Lysosomal Biogenesis and Implications for Hydroxychloroquine Disposition^S

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ABSTRACT

Lysosomes act as a cellular drug sink for weakly basic, lipophilic (lysosomotropic) xenobiotics, with many instances of lysosomal trapping associated with multiple drug resistance. Lysosomotropic agents have also been shown to activate master lysosomal biogenesis transcription factor EB (TFEB) and ultimately lysosomal biogenesis. We investigated the role of lysosomal biogenesis in the disposition of hydroxychloroquine (HCQ), a hallmark lysosomotropic agent, and observed that modulating the lysosomal volume of human breast cancer cell lines can account for differences in disposition of HCQ. Through use of an in vitro pharmacokinetic (PK) model, we characterized total cellular uptake of HCQ within the duration of static equilibrium (1 hour), as well as extended exposure to HCQ that is subject to dynamic equilibrium (>1 hour), wherein HCQ increases the size of the lysosomal compartment through swelling and TFEBinduced lysosomal biogenesis. In addition, we observe that pretreatment of cell lines with TFEB-activating agent Torin1 contributed to an increase of whole-cell HCQ concentrations by

1.4- to 1.6-fold, which were also characterized by the in vitro PK model. This investigation into the role of lysosomal volume dynamics in lysosomotropic drug disposition, including the ability of HCQ to modify its own disposition, advances our understanding of how chemically similar agents may distribute on the cellular level and examines a key area of lysosomal-mediated multiple drug resistance and drug-drug interaction.

SIGNIFICANCE STATEMENT

Hydroxychloroquine is able to modulate its own cellular pharmacokinetic uptake by increasing the cellular lysosomal volume fraction through activation of lysosomal biogenesis master transcription factor EB and through lysosomal swelling. This concept can be applied to many other lysosomotropic drugs that activate transcription factor EB, such as doxorubicin and other tyrosine kinase inhibitor drugs, as these drugs may actively increase their own sequestration within the lysosome to further exacerbate multiple drug resistance and lead to potential acquired resistance. Downloaded from jpet.aspetjournals.org at ASPET Journals on March 20, 2024

Introduction

Hydroxychloroquine (HCQ) has been investigated in over 50 cancer clinical trials and countless preclinical studies through the past decade as an additive compound to primary chemotherapy treatment. Mechanistically, HCQ acts to inhibit autophagy, which has been identified as a mechanism of tumor cell resistance to chemotherapy and a key pathway to survival within the tumor microenvironment. Although nextgeneration autophagy inhibitors are in preclinical stages of development, HCQ remains the only clinically approved

autophagy inhibitor. Overall, HCQ is generally effective and safe as a treatment but suffers from various pitfalls within the pharmacologic space—specifically in terms of predictability of patient likelihood of response and reliable pharmacokinetic (PK) and pharmacodynamic (PD) metrics. There is a general disconnect between HCQ PK in the tumor and blood, which makes it challenging to be confident that target concentrations of HCQ are achieved within the tumor when optimizing dosing strategies (Barnard et al., 2014). Characterization of HCQ PK properties on a cellular level may provide insight toward tumor-specific factors that would promote this disconnect.

Cellular uptake of HCQ is driven primarily through lysosomal ion trapping due to the weakly basic and lipophilic properties that characterize it as a hallmark lysosomotropic agent. HCQ readily diffuses through neutral spaces but becomes diprotonated in the lysosome, whose acidic environment facilitates accumulation. This results in concentrations significantly higher than other compartments within the cell (Duvvuri and Krise, 2005). Key mechanisms contributing to pharmacokinetics have been considered and consolidated into

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ABBREVIATIONS: AUC, area under the curve; CI, confidence interval; Ctrl, control; CQ, chloroquine; Cy3, Cyanine; DAPI, 4',6-diamidino-2-phenylindole; GSEA, gene set enrichment analysis; HCQ, hydroxychloroquine; MAPE, median absolute performance error; MN, monensin; MPE, median performance error; PD, pharmacodynamic; PE, performance error; PK, pharmacokinetic; Vf_{lys}, Lysosomal Volume Fraction; T1, Torin1; TFEB, transcription factor EB; TKI, tyrosine kinase inhibitor.

in vitro mathematical models to describe cellular uptake of compounds with similar physicochemical properties to HCQ—namely, weakly basic and lipophilic (Ishizaki et al., 2000; Trapp et al., 2008; Kornhuber et al., 2010; Schmitt et al., 2019). These models assume the cell behaves as a static system of compartments, but recent evidence suggests that lysosomotropic compounds induce changes in the lysosome itself.

Lysosomes swell in response to chloroquine (CQ) exposure, which is mechanistically and pharmacokinetically similar to HCQ. Near clinically relevant doses of CQ have been observed to cause a 3-fold increase in relative lysosomal volume after multiple days of exposure to the drug (King et al., 2016), and recruitment of galectin-3 to sites on these swollen lysosomes, indicating membrane damage (Gallagher et al., 2017). In addition to an increase in lysosomal size, an increase in lysosomal number has also been reported in response to a panel of lysosomotropic agents, including CQ, over the course of 4 and 24 hours (Lu et al., 2017). This increase in lysosomal number was linked to an increase in nuclear translocation of master lysosomal biogenesis transcription factors TFEB, transcription factor binding to IGHM enhancer 3 (TFE3), and melanocyte inducing transcription factor (MITF). TFEB activation of the coordinated lysosomal expression and regulation (CLEAR) network of genes has been described as the master regulation system of lysosomal biogenesis (Settembre et al., 2011, 2012). Lysosomotropic agents cause activation of this network by inducing lysosomal stress, and it has been shown to activate with as little as 90 minutes of exposure to CQ (Zhitomirsky et al., 2018). The results of lysosomal swelling and TFEB activation by lysosomotropic agents may be linked to their long-term tissue PK. Extended dosing of HCQ is characterized by a consistent increase in HCQ concentrations in rat tissue over the course of 3 months of daily dosing. In some tissues (spleen, kidney, liver, heart) HCQ concentrations continued to increase out to the latest time point measured (at 7 months) (McChesney, 1983). This long-term PK profile is noted by McChesney to be similar to other lysosomotropic drugs, such as chlorpromazine.

Because of the frequent, high doses of HCQ used in cancer clinical trials, we characterized this lysosomal adaptation response on a cellular level and investigated the extent to which it may influence cellular disposition of the drug. The data described herein suggest that baseline cell uptake of HCQ is associated with different baseline lysosomal profiles between cell lines, which were integrated into a lysosome cellular PK mathematical model (Trapp et al., 2008; Kornhuber et al., 2010) to accurately model whole-cell HCQ PK to chemical kinetic equilibrium. Small but consistent increases in whole-cell uptake of HCQ over an extended time period were also observed and were attributed to an increasing volume of the lysosomal compartment. The in vitro PK profile of HCQ was thus broken up into a short-term kinetic equilibrium portion, within 1 hour of exposure, and a long-term dynamic portion, out to 24 hours of exposure, for all cell lines. These data were used in conjunction with observed differences in baseline lysosomal profiles to further characterize the cellular PK of HCQ by accounting for a dynamic lysosomal compartment.

