Contents lists available at ScienceDirect



Bioorganic & Medicinal Chemistry Letters

journal homepage: www.elsevier.com/locate/bmcl



Prenyloxyphenylpropanoids as a novel class of anticonvulsive agents

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ARTICLE INFO

Article history: Received 15 June 2009 Revised 22 July 2009 Accepted 23 July 2009 Available online 26 July 2009

Keywords: Mes test Neuroprotective activity Prenyloxyphenylpropanoids

ABSTRACT

In this study, we synthesized some natural and semi-synthetic prenyloxyphenylpropanoids (e.g., acetophenones, benzoic and cinnamic acids, chalcones, and coumarins), and we assessed their in vivo neuroprotective activity, using the mouse maximal electroshock-induced seizure model (MES test). 7-Iso-pentenyloxycoumarin and (2*E*)-3-{4-[(3-methylbut-2-enyl)oxy]phenyl}prop-2-enoic acid, administered ip at a dose of 300 mg/kg, suppressed MES-induced seizures in mice in a time- and dose-dependent manner.

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Neurodegenerative syndromes represent a widely differentiated group of diseases affecting the nervous system, including the brain, the spinal cord, and peripheral nerves, arising from several causes, many of which are still unknown.¹ Due to their incidence, morbidity, and mortality, neurodegenerative diseases represent nowadays a huge problem from a medical, social, and financial point of view for the human society all over the world. Pathologically, neurodegeneration result from abnormalities in the functionality of specific regions of the central and the peripheral nervous systems and/or specific populations of neurons. Depending on which cluster of cells begins to undergo pathological changes, this will lead to the clinical phenotype of that particular illness. Recent studies allowed to identify a certain number of genes responsible for several neurodegenerative diseases. Moreover, during last years, valuable animal models were developed to study factors and mechanism underlying the etiology and pathogenesis of these syndromes. Currently used therapeutic approaches are able to treat the symptoms for a limited period of time but, as progression of the disease advances, these treatments become ineffective. Different therapeutic approaches were proposed, depending on which biological target is triggered. Drugs commonly used in such situations are able to restore the imbalance in neurotransmitters functionality that characterize the most part of neurodegenerative diseases. Explicative examples to this aim are (a) the use of L-DOPA in

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combination with DOPA-decarboxylase inhibitors in the treatment of Parkinson's disease and (b) the use of cholinomimetics and acetylcholinesterase inhibitors in the therapy of Alzheimer's disease. Other drugs that were proposed in the therapeutic management of neurodegenerative disorders include inhibitors of key enzymes like COX-2,² MAO-B,³ *i*NOS,⁴ and caspase,⁵ heavy metal chelators,⁶ radical and reactive oxygen species scavengers,^{7,8} statins,⁹ and finally the most recent approaches are genetic¹⁰ and stem-cell¹¹ based therapies.

In recent years natural products have been re-discovered as a valuable source of novel agents exerting positive effects in the treatment of cancer and microbial infections, against which synthetic drugs failed due to progressively increasing pharmacological resistance. In addition several natural compounds were seen in the last decade to efficiently interfere at different levels with the development and progress of many neurodegenerative disorders. These include ferulic acid,¹² oleamide,¹³ polyphenols,¹⁴ tocopherols,¹⁵ curcumin,¹⁶ folate,¹⁷ and coumarins.¹⁸

In the last five years our research group studied extensively chemical and biological properties of secondary metabolites of phenylpropanoid biosynthetic origin, and containing a sesquiterpenyl, monoterpenyl, and semiterpenyl chains attached to a phenol group. These represent quite a rare group of natural products.¹⁹ As a continuation of our studies aimed at evaluating pharmacological properties of prenyloxyphenylpropanoids, we wish to report herein the activity of natural compounds belonging to selected subcategories (e.g., acetophenones, benzoic and cinnamic acids, chalcones, and coumarins) as in vivo neuroprotective

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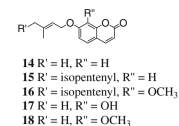
anticonvulsant agents using the mouse maximal electroshock-induced seizure model (MES test). The following secondary metabolites were synthesized and evaluated: the acetophenones 1-{2hydroxy-4-[(3-methylbut-2-enyl)oxy]phenyl}ethanone (1) and 1-(4-{[(2*E*)-3,7-dimethylocta-2,6-dienyl]oxy}-2-hydroxyphenyl)ethanone (2), both isolated from *Melicope* spp.²⁰ and *Euodia merrilli* Kanehira & Sasaki ex Kanehira²¹ (Fam. Rutaceae); the vanillic acid derivatives 3-methoxy-4-[(3-methylbut-2-enyl)oxy]benzoic acid (3) and 4-{[(2E)-3,7-dimethylocta-2,6-dienyl]oxy}-3-methoxybenzoic acid (4), both obtained as methyl esters from the liverwort Trichocolea lanata (Ehrh.) Dumm. (Fam. Trichocolaceae);²² the pcoumaric and ferulic acids derivatives (2E)-3-{4-[(3-methylbut-2enyl)oxy]phenyl]prop-2-enoic acid (5), isolated from Esenbeckia hieronymi (Fam. Rutaceae),23 boropinic acid (6), extracted from Boronia pinnata Sm. (Fam. Rutaceae),²⁴ (2E)-3-(4-{[(2E)-3,7-dimethylocta-2,6-dienyl]oxy}phenyl)prop-2-enoic acid (7), and (2E)-3-(4-{[(2*E*)-3.7-dimethylocta-2.6-dienylloxy}-3-methoxyphenyl)prop-2-enoic acid (8), both obtained from Acronychia baueri Schott. (Fam. Rutaceae),²⁵ 3-{4-[(3-methylbut-2-enyl)oxy]phenyl}propanoic acid (9) and 3-(4-{[(2E)-3,7-dimethylocta-2,6-dienyl]oxy}phenyl)propanoic acid (10), both isolated as methyl esters from Zanthoxylum pistaciiflorum Hayata (Fam. Rutaceae);²⁶ the chalcones cordoin (**11**), extracted from *Millettia erythrocalyx* Gagnep.,²⁷ Cordoa placa,²⁸ and Derris spp. (Fam. Leguminosae),²⁹ p-hydroxycordoin (**12**), isolated from *C. placa*,³⁰ and 4'-geranyloxyisoliquiritigenin (**13**), obtained from *Millettia* spp.;³¹ finally the coumarins 7-isopentenyloxycoumarin (14), auraptene (15), widespread in the genus Citrus (Fam. Rutaceae), collinin (16), isolated from Zanthoxylum schinifolium Siebold & Zucc. (Fam. Rutaceae), 7-hydroxy-8isopentenyloxycoumarin (17), extracted from Melampodium divaricatum DC. (Fam Asteraceae), and lacinartin (18), obtained from Z. schinifolium, Artemisia and Diosma spp. (Fam. Rutaceae).³²

