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# Research paper

# Identification of a novel DGK $\alpha$ inhibitor for XLP-1 therapy by virtual screening



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#### ABSTRACT

As part of an effort to identify druggable diacylglycerol kinase alpha (DGK $\alpha$ ) inhibitors, we used an insilico approach based on chemical homology with the two commercially available DGK $\alpha$  inhibitors R59022 and R59949. Ritanserin and compound AMB639752 emerged from the screening of 127 compounds, showing an inhibitory activity superior to the two commercial inhibitors, being furthermore specific for the alpha isoform of diacylglycerol kinase. Interestingly, AMB639752 was also devoid of serotoninergic activity. The ability of both ritanserin and AMB639752, by inhibiting DGK $\alpha$  in intact cells, to restore restimulation induced cell death (RICD) in SAP deficient lymphocytes was also tested. Both compounds restored RICD at concentrations lower than the two previously available inhibitors, indicating their potential use for the treatment of X-linked lymphoproliferative disease 1 (XLP-1), a rare genetic disorder in which DGK $\alpha$  activity is deregulated.

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# 1. Introduction

The atypical lipid kinases DGK (diacylglycerol kinases) catalyze the phosphorylation of diacylglycerol (DAG) to phosphatidic acid (PA) (Fig. 1) [1].

Both these molecules act as second messengers regulating a plethora of biological processes such as cell proliferation and motility, immune responses, glucose metabolism, and cardiovascular responses [2].

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DGKs permit to control the cellular concentration of these two bioactive lipids, terminating the DAG signaling and increasing the concentration of PA [3], and it is not surprising that an up- or downregulation of DGKs has been linked with several pathological conditions such as cancer, epilepsy, Parkinson's disease, bipolar disorders, cardiac hypertrophy, type II diabetes, and hypertension [4]. Among the ten human isoforms of DGK ( $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\varepsilon$ ,  $\zeta$ ,  $\eta$ ,  $\theta$   $\iota$ , and  $\kappa$ ) [5], the most studied one is the  $\alpha$  isoform which is particularly expressed in the central nervous system, spleen, thymus, T cells, and lungs. Interestingly, it has been shown that the inhibition of the α form can be a valuable strategy for the discovery of novel anticancer compounds [6]. Indeed, this enzyme is overexpressed in several malignancies, particularly in hepatocellular carcinoma and melanoma cells, showing pro-tumoral and anti-immunogenic properties, and its inhibition damages cancer cells viability, blocks the angiogenetic process and, remarkably, it increases the T cells activation [7,8].

Another disease linked with overactivation of DGKα is the X-

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R1: R2= fatty acids

Fig. 1. DAG is phosphorylated by diacylglycerol kinases to phosphatidic acid.

linked lymphoproliferative disease type 1 (XLP-1) a primary immunodeficiency of genetic origin characterized by dysgammaglobulinaemia and poor CD8+ cytotoxic T cell function, with some patients developing lymphomas of the bowel, peritoneum, and brain [9]. Most importantly, XLP-1 patients face significant morbidity and mortality in response to Epstein-Barr virus (EBV) infection, which causes marked lymphoproliferative disease, featuring fulminant infectious mononucleosis (FIM), cytopenia, liver failure, hepatosplenomegaly, and/or hemophagocytic lymphohistiocytosis [10]. XLP-1 is due to loss of function mutations of the 128-amino acid signaling lymphocytic activation molecule (SLAM)—associated protein (SAP) adaptor molecule. Loss of SAP in XLP-1 patients impairs both adaptive and innate immunity, resulting in several immunologic alterations. Notably, a block in NKT-cell development [11]; decreased NK and CD8<sup>+</sup> cytotoxicity, specifically toward EBV-infected B-cell targets [12]; impaired CD4<sup>+</sup> T helper function and B cell conjugation, resulting in reduced germinal center formation and ineffective antibody production [13] and resistance to restimulation induced cell death (RICD), a homeostatic mechanism preventing excessive T cell proliferation [14]. Interestingly, the signaling alterations reported in SAP deficient cells are consistent with a reduced DAG signaling due to increased DGKα activity [15]. Indeed, TCR signaling inhibits DGKα through a signaling pathway that requires PLC activity and calcium but also the expression of SAP [15]. Thus, T cells from XLP-1 patients, which lack functional SAP, feature deregulated DGKα activity, which is a main contributor of the defective lymphocyte responses observed both in XLP-1 patients and in SAP KO mice. Indeed, pharmacological inhibition of DGKα restores RICD in lymphocytes from XLP-1 patients and prevents the excessive CD8<sup>+</sup> T cell expansion and interferon-y production that occur in SAP-deficient mice after lymphocytic Choriomeningitis virus infection [16]. Collectively, these data highlight DGK $\alpha$  as a viable therapeutic target to reverse the life-threatening EBV-associated immunopathology that occurs in XLP-1 patients.

To date, a handful of DGKs inhibitors have been discovered, but some drawbacks are associated with their use as drugs such as low potency, lack of selectivity, off-target effects and poor pharmacokinetic properties. In Fig. 2 are reported the best documented DGK inhibitors, which can be gathered in three main groups: i) allosteric inhibitors; ii) DAG analogues and iii) competitive inhibitors at the ATP binding site.

In the class of allosteric inhibitors, the two compounds R59949 and R59022 are the most important discovered to date [17,18], show selectivity for DGK $\alpha$  [19] and they have been important pharmacological tools for studying the role of DGKs in cells. The IC50 values of R59022 and R59949 are dependent by assay conditions: using whole cell lysates of DGK $\alpha$  overexpressing Cos-7 cells in a nonradioactive assay method, the IC50 values of R59022 and R59949 were approximately 25  $\mu$ M and 18  $\mu$ M, respectively [19,20]. In the original reports, however, R59022 featured an IC50 of 3.8  $\mu$ M on erythrocyte ghosts while R59949 displayed an IC50 of 0.3  $\mu$ M on platelet membranes [17,18]. It has been postulated that R59949

binds to the catalytic domain of DGKα in synergy with Mg-ATP without affecting DAG binding. Putatively R59949 acts by binding in proximity to the Mg-ATP site or promoting a conformational change of the enzyme that exposes both Mg-ATP and R59949 binding sites [20]. However, their use in vivo is severely limited by poor pharmacokinetic properties, and several off-target activities (https://www.ebi.ac.uk/chembl/). Despite these drawbacks and its short half-life. R59022 has been used in mice with positive results in glioblastoma models and in XLP-1 [16,21]. Based on structural similarity with R59022, ritanserin was recently characterized as an active non-competitive inhibitor of DGKa with an IC50 around 20 μM [22]. Structural studies support a contiguous ligand-binding site composed of C1, DAGKc, and DAGKa domains in the DGKα active site for this class of inhibitors [23,24]. Both ritanserin, and R59022/R59949 are also antagonists of serotonin receptors. For example, ritanserin was indeed originally described as a selective 5-HT2A (Ki = 0.45 nM) and 5-HT2C receptor (Ki = 0.71 nM) antagonist and underwent clinical trials for the treatment of schizophrenia and other mental diseases [25].

