Differential effects of the essential oils of *Lavandula luisieri* and *Eryngium duriaei* subsp. *juresianum* in cell models of two chronic inflammatory diseases

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**Abstract**

**Context:** Effective drugs to treat osteoarthritis (OA) and inflammatory bowel disease (IBD) are needed.

**Objective:** To identify essential oils (EOs) with anti-inflammatory activity in cell models of OA and IBD.

**Materials and methods:** EOs from *Eryngium duriaei* subsp. *juresianum* (M. Lainz) M. Lainz (Apiaceae), *Laserpitium eliasii* subsp. *thalictrifolium* Sennen & Pau (Apiaceae), *Lavandula luisieri* (Rozeira) Rivas-Martínez (Lamiaceae), *Othanthus maritimus* (L.) Hoff. & Link (Asteraceae), and *Thapsia villosa* L. (Apiaceae) were analyzed by GC and GC/MS. The anti-inflammatory activity of EOs (5–200 μg/mL) was evaluated by measuring inducible nitric oxide synthase (iNOS) and nuclear factor-κB (NF-κB) activation (total and phosphorylated IκB-α), in primary human chondrocytes and the intestinal cell line, C2BBe1, stimulated with interleukin-1β (IL-1β) or interferon-γ (IFN-γ), IL-1β and tumor necrosis factor-α (TNF-α), respectively.

**Results:** The EO of *L. luisieri* significantly reduced iNOS (by 54.9 and 81.0%, respectively) and phosphorylated IκB-α (by 87.4% and 62.3%, respectively) in both cell models. The EO of *E. duriaei* subsp. *juresianum* caused similar effects in human chondrocytes, but was inactive in intestinal cells, even at higher concentrations. The EOs of *L. eliasii* subsp. *thalictrifolium* and *O. maritimus* decreased iNOS expression by 45.2 ± 8.7% and 45.2 ± 6.2%, respectively, in C2BBe1 cells and were inactive in chondrocytes. The EO of *T. villosa* was inactive in both cell types.

**Discussion and conclusion:** This is the first study showing anti-inflammatory effects of the EOs of *L. luisieri* and *E. duriaei* subsp. *juresianum*. These effects are specific of the cell type and may be valuable to develop new therapies or as sources of active compounds with improved efficacy and selectivity towards OA and IBD.

**Introduction**

An increasing number of reports describes anti-inflammatory properties of many natural products and compounds. Nonetheless, very few studies have been dedicated to evaluate their therapeutic potential on chronic inflammatory diseases. These remain important therapeutic targets due to their high prevalence and lack of effective therapies. Indeed, most studies concerning anti-inflammatory properties of natural products have been performed in cell or animal models of acute inflammation. These models do not entirely reflect pathological mechanisms and cells involved in chronic inflammatory diseases.

Among natural products, essential oils (EOs) are particularly interesting to look for compounds with pharmacological activities, since they are complex mixtures of a huge diversity of low molecular weight (<300 Da) lipophilic compounds. These characteristics may represent favorable pharmacokinetic properties (Miguel, 2010).

Recent reviews on the anti-inflammatory potential of EOs fully highlight that these plant extracts and some of their components are useful therapeutic alternatives, modulating several molecular targets of the acute inflammation cascades, namely in cells of the immune system, like monocytes and macrophages (Adorjan & Buchbauer, 2010; Miguel, 2010). There is, however, an evident lack of information on the potential of EOs as modulators of chronic inflammatory diseases, especially involving cells unrelated to the immune system. Exceptions are two reports from our group (Neves et al., 2010; Rufino et al., 2014) describing anti-inflammatory effects of the EO of *Juniperus*...
oxycedrus L. subsp. oxycedrus (Cupressaceae) and α-pinene in a cell model of osteoarthritis (OA).

