Chapter 22
Intestinal Crypt Organoids as Experimental Models

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Abstract  When it comes to studying the effect of food bioactives on gut health, one of the essential steps that needs to be assessed is characterizing specific effects of the bioactives on the physical barrier of the lumen, the gastrointestinal tissue. In addition to studying the effects on transport function (e.g. by using Ussing chambers or cell culture systems), it is of great interest to evaluate the effects on morphology, cell biology, gene expression, and relevant functions of different cell types that are resident in the gastrointestinal (GI) tract. An ideal near-physiological model should contain a mixture of different GI epithelial cells (e.g. Paneth cells, goblet cells, absorptive and hormone secretive epithelial cells), which can be cultured indefinitely. Recently, the culture and applications of long-term primary multi-cellular cluster structures gastrointestinal organoids (or enteroids) have been demonstrated, and within the last 5 years the number of researchers that commonly use this tissue culture model has increased rapidly. This multi-cellular system may be a promising addition for existing ex vivo and alternative for animal models for testing effects of food bioactives on the intestinal tissue, and could provide a model for pre-screening of compounds prior to moving to the large scale testing systems. Moreover, intestinal organoids can be cultured from different species (e.g. human, pig and mouse). In this chapter we will focus on organoids cultured from mouse and pig crypt cells. We will give a short overview on how to isolate, culture, incubate, and apply them in different research fields.

Keywords  Ex vivo • Intestinal epithelium • Paneth cells • Crypts • Lgr5 • Stem cells
22.1 Gastrointestinal Organoids

Several attempts have been made to isolate the proliferative regions of the intestinal epithelium from different species (e.g. mouse, porcine, human) (Pageot et al. 2000; Agopian et al. 2009). Proliferative regions of the intestinal epithelium consist of the transit amplifying cells, which are undifferentiated progenitor cells that can differentiate into all different epithelial cells of the intestinal epithelium. Initially cultured intestinal tissue contained crypt and villus domains, resembling the intestinal epithelial architecture present in vivo. However, these cultured intestinal tissue pieces from the intestinal epithelial could be maintained in culture for up to several days at maximum, and thus were not self-renewing.

The need for a self-sustaining, long-term GI model remained. The big breakthrough came when Dr. Hans Clevers and colleagues, for the first time, identified the intestinal stem cells by thorough lineage tracing experiments (Barker et al. 2007). These experiments were performed over a period of 60 days in mice, and confirmed the presence of Lgr5 (marker for adult stem cells)-positive stem cells in small intestinal and colonic crypts. These stem cells are able to differentiate into all intestinal epithelial cells (enterocytes, Paneth cells, Goblet cells, enteroendocrine cells, but also stem and progenitor cells), as shown in vivo and in culture where these Lgr5-positive cells formed so-called ‘mini-guts’ or ‘organoids’ (Sato et al. 2009) (Fig. 22.1).

![Image of intestinal organoid]

**Fig. 22.1** Schematic image of an intestinal organoid. All epithelial cell types normally present in vivo are also present in the cultured intestinal organoids, as indicated by different colored cells. Figure adapted from Roeselers et al. (2013)
Secretion of mucins into the luminal space and lysozyme production by Panneth cells illustrate the functionality of this epithelium (McCracken et al. 2011).

The discovery of these stem cells opened a new world of the possibilities for GI culture methods, by manipulation of these cells. The great potential of these now self-sustaining ‘mini-guts’ in the field of transplantation and regenerative medicine was immediately evident, and soon engraftment of intestinal organoids in mice was successfully performed (Yui et al. 2012; Shaker and Rubin 2012). So far, the human organoid transplantation has not been performed in humans yet, although human intestinal organoids can be cultured and maintained in culture for some time now (Sato et al. 2011a).

Recently human organoids cultured from induced pluripotent stem cells (iPSCs) have been transplanted into mice (in vivo), where intestinal adaptive capacity as well as epithelial function (peptide uptake and permeability) have been demonstrated in the transplants (Watson et al. 2014). In addition, Clevers and colleagues (University Medical Center Utrecht, the Netherlands), proposed treatment for children suffering from a rare gastrointestinal disease (microvillus inclusion disease) by transplantation of stem-cell derived organoids (Wiegerinck et al. 2014), but the outcomes of these studies have not been published yet. Intestinal organoids have been applied to study cystic fibrosis by using primary organoids from cystic fibrosis patients. This model facilitates diagnosis, functional studies, drug development and personalized medicine approaches in cystic fibrosis (Dekkers et al. 2012).

