Investigation of the Effect of Hyperforin and Hypericin on Inflammatory Response in RAW 264.7 Macrophages

Hiperforin ve Hiperisinin RAW 264.7 Makrofajında İnflamatuar Yanıt Üzerine Etkisini İncelenmesi

Mehmet Berköz¹, Oruc Allahverdiyev², Metin Yıldırım³

¹Department of Pharmaceutical Biotechnology, Faculty of Pharmacy, Van Yuzuncu Yil University, Van, Turkey
²Department of Pharmacology, Faculty of Pharmacy, Van Yuzuncu Yil University, Van, Turkey
³Department of Biochemistry, Faculty of Pharmacy, Mersin University, Mersin, Turkey

ABSTRACT

Objective: This study aims to determine the anti-inflammatory property of hypericin and hyperforin using in vitro model.

Materials and Methods: In this study, 0, 25, 50, 75 and 100 nM hyperforin and 0, 2.5, 5, 7.5 and 10 μM hypericin were treated to lipopolysaccharide (LPS) stimulated RAW 264.7 macrophages. We aimed to display the anti-inflammatory effects of hypericin and hyperforin via measuring prostaglandin E₂ (PGE₂) and nitric oxide (NO) production and cyclooxygenase-2 (COX-2) and inducible nitric oxide synthase (iNOS) gene expression in LPS induced RAW 264.7 macrophages.

Results: Our study has shown that all applied concentration of hyperforin and hypericin decreased nitric oxide (NO) production and inducible nitric oxide synthase (iNOS) and cyclooxygenase-2 (COX-2) gene expression in RAW 264.7 macrophages, significantly. Similarly, all applied concentration of hyperforin and hypericin decreased prostaglandin E₂ (PGE₂) production but this decrease was statistically significant only in groups treated with 75 and 100 nM hyperforin and 7.5 and 10 μM hypericin.

Conclusion: This indicates that hyperforin and hypericin inhibited NO and PGE₂ production and iNOS and COX-2 gene expression in RAW 264.7 macrophages in a dose-dependent manner. Inhibition of NO and PGE₂ production by hyperforin and hypericin is a result of the inhibition of iNOS and COX-2 gene expression. For this reason, it is possible to say that hypericin and hyperforin can be used to reduce the inflammatory response.

Key Words: Hypericin, hyperforin, RAW 264.7, nitric oxide, prostaglandin E₂

ÖZET

Amaç: Bu çalışma in vitro model kullanarak hiperforin ve hiperisinin anti-inflamatuar özelliğini belirlemeyi amaçlamaktadır.

Gereç ve Yöntem: Bu çalışmada, lipopolisakkarit (LPS) ile uyurulmuş RAW 264.7 makrofajları, 0, 25, 50, 75 ve 100 nM hiperforin ve 0, 2,5, 5, 7,5 ve 10 μM hiperisin ile muamele edilmişdir. LPS ile indüklennmiş RAW 264.7 makrofajlardında, prostaglandin E₂ (PGE₂) ve nitrik oksit (NO) üretimi ile siklooksijenaz-2 (COX-2) ve induklenber nitrik oksit sentaz (iNOS) gen ekspresyonunu ölçürek hiperforin ve hiperisinin antienflamatuar etkilerini göstermeyi amaçlamaktadır.

Bulgular: Yaptığımız çalışma, uygulanan tüm hiperforin ve hiperisinin konsantrasyonlarının, RAW 264.7 makrofajlarında nitrik oksit (NO) üretimini ve indüklenebilir nitrik oksit sentaz (iNOS) ve siklooksijenaz 2 (COX-2) gen ekspresyonlarını önemli ölçüde azalttığını göstermektedir. Benzer bir şekilde, uygulanan tüm hiperforin ve hiperisinin konsantrasyonlarının prostaglandin E₂ (PGE₂) üretimini azalttı ancak bu düşüşün sadece 75 ve 100 nm hiperforin ile 7,5 ve 10 μM hiperisin uygulanan gruplarda istatistiksel olarak anlamlı olduğu görülmüştür.

