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Enhancing the anti-inflammatory activity of chalcones by tuning the Michael acceptor site†

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Inflammatory signaling pathways orchestrate the cellular response to infection and injury. These pathways are known to be modulated by compounds that alkylate cysteinyl thiols. One class of phytochemicals with strong thiol alkylating activity is the chalcones. In this study we tested fourteen chalcone derivatives, α -X-substituted 2',3,4,4'-tetramethoxychalcones (α -X-TMCs, X = H, F, Cl, Br, I, CN, Me, *p*-NO₂-C₆H₄, Ph, *p*-OMe-C₆H₄, NO₂, CF₃, COOEt, COOH), for their ability to modulate inflammatory responses, as monitored by their influence on heme oxygenase-1 (HO-1) activity, inducible nitric oxide synthase (iNOS) activity, and cytokine expression levels. We confirmed that the transcriptional activity of Nrf2 was activated by α -X-TMCs while for NF- κ B it was inhibited. For most α -X-TMCs, anti-inflammatory activity was positively correlated with thiol alkylating activity, *i.e.* stronger electrophiles (X = CF₃, Br and Cl) being more potent. Notably, this correlation did not hold true for the strongest electrophiles (X = CN and NO₂) which were found to be ineffective as anti-inflammatory compounds. These results emphasize the idea that chemical fine-tuning of electrophilicity is needed to achieve and optimize desired therapeutic effects.

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Introduction

The expression of anti-inflammatory proteins, such as heme oxygenase-1 (HO-1), and proinflammatory proteins, such as inducible NO synthase (iNOS), can be modulated by protein thiol modifications. In particular, the oxidation of protein thiols or their alkylation by electrophiles, especially α , β -unsaturated carbonyl compounds (often referred to as Michael acceptors since they are able to undergo 1,4-additions with thiols, see Fig. 2A)¹ leads to either activation or inhibition of individual transcription factors. Alkylation of Keap1² releases the transcription factor Nrf2 and thus induces HO-1 expression (Fig. 2B). The inactivation of NF- κ B³ by Michael

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acceptors⁴ inhibits expression of proinflammatory proteins (*e.g.* iNOS, TNF, IL-6, IL-8) and appears to be mediated by either inhibition of I κ B α kinase (IKK) (Fig. 2C, Inhibition I)⁵ or by alkylation of NF- κ B cysteinyl residues within the DNA-binding domain⁶ (Fig. 2C, Inhibition II). For example, the natural chalcone xanthohumol was found to covalently modify cysteinyl residues of both NF- κ B p65 and IKK.⁵ However, the activation of Nrf2 and inactivation of NF- κ B by protein thiol modifications may also be caused more indirectly through glutathione (GSH) depletion or oxidation.⁷

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The anti-inflammatory activity of chalcones is of significant biomedical interest. Precise tuning of chalcone thiol reactivity may lead to derivatives with improved therapeutic potential. To modulate the chemical reactivity in thia-Michael additions a substituent can be introduced in the α -position of the α , β -unsaturated carbonyl system (Fig. 2A). Compounds with different substituents in the α -position can then be screened for desired biological activities.⁸ We previously established a panel of fourteen α -X-substituted 2',3,4,4'-tetramethoxychalcones (α -X-TMCs, Fig. 1) exhibiting a wide range of chemical reactivity. We observed that increased chalcone thiol reactivity correlated with stronger induction of HO-1 and stronger repression of iNOS.^{8b} Using a selection of cell-based assays, we now further demonstrate that α -X-TMCs differentially affect the Nrf2 and NF- κ B inflammatory response pathways within living cells.

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[†]Electronic supplementary information (ESI) available: Fig. S1: inhibition of NO production and Table S1: C10 values of Nrf2 induction. See DOI: 10.1039/c4ob02301c

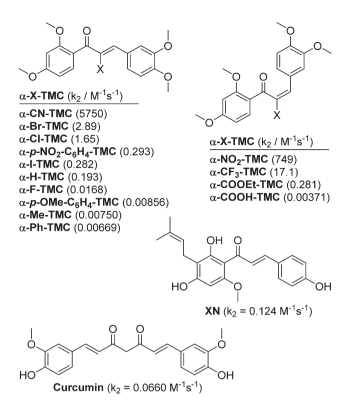


Fig. 1 Structures of α -X-2',3,4,4'-tetramethoxychalcones (α -X-TMCs) tested in this study together with the natural products xanthohumol (XN) and curcumin which served as positive controls. Second-order rate constants (k_{22}) for the reaction with cysteamine are given in parentheses.⁸