Materials and Methods

Cell Lines. Human breast cancer cell lines, including MDA-MB-231, MDA-MB-468, T47D, and MCF7 human breast cancer cell lines,

were purchased from Cell Culture Services at University of Colorado Anschutz Medical Campus (Aurora, CO). For cell pharmacokinetic experiments, all cell lines were stably transfected with Incucyte NucLight Red Lentivirus (4476; Sartorius, Gottingen, Germany) prior to experimentation.

Cell lines were cultured in Dulbecco's modified essential medium (10-017-CV; Corning) supplemented with 10% fetal bovine serum (PS-FB3; Peak Serum), 1% penicillin/streptomycin (30-002-CI; Fisher, Hampton, NH), and 1% sodium pyruvate (25-000-CI; Fisher) at 37°C and 5% $\rm CO_2$ in a humidified incubator.

Chemicals and Reagents. HCQ sulfate was purchased from Sigma-Aldrich (St. Louis, MO). Monensin (MN) was purchased from VWR (Radnor, PA). Torin1 (T1) was purchased from Fisher. Hoechst 33342 20 mM solution (62249) and CellLight Lysosomes-GFP, Bac-Mam 2.0 (C10507) were purchased from ThermoFisher (Waltham, MA). pyrene-benzothiazolium dye 1b (ETP), a pH-insensitive lysosomal lumen dye applicable for live and fixed cell imaging, was received as a gift from Dr. Yi Pang at the University of Akron (Abeywickrama et al., 2019).

Cell Uptake Experiments. Four human breast cancer cell lines were used to perform experiments. The first two, MBA-MB-231 and MDA-MB-468, are mechanistically sensitive to autophagy inhibition through gene knockdown (Maycotte et al., 2014; Towers et al., 2019) and pharmacologically sensitive to hydroxychloroquine. The second two, MCF7 and T47D, are insensitive to both methods (Maycotte et al., 2014)

Cell lines were cultured in standard Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% FBS, 1% penicillin streptomycin (PS), and 1% sodium pyruvate in an incubator at 37°C and 5% CO $_2$. T47D and MCF7 cell lines required supplementation of 8 $\mu \rm g/ml$ of insulin to maintain growth. For acidic media experiments with MDA-MB-231 cells, Dulbecco's Modified Eagle Medium (DMEM) without sodium bicarbonate was purchased, and bicarbonate was added to achieve an incubator-conditioned media pH. Culture media buffers other than sodium bicarbonate, such as HEPES, were purposely omitted, as HEPES has been shown to activate the micropthalmia/transcription factor E (MiT/TFE) network, resulting in lysosomal biogenesis (Tol et al., 2018).

Cell lines were plated, allowed to adhere overnight, and treated at roughly 70% confluence. All cell lines stably expressed NucLight Red, and cell counts were measured by counting red fluorescent nuclei on an Incucyte ZOOM (Sartorius) prior to treatment with drug. For 4- and 24-hour time points, cells were counted immediately prior to harvesting, as the doubling time for these cell lines is between 16 and 20 hours. Experiments were not extended past 24 hours to avoid cell line growth inhibition in response to HCQ. For treatment, culture media was aspirated and replaced with fresh, incubator-conditioned media containing 10 µM HCQ and placed in a 37°C incubator until the designated time point. At the time point, cell media containing HCQ were aspirated, and cells were washed with Hank's balanced salt solution (HBSS) and then trypsinized for 5 minutes. Trypsinized cells were washed off the plate with additional culture media at a ratio of 4:1 media:trypsin. Cells were then pelleted at 1600 rpm for 5 minutes, media were carefully aspirated, and the cell pellet was reconstituted in MilliQ-water and frozen until liquid chromatography with tandem mass spectrometry (LC/MS-MS) extraction and analysis.

(LC/MS-MS) analysis follows methods described in Barnard et al. (2014); briefly, cell pellets were vortexed and lysed for 10 minutes in a sonication water bath. Acetonitrile was then added to the cell pellets to yield a final solution of 50% acetonitrile + 50% MilliQ-H₂O. Cells were vortexed for 10 minutes and then centrifuged at 13,300 rpm for 10 minutes. Supernatant was transferred to polypropylene inserts. CQ was used as an internal standard. Concentrations of supernatant obtained on LC/MS-MS were normalized to the cell counts obtained for each sample from an Incucyte ZOOM, and samples were then normalized to cell volume, obtained for each cell line from suspended cell diameter measured on a Countess system (n=4), to obtain HCQ concentration (micromolars) in each cell.

For experiments with MN, cells were pretreated with 25 μM MN for 30 minutes prior to adding HCQ. After 30 minutes, 10 μM HCQ was added for the duration of the 1-hour exposure. For experiments with T1, cells were pretreated with 250 nM Torin1 for 16 hours prior to adding HCQ. All cell uptake experiments were performed at least in triplicate.

Lysosomal Imaging. For all imaging experiments, cells were plated at 10,000 cells per well on an eight-well glass chamber slide (C8-1.5H-N; Cellvis) and allowed to adhere overnight.

To determine lysosomal volume, live cells were treated with 1 μM ETP (Abeywickrama et al., 2019) and incubated for 30 minutes. Excess dye was removed by washing twice with warm media and incubated with 10 µM of Hoescht 33342 for 10 minutes. Cells were then washed twice with PBS and fixed for 10 minutes in 2% paraformaldehyde (PFA). PFA was washed off twice with PBS, and cells were imaged on an Olympus IX83 microscope with a Plapon 60× objective, numerical aperture (NA) = 1.42, on Cy3 and DAPI confocal channels. In total, 43 z-stacks of 0.24-µm step size were captured on Cy3 and DAPI channels. Lysosomal volume fraction of cells was determined using an in-house image processing pipeline, which is described briefly as follows: raw confocal image stacks were converted to tagged image file format (TIF) files and imported into Python version 3.7.6. A reference image was created by compressing the stack by its maximum pixel intensity. Each slice of each image stack was analyzed to determine the peak signal-to-noise ratio relative to the reference image using the sewar module (https://pypi.org/project/sewar/). A comma separated value (CSV) file containing the top 10 peak signal-to-noise ratio slices for each image stack was generated and imported into a FIJI macro that calculated the Cy3 (lysosome) area for each raw VSI image slice based on an algebraic threshold of mean intensity. The area for each slice was multiplied by the step size (0.24 μm) to generate a volume, and then volumes from the 10 slices per image stack were summed to generate a lysosomal volume for the image. The lysosomal volume was normalized to the cell count of each image, and the cell count was normalized to the experimentally determined cell volume for each cell line to generate a lysosomal volume fraction per cell. In total, 5 to 10 images were obtained for each experimental replicate, and lysosomal volume fraction is presented as the mean \pm S.D. of each replicate (n = 3) for each cell line. Imaging of cell lines for GFP-lysosomal associated membrane protein 1 (LAMP1) expression followed the same imaging process, except cells were incubated with 1 µl of the baculovirus overnight prior to treatment with HCQ.

