Substitution of Pro165 in Transmembrane 4 of PfCRT Abolishes Lysosome Acidification Function in Stably Transfected HEK-293F Cells

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Authors’ contributions

This work was carried out in collaboration between both authors. Both authors read and approved the final manuscript.

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ABSTRACT

The Plasmodium falciparum chloroquine resistance transporter (PICRT) is localised on the parasite digestive vacuole, an organelle that maintains an acidic lumen. Here we demonstrated the isolation of HEK-293F cells stably expressing wild type 3D7 and mutant Pfcrt alleles. Immuno-fluorescence staining of HEK-293F transfectants confirms the localization of Pfcrt alleles to the lysosomal vesicles. Moreover, cells expressing mut-PICRTETSE showed greater lysosomal acidification as demonstrated by the dramatic increase in the accumulation of two weak bases, acridine orange and CytPainter LysoOrange dyes. Furthermore, HEK-293 cells stably expressing mut-PICRTETSE with a single substitution of proline 165 in transmembrane 4 completely inhibited the accumulation of weak bases. Taken together, our results demonstrate the role of Pro165 in PICRT lysosomal acidification function in HEK-293F cells.

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1. INTRODUCTION

The human malaria is a major public health problem in tropical countries with Plasmodium falciparum infections accounting for the majority of the total malaria cases [1]. The progression of antimalarial treatment from the natural product quinine to chloroquine (CQ) extended the effectiveness of antimalarials for nearly 20 years [2]. However, the rise of CQ resistant parasites, largely due to mutations in Plasmodium falciparum Chloroquine Resistant Transporter (pfcrt), was a major setback given the safety and efficacy of this drug [3]. Several mutant alleles of pfcrt have been identified in CQ-resistant field isolates [4], including the key CQ-resistance causative mutation at position 76 (Lys76Thr) of PfCRT [5]. PfCRT encodes a 424 amino acids polypeptide, with 10 transmembrane (TM) helices, localized to the digestive vacuole (DV) of the parasite [6]. A recent report, using single-particle cryo-electron microscopy, revealed the three-dimensional structure of PfCRT at a resolution of 3.2Å [7]. PfCRT structure revealed eight transmembrane helices (TMs 1-4 and 6-9) forming the central cavity of 25Å diameter with the resistance causative mutations localized to TM helices of the central cavity [8]. Transmembrane helices 4 and 10 are positioned outside the central cavity and thought to contribute to the “open and close” conformations on the vacuolar side; while TMs 4 and 9 play the same role on the opposite side of the membrane in a “rocker-switch” motion [9].

Based on PfCRT resolved 3D structure, it was of interest to examine the effects of substituting two highly conserved proline residues (Pro165 and Pro354) in PfCRT TMs 4 and 9 on its lysosomal acidification activity in HEK-293F cells. In this report, we show the effects of specific substitutions in mut-pfcrt allele on its in-situ lysosomal acidification function and propose that this function of mut-PfCRT in HEK-293 cells can act as a segregate assay for its CQ-resistance role in P. falciparum.

2. MATERIALS AND METHODS

2.1 Cloning of Codon Optimized PFCR Mutants

Wild-type pfcr (3D7) encoding the CQ sensitive gene sequence obtained from plASmo DB (gene ID PF3D7_1133400) was human codon optimised and commercially synthesized (Genscript®, USA). The full-length pfcr coding sequence was excised from pUC57 vector and re-ligated into pCDNA3.1+ (Invitrogen™) at restriction sites BamHI and EcoRI (New England Biolabs Inc.) with a Kozak consensus sequence included. Pfcr allele encoding mutations shown to confer CQ resistance (mut-PfCRT^ETSE), as described earlier by Summers et al. [4]; D17 variant) was constructed by ligating 3 fragments, amplified using the Phusion High-Fidelity DNA Polymerase (Life Technologies™), possessing 4 point mutations (e.g., N75E, K76T, A220S & Q271E) into the humanised 3D7-pfcr (wt-PfCRT^NKAQ) gene sequence using the Gibson assembly approach [10]. The latter approach was also used to introduce mutations at positions Pro165 to Ala (P165A) and Pro354 to Ala (P354A) in D17-pfcr allele (or mut-P165A-PfCRT^ETSE and mut-P354A-PfCRT^ETSE). For primers sequences and reaction conditions see tables I and II (Supplementary material).
2.2 Tissue Culture and Selection of Stable Transfectants

