Suppression of Trichothecene-Mediated Immune Response by the *Fusarium* Secondary Metabolite Butenolide in Human Colon Epithelial Cells

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Butenolide (BUT, 4-acetamido-4-hydroxy-2-butenoc acid gamma-lactone) is a secondary metabolite produced by several *Fusarium* species and is co-produced with the major trichothecene mycotoxin deoxynivalenol (DON) on cereal grains throughout the world. BUT has low acute toxicity and only very limited occurrence and exposure data are available. The intestinal epithelium represents the first physiological barrier against food contaminants. We aimed to elucidate the intestinal inflammatory response of the human, non-cancer epithelial HCEC-1CT cells to BUT and to characterize potential combinatory interactions with co-occurring trichothecenes, such as DON and NX-3. Using a reporter gene approach, BUT (≥ 5 µM, 20 h) was found to decrease lipopolysaccharide (LPS; 10 ng/mL) induced nuclear factor kappa B (NF-κB) activation in a dose-dependent manner, and in combinatory treatments BUT represses trichothecene-induced enhancement of this important inflammatory pathway. Analysis of transcription and secretion levels of NF-κB-dependent, pro-inflammatory cytokines, revealed a significant down-regulation of IL-1β, IL-6, and TNF-α in IL-1β-stimulated (25 ng/mL) HCEC-1CT cells after BUT exposure (10 µM). Trichothecene-induced expression of pro-inflammatory cytokines by the presence of 1 µM DON or NX-3 was substantially suppressed in the presence of 10 µM BUT. The emerging mycotoxin BUT has the ability to suppress NF-κB-induced intestinal inflammatory response mechanisms and to modulate substantially the immune responsiveness of HCEC-1CT cells after trichothecene treatment. Our results suggest that BUT, present in naturally occurring mixtures of *Fusarium* fungal metabolites, should be increasingly monitored, and the mechanism of inhibition of NF-κB that might affect the pathogenesis or progression of intestinal inflammatory disorders, should be further investigated.

Keywords: mycotoxin, deoxynivalenol, NF-κB, intestinal inflammation, combinatory effects, HCEC-1CT, DON, NX-3
INTRODUCTION

Mycotoxins are toxic secondary metabolites of low-molecular-weight produced by a wide diversity of filamentous fungi. Contaminating crops pre- or post-harvest, mycotoxins may enter the food and feed chains and pose a risk to human and animal health (1). As mycotoxin-producing fungi, such as Fusarium species, are able to generate simultaneously a wide variety of secondary metabolites, co-exposure to mixtures of toxic fungal compounds is evident and the evaluation of potential toxicological interactions is therefore of great importance for comprehensive risk assessment.

Butenolide (BUT, 4-acetamido-4-hydroxy-2-butenoic acid gamma-lactone, Figure 1A) is a secondary metabolite produced by various Fusarium species, such as F. sporotrichioides and F. graminearum. It was already isolated from F. tricinctum-colonized tall fescue grass in the 1960s, causing a livestock mycotoxicosis called “fescue foot” (2–4). Epidemiologically, BUT is considered one of the mycotoxins implicated in the etiopathogenesis of Kashin-Beck disease, an endemic, chronic and degenerative osteoarthropathy and of Keshan disease, an endemic cardiomyopathy, in China (5–7). In regard to occurrence and exposure, scientific data are lacking. Abia et al. (8) reported BUT in co-occurrence with trichothecene mycotoxins in up to 100% of the tested maize samples, resulting in mean contamination levels of 131 µg/kg, respectively. A multi-mycotoxin analysis of Norwegian grain samples showed median BUT concentrations of about 200 µg/kg, whereas the highest concentration of 3,370 µg/kg was detected in oats (9). Further studies revealed BUT contamination levels at mean and maximum concentrations of 299 and 798 µg/kg in naturally Fusarium contaminated oat samples (10), as well as 23 and 1,490 µg/kg in feed and feed raw materials, respectively.

 Abbreviations: ACTB1, β-actin; BUT, butenolide; COX-2, cyclooxygenase-2; d, days; DON, deoxynivalenol; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; HKLM, heat killed Listeria monocytogenes; HO-1, heme oxygenase; IL-1α, interleukin-1α; IL-1β, interleukin-1β; IL-6, interleukin-6; IL-8, interleukin-8; iNOS, inducible nitric oxide synthase; JNK/SAPK, c-Jun N-terminal kinases/stress-activated kinases; LC-UV, liquid chromatography-ultraviolet; LPS, lipopolysaccharide; MAPKS, mitogen-activated protein kinases; MCP-1, monocyte chemotactic protein 1; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; Nrf2/ARE, nuclear factor E2-related factor 2/antioxidant responsive element; qRT-PCR, qualitative reverse transcription polymerase chain reaction; ROS, reactive oxygen species; RT, room temperature; SE, standard error; SRB, sulforhodamine B; TNF-α, tumor necrosis factor-alpha.