For drug treatment conditions, single plane images were captured at $20\times$ using the same microscope but an Uplsapo $20\times$ objective, numerical aperture (NA) = 0.75, on Cy3 and DAPI light-emitting diode (LED) channels. To determine a change in lysosome content of the cell, cells were treated with the same drug treatment conditions as described in the *Cell Uptake Experiments* section. Chamber slides were prepared in the same way as described above. Images were quantitated using a FIJI macro with an algebraic threshold of the mean intensity, and data were calculated as Cy3 area/DAPI area. Roughly 10 images were obtained for each experimental replicate, and data are reported as the mean \pm S.D. of each replicate (n=4). Statistical significance was calculated as a one-way ANOVA with Dunnett's multiple comparisons test relative to the control, with significance as P<0.05.

TFEB Activation. TFEB activation by HCQ was determined with two different kinds of methods, including gene set enrichment analysis (GSEA) of the TFEB-regulated Coordinated Lysosomal Expression and Regulation (CLEAR) network and through a transcription factor EH-EB activity assay (TFEH-TFEB; RayBiotech). Gene expression microarray data were prepared for all four human breast cancer (hBC) lines following cell culture preparation steps as outlined in the Cell Uptake Experiments section. Cell lines were treated with 20 μM HCQ for 24 hours prior to harvesting for gene expression preparation in triplicate. Prepared cell extraction homogenate was run on an HG_U133_Plus_2 chip. Cell intensity (CEL) file

data were extracted and prepared for GSEA using the Affymetrix Bioconductor package in R and normalized using Robust Multi-array Average (RMA). Expression data were then analyzed using the GSEA 4.0.3 software provided by the Broad Institute. Gene sets from the four cell lines, treated with HCQ or vehicle, were analyzed in GSEA using a gene set representative of TFEB lysosome targets (Palmieri et al., 2011) and run in tandem with the Hallmarks of Cancer gene set data base version 7.1 provided in the GSEA software. The run was set for 1000 permutations following a "gene_set" permutation type. Data are reported as the normalized enrichment score (NES) value, and significance was determined by false discovery rate (FDR) q-value generated in the run. Raw cell intensity (CEL) files are available in the supplement. TFEB nuclear activation was performed in duplicate and was determined using the human TFEB transcription factor activity assay following the manufacturer's instructions. TFEB activity data are expressed as means ± S.D. and analyzed by one-way ANOVA with Dunnett's multiple comparison test relative to control, with significance as P < 0.05.

Cell Pharmacokinetic Mathematical Modeling and Simulation. To connect whole-cell HCQ PK with the perturbations observed in the lysosomal compartment, HCQ whole-cell uptake was simulated using a previously published base model of lysosomal uptake (Trapp et al., 2008; Kornhuber et al., 2010). Lysosomal volume fraction for each cell line was determined by imaging methods described above. Extracellular pH was determined by measuring the pH of incubator-equilibrated media in a pH meter (7.6) and then testing the change in cellular uptake of HCQ at the 1-hour time points in media of decreased pH (7.0). The influence of the lysosome and cytosolic pH was determined experimentally by pretreating MDA-MB-231 cells with MN, a lysosome-selective ionophore (Grinde, 1983), and simulating the experimental uptake of HCQ in cells after pretreatment. The model values of cytosolic pH of roughly 7.0 and lysosomal pH of 5.0 worked well with the cell lines used, which are similar to what has been previously reported for MCF7 cytosolic pH (Belhoussine et al., 1999) and MCF7 and MDA-MB-231 lysosomal pH values (Montcourrier et al., 1994). It is noted that the range of the pH values from different literature reports is quite broad, with lysosomal pH values reported around 4.2 for MDA-MB-231 cells (Ndolo et al., 2012), 4.5 for MCF7 and MDA-MB-231 cells (Wang et al., 2019), and 5.1 for MCF7 and MDA-MB-231 cells (Montcourrier et al., 1994). Cytosolic pH exhibited a similar trend from literature, with reported values around 6.9 for MCF7 (Belhoussine et al., 1999), 7.3 for MCF7 (Persi et al., 2018), and 7.4 for MDA-MB-231 and MCF7 (Wang et al., 2012). From the studies using multiple human breast cancer lines, which were also used in this work, the lysosomal and cytosolic pH were not very different between each cell line, so we chose to use fixed values for these parameters across all four cell line models. Additional factors that could potentially play a role in cellular uptake and clearance of HCQ, such as acid-phospholipid binding and endo/lysosomal turnover, were not investigated between cell lines.

Evaluation of Model Fit. Model fit versus experimental data were evaluated by comparing the experimental $AUC_{0-1\ hour}$ or $AUC_{0-24\ hour}$ to the simulated value for each cell line. Additionally, model accuracy was analyzed by calculating the performance error (PE), the median performance error (MPE%) to investigate positive/negative bias, and the mean absolute performance error (MAPE%) to determine overall model fit (Gustafsson et al., 1992). These calculations included all time points relevant to the parent figure.

Computer Simulation and Software. Intracellular PK model simulation was done in MATLAB version R2020a (Mathworks, Natick, MA), and the intracellular compartmental system was solved using the ode45 package. Image selection was done in Python version 3.7.6 using sewar and tqdm packages. Image analysis was done in ImageJ version 1.52p using methods described above.

Results

Whole-Cell Uptake of HCQ Is Proportional to Basal Lysosomal Volume Fraction. The in vitro PK of HCQ was assessed in four human breast cancer (hBC) cell lines—MDA-MB-231, MDA-MB-468, T47D, and MCF7—by collecting total cell homogenate at 1, 5, 15, 30, 60, 240, and 1440 minutes after incubating with 10 µM HCQ. This concentration of HCQ was chosen because it is within the upper range of clinically achievable plasma C_{max} concentrations. HCQ concentration within the whole cell was determined by normalizing total HCQ in the homogenate to the total cell volume in the homogenate outlined in Supplemental Fig. 1. We observed different concentration levels of HCQ for each cell line, which mostly followed the ranking of MDA-MB-231 > MDA-MB-468 > T47D > MCF7 (Fig. 1A). For all four hBC lines, the HCQ total cellular concentrations increased the longer the cell lines were exposed to the drug, up to total cellular concentrations peaking between 0.58 and 2.67 mM, which were 58-267 times the extracellular concentration of HCQ, and are consistent with steady-state partition coefficients of HCQ tissue:plasma from previous in vivo studies (McChesney et al., 1967; McChesney, 1983; Wei et al., 1995).

