BENZIMIDAZOLE SCHIFF BASES MICROWAVE ASSISTED SYNTHESIS AND THE EFFECT ON LEUKEMIA CELLS WITH FLOW CYTOMETRY

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The present paper introduces a simple and efficient method for the synthesis of a 2-aryl-substituted benzimidazole by heterocyclization of 1,2-phenylenediamine and 5-aminoisophthalic acid in the presence of polyphosphoric acid as catalyst under solvent-less conditions, which produced good yield of corresponding benzimidazoles in a short reaction time. Two new bisbenzimidazoles and tetrakisbenzimidazole Schiff bases were synthesized by reaction between 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA) and terephthalaldehyde or 4-formylbenzoic acid. The condensations proceed in short time to give products which, in certain instances, are not readily attainable by conventional condensation techniques. Due to its good performance, the microwave device used in household was preferred. The structures of the compounds were assigned by FT-IR, 1H NMR, 13C NMR and elemental analysis. The effects of benzimidazoles and Schiff bases on the cancerous cells have been known. Synthesized by bisbenzimidazole and bisbenzimidazole/tetrakisbenzimidazole-Schiff bases derivatives 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA), N,N’-(1,4-phenylenebis(methan-1-yl-1-ylidene))bis(3,5-di(1H-benzo[d]imidazol-2-yl)aniline) (PM-BIMA), (E)-4-((3,5-di(1H-benzo[d]imidazol-2-yl)phenylimino)methyl)benzoic acid (BIM-PMBA) with flow cytometric analysis was performed, blastic cells containing 90%. We measured rates of cell death in cells blast on the matter.

INTRODUCTION

Benzimidazoles and their derivatives are important classes of heterocyclic molecules in several field of organic chemistry.1-5 Bisbenzimidazole and bisbenzimidazole/tetrakisbenzimidazole-Schiff bases derivatives are key components in a large number of bioactive compounds of both natural and synthetic origin.6-12 They are a common heterocyclic scaffold in biologically active and medicinally significant compounds13,14 and are found in many natural products.15 Moreover, these groups of heterocyclic compounds exhibit a wide range of pharmacological properties, which include antiviral,16 anticancer,17-19 antimicrobial20-23 antitumor,24-25 antifungal,26-27 anti-inflammatory activity,28-29 anti-parasitic,30,31 antibacterial,32 and many others properties. Benzimidazoles have shown exciting and diverse biological and pharmaceutical properties and are therefore considered privileged structures in the design and discovery of new drugs.33

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The compounds with imine group in their structure, known as Schiff bases, are synthesized by the condensation reaction of the amines with the carbonyl groups. Schiff bases of benzimidazole have been reported with remarkable antibacterial and antimicrobial activities. Traditionally, bisbenzimidazoles were most often been prepared from the reaction of o-phenylenediamine with 5-aminoisophthalic acid under harsh dehydrating conditions, using strong acids. However, Microwave-assisted organic synthesis (MAOS) continues to affect synthetic chemistry significantly by enabling rapid, reproducible experiments. Particularly, the use of milder reagents has improved both the yield and purity of this reaction. Today, cancer patients are being treated with radiotherapy, chemotheraphy and surgical interventions. It is known that the procedures applied in this process are heavy for the patient. Since chemotherapy drugs kill both live and cancer cells the patient's immune system weakens. Therefore, the effect of bisbenzimidazole/tetrakisbenzimidazole-Schiff bases on cancer cells being well known, we have planned to obtain Schiff bases from benzimidazole in order to protect live cells, and to study the effect of leukemia cells on flow cytometry. In the patients diagnosed with leukemia, the appropriate panels were studied by applying the synthesized substances as a 10% solution to cells and the measurements were recorded in the device. The present study reports a method for achieving bisbenzimidazole/tetrakisbenzimidazole-Schiff base systems formed by condensation reaction 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA). Herein, we reported the synthesis BIMA and its bisbenzimidazole/tetrakisbenzimidazole-Schiff base as a new template. We envisioned that microwave irradiation would enhance this chemistry and expand the chemistry scope.

