A Multicompartment Human Kidney Proximal Tubule-on-a-Chip Replicates Cell Polarization–Dependent Cisplatin Toxicity

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ABSTRACT

Drug-induced kidney injury is a major clinical problem and causes drug attrition in the pharmaceutical industry. To better predict drug-induced kidney injury, kidney in vitro cultures with enhanced physiologic relevance are developed. To mimic the proximal tubule, the main site of adverse drug reactions in the kidney, human-derived renal proximal tubule epithelial cells (HRPTECs) were injected in one of the channels of dual-channel Nortis chips and perfused for 7 days. Tubes of HRPTECs demonstrated expression of tight junction protein 1 (zona occludens-1), lotus lectin, and primary cilia with localization at the apical membrane, indicating an intact proximal tubule brush border. Gene expression of cisplatin efflux transporters multidrug and toxin extrusion transporter (MATE) 1 (SLC47A1) and MATE2-k (SLC47A2) and megalin endocytosis receptor increased 19.9 ± 5.0, 23.2 ± 8.4, and 106 ± 33-fold, respectively, in chip cultures compared with 2-dimensional cultures. Moreover, organic cation transporter 2 (OCT2) (SLC22A2) was localized exclusively on the basolateral membrane. When infused from the basolateral compartment, cisplatin (25 µM, 72 hours) induced toxicity, which was evident as reduced cell number and reduced barrier integrity compared with vehicle-treated chip cultures. Coexposure with the OCT2 inhibitor cimetidine (1 mM) abolished cisplatin toxicity.

In contrast, infusion of cisplatin from the apical compartment did not induce toxicity, which was in line with polarized localization of cisplatin uptake transport proteins, including OCT2. In conclusion, we developed a dual channel human kidney proximal tubule-on-a-chip with a polarized epithelium, restricting cisplatin sensitivity to the basolateral membrane and suggesting improved physiologic relevance over single-compartment models. Its implementation in drug discovery holds promise to improve future in vitro drug-induced kidney injury studies.

SIGNIFICANCE STATEMENT

Human-derived kidney proximal tubule cells retained characteristics of epithelial polarization in vitro when cultured in the kidney-on-a-chip, and the dual-channel construction allowed for drug exposure using the physiologically relevant compartment. Therefore, cell polarization–dependent cisplatin toxicity could be replicated for the first time in a kidney proximal tubule-on-a-chip. The use of this physiologically relevant model in drug discovery has potential to aid identification of safe novel drugs and contribute to reducing attrition rates due to drug-induced kidney injury.

Introduction

The proximal tubule epithelium is the tissue within the kidney that is most prone to drug-related adverse effects. Renal proximal tubule epithelial cells (RPTECs) are tasked with the active excretion of waste products, including urea and uremic toxins and reabsorption of essential molecules, including water, salts, glucose, amino acids, and proteins (Nigam et al., 2015). To this end, RPTECs express specialized transmembrane drug transporter proteins, facilitating influx from the kidney interstitium and efflux to the glomerular filtrate, which is collectively referred to as transcellular transport, clearing compounds from the internal circulation. Imbalances between influx and efflux, however, can render RPTECs vulnerable to drug-induced toxicity (Konig et al., 2013; Nigam et al., 2015; Nieskens and Sjogren, 2019).

The main adverse effect of the chemotherapeutic cisplatin is acute kidney injury, which occurs in approximately 30% of patients (Hartmann et al., 1999). Cisplatin is taken up by organic cation transporter 2 (OCT2) (SLC22A2), which is located on the basolateral membrane of RPTECs (Ciarimboli et al., 2005; Yonezawa et al., 2005; Filipski et al., 2008; Ciarimboli et al., 2010) and excreted by multidrug and toxin extrusion transporters.
transporter (MATE) 1 (SLC47A1) and to a lesser extent MATE2-k (SLC47A2), which is located on the apical membrane of RPTECs (Yonezawa et al., 2006; Nakamura et al., 2010; Li et al., 2013). The toxic potential of cisplatin correlates directly to its accumulation, which is in turn determined by the activity of cation influx and efflux transporters (Li et al., 2013). The physiologic relevance and predictive value of in vitro proximal tubule models for nephrotoxic drugs is therefore in part dependent on the ability to form distinct apical and basolateral membranes, which is referred to as polarization, which enables epithelial barrier formation and correct membrane localization of drug transporters similar to in vivo (Ito et al., 2005; Stoops and Caplan, 2014).

