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Synthesis of 2-Prenylated Alkoxylated Benzopyrans by Horner-Wadsworth-Emmons olefination with PPARα/γ Agonist Activity

Ainhoa García,† Laura Vila,†,‡ Paloma Marín,† Álvaro Bernabeu,† Carlos Villarroel-Vicente,†,‡ Nathalie Hennuyer, Bart Staels, Xavier Franck,§ Bruno Figadère, Nuria Cabedo,†,‡,* and Diego Cortes†,*

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ABSTRACT: We have synthesized series of 2-prenylated benzopyrans as analogs of the natural polycerasoidol, a dual PPAR α/γ agonist with anti-inflammatory effect. The prenylated side chain consists of five- or nine-carbons with an α -alkoxy- α , β -unsaturated ester moiety. Prenylation was introduced via Grignard reaction, followed by Johnson-Claisen rearrangement, and the α -alkoxy- α , β -unsaturated ester moiety by the Horner-Wadsworth-Emmons reaction. Synthetic derivatives showed high efficacy to activate both hPPAR α and hPPAR γ as dual PPAR α/γ agonists. These prenylated benzopyrans emerge as lead compounds potentially useful for preventing cardiometabolic diseases.

KEYWORDS: Prenylated benzopyrans, Horner-Wadsworth-Emmons reaction, PPARa/y activity

Among nuclear receptor family, peroxisome proliferator activated receptors (PPARs) are transcription factors activated by ligands, which are implicated in numerous cell functions including glucose and lipid metabolism, and inflammation.1 Dual PPARα/γ activators can improve atherogenic dyslipidemia and insulin resistance. Polycerasoidol is a transδ-tocotrienolic acid analog² isolated from *Polyalthia cerasoides* (Annonaceae).^{3,4} Polycerasoidol, containing a 6-chromanol nucleus and a 2-prenylated side chain, is biosynthesized from homogentisate and geranylgeranyl pyrophosphate via the shikimate and the mevalonate pathway, respectively. Pharmacologically, it displays dual PPARα/γ agonist activity and ameliorates inflammation of dysfunctional endothelium.⁵ In structural terms, this natural benzopyran possesses a benzopyran nucleus (lipophilic tail), a prenylated chain (flexible linker), and a carboxylic acid (polar head), as other natural and synthetic PPARα and/or PPARγ agonists.⁶ The structure-activity relationships (SAR) of polycerasoidol and semisynthetic analogs revealed that the 6-oxygenated dihydrobenzopyran core and the linker hydrocarbon chain of at least a five-carbon length are essential moieties to activate PPARα and/or PPARγ. Tit is noteworthy that chroman-6-ol pharmacophore is present in troglitazone, the first PPARy agonist belonging to the class of thiazolidinediones, which was approved as anti-diabetic agent.^{8,9} Currently, rosiglitazone and pioglitazone are selective PPARy agonists used to manage hyperglycemia in type 2 diabetes, while saroglitazar and lobeglitazone are PPARa/y activators (agonists) approved against diabetes in India and South Korea, respectively.9 In order to find new active and safer PPAR activators, we describe the synthesis and PPAR activity of 2-prenylated benzopyrans that bear an α -alkyloxy- α , β -unsaturated ester as a trioxygenated polar head on the prenylated side chain.

Figure 1. Bioactive prenylated benzopyrans.

We first prepared the benzopyran-4-one nucleus (γbenzopyrone 3) by conventional methods. 10 The reaction mechanism consists of an aldol condensation between the enolate from an ortho-hydroxyacetophenone and the carbonyl group from an alkyl ketone. This is followed by an intramolecular cyclisation via Michael addition, which is promoted by the ortho-phenol group.11 Thus, 2,5dihydroxyacetophenone (1) and ethyl levulinate (2) in the presence of pyrrolidine gave the chroman-4-one 3 as a racemic mixture in a single step with a good yield (80 %) (Figure 2).^{7,10} y-Benzopyrone 3 was reduced under Clemmensen conditions and the phenol hydroxyl group at 6-position was protected by O-benzylation to give compound 4 with a good overall yield. The controlled reduction of the ester function by DIBAL-H at -78°C gave the aldehyde 5 with a 92 % yield. In a first approach, we synthesized five-carbon side chain alkoxylated prenylated benzopyrans (series 1) from aldehyde intermediate 5 by Horner-Wadsworth-Emmons olefination. For this purpose, appropriate

[†] Departamento de Farmacología, Facultad de Farmacia, Universidad de Valencia, 46100 Burjassot, Valencia, Spain

[‡] Institute of Health Research-INCLIVA, University Clinic Hospital of Valencia, 46010 Valencia, Spain

¹ Université Lille, Inserm, CHU Lille, Institut Pasteur Lille, U-100-EGID 59019 Lille, France

[§] Normandie Univ, CNRS, INSA Rouen, UNIROUEN, COBRA (UMR6014 & FR 3038), 76000 Rouen, France

