Capsaicin and Piperine as Functional Excipients for Improved Drug Delivery across Nasal Epithelial Models

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ABSTRACT

The fruit from various pepper plants has been employed for the seasoning of food, as perfuming agents, and also as traditional medicines. Phytochemicals isolated from different pepper species have been found to modulate the pharmacokinetics of orally administered drugs. This study investigated the possibility to apply capsaicin and piperine (extracted alkaloids) as modulators for drug delivery across the nasal epithelium. Both a nasal epithelial cell line (RPMI 2650) and excised sheep nasal tissue were used as models to investigate the effects of the selected pepper compounds on drug permeation. FITC-dextran 4400 (MW 4400 Da) was used as a large molecular weight marker compound for paracellular transport, while rhodamine 123 was used as a marker compound that is a substrate for P-glycoprotein-mediated efflux. From the permeation results, it was clear that capsaicin inhibited P-glycoprotein efflux to a larger extent, while piperine showed drug permeation enhancement via other mechanisms. The cell cytotoxicity studies indicated that capsaicin was noncytotoxic up to a concentration of 200 µM and piperine up to a concentration of 500 µM as indicated by cell viability above 80%. The histological analysis of the excised nasal tissue and cultured RPMI 2650 cell layers indicated that some damage occurred after treatment with 200 µM capsaicin, but no changes were observed for piperine up to a concentration of 50 µM.

ABBREVIATIONS

ALI	air-liquid interface
FD4	FITC-dextran 4400
KRB	Krebs-Ringer bicarbonate buffer
LCC	liquid-covered culture
MEM	modified Eagle's medium
P _{app}	apparent permeability coefficient
P-gp	P-glycoprotein
PTP	polyethylene terepthalate
R123	rhodamine 123
TEER	transepithelial electrical resistance
TRPV1	transient receptor potential vanilloid 1

Introduction

Chili pepper (family *Solanaceae*) and black pepper (family *Piperaceae*) are annual herbs grown in warm climates. The production thereof is globally regarded as one of the most important economical and agricultural activities in developed and developing countries. The fruits of these plants are widely used in the flavoring of cuisine, especially in Asia and South America, and have also found applications in the cosmetic industry and for medicinal uses [1, 2].

Capsaicin (**>** Fig. 1A) is one of the main capsaicinoids uniquely present in the fruit of plants from the genus *Capsicum*. These plants have been known for their medicinal value, with Mayan pharmacopoeia reporting the application of chili peppers in respiratory and bowel ailments, relief of earaches, and in the treatment of wounds [3]. It is now also known that capsaicin is able to inhibit melanoma proliferation, induce apoptosis in human mammary glands, and is capable of inhibiting P-gp efflux transporters in the

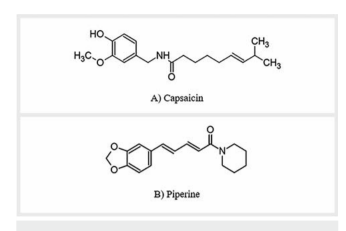


Fig. 1 Chemical structures of (A) capsaicin (MW = 305.41 g/mol) and (B) piperine (MW = 285.34 g/mol).

Caco-2 (human colon adenocarcinoma) and KB-C2 (human papillomavirus-related endocervical adenocarcinoma) cell lines [4]. In addition, it was proven that capsaicin presents with analgesic, antimicrobial, and antioxidant effects [1, 3].

Black pepper (*Piper nigrum* L.) and long pepper (*Piper longum* L.) belong to the genus *Piper* for which piperine (▶ **Fig. 1B**) is the major alkaloid and is responsible for the pungent taste of the fruits of both. Piperine has long been used as a traditional medicine, preservative, and insecticide in addition to the more commonly known seasoning and perfuming applications. Piperine has also been proven to be effective in improving the pharmaco-kinetics of poorly bioavailable drugs and other medicinal herbs through inhibiting P-gp and CYP450 metabolism in intestinal epithelial cells [2, 5–7].

The effects of capsaicin and piperine on the permeation of compounds have been tested in different cell models (i.e., Caco-2 and MDCK II cells) for potentially improved delivery of drug compounds, but they have not yet been explored for improved nasal delivery of drug compounds. Nasal delivery of drugs holds certain benefits for the systemic delivery of drugs, including a rich blood supply and an epithelium that is relatively more permeable in comparison with other areas in the body, and it circumvents first-pass metabolism. Furthermore, with a relatively large surface area (150 cm²) and volume (15–20 mL), the nasal cavity provides an alternative route for self-administration of poor orally bioavail-able drug products that would have otherwise been administered via the parenteral route [8–10].