HEK-293F cells were cultivated and maintained in DMEM culture media (Gibco™) containing 10% (v/v) fetal bovine serum (FBS) at 37°C. Cells were grown to 70-80% confluency prior to transfection. Transfections, using pCDNA3.1+ empty plasmid, full-length wt-PfCRTNKAQ, mut-PfCRTETSE, mut-P165A-PfCRTETSE and mut-P354A-PfCRTETSE containing plasmids, were linearized with MfeI restriction enzyme (New England Biolabs Inc.) and transfected into HEK-293F cells using Lipofectamine 2000 (Invitrogen™), following manufacturer’s instructions. Clones from several transfections were selected in 800 µg/ml G418 in 10 % (v/v) FBS/DMEM and allowed to proliferate for several weeks prior to freezing aliquots from each clone in liquid nitrogen. Two clones for wt-PfCRTNKAQ (clone 2 and 4), two clones for mut-PfCRTETSE (clones 8 and 9) and one clone for the mut-P165A-PfCRTETSE were isolated and allowed to proliferate continuously in the presence of G418 prior to characterization for PfCRT (wt or mutant) expression and drug transport studies. Unfortunately, it was not possible to obtain additional clones for mut-P165A-PfCRTETSE or any clones for mut-P354A-PfCRTETSE, after several transfection and selection attempts.

2.3 Protein Extraction and Western Blotting

For each clone of HEK-293F transfectants, 3X10⁶ cells were washed in PBS and extracted in RIPA buffer (50 mM Tris-HCl pH 7.4, 1% (v/v) NP40, 0.25% (w/v) sodium deoxycholate in the presence of protease inhibitors). Protein extracts were resolved on 10% SDS-PAGE [11] and transferred to PVDF membranes [12]. Membranes were probed for PfCRT expression using rabbit anti- serum raised against PfCRT N-terminal sequence (1-58 polypeptide) at 1:6000 (v/v) dilution (Baakdah F. and Georges E., unpublished results). All PVDF membranes were probed with anti-tubulin monoclonal antibody (1 µg/ml) for equal protein loading. Following overnight incubation at 4°C with primary antibodies, PVDF membranes were washed with PBS and incubated with horse radish peroxidase (HRP)-conjugated secondary antibodies (1:3000 (v/v)) diluted in 3% (w/v) milk/PBS. Membranes were washed with PBS and developed using Thermo Scientific™ SuperSignal™ West Pico Chemiluminescent Substrate. Images were captured with ChemiDoc imaging system from BIO-RAD Inc.

2.4 Indirect immunofluorescence Assay

Cells were cultivated in tissue culture onto poly L-lysine coated cover slips. The cover slips containing cells were fixed with 4% paraformaldehyde-PBS (Electron microscopy sciences) for 30 minutes at room temperature (RT). Fixed cells were washed once in PBS then quenched with 0.15% (w/v) glycine-PBS for 10 minutes at RT. Cells were washed and permeabilized with 0.1% (v/v) TritonX-100/PBS (Bio Basic Inc.) for 10 minutes at RT. Permeabilized cells were washed three times with PBS and blocked with 1% (v/v) goat serum in PBS for 1hr at 4°C. Cells were incubated with PfCRT antiserum and/or anti-sirtuin1 or anti-LAMP1 monoclonal antibodies (DSHB, at the University of Iowa, Department of Biology, USA) at 4°C overnight followed by PBS wash and probing with secondary antibody Alexa-fluor 594-conjugated goat anti-rabbit (Life Technologies™) for anti-PfCRT and Alexa-fluor 488-conjugated goat anti-mouse for the primary antibodies for 45min. The cover slips were washed three times in PBS, dipped once in distilled water, air dried and mounted on microscope slides using fluor mount G mounting medium (Southern Biotech). Images were captured using confocal microscopy (Carl Zeiss GmbH, Jena, Germany).

