Human small intestinal organotypic culture model for drug permeation, inflammation, and toxicity assays

Jan Markus¹ · Tim Landry² · Zachary Stevens² · Hailey Scott² · Pierre Llanos² · Michelle Debatis² · Alexander Armento² · Mitchell Klausner² · Seyoum Ayehunie²

Received: 3 July 2020 / Accepted: 23 October 2020 / Published online: 25 November 2020 / Editor: Tetsuji Okamoto
© The Society for In Vitro Biology 2020

Abstract
The gastrointestinal tract (GIT), in particular, the small intestine, plays a significant role in food digestion, fluid and electrolyte transport, drug absorption and metabolism, and nutrient uptake. As the longest portion of the GIT, the small intestine also plays a vital role in protecting the host against pathogenic or opportunistic microbial invasion. However, establishing polarized intestinal tissue models in vitro that reflect the architecture and physiology of the gut has been a challenge for decades and the lack of translational models that predict human responses has impeded research in the drug absorption, metabolism, and drug-induced gastrointestinal toxicity space. Often, animals fail to recapitulate human physiology and do not predict human outcomes. Also, certain human pathogens are species specific and do not infect other hosts. Concerns such as variability of results, a low throughput format, and ethical considerations further complicate the use of animals for predicting the safety and efficacy xenobiotics in humans. These limitations necessitate the development of in vitro 3D human intestinal tissue models that recapitulate in vivo–like microenvironment and provide more physiologically relevant cellular responses so that they can better predict the safety and efficacy of pharmaceuticals and toxicants. Physiological relevance has been made in the development of in vitro intestinal models (organoids and 3D-organotypic tissues) using either inducible pluripotent or adult stem cells. Among the models, the MatTek’s intestinal tissue model (EpiIntestinal™ Ashland, MA) has been used extensively by the pharmaceutical industry to study drug permeation, metabolism, drug-induced GI toxicity, pathogen infections, inflammation, wound healing, and as a predictive model for a clinical adverse outcome (diarrhea) to pharmaceutical drugs. In this paper, our review will focus on the potential of in vitro small intestinal tissues as preclinical research tool and as alternative to the use of animals.

Keywords Intestinal tissue model · Drug absorption · Metabolism · Drug-induced toxicity · Nanotoxicity · Inflammation · Wound healing

Introduction
The gastrointestinal tract (GIT) represents one of the largest body surfaces. It is exposed to food, fluid, medicaments, and toxicants and is an important gateway tissue that plays a crucial role in drug absorption and metabolism, food digestion, and nutrient uptake. Epithelial cells of the small intestine are organized into structures called crypts of Lieberkühn and finger-like protrusions called villi (van der Flier and Clevers 2009). The crypts harbor Paneth cells (Sato et al. 2009, Gassler 2017) and LGR5+ (leucine-rich-repeat-containing G protein-coupled receptor 5, also known as Gpr49) stem cells (Barker et al. 2007), and the LGR5+ stem cells give rise to transiently amplifying cells of the villi (Mahe et al. 2017). The villi consist of absorptive enterocytes and mucus-producing goblet cells (Bland et al. 1995). Sawant-Basak et al. 2018, hormone producing enteroendocrine cells (Gribble and Reimann 2019), microbial antigen shuttling M-cells (Corr et al. 2008), and chemosensor tuft cells (Ting and von Moltke 2019). The GIT is also one of the major entryways for many human pathogens and thereby plays an important role in microbial recognition and antimicrobial defense. In
fact, dysfunction of the barrier of the intestinal epithelium is a hallmark of intestinal inflammatory diseases (König et al. 2016). This functional diversity is backed by the complex segmentation, structure, and cellular phenotypes that are present in the gastrointestinal tract.

Historically, a variety of animal models have been used for the preclinical safety evaluation of drugs, intestinal injury, infection, and inflammation as summarized by others (Waterston et al. 2002; Lin and Hackam 2011; Jiminez et al. 2015). Currently, two animal species such as a rodent and a non-rodent (e.g., rats and dogs) are used for short-term (up to 1-mo duration) preclinical safety evaluation of biotechnology-derived pharmaceuticals. If the toxicity profile in the two species is comparable in the short term, the use of only one species for subsequent long-term toxicity studies can be acceptable following regulatory guidelines (https://www.fda.gov/media/72028/download; https://www.ema.europa.eu/en/ich-s6-r1-preclinical-safety-evaluation-biotechnology-derived-pharmaceuticals). Rodents are widely used to study human diseases because of their relatively low maintenance requirements, their rapid reproduction rates, and the availability of resources such as antibodies (Gonzalez et al. 2016). However, rodent models frequently fail to fully mimic clinical signs of human diseases and inflammatory responses (Pizarro et al. 2011). Due to similarities in the anatomical structure of the gastrointestinal tract to that of the human intestine, pigs have been used for intestinal injury, inflammation, inflammatory bowel disease (IBD) modeling, and drug-induced gastrointestinal toxicity studies (Walters et al. 2012). Even though pigs and non-human primates provide valuable data on intestinal inflammation and disease conditions, their use for routine intestinal studies is limited due to the high cost associated with their care, potential hazards of carrying highly virulent zoonotic agents, and ethical considerations (Ideland 2009; Coors et al. 2010). Additionally, animal models are low throughput and they do not adequately recapitulate human physiology (Mathur et al. 2017). Therefore, pharmaceutical and academic researchers recognize the need for 3D human intestinal tissue models for preclinical studies since such models are less time-consuming, more cost-effective, and adaptable to high throughput screening (HTS). As reviewed below, these models can be physiologically relevant predictors of human responses including drug absorption and metabolism, drug-induced GI toxicity, inflammation, disease, and intestinal restitution.

Structure and Cellular Phenotypes of Small Intestinal Epithelium The small intestine constitutes a complex organ system due to its rapid self-renewal time (5–7 d), cellular composition, numerous functions, and the unique dynamic nature of the villi. Functionally, the small intestine is important since ~90% of the absorption in the digestive tract occurs in this organ (Balimane and Chong 2005). Anatomically, it is divided into three segments, the duodenum, jejunum, and ileum. It is the longest part of the alimentary canal, approximately 3 m (10 ft) in length, and has a smaller diameter (~2.5 cm) compared with the large intestine (https://opentextbc.ca/anatomyandphysiology/chapter/23-5-the-small-and-large-intestines/). The epithelium of the small intestine differentiates to form finger-like structures on its apical surface to form the villi (Mahe et al. 2017) which are ~0.5 μm tall. Each villus has hair-like projections called microvilli that pack together to form the brush border. The villi and microvilli increase the intestinal surface area to enhance absorption by 30–600-fold (Kiela and Ghishan 2016), and they play a significant role in nutrient, fluid, drug uptake, solute transport, and host defense (Crawley et al. 2014). At least 22 enzymes and 19 drug transporters are localized in the brush border of the small intestine (Holmes and Lobley 1988). In addition to the villi, the small intestine is organized into folded structures, Kerckring folds, that triple the surface area for rapid intestinal absorption. These folds also slow down the flow of chyme in the gut (Igam et al. 2019).

Within the small intestine, different cell types co-exist in a 3-dimensional space (summarized in Table 1) in which they exchange biochemical and mechanical cues (cellular “cross-talk”) that help determine tissue properties such as tissue differentiation and barrier function. The ability of tissue models to mimic such spatial organization and recapitulate the aforementioned interactions makes them more physiological and imparts functionality resembling the in vivo counterparts.