In the current study, the effects of capsaicin and piperine were investigated on the permeation of two model compounds in two nasal epithelial models. The RPMI 2650 cell line is currently the only commercially available human nasal epithelial cell line. When grown under the correct ALI conditions, RPMI 2650 cells have been found to express P-gp, multidrug resistance proteins, and tight junctions. Furthermore, even though the model presents with multilayered cells, the TEER and permeability properties are comparable to that of the human nasal mucosa [8, 11, 12]. The excised sheep nasal mucosa model has been proven to be well suited for permeation and metabolism studies related to the nasal delivery of compounds. This model is more cost-effective than comparable *in vitro* and *in vivo* models, and sheep nasal tissue is easily accessible from abattoirs. The sheep nasal model has been found to closely resemble nasal absorption in humans [13]. In addition, comparable TEER values have been reported between sheep and human nasal epithelium, and the sheep nasal tissue also expresses P-gp efflux transporters [14, 15].

Results and Discussion

Validation results of the analysis by fluorescent spectroscopy of marker compounds used in the permeability studies are shown in **Table 1**. R123 was used as a marker for efflux-mediated transport, while FD4 was used as a macromolecular marker for paracellular transport.

Linearity and purity of the two pepper extracts, capsaicin and piperine, that were evaluated/screened for possible pharmacokinetic interactions were determined using HPLC analysis (**> Table 2**). Linearity was determined using reference standards of each compound, whereas purity is expressed as a percentage of raw material in relation to the reference standards. The concentrations of the reference standard were 250 µg/mL for capsaicin and 150 µg/mL for piperine.

The P_{app} values of FD4 across RPMI 2650 epithelial cell layers in the apical-to-basolateral direction are shown in **Fig. 2** in the absence (FD4 alone, control group) and presence of different concentrations of capsaicin (50, 100, and 200 μ M) and piperine (10, 25, and 50 μ M).

From > Fig. 2, it is clear that capsaicin increased the FD4 permeation (Papp) across RPMI 2650 nasal epithelial cell layers at all the concentrations applied, which was to some extent inversely proportional to the capsaicin concentration. However, the permeation of FD4 was statistically significantly higher than the control (FD4 alone) only at the lowest concentration of capsaicin (50 µM) applied. In general, these findings in the nasal epithelial cell line are in coherence with previous findings obtained for capsaicin across different epithelial models such as the Caco-2 cell model [16, 17] and the MDCK II model [18, 19]. In both these previous sets of studies, it was found that capsaicin reduced TEER and increased paracellular permeation of different compounds by means of tight junction modulation. It was suggested that capsaicin can open tight junctions via a unique and reversible dual-action mechanism. Firstly, capsaicin caused an influx of Ca^{2+} into the cells, which led to cofilin dephosphorylation that directly induced depolymerization of F-actin leading to the opening of tight junctions [1, 16]. Secondly, the total concentration of occludin expression was reduced, while a redistribution of actin at the bicellular tight junctions was seen [17, 19]. These two mechanisms seemed to occur independently from one another, but both contributed to the increased permeability through the paracellular transport pathway.

In this study, capsaicin had a negligible effect on the TEER (data not shown) of the RPMI 2650 nasal epithelial cell layers. This phenomenon may be explained by the fact that RPMI 2650 cells grow in multilayers (please refer to histology below) and not in monolayers like other epithelial cell models (e.g., Caco-2 and MDCK II cells). The study by Shiobara et al. [19] concluded that capsaicin acted differently on actin at bicellular tight junctions than at tri-

Table 1 Fluorescent spectroscopy validation parameters or	of marker compounds used for permeation studies.
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	R123			FD4	FD4		
Linearity	$r^2 = 0.999$	$r^2 = 0.999$			r ² = 0.999		
Limit of detection	0.002 µM	0.002 µM			0.005 μg/mL		
Limit of quantification	0.005 µM	0.005 μM			0.016 µg/mL		
	5 µM	5 μΜ 2.5 μΜ 0.125 μΜ			62.5 µg/mL	12.5 µg/mL	
Accuracy	98.21%	98.14%	116.64%	95.77%	98.27%	104.67%	
Intraday precision (%RSD)	4.97%	2.37%	6.45%	2.77%	3.69%	2.63%	
71 (7							

Table 2 Linearity and purity results of the selected pepper extracts (i.e., capsaicin and piperine) as determined by HPLC.

	Capsaicin	Piperine
Linearity	r ² = 0.999	r ² = 0.999
Purity	91.82%	97.62%

cellular tight junctions. In a multilayer model, such as the RPMI 2650 cells grown on insert membranes, there will be a larger amount of tricellular tight junctions than bicellular tight junctions as compared to that found in monolayer cell models. A probable explanation for the inverse proportion in FD4 permeation as a function of capsaicin concentration across the multilayered RPMI 2650 cells could be attributed to an overall increase in tricellular tight junctions found in the model. Due to this, a saturation of the interactions between the capsaicin and bicellular tight junctions may have already been reached at the lowest concentration (50 μ M) applied and higher concentrations therefore did not cause any higher permeation enhancement effect.

