Evaluation of Caco-2 and human intestinal epithelial cells as in vitro models of colonic and small intestinal integrity

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1. Introduction

The small intestinal epithelium has two major functions, namely, absorption of nutrients and forming a tight barrier against the potentially harmful components we are exposed to via food. The desire to understand the human intestinal epithelium in more detail has led scientists to mimic the small intestinal microenvironment in vitro models that resemble small intestine physiology [1,2]. Commercially available cell lines derived from the gut, such as Caco-2, HT-29 and T84 cell are immortalized, which facilitates handling of cells under laboratory settings and turns these cell lines into powerful models of intestinal permeability [1,3–5]. These in vitro intestinal systems, such as the Caco-2 cell line, are widely accepted as models to study permeability, physiological processes, and disease mechanisms.

After growth of Caco-2 cells to confluence in vitro, the cells will begin to differentiate and phenotypically resemble enterocytes. Nonetheless, differences between Caco-2 cells and small intestinal epithelial cells are observed as Caco-2 cells lack specific transporter expression and metabolizing enzymes [1,6,7], which emphasizes how Caco-2 cells do not fully represent small intestinal physiology. Furthermore, the cell-cell tightness of the cell line has been reported to be significantly stronger compared to the cell-cell tightness of the small intestine in vivo [8]. This implies a decreased paracellular flux and lower absorption of the Caco-2 cell line epithelium compared to the small intestinal epithelium, in vivo. Also, adult-derived human intestinal epithelial cell (HIEC) monolayers were better at predicting absorption of certain test drugs compared to Caco-2 cells [4,9,10].

The importance of recapitulating major features of the human intestinal epithelium in the best possible way has challenged research groups to develop primary cell lines with characteristics that better resemble the small intestinal physiology compared to the immortalized cell lines. Beaulieu JF isolated and characterized small intestinal HIECs from a 17–19 week old human fetus, called HIEC-6 [7,11]. The HIEC-6 cell line has been thoroughly characterized and, besides the undifferentiated epithelial morphology, it also expresses crypt features and intestinal cell markers such as brush border enzymes and keratins [2,12]. Different research groups have studied this model for regulation of cell proliferation, mechanism of FAS-induced apoptosis, lipid metabolism

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https://doi.org/10.1016/j.bbrep.2022.101314
Received 2 May 2022; Received in revised form 12 July 2022; Accepted 12 July 2022
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and drug absorption [9], however, to the best of our knowledge the epithelial barrier function of HIEC-6 cells has not been well characterized.

A widely used and well-established method to evaluate the integrity and tightness of epithelial monolayers is to measure transepithelial electrical resistance (TEER) across the cell layer. TEER has previously provided insights to the integrity of tight junctions (TJs) between neighboring cells [4,13]. This can be elegantly done via a cellZscope, which is an advanced automated system conducting repeated real-time TEER measurements.

TJs are dynamic structures that are located apically on the epithelial cells creating “kissing points” to a corresponding protein present on the neighboring cell. The TJ proteins represented in the small intestine, Occludin and Claudin, are bound to the cytoskeleton via scaffolding proteins such as Zonula occcludens-1 (ZO-1) and this assembly plays a pivotal role in regulation of barrier function [13-17]. Based on the physiology and functional properties of the gut, these proteins can be localized. For instance, most the claudin family (CLDN 3, 4, 7 and 8) can be found in villus and crypt cells of small intestine [18,19]. Ideally, a combination of methods should be used to evaluate permeability and localize TJ expression when using in vitro models. Immunofluorescence is a technique often used to visualize junction markers intracellularly through fluorophores [20,21]. A mixed approach has already been employed on Caco-2 systems, offering a more complex answer to how the cells are affected by challenges and stressors [22,23].

Likewise, interactions between epithelial cell lines and probiotic bacteria have been extensively studied [22-24] By definition, probiotic bacteria are “live microorganisms which when administered in adequate amounts confer a health benefit on the host” [25]. In specific cases, probiotic bacteria have been shown to strengthen barrier integrity and increase gene expression of TJ-related structures [23,26].