Cell Line–Based Models Caco-2 Cells: Since the 1970s, the Caco-2 cell–based assays are considered as a model for enterocytes (Fogh et al. 1977; Grasset et al. 1984; Roussel et al. 1985) and have been used as the gold standard for drug absorption studies, even though the cells originated from a colon carcinoma. Caco-2 cells have been used alone, or in combination with other cell types such as methotrexate-treated HT29 colon adenocarcinoma cells, as an in vitro model of the small intestine to investigate drug absorption, inflammation, nutrient uptake, and toxicity in the gut (Walter et al. 1996; Hilgendorf et al. 2000, and Kleiveland 2015)) and have been used for drug absorption, distribution, metabolism, and excretion (ADME) studies (Sun et al. 2008). However, the Caco-2 cells form a non-physiological barrier due to paracellular junctions that are much tighter and less permeable, rendering the Caco-2 cultures more similar to the colon than to the human small intestine (Srinivasan et al. 2015). Depending on the laboratory, the clonal type, and passage differences the transepithelial electrical resistance (TEER) values for the Caco-2 cultures varies from 300 to 2400 Ω cm², compared with TEER values of 12–120 Ω cm² for human small intestine tissue (Briske-Anderson et al. 1997; Gupta et al. 2013; Takenaka et al. 2014) and 100 Ω cm² reported for intestinal explants (Artursson et al. 1993). These
Table 1. Cellular phenotypes of the small intestine epithelium

| Cellular phenotypes | Function | Marker/stain | Reference |
|---------------------|----------|--------------|-----------|
| Enterocytes         | Most common cell type in the surface epithelium, responsible for digestion and absorption of nutrients, forms the intestinal barrier. | Villin, alkaline phosphatase | Sawant-Basak et al. (2018) |
| Goblet cells        | Secrete mucus which entraps bacteria and prevents their translocation into the intestinal epithelium. | Periodic acid–Schiff (PAS) and MUC-2 | Sawant-Basak et al. (2018) |
| Paneth cells        | Contribute to crypt morphogenesis and intestinal homeostasis, the intestinal microbiome (by secreting antimicrobial peptides such as defensins), and crypt fission. Associated with intestinal diseases including ileal Crohn’s disease. Originate from intestinal stem cells. Found at the bottom of small intestine crypts. | Lysozyme | Sato et al. (2009); Gassler (2017) |
| Enteroendocrine cells | Produce a range of hormones for chemo-sensing that have key roles in food absorption, insulin secretion, and appetite. Scattered along the length of the intestinal epithelium. | Anti-synaptophysin | Gribble and Reimann (2019) |
| Tuft cells          | Play chemo-sensor role, communicate with neurons, police entry of parasites. Help eliminate gut pathogens by releasing interleukin-25 (IL-25), which stimulates mucus-producing goblet cells, recruits immune cells, and leads to muscle contractions. | DCLK1 | Ting and von Moltke (2019) |
| M-cells             | Highly specialized to take up intestinal microbial antigens and deliver them to gut-associated lymphoid tissue (GALT) for efficient mucosal and systemic immune responses. Subset of intestinal epithelial cells with reduced brush border and lack of enzymatic activity. | Lack of alkaline phosphatase staining; Transmission electron microscopic observation of apical epithelial cells that lack brush border | Cott et al. (2008) |
| Lgr5(+) stem cells  | Proliferative stem cells located at base of intestinal crypts, which give rise to TA cells and Paneth cells | Rabbit/mouse Anti-LGR5 antibody; Lgr5 expression reporters (Lgr5-GFP)/lineage tracing | Barker et al. (2007); Sato et al. (2009); Dame et al. (2018) |
| Quiescent stem cells | Relatively quiescent intestinal stem cells capable of crypt repopulation upon injury | Lrig1 antibody/lineage tracing | Powell et al. (2012) |

Elevated barrier properties are due to the fact that the Caco-2 lines originated from the large intestine and also have an average tight junction pore radius of 3.7 Å, compared with 8–13 Å for native human small intestine (Tavelin et al. 2003). Caco-2 cell–based assays have other limitations such as (a) weak expression of important intestinal metabolic enzymes such as cytochrome P450 (CYP) 3A (Eric Le Ferrec et al. 2001), (b) lack of the crypt-villus axis which is important for fluid and ion transport, and (c) the absence of mucus-producing cells (Huang and Adams 2003). Another drawback of Caco-2 cell–based assays is that the high passages used in many laboratories induce variable expression levels of differentiation markers and transporters (Briske-Anderson et al. 1997; Behrens et al. 2004). The heterogeneity of the cell line, variability of the clones used by different researchers, and the passage numbers make it difficult to generate reproducible results among research laboratories.

To overcome some of the limitations of Caco-2 cultures, there have been numerous attempts to clone and re-clone Caco-2 cells to select clones with increased and enhanced drug transport rates (Woodcock et al. 1991). Even though Caco-2 cells have been used as a model for pharmacological and toxicological studies, the expression of specific transporters and ion channel genes often differs from that of the human small intestine which complicates the interpretation of results (Anderle et al. 2003). Although Caco-2 is widely used for ADME studies in many labs, the model is composed only of enterocyte-like cells and lacks additional functional cells types. To compensate, a mixed culture of Caco-2 and methotrexate-treated HT29 colon adenocarcinoma cells, which are known to produce mucus, has been developed to mimic an enterocyte and goblet cell co-culture system. To further increase functional complexity of Caco-2 cultures, researchers have also used triple co-cultures of Caco-2 cells, HT29 cells, and the Raji B cell line (to mimic M-cells) for pharmacokinetic studies (Araújo and Sarmento 2013; Lozoya-Agullo et al. 2017). However, these improvements increase the variability of results obtained by different laboratories.

**Primary Human Three-Dimensional (3D) Tissue Models** An in vitro small intestinal 3D tissue model needs to closely resemble the human intestinal epithelium structurally and...
phenotypically by comprising the different cell types of the intestinal epithelium. The model should show architectural similarity, physiological properties, and the functions of the human counterpart. In addition, the tissue model should be able to be cultured for long periods (Costa and Ahluwalia 2019). The most common primary human 3D intestinal tissue models include explants, organoids, and organotypic intestinal tissues as discussed in the following sections.

Intestinal Explants: The use of porcine intestinal tissue as a predictive model for human intestinal absorption has been demonstrated by the Netherlands Organization for Applied Scientific research (TNO). Recently, TNO has also utilized ex vivo human intestinal tissue in a newly developed InTESTime™, two-compartment disposabele device for drug absorption studies (Stevens et al. 2019). These models have been utilized to rank order compound permeability and to study metabolism at TNO. Even though explant tissues from multiple donors reflect the variability of the human population, the scarcity of normal human tissue and their short survival time ex vivo make their use infeasible for meeting the needs of modern-day drug development programs, which need to screen large numbers of compounds for safety and efficacy. Also, donor-to-donor variability makes the results from explant studies difficult to utilize in a predictive manner.

