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Synthesis and Characterization of Novel Five-Membered Heterocycles and Their Activity against Candida Yeasts

Hiba Maher Tawfeeq,1 Rasim Farraj Muslim,2 Obaid Hasan Abid3 and Mustafa Nadhim Owaid2,4

1 Department of Chemistry, College of Education for Pure Sciences, University Of Anbar, Anbar 31001, Iraq
2 Department of Ecology, College of Applied Sciences, University Of Anbar, Anbar 31007, Iraq
3 Department of Scientific Affairs and Graduate Studies, University of Fallujah, Anbar, Iraq
4 Department of Heet Education, General Directorate of Education in Anbar, Ministry of Education, Anbar 31007, Iraq

* Corresponding author: E-mail: dr.rasim92hmts@gmail.com

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Abstract

Some new tetrazole derivatives were prepared by the reaction between the prepared azomethine compounds I6–I10 with sodium azide in anhydrous tetrahydrofuran (THF) with a few drops of distilled water and under reflux conditions. Azomethine compounds were prepared by thermal condensation reactions of aromatic aldehydes with primary aromatic amines. The prepared compounds (tetrazole derivatives) were screened for their antibacterial activity (by disc diffusion method). Compound I6 is the most active derivative that has recorded a significantly (p<0.01) stronger influence to inhibit the growth of Candida zeylanoides with an average zone of inhibition of 26.0 mm. Derivatives I7 and I9 showed the lowest zone of inhibition of 8.0 mm against Candida zeylanoides. This study may be helpful in designing more potential anticandidal agents for therapeutic use in the future.

Keywords: Candida sp.; pharmaceutical; azomethine; sodium azide; biological activity.

1. Introduction

Azomethine compounds discovered by Hugo Schiff in 1864, can be prepared by different methods, one of the more important being the condensation reaction between primary amine with aldehyde.1 Azomethine compounds contain the N=C group.2 Some of the azomethine compounds are used as antibacterial agents.3,4 The structure of azomethine compounds usually includes a phenyl or aryl group with the double bond between the carbon atom and the nitrogen atom.5,6 (Scheme 1)

The reaction of triazole diamine compound with 4-bromobenzaldehyde in the presence of glacial acetic acid gave the next product.7 (Scheme 2)

Tetrazole derivatives are heterocyclic compounds containing four nitrogen atoms and one carbon atom within one ring.8 Tetrazoles as a group of heterocyclic compounds appear in IR spectra as broad signals; having peculiar biological activities.9,10 Tetrazole derivatives have a special structure and can display anti-bacterial properties, such as antiviral and anti-allergic.11–13 There are several methods to prepare tetrazole derivatives, and each method depends on the constituents of the reaction.14 Recently, there were many various types of compounds (including such containing a metal centre coordinated with suitable ligands) tested against Candida albicans, with a varying degree of success.15–17 An example of one of tetrazole derivatives is the product from the reaction between azomethine compound (biphenyl bis
(1-(pyridin-4-yl) methanimine)) with sodium azide. (Scheme 3)

Many of tetrazole derivatives can be prepared by the reaction between a different aldehyde and different amines. (Scheme 4)

In this study, tetrazole derivatives derived from the reaction of the prepared azomethine compounds with sodium azide were evaluated for their biological activity against four types of Candida yeasts. The products were identified by their melting points, FT-IR and \(^{1}H\) NMR spectra.

### 2. Experimental

#### 2.1. Apparatuses

The measurement of melting point was conducted by the electrothermal melting point apparatus. IR spectra were recorded at room temperature in the range of 400–4000 cm\(^{-1}\) by a Fourier transform infra-red Spectrophotometer Model Tensor 27 Bruker Co., Germany. The \(^{1}H\) NMR spectra were recorded on Bruker Ac-300 MHz spectrometer.