Results and discussion

α-X-TMCs induce HO-1 activity in RAW264.7 macrophages

Using our recently developed ELISA-based HO-1 assay,⁹ α -X-TMCs were tested for their ability to induce HO-1 activity in RAW264.7 macrophages (Fig. 3). All α-X-TMCs were used at a concentration of 1 μ M, except for α -CF₃-TMC, which is cytotoxic at this concentration^{8b} and therefore was used at 0.5 µM. HO-1 activity was determined after 3, 6 and 24 h of treatment with α -X-TMCs. Most of the α -X-TMCs stimulated HO-1 activity weakly, with a maximal increase of 2.7 fold after 6 h of treatment in the case of α -CF₃-TMC. The derivatives α -CF₃-TMC, α-Br-TMC, α-H-TMC and α-Cl-TMC exhibited the most significant effect (2.1-2.7 fold of control). Induced HO-1 activity peaked at 6 h, with the exception of α -I-TMC which induced even higher activity at 24 h (1.9 fold induction). Overall, these findings agree well with our previous study, which showed that α-X-TMCs with intermediate to high electrophilicity (*i.e.* α -CF₃-TMC, α -Br-TMC, α -Cl-TMC and α -H-TMC) efficiently induce HO-1 protein expression at the same concentrations (0.5, 1 µM).^{8b} Likewise, as seen previously,^{8b} the strongest electrophiles, α -CN-TMC and α -NO₂-TMC did not cause significant induction of HO-1 activity at 1 µM within the same time frame. This may be due to a direct reaction with GSH,¹⁰ potentially competing with the modification of protein thiols.

α-X-TMCs induce Nrf2 activity in AREc32 cells

Having observed induction of HO-1 activity by α -X-TMCs (Fig. 3), we then asked whether this effect was caused by the activation of Nrf2, a key transcription factor for HO-1

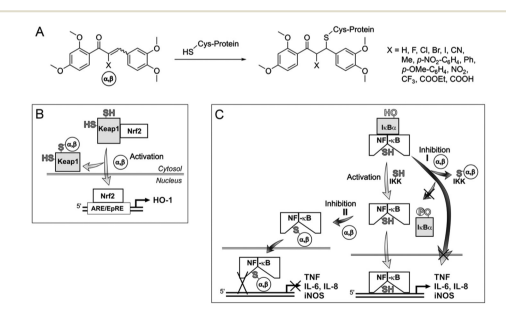


Fig. 2 Influence of α ,β-unsaturated carbonyl compounds, acting as Michael acceptors, on transcription factors involved in inflammation. (A) Michael addition leading to protein-electrophile adducts; possible influence of Michael acceptors (α , β) on the Keap1-Nrf2 pathway (B) and on the NF- κ B pathway (C). α , β , α , β -unsaturated carbonyl compound; Keap1, Kelch-like ECH-associated protein 1; Nrf2, nuclear factor-erythroid 2-related factor 2; ARE/EpRE, antioxidant response element/electrophile response element; HO-1, heme oxygenase-1; I κ B α , inhibitory factor κ B α ; NF- κ B, nuclear factor κ -B kinase; IL-6, IL-8, interleukin-6, interleukin-8; iNOS, inducible NO synthase; TNF, tumor necrosis factor.

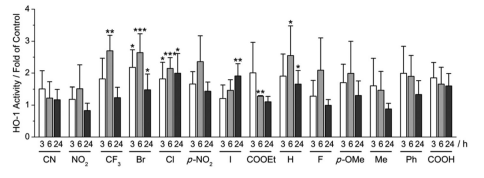


Fig. 3 Influence of α -X-TMCs on HO-1 activity. RAW264.7 cells were treated with different α -X-TMCs at 1 μ M (0.5 μ M for α -CF₃-TMC) for the indicated times (3, 6 and 24 h). Labels indicate the substituent X in the α -position of the α , β -unsaturated carbonyl moiety of the chalcone (p-NO₂ is short for p-NO₂-C₄H₆ and p-OMe for p-OMe-C₄H₆). HO-1 activity was determined by an ELISA-based bilirubin quantification assay. At least three independent measurements were performed. Levels of significance: ***, p < 0.001; **, p < 0.01; *, p < 0.05.

