Salsalate, but not metformin or canagliflozin, slows kidney cyst growth in an adult-onset mouse model of polycystic kidney disease

Wouter N. Leonharda,1, Xuewen Songb,e,1, Anish A. Kanhaia, Ioan-Andrei Iliutab,e, Andrea Bozovicf, Gregory R. Steinbergd, Dorien J.M. Petersa,e,2, York Pei b,e,2

a Department of Human Genetics, Leiden University Medical Centre, Leiden, the Netherlands
b Division of Nephrology, University Health Network, Toronto, Ontario, Canada
c Division of Nephrology, University Health Network, Toronto, Ontario, Canada
d Laboratory Medicine and Pathobiology, University Health Network, Toronto, Ontario, Canada
e Centre for Metabolism, Obesity and Diabetes Research, McMaster University, Hamilton, Ontario, Canada
f Division of Nephrology, University of Toronto, Toronto, Ontario, Canada
g Laboratory Medicine and Pathobiology, University of Toronto, Toronto, Ontario, Canada

1. Introduction

Autosomal dominant polycystic kidney disease (ADPKD) is the most common inherited kidney disease worldwide with a life-time risk of at least 1/1000 in the general population [1]. Mutations of two genes, PKD1 and PKD2, account for 75–85% and 15–25% of the genetically resolved

Cases [2–5]. Disease progression of ADPKD is highly variable in part due to gene locus and allelic effects with the most severe disease associated with PKD1 protein-truncating mutations, intermediate disease severity with PKD1 non-truncating mutations, and mild disease with PKD2 mutations [3–5]. Progressive increase in cyst number and size with age results in distortion of the normal kidney architecture and ultimately, end stage renal disease (ESRD) in a majority of patients [6]. Additionally, non-kidney related complications such as intracranial arterial aneurysms, polycystic liver, and heart valve defect also contribute to the morbidity and mortality of this disease [7]. Overall, ADPKD accounts for 5–10% of ESRD in the developed countries.

The pathobiology of ADPKD is not well understood but a “threshold model” of cystogenesis is supported by recent studies [7–9]. Under this...
Autosomal dominant polycystic kidney disease (ADPKD) is the most common inherited kidney disease worldwide. Previous preclinical studies showed that metformin slowed PKD progression by activating AMPK; however, translating these findings to patients might require a higher oral metformin dose than can be achieved clinically.

To optimize AMPK-based therapeutic in ADPKD, we tested oral metformin-based combination therapies for their additive (metformin plus canagliptin) and synergistic (metformin plus salsalate) effects and each drug individually in a conditional Pkd1 knockout mouse model. Using dosages expected to yield clinically relevant serum drug levels, we found only salsalate (a pro-drug dimer of salicylate) slowed PKD progression by improving mitochondrial function and reducing inflammation.

We measured serum metformin levels by an in-house assay we developed using liquid chromatography tandem mass spectrometry (LC-MS/MS) at the Clinical Biochemistry Laboratory of the Toronto General Hospital (see Supplementary Information). We measured serum salicylate levels using a commercially available kit (Neogen Corporation) and protein kinase (AMPK). AMPK is a highly conserved and ubiquitously expressed heterotrimeric enzyme complex (consisting of α, β and γ isoforms) that functions as a sensor of cellular energy status [25]. By expressing the drug-binding domain of the AMPK α1 isoform [25,33], importantly, previous studies have indicated that given their distinct mechanisms for activating AMPK, there are synergies between metformin and salsalate therapy for treating fatty liver disease and suppressing mTOR in cancer cells [34,35]. All three drugs have excellent safety profile and may be appropriate for drug repurposing to treat ADPKD [36]. To examine the additive (metformin plus canagliptin) or synergistic (metformin plus salsalate) effect of metformin combination therapy, we tested in this study the efficacy of oral treatment of (i) metformin, (ii) canagliptin, (iii) salsalate, (iv) metformin plus canagliptin, (v) metformin plus salsalate vs. (vi) untreated mutant control in an adult-onset Pkd1 conditional deletion mouse model.