Sonuç: Bu durum, hiperforin ve hiperisinin doza bağımlı bir şekilde RAW 264.7 makrofajlarındaki NO ve PGE₂ üretimini ve iNOS ve COX-2 gen ekspresyonunu inhibe ettiği göstermektedir. Bu nedenle, hiperforin ve hiperisinin inflmatuar yanıt azaltmak için kullanlabileceği söylenmek mümkündür.

Anahtar Kelimeler: Hiperisin, hiperforin, RAW 264.7, nitrik oksit, prostaglandin E₂

Introduction

Acute inflammatory response represents an initial protective mechanism in the body (1,2). However, excessive and chronic inflammation results in severe damage of cells and tissues (3). Emerging evidences support the hypothesis that chronic inflammation plays a critical role in various

*Sorumlu Yazar: Öğr. Göz. Dr. Mehmet Berköz, Yüzüncü Yıl Üniversitesi Eczacılık Fakültesi, Farmasötik Biyoteknoloji Anabilim Dalı, Zeve Kampüsü, VAN, GSM: +90 (536) 719 71 24, E-posta Adresi: mehmet_berkoz@yahoo.com

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pathological conditions, including atherosclerosis, autoimmune disorders, neurodegenerative diseases, and inflammation related various human cancers (4,5).

Nitric oxide (NO) is enzymatically formed from the terminal guanidine-nitrogen of L-arginine by a group of enzymes called nitric oxide synthesis (NOS) (6). The enzymes are classified into three isoforms; endothelial NOS (eNOS), neuronal NOS (nNOS) and inducible NOS (iNOS). eNOS is a constitutive enzyme present primarily in endothelium. nNOS is a neurally associated constitutive NOS. iNOS is an inducible enzyme the expression of which in macrophages, neutrophils, endothelium and epithelium is induced by LPS and several cytokines. iNOS can produce relatively large concentrations of NO for a long time and therefore, has been associated with pathophysiological events (7,8).

Cyclooxygenases (COXs) are the key enzymes required for the synthesis of prostaglandins (PGs) from arachidonic acid. Cyclooxygenases has three known forms; COX-1, COX-2 and COX-3. COX-1 is constitutively expressed in the gastrointestinal tract and is considered to be responsible for maintaining mucosal integrity. COX-2 is induced by pro-inflammatory mediators and involved in the amplification of inflammation and pain. On the other hand, COX-3 enzyme is expressed only in the brain. COX-2 expression could be induced in neutrophils, monocytes and endothelial cells by many cytokines including IL-1 and TNF-α (9). When COX-2 expression is induced in cells, a proportionately larger amount of PGE$_2$ is produced by non-enzymatic conversion. For this reason, COX products, mainly PGE$_2$ are well-known mediators of the cardinal features of inflammation-pain, oedema, erythema and warmth (10).

Hypericum perforatum L. is a member of Hypericaceae family. It is a herbaceous perennial plant native to Asia and Europe. At present, they are either naturally occurring on, or which have been introduced to, every continent in the world, except Antarctica. It is a beneficial plant used in America and Europe for medical properties (11). It contains lots of biologically active compounds including: naphthodianthrones (hypercin and pseudohypercin), prenylated acylphloroglucinols (hyperforin and adhyperforin), flavonoids (quercetin, hyperoside, rutin, and quercitrin), essential oil rich in sesquiterpenes, and xanthones (1,3,6,7-tetrahydroxyanthone) (12). In medicine, extracts of Hypericum have an extensive range of pharmacological properties, the best important utilization of Hypericum perforatum L. is using for treatment of mild to moderate depression and currently it has an important role in the treatment of depressive disorder taking into account of these pharmacological activities. Hypericum perforatum L. preparations appear as one of the major herbal dietary supplements in the whole world, also it has antiviral, antitumoral, antimicrobial, antioxidant and anti-inflammatory activities and it is used for treatment of wound healing (11,12).

