Pharmacological and Genetic Targeting of the PI4KA Enzyme Reveals Its Important Role in Maintaining Plasma Membrane Phosphatidylinositol 4-Phosphate and Phosphatidylinositol 4,5-Bisphosphate Levels*

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Background: PI4KA is a critical host factor for replication of hepatitis C virus in liver and a potential therapeutic target.

Results: PI4KA inhibitors prevent the maintenance of PtdIns(4,5)P2 pools during strong PLC activation.

Conclusion: PI4KA plays a critical role in maintaining plasma membrane phosphoinositide pools.

Significance: Safe pharmacological targeting of PI4KA is not feasible.

A kinase essential for hepatitis C virus replication and hence is a target for drug development. PI4KA has also been linked to endoplasmic reticulum exit sites and generation of plasma membrane phosphoinositides. Here, we developed highly specific and potent inhibitors of PI4KA and conditional knock-out mice to study the importance of this enzyme in vitro and in vivo. Our studies showed that PI4KA is essential for the maintenance of plasma membrane phosphatidylinositol 4,5-bisphosphate pools but only during strong stimulation of receptors coupled to phospholipase C activation. Pharmacological blockade of PI4KA in adult animals leads to sudden death closely correlating with the drug’s ability to induce phosphatidylinositol 4,5-bisphosphate depletion after agonist stimulation. Genetic inactivation of PI4KA also leads to death; however, the cause in this case is due to severe intestinal necrosis. These studies highlight the risks of targeting PI4KA as an anti-hepatitis C virus strategy and also point to important distinctions between genetic and pharmacological studies when selecting host factors as putative therapeutic targets.

Phosphatidylinositol 4-kinase type IIIα (PI4KA) is a host factor essential for hepatitis C virus replication and hence is a target for drug development. PI4KA has also been linked to endoplasmic reticulum exit sites and generation of plasma membrane phosphoinositides. Here, we developed highly specific and potent inhibitors of PI4KA and conditional knock-out mice to study the importance of this enzyme in vitro and in vivo. Our studies showed that PI4KA is essential for the maintenance of plasma membrane phosphatidylinositol 4,5-bisphosphate pools but only during strong stimulation of receptors coupled to phospholipase C activation. Pharmacological blockade of PI4KA in adult animals leads to sudden death closely correlating with the drug’s ability to induce phosphatidylinositol 4,5-bisphosphate depletion after agonist stimulation. Genetic inactivation of PI4KA also leads to death; however, the cause in this case is due to severe intestinal necrosis. These studies highlight the risks of targeting PI4KA as an anti-hepatitis C virus strategy and also point to important distinctions between genetic and pharmacological studies when selecting host factors as putative therapeutic targets.

Phosphatidylinositol 4-kinases (PI4Ks)3 have long been viewed as enzymes that produce PtdIns(4)P as an intermediate for PtdIns(4,5)P2 synthesis in the plasma membrane (PM). PtdIns(4,5)P2 is one of the most important regulatory lipids in the PM. It is a precursor of inositol 1,4,5-trisphosphate and diacylglycerol, generated by PLC enzymes, and of phosphatidylinositol 3,4,5-trisphosphate produced by PI3Ks. PtdIns(4,5)P2 also serves as a regulator of ion channels and transporters and interacts with proteins that regulate both endocytosis and exocytosis (1). Therefore, PtdIns(4,5)P2 synthesis is essential for cells and so are the PI4Ks that produce PtdIns(4)P to be converted to PtdIns(4,5)P2. However, the identification of four distinct mammalian PI4Ks and subsequent studies on the biology of these enzymes clearly indicated that the different PI4K enzymes and PtdIns(4)P have functions unrelated to PtdIns(4,5)P2 synthesis (2–4). Yeast studies have shown that the Pik1 enzyme, the homologue of mammalian PI4KB, is important for secretion from the Golgi (5, 6), and the Golgi localization of the PI4KB enzyme, as well as the fluorescent reporters detecting PtdIns(4)P, firmly established the importance of this lipid in Golgi function in mammalian cells (7). Less clear is the role of the other type III PI4K, PI4KA. Yeast studies showed that its orthologue, Stt4, is important for the production of PtdIns(4,5)P2 (8), and it was also found that the PI4KA enzyme is responsible for maintenance of the PM PtdIns(4,5)P2 pool in mammalian cells (9, 10). The yeast homologue, Stt4, is localized to the PM and organized into signaling domains (11). Although the mammalian PI4KA is primarily located in the endoplasmic reticulum (2, 12), recent studies revealed that the EFR3 and TTC7 proteins together recruit the PI4KA enzyme to the PM (13). Part of the reasons why the functions of PI4KA have been so difficult to understand is the lack of specific inhibitors of its enzymatic functions. The only way to study the importance of the enzyme was to use siRNA-mediated gene silencing. This procedure takes several days, and assuming critical roles of these enzymes (both Pik1 and Stt4 mutant yeast strains are practically unviable), cells with efficient knockdown may be eliminated, and the surviving cells may have enough enzyme left to support essential func-
tions. In addition, conclusions from siRNA studies may be con-
confounded by functions prescribed to domains found outside the
enzymatic modality, which may be involved in scaffolding or
other protein-protein interactions, particularly in the context of
infection (14).