2. Materials and methods

2.1. Mouse experimental protocol

The experimental design of our study is shown in Fig. 1. All the study groups were run concurrently in one large experiment. We used the iKspPkd1fl/fl conditional knock-out (KO) mice on a C57BL6/J congenic background for the testing [10,37]. Twenty male mice/group were treated with tamoifin (150 mg/kg, by oral gavage) at days P18 and P19 to inactivate Pkd1 which will lead to renal failure at around 4 months of age. Drug treatment began on day P40. Metformin (AK-ScientiFL, # I506) was administered orally in drinking water at 1.5 mg/mL (or ~300 mg/kg/day); canagliptin (MedChemExpress, # HY-10451) was administered orally at 10 mg/kg/day (62.5 mg/kg of food pellets (RM 3(E)), Special Diet Services); and salsalate (AK-Scientific, # F817) was administered orally at 400 mg/kg/day (2.5 g/kg of food pellets, Special Diet Services). As control, one group of mice received food pellets generated by the same protocol, but without any drug. Blood urea nitrogen (BUN) was measured weekly starting at day P75 using 30 μL blood samples from the tail vein (Reflotron technology, Roche). All mice with a BUN >20 mmol/L were considered to have ESRD and sacrificed; BUN of WT mice was ~9 mmol/L. When ~50% of the untreated mice reached ESRD at 111–115 days of age, all mice from the control and other treatment groups were sacrificed within 5 days. For mice with BUN >20 mmol/L, the time to ESRD was calculated by linear interpolation of the ages between the last two BUN measurements. The age at ESRD or at censoring (i.e. mice without ESRD at the time of their last BUN measurement between 111 and 115 days of age) was used for Kaplan Meier survival analysis. All animal experiments were approved by the Animal Experiment Committee of the Leiden University Medical Centre and the Commission Biotechnology in Animals of the Dutch Ministry of Agriculture, and performed in accordance to Directive 2010/63/ EU for animal experiments.

2.2. Measurement of drug and cAMP levels

We measured serum metformin levels by an in-house assay we developed using liquid chromatography tandem mass spectrometry (LC-MS/MS) at the Clinical Biochemistry Laboratory of the Toronto General Hospital (see Supplementary Information). We measured serum salicylate levels using a commercially available kit (Neogen Corporation) and gastrointestinal intolerance (in ~30% of patients) coupled with a high hepatic first-pass effect (i.e. liver concentrating a high percentage of drug after gut absorption) raise concerns that the maximal oral dose of metformin (i.e. 2.0 g/day) used clinically may not be sufficient for therapeutic AMPK activation in the kidney [29,30]. Thus, there is a need to develop more effective AMPK-based therapeutics which can be translated to treat patients with ADPKD.

In this study, we explored whether the addition of a second AMPK-activating drug to metformin would enhance its therapeutic effects for experimental treatment of ADPKD. One of these drugs was a recently approved type 2 diabetic medication, canagliptin, also an indirect AMPK activator like metformin [31,32]. In addition, we also examined the effects of salsalate, a prodrug dimer of salicylate, which in contrast to metformin and canagliptin, activates AMPK through direct interactions with the drug-binding domain of the AMPK α1 isoform [25,33].
kidney tissue cAMP using the Direct cAMP ELISA kit (Enzo Life Sciences), according to the manufacturer instructions. Serum drug levels were measured in individual mutant mice after at least one month of treatment in the metformin group \((n = 20)\) and salsalate group \((n = 21)\) and at the time of sacrifice in the metformin group only \((n = 19)\). Canagliflozin was not measured directly, but was expected to induce osmotic diuresis and hence, increased water intake was used as a surrogate marker of the drug effect. Indeed, we observed \(\sim 1.5\) fold increased water intake in canagliflozin-treated mice.

2.3. Histological studies of kidney tissues

Kidneys were fixed in 4% buffered formaldehyde solution and embedded in paraffin. Periodic acid–Schiff (PAS) and Sirius Red staining was performed using standard protocols. F4/80 staining was used to detect macrophages. In brief, proteinase K antigen retrieval was used followed by a 20 min \(0.12\%\) \(\text{H}_2\text{O}_2\) incubation step to block endogenous peroxidase. The sections were then incubated with rat anti-F4/80 (1:100; Serotec), followed by a second incubation step with anti-rat ImmPRESS™ (Vectorlabs). Immune reactions were revealed using diaminobenzidine as a chromogen and counterstained with hematoxylin, dehydrated, and mounted. Quantifications were done using Photoshop software as described previously [37]. Colour palettes were designed to select pixels specific for those within cysts, those of the red Sirius Red signal (large arteries were excluded from the analyses), or those of the brown F4/80 signal. The total number of pixels from the entire section (without cysts) and the total number of pixels from either the Sirius red or F4/80 signal were used to calculate the percentage of Sirius Red or F4/80 staining within the entire kidney section.

2.4. Gene expression and bioinformatics analysis

Using the Mouse Gene 2.0 ST Arrays (Affymetrix), we performed global gene expression profiling of 30 kidney samples (wild-type,
n = 7; KO mice, n = 12, and KO mice treated by salsalate, n = 11) which were selected around the median of the cystic index from each group. Microarray gene expression studies were performed by the Centre for Applied Genomics Core at the Hospital for Sick Children (Toronto, Ontario, Canada). QC measures and filtering are detailed in the Supplementary Information and in Figs. S1, S4, and S5. Differentially expressed genes were split into two groups of up-regulated and down-regulated genes. The lists of up-regulated and down-regulated genes were each used for gene set enrichment analysis (GSEA). We used Enrichr as the primary tool for GSEA [38]. We used Benjamini-Hochberg (false discovery rate) FDR-adjusted p-values and Z-scores computed by Enrichr for the ranking of each gene set. Hierarchical clustering analysis and heatmap were performed using the DNA-Chip Analyzer (dChip) software package [39].