2.5 Dye Accumulation Assay

Cells were detached using 0.25% (w/v) trypsin in 0.5mM (w/v) EDTA solution. A total of 3 X 10⁶ cells, in two sets for each cell line, were washed in HEPES-Hanks Balanced Salt Solution (HBSS) buffer (20mM (w/v) HEPES, pH 7.2 and 1X HBSS), filter sterilized and preheated at 37°C. One set was re-suspended in 1ml of 30mM NH₄Cl HHBS buffer pH 7.2, while the second set was re-suspended in 1ml HHBS, and placed for 30minutes at 37°C. The two dyes (acridine orange (AO), Invitrogen™) or CytiPainter LysoOrange (LO) Indicator reagent, abcam® were added to cells (4µM for AO and 2units/ml for LO) and incubated for one hour at 37°C. After centrifugation, the resulting cell pellets of set 1 were re-suspended in 10ml 30mM NH₄Cl HHBS buffer and set 2 were re-suspended in 10ml HHBS buffer for 1hr at 37°C. The final cell pellets were re-suspended in 1ml wash buffer and aliquoted into a round bottom black 96 well plate. Fluorescence was measured using Synergy H4
hybrid spectrophotometer from BioTek® and Gen5™ microplate reader and image software, according to each dye respective wavelength (AO Eg 485/Er 530 and LO Eg 542/Er 565). The results for dye accumulation values were adjusted for PICRT expression and equal loading based on tubulin levels minus dye accumulation in vector transfected HEK-293F cells.

2.6 Homology Modeling

The SWISS-MODEL tool [13] was used to generate the PDB file for mut-P165A-PICRTETSE using model-template PDB 6ukj [14] and the homology models were built and edited in PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC.

2.7 Statistical Analysis

Plotted results were analysed by one way ANOVA using GraphPad Prism version 8.4.0 (671).

3. RESULTS AND DISCUSSION

PICRT-mediated drug transport studies in P. falciparum have been hampered by the complexity of measuring meaningful drug transport across several membranes in infected erythrocytes [15]. Hence, efforts to establish a simple in-situ heterologous expression system for PICRT in HEK-293F cells stably expressing wild-type and mutant forms of PICRT would greatly facilitate PICRT structure-functions studies. Figure 1A shows a schematic of PICRT secondary structure with 10 transmembrane domains and the positions of amino acid residues mapped onto the wild-type PICRT (encoding N75, K176, A220 & Q217ETSE) or wt-PICRTNKAQ), mutant-PICRT encoding four mutations shown previously to be sufficient to confer resistance to CQ (N75E, K76T, A220S & Q217E; or mut-PICRTETSE), and mutant-PICRTETSE containing an additional proline to alanine mutation at position 165 (mut-P165A-PICRTETSE). Human codon optimized full-length constructs of wt-PICRTNKAQ, mut-PICRTETSE or mut-P165A-PICRTETSE were linearized and transfected into HEK-293F cells. Cells were selected with 800µg/ml G418, and isolated clones were allowed to proliferate and extracted for protein analyses. Figure 1B shows Western blot results of HEK-293F extracts from cells transfected with vector alone (Vc1), vector containing wt-PICRTNKAQ (clones C2 and C4), mut-PICRTETSE (clones C8 and C9) and mut-P165A-PICRTETSE then probed with PICRT N-terminal-specific antibody (PICRT antiserum). The results in lanes 2-5 of figure 1B show a 42kDa polypeptide encoding wt-PICRTNKAQ, mut-PICRTETSE and mut-P165A-PICRTETSE expressed to varying levels. Interestingly, and consistently, total protein extracts from mut-PICRTETSE clones expressed significantly less PICRT protein than extracts from wt-PICRTNKAQ transfectants (Fig. 1B, lanes 2-3 versus 4-5). Moreover, the mut-P165A-PICRTETSE polypeptide consistently migrated slightly faster on SDS-PAGE than wt-PICRTETSE or mut-PICRTETSE (Fig. 1B, lanes 2-5 versus 6). In addition, lanes 2, 3 and 6 revealed additional polypeptides migrating with molecular masses.

with molecular masses of ~52kDa and ~90kDa with varying intensities proportionate to the intensity of the 42kDa band suggesting that they may represent a post-translationally modified PICRT (e.g., 52kDa) or a PICRT homodimer (e.g., 90kDa) [17]. The latter speculations are strengthened further since both the 52kDa and 90kDa polypeptides in lanes 2 and 3 (wt-PICRTNKAQ) appear to migrate slightly faster in lane 6 (mut-P165A-PICRTETSE) mimicking the slight increase in mobility of the 42kDa polypeptide. The significance of the 52kDa and 90kDa PICRT proteins is presently not clear but appears to be present at lower levels than the 42kDa form of PICRT.