An unexpected and much needed stimulus in PI4K research
was brought about by the finding that PI4KA is an essential host
factor for hepatitis C virus (HCV) replication, reported in sev-
eral simultaneous studies using RNAi screens (15–19). In
parallel studies, it was also shown that several small RNA
enteroviruses reprogrammed the endoplasmic reticulum–Golgi
trafficking machinery to serve their replication needs and used
another PI4K, PI4KB, in the process (14, 20). These studies
offered new possibilities to counter the establishment of viral
replication machinery using PI4K inhibitors. The important
feature of these studies was that viral replication was almost
completely abolished by even partial PI4K knockdown, which
elicited no apparent cellular toxicity in vitro (21). These
findings have confirmed our previous conclusions that even small
amounts of the enzyme can fulfill its “housekeeping” functions
(9). Because of this difference for the enzyme for viral replica-
tion versus normal cellular functions, some viral studies have
concluded that PI4Ks are not important for the cell, and they
can be targeted without any problem as an antiviral therapeutic
strategy. These conclusions started to be challenged by reports
showing deleterious effects of PI4KA genetic inactivation (22).

In this study, we report on the characterization of a set of
compounds that selectively inhibit PI4KA and interfere with
HCV replication. We show that these compounds inhibit the
synthesis of PtdIns(4,5)P2 in the PM and impair the maintenance
of PtdIns(4,5)P2 levels under strong PLC activation. Curiously,
the potency of these compounds to inhibit purified PI4KA in vitro
and to inhibit PtdIns(4)P synthesis in the PM in cells
shows significant variations raising questions about the ability
of the compounds to reach the relevant cellular compartments
despite similar chemistries. Importantly, the inhibitory effects
on PtdIns(4)P in the PM and on PtdIns(4,5)P2 levels in PLC-
stimulated cells were closely correlated. Toxicity studies in ani-
mals showed that the most potent small molecule inhibitors of
PtdIns(4)P synthesis and PtdIns(4,5)P2 maintenance caused
sudden death when applied at high doses with symptoms remi-
niscent of cardiovascular collapse. These may reflect the ability
of the compound to inhibit PtdIns(4,5)P2 maintenance during
G1-coupled receptor signaling that is essential for maintaining
vascular tone. Finally, genetic inactivation of the PI4KA enzyme
in adult animals with a tamoxifen-induced conditional knock-
out mouse caused a lethal gastrointestinal phenotype that was
different from the acute drug-induced toxicity. These differ-
cences will require further studies to be fully understood but
highlight the need for both types of approaches to anticipate the
results of pharmacological interventions on the biology of
whole animals.

EXPERIMENTAL PROCEDURES

Materials—Angiotensin II (human octapeptide) was from
Bachem (Torrance, CA). Wortmannin was purchased from
Calbiochem. Allother chemicals were of the highest analyt-
ic grade. [γ-32P]ATP (6000 Ci/mmol) was purchased from
PerkinElmer Life Sciences. myo-[3H]Inositol (30–80 Ci/mmole)
was from Amersham Biosciences and American Radiolabeled
Chemicals (St Louis, MO). The monoclonal anti-HA antibody
(HA.11) was from Covance; the PtdIns(4)P antibody was from
Echelon (Salt Lake City, UT; catalogue no. Z-P004), and the
rabbit polyclonal anti-GST was from Millipore (Billerica, MA;
catalogue no. AB3282).

Synthesis of PI4KA Active Compounds—A1 was synthesized
as described in Ref. 23, and the synthesis of C1, F1, H1, and J1
was according to Ref. 24. The synthesis of M1 was described in
Ref. 25, and G1 was synthesized according to the procedure
detailed in Ref. 26.

Transfection of Cells for Microscopy—COS-7 cells or
HEK293-AT11 cells (a HEK293 cell line stably expressing the rat
AT1a angiotensin receptor) were used. Cells (50,000 cells/well)
were plated onto 25-mm-diameter circular glass coverslips in
6-well plates, and plasmid DNAs (0.5–1 μg/well) were trans-
fected with the PLC61-PH-GFP or Tubby domain-GFP fusion
constructs (27) using the Lipofectamine 2000 reagent (Invitro-
gen) and Opti-MEM (Invitrogen) following the manufacturer’s
instructions.

Live Cell Imaging—After 20–24 h of transfection, cells were
washed on glass coverslips with a modified Krebs-Ringer solu-
tion, containing 120 mM NaCl, 4.7 mM KCl, 1.2 mM CaCl2, 0.7
mM MgSO4, 10 mM glucose, 10 mM Na-Hepes, pH 7.4, and the
coverslip was placed into a metal chamber (Atto, Invitrogen)
that was mounted on a heated stage (35 °C or room tempera-
ture for HEK293-AT1 cells). Cells were incubated in 1 ml of the
Krebs-Ringer buffer, and the stimuli were dissolved and added
in 200 μl of warm buffer removed from the cells. Cells were
examined in an inverted microscope. Confocal images were
obtained with a Zeiss LSM 510-META laser confocal micro-
scope (Carl Zeiss Microlmaging, Inc.) using a 63× oil-immern-
sion objective equipped with an objective heater (Biotech).

