Canagliflozin mediated dual inhibition of mitochondrial glutamate dehydrogenase and complex I: an off-target adverse effect

Philipp F. Secker, Sascha Beneke, Nadja Schlichenmaier, Johannes Delp, Simon Gutbier, Marcel Leist and Daniel R. Dietrich

Abstract
Recent FDA Drug Safety Communications report an increased risk for acute kidney injury in patients treated with the gliflozin class of sodium/glucose co-transport inhibitors indicated for treatment of type 2 diabetes mellitus. To identify a potential rationale for the latter, we used an in vitro human renal proximal tubule epithelial cell model system (RPTEC/TERT1), physiologically representing human renal proximal tubule function. A targeted metabolomics approach, contrasting gliflozins to inhibitors of central carbon metabolism and mitochondrial function, revealed a double mode of action for canagliflozin, but not for its analogs dapagliflozin and empagliflozin. Canagliflozin inhibited the glutamate dehydrogenase (GDH) and mitochondrial electron transport chain (ETC) complex I at clinically relevant concentrations. This dual inhibition specifically prevented replenishment of tricarboxylic acid cycle metabolites by glutamine (anaplerosis) and thus altered amino acid pools by increasing compensatory transamination reactions. Consequently, canagliflozin caused a characteristic intracellular accumulation of glutamine, glutamate and alanine in confluent, quiescent RPTEC/TERT1. Canagliflozin, but none of the classical ETC inhibitors, induced cytotoxicity at particularly low concentrations in proliferating RPTEC/TERT1, serving as model for proximal tubule regeneration in situ. This finding is testimony of the strong dependence of proliferating cells on glutamine anaplerosis via GDH. Our discovery of canagliflozin-mediated simultaneous inhibition of GDH and ETC complex I in renal cells at clinically relevant concentrations, and their particular susceptibility to necrotic cell death during proliferation, provides a mechanistic rationale for the adverse effects observed especially in patients with preexisting chronic kidney disease or previous kidney injury characterized by sustained regenerative tubular epithelial cell proliferation.

Introduction
Canagliflozin is a member of the gliflozin group of pharmaceuticals indicated for treatment of type 2 diabetes mellitus (T2DM). Gliflozins are inhibitors of members of the sodium-coupled glucose co-transporters (SGLT; SLC5A gene family) and primarily target SGLT2 expressed in renal proximal tubule epithelial cells (RPTECs) of the kidney. SGLT2 is responsible for the bulk of renal glucose reabsorption, while the SGLT1 isoform, expressed in the pars recta of the renal proximal tubule, is a high-affinity/low-capacity transporter, responsible for the uptake of the remaining glucose and galactose molecules in the primary urine. SGLT1 is also expressed in the brush border membrane of the small intestine. Two inherited human disorders of sodium-coupled glucose transport, i.e., intestinal glucose-galactose malabsorption (GGM), involving SGLT1 gene mutations, and familial renal glucosuria (FRG), involving mutations of the SGLT2 gene, are known to date. Neither GGM nor FRG disorders are accompanied by serious health issues for the affected
individuals, nor have they been specifically associated with intestinal or renal pathology\textsuperscript{2}. Hence, the inhibition of renal SGLT2 was considered useful for treatment of T2DM, which was supported by studies with the natural compound phlorizin, a metabolically unstable and unspecific inhibitor of SGLT2 and SGLT1\textsuperscript{3}. Accordingly, analogs of phlorizin, yet with higher selectivity of SGLT2 over SGLT1\textsuperscript{4} and increased stability and bioavailability, were developed to increase urinary clearance of blood glucose. Three such SGLT2 inhibitors, canagliflozin (Invokana\textsuperscript{4}), dapagliflozin (Forxiga\textsuperscript{5}) and empagliflozin (Jardiance\textsuperscript{6}), are currently approved by the Food and Drug Administration (FDA) and the European Medicines Agency (EMA) for treatment of T2DM. The pharmacology of SGLT2 inhibition is generally regarded as safe, mainly because of the low risk of hypoglycemia and in conjunction with the benign conditions of GGM and FRG patients. However, recent FDA Drug Safety Communications do suggest that canagliflozin, and to a lesser extent dapagliflozin, could be nephrotoxic in patients with preexisting chronic kidney disease or previous kidney injury\textsuperscript{5} and that gliflozin use is associated with an increased risk of diabetic ketoacidosis\textsuperscript{6}.

Consequently, we compared the cytotoxicity of dapagliflozin, empagliflozin and canagliflozin in quiescent and proliferating human RPTEC/TERT1 cells and investigated the potential direct interference of gliflozins with RPTEC/TERT1 energy metabolism. RPTEC/TERT1 cells were derived from primary human RPTECs immortalized by transfection with telomerase\textsuperscript{7}, which largely retained their expression profile and functionality\textsuperscript{8,9}. Via cultivation for 10 days after reaching confluence, these cells can be converted to a differentiated cell monolayer\textsuperscript{9}, displaying functional and morphological changes that mimic the healthy proximal tubule epithelium in situ. RPTEC/TERT1 cells cultured under proliferating conditions served as model for tubule epithelial cell regeneration\textsuperscript{10}.

We found that canagliflozin, but not dapagliflozin or empagliflozin, exhibited an off-target, and thus SGLT2-independent adverse effect, characterized by the dual inhibition of glutamate dehydrogenase (GDH) and complex I of the mitochondrial electron transport chain (ETC) at pharmacologically relevant concentrations. This combined ETC and GDH inhibition obstructed glutamine input into the tricarboxylic acid (TCA) cycle (i.e. glutamine anaplerosis). As proliferating cells are much more dependent on anaplerosis, this dual inhibition explains why canagliflozin is significantly more toxic for proliferating than for quiescent cells and considerably more potent than classical ETC inhibitors. Thus, our findings demonstrate that canagliflozin interferes with essential energy pathways in glutamine-dependent human cells. This offers a novel mechanistic explanation for the nephrotoxicity reported in patients with increased regenerative cell proliferation, e.g., observed subsequent to acute kidney injury or in chronic kidney disease.