2.5. Western blot analysis

Protein extraction from whole kidney tissues for Western blot analysis are detailed in Supplementary Information. After collecting the supernatants from the kidney lysates, protein extracts were quantitated using Protein Assay Kit (Bio-Rad) and then denatured in Laemmli sample buffer. Proteins were separated on TGX gradient gels (Bio-Rad) and then transferred onto PVDF membranes (Bio-Rad) by using Trans-Blot Turbo System (Bio-Rad). The membranes are incubated in blocking buffer (150 mM NaCl, 20 mM Tris, 5% skim milk, 0.1% Tween 20) for 60 min, and then incubated with primary antibodies at 4 °C overnight. Subsequently, primary antibody binding was detected with horseradish peroxidase (HRP)-conjugated anti-rabbit, or anti-mouse secondary antibodies, and proteins were visualized with an enhanced chemiluminescence detection kit (ECL, Bio-Rad or Cell Signalling Technology). For determination of phospho/total protein levels, immunoblots were first probed for phospho levels, then stripped at RT for 30 min using Re-Blot Plus Mild Antibody Stripping Solution (EMD Millipore) and re-probed overnight to detect total levels. Protein expression was quantitated by densitometric analysis with Imagej software (NIH, http://rsbweb.nih.gov/ij/) according to the guidelines. All the primary and secondary antibodies used for this study are listed in Supplementary Information.

2.6. Quantification of mitochondrial DNA copy number

Total DNA containing mitochondrial and nuclear DNA (nDNA) was isolated by using QiAamp DNA Mini Kit (Qagen). We performed quantitative real-time PCR using Power SYBR® Green PCR Master Mix (Applied Biosystems) with primers specific to a mitochondria-encoded gene, 16S ribosomal RNA (16 s rRNA) (forward: 5′-CCGCAAGGGAA AGATGAAAGAC-3′; reverse: 5′-TCGTGGTCGCTTTGGGTT-3′) and a nuclear-encoded gene, B2m (forward: 5′- GCCAGATCTGAAGTAC-3′; reverse: 5′- GTGAGCAGGATATAAGACCCC-3′), as previously described [40]. Mitochondrial DNA (mtDNA) copy number was estimated by the ratio of 16S rRNA/B2m.

2.7. Statistical analysis

Statistical analysis was performed using GraphPad Prism 7 software (San Diego, CA, USA). All results are expressed as dot-blots with median and interquartile ranges or mean ± SEM. Comparisons involving more than two groups were performed by one-way ANOVA followed by Tukey’s or Dunnett’s test post hoc; p-values corrected for multiple comparisons were reported. The Log Rank (Mantel-Cox) test was used for

Fig. 3. Salsalate reversed multiple pathogenic mediators of cystic kidney disease in iKsp-Pkd1mice. Western blot analyses showed that kidneys from mutant Pkd1 mice (compared to WT mice) displayed increased expression of protein markers for (a) mTORC1 activity (i.e. levels of pS6K), (b) cell proliferation (i.e. PCNA), (c) cell cycle progression (i.e. CDK2), and (d) cAMP; all these changes were significantly attenuated by salsalate treatment. Data presented as mean ± SEM. For kidney cAMP levels, WT (n = 7), KO (n = 8), SAL (n = 9).
the Kaplan Meier survival analyses. The fibrotic index and F4/80 index between salsalate-treated and untreated Pkd1 mice were tested using unpaired Student’s t-tests.

3. Results

3.1. Salsalate, but not metformin or canagliflozin, slowed polycystic kidney disease

The iKsp-Pkd1del mice were treated with clinically relevant drug doses according to experimental protocol (Fig. 1; also see Materials and Methods). When 46% of untreated Pkd1 mutant (‘disease control’) mice had reached ESRD by 111–115 days of age, all the mice from all study groups were sacrificed at that time. The mean serum metformin levels measured by LC-MS/MS in the mutant mice after at least one month of treatment (n = 20) and at the time of sacrifice (n = 19) were 14.6 (90%CI: 18.6–21.4) and 17.2 (90%CI: 14.1–20.3) μM, respectively. The median serum salicylate levels measured by ELISA in the mutant mice after at least one month of salsalate treatment (n = 21) was 222 (IQR: 108–372) μM. By comparison, the mean serum metformin and salicylate levels reported in diabetic patients on steady state clinical doses were 10 μM and 1350 μM, respectively.29,30,41 Moreover, in a pilot study of 5 PKD patients on 1.5–2 g/day of oral metformin, we found a median serum metformin level of 9.18 μM, respectively.29,30,41

