause of Gaucher disease) to the lysosome (30). Therefore, both positive and negative manipulations of the PI4KA enzyme hold significant promise in understanding and possibly treating a whole range of lysosomal disorders.

Finally, the most recent research unveiled an important role of intermembrane gradients of PI4P, the lipid product of PI4Ks, as a source of energy to drive nonvesicular lipid transport between adjacent membranes in organelle contact sites (31). Several lipid transport proteins were found to bind PI4P as one of their cargoes, and, depending on the organelles involved, the PI4P is provided by distinct PI4K enzymes. This placed PI4Ks in the center of lipid metabolism, and, hence, membrane biogenesis and studies are emerging that suggest that defects in these lipid transport processes may have close links to neurodegenerative diseases, such as ALS (32), Parkinson (33), or demyelinating neuropathies (34).

Based on all of these data linking type II PI4Ks to many physiological processes relevant to a variety of diseases, it would be desirable to find inhibitors for PI4KA, the class of PI4Ks for which inhibitors are not available as yet. A recent study described an inhibitor (PI-273) of PI4KA (35), which showed promise in inhibiting cancer cell growth. That study used docking-based and ligand-based screening strategy to identify PI-273, which was found to interfere with substrate binding or substrate access and not with ATP binding (35). Our studies have used a different strategy, namely, the screening of a small molecule library of compounds using a recombinant PI4KA. The inhibitors identified in our study have a different mode of action, interfering with ATP binding rather than with the binding of the lipid substrate.
energy transfer (BRET) probe was designed to express a Rab7-directed Venus together with an S-luciferase-fused PHP reporter from a single plasmid such that PI4P made on endosomes would excite the endosome-targeted Venus via bioluminescence energy transfer. For this, a pmRFP1C-Sluciferase-P4M2x-T2A-Venus-Rab7 plasmid was created in multiple steps. First, a Venus-Rab7 construct was made using the iRFP-FRB-Rab7 (described in (37)) by replacing the iRFP-FRB part of this construct with Venus (pmVenus-C1) using the NheI-HindIII fragments of the two plasmids. In the second step, the Sluciferase-P4M2x-T2A sequence was created by PCR using a forward primer containing a NheI site and a reverse primer containing the T2A sequence and an AgeI site. The L10-Venus-T2A-Sluciferase-P4M-2x plasmid was used as template after removing an internal NheI site from this original construct (described in (38)). The PCR fragment was cloned into TOPO vector and the NheI-AgeI-digested insert was placed in the mVenus-Rab7 plasmid digested with NheI-AgeI to get the final construct.

Cell culture and transfection

The HEK293-AT1 cell line stably expressing the rat AT1α angiotensin receptor was described previously (12). The cell line was regularly tested for Mycoplasma contamination using InvivoGen mycoplasma detection kit each time after thawing and treating with Plasmocin prophylactic (InvivoGen) at 500 μg/ml for 1 week. The subsequent passages were maintained at 5 μg/ml of Plasmocin. HEK293-AT1 cells were cultured in DMEM (high glucose) containing 10% (v/v) FBS, 100 μg/ml penicillin, and 100 μg/ml streptomycin. COS-7 cells (also tested for Mycoplasma) were cultivated in DMEM containing 10% FBS, 100 μg/ml penicillin, and 100 μg/ml streptomycin. For confocal microscopy, COS-7 cells (200,000 cells/dish in 2 ml) were seeded onto glass-bottom dishes containing 10% (v/v) FBS, 100 μg/ml penicillin, 100 μg/ml streptomycin. For immunoprecipitation of PI4K2A-mRFP, COS-7 cells were precoated with PLL, transfected, and subjected to BRET analysis. For the robotic screening, the ADP-Glo assay using the ADP-Glo™ kinase assay kit from Promega was miniaturized and optimized on 1,536-well white solid bottom plates (Greiner Bio-one). The substrate, 1.2 mM ATP, was prepared in the assay buffer [40 mM Tris (pH 7.5), 20 mM MgCl₂, 1 mM EGTA, 0.2% Triton X-100, 0.5 mM DTT, 0.5 mg/ml BSA] and PI4K2A enzyme, diluted also in the assay buffer to 4 μg/ml, were dispensed onto the plates in equal volumes (1 μl/well), each with 1 μl of kinase detection buffer were added to the plates and incubated for an additional 40 min. All incubation steps were performed at ambient temperature. The resulting luminescence signal was measured on a Viewlux plate reader (PerkinElmer). Adenosine, a partial nonspecific inhibitor, was used as a positive control at 100 μM final concentration to assay the assay’s quality and plate-to-plate reproducibility. We also used “no enzyme” control to gauge the assay’s maximal potential inhibition (IC₅₀).