Measurements of Receptor-stimulated PtdIns(4,5)P2 Kinetics
Using Tubby Domain Translocation—HEK293-AT1 cells were
transfected with the Tubby domain-GFP construct. After 24 h,
cells were mounted in the Atto chambers and imaged in a Zeiss
LSM 510-META confocal microscope at room temperature.
After stimulation with 100 nM AngII, the translocation of the
Tubby domain-GFP fusion construct to the membrane was
monitored in time-lapse imaging. The translo-
cation of the construct from the membrane to the cytosol was
quantified by measuring cytosolic GFP intensity in regions of
interest outside the nucleus and plotted against time using the
Zeiss image processing software. Intensity curves were normal-
ized to prestimulatory values, and they were averaged from a
large number of cells in recordings obtained in several dishes in
multiple independent experiments.

Measurements of Receptor-stimulated PtdIns(4,5)P2 Kinetics
using FRET with the PLC61-PH Domain—HEK293-AT1 cells
were transfected with the CFP- and YFP-tagged PLC61-PH
constructs for 24 h. After transfection, cells were mounted in
the Atto chambers and imaged in an Olympus IX70 microscope
equipped with a Micromax 1024BF (Princeton Instruments)
camera, a Lambda-DG4 illuminator (Sutter), and a beam-split-
ter (Optical-Insights) that contained the filter set for recording
YFP and CFP fluorescence. Data acquisition and processing
were done with the MetaFluor software. The ratios of YFP to
PI4KA Inhibition

CFP for individual cells were calculated normalizing to their initial values before stimulation. Recordings from several cells in each dish and several dishes from separate experiments were pooled and averaged.

Analysis of myo-[\(^{3}H\)]inositol- or \[^{32}P\]P|Phosphate-labeled Lipids—HEK293-AT1 cells plated on 12-well plates (at a density of 30,000 cells/ml) were labeled with myo-[\(^{3}H\)]inositol (20 \(\mu\)Ci/ml) in 1 ml of inositol-free DMEM supplemented with 2% dialyzed FBS for 24 h or with 2 \(\mu\)Ci/ml \[^{32}P\]P|phosphate for 3 h in phosphate-free DMEM supplemented with 2% dialyzed FBS. Cells were treated with various concentrations of the drugs (dissolved in DMSO) for 10 min and stimulated with angiotensin II (100 nM) for the indicated times or left unstimulated. Reactions were terminated by the addition of ice-cold perchloric acid (5% final concentration), and cells were kept on ice for 30 min. After scraping and freezing/thawing, the cells were centrifuged, and the cell pellet was processed to extract the phosphoinositides by an acidic chloroform/methanol extraction followed by thin layer chromatography (TLC) essentially as described previously (10, 28). TLC plates were sprayed with Enhance solution (Perkin-Elmer Life Sciences) and were subjected to autoradiography (TLC) essentially as described previously (10, 28). Densitometric analysis.

In Vitro PI Kinase and PIP 5-Kinase Measurements—Enzymes were prepared from COS-7 cells expressing the respective kinases epitope-tagged with an HA, FLAG, or Myc tag at their N termini. Proteins were immunoprecipitated from the cell lysates and after several washes, their activity was measured on agarose beads. The activities of PI4Ks were measured as incorporation of radioactivity from \[^{32}P\]ATP into organic solvent-extractable material (32). The standard reaction mixture for PtdIns 4-kinase (50 \(\mu\)l final volume) contained 50 mM Tris/HCl, pH 7.5, 20 mM MgCl\(_2\), 1 mM EGTA, 1 \(\mu\)M PtdIns(4,5)P\(_2\), 4 mM ATP, and 0.8 \(\mu\)g/ml anti-PI4K antibody in the continued presence of anti-GST in the reaction mixture.

Immunostaining for PM PtdIns(4)P and PtdIns(4,5)P:\_2—A modified protocol (29, 30) was used. COS-7 cells were grown to \sim 50% confluence on poly-L-lysine-coated 8-well test slides and then treated for 10 min with the indicated concentration of PI4K inhibitor in 0.1% DMSO/phenol red-free DMEM in a total volume of 25 \(\mu\)l at 37 \(^\circ\)C, 10% CO\(_2\) in a humidified atmosphere. The cells were then rapidly fixed by the addition of a further 25 \(\mu\)l of PIPES buffer (137 mM NaCl, 2.7 mM KCl, 20 mM Na-PIPES, pH 6.8) containing 0.4% (v/v) formaldehyde and 0.4% (w/v) glutaraldehyde and incubated for 10 min at room temperature before placing the cells on ice for all subsequent steps. Autofluorescence was quenched by three 2-min washes in freshly prepared 1% (v/v) sodium borohydride in PIPES buffer. Cells were then stained for 30 min sequentially in ice-cold blocking buffer (PIPES buffer supplemented with 0.5% saponin and 5% normal goat serum) containing 1.25 \(\mu\)g/ml anti-PtdIns(4)P IgM and 50 \(\mu\)g/ml GST-P-PLC\(_{\delta 1}\), followed by 2.5 \(\mu\)g/ml rabbit polyclonal anti-GST in the continued presence of anti-PItdIns(4)P. After two rinses in ice-cold PIPES buffer, cells were stained for a further 30 min in ice-cold blocking buffer containing 5 \(\mu\)g/ml each of AlexaFluor647-conjugated anti-rabbit IgG and AlexaFluor555-conjugated anti-mouse IgM, as well as 5 units/ml AlexaFluor488-conjugated phalloidin (Life Sciences). Finally, cells were rinsed four times in ice-cold PIPES buffer, post-fixed for 10 min in ice-cold 4% (v/v) formaldehyde in PIPES buffer, rinsed three times in PIPES buffer, once in water, and mounted in Pro-Long Gold containing DAPI (Life Sciences).