The class of DAG analogues has not produced potent and pharmacologically useful DGK inhibitors to date with dioctanoylethylene glycol  $(Ki = 58 \mu M)$ 1-monooleolylglycerol and  $(Ki = 91 \mu M)$  the two most potent inhibitors found [26,27]. To note that phorbol esters, well-known DAG mimetic on PKC, are active only with the  $\beta$  and  $\gamma$  isoform, while they have no activity on the other eight isoforms [28,29]. Finally, it is important to highlight the problem of identifying ATP competitive inhibitors of DGKs. The reason lavs in their different ATP binding pocket compared to serine-tyrosine kinases but also to lipid kinases such as PI3Ks. The different architecture of the ATP binding pocket is exemplified by the total lack of activity of the pan-kinase inhibitor staurosporine on DGKs [28]. Very recently, a compound named CU-3 was identified via high throughput screening. In a non-radioactive assay method, CU-3 was able to inhibit DGK $\alpha$  with an IC<sub>50</sub> value of 0.6  $\mu$ M in DGKα overexpressing Cos-7 cells. Moreover, CU-3 acts as a competitive inhibitor of DGK $\alpha$  that reduces affinity of DGK $\alpha$  for ATP, but not diacylglycerol or phosphatidylserine [30].

The goal of this work was therefore to identify novel druggable inhibitors of DGKα, for the treatment of XLP-1 using a virtual screening approach. As there is no detailed structural information on DGKα catalytic domain, we searched for new inhibitors using R59022 and R59949 as templates to explore the chemical space via 2D/3D virtual screening approaches. Herein, we report that out of 127 compounds tested, ritanserin and AMB639752 were more active than the lead inhibitors R59022 and R59949. We also show that both compounds can restore TCR pro-apoptotic signaling and RICD in SAP-deficient T cells, suggesting their use for treatment of EBV-induced FIM in XLP-1 patients. Moreover, AMB639752 does not perturb serotonin signaling indicating an increased selectivity.

#### 2. Result and discussion

# 2.1. In silico drug discovery

Starting from the structure of R59949 and R59022, we commenced with a fast in-silico molecular comparison method to calculate molecular similarity of target substances to both reference compounds (2D-similarity-based virtual screening). In brief, the 2D fingerprint-based methods encode the structural features of molecules as bit strings, whereby each bit indicates the presence or absence of pre-defined structural patterns such as atom sequences, electronic configurations, atom pairs, and ring systems. We used the PubChem library and selected molecules with a Tanimoto coefficient  $\geq$ 0.75 in comparison with R59949 (reference compound). Five compounds emerged from this screening (R59022, altanserin,

# Allosteric inhibitors R59949 (IC<sub>50</sub>= 18 µM) R59022 (IC<sub>50</sub>= 25 µM) ritanserin (IC<sub>50</sub>= 20 µM) OH OAc calphostin C (IC<sub>50</sub>= 40 µM) cochlioquinone A (Ki= 2.26 µM) stemphone (Ki= 3.32 µM) **DAG** analogues OH dioctanoylethylene glycol (Ki= 58 µM) 1-monooleolylglycerol (Ki= 91 µM) ATP competitive inhibitors CU-3 ( $IC_{50}$ = 0.6 $\mu$ M)

Fig. 2. DGKs inhibitors: state of the art.

ritanserin, ketanserin, pimozide) (Fig. 3A). As all the identified compounds were already evaluated as serotonin receptor antagonists, we decided to select other 24 serotonin receptor antagonists as negative control for DGK $\alpha$  inhibition (Supplementary Table S1). Of those, only 21 compounds resulted commercially available (R59022, ritanserin, ketanserin, pimozide and 17 negative controls) and were included in the subsequent screen. Subsequently, we used an in-silico 3D approach using R59022 and R59949 as query ligands to identify in the ZINC database<sup>24</sup> compounds with similar three-dimensional shape and electrostatic properties compared with the two standards.

Of note, our query compounds contain several rotatable single bonds and their bioactive conformations are unknown. Therefore, in our shape-based virtual screening strategy we explored a range of possible conformations of templates by screening against the most stable conformations of both R59022 (9 conformers) and R59949 (6 conformers) generated with Omega [31]. After generation of conformers for each ZINC compound in the "drug-now" subset, the 3D shape comparison between R59s and the molecules in the ZINC library was performed by ROCS [32], in which shape is approximated by atom-centered overlapping Gaussians and calculates the maximal intersection of the volume of two molecules.

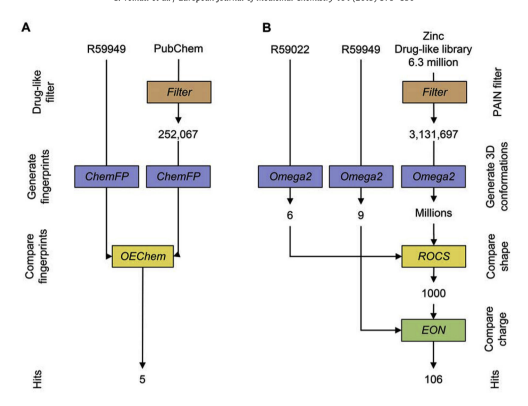


Fig. 3. Virtual screening protocols details. A) Flow chart outlining the protocol used in the 2D-Similarity-based virtual screening. B) Flow chart outlining the protocol used in the 3D-Similarity-based virtual screening.

The top 1000 'hits' were re-ranked for similarity to R59s based on electrostatics with the program EON [33]. Electrostatic rank was based on an electrostatic Tanimoto score, ranging from 0 to 2, where 2 represents an exact match of both shape and functional groups between two molecules. We selected the top 150 compounds and purchased the 106 compounds that were commercially available (Fig. 3B). Full details for these compounds including their structures, Tanimoto ranks and chemical identification numbers are provided in Supplementary Table S2.

#### 2.2. Biological results

To identify the active compounds between those emerged from the two in-silico screenings, all the candidates were dissolved in DMSO and tested at a concentration of 100 µM for the ability to inhibit DGKα using equal amounts of DMSO as control. Assay uses OST-DGKα overexpressing cell lysates in presence of saturating exogenous DAG and ATP [34]. Due to the poor solubility, risperidone and AMB9107756 were not included in this assay, resulting in 127 compounds tested. Each inhibitor was tested in duplicate, and DGKα activity was expressed as percentage of residual DGKα activity compared to DMSO control in the same assay. Out of the 127 compounds, we identified 17 compounds capable of reducing OST-DGK $\alpha$  activity at least by 25% (Table 1). As expected, our reference inhibitors R59022 and R59949 featured 73% and 72% inhibition respectively, confirming the quality of data obtained. Interestingly, ritanserin and above all the new entry AMB639752 inhibited OST-DGKα better than R59022 and R59949, thus representing novel DGK $\alpha$  inhibitors. In order to confirm the structure and purity of AMB639752, NMR structural characterization, mass analysis and elemental analysis were carried out, verifying the correctness of the structure (see "Experimental protocols" section).

