Synthesis, Characterization and Cytotoxicity Evaluation of New Compounds from Oxazol-5(4H)-ones and Oxazoles Class Containing 4-(4-Bromophenylsulfonfonyl)phenyl Moiety

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The interest in the chemistry of the saturated azlactones - which are internal anhydrides of α-amino acids - is due to their usefulness as intermediate in the synthesis of different heterocyclic compounds or modified α-amino acids or their derivatives [12]. Also, 1,3-oxazol-5(4H)-ones have been reported to present antimicrobial [13], antitumoral [14], antiviral activities [15] etc.

Further, diaryl sulfone derivatives (e.g. Dapsone, Amidapsone, Acedapsone, Promanide, Solasulfone, Sulfoxone, Diuciphone) were also found to possess antibacterial, antiviral, antituberculosis, and antioxidant action [16]. In the view of these reports, the diaryl sulfone moiety was incorporated into various heterocyclic systems with potential biological activity [17-19].

Based on all above considerations and also in continuation of our researches [20,21], in this work is reported the synthesis and characterization of new heterocyclic compounds from oxazol-5(4H)-ones and oxazoles class wherein the 2-aryl group is 4-(4-bromophenylsulfonfonyl)phenyl and of their acyclic intermediates, with the aim to obtain potent biologically active compounds. The synthesized compounds were tested for cytotoxic activity using Daphnia magna bioassay. The method is simple, rapid, and can predict the biological effect [22-26].

Keywords: N-acyl-α--amino acid, 1,3-oxazol-5(4H)-one, α-acylamino ketone, 1,3-oxazole, cytotoxicity

Heterocyclic compounds containing 1,3-oxazol-5(4H)-one and 1,3-oxazole ring are important targets in synthetic and medicinal chemistry, because of their applications as active substances.

1,3-Oxazoles are substructures of various biologically active natural products, pharmaceuticals, and synthetic intermediates. Thus, the 1,3-oxazole nucleus is an important pharmacophore in modern drugs, due to having a wide spectrum of biological activities [1], such as anti-inflammatory (e.g. Oxaprozin, Ramazarit, Ditasol, Isaxamole, Tioxaprofene, Tilmacoxib) [2], analgesic (e.g. Oxaprozin) [3], antibacterial, antifungal (e.g. Sulfamoxole, Sulfaguanole) [4], anti-diabetic (e.g. Aleglitazar, Farglitazar, Darglitazone, Muraglitazar, Imiglitazar) [5], antitumoral (Mubritinib) [6], antioxidant [9], and HIV-inhibitor effect [10]. Moreover, various natural products of peptide origin containing oxazole ring are active substances which exhibit several pharmacological properties [11], including antitumoral (e.g. Telomestatin, Thiangazole, Diazonamide A, Micalolide A, Leucascandrolide A), analgesic (e.g. Hennoxazole A), antifungal (e.g. Rhizoxin D, Phorboxazole A and B, Leucascandrolide A, Bengazole A, Mycalolide A), antibacterial (e.g. Pristinamycin IIb or Virginiamycin M), Bistratamide, Sulfonycin I, Griseoviridin, Madumycin II, Oxafoxazole, Microcin B17, Promothiocin A, Flopristin), antiviral (e.g. Hennoxazole A, Thiangazole, Phenoxan), antimycobacterial (e.g. Texaline), antileukemia (e.g. Ulapualide A), and anticonvulsant (e.g. Pimprinpine).

The method is simple, rapid, and can predict the biological effect [22-26].

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Experimental part

Chemistry

Melting points were determined on a Böetius apparatus and are uncorrected. FT-IR spectra were recorded in KBr pellets on a Bruker Vertex 70 spectrometer; intensity of IR bands are given as: weak (w), medium (m), strong (s), and very strong (vs). UV-Vis spectra were registered in methanolic solution (2.5.10^{-5} M) on an Analytik Jena AG Specord 40 spectrophotometer. NMR spectra were recorded on a Varian Gemini 3000BB spectrometer at 300 MHz for ^1H-NMR and 75 MHz for ^13C-NMR using DMSO-d_6 or CDCl_3 as solvents; chemical shifts (δ) are reported in ppm relative to tetramethylsilane (TMS) as internal standard and coupling constants (J) are expressed in Hz. For multiplicity of signals in ^1H-NMR spectra, following abbreviations were used: singlet (s), broad singlet (br s), doublet (d), broad doublet (br d), doublet of doublets (dd), triplet (t), broad triplet (br t), triplet of triplets (tt), quartet (q), quintet (quint), and multiplet (m). Mass spectra (ESI-MS/MS) were recorded on a Varian 1200 LC-MS/MS high performance liquid chromatograph coupled with a triple quadrupole mass spectrometer with electrospray interface (ESI), by positive and/or negative ionization. GC-EI-MS analysis was carried out using a Fisons Instruments GC 8000 with electron impact quadrupole and MD 800 mass spectrometer detector. Compounds purity was checked by RP-HPLC using a Beckman System Gold 126 liquid chromatograph, equipped with a System Gold 166 UV-Vis detector; retention time (t_R) of compounds in min is reported. Contents of C, H, N, and S were determined using a Costech ECS 4010 micro elemental analyzer.

