evista de Fitoterapia

Primer Congreso Iberoamericano de Fitoterapia





LIBRO RESÚMENES

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FIGURA 1. Echinacea purpurea. Foto: Bernat Vanaclocha.

PLO3 Recent progress in the research on traditional herbal medicinal products

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Abstract

In the European Union, the use of traditional herbal medicinal products has recently been regulated in Directive 2004/24/EC. According to this regulation, clinical studies and pre-clinical tests are not obligatory, but quality needs to be demonstrated in any individual case.

Echinacea and butterbur (Petasites) will be used as examples for demonstrating the progress in medicinal plant research. Alkamides, the major lipophilic constituents of Echinacea, have recently been found to be rapidly absorbed after oral application. Using LC-MS their pharmacokinetics have been studied and ex-vivo effects have been measured. Alkamides have also been shown to bind to cannabinoid receptors (CB2) which may represent a molecular mechanism of action of Echinacea. Extracts of the rhizomes of *Petasites hybridus* have been shown to inhibit 5-lipoxygenase and cyclooxygenase-2 and COX-2 expression. They are useful for the prevention of migraine and for the treatment of asthma and seasonal allergic rhinitis.

Key words: Traditional herbal medicinal products, *Echinacea angustifolia*, *Echinacea pallida*, *Echinacea purpurea*, *Petasites hybridus*.

Resumen

En la Unión Europea, el uso de medicamentos tradicionales a base de plantas medicinales ha sido regulado recientemente en la Directiva 2004/24/EC. De acuerdo con esta directiva, los estudios clínicos y los ensayos preclínicos no son obligatorios, pero la calidad requiere ser demostrada de forma individual en cada caso. La equinácea y el petasites pueden tomarse como ejemplos para demostrar el avance en la investigación sobre plantas medicinales.

Se ha demostrado recientemente que las alquilamidas, principales constituyentes lipofílicos de la equinácea, se absorben rápidamente tras su administración oral. Utilizando LC-MS, se ha estudiado su farmacocinética y se han medido sus efectos ex vivo. También se ha demostrado que las alquilamidas tienen la capacidad de unirse a los receptores canabinoides (CB2), lo cual puede constituir un mecanismo de acción molecular de la equinácea. Los extractos de rizoma de *Petasites hybridus* inhiben la 5-lipoxigenasa y la cicloxigenasa-2, así como la expresión de esta última. Son útiles en la prevención de la migraña y en el tratamiento del asma y de la rinitis alérgica estacional.

Key words: Medicamentos tradicionales a base de plantas, *Echinacea angustifolia*, *Echinacea pallida*, *Echinacea purpurea*, *Petasites hybridus*.



In the European Union, the use of traditional herbal medicinal products has recently been regulated in Directive 2004/24/EC ⁽¹⁾. According to this regulation, clinical studies and pre-clinical tests are not obligatory for the registration of herbal products, as long as the efficacy of them is plausible on the basis of long-standing use and experience. However, quality needs to be demonstrated in any case. Nevertheless, clinical and pharmacological studies are necessary for a rational use and to convince the medical profession that these products have a reasonable activity.

Echinacea species (Asteraceae) have a long history of medicinal use. Echinacea angustifolia DC., Echinacea pallida (Nutt.) Nutt. and Echinacea purpurea (L.) Moench are frequently used as herbal immunomodulators. Traditionally, roots and aerial parts of these plants have been used to treat wounds as well as insect and snake bites. Today echinacea is mainly used for infections of the upper respiratory tract. In order to obtain consistent quality of batches, HPLC analysis of the major constituents, alkamides and caffeic acid derivatives is necessary (2). Determination of the active polysaccharides/glycoproteins would be useful as well. Specific ELISA methods have become recently available for that purpose (3).

Echinacoside and cynarine are the major polar constituents in the roots of *E. angustifolia* and frequently used for the standardization of corresponding *echinacea* preparations. Also in the current draft of the USP/NF monograph of *E. angustifolia* roots, echinacoside and cynarine are the compounds which have to be determined in an assay for total phenolics. The concentration of these compounds may change in certain preparations during storage (FIGURE 2). Cichoric acid is the major polar caffeic acid derivative in the roots of *Echinacea purpurea* and in the aerial parts of echinacea roots. Recent investigations have shown that these caffeic acid derivatives

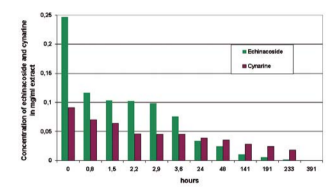


FIGURE. 2. Content of echinacoside and cynarine in 60 % EtOH *Echinacea angustifolia* root extract, during storage at 4°C over 16 days ⁽⁵⁾.

are rapidly oxidised by polyphenol oxidase present in the plant material ^(4, 5). Therefore, it is necessary to control this enzymatic activity in order to obtain products with consistent content of caffeic acid derivatives. Alkamides, the major lipophilic constituents, can be found in high concentrations in the aerial parts of *E. purpurea*, *E. pallida* and *E. angustifolia*, and in the roots of *E. purpurea* and *E. angustifolia*. They can be analysed by DAD-HPLC and, more sensitive and more specifically by LC-MS.