Previous heterologous expression of wild-type HB3 and Dd2 mutant-picrt alleles into HEK-293 cells showed lysosomal localization, an equivalent organelle to the DV in P. falciparum [18]. To determine the subcellular localization of PICRT in HEK-293F cells stably expressing wt-PICRTNKAQ and mut-PICRTETSE, cells were subjected to double immunofluorescence staining (IFA) with anti-PICRT and anti-LAMP1 (established lysosomal protein [19]) or anti-sirtuin 1 (a nuclear protein [20]) antibodies, respectively. Figure 2 shows punctate staining with anti-PICRT and anti-LAMP1 in the cytosol and outside the cell nuclei that perfectly overlap. By contrast, and as expected, anti-sirtuin 1 stained cell nuclei and did not overlap with anti-PICRT staining (Fig. 2). Similar results were observed for mut-PICRTETSE and mut-P165A-PICRTETSE (not shown). It is interesting that, unlike in HEK-293F and P. falciparum, PICRT expressed in yeast and Xenopus laevis oocytes was shown to localize to the plasma membrane [21]. Differences in PICRT subcellular localization...
between yeast and *Xenopus laevis* oocytes versus *P. falciparum* and HEK-293F cells is not entirely clear but may be due to differential post-translational modification. With respect to the latter, it has been suggested that phosphorylation at threonine 416 in PICRT C-terminal mediates its sorting to the DV membrane in *P. falciparum* [22]. Hence, it will of interest to determine if PICRT is post-translationally modified in HEK-293 cells and if mutations of such post-translationally modified sites alter its subcellular localization. Given earlier findings [23] relating to increased lysosomal acidification in HEK-293 cells transiently expressing mutant Dd2-associated pfCRT allele, we made use of two lysosome fluorescent dyes (i.e., AO and LO) to validate this function of mutant-PfCRT using HEK-293F transfectants stably expression PfCRT [24].

**Fig. 1.** Stable expression of wild-type and mutant-PfCRT in HEK-293F cells. Panel A shows a schematic representation of full-length PICRT as a solid bar with transmembrane domains indicated as blue filled rectangles (1-10). The positions of various amino acid residues found in wild-type 3D7 PICRT (wt-PICRTNKAQ), mutant-PICRT (mut-PICRTETSE) and Pro165Ala substituted mutant-PICRTETSE (mut-P165A-PICRTETSE) are localized onto PfCRT schematic using single letter code. Panel B shows cell lysates from HEK-293F cells transfected with vector alone (Vc1; lane 1), vector containing wt-PICRTNKAQ (clones C2 and C4; lanes 2 and 3), mut-PICRTETSE (clones C8 and C9; lanes 4 and 5) and mut-P165A-PICRTETSE (lane 6) were resolved on 10% SDS-PAGE, transferred to PVDF membrane, and probed with PfCRT antiserum and tubulin monoclonal antibody. PfCRT polypeptide migrated with an apparent molecular mass of 42kDa seen in all HEK-293F transfectants (lanes 2-6), except in vector transfected HEK-293F (Vc1; lane 1). The tubulin polypeptide band, loading control, is detected in all samples (lanes1-6)
Fig. 2. Immuno-fluorescence staining of PfCRT in HEK-293F cells. PfCRT-HEK clones were prepared for IFA and images were taken using confocal microscopy. Panels C and G show cells expressing wt-PfCRT\textsuperscript{NKAQ} when stained with PfCRT antiserum; while Panels B and F show cells stained with anti-sirtuin1 or anti-LAMP1, respectively. Panels D and H show the overlay of Panels B+C and D+G, respectively. Panels A and E show the phase contrast images of cells expressing wt-PfCRT\textsuperscript{NKAQ}. The scale bar on each panel reads 10μm.