Synthesis and characterization of compounds

The synthetic method used in this approach consisted in N-acylation of α-alanine with benzoyl chloride 2 by Steiger procedure to the N-acyl-α-alanine 3, followed by cyclization of this compound to the corresponding saturated azlactone 4. Friedel-Crafts acylation of aromatic hydrocarbons (benzene, toluene, m-xylene, mesitylene) with 1,3-oxazol-5(4H)-one (2-oxazolin-5-one) 4 or N-acyl-α-alanyl chloride 5, in the presence of anhydrous aluminum chloride, yielded the corresponding α-acylamino ketones 6a-d. These intermediates were converted into 1,3-oxazoles 7a-d by Robinson-Gabriel cyclocondensation with phosphorus oxychloride or concentrated sulfuric acid in the presence of acetic anhydride (scheme 1). The structures of new synthesized compounds 3-7 were established unequivocally by FT-IR, UV-Vis, MS, ^1H-NMR, ^13C-NMR spectra and elemental analysis.

Scheme 1. Synthesis of the compounds

Synthesis of 2-[4-(4-bromophenylsulfonyl)benzamido]propanoic acid 3
α-Alanine (1.78 g, 20 mmol) was dissolved in 20 mL of 1N NaOH solution. This solution was cooled to 0-5°C and then two solutions were added simultaneously dropwise under continuous stirring for 30 min, as follows: a solution of crude acyl chloride 2 (7.19 g, 20 mmol) in 45 mL anhydrous CH₂Cl₂ and 10 mL of 2N NaOH solution, respectively. After 1 h stirring at room temperature, the aqueous layer was separated and then acidified with 2N HCl. The precipitate was filtered off and recrystallized from water as white needle-shaped crystals; yield 96%; m.p. 197-198°C; UV-Vis (CH₂OH, λ nm) (lg ε): 202.6 (4.46), 226.4 (4.08), 252.0 (4.36); FT-IR (KBr, ν cm⁻¹): 3377s, 3088m, 3066m, 2989m, 2952m, 2872w, 1708vs, 1644vs, 1577s, 1478s, 1462m, 1352vs, 1233v, 1158vs, 835s, 571vs; 1H-NMR (DMSO-d₆, δ ppm, J Hz): 3.03 (d, 7.4, 3H, H-18), 7.84 (d, 8.8, 2H, H-13, H-17), 8.05 (d, 8.5, 2H, H-7, H-11), 8.14 (d, 8.5, 2H, H-8, H-10); 13C-NMR (DMSO-d₆, δ ppm): 13.11 (C-18), 136.61 (C-2), 127.79 (C-5), 132.79 (C-9), 142.07 (C-2), 120.11 (C-6), 131.54 (C-15), 133.00 (C-14, C-16), 139.92 (C-12), 145.03 (C-9), 160.28 (C-2), 178.09 (C-5); -MS/MS (m/z, rel. abund. %): 412 (79Br)/414 (81Br) [M+H]+; 394/396 (19.8/17.6) [M+H-H₂O]+; 366/368 (66.2/70.6) [M+H-H₂O-CO]+; 323/325 (100, BP) [BrC₆H₄SO₂]+; 203/205 (40.5/50.6) [BrC₆H₄SO₂C₆H₄]+; 155/157 (25.2/28.4) [BrC₆H₄SO₂]+; 104 (25.2) [C₄H₂]+×; 76 (31.6) [C₆H₄CHNH]+×; 50 (8.7) [C₄H₂]+×; 39 (13.3) [C₆H₄]+×; tR 30.62 min; -EI-MS (m/z, rel. abund. %): 394 (39%) [Br]/396 (41%) [Br] [M⁻]; 368/366 (66/62.7) [M⁻H₂O]+×; 349/351 (27/28.4) [M⁻CO]+×; 323/325 (88/100, BP) [M⁻CO-C₆H₄]+×; 203/205 (40.5/50.6) [BrC₆H₄SO₂]+×; 155/157 (25.2/28.4) [BrC₆H₄SO₂]+×; 104 (25.2) [C₆H₄CHNH]+×; 50 (8.7) [C₄H₂]+××; 39 (13.3) [C₆H₄]+××; tR 30.62 min; -RP-HPLC (CH₂OH:H₂O = 30:70, 1 mL/min, 250 nm): purity 90.78%, tR 4.63 min; Anal. (%): Calcd. for C₁₆H₁₄BrNO₅S (394.24 g/mol): C, 46.61; H, 3.42; N, 3.40; S, 7.78. Found: C, 46.68; H, 3.40; N, 3.45; S, 7.76. Synthesis of 2-[4-(4-bromophenylsulfonyl) benzamido] propanoyl chloride 5