expression. Nrf2 activation was quantified in human breast cancer MCF7 cells expressing a luciferase reporter construct under the control of the human ARE (antioxidant response element) core sequence (AREc32 cells).¹¹ The α -X-TMCs were applied for 24 h at concentrations ranging from 1.56 to 100 µM. In parallel, cytotoxicity was assessed by sulforhodamine B (SRB) staining.¹² Only compound concentrations causing less than 50% cytotoxicity were included in the data analysis. Values were normalized to the established Nrf2inducer xanthohumol (XN) which in our assay activated Nrf2 maximally at 25 μ M with 42 ± 11 fold induction (taken for normalization as 100%) relative to the solvent control (Fig. 4). The best Nrf2 activators were α -CF₃-TMC (219% of XN at 3.13 μ M), α -Br-TMC (317% of XN at 6.25 μ M) and α -Cl-TMC (180% of XN at 6.25 μ M), which mirrors their effectiveness in inducing HO-1 expression and activity. Similar values of

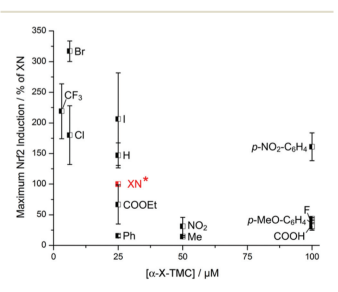


Fig. 4 Maximum Nrf2 induction of α -X-TMCs relative to the maximum Nrf2 induction of the positive control xanthohumol (XN*) (set to 100%). Activities were tested by incubating α -X-TMCs at different concentrations (1.56–100 μ M) for 24 h with AREc32 cells. α -CN-TMC showed no effect and thus was not included. Data points represent the mean of 2–3 independent measurements.

maximal Nrf2 induction showed α -H-TMC (147% of XN) and α -I-TMC (206% of XN), however at a higher concentration (25 μ M). Weak electrophiles (α -COOH-TMC, α -*p*-OMe-C₆H₄-TMC, α -F-TMC, and α -Ph-TMC) were less effective than XN (16–44%), even at higher concentrations (25, 50 or 100 μ M). Overall, and similar to HO-1 activity, k_2 values (Fig. 1)⁸ correlated with the ability of α -X-TMCs to induce Nrf2 activity (Fig. 4). In agreement with our previous results on HO-1 expression,^{8b} the α -X-aromatic compound α -*p*-NO₂-C₆H₄-TMC activated Nrf2 only at very high concentrations (161% of XN at 100 μ M). While α -NO₂-TMC was slightly active (31% of XN at 50 μ M), α -CN-TMC was completely inactive, again suggesting that the two strongest electrophiles in our panel are ineffective in inducing Nrf2 dependent gene expression.

Some of the compounds enhanced Nrf2 activity by 10 fold at a very low concentration, as quantified by their C10 values (Table S1[†]). α -CF₃-TMC (C10 = 0.495 μ M) was most potent, followed by α -Br-TMC (1.39 μ M), α -Cl-TMC (2.46 μ M), α -I-TMC (5.94 μ M), α -H-TMC (7.23 μ M) and XN (10.4 μ M). C10 values strongly correlated with maximal Nrf2 activation (Fig. 4) and with thiol reactivity k_2 values (Fig. 1).

The unexpectedly high C10 values of α -*p*-NO₂-C₆H₄-TMC (14.8 μ M) and α -COOEt-TMC (18.3 μ M) may indicate additional intracellular interactions caused by the substituent X, potentially interfering with the electrophilic reactions that lead to Nrf2 activation.

α -X-TMCs inhibit iNOS-dependent NO production in LPS-stimulated RAW264.7 macrophages

Next we asked to what extent the different α -X-TMCs inhibit NF+ κ B-dependent gene expression. To this end we quantified iNOS-dependent NO production by LPS-stimulated RAW264.7 macrophages using the Griess assay¹³ (Table 1). In our previous study we already observed a significant correlation between k_2 values and the ability of α -X-TMCs to inhibit iNOS activity.^{8b} This correlation was established for the series X = CF₃, Br, Cl, I, H, and F, which are the α -X-TMCs with intermediate to strong, but not very strong electrophilicity, and for which X is not expected to engage in extra H bonding or π -interactions.