Current knowledge around primary human small intestine cell models and their application for assessing barrier function is limited which encouraged us to investigate TEER and TJ expression in HIEC-6 cells as a potentially alternative model. Additionally, host-microbe interaction was studied by evaluating the junction strengthening effect of well-established probiotic bacteria.

2. Material and methods

2.1. Cell culture

HIEC-6 cells (ATCC-CRL 3266) were maintained on OptiMEM 1 Reduced Serum (Gibco 31985) and supplemented with 4% fetal bovine serum (FBS, Gibco 10270), 20 mM HEPES (1 M, Thermofisher CAT no. 15630080), 10 mM GlutaMAX (Gibco 35050), 10 ng/mL Epidermal Growth Factor (EGF, Sigma SRP3027) reconstructed with 5% Trehalose (Sigma T0167); Penicillin-Streptomycin solution (10000 units Penicillin G Sodium Salt and 10 mg/mL streptomycin sulfate) (Sigma, 69-57-8, 3810-74-0) hereafter referred as complete OptiMEM. Cells were grown in T175 cm² plastic culture flasks (Thermo Scientific 15991) in a humidified atmosphere 37 °C with 5% CO₂. Cells were used as passage number 5 to 15 for the assay.

2.2. Bacteria culture

Lactcaseibacillus rhamnosus GG (formerly known as Lactobacillus rhamnosus GG, hereafter referenced by the use of the trademark LGG®), Bifidobacterium animalis subsp. lactis, BB-12, hereafter referenced by the use of the trademark BB-12® and Lactcaseibacillus paracasei subsp. paracasei CRL 431 (formerly known as Lactobacillus paracasei subsp. paracasei CRL 431) and hereafter referenced by the use of the trademark L.CASEI 431®) were supplied by Chr. Hansen A/S, Denmark. Lactcaseibacillus strains were inoculated from frozen stock and grown overnight at 37 °C in Man-Rogosa-Sharpe (MRS) medium pH 6.5, whereas BB-12% was cultured anaerobically in anaerobic boxes with Anaerogen™ pads (Thermo Scientific, AN0035A) at 37 °C in MRS supplemented with 0.05% cysteine hydrochloride monohydrate (CyHCl, CAS 7048-04). For TEER studies, 10-fold dilution series up to 1 × 10⁻⁷ were grown overnight at 37 °C.

After overnight incubation, bacterial cultures at late exponential/early stationary phase were selected based on their optical density at 600 nm wavelength (OD600). Bacterial cultures were centrifugated at 3500 g for 10 min at 19 °C, and the resulting pellets were washed in Hank’s Balanced Salt Solution (HBSS; Gibco, 14175) at 37 °C and suspended in the same media. Washing was repeated twice with a final resuspension in antibiotic-free complete OptiMEM or DMEM for HIEC-6 and Caco-2 cells, respectively. Subsequently, bacterial suspensions were adjusted to OD600 of 4.0.

2.3. Transepithelial electrical resistance (TEER)

Three TEER experiments were performed to characterize barrier function and expression of TJ proteins in HIEC-6 and Caco-2 cells and to investigate the interaction with lactobacilli and bifidobacteria.

In the first experiment, the aim was to investigate if the TEER readouts varied following different number of days in culture. Briefly, upon HIEC-6 and Caco-2 cells reached 80% confluence were treated with TrypLE Express Enzyme (Gibco, 12604) and seeded apically at 1 × 10⁴ cells/well and 1 × 10⁵ cells/well onto 12-well, 12 mm Transwell® with 0.4 µm pore polyester membrane insert (Corning; 3460) respectively; as previously described by Tatenaka et al. 2014. Growth media was renewed every second day with complete OptiMEM and DMEM, correspondingly. A TEER assay was performed after 6, 8, 10 and 12 days in culture for the HIEC-6 cells whereas Caco-2 were grown for 15, 18, 21 and 24 days in culture. The day prior to the TEER experiment, transwell inserts were transferred to the CellZscope2 (Nanoanalytics, Munster, Germany) and kept in similar conditions as aforementioned HIEC-6. Cell passage numbers ranging from 9 and 20 were used for the experiments.
baseline TEER in each well and served as a baseline and a quality control of a stable electrical resistance. After hourly overnight readings, the CellZscope2 was paused and 100 μL of apical medium was replaced with 100 μL of OD adjusted bacterial cell suspension or ABx alone. The final OD₀₄₅₀ of the bacterial suspensions was 0.5. Each condition was measured in triplicate. The CellZscope2 was transferred back to the incubator and readings were resumed for a total of 18 h. Changes in TEER during bacterial stimulation were calculated relative to the latest measurement were set to 100%). Area under curve (AUC) was calculated for each condition.