The pharmacokinetic properties of echinacea alkamides have recently been studied using ion trap SEI MS/MS ^(6,7). The studies have demonstrated that echinacea alkamides are detectable in human blood already 10 minutes after oral application (FIGURE 3).

There is also evidence that alkamide containing echinacea preparations trigger effects on the pro-inflammatory cytokine TNF- α and chemokine IL-8 from a recent ex vivo study. (7) Gertsch et al. (8) could demonstrate modulation of TNF- α gene expression and multiple signal transduction pathways for echinacea alkamides and postulated a mechanism related to cannabinoid receptors. In vitro kinetic experiments measuring both TNF- α mRNA and protein levels over a time-span of 39 h after a co-incubation with LPS and Echinacea purpurea tincture (EchinaforceTM) have also been performed. LPS-stimulated TNF- α protein expression was modulated by the Echinacea purpurea tincture, resulting in a significant inhibition (\sim 40 %) during the first 20 h, and subsequent stimulation of TNF- α protein expression. Because of the structural similarity of echinacea alkamides and anandamide (AEA), the endogenous ligand of CB receptors, receptor binding studies with alkamides to rodent CB1 and CB2 receptors were conducted in parallel. It could be demonstrated that echinacea alkamides have in fact high affinity to CB receptors (FIGURE 4) (9). Pentadeca-2E,9Z-diene-12,14-diynoic acid isobutylamide showed the highest affinity for CB1 with a Ki value of 2.0 μ M, while Tetradeca-2E-ene-10,12-diynoic acid isobutylamide with a Ki of 1.9 μ M was the most selective and most affine ligand for CB2. Most of the echinacea



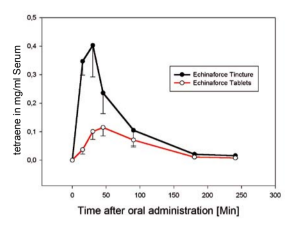
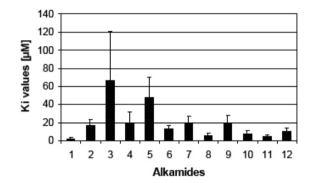


FIGURE 3. Comparison of AUC from Dodeca 2E,4E,8Z,10E/Z-tetraenoic acid isobutylamides in serum after oral administration of 4 ml Echinaforce™ tincture and 12 Echinaforce™ tablets (7).



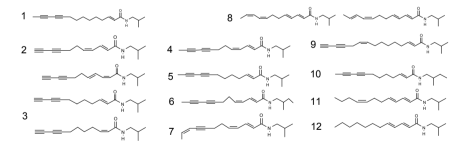


FIGURE 4. Selectivity of alkamides (1-12) from *Echinacea angustifolia* for CB2 receptor from mouse membranes, obtained by a standard receptor binding assay using a H.-CP-55,940 as the radioligand and reported as mean Ki values μ M with corresponding 95 % confidence intervals determined from at least three independent experiments ⁽⁹⁾.

alkamides showed affinities to CB2 receptors with Ki values lower than 20 μ M, some only five times less active than anandamide ⁽⁹⁾. The most recent evidence of CB2-receptor-binding has recently been demonstrated by Raduner et al. ⁽¹⁰⁾. At concentrations below 100 nM, dodeca-2E,4E,8Z,10Z- tetraenoic acid isobutylamide and dodeca-2E,4E-dienoic acid isobutylamide potently displaced the radioligand with Ki values of 57 \pm 14 nM and 60 \pm 13 nM, respectively.

Obviously alkamides from echinacea, as well as anandamide influence the cytokine milieu in human whole blood at low nM concentrations. Moreover, echinacea alkamides can exert both anti- but also pro-inflammatory effects in human blood, depending on the stimulus applied, drug concentration used, and degree of unsaturation of the lipophilic tail of the specific alkamide and can be considered as a new class of cannabinomimetics.



Although the outcome clinical studies with echinacea preparations is not consistent (11, 12), and the evidence may differ from product to product, a recent Cochrane review (13) came to the conclusion, that especially preparations based on the aerial parts of *Echinacea purpurea* have some evidence be effective for this purpose. According to a meta-analysis with three selected experimental infection studies, the likelihood of experiencing a clinical cold was 55 % higher with placebo than with echinacea (14).