Method 1. Anhydrous AlCl₃ (2.0 g, 15 mmol) was added portionwise under stirring at room temperature to the crude azlactone 4 (1.97 g, 5 mmol) in excess of dry aromatic hydrocarbon (25 mL). The reaction mixture was stirred for 20 h and then poured over 100 mL ice-water with 5 mL concentrated HCl. The precipitate of crude product was filtered off and washed on the filter with a small amount of cool ethanol. The product was obtained as white crystals; 98% yield; m.p. 162-164°C (cyclohexane); UV-Vis (CH₂OH, λ nm) (lg ε): 202.6 (4.47), 228.2 (4.08), 252.9 (4.36); FT-IR (KBr, ν cm⁻¹): 3092m, 3069m, 2989m, 2943m, 2874w, 1820s, 1650vs, 1597m, 1573s, 1472m, 1331vs, 1303s, 1295s, 1256s, 1163vs, 1046s, 846s, 574s; -MS/MS (m/z, rel. abund. %): 156 (88.1/100, BP) [M⁻CO]+×; 145/147 (25.2/28.4) [M⁻CO-C₆H₄]+×; 104 (25.2) [C₆H₄CHNH]+×; 50 (8.7) [C₄H₂]+××; tR 30.62 min; -EI-MS (m/z, rel. abund. %): 394 (39%) [Br]/396 (41%) [Br] [M⁻]; 368/366 (66/62.7) [M⁻H₂O]+×; 349/351 (27/28.4) [M⁻CO]+×; 323/325 (88/100, BP) [M⁻CO-C₆H₄]+×; 203/205 (40.5/50.6) [BrC₆H₄SO₂]+×; 155/157 (25.2/28.4) [BrC₆H₄SO₂]+×; 104 (25.2) [C₆H₄CHNH]+×; 50 (8.7) [C₄H₂]+××; tR 30.62 min; -RP-HPLC (CH₂OH:H₂O = 30:70, 1 mL/min, 250 nm): purity 90.78%, tR 4.63 min; Anal. (%): Calcd. for C₁₆H₁₂BrNO₄S (394.24 g/mol): C, 48.74; H, 3.07; N, 3.55; S, 8.13. Found: C, 48.85; H, 3.01; N, 3.49; S, 8.19. Synthesis of 2-[4-(4-bromophenylsulfonyl) benzamido] propanoyl chloride 5

Method 2. 2-[4-(4-Bromophenylsulfonyl) benzamido] propanoic acid 3 (2.27 g, 5.5 mmol) was refluxed with 25-fold molar excess of thionyl chloride (10 mL) on a water bath until emission of sulfur dioxide and hydrogen chloride gas ceased. Unreacted SOCl₂ was removed to dryness by distillation under reduced pressure on a water bath. The yellow crystalline crude product was used without further purification; 98% yield; m.p. 156-158°C; FT-IR (KBr, ν cm⁻¹): 3345m, 3090m, 3067m, 2989m, 2951m, 2843w, 1826s, 1788s, 1651vs, 1599m, 1573vs, 1472m, 1524s, 1326vs, 1292v, 1160vs, 851m, 886m, 575vs.
longer produced (≈20 h) and then the reaction mass was poured over 100 mL mixture of acidulated (HCl) ice-water. After extraction in CH₂Cl₂, the organic layer was washed with 5% NaHCO₃ solution, then with water and dried over Na₂SO₄. Evaporation of the solvent mixture under reduced pressure and recrystallization of crude products led to colourless solids.

**Fig. 3.** The general structure of compounds 6 with atom numbering (for NMR assignments)

**4-(Bromophenylsulfonyl)-N-(1-oxo-1-phenylpropan-2-yl)benzamide 6a,** obtained by reaction with benzenel; 96% yield (method 1), 80% yield (method 2); m.p. 127-129°C (cyclohexane).