Table 1
Top ranked dysregulated Reactome pathways enriched in Pkd1 mutant mouse kidneys but attenuated by salsalate treatment.

| Top 20 down-regulated Reactome pathways enriched in iKsp-Pkd1del mouse kidneys but attenuated by salsalate treatment | Top down-regulated Reactome pathways enriched in iKsp-Pkd1del (vs. WT) mice | Top up-regulated Reactome pathways in salsalate-treated (vs. untreated) iKsp-Pkd1del mice |
|---|---|---|
| **Metabolism** | **Adjusted P-value** | **Z-score** | **Adjusted P-value** | **Z-score** |
| The citric acid (TCA) cycle and respiratory electron transport | 6.49E–82 | −2.25 | 1.88E−69 | −2.25 |
| Respiratory electron transport, ATP synthesis by chemiosmotic coupling, and heat production by uncoupling proteins | 6.94E–67 | 1.97 | 7.30E–50 | −1.97 |
| Complex I biogenesis | 1.12E–51 | 1.98 | 5.43E–38 | −1.98 |
| Metabolism of lipids and lipoproteins | 8.07E−46 | 1.95 | 1.15E−37 | −1.95 |
| Pyruvate metabolism and Citric Acid (TCA) cycle | 3.61E−31 | 2.02 | 3.33E−23 | −2.02 |
| Metabolism of amino acids and derivatives | 1.12E–14 | −2.13 | 6.39E−22 | −2.13 |
| Fatty acid, triacylglycerol, and ketone body metabolism | 5.32E−14 | −2.15 | 2.12E−16 | −2.15 |
| Branched-chain amino acid catabolism | 1.16E–19 | −2.05 | 1.27E−12 | −2.05 |
| Mitochondrial protein import | 2.34E−09 | −1.40 | 5.04E–09 | −1.40 |
| Mitochondrial Fatty Acid Beta-Oxidation | 9.13E–13 | −1.84 | 3.08E–08 | −1.84 |
| Mitochondrial translation elongation | 1.40E–15 | −1.99 | 3.08E–08 | −1.99 |
| Peroxisosomal lipid metabolism | 3.75E−14 | −1.99 | 3.68E–08 | −1.99 |
| Glyoxylate metabolism and glycine degradation | 2.37E−11 | −1.54 | 1.38E–07 | −1.54 |
| Histidine, lysine, phenylalanine, tyrosine, proline and tryptophan catabolism | 5.41E−12 | −1.06 | 3.56E–07 | −1.06 |
| Mitochondrial translation initiation | 3.82E−27 | −2.07 | 4.45E–07 | −2.07 |
| Top 20 up-regulated Reactome pathways enriched in iKsp-Pkd1del mouse kidneys but attenuated by salsalate treatment | **Adjusted P-value** | **Z-score** | **Adjusted P-value** | **Z-score** |
| **Immune system** | 1.01E−28 | −2.23 | 3.12E–35 | −2.23 |
| Extracellular matrix organization | 6.10E−24 | −2.09 | 1.90E–33 | −2.09 |
| Hemostasis | 1.44E−18 | −2.10 | 1.51E–23 | −2.10 |
| Signalling by Rho GTPases | 1.42E−25 | −2.20 | 4.45E−22 | −2.20 |
| Cytokine signalling in immune system | 2.67E−17 | −2.34 | 1.91E−17 | −2.34 |
| RHO GTPase effectors | 2.92E−23 | −2.44 | 3.04E–17 | −2.44 |
| Axon guidance | 9.22E−19 | −2.23 | 8.48E–17 | −2.23 |
| Cell cycle | 9.37E−14 | −2.28 | 8.95E–17 | −2.28 |
| Developmental biology | 2.90E−12 | −2.27 | 1.77E–16 | −2.27 |
| Innate immune system | 1.99E−25 | −2.44 | 1.66E–16 | −2.44 |
| Intercellular matrix interactions | 5.69E−15 | −2.36 | 2.16E–16 | −2.36 |
| Platelet activation, signalling and aggregation | 6.19E−11 | −1.83 | 3.03E–15 | −1.83 |
| Cell-cell communication | 2.49E−12 | −2.06 | 5.11E–15 | −2.06 |
| Non-integrin membrane-ECM interactions | 1.02E−09 | −1.85 | 1.54E–13 | −1.85 |
| Mitotic prometaphase | 8.67E−10 | −1.81 | 5.97E–13 | −1.81 |
| RHO GTPases activate forms | 2.91E−14 | −1.97 | 7.82E–12 | −1.97 |
| Interferon signalling | 1.06E−12 | −1.94 | 1.92E–11 | −1.94 |
| Platelet degranulation | 4.27E−11 | −1.97 | 2.10E–11 | −1.97 |
| Response to elevated platelet cytosolic Ca2+ | 3.35E−07 | −1.58 | 7.99E–11 | −1.58 |
| 2.03E−07 | −1.61 | 9.23E–11 | −1.61 |