2.4. Immunofluorescence of tight junctions (occludin, Zonula occludens-1 and Claudin-1)

Following TEER measurements of experiment 2 and 3, the HIEC-6 and Caco-2 transwells were transferred to 12-well plates containing Phosphate Buffer Solution (PBS, Sigma A806552). Each transwell was carefully washed once with PBS and the cell membranes were fixed with 4% formaldehyde (VWR 1.004.968.350) for 6 min at room temperature (RT) and washed three times in PBS. Treated membranes were then removed from the transwells using a scalpel and transferred into a 48 well-plate (Costar 3548) with PBS. Subsequently, cell permeabilization was ensured by treating the membranes with 0.2% Triton X-100 (Sigma A9934) for 5 min at RT followed by a washing cycle with PBS. Blocking was done by treating the membranes for 1 h with 2% Bovine Serum Albumin (BSA, Sigma A5882). Primary antibodies were diluted in PBS as described in Table 1 and the treated membranes were incubated overnight at 4 °C on a platform rocker (14 inclination with 5 s intervals) (Mimetas, Organo-flow L-191000059). The membranes were washed three times with 0.25% BSA and 0.1% Triton X-100 in PBS (washing buffer) and then incubated for 2 h in the dark at RT with secondary antibodies (diluted in washing buffer according to descriptions in Table 1). Following incubation with the antibodies, the slides were washed twice with PBS and labelled with nuclei staining; 4’,6-diamidino-2-phenylindole (DAPI, Life Technologies D1306), by incubation for 5 min, protected from light. The slides were washed three times with deionized water. Upon immunofluorescence protocol completion, membranes were transferred onto 18 mm slides (VWR deionized water). Upon immunofluorescence protocol completion, membranes were transferred onto 18 mm slides (VWR European 631–1550), mounted with fluorescence mounting medium (Agilent S3023) and coverslips (Karl Hecht GmbH, 4100109). Staining was visualized via a fluorescence microscope EVOS FL Auto 2D (Invi- 

3. Statistics

The impact of days after seeding and seeding number were tested statistically by using the one-way ANOVA test on the 18 h timepoints for Caco-2 and HIEC-6 cells, separately. In the co-incubation experiment, cell cultures exposed to bacterial suspension were compared to control wells via the one-way ANOVA test. Statistics were calculated using the software GraphPad Prism (v. 9.3.1 GraphPad Software, LLC.)

4. Results

4.1. Transepithelial electrical resistance (TEER)

In order to determine the best incubation period to achieve a tight cellular monolayer, HIEC-6 cells were incubated for 6, 8, 10, 12 days on transwells at a seeding density of 10⁴ cells/well whereas Caco-2 cells were seeded at 10⁵ cell/well grown for 15, 18, 21 and 24 days on transwells after seeding. As shown in Fig. 1, TEER measurements were not significantly affected by different incubation times in either of the two cell lines. Although not significantly different, Caco-2 cells showed higher TEER mean values after 18 days and 21 days of incubation compared to 15 days and 24 days in culture. Initial peaks indicate cells adapting to the CellZscope2 and incubator’s atmosphere during the first measurements. The low TEER measurements of the HIEC-6 monolayer led us to further investigate whether seeding densities could affect the TEER readings.

Since few studies have been conducted using HIEC-6 cells, the incubation period was extended until day 15 and two TEER measurements of 18 h each were performed at incubation day 12 and 15. In order to standardize HIEC-6 protocol and determine best seeding concentration to obtain a tight monolayer, we performed a dose-dependent seeding experiment. Cells were seeded on transwells using 1 × 10⁵ to 1 × 10⁷ cells/well. Interestingly, none of the seeding densities increased TEER values regardless of the number of days in culture (Fig. 2). Even though the higher densities of seeded cells ranging from 1 × 10⁴ to 1 × 10⁷ reached confluency, TEER values remained comparable to control conditions (transwells without cells) and to the less confluent monolayers (Fig. 2). No significant differences in TEER readings were found between cell densities or incubation days compared to controls. TEER mean values were between 32.24 ± 2.8 Ω × cm² and 31.5 ± 5.01 Ω × cm² on day 12 and 15 respectively, which were similar to TEER measurements from the initial experiment (Fig. 1).