Chronic inflammatory diseases, in general, lead to the upregulation of a series of enzymes and signaling molecules that bring about the characteristic inflammation and tissue destruction. Likewise, OA and inflammatory bowel disease (IBD) share many features and mechanisms and are largely dependent on the transcription factor, nuclear factor-κB (NF-κB). In chronic inflammatory diseases, the onset and perpetuation of inflammatory responses and tissue destruction are primarily dependent on the transcription factor, nuclear factor-κB (NF-κB) (Tak & Firestein, 2001). This promotes the expression of inflammation-related genes, including cytokines (IL-1β, TNF-α, IL-6, etc.) and enzymes. One of these, the inducible nitric oxide synthase (iNOS), produces large amounts of nitric oxide (NO), a potent and destructive inflammatory mediator that plays an important role in the development and progression of OA and IBD. For instance, our previous study showed that iNOS expression, NO production, and NF-κB activity are spontaneously increased in chondrocytes isolated from OA patients compared with those isolated from non-affected human cartilage (Rosa et al., 2008). Increased mucosal and plasma NO concentrations associated with augmented iNOS activity have also been demonstrated in active IBD (Quenon et al., 2013). Moreover, NF-κB promotes the expression of specific proteases that degrade the extracellular matrix, causing the characteristic tissue destruction of OA and IBD (Goldring & Otero, 2011; Jobin & Sarton, 2000; Wielockx et al., 2004).

Therefore, this transcription factor constitutes an attractive target for anti-inflammatory therapeutic interventions both in OA and in IBD (Berenbaum, 2004; Goldring & Otero, 2011; Jobin & Sarton, 2000; Marcu et al., 2010). Hence, we proposed to study the ability of EOs to inhibit NF-κB activation in cell models of these diseases.

To address this purpose, EOs to be screened were selected considering two major criteria: (i) to collectively ensure the highest diversity of compounds from the chemical classes usually found in EOs; and (ii) the availability of ethnopharmacological information or previous evidence of anti-inflammatory activity. For this, several EOs, isolated at laboratory from native or endemic species of the Iberian flora, were first fully characterized by GC-MS analysis, as described below, the EOs were stored at −70°C in hermetically sealed amber glass vials. α-Pinene (purity ≥ 98%) was purchased from Sigma Chemical Co. (St. Louis, MO).

### Materials and methods

#### Essential oils

EOs from the aerial parts of Eryngium duriaeii subsp. juresium (M. Lainz) M. Lainz (Apiaceae), from the aerial parts of Lasertepium eliasi subsp. thalictroidium Sennen & Pau (Apiaceae), from the leaves and flowers of Lavandula luister (Rozeira) Rivas-Martinez (Lamiaceae), from the aerial parts of Orthanthus maritimus (L.) Hoff. & Link (Asteraceae), and from the aerial parts of Thapsia villosa L. (Apiaceae) were picked from the collection of plant extracts of the Faculty of Pharmacy, University of Coimbra. Plant materials, collected at different locations in the center region of Portugal, were identified by plant taxonomists (Ana Cristina Tavares, PhD and Celia Cabral, PhD, University of Coimbra). Voucher specimens of plant material were deposited at the Herbarium of the Botanic Garden, University of Coimbra (COI) or at the Herbarium of the Faculty of Pharmacy, University of Coimbra.

All EOs were prepared at laboratory by water distillation using a Cleveenger-type apparatus (EDQM, 2007). After chemical analysis, as described below, the EOs were stored at −70°C in hermetically sealed amber glass vials. α-Pinene (purity ≥ 98%) was purchased from Sigma Chemical Co. (St. Louis, MO).

#### Analysis of essential oils

The composition of each EO was established immediately after extraction following a combined methodology of GC and GC/MS. Analytical GC was performed in a Hewlett-Packard 6890 (Agilent Technologies, Palo Alto, CA) gas chromatograph with a HP GC ChemStation Rev. A.05.04 data handling system (Agilent Technologies, Palo Alto, CA), equipped with a single injector and two-flame ionization detectors (FID). A graphpak divider (Agilent Technologies Palo Alto, CA, part no. 5021-7148) was used for simultaneous sampling to two Supelco (Supelco, Bellefonte, PA) fused silica capillary columns with different stationary phases: SPB-1 (polydimethylsiloxane 30 m × 0.20 mm i.d., film thickness 0.20 μm) and SupelcoWax-10 (polyethylene glycol 30 m × 0.20 mm i.d., film thickness 0.20 μm). Oven temperature program: 70–220°C (3°C/min), 220°C (15 min); injector temperature: 250°C; carrier gas: helium adjusted to a linear velocity of 30 cm/s; split ratio 1:40; detectors temperature: 250°C. GC-MS was performed in a Hewlett-Packard 6890 gas chromatograph (Agilent Technologies, Palo Alto, CA) fitted with a HP1-fused silica column (polydimethylsiloxane 30 m × 0.25 mm i.d., film thickness 0.25 μm), interfaced with an Hewlett-Packard mass selective detector 5973 (Agilent Technologies, Palo Alto, CA) operated by HP Enhanced ChemStation software (Agilent Technologies, Palo Alto, CA), version A.03.00. GC parameters were as described above; interface temperature: 250°C; MS source temperature: 230°C; MS quadrupole temperature: 150°C; ionization energy: 70 eV; ionization current: 60 μA; scan range: 35–350 units; scans/s: 4.51.