RESULTS AND DISCUSSION

3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA) was prepared by the reaction of 1,2-phenylenediamine with 5-aminoisophthalic acid (Scheme 1, Fig. 1). The structural formula of the 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA) was verified by elemental analysis, 1H NMR and FT-IR. The band at 1665 cm\(^{-1}\) for the C=O stretch in the FT-IR spectrum disappeared and appeared at 1686–1673 cm\(^{-1}\) for azomethine C=N group PM-BIMA and BIM-PMBA, respectively. In the ligand BIM-PMBA, the band at 3055 cm\(^{-1}\) can be assigned to the -OH group vibrations. To identify the structures of BIMA, PM-BIMA and BIM-PMBA, the 1H NMR spectra were recorded in DMSO-d6. The 1H NMR data were also in good correlation with the structures of the synthesized compounds. The 1H NMR spectrum signals of ligands BIM-PMBA and PM-BIMA at \(\delta\) (ppm), 10.05-10.13 ppm, correspond to the -NH proton, respectively (Fig. 2 and Fig. 3). The signal for HC=O group disappeared and gave the signal at higher frequency for the -CH=N and –OH proton in the 1H NMR spectrum. The signals of 13C NMR spectrum of BIM-PMBA (Fig. 2) were in a good agreement with the proposed formula. The "benzimidazole Schiff bases" prepared in this way was obtained in nearly quantitative yield and high purity.

Flow cytometry was conducted to investigate cell death of bisbenzimidazole and bisbenzimidazole/tetrakisbenzimidazole-Schiff base derivatives on CD34 including cancer cells. As shown in Fig. 4, Fig. 5 and Fig. 6 according to measurements. The cavity between the benzimidazole ligands is thought to be suitable for the diameter of the leukemia cells (Scheme 3).

First tube shows the values that are leukemia blast cells. Vital CD34 cells/microliters: 18039.65/microliters and Viable CD45 cells/microliters: 33901.39/microliters. If we take this as the standard, we can observe the exchange of added substances to tube two and tube three on the blast cells. Added to the second tube "PM-BIMA"s effect on the Viable CD34 cells/microliters:974.36/microliters and Vital CD45 cells/microliters:40696.39/microliters (Table 1).
Scheme 1 – Proposed structures of the microwave-assisted bisbenzimidazole formation (BIMA).

Fig. 1 – $^1$H NMR Spectrum of 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA).
Scheme 2 – Proposed structures of the microwave-assisted bisbenzimidazole/tetrakisbenzimidazole Schiff bases.
Fig. 2 – $^1$H-$^{13}$C NMR Spectrum of 4-((3,5-di(1H-benzimidazol-2-yl)phenylimino)methyl)benzoic acid (BIM-PMBA).

Fig. 3 – $^1$H NMR Spectrum of N,N’-(1,4-phenylenebis(methan-1-yl-1-ylidene))bis(3,5-di(1H-benz[d]imidazol-2-yl)aniline) (PM-BIMA).
Scheme 3 – The bisbenzimidazole/tetrakisbenzimidazole-Schiff bases and leukaemia cells.

Table 1

<table>
<thead>
<tr>
<th>Gate Statistics AML Sample</th>
<th>AML Sample PM-BIMA</th>
<th>AML Sample BIM-PMBA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alive CD34 Cells/µL</td>
<td>18039.65</td>
<td>974.36</td>
</tr>
<tr>
<td>Alive CD45 Cells/µL</td>
<td>33901.39</td>
<td>40696.39</td>
</tr>
<tr>
<td>Total CD34 Cells/µL</td>
<td>24367.57</td>
<td>12034.09</td>
</tr>
<tr>
<td>The alive percentage of CD45</td>
<td>73.87 %</td>
<td>8.08 %</td>
</tr>
<tr>
<td>The alive percentage of CD34, CD45</td>
<td>0.04 %</td>
<td>0.04 %</td>
</tr>
</tbody>
</table>

Fig. 4 – The standard values of AML sample (Tube 1).
Added to the third tube “BIM-PMBA”’s effect on the Viable CD34 cells/microliters: 7015.67/microliters and Viable CD45 cells/microliters: 20163.89/microliters (Table 1).