As acidic organelles are suggested to account for a majority of cellular distribution of diprotic lipophilic compounds, we investigated the total lysosomal volume fraction of each cell line using the microscope imaging methods outlined in Supplemental Fig. 2. We observed lysosomal percent volume of total cellular volume ranging from 0.50% \pm 0.19% in MCF7 cells to roughly 3.67% \pm 1.09% in MDA-MB-231 cells (Fig. 1B), which followed roughly the same trend as total cellular uptake of HCQ based on AUC $_{0-24\ hour}$ (Fig. 1C). Overall, mean total cell uptake concentrations of HCQ showed a significant

Pearson correlation with lysosomal volume fraction for time points 5, 15, 30, 60, 240, and 1440 minutes (Fig. 1D). The 1-minute time point was not significant, likely due to insufficient time for HCQ to diffuse into the lysosomal compartment. Total exposure to HCQ, as calculated by $AUC_{0-24\ hour}$, also had a strong correlation ($P=0.032,\ R^2=0.936$) with cellular lysosomal volume fraction (Fig. 1E).

Basal Lysosomal PK Model of HCQ Accounts for Uptake by Adjusting Lysosomal Volume Fraction. To further investigate the in vitro PK of HCQ in the four hBC lines, we used a base PK model of lysosomotropic drug uptake (Trapp et al., 2008; Kornhuber et al., 2010) and modified it to be specific to the four cell lines in the study with experimentally determined lysosomal volume fractions. HCQ in vitro PK was modeled as a diffusion-based three-dimensional system in which net flux of HCQ across membranes is driven by the permeability of each ionization state, and ionization state (0, 1⁺, 2⁺) is a function of pH described by the Henderson-Hasselbalch activity ratio. The model consisted of three compartments representative of in vitro settings: culture media, cytosol, and lysosomes (Fig. 2A). Trafficking between compartments was driven by net flux. The lysosomal compartment had a dynamic pH (\Delta pH) term that was a function of lysosomal HCQ concentration, as HCQ is suggested to neutralize the acidic lysosomal pH. Model parameters are listed in Supplemental Table 1.

By adjusting the lysosomal volume fractions for each cell line, we simulated the in vitro PK experiment of HCQ administered at 10 μ M for each cell line out to 1 hour (Fig. 2B). For all cell lines except T47D, the experimentally derived mean lysosomal volume fraction allowed the simulation to capture the mean concentration of HCQ with a strong fit. The simulation was also

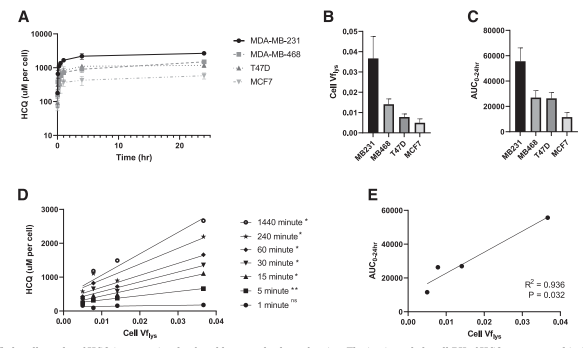


Fig. 1. Whole-cell uptake of HCQ is proportional to basal lysosomal volume fraction. The in vitro whole-cell PK of HCQ was assessed in four hBC cell lines (MDA-MB-231, MDA-MB-468, T47D, and MCF7) at 1, 5, 15, 30, 60, 240, and 1440 minutes after incubating with 10 μ M HCQ (A). Cell lysosomal volume fraction was calculated as outlined in Supplemental Fig. 2 (B). Lysosomal volume fraction closely followed the ranking of AUC_{0-24 hour} of the whole-cell PK data (C). Mean whole-cell HCQ concentrations at each time point, except 1 minute, were significantly correlated with cellular lysosome volume fraction (D). This correlation was also significant when comparing the mean AUC_{0-24 hour} of each cell line to the cellular lysosome volume fraction (E). Applicable data are shown as means \pm S.D., and significance is defined as P < 0.05 (*) and P < 0.01 (**).

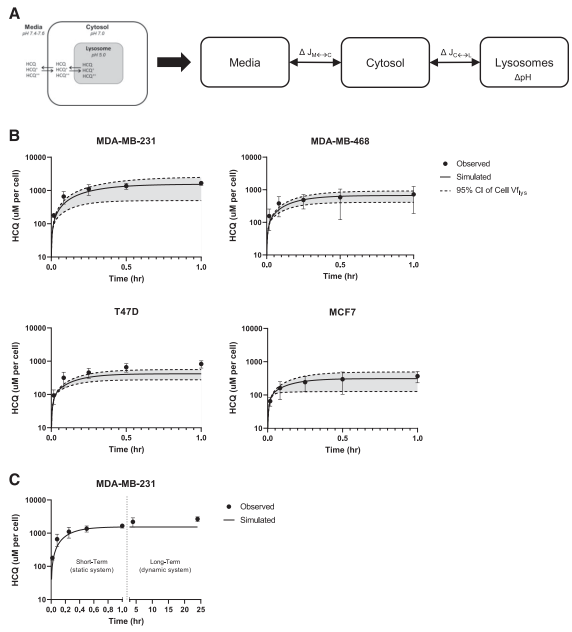


Fig. 2. Basal lysosome PK model of HCQ accounts for uptake by adjusting lysosomal volume fraction. To further investigate the in vitro PK of HCQ in the 4 hBC lines, we used a previously published base model of lysosomotropic drug uptake into cells. The compartmental model is outlined, in which compartments considered are culture media, cytosol, and lysosome, with the lysosome contained within the cytosolic compartment (A). Mathematically, the three compartments are separate; HCQ diffuses freely from media (M) \leftrightarrow cytosol (C) \leftrightarrow lysosomes (L), and diffusion is represented by a net flux term (Δ J) that is the sum of permeability of each HCQ ionization state (neutral, +, ++). The lysosomal compartment has a dynamic pH feedback term that was representative of the proposed mechanism of HCQ to increase the pH of the lysosome based on free drug concentration. The model was used to simulate HCQ uptake in each cell line based on lysosomal volume fractions and other parameters in Table 1. The output of the model simulation for each cell line is shown as the observed mean \pm 95% CI bounds in conjunction with the simulated mean \pm 95% CI bounds of the cellular lysosomal volume fraction (B). The model only considered time points out to the 1st hour, which were labeled as short-term static equilibrium. In the experimental data, it is observed that concentrations continue to increase after 1 hour, which is characterized as the long-term dynamic system (C), and this is investigated in later figures.

run using the 95% CI bounds for the experimentally derived lysosomal volume fractions to investigate variability due to this parameter. The simulation suggested that the system reaches equilibrium around 30 minutes; however, the experimental PK data for all four cell lines suggests that HCQ concentrations in the cell continue to increase steadily at 1, 4, and 24 hours (Fig. 2C). Early time points were characterized by chemical kinetic equilibrium, in which the cellular physiologic parameters are static, followed by later time points

when the system physiologic parameters are dynamic. The dynamic system is investigated further in later figures. The model fit for each of the cell lines out to 1 hour is characterized by the ratios of simulated:observed AUC (Table 1), as well as measures of predictive performance PE, MAPE%, and MPE% (Supplemental Table 2). Using a previously determined range of 0.5–2.0 as acceptable ratios for AUC of simulated: experimental, the model captured MDA-MB-231, MDA-MB-468, and MCF7 very closely (0.936, 0.995, 0.980). T47D was