Small Intestinal Organoid Models: In recent years, progress has been made in the development of models generated from normal primary cells isolated from small intestine tissues which are more relevant for modeling the complex biological processes of the native tissue. The key innovation was the development of methods for cultivating mouse and human intestinal organoids from stem cells isolated from intestinal crypts (Sato et al. 2009; Spence et al. 2011). The importance of Wnt signaling was identified and a cocktail of growth factors was developed to allow the maintenance and expansion of Lgr5+ stem cells and further formation of enteroids (intestinal organoids). Also, analogous intestinal organoids were developed from embryonic stem cells or induced pluripotent stem cells (Ogaki et al. 2015). Both types of organoids contain the major cell types found in the small intestine, including enterocytes, Paneth cells, goblet cells, and enteroendocrine cells. These organoids recapitulate many of the normal processes of the intestinal mucosa and allow study of phenomena such as intestinal toxicity, immune reactions, and interactions with microbiota (Leslie et al. 2015; Lu et al. 2017; Bar-Ephraim et al. 2020). Organoids can also be generated from patients with various pathological conditions, such as inflammatory bowel disease or from individuals with various genetic backgrounds (Dekkers et al. 2013; VanDussen et al. 2015). Organoids from individuals with cells of varying genetic make-up opens a plethora of possibilities for designing of idiosyncratic therapies and personalized medicine (reviewed in Park et al. 2018; Lehe et al. 2019). Nonetheless, there are disadvantages associated with the physical organization of organoids. The villi face inward which means that direct access of test compounds to the apical surface of the enterocytes to mimic luminal exposure is not possible. Likewise, apical-to-basolateral drug permeation studies cannot be performed or depend on microinjecting the test articles into the organoid. The need for microinjection can be avoided by the use of recently developed “apical-out” enteroids following manipulating extracellular membrane (ECM) components in the culture system which resulted in enteroid polarity reversal (Co et al. 2019). Such apical-out organoids may allow to examine (a) drug–drug interactions following co-administration of drugs, (b) drug metabolism on the exposure site (apical surface), and (c) pathogen exposure. However, this polarity reversal was achieved in a suspension cell culture system and it is not clear if these organoids will form a continuous epithelial structure in Transwell inserts. Others have also addressed this limitation by converting colon spheroids 2D cultures into Transwell inserts. Following dissociation of 2D systems into single cells, the cells were seeded onto Transwell® inserts. However, these cultures did not form villi or similar secondary structures and their TEER was relatively high (~400 Ω cm²) (VanDussen et al. 2015) which is more similar to the colon epithelium than the small intestine.

A group from the Wyss Institute, Harvard University, has developed human small intestine-on-a-chip tissue containing epithelial cells that were isolated from intestinal duodenal biopsies. The primary epithelial cells were expanded as 3D organoids, dissociated, and cultured on a porous membrane within a microfluidic device with human intestinal microvascular endothelial cells (Kasendra et al. 2018). The intestinal tissue model was shown to form villi-like projections lined by polarized epithelial cells that undergo multi-lineage differentiation similar to organoids. Positive aspects of the system include an open lumen apical surface, an interface with the endothelium, and media flow. Transcriptomic analysis also indicated that the intestine-on-a-chip more closely mimics the human duodenum when compared with the duodenal organoids. However, the model is not currently commercially available to the broader scientific community.

3D Organotypic Tissue Models: While organoids exhibit an inward growth of the luminal surface, 3D organotypic tissue models have an open luminal surface which makes them ideal for topical application of test compounds mimicking in vivo oral exposure. Organotypic models are able to recreate the architectural features and physiology of native human intestinal tissue and hence are more relevant than Caco-2 or animal models. Besides the ethical advantages of organotypic models over the live animals, these tissues have better biological relevance in terms of translational ability of human responses and their potential to simulate specific conditions such as drug absorption, metabolism, pathogen infection, and inflammation. These organotypic tissue models allow study of the molecular aspects of each process and help
“deconstructing” biological processes to study them with variable levels of complexity. For example, one can study the effect of a substance on isolated intestinal mucosa, intestinal mucosa together with immune cells, or combine the intestine with a liver-on-a-chip model and observe what happens once the test compound undergoes metabolic changes. The current in vitro intestinal tissue models utilize either cell lines or primary intestinal cells.

Human Primary Cell–Based Organotypic Small Intestinal Tissue Models: The development of a human primary cell–based tissue model to accurately predict drug safety and efficacy remains a major challenge for the pharmaceutical industry (Li 2005). However, the availability of MatTek’s small intestinal tissue model (EpiIntestinal™, MatTek Corporation, Ashland, MA) is a significant development in studying ADME of pharmacological drugs (Ayehunie et al. 2018). Two types of EpiIntestinal models have been developed by MatTek.

Organotypic epithelial tissue models are produced by seeding small intestinal epithelial cells onto cell culture inserts (MatTek Corp.) and culturing at air liquid interface (ALI) for up to 14 d to form a partial-thickness intestinal tissue model. During this culture period, the cultures stratify, differentiate, and form a distinct apical-basolateral polarity. The polarized organotypic small intestinal partial thickness tissues form “villi-like” structures and express the efflux transporters such as P-glycoprotein (P-gp) breast cancer resistance protein (BCRP) (Ayehunie et al. 2018).

A more complex organotypic tissue model is the full-thickness intestinal tissue which comprises intestinal epithelial cells and intestinal fibroblasts. To reconstruct the full-thickness tissue, a mixture of primary epithelial cells and fibroblasts is seeded onto the microporous membrane of tissue culture inserts under submersed and then at ALI conditions for 14 d. In this model, the epithelial cells and fibroblasts self-assembled in the correct orientation and form a distinct and polarized tissue structure with an apical epithelial architecture on top of a fibroblast substrate (Fig. 1). Unlike the partial-thickness intestinal models, these cultures show deposition of extra-cellular matrix (ECM) proteins of collagen IV and fibronectin that emanated from the epithelial-fibroblast co-culture system (Peters et al. 2019).

The structure of EpiIntestinal tissue shows a well polarized geometry and the tissue forms villi (Fig. 1), microvilli, brush borders (Fig. 2), and tight junctions that mimic the in vivo counterpart. These tissues can be cultured for up to 42 d at ALI and can have utility for repeat dose applications (Peters et al. 2019). These organotypic small intestinal tissues can be reconstructed in 24 single-well cell culture inserts or in a 96-well plate format for high throughput applications. Even though, the standard EpiIntestinal model is reconstructed using epithelial cells from the ileum section of the gut, organotypic tissues from the duodenum and jejunum segments of the small intestine have also been developed (Ayehunie, unpublished). The availability of the models from the different segments of the small intestine will help address questions regarding the contribution of each segment in drug pharmacokinetics studies.

One of the attractive features of the organotypic tissue models is that they express efflux transporters and drug-metabolizing enzymes that mimic in vivo intestinal tissues. For instance, RT-PCR analysis showed that the tissue models expressed drug-metabolizing enzymes (e.g., CYP3A4, CYP3A5, CYP2C9, and CYP2C19 and GSTs) and the efflux transporters (e.g., P-gp/MDR1, MRP1, MRP-2, and BCRP), which are known to be present on walls of human intestinal tissue. The activity of the drug transporters and drug-metabolizing enzymes, MDR1, P-gp, BCRP, CYP3A4, CYP2J2, UDP-glucuronosyltransferases (UGT), and carboxylesterases (CES) on the intestinal organotypic tissue models was confirmed using functional assays with selective substrates and inhibitors. In this study, the authors concluded that EpiIntestinal tissues provide a more holistic model for the investigation of drug absorption and metabolism in human gastrointestinal tract compared with Caco-2 cells (Cui et al. 2020).