**Results**

**Canagliflozin increased aerobic glycolysis and triggered cytotoxicity in RPTEC/TERT1 cells**

RPTEC/TERT1 cells can be cultured to a contact-inhibited monolayer reflecting primary RPTECs\textsuperscript{7,8}, from here on referred to as differentiated RPTEC/TERT1. We exposed proliferating (day 1 after seeding) and differentiated cells (day 16 after seeding) for 24 h to canagliflozin, dapagliflozin or empagliflozin at concentrations reflecting those observed in clinical and preclinical (rodent) settings\textsuperscript{11–16}. Subsequently, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) reduction (cell viability) and lactate dehydrogenase (LDH) leakage (cytotoxicity) was quantified. Of the gliflozins tested, only canagliflozin showed cytotoxicity (Fig. 1a–d), and was more cytotoxic in proliferating than in quiescent cells (Fig. 1c, d). Necrotic cell death was confirmed by morphological changes and absence of annexin V staining (Fig. S1A). Canagliflozin did not induce autophagy as shown via lysosomal compartment staining with LysoID and immunobots of LC3B conversion (Fig. S1B). Since the MTT assay depends on cellular redox status maintained by mitochondrial activity, we analyzed the cellular glycolytic rate, commonly increased when mitochondrial function is impaired. In differentiated cells, canagliflozin, but not the other two gliflozins, significantly raised cellular glucose to lactate conversion (Fig. 1e). Moreover, canagliflozin increased both total lactate secretion (Fig. S2A) and glucose consumption (Fig. S2B), a characteristic response observed with mitochondrial impairment.

**Canagliflozin specifically inhibited ETC complex I**

Increased glucose to lactate conversion is often the consequence of a blocked mitochondrial electron transport chain\textsuperscript{17,18}. To better characterize the presumed canagliflozin-induced impairment of mitochondrial function, we measured cellular oxygen consumption rates (OCR) and extracellular acidification rates (ECAR) of gliflozin-treated differentiated RPTEC/TERT1 cells (Fig. 2a, b). Canagliflozin at 50 \( \mu \)M, but not at 10 \( \mu \)M, significantly decreased basal respiration, mitochondrial adenosine triphosphate (ATP) production, maximal and spare respiration, but not proton leak and nonmitochondrial respiration (Fig. 2c). Concomitantly, ECAR was increased, suggestive of a direct glycolytic response (Fig. 2b). Equinolar dapagliflozin or empagliflozin had no effect. Plasma membrane permeabilization with digitonin allowed for delivery of specific metabolic substrates while maintaining mitochondrial integrity (Fig. S3). Accordingly, we determined that canagliflozin
completely blocked mitochondrial respiration of the complex I substrates pyruvate, fatty acids and glutamine (Fig. 2d), but did not affect respiration from complex II and IV (Fig. 2e, f). To corroborate complex I-specific inhibition by canagliflizin, we prepared mitochondrial-enriched fractions of RPTEC/TERT1 cells and quantified rotenone-sensitive NADH oxidation (equivalent to complex I activity) as well as atpenin A5-sensitive succinate oxidation (equivalent to complex II activity). As expected, canagliflizin concentration-dependently reduced NADH oxidation (Fig. 2g) but did not affect succinate oxidation (Fig. 2h), thereby confirming canagliflizin-mediated impairment of the mitochondrial ETC via complex I inhibition.

**Canagliflizin generated a unique metabolic profile of amino acid accumulation**

ETC inhibition is well known to harbor an anti-proliferative potential, primarily via reduction of cellular NAD$^+$/NADH ratio and thus depleting aspartate and citrate levels required for anabolism$^{19}$. Cytotoxicity, i.e., cell death induction, in contrast, is rarely observed for...
classical ETC inhibitors under standard culture conditions and is accompanied by ATP depletion or reactive oxygen species production. Accordingly, none of the tested ETC inhibitors—rotenone (complex I), atpenin A5 (complex II), antimycin A (complex III) and oligomycin A (complex V)—were cytotoxic in proliferating RPTEC/TERT1 cells (Fig. 3a), despite the use of concentrations ensuring maximal inhibition as indicated by near-complete ATP depletion (Fig. 3b), massive increase in glycolytic rate (Fig. 3c) and—in case of rotenone and antimycin A—pronounced superoxide production (Fig. 3d). Testimony of its effect on complex I, canagliflozin mildly increased cellular superoxide levels (Fig. 3e) but had no effect on cellular ATP levels (Fig. 3b). We thus concluded that canagliflozin must impact on an additional pathway, beyond the inhibition of complex I activity, to explain the observed cytotoxicity. We uncovered this additional pathway by comparing the steady-state metabolic alterations induced by gliflozins with those of established mitochondrial inhibitors (Fig. 3f), i.e., UK5099, targeting mitochondrial pyruvate transporter, and CPI-613, an inhibitor of pyruvate dehydrogenase (PDH) and α-ketoglutarate dehydrogenase (OGDC), in addition to the already introduced ETC inhibitors. Similar to canagliflozin, UK5099 and CPI-613 increased glucose/lactate conversion rates (Fig. 3c), but did not significantly affect ATP levels (Fig. 3b). The latter could suggest that canagliflozin could also be inhibitive at the level of pyruvate import and pyruvate dehydrogenase, in the glutamine to α-ketoglutarate (αKG) conversion or further down in the TCA cycle (Fig. 3f). As amino acids are key metabolites central to different metabolic pathways, e.g., glycolysis or the TCA cycle (Figs. 3f and 4a), we analyzed the gliflozin-induced alterations within the intracellular amino acid (AA) pool of differentiated RPTEC/TERT1 cells. Canagliflozin, atpenin, UK5099 and CPI-613 treatment dramatically altered the cellular amino acid pool, whereas no such alteration was observed with the typical ETC inhibitors or the other two gliflozins (Fig. 4b and Fig. S4). Strikingly, canagliflozin was the only treatment resulting in a pronounced increase in l-glutamate, l-glutamine and l-alanine (Fig. 4b,c). Glutamine is deaminated to glutamate (Fig. 3f), which can be further metabolized to αKG either via transamination, producing other amino acids such as serine, alanine or aspartate, or via deamination, thereby releasing ammonium (Fig. 4a). The αKG produced is fed into the TCA cycle where it is primarily oxidatively metabolized to produce oxaloacetate (OAA) and ATP, or reductively to citrate. The fact that glutamine and glutamate together accounted for more than 50% of the total free amino acids in control cell lysates (Fig. S5) highlight the importance of the latter metabolic route. The reduction of aspartate, the only amino acid being decreased upon canagliflozin exposure, is in line with data obtained for the ETC inhibitors rotenone (complex I) and antimycin A (complex III). Decreasing the NAD+/NADH ratio by ETC inhibition reduces OAA and thus the level of its transamination product aspartate. Atpenin A5 truncates the TCA cycle at complex II, thereby causing a characteristic complete depletion of aspartate via depletion of OAA. UK5099, in contrast, cuts off the pyruvate supply to mitochondria, resulting in a concomitant OAA and aspartate accumulation. Both depletion or accumulation of OAA and aspartate resulted in an increased dependence on glutaminolysis for ATP production, as indicated by the severely depleted glutamine and glutamate pools observed (Fig. 4c). In addition, canagliflozin, atpenin A5 and UK5099 exposure increased concentrations of the cytosolic (glycolytic) transamination products serine and glycine (Fig. 4c), most likely as a means to increase αKG levels for oxidation in the TCA cycle (Fig. 4a). However, only canagliflozin raised alanine levels, suggesting an increased activity of alanine aminotransferase (ALT). Overall, quantification of cellular amino acids revealed a strong alteration in cellular amino acid metabolism by canagliflozin, which is distinct from classical ETC inhibitors or UK5099 and atpenin A5, as is shown by the principal component analysis (Fig. 4d), and is suggestive of an impaired utilization of glutamine to yield αKG and other metabolites, and thus an impaired glutamine catabolism.
Canagliflozin inhibited glutamate dehydrogenase