Recently, we have demonstrated that intestinal organoids provide a powerful model for host–microbiome interaction studies (Lukovac et al. 2014). In this study we observed that specific gut bacteria and their metabolites differentially modulate epithelial transcription in mouse organoids (Lukovac et al. 2014).

### 22.2 General Protocol

Isolation of mouse intestinal crypts and organoid culture, adapted from (Sato and Clevers 2013) is schematically represented in Fig. 22.2. A detailed protocol of the culture protocol is described below.

#### 22.2.1 Small Intestinal Crypt Isolation

Thaw aliquots of Matrigel™ Basement Membrane Matrix (BD Biosciences) on ice before isolation and pre-incubate a 24-well plate in a CO₂-incubator (5 % CO₂, 37 °C). Matrigel™ is a solubilized gelatinous protein mixture and is derived from the Engelbreth–Holm–Swarm (EHS) mouse sarcoma cells. The extract resembles the complex extracellular environment found in many tissues and is commonly used as a substrate for culturing cells. At room temperature, Matrigel™ polymerizes into a three dimensional structure that is useful for both cell culture and studying cellular processes in three dimensions, including cell migration.
Isolate a piece of mouse small intestine (at least 5 cm) and open longitudinally. Wash the intestine with ice cold PBS until most of the luminal contents are cleared and scrape off the villi using a coverslip. After another washing with ice-cold PBS, cut the intestine into 2–4 mm pieces with scissors and transfer them to a 50-mL tube. Add 30 mL ice-cold PBS and wash the fragments by gently pipetting the fragments up and down with a 10 mL pipette and discard supernatant after settling down. Repeat this step 5–10 times until the supernatant is almost clear. Next, add 30 mL of ice-cold crypt isolation buffer (5 mM EDTA in PBS; one can also use higher EDTA concentration, up to 10 mM if yields are low) and gently rock the tube at 4 °C for 30 min. Keep pipetting up and down the tissue in solution every 10 min in order to release as much epithelium as possible. After settling down the fragments, the supernatant can be removed and 20 mL of ice-cold PBS can be added to wash the fragments using a pipet. Repeat this procedure when the fragments are settled down, in order to release most of the crypts, that will also settle down after some time. Villous fractions present in the supernatant are discarded and crypts fractions are passed through a 70-mm cell strainer and collected into 50 mL tube(s), after adding 5% FBS to a crypt solution. Spin down the crypt fractions at 300×g 5 min and resuspend the pellets in 10 mL of ice-cold basal culture medium (advanced Dulbecco’s modified Eagle medium/F12 supplemented with penicillin/streptomycin, 10 mmol/L HEPES, Glutamax, 1×N2, 1×B27, and 1 mmol/L N-acetylcysteine). The suspension is transferred to a 15 mL Falcon tube and centrifuged at 150–200×g for 2 min, to remove single cells (mostly lymphocytes), which end up in the supernatant.
This washing step should be repeated 2–3 times until most single cells are cleared. After the washing steps, the number of crypts can be calculated (take 10 \( \mu \)L of the crypts suspension and count numbers of crypts using a heamocytometer by inverted microscopy; the total number of crypts = the number of crypts counted \( \times 1,000 \)). It should be noted that at this point it is difficult to distinguish true crypts from other epithelial debris, hence the count is only a rough estimate.

22.2.2 Small Intestinal Organoid Culture

Centrifuge small intestinal crypts, and remove supernatant as much as possible to avoid dilution of Matrigel\textsuperscript{TM} in the next step. Keep the tube at 4 °C and resuspend the crypts pellet in Matrigel\textsuperscript{TM} (200–500 crypts/50 \( \mu \)L matrigel). Pipet 50 \( \mu \)L of the crypt-Matrigel\textsuperscript{TM} suspension into the pre-warmed 24-well plate. The suspension should be applied on the center of the well so a hemispherical droplet can be formed. Transfer the plate back into the CO\(_2\) incubator (5 % CO\(_2\), 37 °C) as soon as possible after the seeding. Allow the Matrigel\textsuperscript{TM} to solidify for 5–10 min and add 500 \( \mu \)L complete organoid culture medium per well. For preparation of the organoid culture medium (e.g. amount of growth factors) check Table 22.1. Place the 24-wells plate in a CO\(_2\)-incubator and refresh the culture medium every 4–6 days.