The most common compounds in Hypericum perforatum L. are hypercin and hyperforin. Hypercin has been at the forefront of many studies investigating anti-viral, anti-cancer, anti-depressive and anti-microbial activities. On the other hand, hyperforin has anti-depressive, anti-bacterial, anti-tuberculosis and anti-fungal activities (13). Although many biological activities of hypercin and hyperforin are well known, what role it plays in the inflammatory response is not yet known (13,14).

This study was designed to estimate the inhibitory effects of hyperforin and hypercin on inflammatory response by LPS stimulation in RAW 264.7 macrophages via measuring the production of NO and PGE$_2$ and mRNA expression of iNOS and COX-2.

Materials and Methods

Cell culture

RAW 264.7 macrophages were purchased from American Type Culture Collection (ATCC) and the cells were incubated at 37°C and 5% CO$_2$. All the chemicals used in our study were purchased from Sigma-Aldrich and Merck companies and all products were in cell culture purity. The cells were passaged when they were cultivated at a density of 70–80% full of the dish. Medium was removed and 1× Phosphate buffered saline (PBS) was used to wash the dish. We subsequently tapped the dish and used fresh medium to wash down the cells. The fallen cells were collected in centrifuge tube, and we centrifuged the tube with 1000 rpm for 5 minute. After that, the supernatant was sucked up and the pellet was dispersed by fresh medium. Finally, the cells were seeded in dishes or wells with a proper density.

After the cells grew stably and reached at a density of 2·3·10$^6$ cells/mL, the medium was removed and 1× PBS was used to gently wash the dish. The medium was replaced serum-free DMEM. The cells were induced by incubation in medium containing 0.1 µg/mL LPS (E. coli 0127: E8).
Hyperforin and hypericin were dissolved in DMEM. These dissolution compounds were treated cells together with LPS.

**Cell viability assay**

The cellular toxicity of hyperforin and hypericin were determined by \( (3\cdot4,5\text{-Dimethylthiazol-2-yl})\cdot2,5\text{-Diphenyltetrazolium Bromide (MTT) assay} \) (15). After 24 hr of incubation with the cells, hyperforin and hypericin were removed by washing the cultured cells three times with medium. Then, the cells were collected by centrifugation and the pellets were suspended in fresh medium. The MTT solution was prepared using phosphate buffered solution (PBS) at 5 mg/mL concentration. MTT solution was added to the culture plate with cells and then incubated at 37°C for 4 hours. The solution of 0.4 N HCl in isopropyl alcohol was added to the culture well to dissolve the formed tetrazolium salt. The results were measured by ELISA which read the absorption at 540 nm in ELISA microplate reader (Thermo Fisher Scientific, Watham, MA, USA).

The experiment was repeated by three times. Medium without cells was used as blank.

**Determination of NO production**

Nitrite and peroxynitrite levels in the cell supernatant were measured using the Griess method (16) to determine NO level in the cell culture medium. Nitrite and peroxynitrite are stable end products of NO in vitro system. By this way, the level of NO synthesis by iNOS can be evaluated. Griess reagent was prepared by mixing 1 volume of 1 % sulfanilamide solution and 1 volume of 0.1% N-l-naphthyl ethenediamine dihydrochloride prepared in 2.5% H₃PO₄. Tested cells were treated by 0.1 µg/mL LPS. The tested compounds were added in gradient concentrations into the medium with cultured cells before the adding of the stimulants. After 24 hr, 48 hr and 72 hr of stimulation, 50 µL of supernatant of the culture wells was taken and mixed with 50 µL freshly prepared Griess reagent. After incubation at room temperature for 10 minutes, the ELISA plate was centrifuged for 5 minutes at 1000 rpm. The results were obtained from ELISA by reading the absorption at 540 nm. This reaction was repeated three times for each sample. Accumulation of nitrite was calculated as nmol of nitrite per 10⁶ cells from three independent experiments. The culture medium without cells was used as blank. Sodium nitrate (NaNO₂) was diluted to 3 µM, 25 µM, 50 µM and 100 µM and used as standard solutions.