Identified CD34 cells in AML patients are blast cells. CD45 cells are antibodies that must be presented in every human body. Total cells death is determined with 7AAD also.

EXPERIMENTAL

Materials and Methods

All the chemicals and solvents used for this work were purchased from Sigma Aldrich. LG-health wave microwave system (MG-607APR, 230V—50Hz) was used and the output of microwave power is mentioned as percent intensity i.e. (20%, 40%, 60%, 100%). Melting points were measured using an Optimelt Automated Melting Point System (Digital Image Processing Technology) SRS apparatus. Elemental analyses (CHN) were performed using a Leco, CHNS-932 model analyzer. 1H NMR spectra were recorded at room temperature with a Varian, 400 MHz spectrometer using TMS as an internal standard. FT-IR spectra were recorded with a Perkin-Elmer Spectrum 100 with Universal ATR Polarization Accessory. The pH values were measured with a WTW pH 537 pH meters. Flow cytometry was used to influence the substance of blast cells. A reaction mixture of o-phenylenediamine and appropriate acid was subjected to microwave irradiation at 350W for 25 minutes. Completion of the reaction was monitored by TLC.

Preparation of Microwave Assisted 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA). Equimolar amounts of o-phenylenediamine and the 5-aminoisophthalic acid were mixed with a sufficient quantity of polyphosphoric acid to give a stirrable paste. The mixture was heated using microwave irradiation technique at 350W for 25 minutes, permitted to cool, and poured in a thin stream into a large
volume of rapidly stirred water. The insoluble residue was collected by filtration, washed with a small amount of water and resuspended in an excess of 10% sodium carbonate solution. The alkaline slurry was filtered and the product washed thoroughly with water. The product was recrystallized from alcohol or an aqueous alcohol mixture after treatment with a small amount of activated charcoal. Completion of the reaction was monitored by TLC. (BIMA); Yield: 60%. The CHN analyses of the compound corresponded to the molecular formula C_{20}H_{15}N_{5}. Elemental analysis was found as: C, 73.83; H, 4.62; N, 21.49%. Calc.: C, 73.83; H, 4.65; N, 21.52%. FT-IR (cm⁻¹): 3048 (NH), 1606 (C=C), 1450 (hetero-ring). ¹H NMR (400 MHz, DMSO-d₆) (δ ppm): 8.19 (s, H, CH₉), 7.62–7.61 (d, 4H, CH₉), 7.52 (s, 2H, CH₉), 7.72–7.21 (d, 4H, CH₉), 5.61 (s, 2H, NH₂).