**UV-Vis (CH₃OH, λ nm) (lg e):** 203.5 (4.46), 251.1 (4.35); FT-IR (KBr, ν cm⁻¹): 3475, 3088, 3063, 2983, 2937, 2876, 1693, 1650, 1598, 1573, 1521, 1485, 1450, 1357, 1342, 1295, 1189, 1159, 857, 775;

**1H-NMR (DMSO-d₆, δ ppm, J Hz):** 1.36 (d, 7.0, 3H, H-18); 5.50 (quint, 7.0, 1H, H-4), 7.53 (br t, 7.4, 2H, H-21, H-23), 7.64 (br t, 7.4, 1H, H-22), 7.85 (d, 9.1, 2H, H-14, H-16), 7.91 (d, 9.1, 2H, H-13, H-17), 7.99 (dd, 1.7, 7.4, 2H, H-20, H-24), 8.04 (d, 8.8, 2H, H-7, H-11), 8.08 (d, 8.8, 2H, H-8, H-10), 9.10 (d, 7.0, 1H, NH);

**13C-NMR (DMSO-d₆, δ ppm):** 13.68 (7.0, 3H, CH₃), 19.85 (7.0, 3H, CH₃), 21.10 (CH₃), 50.61 (C-4), 126.09 (C-23), 127.56 (C-8, C-10), 128.85 (C-8), 129.81 (C-9, C-11), 132.49 (C-20, C-24), 133.03 (C-14, C-16), 134.91 (C-19), 137.64 (C-20), 138.60 (C-6), 139.84 (C-22), 141.18 (C-14), 142.74 (C-9), 164.69 (C-2), 202.27 (C-5);

**RP-HPLC (CH₃OH:H₂O = 60:40, 1 ml/min, 250 nm):** purity 97.46%; t = 5.88 min;

Anal. (%): Calcd. for C₂₃H₂₀BrNO₄S (486.38 g/mol): C, 57.60; H, 4.43; N, 2.80; S, 6.41. Found: C, 57.68; H, 4.43; N, 2.88; S, 6.59.

**4-(Bromophenylsulfonyl)-N-(1-oxo-1-p-tolylpropan-2-yl)benzamide 6b,** obtained by reaction with toluene; 97% yield (method 1), 86% yield (method 2); m.p. 140-143°C;

**UV-Vis (CH₃OH, λ nm) (lg e):** 203.9 (4.46), 253.1 (4.35); FT-IR (KBr, ν cm⁻¹): 3395, 3093, 3066, 2980, 2937, 2876, 1693, 1650, 1598, 1573, 1521, 1485, 1450, 1357, 1342, 1295, 1189, 1159, 857, 775;

**1H-NMR (DMSO-d₆, δ ppm, J Hz):** 1.38 (d, 7.0, 3H, H-18); 5.50 (quint, 7.0, 1H, H-4), 7.55 (br t, 7.4, 2H, H-21, H-23), 7.64 (br t, 7.4, 1H, H-22), 7.85 (d, 9.1, 2H, H-14, H-16), 7.91 (d, 9.1, 2H, H-13, H-17), 7.99 (dd, 1.7, 7.4, 2H, H-20, H-24), 8.04 (d, 8.8, 2H, H-7, H-11), 8.08 (d, 8.8, 2H, H-8, H-10), 9.10 (d, 7.0, 1H, NH);

**13C-NMR (DMSO-d₆, δ ppm):** 13.68 (7.0, 3H, CH₃), 19.85 (7.0, 3H, CH₃), 20.23 (CH₃), 50.61 (C-4), 127.68 (C-8, C-10), 128.28 (C-21, C-23), 128.80 (C-8), 128.85 (C-7), 129.81 (C-9, C-11), 132.49 (C-20, C-24), 133.03 (C-14, C-16), 134.91 (C-19), 138.63 (C-6), 139.96 (C-12), 142.89 (C-14), 164.73 (C-2), 198.90 (C-5);

**RP-HPLC (CH₃OH:H₂O = 60:40, 1 ml/min, 250 nm):** purity 97.55%; t = 6.97 min;

Anal. (%): Calcd. for C₂₄H₂₂BrNO₄S (500.40 g/mol): C, 57.60; H, 4.43; N, 2.80; S, 6.87; Found: C, 57.65; H, 4.37; N, 2.81; S, 6.36.

**General procedures for the synthesis of 5-aryl-2-[4-(bromophenylsulfonyl)phenyl]-4-methyloxazoles 7**

**Method 1.** The crude N-[1-aryl-1-oxopropan-2-yl]-4-(bromophenylsulfonyl)benzamides 6 (10 mmol) were refluxed in 20 mL phosphorus oxychloride for 4 h. The excess of POCl₃ was removed under vacuum. After cooling, the oily residue was treated with a mixture of ice-water and extracted twice with 20 mL CH₂Cl₂. The organic layers were combined and washed several times with 5% NaHCO₃ solution, then with water and dried (Na₂SO₄). After evaporation of the solvent, the crude products were recrystallized from ethanol as colourless needle-shaped crystals.