* P-values computed by Fisher exact test and adjusted for multiple testing using the Benjamini-Hochberg method.
improvement in kidney survival (Fig. 2a) and reduction in cystic kidney disease severity (Fig. 2b-d). However, the therapeutic effects of salsalate vs. metformin plus salsalate did not differ, suggesting that the therapeutic effects in the latter group was due to salsalate alone. Of note, the therapeutic effect of salsalate was very strong, and comparable to that by oral tolvaptan treatment using the same model and protocol in a separate study (manuscript in preparation). By contrast, treatment with metformin, canagliflozin, alone or together was not effective (Figs. 2, S2). A concern of metformin treatment is its potential for gastrointestinal side-effects including anorexia and diarrhea which can cause volume depletion. However, our metformin-treated mice appeared well and gained weight similar to mice from the other experimental groups throughout the study (Fig. S3). Moreover, there was an excellent correlation between BUN and 2K/2B (Fig. S4) suggesting the loss of GFR was related to the cystic disease burden. Since treatment with metformin or canagliflozin was not effective, we focused our mechanistic study on salsalate.

3.2 Salsalate attenuated pathogenic mediators in Pkd1 mutant mouse kidneys

Aberrant mTORC1 activation [16–22] and increased cAMP signalling [7,11,13] in cystic tissues are two well-validated pathogenic mechanisms driving cyst growth in ADPKD. In turn, both of these pathogenic mechanisms have been shown to increase cyst epithelial cell proliferation while increased cAMP signalling also drives cyst fluid secretion [11]. By Western blot analyses, we found that Pkd1 mutant mouse kidneys (compared to wild type) displayed increased markers for

### Table 2

Top ranked dysregulated biological processes enriched in iKsp-Pkd1<sup>del</sup> mutant mouse kidneys but attenuated by salsalate treatment.

| Top 20 down-regulated biological processes enriched in iKsp-Pkd1<sup>del</sup> mouse kidneys but attenuated by salsalate treatment | Top down-regulated biological processes enriched in iKsp-Pkd1<sup>del</sup> (vs. WT) mice | Top up-regulated biological processes in salsalate-treated (vs. untreated) iKsp-Pkd1<sup>del</sup> mice |
|--------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|
| Respiratory electron transport chain                                                                                 | Adjusted P-value<sup>*</sup> 6·90E–34  4·96E–28  1·36  Z-score 4·96E–28  1·36 | Adjusted P-value<sup>*</sup> 6·35E–30  7·48E–28  1·90  Z-score 7·48E–28  1·90 |
| Mitochondrial ATP synthesis coupled electron transport                                                               | 3·35E–30  1·90  4·74E–25  1·90                                                                 |
| Fatty acid beta-oxidation                                                                                              | 2·82E–23  1·52  1·12E–21  1·53                                                                 |
| Fatty acid catabolic process                                                                                           | 4·98E–25  1·82  2·98E–19  1·83                                                                 |
| Mitochondrial respiratory chain complex assembly                                                                       | 1·44E–29  1·78  2·98E–19  1·78                                                                 |
| Mitochondrial respiratory chain complex biogenesis                                                                   | 9·04E–27  1·58  8·73E–19  1·58                                                                 |
| NADH dehydrogenase complex assembly                                                                                   | 9·04E–27  1·57  8·73E–19  1·57                                                                 |
| Mitochondrial respiratory chain complex I assembly                                                                    | 9·04E–27  1·41  8·73E–19  1·41                                                                 |
| Mitochondrial electron transport, NADH to ubiquinone                                                                   | 2·54E–20  1·72  2·41E–18  1·72                                                                 |
| Fatty acid oxidation                                                                                                   | 1·84E–19  1·81  1·08E–14  1·81                                                                 |
| Monocarboxylic acid metabolic process                                                                                  | 1·56E–11  1·64  7·54E–11  1·65                                                                 |
| Fatty acid catabolic process                                                                                           | 6·96E–12  1·82  1·31E–10  1·83                                                                 |
| Cellular respiration                                                                                                   | 2·41E–15  1·24  4·59E–10  1·24                                                                 |
| Cellular amino acid catabolic process                                                                                    | 1·59E–11  1·68  7·19E–10  1·69                                                                 |
| Branched-chain amino acid catabolic process                                                                           | 1·79E–11  1·77  2·53E–09  1·78                                                                 |
| Branched-chain amino acid metabolic process                                                                           | 1·80E–09  1·74  5·34E–09  1·75                                                                 |
| Fatty acid metabolic process                                                                                           | 5·62E–10  1·31  5·34E–09  1·32                                                                 |
| Fatty acid beta-oxidation using acyl-CoA dehydrogenase                                                                 | 4·92E–07  2·31  1·06E–07  2·32                                                                 |
| Coenzyme metabolic process                                                                                             | 8·55E–07  1·44  1·39E–07  2·48                                                                 |
| Mitochondrial transport                                                                                               | 4·26E–18  1·33  5·67E–07  1·33                                                                 |