Surprisingly, in this assay, more than 60% of the compounds

stimulated DGK $\alpha$  activity, with some compounds enhancing the enzyme activity by several folds. We can confirm that the most active one, fluoxetine, significantly enhances DGK $\alpha$  activity *in vitro* (Fig. S1). As reported for the parental compounds, these molecules may enhance affinity for Mg-ATP or promote an active enzyme conformation by inserting in an allosteric pocket [23,24]. However, for the moment, we did not investigate these compounds further.

# 2.3. Potency and isoform specificity of ritanserin and AMB639752

To measure the inhibitor potency, we determined the IC $_{50}$  values for active compounds by measuring the residual OST-DGK $\alpha$  activity over a dose range of inhibitor concentrations (0.1, 0.3, 1.0, 3.0, 10.0, 30.0 and 100.0  $\mu$ M). For R59022 and R59949 we measured IC $_{50}$  values of 19.7  $\pm$  3.0  $\mu$ M and 10.6  $\pm$  3.2  $\mu$ M respectively, comparable to previous reports using similar assay conditions [19,20]. Conversely, we found IC $_{50}$  values of 16.7  $\pm$  4.6  $\mu$ M for ritanserin and 4.3  $\pm$  0.6  $\mu$ M for AMB639752 (Fig. 3). Those results indicate a superior potency of AMB639752 with respect to both ritanserin and the template compounds.

127 molecules were dissolved in DMSO and tested at 100  $\mu$ M for the capacity to inhibit DGK $\alpha$  in homogenates of OST-DGK $\alpha$  overexpressing MDCK cells. Dose response of the most active compounds along with their IC50 values is shown. Data from at least three independent experiments performed in triplicate.

Concerning similarity to R59s, AMB639752 is different with respect to their chemical scaffold, but similar in three-dimensional shape, as demonstrated by their shape Tanimoto's ranks (0.623, Supplementary Information Table S2). AMB639752 is also electrostatically similar to R59949, with an electrostatic Tanimoto of 0.745 (Fig. 4).

To check the isoform specificity of AMB639752, we tested it along with R59s, for the ability to inhibit DGK $\alpha$ , DGK $\zeta$  (the other

**Table 1** Compounds that inhibit DGKα at least by 25%.

Sl.No	Code	Name	Residual DGK $\alpha$ activity at 100 $\mu$ M (% of control)
1.	Amb639752		4
2.	R103	ritanserin	17
3.	R59022	diacylglycerol kinase inhibitor I	27
4.	R59949	diacylglycerol kinase inhibitor II	28
5.	Amb3494010		39
6.	Amb19541048		49
7.	Amb8223059		50
8.	Amb11070312		57
9.	H0972	haloperidon	62
10.	Amb242047		64
11.	Amb20056316		67
12.	Amb8244320		68
13.	Amb110991		68
14.	E0925	ebastine	69
15.	Amb252835		72
16.	U0085	urapidil	75
17.	Amb19382197		75

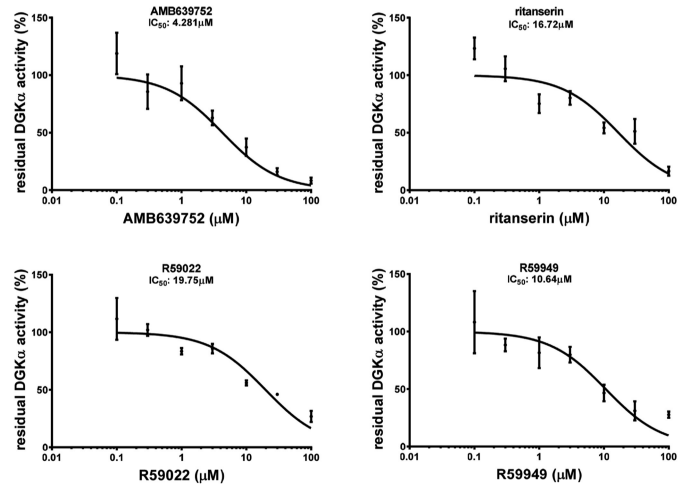


Fig. 4. Biochemical screen for novel DGKα inhibitors.

major DGK isoform expressed in lymphocytes), and the more distantly related DGK $\theta$ . Also at the highest concentration of 100  $\mu$ M, R59022, R59949 and AMB639752 are highly specific against DGK $\alpha$  as they completely inhibit DGK $\alpha$  (68%, 81%, 93% respectively, Fig. 5A) while they do not have significant effects on DGK $\zeta$  (Fig. 5B) and DGK $\theta$  (Fig. 5C). Similar results were reported for ritanserin [22] indicating that this class of inhibitors is highly specific for DGK $\alpha$ .

# 2.4. Activity of compounds on serotonin receptors

As R59022, R59949 and ritanserin feature a dual activity as DGK $\alpha$  inhibitors and serotonin receptor antagonists [22], we investigated whether also AMB639752 affects serotonin signaling.

To this purpose, we measured the effect of serotonin on PMAinduced oxidative burst in human monocytes. As previously

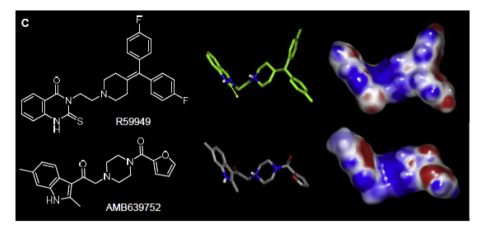


Fig. 5. 2D, 3D and shape of R59949 and Amb639752 chemical structures. Electrostatic surfaces are color coded (red for negative and blue for positive).

shown,  $1 \,\mu\text{M}$  serotonin reverses the oxidative burst to control values [35]. When tested at  $10 \,\mu\text{M}$ , the known serotonin receptor antagonists ritanserin and ketanserin impaired serotonin action, while pure DGK $\alpha$  inhibitors such as CU-3 had no effect (Fig. 7).