TABLE 1 Static model metrics (AUC)

AUC Summary (0 to 1 h) Cell Line	Experimental (Exp) Mean	Simulated (Sim) Mean	Ratio Mean Sim/Exp
MDA-MB-231 MDA-MB-468 T47D	1239 551 588	1160 549 354	0.936 0.995 0.602
MCF7	276	270	0.980

underpredicted but still fell within the acceptable range (0.602). In comparing the model PE, the model underpredicted early time points (1, 5 minutes) for all cell lines except MCF7. Time points between 15 and 60 minutes were predicted with good accuracy, aside from T47D. MAPE%, a metric of model accuracy, suggests that the model predicts individual time points in MDA-MB cell lines with about 30% error, T47D around 60%, and MCF7 about 6.5%. Overall, the model underpredicts concentrations (MPE%) by 18.4 and 8.6% in the MDA-MB cell lines, although this is biased by early time point error, 64% in T47D, and overpredicts by only 1.3% in MCF7. Observations from these metrics suggest that T47D cell uptake of HCQ may be better explained by the upper CI bound of its lysosomal volume fraction (1.16%) rather than its mean (0.78%). Model metrics also suggest that early time points (1, 5 minutes) in MDA-MB cell lines were either underpredicted based on uptake kinetics or there was time-based experimental error exacerbated by early time point sampling.

We also investigated the influence of the pH parameters on total cellular uptake of HCQ in MDA-MB-231 cells. We measured the pH of incubator-conditioned culture media (5% CO₂, 37°C) used in the previous experiments at 7.6 and made acidic culture media at an incubator-conditioned pH of 7.0. As stated in the *Materials and Methods* section, we opted out of using culture media buffers other than sodium bicarbonate, such as HEPES, as it has been suggested to activate the micropthalmia/transcription factor E (MiT/TFE) pathway, which ultimately results in lysosomal biogenesis (Tol et al., 2018). As the model accurately captured MDA-MB-231 cell lines under neutral conditions, they were exposed to 1 or 10 µM HCQ in the conditioned culture media for 1 hour, and the resulting decrease in total cellular uptake of HCQ associated with decreasing media pH was quite dramatic. We observed a 6.4-fold decrease in HCQ uptake at the treated concentrations when switching the media pH from 7.6 to 7.0 such that in most cases the 10 μM HCQ uptake at pH of 7.0 was nearly equivalent to the 1 µM HCQ uptake at pH of 7.6 (Supplemental Fig. 3A). The shape of the simulated kinetics uptake curve was not visibly modified by extracellular pH out to the 1-hour time point tested (Supplemental Fig. 3B). Interestingly, lysosomotropic drug partition into cells and tissues has been suggested to be significantly reduced by acidic extracellular pH (Wojtkowiak et al., 2011), causing a subsequent resistance of cells to growth inhibition by HCQ. We tested the effect of acidic culture media in blunting HCQ growth inhibition in MDA-MB-231 cells and observed an almost complete reduction in growth inhibition (Supplemental Fig. 3C), similar to the trend described with CQ (Pellegrini et al., 2014). In addition to testing the influence of extracellular pH directly, we also investigated the contribution of lysosome and cytosol pH gradient indirectly by treating cells

with MN prior to HCQ to eliminate the gradient. In MDA-MB-231 cells, pretreatment with 25 μM MN for 30 minutes prior to HCQ caused a reduction in total cellular uptake of HCQ by 93.2% at 1 hour. We simulated this experiment using the model by changing the lysosomal pH to 7.0 and comparing with the model with lysosomal pH of 5.0 (Supplemental Fig. 3D). Overall, the model captured the decrease in total cellular uptake well, with simulated MN(+):MN(-) concentration ratio of 0.078 versus observed MN(+):MN(-) concentration ratio of 0.068.

HCQ Increases the Size of the Lysosomal Compartment. Although the model captures early time point in vitro PK (<1 hour) of HCQ, as well as variabilities in influential model parameters along the pH gradient, it fails to capture later time points, as the simulation suggests the system reaches equilibrium when the observed concentrations actually continue to gradually increase out to 24 hours (Fig. 2C). Recent reports suggest that many weakly basic lipophilic compounds, including CQ, activate lysosomal biogenesis by activating TFEB (Lu et al., 2017; Zhitomirsky et al., 2018; Zhao et al., 2020), which ultimately increases the lysosomal volume fraction of treated cells and should allow them to sequester even more drug than their original baseline (Ruzickova et al., 2019). CQ has been observed to activate TFEB and increase lysosomal content of the cell, so we investigated the capability of HCQ to do the same. Treatment of all four hBCs with either HCQ for 24 hours or TFEB activator Torin1 for 16 hours caused a significant increase in TFEB activity in the nucleus (Fig. 3A), suggesting it is released from the lysosome and translocated to the nucleus after HCQ treatment. In addition, gene microarray data on all four cell lines treated with HCQ or vehicle for 24 hours showed a significant enrichment in expression of the TFEB-associated direct targets in lysosomal function (Fig. 3B) as characterized by the Ballabio group (Palmieri et al., 2011). Treatment of all four cell lines with HCQ or Torin1 caused an increase in the relative volume of the lysosome compartment. MCF7 cells had the most visually distinct increase in lysosomal burden with both drugs (Fig. 3D). Quantitation of the increase in lysosomepositive area per nuclei suggests a relative increase of 1.59-to 14.93-fold (average 5.90-fold) in the cell lines with 24-hour treatment with HCQ and a 2.25- to 10.7-fold (average 5.55fold) increase with Torin1 (Fig. 3C). This increase in lysosomal burden by HCQ may have some contribution independent of TFEB as well, as CQ has been shown to induce swelling in lysosomes at higher doses, which is visually apparent in images comparing Torin1 increase in lysosomal burden versus HCQ increase in lysosomal burden in MCF7 (Fig. 3D) and the other three cell lines (Supplemental Fig. 4). To further investigate lysosomal swelling by HCQ, we transfected MDA-MB-468, T47D, and MCF7 cells with a GFP baculovirus for lysosomal associated membrane protein 1, as it more clearly defines the membranes of individual lysosomes, and we imaged cells after treatment with $10 \mu M$ HCQ for 24 hours. MDA-MB-231 cells would not express the virus and were omitted from this experiment. In the three cell lines, we observed a dramatic increase in size of some lysosomes after HCQ exposure (Supplemental Fig. 5).