ADP-Glo assay for testing selected compounds

The ADP-Glo assay was performed using the ADP-Glo™ kinase assay kit from Promega with slight modifications in the total reaction volumes. The enzyme assay for the COS-7-expressed immunosolated PI4K2A enzyme was done in a total volume of 200 μl. PI4K2A-bound Dynabeads (25 μl) suspended in kinase assay buffer were transferred in per well of a 96-well white plate and preincubated with 1 μl of inhibitor candidates (final concentration, 10 μM) at room temperature for 10 min. Following preincubation with inhibitors, 25 μl of kinase assay buffer containing 0.6 μM ultrapure ATP and 1.6 mM P1 were added in each well and the reaction was continued at room temperature for 1.5 h. The reaction was stopped by adding 50 μl of ADP-Glo reagent, and further incubated for 40 min followed by addition of 100 μl of kinase detection reagent and incubation for 40 min in the dark at room temperature. The luminescence was measured using a Tristar2 LB 942 multimode microplate reader (Berthold Technologies).

In situ PI4K2A kinase assay with COS-7 cells

For determining the extent of inhibition on PI4K2A exerted by the candidate inhibitors in COS-7 cells, PI4K2A-mRFP was transfected to COS-7 cells and protein was expressed overnight. The transfected cells were incubated in 1 ml of modified Krebs-Ringer
buffer [120 mM NaCl, 4.7 mM KCl, 0.7 mM MgSO₄, 10 mM HEPES (pH 7.3), 1.802 g D+ glucose, 2 mM CaCl₂, with a final pH adjusted to 7.4] with 1 μM of PI4K2A inhibitors in a 37°C water bath for 20 min. After 20 min, 10 μM of wortmannin was added to each well following reincubation at 37°C for another 10 min. The medium was replaced with 500 μl of permeabilization buffer [110 mM KCl, 10 mM NaCl, 5 mM MgCl₂, 20 mM HEPES (pH 7.3), 2 mM EGTA, 0.05% BSA (v/v)] containing 15 μg/ml of digitonin, 0.3 mM freshly prepared ATP, 1 μM inhibitors, 10 μM of wortmannin, and 15 μl of 10 μCi/ml of P³² labeled ATP. The cells were incubated in a 37°C water bath for 30 min. The reaction was stopped by quickly replacing the permeabilization buffer with 500 μl of ice chilled 1x PBS containing 5% perchloric acid followed by incubation on ice for 30 min. Cells were scraped and transferred to Eppendorf tubes, which were centrifuged at 16,000 g for 5 min at room temperature. The supernatant was discarded, and the pellet was resuspended in 800 μl of 1 N HCl and transferred to 15 ml polypropylene tubes. The phospholipids were extracted following the lipid extraction protocol (40).