Images were acquired on a Zeiss LSM 780 using a 20 \times 0.8 NA air objective with the confocal pinhole fully opened to capture fluorescence from the entire depth of the cells. For quantitative analysis, the images were automatically analyzed by using an automated pipeline that used the open access package CellProfiler (31) and by using a pipeline that used DAPI and phalloidin staining to identify and segment individual cells, as described previously (30). For each experiment, background fluorescence intensity (measured for cells whereby primary lipid probes were omitted) was subtracted. Individual intensity values were then normalized to the mean intensity value for the 0.1% DMSO control condition. These normalized intensity values were pooled to create the grand mean from independent experiments. Data were plotted in Prism 5 (Graphpad), and fit with the function shown in Equation 1,

\[ I = I_{\text{min}} + \frac{(I_{\text{max}} - I_{\text{min}})}{1 + 10^{x - \log IC_{50}}} \]  

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where \(I\) is the normalized mean pixel intensity, and \(x\) is \log[\text{inhibitor}].

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where \(I\) is the normalized mean pixel intensity, and \(x\) is \log[\text{inhibitor}].
6-kb long homology arm were placed 5′ and 3′, respectively, of a loxP/FRT-flanked neomycin resistance cassette in the 3′–5′ orientation, together with the “target” region containing a further loxP element positioned 3′ to exon 48. Homologous recombination in neomycin-resistant ES cells was confirmed at the 5′ and 3′ ends by Southern blot analysis using NsiI-digested ES cell-derived genomic DNA and probes external to the homology arms, and the presence of the 3′ loxP element was confirmed by PCR (data not shown). Correctly targeted ES cell clones were subsequently injected into C57BL/6-derived blastocysts. Resultant male chimeras were crossed with C57BL/6 female mice carrying an Flp recombinase transgene to remove the neomycin resistance cassette, and the resulting mice heterozygous for the Pi4ka conditional allele were bred to wild-type mice to remove the Flp recombinase transgene. Standard breeding procedures were then used to generate study populations of mice homozygous for the Pi4ka conditional allele and carrying the R26MCM transgene. The latter transgene expresses the MerCreMer cDNA (34) from the Rosa26 locus (35) and enables tamoxifen-inducible, ubiquitous Cre recombinase-mediated deletion of the floxed region of conditional alleles in mice. Littermates homozygous for the conditional Pi4ka allele were used as controls.

To induce MerCreMer recombinase activity, study mice were dosed orally once daily for 5 consecutive days with tamoxifen (1 mg/5 g body weight) dissolved in corn oil; control groups were dosed likewise with corn oil. A standard washout period of 17 days was applied to all recombinase activation studies to
**TABLE 1**

Inhibitory potencies of selected PI4KA inhibitors on PI3K and PI4K used in this study as measured with ADP-Glo or by \( [\gamma-^{32}\text{P}]\text{ATP} \) in PI kinase assays (boldface).

| Compound | PI4KA IC\(_{50}\) [-log(M)] | PI4KB IC\(_{50}\) [-log(M)] | PI3KA IC\(_{50}\) [-log(M)] | PI3KB IC\(_{50}\) [-log(M)] | PI3KG IC\(_{50}\) [-log(M)] | PI3KD IC\(_{50}\) [-log(M)] | PI4K2A IC\(_{50}\) [-log(M)] | PI4K2B IC\(_{50}\) [-log(M)] |
|----------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| A1       | 9.8/8.5                  | 7.7/7.2                  | 7.3                      | 7                        | 7.8                      | 7.1                      | \(<5\)                    | \(<5\)                    |
| C1       | 9.1/8.8                  | 7.2/6.8                  | 8.1                      | 7.1                      | 8.2                      | 8.6                      | \(<5\)                    | \(<5\)                    |
| F1       | 8.0/7.8                  | 5.9/6.0                  | 5.8                      | 5.9                      | 5.9                      | 6.4                      | \(<5\)                    | \(<5\)                    |
| G1       | 8.7/8.6                  | 5.9/6.2                  | 7.2                      | 7.6                      | 7.5                      | 8.0                      | \(<5\)                    | \(<5\)                    |
| H1       | 7.7/7.5                  | 7.6/7.6                  | 5.9                      | 5.8                      | 5.9                      | 5.7                      | \(<5\)                    | \(<5\)                    |
| Wm       | 6.4                      | 6.5                      | ND                       | ND                       | ND                       | ND                       | ND                       | ND                       |

**FIGURE 2. Inhibition of HCV replication by inhibitors of PI4KA.** HCV replication was tested as described previously (33). A shows a tight correlation between the potencies of drugs against PI4KA and inhibition of HCV replication using two different HCV strains (r\(^2\) = 0.758, p < 0.0001, and r\(^2\) = 0.797, p < 0.0001, for HCV1A and HCV1B, respectively). B shows that no correlation was found with the PI4KB potencies and HCV replication with the same set of compounds (r\(^2\) = 0.149, p = 0.114, and r\(^2\) = 0.057, p = 0.341, for HCV1A and HCV1B, respectively). Compounds selected for future analysis are labeled in A.

allow for any initial effect of dosing tamoxifen and corn oil to subside. All animal procedures were reviewed and approved by the GlaxoSmithKline Animal Care and Use Committee and were performed in accredited facilities in accordance with institutional guidelines and the Guide for the Care and Use of Laboratory Animals (Institute of Laboratory Animal Resources, National Research Council).