[∆] BioCIS, CNRS, Université Paris-Saclay, Châtenay-Malabry, France

alkylphosphonates, e.g., ethyl 2-(diethoxyphosphoryl)-2-ethoxyacetate (**8a**) and ethyl 2-(diethoxyphosphoryl)-2-(2,2,2-trifluoroethoxy)acetate (**8b**), were previously prepared from (ethoxyacetate) diethoxyphosphorane and TsN_3 (**6**) in the presence of NaH. ¹² Then, aldehyde intermediate **5** was reacted with phosphonate **8a** or **8b** to afford the α -alkyloxy- α , β -unsaturated esters of prenylated benzopyrans **9** (20 %, Z/E= 60:40) or **10** (84 %, Z/E= 35:65), respectively. The saponification of ethyl ester (**10**) quantitatively afforded carboxylic acid (**11**, Z/E= 40:60) (Scheme 1).

Scheme 1. Synthesis of prenylated benzopyrans **9-11** (series 1).^a

^a Reaction conditions: (a) Pyrrolidine, EtOH, 60 °C, molecular sieve 3Å, 24h, 80 %; (b) Zn/HCl, AcOH-H₂O, rt, 2h; (c) ClCH₂C₆H₅, K₂CO₃, EtOH, 60 °C, 5h, 90 %; (d) DIBAL-H, CH₂Cl₂, -78 °C, N₂, 20 min, 92 %; (e) 0 °C, acetone, 2h, 81 %; (f) 60 % NaH, THF, 0 °C, N₂, 16h, 65 %; (g) EtOH, Rh(OAc)₂, toluene, 45 °C, overnight, 8a (R= CH₂CH₃), 40.2 % or 8b (R= CH₂CF₃), 51 %; (h) Phosphonate 8a or 8b, THF, NaH, 0 °C, N₂, 1h + 5, THF, rt, overnight: 9, 20 % or 10, 84 %.; (i) 20 % KOH, reflux, 5h, 99 %.

In a second approach, we synthesized the nine-carbon side chain O-alkoxylated prenylated benzopyrans (series 2) 15, 16 and 17 from aldehyde intermediate 5. The prenylated side chain at the 2-position of the dihydrobenzopyran nucleus was elongated with a sequence of Grignard reaction, Johnson-Claisen rearrangement and Horner-Wadsworth-Emmons olefination (Scheme 2). The aldehyde synthon 5 was treated with isoprenylmagnesium bromide as the vinyl Grignard reagent, followed by Johnson-Claisen rearrangement of allylic alcohol 12 using ethyl orthoacetate to produce unsaturated ester 13 with a 50 % yield in the last two steps. Ester 13 was subjected to a controlled reduction using DIBAL-H at -78°C to give the aldehyde intermediate 14 in 89 % yield. The α-alkoxy-α,βunsaturated ester on the prenylated side chain was introduced by a Horner-Wadsworth-Emmons reaction. 13 Thus, aldehyde 14 was treated with phosphonates 8a and 8b to afford esters 15 (85

%) and 16 (82 %), respectively. ¹⁴ It is noteworthy that ester 15 was obtained as *Z*-alkene isomer exclusively. Once again, the saponification of ethyl ester 15 quantitatively yielded carboxylic acid 17 (Scheme 2). In addition to *O*-benzyloxy benzopyrans bearing a nine-carbon prenylated alkoxy side chain (series 2), we accomplished the synthesis of its *O*-propyloxy and *O*-*p*-fluorobenzyloxy benzopyran analogs. According to the second approach followed to prepare ester 15, but starting from the chroman-4-one 3, we synthesized *O*-propyloxy ester 18 and *O*-*p*-fluorobenzyloxy ester 19 (Scheme 3).

Scheme 2. Synthesis of prenylated benzopyrans **15-17** (series 2).^a

^a Reaction conditions: (a) CH₃C(MgBr)=CH₂, THF, -78 °C, N₂, 1h, 84 %; (b) MeC(OEt)₃, isobutyric acid, 140 °C, 2h, 50 %; (c) DIBAL-H, CH₂Cl₂, -78 °C, N₂, 20 min, 89 %; (d) Phosphonate **8a** or **8b**, THF, NaH, 0 °C, N₂, 1h + **14** in THF, rt, overnight: **15** (R= CH₂CH₃), 85 % or **16** (R= CH₂CF₃), 82 %; (e) 20 % KOH, reflux, 5h, 99 %.

Scheme 3. Synthesis of prenylated benzopyrans 18 and 19.^a

^a Reaction conditions: (a) See reagents and conditions described in Scheme 1 for synthesis of **5**, and Scheme 2 for synthesis of **15** and **16**.