4.2. Tight junction identification

The aim was to further understand the lack of increase of TEER with increased seeding cell number and prolonged incubation. Therefore, the tight junction expression pattern was evaluated according to cell seeding density after 18 days in cell culture.

Occludin and Zonula Occludens-1 (ZO-1) are extensively investigat- jacketed junctional complexes [15,27–29], therefore these proteins were selected in order to be localized within HIEC-6 cells. ZO-1 was expressed along the cell membrane as seen in Fig. 3 regardless of the cell density. Occludin, on the other hand, was not detected in any of the membranes (data not shown).

Moreover, based on the bioimaging results, variations in the morphological and structural features of HIEC-6 cells were observed. At the highest density (1 × 10⁵), cell silhouettes became geometrical rather than preserving their elegant, elongated structure which was observed at lower densities (≤10⁴) (Fig. 3), nonetheless, TJ localization remained unaffected and was evenly distributed. Despite cells reaching confluency after 18 incubation days, cellular gaps were still present in the mono- 

4.3. Claudin-1 expression

Because of the limited capacity of HIEC-6 cells to express the Occludin protein, we aimed to identify a transmembrane junction strand. Claudins have been demonstrated to constitute part of TJ [15] and multiple family members have been identified; i.e, claudin-1/2/3/4/7, however, in this study only Claudin-1 was examined.

Table 1

| Antibody/protein | Function | Dilution | Product ID |
|------------------|----------|----------|------------|
| Occludin         | Tight junction marker Primary antibody | 1:100 | Thermofisher 61-7300 |
| Zonula Occludens-1 | Tight junction marker Primary antibody | 1:200 | All Diagnostics Sc-133256 |
| Claudin-1 (A-9)  | Barrier function & TJ Primary antibody | 1:100 | Santa Cruz Biotechnology Sc-166338 |
| Goat anti-mouse, Alexa flour 488 | Secondary antibody | 1:1000 | Life Technologies A-11001 |
| Donkey anti-rabbit, Alexa flour 555 | Secondary antibody | 1:1000 | ThermoScientific A31572 |
As observed in Fig. 5, Claudin-1 was uniformly identified in HIEC-6 cells at both seeding densities tested ($1 \times 10^2$ and $1 \times 10^7$). The immunofluorescence staining was an appreciable strand along the cells, indicating that this protein is expressed within the cytoplasm.

4.4. Bacterial co-incubation

Caco-2 and HIEC response to bacterial co-incubation was also explored. Three different bacterial strains all representing industrially...
produced probiotic strains and representing different species were used to explore TEER response and TJ expression. Fig. 6 shows LGG®, L. CASEI 431® and BB-12® co-incubated with Caco-2 and HIEC-6, respectively. LGG® and L.CASEI 431® both induced high TEER readings compared to baseline, whereas BB-12® only showed limited effect on TEER when co-incubated with Caco-2 cells. In HIEC cells, no response to either of the bacteria was observed, this is consistent with the previous finding that showed confluent cells that were not tight enough to generate a significant TEER reading.

In Fig. 7 staining of the HIEC-6 and Caco-2 cell membranes are shown after bacterial co-incubation. In the Caco-2 cells, ZO-1 showed a consistent spiderweb-like staining pattern on the cell periphery that did not change upon bacterial exposure. In contrast, ZO-1 staining in HIEC-6 appeared different according to which bacterium was used for co-incubation compared to control, in which labeling appears uniformly within the cytoplasm. Upon LGG® exposure, ZO-1 staining was localized intercellularly especially close to the cell periphery. Interestingly, HIEC-6 showed a change in morphology to an elongated structure and more intense homogenous fluorescence after BB-12® co-incubation compared to control. No changes were observed on HIEC-6 co-cultured with L. CASEI 431® compared to control.