These organotypic tissue models allow the evaluation of lot-to-lot reproducibility using endpoints such as transepithelial electrical resistance (TEER) and the LY leakage assay, both of which monitor the barrier integrity of the organotypic intestinal tissues. For instance, TEER values of 90–300 Ω cm² and LY leakage < 4% were applied as QC acceptance criteria for EpiIntestinal tissues (MatTek Corporation). The mean TEER values for the partial thickness, full-thickness 24-well format, and 96-well formats were shown to be 150 ± 8.6 Ω cm² (N = 129 lots), 175 ± 12.3 Ω cm² (N = 201 lots), and 162 ± 10.8 Ω cm² (N = 68 lots), respectively, which is close to physiological values.

Use of 3D Organotypic Intestinal Tissue Models to Predict Drug Absorption and Metabolism: The utility of the EpiIntestinal model to rank order the permeability of three model drugs, representative of low (< 50%, talinolol), moderate (50–84%, ranitidine), and high (≥ 85%, warfarin) absorption in humans, was demonstrated (Ayehunie et al. 2018). Drug permeation data obtained from EpiIntestinalFT and Caco-2 cells were also compared with historical data of % fraction absorbed in humans. The results showed that the EpiIntestinalFT data correlated better with human absorption data (r² = 0.91) compared with Caco-2 cells (r² = 0.71) (Ayehunie et al. 2018). The study was expanded further to analyze the permeation of 28 drugs with diverse absorption properties. Permeability coefficients determine with the EpiIntestinal tissues were compared with historical data for the fraction-absorbed (Fa) values observed in humans (Table 2). Using an apparent permeability coefficient (Papp) value of 5 × 10⁻⁶ as a cutoff, the EpiIntestinal tissues were able
to classify test articles as high ($P_{\text{app}} > 5 \times 10^6$) permeability or low permeability drugs ($P_{\text{app}} < 5 \times 10^6$), which correlated well with historical absorption data in humans (Table 2; Ayehunie et al. 2018). Overall, the reconstructed intestinal tissue model was able to differentiate flux of high vs low permeability compounds, and monitor drug–drug interactions, and metabolism (Ayehunie et al. 2018; Cui et al. 2020).

In another study, the effect of enhancers on the permeation of low permeability drugs was demonstrated using the EpiIntestinal tissue model. Complexing the low permeability, idiopathic pulmonary fibrosis drug, Nintedanib, with cyclodextrin was shown to significantly increase absorption and bioavailability across the intestinal barrier (Vaidyaa et al. 2019). Recently, Marrella et al. (2020) utilized in a flow system to monitor the absorption of two non-metabolized sugars, lactulose and mannitol, in EpiIntestinal tissues grown under standard (healthy) and pathological conditions (EGTA-induced barrier disruption). The ratio of lactulose-to-mannitol in urine samples is a clinical test used to assess disorders characterized by changes in gut permeability (Johnston et al. 2000), and this kinetics was well recapitulated in the closed circuit hosting EpiIntestinal tissue (Marrella et al. 2020).

Since metabolism and facilitated transport are key components in the assessment of oral drug absorption and bioavailability, drug-metabolizing enzymes and transporters play an important role in determining the pharmacokinetics, safety, and efficacy profiles of drugs (International Transporter Consortium et al. 2010). In this regard the in vitro 3D organotypic intestinal tissue models are emerging as a
predictive tool to examine drug bioavailability, drug–drug interaction, and drug biotransformation. For instance, apical (A–B) exposure of the EpiIntestinal tissues to the drugs (10 μM of midazolam or fosphenytoin for 2 h) resulted in conversion of the parent drug Midazolam to its metabolite, alpha-hydroxymidazolam (6.5% conversion), and the parent drug Fosphenytoin to its metabolite, Phenytoin (57.2% conversion), verifying the functionality of the Cytochrome P450 (CYP) 3A4 enzyme (Ayehunie, manuscript in preparation). Recently, relevant drug transporters and drug-metabolizing enzymes, including MDR1 (P-gp), BCRP, CYP3A4, CYP2J2, UDP-glucuronosyltransferases (UGT), and carboxylesterases (CES), were detected in functional assays with selective substrates and inhibitors (Cui et al. 2020). In this study, the selective BCRP inhibitor Ko-143 (3 μM) strongly reduced the efflux of rosuvastatin by 32.2-fold compared with the no inhibitor control. Additionally, a substantial amount of 1-hydroxymidazolam, the CYP3A4-selective metabolite of midazolam, was detected in EpiIntestinal tissues and this conversion was suppressed by the addition of the selective CYP3A4 inhibitor, CYP3cide. In comparison, only a negligible amount of the metabolite was detected in Caco-2 cultures. In order to profile the metabolism capacity of EpiIntestinal tissues the investigators further study the metabolic pathway of the double pro-drug Dabigatran etexilate. Dabigatran etexilate is known to be first hydrolyzed by CES1 into BIBR 1087 and the formation of the active drug BIBR 953 from the intermediate metabolite BIBR 1087 is catalyzed by CES2 (predominantly expressed in human intestine). Interestingly, Dabigatran etexilate was metabolized to

| Test article | Mean A → B $P_{app}$ ($10^{-6}$ cm s$^{-1}$) St dev | Mean B → A $P_{app}$ ($10^{-6}$ cm s$^{-1}$) St dev | Efflux ratio | Reproducibility | Human (historical values) | BCS classification |
|--------------|-----------------------------------------------|-----------------------------------------------|--------------|----------------|-----------------------|------------------|
| Carbamazepine | 19.7 1.9 | 32.6 1.7 | 1.7 | N = 3 | 97 | High |
| Citalopram | 15.9 1.3 | 25.0 1.6 | 1.6 | N = 2 | 90 | High |
| Digoxin | 1.1 0.6 | 3.6 3.1 | 3.1 | N = 2 | 81 | Low/efflux substrate |
| Midazolam | 11.1 0.4 | 34.1 3.1 | 3.1 | N = 2 | 90 | High |
| Metoprolol | 8.4 2.4 | 22.6 2.7 | 2.7 | N = 5 | 95 | High |
| Metronidazole | 14.0 0.4 | 18.8 1.3 | 1.3 | N = 3 | 80 | Low/high |
| Mycophenolate | 12.8 2.3 | 10.7 0.8 | 0.8 | N = 2 | 94 | High |
| Naproxen | 15.5 3.8 | 19.4 2.6 | 2.6 | N = 2 | 98 | High |
| Omeprazole | 12.2 5.2 | 24.9 2.0 | 2.0 | N = 3 | 88 | High |
| Propranolol | 8.4 4.4 | 25.9 3.1 | 3.1 | N = 3 | 90 | High |
| Quinidine | 8.0 2.4 | 17.0 2.1 | 2.1 | N = 3 | 80 | High |
| Verapamil | 5.7 2.8 | 25.6 4.5 | 4.5 | N = 3 | 100 | High |
| Warfarin | 18.8 4.8 | 9.1 0.5 | 0.5 | N = 4 | 98 | High |
| Aцикловир | 0.3 0.0 | 1.1 3.4 | 3.4 | N = 3 | 30 | Low |
| Amoxicillin | 0.6 0.4 | 1.1 1.8 | 1.8 | N = 2 | 77 | Low/high |
| Atenolol | 0.7 0.4 | 1.1 1.5 | 1.5 | N = 5 | 50 | Low/moderate |
| Cиметидин | 1.9 0.2 | 1.5 0.8 | 0.8 | N = 3 | 62 | Low |
| Ethambutol | 0.8 0.7 | 0.6 0.8 | 0.8 | N = 3 | 75 | Low |
| Enalapril | 0.5 0.3 | 0.7 1.4 | 1.4 | N = 5 | 40 | Low/moderate |
| Erythromycin | 0.6 0.4 | 2.5 4.5 | 4.5 | N = 3 | 35 | Low |
| Furosemide | 0.7 0.3 | 4.9 7.0 | 7.0 | N = 3 | 61 | Low/moderate |
| Metformin | 1.3 0.5 | 1.3 1.0 | 1.0 | N = 4 | 71 | Low/moderate |
| Methotrexate | 0.8 0.5 | 0.7 0.9 | 0.9 | N = 5 | 20 | Low/Moderate |
| Ramitidine | 1.1 0.6 | 1.3 1.2 | 1.2 | N = 3 | 61 | Low/Moderate |
| Raloxifine | 1.9 0.1 | 3.1 1.7 | 1.7 | N = 2 | 60 | Low |
| Rosuvastatin | 0.4 0.1 | 7.0 16.7 | 16.7 | N = 2 | 20 | Low |
the active drug, BIBR953, through the intermediate BIBR 1087 by EpiIntestinal tissues similar to that observed in vivo, suggesting the involvement of functional CES2 esterase enzyme. These results indicate that the EpiIntestinal tissues mimic drug metabolism in the human small intestine and likely will be more predictive than Caco-2 cells which lack key drug-metabolizing enzymes found in native tissue. It is anticipated that the 3D human organotypic small intestinal model can play a role in narrowing the gap from preclinical to clinical translation of results.