**Method 2.** The N-[1-(aryl-1-oxopropan-2-yl)-4-(bromophenylsulfonyl)benzamides 6 (10.51 mmol) were dissolved in 40 mL ethyl acetate. Acetic anhydride (3 mL, 31.75 mmol) and 98% sulfuric acid (0.17 mL, 3.19 mmol) in 2.5 mL ethyl acetate were added. The reaction mass was heated at reflux for 3 h. After cooling to room temperature, a 2.52N NaOH solution (25 mL) was added. The reaction mixture was heated at reflux for another 30 min and then cooled to room temperature. The obtained precipitate was filtered off and washed with cool 1N HCl, then with cool 10% NaCl solution and finally, with cool water. The layers of filtrate were separated and the organic layer was washed with 1N HCl, then with 10% NaCl solution, dried over Na₂SO₄, and evaporated under vacuum, leaving a second fraction of crude product. The high purity colourless crystals of title compounds were obtained after purification.
FT-IR (KBr, ν cm⁻¹): 3092m, 3065w, 2923m, 2861w, 1601s, 1572s, 1495s, 1469m, 1484m, 1328v, 1291s, 1279s, 1159v, 1096v, 843s, 571v;
1H-NMR (CDCl₃, δ ppm, / Hz): 2.26 (s, 3H, CH₃), 2.33 (s, 3H, CH₃), 2.38 (s, 3H, CH₂), 7.09 (br d, 8.0, 1H, H-23), 7.14 (br s, 1H, H-21), 7.25 (d, 8.0, 1H, H-24), 7.65 (d, 8.5, 2H, H-14, H-16), 7.82 (d, 8.5, 2H, H-13, H-17), 7.99 (d, 8.8, 2H, H-7, H-11), 8.16 (d, 8.8, 2H, H-8, H-10);
13C-NMR (CDCl₃, δ ppm): 125.3 (C-18), 20.40 (CH₃), 21.38 (CH₂), 126.78 (C-23), 126.90 (C-8, C-10), 128.41 (C-7, C-11, C-12), 128.82 (C-15), 129.41 (C-13, C-17), 129.97 (C-24), 131.86 (C-21), 132.36 (C-6), 132.87 (C-14, C-16), 135.07 (C-19), 137.50 (C-20), 139.68 (C-22), 140.77 (C-12), 141.12 (C-9), 147.89 (C-4), 150.07 (C-5), 175.73 (C-22);
+ESI-MS/MS (m/z, rel. abund. %): 482 [M+H⁺]; 466/468 [M+H⁺]+; 323/325 [BrCH₂SO₂⁺]; 279 [M+H⁺+Br⁺]; 263 (100, BP) [M⁺+Br⁺]; 248 [M⁺+H⁺+Br⁺]; 158 [M⁺+Br⁺];
RP-HPLC (CH₂Cl₂:H₂O = 70:30, 1 mL/min, 335 nm): purity 96.80%, tᵣ 5.68 min;
Anal. (%): Calcd. for C₁₉H₁₈BrNO₃S: C, 59.76; H, 4.18; N, 2.90; S, 6.46; Found: C, 59.82; H, 4.11; N, 2.84; S, 6.68.

2-[4-(3-Bromophenylsulfonyl)phenyl]-5-mesityl-4-pentyloxazole 7d
89% yield (method 1), 92% yield (method 2); m.p. 155-157°C (ethanol);
UV-Vis (CH₃OH, λ nm) (lg ε): 204.4 (4.47), 249.3 (4.21), 324.2 (4.13);
FT-IR (KBr, ν cm⁻¹): 3089w, 3066w, 2921m, 2863w, 1600s, 1573s, 1499m, 1472m, 1455m, 1325m, 1290s, 1159s, 1101s, 845m, 575v;
1H-NMR (CDCl₃, δ ppm, / Hz): 2.27 (s, 3H, CH₃), 2.29 (s, 3H, CH₃), 2.38 (s, 3H, CH₂), 7.10 (s, 1H, H-23), 7.13 (s, 1H, H-21), 7.66 (d, 8.5, 2H, H-14, H-16), 7.83 (d, 8.5, 2H, H-13, H-17), 7.99 (d, 8.8, 2H, H-7, H-11), 8.17 (d, 8.8, 2H, H-8, H-10);
13C-NMR (CDCl₃, δ ppm): 12.57 (C-18), 19.31 (CH₃), 19.68 (CH₂), 19.88 (CH₂), 126.83 (C-8, C-10), 128.37 (C-7, C-11, C-12), 128.82 (C-15), 129.36 (C-13, C-17), 130.96 (C-23), 132.28 (C-6, C-14, C-16), 134.19 (C-19), 138.30 (C-20, C-24), 139.13 (C-22), 140.58 (C-12), 141.91 (C-9), 147.90 (C-4), 157.95 (C-5), 175.70 (C-22);
RP-HPLC (CH₂Cl₂:H₂O = 70:30, 1 mL/min, 335 nm): purity 90.58%, tᵣ 6.02 min;
Anal. (%): Calcd. for C₁₉H₁₈BrNO₃S: 496.42 g/moI: C, 60.49; H, 4.47; N, 2.82; S, 6.46; Found: C, 60.54; H, 4.47; N, 2.85; S, 6.42.