5. Discussion

Previous investigation of HIEC monolayers indicate superior drug absorption prediction [30] and better physiological representation of small intestinal conditions compared to Caco-2 cells [9]. In this study, we evaluated HIEC-6 cells as a model of intestinal barrier function. Previously, Perreault and Beaulieu [31] showed ZO-1 expression and desmosome protein ZK-31 in this cell line, yet its permeability and transepithelial barrier capacity has not been fully investigated. Our barrier integrity experiments did not indicate any increased TEER when using HIEC-6 cells as monolayers compared to control (no cells), regardless of high cell density and a broad range of incubation days. Our TEER readings for the HIEC cells were similar to the TEER readings for the control with no cells, strongly indicating that measurements correspond to resistance across the transwells and not across the cell layer itself. In the publication by Takenaka et al. [9], the HIEC cells were also seeded at a density of $1 \times 10^5$ cells/well on 24-well fibrillar collagen–coated inserts and at a density of $6.3 \times 10^4$ cells/well on 12-well noncoated membrane inserts, and TEER was measured daily. After day 5, TEER readings reached a plateau at approximately $100 \Omega \times cm^2$ that lasted at least until day 11. We were not able to reproduce the tight monolayer formed by the HIEC cells and the increase of TEER as previously found [9]; even though we did prolong growth and differentiation phase on the membranes and increase the number of cells seeded. In support of this, probiotic bacteria were not able to induce an increase in TEER readings. In vivo, the human small intestine has lower TEER compared to the colon whereas ex vivo measurements using the Ussing chamber showed values of $(45 \pm 21 \Omega \times cm^2)$, $(34 \pm 12 \Omega \times cm^2)$ and $(37 \pm 4.36 \Omega \times cm^2)$ for duodenum, jejunum and ileum, respectively, however not all of the samples were derived from healthy subjects [32]. This suggests a less tight epithelial layer allowing for increased passage of macromolecules. In the HIEC cells of human embryonic origin, the missing TEER development across the cell membrane and the notorious holes observed via immunofluorescence suggest an immature epithelial cell line. This is in line with the impaired gut barrier function observed during early life in many mammalian species, which allows for passive transmission of immunoglobulins from the mother’s milk [33], for...

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**Fig. 3.** Zonula Occludens-1 (red) and nuclei (blue) identification by immunofluorescence at different HIEC-6 cell seeding densities after 18 days of incubation. Magnification 10×. Scale bar is 275 µm. Control is cell-free Transwell insert with OptiMEM media only and the negative control (secondary antibody alone) did not exhibit any fluorescence (data not shown). (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

**Fig. 4.** 18-days-old HIEC-6 monolayer at $10^7$ cell concentration. White arrow indicate inter-cellular gap. Magnification 40×. Scale bar 75 µm.
example. Interestingly, the HIEC cells seemed to differentiate and showed an epithelial-like phenotype after 18 days in culture seeded at a density of $1 \times 10^7$ (Fig. 5) compared to cells seeded at a lower density, indicating that cell-cell contact may lead to physiological cell changes.

To better understand HIEC-6 transepithelial resistance, we used an immunofluorescence technique on TJ proteins. TJ features have been identified on Caco-2 cells [34] and Occludin and ZO-1 are well established junction proteins that have been localized on these colonic cells [35,36]. In contrast, little is known about TJ expression in HIEC cells. Our fluorescence results indicate that this cell line does not express Occludin, which could explain the lack of TEER since this protein had been thought of as essential part for anchoring adjacent cells [16,17]. However, studies on Occludin$^{-/-}$ mice showed that its absence did not impair barrier function [37,38] whereas it did cause alterations on paracellular flux mainly in gastric mucosa [39]. In this study, HIECs expressed ZO-1, which is in line with previous findings in which ZO-1 was expressed in differentiated epithelial cells from Occludin-deficient embryonic stem cells, and thus ZO-1 seems essential to form TJ in embryonic tissue [15,27]. ZO-1 is responsible for anchoring Occludin and Claudin to Actin in the cytoskeleton [16] through the PDZ-domain, and this finding led to the identification of the Claudin family as one of the backbones of the TJ networks [27,37,40].