▶ Fig. 2 indicates that piperine increased FD4 permeation pronouncedly across the RPMI cell layers, albeit not statistically significantly (p > 0.05). The effect by piperine on FD4 permeation across the RPMI 2650 cell layers did not increase with an increase in piperine concentration over the concentration range investigated (10-50 µM). It was previously proposed that piperine led to polarized paracellular opening of the blood-brain barrier [20] and increased drug absorption by fluidizing the brush border membrane while increasing microvilli length [21]. Furthermore, TEER measurements during the current permeation studies indicated a 8.99, 8.57, and 18.69% decrease in the presence of 10, 25, and 50 µM piperine, respectively. Since piperine only had a slight decreasing effect on TEER of the RPMI nasal epithelial cell layers, it may therefore have increased FD4 permeation across the RPMI 2650 nasal epithelial cell layers, most probably by a combination of the mechanisms previously proposed. Similar to that observed with capsaicin, piperine also exhibited the maximum effect at the lowest concentration applied and this effect did not increase further with an increase in concentration. Once again, this may be explained by a saturation of the interaction between piperine and the tight junctions at the lowest concentration applied.

The P_{app} values of the efflux substrate, R123, across RPMI 2650 nasal epithelial cell layers in the apical-to-basolateral (absorptive) and basolateral-to-apical (secretory) directions are shown in **Fig. 3**. The P_{app} values for R123 are shown in the absence (R123 alone, control group) and presence of different concentrations of capsaicin (50, 100, and 200 μ M) as well as piperine (10, 25, and 50 μ M).

Capsaicin increased the permeation of R123 across nasal epithelial RPMI 2650 cell layers in the absorptive direction compared to the control in a concentration-dependent manner, which was statistically significant ($p \le 0.05$) at the highest concentration (200 µM) applied (> Fig. 3). Although the secretory permeation was slightly higher than that of the control group, it decreased with an increase in capsaicin concentration and the effect is clearly visible in the efflux ratio values (> Table 3). The slightly higher permeation in the secretory direction may be explained by other mechanisms, but this was overshadowed by efflux inhibition at higher concentrations of capsaicin. From the permeation results obtained for both FD4 and R123 in the presence of capsaicin, it seems that capsaicin exhibited a larger effect on efflux-related permeation than paracellular permeation. This efflux inhibition effect of capsaicin on nasal epithelial cells correlates well with the efflux inhibition exhibited by capsaicin in other in vitro epithelial cell models such as Caco-2 cells [3] and KB-C2 cells [22]. Additionally, it has also been proven with in vivo models [23].

In the presence of piperine, both absorptive and secretory permeation of R123 increased in a concentration-dependent manner together with a decrease in the efflux ratio value at the highest concentration applied (**> Table 3**). It was previously shown in the intestinal epithelium that piperine caused efflux inhibition when applied together with a P-gp transporter substrate [6, 24, 25]. However, the results of this study indicated that piperine had a larger modulatory effect on paracellular permeation than on efflux inhibition across the nasal epithelial cells (although efflux inhibition was not explicitly ruled out as seen by the reduced efflux ratio value at the highest concentration). Since R123 is also able to diffuse paracellularly [26], the potential opening of tight junctions by the coapplied piperine correlated well with the increase in P_{app} values of R123.

The effect of capsaicin and piperine on the permeation of model compounds was also evaluated across excised sheep nasal tissues. The P_{app} values of FD4 across excised sheep nasal epithelial tissues in the apical-to-basolateral direction are shown in **Fig. 4** in the absence (FD4 alone, control group) and presence of differ-

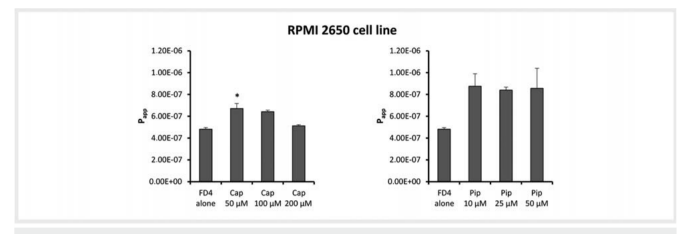


Fig. 2 P_{app} values of FD4 in the absence (control) and presence of different concentrations of capsaicin (Cap) and piperine (Pip) across RPMI 2650 nasal epithelial cell layers in the absorptive direction (* $p \le 0.05$, Kruskal-Wallis test followed by Dunn's post hoc test).

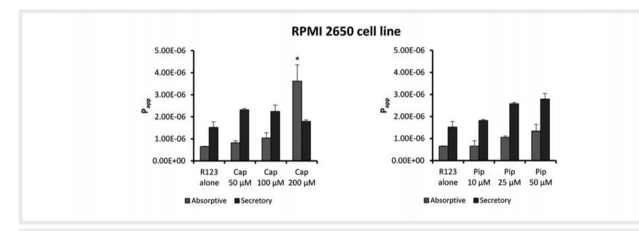


Fig. 3 P_{app} values of R123 in the absence (control) and presence of different concentrations of capsaicin (Cap) and piperine (Pip) across RPMI 2650 nasal epithelial cell layers in the absorptive and secretory directions (*p < 0.05, Kruskal-Wallis test followed by Dunn's post hoc test).

Table 3 Efflux ratio values of R123 in the absence (R123 alone, control) and presence of different concentrations of capsaicin and piperine across RPMI 2650 nasal epithelial cell layers.