Butterbur (*Petasites hybridus*) products enjoy increased interest due to a number of recently published doubleblind and placebo-controlled clinical trials in migraine prevention and treatment of seasonal rhinitis. due to the toxicity of pyrrolizidine alkaloids, only lipophilic supercritical carbon dioxide extracts of the rhizomes and the leaves are used as medicinal products today.

In vitro experiments have identified a group of sesquiterpenes, mainly petasin and iso-petasin as the pharma-cologically active components from the *Petasities hybridus* plant. Those compounds have spasmolytic as well as anti-inflammatory activities. Petasin and isopetasin relax smooth muscle and tracheal rings *in vitro* through effects involving calcium channels and calcium mobilization $^{(15, 16)}$. Both compounds have also been demonstrated to possess anti-inflammatory effects by inhibiting leukotriene synthesis and the cyclooxygenase-2 enzyme $^{(17, 18)}$. Recently, several clinical trials have been published with a special CO_2 extract made from the rhizomes of *Petasities hybridus* for migraine prevention $^{(19:21)}$. An open trial in asthma patients suggests that this extract might also be effective in improving lung function, reducing corticoid use and asthma symptoms $^{(22)}$. These trials were conducted with an extract containing a minimum of 15% petasin/isopetasin (Petadolex®, Weber & Weber, Germany). In addition, a different special CO_2 extract made from butterbur leaves has been shown to be effective in randomized controlled trials for the treatment of seasonal allergic rhinitis $^{(23, 24)}$. That extract is standardized similarly to 8 mg of total petasines per tablet (20-40 mg Ze339-extract Tesalin®, Zeller AG, Switzerland).

Due to the new popularity of butterbur, many different butterbur products have appeared on the market, especially in the USA. We have analysed the most common products in terms of petasin/isopetasin content by HPLC-DAD/MS. TABLE 1 summarizes the products analysed and the determined petasin/isopetasin content. From 6 tested products only 2 products contain what is specified on the label. Using underdosed products may lead consumers to the erroneous conclusion, that herbal drugs are ineffective. Therefore, efficient quality control and GMP is needed also for consistent quality and reliable activity of dietary supplements.

Product	Petasin+Isopetasin per unit* as specified by manufacturer (mg)	Petasin+Isopetasin per unit* as determined by HPLC-DAD/ MS (mg)
Petadolex®, Enzymatic Therapy (softgels) Lot: 40242 Lot: 42241	7.5 7.5	10.51 14.89
MigraSolve Petadolex®, Rainbow Light (softgels) Lot: K5093A Lot: I5154A	7.5 7.5	13.22 12.22
Continence, Solaray® (capsules) Lot: 091901 Lot: 091512	7.5 7.5	0.0058 0.0254
Butterbur, Solaray® (vegetarian capsules) Lot: 101402 Lot: 092609	7.5 7.5	0.0123 0.0258
Butterbur, NOW Foods® (Vcaps®) Lot: 742589 1804 Lot: 722027 0906	11.25 11.25	0.0179 0.0182
MigraDefense, KAL® (RapidSolv® tablets) Lot: 101503 Lot: 091009 Lot: 091807	7.5 7.5 7.5	0.0185 0.0285 0.0156

TABLE 1. Quantitative analysis of petasin and isopetasin content in various commercially available butterbur products (25). *Unit is one tablet, capsule or softgel.