Kidney in vitro models are valuable tools for preclinical investigation of drug-induced toxicity but inherently suffer from partial dedifferentiation (Lash et al., 2006, 2008; Brown et al., 2008). This reduces tight junction expression and epithelial barrier function and limits replication of cell polarization-dependent drug-induced toxicity (Lash et al., 2018). By recapitulating the microenvironment of the physiologic proximal tubule, recently developed kidney-on-a-chip models, which are also known as kidney microphysiologic systems, aim to enhance proximal tubule cultures toward their in vivo phenotype (Jang et al., 2013; Weber et al., 2016; Vedula et al., 2017; Vriend et al., 2018). Epithelial polarization has been confirmed in these chip systems by tight junction formation and the presence of primary cilia (Jang et al., 2013; Jansen et al., 2015; Weber et al., 2016; Vedula et al., 2017; Vormann et al., 2018; Vriend et al., 2018) as well as proof-of-concept studies indicating that transepithelial transport of anionic and cationic organic compounds is feasible (Weber et al., 2016; Jansen et al., 2019; van der Made et al., 2019; Stahl et al., 2020). Kidney-on-a-chip models have been used to study drug-induced toxicity (Jang et al., 2013; Sakolish et al., 2018; Suter-Dick et al., 2018; Weber et al., 2018; Vormann et al., 2018; Maass et al., 2019) but crucially have not previously been applied to study the implications of epithelial polarization and localization of drug transporters for replication of membrane-dependent drug toxicity in vitro.

The aim of this study was to develop a human-derived kidney proximal tubule-on-a-chip that is capable of replicating polarization-dependent nephrotoxicity using cisplatin as model compound. Polarization of the proximal tubule epithelium was demonstrated by apical localization of tight junctions and primary cilia and basolateral localization of OCT2. Moreover, correct polarized localization and function of cisplatin toxicity can only be observed if perfused from the basolateral compartment and not when perfused from the apical compartment. Polarization is crucial for the physiologic relevance of an in vitro model, and when implemented at a fitting stage of drug discovery, this kidney-on-a-chip has the potential to aid the selection of drugs with the right safety profile, improve safety translation, and contribute to bringing safer drugs to the patients.

Materials and Methods

Cell Culture. Cryopreserved human-derived renal proximal tubule epithelial cells (HRPTECs) (batch RPT101030, male, mycoplasma negative) were purchased from Biopredic (Rennes, France). To increase homogeneity, cells of passage 2 were seeded at 10⁶ cells/ml. Cells were cultured in Dulbecco’s modified Eagle’s medium/Ham’s F12 with GlutaMAX supplement (Life Technologies, Paisley, UK) supplemented with 10 ng/ml insulin, 10 ng/ml transferrin, 10 ng/ml selenium, 24 ng/ml hydrocortisone, 10 ng/ml epithelial growth factor, 4 pg/ml tri-iodothyronine (Sigma, St. Louis), 1% fetal calf serum (Life Technologies), and 1% penicillin/streptomycin (Life Technologies), with the latter used only during seeding. For regular culture plates, the medium was refreshed every 2 to 3 days, whereas medium was continuously perfused in the microphysiologic chips at a rate of 1 μl/min.

Microphysiologic Chip Preparation. Dual-channel microphysiologic chips (dual-channel chip 001), reservoir kits (reservoir kits 001), and all other perfusion equipment was obtained from Nortis, Inc. (Woodinville, WA) and handled according to the manufacturer’s instructions. Briefly, the matrix compartment of each chip was first washed with 3 ml ethanol (99.5%; CCS Healthcare, Borlänge, Sweden) and dried by aspiration for 1.5 minutes. Next, rat tail-derived collagen 1 (7 mg/ml; Corning Life Sciences) was freshly supplemented with Dulbecco’s phosphate-buffered saline (1×), phenol red (0.004 mg/ml), HEPES (25 mM), and genipin (0.2 mM); final concentrations are listed. Finally, NaOH and filter-sterilized deionized water were added to obtain a pH of 8.0–8.5. The matrix was subsequently injected into the matrix compartment of each chip and incubated at 4°C for 1 to 2 hours, which was followed by overnight incubation at 37°C for polymerization. The two aligning fiber pins were removed from the chips the next day to form the tubular collagen lumen, with both connecting their respective prefabricated polydimethylsiloxane circuits. The circuits were perfused overnight with culture medium using the Nortis pump setup at 1 μl/min. The next day, only one channel was injected (2 × 2.5 μl) with freshly prepared collagen IV (10 μg/ml, Sigma) solution in Dulbecco’s phosphate-buffered saline (1×), to allow for coating of the lumen and incubated at 37°C and 5% v/v CO2. After 1 hour, a suspension of cryopreserved HRPTECs in culture medium was injected (2 × 2.5 μl) into the collagen IV-coated channel at a density of 10 × 10⁶ cells/ml and subsequently incubated at 37°C and 5% v/v CO2. The matrix ports were closed after 4 hours to prevent drying. Perfusion of both circuits was reinitiated after overnight incubation, enabling the remaining unattached cells to flush from the lumen. Medium was continuously perfused through both circuits in the microphysiologic chips at a rate of 1 μl/min (fluid shear stress of 0.9 dyne/cm²) for 7 days, allowing the cells to form a confluent tube-shaped monolayer.