In summary, the drug-metabolizing enzymes and drug transporters expressed in the 3D-human intestinal tissue models make them appropriate for studies involving (a) bidirectional drug transport from apical (A) to basolateral (B) or B-to-A, (b) drug–drug interaction, and (c) drug metabolism studies. Compared with animal studies, the in vitro intestinal models will be more cost-effective and will allow for high throughput screening. These models will be useful for optimizing compound formulations and predicting metabolism and bioavailability of orally administered drugs at an early stage of the drug development process. Additional evaluation by transcriptomic, proteomic, metabolomic, and functional endpoints will increase confidence in the use of intestinal models for pharmacokinetic studies (Sawant-Basak et al. 2018).

Drug Toxicity Studies Drug-induced gastrointestinal toxicity (DIGT) is among the most frequently occurring adverse effects in clinical studies (Peters et al. 2020). To date, no specific mechanistic diagnostic/prognostic biomarkers or translatable preclinical models of DIGT exist (Carr et al. 2017). Therefore, predicting toxicity of candidate and investigational drugs is of paramount importance in the preclinical stage of drug development (Stevens and Baker 2009). Currently, preclinical toxicity testing heavily depends on the animal models. However, animal models, in particular rodents, show very low predictability for DIGT adverse effects. Better predictions can be made in canine and non-human primate models; however, due to cost, these species are often only used in later stages of drug development (Olson et al. 2000). To overcome challenges of determining DIGT at the preclinical stage, in vitro models can be used as a promising alternative to animal testing (Peters et al. 2019). For example, AstraZeneca tested a validation set of 31 widely prescribed drugs with the Epilntestinal tissue model. Using TEER as a measure of barrier function, a threshold value was established that distinguished between drugs that induced clinical diarrhea from those which were well-tolerated. The in vitro study gave a predictive accuracy of 80% which matches the translation accuracy of in vivo studies in higher-order species. The predictivity was much higher for parallel experiments performed with Caco-2-based model (Peters et al. 2019). In other experiments on a limited set of other drugs which had failed in clinical trials, the EpiIntestinal TEER–based assay more accurately predicted the maximum tolerated dose than was predicted based on 1 mo-long animal studies (rats and dogs). The authors concluded that the EpiIntestinal tissue model is the first in vitro assay with validated predictivity for diarreaha-inducing drugs and suggested its potential utility for lead optimization, dose schedule exploration, and clinical translation (Peters et al. 2019).

Nanoparticle Toxicity Recently, there has been a dramatic increase in the use of engineered nanoparticles (ENPs) in a broad range of products. ENPs introduced into the body through food, drink, pharmaceutical formulations, or inadvertently lead to the increased exposure of all body systems including the GI tract, where there is a very limited amount of toxicological data (McCauley and Wells 2017). In vitro intestinal tissues represent a system that might allow rapid prescreening of the ingestible nanomaterial particles. Studies aimed at observing the toxic properties of copper nanoparticles such as (CuO and CuSO4) on EpiIntestinal tissues revealed a significant decrease in tissue viability of 30% and 75% for doses of 40 μg/mL and 80 μg/mL, respectively, when exposed to 24 h time period (Henson et al. 2019). The authors also showed that EpiIntestinal tissues require higher concentrations of CuO to elicit a cytotoxic effect by 24 h compared with rat intestinal epithelial cell line (IEC-6, 2D intestinal model).

Recent experiments in our laboratory examined the effects of cupric(II) oxide (CuO) (50 nm), zinc oxide (ZnO, 35–45 nm), titanium oxide (TiO2, 40 nm), and silver (20 nm, 30–50 nm, and 80–100 nm) nanoparticles on the Epilntestinal tissues following 24 h exposure to different concentrations of these ENPs by monitoring (1) barrier integrity (TEER), (2) tissue viability (MTT assay), (3) oxidative stress (8-isoprostane release), and (4) inflammatory response (IL-8). The results showed a dose-dependent reduction of the tissue barrier and viability following exposure to CuO, ZnO, and single wall carbon nanotubes (SWCNT). The analysis of culture supernatants collected at 24 h also showed a dose-dependent release of IL-8 for CuO and ZnO and the oxidative stress indicator 8-isoprostane for CuO (Ayeunie, manuscript in preparation). Silver nanoparticles showed no acute adverse effects on the intestinal microtissues in vitro, which was consistent with in vivo observations (Burdus et al. 2018). In summary, the Epilntestinal tissue model appears useful as a preclinical screening tool to examine the toxicity profile of ingested nanoparticles and to improve the design of nanoparticle-based therapeutic formulations or personal care products.

Intestinal Inflammation: Epithelial tissues exposed to chemicals, drugs, ligands, antigens, and cytokines have been shown to undergo quantitative changes in inflammatory responses and epithelial permeability. These inflammatory
responses could (1) exacerbate cytokine/chemokine release, (2) alter drug-metabolizing enzymes and ABC transporter expression levels, and/or (3) compromise the epithelial barrier by affecting tight junctions. To examine the effect of cytokines on innate immune responses, the EpiIntestinal tissues were treated with IFN-γ, IL-17A/F, or IL-17A/F plus IFN-γ and changes in gene expression levels for the proinflammatory cytokines, CCL-20, IL-8, TNF-α, and CXCL5, were monitored. The results showed increases in IL-8, TNF-α, and CXCL5 in a synergetic manner (Fig. 3). In another study, the impact of IL-6 on drug-metabolizing enzymes and transporters was investigated using the EpiIntestinal tissue model. mRNA expression levels of inflammatory response receptors, drug-metabolizing enzymes (phase I and phase II), drug transporters, and nuclear transporters were impaired when the EpiIntestinal tissues were exposed to the proinflammatory cytokine IL-6 (5–20 ng/mL) for 72 h (Simon et al. 2019). The authors showed that exposure of EpiIntestinal tissues to IL-6 can have downstream effects in reducing CYP450 mRNA expression such as CYP2C19, CYP2C9, and CYP3A4 by 40–50% with a decrease in activity level of 20–75%. Such observations may have important implications in relation to the effect of released inflammatory cytokines in presystemic metabolism and dosage application.