	R123 alone	Capsaicin			Piperine		
		50 µM	100 µM	200 µM	10 µM	25 µM	50 µM
Efflux ratio value	2.33	2.83	2.17	0.50	2.75	2.44	2.08

ent concentrations of capsaicin (50, 100, and 200 $\mu M)$ and piperine (10, 25, and 50 $\mu M).$

Capsaicin mediated a slight increase in the permeation of FD4 across the excised nasal epithelial tissues at the concentrations applied in this study, as shown in **Fig. 4**, and no statistically significant differences were evident. This indicated that the tight junction modulation effect of capsaicin to increase the paracellular permeation of FD4, a macromolecular model compound, was much lower in the excised sheep nasal tissue model compared to the nasal epithelial RPMI cell model (**Fig. 2**). Piperine increased FD4 permeation in the absorptive direction to a larger extent than capsaicin, but it was still not statistically significant. The less pro-

nounced effect of the selected pepper extracts on FD4 permeation across the excised sheep nasal tissue as compared to the nasal epithelial RPMI 2650 cell layers may be explained by the differences in the inherent properties of the two models. In contrast to nasal epithelial cell models (e.g., RPMI 2650), animal excised tissues have ciliated cells and goblet cells, and mucus can be present [15]. Mucus plays an important role in the permeation of compounds, especially large molecules such as FD4, across biological membranes.

Both capsaicin and piperine caused pronounced concentration-dependent increases in the absorptive permeation of R123 and mediated a reduction in R123 permeation in the secretory

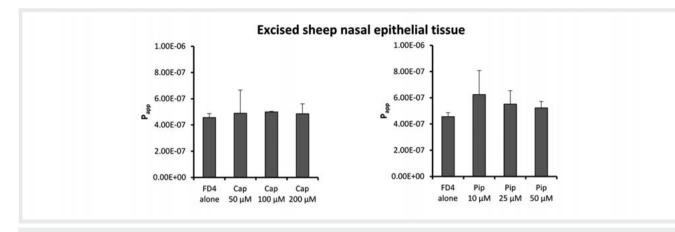


Fig. 4 P_{app} values for FD4 in the absence (FD4 alone, control) and presence of different concentrations of capsaicin (Cap) and piperine (Pip) across excised sheep nasal epithelial tissue in the absorptive direction.

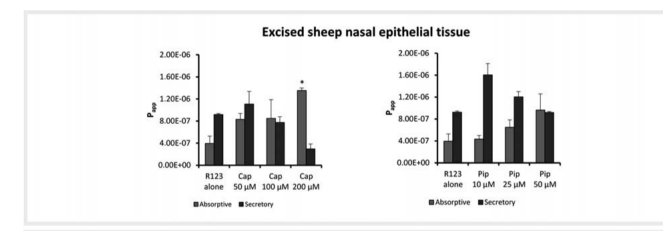


Fig. 5 P_{app} values of R123 in the presence of capsaicin (Cap) and piperine (Pip) across sheep nasal mucosa in absorptive and secretory directions (*p \leq 0.05, Kruskal-Wallis test followed by Dunn's post hoc test).

Table 4 Efflux ratios of R123 in the absence (control) and presence of different concentrations of capsaicin and piperine across excised sheep nasal epithelial tissue.

	R123 alone	Capsaicin			Piperine		
		50 µM	100 µM	200 µM	10 µM	25 µM	50 µM
Efflux ratio	s 2.34	1.33	0.91	0.22	3.70	1.86	0.95

direction across excised sheep nasal epithelial tissues (**>** Fig. 5). When the experimental transport values are compared to that of the controls it is evident that the efflux ratio values had decreased (**>** Table 4). These results clearly show that both capsaicin and piperine exhibit P-gp-related efflux inhibitory properties in excised nasal epithelial mucosa, which means that the absorption of efflux transporter substrates can be improved by coadministration of these pepper extract components. The improved absorption of efflux transporter substrates in the presence of capsaicin and piperine suggests that these substrates can be administered via the nasal route at lower concentrations and still attain similar blood plasma levels.

It has been shown previously that piperine has the ability to inhibit CYP3A4 metabolism, which has been proven in a variety of *in vitro* and *in vivo* models [2,6,25]. This has also been shown for capsaicin in an *in vivo* model [23]. The metabolism inhibition effects of these pepper extracts should also be investigated after nasal administration in future studies to elucidate if the metabolism inhibitory effects contribute to their drug absorption enhancing effects.

MTT assays were employed on the RPMI 2650 cell line to evaluate the cytotoxicity effects of capsaicin and piperine on nasal epithelial cells. This is an important consideration in order to establish if the selected pepper extract components have cytotoxic-

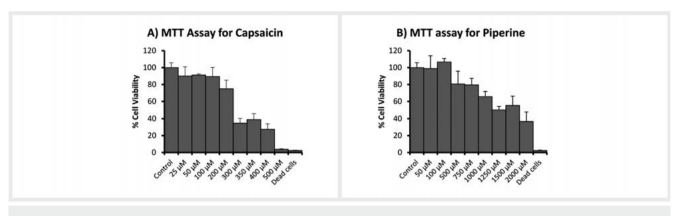


Fig. 6 Percentage cell viability as determined with an MTT assay on RPMI 2650 cells after 24 h exposure to a range of (A) capsaicin and (B) piperine concentrations.

ity effects that can cause damage at the concentrations where they are effective as drug absorption enhancers across nasal epithelial surfaces.