Compound Exposure, Cell Count, Immunofluorescence Staining, and Barrier Integrity Evaluation. In preparation for compound exposure in microphysiologic chips, medium of the apical channel inflow reservoirs was supplemented with dextran 3000–Alexa Fluor 680 (0.02 mg/ml; Life Technologies). After 1 day, the outflow reservoirs of both the apical channels and basolateral channels were sampled for baseline values. Next, the medium in the apical channel or basolateral channel inflow reservoirs was replaced by medium containing cisplatin (25 μM; Sigma) with or without cimetidine (1 mM; Sigma) depending on the experimental condition. Outflow reservoirs of both channels were sampled at 24, 48, and 72 hours. After 72 hours exposure, nuclear stain was performed by disconnecting chips from the perfusion system and perfused with HBSS (1×, 37°C) supplemented with Hoechst 33342 (1:1000; Life Technologies) at 1 μl/min for 2.5 hours. Benchtop perfusions were performed at room temperature using 5-ml syringes (Henke Sass Wolff, Tuttinglen, Germany) and CMA 400 syringe pump (CMA Microdialysis, Kista, Sweden). Chips were imaged immediately at preset excitation and emission wavelengths (Supplemental Table 1) at 20× magnification using ImageXpress Micro Confocal High-Content Imaging System (Molecular Devices, San Jose). The same exposure times were used to establish a baseline in control chips and applied to treatment groups equally. After imaging, chips were treated further for immunofluorescence staining. Counting of nuclei was performed on a single representative section, and single z-slice was focused on the bottom of each tube using “find maxima” in Fiji (version 2.0.0) and a noise tolerance of 600 or 2500 for Hoechst 33342 (depending on the background). Values were expressed as percentage of vehicle, and results were plotted with GraphPad Prism (version 8.01; GraphPad Software). Fluorescence intensity of labeled dextran was evaluated at excitation and emission wavelengths of 670 and 720 nm, respectively, using the Clarioskoe microplate reader (BMG Labtech, Ortenberg, Germany). Values were expressed as fold change compared with baseline, and results were plotted with GraphPad Prism (version 8.01; GraphPad Software).

For immunofluorescence staining, chips were disconnected from the perfusion system and first perfused with HBSS (1×, 37°C) at 1 μl/min for 2.5 hours. Next, chips were perfused with formaldehyde (4%; WWR, Spanga, Sweden) at 1 μl/min for 1.5 hours, which was followed by wash buffer at 10 μl/min for 1 hour. Wash buffer consisted of bovine serum albumin (2%; Sigma) and Triton × 100 (1%; Sigma) in HBSS. Primary antibody was diluted to the indicated ratio in wash buffer (Supplemental Table 1) and subsequently manually injected (200 μl) into the chips, which was followed by overnight incubation at 4°C. Next day, chips

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were first perfused with wash buffer at 10 μl/min for 1 hour, which was followed by manual injection (200 μl) of secondary antibody (Supplemental Table 1) supplemented with Hoechst 33342 (1:1000; Life Technologies) and optionally phalloidin–Alexa Fluor 647 (1:100; Life Technologies). The chips were perfused with wash buffer at 10 μl/min for 1 hour before imaging at preset excitation and emission wavelengths (Supplemental Table 1) at 20× magnification using ImageXpress Micro Confocal High-Content Imaging System (Molecular Devices).

The intensity profile of the images was automatically rescaled to correct for background for visualization only, and image montages were made using Fiji (version 2.0.0). For 3-dimensional (3D) reconstruction of tight junctions, the spinning disk confocal function (60-μm pinhole) was enabled before images were taken every 2 μm over 120 μm in the z-direction. Reconstruction was performed using “3D Project” in Fiji with “brightest point method,” 2-μm spacing between images, and turning the resulting tube to an angle of 30°.

**Gene Expression Analysis.** For gene expression analysis in microphysiologic chips, cells were disconnected from the perfusion system and first perfused with HBSS (1×, 37°C) at 1 μl/min for 2.5 hours. Next, cells were perfused with RLT lysis buffer (Qiagen, Hilden, Germany) at 5 μl/min for 30 minutes, and the perfusate was collected. After a static incubation for 30 minutes, cells were perfused again with RLT lysis buffer (Qiagen) at 5 μl/min for 40 minutes, thus collecting the perfusate. For gene expression analysis in regular monolayer cultures, cells seeded in 12-well plates were first washed 3× with HBSS (1×) and directly lysed in RLT lysis buffer (0.5 ml/well; Qiagen). RNA was isolated using the RNeasy Mini kit (Qiagen) according to the instructions as provided by the manufacturer. High Capacity cDNA Reverse Transcription kit (Applied Biosystems, Foster City) was used to synthesize cDNA at a final concentration of 6 ng/μl according to the instructions as provided by the manufacturer. Expression levels of mRNA were determined using gene-specific primer probe sets (Supplemental Table 2) and Taqman Fast Advanced Master Mix (Applied Biosystems) using the QuantStudio Flex 7 (Applied Biosystems) at a final concentration of 1.8 ng/μl cDNA and analyzed using Quantstudio Real Time PCR Software (version 1.3; Applied Biosystems). Expression levels in the microphysiologic chips were expressed as ΔΔCt, compared with regular monolayer cultures using GAPDH as reference gene. Results were plotted with GraphPad Prism (version 8.01; GraphPad Software).