FIGURE 1 | Chemical structures of the investigated secondary metabolites of Fusarium: (A) BUT, (B) DON and (C) NX-3.
(24–26). Regarding non-transformed, intestinal epithelial cells, IPEC-J2 porcine cells showed an upregulation of interleukin-1α (IL-1α), IL-1β, interleukin-6 (IL-6), IL-8, TNF-α and monocyte chemotactic protein 1 (MCP-1) after DON-exposure (2 μM, 48 h), whereas at 0.5 μM only IL-1β, IL-6 and IL-8 were induced (27). However, the modulatory effects of DON on the inflammatory response of human, non-cancer, colonic epithelial cells, especially focusing on the important inflammatory signaling pathway NF-κB, have not been assessed yet.

Whereas, DON can be considered one of the most studied mycotoxins worldwide, only little is known about the recently discovered type A trichothecene NX-3 (Figure 1C). NX-3, structurally identified as an analog of DON, lacking the carbonyl moiety at position C8, represents an in planta de-acetylation product of the *Fusarium* secondary metabolite NX-2, which is produced by an emerging population of *F. graminearum* in North America known as so-called “Northland”- or N-strains (28, 29). A first toxicological characterization revealed NX-3 to inhibit in vitro plant- and mammalian-derived ribosomes to a similar extent as DON (28, 30). Furthermore, both trichothecenes caused comparable cytotoxic effects in different mammalian cancer and non-cancer cell lines (12, 30). Comparing the impact of NX-3 and DON on the onset of oxidative stress in HepG2 cells, only marginal differences were identified at the selected end-points (12). As DON has been reported to alter inflammatory signaling processes in human intestinal epithelial cells (21), similar effects can be expected after exposure to the novel *Fusarium* mycotoxin NX-3. The assessment of specific effects in this regard might be crucial for hazard characterization and have not been assessed so far.

BUT, considered as moderately toxic, is rarely monitored and not regulated by any food safety authority, despite the fact that it can co-occur in high concentrations with DON (9). Therefore, we have assessed already in a previous study the ability of BUT to contribute to the toxicity of mixtures with trichothecene mycotoxins in HepG2 cells (12). So far, DON is well-characterized with respect to its pro-inflammatory properties and several in vitro studies have already investigated the interaction of DON with other known mycotoxins on the immune response. However, a potential immunomodulatory contribution of less- or non-toxic co-occurring secondary metabolites, such as BUT, has not been taken into account so far and is the focus of the present study. In a first step, using a reporter gene approach, we assessed the impact of BUT, DON, NX-3 and respective combinations on the inflammatory key signaling pathway NF-κB. As occurrence studies detected BUT and DON in various molar ratios (between 0.06 and 72.8 BUT/DON), equimolar as well as 10-fold molar excess of BUT were assessed in this study trying to mimic different exposure scenarios. Furthermore, we analyzed mRNA expression and protein secretion levels of four pro-inflammatory cytokines performing qRT-PCR and Magnetic LumineX® experiments, respectively, in order to obtain a more detailed characterization of the inflammatory response of the applied non-cancer intestinal cell model HCEC-1CT.

**MATERIALS AND METHODS**

**Chemicals and Reagents**

BUT was synthesized and purified according to Burkhardt et al. (31), as described previously by Woelflingseder et al. (12). DON was purchased from Romer Labs (Tulln, Austria). NX-3 was produced and purified as deacetylation product of NX-2 using preparative high-performance liquid chromatography (purity >99% according to LC-UV at 200 nm) as published by Varga et al. (28). All substances were dissolved in water (LC-MS grade) to obtain stock solutions of 10 mM, which were aliquoted and stored at −20°C. Cell culture media, supplements and materials were purchased from GIBCO Invitrogen (Karlsruhe, Germany), Sigma-Aldrich (Munich, Germany), and Sarstedt AG & Co (Nuembrecht, Germany).

**Cell Culture and Treatment**

The human monocytic cell line THP1-Lucia™ NF-κB, derived from the human THP-1 monocyte cell line by stable integration of an NF-κB inducible Luciferase reporter construct, was purchased from InvivoGen (San Diego, USA). The non-tumorigenic human colon epithelial cell line HCEC-1CT™ (32, 33) was kindly provided by Prof. Jerry W. Shay (UT Southwestern Medical Center, Dallas, TX, USA). THP1-Lucia™ were cultured in RPMI 1640 medium, supplemented with heat-inactivated 10% (v/v) fetal bovine serum and 1% (v/v) penicillin/streptomycin (100 U/mL). THP1-Lucia™ were treated alternately with zeocin and normocin (100 μg/mL; Invivogen, USA). HCEC-1CT™ cells were cultivated in Dulbecco’s Modified Eagle’s Medium (high glucose) supplemented with 2% (v/v) Medium 199 (10X), 2% (v/v) GE Healthcare™ HyClone™ Cosmic Calf™ Serum, 20 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, 50 μg/mL gentamicin, insulin-transferrin-selenium-G (10 μg/mL; 5.5 μg/mL; 6.7 ng/mL), recombinant human epidermal growth factor (18.7 ng/mL) and hydrocortisone (1 μg/mL). Both cell lines were maintained in humidified incubators at 37°C and 5% CO₂ and sub-cultured every 3–4 d. Cells were routinely tested for the absence of mycoplasma contamination and used for experiments at passages 12–18. BUT, DON, NX-3 and their combinations were added to the incubation solutions, resulting in a final solvent concentration of 2% (v/v) water (LC-MS grade). In order to ensure data comparability, combinatory treatments were always performed in parallel to the individual substances using the same passage of cells on the same microtiter-plate.