Indeed, accumulation of glutamine, glutamate, most transamination products of glutamate as well as the striking accumulation of alanine could potentially be explained by reduced deamination of glutamate via specific inhibition of GDH by canagliflozin (Fig. 4b). We thus utilized mitochondrial enriched fractions of RPTEC/TERT1 cells and analyzed the enzymatic activities of all mitochondrial enzymes directly involved in feeding glutamine into the TCA cycle, either in the presence or absence of the three gliozins. Canagliflozin and to much lower extent dapagliflozin and empagliflozin reduced GDH, but not ALT, aspartate aminotransferase (AST) or α-ketoglutarate dehydrogenase complex (OGDC) activity (Fig. 4e). Moreover, canagliflozin-mediated GDH inhibition was also demonstrated with purified bovine liver GDH, which was not inhibited by the other two gliozins (Fig. 4f), confirming that the unique canagliflozin-induced cellular amino acid profile likely originated from inhibition of mitochondrial GDH.
Fig. 4 (See legend on next page.)
mitochondrial but not glycolytic ATP production (Fig. 5a).

Yang et al.\textsuperscript{26} demonstrated that inhibition of mitochondrial pyruvate uptake by UK5099 altered mitochondrial metabolism towards increased glutaminolysis and induced mitochondrial pyruvate synthesis from glutamine via GDH and the TCA cycle. An additional block of GDH activity caused synthetic lethality in their model. Correspondingly, we found diminished glutamine and glutamate levels under UK5099 treatment indicative of elevated glutaminolysis (Fig. 4c). Based on canagliflozin’s potential to inhibit GDH, we hypothesized that combining canagliflozin with UK5099 would strongly impair mitochondrial energy production. Indeed, while UK5099 alone had no effect, the combination with canagliflozin completely depleted cellular ATP levels (Fig. 5a), starting already 1 h after exposure (Fig. 5b). A reduction of ATP levels by combination of UK5099 and canagliflozin was also observed in medium containing 5 mM but not 20 mM D-glucose, demonstrating the importance of aerobic glycolysis to maintain cellular ATP levels under this condition (Fig. S6).

**Prevention of mitochondrial pyruvate transport sensitized cells towards canagliflozin**

As canagliflozin inhibited ETC complex I and GDH, we determined how specific metabolic settings affect the toxicity of canagliflozin in RPTEC/TERT1 cells. Canagliflozin treatment for 24 h did not result in decreased ATP levels in differentiated cells, suggesting incomplete complex I inhibition with sufficient compensatory glycolytic ATP production. To corroborate this assumption, we exchanged glucose for galactose, thus limiting glycolysis. Under the latter conditions, canagliflozin reduced cellular ATP levels by 32% (Fig. 5a). Addition of 2-deoxyglucose (2-DG), a competitive glycolytic inhibitor, reduced cellular ATP levels by 30%, and in combination with canagliflozin, decreased ATP by an additional 37% down to 30% of control, suggesting that canagliflozin indeed limits mitochondrial but not glycolytic ATP production (Fig. 5a).

**Combined GDH and ETC inhibition by canagliflozin was antiproliferative**

The TCA cycle is a critical source for intermediates required for amino acid, fatty acid and steroid synthesis during cell homeostasis but more importantly during cell proliferation and differentiation. Refueling the TCA cycle (anaplerosis) is thus of key importance for proliferating cells, as would be the case, e.g., in the regenerative phase after kidney injury, or during chronic exposure to toxins resulting in continuously high proliferation rates\textsuperscript{10}. In the TCA cycle, glutamine is the most important anaplerotic precursor\textsuperscript{27}. Beyond increased glutamine catabolism, proliferating cells show high glycolytic rates to support ribose production as well as serine and glycine synthesis required for biosynthetic reactions.\textsuperscript{28} Correspondingly, proliferating RPTEC/TERT1 cells showed almost equal
consumption of glucose and glutamine, and compared to
differentiated cells, overall consumption is increased 5–6-
old (Fig. 6a). Since no net excretion of glutamate was
observed, it can be assumed that most of the glutamine
consumed is indeed utilized for anabolic reactions.