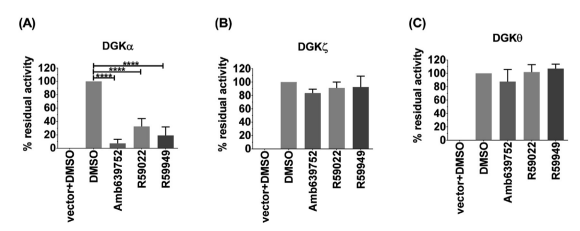
These data indicate that this assay is sensitive to perturbations in serotonin signaling independently of activity against DGK $\alpha$ . AMB639752, despite sharing surface similarity with R59949 and ritanserin, did not affect serotonin action demonstrating that it is not a serotonin receptor antagonist. This interesting observation suggests a better pharmacological profile for AMB639752, which inhibits DGK $\alpha$  without off-target effects on serotonin receptors.

#### 2.5. DGK $\alpha$ specific inhibitors restores RICD in SAP deficient T cells

The defective RICD observed in T cells from XLP-1 patients was rescued by silencing DGK $\alpha$  expression or by pre-treatment with DGK $\alpha$  inhibitors R59949 or R59022 at 5–10  $\mu$ M [16]. Interestingly, R59022 also showed beneficial effects in an *in vivo* model of XLP-1, but its poor pharmacological proprieties make its use in human patients unlikely. We therefore tested the effect of ritanserin and AMB639752 on RICD sensitivity of SAP-deficient T cells. As additional controls, we also included ketanserin to evaluate the contribution of serotonin antagonism to the effects observed and CU-3 as representative of a different class of molecules inhibiting DGK $\alpha$  by competing with ATP. To model XLP-1 T cells *in vitro*, we previously silenced SAP in Jurkat cells [15], a widely used cellular

model for RICD mediated by FAS induction [36]. RICD was induced by treatment with immobilized anti-CD3 antibody (OKT3, 50 μg/ ml) or PHA (50 μg/ml) [37]. As expected, SAP-silenced Jurkat cells showed reduced RICD compared to cells transfected with a control siRNA, making this an easy model for testing our inhibitors (Fig. 8B and C). The inhibitors were not overly toxic to Jurkat cells as 24-h treatment at 1 µM doses did not decrease the viability of Jurkat cells, apart for a modest toxicity of CU-3 (Fig. 8A). At this concentration, ritanserin, AMB639752 and CU-3 significantly promoted RICD in SAP-silenced Jurkat cells, restoring death to levels noted in Jurkat control cells after either anti-CD3 (Fig. 8B) or PHA (Fig. 8C) treatment. Conversely, ketanserin only had a modest effect (Fig. 8B and C). These data indicate that the new DGKα inhibitors identified by our screen can restore RICD in this cell model of XLP-1. Furthermore, our results suggest these effects result from inhibiting DGKa independently of eventual activity on serotonin receptors.

To assess inhibitor efficacy in a more physiological context, we also modelled XLP-1 by silencing SAP in primary peripheral blood T lymphocytes (PBL) and re-stimulating them with increasing doses of anti-CD3 antibody (OKT3 1–100 ng/ml, 24 h). In these assay conditions, 30-min pre-treatment with R59949 or R59022 restored RICD sensitivity in SAP silenced cells at concentrations of 5–10  $\mu$ M [16]. Also in this setting, ritanserin does not influence RICD induced by CD3 triggering in control siRNA-transfected cells, indicating that it is neither affecting TCR signaling nor is cytotoxic to primary cells



**Fig. 6.** AMB639752 is DGK $\alpha$  specific. A, B, C) 293T cells were transfected with different DGK isoforms (A- DGK $\alpha$ , B- DGK $\alpha$ , C- DGK $\theta$  respectively) or empty vectors and homogenized. AMB639752, R59022 and R59949 were tested at 100 μM for their capacity to inhibit the DGK activity of the different homogenates. Data are means  $\pm$  S.E.M. of at least three independent experiments performed in triplicate.

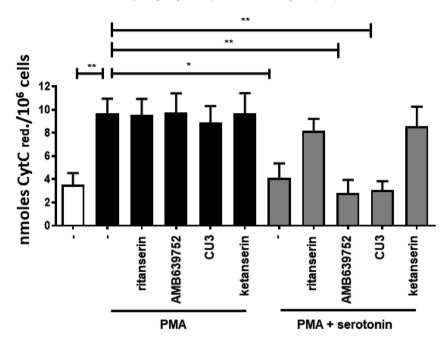


Fig. 7. AMB639752 does not affect serotonin signaling. Human monocytes were pre-incubated for 1 h with the indicated drugs ( $10 \,\mu\text{M}$ ) in absence or presence of serotonin ( $1 \,\mu\text{M}$ ) and then stimulated with PMA 1  $\mu$ M for 30min ( $\square$  control unstimulated cells,  $\blacksquare$  PMA treated cells,  $\blacksquare$  PMA and serotonin stimulated cells). Results are expressed as nmoles of reduced cytochrome C/ $10^6$  cells. Data are means  $\pm$  S.E.M. of 6 independent experiments performed in triplicate.

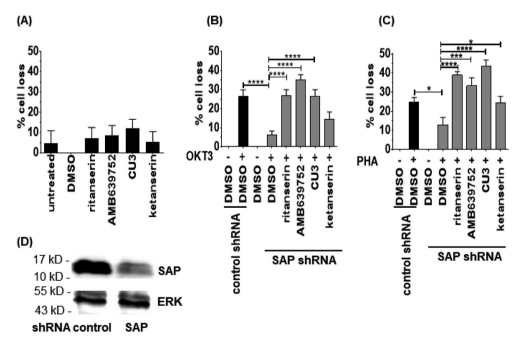


Fig. 8. DGKα inhibitors rescues RICD in SAP deficient Jurkat cells. Jurkat T cells (control shRNA  $\blacksquare$  and SAP shRNA  $\blacksquare$ ) were treated with ritanserin, AMB639752, CU-3 or ketanserin (1  $\mu$ M) and plated on a 96 well flat bottom plate previously coated with nothing (**A**), anti CD3 (OKT3 50  $\mu$ g/ml, **B**) and PHA (50  $\mu$ g/ml, **C**). After 24 h of incubation, live cells were counted using the trypan blue exclusion assay and expressed as % of cell loss respect to untreated controls. The data are the means of  $\pm$ SEM of at least five experiments performed in triplicate. **D**) Western blot confirming the silencing of SAP in SAP shRNA Jurkat cells compared with control shRNA Jurkat cells.

at concentrations between 1 and  $10\,\mu\text{M}$ . Conversely, ritanserin completely rescued the apoptotic defect in SAP-deficient T cells at concentrations as low as  $1\,\mu\text{M}$  (Fig. 9A). We used ritanserin  $10\,\mu\text{M}$  as reference for further sets of experiments (we always compare controls and SAP silenced lymphocytes from the same donors as there is a large individual variability in RICD sensitivity). AMB639752 demonstrated similar activity, not affecting RICD sensitivity in control lymphocytes, but fully rescuing the RICD

defect of SAP-deficient cells also at 1  $\mu$ M concentration and with a potency equal to 10  $\mu$ M ritanserin (Fig. 9B). When we tested the ATP competitive DGK $\alpha$  inhibitor CU-3, we observed a modest induction of RICD in control-silenced lymphocytes that is absent in the other molecules tested. In SAP-silenced cells, CU-3 improved RICD sensitivity approaching ritanserin levels when used at concentrations  $\geq$ 5  $\mu$ M, (Fig. 9C). These observations suggest that CU-3 has a narrower therapeutic window with some likely cytotoxicity at