**NF-κB Reporter Gene Assay in THP1-Lucia™ Cells**

BUT, DON, NX-3 and 10:1 and 1:1 combinations of BUT with the trichothecenes were prepared in reaction tubes to be further diluted 1:100 by the addition of the THP1-Lucia™ cell suspension. 10:1 and 1:1 ratios were selected due to the high variability of secondary metabolite concentrations in the available occurrence studies (8, 9, 34) and due to previously performed cell viability experiments in Woelflingseder et al. (12). Cellular concentration was determined by trypan blue.
exclusion, cells were centrifuged for 5 min at 250 × g and resuspended at a concentration of 1 × 10^6 cells/mL in fresh, pre-warmed growth medium. Cell suspension was added to the respective toxin preparations into the reaction tubes. They were gently mixed and transferred as technical duplicates into a 96-well plate (100 µL/well). After 2 h of incubation at 37°C and 5% CO₂ in the humidified incubator, LPS (1 µL/well) at a final concentration of 10 ng/mL was added to the cells and incubated further for 18 h. 2% (v/v) water (LC-MS grade) with and without LPS treatment were used as solvent control. As recommended by the supplier, heat killed *Listeria monocytogenes* (HKLM) at a concentration of 20 × 10^6 cells/well were applied as positive control for TLR activity. Following in total 20 h of incubation, well plate was centrifuged (250 × g, 5 min) and 10 µL/well of the supernatant were collected. Subsequently, the reporter gene activity assay (secreted luciferase) was performed according to the manufacturer's protocol. QUANTI-Luc™ solution (InvivoGen, US), containing the recommended coelenterazine substrate, was added for luminescence measurements.

In parallel cellular metabolic activity was monitored by the alamarBlue® assay (for details see 2.6). Therefore, after supernatant collection for the measurements of luciferase activity, 10 µL of alamarBlue® reagent (Thermo Fisher Scientific, Massachusetts, USA) were added to the cells and incubated for 2 h. Subsequently, after another centrifugation of the plate (250 × g, 5 min) 70 µL/well were transferred to a black 96-well plate and fluorescence intensity was measured at 530/560 nm (excitation/emission). Both luciferase activity and fluorescence intensity measurements were performed on a Synergy™ H1 hybrid multi-mode reader (BioTek, Bad Friedrichshall, Germany) assessing at least five independent experiments in technical duplicates. Modulation of NF-κB-dependent luciferase expression was referred to the respective THP1-Lucia™ cell viability determined as alamarBlue®-fluorescence intensity.

### Quantitative Analysis of Cytokine Gene Transcription in HCEC-1CT Cells

In HCEC-1CT cells gene transcription of the four cytokines TNF-α, IL-1β, IL-8, and IL-6 was analyzed by quantitative reverse transcription polymerase chain reaction (qRT-PCR). Cells were seeded in 12-well plates (35,000 cells/well) and allowed to grow for 48 h. Preliminary experiments revealed insufficient pro-inflammatory stimulation of HCEC-1CT cells by LPS-treatment. Thus, IL-1β (25 ng/mL), generally applied as inflammatory trigger in Caco-2 cells (35, 36), was used in all HCEC-1CT cell experiments as inflammatory stimulus. Cells were incubated with BUT, DON, NX-3 and respective combinations in total for 5 h, consisting of 2 h of toxin incubation, followed by IL-1β co-treatment (25 ng/mL) for additional 3 h. Subsequently, total RNA was extracted using Maxwell® 16 Cell LEV Total RNA Purification Kits (Promega, US) and reversed transcribed into complementary DNA (cDNA) by QuantiTect® Reverse Transcription Kit (Qiagen) according to the supplier's protocols. Afterwards, cDNA samples were amplified in technical duplicates in presence of gene specific primers (QuantiTect® Primer Assays, Qiagen) and QuantiTect® SYBR Green Master Mix (Qiagen) using a StepOnePlus® System (Applied Biosystems, Foster City, USA). Therefore, the following primer assays purchased from Qiagen (Hilden, Germany) were used: β-actin (ACTB1, Hs_ACTB_1_SG, QT00095431); glyceraldehyde 3-phosphate dehydrogenase (GAPDH, Hs_GAPDH_1_SG, QT00079247); TNF-α (Hs_TNF_1_SG; QT00029162); IL-1β (Hs_IL1B_1_SG, QT00021385); IL-8 (Hs_CXCL8_1_SG, QT00003322); IL-6 (Hs_IL6_1_SG, QT00083720). Qiagen does not provide detailed information on the sequences of the QuantiTect® Primer Assays. However, information on the approximate location of primers can be viewed on the Product Detail pages of the Qiagen GeneGlobe database. A universal PCR protocol was applied including enzyme activation at 95°C for 15 min, 40 cycles of 15 s at 94°C, 30 s at 55°C and 30 s at 72°C. In order to check for primer specificity melting curve analysis was performed at the end of every PCR experiment. StepOnePlus® software (version 2.3, Applied Biosystems, USA) was used for fluorescence signal quantification and further data analysis. Of each tested sample at least five biologically independent experiments were performed in technical duplicates. Presented transcript data was normalized to the mean of transcript levels of endogenous control genes (ACTB1, GAPDH) applying the ΔΔCt-method (37) for relative quantification.