Furthermore, proliferation of RPTEC/TERT1 cells
depended on availability of both glucose and glutamine,
whereas absence of either resulted in cytostasis (Fig. 6b).
In order to characterize the impact of canagliflozin on
proliferation, we exposed proliferating cells to canagliflozin in medium containing 4 mM L-glutamine and 20
mM D-glucose to avoid glucose limitation, and quantified
cell numbers after 72 h. Canagliflozin exposure resulted in
a concentration-dependent reduction of proliferation,
whereby 20–30 µM canagliflozin were cytostatic and
concentrations >30 µM were cytotoxic (Fig. 6c). Strik-
ingly, the GDH inhibitor epigallocatechin gallate (EGCG)29 also caused cytotoxicity, demonstrating that
GDH activity is crucial for this cell line during prolifera-
tion (Fig. 6c). In contrast, neither dapagliflozin nor
empagliflozin had any effect on cell proliferation when
tested to up to 100 µM (Fig. S7). Moreover, complete ETC
inhibition with rotenone, antimycin A or oligomycin A,
and TCA truncation with atpenin A5 significantly
reduced proliferation but did not cause cell death during
72 h of incubation (Fig. 6d), supporting the hypothesis
that GDH inhibition by canagliflozin is responsible for cell
death in RPTEC/TERT1 cells. Final confirmation of this
was obtained by co-treating proliferating cells with ETC
inhibitors or UK5099 in combination with increasing
concentrations of canagliflozin. Addition of rotenone or
atpenin A5 did not alter cytotoxicity of canagliflozin,
corroborating that the mechanism underlying the
observed cytotoxicity is independent of complex I and II
activity. Therefore, canagliflozin-mediated toxicity cannot
originate from ETC inhibition (Fig. 6e). However, under
conditions of increased glutaminolysis induced by
UK5099, RPTEC/TERT1 cells were sensitized towards
canagliflozin but not towards rotenone-mediated inhibi-
tion (Fig. 6e and Fig. S8). With these data we were able to
demonstrate that the antiproliferative and cytotoxic
activity of canagliflozin in RPTEC/TERT1 cells is medi-
ated mainly by inhibiting GDH-dependent glutamine
anaplerosis but not ETC complex I inhibition.

Discussion
Recent FDA Drug Safety communications warned of
gliflozin treatment-associated diabetic ketoacidosis and an
increased risk of acute kidney injury primarily in con-
junction with canagliflozin but also with dapagliflozin
T2DM treatment, specifically relevant for patients with
preexisting chronic kidney disease or a history of kidney
injury5,6. These patients present with a major reduction of
physiologically functional nephrons, sustained regen-
ervative proliferation and increased areas of non-functional
scar tissue30. Concomitantly, the remaining intact
nephrons are challenged not only by having to maintain
renal functionality but also, under the condition of con-
current T2DM treatment with gliflozins, with an
increased in situ (per proximal epithelial cell) concentration of gliflozins, a fact often noted also for many other therapeutics, which can be countered by dosing adjustment31. Accordingly, the typical Cmax for gliflozins calculated from the average T2DM patient may be largely surpassed in patients with preexisting kidney injury, chronic kidney disease32 or elderly patients subject to polypharmacy33–35. Most of the gliflozin-associated cases of acute kidney injury and ketoacidosis occur under canagliflozin treatment5,6, which suggests a compound-specific rather than a class effect.

Despite their capacity for inhibiting SGLT2, gliflozins did not result in glucose limitation in the human proximal tubule epithelial cell line RPTEC/TERT1 in vitro, even at concentrations that were close to or exceeded the Cmax observed under therapeutic conditions. On the contrary, glucose uptake and utilization was enhanced upon canagliflozin exposure (Fig. S2B), thus suggesting that the observed findings are SGLT independent. By quantification of mitochondrial activities combined with a metabolomics approach, we discovered that at similar concentrations canagliflozin impaired two discrete mitochondrial enzymatic functions, namely GDH and ETC complex I. The latter inhibition and their downstream effects were observed at concentrations of 10–50 µM, thus reflecting the clinical Cmax (10 µM) and the concentrations (approximately 65 µM) employed in the animal experiments13. The canagliflozin-mediated inhibition of complex I activity in cancer cell lines was recently published by Villani et al.36 thus corroborating the off-target findings we observed in human renal epithelial cells. Hawley et al.37 demonstrated that canagliflozin but not dapagliflozin or empagliflozin activated AMP-activated protein kinase pathway in vivo via mitochondrial dysfunction, independent of its effect on glucose uptake, also supporting our interpretation that the canagliflozin findings we report here are SGLT independent, substance specific and not a class effect.

Remarkably, GDH and ETC activity have a well-documented interplay with a direct consequence for glutamine anaplerosis. Under physiological conditions ETC activity maintains a high NAD+/NADH ratio, allowing for extraction of the glycolytic metabolite 3-PG for serine and glycine synthesis by glutamate transamination. Pyruvate and glutamate can be used by mitochondrial ALT to produce alanine and α-ketoglutarate. OAA levels are high allowing for AST activity. Deamination of glutamate by GDH is dispensable for many cells under this conditions. ETC inhibition reduces NAD+/NADH ratio forcing pyruvate to lactate conversion to maintain glycolytic flux and reducing OAA levels. Thereby, 3-PG, pyruvate and OAA become unavailable for transamination and cells depend on GDH activity to secure glutamine anaplerosis. By inhibiting ETC complex I and GDH, canagliflozin interferes with both partially redundant pathways allowing glutamine anaplerosis. This results in glutamine and glutamate accumulation and depletion of TCA metabolites leading to proliferation arrest and finally cytotoxicity.
glycolysis) and decreasing OAA availability. Hence, similar to glucose restriction under ETC inhibition, substrates for transamination reactions are unavailable, resulting in GDH dependence for glutamate to αKG conversion (Fig. 7b). Indeed, the GDH inhibitor EGCG was reported to exhibit cytotoxicity when combined with either glucose deprivation or UK5099, prohibiting pyruvate transamination. Consequently, canagliozin, by inhibiting both the ETC and GDH, reduced the availability of glutamine for the TCA cycle, which resulted in glutamine and glutamate accumulation in differentiated cells and cytotoxicity in proliferating cells, especially as the latter are depleted of anabolic precursors (Fig. 7c).