Data from another study suggested that IL-22 may contribute to tissue inflammation in certain mouse models (Kamanaka et al. 2011). Using the EpiIntestinal model, the function of IL-22 on the regulation of intestinal epithelium

Figure 3. Cytokine/chemokine gene expression levels following treatment of EpiIntestinal tissues with T cell cytokines (interferon gamma and IL-17 A/F).

Figure 4. Migrating epithelial cells stained with cytokeratin-19 (red), fibroblasts stained for vimentin (green), and nuclei stained with DAPI (blue) after wounding of full-thickness intestinal tissues (EpiIntestinalFT) with 2 mm biopsy punch and cultured for 3 d. Note: On day 3, the fibroblasts are at the leading edge of the resealing wound.
barrier was examined. The investigators found that exposure of EpiIntestinal tissues to IL-22 (100 ng/mL) for 24 h significantly induced the expression of Claudin-2, a cation–pore/channel-forming tight junction protein through the Janus kinase-signal transducer and activator of transcription (JAK/STAT) signaling pathway. Upregulation of the pore forming claudin-2 by IL-22 led to reduced transepithelial TEER and increased paracellular intestinal epithelial permeability of ions (Wang et al. 2017). The researchers also noted that IL-22 signals exclusively through the basolateral side of the polarized tissues.

Repeated epithelial damage or injury is also implicated in intestinal disorders including inflammatory bowel disease. Rapid closure or resealing of wounds has key physiological importance and is a natural defense mechanism against such damage. The process of intestinal epithelial restitution and the various factors that contribute to gut homeostasis and intestinal epithelial wound healing following injury have been reviewed in detail by Lizuka and Konno (2011) using rat IECs (IEC-6) cell culture system. In this system, the epithelial cells found in the wounded area are thought to have migrated into the wound in a process of restitution which involves reorganization of the actin cytoskeleton. To recapitulate in vivo wound repair, the EpiIntestinal tissue was wounded using a 2-mm biopsy punch. The wounded tissues were maintained in culture and time dependent healing was noted. On day 2 post-injury, epithelial cells were found at the leading edge of the wound and on day 3, fibroblasts had

| Parameter                          | Organoid                        | Organotypic tissue models                                      |
|-----------------------------------|---------------------------------|-----------------------------------------------------------------|
| Size and shape                    | Undefined                       | Defined tissue diameter                                        |
| Culture condition and Epithelial  | Embedded in Matrigel; inward   | Wall-to-wall polarized tissue growth with accessible apical “luminal” and basolateral surface |
| differentiation                   | villi growth                    | Allows access to apical and basal sides for drug permeation studies |
| Drug application                  | Access to apical side is difficult (microinjection needed) | Quantified by TEER measurement and leakage experiments such as Lucifer yellow leakage |
| Barrier integrity measurement     | Not quantifiable                | Can be modeled                                                 |
| Intestinal restitution/wound healing | Difficult to model             | Standardized                                                   |
| Quality control data              | Not standardized                | Difficult                                                       |
| Patient-specific disease modeling | Possible                       | Majority expressed                                             |
| Intestinal cell phenotypes        | Expressed                       | Yes                                                            |
| Host–pathogen interaction studies | Yes                             | Up to 6 wk                                                     |
| Long-term culture                 | A year or more                  | Relatively inexpensive                                         |
| Cost                              | Expensive (due to growth factors) |                                                                |

Figure 5. Histology of EpiIntestinalFT showing tissue restitution over a 6-d period after wounding. Initial migration of leading cells over the wounded section of the tissue was noted on day 2 post-wound, black arrow) and complete resealing of the wound and tissue differentiation occur on day 6.
expanded into the wound and epithelial cells had started to grow on the fibroblasts. On day 4, the wound was re-epithelialized. By day 6, there was a complete intestinal restitution and differentiation of epithelial layer. The progress of the healing process was monitored by immunohistochemical staining and histological observation (Figs. 4 and 5).

Infectious Disease Modeling: Enteric pathogens which are the causative agents of diarrheal diseases are a significant public health burden in that they often result in hospitalization of the elderly (> 65 yr old) in developed nations and cause sickness or death in children and adults in developing countries (Phalipon and Sansonetti 2007; Schiller 2009). Humans and/or higher primates are the natural hosts for certain enteric pathogens (e.g., *Shigella flexneri* and the Norovirus; Ettayebi et al. 2016; Todd and Tripp 2019). Due to the cost and ethical issues associated with the use of higher primates, human intestinal tissues are ideal models to decipher cell-pathogen interactions and mechanisms of infection. In a recent study, the EpiIntestinal model was used to investigate the pathophysiology and mechanisms by which *S. flexneri* invades the intestinal epithelium. *S. flexneri* affects the columnar architecture of the villi of intestinal tissues accompanied by F-actin organizational disruption associated with cofilin activation (Maldonado-Contreras et al. 2017). Mechanistically, *S. flexneri* infection secretes serine protease A (SepA) which is responsible for the disruption of the intestinal epithelial barrier via the LIM kinase 1 (LIMK1) pathway (Maldonado-Contreras et al. 2017) and which in turn is involved in negative inhibition of actin-remodeling proteins such as cofilin. Insights gained from these studies demonstrated the use of the in vitro intestinal tissues as a valuable enteric bacterial infection model to study molecular mechanisms or pathways involved in viral infection and to screen the safety and efficacy of candidate drugs against enteric pathogens. Most importantly, a tissue model that can be infected with human pathogens such as *S. flexneri* (causative agent of Shigellosis) is critically needed since humanized mouse models or xenotransplants of human fetal intestine to mice poorly reflect human responses and are time-consuming, expensive, and ethically problematic.

Although the main route of SARS-CoV-2 (causative agent of COVID-19 disease) transmission is generally accepted to be through the respiratory tract, emerging reports indicate that the intestinal tissue could be an important target organ that plays a role in SARS-CoV-2 infection and transmission. A review by He et al. (2020) showed potential existence of a “gut-lung” axis in SARS-CoV-2 infection and pointed out that patients with gastrointestinal (GI) symptoms showed increased disease severity. Clinical studies show diarrhea to be one of the symptoms in patients infected with SARS-CoV-2 at a rate ranging from 2 to 50% of cases, indicating a potential role of the small intestine in COVID-19 infection and transmission (D’Amico et al. 2020). Recent studies have shown that the EpiIntestinal tissue model strongly expresses the ACE2 receptor along with TMPRSS2, both of which are essential for SARS-CoV-2 entry to target tissues (Wang et al. 2020). Clevers group and others also reported productive infection of SARS-CoV-2 in ACE2+ mature enterocytes in human small intestinal organoids/enteroids, and infection was facilitated by TMPRSS2 and TMPRSS4, by promoting SARS-CoV-2 spike fusogenic activity which may promote virus entry into host cells (Lamers et al. 2020; Zang et al. 2020). Hence, the role of the small intestinal organotypic and organoids in identifying biomarkers to predict SARS CoV-2 infection and transmission or disease severity needs further investigation.