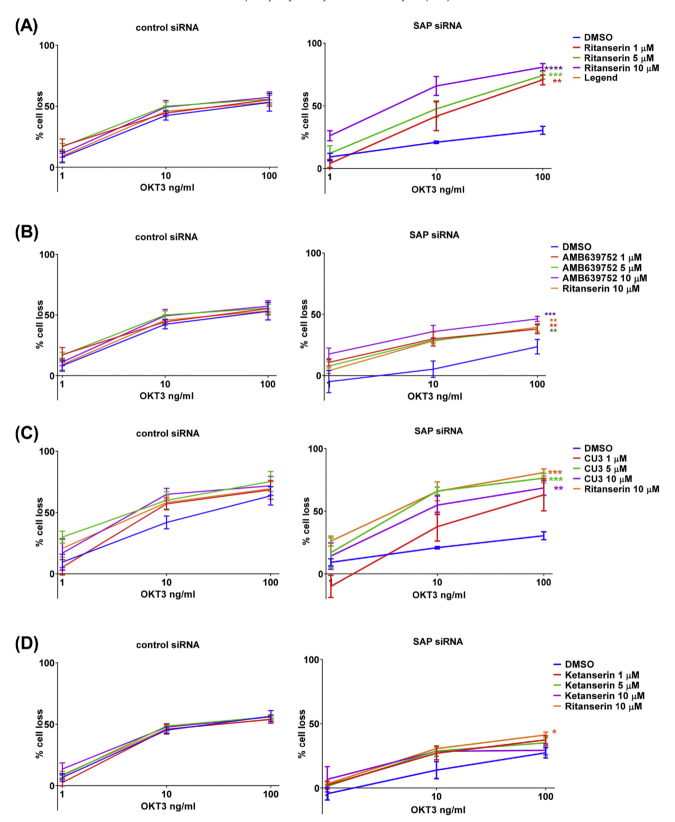


Fig. 9. DGK $\alpha$  inhibitors rescues RICD in SAP deficient T cells. Lymphocytes from normal subjects were transfected with control or SAP specific siRNA. A, B, C, D) After 4 days, the cells were restimulated with increasing doses of CD3 agonist OKT3 (1, 10, 100 ng/ml) in presence or absence of (A) ritanserin, (B) AMB639752, (C) CU-3, (D) ketanserin (1, 5, 10 μM). 24 h later, the % of cell loss was evaluated by PI staining. Data are the mean  $\pm$  SEM of at least three independent experiments performed in triplicate.

higher doses. Finally, we tested ketanserin in our RICD assays to verify if serotonin receptor antagonism could contribute to the observed effects. Interestingly, ketanserin had no effect on RICD in control cells, but partially increased RICD in SAP-silenced T cells, although this effect was not statistically significant (Fig. 9D). These findings are in line with our Jurkat cell experiments, suggesting that inhibition of DGK $\alpha$  is an efficient strategy to rescue of RICD in XLP-1 models, while serotonin antagonism plays a minor role.

In summary, these data confirm that ritanserin and AMB639752 rescue RICD susceptibility in T cell models of XLP-1. Our observations are consistent with those on DGK $\alpha$ -silenced cells [16], indicating that DGK $\alpha$  is a relevant pharmacologic target for restoring RICD and immune homeostasis in XLP-1 patients.

2.6.  $DGK\alpha$  specific inhibitors potentiates TCR signaling and induces expression of the pro-apoptotic nuclear receptors Nur77 and Nor1 in SAP deficient T cells

Silencing DGK $\alpha$  expression or pre-treatment with DGK $\alpha$  inhibitors R59949 or R59022 boosts RICD in SAP-deficient T cells by selectively restoring TCR-induced upregulation of the pro-apoptotic orphan nuclear receptors NUR77 (NR4A3) and NOR1 (NR4A1) [16]. To verify if ritanserin and AMB639752 acts through the same mechanism, we measured TCR-dependent induction of NR4A1 and NR4A3 in SAP-silenced activated T cells. Indeed, ritanserin potentiated TCR-induced NR4A1 expression in control cells and to a lesser extent, in SAP silenced cells. In contrast, AMB639752 was more active in rescuing NR4A1 and NR4A3 in SAP-silenced activated T cells (Fig. 9). These observations suggest that both ritanserin and especially AMB639752 restore RICD by promoting the expression and apoptotic activity of NR4A1 and NR4A3 (see Fig. 10).

### 3. Conclusion

Summarizing, the purpose of this project was to discover novel inhibitors against  $DGK\alpha$  suitable for future clinical trials in XLP-1 patients. To find potentially drug candidates, we explored insilico the chemical space of ZINC by using the two commercially available inhibitors (R59022 and R59949) as templates. We assembled a library of 127 compounds using this approach, including both uncharacterized molecules and a series of compounds already in clinical use or development.

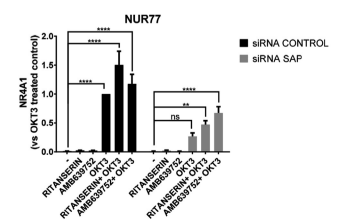
By screening this library, we found 17 active inhibitors, with two more active than the lead compounds: the compound AMB639752 and ritanserin. We defined the IC $_{50}$  values of ritanserin (16.7  $\pm$  4.5  $\mu$ M) and AMB639752 (4.3  $\pm$  0.6  $\mu$ M) for inhibiting DGK $\alpha$ . Thus, IC $_{50}$  value of AMB639752 compares favourably with ritanserin, R59022 (19.7  $\pm$  3.0  $\mu$ M) and R59949 (10.6  $\pm$  3.2  $\mu$ M) in the same assay. Interestingly 60% of the compounds in the library acted as activators and not inhibitors of DGK $\alpha$  (see supporting information). These compounds might bind to the same pocket as R59949 and increase affinity for Mg-ATP or stabilize an active conformation. Further studies will explore whether these DGK $\alpha$  activators may offer interesting opportunities for the control of DAG metabolism in pathological settings such as inflammation.