The transactivation studies ¹⁵ of the synthesized benzopyrans were carried out and compared to the maximal efficacy of WY-14,643 (at 10 μ M) or rosiglitazone (at 1 μ M) as hPPAR α and hPPAR γ reference compounds, respectively. At

the 10 μ M dose, compounds 10, 15, 16 and 18 were moderate hPPAR activators for both receptors, while compounds 11, 17 and 19 showed high efficacy as dual hPPAR α/γ agonists. Indeed, compound 11 showed higher efficacy to activate hPPAR α than PPAR γ (α/γ ratio= 1.73), and 17 displayed slight selectivity towards hPPAR γ (α/γ ratio= 0.64). Therefore, the elongation of the side chain from five- to nine-carbons is beneficial to activate hPPAR γ . In agreement with previous docking analysis of polycerasoidol, the carboxylic moiety at the C-9′ position of 17 plays a key role as an anchoring point to bind PPAR γ receptor.⁵

Compound 9

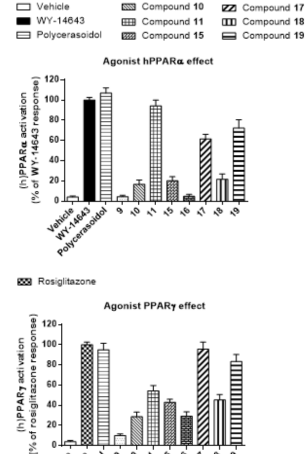


Figure 2. hPPAR α and hPPAR γ transactivation assays. Synthesized benzopyrans were tested at 10 μ M, and WY-14,643 (10 μ M) and rosiglitazone (1 μ M), as reference compounds for α and γ , respectively.

In conclusion, we efficiently prepared new series of the 2-prenylated O-alkoxylated benzopyrans possessing the α -alkoxy- α , β -unsaturated moiety on the prenylated chain by the Horner-Wadsworth-Emmons reaction. Synthetic derivatives were efficient in activating both hPPAR α and hPPAR γ as dual PPAR α/γ agonists. These prenylated benzopyrans emerge as lead compounds that might be potentially useful for preventing cardiometabolic diseases.

AUTHOR INFORMATION

Corresponding Authors

Diego Cortes – Email: dcortes@uv.es Nuria Cabedo – Email: ncabedo@uv.es

Notes

EEE Compound 16

The authors declare no competing financial interest.

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ABBREVIATIONS

PPARs, Peroxisome proliferator-activated receptors; SAR, structure-activity relationships; DIBAL-H, Diisobutylaluminium hydride; TsN₃, Tosyl azide.

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- tion (hexane/ EtOAc 80:20) yielded 64.5 mg of 15 (85 %) as a colorless oil: 1H NMR (300 MHz, CDCl₃): δ 7.36 (m, 5H, H-2" to 6"), $6.75 \text{ (dd, } J = 7.1, 2.3 \text{ Hz, } 1H, H-7), } 6.72 \text{ (d, } J = 7.1 \text{ Hz, } 1H, H-8), }$ 6.69 (d, J = 2.3 Hz, 1H, H-5), 6.23 (t, J = 7.4 Hz, 1H, H-7'), 5.16(m, 1H, H-3'), 4.99 (s, 2H, OCH₂Ar), 4.21 (q, J = 7 Hz, 2H, $COOCH_2CH_3$), 3.85 (q, J = 7 Hz, 2H, OCH_2CH_3), 2.73 (t, J = 6.8Hz, 2H, CH₂-4), 2.34 (q, J = 7.4 Hz, 2H, CH₂-6'), 2.09 (m, 4H, CH₂-2', CH₂-5'), 1.81 (m, 2H, CH₂-3), 1.62 (m, 2H, CH₂-1'), 1.60 (s, 3H, CH₃-10'), 1.30, 132 (2t, J = 7 Hz, 3H, OCH₂CH₃, COOCH₂CH₃), 1.27 (s, 3H, CH3-11'). ¹³C NMR (75 MHz, CDCl₃): δ 164.6 (CO), 152.5 (C-6), 148.5 (C-8a), 145.4 (C-8'), 137.9 (C-1"), 134.5 (C-4'), 129.1 (CH-7'), 128.9, 128.2, 127.9 (CH-2" to CH-6"), 125.4 (CH-3'), 122.0 (C-4a), 116.4 (CH-7), 115.5 (CH-5), 114.7 (CH-8), 76.0 (C-2), 71.0 (OCH₂Ar), 68.4 (OCH₂CH₃), 61.1, (COOCH₂CH₃), 40.3 (CH₂-5'), 39.7 (CH₂-1'), 31.1 (CH₂-3), 25.4. (CH₂-6'), 24.5 (CH₃-11'), 22.8 (CH₂-4), 22.6 (CH₂-2'), 16.2 (CH₃-10'), 15.8 (COOCH2CH3), 14.9 (OCH2CH3). HREIMS m/z 492.2869 [M]+ (492.2875 calcd. for $C_{31}H_{40}O_5$) (100%).
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