**BRET assay**

For measuring the dynamic change of the endosomal PI4P level, vector (Sluc-P4M2X-T2A-Venus-Rab7)-transfected cells were subjected to BRET measurement. After 24 h of transfection, HEK-AT1 cells were rinsed once with modified Krebs-Ringer buffer [120 mM NaCl, 4.7 mM KCl, 0.7 mM MgSO₄, 10 mM HEPES (pH 7.3), 1.802 g D+ glucose, 2 mM CaCl₂, with a final pH adjusted to 7.4]. Cells were preincubated in 50 μl/well Krebs-Ringer buffer for another 30 min (1 min/cycle). Subsequent BRET measurements were done at room temperature. The cells were first subjected to a baseline measurement for 4 min (1 min/cycle) with 40 μl of luciferase substrate coelenterazine h added per well followed by addition of 10 μl/well Krebs-Ringer buffer containing DMSO or candidate inhibitors (final concentration 30 or 10 μM), and BRET measurement was conducted for 10 min. After 10 min, 10 μl of Krebs-Ringer buffer containing DMSO or A1 (final concentration 10 mM) were added per well, and BRET measurements were continued for additional time as indicated (1 min/cycle). In experiments requiring pretreatment with A1, following baseline measurements for 4 min, further measurement was conducted initially with A1 for 30 min and then with inhibitors mixed in 10 μl of Krebs-Ringer buffer for another 30 min (1 min/cycle). For BRET analysis of endosomal PI4P levels, BRET values were calculated from measurements after adding coelenterazine h. A Tri-star2 LB 942 multimode microplate reader (Berthold Technologies) equipped with 540/40 nm (Venus fluorescent measurement) and 475/20 nm (luciferase measurement) emission filters was used for BRET. Because of the robustness and reproducibility of this assay, testing the individual compounds for screening purposes was done in a single experiment, but the NC03 compound was always included as a positive control. Assays for compounds that showed inhibition were repeated one more time to confirm their activity.

**Molecular docking studies**

Crystal structure of PI4K2A was retrieved from the Protein Data Bank (4HNE). Docking of small molecule inhibitors to the active site of protein was performed using the MOE dock program (https://www.chemcomp.com/MOE-Structure_Based_Design.htm). The ligand-induced fit protocol was applied and the binding affinity was evaluated using the GBVI/WSA score. The best binding model with the lowest binding free energies was further energetically minimized using the MOE program. The PyMOL program was used to make the final graphs.

**RESULTS**

**High-throughput screening for the discovery of small molecule PI4K2A inhibitors using a PI4K2A activity assay**

The ADP-Glo assay described earlier by the Tai laboratory (41) was used as a starting point to develop a miniaturized version for high-throughput screening (HTS) applications. To implement the HTS screen, the PI4K ADP-Glo™ kinase assay (Promega, V9102) was miniaturized to a 1,536-well format to enable a large scale dose response quantitative HTS (qHTS) using an automated robotics platform. For the optimization of the assay, reagent concentrations, addition volumes, and reaction/incubation times were modified to obtain a Z-factor of >0.5 (see Fig. 1). The optimized 1,536-well plate assay protocol is shown in Table 1. Following establishment of 1,536 assay conditions, the qHTS assay was further validated by screening the LOPAC®1280 (Sigma-Aldrich) collection of biologically active compounds.

![Fig. 1. Summary of LOPAC collection validation screening with the recombinant PI4K2A enzyme produced in bacteria using ADP-Glo assay. A: Scatter plots of plates treated with either DMSO (left) or compounds at the highest dose (50 μM) (right) on one single 1,536-well plate. Each plate contains 48 columns (x axis), each column contains 32 wells. Different compounds were added to each well in columns 5–48 as a single point at a single concentration. The seven concentrations of the drugs were tested in plates C1–C7. The controls were added onto columns 1–4 as follows: columns 1 and 4, DMSO (EC0); column 2, no enzyme (EC100); and column 3, adenosine at 100 μM. B: In each of the plates, including those where columns 5–48 were treated with inhibitors, the DMSO (EC0) and no enzyme (EC100) values were used to calculate the signal to background (S/B) ratios (open circles) and calculate the Z-factor (closed circles). C: Dose response curves of selected robust inhibitors.](https://www.chemcomp.com/MOE-Structure_Based_Design.htm)
Seven concentrations of each compound were assayed from 4.8 nM to 76 μM, with 5-fold dilutions (plates C1–C7, each containing a single concentration of the inhibitors). One assay plate with plain DMSO was used to determine the assay robustness in the absence of compounds (Fig. 1A). A total of eight assay plates were tested in the LOPAC pilot screen and the median assay parameters per plate were 10-fold signal-to-background ratio and the median \( Z' \)-factor was 0.85 calculated from the DMSO (EC0) and no enzyme (EC100) columns in each plate (Fig. 1B), thus further validating the reliability of this assay in 1,536-well format. Figure 1A shows scatter plots for two assay plates from the LOPAC screen, one testing a plate with DMSO and another with the highest dose of LOPAC compounds (single measurement for each compound). Using the curve class classification algorithms developed at the National Center for Advancing Translational Sciences Chemical Genomics Center for hit selection from dose response HTS (42), 17 compounds were determined to be high quality inhibitors of the PI4K ADP-Glo assay (>1% hit rate). The assay was also able to detect 11 compounds as potential activators of PI4K2A (not shown).