| Top 20 up-regulated biological processes enriched in iKsp-Pkd1<sup>del</sup> mouse kidneys but attenuated by salsalate treatment |
|--------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|
| Extracellular matrix organization                                                                                       | Adjusted P-value<sup>*</sup> 2·11E–27  1·76E–35  1·65  Z-score 1·76E–35  1·65 |
| Cellular response to cytokine stimulus                                                                                 | 5·98E–20  1·17  6·33E–23  1·17                                                                 |
| Regulation of cell migration                                                                                           | 1·98E–16  1·27  8·07E–22  1·27                                                                 |
| Cytokine-mediated signalling pathway                                                                                    | 1·05E–23  1·34  1·42E–21  1·34                                                                 |
| Neutrophil mediated immunity                                                                                            | 2·98E–08  1·90  1·11E–15  1·90                                                                 |
| Neutrophil degranulation                                                                                               | 4·15E–08  1·96  1·53E–15  2·01                                                                 |
| Response to cytokine                                                                                                   | 1·22E–12  1·28  1·53E–15  1·28                                                                 |
| Neutrophil activation involved in immune response                                                                       | 4·15E–08  1·23  2·61E–15  1·25                                                                 |
| Regulation of apoptotic process                                                                                        | 2·08E–11  1·77  1·26E–14  1·78                                                                 |
| Positive regulation of gene expression                                                                                | 1·47E–13  1·66  2·87E–13  1·66                                                                 |
| Transmembrane receptor protein tyrosine kinase signalling pathway                                                       | 5·92E–14  1·74  1·75E–12  1·73                                                                 |
| Toll-like receptor signalling pathway                                                                                    | 2·55E–10  1·29  1·75E–12  1·29                                                                 |
| Regulation of cell proliferation                                                                                        | 6·56E–13  1·13  2·57E–12  1·12                                                                 |
| Protein phosphorylation                                                                                               | 1·50E–11  1·09  1·77E–11  1·09                                                                 |
| Positive regulation of transcription, DNA-templated                                                                    | 8·62E–16  1·71  2·36E–11  1·71                                                                 |
| Cellular response to mechanical stimulus                                                                               | 2·90E–10  2·15  3·14E–11  2·15                                                                 |
| Positive regulation of cell migration                                                                                  | 4·24E–08  1·87  3·14E–11  1·90                                                                 |
| Pattern recognition receptor signalling pathway                                                                       | 1·17E–06  1·45  3·14E–11  1·47                                                                 |
| Platelet degranulation                                                                                                 | 5·35E–06  1·38  3·14E–11  1·42                                                                 |
| Positive regulation of cell proliferation                                                                             | 5·55E–09  1·54  3·98E–11  1·54                                                                 |

*P-values computed by Fisher exact test and adjusted for multiple testing using the Benjamini-Hochberg method.
mTORC1 activity (i.e. pS6K), cell proliferation (i.e. PCNA), and cell cycle progression (i.e. CDK2), as well as increased levels of cAMP; all of these changes were significantly attenuated by salsalate treatment (Fig. 3). However, we were unable to directly demonstrate increased AMPK activity which might be due to a number of technical issues (see Supplementary Information).

3.3 Salsalate treatment attenuated defective metabolism in Pkd1 mutant mouse kidneys

For a more comprehensive analysis of the molecular alterations, we performed gene expression profiling. Tables 1 and 2 highlight the key findings. Of interest, the top 20 down-regulated Reactome pathways and biological processes enriched in Pkd1 mutant mouse kidneys but attenuated by salsalate treatment were all related to metabolism. These results confirm our previous data indicating a generalized defect in metabolism in Pkd1 mutant kidneys with impaired fatty acid oxidation, amino acid catabolism and oxidative phosphorylation [42], all these changes were attenuated by salsalate (Fig. 4a). Reduced AMPK activity in cystic tissues is expected to decrease the activity of PGC-1α, a master regulator for the transcriptional factors PPARα, ERRα, and ERRγ; all play a critical role in fatty acid oxidation and mitochondrial biogenesis [43]. By Western blot analysis, we confirmed reduced PGC-1α expression in Pkd1 cystic kidneys, which was attenuated by salsalate (Fig. 4b). Our in-silico analysis also predicted Ppara and its heterodimeric partner, Rxr, as the top-ranked transcriptional factors inhibited in Pkd1 mutant kidneys, but active in the salsalate treatment group (Table S1). Consistent with defective mitochondrial biogenesis in ADPKD [44–46], we found an inverse correlation between relative mitochondria copy number (using mtDNA/nDNA as a readout) and kidney weight/body weight in Pkd1 cystic kidneys ($r = -0.8, p < 0.0001$) (Fig. 4c); the salsalate treated mice had higher mtDNA/nDNA ratios consistent with their milder PKD phenotype (Fig. 4d).