**Transcellular 4-Di-1-ASP Transport Assay.** To evaluate transcellular transport capability of cations in microphysiologic chips, cells were disconnected from the perfusion system and first perfused with HBSS (1×, 37°C) supplemented with HEPES (10 mM; Life Technologies) at 1 μl/min for 3 hours. Next, perfusion of the basolateral channel was switched to HBSS-HEPES containing 4-di-1-ASP (100 μM; Life Technologies) with or without cimetidine (1 mM; Sigma) at 2 μl/min. The apical channel was perfused with HBSS-HEPES. Accumulative perfusate samples were collected from 6 hours after initiation of perfusion for 14 hours. Finally, 100 μl of each sample was transferred to a 96-well plate and fluorescence intensity of 4-di-1-ASP was evaluated at excitation and emission wavelengths of 485 and 590 nm, respectively, and a gain of 1600 using the Clariostar microplate reader (BMG Labtech). Values were expressed as arbitrary units or blank-subtracted and normalized against uninhibited control. Results were plotted with GraphPad Prism (version 8.01; GraphPad Software).

**Lactate Dehydrogenase Evaluation.** Lactate dehydrogenase (LDH), as measure for cell-membrane integrity in regular 2-dimensional (2D) cultures, was evaluated from supernatant (10 μl) using the Pierce LDH Cytotoxicity Assay Kit (Thermo Fisher) according to the instructions as provided by the manufacturer. Reactions were run for 30 minutes, and absorbance was evaluated at 490 and 680 nm with the ClarioStar microplate reader (BMG Labtech). Values at 680 nm were subtracted from 490 nm, and results were plotted with GraphPad Prism (version 8.01; GraphPad Software).

**Cisplatin Exposure Measurements.** To estimate the concentration cisplatin cultured cells are exposed to when cisplatin is perfused from the basolateral and apical compartments, chips were prepared as described earlier while cell injection was omitted to eliminate any drug-transporter or metabolic-related effects. Basolateral exposure was evaluated by perfusing one channel with cisplatin (25 μM; Sigma) and the other with medium and collecting perfusate from the medium channels. Apical exposure was evaluated by perfusing only one channel with cisplatin (25 μM; Sigma) and collecting from the same, whereas the parallel channel was not perfused. Perfusions were performed using the Norris pump setup at 1 μl/min, and samples were taken every 12 hours for 72 hours. Samples were processed first by passing them through a Microcon 30-kDa microcentrifuge filter unit (Ultracel-30 membrane; Merck-Millipore, Burlington, MA) using centrifugation for 10 minutes at 16,000g and 4 °C, dilute (20 μl), diluted with internal standard, and derivatized with diethyliodiocarbamate (5% diethyliodiocarbamate in sodium hydroxide solution; Sigma). Samples were diluted with acetonitrile/water (1/1 v/v), and cisplatin content was evaluated by high-pressure liquid chromatography coupled to tandem mass spectrometry. Separation of derivatized cisplatin was achieved using high-pressure liquid chromatography with an Acquity BEH C18 column (Waters, Milford, MA) and acetonitrile/acetic acid (100:0.2 v/v) and ammonium formate (20 mM) mobile phase run under gradient conditions. Detection was via an API5000 mass spectrometer (Sciex, Framingham, MA) in positive TurbolonSpray mode, and results were plotted with GraphPad Prism (version 8.01; GraphPad Software).

**Results**

HRPTECs Cultured in the Kidney Proximal Tubule-on-a-Chip Were Polarized with Tight Junctions, and Primary Cilia Localized on the Apical Brush Border and OCT2 Were Strictly Localized on the Basolateral Membrane. To increase the physiologic relevance of the culture environment, HRPTECs were cultured in Nitos dual-channel microphysiologic chips. These consist of two parallel collagen IV–coated hollow channels inside a larger compound-permeable collagen I matrix that allows for independent medium perfusion through separate channel circuits (Fig. 1A). When HRPTECs were injected into a single channel and perfused with culture medium at 1 μl/min (fluid shear stress of 0.9 dyne/cm²), cells attached directly to collagen IV, and 3D tube formation was observed over the course of 7 days (Fig. 1B). The remaining empty channel was also perfused, generating a culture system with distinct apical and basolateral compartments, thus representing the tubule lumen and the kidney interstitium, respectively. To investigate cellular polarization, HRPTECs were subsequently stained with antibodies for confocal fluorescence imaging at the positions indicated in Fig. 1E. Tubes demonstrated a characteristic epithelial expression pattern of tight junction protein highlighting cell-cell interactions with focused expression at the apical membrane (Fig. 1C). A 3D reconstruction of the tight junction pattern using confocal imaging confirmed an elongated, tube-shaped tissue structure (Fig. 1D). In addition, tubes showed apical localization of lotus lectin, the presence of primary cilia protruding from the apical membrane into the tubule lumen (by staining for acetylated α tubulin), and strict basolateral localization of OCT2 (Fig. 1C), which together demonstrated an intact tubule brush border that is tight and polarized. Finally, phalloidin was used to demonstrate that F-actin is mainly located at the cell borders and the basolateral membrane, which is a typical distribution for epithelial cells (Fig. 1C).