22.2.3 Small Intestinal Organoid Passage

Crypt organoids can be passaged 7–14 days after seeding. Place the plate with organoids on ice to allow the matrigel to dissolve. Remove the culture medium and gently break up the Matrigel\textsuperscript{TM} by pipetting with a 100–1,000 \( \mu \)L-pipette. Resuspend the organoids in 1–2 mL basal culture medium and transfer them to a 15 mL-tube. Gently disrupt the organoids using a fire-polished Pasteur pipette to remove released dead cells and single cells. Pellet everything by centrifuging at 300\( \times g \) for 5 min, and wash the organoids afterwards with 10 mL basal culture medium. Pellet again at the same speed and add the correct amount of Matrigel\textsuperscript{TM}, namely 50 \( \mu \)L per well that you would like to plate. Count the “crypts/immature organoids” as described before (by hemocytometer) and pipet the Matrigel\textsuperscript{TM} suspension with crypts (50 \( \mu \)L per well) into a new 24 culture plate and place the plate back in CO\(_2\) incubator (5 % CO\(_2\), 37 °C) to culture the organoids.

22.3 Monitoring Viability

Organoid culture and development across time after isolation, and post-passage can be verified using microscopy. Mainly, the dead cells of the organoid tissue will be visible in the central part of the organoid, corresponding to the intestinal lumen.
in vivo (these aggregates of dead cells can be visualized by microscopy as shown in Fig. 22.3 at day 5–7). Figure 22.3 shows a very simplified summary of a mouse intestinal crypt development into a mature organoid structure. Additional conformation of the viability of the organoids can be performed upon by the use of the LDH toxicity assay or the immunofluorescent staining procedures for viability markers (e.g. apoptotic caspase 3 staining), but the organoids have to be sampled destructively in order to perform this assay, thus this step can be only applied at the final stages of specific experiments. It should be noted that once growth, normal cell proliferation and differentiation stops, and organoids start to die, that this will immediately become visible, thus it is essential to monitor the organoid growth and proliferation on a daily basis.

Culture and maintenance of intestinal organoids, relies on the presence and activity of mature (intestinal) stem cells. Even though intestinal organoids can be isolated

| Table 22.1 Concentrations of growth factors and other compounds added to the basal culture medium used for mouse and pig organoid culture |
|---|---|
| Reagent (supplier) | Mousea | Pig |
| Advanced DMEM/F12 (ADF) (Invitrogen, Cat #11320-082) | 1× | 1× |
| HEPES buffer (Sigma-Aldrich, Cat #83264-100ML-F) | 10 mM | 10 mM |
| GlutaMax (Invitrogen, Cat #35050-061) | 2 mM | 2 mM |
| Penicillin/Streptomycin (Invitrogen, Cat #15070-063) | 0.5 U/mL (100×) | 0.5 U/mL (100×) |
| N2 Supplement (Invitrogen, Cat #17502-048) | 1× (stock: 100×) | 1× (stock: 100×) |
| B-27 Supplement Minus Vitamin A (Invitrogen, Cat #12587-010) | 1× (stock: 50×) | 1× (stock: 50×) |
| Mouse Recombinant EGF (Invitrogen, Cat #PMG8041) | 50 ng/mL | 50 ng/mL |
| Mouse Recombinant Noggin (Peprotech, Cat #250-38) | 100 ng/mL | 100 ng/mL |
| Human Recombinant RSPO1 (R&D Systems, Cat #4645-RS-025)b | 500 ng/mL | 500 ng/mL |
| Wnt3a (Cell Guidance Systems, Cat # GFM77) | – | 100 ng/mL |
| Gastrin (Sigma-Aldrich, Cat # SCP0152) | – | 15 nM |
| Nicotinamide (Sigma-Aldrich, Cat # N0636) | – | 10 mM |
| P38 Mapk inhibitor SB202190 (Sigma-Aldrich, Cat # S7067) | – | 10 μM |
| TGFβ type I receptor inhibitor A83-01 (Tocris, Cat # 2939) | – | 600 nM |
| N-Acetylcysteine (Sigma-Aldrich, Cat # A9165-5G) | – | 12.5 mM |
| Primocin (Invitrogen, Cat # ant-pm-1) | – | 100 μg/mL |