Cytotoxicity evaluation
The Daphnia magna bioassay was performed under constant temperature and light conditions (at 25 ± 1°C, in the dark) using a Sanyo MLR-351-H, USA climatic chamber.
The determinations were made in duplicate against algal (positive control) and 1% DMSO (negative control). The experiment was carried out according to the protocol previously described [27-29]. From each compound, six concentrations ranging from 5 to 200 μg/ml were tested. The lethality curves were plotted using the logarithm of concentrations and against lethality percentage, L (%), recorded at 24, 48 and 72 h. The prediction was performed with the GUSAR software application.

Results and discussions
Chemistry
In the light of the above importance of oxazol-5(4H)-ones and oxazoles, is seems of interest to synthesize new
heterocyclic compounds from these classes and their acyclic intermediates using the reaction sequences from scheme 1. The key precursor, 4-(4-bromophenylsulfonyl) benzoic acid 1, and corresponding acyl chloride 2 were already described in literature [30,31]. Compound 1 was synthesized by Friedel-Crafts reaction between bromobenzene and 4-methylbenzene-1-sulfon chloride (p-tosyl chloride) in the presence of anhydrous AlCl3 at reflux, followed by oxidation of 4-(4-bromophenylsulfonyl)-1-methylbenzene with chromium trioxide in glacial acetic acid at reflux [30]. The acid 1 was then converted by reaction with SOCl2 into 4-(4-bromophenylsulfonyl)benzoyl chloride 2 [20,21] which was used without further purification for N-acylating α-alanine according to Steiger’s procedure in order to obtain 2-[4-(4-bromophenylsulfonyl)benzamido]propanoic acid 3. This compound was then cyclodehydrated to the corresponding azlactone 4 by two methods using either ethyl chloroformate in the presence of N-methylmorpholine in methylene chloride at room temperature or acetic anhydride at reflux. Cyclization in basic medium may be considered to take place according to the similar mechanism to that we previously described for other 2,4-disubstituted-5(4H)-oxazolone [20].

The N-acylated amino acid 3 was also converted through a nucleophilic substitution reaction with excess of thionyl dichloride at reflux into the corresponding acyl chloride 5.

The AlCl3-catalyzed acylaminoacylation of the aromatic hydrocarbons (in excess both as reactant and solvent) with 5(4H)-oxazolone 4 was carried out at ambient temperature and led to the α-acylamino ketones 6, with a high regioselectivity and at excellent yields - which increase in the order: benzene, toluene, m-xylene, in agreement with the increasing nucleophilicity of these substrates and the stability of the corresponding Wheland intermediate in electrophilic aromatic substitution (EAS). The proposed ring opening reaction mechanism is formerly indicated by us in the literature [21]. Compounds 6 have also obtained by Friedel-Crafts acylation of aromatic hydrocarbons with 2-[4-(4-bromophenylsulfonyl)benzamido]propanoyl chloride 5, but the reaction yields were lower. These results indicate that 5(4H)-oxazolones are better N-acylating reagents than N-acyl-α-aminocarboxyl chloride.

In the Robinson-Gabriel synthesis conditions, by using phosphorl trichloride or concentrated sulfuric acid in the presence of acetic anhydride in ethyl acetate, the above N-(1-aryl-1-oxopropan-2-yl)-4-(4-bromophenylsulfonyl) benzamides 6a-c were cyclodehydrated affording 2,5-diaryl-4-methylxoxazoles 7a-c in very good yields.

Generally, intermediate compounds from α-acylamino ketones class (6a-c) were isolated as pure colourless crystals and characterized physico-chemically, with the exception of 4-[4-(4-bromophenylsulfonyl)-N-(1-mesityl-1-oxopropan-2-yl)benzamido 6d (obtained by reaction with mesitylene), which could not be isolated in pure form, but which has been used in crude state in the synthesis of the corresponding oxazole 7d.