Components of each EO were identified considering their retention indices (RIs) on both SPB-1 and SupelcoWax-10 columns, and their mass spectra. RIs, calculated by linear interpolation relative to retention times of C₈–C₂₃ n-alkanes (Vandendool & Kratz, 1963), were compared with those of authentic samples or data available in digital banks (Acree,
2004; El-Sayed, 2012; Linstrom, 2013). Acquired mass spectra were compared with reference spectra from the laboratory database, Wiley/NIST database (EDQM, 2007) and literature data (Adams, 1995; Joulain, 1998). Relative amounts of individual components were calculated based on GC raw data without further correction.

Before beginning the cell assays, each EO was re-analyzed and the composition was compared with that obtained on the first analysis. No significant differences, either qualitative or quantitative, were observed. Table 1 shows the composition of each essential oil obtained in the second analysis, performed immediately before the pharmacological assays. For these assays, the EOs were diluted in dimethylsulfoxide (DMSO) and then dispersed in the culture medium to achieve final concentrations of 5–200 μg/mL. The final DMSO concentration did not exceed 0.1% (v/v).

**Cartilage samples and cell cultures**

Human chondrocytes were isolated by enzymatic digestion (Rosa et al., 2009) of knee cartilage from the distal femoral condyles of multi-organ donors (20–67 years old, mean = 48.5, n = 17) and, with informed consent, of patients (59–71 years old, mean = 64.0, n = 4) undergoing total knee arthroplasty at the Orthopedic Department of the University and Hospital Center of Coimbra (CHUC). The cartilage samples presented variable degrees of degradation, ranging from intact to severely damaged. All procedures were approved by the Ethics Committee of CHUC (protocol approval numbers 8654/DC and HUC-13-05).

Chondrocyte cultures were established from non-pooled cartilage samples. Before each experiment, the cells were serum-starved for at least 6 h and maintained thereafter in serum-free culture medium. The human colorectal adenocarcinoma cell line, C2BBe1 (ATCC CRL-2102), was cultured as recommended by the American Type Culture Collection. Before treatments, C2BBe1 cells were cultured for 7–9 d until reaching a hyperconfluent state to induce differentiation.

Primary chondrocyte and C2BBe1 cultures were treated with 10 ng/mL IL-1β (Peprotech EC, London, UK) or a cytokine mixture (CytMix, 1000 U/mL IFN-γ, 10 ng/mL IL-1β and 10 ng/mL TNF-α, Peprotech EC), respectively, in the presence or absence of the EOs or the positive control compound, Bay 11-7082 (EMD Millipore, Billerica, MA), for the periods indicated in figure legends. In all cases, the EOs, α-pinene, or Bay 11-7082 were added to the cell cultures 30 min before the respective pro-inflammatory stimulus.