Table 1InhibitionofNOproductioninLPS-stimulatedRAW264.7macrophages by α -X-TMCs. The α -X-TMCs are listed in orderof relative thiol reactivity (k_2 values, decreasing from top to bottom)

α-X-TMC	Toxicity limit ^a [µM]	Maximum inhibition of NO production [%] (conc [µM]) ^b	IC ₅₀ (NO production) [μM]
CN	25	$16.4 \pm 8.7 (1)$	nd
NO_2	10	$26.9 \pm 7.9(10)$	nd
CF ₃	0.5	$84.6 \pm 8.9(0.5)$	0.120 ± 0.062
Br	1	$67.4 \pm 11.7(1)$	0.640 ± 0.179
Cl	1	$47.2 \pm 14.6(1)$	0.992 ± 0.482
$p-NO_2-C_6H_4$	10	35.8 ± 2.2 (10)	nd
Ī	5	$72.2 \pm 9.7(5)$	3.15 ± 0.70
COOEt	10	66.8 ± 8.4 (10)	6.65 ± 0.53
Н	10	$96.9 \pm 11.9(10)$	4.44 ± 1.29
F	25	$86.6 \pm 18.9(25)$	12.9 ± 5.6
p-OMe-C ₆ H ₄	10	ns	nd
Me	10	$-34.7 \pm 15.2 (5)$	nd
Ph	25	27.9 ± 13.4 (25)	nd
COOH	25	$20.8 \pm 3.7 (25)$	nd

^{*a*} Toxicity of chalcones was measured with the MTT assay in the absence of LPS (no differences were found in the presence of LPS). Concentrations causing cell viabilities > 80% after 24 h incubation were assigned as non-toxic. ^{*b*} Stimulation with α -X-TMCs for 24 h in the presence of 10 ng mL⁻¹ LPS. ns, non-significant (no significant NO inhibition observed); nd, not determined.

Using different non-toxic concentrations we determined the IC₅₀ values for the inhibition of NO production (Table 1). With the exception of α -Me-TMC and α -*p*-OMe-C₆H₄-TMC, all α -X-TMCs inhibited NO generation. α -*p*-OMe-C₆H₄-TMC was completely inactive, while α -Me-TMC even enhanced NO production by 35%. Most compounds revealed a concentration-dependent inhibitory effect on NO production, with the exception of α -CN-TMC (see ESI Fig. S1†). While the strongest inhibition of NO release was achieved by treatment with 10 μ M α -H-TMC (97% inhibition, Table 1), the most potent chalcone was α -CF₃-TMC causing 85% inhibition at a concentration of 0.5 μ M.

For seven α -X-TMCs it was possible to calculate IC₅₀ values for the inhibition of NO production. These correlated very well with the corresponding k_2 values, in the following order: α -CF₃-TMC (IC₅₀ = 120 nM) < α -Br-TMC (640 nM) < α -Cl-TMC (992 nM) < α -I-TMC (3.15 μ M) < α -H-TMC (4.44 μ M) < α -F-TMC (12.9 μ M). The IC₅₀ value for α -COOEt-TMC (6.65 μ M) did not match the expected correlation. In addition, α -*p*-NO₂-C₆H₄-TMC was considerably less active than one would have predicted from its k_2 value. However, these observations are in agreement with our previous study and may be explained by electrophilicity-independent intracellular interactions or reactions. The relatively weak inhibition of NO production by α -CN-TMC and α -NO₂-TMC (Table 1) could result from their very strong electrophilicity, possibly causing their inactivation.

α -X-TMCs inhibit TNF secretion by LPS-stimulated primary human macrophages

Having observed consistent effects in different assays, we selected four α -X-TMCs with anti-inflammatory activity (X = CF₃, Br, Cl, and H) and two inactive α -X-TMCs (X = CN, NO₂) for further study. We wanted to ascertain that anti-inflamma-

tory effects can also be observed with primary human cells, not just with cell lines. Thus we tested the influence of the compounds on TNF secretion by LPS-stimulated primary human macrophages (Fig. 5A).

TNF levels in the cellular supernatant were measured by ELISA after 24 h. Using α -X-TMCs at a concentration of 5 μ M, the following percentages of inhibition were found: α -Br-TMC 76%, α -CF₃-TMC 70%, α -Cl-TMC 30%, α -H-TMC 25%, α -CN-TMC 17%, and α -NO₂-TMC 0%. At a concentration of 1 μ M only α -Br-TMC (35%) and α -Cl-TMC (14%) inhibited TNF release. The α -CF₃-TMC response was highly variable amongst the four donors tested and therefore appeared less pronounced. We confirmed that decreases in TNF production were not caused by cell death (data not shown). This result demonstrated once again that α -Br-TMC, α -CF₃-TMC and, to