Our data also showed that RPTEC/TERT1 strongly rely on GDH activity, even in presence of sufficient levels of glucose. In fact, proliferating RPTEC/TERT1 consumed remarkably high amounts of glutamine, i.e., comparable to the consumption of glucose, without secreting glutamate (Fig. 6a). The results obtained with the GDH inhibitor EGCG support the finding that proliferating RPTEC/TERT1 cells are exceptionally sensitive to GDH inhibition (Fig. 6c) when compared to cell types used in other studies, where an effect of EGCG was only observed in combination with inhibitors of glycolysis or mitochondrial pyruvate transport.

Indeed, proximal tubule epithelial cells under physiological conditions in vivo showed high GDH activity and exhibited very little glycolytic activity, producing OAA and acetyl-coenzyme A for citrate synthesis primarily from glutamine and β-oxidation of fatty acids, respectively. In addition to its role in anaplerosis, GDH is also critical for renal regulation of pH homeostasis via the production of ammonium (Fig. 4b and Fig. 7). Indeed, glutaminolysis and GDH activity can be elevated to up to fivefold in the proximal tubule epithelium in response to metabolic acidosis. Thus, the observed GDH inhibition by canagliozin provides a logical explanation for the reported increased incidence of ketoacidosis observed primarily under canagliozin treatment. In summary, our results revealed a hitherto unknown mechanism of mitochondrial interference by canagliozin that provides for a novel explanation of canagliozin-induced acute kidney injury and ketoacidosis by inhibition of ETC complex I and GDH.

Materials and methods

Materials
Canagliozin, dapagliozin, empagliozin, antimycin A, oligomycin A and CPI-613 were from Selleckchem (Munich, Germany). Rotenone, UK5099, thienoyltrifluoroacetone (TTFA) and EGCG were from Sigma-Aldrich (Munich, Germany) and aptenin A5 was purchased from Enzo Life Sciences (Lörrach, Germany).

Cell culture
RPTEC/TERT1 cells (Everycyte GmbH, Vienna, Austria) were cultured in a 1 to 1 mixture of Dulbecco’s modified Eagle’s medium (Life Technologies, Darmstadt, Germany) and Ham’s F-12 nutrient mix (Life Technologies) with 5 mM d-glucose final, supplemented with 2 mM GlutaMAX (Life Technologies), 5 µg/ml insulin (Sigma-Aldrich), 5 µg/ml transferrin (Sigma-Aldrich), 5 ng/ml sodium selenite, 100 U/ml penicillin, 100 µg/ml streptomycin, 10 ng/ml EGF (Sigma-Aldrich) and 36 ng/ml hydrocortisone (Sigma-Aldrich). Cells were subcultured after establishing contact-inhibited monolayer and reseeded at 30% confluence. Proliferating cells were used at day 1 (30–40% confluence). Differentiated cells were cultured for 16 days before use.

Cell viability and cytotoxicity
Cells were treated with compounds in the presence of 0.5% dimethyl sulfoxide (DMSO), 0.5% DMSO alone (solvent control) or without any treatment (non-treated) as indicated. To quantify cell viability, treatment was exchanged for fresh medium containing 0.5 mg/ml MTT followed by incubation at 37 °C for 45 min. MTT was removed and cells were lysed in 95% isopropanol and 5% formic acid. Formazan absorption was measured at 550 nm using a Tecan M200pro microplate reader (Tecan, Männedorf, Switzerland). Values obtained for dead cells (0.1% Triton X-100 treated) were subtracted from all other values and data were expressed as percentage of non-treated cells. Cell death was analyzed separately by quantification of LDH activity in supernatant and cell lysate. Treatment medium was harvested and cells were lysed in 0.1% Triton X-100/phosphate-buffered saline (PBS). LDH activity was quantified as described in ref. using a microplate reader. The %LDH leakage was expressed as LDH activity in treatment supernatant divided by the combined activity in supernatant and lysate. LDH leakage obtained for solvent control (0.5% DMSO) was normalized to 0%. Basal LDH leakage was ~15% for proliferating and ~6% for differentiated RPTEC/TERT1.

Extracellular metabolite quantification
Lactate was quantified exactly as published by Limonciel et al. Glucose content was quantified using the glucose oxidase (GOD), peroxidase (POD) reaction coupled to 2,2’-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid) (ABTS). Briefly, 10 µl medium samples were mixed with 90 µl reaction solution containing 100 U/ml GOD (Sigma-Aldrich), 0.25 U/ml POD (Sigma-Aldrich) and 1.5 mg/ml ABTS (Sigma-Aldrich) in 0.1 M Na2HPO4 pH 6.5 followed by incubation for 40 min at room temperature. Absorption at 420 nm was quantified using a plate reader and glucose content was calculated according to a d-glucose calibration curve (0–2 mM). Glutamate was
quantified using a modification of the lactate assay, replacing LDH (Sigma-Aldrich) by GDH (Sigma-Aldrich) and using a glutamate calibration curve (0–12.5 mM). Glutamine was deaminated to glutamate prior to quantification. Briefly, samples were acidified by addition of 0.25× volumes of 500 mM acetic acid pH 5 followed by addition of 0.25× volumes of 5 U/ml glutaminase (Sigma-Aldrich) and incubation for 1 h at 37 °C. Samples were neutralized by addition of 0.25× volumes 1 M Tris base pH 8.3 and glutamate content was quantified as described. Samples incubated in the absence of glutaminase served as background reading for basal glutamate content. A glutamate calibration curve (0–12.5 mM) incubated with glutaminase was used for quantification.