Dynamic Lysosomal Volume Accounts for Simulation Error. The original model with a static lysosomal volume compartment was able to capture total cell uptake of HCQ out to 1 hour but significantly underpredicted 4- and 24-hour time

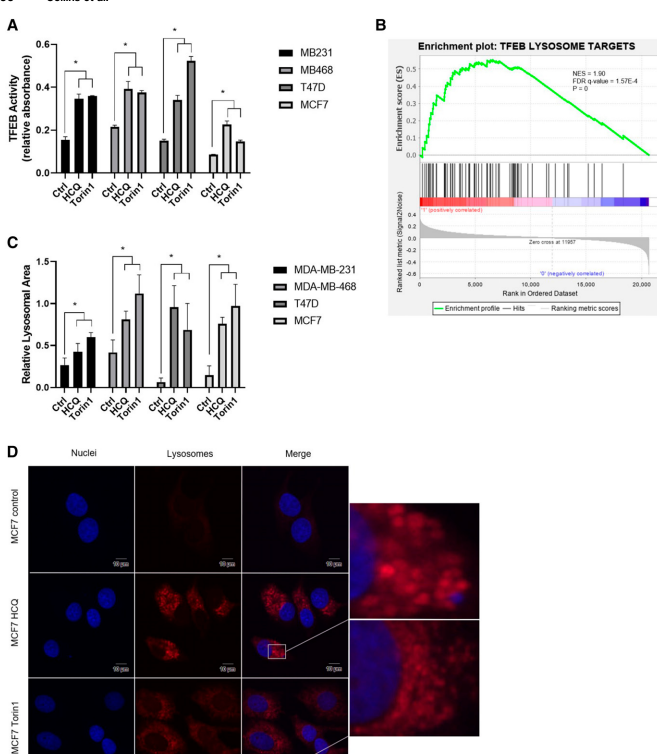


Fig. 3. HCQ increases the size of the lysosomal compartment. We investigated the ability of HCQ to change the size of the lysosomal compartment. TFEB activity in the nucleus was significantly increased in all cell lines treated with 10 μ M HCQ for 24 hours and with 250 nM Torin1, a molecular TFEB-activating agent, for 16 hours (A). GSEA of all four cell lines treated with 20 μ M HCQ for 24 hours also caused a significant enrichment of TFEB lysosome targets, as determined by the normalized enrichment score (NES) and false discovery rate (FDR) (B). Fluorescence microscopy imaging of all four cell lines with ETP showed a significant increase in lysosome accumulation within all cell lines treated with 10 μ M HCQ for 24 hours or 250 nM Torin1 for 16 hours (C). Representative images of MCF7 cells imaged with ETP under the treatment conditions are shown in (D). Zooming in on MCF7 cells treated with HCQ shows a visual increase in lysosomal size in comparison with the Torin1 treatment. Significance is defined as P < 0.05 (*). To prepare figures for publication, the raw image threshold was adjusted to the same upper and lower bounds across the entire image for all images shown.

points. To account for this error, we incorporated lysosomal biogenesis into the model by adding a dynamically growing lysosomal volume fraction represented by the proposed mechanism outlined in Fig. 4. The growing compartment is represented as a linear increase in lysosomal volume fraction as a function of time: $Vf_{lys}(t) = Vf_{lys_0} \cdot (1 + x \cdot t)$. We found that a 1.5- to 3-fold total increase in lysosomal volume fraction by 24 hours in the dynamic lysosomal model accounted for the underprediction of 4- and 24-hour time points predicted by the static lysosomal model (Fig. 5A). The simulated 1.5- to 3-fold increase in lysosomal volume fraction over 24 hours of HCQ exposure was within the 95% CI of the experimentally observed increase due to HCQ in the four cell lines shown in Fig. 3C, except for T47D. As such, for T47D, we tested an increase of 2.5fold in lysosomal volume starting from the upper bound of the 95% CI (1.155%), which resulted in a better model fit to 24 hours. Incorporation of this time-based linear increase did not significantly affect time points before 1 hour (Fig. 5B).

The model fit, as characterized by the ratios of simulated: observed AUC_{0-24 hour}, is shown in Table 2, and measures of predictive performance PE, MAPE%, and MPE% are shown in Supplemental Table 3. Overall, the model captured the early time points of the cell lines with good performance, as the AUC and ratios were close to 1. PE followed the same trend to the static model for time points 0–1 hour, with 4- to 24-hour time points being slightly underpredicted for all cell lines except MCF7 and T47D at 24 hours. MAPE% suggests the model fits overall time points for all cell lines between 7.53% and 45.21%, and MPE% suggests a slight underprediction bias for the model. Overall, this experimental data and model simulation suggest that HCQ creates a positive-feedback loop within the cell lysosomal compartment, causing it to increase its own distribution within the cell over time.

Torin1 Increases Size of Lysosomal Compartment, Which Increases Cell Uptake of HCQ. Although HCQ appears to be able to modify its own distribution, we investigated the capability of direct modification of the cell lysosomal volume fraction to increase HCQ uptake capacity. In Fig. 4D, we observed that Torin1 increased the relative lysosomal burden by 2.25- to 10.7-fold. In this experiment, we repeated the HCQ cell uptake experiments out to 24 hours but first pretreated cells with Torin1(+) or (-) for 16 hours prior to

administering HCQ. Pretreatment with Torin1 resulted in a mean increase in cellular uptake of HCQ at all time points in all cell lines of 1.4- to 1.6-fold (Fig. 6A). To test this in the context of the in vitro PK model, we changed the cell line base lysosomal volume fraction at time 0 based on the mean fold increase of lysosomal fraction by Torin1 from Fig. 3 and simulated the dynamic lysosome PK model (Fig. 6A). The dynamic lysosome PK simulation suggested that this change in initial lysosome starting fraction accounts for the increased uptake of HCQ in the cells. For the new data set, the MDA-MB-468 T1(-) uptake data were simulated against the lower CI basal Vf_{lvs} (0.835%) and the T47D against the mean starting Vf_{lvs} (0.783%) from Fig. 1B. In MDA-MB-231 and 468 cells, the mean increase of lysosomes by Torin1 accounted for the increased cell uptake of HCQ in the simulation. In T47D and MCF7, the experimental mean increase in lysosomes by Torin1 from Fig. 3C was very high, so the ratio from the lowest replicate was used to show the increase the model would account for. Comparison of observed versus simulated AUC_{0-24 hour} are shown in Table 3. All simulation:experimental ratios except the T47D and MCF7 mean T1 were within the range of 0.5-2.0; however, T47D and MCF7 simulations using the lowest T1:Ctrl lysosome ratio from a replicate did fall within this range. Model performance metrics are shown in Supplemental Table 4. The model for each cell line followed similar PE trends to the earlier versions of the models. MAPE % for all cell lines suggests that the model fits overall time points between 15.48% and 61.69%, and MPE% suggests an even spread of over- and underprediction of the model across the cell lines. Overall, the cell PK model of HCQ supports that steadily increasing long-term whole-cell concentrations, as well as the increase in concentration caused by Torin1 pretreatment, are linked directly with the increase in size of the lysosomal compartment observed experimentally.