**Conclusion**

The need for predictive in vitro intestinal models for drug absorption, metabolism, inflammation, drug-induced GI toxicity, disease modeling, and intestinal injury repair has led to the development of a variety of models that utilize animals, cell lines, ex vivo tissues, organoids, and organotypic small intestinal tissues. Emerging data show that organoids and organotypic tissue culture have polarized tissue architecture and physiological features and functions that recapitulate many of the in vivo small intestinal properties. Among the models, the EpiIntestinal tissues are found to be highly reproducible and are widely used by industry and academic institutions for basic and applied research. Although recent developments in intestinal organoid cultures have shown promise in translational research, the organoid culture format has limitations compared with 3D organotypic cultures (Table 3).

Recently, the EpiIntestinal tissue model was able to identify with > 80% accuracy the commonly prescribed drugs that are known to induce clinical diarrhea. This predictive accuracy matches the translation accuracy of in vivo studies in higher-order species. Due to the presence of efflux transporters and drug-metabolizing enzymes, the organotypic intestinal tissue models can be used to predict drug absorption and metabolism to mimic responses of the human gastrointestinal tract and these models are more holistic compared with Caco-2 cells which lack major drug-metabolizing enzymes.

**Funding** Previously unpublished results presented herein for the EpIIntestinal tissue model were partially supported by grants by the National Institute of Health (NIH) from the National institute of General Medicinal Science (NIGMS, R44GM108164) and National Institute of Environmental Health Sciences (NIEHS, R43ES030648).

**References**

Anderle P, Rakhmanova V, Woodford K, Zerangue N, Sadée W (2003) Messenger RNA expression of transporter and ion channel genes in...
useful screening test for adult coeliac disease. Ann Clin Biochem 37: 512–519

Kamanaka M, Huber LA, Zeneewicz N, Gagliani C, Rathinam W et al (2011) Memory/effector (CD45RB(lo)) CD T cells are controlled directly by IL10 and cause IL-22-dependent intestinal pathology. J Exp Med 208:1027–1040

Kasendra M, Tovagliari A, Sontheimer-Phelps A, Jalili-Firoozinezhad S, Bein A, Chalakiadaki A, Scholl W, Zhang C, Rickner H, Richmond C, Li H, Breault DT, Ingeber DE (2018) Development of a primary human small intestine-on-a-chip using biopsy-derived organoids. Sci Rep 8:2871

Kiela PR, Ghishan FK (2016) Physiology of intestinal absorption and secretion. Best Pract Res Clin Gastroenterol 30:145–159

Kleieveld CR (2015) Co-cultivation of Caco-2 and HT-29MTX. In: Verhoeckx K et al (eds) The impact of food bioactives on health. Springer, Cham. https://doi.org/10.1007/978-3-319-16104-4_13

König J, Wells J, Patrice D, Carter PD, Garcia-Rodenas CL, McDonald T, Mercenier A, Whyte J, Freddy Troost F, Brummer R-J (2016) Human intestinal barrier function in health and disease. Clin Transl Gastroenterol 7:e196

Lamers MM, Beumer J, van der Vaart J, Knoops K, Puschhof J, Breugem TJ, Ravelli RBG, Ivan Schayck JP, Myktyyn AZ, Duimel HQ, Donselaar E, Rieseobach S, Kuipers HJJ, Schipper D, van der Wetering WJ, de Graaf M, Koopmans M, Cuppen E, Peters PJ, Haagmans BL, Hans CH (2020) SARS-CoV-2 productively infects human gut enterocytes. Science 369:50–54

Lehle AS, Farin HF, Marquardt B, Michels BE, Magg T, Li Y, Liu Y, Ghalandary M, Lamans K, HolleziCK S, RohlfM S, Conca R, Walz C, Weiss B, Lev A, Simon AJ, Groll O, Gaedt MM, Hornung V, CLEVERS H, Yazbeck N, Hanna-Wakim R, Shouval DS, Warner N, Somech R, Muise AM, SNAPPER SB, Butler P, Koletsko S, Klein C, Kotlarz D (2019) Intestinal inflammation and dysregulated immunity in patients with inherited caspase-8 deficiency. Gastroenterology 156:275–278

Leslie JL, Huang S, Opp JS, Nagy MS, Kobayashi M, Young VB, Spence JR (2015) Persistence and toxin production by Clostridium difficile within human intestinal organoids result in disruption of epithelial paracellular barrier function. Infect Immun 83:138–145

Li AP (2005) Preclinical in vitro screening assays for drug-like properties. LeHle AS, Farin HF, Marquardt B, Michels BE, Magg T, Li Y, Liu Y, Ghalandary M, Lamans K, HolleziCK S, RohlfM S, Conca R, Walz C, Weiss B, Lev A, Simon AJ, Groll O, Gaedt MM, Hornung V, CLEVERS H, Yazbeck N, Hanna-Wakim R, Shouval DS, Warner N, Somech R, Muise AM, SNAPPER SB, Butler P, Koletsko S, Klein C, Kotlarz D (2019) Intestinal inflammation and dysregulated immunity in patients with inherited caspase-8 deficiency. Gastroenterology 156:275–278

Leslie JL, Huang S, Opp JS, Nagy MS, Kobayashi M, Young VB, Spence JR (2015) Persistence and toxin production by Clostridium difficile within human intestinal organoids result in disruption of epithelial paracellular barrier function. Infect Immun 83:138–145

Li AP (2005) Preclinical in vitro screening assays for drug-like properties. Drug Discov Today Technol Actions Summer 2:179

Lin J, Hackman DJ (2011) Worms, flies and four-legged friends: the ap-
Stevens LJ, van Lipzig MMH, Erpelink SLA, Pronk A, van Gorp J, Wortelboer HM, van de Steeg E (2019) A higher throughput and physiologically relevant two-component human ex vivo intestinal tissue system for studying gastrointestinal processes. Eur J Pharm Sci 137:104989

Sun H, Chow EC, Liu S, Du Y, Pang KS (2008) The Caco-2 cell monolayer: usefulness and limitations. Expert Opin Drug Metab Toxicol 4:395–411

Takenaka T, Harada N, Kuze J, Chiba M, Iwao T, Matsunaga T (2014) Human small intestinal epithelial cells differentiated from adult intestinal stem cells as a novel system for predicting oral drug absorption in humans. Drug Metab Dispos 42:1947–1954

Tavelin S, Taipalensuu J, Söderberg L, Morrison R, Chong S, Artursson P (2003) Prediction of the oral absorption of low permeability drugs using small intestine-like 2/4A1 cell monolayers. Pharm Res 20: 397–405

Ting H-A, von Molike J (2019) The immune function of tuft cells at gut mucosal surfaces and beyond. J Immunol 202:1321–1329

Todd K, Tripp R (2019) Human norovirus: experimental models of infection. Viruses 11:2(1)