Seahorse extracellular flux analysis
RPTEC/TERT1 cells were seeded in in Seahorse XFe24 culture plates (Agilent, Waldbronn, Germany) and cultured for 16 days. Cells were analyzed using the Seahorse XF Cell Mito Stress Kit with gilofinos or DMSO being injected via port A as indicated. Cell permeabilization for assessing activity of individual ETC complexes was performed as published by Salabei et al. In brief, cells were exposed to 50 µM canagliflozin or 0.5% DMSO (sub-cytaneous (s.c.)) in mannitol and sucrose (MAS) buffer followed by injection of 25 µg/ml digitonin and 1 mM adenosine diphosphate (ADP) in combination with specific substrates for complex I or II (see figure legends) via port A. Coupling of mitochondria was assessed by addition of oligomycin A (1 µM) via port B and subsequent addition of rotenone (1 µM) and antimycin A (1 µM) via port C. Complex IV-dependent respiration was assessed by subsequent addition of tetramethylphenylenediamine (TMPD; 0.5 mM) and ascorbate (2 mM). All concentrations indicated were final concentrations used.

Preparation of mitochondria-enriched fractions
RPTEC/TERT1 cells (~3×10^6) were harvested by trypsinization and lysed using a glass-teflon homogenizer. Liquid phase after centrifugation for 10 min at 600 × g and 4 °C was collected and centrifuged for 10 min at 11,000 × g and 4 °C. Pellet was washed with hypotonic buffer (25 mM potassium phosphate (KPP), 5 mg MgCl₂, pH 7.2) and centrifuged again. Resulting pellet was dissolved in hypotonic buffer and subjected to three freeze/thaw cycles in liquid N2 prior to enzymatic activity assessments.

Biochemical quantification of mitochondrial activities
Enzymatic activities were quantified in mitochondrial-enriched fractions. Complex I and II activities were quantified as previously published with following modifications. Briefly, 25 µl sample (0.7 µg/µl protein) was mixed with 200 µl reaction solution containing 2.5 mM fatty acid bovine serum albumin (BSA; Sigma-Aldrich), 130 µM NADH (Roth, Karlsruhe, Germany), 2 mM KCN (Sigma-Aldrich), 3.6 µM antimycin A, 65 µM ubiquinone 1 (Sigma-Aldrich) and 60 µM 2,6-Dichlorophenol-indophenol (DCIP; Roth) in 25 mM KPP pH 7.5 with or without compound to be tested. The linear phase of the absorption decrease was determined for 3–5 min using a microplate reader. Complex I-independent activity was determined by addition of 5 µM rotenone and was subtracted from all other conditions. Activities were normalized to 0.5% DMSO-treated samples. Complex II activity was determined using same sample volume and readout as for complex I and KPP buffer containing same BSA, KCN, antimycin A, rotenone, ubiquinone 1, DCIP and 20 mM sucinate instead of NADH. Complex II-independent activity was determined by addition of 1 µM atepinin A5. ALT and AST activity were quantified using commercial kits (Sigma, #MAK052 and #MAK055) and 70 µg or 12.5 µg total protein, respectively. GDH activity was quantified using 50 µg protein in a 200 µl reaction containing 5 mM αKG, 1.25 mM NADH, 5 µM rotenone, 3.6 µM antimycin A, 10 mM NH₄Cl and 5 mM MgCl₂ in KPP buffer pH 7.2 with or without gilofinos. The decrease in NADH absorption at 340 nm was observed for 30 min using a microplate reader. GDH specificity was determined by subtracting activity of a parallel reaction lacking αKG. Purified bovine liver GDH (Sigma, #G2626) was used at 0.01 U per 200 µl reaction. OGDC activity was determined as previously published with following modifications. Briefly, 100 µl sample (0.25 µg/µl protein) was mixed with 100 µl reaction solution containing 10 mM KPP pH 7.4, 200 µM EDTA, 2 mM CaCl₂, 2 mM MgCl₂, 100 µM thiamine pyrophosphate (Sigma-Aldrich), 1 mM dithiothreitol, 150 µM coenzyme A (Sigma-Aldrich), 1.6 mM βNAD (Roth), 5 mM αKG, 5 µM rotenone and 3.6 µM antimycin A. Increases of NADH absorption at 340 nm was monitored for 15 min using a plate reader.

Superoxide quantification
Superoxide was measured in differentiated RPTEC/TERT1 cells cultured in 96-well plates using a Superoxide detection Kit (Enzo Life Sciences) according to the manufacturer’s instructions using a microplate reader.

ATP quantification
After treatment as indicated in 96-well plates, differentiated RPTEC/TERT1 cells were washed with PBS and lysed in 100 µl 50% ice-cold methanol for 10 min at −20 °C. Samples were diluted 1:5 with H₂O and 10 µl samples were mixed with 90 µl ATP determination kit reagent (Thermo Fisher) and incubated for 7 min prior reading of luminescence using a microplate reader. ATP levels of exposed cells were normalized to 0.5% DMSO-treated cells.
Cellular AA quantification
RPTEC/TERT1 cells were seeded, reached confluency at 6 days after seeding and were then differentiated for additional 10 days in 6 cm dishes (yielding 6 × 10⁶ cells) followed by treatment as indicated for 24 h. Cells were washed with PBS and scraped on ice in 1.4 ml 50% (V/V) ice-cold methanol/water. Samples were transferred to clean 1.5 ml tubes and shaken for 30 min at 4 °C followed by centrifugation for 10 min at 16,000 × g at 4 °C. Liquid phase was collected and evaporated using a vacuum concentrator followed by solubilization in 150 µl 2% 5-sulfosalicylic acid. AAs were quantified using a Sykam S433 AA analyzer (Sykam, Fürstenfeldbruck, Germany)⁵⁴. Briefly, AAs were separated by high-performance liquid chromatography and subsequent post-column derivatization with ninhydrin. Samples were directly injected in a volume of 100 µl. Chromatography was performed using a lithium-based anion exchange column loaded with spherical polystyrene resin (7 µm diameter, 10% crosslinks, cat. no. 5125022). Elution was performed using buffers with increasing pH and ion strength (pH 2.9 to pH 12; buffer concentration 0.12 M to 0.45 M), supported by a temperature gradient. Absorbance of the reaction products was quantified at 440 nm (intermediate product; quantifies cysteine and proline) or 570 nm (quantifies all other AAs). AA concentrations were determined relative to a reference standard using the area under the peak method in the ChromStar 7 software (SCPA, Weyhe-Leebste, Germany).