In addition, we ran the same experiment in MDA-MB-231 cells, except cells were pretreated with T1, T1 + MN, or MN alone prior to treatment with HCQ for 1 hour. We observed that the ratio of T1:Ctrl was not statistically significant from the ratio of MN-T1:MN-Ctrl (1.6), suggesting that this increase in cell uptake of HCQ is due to only the increase in lysosomal volume fraction induced by T1 (Fig. 6C). In MDA-MB-231, model simulation of the T1(+) versus T1(-) scenarios

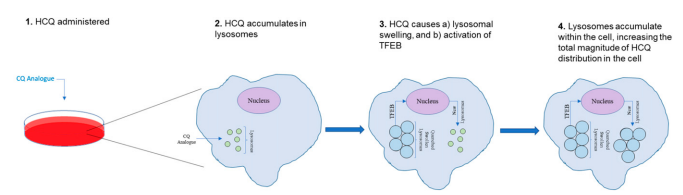


Fig. 4. HCQ increases the size of the lysosome compartment, which should increase its whole-cell drug uptake. This figure outlines the process by which HCQ increases its volume of distribution within the cell. 1) HCQ is initially added to the cell culture, after which 2) it immediately begins to accumulate within the lysosomes of the cells until it reaches chemical equilibrium between 30 and 60 minutes. 3) After reaching chemical equilibrium, the lysosomes begin to swell and simultaneously activate TFEB. 4) TFEB triggers the formation of new lysosomes, which undergo the same swelling process under static extracellular concentrations of HCQ. This hypothesized mechanism describes the rapid chemical equilibrium of HCQ within the cell, until it starts causing a dynamic increase in the size of the lysosome compartment, ultimately leading to this positive-feedback loop in which HCQ increases its own cellular volume of distribution.

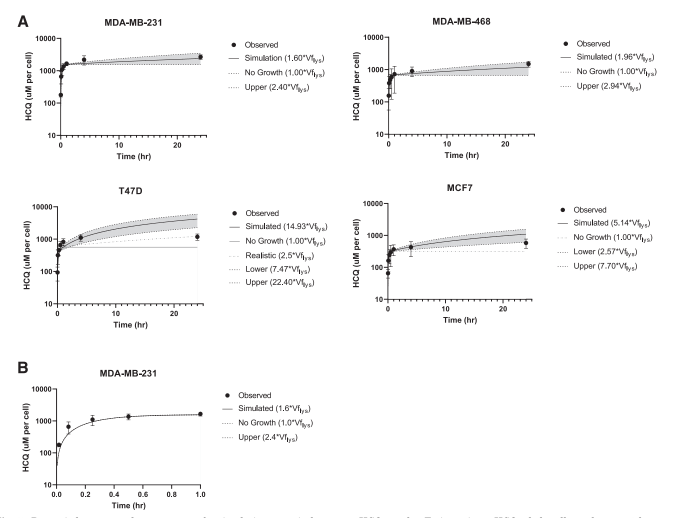


Fig. 5. Dynamic lysosome volume accounts for simulation error in long-term HCQ uptake. To investigate HCQ whole-cell uptake past 1 hour, we incorporated a growing lysosomal component into the model for all cell lines. Lysosomal growth is represented by a linear increase as a function of time, out to a maximum lysosomal volume of the fold increase observed with HCQ treatment of each cell line in Fig. 3C. The dynamic system model is shown (A) for MDA-MB-468, T47D, and MCF7, respectively. MDA-MB-231 and MDA-MB-468 simulated uptake is shown as the mean increase in experimental lysosomal volume after HCQ treatment, the upper value of the lysosomal volume, and with no growth incorporated. T47D is shown as the mean, upper, and lower increase in lysosomal volume as well as the value we would expect based on the PK data (realistic) and no growth. MCF7 is shown as the mean, upper, lower, and no growth. It should be noted that the time-based increase in lysosomal volume in the dynamic model does not affect earlier time points and thus would not affect model fit vs. the static model out to 1 hour (B).

represented an increase in total cellular uptake of HCQ but also showed that the lysosomal concentrations in both conditions was roughly the same (Fig. 6D). This suggests that even though the quantifiable concentration of HCQ from a PK study (whole-cell or tissue homogenate) may change, the concentration in the lysosome may be the same. This observation could have implications for PK:PD correlations, as the lysosome is the target site of HCQ for autophagy inhibition.

TABLE 2 Dynamic model metrics (AUC)

AUC Summary (0–24 h) Cell Line	Experimental (Exp) Mean	Simulated (Sim) Mean	Mean Sim/Exp
MDA-MB-231	55,660	46,753	0.84
MDA-MB-468	26,922	22,085	0.82
T47D	26,340	56,178	2.13
T47D (realistic)		21,198	0.80
MCF7	11,553	16,823	1.46
MCF7 (lower)		10,975	0.95

Discussion

Lysosomes have been observed as a drug sink for many lysosomotropic agents, including many prescribed drugs, such as doxorubicin, vinblastine, and imipramine, and a variety of TKIs, such as gefitinib (Skoupa et al., 2020), imatinib (Burger et al., 2015), and sunitinib (Ruzickova et al., 2019). In a recent study by Ruzickova et al., the lysosomotropic agent sunitinib was shown to be capable of increasing overall lysosomal sequestration of the TKIs tested, although this effect was insufficient in reducing sensitivity to the TKIs by the hypothesized mechanism of sequestering them away from their mechanistic target sites as the concentrations tested in culture maintained a static extracellular concentration. The in vitro PK model of HCQ supports this finding and would suggest the same observation: the direct increase of lysosomal volume fraction by pretreating with Torin1 (Fig. 6C) barely increases HCQ concentration in the lysosome even though the whole-cell concentration is higher. This observation sheds light onto one commonly suggested hypothesis for