AMB639752 maybe a promising molecule for a drug optimization program, as it potently inhibits DGKα without influencing serotonin signaling on the contrary of ritanserin. AMB639752 was highly active in cell-based assays, with negligible cytotoxicity at concentrations used for the two commercially available DGK inhibitors. Moreover, it did not alter RICD sensitivity of control (SAPsufficient) lymphocytes. Both ritanserin and AMB639752 restore RICD in SAP-deficient cells at 1, 5 and 10 µM concentration, consistent with DGKa inhibition. At these doses, neither compound displayed toxicity nor impaired TCR signaling. Both drugs restored the induction of key pro-apoptotic molecules NUR77 and NOR1 in SAP-deficient cells, but AMB639752 was far more active. This indicates that similarly to R59s and DGKa silencing, ritanserin and AMB639752 are not simple TCR signal enhancers, but selectively influence SAP-dependent downstream signaling outcomes that promote RICD. Altogether, these observations support our initial hypothesis that excessive DGKα activity contributes to XLP-1 pathogenesis and suggest that we are moving in the right direction to find new compounds for countering life threatening disease complications in this disease (e.g. EBV-induced FIM).

# 4. Experimental protocols

#### 4.1. Structural confirmation for AMB639752

AMB639752 or 1-(2,5-dimetil-1*H*-indol-3-yl)-2-(4-(furan-2-carbonil)piperazin-1-yl)etan-1-one: yellowish solid;  $^1H$  NMR (300 MHz, CDCl<sub>3</sub>):  $\delta=8.70$  (br s, 1H), 7.74 (d, J=8.2 Hz, 1H), 7.47 (s, 1H), 7.26 (s, 1H), 7.13 (s, 1H), 7.07 (d, J=8.2 Hz), 6.98 (br d, 1H), 6.47 (br d, 1H), 3.92 (br s, 4H), 3.85 (s, 2H), 2,79 (br s, 4H), 2,74 (s, 3H), 2.44 (s, 3H); MS (ESI):  $m/z=366\,[{\rm M}+{\rm H}]^+;$  elemental analysis calcd (%) for C<sub>21</sub>H<sub>23</sub>N<sub>3</sub>O<sub>2</sub>: C 69.02, H 6.34, N 11.50: found: C 69.23, H 6.58, N 11,33.



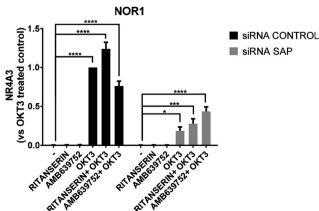


Fig. 10. DGKα inhibitors rescues Nur77 and Nor1 expression in SAP deficient T cells. Lymphocytes from normal subjects were transfected with control or SAP specific siRNA. After 4 days, the cells were restimulated with the CD3 agonist OKT3 ( $10 \mu g/ml$ ) in presence or absence of ritanserin or AMB639752 at  $10 \mu M$  concentration. 4 h later, the expression of NR4A1 and NR4A3 evaluated by quantitative PCR using GUSB as a reference. Data are the mean  $\pm$  SEM of at least four independent experiments.

#### 4.2. Cell lines

Madin-Darby canine kidney cells (MDCK) stably expressing One Strep Tag DGK $\alpha$  (OST-DGK $\alpha$ ) were prepared by infecting MDCK cells with a vector expressing an inducible OST tagged DGK $\alpha$  constructs [38]. MDCK cells infected with empty vector were used as controls. MDCK cells were cultured in MEM (Minimal Essential Medium) with 5% FBS (Fetal Bovine Serum) and 1% antibiotic-antimycotic solution. Routinely, cells were split every 3–4 days with Trypsin-EDTA 0.25% in standard 100 mm dishes.

Human embryonic kidney 293T cells (10 cm<sup>2</sup> plates) were cultured in RPMI with 10% FBS and 1% penicillin/streptomycin.

Generation of Jurkat shRNA SAP and the Jurkat shRNA control was described in Ref. [15]. Jurkat cells were cultured in RPMI-GlutaMAX (Life Technologies) with 10% FBS (Lonza) and 1% penicillin/streptomycin (Life Technologies).

# 4.3. Primary cells

Peripheral blood mononuclear cells (PBMCs) were isolated from healthy anonymous human buffy coats (provided by the Transfusion Service of Ospedale Maggiore della Carità, Novara, Italy) by Ficoll-Paque PLUS (GE Healthcare) density gradient centrifugation, washed, and resuspended at  $2\times10^6$  cell/ml in RPMI-GlutaMAX containing 10% heat inactivated FCS, 2 mM glutamine, and 100 U/ml of penicillin and streptomycin. T cells were activated with 1 µg/ml anti-CD3 (clone OKT3) and anti-CD28 (clone CD28.2) antibodies. After 3 days, activated T cells were washed and cultured in medium supplemented with 100 IU/ml rhIL-2 (Peprotech) at 1.2  $\times$  106 cells/mL for >7 days by changing media for every 2–3 days.

Human monocytes were isolated from healthy anonymous human buffy coats (provided by the Transfusion Service of Ospedale Maggiore della Carità, Novara, Italy) by the standard technique of dextran sedimentation and Histopaque (density =  $1.077 \, \mathrm{g \, cm^3}$ , Sigma-Aldrich, Milano, Italy) gradient centrifugation ( $400 \times g$ , 30min, room temperature) and recovered by fine suction at the interface, as described previously [35]. Purified monocytes populations were obtained by adhesion (90 min,  $37\,^{\circ}$ C,  $5\%\,^{\circ}$ CO2) in serum-free RPMI 1640 medium (Sigma-Aldrich, Milano, Italy) supplemented with 2 mM glutamine and antibiotics. Cell viability (trypan blue dye exclusion) was usually >98%.

#### 4.4. Molecular modelling

All molecular modelling studies were performed on a Tesla workstation equipped with two Intel Xeon X5650 2.67 GHz processors and Ubuntu 11.10 (www.ubuntu.com). The protein structures and 3D chemical structures were generated in PyMOL [The PyMOL Molecular Graphics System, version 1.3; Schrödinger LLC: 2010].

2D-Similarity-based virtual screening. The collection of compounds was retrieved from PubChem (February 2nd, 2013), this searchable database includes chemical structures available from published literature [39]. The structures were downloaded and filtered using software FILTER [FILTER, version 2.0.2; OpenEye Scientific Software: Santa Fe, NM. http://www.eyesopen.com]: molecular weight <500 Da and >150, logP  $\leq$ 5, number of rotatable bonds  $\leq$ 7, polar surface area <150, number of hydrogen bond donor  $\leq$ 5 and number of hydrogen bond acceptor  $\leq$ 10. Using these criteria 252,067 structures were retrieved. First, the ChemFP Substructure fingerprints were generated for the compounds using chemfp v1.1 [chemfp, version 1.1; Dalke Scientific: Sweden, 2013. http://chemfp.com], with the OEChem [OEChem TK, Python version 1.7.7; OpenEye Scientific Software: Santa Fe, NM. http://www.eyesopen.com], was used to select compounds that are similar to

R59949 by at least a Tanimoto cutoff of 0.75; 5 compounds were selected (Table 1). In order to qualitatively validate the approach 24 other compounds with similar biological activity, not selected by our method, were used as negative control (Supplementary Table S1).