### Profiling of Pro-inflammatory Cytokine Secretion

Cytokines released from HCEC-1CT cells into cell culture supernatants after 5 h of incubation were measured using a Magnetic Luminex® assay (R&D SYSTEMS, Minneapolis, USA). Cell culture supernatant samples were collected after the incubation for cytokine gene transcription (details for incubation see 2.4.) and stored until the measurement at −80°C. After 2-fold sample dilution for TNF-α analysis and 40-fold dilution for the analysis of IL-8 and IL-6, the assay was performed according to the manufacturer's protocol. Briefly, in a 96-well plate 50 µL of standard or sample were mixed with 50 µL microparticle cocktail per well. After 2 h of incubation at room temperature (RT), plate was washed three times with 100 µL wash buffer per well. Afterwards, samples were incubated with 50 µL per well of a biotin-antibody cocktail for 1 h at RT. Again, three washing steps were performed prior to 30 min incubation with 50 µL per well of streptavidin-PE solution. Finally, samples were rinsed again three times with wash buffer and resuspended in 100 µL per well prior to the read-out on a Bio-Plex® 200 multiplex array reader (Bio-Rad Laboratories, Hercules, USA). Using the Bio-Plex Manager™ software a standard curve was created for each analyte by performing a five-parameter logistic curve-fit, average blank-values were subtracted and respective cytokine concentrations were calculated. The assay was performed in technical duplicates, analyzing at least three independent biological replicates.
Determination of Cellular Metabolic Activity and Protein Content (AlamarBlue® and Sulforhodamine B Assay)

In order to rule out combinatory effects and cytotoxicity on gene transcription analysis, the impact of BUT, DON, NX-3 and their combinations on cell viability after 5 h of incubation was determined. Therefore, two cell viability assays, the alamarBlue® and the sulforhodamine B assay (SRB) according to Skehan et al. (38) were performed. HCEC-1CT cells were seeded into 96-well plates (2,500 cells/well) and allowed to grow for 48 h. Cells were then incubated for a total of 5 h, consisting of 2 h of toxin pre-incubation, followed by co-treatment with 25 ng/mL IL-1β for 3 h. As BUT was combined in a constant 10:1 ratio with DON and NX-3, the following concentrations and combinations were applied, obtained by 1:100 dilutions of respective stock solutions: 0.1–100 µM for BUT; 0.01–10 µM for DON/NX-3. After 4 h of incubation, 10 µL alamarBlue® reagent (Thermo Fisher Scientific, Massachusetts, USA) were added to the cells and incubated for further 1.25 h. Subsequently, 70 µL/well were transferred to a black 96-well plate and fluorescence intensity was measured at 530/560 nm (excitation/emission) using a Synergy H1 hybrid multi-mode reader (BioTek, Bad Friedrichshall, Germany). Afterwards, cells were rinsed with pre-warmed phosphate buffered saline and fixed by the addition of 1% (v/v) acetic acid solution to remove remaining staining and Sulforhodamine B assay according to Skehan et al. (38) were performed. HCEC-1CT cells were seeded into 1% (v/v) acetic acid solution and then washed twice with water and measured data of at least five independent, biological replicates performed in technical duplicates, was referred to the IL-1β-stimulated solvent control.

Data Visualization and Statistical Analysis

For statistical analysis and data visualization the software Origin 2018 (Northampton, USA) was used. In case of the NF-κB reporter assay and the respective cell viability assessment in THP1-Lucia™ cells at least five independent biological replicates were performed in technical duplicates. In HCEC-1CT cells, at least five biologically independent cell preparations were assessed by qRT-PCR analysis in technical duplicates, whereas only three independent biological replicates were performed in technical duplicates in case of the Magnetic Luminex® assay. Cell viability assessment in HCEC-1CT cells by the alamarBlue® and sulforhodamine B assay was conducted in at least five independent biological replicates performed in technical duplicates. In the reporter gene and cell viability experiments respective data were tested for normality of the distribution with the Shapiro-Wilk test. Significant differences between concentrations of the same compound with the respective lowest tested concentration were calculated by one-way ANOVA, followed by Bonferroni’s post-hoc tests (p < 0.05), whereas significant differences between the DON or NX-3 and the respective combinations with BUT were calculated with Student’s t-test. Regarding the gene transcription and cytokine secretion data, significant differences were calculated with Student’s t-test.