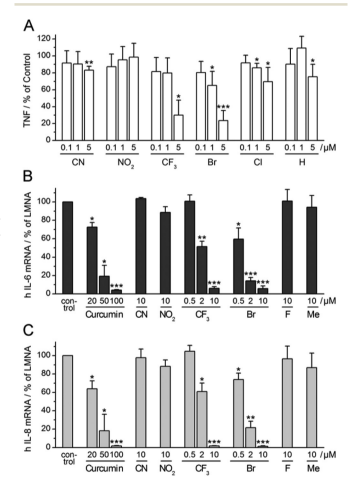


Fig. 5 Influence of selected α -X-TMCs on LPS-induced cytokine expression. (A) TNF production of LPS-stimulated (100 ng mL⁻¹) human primary macrophages after 24 h incubation without (control) or with α -X-TMCs; independent experiments with cells from four individual donors were performed. TNF was determined in cellular supernatants by (B) ELISA. (C) RT-PCR analysis of IL-6 and IL-8 mRNA levels in human HeLa cells relative to LMNA (human Lamin A/C). HeLa cells were pre-incubated with α -X-TMCs or DMSO (control) for 45–60 min prior to a 60 min stimulation with 30 ng mL⁻¹ recombinant human TNF: (B) IL-6, (C) IL-8. Data represent four (A) or three (B), (C) independent experiments. Levels of significance: ***, p < 0.001; **, p < 0.01; *, p < 0.05.

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lesser extent, α -Cl-TMC interfere with proinflammatory pathways – in this case inhibiting TNF release.

α -X-TMCs inhibit expression of IL-6 and IL-8 in TNF-stimulated HeLa cells

Expression of the cytokines interleukin-6 (IL-6) and interleukin-8 (IL-8) is stimulated by NF-kB transcriptional activity and therefore predicted to be influenced by treatment with α -X-TMCs. We tested the two strongest (α -CN-TMC, α -NO₂-TMC), two moderately strong (α -CF₃-TMC and α -Br-TMC) and two poor to very poor electrophiles (α -F-TMC and α -Me-TMC). As expected, using TNF-stimulated HeLa cells, we found a strong reduction of IL-6 and IL-8 mRNA levels upon treatment with either α -CF₃-TMC or α -Br-TMC (Fig. 5B and C). At 10 μ M both compounds almost completely abolished the induction of IL-6 and IL-8 (94-99% inhibition). Whilst α-Br-TMC still showed a pronounced effect at 0.5 μ M, α -CF₃-TMC gave similar inhibition ratios at 2 µM, but not below. Both compounds were about 10 fold more active than the established NF-kB inhibitor curcumin¹⁴ which served as a positive control and reference (Fig. 5B and C).

α-Br-TMC inhibits TNF-induced nuclear translocation of NF-κB p65 in HeLa cells

Having observed that NF- κ B-dependent transcription of IL-6 and IL-8 was inhibited by α -CF₃-TMC and α -Br-TMC, we aimed to confirm that this effect was due to the inhibition of NF- κ B activation. We therefore monitored nuclear translocation of NF- κ B in TNF stimulated HeLa cells in the presence or absence of 10 μ M α -Br-TMC.

While cytosolic levels of NF- κ B p65 were largely unaffected by α -Br-TMC or TNF, TNF-induced nuclear translocation of NF- κ B p65 was significantly reduced after pre-treatment with α -Br-TMC (Fig. 6). This result demonstrated that α -Br-TMC inhibits the NF- κ B activation pathway upstream of nuclear translocation.

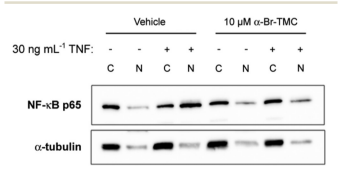


Fig. 6 Inhibition of TNF-induced nuclear translocation of NF- κ B p65 by α -Br-TMC. Western blot analysis of cytosolic (C) and nuclear (N) extracts stained with an anti-NF- κ B p65 antibody. HeLa cells were pre-incubated with α -Br-TMC or DMSO (vehicle) for 45 min prior to a 15 min stimulation with 30 ng mL⁻¹ recombinant human TNF. One representative blot out of three independent experiments is shown. Immunostaining of α -tubulin served as a quality control for cellular fractionation.