**Animal Studies Using PI4KA Inhibitors**—A 14-day oral toxicity study was performed in mice with the F1 compound. The compound was formulated as a solution in 30% solutol, 70% polyethylene glycol and administered to mice at a dose volume of 10 ml/kg/day. It was given to male mice (8/group) at doses of 3, 10, \(^4\) 20, and 40 \(^4\) mg/kg/day twice daily and 6 h apart for 7\(^4\) or 14 days by oral gavage. These doses were based on the in vitro protein adjusted EC\(_{50}\) (0.16 \(\mu\)M) and a preliminary dose escalation study at doses of 3, 10, and 50 mg/kg (data not shown). All living animals were humanely euthanized upon completion of the study, and an exhaustive histopathology was performed, and sections were examined for any abnormalities.

**RESULTS**

**Synthesis of Potent PI4KA-selective Compounds of Different Chemotypes**—High throughput screening for inhibitors that inhibit HCV replication and also potently inhibit PI4KA has been the starting point in the generation of more selective PI4KA inhibitors (see Ref. 33 for more details). The first iteration of these compounds showed potent ATP-competitive inhibition of PI4KA but little selectivity against PI4KB and class I PI3Ks (33). Several rounds of structure-activity relationship studies using compounds with chemical substitutions in the original hits yielded inhibitors with high potency and selectivity (33). These were evaluated for their ability to inhibit PI4KA (using ADP-Glo assay and HCV replication. These compounds were also tested against a whole set of PI kinases (Table 1) and subjected to KiNativ in situ kinase profiling to determine their effects on protein kinases. These analyses showed that several compounds had good selectivity and high potency against PI4KA with no effect on protein kinases. The antiviral effect showed very close correlation with the inhibitory potency against PI4KA (\(r^2 = 0.758, p < 0.0001\), and \(r^2 = 0.797, p < 0.0001\), for HCV1A and HCV1B, respectively) (Fig. 2A). In contrast, the potencies of the same compounds against PI4KB showed no correlation with HCV inhibition (Fig. 2B). A selected set of compounds (Fig. 3) was further tested in conventional PI4K lipid kinase assays based on PtdIns phosphorylation using \(^{32}\)P-labeled ATP and epitope-tagged PI4Ks expressed in COS-7 cells and immunoisolated. These experiments confirmed the IC\(_{50}\) values determined in the nonradioactive assays for PI4KA and PI4KB (Fig. 4 and Table 1). The compounds were also tested on the type II PI4Ks and on the three major forms of PIP.

\(^4\) Because of low exposures on day 1, the 10 mg/kg/day dose was raised to 40 mg/kg/day on days 8–13.
5-kinases (α, β, and γ) and were found to have no effect at a concentration of 10 μM (Table 1).

Inhibition of PI4KA Limits PtdIns(4)P Production in the PM—Next, we moved to intact cells to test the effects of a selected set of compounds on PtdIns(4)P levels using HEK293 cells stably expressing the AT1 angiotensin receptors (HEK-AT1). We used [3H]inositol labeling for 24 h to label the cells stably expressing the AT1 angiotensin receptors (HEK-AT1). We used [3H]inositol labeling for 24 h to label the cells stably expressing the AT1 angiotensin receptors (HEK293-AT1 cells and stimulation with angiotensin II (AngII). We used several approaches to follow PtdIns(4,5)P2 reporters (37). As shown in Fig. 7A, in 32P-labeled cells pretreatment with 100 nM A1 for 10 min completely depleted PtdIns(4,5)P2 prior to AngII stimulation when compared with untreated control cells. As observed before, control cells showed a typical rapid depletion of PtdIns(4,5)P2 upon stimulation. There was no effect of AngII treatment on the already depleted PtdIns(4,5)P2 pools in cells pretreated with the A1 compound (Fig. 7A). As expected, stimulation with AngII rapidly depleted PtdIns(4,5)P2 followed by a partial replenishment of these pools in control cells (Fig. 7B). In contrast, AngII stimulation depleted PtdIns(4,5)P2 even more completely, and no replenishment was observed in A1-treated cells (Fig. 7B).

The effects of the inhibitors were further tested on PtdIns(4,5)P2 kinetics using different approaches. The PH domain of PLCβ1 and the Tubby domain of the Tubby protein fused to fluorescent proteins have been used as probes for PtdIns(4,5)P2 in the membrane (27). To quantitate the lipid changes, we used both FRET analysis (in the case of the PH domain) (38) or increased intensity of fluorescence in the cytosol followed in confocal microscopy (in the case of the Tubby domain; note the inverted scale of the ordinate!). Both of these measurements showed the transient decrease in the level of PtdIns(4,5)P2 in the PM after AngII stimulation in control cells (Fig. 7, C and D, blue traces) and the severely impaired resynthesis of PtdIns(4,5)P2 in the PM in the A1-pretreated cells (Fig. 7, C and D, red traces). These results collectively showed that under acute and complete inhibition of PI4KA, cells are unable to maintain PtdIns(4,5)P2 levels against a robust PLC activation. However, PtdIns(4,5)P2 levels can be maintained in quiescent cells even when the PM PtdIns(4)P pools are largely reduced.