3.4 Salsalate treatment attenuated inflammation in Pkd1 mutant mouse kidneys

Recent studies have documented that kidney tissue inflammation can promote progression of experimental ADPKD. Specifically,
pharmacological and genetic manoeuvres depleting kidney tissue macrophage infiltration were shown to attenuate cystic disease severity [47,48]. Interestingly, multiple top upregulated Reactome pathways and biological processes enriched in Pkd1 cystic kidneys but attenuated by salsalate were involved in immunity (Tables 1 and 2). By gene expression analysis, we found an increased expression of markers for damage-associated molecular patterns (DAMPs) and their receptors [49,50], macrophages, inflammation and fibrosis in Pkd1 cystic kidneys; all of these changes were attenuated by salsalate treatment (Fig. 5a). NFκB is a key regulator of innate immunity [49] and its p65 (Rela) subunit was predicted to be a top-ranked activated transcriptional factor in Pkd1 mutant kidneys, inhibited by salsalate (Table S1). We confirmed this finding by Western blot analysis (Fig. 5b). Consistent with our gene expression data, we found increased fibrosis and macrophage infiltration in Pkd1 mutant kidneys, which were attenuated by salsalate treatment (Figs. 5c, d and 6).

4. Discussion

Given the concerns that therapeutic AMPK activation in the kidney might require a higher dose of oral metformin than can be achieved clinically [26,27,29,30], we tested metformin-based combination therapies for their additive (i.e. metformin plus canagliflozin) and synergistic (i.e. metformin plus salsalate) effects in an adult-onset Pkd1-deletion mouse model using dosages expected to yield clinically relevant drug levels [29–33]. We found that only salsalate (a prodrug dimer of salicylate) was effective in slowing PKD progression with serum salicylate levels at the lower clinical therapeutic range (i.e. 0.7–2.2 mmol/L) [41,51]. By contrast, metformin was not effective despite serum drug levels in the clinical therapeutic range (i.e. 10 μmol/L) [29,30]. Comparing our study to that by Takiar et al., which reported that intra-peritoneal metformin treatment was effective in slowing PKD [19], both studies used the same daily dose (i.e. 300 mg/kg). However, the bioavailability of oral metformin is only ~50% [26,27]. More recently, oral metformin treatment was reported to slow PKD progression in a miniature pig model using a dose of 41.7 mg/kg/day [52]. However, the estimated human equivalent dose used for this study is 2.77 g/day, which is higher than the usual dose (2 g/day) used clinically (see supplementary data) [53]. Differences in animal models and drug dosing in these studies may account for their discrepant results. However, serum metformin levels were not measured in these studies rendering them difficult to compare to our study. Preclinical studies that employ drug dosages modelling the clinical setting are more likely to have clinical relevance.

Increased mTORC1 [16–20] and cAMP signalling [11–13] in cystic tissues are two major pathogenic mechanisms driving cyst growth in ADPKD. In turn, activation of these signalling pathways have been shown to increase cystic epithelial cell proliferation in virtually all models of PKD while increased cAMP signalling also drives cyst fluid secretion [11]. In this context, our results are consistent with upregulation of these signalling pathways in the Pkd1 mutant mouse kidneys which was attenuated by salsalate treatment. Our findings suggest that salsalate may have an additive therapeutic effect with tolvaptan by targeting cAMP signalling [7,11–13].

Recent studies have shown that Pkd1 cystic tissues display metabolic reprogramming consistent with a shift of the normal utilization of glucose through mitochondrial oxidative phosphorylation to aerobic glycolysis (a.k.a. the “Warburg effect”) [20,21,54]. Instead of fully oxidizing one glucose molecule in the mitochondria to yield 36 ATP,
cystic tissues prefer a metabolic pathway that generates only 4 ATP and 2 lactate molecules; the latter are then used to build macromolecules (nucleotides, proteins, lipids) to support cell proliferation [54]. In addition, defective fatty acid β-oxidation and altered mitochondrial function have been observed both in vitro and in animal models of ADPKD [44,55,56]. Indeed, the top 20 most down-regulated gene pathways in Pkd1 mutant kidneys involved multiple metabolic processes ranging from tricarboxylic acid cycle, catabolism of fatty acids and amino acids, oxidative phosphorylation, to mitochondrial biogenesis; these changes suggest a generalized defective metabolism in Pkd1 mutant kidneys which was attenuated by salsalate treatment. Consistent with the findings by Lakhia et al. [57], our data suggest that decreased activity of PPARα and PGC-1α (co-activator of PPARα) might contribute to the defective fatty acid oxidation and mitochondrial biogenesis in Pkd1 cystic tissues. Of interest, fenofibrate, a PPARα agonist, was found in the former study to attenuate cystic kidney and liver disease in Pkd1KO mice [57]. However, the clinical use of this drug is limited by its potential nephrotoxicity, especially for patients with chronic kidney disease [58].