Proliferation assays
Cells were plated at 6 × 10⁴ per cm² in 24-well plates and allowed to adhere overnight. Cells were treated in 950 µl culture medium for 3 days. At the time of treatment, cells on a satellite plate were fixed with 4% paraformaldehyde to determine baseline cell number. After 3 days, cells were washed with PBS, fixed and Hoechst 33342 stained. Number of nuclei was determined using a CellInsight automated microscope (Thermo Fisher).

Statistical analysis
Unless otherwise indicated data are presented as mean ± SEM. Sample size (n) indicates the number of independent experiments performed. Statistical significance was determined using GraphPad Prism Version 5.04, statistical tests as indicated in the figure legends. Principal component analysis and heatmap visualization was carried out using ClustVis⁵⁴.

Acknowledgements
We thank Sophia Hochrein, Elia Pilgram and Sabine Drewitz for technical support and Lisanne Luks and Marcia Maier for helpful discussions. P.F.S., J.D. and S.G. were supported by the University of Konstanz and the Deutsche Forschungsgemeinschaft (RTG 1331). The project was additionally supported by a research grant from Boehringer Ingelheim no. 15958812 to D.R.D.

Authors’ contributions
P.F.S. planned, designed and carried out most experiments, analyzed, interpreted and visualized the data, wrote the initial draft and edited the manuscript. S.B. planned experiments, discussed data and edited the manuscript. S.B. and N.S. carried out additional experiments required for the revision of the manuscript. J.D. was involved in amino acid quantification, analyzed and discussed obtained data and edited the manuscript. S.G. was involved in planning and conducting Seahorse experiments and discussed obtained data. M.L. evaluated data and edited the manuscript. D.R.D. developed and supervised the project, evaluated data and edited the manuscript.

Conflict of interest
The authors declare that they have no conflict of interest.

Publisher’s note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Supplementary Information accompanies this paper at (https://doi.org/10.1038/s41419-018-0273-y).

Received: 15 August 2017 Revised: 20 December 2017 Accepted: 27 December 2017

Published online: 14 February 2018

References
1. Chao, E. C. & Henry, R. R. SGLT2 inhibition—a novel strategy for diabetes treatment. Nat. Rev. Drug. Discov. 9, 551–559 (2010).
2. Wright, E. M., Hirayama, B. A. & Loo, D. F. Active sugar transport in health and disease. J. Intern. Med. 261, 32–43 (2007).
3. Ehrenkranz, J. R., Lewis, N. G., Kahn, C. R. & Roth, J. Phlorizin: a review. Diabetes Metab. Res. Rev. 21, 31–38 (2006).
4. Grempler, R. et al. Empagliflozin, a novel selective sodium glucose cotransporter-2 (SGLT-2) inhibitor: characterisation and comparison with other SGLT-2 inhibitors. Diabetes Obes. Metab. 14, 83–90 (2012).
5. U.S. Food and Drug Administration. FDA Drug Safety Communication: FDA strengthens kidney warnings for diabetes medicines canagliflozin (Invokana, Invokamet) and dapagliflozin (Farxiga, Xigduo XR). [Available from: https://www.fda.gov/Drugs/DrugSafety/ucm505860.htm].
6. U.S. Food and Drug Administration. FDA Drug Safety Communication: FDA warns that SGLT2 inhibitors for diabetes may result in a serious condition of too much acid in the blood 2015 [Available from: https://www.fda.gov/Drugs/DrugSafety/ucm594683.htm].
7. Wesser, M. et al. TERT1 alone immortalizes epithelial cells of renal proximal tubules without changing their functional characteristics. Am. J. Physiol. Ren. Physiol. 295, F1365–F1375 (2008).
8. Aschauer, L. et al. Delineation of the key aspects in the regulation of epithelial monolayer formation. Mol. Cell Biol. 33, 2535–2550 (2013).
9. Aschauer, L., Carta, G., Vogelsang, N., Schlatter, E. & Jennings, P. Expression of xenobiotic transporters in the human renal proximal tubule cell line RPTC/TERT1. Toxicol. In Vitro. 30, 95–105 (2015). 1 Pt A.
10. Berger, K. et al. Origin of regenerating tubular cells after acute kidney injury. Proc. Natl. Acad. Sci. USA 111, 1533–1538 (2014).
11. Bogdanffy, M. S. et al. Nonclinical safety of the sodium-glucose cotransporter 2 inhibitor empagliflozin. Int. J. Toxicol. 33, 456–449 (2014).
12. Devine, D. et al. Pharmacokinetics and pharmacodynamics of canagliflozin, a sodium glucose co-transporter 2 inhibitor, in subjects with type 2 diabetes mellitus. J. Clin. Pharmacol. 53, 601–610 (2013).
13. Marmidi, R. N. et al. Carbohydrate malabsorption mechanism for tumor formation in rats treated with the SGLT2 inhibitor canagliflozin. Chem. Biol. Interact. 221, 109–118 (2014).
14. Rebly, T. P. et al. Carcinogenicity risk assessment supports the chronic safety of dapagliflozin, an inhibitor of sodium-glucose co-transporter 2, in the treatment of type 2 diabetes mellitus. Diabetes Ther. 5, 73–96 (2014).
15. Scheen, A. J. Pharmacokinetic and pharmacodynamic profile of empagliflozin, a sodium glucose co-transporter 2 inhibitor. Clin. Pharmacokinet. 53, 213–225 (2014).
16. Timmerstein, M. et al. Nonclinical toxicology assessments support the chronic safety of dapagliflozin, a first-in-class sodium-glucose cotransporter 2 inhibitor. Int. J. Toxicol. 32, 336–350 (2013).