**Gene Expression of MATE1 and MATE2-k Efflux Drug Transporters Increased in HRPTECs Cultured in the Kidney Proximal Tubule-on-a-Chip Compared with Regular 2D Cultures.** Next, gene expression of several drug transporters and proximal tubule markers in chip-cultured HRPTECs was compared with regular 2D HRPTEC cultures (Fig. 2; Table 1). Expression of MATE1 (SLC47A1) and MATE2-k (SLC47A2), which are both responsible for cisplatin efflux, increased 19.9 ± 5.0— and 23.2 ± 8.4-fold (mean ± S.D., n = 5), respectively, when HRPTECs were cultured in the chips as compared with regular 2D cultures (Fig. 2; Table 1). Cisplatin-uptake transporters OCT2 (SLC22A2) and high-affinity copper-uptake transporter 1 (SLC31A1)
were expressed at similar levels in chip-cultured and regular 2D-cultured HRPTECs (Fig. 2; Table 1). In addition, chip-cultured HRPTECs increased expression of megalin endocytosis receptor (LRP2) with $10^6$-fold (mean ± S.D., $n=5$) and expression of efflux transporter breast cancer resistance protein (ABCG2) with $4.3 \pm 0.4$-fold (mean ± S.D., $n=3$), whereas the expression of efflux transporter P-glycoprotein (ABCB1) was decreased to $0.32 \pm 0.06$-fold (mean ± S.D., $n=5$) (Fig. 2; Table 1). Interestingly, organic anion transporter 1 (SLC22A6) was expressed exclusively in chip-cultured HRPTECs ($\Delta C_t$ of $11.8 \pm 0.8$ relative to GAPDH, $n=5$, Table 1) and was not detected in regular 2D cultures. Gene expression of organic anion transporter 3 (SLC22A8) was, however, not detected in either regular 2D or chip-cultured HRPTECs (Table 1).

A Trend toward Transepithelial Transport of Fluorescent Organic Cation 4-Di-1-ASP Was Observed in the Kidney Proximal Tubule-on-a-Chip. To investigate the concerted activity of cation influx and efflux transport mechanisms in the human kidney proximal tubule-on-a-chip, transcellular transport of 4-di-1-ASP was evaluated. The fluorescent organic cation 4-di-1-ASP is reported to be a substrate for both OCT2 and MATE1/2-k drug transporters covering the cation transcellular transport axis (Biermann et al., 2006; Wittwer et al., 2013). To this end, 4-di-1-ASP (100 μM) was perfused into the basolateral compartment while buffer was perfused into the apical compartment so that fluorescence intensity of accumulative apical perfusate (14 hours) reflected compound transfer (Fig. 3A). As expected, the apical fluorescence signal was reduced compared with the
Polarization Determines Kidney-on-a-Chip Cisplatin Toxicity

Resistance protein (mediated inhibition of 4-di-1-ASP transfer was observed in every inhibitor cimetidine (1 mM, Fig. 3B). A trend toward cimetidine-evaluated by supplementing 4-di-1-ASP with competitive OCT2 showed significantly higher MATE1 (cultured in the chip compared with regular 2D cultures. Gene expression analysis of gene expression levels in human-derived kidney proximal tubule cells for multiple comparisons using the Holm-Sidak method). n = 3 (biologic replicates after experimental design); n = 4 (AQP1; due to technical error); and n = 3 (ABCG2, SLC5A2, SLC22A1, SLC22A4, SLC22A5, SLC31A1, SLCO4C1; due to inclusion after experimental design) performed with one experimental replicate with 18 comparisons in total, and **p < 0.01, ***p < 0.001, and ****p < 0.0001 (corrected for multiple comparisons using the Holm-Sidak method). AQP1, aquaporin 1; CUBN, cubilin; GGT1, gamma-glutamyltransferase 1; HPRT1, hypoxanthine phosphoribosyltransferase 1; TJP1, tight junction protein 1.

basolateral signal because it reflects HRPTEC epithelial barrier integrity (Supplemental Fig. 1). The active transport component was evaluated by supplementing 4-di-1-ASP with competitive OCT2 inhibitor cimetidine (1 mM, Fig. 3B). A trend toward cimetidine-mediated inhibition of 4-di-1-ASP transfer was observed in every individual experiment [66%, 30%, and 17%; with an average of 38% ± 25% (mean ± S.D., n = 3, Fig. 3B)] but was not statistically significant (paired Student’s t test, P = 0.18).