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References

- 1. Silano M, De Vincenzi M, De Vincenzi A, Silano V. The new European legislation on traditional herbal medicines: main features and perspectives. Fitoterapia 75, 107 116 (2004).
- 2. Bauer R In. Immunomodulatory Agents from Plants. Wagner H (ed.). Basel: Birkhäuser Verlag, 1999 pp. 41-88..
- 3. Classen B, Mau SL, Bacic A. The arabinogalactan-proteins from pressed juice of Echinacea purpurea belong to the "hybrid" class of hydroxyproline rich glycoproteins, Planta Med 2005; 71: 59–66.
- 4. Nüsslein B, Kurzmann M, Bauer R, Kreis W. Enzymatic Degradation of Cichoric Acid in Echinacea purpurea Preparations. J Nat Prod 2000; 63: 1615 1618.
- 5. Wölkart K, Gangemi JD, Turner RB, Bauer R. Enzymatic Degradation of Echinacoside and Cynarine in Echinacea angustifolia Root Preparations. Pharm Biology 2004; 42(6): 443-448.
- 6. Woelkart K, Koidl C, Grisold A, Gangemi JD, Turner RB, Marth, E, Bauer R. Bioavailability and pharmacokinetikcs of alkamides from the roots of Echinacea angustifolia in humans. J Clin Pharmacol. 2005; 45: 683-689.
- 7. Woelkart K, Marth E, Raggam R, Suter A, Schoop R, Koidl C, et al.. Bioavailability and pharmacokinetic studies on Echinacea purpurea preparations and their interaction with the immune system. Int J Clin Pharm Ther 2006; 44: 401-408.
- 8. Gertsch J, Schoop R, Kuenzle U, Suter A. Echinacea alkylamides modulate TNF- gene expression via cannabinoid receptor CB2 and multiple signal transduction pathways. FEBS Lett 2004; 577: 563-569.
- 9. Wölkart K, Xu W, Pei Y, Makriyannis A, Picone RP Bauer R. The endocannabinoid system as a target for alkamides from Echinacea angustifolia roots. Planta Medica 2005; 71: 701-705.
- 10. Raduner S, Majewska A, Chen JZ, Xie XQ, Hamon J, Faller B, et al. Alkylamides from Echinacea are a new class of cannabinomimetics CB2-Receptor dependent and independent immunomodulatory effects. J Biol Chem 2006; 281:14192-14206.
- 11. Turner RB, Bauer R, Wölkart K, Hulsey TC, Gangemi JD. Evaluation of Chemically Defined Preparations of Echinacea angustifolia roots for Prevention and Treatment of Experimental Rhinovirus Infections. New Engl J Med 2005; 353: 341-348.
- 12. Goel V, Lovlin R, Chang Ch, Slama JV, Barton R, Gahler R, et al. A proprietary extract from the Echinacea plant (Echinacea purpurea) enhances systemic immune response during a common cold. Phytother Res 2005; 19 (8): 689-694.
- 13. Linde K, Barrett B, Wölkart K, Bauer R, Melchart D. Echinacea for preventing and treating the common cold. Cochrane Database Syst Rev 2006 Jan 25; (1):CD000530.
- 14. Schoop R, Klein P, Suter A, Johnston SL. Echinacea in the Prevention of Induced Rhinovirus Colds: A Meta-Analysis. Clin Ther 2006: 28: 1-10.
- 15. Käufeler R, Polasek W, Brattström A, Kötter U. Efficacy and safety of butterbur herbal extract Ze 339 in seasonal allergic rhinitis: postmarketing surveillance study. Adv Therapy 2006; 23: 373-384.
- 16. Kälin P, Sulger Büel E. Gemeine Pestwurz (Petasites hybridus). Porträt einer Arzneipflanze: Geschichte, Pharmakologie und Klinik. Schweiz Zschr Ganzheits Medizin 2002;14 (5): 267-274.
- 17. Bickel D, Röder T, Bestmann HJ, Brune K. Identification and characterization of inhibitors of peptido-leukotriene-synthesis from Petasites hybridus. Planta Med 1994; 60: 318-322.
- 18. Fiebich BL, Grozdeva M, Hess S, Hüll M, Danesch U, Bodensieck A, Bauer R. Petasites hybridus extracts in vitro inhibit COX-2 and PGE2 release by direct interaction with the enzyme and by preventing p42/44 MAP kinase activation in rat primary microglial cells. Planta Med 2005; 71:12-9.
- 19. Diener HC, Rahlfs VW, Danesch U. The first placebo-controlled trial of a special butterbur root extract for the prevention of migraine: reanalysis of efficacy criteria. Eur Neurol 2004; 51: 89-97.
- 20. Lipton RB, Göbel H, Einhäupl KM, Wilks K, Mauskop A. Petasites hybridus root (butterbur) is an effective preventive treatment for migraine. Neurology 2004; 63: 2240-4.
- 21. Pothmann R, Danesch U. Migraine prevention in children and adolescents. Results of an open study with a special butterbur root extract. Headache 2005; 45:196-202.
- 22. Danesch U. Petasites hybridus (Butterbur root) extract in the treatment of asthma an open trial. Altern Med Rev 2004; 9: 54-62.
- 23. Schapowal A. Petasites Study Group. Randomised controlled trial of butterbur and cetirizine for treating seasonal allergic rhinitis. BMJ 2002; 19; 324: 144-6.
- 24. Schapowal A; Petasites Study Group. Butterbur Ze339 for the treatment of intermittent allergic rhinitis: dose-dependent efficacy in a prospective, randomized, double-blind, placebo-controlled study. Arch Otolaryngol Head Neck Surg 2004; 130: 1381-6.
- 25. Bodensieck A, Wölkart K, Bauer R. Commercially available butterbur products often lack petasins. Herbalgram 2006 (in press).