**Preparation of Microwave Assisted N,N’-(1,4-phenylenebis(methan-1-yl-1-ylidene))bis(3,5-di(1H-benzo[d]imidazol-2-yl)aniline) (PM-BIMA) and 4-(3,5-di(1H-benzoimidazol-2-yl)phenylimino)methyl)benzoic acid (BIM-PMBA).** The Schiff bases, PM-BIMA and BIM-PMBA were obtained as solids by refluxing a mixture of (BIMA) (7.8 g; 24 mmol) and terephthalaldehyde (1.60 g; 12 mmol) or (BIMA) (3.25 g; 10 mmol) and 4-formylbenzoic acid (1.50 g; 10 mmol) in ethanol (35 mL) which was heated using microwave irradiation technique at 350W for about 25 minutes. The solid was washed with ether and recrystallized from ethanol and dried under vacuum. (PM-BIMA); Yield: 65%; m.p.: 310 °C; The CHN analyses of the compound corresponded to the molecular formula C_{30}H_{28}N_{10}. Elemental analysis was found as: C, 76.94; H, 4.34; N, 18.68%. Calc.: C, 76.99; H, 4.31; N, 18.70. FT-IR (cm⁻¹): 2816 (NH), 1686 (CH=NH), 1498 (C=N heterocyclic). ¹H NMR (400 MHz, DMSO-d₆) (δ ppm): 10.13 (s, 4H, NH), 9.03 (s, 2H, CH=N), 8.30 (m, 4H, CH₉), 8.10 (s, 4H, CH₉), 7.70 (m, 8H, CH₉), 7.50 (s, 2H, CH₉), 7.61–7.03 (m, 8H, CH₉). (BIM-PMBA) Yield: 70%; m.p.: 298 °C; The CHN analyses of the compound corresponded to the molecular formula C_{36}H_{25}N_{12}O_{2}. Found: C, 73.50; H, 4.22; N, 15.28; O, 6.93%. Calc.: C, 73.51; H, 4.19; N, 15.31; O, 6.91 %. FT-IR (cm⁻¹): 3055 (OH), 3290 (NH), 1673 (CH=NH), 1600 (C=C), 1538 (C=N heterocyclic), 1281 (C=O). ¹H NMR (400 MHz, DMSO-d₆) (δ ppm): 10.05 (s, 2H, NH), 8.95 (s, H, CH=N), 8.20 (s, H, CH₉), 8.12–8.10 (d, 2H, CH₉), 7.96–7.94 (d, 2H, CH₉), 7.92–7.90 (d, 2H, CH₉), 7.63–7.61 (m, 4H, CH₉), 7.22–7.20 (m, 4H, CH₉). ¹³C NMR (100 MHz, DMSO-d₆, 25 °C) (δ ppm): 167.46, 152.64, 152.02, 150.86, 149.92, 139.75, 136.38, 134.00, 132.42, 130.35, 130.33, 129.98, 129.39.

**The effects of substances on blast cells by flow cytometry:** Materials were solved in ethanol in three different tubes. On the other hand, we used the blood samples containing 90% blasts. We have prepared three different flow cytometry tubes. We’ve also added 20 µl “CD34/CD45/7AAD” surface markers into tubes each containing 100 µl of blood. We waited 20 minutes. We prepared lysing solution (20 µl lysing solution/1000 µl of water) and we added it to the tubes (2000 µl). We left 10 minutes of incubation in the dark. Then we washed the cells and divided it into three tubes: (1) tube; We studied the flow cytometry device, we recorded values. This value refers to the patient’s percentage of “CD34” (Blast cells) before the addition of substances (Fig. 4); (2) tube; “PM-BIMA (10%)” have added, after five minutes we did the measurement in flow cytometry, we recorded values (Fig. 6); (3) tube; “BIM-PMBA (10%)” have added, after 5 minutes we did the measurement in flow cytometry, we recorded values (Fig. 6).

**CONCLUSIONS**

In this study, N,N’-(1,4-phenylenebis(methan-1-yl-1-ylidene))bis(3,5-di(1H-benzo[d]imidazol-2-yl)aniline) (PM-BIMA) and (E)-4-((3,5-di(1H-benzo[d]imidazol-2-yl)phenylimino)methyl)benzoic acid (BIM-PMBA) were synthesized by the reaction of 3,5-di(1H-benzo[d]imidazol-2-yl)aniline (BIMA) with terephthalaldehyde and 4-formylbenzoic acid. Compounds (PM-BIMA) and (BIM-PMBA) were originally synthesized. The structures of the compounds were assigned by FT-IR, ¹H NMR and elemental analysis. If we look at the flow cytometry measurements, the second tube added “PM-BIMA”s molecular structure, size and the benzimidazole activity, CD34 cells were quickly killed. Four kinds of benzimidazole groups and two Schiff bases were included. The third tube added “BIM-PMBA” benzimidazole effect is half up to the “PM-BIMA”. In the substances, as the benzimidazole and Schiff base increase, the effect on the blast cells also increases. We also determined that by adding “FM-BIFA” material into the second tube, the analyses with flow cytometer, the CD45 cells that must have been in all alive were increased. This material killed the blast cells and increased alive cells.

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