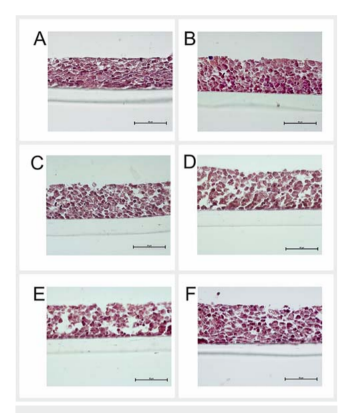
According to the ISO 10993-5:2009 standards determined for "Tests for in vitro cytotoxicity" [27], from > Fig. 6A, one can see that capsaicin is not cytotoxic to RPMI 2650 cells up to a concentration of 100 µM (cell viability above 80%), weakly cytotoxic at a concentration of 200 µM (cell viability between 60-80%), and highly cytotoxic from 300 µM and above (cell viability below 40%). Piperine, on the other hand (> Fig. 6B), is considered noncytotoxic to RPMI 2650 cells up to a concentration of 750 µM. The difference in toxicity between the two pepper extract components can potentially be related to the pungency of the extracts measured in Scoville heat units. The Scoville heat unit scale is used to depict the relative pungency of a pepper. Pure capsaicin possesses a pungency of 16×10⁶ Scoville heat units [1], which is much higher than that of piperine at 2×10^5 Scoville heat units [5]. Furthermore, capsaicin and piperine are both proven agonists of the TRPV1 ion channel, which is responsible for the transmittance of multiple noxious stimuli. It is known that the pungency (i.e., the total Scoville heat units) of TRPV1 agonists is directly proportionate to the activation of the receptor [28, 29]. Caterina et al. [28] found that capsaicin kills both neuronal and non-neuronal cells that express vanilloid receptors through cytoplasmic swelling, an amalgamation of the contents with eventual lysis due to increased receptor activation.

Histological analysis of both RPMI 2650 cell layers grown with the ALI technique and excised sheep nasal epithelial tissues were conducted in the absence (control) and presence of capsaicin and piperine to establish whether any physical changes occurred in the cells or tissues during the permeation experiments.

The histological investigation revealed that RPMI 2650 cells cultured under ALI conditions on porous membranes formed multilayered epithelial surfaces without differentiation into different types of cells that are found in the normal nasal epithelial mucosa. There is no evidence of columnar cells with any specialized modifications, such as goblet and ciliated cells. All cells have a round or slightly elliptical shape. Nuclei are dense without heterochromatin and euchromatin segregation, which is usually associated with the absence of secretory activity. Indeed, no mucus secretion was observed, which was confirmed by the absence of Alcian Blue staining. On some areas of the RPMI epithelial cell layers, spaces are visible between the cells, resulting in a less dense aggregation of cells. Exposure to KRB for 180 min as well as 50 µM capsaicin and 50 µM piperine did not show any signs of adverse effects on the cell layers in any way (► Fig. 7B, C, F). Signs of cell loosening or detachment started to appear after exposure to 100 µM capsaicin for 180 min (► Fig. 7D), while exposure to 200 µM capsaicin for 180 min resulted in damage to the cell layers, evidenced by the formation of sinusoids (► Fig. 7E), which is in coherence with MTT results.

The excised sheep nasal epithelial tissues presented as pseudostratified ciliated columnar epithelium with mucus-producing goblet cells present. Epitheliocytes have an elongated shape with cilia on the apical side (free surface) of the tissue. Nuclei are loose with visible nucleoli that indicate an active synthesis process in the tissue cells. Underneath the ciliated epithelial layer, connective tissue with serous glands and blood vessels are visible (> Fig. 8A). After exposure to KRB, all the epithelial tissue structures remained intact with no obvious signs of tissue damage (> Fig. 8B). Furthermore, the underlying tissue layers including the connective tissue, blood vessels, and serous glands did not show signs of necrosis, edema, or hemorrhage after 180 min exposure to $200\,\mu\text{M}$ capsaicin and 50 µM piperine. However, the epithelial layer was altered after exposure to $200\,\mu\text{M}$ capsaicin with excessive cell loss (> Fig. 8C). The columnar cell layer appears to be separated from the rest of the tissue distal to the germ layer. Incubation of sheep nasal tissue with piperine 50 µM did not have any adverse effects on the excised sheep nasal epithelium and all structures appeared to be the same as in the intact control tissue (> Fig. 8 D).