The proposed mechanism for synthesis of 2,5-diaryl-4-methylxoxazoles 7 from 2-aza-1,4-diaryl-3-methyl-1,4-butanediones 6 in the presence of excess phosphorus oxychloride occurs via the enolized forms I and then through the ester-dichloride intermediates of phosphoric acid II. The leaving group dichlorophosphate, -POCl2, is replaced by chloride anion by bimolecular nucleophilic substitution mechanism in order to form chlorinated compounds III. The unstable chloride anions IV are then obtained, which lead to the formation of the compounds with oxazole ring 7 by intramolecular nucleophilic addition accompanied by elimination of chloride anion (scheme 2a). In acid medium, the reaction mechanism for obtaining 1,3-oxazoles 7 involves the protonation of compounds 6 with the formation of two electrophilic structures in resonance: oxonium ions V and corresponding carbocations VI. The carbocations VI were then deprotonated simultaneously with the cyclization by nucleophilic attack at C-4, leading to the corresponding unstable hemiketals (2,5-diaryl-5-hydroxy-4-methyl-4,5-dihydroxazoles) VII. An intramolecular dehydration reaction of these intermediates affords heterocyclic compounds from 1,3-oxazoles class 7 (scheme 2b). This mechanism is in accordance to literature data [32], based on 18O-labeling, which indicated that the amidic oxygen from acyclic intermediates is maintained in the oxazole ring and the ketonic oxygen is removed as water.

The chemical structures of the new compounds are confirmed due to their spectral (UV-Vis, IR, 1H-NMR, 13C-NMR, MS) and elemental analysis.

Generally, the electronic absorption spectra of the new compounds presented a sharp band at λ 202.6-204.4 nm (E band) and an absorption at λ 249.3-255.5 nm (B band). In addition, the compounds 3 and 4 show a third absorption maximum of weak intensity at λ 226.4 nm, 228.2 nm, respectively (K band). The presence of an additional intense absorption maximum at higher longest-wavelengths, λ 324.2-341.9 nm, is observed in the UV spectra of the new oxazoles 7 compared with those of acyclic precursors 6. This bathochromic shift is due to extending of conjugation by formation of oxazole chromophore.

Presence of the characteristic absorption bands in IR spectra of the synthesized products provides useful information for determining the structure of newly compounds 3-7. Thus, 2-[4-(4-bromophenylsulfonyl) benzamido]propanoic acid 3 and N-(1-aryl-1-oxopropan-2-yl)-4-(4-bromophenylsulfonyl) benzamides 6 exhibited the following characteristic absorption bands at wavenumbers: 3347-3396 cm⁻¹ for N-H stretching, ν(N-H), at 1686-1708 cm⁻¹ due to carbonyl absorption, ν(C=O), and at 1644-1666 cm⁻¹ due to amidic carbonyl group stretching vibration, ν(C=O) (amide I band). Characteristic of these compounds is also the amide II band, assigned to deformation vibration of N-H group, μ(N=O), present in the region 1523-1537 cm⁻¹. Amide III band (ν(C=O)-ν(N-H)) is strong and very broad, extending from 3000 cm⁻¹ to 3000 cm⁻¹. This absorption overlaps the medium sharper C-H stretching peaks, which are extending beyond the O-H envelope.

Evidence for the obtaining of acyl chloride 5 are presence in IR spectrum of two strong absorption bands due to ν(C=O) at 1738 cm⁻¹ (fundamental vibration) and 1788 cm⁻¹ (Fermi resonance band), and a medium band due to ν(C=O) at 1662 cm⁻¹.

The IR spectra of heterocyclic compounds 4 and 7 were clearly distinguished from those of corresponding acyclic intermediates 3 and 6, respectively by having different characteristic wavenumbers, in agreement with the literature data [20,21]. Thus, in IR spectrum of azlactone 4, the absorption band due to the valence vibration of carbonyl group is shifted at 1719 cm⁻¹, while the ν(N=O), ν(C=O), ν(C=O) (amide I band), and ν(C=O) (amide II band) at 1662 cm⁻¹. Also, the IR spectra of oxazoles 7 revealed the absence of signals in the N-H vibrations.
...and C=O regions. The peaks at 1650 cm\(^{-1}\) (from 4), and in the range 1594-1601 cm\(^{-1}\) (from 7) were assigned to the C=N stretching vibration of these new heterocycles.

The formation of compounds 3, 4, 6 and 7 was further confirmed by the \(^1\)H-NMR spectra. Assignments of the signals are based on the chemical shift and intensity pattern. Furthermore, the 2D \(^1\)H-\(^1\)H COSY experiments allow unambiguous assignments.

The \(^1\)H-NMR spectra of the compounds 3 and 6 exhibited a doublet attributed to secondary amide proton at a chemical shift between 8.95-9.10 ppm.