Due to the absence of Occludin expression in the HIEC cells and the desire to better understand the TJ composition in these cells, we investigated Claudin-1 protein expression in HIEC-6. TJ assembly begins basolaterally with ZO-1 binding adherens junctions henceforth recruiting primordial junctions ($\alpha$ and $\beta$ catenins) as junctions continue to mature. ZO-1 then migrates upwards until maturation and Occludin interacts with junctional adhesion molecules-A, culminating in accumulation of Claudin apically [15,37]. Although fetal small intestinal cells at 10–12 weeks of gestation have been reported to express most of the differentiated features observed in cells from adults [11]. HIEC cells represent cryptcells that are of a proliferative and undifferentiated nature [7]. In the current study, we substantiate these findings by underlining that these cells are not at a maturation stage in which they can generate a TEER across the cell layer and a tight paracellular monolayer.

In contrast, Caco-2 monolayers, originating from colonic tissue, are widely used as a model of intestinal integrity and seem to generate higher TEER readings, although large interexperimental variation exist due to the different methodologies used, such as chopsticks, CellZscope, Millicell-ERS system and EVOM Endohm [4,9,32,41].

Interestingly, previous studies have shown that barrier function may be significantly compromised or strengthened by co-incubation with bacteria, bacterial components or metabolites. Several pathogens have shown detrimental effects on barrier function by direct invasion of the epithelial cells and disruption of the TJs, as observed by Salmonella typhimurium and enterotoxigenic Escherichia coli (ETEC) [42].

Probiotic bacteria, including lactobacilli and bifidobacteria have been acknowledged to affect barrier function via various mechanisms. For instance, the cell wall polysaccharide fraction of B.breve has shown anti-ulcer and anti-erosion effects in gastric models of acetic acid and

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**Fig. 5.** Immunofluorescence staining of HIEC-6 cells for Zonula occludin-1, Claudin-1 and nuclei when seeded in high density; $10^7$ cells/well (top row) and low density; $10^2$ cells/well (bottom row). Magnification 20×. Scale bar 125 μm.

**Fig. 6.** TEER readings normalized to baseline reading immediately before adding bacteria; LGG®; BB-12®; L.CASEI 431® and control (media only) for A) Caco-2 cells and B) HIEC-6 cells.
ethanol-induced erosions [43]. Furthermore, *B. breve* has been associated with expression of Tad pili that stimulate colonic epithelial cell proliferation [44] that may support barrier function further. Additionally, secreted molecules including metabolites such as short chain fatty acids and indoles acting via the Aryl Hydrocarbon receptor [24, 45] improve gut barrier function. The LGG® strain was reported to secrete peptides; p40 and p75, responsible for strengthening the barrier integrity and protect against H2O2-induced damage [23, 46]. In this study, three probiotic bacteria were assessed as co-culture with Caco-2 cells, and all three showed increased TEER measurements compared to control, although to various degrees. LGG® was previously reported to induce increases in TEER when using fresh cultures [41], freeze dried product [47] and when present in a yogurt matrix [48]. On the other hand, BB-12® generated lower TEER readings compared to LGG® and L.CASEI 431® which has been previously reported [41]. This could be due to lower survival rate of BB-12® compared to the *Lactobacillus* strains that are generally more aerotolerant.

6. Conclusions

The data from this study suggest that despite HIECs being of non-immortalized gut epithelium origin, the cells do not form a confluent monolayer and therefore transepithelial electrical resistance cannot be measured across the cell layer regardless if the cells are stimulated with probiotic bacteria. The absence of Occludin in the HIECs supports the more immature profile compared to Caco-2 cells. In conclusion, HIECs do not seem suitable as a model for mimicking gut barrier function in vitro.

Declarations of competing interest

Both authors Silvia Lopez-Escalera and Anja Wellejus are employed at Chr. Hansen A/S.

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Acknowledgement

We thank Dr. Jane Thompson for support and proofreading this manuscript. This project has received funding from the European Union’s Horizon 2020 research and innovation program under Marie Skłodowska-Curie grant agreement No 813781. All probiotic strains tested in this study; L.CASEI 431®, LGG® and BB-12® are trademarks of Chr. Hansen A/S.
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