The results in figure 3A show the relative accumulation of AO in HEK-293F cells stably transfected with wt-PfCRT\textsuperscript{NKAQ} (clones C2 and C4) and mut-PfCRT\textsuperscript{ETSE} (clones C8 and C9) relative to vector transfected cells adjusted to PfCRT protein expression levels. These results show dramatic increase in AO accumulation in clones C8 and C9 expressing mut-PfCRT\textsuperscript{ETSE} relative to clones C2 and C4 which express wt-PfCRT\textsuperscript{NKAQ}. This increased AO accumulation seen in clones C8 and C9 is likely due to the enhanced acidification of the lysosomal vesicles by mut-PfCRT\textsuperscript{ETSE} (Fig. 3A). Moreover, short-term pre-incubation of cells expressing both wt-PfCRT\textsuperscript{NKAQ} and mut-PfCRT\textsuperscript{ETSE} in 30mM NH\textsubscript{4}Cl completely abolished the accumulation of AO in all cells relative to vector transfected HEK-293F cells, confirming the role of differential proton gradient on AO accumulation (Fig. 3A). Similar results were seen using a different lysosome targeting dye, LO (Fig. 3B), confirming the effect of mut-PfCRT\textsuperscript{ETSE} on the acidification of lysosomes in HEK-293F cells (clones C8 and C9), relative to wt-PfCRT\textsuperscript{NKAQ} expressing clones (C2 and C4). The mutations in PfCRT selected in this study (i.e., mut-PfCRT\textsuperscript{ETSE}) were the same mutations that increased the uptake of CQ in Xenopus laevis oocytes, but are three fewer mutations than Dd2-PfCRT [25] used in an earlier study to demonstrate lysosomal acidification in HEK-293 cells [26]. Hence, we speculate that there may be a correlation between PfCRT ability to transport CQ into Xenopus laevis oocytes (which is consistent with CQ resistance based on PfCRT’s orientation in the DV membrane) and lysosome acidification in HEK-293 cells [27]. The latter speculation is supported by earlier observation in Dictyostelium discoideum, whereby mutant-PfCRT expelled CQ and caused a significant intra-vesicular acidification of the acidic vesicles [28]. Moreover, it was previously shown that mutant alleles of pfcrt from CQR strains (e.g., 7G8 and Dd2) conferred increased acidification of the DV relative to CQS strains [24].

To confirm the direct role of mut-PfCRT in lysosomal acidification in HEK-293 cells and to determine the effects of specific amino acids substitutions on PfCRT segregate function in HEK-293F cells (i.e., lysosomal acidification), it was of interest to examine the effects of substituting two proline residues (Pro165 and Pro354) on mut-PfCRT function.
Fig. 3. Accumulation of AO and LO in HEK-293F cells stably expressing wild-type and mutant-PfCRT. HEK-293F cells stably expressing wt-PfCRT<sup>NKAQ</sup> (clones C2 and C4), mut-PfCRT<sup>ETSE</sup> (clones C8 and C9) and mut-P165A-PfCRT<sup>ETSE</sup> were incubated with AO (Panel A) or LO (Panel B) without and with 30mM NH<sub>4</sub>Cl, respectively. The relative accumulation of AO and LO, in arbitrary fluorescence units, is shown on the Y-axis. Results represent experiments from three independent repeats done in triplicates. Statistical significance (****P value <0.0001) was analysed by one way ANOVA using GraphPad Prism.