A histometrical analysis was conducted on both the RPMI 2650 and excised sheep nasal epithelial models, and the values of the parameters that were measured are shown in **Table 5**. The thickness of the whole excised sheep mucosal tissues ranged between 553.4 ± 65.8 and 683.6 ± 336.1 μ m, while the thickness of the sheep nasal epithelial layer alone was 52.9 ± 4.4 μ m. This compares well with the RPMI 2650 cell layers, where the epithelial thickness ranged between 47.7 ± 2.5 and 58.5 ± 4.34 μ m. The epithelial thickness and average epitheliocyte nuclei areas were comparable between the intact control and after piperine expo-



▶ Fig. 7 Representative histological sections of RPMI 2650 cell layers grown at ALI. A Intact RPMI 2650 cell layer before permeation experiment. B RPMI 2650 cell layer after 180 min exposure to KRB, i.e., in the absence of capsaicin and piperine (control). C RPMI 2650 cell layer after 180 min exposure to 50 µM capsaicin. D RPMI 2650 cell layer after 180 min exposure to 100 µM capsaicin. E RPMI 2650 cell layer after 180 min exposure to 200 µM capsaicin. F RPMI 2650 cell layer after 180 min exposure to 50 µM piperine. Scale bar = 50 µm.

sure for both the excised sheep tissue and RPMI 2650 cell models. However, the thickness of the epithelial layer and the nucleus area of epitheliocytes was dramatically decreased after exposure to capsaicin.

Materials and Methods

Cell lines and chemicals

The RPMI 2650 (nasal septum carcinoma) cell line was purchased from the ATCC. FITC-dextran (MW 4400, FD4), R123, capsaicin (purity > 91% by HPLC against a reference standard), piperine (purity > 97% by HPLC against a reference standard), hematoxylin-eosin, and 1% Alcian Blue were purchased from Sigma-Aldrich. Krebs-Ringer bicarbonate buffer was prepared according to Sigma-Aldrich's product information sheet [30]. PBS, MEM, and FBS were purchased from Thermo Fischer, while L-glutamine, Pen-Strep, nonessential amino acids, and trypsin-EDTA (0.25% w/v) were purchased from Whitehead Scientific. MTT, hematoxylin solution (GHS132), Eosin Y solution, and aqueous (HT110216) and Alcian Blue solution (B8438) were purchased from Sigma-Aldrich. ThinCert 12-well plates (catalogue number 665110) and PTPcoated inserts (catalogue number 665641) were purchased from Separation Scientific. Sheep (Merino, Dorper or Île-de-France subject to availability) nasal tissue was collected from a local abattoir in Potchefstroom, South Africa.

Chemical characterization of capsaicin and piperine raw materials

The purity of the purchased piperine and capsaicin raw materials was determined by means of liquid chromatography analysis using an Agilent 1100 series HPLC equipped with a gradient pump, autosampler, UV detector, and OpenLab CDS Chemstation Rev. C.01.07 SR3 data acquisition and analysis software (Agilent Technologies). HPLC conditions were adapted from the literature and were as follows: USP L1, Venusil XBP C18 column, 150×4.5 mm, $5 \mu m$ (Agela Technologies) with the mobile phase as ACN/H₂O with 0.1% H₃PO₄ 60:40 at a flow rate of 1.0 mL/min for both capsaicin and piperine. Both injection volumes were set at 10 µL per sample. For capsaicin, the detection wavelength was set at 225 nm and it eluted at a retention time of \pm 5.28 min [31], whereas a wavelength of 325 nm was used for piperine with a retention time of \pm 5.18 min [5].

The linearity of capsaicin and piperine reference standards were determined first followed by a purity analysis. The purity of the raw materials was determined against the reference standard for each extract using Equation 1 where the reference standard concentrations were $250 \,\mu$ g/mL for capsaicin and $150 \,\mu$ g/mL for piperine.

% Purity of pepper ectract $=\frac{\text{experimental concentration}}{\text{reference standard concentration}} \times 100$ (Eq. 1)

Fluorescence spectroscopic analysis of FITC-dextran and rhodamine 123

Fluorescence spectroscopic analysis was used to determine FD4 and R123 in the permeation samples and the analyses were done using a Spectramax Paradigm (serial nr. 33270–1142) multimode detection platform plate reader. FD4 was analyzed with excitation and emission wavelengths set at 485 nm and 525 nm [10, 32], respectively, while R123 was analyzed using 480 nm excitation and 520 nm emission wavelengths [33, 34]. The fluorescence analytical methods were validated for linearity, accuracy, and precision.

Cell culturing

RPMI 2650 cell layers were cultured on ThinCert insert membranes by means of the ALI technique using previously published methods [8, 11, 35] with some slight modifications.

Briefly, RPMI 2650 cells were first grown in culture flasks using MEM fortified with 10% FBS and 1% each of PenStrep, nonessential amino acids, and L-glutamine. Cells were maintained in a humidified 5% CO₂ atmosphere at 37 °C with media changes every 2 days. After confluency of approximately 90% was reached in the culture flasks, cells were washed twice with 10 mL PBS and then trypsinized with 0.25% (w/v) trypsin-EDTA for 5 min in an incubator. Cells were then counted using a hemocytometer and seeded into PTP-coated ThinCert inserts (113.1 mm² culture surface, 0.4 µm pore size, $2 \times 10^6 \cdot cm^{-2}$ pore density) at a density of

 6×10^5 cells/cm². Seeding was done with cells between the passages of 25 and 47.