### A HTS identifies PI4K2A inhibitor candidates

A HTS was then performed with ~400,000 compounds from small molecule diversity collections. Sytravon, NPC, and MLPCN collections were screened at the top two doses (76 and 15 μM final concentration) using a bacterially expressed human PI4K2A enzyme (39). Based on the screening results, we identified ~580 compounds with >50% inhibitory activity. These compounds were retested at seven doses in 1:3 serial dilution to confirm their activity. Further, they were counter-screened with ADP-Glo™ reagents in the absence of the enzyme to exclude artificially luminescent compounds, and with PI4KB, another structurally unrelated PI4K. The list was further narrowed by eliminating compounds that were known inhibitors of protein kinases or were deemed structurally unsuitable for further development, yielding 14 inhibitors shortlisted for further studies. The workflow of HTS and confirmation is shown in Fig. 2. The structures of confirmed hits are shown in Fig. 3.

### PI4K2A inhibitors also inhibit the mammalian expressed palmitoylated enzyme

The kinase activity of the human PI4K2A is largely increased when a cysteine-rich sequence (174-CCPCC-178) within the kinase domain is palmitoylated in the vertebrate enzyme providing strong membrane association (44). The palmitoylation was absent in the bacterially expressed enzyme used for the high-throughput screen, as the E. coli lacks the necessary enzyme needed for palmitoylation. Hence, we tested all of the 14 candidates using a human PI4K2A expressed and isolated from COS-7 cells. [In earlier studies, we showed palmitoylation of the expressed GFP-fused enzyme (45).] For this, PI4K2A-mRFP was expressed in COS-7 cells and immunoprecipitated in a two-step pull down assay using a biotin-conjugated anti-mRFP antibody along with anti-streptavidin-conjugated Dynabeads.
The precipitated beads capturing the PI4K2A enzyme were tested for kinase activity using the in vitro ADP-Glo enzyme assay along with no enzyme or no substrate controls. The results showed a strong correlation between the potency of inhibitors between the enzymes isolated from COS-7 cells and bacteria, indicating that the candidates can inhibit the palmitoylated enzyme in a same manner as well (Fig. 4A, B).

**Inhibitory effect of the compounds in in situ kinase activity assays**

To evaluate the effect of the candidate inhibitors on the enzyme within the cells, in situ kinase assays were performed in permeabilized COS-7 cells using radiolabeled γ-32P-labeled ATP followed by phosphoinositide extraction, TLC analysis, and quantification. To eliminate the contribution of the activity of PI4KA and PI4KB enzymes that generate a large portion of PI4P within the cells, treatment of the cells with 10 μM of wortmannin for 10 min was used before treatment with the candidate inhibitors (10 μM) for 30 min in cell permeabilization buffer to ensure proper penetration of the inhibitors and ATP. Figure 5A shows a representative TLC from one of these experiments and the summary of two experiments is shown in Fig. 5B.

This analysis indicated a substantially lower potency of the compounds than what was observed with the isolated proteins and also revealed that some inhibitors also inhibited the labeling of phosphatidic acid, suggesting that they inhibited DG kinase activities. Based on these results, two compounds, code-named NC02 and NC03 (see Fig. 3), were selected for further investigations.

**The PI4P pools in Golgi and endosomes decrease upon treatment with NC03**

The effects of the two selected inhibitors were then analyzed in live cell confocal imaging of COS-7 cells transfected with GFP-P4M, a reporter showing the different cellular PI4P pools (37). PI4P is produced by different PI4Ks in specific intracellular compartments and, based on previous studies on the localization of the enzyme (6), PI4K2A was expected to be primarily responsible for the endosomal pool and partly responsible for the Golgi pool of PI4P. As shown in Fig. 6, 10 μM of NC03 elicited a rapid reduction in the Golgi and endosomal PI4P pool. However, NC02 had no noticeable effects (not shown).