These data confirmed our earlier observation that PtdIns(4,5)P2 levels can be maintained in quiescent cells at a wide range of PtdIns(4)P concentrations (30). However, our previous studies also showed that wortmannin (Wm) or phenylarsine oxide treatment prevents the resynthesis of PtdIns(4,5)P2 during PLC activation (10, 30, 36). Although these inhibitors are not selective, their effect was attributed to inhibition of PI4KA in those studies. To test if selective PI4KA inhibitors indeed prevent the maintenance of PtdIns(4,5)P2 during PLC activation, we used HEK293-AT1 cells and stimulation with angiotensin II (AngII). We used several approaches to follow PtdIns(4,5)P2. First, we used cells labeled with [32P]phosphate for 3 h prior to stimulation with AngII, a condition we have previously used in these cells (10). Second, we used the PLCβ1-PH domain in confocal studies to monitor PtdIns(4,5)P2 during stimulation. These protein domains are widely used as PtdIns(4,5)P2 reporters (37). As shown in Fig. 7A, in 32P-labeled cells pretreatment with 100 nM A1 for 10 min completely depleted PtdIns(4,5)P2 prior to AngII stimulation when compared with untreated control cells. As observed before, control cells showed a typical rapid depletion of PtdIns(4,5)P2 upon stimulation. There was no effect of AngII treatment on the already depleted PtdIns(4,5)P2 pools in cells pretreated with the A1 compound (Fig. 7A). As expected, stimulation with AngII rapidly depleted PtdIns(4,5)P2 followed by a partial replenishment of these pools in control cells (Fig. 7B). In contrast, AngII stimulation depleted PtdIns(4,5)P2 even more completely, and no replenishment was observed in A1-treated cells (Fig. 7B).

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**FIGURE 3. Structure of PI4KA inhibitors most characterized in this study.**
PI4KA Inhibitors Show a Wide Range of Potencies When Tested in Cellular Settings—Several PI4KA inhibitors of a similar chemotype were then tested to determine their ability to inhibit PtdIns(4,5)P2 synthesis during PLC activation. For this, we used HEK-AT1 cells labeled with [32P]phosphate for 3 h and stimulated with AngII for 10 min after a 10-min pretreatment with DMSO or varying concentrations of the selective PI4KA inhibitors. Because PtdIns(4,5)P2 levels return to about 75% of their initial value by 10 min of stimulation (see Fig. 7) and this replenishment of the pool is almost completely prevented by PI4KA inhibition (Fig. 7, B–D), this protocol provided a sensitive means to assess the potency of the inhibitors. As shown in Fig. 8, a large variation was observed between the potencies of the inhibitors to inhibit PtdIns(4,5)P2 resynthesis. The most potent inhibitor was A1 with an IC50 of about 3 nM, whereas the least active of these drugs was H1 with an IC50 of 30 nM. This analysis was then extended with additional compounds for which the IC50 for PI4KA and PI4KB inhibition was determined with the [32P]ATP-based PI4K assays. When the IC50 values for the in vitro kinase assays were plotted against the IC50 for PtdIns(4,5)P2 inhibition, it was apparent that, except for A1, most inhibitors were significantly less potent in the cellular assay. This finding was partially explained by the fact that even a fraction of the kinase activity can support PtdIns(4,5)P2 synthesis as concluded by earlier studies (10). However, it did not explain the fact that significant differences in cellular potencies were found within compounds that inhibited PI4KA in vitro with almost identical potencies (see J1, M1, and H1). This was not due to additional inhibition of PI4KB, because H1, which inhibited PI4KA and PI4KB equally in vitro and more potently than Wm, was less potent than Wm in the cellular assay. Part of the poor cellular potency must lie in the bioavailability of the compounds in the cellular compartment where they need to act, and indeed, the compounds that deviated the most from their in vitro potencies for PtdIns(4,5)P2 synthesis were also the outliers in their ability to inhibit HCV replication (see Fig. 1). Comparison of the potencies of the compounds on HCV replication and PtdIns(4,5)P2 synthesis also confirmed previous conclusions that viral replication is more sensitive to kinase inhibition, whereas PtdIns(4,5)P2 synthesis can be maintained even at a fraction of the total kinase activity.

Cells Tolerate PI4KA Inhibition for Prolonged Periods without Significant PtdIns(4,5)P2 Depletion—Given the importance of PI4KA in the maintenance of plasma membrane PtdIns(4,5)P2 levels during robust agonist stimulation, we expected that cells would not tolerate prolonged treatments with PI4KA inhibitors. Also, we anticipated that cells growing in culture and subjected to an increased PLC activity and membrane remodeling would eventually get depleted in PtdIns(4,5)P2. However, surprisingly, HEK293 or COS-7 cells treated with A1 (30 nM) for 1

![Inhibition of immunoisolated PI4KA and PI4KB by a selected set of PI4KA inhibitors in vitro.](image-url)