RESULTS

Impact on NF-κB Signaling Pathway Activity

In order to elucidate the role of BUT, DON, NX-3 and their combinations on NF-κB signaling pathway activation, THP1-Lucia™ cells, carrying a NF-κB-inducible Luc reporter construct, were grown 20 h in the presence of respective Fusarium secondary metabolites. After 2 h incubation, cells were further exposed to a pro-inflammatory IL-1β-stimulus (25 ng/mL). In a first experiment equimolar concentrations of BUT and the trichothecenes over a concentration range of 0.01–10 µM were tested (Figure 2A). The data show the percentage change of reporter gene activity of treatment (LPS + mycotoxin) in relation to LPS treatment alone. Compared to the solvent control already LPS alone triggered a 270-fold increase in luciferase activity, which was further enhanced by the co-treatment with the trichothecenes. Serving as positive control, heat killed Listeria monocytogenes cells (HKLM) induced the reporter stronger than LPS alone. Whereas, the lowest tested BUT concentrations (0.01, 0.1, and 1 µM) did not modulate NF-κB activity, at 5 µM a reduced luciferase signal was detected, reaching statistical significance at 10 µM. Both trichothecenes, DON and NX-3 caused comparable effects, so that at 1 µM a significant induction of the luciferase signal up to 183 ± 48% (DON) and 155 ± 42% (NX-3) of the stimulated solvent control were observed. These inductive effects were still measurable with 5 µM DON or NX-3. However, at 10 µM of either toxin a decrease of the signal to the solvent control level was determined, in line with a substantial loss in metabolic activity seen in the alamarBlue® assay (Figure 2B). In comparison to the effects induced by DON or NX-3 as single compounds, equimolar combinations of BUT and the trichothecenes at concentrations ≥ 1 µM, resulted mostly in a lower luciferase signal. Thus, on the one hand 1 µM DON and NX-3 led to a potent induction of the NF-κB signaling pathway and on the other hand 10 µM BUT significantly reduced the signal to 57 ± 10% without any cytotoxicity involved.

Even though, only limited BUT occurrence data are available, studies in oats reported a molar BUT/DON occurrence ratio for mean and maximum contamination levels of 3.2 and 8, respectively (9). Analyzing feed and feed components, Streit et al. (11) reported huge variations in the molar BUT/DON ratio (0.4–72). Therefore, trying to mimic different realistic occurrence scenarios, further experiments were performed combining BUT also at a 10-fold molar excess with the trichothecene mycotoxins (Figure 3). Fifty and hundred micromolar BUT, only the latter causing a significant decrease in cell viability, reduced the NF-κB activity to 0.3 ± 0.2% and 0.1 ± 0.1%, respectively. In 10:1 combination with 1 µM DON or NX-3, the presence of BUT reduced the luciferase signal from 195 and 196% to a signal even.
below the control level (96 and 91%). At higher concentrations, cytotoxic effects need to be taken into account. Interestingly, NF-κB activity was completely suppressed in mixtures containing 5 and 10 μM DON or NX-3. As the ingestion of highly contaminated cereal samples [up to 2,260 μg/kg (9–11)] is expected to cause at least intestinal BUT concentrations at low micromolar ranges (12), 10 μM BUT and 1 μM DON/NX-3 were selected as insightful and significant concentrations for the experiments focusing especially on the effects in human colon epithelial cells.

Modulatory Effects of BUT on Trichothecene-Induced Cytokine Gene Transcription

NF-κB signaling is involved in transcriptional induction of many cytokines, chemokines, and additional inflammatory mediators, such as interleukins, lymphokines, and tumor necrosis factors (39–41). As BUT was found to suppress significantly LPS-induced NF-κB activation in the THP1-Lucia™ cell line and to further inhibit NF-κB induction after trichothecene exposure, we aimed to elucidate the impact
Impact of BUT and Its Combinations With DON and NX-3 on Cytokine Secretion

Quantifying cytokine secretion levels by a magnetic Luminex® assay, we aimed to investigate, whether the reduced cytokine gene transcription of TNF-α, IL-8, and IL-6 after BUT-treatment, is also reflected on the protein level (Figure 5). Ten micromolar BUT significantly reduced TNF-α and IL-6 expression levels to 62 and 81% of the IL-1β stimulated solvent control samples, respectively. Protein levels of IL-8 were slightly induced after BUT treatment. Trichothecene treatments (1 µM) did not cause significant alterations of the cytokine secretion levels of TNF-α, IL-8, and IL-6 compared to the IL-1β stimulated control levels. However, combinatory treatments of BUT with DON or NX-3 showed a similar pattern as already observed after BUT treatment: TNF-α and IL-6 expression levels were both significantly reduced to 67% in case of DON and to 69 and 68%, respectively, after NX-3-exposure. In regard to IL-8 secretion also in the combinatory treatments no significant modulations were observed. Comparing the different cytokine concentrations after IL-1β stimulation (Table 1) substantial differences were found. Whereas, TNF-α was present at very low concentrations of about 50 pg/mL, IL-6 was detected at concentrations around 2,700 pg/mL. IL-8 was the most abundant cytokine, occurring at concentrations around 46 ng/mL. IL-1β stimulation (Table 1) caused an up to 100-fold increase in cytokine secretion. Non-stimulated solvent control samples