Culturing in the 12-well ThinCert culture plates took place for a total of 21 days with media changes every 2 days. Cells were cultured under LCC conditions for the first 2 days, allowing cells to adhere to the membrane. LCC conditions entail media being present at both the apical (0.8 mL) and basolateral (approximately 4 mL) side of the membrane. After 2 days, during the first media change, cells were lifted to ALI conditions. ALI conditions entail media (approximately 4 mL) to only be added to the basolateral side of the membrane for a period of 19 days. After a total of 21 days, the inserts were carefully transferred to 12-well ThinCert transport plates, the TEER across each cell layer on insert membrane was measured, and permeation studies were then initiated.

Collection and excision of sheep nasal epithelial tissue

Approval from the North-West University's animal ethics committee (AnimCare) was obtained for the use of excised sheep nasal tissue in the permeation experiments (ethics approval certificate nr. NWU-00285-17-A5). The removal of sheep nasal mucosal tissue was accomplished via a method that was adapted from previously published methods for excision of nasal epithelial tissues from different animals [15, 36–38].

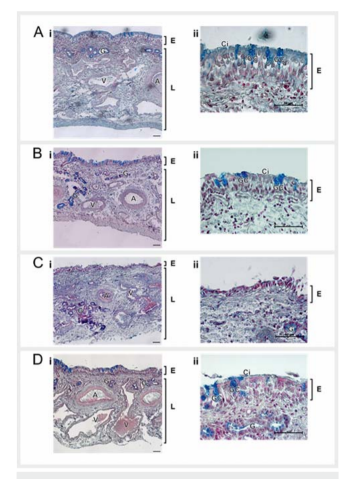
After slaughter, the anterior portion of the skull (i.e., the snout) was removed via a longitudinal incision along the frontal plane, anterior to the eyes. The skin was then removed from the snout via dissection, and the snout specimen was rinsed and sub-merged in ice-cold KRB and transported to the laboratory. In the laboratory, a vertical incision was made along the septal midline in order to remove the septum. After separation into two halves, a vertical incision was made through the lateral wall of the snout, just posterior to the *incisura nasoincisiva* in order to isolate the nasal conchae. Hereafter, the nasal epithelial tissue layer was separated from the conchae cartilage via blunt dissection.

The pieces of epithelium were laid out on moistened filter paper in order to facilitate easier handling and to prevent damage to the epithelial cells. Strips of tissue were then cut in widths of approximately 1 cm each and mounted in Sweetana-Grass diffusion chambers after filter paper removal.

Ex vivo permeation studies

All permeation studies of R123 and FD4 in the absence (control groups) and presence of capsaicin or piperine were done in both of the nasal epithelial models (i.e., RPMI 2650 cell layers grown on insert membranes and excised sheep nasal tissues mounted in Sweetana Grass diffusion chambers) in triplicate.

For the RPMI 2650 cell model, succeeding the 21 days in culture, permeation studies were slightly adapted from previously published methods [8,10,12]. Firstly, the PTP-coated ThinCert inserts were transferred from 12-well culture plates to 12-well transport plates while maintaining sterile conditions. A volume of 0.8 mL preheated (37° C) KRB was added to the apical compartment and 1.8 mL KRB to the basolateral compartment. The 12-well transport plates were then left to equilibrate for 15 min in an incubator. For permeation studies in the absorptive (apical-to-basolateral) direction, 0.8 mL of the selected marker compound (i.e., either FD4 or R123) in the presence of the selected pepper



▶ Fig. 8 Representative histological sections of excised sheep nasal epithelial tissues with column i indicating the whole excised tissue and column ii only the epithelial layer at a higher magnification. A Intact excised sheep nasal epithelial tissue before permeation experiment. B Excised sheep nasal epithelial tissue after 180 min exposure to KRB. C Excised sheep nasal epithelial tissue after 180 min exposure to 200 µM capsaicin. D Excised sheep nasal epithelial tissue after 180 min exposure to 180 µM capsaicin. D Excised sheep nasal epithelial tissue after 180 min exposure to 50 µM piperine. Legend: A – arteries, Ci – cilia, E – epithelial layer, G – serous glands, GC – goblet cells, L – *lamina propria*, V – veins. Scale bar = 50 µm.

extract component (i.e., either capsaicin or piperine) suspended in KRB was applied and incubated on the apical side of the membrane. Samples of $180 \,\mu$ L were withdrawn from the basolateral side at the time intervals of 5, 10, 15, 30, 60, 90, 120, and 180 min and immediately replaced with an equal volume of fresh KRB after each withdrawal. Similarly, for permeation studies in the secretory (basolateral-to-apical) direction, 1.8 mL test solution was applied and incubated on the basolateral side. Samples of 180 μ L were withdrawn from the apical side at the time intervals of 5, 10, 15, 30, 60, 90, 120, and 180 min and immediately replaced with an equal volume of fresh KRB after each withdrawal.