#### 2.2. Preparation of Azomethine Compounds I\(_{1}\)–I\(_{5}\)

Azomethine compounds were prepared according to the literature procedure, in an equimolar mixture 0.02 mole of aldehydes and 0.02 mole of amines and trace of acidic catalyst in 25 mL absolute ethanol were reacted at reflux temperature for 4 hours, whereby a crystalline solid separated out. The products were filtered off and recrystallized from absolute ethanol.

#### 2.3. Preparation of Tetrazole Derivatives I\(_{6}\)–I\(_{10}\)

A mixture of azomethine compounds (0.01 mol) and sodium azide (0.01 mol) was dissolved in 20 mL of THF and 2 mL distilled water and refluxed for 4 hours and left to stand for 24 hour at room temperature, then the solid product separated out. \(^{21}\) The products were filtered off and recrystallized from absolute ethanol as shwon in Table 2.
2. 4. Activity against Candida Yeasts

This test was carried out in vitro to investigate the inhibitory effects of the prepared tetrazole derivatives using well diffusion method on Muller-Hinton agar. This experiment was done as mentioned by Owaid et al. with some modifications.22 Four milligrams of the prepared tetrazole derivatives were applied separately in 6 mm-well. After 18 hour at 37 °C, the zone of inhibition was measured using the ruler in millimeters.

3. Results and Discussion

Tables 1 and 2 show structural formulae, names, yields, melting points and color of all prepared compounds I₁–I₁₀. The best yield achieved for the prepared azomethine compounds was for compounds I₃ 82% and I₄ 84%, while the lowest yield was for compound I₁ 78% and the best yield of the prepared tetrazole derivatives was for I₁₀ 92%, while the lowest yield was for I₆ 76%. The highest melting point for azomethine compounds was for I₅ 240–242 °C.
pound I₂, the lowest melting point was for compound I₁, while the highest melting point of the prepared tetrazole derivatives was for compound I₇, the lowest melting point was for compound I₆. The different colors and melting points of the products compared with the raw material are initial evidence of interaction.

3. 1. Azomethine Compounds I₁–I₅

Azomethine compounds were prepared from commercially available aldehydes with primary amines and identified by their melting points and FT-IR spectra. Table 3 shows the appearance of the stretching absorption bands of the characteristic groups of the resulting group (C=N) at 1609–1681 cm⁻¹ beside the characteristic bands of the residual groups in the structure, being indicative of the formation of the products.²³,²⁴, 3, 9a- Tetrahydrobenzo [e] [1,3] oxazepin - 5(5aH

<table>
<thead>
<tr>
<th>Compound</th>
<th>C≡N C=C C–H C–H</th>
<th>FT-IR, ν (cm⁻¹)</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Aromatic</td>
<td>Aromatic</td>
<td>C–H Aliphatic Symmetric</td>
</tr>
<tr>
<td>I₁</td>
<td>1681</td>
<td>1601</td>
<td>3042 3105 2969 2849 NO₂ C=N pyrimidine 1634</td>
</tr>
<tr>
<td>I₂</td>
<td>1609</td>
<td>1580</td>
<td>3042 3113 2972 NO₂ O–H 3491, NH 3278</td>
</tr>
<tr>
<td>I₃</td>
<td>1609</td>
<td>1581</td>
<td>3041 3089 – – NO₂ N–H 3283, C–Cl 823</td>
</tr>
<tr>
<td>I₄</td>
<td>1611</td>
<td>1581</td>
<td>3042 3087 – – NO₂ N–H 3263</td>
</tr>
<tr>
<td>I₅</td>
<td>1615</td>
<td>1592</td>
<td>3042 3112 – – NO₂ O–H 3422, N–H 3257</td>
</tr>
</tbody>
</table>

The general equation (Scheme 5) represents the main reaction through which the prepared azomethine compounds were obtained. The mechanism of azomethine compounds formation was thoroughly studied and established by many authors in the literature.²⁵,²⁶

![Scheme 5. The main reaction of azomethine compounds](image)