**Cell viability assay**

The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium (MTT; Sigma, St. Louis, MO) reduction assay, a quantitative colorimetric method based on the reduction of the MTT salt by the mitochondrial enzymes of viable cells (Mosmann, 1983), was used to determine the cytotoxicity of the EOs in the cell models used and to select non-cytotoxic concentrations of each EO. Briefly, the cells were incubated with each EO in concentrations ranging from 5 to 200 μg/mL, for 24 h in the presence or absence of the inflammatory stimulus. Then, the culture medium was replaced with fresh medium containing 5 μg/mL MTT and the cells further incubated for 30 min. The resulting dark blue crystals of formazan were then dissolved in acidified isopropanol and the absorbance of the corresponding solution, which is directly proportional to the number of living cells, was measured in an automatic plate reader (SLT, St Anton, Austria) set at a test wavelength of 570 nm and a reference wavelength of 620 nm. The absence of cytotoxic effects in each cell model was defined as the concentration of EO eliciting an absorbance reading of at least 80% of the value measured in the respective control untreated cells or cells treated with the inflammatory stimulus for the same time period.

**Nitric oxide production**

The concentration of nitrite, which reflects NO production, was measured in the cell-free supernatants collected from chondrocyte or C2BBe1 cell cultures treated for 24 h with IL-1β or CytMix, respectively, in the presence or absence of different concentrations of each EO. Nitrite concentration was measured using the spectrophotometric method based on the Griess reaction (Green et al., 1982).

**Western blot**

Total and cytoplasmic cell extracts were prepared and subjected to western blots as described previously (Rosa et al., 2009). The membranes were probed with the following antibodies: mouse monoclonal anti-human iNOS (R&D Systems, Minneapolis, MN), rabbit polyclonal anti-human IκB-α or mouse monoclonal anti-human phospho-IκB-α (Cell Signaling Technology, Inc., Beverly, MA), and anti-rabbit or anti-mouse alkaline phosphatase-conjugated secondary antibodies (GE Healthcare, Little Chalfont, UK). Mouse anti-human β-tubulin (Sigma, St. Louis, MO) or anti-human actin monoclonal antibodies (EMD Millipore Corporation, Billerica, MA) were used to detect β-tubulin or actin as loading controls. Immune complexes were detected with the Enhanced ChemiFluorescence reagent (GE Healthcare, Little Chalfont, UK). The results were normalized by calculating the ratio between the intensities of the bands corresponding to the protein of interest and the protein used as a loading control.

**Statistical analysis**

Results are presented as mean ± SEM. Statistical analysis was performed using the GraphPad Prism (version 5.00, GraphPad Software Inc., San Diego, CA). SPSS software (version 17.0, SPSS Inc., Chicago, IL) was used to assess the normality (Kolmogorov–Smirnov test) and homogeneity of variances to determine whether the conditions required to apply parametric tests were satisfied. As in all cases such conditions were observed, the statistical analysis was performed using the paired *t*-test for comparison of each condition with its respective control and one-way ANOVA for comparison of all conditions. Results were considered statistically significant at *p* < 0.05.
Table 1. Classes of compounds and major constituents of the essential oils tested.

<table>
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<tr>
<th></th>
<th>L. luisieri</th>
<th>E. duriae subsp. juresianum</th>
<th>O. maritimus</th>
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Results

Composition of the essential oils

Chemical characterization of the EOs of L. luisci, E. duriae subsp juresianum, O. maritimus, L. eliasi, and T. villosa allowed the identification and quantification of 94, 25, 30, 42, and 14 compounds, respectively, comprising at least 85% of the respective composition. Table 1 summarizes the composition of each EO, indicating the relative amounts of grouped components and the most representative compounds (≥2%).

Thirty-eight compounds, comprising seven monoterpen hydrocarbons, 12 oxygen-containing monoterpenes, including four neryl derivatives, eight sesquiterpene hydrocarbons, 12 oxygen-containing sesquiterpenes, the phenylpropanoid, methyleugenol, and the aliphatic, n-nonane, were found in concentrations over 2.0% in, at least, one EO.

Evaluation of cytotoxicity and selection of non-cytotoxic concentrations of the essential oils

Neither the EOs nor the α-pinene was cytotoxic to C2BBe1 cells in concentrations up to 200 μg/mL, either in the presence or in the absence of the pro-inflammatory cytokine mixture (Figure 1C and D). On the contrary, in human chondrocytes, only the EO of T. villosa had no significant cytotoxic effects at that concentration (Figure 1A and B). In concentrations up to 50 μg/mL for 24 h, the EOs of E. duriae subsp, juresianum, L. eliasi, and L. luisci did not affect chondrocyte viability either in the presence or in the absence of IL-1β. The EO of O. maritimus showed no cytotoxicity at concentrations up to 100 μg/mL (Figure 1A and B). Therefore, subsequent experiments were performed using the non-cytotoxic concentrations identified for each EO in each cell model, as shown in Figure 1.