Cisplatin-Induced Toxicity in the Kidney Proximal Tubule-on-a-Chip Exclusively when Exposed to the Basolateral Membrane, and Toxicity Was Abolished by the OCT2 Inhibitor Cimetidine. When HRPTECs were cultured in regular 96-well plates, cisplatin (12.5–50 µM) induced toxicity, as demonstrated by a time- and concentration-dependent increase of LDH in the supernatant (Supplemental Fig. 2). As expected, toxicity was attenuated by coincubation with cimetidine (1 mM), indicating its dependence on OCT2 influx activity (Supplemental Fig. 2). In contrast to regular 2D cultures, the dual-channel chip layout allows for selective exposure of the proximal tubule apical or basolateral membrane, which is relevant when mimicking physiologic exposure to therapeutics in vitro. When cisplatin (25 µM) was perfused for 72 hours from the basolateral channel, mimicking exposure through the clinically relevant interstitium and internal circulation, the tight junction pattern on the bottom slice of the tube was partly disrupted (Fig. 4A), and cell count was reduced to 41.2% ± 6.6% (mean ± S.D., n = 5) compared with vehicle control (Fig. 4B). This resulted in a concomitant 8.7 ± 6.3-fold (mean ± S.D., n = 4) increase in paracellular diffusion of dextran 3000 across the epithelium from the apical compartment into the basolateral compartment (Fig. 4C). As expected, coincubation with the OCT2 inhibitor cimetidine (1 mM) abolished all toxic effects, demonstrating that cisplatin-induced toxicity is dependent on OCT2 influx activity (Fig. 4, A–C). The actual concentration of cisplatin reaching the cells through basolateral exposure was measured by omitting cell injections, perfusing the basolateral channel with cisplatin (25 µM), and collecting the perfusate from the apical channel (Fig. 5A). The resulting concentration of cisplatin was 3.4 ± 0.05 µM (at 24 hours, mean ± S.D., n = 3, Fig. 5C), which is similar to clinical therapeutic total plasma concentrations reported in literature, although this is dependent on the dosing regimen applied (Himmelstein et al., 1981; Ikeda et al., 1998; Petrillo et al., 2019). In contrast, when cisplatin was perfused from the apical channel mimicking exposure through the tubule lumen, toxicity was significantly lower in dual-channel chip compared with regular 2D cultures. Gene expression levels of drug transporters, endocytosis receptors, tight junction protein, and proximal tubule markers in regular 2D cultures and Nortis chip cultures were compared.

Table 1

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<tr>
<th>Gene</th>
<th>Ct 2D HRPTECs</th>
<th>Nortis HRPTECs</th>
<th>−ΔCt 2D HRPTECs</th>
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<td>19.7 ± 1.5</td>
<td>0.0 ± 1.1</td>
<td>0.7 ± 1.0</td>
</tr>
</tbody>
</table>

AQP1, aquaporin 1; CUBN, cubilin; GGT1, gamma-glutamyltransferase 1; HPRT1, hypoxanthine phosphoribosyltransferase 1; TJP1, tight junction protein 1.

*Mean ± S.D.

Not detected.
toxicity was not observed (Fig. 4, A–C). The concentration of cisplatin through apical exposure was evaluated by perfusing the apical channel with cisplatin (25 μM) and collecting perfusate from the apical channel (Fig. 5B), resulting in an actual exposure of 20.2 ± 1.3 μM (at 24 hours, mean ± S.D., n = 3, Fig. 5C). So, even though apical cisplatin exposure was more than 5-fold higher than the basolateral exposure, it did not induce toxicity. These results confirm that the transporters mediating cisplatin toxicity were uniquely expressed on the basolateral membrane, supporting the epithelial polarization of this proximal tubule-on-a-chip.

Discussion

Achieving a safety profile for candidate drugs lacking renal adverse effects prior to the clinical phase of development requires predictive and physiologically relevant in vitro proximal tubule epithelial models.

Fig. 3. Transepithelial transfer of fluorescent organic cation 4-di-1-ASP in the proximal tubule-on-a-chip. (A) Fluorescent cation 4-di-1-ASP (100 μM) was perfused at 2 μl/min into the basolateral compartment of the proximal tubule-on-a-chip with or without competitive OCT2 inhibitor cimetidine (1 mM), whereas HBSS-HEPES buffer was perfused into the apical compartment. (B) Fluorescence intensity of 4-di-1-ASP in accumulative apical perfusate (8612 ± 5208) was not significantly reduced (P = 0.18) upon cocultivation with cimetidine (1 mM) [4956 ± 3616, mean ± S.D., fluorescence intensity of the apical perfusate without cimetidine compared with cimetidine by paired Student’s t test, n = 3 (with the number of biologic replicates set in advance), performed with one experimental replicate, one comparison in total].