3. 2. Tetrazole Derivatives I₆–I₁₀

In this work, the preparation of tetrazole compounds was achieved by the reaction between prepared azomethine compounds (I₁–I₅) with sodium azide. The resulting products were identified using the melting points, FT-IR and ¹H NMR spectra. Table 4 shows characteristic stretching absorption bands at 1219–1286 cm⁻¹, 1007–1083 cm⁻¹ and 1487–1509 cm⁻¹ indicative of C–N, N–N and N=N bonds of tetrazol rings beside the characteristic stretching absorption bands of the residual groups in their structure.²⁷

The ¹H NMR spectrum of compound I₇ (Fig. 1, in solvent DMSO-d₆) showed the following signals: singlet at δ 1.76 indicating the presence of 3H as an methoxy group (OCH₃), singlet at δ 3.83 indicating the presence of 1H as an NH group (NH outside of the tetrazole ring), singlet at δ 8.55 indicating the presence of 1H as another NH group (NH inside of the tetrazole ring), singlet at δ 9.33 indicating the presence of 1H as one CH group (N-CH), singlet at δ 11.56 indicating the presence of 1H as one hydroxy group (OH), multiplet at δ 8.96–6.99 indicating the presence of 6H of aromatic protons. ¹H NMR spectrum of compound I₈ (Fig. 2) shows the following signals: singlet at δ 3.33 indicating the presence of 1H as an NH group (NH outside of the tetrazole ring), singlet at δ 8.71 indicating the presence of 1H as another NH group (NH inside of the tetrazole ring), singlet at δ 11.71 indicating the presence of 1H as one CH group (N-CH), multiplet and doublet of doublet at δ 8.41–7.56 indicating the presence of 7H of aromatic protons.²⁷ Other chemical shifts for compounds I₆, I₉ and I₁₀ are given in Table 5.
Table 4. FT-IR of tetrazole derivatives $I_6$–$I_{10}$

<table>
<thead>
<tr>
<th>Compound</th>
<th>N–H (cm$^{-1}$)</th>
<th>N–N</th>
<th>N=N</th>
<th>C–N (cm$^{-1}$)</th>
<th>C=O (cm$^{-1}$)</th>
<th>C–H Aromatic (cm$^{-1}$)</th>
<th>C–H Symmetric (cm$^{-1}$)</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I_6$</td>
<td>3383</td>
<td>1007</td>
<td>1500</td>
<td>1286</td>
<td>1606</td>
<td>3044</td>
<td>2976</td>
<td>2849</td>
</tr>
<tr>
<td>$I_7$</td>
<td>3280</td>
<td>1020</td>
<td>1509</td>
<td>1273</td>
<td>1614</td>
<td>3113</td>
<td>2972</td>
<td>2880</td>
</tr>
<tr>
<td>$I_8$</td>
<td>3266</td>
<td>1083</td>
<td>1489</td>
<td>1219</td>
<td>1615</td>
<td>3088</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>$I_9$</td>
<td>3299</td>
<td>1083</td>
<td>1487</td>
<td>1269</td>
<td>1585</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>$I_{10}$</td>
<td>3258</td>
<td>1074</td>
<td>1488</td>
<td>1276</td>
<td>1612</td>
<td>3113</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Table 5. The $^1$H NMR spectral data of tetrazole derivatives $I_6$–$I_{10}$ (in DMSO-$d_6$).