**Development of a BRET assay to monitor PI4P generation on endosomes**

Quantification of the changes in the PI4P level in the endosomes from imaging data is challenging because of...
the heterogeneity of cell expression of the reporter and the requirement for the quantitation from a high number of cells to have a robust signal. This method, therefore, is not practical for the analysis of a high number of compounds for quantitative comparisons. The high content cell imaging assay was used to assess the ability of the compounds to modulate endosomal and Golgi-associated GFP-P4M signals, but it was not robust enough to recognize and quantify the signals reliably in these compartments. A BRET assay was therefore developed to measure the PI4P pool in the various endosomes in a live cell-based assay to enable the testing of the effects of the compounds and their dose-dependence for determination of potency and efficacy. Initial BRET measurements were designed to monitor PI4P levels in Rab7-positive late endosomes. For this, we coexpressed the tandem P4M domain of the Legionella SidM protein fused to the Renilla luciferase and the Venus fluorescent protein fused to Rab7. To achieve an equal ratio of coexpression, the luciferase-fused probe and the Rab7-tagged Venus were expressed from a single plasmid separated by the viral T2A peptide sequence. The presence of PI4P in Rab7 endosomes brings the luciferase in proximity to the Venus protein and, in the presence of the coelenterazine substrate, energy-transfer takes place that can be

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**Fig. 5.** Effect of the inhibitory compounds on PI4P and PI(4,5)P2 synthesis tested by an in situ kinase assay. A: Picture of the TLC plate from a representative experiment showing the 32P-labeled lipids. COS-7 cells were permeabilized and treated with respective inhibitors prior to adding [32P]ATP. The results show reduction in PI4P and PI(4,5)P2 labeling upon treatment with 10 μM of NC02 or NC03. Note the effect of NA045 on phosphatidic acid (PA) labeling indicating an off-target inhibitory effect on diacylglycerol kinases. B: Quantification of total PI4P and PI(4,5)P2 from three (gray columns) and two (black columns) independent experiments performed in duplicate. Mean values ± SEM or range are shown.
monitored in a plate reader giving a measurement from whole cell population (46). To increase the availability of the P4M2x-SLuc component in the cytosol, we used an inhibitor of the PI4KA enzyme, which causes elimination of the large PM pool of PI4P, making more of the lipid probe available in the cytosol to find the endosomal PI4P pools (Fig. 7). It is important to note that the PI4KA inhibitor, A1, does not inhibit PI4K2A or PI4K2B (47). Similar experiments performed in identical assays designed for other endosomal pools (namely, Rab4, Rab5, and Rab11) showed very small, if any, PI4P present in those endosomes and that genetic inactivation of PI4K2A eliminated ~80% of the Rab7-associated PI4P assessed by this method (T. Baba et al., unpublished observations). These data have suggested that the BRET assay assessing the Rab7-associated PI4P was a suitable platform to test PI4K2A inhibitors. After verifying the method, all 14 inhibitors were tested using this assay. The results indicated that the NC03 compound decreased PI4P production in the Rab7 compartment by 40–50%. In spite of its potency in the enzyme assays, the NC02 compound failed to show any inhibition in intact cells. This suggested that this compound may have a problem mononuclear in a plate reader giving a measurement from whole cell population (46). To increase the availability of the P4M2x-SLuc component in the cytosol, we used an inhibitor of the PI4KA enzyme, which causes elimination of the large PM pool of PI4P, making more of the lipid probe available in the cytosol to find the endosomal PI4P pools (Fig. 7). It is important to note that the PI4KA inhibitor, A1, does not inhibit PI4K2A or PI4K2B (47). Similar experiments performed in identical assays designed for other endosomal pools (namely, Rab4, Rab5, and Rab11) showed very small, if any, PI4P present in those endosomes and that genetic inactivation of PI4K2A eliminated ~80% of the Rab7-associated PI4P assessed by this method (T. Baba et al., unpublished observations). These data have suggested that the BRET assay assessing the Rab7-associated PI4P was a suitable platform to test PI4K2A inhibitors. After verifying the method, all 14 inhibitors were tested using this assay. The results indicated that the NC03 compound decreased PI4P production in the Rab7 compartment by 40–50%. In spite of its potency in the enzyme assays, the NC02 compound failed to show any inhibition in intact cells. This suggested that this compound may have a problem.
entering the cells, an assumption that was tested in further analysis (see below). Several modifications of the original NC03 compound failed to improve its potency or efficacy. The trimethoxy substitution in the bending phenyl ring seemed to be particularly important. All attempts to reduce or change the number of substituents in this ring resulted in a reduction and/or elimination of potency (supplemental Fig. S1). Importantly, there was no structural similarity between PI-273 (35) and any of the active compounds identified in this study.