Cells pre-exposed for 2 h to BUT (10 µM) and then further challenged for 3 h with IL-1β (25 ng/mL), showed a significantly reduced mRNA expression of TNF-α, IL-1β, and IL-6. Whereas IL-8 was not altered significantly in its gene transcription compared to the IL-1β stimulated solvent control samples, only 0.4-, 0.6-, and 0.7-fold expressions of TNF-α, IL-1β, and IL-6 transcripts were reported, respectively. DON and NX-3 (1 µM), known for their pro-inflammatory properties, caused a substantial increase in TNF-α, IL-8, and IL-6 mRNA expression, while IL-1β gene transcription was found to be significantly elevated only after NX-3 treatment. Finally, combinations of BUT (10 µM) with DON or NX-3 (both 1 µM) were analyzed in the qRT-PCR experiments. For both trichothecenes a similar pattern of cytokine mRNA levels was determined. TNF-α, induced after DON and NX-3 treatment 2.2- and 1.9-fold, was reduced to 0.9 and 0.8-fold relative transcription, respectively, in combinatory treatments with BUT. While a similar reduction to the IL-1β stimulated solvent control level was found for IL-6, an even stronger decrease was determined in case of IL-1β, which was reduced to relative transcription levels of approximately 0.6. However, once more, IL-8 transcription levels were not affected by BUT in the combinatory treatments.

FIGURE 4 | Gene transcription levels of TNF-α, IL-1β, IL-8, and IL-6 in HCEC-1CT cells after 2 h pre-incubation with BUT (10 µM), DON (1 µM), NX-3 (1 µM) and respective 10:1 combinations, followed by 3 h co-stimulation with IL-1β (25 ng/mL) measured by qRT-PCR. Transcription data are normalized to the gene expression levels of two endogenous control genes (ACTB1 and GAPDH) and referred to the stimulated solvent control levels. Data are expressed as mean values ± SD of at least six independent experiments performed in technical duplicates. Significant differences to the stimulated solvent control are indicated with *p < 0.05, **p < 0.01 and ***p < 0.001, whereas significant differences of the combinatory data to the respective DON and NX-3 data are indicated with ◦p < 0.05 and ◦◦◦p < 0.001.
showed cytokine concentrations of 0.47 pg/mL TNF-α, 815 pg/mL IL-8, and 22 pg/mL IL-6 in the supernatants.

**Assessment of Cell Viability After Exposure to BUT and Its Combinations With DON and NX-3**

In order to rule out potential cytotoxicity to interfere in the qRT-PCR experiments, cell viability was assessed after 5 h exposure. In the alamarBlue® assay cellular metabolic activity was elucidated, whereas in parallel the cellular protein content was determined in the SRB assay (Figure 6). BUT decreased significantly the metabolic activity at 50 and 100 µM after 2 h pre-incubation and 3 h co-exposure with IL-1β. However, in the SRB assay no significant effects of BUT on the protein content of HCEC-1CT cells were determined.

At the highest concentrations tested (5 and 10 µM), DON and NX-3 slightly reduced the fluorescence signal in the alamarBlue® assay, even though NX-3 showed the tendency to be less potent. However, these effects were found to be more pronounced in the SRB assay reducing the cellular protein content in case of DON to 74 ± 21% (5 µM) and 76 ± 15% (10 µM) of the IL-1β challenged solvent control and to 83 ± 20% (5 µM) and 80 ± 1% (10 µM) after NX-3 incubation.

Mixtures of BUT with the trichothecenes at a constant molar ratio of 10:1, caused a slight loss in metabolic activity at 10/1 µM. Higher concentrations (50/5 and 100/10 µM) resulted in a more pronounced effect, reducing the metabolic activity to about 76 and 66%. However, in all tested combinations the impact of DON and NX-3 was found similar. Regarding the cellular protein content, up to the highest tested concentrations only marginal effects were observed in the SRB assay.

**DISCUSSION**

As the intestinal epithelium represents an important defense barrier against food contaminants, numerous studies investigated the impact of *Fusarium* mycotoxins on intestinal cells and
mediated immune and inflammatory response mechanisms (26, 27, 42, 43). Although the most prevalent Fusarium mycotoxin, DON, often co-occurs with many other co-regulated secondary fungal metabolites, data describing potential combinatorial effects on intestinal inflammation are lacking. Especially fungal compounds of low acute toxicity (4, 44), however occurring at high concentrations, such as BUT (8–11), are rarely taken into account.