Effect of the EOs on cytokine-induced iNOS expression and NO production

Treatment of human chondrocytes (Figure 2) and C2BBe1 cells (Figure 3) with IL-1β or CytMix, for 24 h, strongly induced the expression of iNOS. Accordingly, human chondrocytes stimulated with IL-1β produced almost six-fold more NO than control cells, as shown by the concentration of nitrite in the respective culture medium (15.6 ± 1.9 μM and 2.7 ± 0.8 μM, respectively, p < 0.001, Figure 4).

Surprisingly, no differences in nitrite concentration were detected in C2BBe1 cell cultures treated with the CytMix for various periods (6–48 h) in comparison with untreated cells (data not shown). As measurement of NO production was used just as a rapid screening assay and iNOS protein was detected in C2BBe1 cells treated with 200 μg/mL of the EOs of L. luisci and E. duriae subsp. juresianum, respectively. Moreover, total IxB-α levels in chondrocytes treated with 200 μg/mL of the EO of E. duriae subsp. juresianum were not significantly different from those observed in control cells, indicating that this concentration completely prevented the response induced by IL-1β.

In C2BBe1 cells, the EO of L. luisci, at a concentration of 200 μg/mL, achieved the highest inhibition of iNOS expression (81.0 ± 5.2%). At a concentration of 50 μg/mL, iNOS levels were similarly reduced in C2BBe1 cells and in chondrocytes (Figure 3). The EOs of L. eliasi and O. maritimus significantly reduced iNOS protein levels (by 45.2 ± 8.7% and 45.2 ± 6.2%, respectively) in C2BBe1 cells, but were much less effective than the EO of L. luisci (Figure 3). Noticeably in these cells, the EO of E. duriae subsp. juresianum had no significant effect on iNOS expression (Figure 3), even at a concentration of 100 μg/mL which is several fold higher than those tested in human chondrocytes. Neither the EO of T. villosa nor the α-pinene had any significant effect (Figure 3).

Effect of the EOs of L. luisci and E. duriae subsp. juresianum on NF-κB activation

In response to appropriate stimuli, the NF-κB inhibitory protein, IxB-α, is phosphorylated and subsequently degraded, which releases NF-κB and allows its translocation to the nucleus to induce the expression of target genes (Hayden & Ghosh, 2008; O’Dea & Hoffmann, 2009). To assess NF-κB activation, we evaluated the cytoplasmic levels of phosphorylated and total IxB-α.

The EOs more effective in inhibiting iNOS expression and NO production were selected to evaluate their ability to inhibit IL-1β-induced NF-κB activity in human chondrocytes and C2BBe1 cells. In human chondrocytes, the EOs of L. luisci and E. duriae subsp. juresianum, at the concentration of 200 μg/mL, completely inhibited IL-1β-induced IxB-α phosphorylation, achieving an effect similar (p = 0.82 and p = 0.25, respectively) to that elicited by Bay 11-7082, a specific inhibitor of IxBα-phosphorylation (Mendes Sdos et al., 2009) used as a positive control (Figure 5). Total IxB-α levels relative to those in control cells increased from 4.3 ± 1.5% (n = 4) in IL-1β-treated chondrocytes to 47.9 ± 10.7% (n = 4) and 55.6 ± 9.9% (n = 4) in cells treated with 200 μg/mL of the EOs of L. luisci and E. duriae subsp. juresianum, respectively. Moreover, total IxB-α levels in chondrocytes treated with 200 μg/mL of the EO of E. duriae subsp. juresianum were not significantly different from those observed in control cells, indicating that this concentration completely prevented the response induced by IL-1β.