Permeation studies across excised sheep nasal epithelial tissue in the Sweetana-Grass diffusion chambers were done in a similar fashion. After assembly of the diffusion chambers, 7 mL preheated (34 °C) KRB were incubated at both sides of the membrane, connected to a heating block and carbogen (95% O_2 :5%

	Intact control	Capsaicin, 200 µM	Piperine, 50 µM		
	Excised sheep tissue				
Excised mucosa thickness (µm)	683.6 ± 336.1	656.3 ± 93.1	553.4 ± 65.8		
Epithelial thickness (µm)	52.9 ± 4.4	18.8 ± 7.4*	47.5 ± 9.1		
Average epitheliocyte nucleus area (µm²)	36.1 ± 6.6	22.2 ± 8.2*	35.1 ± 7.3		
	RPMI 2650 cell layers grown with ALI technique				
Epithelial thickness (µm)	53.3 ± 8.1	47.7 ± 2.5*	58.5 ± 4.34		
Average epitheliocyte nuclei area (µm²)	14.3 ± 7.2	10.8 ± 3.7*	13.1 ± 5.4		

► **Table 5** Parameters of epithelia of two different nasal tissue models (*statistically significant differences compared to the control group with p ≤ 0.01).

CO₂) supply to equilibrate for 15 min. Permeation studies in the absorptive direction contained 7 mL test solution in the apical chamber and 7 mL KRB in the basolateral chamber, while for secretory permeation studies, the 7 mL test solution was applied to the basolateral chamber and 7 mL KRB in the apical chamber. Samples were withdrawn at the same volume and time intervals as described for the permeation studies on the RPMI 2650 cell model.

Following analysis of samples using a Spectramax Paradigm plate reader for either FD4 or R123 concentrations, the percentage transport (Eq. 2) and the P_{app} values were calculated (Eq. 3).

$$\% \text{ Transport } = \frac{\text{Conc. value at specific time}}{\text{Conc. value of donor solution}} \times 100 \quad \text{(Eq.2)}$$

$$P_{app} = \frac{dQ}{dt} \left(\frac{1}{A.60.C_0} \right)$$
(Eq.3)

Where P_{app} is the apparent permeability coefficient (cm × s⁻¹), $\frac{dQ}{dt}$ is the permeability rate per minute, A is the diffusion area of the membrane (cm²), and C₀ is the initial concentration of each marker compound used.

The efflux ratio of the P-gp substrate, R123, was also calculated in order to determine the extent to which peppers are able to modulate efflux proteins (Eq. 4).

$$ER = \frac{P_{app} (secretory)}{P_{app} (absorptive)}$$
(Eq.4)

Cytotoxicity and histology

An MTT assay was performed to determine cytotoxicity on RPMI 2650 cells based on previously published [9, 39] methods. Cells were seeded in a clear bottom 96-well plate at a density of 3×10^5 cells/cm² and left to attach for 24 h. Both capsaicin (25–500 µM) and piperine (50–2000 µM) were prepared in MEM, added to the cells after attachment, and left to incubate at $37 \,^{\circ}$ C in a 5% CO₂ humidified atmosphere for 24 h. Following drug exposure, the cells were washed twice with 100 µL PBS. Thereafter, 180 µL non-additive MEM were added to the cells, followed by 20 µL of 5 mg/mL MTT in PBS. Plates were then returned to the incubator for 4 h. Following the assay, the media was aspirated from all wells and crystals were dissolved in 200 µL DMSO. The

plates were then placed on an orbital shaker for 1 h and absorbance was measured at two wavelengths, 560 and 630 nm, for cell signal and background noise, respectively, using a Spectramax Paradigm[®] plate reader. All experiments were done in triplicate.

Histological analysis was conducted to determine the effects of the pepper extract components on both the RPMI 2650 cell layers and the excised sheep nasal epithelial tissues following exposure during the permeation studies in triplicate. Excised sheep nasal tissue samples before permeation (intact control) and after the 3 h permeation studies were fixed in 10% buffered formalin at + 4 °C for 24 h. RPMI 2650 ALI cell layers on ThinCert insert membranes before permeation (intact control) and after the 3 h permeation studies were washed with PBS and fixed in 10% buffered formalin at room temperature for 1 h. The samples were then dehydrated in a graded series of ethanol and embedded in paraffin, sliced into 4-5 µm sections, stained with hematoxylin-eosin and 1% Alcian Blue, and then examined under a Nikon E800 compound microscope using 60× objective. The sheep mucosa thickness, epithelial thickness, and nucleic area of the epitheliocytes were measured from five different sections per sample using NIS-Elements. Version 4.05 software.

Statistical analysis

Statistical analysis was performed using Statistica v. 13.3 (StatSoft, Inc.). Data are presented as the mean \pm SD. Data analysis of the P_{app} values for all test groups and in both models as well as histological parameters was performed using Kruskal-Wallis tests for comparison of multiple groups followed by Dunn's test for comparison of two groups. P \leq 0.05 was considered to represent a statistically significant difference.

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Conflict of Interest

The authors declare no conflict of interest.

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