To the best of our knowledge, so far effects of BUT on the activation of the NF-κB signaling pathway in intestinal epithelial cells have not been reported. The transcription factor NF-κB is considered one of the key regulators of inflammatory immune responses and its activation alters the expression of several inflammatory mediators, including cytokines and enzymes like cyclooxygenase-2 (COX-2) or inducible nitric oxide synthase (iNOS) (41). Using a reporter gene approach, we elucidated, applying not-cytotoxic BUT concentrations (<50 µM), significant reductions in NF-κB activity (Figures 2, 3). Previously, various butenolide-based structures, belonging to the family of α, β-unsaturated lactones, known as furanones, but decorated with different moieties, partly showed anti-inflammatory activity (45–47). While Ali et al. (46) reported suppressive effects of novel butenolide-based benzyl pyrroles (20 µM) on TNF-α, COX-2, and NF-κB protein expression of RAW 264.7 cells after 24 h incubation, in silico docking studies revealed other butenolide-derivatives as potential COX-2 inhibitors (45). Recently anti-inflammatory capacities of 10 butenolide-derivatives, isolated from the coral-derived fungus Aspergillus terreus were elucidated. Three compounds, tested at a concentration of 20 mM, showed remarkable inhibitory effects against nitric oxide production in the LPS-stimulated murine macrophage cell line, RAW 264.7 (47).

Even though preliminary studies described anti-inflammatory properties of various butenolide-derivatives, little is known about the underlying mechanism of action. However, as the Fusarium secondary metabolite BUT was found to significantly activate the Nrf2/ARE signaling pathway at concentrations between 1 and 50 µM (12) and as Nrf2 and NF-κB are considered two interplaying key pathways regulating the cellular redox status and responses to inflammatory triggers, their molecular cross-talk might play an important role in BUT-induced toxicities. One possible hypothesis to be tested in future work is that BUT, as a Michael acceptor, potentially reacts with the critical cysteine thiolate groups of the Nrf2-repressor protein Kelch-like ECH-associated protein 1, releasing Nrf2 and inducing the transcription of ARE-mediated antioxidative and cytoprotective enzymes, such as heme oxygenase-1 (HO-1), NAD(P)H dehydrogenase (quinone 1), γ-glutamate-cysteine ligase, glutathione peroxidase, and glutathione S-transferases (48–50). Numerous studies dealing with Nrf2-activating compounds reported respective anti-inflammatory potential, including reduced COX-2, TNF-α, iNOS, and IL-1β production (51, 52). Furthermore, increases in glutathione levels, as well as in HO-1 activity were found to decrease or to even inhibit NF-κB activity (52–54). Hence, it can be speculated that BUT inhibits NF-κB activity in a negative feedback loop due to a pronounced Nrf2/ARE signaling pathway activation.

Combining BUT, as an NF-κB-suppressor, at equimolar concentrations or 10-fold excess of BUT with the co-occurring mycotoxins DON and NX-3, both causing an increase in NF-κB activity, mediated a reduction of the luciferase signal down to solvent control level (Figures 2, 3). Whereas, the impact of NX-3 on the activation of the NF-κB signaling pathway has not been addressed in literature so far, DON-induced NF-κB translocation has been reported already several times (55, 56). This trichothecone-induced inflammatory response was further enhanced by co-treatment with LPS and interferon-γ of...
non-tumorigenic rat IEC-6 cells, involving TNF-α production, iNOS and COX-2 expression, ROS release and inflammasome activation (56). In intestinal explants and human Caco-2 cells an induction of gene expression and secretion of the NF-κB-dependent, pro-inflammatory cytokine IL-8 was observed after exposure to DON (26, 43). This interleukin is considered an early marker of inflammatory processes, acting as a potent chemo-attractant for leukocytes and T-lymphocytes underlying gut epithelial cells (57). In our study both trichothecenes (1 μM) increased mRNA levels of IL-8, as well as of TNF-α and IL-6 (Figure 4). BUT, in contrast, significantly reduced transcript concentrations of TNF-α, IL-1β, and IL-6, whereas no suppression was found in case of IL-8. Accordingly, in the combinatorial cell treatments, a suppression of TNF-α, IL-6, and IL-1β transcripts was found, of which the latter was even significantly below the stimulated solvent control level. IL-8 was not modified by the presence of BUT, showing similar mRNA levels as in the trichothecene-treated samples. Going one step further, having a look at potential modifications of the protein expression levels (Figure 5), surprisingly the induction in the transcription levels of all four pro-inflammatory cytokines due to the presence of DON and NX-3 was not reflected in the cytokine expression. Van De Walle et al. (26), applying similar DON concentrations (1 and 10 μM) on Caco-2 cells for 48 h in presence of IL-1β (25 ng/mL), reported a dose-dependent increase of the IL-8 secretion levels, in accordance with the findings of Moon et al. (24) in human intestinal 407 cells after 12 h incubation. Corresponding to the substantial inhibition of the NF-κB signaling pathway observed in the reporter gene assay, BUT, alone, as well as in the combinatorial treatments significantly reduced TNF-α and IL-6 cytokine expression. However, it remains unclear, why BUT has no inhibitory effect on IL-8 transcription and expression, even though a substantial suppression of the NF-κB activity leading to a decrease in TNF-α, IL-6, and IL-1β transcription and secretion was detected. Cytokine secretion is multifactorially regulated. Having a closer look into the IL-1 signaling pathway, IL-1β-stimulation causes not only post-receptor activation of NF-κB, but also of MAPKs such as p38 or c-Jun N-terminal kinases/stress-activated kinases (JNK/SAPK), cooperatively inducing the transcription of target genes like IL-6, IL-1β, and IL-8 (58, 59). Furthermore, the differences in the cytokine concentrations determined in the Magnetic Luminex® assay need to be taken into account (Table 1). IL-8 was present in the supernatant of IL-1β-stimulated samples at concentrations around 50,000 pg/mL, whereas IL-6 and TNF-α were detected at 20- or even 1,000-fold lower concentrations. Stimulating for 16 h, Schuerer-Maly et al. (60) reported similar IL-8 concentrations in HT-29, while in the supernatants of Caco-2 cells only 4,000 pg/mL were detected. Even though different exposure procedures render such direct comparisons difficult, the comparably high expression of IL-8 in our samples, leads to the assumption that some cellular signaling may be induced in HCEC-1CT cells substantially activating IL-8 expression and secretion. Thus, potential BUT-induced inhibiting effects on IL-8, might be overlaid by such co-occurring mechanisms.