**FIGURE 4.** Inhibition of immunoisolated PI4KA and PI4KB by a selected set of PI4KA inhibitors in vitro. Enzymes were prepared from COS-7 cells expressing the respective kinases epitope tagged with an HA tag at their N termini. Proteins were immunoprecipitated from the cell lysates, and after several washes their activity was measured on agarose beads. The activities of PI4Ks were measured as incorporation of radioactivity from [γ-32P]ATP into organic solvent-extractable material. Activities were expressed as percent of DMSO-treated controls. Means ± range from duplicate determinations are shown from a representative experiment. Blue, PI4KA; red, PI4KB.
PI4KA Inhibitors Vary in Their Adverse Effects in Whole Animals—At this point only a 14-day oral toxicity study was performed in mice to test potential adverse effects of PI4KA inhibition in whole animals. It was performed with the F1 compound, which is the same chemotype as A1 but has been proven to have markedly better pharmacokinetic properties required to evaluate this chemotype in vivo. The compound was formulated as a solution in 30% solutol, 70% polyethylene glycol and administered to mice at a dose volume of 10 ml/kg/day. It was given to male mice (8/group) at doses of 0 (vehicle), 3, 10, 20, and 40 mg/kg/day twice daily and 6 h apart for 7 or 14 days by oral gavage. These doses were based on the in vitro protein adjusted EC90 (0.16 μM) and a preliminary dose escalation study at doses of 3, 10, and 50 mg/kg (data not shown).

One mouse given 3 mg/kg/day and another mouse given 20 mg/kg/day were found dead on days 12 and 8, respectively, with no clinical signs noted. Four mice given 40 mg/kg/day were either found dead or were humanely euthanized in a moribund condition between days 11 and 14. Clinical signs for these mice included decreased activity, loss of skin elasticity, rough coat, eyes partially closed, hunched posture, cold to the touch, slow breathing, loose/watery feces, wet skin/fur, and staining in the urogenital/anal area. One animal had limited use of its forelimbs on day 12 prior to becoming moribund. All of these animals had slight body weight loss from day 1 to day 11 (0.78×) when compared with their pretest bodyweights. Clinical signs in animals that survived to terminal necropsy included loss of skin elasticity, loose/watery feces, decreased activity, rough coat, hunched posture, and brown/orange staining of the anal area. Animals treated with F1 at 40 mg/kg/day exhibited morbidity and mortality and test article-related changes in the stomach, thymus, spleen, and clinical pathology parameters. These included nonglandular gastric lesions present in all animals given 40 mg/kg/day characterized by combinations of erosion/ulceration, squamous epithelial cell hyperplasia and hyperkeratosis, lamina propria edema, a mixed inflammatory cell infiltrate, and occasional lamina propria vessels with medial degeneration/necrosis (data not shown). Individual mice at 40 mg/kg/day also had a gastric ulcer in the pylorus and multifocal foveolar cell necrosis in the glandular stomach. It is important to note that F1 does not cross the blood-brain barrier, which may explains the lack of CNS symptoms.

Several other potent PI4KA inhibitory compounds were then tested in animal studies for adverse effects. They all caused sudden death before any pathology could develop. These included C1, M1, and J1. These animals all appeared to die from a cardiovascular collapse. The toxicity of these compounds correlated with their ability to inhibit PtdIns(4,5)P2 resynthesis in the cellular assay. Because these compounds severely affected Gq signaling based on cellular data, it is expected that they cause pleiotropic effects in whole animals. These will require a lot more detailed analysis, including ruling out off-target effects.

Genetic Ablation of Pi4ka in Adult Animals Causes Lethality—To evaluate the importance of the kinase in whole animal models, cKO mice were generated by targeting exon 48 in the Pi4ka gene. Deletion of this exon generates a catalytically inactive truncated protein. (The targeting strategy is shown in Fig. 1, and a detailed description of the procedure is found under “Experimental Procedures.”) Homozygous mice were bred with a mouse line that carried a ubiquitous tamoxifen-inducible Cre transgene (34, 35) to generate homozygous conditional alleles. These animals were humanely euthanized, and cKO mice were dosed orally once daily for 5 consecutive days with tamoxifen (or treated identically without tamoxifen) followed by a 17-day waiting period.

Pi4ka cKO mice were found moribund by day 7 post-tamoxifen treatment. These animals were humanely euthanized, and histopathology exhibited epithelial cell degeneration/necrosis (Fig. 9) with evidence of regeneration in the stomach, small intestine, and large intestine. Tamoxifen pharmacokinetics showed high levels in the gut and other organs, with little exposure in the CNS (data not shown). No gastrointestinal abnormalities were detected in animals in the corn oil (vehicle) control, tamoxifen-treated (not cKO), and cKO mice not treated
with tamoxifen groups. These findings were similar to those reported in a recent cKO pi4ka mouse study (22).

**DISCUSSION**

These studies were designed to elucidate the effects of PI4KA inhibition in vivo and in vitro. Synthesis of selective PI4KA inhibitors has been motivated by the prospect that such inhibitors might be useful to fight HCV infection. RNAi studies identified PI4KA as a mandatory host factor in HCV replication, and those studies suggested that PI4KA down-regulation is reasonably tolerated by cultured cells (15, 17, 39, 40). In addition, it has been reported that HCV infection can induce the expression of PI4KA and increase PtdIns(4)P levels in humans (39). However, cellular studies using RNAi-mediated knockdown indicated that PI4KA is responsible for the maintenance of the PM PtdIns(4,5)P2 pools during strong PLC activation (10) and predicted that complete blockade of this enzyme may interfere with a number of signaling processes that require the presence of these lipids in the PM. The conclusion drawn from these studies was that cells tolerate a significant level of knockdown of PI4KA because even a small amount of enzyme is able to maintain its housekeeping functions, whereas HCV replication is strongly inhibited even by a 50% reduction of the level of this enzyme (15, 17, 39, 40). This has lent strong support to the notion that there is a therapeutic window for inhibiting HCV replication without affecting cellular functions.