**Determination of PGE₂ production**

Prostaglandins (PGs) are lipid compounds derived from arachidonic acid. Like all PGs, PGE₂ can be produced by several tissues including the gut, the uterus, blood vessels, bladder, placenta, brain and cells of the immune system. In some pathologic conditions such as tissue injury, inflammation and many cancers the production of PGE₂ is increased in bodily fluids. The principle of PGE₂ immunoassay is that PGE₂ from culture medium competes with horseradish peroxidase (HRP) - labeled PGE₂. After both PGE₂ and HRP-labeled PGE₂ bind to the antibody sites, a substrate solution is added to evaluate the bound enzyme activity. The concentration of PGE₂ present in the sample is inversely proportional to the intensity of the color at the end of the immunoassay.

**Gene expression analysis**

**RNA extraction:** Culture medium was removed after 4 or 5 hours of incubation. 300 µL lysis buffer was added per 10⁶ cells. Then, 100 µL of protein/DNA precipitating buffer was mixed and centrifuged to remove the protein and DNA. Supernatant was removed to new eppendorf tube and mixed with 300 µL isopropanol. The tube was put in -70 °C freezer for more than 24 hours to precipitate RNA. RNA pellet was washed by 70% ethanol and then dissolved in DEPC-water. For tissue samples, TriReagent (Sigma) was used to homogenize the tissue samples in homogenizers. Chloroform was added to remove DNA and protein. After centrifugation, the water phase was removed to a new tube and 1 volume isopropanol was mixed to precipitate RNA. RNA was washed by 70% ethanol to remove isopropanol and dissolved into DEPC-water. The concentration of extracted RNA was quantitated by measuring the absorption in OD₂₆₀ and OD₂₈₀ in spectrophotometer. The ratio of OD₂₆₀/OD₂₈₀ is between 1.6 to 1.8 for samples were used for the following experiments.

**cDNA synthesis:** Total RNA extracted from cultured cells and tissue samples were used as template for reverse transcription. 300 ng total RNA was added to 25 µL total volume of reaction mix containing 0.6 mM dNTPs (Promega), 10 units Rnase inhibitor (Clonetech), Oligo(dT)₁₅ primer (Promega) and 100 units of MMLV-RT (Promega). The whole reaction mix was incubated for 20 minutes at 42°C. Then, the activity of reverse transcriptase was stopped via heating at 94°C for 5 minutes. The synthesized cDNA was chilled on ice immediately for PCR reaction or stored at -70°C for later analysis.

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**Real time PCR Analysis:** The production of PCR products was determined by measuring the SYBR Green fluorescence signal. SYBR Green DNA enters small, large cavities in double strands. The unbound dye in the solution emits very low fluorescence. However, as the binding of SYBR Green to DNA increases, the fluorescence emission at a wavelength of 530 nm rises. During PCR, the increase in SYBR Green fluorescence composition increases with the proportion of directly produced double-stranded DNA. Real time quantitative PCR was applied in the direction of the manufacturer using the Invitrogen Universal EXPRESS SYBR Greener qPCR SuperMixes, Two-Step qRT-PCR kit on the Roche LightCycler 480 II device and primers and probes suitable for the gene region (17). In our study, β-actin gene was used as the house-keeping gene. Concentration values of iNOS and COX-2 gene expression of each sample were proportional to the concentration of β-actin of the same sample. The obtained values were multiplied by 100 and gene expression levels were calculated for each sample and the results obtained were compared. Relative quantitation was also used when expression changes were calculated and the expression of iNOS and COX-2 genes in LPS and different concentration of hyperforin and hypericin treated RAW 264.7 macrophages was expressed as folds, assuming gene expression values 1 in no-treatment controls.

**Statistical analysis**

All experiments were repeated three times. The values were determined as mean ± standard deviation (SD). SPSS (version 15.0 for windows) was used for determining the treatment effects by one-way ANOVA. The difference or correlation was considered to be statistically significant if p <0.05 in the obtained results.