Tissue injury can initiate an inflammatory response through the actions of damage-associated molecule patterns (DAMPs) which comprise a heterogeneous group of molecules released during cell necrosis, tissue repair and remodelling. In turn, DAMPs can trigger innate immunity by activating Toll-like receptors, purogenic receptors, or NLRP3 inflammasome [49]. Recent studies have shown that depletion of kidney macrophage infiltration by pharmacological and genetic manoeuvres attenuated Pkd1 cystic disease severity [47,48]. We found that Pkd1 cystic kidneys were enriched with increased expression of genes encoding for multiple necrosis-related DAMPs, alarmins and pathway proteins (e.g. multiple toll-like receptors (TLRs), Myd88, Nfκb; Ripk3 and Mlkl) for innate immunity; all of these changes were attenuated by salsalate treatment [50]. Taken together, our data are consistent with an emerging body of literature suggesting necroinflammation can provide an auto-amplification loop of necrosis and inflammation that mediates progression of chronic kidney diseases [50]. Thus, salsalate treatment may confer a therapeutic effect for other kidney diseases beyond ADPKD.

There are limitations or potential concerns in our study. First, our selection of samples closest to the median PKD severity for the gene expression studies might create bias by not including the outliers; nevertheless, this approach provides a valid assessment for the “average effect” of each study group. Second, metformin might result in volume depletion due to anorexia and diarrhea and a functional (i.e. renal vasocstriction rather than structural damage) cause for increased BUN. However, our metformin-treated mice appeared well and gained weight similar to mice from other experimental groups throughout the study. There was also an excellent correlation between BUN and 2KW/BW which would not be expected if the rise of BUN was due to a functional cause. Nevertheless, there was a trend toward a steeper slope in the BUN vs. 2KW/BW correlation in the metformin-treated versus untreated mutant mice (Fig. S4), so a small functional effect on BUN could not be excluded. Lastly, since untreated mutant mice that reached ESRD were sacrificed earlier than the salsalate-treated mutant mice, there is a bias in our study design to underestimate the salsalate treatment effects when 2KW/BW and cystic index were used as the outcome readouts.

In conclusion, using dosages expected to yield clinically relevant serum drug levels we found that salsalate, but not metformin or canagliflozin, was highly effective in attenuating renal cystic disease in an adult-onset Pkd1 mouse model. Of note, salsalate is a pro-drug which is released as two molecules of salicylate in the small intestine. Compared to aspirin (acetyl-salicylic acid), salicylate displays weak cyclooxygenase enzyme inhibitory activities at clinical dose and is associated with minimal bleeding risk and fewer GI side-effects [59,60]. Our observations of metabolic reprogramming and inflammation in Pkd1 cystic kidneys (as compared to wild-type control) which were attenuated by salsalate treatment are consistent with the putative mechanisms of action of salicylate to directly activate AMPK [33], uncouple oxidative phosphorylation [51], and strongly inhibit NFκB [59]. The oral bioavailability of salicylate in humans is ~80%; most of the salicylate is metabolized by the liver and excreted by the kidneys. With an excellent safety profile, salsalate, at concentrations capable of activating AMPK, has been used for over five decades for the treatment of arthritis and more recently re-purposed for testing in phase 2 clinical trials of type 2 diabetes and cardiovascular disease [59,60]. These features, together with the results of our study, suggest that salsalate is a highly promising candidate for drug repurposing and clinical testing in ADPKD.

**Author contributions**

W.N.L., X.W.S., I.A.I., G.S., D.J.M.P., and Y.P. designed the study; W.N.L., X.W.S. and A.A.K. performed the study, analyzed the data, produced the figures and drafted the paper; A.B. developed the assays for and measured serum metformin levels; all the authors have read, edited and approved the final version of the manuscript.

**Declaration of Competing Interest**

Y.P. has served as consultant and received honoraria from Otsuka and Vertex Pharmaceutical. D.P. has served as consultant and received honorarium from Mironid. All other authors have nothing to disclose.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2019.08.041.

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Author/s:
Leonhard, WN; Song, X; Kanhai, AA; Iliuta, I-A; Bozovic, A; Steinberg, GR; Peters, DJM; Pei, Y

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