In C2BBe1 cells, the EO of L. luisci, at concentrations of 50 and 200 μg/mL, significantly reduced IxB-α phosphorylation to 77.3 ± 8.7% (n = 7) and to 37.7 ± 3.0% (n = 7), respectively, of the CytMix-induced response (Figure 6). Total IxB-α levels increased from 11.7 ± 1.2% in CytMix-treated cells to 30.4 ± 4.3% (n = 8) in L. luisci-treated cells relative to the control, indicating that the EO effectively decreased CytMix-induced NF-κB activation. Moreover, 200 μg/mL of this EO inhibited IxB-α phosphorylation to even a larger extent than the positive control, Bay 11-7082 (p = 0.028), while IxB-α degradation was similarly inhibited (p = 0.31).

Discussion

The results presented show clear differences in the ability of the EOs tested to inhibit relevant mediators of inflammation.
Figure 1. Viability of human chondrocytes (A and B) and C2BBe1 cells (C and D) treated with the EOs for 24 h in the absence (A and C) or presence (B and D) of the respective pro-inflammatory stimulus. Each column represents, at least, four independent experiments. The dotted line represents the limit below which cell viability is impaired. $p < 0.05$ and $$p < 0.01$ relative to the respective control (untreated) cells.
in the two cell models of OA and IBD used. Such differences are evident among distinct EOs, as well as comparing each one in the two cell models. Indeed, only the EO of *L. luisieri* was capable of significantly inhibiting inflammatory markers (iNOS expression and NF-κB activation) both in human chondrocytes and intestinal C2BBe1 cells. These results confirm our previous observation that this EO inhibits IL-1β-induced NO production in human chondrocytes (Neves et al., 2010). On the other hand, the EO of *E. duriaei* subsp.* juresianum did not significantly inhibit iNOS expression in intestinal cells over a wide range of concentrations, but was effective in human chondrocytes, even at much lower concentrations. Conversely, the EOs of *O. maritimus* and *L. eliasii* showed some activity in intestinal cells, but were inactive in chondrocytes. These results indicate that the EOs of *L. luisieri* and *E. duriaei* subsp. *juresianum* have cell type-specific anti-inflammatory effects.

To our knowledge, this is the first study showing anti-inflammatory effects of the EOs of *L. luisieri* and *E. duriaei* subsp. *juresianum*. Indeed, EOs from *Lavandula* ssp, namely *L. angustifolia*, *L. stoechas*, *L. multifida*, and *L. viridis*, have demonstrated antibacterial, antifungal, analgesic, and anti-inflammatory effects (Amira et al., 2012; Ghelardini et al., 1999; Kirmizibekmez et al., 2009; Zuzarte et al., 2011a,b), while the EO of *L. luisieri* has only been reported to have antifungal effects (Zuzarte et al., 2012). On the other hand, EOs from *Eryngium* ssp, including *E. duriaei* subsp. *juresianum*, have been shown to have anti-fungal and anti-bacterial properties (Cavaleiro et al., 2011; Celik et al., 2011), whereas anti-inflammatory effects were only reported for aqueous or alcoholic extracts from *Eryngium* species (Dawilai et al., 2013; Kupeli et al., 2006) not including *E. duriaei* subsp. *juresianum*.

α-Pinene from a commercial source (purity ≥ 98%) was unable to decrease iNOS expression in intestinal cells, while
Figure 3. Effect of EOs on iNOS protein expression in C2BBe1 cells left untreated (Ctrl) or treated with CytMix, for 24 h, after pre-treatment with each EO. The images shown are representative of, at least, three independent experiments. *p < 0.05, **p < 0.01, and ***p < 0.001 relative to cells treated with CytMix.

Figure 4. Effect of EOs on IL-1β-induced NO production in human chondrocytes left untreated (Ctrl) or treated with IL-1β, 10 ng/mL, for 24 h, after pre-treatment with each EO. Each column represents, at least, four independent experiments. *p < 0.05, and ***p < 0.001 relative to IL-1β-treated cells.
our previous studies showed that both commercial α-pinene (Rufino et al., 2014) and the EO of *J. oxycedrus* subsp. *oxycedrus* (76.4% α-pinene) and one of its fractions (93% α-pinene) (Neves et al., 2010) were effective in human chondrocytes. Interestingly, the EOs of *L. luisieri*, *O. maritimus*, and *L. eliasii*, the only ones containing significant amounts of α-pinene (2.3, 6.7, and 30.5%, respectively), were all effective in intestinal cells, but only that of *L. luisieri* was also effective in human chondrocytes, even though it presents the lowest α-pinene content. Taken together, these results suggest that α-pinene is unlikely the compound responsible for the inhibitory activities of the EOs of *L. luisieri*, *O. maritimus*, and *L. eliasii* observed in this study.