Fig. 4. Membrane-dependent cisplatin-induced toxicity in the human-derived kidney proximal tubule-on-a-chip. Cisplatin (25 μM, 72 hours) induced toxicity when perfused from the basolateral compartment but not the apical compartment and is abolished by coperfusion with OCT2 inhibitor cimetidine (1 mM) as evaluated by (A) disruption of tight junctions at the bottom and middle planes of chip-cultured HRPTECs, (B) reduced nuclei count, and (C) reduced epithelial barrier integrity according to the figure in (D). Analysis of (A) (n = 5 (vehicle; 25 μM basolateral; 25 μM apical) and n = 4 (25 μM basolateral, cimetidine) with all biologic replicates set in advance, performed with one experimental replicate, and representative images were selected, original magnification, 20×). (B) Mean ± S.D., Hoechst count in the “bottom” images compared with vehicle by one-way ANOVA with Dunnett’s multiple comparisons test, n = 5 (vehicle; 25 μM basolateral; 25 μM apical) and n = 4 (25 μM basolateral, cimetidine) with all biologic replicates set in advance, performed with one experimental replicate, with three comparisons in total; ****P < 0.0001 (multiplicity-adjusted P values). (C) Mean ± S.D., dextran 3000–Alexa Fluor 680 fluorescence intensity sampled from the basolateral channel perfusate compared with time-matched vehicle by two-way ANOVA with Dunnett’s multiple comparisons test, n = 4 (vehicle; 25 μM basolateral; 25 μM apical) and n = 3 (25 μM basolateral, cimetidine) with one biologic replicate excluded because of technical error, performed with one experimental replicate, with 12 comparisons in total; ****P < 0.0001 (multiplicity-adjusted P values). Cim, supplemented with cimetidine (1 mM).
Here, we developed a human-derived kidney proximal tubule-on-a-chip that demonstrates epithelial polarization with intact tight junctions and primary cilia on the apical brush border membrane and strict localization of OCT2 on the basolateral membrane. Exclusive OCT2-dependent sensitivity to the nephrotoxic drug cisplatin was shown when the tube was exposed via the basolateral compartment, whereas toxicity was not observed when exposed to more than 5-fold higher concentration via the apical compartment. Therefore, the importance of polarized localization of cation drug uptake transporters to more accurately replicate drug-induced kidney toxicity was demonstrated for the first time in a chip-based proximal tubule model.

Influx of cisplatin into renal proximal tubule epithelial cells and the nephrotoxicity this results in have been shown to be mediated by OCT2 (Ciarimboli et al., 2005, 2010; Yonezawa et al., 2005, 2006) and high-affinity copper-transport protein 1 (SLC31A1) (Pabla et al., 2009). OCT2 is a facilitated diffusion transporter driving the influx of cationic compounds across the basolateral membrane using the inward-directed negative membrane potential (Okuda et al., 1999b; Budiman et al., 2000). The current study showed attenuation of cisplatin toxicity by cimetidine in both regular 2D HRPTEC cultures and chip-cultured HRPTECs, demonstrating OCT2 activity is the main contributing factor to cisplatin toxicity and confirming previous results in vitro (Yonezawa et al., 2005, 2006; Sato et al., 2008) and in vivo (Ciarimboli et al., 2010; Katsuda et al., 2010). It is worth noticing, however, that the cimetidine concentration used is higher than a typical pharmacological concentration, as the $C_{\text{max}}$ is reported to be 16 $\mu$M (Schmidt et al., 1998).

Moreover, the current study elegantly demonstrates that when HRPTEC cultures are correctly polarized, form tight junctions, and concentrate OCT2 strictly on the basolateral membrane, cisplatin only induces toxicity when the basolateral membrane is exposed and at clinically relevant total-plasma concentrations (Himmelstein et al., 1981; Ikeda et al., 1998; Petritello et al., 2019). Although the toxicity data presented in the current manuscript support OCT2-mediated cisplatin transport, future intracellular accumulation and transcellular transport studies using cisplatin as substrate may provide further insights into actual handling of this nephrotoxicant by the proximal tubule-on-a-chip. Cisplatin toxicity has previously been evaluated in a few chip-based models. Jang et al. (2013) showed that cisplatin exposed basolaterally induced toxicity in their 2D chip-based model, but the concentration used (100 $\mu$M) extends beyond the therapeutic concentration, and cellular cisplatin exposure was not evaluated (Himmelstein et al., 1981; Jang et al., 2013). In other studies, both compartments were used simultaneously, which prohibited any evaluation of the impact of epithelial polarization on cisplatin sensitivity (Suter-Dick et al., 2018; Vormann et al., 2018). In contrast to the current study, single-channel Nortis proximal tubule-on-a-chips were used to show that apical cisplatin exposure (1 $\mu$M) increased concentrations of Kidney Injury Molecule-1 in the supernatant, whereas LDH levels (reflecting cytotoxicity) were not affected (Sakolish et al., 2018; Maass et al., 2019). Although we can only speculate about the reason for the discrepancy compared with the current study, it may be explained by differences in exposure times and endpoints used, assays, cell source, and media components. An earlier study employing Lilly Laboratories Cell-Porcin Kidney 1 cells cultured on porous membrane inserts showed that basolateral treatment with cisplatin (300 $\mu$M) induced a higher degree of cytotoxicity compared with apical treatment, thus confirming the current findings, although the applied concentration was not regarded as clinically relevant, and localization of OCT2 was not investigated (Okuda et al., 1999a).

In contrast to the cisplatin parent compound, proximal tubule uptake of its metabolites is suggested to be mediated through alternative mechanisms (Townsend et al., 2009). Extrarenal glutathione conjugation generates water-soluble molecules capable of being filtered by the glomerulus, therefore mainly exposing the apical membrane of proximal tubule cells. Interestingly, inhibition of $\gamma$-glutamyl transpeptidase or cysteine s-conjugate $\beta$-lyase, which are vital enzymes in the conjugation pathway, or shifting the equilibrium toward oxidized glutathione can both reduce proximal tubule toxicity in mice, indicating that cisplatin-glutathione contributes to nephrotoxicity even when exposure is predominantly apical (Townsend and Hanigan, 2002; Jenderny et al., 2010). Extrarenal metabolite formation is not incorporated in the proximal tubule-on-a-chip described in the current study. However, it could be extended with a liver-on-a-chip compartment, making it suitable to study the contribution of metabolites to drug-induced nephrotoxicity, as shown earlier for ifosfamide and aristolochic acid (Choucha-Snouber et al., 2013; Chang et al., 2017).