**Results**

**Cytotoxicity of hyperforin and hypericin in RAW 264.7 macrophages**

Prior to evaluation of the NO and PGE₂ inhibitory effect of hyperforin and hypericin, we first examined its cytotoxic effect in RAW 264.7 macrophages using MTT assays (Fig. 1a and 1b). While, hyperforin did not influence the survival of RAW 264.7 macrophages at concentrations of 25, 50, 75 and 100 nm, hypericin did not influence the survival of RAW 264.7 cells at concentrations of 2.5, 5, 7.5 and 10 µM. Thus, these concentrations were used in subsequent experiments.

**Effects of hyperforin and hypericin on NO and PGE₂ production in RAW 264.7 macrophages**

We investigated the potential anti-inflammatory effect of hyperforin and hypericin on NO and PGE₂ production in RAW 264.7 macrophages. As shown in Fig. 2a and 2b, NO production and in Fig. 2c and 2d, PGE₂ production was substantially higher in LPS-treated cells than in the untreated control cells (p<0.001). The addition of 25, 50, 75 and 100 nm hyperforin caused 42.68%, 47.49%, 56.37% and 58.11% reduction and the addition of 2.5, 5, 7.5 and 10 µM hypericin caused 37.69%, 41.66%, 44.69% and 53.63% reduction in LPS-induced NO production, respectively (Fig. 2a and 2b). While, 25, 50, 75 and 100 nm hyperforin caused 8.14%, 10.04%, 31.24% and 56.72%
reduction in LPS-induced PGE$_2$ production, 2.5, 5, 7.5 and 10 µM hypericin caused 5.98%, 16.45%, 38.53% and 53.37% reduction in LPS-induced PGE$_2$ production, respectively (Fig 2c and 2d).

**Effect of hyperforin and hypericin on iNOS and COX-2 mRNA expression in LPS-stimulated RAW 264.7 cells**

RT-PCR was performed to determine whether the inhibitory effects of hyperforin and hypericin on NO and PGE$_2$ production were related to the mRNA expression of iNOS and COX-2. As shown in Fig. 3a and 3b, iNOS mRNA expression and in Fig. 3c and 3d, COX-2 mRNA expression was substantially higher in LPS-treated cells than in the untreated control cells (p<0.001). The addition of 25, 50, 75 and 100 nm hyperforin caused 48%, 59%, 64% and 69% reduction and the addition of 2.5, 5, 7.5 and 10 µM hypericin caused 37%, 42%, 48% and 57% reduction in LPS-induced iNOS mRNA expression, respectively (Fig. 3a and 3b). While, 25, 50, 75 and 100 nm hyperforin caused 44%, 59%, 64% and 78% reduction in LPS-induced COX-2 mRNA expression, 2.5, 5, 7.5 and 10 µM hypericin caused 42%, 49%, 56% and 65% reduction in LPS-induced COX-2 mRNA expression, respectively (Fig. 3c and 3d).

**Discussion**

Most of the previous studies on *Hypericum perforatum* have focused on their chemical constituents, anti-depressive, anti-microbial, and anti-cancer effects. Traditional use of *Hypericum perforatum* includes wound healing and resolving infection, which could result from potential anti-inflammatory properties (13,14). The in vitro cell culture studies have been done to evaluate various *Hypericum perforatum* preparations and their contents against inflammatory mediator production (13,14). Bezakova et al. (18) reported inhibition of 12-lipoxygenase by hypericin and pseudohypericin in a *Hypericum perforatum* extract. This inhibition could limit the substrate supply for lipid inflammatory mediator synthesis. Tedeschi et al. (19) treated A549 human alveolar epithelial cells, DLD-1 human colon carcinoma cells and
Fig. 3. Effects of hyperforin and hypericin on LPS-induced iNOS and COX-2 gene expression in RAW 264.7 macrophages. iNOS gene expression of cells pretreated with different concentrations of hyperforin (a) and hypericin (b) for 1 h, then with LPS (1 μg/mL), and incubated for 48 h. COX-2 gene expression of cells pretreated with different concentrations of hyperforin (c) and hypericin (d) for 1 h, then with LPS (1 μg/mL), and incubated for 48 h. The results are expressed as mean ± SD of five independent experiments.