The \(^1\)H-NMR spectra of compounds 4 and 7 contain two sub-spectra characteristic of the 5(4H)-oxazolone and oxazole ring, respectively and of the diarylsulfone moiety. The signal attributed to the one proton of the NH group from acyclic precursors 3 and 6 is absent in the \(^1\)H-NMR spectra of corresponding heterocycles 4 and 7, respectively and this proves that these new compounds have been obtained.

In the \(^1\)H-NMR spectra of the compounds 3 and 6, the methine proton from C-4 appears as a quintet at 4.41 ppm (3) and 5.26-5.50 ppm (6), while for azlactone 4 was observed at 4.49 ppm as a quartet and in the case of oxazoles 7 it is absent.

Evidence for the formation of the oxazoles 7 was provided by their \(^1\)H-NMR spectra, which revealed a downfield shift in the signal attributed to the three protons (H-18) of the methyl group in 4-position from \(\delta\) 1.32-1.39 ppm in \(\alpha\)-acylamino ketones 6 as a doublet (because of vicinal couplings with H-4) to 2.38-2.50 ppm in oxazoles 7 as a singlet, due to the stronger deshielding effect of oxazole ring compared to that of the C=O and CONH groups from acyclic intermediates 6. Also, the methyl doublet in azlactone 4 showed a discernible downfield shift of 0.21 ppm relative to the acyclic precursor 3, due to the stronger deshielding effect of oxazolone ring compared to that of the COOH and CONH groups from compound 3.

The signals in \(^13\)C-NMR spectra are also in good agreement with the proposed structures for the newly synthesized compounds. The assignment of the signals in \(^13\)C-NMR of 3, 4, 6 and 7 resulted from the 2D \(^1\)H-\(^13\)C HETCOR experiments.

The chemical shift of the C-4 atom from N-acyl-a-amino acid 3 at 48.36 ppm is downfield after intramolecular cyclodehydration to 5(4H)-oxazolone 4 with 13.03 ppm. Also, in the oxazoles 7 the C-4 signal was more deshielded with \(=\) 96 ppm (\(\delta\) 147.25-147.90 ppm) by comparison of the signal of the same atom from 6 (\(\delta\) 50.34-52.61 ppm) and this confirmed that cyclization of the \(\alpha\)-acylamino ketones 6 took place. It can be noticed the apparition of the downfield signal attributed to the C-2 at \(\delta\) 175.70-176.04 ppm from the oxazole nucleus, while the carbon atom signal attributed to the amidic carbonyl group from intermediates 6 (in the range 198.22-202.27 ppm) revealing an upfield shift for this carbon in the oxazole structure, which is a further indication that the oxazole formation had taken place.
Furthermore, an additional support for the assigned structures of new compounds 3, 4, 6c and 7c was obtained by recording their mass spectra by LC-ESI-MS/MS analysis.

**Cytotoxicity evaluation**

The results of *Daphnia magna* bioassay are presented in table 1. LC50 could not be calculated for any of the tested compounds at 24 and 48 h due to an L% below 10%. At 72 h, the highest toxicity was induced by compound 6b, followed by 6a, 7d and 6c. Compound 4 showed toxicity comparable with 7a, whereas compound 3 induced an approximately 3-fold lower toxicity than compound 4. As expected, no lethality was recorded for α-alanine during the experiment. Compounds belonging to 7-series presented a higher toxicity as opposed to the 6-series compounds. The predicted values of LC50 for all newly synthesized compounds showed a high toxicity. However, the prediction was confirmed only for compound 6b and in a lesser extent for compound 6a.

**Conclusions**

Ten newly compounds from *N*-acyl-α-amino acid, *N*-acetyl-α-amino acid chloride, 1,3-oxazol-5(4H)-one-α-acylamino ketone and 1,3-oxazole class were synthesized and characterized. The new azlactone 4 has been obtained by the reaction of acyl chloride 2 with α-alanine, followed by cyclodehydration of the new *N*-acyl-α-alanine 3. The new α-acylamino ketones 6 have been obtained by treatment of 5(4H)-oxazolone 4 or new *N*-acyl-α-alanyl chloride 5 with aromatic hydrocarbons under Friedel-Crafts reaction conditions. Finally, by refluxing these intermediates 6 with phosphorous oxychloride or sulfuric acid in the presence of acetic anhydride, the intramolecular ring closure occurred with formation of the new oxazoles 7. The structure of compounds was confirmed by elemental analysis and different spectral methods.

The newly synthesized compounds 3, 4, 6a-c, 7a-d have been investigated for their biological activity on *Daphnia magna*. Compounds 6a and 6b showed the highest toxicity, comparable with the predictions performed using GUSAR software. However, further studies are needed in order to investigate the mechanism of action and the therapeutic potential of the compounds.

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