Similar considerations can be made about other compounds present in two or more of the EOs tested. Limonene, for instance, represents 58.8% of the EO of *T. villosa*, but only 2.7% of the EO of *L. eliasii*. Nevertheless, this EO was effective in intestinal cells while that of *T. villosa* was inactive in both cell types.

In contrast, the EOs of *L. luisieri* and *E. duriaeie* subsp. *juresianum* have distinct compositions, the first being rich in oxygenated monoterpenes, while the second is mainly composed of oxygenated and non-oxygenated sesquiterpenes. Nonetheless, they were both effective in reducing markers of inflammation in human chondrocytes, suggesting that distinct compounds are involved in the observed activities of these EOs.

Moreover, we cannot discount the possibility that different components of the EOs act in synergy or in antagonism to modulate their overall activity, contributing to the apparent discrepancies described above. This possibility is even more plausible as some of these compounds, like limonene (Chi et al., 2012), have been shown to exert anti-inflammatory effects in various cell and animal models.

Taken together, the results presented indicate that two of the EOs studied, those of *L. luisieri* and *E. duriaeiae* subsp.
juresianum, have significant cell type-specific anti-inflammatory effects that can be useful for the development of tissue-selective anti-inflammatory therapies.

The results obtained also show that the EOs of *L. luissieri* and *E. duriae* subsp. *juresianum* which were the most effective in inhibiting iNOS expression in human chondrocytes and/or intestinal epithelial cells also decreased cytokine-induced NF-κB activation. Since NF-κB is essential for iNOS expression (Taylor et al., 1998), these results strongly suggest that the observed inhibition of iNOS is mediated, at least in part, by the inhibitory effects of these EOs on NF-κB activation. The concentrations of the EOs of *L. luissieri* and *E. duriae* subsp. *juresianum* effective in inhibiting IL-1β-induced NF-κB activation in human chondrocytes were substantially higher than those that inhibited iNOS expression and NO production, suggesting that other mechanisms may contribute to the inhibitory activity of these EOs, at least in human chondrocytes. Since none of the EOs showed NO scavenging activity (data not shown), this can be discarded as a potential contributing mechanism. Other possibilities include direct inhibition of iNOS activity and inhibition of other signaling pathways that are required for iNOS expression in human chondrocytes (Mendes et al., 2002). More studies are required to identify the specific molecular targets of the EOs of *L. luissieri* and *E. duriae* subsp. *juresianum*. On the contrary, since EOs are complex mixtures of chemically diverse compounds, it is possible that distinct components have different targets, so that the effects observed result from the combined actions of individual compounds. Future work will be directed at fractionating each of these EOs and elucidating the chemical composition and pharmacological activity of each fraction in order to identify the active compound(s) and their specific molecular targets, as well as potential pharmacological interactions.

In summary, this study shows for the first time that the EOs of *L. luissieri* and *E. duriae* subsp. *juresianum* efficiently inhibit NF-κB activation and the expression of its target genes, namely iNOS, in cells unrelated to the immune system. Moreover, these EOs display differential effects in relevant cell models of OA and IBD and in response to distinct inflammatory stimuli. These properties may be of great value in the development of new therapies with improved efficacy and selectivity towards distinct chronic inflammatory diseases, namely OA and IBD.

**Declaration of interest**

The authors report that there are no declarations of interest. This work was supported by Grants CENTRO-07-ST24-FEDER-002006, PEst-C/SAU/LA0001/2011, Pest-OE/SAU/UI0177/2011, and PTDC/EME-TME/113039/2009 and the PhD fellowship, SFRI/BD/47470/2008, to Rufino A. T. from FEDER through the programs COMPETE and QREN and by national funds through the Portuguese Foundation for Science and Technology (FCT).

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**Pharm Biol, Early Online:** 1–11