$$R = H$$

$$R = isopentenyl$$

3 R' = COOH, R" = OCH₃, R''' = H 4 R' = COOH, R" = OCH₃, R''' = isopentenyl 5 R' = CH=CH-COOH, R" = H, R''' = H 6 R' = CH=CH-COOH, R" = OCH₃, R''' = H 7 R' = CH=CH-COOH, R" = H, R''' = isopentenyl 8 R' = CH=CH-COOH, R" = OCH₃, R''' = isopentenyl 9 R' = CH₂CH₂-COOH, R" = H, R''' = H 10 R' = CH₂CH₂-COOH, R" = H, R''' = isopentenyl \bigcirc

Compounds **3–8** and **14–16** were synthesized as already reported.³³ Acids (**9**) and (**10**) were obtained by the same environmentally friend procedure reported for compounds **3–8** in 77% and 83% yield, respectively, and using commercially available 3-(4-hydroxyphenyl)propionic acid as the starting material. Acetophenones (**1**) and (**2**) were both synthesized in a single step procedure,



in 76% and 68% yields, respectively, from 2,4-dihydroxyacetophenone by selective alkylation with 3,3-dimethylallyl bromide and geranyl bromide promoted by 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), followed by crystallization from *n*-hexane (Fig. 1).³⁴

Cordoin (11) was obtained in 93% yield from acetophenone (1) and benzaldehyde by an aldol condensation in a basic hydroalcoholic medium, followed by acidic work-up, and crystallization from n-hexane (Fig. 2).

The two other chalcones, *p*-hydroxycordoin (**12**), and 4'-geranyloxyisoliquiritigenin (**13**) were obtained from acetophenones (**1**) and (**2**), and *p*-hydroxybenzaldehyde in 32% and 35% yield, respectively, by a slight modification of the synthetic scheme of cordoin (**11**) (Fig. 3).³⁵

Finally, coumarins (**17**) and (**18**) were obtained by the same procedure reported for compounds **14–16** in 84% and 62% yields, respectively.³⁴

Compounds **1–18** were then evaluated in vivo for their neuroprotective anticonvulsant properties, using the mouse maximal electroshock-induced seizure model (MES test). Procedures involving animals and their care were carried out in accordance with current European Community and Polish legislation on animal experimentation. Additionally, all efforts were made to minimize animal suffering and to use only the number of animals necessary to record reliable scientific data. The experimental protocols and procedures described in this manuscript were approved by the Local Ethics Committee at the Medical University of Lublin (Licenses no.: 20/2008 and 26/2008), and complied with the European Communities Council Directive of 24 November 1986 (86/609/EEC). All compounds were administered intraperitoneally (ip) at a dose of 300 mg/kg. Results from the mouse MES model are reported in Table 1.

The anticonvulsant effects were considered of pivotal importance if at least 50% of the animals tested were protected against MES-induced seizures. As shown in Table 1, a well defined and distinguished pattern of results was recorded. The results partly confirm data already obtained in an in vitro study in which 7-isopentenyloxycoumarin was found to protect neuronal cells against NMDA-induced excitotoxicity.³⁵ Results obtained herein indicated that the compounds (**5**), (**8**), (**14**), and (**17**) considerably protected the animals against MES-induced seizures, possessing the definite anticonvulsant properties, at the respective pretreatment times in the mouse MES model. In particular for 7-isopentenyloxycoumarin (**14**) and (2*E*)-3-{4-[(3-methylbut-2-enyl)oxy]phenyl}prop-2-enoic acid (**5**) the maximum of anticonvulsant activity was reached within a very short time after ip administration, leading to a dose and time-dependent protection of all animals treated. The dose-

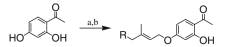


Figure 1. Reagents and conditions: (a) 3,3-dimethylallyl bromide [geranyl bromide] (1 equiv), DBU (1 equiv) acetone, rt, 24 h; (b) crystallization.

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