Cisplatin efflux from the proximal tubule is mainly mediated by MATE1 (Yonezawa et al., 2006; Nakamura et al., 2010; Li et al., 2013).
The current study showed significantly increased gene expression of MATE1 and MATE2-k in chip-cultured HRPTECs compared with regular 2D cultures, which suggests a more physiologically relevant phenotype. However, immunofluorescence did not succeed in detecting specific staining of MATE1 or MATE2-k in the proximal tubule-on-a-chip (unpublished data), which was most likely because of poor specificity of the primary antibodies, thus requiring extended optimization. Therefore, localization of these transporters remains to be elucidated in future studies. Treating HRPTECs with bardoxolone methyl, an inducer of signaling by the nuclear factor erythroid 2–related factor 2 pathway, increases MATE1 gene expression and reduces cisplatin sensitivity (Atilano-Roque et al., 2016). Interestingly, nuclear factor erythroid 2–related factor 2 has also been shown to regulate MATE2-K in response to fluid flow in proximal tubule cells (Fukuda et al., 2017), sparking speculation that Nrf signaling might explain upregulation of MATE gene expression observed in the current study.

Transepithelial transport activity is indicative of a polarized epithelial phenotype, as was previously shown for organic anions (Weber et al., 2016; Jansen et al., 2019; van der Made et al., 2019). The current study did not demonstrate a statistically significant inhibition of 4-di-1-ASP transcellular transport using cimetidine and therefore concerted activity of OCT2 and MATE1/2-k. However, the observed trend in inhibition is in line with the findings by Stahl et al. (2020), who demonstrated transepithelial transport of cation drug metformin in a proximal tubule-on-a-chip using the same chip platform, although a different tissue donor was employed. Differences in methods might explain this discrepancy because radioactively labeled metformin was used as cation substrate combined with inhibitor imipramine, and this substrate and inhibitor combination reflects a different inhibition potential. In addition, increased sensitivity of detection due to radioactive labeling compared with the fluorescent methods used in the current study potentially allows detection of inhibition across a wider range of substrate concentration (Stahl et al., 2020). In turn, variations in substrate transport might reflect cell-to-chip differences in barrier integrity, increasing the diffusion component of substrate transport across the epithelial tube and highlighting that advanced chip-based culture models remain technically challenging.

MATE-mediated transport is driven by a concentration gradient of H⁺ over the brush border membrane (Tsuda et al., 2007; Sato et al., 2008; König et al., 2011). It has been shown that evaluation of cisplatin transport by MATE1 in vitro requires artificially acidifying the cytoplasm and reversing the naturally occurring H⁺ gradient and direction of MATE1-mediated transport (Nakamura et al., 2010). Therefore, transepithelial transport activity and MATE-mediated efflux of cisplatin in chip-cultured HRPTECs may be investigated in future studies by increasing the acidity of the luminal chip channel, providing a more physiologically relevant driving force. In addition, genetic knockdown or knockout approaches would be a suitable future strategy to provide more direct evidence of organic cation transport– and MATE-mediated transport function in the currently developed proximal tubule-on-a-chip.

The current study found significantly increased gene expression of LRP2 coding for the megalin endocytosis receptor in chip-cultured HRPCECs compared with regular 2D HRPTEC cultures. Megalin is expressed on the apical brush border of proximal tubule cells and is responsible for reabsorption of low-molecular-weight proteins and albumin from the glomerular filtrate (Nielsen et al., 2016). Increased uptake of megalin substrate bovine serum albumin has been described in response to changes in fluid flow (Ferrell et al., 2012, 2018; Jiang et al., 2013; Raghavan et al., 2014). More recently, endocytosis activity has also been implicated in the uptake of antisense oligonucleotides into the proximal tubule epithelium (Janssen et al., 2019). Therefore, future studies may investigate whether the proximal tubule model developed here is suited to study antisense oligonucleotide–induced renal adverse effects or the effects of fluid flow on receptor-mediated endocytosis.

In conclusion, the current study presented the first kidney proximal tubule-on-a-chip in which a nephrotoxic drug has been exposed through both the clinically relevant and nonrelevant compartments to demonstrate the contribution of epithelial polarization and membrane localization of drug influx mechanisms in achieving drug sensitivity at clinically relevant concentrations. Therefore, the developed in vitro model displays increased physiologic relevance compared with single-compartment models and could be applied in the future to improve preclinical prediction of drug-induced kidney toxicity and eventually reduce kidney-related adverse effects of candidate drugs in the clinic.

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Authorship Contributions

Participated in research design: Nieskens, Kelly, Sjögren.
Conducted experiments: Nieskens.
Performed data analysis: Nieskens, Sjögren.
Wrote or contributed to the writing of the manuscript: Nieskens, Persson, Kelly, Sjögren.

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