3D-Similarity-based virtual screening. The collection of compounds was retrieved from ZINC (February 19th, 2014), this searchable database includes chemical structures available from vendors [40,41]. The "drug-now" subset, that contains molecules with drug-like properties (molecular weight <500 Da and >150, logP <5, number of rotatable bonds <7, polar surface area <150, number of hydrogen bond donor ≤5 and number of hydrogen bond acceptor <10), was downloaded. Using these criteria, 6,366,032 structures were retrieved. An in-house Perl script was used in order to standardize charges, enumerate ionization states, generate tautomers at physiological pH range and remove PAINS substructures 62 using the FILTER and QUACPAC software from OpenEye (FILTER, version 2.5.1.4; OpenEye Scientific Software: Santa Fe, NM, http:// www.eyesopen.com; QUACPAC, version 1.6.3.1; OpenEye Scientific Software: Santa Fe, NM, http://www.eyesopen.com). 3,131,697 structures were retained. The latter operation was followed by a 3D structure optimization and conformers generation using OMEGA2 software (OMEGA, version 2.4.6; OpenEye Scientific Software: Santa Fe, NM. http://www.eyesopen.com) [31,42]. A three dimensional structures of R59022 and R59949 were built: we used OMEGA2 software to generate 9 conformers of R59022 and 6 conformers of R59949. A RMSD cutoff of 0.75 Å was used in this step. Successful candidates were filtered using shape matching. ROCS software was used for 3D shape comparisons (ROCS, version 2.4.1; OpenEye Scientific Software: Santa Fe, NM. http://www. eyesopen.com) [32]. The top 1000 structure were considered for electrostatic comparisons, EON was used (EON, version 2.0.1; OpenEye Scientific Software: Santa Fe, NM. http://www.eyesopen. com) [33]. The best compounds were selected based on TanimotoCombo values (ShapeTanimoto + ElectrostaticTanimoto). At the end the best 150 compounds were considered, 106 compounds were commercially available and were retrieved from Ambinter vendor (Supplementary Table S2).

All these compounds, including the two known inhibitors were dissolved in DMSO at a concentration of 10 mM AMB9107756 remains undissolved and due to solubility problems ebastine (E0925) was dissolved at 5 mM, risperidone (R3030) at 1 mM concentrations respectively.

ATP $\gamma$ 32P is from PerkinElmer, cell culture reagents are from Thermo Fisher, all other chemicals were from Sigma Aldrich. CU-3 is a king gift of F. Sakane (Chiba University, Japan).

### 4.5. Preparation of DGK $\alpha$ enriched homogenates

Large cultures for enzyme preparation were done by plating  $5\times 10^6$  cells in 245 mm² dishes. Once they reached nearly 70% confluence, cells were treated with doxycycline (1 µg/ml, 2 days). After 2 days of treatment, each plate was washed in cold PBS and cells homogenized in 5 ml of homogenate buffer (25 mM Hepes (pH 8), 20% glycerol, 135 mM NaCl, 5 mM ethylenediaminetetraaceticacid (EDTA), 1 mM ethylene glycol-bis(beta-aminoethyl ether)-N,N,N',N'-tetraacetic acid (EGTA), 1 mM sodium orthovanadate and protease inhibitor cocktail) for each dish. Cells were collected with a rubber scraper, homogenized by passing them through a 29G-needle syringe 20 times and stored in aliquots at  $-80\,^{\circ}$ C. Presence of OST-DGK $\alpha$  was confirmed by western blotting and enzyme assay, transduced DGK $\alpha$  has an activity >100 folds the endogenous DGK.

#### 4.6. Preparation of DGK $\zeta$ and DGK $\theta$ enriched homogenates

293T cells were transiently transfected with indicated DGK isoform plasmid DNA using Lipofectamine 3000, Invitrogen (Carlsbad, CA). 48 h after transfection, cells were harvested and homogenized with a 29G-needle using 500  $\mu l/plate$  of homogenate buffer for each dish and immediately stored in aliquots at  $-80\,^{\circ}\text{C}.$  Cells transfected with empty vector were used as controls, overexpressed DGK has an activity >50 folds the endogenous.

#### 4.7. DGK assay

DGK activity was assayed by measuring initial velocities (5 min at 27 °C) in presence of saturating substrate concentration. Reaction mixture comprises enzyme (24.5  $\mu$ l of homogenate), inhibitor or DMSO (0.5  $\mu$ l), ATP solution (10  $\mu$ l) and DAG solution (15  $\mu$ l).

ATP solution contains and 0.25  $\mu$ Ci/ $\mu$ l (9.25 KBq/ $\mu$ l) of  $\gamma$ 32P labelled radioactive ATP along with 25  $\mu$ M normal unlabeled ATP, 50  $\mu$ m MgCl2, 5  $\mu$ m sodium orthovanadate and water to make up the final volume.

DAG solution contains 1,2-dioleoyl-sn-glycerol (3  $\mu g/\mu l$ ) and 1 mM EGTA in 25 mM Hepes pH 8. DAG is stored in CH3OH:CHCl3 1:1 solution at 10  $\mu g/m l$ , the required amount was dried using  $N_2$  or vacuum centrifugation and sonicated extensively in ice could buffer

The reaction was stopped after 5 min by adding 200  $\mu$ l of freshly prepared 1M HCL and lipid extracted by adding 200  $\mu$ l of CH<sub>3</sub>OH:CHCl<sub>3</sub> 1:1 solution and vortexing for 1 min. The two phases were separated by centrifugation (12000 RCF for 2 min). 25  $\mu$ l of the lower organic phase was spotted in small drops on silica TLC plates (previously coated with (1.3% potassium oxalate and 5 mM EDTA in methanol and dried). Mobile phase was CHCl<sub>3</sub>:CH<sub>3</sub>OH:H<sub>2</sub>O:25% NH<sub>4</sub>OH (60:47:11:4).

TLC was run 10 cm and dried before radioactive signals were detected by GS-250 molecular imager and was quantified by quantity one (Bio-Rad) software assuring the absence of saturated spots.

Percentage residual activity was calculated as follows:

(OST-DGK $\alpha$  homogenate with inhibitor — vector homogenate)/ (OST-DGK $\alpha$  homogenate with DMSO — vector homogenate) \*100.

# 4.8. Superoxide anion $(O^{2-})$ production

All the experiments were performed in triplicate using cells isolated from each single donor.