The rationale for selecting Pro<sup>165</sup> and Pro<sup>354</sup> was based on several points: a) both residues are highly conserved in Plasmodium orthologues of PfCRT, b) their localization in TM helices 4 and 9 is likely to affect helical distortion and flexibility of these domains, and c) proline residues in α-helical TMDs play structural and/or functional roles in polytopic membrane proteins whereby the geometry and the limited hydrogen bonding of this residue introduce a molecular joint or hinge that could be important for function [29]. Therefore, proline residues in TM helices can have important structural and/or functional roles in membrane proteins. Moreover, substitution of proline by alanine, an amino acid with the highest helix propensity, contrary to proline, should have minimal effects on PfCRT structural integrity and would eliminate the possible molecular hinge introduced by proline [28]. The results in figure 3A and 3B show the effect of Pro165Ala on AO...
and LO accumulation in HEK-293 cells expressing mut-P165A-PICRTETSE. The substitution of Pro165 for Ala-165 in TMD4 of mut-PICRTETSE completely abolished its lysosome acidification function to the same level as wt-PICRTNKAQ as measured by the accumulation of AO and LO in HEK-293F cells (Fig. 3A and 3B, respectively). Furthermore, treatment of cells with NH₄Cl caused further decrease in AO and LO accumulation, relative to control vector transfected HEK-293F cells (Fig. 3A and 3B, respectively). It is interesting to note that unlike mut-PICRTETSE HEK-293F transfectants, both Pro165Ala-PICRTETSE and wt-PICRTNKAQ HEK-293F transfectants expressed higher levels of PICRT which could be related to their inability to function as lysosomal acidifiers and as such higher expression levels are tolerated. Although it is not entirely clear how substituting Pro165 for Ala inhibits mut-PICRTETSE ability to acidify the lysosomes, it has been suggested that residues in TMD 4 and 9 may contribute to the “open and close” conformations at the vacuolar side of the DV, while residues in TMD 5 and 10 may play the same role on the opposite side of the membrane in a “rocker-switch” movement [9]. Based on the resolved structure of 7G8 mutant-PICRT [30], Pro165 is localized midway in TM helix 4 facing outside the central cavity formed by TMDs 1-4 and 6-9, but in close contact with TM helix 5 (Fig. 4). In support of the latter speculation, figure 4 shows a slight shift in TM4 helix with Ala165 versus Pro165 which may affect PICRT proposed “rocker-switch” movement and consequently its function. Similarly, P354 is also localized roughly midway of TM helix 9 facing outside pore cavity, and in close vicinity of TMD 10 (Fig. 4).

Fig. 4. PICRT homology modeling with P-165-A substitution in mut-PICRT. The homology modeling of PICRT was done based on PICRT-7G8-Pro165 TMD’s crystal structure (PDB 6UKJ). In blue are the TMDs of mut-PICRT-7G8 overlapping with the gray TMDs of mut-P165APICRTETSE. In yellow is the proline (yellow arrow) and in red is the alanine (red arrow)
Together, it is likely that TMD 4 and 9 modulate PICRT transport function through their interactions with TMD 5 and 10. Unfortunately it was not possible to test the effect of Pro-365 substitution on PICRT function. Moreover, it would be of interest to test the role of Pro165 substitution on the ability of mutant-PICRT to confer resistance to CQ in the parasite. Given that PICRT is likely to continue performing its normal function in CQ-resistant strains, it would be equally important to test if substituting Pro165 with Ala affects both the wt-type and mutant-PICRT functions in the parasite’s DV.

PICRT is a member of the drug/metabolite transporter superfamily and may act as a proton-coupled transporter as other members of the family [31]. However, it was reported that parasite strains encoding CQ-resistant mutants of PICRT allow leakage of protons from the DV [32]. The latter results are at odds with our findings and those by Reeves et al. [16]. However, it is likely that in HEK-293 cells, mut-PICRTETSE does not transport the same substrate in the parasite DV. Moreover, mut-PICRTETSE in HEK-293F does not seem to mediate the efflux of CQ from lysosomes (results not shown and [16]) and consequently, proton-leakage from the parasite DV appears to be associated with CQ-efflux. We speculate that mut-PICRTETSE expressed in HEK-293F may not undergo similar post-translational modification to that in P. falciparum and this could affect its CQ-mediated drug resistance and transport functions. In line with the latter possibility, a recent report demonstrating a 50% reduction in CQ resistance following the substitution of Ser33 to Ala, but not Asp phospho-mimic, in Dd2 P. falciparum isolate [30].

4. CONCLUSION

The results of this study describe the use of HEK-293F human cell line stably expressing a heterologous malaria membrane protein that localizes to the lysosomal membrane, an equivalent organelle to the digestive vacuole in P. falciparum, hence establishing PICRT-HEK-293F cells as a heterologous expression system to study PICRT structure-function. Moreover, consistent with an earlier study, we demonstrate the ability of mutant PICRT (mut-PICRTETSE, a chloroquine competent transporter) can cause the acidification of lysosomes in HEK-293F cells. Lastly, we show for the first time that mutations of certain amino acids (e.g., Pro165 to Ala) in transmembrane 4 of PICRT abolishes the ability of mut-P165A-PICRTETSE to acidify the lysosomal compartment in HEK-293F cells. Work in progress is focused on establishing HEK-293F cells stably expressing wild-type and mutant PICRT as a segregate system to study the mechanism of PICRT mediated CQ-resistance.

DISCLAIMER

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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