For the protocol used for the isolation and cultivation of human intestinal organoids we refer to Sato et al. (2011a, b)

aThe mouse organoid culture protocol was adapted from Sato et al. (2009)

bInstead of using the commercially available RSPO1, one can culture transfected HEK293 cells, which carry a plasmid encoding mRSPO1-Fc (construct developed by Dr. Calvin Kuo at Stanford University, La Jolla, CA). For the protocol for HEK293 RSPO1 production, we refer to Carmon et al. (2011) and the ATCC website: http://www.lgcstandards-atcc.org/en.aspx
and cultured from human induced pluripotent stem cells (iPPS), as described by McCracken et al. (2011), most studies culture organoids from mature stem cells from the intestinal crypt regions. The reason for this is that this method described is relatively elaborate and it takes much longer to mature the intestinal organoids in culture compared to the other method based on the organoid culture from adult intestinal cells. For a detailed protocol of iPPS-based culture of the intestinal tissue we refer to McCracken et al. (2011). As mentioned previously, lineage studies confirmed the stem cell hierarchy in intestinal organoids, either by fast cycling Lgr5+- or slowly cycling Bmi1+-stem cell activity (Sangiorgi and Capecchi 2008; Sato et al. 2009). Stem cell-based organoid cultures are an attractive model due to the presence of the self-renewing capacity; they can be passaged every 5–7 days. In addition, the structure and hierarchy of the mature organoids (after 5–7 days of culturing), highly resembles the in vivo intestinal epithelium (Fig. 22.1). Similar to the intestinal epithelium, stem cells and highly proliferative transit amplifying cells reside in the crypt-like domain of the organoids (Fig. 22.1). Differentiated cell types reside in the villus-region of the organoids and include enterocytes, goblet cells, Paneth cells, and enteroendocrine cells. These cells can be identified using immunohistochemistry with intestinal epithelial cell-specific markers.

22.4 General Advantages and Limitations of Intestinal Organoid Systems

Despite the described significant advances in culturing organoids, this system has some limitations. A major advantage of intestinal models in which an apical to basolateral polarity is preserved is the possibility to characterize epithelial transport mechanisms and barrier integrity upon exposure to microorganisms and microbial components. Indeed intestinal organoids consist of polarised, columnar epithelia. However, the inwards orientation of the epithelia (i.e. villi protrude into the lumen

![Fig. 22.3 Phase contrast images of a developing mouse intestinal crypt into a organoid in culture. (a) immature organoid in culture 0.5–1 day after crypt isolation; (b) mature organoid in culture 3–5 days after crypt isolation; (c) mature organoid in culture 5–14 days after crypt isolation, ready to be passaged (note the accumulated debris of dead cells in the lumen of the organoid). This morphology is representative for other species as well [image adapted from Lukovac et al. (2014)]
of the organoids) makes the apical side relatively inaccessible for direct experimental stimulation (Fig. 22.1).

Also, the organoids lack several essential components of the living digestive tract, such as the enteric nervous system, the vascular system, lymphatic systems and functional adaptive and innate immune systems. Additionally, the 3D architecture is not as regular as seen in vivo, and the crypt-villus structures are variable in size and shape from one organoid to another.

Regardless of these drawbacks, this model offers great experimental utility for understanding and modeling human intestinal development, physiology, and host–microbe interactions. Indeed, this model has several other advantages, such as a long-term culturing, broad range host species and GI tract compartments (stomach, ileum, colon) and gene manipulation possibilities (Koo et al. 2011). Moreover, this system provides a viable starting point for the development of screening platforms for pharmaceutical and food compounds.

Further expansion of this system will lead to broader application range and expansion of the read out possibilities. One example is transition of the existing system towards a structure with the apical/luminal compartment exposed on the outer site of the organoids, which will make the apical site of the organoids experimentally more accessible. Furthermore, this system might be a promising tool in studies on epithelial interactions with food bioactive in individual subjects, as it has been demonstrated that organoids can be cultured from human intestinal biopsy samples (Sato et al. 2011b).

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