Modifying NC02 to increase its cell penetration

The NC02 hit consists of three parts: a modified coumarin core, a 5-hydroxytryptophane, and a short acyl linker connecting the two. In the course of the project, the coumarin core was altered ranging from slight modifications (e.g., shortening or elongating its alkyl sidechain) to more substantial modifications, such as introduction of a different heterocycle of similar size (e.g., quinolinone or quinoxazolione). The 5-hydroxytryptophane was substituted with other amino acids or their residues (e.g., tyrosine, tryptophane, tryptamine), and the connecting linker was modified in length (one to five atoms) and type of connection at either of the ends (amide, amine, ether, etc.). Over 60 new compounds were prepared; however, none surpassed the parent NC02 compound in biological activity (a selected group is shown in supplemental Fig. S2). Incidentally, during these efforts to improve the lead structure, we prepared a few simple esters (methyl, ethyl, isopropyl) and amides (methyl, dimethyl) of the NC02 hit. Although these compounds proved to possess similar activity in the soluble enzyme assays, due to the lack of the negative charge of the carboxy function, they were found superior in terms of cell penetration (Fig. 8). The preparation of esters was performed very similarly to their original template, only by using different esters of 5-hydroxytryptophane. Amides were prepared by the treatment of a corresponding methylester with methylamine or dimethylamine.

DISCUSSION

This study was aimed at identifying small molecular inhibitors of PI4K2A. The role of this enzyme and its sister enzyme, PI4K2B, in cellular functions is the least understood among the four PI4K enzymes. While most studies found PI4K2A responsible for the majority of the endosomal PI4P pools (48) based on its structural similarities to PI4K2B (19), a functional redundancy between these enzymes likely takes place. A small molecule inhibitor would most probably target both enzymes allowing analysis of their cellular functions under acute inhibition, rather than after their prolonged genetic inactivation.

Our efforts have identified compounds that can be potential starting points for further studies that optimize them. In spite of considerable efforts, we could not modify any of our best hits to increase their potencies and, therefore, we did not pursue further studies at this stage to analyze the specificity of these hits against the larger kinome.

Clearly, these inhibitors are specific among the PI4Ks to the type II isoforms, but they may hit other kinases. For the same reasons, we did not attempt to perform any biological studies until compounds with better potency are found that would be worthy of further analysis. Such efforts are being pursued in our laboratories.

A major advance in this study was the development of a method that allows the analysis of PI4K2A inhibitors in a cellular context using the BRET principle in a plate reader format. This has proven to be an extremely useful tool to test larger numbers of inhibitors without the need of cumbersome lipid analyses. This tool should facilitate testing potential PI4K2A inhibitors in the future.

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Fig. 8. Modifications in NC02 increased cell penetration resulting in enhancement of the cell potency. BRET analysis was performed as described in the legend to Fig. 7. Following 4 min of the baseline BRET signal measurements, the cells were treated with NC02 or NC03 (10 μM) (A) or the derivatives NC02-770 and NC02-567 (10 μM) (B) for 15 min. Cells were then treated with A1 (10 nM) and the BRET signal was measured for 30 min. DMSO was used as a control. Note the lack of effect of NC02 in the cellular assay in spite of its potent inhibitory effect on the enzyme. C: Chemical modifications improved the cell permeability of NC02. Mean ± SEM is shown from one of two similar experiments performed in triplicate. D: Predicted binding model of NC03 at the active site of PI4K2a. Both the surface representation and the ribbon model are shown. In the ribbon model, the protein is shown in blue, while the key residues interacting with the inhibitor are shown as sticks (green). NC03 bound at the active site is depicted as sticks. Three hydrogen bonds are predicted to be formed with residues S134, F263, and V264 at the active site.
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REFERENCES