*p<0.05 as compared with the control cells, and †p<0.05, ‡p<0.001 as compared with the LPS-only treated cells. ANOVA and Turkey’s post-hoc test were used to determine the significances of differences.

ECV304 human cells with various doses of a commercially available Hypericum perforatum extract. Expression of iNOS gene, its catalytic product NO, as well as STAT-1 protein were found to be decreased at a dose as low as 10 μg/mL. The authors also noted that direct inhibition of JAK2 was the mechanism, instead of alteration of NF-κB. Some studies found immune-stimulatory activity of Hypericum perforatum constituents. Zhou et al. (20) treated human intestine epithelial cells and hepatocytes with hyperforin and found increased IL-8 mediated by MAPK activation. Hypericin has also been shown to promote the expression of COX-2 protein at respectively low concentrations of 125 nM and 150 nM in HeLa and T24 cells.

Animal studies were conducted to test the in vivo anti-inflammatory potential of Hypericum perforatum extract and their major active constituents. Abdel-Salam (21) evaluated the impact of orally administered Hypericum perforatum extract (50-300 mg/kg body weight) on rats with carrageen-induced edema, electrically or hot plate induced nociception, and pylorus-ligation. The results indicated strong inhibition of the pain associated with inflammation by the extract. Hu et al. (22) used irinotecan to induce intestinal inflammation and diarrhea in rats, with or without an 8 day treatment regimen of 400 mg/kg Hypericum perforatum extract. Those rats that received treatment had less apoptosis in the intestine, with lower TNF-α mRNA expression. Menegazzi et al. (23) reported that Hypericum perforatum extract treatment protected mice against zymogen-induced multi-organ failure by reducing iNOS expression and scavenging NO.

Zdunic et al. (24) studied the anti-inflammatory and gastroprotective activity of Hypericum perforatum oil extracts in rat models. The results suggested flavonoids, specifically quercetin and amentoflavone, contributed to the observed inhibition of paw edema and intestine mucosa damage. Paterniti et al. (25) comprehensively investigated the impact of a Hypericum perforatum extract during periodontitis. The extract used contained 0.34% hypericin, 4.1% hyperforin, 5% flavonoids, and 10% tannins. 2 mg/kg daily oral dose was applied to mice and lasted eight
days. By the end of the study, iNOS expression, NF-κB activation, IL-1β and ICAM-1 levels were inhibited in the treatment group, with less severe tissue damage and alveolar bone loss than was observed in the control group.

Our study has shown that all applied concentrations of hyperforin and hypericin decreased LPS-induced NO production and iNOS and COX-2 gene expression in RAW 264.7 macrophages, significantly. Similarly, all applied concentration of hyperforin and hypericin decreased LPS-induced PGE₂ production but only 75 and 100 nm hyperforin and 7.5 and 10 μM hypericin decreased PGE₂ production significantly. This indicates that hyperforin and hypericin inhibited LPS-induced NO and PGE₂ production and iNOS and COX-2 gene expression in RAW 264.7 macrophages in a dose-dependent manner, indicating that the inhibition of NO and PGE₂ production by hyperforin and hypericin is a result of the inhibition of iNOS and COX-2 gene expression.

In conclusion, the present study showed that hyperforin and hypericin isolated from Hypericum perforatum inhibits inflammation via suppressing iNOS and COX-2 gene expression and NO and PGE₂ production. For this reason, it is possible to say that hypericin and hyperforin can be used to reduce the inflammatory response, but molecular mechanisms underlying the anti-inflammatory potential of Hypericum perforatum and its active constituents were not well-characterized. Although gene transcription profile change under Hypericum perforatum treatment was described using murine cells and yeast, the regulatory pathways that lead to the transcriptome change remains yet to be revealed. Future studies on the anti-inflammatory activity of all the contents of Hypericum perforatum need to address the connection between multiple phytochemical components, bioactivity, the molecular mechanisms, and probably the overall outcome in animal models that better mimic actual inflammatory diseases.

References


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