Regarding cytotoxic effects, after 5 h of incubation the concentrations used in these experiments only marginally affected cellular viability. In the combinatorial treatments cellular metabolic activity was slightly modulated, whereas no effects on the cellular protein content were observed. Various in vitro and in vivo studies investigated BUT-induced toxicities, showing the compound’s potential to trigger oxidative stress, to increase intracellular ROS production, lipid peroxidation and to cause oxidative DNA damage (4–7, 13, 14, 61–64). Whereas, most previous studies administered comparably high concentrations of BUT, we could recently show in combinatorial treatments with DON that BUT concentrations in the low micromolar range contributed significantly to the overall cytotoxicity of respective mixtures (12). Beyond that, pre-incubations with BUT for 1 or 3 h were sufficient to enhance DON-induced cytotoxic effects (24 h) in HepG2 cells (12).

In conclusion, our results provide first evidences that the Fusarium secondary metabolite BUT, substantially alters the inflammatory response of the human intestinal epithelial cell line HCEC-1CT. NF-κB activity was significantly suppressed in the presence of BUT, resulting in downstream consequences such as reduced transcription and expression of pro-inflammatory cytokines. As BUT co-occurs in high abundance with mycotoxins such as DON and NX-3 (8–11), known to act as activators of the NF-κB signaling pathway and to increase the expression of inflammatory genes in the intestine (26, 43), we analyzed the intestinal inflammatory response following combined exposure to BUT and DON or NX-3. BUT, in 10-fold molar excess substantially suppressed the trichothecene-induced NF-κB activation, as well as the transcription and secretion of important pro-inflammatory cytokines. Thus, a required inflammatory defense response of the intestinal barrier against trichothecene mycotoxins might be partly suppressed by the presence of BUT in food and feed. A relevant question is therefore whether such a situation may occur in naturally contaminated grain. A survey on 83 animal feed and feed components reported median concentrations of 23 and 122 μg/kg of BUT and DON, respectively (11). This suggests that on average the molar ratio BUT/DON is rather 0.4 than 10, and the observed interaction may not be very relevant. Yet, if individual samples are considered, a different picture emerges. In 3 of six corn cob samples, listed individually, the BUT/DON molar ratio was 29.2, 72.8, and 7.3, suggesting that an interference of BUT with DON-induced inflammatory responses could be relevant after all. Comparing median and maximum contamination levels of BUT (median: 231 μg/kg; maximum: 2,260 μg/kg) and DON (median: 150 μg/kg; maximum: 636 μg/kg) in barley samples analyzed by Uhlig et al. (9) molar BUT/DON ratios of 3.2 and 8 have been detected. Determination of BUT in mycotoxin surveys seems highly warranted to close the gap in occurrence data. In contrast to animal feed, grain used for food is typically not consumed raw, and the stability of BUT during food processing is also unknown.

Our observation of combinatorial effects between trichothecenes and BUT argue for further research into occurrence and intake of BUT, and a more detailed analysis of the underlying mechanism of action, in order to evaluate how BUT modulates the pathogenesis of intestinal inflammatory diseases and whether it should be considered in the risk assessment of naturally occurring mixtures.
DATA AVAILABILITY STATEMENT

All datasets presented in this study are included in the article.

AUTHOR CONTRIBUTIONS

LW was involved in designing the study, measuring the NF-κB reporter gene assay, qRT-PCR, cell viability and magnetic Luminex® assays, carried out statistical analysis, and wrote the manuscript. GA was involved in coordination of the project and refined the manuscript. DM was involved in designing and supervising the project and refined the manuscript. All authors contributed to the article and approved the submitted version.

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Woelflingseder et al.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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