Chemicals and reagents

 α -X-TMCs were prepared as described previously.⁸ Compounds were purchased from the following commercial sources and used without further purification: curcumin (B6938), hemin and gelatin (from cold water fish skin) from Sigma (Germany), NADPH from AppliChem (Germany), OPD (*ortho*-phenylenediamine dihydrochloride) from Acros Organics (Belgium), bilirubin from Frontier Scientific (UK), Triton X-100 from Merck (Germany), xanthohumol (XN) from Carl Roth (Germany) or, alternatively, XN was obtained as hop-derived ethanolic extract with 80% purity (Hallertauer Hopfenverwertungsgesellschaft, Mainburg, Germany) and purified by silica gel column chromatography with *n*-hexane–ethyl acetate as previously described (purity (HPLC) > 99%).¹⁵

Cell viability test (MTT assay) on RAW264.7 cells

Murine macrophage RAW264.7 cells were cultured in RPMI 1640 medium supplemented with 10% heat-inactivated fetal calf serum (FCS) and 2 mM glutamine (Biochrom, Germany) at 37 °C in humidified air containing 5% CO₂. In order to exclude cytotoxicity of the compounds a MTT assay was performed and the effect of the compounds on cell viability compared to control cells was determined. The results were published in a previous study of the α -X-chalcones.^{8b}

ELISA-based HO-1 activity assay on RAW264.7 cells

The assay was performed as previously reported *via in situ* formation of bilirubin by HO-1/BVR activity and quantification of bilirubin by ELISA.⁹

Briefly, RAW264.7 macrophages $(8 \times 10^4 \text{ cells per well})$ were placed in 96-well plates for 24 h and then treated with compounds for 3, 6 and 24 h. Controls received no compounds. After cell lysis (40 mM TRIS-HCl, pH 7.4, 250 mM sucrose, 137 mM NaCl, 10% (v/v) glycerin, 2 mM EDTA, 0.1% (v/v) Triton X-100, complete protease inhibitor cocktail, Roche Diagnostics, Germany) the HO-1 reaction mixture (40 mM TRIS-HCl, pH 7.4, 250 mM sucrose, 0.3 mM NADPH, 1.0 ng BVR (biliverdin reductase, Stressgen, Assay Designs, USA) and 2.5 µM hemin) was applied for 1 h. Bilirubin standards $(0.50-2500 \times 10^{-9} \text{ M bilirubin})$ were prepared in 40 mM TRIS-HCl, pH 7.4, 250 mM sucrose and combined with supernatant of whole cell lysates from control cells. Bilirubin was quantified by using an excess of the anti-bilirubin antibody 24G7 from mouse (Shino-Test, Japan, 0.57 μ g μ L⁻¹ in 1% G-PBS with 0.5 mM sodium salicylate) and subsequently analyzing the unbound 24G7 by ELISA. To trap 24G7 an immunoplate (Nunc, Denmark) coated with a bilirubin-BSA conjugate (0.35 µg protein per well) was utilized. Detection was done using a HRP-conjugated anti-mouse antibody from goat (Rockland, USA; 1:10 000) with a freshly prepared substrate solution (0.40 g mL⁻¹ OPD and 0.40 μ L mL⁻¹ 30% H₂O₂ in citrate buffer, pH 5.0). After quenching with aqueous 3 M H_2SO_4 the absorbance was measured at 492 nm (Multiskan Spectrum, Thermo, Finland). The sigmoidal calibration curve was fit to a

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four parameter logistic equation to determine unknown bilirubin concentrations. HO activity was calculated as pmol bilirubin formed per hour and per milligram of protein (pmol BR h^{-1} mg⁻¹) and expressed as fold HO-1 activity compared to control cells.

AREc32 (human mammary epithelial adenocarcinoma MCF7) cells

AREc32 is a reporter cell line derived from the human mammary epithelial adenocarcinoma MCF7 cells. It contains a luciferase reporter gene construct under the control of eight copies of the Nrf2 binding site ARE.¹¹ AREc32 cells were cultured at 37 °C and 5% CO₂ in a humidified atmosphere in complete DMEM medium supplemented with 10% heat-inactivated FCS, 2 mM glutamine, 100 U mL⁻¹ penicillin and 100 μ g mL⁻¹ streptomycin (medium and all supplements from Gibco, USA), and maintained under antibiotic selection (0.8 mg mL⁻¹ G418 (Sigma, Germany)).