Vaidyaa B, Shuklab SK, Kollurua S, Huena M, Mullac N, Mehrad N, Kanabar D, Balakurthid S, Ayehunie S, Muthb A, Gupta V (2019) Nintedanib-cyclodextrin complex to improve bioactivity and intestinal permeability. Carbohydr Polym 204:68–77

van der Flier LG, Clevers H (2009) Stem cells, self-renewal, and differentiation in the intestinal epithelium. Annu Rev Physiol 71:241–260

VanDussen KL, Marinshaw JM, Shaikh N, Miyoshi H, Moon C, Tarr PI, Vaidyaa B, Shuklab SK, Kollurua S, Huena M, Mullac N, Mehrad N, Kanabar D, Balakurthid S, Ayehunie S, Muthb A, Gupta V (2019) Nintedanib-cyclodextrin complex to improve bioactivity and intestinal permeability. Carbohydr Polym 204:68–77

Van der Flier LG, Clevers H (2009) Stem cells, self-renewal, and differentiation in the intestinal epithelium. Annu Rev Physiol 71:241–260

Walter E, Janich S, Roessler BJ, Hilfinger JM, Amidon GL (1996) HT29-MTX/Caco-2 cocultures as an in vitro model for the intestinal epithelium: in vitro-in vivo correlation with permeability data from rats and humans. J Pharm Sci 85:1070–1076

Walters E, Wolf E, Whyte J, Mao J, Renner S, Nagashima H et al (2012) Completion of the swine genome will simplify the production of swine as a large animal biomedical model. BMC Med Genom 5:55

Wang B, Kovalchuk I, Li D, Ilnytskyy Y, Kovalchuk I, Kovalchuk O (2020) In search of preventative strategies: novel anti-inflammatory high-CBD cannabis sativa extracts modulate ACE2 expression in COVID-19 gateway tissues. Preprints 2020:2020040315. https://doi.org/10.20944/preprints202004.0315.v1

Wang Y, Mumm JB, Herbst R, Kolbeck R, Wang Y (2017) IL-22 increases permeability of intestinal epithelial tight junctions by enhancing claudin-2 expression. J Immunol 199:3316–3325

Waterston RH, Lindblad-Toh K, Birney E, Rogers J, Abril JF, Agarwala P, Agarwala R, Ainscough R, Alexandersson M, An P, Antonarakis SE, Attwood B, Baertsch R, Bailey J, Barlow K, Beck S, Berry E, Birren B, Bloom T, Bork P, Botcherby M, Bray N, Brent MR, Brown DG, Brown SD, Bult C, Burton J, Butler J, Campbell RJ, Carninci P, Cawley S, Chiariomonte F, Chinwalla AT, Church DM, Clamp M, Clee C, Collins FS, Cook LL, Copley RR, Coubold A, Couronne O, Cuff J, Curwen V, Cutts T, Daly M, David R, Davies J, Delahaunty KD, Deri J, Dernitzakis ET, Dewey C, Dickens NJ, Diekhans M, Dodge S, Dubchak I, Dunn DM, Eddy SR, Elnitski L, Eres MD, Eswara P, Eyras E, Felsenfeld A, Fewell GA, Flicek P, Foley K, Frankel WN, Fulton LA, Fulton RS, Furey T, Gage D, Gibbs RA, Glusman G, Gnerre S, Goldman N, Goodstadt L, Grafham D, Graves TA, Green ED, Gregory S, Guigó R, Guyer M, Hardison RC, Haussler D, Hayashizaki Y, Hillier LW, Hinrichs A, Hlevina W, Holzer T, Hsu F, Hua A, Hubbard T, Hunt A, Jackson I, Jaffe DB, Johnson LS, Jones M, Jones TA, Joy A, Kamal M, Karlsson EK, Karolchik D, Kasprzyk A, Kawai J, Keibler E, Kells C, Kent WJ, Kirby A, Kolbe DL, Korf I, Kucherlapati RS, Kulpokas EJ, Kulp D, Landers T, Leger JP, Leonard S, Letunic I, Levine R, Li J, Li M, Lloyd C, Lucas S, Ma B, Maglott DR, Mardis E, Matthews L, Mauceli E, Mayer JH, McCarthy M, McCombie WR, McLaren S, McErsky K, McPherson JD, Meldrim J, Meredith B, Mesirov JP, Miller W, Miner TL, Mongin E, Montgomery KT, Morgan M, Mott R, Mullikin JC, Muzny DM, Nash WE, Nelson JO, Nhan MN, Nicol R, Ning Z, Nusbaum C, O’Connor MJ, Okazaki Y, Oliver K, Overton-Larty E, Pachter L, Parra G, Pepin KH, Peterson J, Pevzner P, Plumb R, Pohl CS, Poliaokh A, Ponce TC, Ponting CP, Potter S, Quail M, Reymond A, Roe BA, Roskin KM, Rubin EM, Rust AG, Santos R, Sapojnikov V, Schultza B, Schultz J, Schwartz MS, Schwartz S, Scott C, Seaman S, Searle S, Sharpe T, Sheridan A, Showkenne R, Sims S, Singer JB, Slater G, Smit A, Smith DR, Spencer B, Stabenau A, Stange-Thommann N, Sugnet C, Suyama M, Tesler G, Thompson J, Torres M, Trevaskis E, Tromp J, Ucla C, Ureta-Vidal A, Vinson JP, Von Niederhausern AC, Wade CM, Wall M, Weber RJ, Weiss RB, Wendt MC, West AP, Wetterstrand K, Wheeler R, Whelan S, Wierzbowski J, Willey D, Williams S, Wilson RJ, Winter E, Worley KC, Wyman D, Yang S, Yang SP, Zdobnov EM, Zody MC, Lander ES (2002) Initial sequencing and comparative analysis of the mouse genome. Nature 415:395–405

Woodcock S, Williamson J, Hassan J, Martin Mackay M (1991) Isolation and characterization of clones from the Caco-2 cell line displaying increased taurocholic acid transport. J Cell Sci 98:323-327

Zang R, Gomez Castro MF, McCune BT, Zeng Q, Rothlauf PW, Sonnek NM, Liu Z, Bruleis KF, Wang X, Greenberg HB, Diamond MS, Ciorba MA, Whelan S, Wierzbowski J, Willey D, Williams S, Wilson RK, Stabenau A, Stange-Thomann N, Sugnet C, Suyama M, Tesler G, Scott C, Seaman S, Searle S, Sharpe T, Sheridan A, Showkenne R, Sims S, Singer JB, Slater G, Smit A, Smith DR, Spencer B, Stabenau A, Stange-Thommann N, Sugnet C, Suyama M, Tesler G, Thompson J, Torres M, Trevaskis E, Tromp J, Ucla C, Ureta-Vidal A, Vinson JP, Von Niederhausern AC, Wade CM, Wall M, Weber RJ, Weiss RB, Wendt MC, West AP, Wetterstrand K, Wheeler R, Whelan S, Wierzbowski J, Willey D, Williams S, Wilson RK, Winter E, Worley KC, Wyman D, Yang S, Yang SP, Zdobnov EM, Zody MC, Lander ES (2002) Initial sequencing and comparative analysis of the mouse genome. Nature 420:397–405

Woodcock S, Williamson J, Hassan J, Martin Mackay M (1991) Isolation and characterization of clones from the Caco-2 cell line displaying increased taurocholic acid transport. J Cell Sci 98:323-327