Monocytes (250.000 cells/well) were treated for 1 h with the indicated drugs (10  $\mu$ M) with or without serotonin (1  $\mu$ M). Then, cells were stimulated with phorbol 12-myristate 13-acetate (PMA; Sigma-Aldrich) 1  $\mu$ M for 30 min. PMA is a well-known stimulus that induces a strong and significant respiratory burst via PKC activation [35]. Superoxide anion production was then evaluated by the superoxide dismutase (SOD)-sensitive cytochrome C (CytC) reduction assay and expressed as nmoles cytochrome C reduced/10^6 cells/30 min, using an extinction coefficient of 21.1 mM. To avoid interference with spectrophotometric recordings, cells were incubated with RPMI 1640 without phenol red, antibiotics and FBS.

### 4.9. RICD assay in Jurkat cells

Treatments with inhibitors were done in 96-well flat-bottom plate, pre-coated with anti CD3 (clone OKT3,  $50 \mu g/ml$ ) or

phytohemagglutinin-P (PHA-P Sigma,  $50\,\mu g/ml$ ) and incubated for 24 h at 4 °C. To test restimulation induced cell death, the plate is washed with medium and then cells were plated at concentration of  $0.5\times10^5$  cells/well in triplicate in RPMI-GlutaMAX medium in presence or absence of  $1\,\mu M$  inhibitor or equivalent volume of DMSO as a negative control. After 24 h incubation, live cells were counted using trypan blue exclusion method and cell death is expressed as % cell loss and calculated as:

% Cell Loss = (100-(number of viable cells in sample/ number of viable cells in control) x 100).

Results were expressed as mean  $\pm$  standard error mean (SEM) of three or more replicate experiments.

#### 4.10. RICD assay for SAP silenced T cells

Activated human PBLs were transfected with 200 pmol of siRNA oligonucleotides specific for the target protein (Stealth Select siRNA; Life Technologies) or a non-specific control oligo (Life Technologies). Transient transfections were performed using Amaxa nucleofector kits for human T cells (Lonza) and the Amaxa Nucleofector II or 4D systems (programs T-20 or EI-115). Cells were cultured in IL-2 (100 IU/ml) for 4 days to allow target gene knockdown. Knockdown efficiency was periodically evaluated by RT-PCR and Western blotting.

Non-specific Stealth RNAi Negative Control Duplexes (12935—300, Life Technologies) were used as a negative control. Sense strand UGUACUGCCUAUGUGUGUGUGUGUAUCA Antisense strand UGAUACAGCAGACAUAGGCAGUACA.

To test restimulation induced cell death, activated T cells ( $10^5$  cells/well) were plated in triplicate in 96-well round-bottom plate and treated with anti-CD3 (clone OKT3) (1, 5, 10, 50, 100, 500 ng/ml) in RPMI-GlutaMAX supplemented with 100 IU/ml rhIL-2 for 24 h. In these assays inhibitors (1, 5, 10  $\mu$ M) were added 30 min before the restimulation with OKT3. 24 h after treatment, cells were stained with 1  $\mu$ g/ml propidium iodide and collected for a constant time of 30 s per sample on a FACScan flow cytometer (FACScaliber or Accuri C6, BD) or Attune Nxt Flow Cytometer (Thermo Fisher Scientific). Cell death is expressed as % cell loss and calculated as:

% cell loss = 100-((number of viable cells in sample / number of viable cells in control) x 100)

Results were expressed as mean  $\pm$  standard error mean (SEM).

### 4.11. Quantitative real-time PCR

Activated human PBLs were prepared and transfected as described above. The RNA concentration and purity were estimated by a spectrophotometric method using NanoDrop 2000c (Thermo Scientific). The RNA was retrotranscribed with High-Capacity cDNA Reverse Transcription Kits (Applied Biosystems) and cDNA quantified by real time PCR (ABI7900 Applied Biosystems) using GUSB as normalizer.

TaqMan gene expression assays were from Life Technologies: Hs00939627\_m1 (glucuronidasi beta, GUSB), Hs00374226\_m1 (NR4A1), Hs00545007\_m1 (NR4A3).

# 4.12. Western blot analysis

To verify the expression of the DGK $\alpha$  in both MDCK cells transfected with OST-Vector and OST-DGK $\alpha$ , and SAP in Jurkat cells transfected with control-shRNA and SAP-shRNA, equal amounts of protein lysates were prepared using 1% NP40 lysis buffer

comprising of 25 mM Hepes, 150 mM Nacl, 5 mM EDTA, 1 mM EGTA, 50 mM NaF, 1% NP40, 10% glycerol, 1 mM orthovanadate along with protease inhibitors at PH 8.0. Proteins were separated on 12% or 15% SDS-PAGE gels and the resolved proteins were then transferred to a polyvinylidene fluoride membrane. The membranes were blocked by incubating with 3% (w/v) dried BSA for 1 h at room temperature. They were then incubated overnight with respective primary antibodies at 4°C with gentle agitation; for detecting DGKα we used anti-DGKα antibody from Abcam (Cambridge, UK) and anti-SAP (SH2D1A-D10G8) from Cell signaling (USA) diluted (1:500) in Tris-buffered saline with detergent Tween 20 (TBST) containing 3% (w/v) BSA. The membranes were then washed 4 times in TBS tween buffer with 15 min time interval. They were then incubated with alkaline HRP-conjugated rabbit or mouse secondary antibody (1:5000) from PerkinElmer (USA) diluted in TBST, for 1 h at room temperature with gentle agitation. After four 15 min washes with TBST, the membranes were developed using the Western Chemiluminescence Substrate (PerkinElmer, USA).

#### 4.13. Quantification and statistical analysis

Data for the screen on OST-DGK $\alpha$  homogenates are the mean of duplicates. The compounds showing inhibitory activity in this assay were tested >4 times and the mean  $\pm$  SEM is reported.

To calculate IC $_{50}$  values of active inhibitors, the inhibitor activity was measured at least three times at 0.1, 1.0, 10.0 and 100.0  $\mu$ M concentration. Data were analysed using [inhibitor] vs. normalized response parameters with least square [ordinary] curve fitting method in GraphPad PRISM 7.0 software mentioning 95% confidence interval and IC $_{50}$  values always greater than 0.0. Graph shows the mean  $\pm$  SEM of inhibitor activity at the indicated concentration. In all the experiments the data were normalized with the controls.

Evaluation of *in vitro* assays across multiple treatments (RICD) were analysed by using one-way ANOVA (Figs. 6–8) and two-way ANOVA (Fig. 9) with multiple comparisons correction using GraphPad PRISM 7.0 software. Error bars are described in figure legends as  $\pm$  SEM or  $\pm$  SD where appropriate. A single, double and triple asterisk denotes significance of a p-value  $\leq$ 0.05,  $\leq$ 0.01 and  $\leq$  0.001 respectively in all experiments.

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# Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejmech.2018.12.061.

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