Cytotoxicity assay on α -X-TMC-stimulated AREc32 cells

Biomass content was assessed by sulforhodamine B (SRB, Sigma, Germany) staining.¹² Briefly, AREc32 cells were seeded and treated as for the luciferase reporter assay. Detached cells in the supernatant were discarded and attached cells were fixed with 10% trichloroacetic acid for 30 min at 4 °C. Then, the plate was rinsed with distilled water, dried and stained for 15 min with a 0.4% solution of SRB in 1% acetic acid, washed several times with 1% acetic acid and dried again. The dye was eluted with 10 mM TRIS base followed by the measurement of OD at 490 nm and 515 nm (SpectraMax M5e Microplate reader, Molecular Devices, USA).

Luciferase reporter assay on α-X-TMC-stimulated AREc32 cells

AREc32 cells were seeded at a density of 9×10^3 per well in 50 μ L growth medium in 384-well plates. After 22 h compounds were added to obtain final concentrations in the range between 1.56–100 μ M. Each concentration was tested in quadruplicate. An equal volume of vehicle was added to the control wells (0.016–1% DMSO). The reporter assay was performed as described previously.¹¹ For normalization all values were divided by vehicle control values, yielding the fold change relative to the control. Further, the luciferase activity was normalized to biomass content.

iNOS inhibition test (Griess assay) on RAW264.7 cells

The produced NO which is accumulated as nitrite in the culture medium was quantified using the Griess reaction. RAW264.7 macrophages (8×10^4 cells per well) were plated in 96-well plates and allowed to attach for 24 h. Stock solutions of compounds were prepared in DMSO (100 mM) and stored at -20 °C. Test concentrations were freshly prepared by diluting the stock solution in culture medium in presence and absence of 10 ng mL⁻¹ LPS and then incubated with the cells for 24 h. The final concentration of DMSO was $\leq 0.025\%$ and the total assay volume was 100 µL. 50 µL of culture medium was mixed with 50 µL of Griess reagent (0.1% NED (*N*-1-naphtylethylene-

diamine dihydrochloride), 1% sulfanilamine, 0.35% phosphoric acid in water) and incubated for 15 min at room temperature. The absorbance was measured at 560 nm (Multiskan Spectrum, Thermo, Finland) at 25 °C and nitrite concentrations were calculated from a standard curve established with serial dilutions of NaNO₂ (0/5/10/25/50 μ M) in culture medium. Nitrite production was expressed as percent compared to control cells incubated with 10 ng mL⁻¹ of LPS.

Quantification of TNF in LPS-stimulated human macrophages treated with α -X-TMCs

Macrophages were differentiated from human monocytes, obtained by leukapheresis of healthy donors, followed by density gradient centrifugation over Ficoll/Hypaque and separation by countercurrent centrifugation (J6M-E centrifuge, Beckmann, Germany) as described previously.¹⁶ The purity (> 85%) of the monocytes was determined by morphology and the expression of CD14 antigen. For the differentiation of macrophages, monocytes were cultured on teflon foils for 7 days at a concentration of 1×10^{6} cells mL⁻¹ in RPMI 1640, 10% human AB-serum (both from PAN Biotech, Germany), 2 mM glutamine and 50 U mL⁻¹ penicillin-50 mg mL⁻¹ streptomycin (all from Gibco/Life Technologies, Germany) at 5% CO2 and 37 °C. Thereafter, macrophages were harvested, counted and 0.2 \times 10⁶ cells in 1 mL incubated overnight in 24-well plates with 100 ng mL⁻¹ LPS (Enzo Life Sciences, Germany) for maturation with or without the addition of α -X-TMCs with X = CN, NO₂, CF₃, Br, Cl, H (0.5, 1, 5 µM). After 24 h, supernatants were harvested, frozen and later analyzed for cytokine production according to the manufacturer's instructions (DuoSet ELISA, R&D Systems, Germany).