17. Howell, N. & Sager, R. Cytoplasmic genetics of mammalian cells: conditional sensitivity to mitochondrial inhibitors and isolation of new mutant phenotypes. Somat. Cell Genet. 5, 833–845 (1979).

18. Harris, M. Pyruvate blocks expression of sensitivity to antimycin A and chloramphenicol. Somat. Cell Genet. 6, 699–708 (1980).

19. Gui, D. Y. et al. Environment dictates dependence on mitochondrial complex I for NAD+ and aspartate production and determines cell sensitivity to metformin. Cell. Metab. 24, 716–727 (2016).

20. Birsoy, K. et al. An essential role of the mitochondrial electron transport chain in cell proliferation is to enable asparagine synthesis. Cell 162, 540–551 (2015).

21. Lee, K. K. & Boelsterli, U. A. Bypassing the compromised mitochondrial electron transport with methylene blue alleviates efavirenz/isoniazid-induced oxidative stress and mitochondria-mediated cell death in mouse hepatocytes. Redox Biol. 2, 599–609 (2014).

22. Li, N. et al. Mitochondrial complex I inhibitor rotenone induces apoptosis through enhancing mitochondrial reactive oxygen species production. J. Biol. Chem. 278, 8516–8525 (2003).

23. Halestrap, A. P. The mechanism of the inhibition of the mitochondrial pyruvate transporter by alpha-cyano-4-nitro-benzoate derivatives. Biochem. J. 156, 181–183 (1976).

24. Zachar, Z. et al. Non-redox-active lipoate derivatives disrupt cancer cell mitochondrial metabolism and are potent anticancer agents in vivo. J. Mol. Med. (Berl.) 89, 1137–1148 (2011).

25. Stuart, S. D. et al. A strategically designed small molecule attacks alpha-ketoglutarate dehydrogenase in tumour cells through a redox process. Cancer Metab. 2, 4 (2014).

26. Yang, C. et al. Glioblastoma cells require glutamate dehydrogenase to survive impairments of glucose metabolism or Akt signaling. Cancer Res. 69, 7968–7973 (2009).

27. Choo, A. Y. et al. Glucose addiction of TSC null cells is caused by failed mTORC1-dependent balancing of metabolic demand with supply. Mol. Cell 38, 487–499 (2010).

28. Hensley, C. T., Wasti, A. T. & DeBerardinis, R. J. Glutamine and cancer: cell biology, physiology, and clinical opportunities. J. Clin. Invest. 123, 3676–3684 (2013).

29. Fendt, S. M. et al. Reductive glutamine metabolism is a function of the alpha-ketoglutarate to citrate ratio in cells. Nat. Commun. 4, 2236 (2013).

30. Balaban, R. S. & Mandel, L. J. Metabolic substrate utilization by rabbit proximal tubule. An NADH fluorescence study. Am. J. Physiol. 254, F407–F416 (1988).

31. Hokom, K. G. & Mandel, L. J. Glycerol and oxidative metabolism in primary proximal tubule cultures. Am. J. Physiol. 257, C333–C340 (1989). 2 Pt 1.

32. Mandel, L. J. Mitochondrial substrates, cellular energy production, and the regulation of proximal tubular transport. Annu. Rev. Physiol. 51, 85–101 (1989).

33. Weiner, I. D. & Verlander, J. W. Renal ammonia metabolism and transport. Comp. Physiol. 3, 201–220 (2013).

34. Burke, K. R., Schumacher, C. A. & Harpe, S. E. SGLT2 inhibitors: a systematic review of diabetic ketoadiposis and related risk factors in the primary literature. Pharmacotherapy 37, 187–194 (2017).

35. Schildknecht, S. et al. Requirement of a dopaminergic neuronal phenotype for toxicity of low concentrations of 1-methyl-4-phenylpyridinium to human cells. Toxicol. Appl. Pharmacol. 241, 23–35 (2009).

36. Limonové, A. et al. Lactate is an ideal non-invasive marker for evaluating temporal alterations in cell stress and toxicity in repeat dose testing regimes. Toxicol. In Vitro 25, 1855–1862 (2011).

37. Salabe, J. K., Gibb, A. A. & Hill, B. G. Comprehensive measurement of respiratory activity in permeabilized cells using extracellular flux analysis. Nat. Protoc. 9, 421–438 (2014).

38. Kirby, D. M., Thorburn, D. R., Turnbull, D. M. & Taylor, R. W. Biochemical assays of TCA cycle enzymes and PDHc. Methods Cell Biol. 80, 93–119 (2007).

39. Reisch, A. S. & Elpeleg, O. Biochemical assays for mitochondrial activity: principles and practice. Methods Cell Biol. 97, 581–599 (2009).

40. Reisch, A. S. & Elpeleg, O. Biochemical assays for mitochondrial activity: principles and practice. Methods Cell Biol. 97, 581–599 (2009).

41. Ermova, L. et al. Switching from astrocytic neuroprotection to neurodegeneration by cytokine stimulation. Arch. Toxicol. 91, 231–246 (2017).

42. Matsuoka, T. & Vilo, J. Clustvis: a web tool for visualizing clustering of multivariate data using principal component analysis and heatmap. Nucleic Acids Res. 43, W566–W570 (2015). W1.