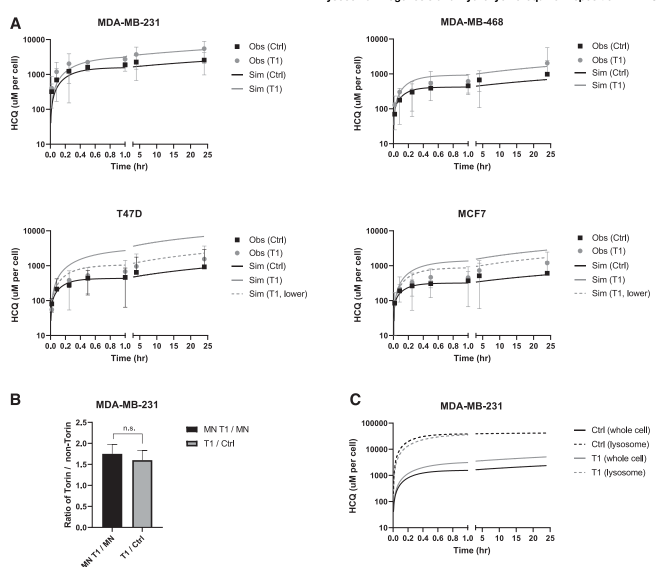


Fig. 6. Torin1 increases the size of the lysosomal compartment, increasing whole-cell uptake of HCQ. Cell lines were pretreated with T1 using the same concentration and time as in Fig. 3. After Torin1 pretreatment, the HCQ PK studies from 1 minute to 24 hours were repeated in Torin1(+) vs. Torin1(-) cells (Obs). Torin1 pretreatment resulted in an average increase of HCQ whole-cell concentrations at all time points by an average of 1.4- to 1.6-fold. The mathematical PK model of HCQ (Sim) was tested at basal lysosomal volume fractions vs. Torin1-modified starting lysosomal volume fractions by multiplying the basal lysosomal volume fraction in each cell line by the mean increase by Torin1 treatment from Fig. 4D (A). MDA-MB-231 [(A), top-left] fit the data well using mean values of Torin1. MDA-MB-468 used the starting lysosomal volume fraction value of the lower 95% CI (0.753%) and mean value of Torin1 increase [(A), top-right]. T47D used the mean starting Vf_{lys} (0.783%) and was tested against the mean Torin1 increase in lysosomes (10.5×) as well as the lowest observed ratio within the replicates (3.8×) [(A), bottom-left]. MCF7 was tested against the mean lysosomal increase by Torin1 and the lowest observed ratio within the replicates (3.8×) [(A), bottom-right]. To test whether the increase in uptake by Torin1 was only due to the increase in lysosomes, MDA-MB-231 HCQ uptake was tested at 1 hour with no pretreatment, Torin1 pretreatment, monensin pretreatment, or Torin1 and monensin pretreatment. No significant change in the ratio between T1:Ctrl and MN + T1:MN HCQ uptake was observed (B) with a two-tailed unpaired t test (P = 0.573). Comparing the simulated lysosomal concentrations vs. the whole-cell uptake concentrations in MDA-MB-231 cells with T1(+) or (-) shows a minimal difference in lysosomal concentration in both scenarios (C).

HCQ growth-inhibition synergism with other lysosomotropic agents in cell culture, which is that HCQ displaces drugs from the lysosome, reducing overall sequestration and increasing concentration at target sites elsewhere within the cell. This work, in conjunction with other recent findings, would suggest that under static extracellular concentrations, this hypothesized mechanism is unlikely, as an overall reduced cellular uptake of lysosomotropic drugs may be due to displacement by HCQ but would not result in a subsequent increase in concentration at the drug's mechanistic target. This does not eliminate this hypothesized mechanism in vivo, though, as extracellular concentrations are not constant.

Many drugs have been shown to activate TFEB in vitro in recent publications (Lu et al., 2017; Zhitomirsky et al., 2018; Zhao et al., 2020), most of which are weakly basic lipophilic agents that would be physicochemically characterized as lysosomotropic. If lysosomotropism is a major factor in the PK of these drugs, as it is for HCQ, then the distribution of these drugs might be altered by their effect on the lysosome as we have observed for HCQ. Our findings emphasize care in cell-level PK studies of drugs with weakly basic lipophilic structures, especially when attempting to correlate concentration with response. This is especially important when using multiple lysosomotropic agents in combination.

TABLE 3
Torin1 dynamic model metrics (AUC)

AUC Summary (0–24 h) Cell Line	Experimental (Exp) Mean	Simulated (Sim) Mean	Mean Sim/Exp
MDA-MB-231 (Ctrl)	56,214	46,753	0.83
MDA-MB-231 (T1)	104,084	99,813	0.96
MDA-MB-468 (Ctrl)	18,490	13,141	0.71
MDA-MB-468 (T1)	29,517	30,276	1.03
T47D (Ctrl)	17,692	15,304	0.87
T47D (T1)	28,044	119,838	4.27
T47D (T1, lower)		39,299	1.40
MCF7 (Ctrl)	12,849	10,284	0.80
MCF7 (T1)	21,471	49,679	2.31
MCF7 (T1, lower)	,	30,585	1.42

Regardless, the possibility of occurrence of this lysosomealtered distribution is supported by in vitro studies in other lysosomotropic agents, whether it be through the activation of lysosomal biogenesis or through lysosomal fusion that is independent of biogenesis (Skoupa et al., 2020), but with limited data to suggest it may occur in vivo. HCQ is somewhat unique in this context because of its long half-life combined with often prolonged dosing regimens, which means that tissue will be exposed to relatively high concentrations of the drug for an extended duration. Extended duration of exposure to tissues might exacerbate the lysosomotropic effects due to HCQ that may not necessarily be prevalent with other drugs that are dosed more short term. We are unaware of any longterm preclinical PK studies involving HCQ that investigate multiple time points aside from the work by McChesney (1983). This study looked at a variety of different tissues in rats that were treated with HCQ 6 days per week for months, with time points at 1, 2, 3, and 7 months. The key observation from this study is that HCQ concentrations continually increase up to 3 months, at which time dosing is discontinued in the first study. In the 7-month study, HCQ only starts to reach tissue steady state between 3 and 7 months in lung, eve, and muscle but continues to gradually increase in concentration for other tissue. This is particularly interesting because the plasma half-life of HCQ in rats has been observed at around 10.6 hours (Moore et al., 2011), suggesting a disconnect between plasma and tissue steady-state values. The study by Wei et al. (1995) also investigated HCQ dosing in rats and sampled multiple time points of blood over the course of 10 weeks, but it only sampled tissue at the 10-week mark. Interestingly, of the three dosing regimens applied to both male and female rats (6 conditions total), only the regimen with 8 mg/kg in male rats appeared to increase in blood concentration after the first time point at 3 weeks, but it remained steady after that. For the rest of the dosing regimens, HCQ appeared to remain constant, which would suggest that steadily increasing tissue concentrations of HCQ would be undetected by the traditional PK sampling of blood in clinical studies. With limited in vivo data available, we would speculate that lysosomal biogenesis plays a role in longterm tissue uptake, although this effect would likely be less pronounced because of overall lower HCQ concentrations.

This work has implications in HCQ PK, both in vitro and in vivo, particularly in the context of the cancer clinical trial in solid tumors by Barnard et al. (2014), in which there was no correlation between HCQ concentrations in the tumor and plasma. Further data are needed to investigate the concentration/time relationship of this effect and possible maximum

values of lysosome volume fractions that are achievable in vitro. For this study, we modeled the lysosomal compartment as increasing linearly with time, although it is more likely that this effect follows a hyperbolic trend—increasing rapidly and leveling off to eventually reach a pseudomaximum. It is also likely that this effect is due to overall exposure to HCQ (AUC) rather than time or concentration alone, but to investigate that, a significant number of time points and concentrations are necessary in conjunction with subsequent PK analysis to maintain a connection between the lysosomal biogenesis mechanism and PK outcome. In addition, a comprehensive PK/PD study connecting increasing tissue concentrations over the long term with an increase in lysosomal volume of tissues is warranted to truly confirm whether this phenomenon is important to consider in vivo.

Authorship Contributions

Participated in research design: Collins, Gustafson.

Conducted experiments: Collins, Witta.

Contributed new reagents or analytic tools: Collins, Coy, Pang. Performed data analysis: Collins.

Wrote or contributed to the writing of the manuscript: Collins, Gustafson.

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