Inhibition of NF-κB activation assay in human HeLa cervical cancer cells

HeLa cells (a kind gift from Daniela Männel, Institute of Immunology, University of Regensburg) were grown in RPMI 1640 supplemented with 10% heat-inactivated FCS and 100 U mL⁻¹ penicillin and 100 µg mL⁻¹ streptomycin (all PAN-Biotech, Germany). The potential inhibitory effect of synthetic chalcones and curcumin on NF-kB activation was measured by incubating HeLa cells for 45-60 min with the indicated concentration of compounds or DMSO as vehicle, prior to stimulation with recombinant human TNF (30 ng mL⁻¹; gift from Daniela Männel) for 15 min (Western blot analysis) or 60 min (gene expression analysis). DMSO was adjusted to 0.01% (Western blot analysis) or 0.1% (gene expression analysis) in all conditions. Following TNF stimulation, cells were washed in cold PBS and harvested for either gene expression or Western blot analysis. For gene expression analysis, cells were lysed in the iScript RT-qPCR sample preparation reagent (Bio-Rad Laboratories, Germany) and subjected to cDNA synthesis using the iScript cDNA Synthesis kit (Bio-Rad Laboratories, Germany), following the manufacturer's instructions. Quantitative PCR was performed on a RotorGene Q (Qiagen, Germany) with SYBR Green I and HotStarTaq (Qiagen, Germany). Data were normalized to human Lamin A/C (LMNA)

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mRNA as previously described,¹⁷ and expressed as relative mRNA levels (% of vehicle or control). Forward and reverse real-time PCR primers specific for human LMNA, IL-6 and IL-8 have been described.¹⁸ Data shown represent the mean of three independent experiments. For Western blot analysis, cells were scrapped off mechanically; nuclear and cytosolic protein fractionation and NF-κB p65 immunoblotting were performed as previously described.¹⁸ Quality of the cytosolic and nuclear extraction was verified by immunoblotting with an antibody directed against the cytosolic marker a-tubulin (Santa-Cruz Biotechnology, Germany; 1:200). Chemoluminescence detection was performed using Amersham ECL Prime (GE Healthcare Life Sciences, Germany) and images were captured on an ImageOuant LAS 4000 mini imaging system (GE Healthcare Life Sciences, Germany). The immunoblot shown is representative of three independent experiments.

Data evaluation/statistical analysis

The data is presented as the mean \pm SD of several independent experiments (as indicated in figure captions) carried out in duplicates, triplicates or quadruplicates depending on the assay. A sigmoidal logistic function was used to fit dose–response curves and determine IC₅₀ values using a Microsoft Excel calculation sheet (Ed50plus, v1.0, April 2000, Mario H. Vargas, Instituto Nacional de Enfermedades, Mexico). Comparison between groups was made using a two-sided paired Student's *t* test. A *p* value < 0.05 was considered statistically significant. Levels of significance: ***, *p* < 0.001; **, *p* < 0.01; *, *p* < 0.05.

Conclusion

The direct covalent modification of protein thiols by alkylating agents can activate Nrf2 and inhibit NF- κ B and hence lead to the dampening of inflammatory responses.⁴ To exploit this phenomenon for therapeutic interventions, fine tuning of electrophilicity is desirable. If the electrophilicity is too low, the compound is unlikely to be effective at concentrations that are pharmacologically accessible. If the electrophilicity is too high, the compound may react non-specifically with a broad range of protein thiols or may be neutralized by glutathione conjugation. In contrast, compounds with intermediate electrophilicity may avoid rapid clearance and target a relatively narrow range of proteins, namely those with exposed and/or activated thiolate residues poised to be involved in protein redox regulation, like those in Keap1 or NF- κ B.

Thus, it appears critical to hit the right thiol reactivity window.¹⁹ We pursued the strategy to first classify electrophiles by *in vitro* kinetics (second order rate constant determination for the reaction with cysteamine), followed by probing the same electrophiles for their cellular effects. A similar approach used theoretical chemistry calculations to predict the electrophilicity of aryl nitriles, successfully explaining structure activity relationships for the inhibition of cysteine proteases.²⁰

Out of fourteen $\alpha\text{-}X\text{-}TMCs,$ three compounds stood out as potent anti-inflammatory agents, namely $\alpha\text{-}CF_3\text{-}TMC,\ \alpha\text{-}Br\text{-}$

TMC and α -Cl-TMC. These were highly efficient in activating Nrf2 and inhibiting NF- κ B, with corresponding effects on their respective transcriptional gene products. Notably, suppression of iNOS activity was observed with IC₅₀ values in the nanomolar range. Although alternative electrophilicity-independent modes of action cannot be strictly ruled out at this point, the inactivity of the most electrophilic compounds α -CN-TMC and α -NO₂-TMC can be rationalized by their predicted rapid reaction with intracellular GSH, which is present at millimolar concentrations.²¹

As a scaffold the chalcones facilitate straight forward access to libraries of fine-tuned electrophiles, allowing detailed studies of the relationship between electrophilicity and biological activity. In future work such electrophiles may be linked to molecules containing non-covalent recognition motifs, thus directing the electrophilic warhead²² to particular protein targets.²³

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