A large number of compounds with various activities against PI4KA did indeed inhibit viral replication, with their antiviral potencies showing close correlation with their PI4KA inhibitory potencies. A selected set of potent PI4KA inhibitory compounds that were subjected to more detailed analysis in cultured cells strongly inhibited PtdIns(4)P synthesis, primarily affecting the PM pool of this lipid. Interestingly, the same treatment had very small if any effect on the PM PtdIns(4,5)P2 pools in quiescent cells. Only when cells were challenged with a strong PLC-activating stimulus did PtdIns(4,5)P2 levels decrease with a greatly diminished ability for these pools to resynthesize. Two recent studies have already noted the dissociation of PtdIns(4,5)P2 from PM PtdIns(4)P. One of these studies showed that PtdIns(4,5)P2 levels can recover after strong PLC activation even when the steady-state level of PtdIns(4)P is greatly reduced by an acutely PM-recruited PtdIns(4)P phosphatase, and only inhibition of PI4Ks impeded PtdIns(4,5)P2 recovery (30). Another study found that mouse embryo fibroblast cells obtained from a different conditional pi4ka knockout mouse showed no signs of PtdIns(4,5)P2 depletion; in fact, these cells appeared to display increased PIP5K activity and accumulation of PtdIns(4,5)P2 in internal membranes (13). This latter feature is highly reminiscent of cells overexpressing PIP5Ks (41) or their activators, such as Arf6 (42). Notably, prolonged pharmacological blockade of PI4KA in this study did not yield the appearance of PtdIns(4,5)P2 in internal membranes in COS-7 cells, and the cells were able to maintain their PM PtdIns(4,5)P2 levels. These observations together suggest that there are additional mechanism(s) by which cells can supply PtdIns(4)P to the PM even when PI4KA is inhibited. Possible sources of PtdIns(4)P could be the endocytic compartments where the lipid is synthesized by type II PI4Ks and where the lipid can reach the membrane during vesicular trafficking as hinted by our previous study (43). Another alternative could be
the production of PtdIns(4,5)P₂ via PtdIns(5)P by the type II PIPKs, a possibility that needs further investigation.

At this stage, the animal studies were only designed to test the toxicity of PI4KA inhibitory compounds during oral administration. These animals displayed a variety of adverse effects in response to the inhibitor. Several animals showed sudden lethality, especially at the highest dose of treatment without any obvious pathology other than mild irritations documented in the upper GI tract. The sudden death suggested a cardiovascular insufficiency that can be caused by dehydoration or a sudden loss of vascular tone. Although one cannot rule out off-target effects, one can speculate that strong activation of Gq-coupled receptors (such as those required to maintain vascular tone) may have caused a severe depletion of PtdIns(4,5)P₂ levels in vascular smooth muscle cells. The inability to replenish PtdIns(4,5)P₂ following stimulation would certainly be deleterious to maintenance of vascular tone or other aspects of cardiovascular homeostasis. Although these toxicity studies were not specifically designed to test these possibilities, they point to a direction that will have to be followed up in future in vivo studies. The animals dosed with a lower level of the compound remained alive throughout the studies and displayed moderate to severe GI abnormalities. This may be due to the highest exposure of these cells during oral administration or a specific role of PI4KA in gut homeostasis. Similarly, the cKO animals were dosed orally with tamoxifen providing highest exposure to the drugs or to Cre induction in the GI tract, and the most severe toxicities were observed in this tissue. The GI abnormalities observed here were similar to those observed in a recent study using a different cKO pi4ka mouse model (22). Interestingly, none of the cKO mice exhibited rapid lethality as was seen with some compounds, but rather they progressed to a moribund condition, suggesting significant differences in the pharmacological and genetic ablation of PI4KA on in vivo toxicity.

More detailed studies are in progress to evaluate the efficiency of gene ablation in the various tissues and the specific consequences of the lack of the proteins. However, even these preliminary studies caution using only genetic ablation studies to determine safety of potential therapeutic targets.

Based on the results of the experiments in cultured cells, one would expect to see defects in many other tissues yielding a variety of symptoms. However, given the fact that cells can...
function with greatly reduced PI4KA levels unless very strongly stimulated via G_{q}-coupled receptors, it is most likely that these animals still maintained some PI4KA expression or activity in tissues other than the GI tract. For example, it is important to point out that the drugs used in these studies do not cross the blood-brain barrier, and also, tamoxifen exposure of the brain was found to be minimal during this regime of tamoxifen administration (data not shown). More studies are in progress to address these questions with different experimental design.

In summary, this study showed that PI4KA is uniquely important in the maintenance of PtdIns(4,5)P_{2} levels during strong PLC activation, but it appears dispensable for normal...
cell growth and maintenance at least for a period of time. It also confirmed previous claims that a fraction of the enzyme or its activity is sufficient to support cellular functions, although HCV replication is more sensitive to inhibition of the enzyme. Although preliminary, the animal studies showed that the cells within the GI tract, especially those showing rapid turnover may require PI4KA for their normal functions. Additionally, acute inhibition of the enzyme with selective small molecules may also exert additional deleterious effects in addition to GI problems such as cardiovascular tone, possibly explaining the sudden death caused by these drugs. These results highlight the significant risks associated with pharmacological targeting of PI4KA. The compounds and animal models generated for these studies will greatly benefit further research on PI4KA to better understand the role of this important enzyme in normal physiology and disease.

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