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Shifts $\delta$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I_6$</td>
<td>Singlet at 1.28 (6H, 2CH$_3$), singlet at 8.68 (H, -NH), singlet at 10.19 (1H, N-CH), singlet at 9.77 (1H, -OH), singlet and doublet of doublet at 8.43–8.08 (5H, aromatic protons).</td>
</tr>
<tr>
<td>$I_7$</td>
<td>Singlet at 1.76 (3H, O-CH$_3$), singlet at 3.83 (1H, NH out), singlet at 8.55 (1H, NH in), singlet at 9.33 (1H, N-CH), singlet at 11.56 (1H, -OH), multiplet at 8.96–6.99 (6H, aromatic protons).</td>
</tr>
<tr>
<td>$I_8$</td>
<td>Singlet at 3.33 (1H, NH outside of the tetrazole ring), singlet at 8.71 (1H, NH inside of the tetrazole ring), singlet at 11.71 (1H, N-CH), multiplet and doublet of doublet at 8.41–7.56 (7H, aromatic protons).</td>
</tr>
<tr>
<td>$I_9$</td>
<td>Singlet at 3.26 (1H, NH outside of the tetrazole ring), singlet at 8.69 (1H, NH inside of the tetrazole ring), singlet at 11.71 (1H, N-CH), multiplet at 8.41–7.79 (7H, aromatic protons).</td>
</tr>
<tr>
<td>$I_{10}$</td>
<td>Singlet at 3.40 (1H, NH out), singlet at 8.88 (1H, NH in), singlet at 10.07 (H, N-CH), singlet at 11.57 (1H, -OH), multiplet and doublet of doublet at 8.59–6.85 fr (7H, aromatic protons).</td>
</tr>
</tbody>
</table>

![Image of 1H NMR Spectrum of $I_7$]
The reaction of the azomethine compounds with sodium azide is given in the equation in Scheme 6.

From the reaction course and the suggested mechanism it may be concluded that the reaction takes place via concerted mechanism (Huisgen 1,3-dipolar cycloaddition).28

3. 3. Activity against Candida Yeasts

Zone of inhibition of some human pathogenic yeasts was determined by the well-diffusion method and used to test the potential of the prepared tetrazole derivatives (I_{6–I_{10}}) as shown in Fig. 3 and 4. Compound I_{6} was found to be the best derivative that has significantly (p<0.01) recorded a stronger influence to inhibit the growth of Candida zeylanoides at an average of the zone of inhibition of 26.0 mm, followed by 24.6–25.6 mm for the rest of the species of Candida. Next, I_{8} derivative recorded zone of inhibition of 11.0 mm toward Candida guilliermondii. Furthermore, I_{7} and I_{9} showed the lowest zone of inhibition of only 8.0 mm against Candida zeylanoides. Additionally, I_{6} derivative recorded zone of inhibition of 11.3 mm against Candida guilliermondii and Candida zeylanoides, respectively. Compound I_{10} did not inhibit the growth of Candida species as shown in Fig. 3. Resistance mechanism depends on which specific paths are inhibited by the drugs.
and if the alternative paths are available to substitute for those paths that the compound has inhibited; in this way the microorganism can modify its pathways and be able to survive by developing resistance.29,30 These results agree with some recent studies which described the synthesis of hybrid heterocycles proving to have in vitro antimicrobial, antibacterial and antifungal activities.

4. Conclusions

The results of FT-IR and $^1$H NMR showed that the five-membered ring compounds were the least obstructed during all preparation processes and that neither light nor humidity affect the prepared compounds, proving that the prepared compounds have an excellent stability. $I_6$ is the best derivative that has significantly ($p < 0.01$) recorded a stronger influence to inhibit the growth of *Candida zeylanoides* at an average zone of inhibition of 26.0 mm. This study may be helpful in designing more potential anticandidal agents for therapeutic use in the future.

5. Acknowledgements

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6. References

Povzetek
Z reakcijo natrijevega azida s predhodno pripravljenimi azometinskimi spojini I₆–I₁₀ smo v brezvodnem tetrahidrofuranu (THF) ob dodatku nekaj kapljic destilirane vode pod pogoji refluksa pripravili nekaj novih derivatov tetrazola. Azometinske spojine smo sintetizirali s termično kondenzacijo aromatskih aldehidov s primarnimi aromatskimi aminami. Pripravljenim spojinam (derivatom tetrazola) smo določili antibakterijske aktivnosti (z metodo difuzije v disku). Spojina I₆ se je izkazala za najbolj aktivni derivat z visoko (p < 0.01) povečanim vplivom zaviranja rasti organizma Candida zeylanoides (s povprečno vrednostjo premera inhibicije 26.0 mm). Derivata I₇ in I₉ pa sta izkazala najslabše inhibitorno delovanje s premerom inhibicije le 8.00 mm proti istemu organizmu (Candida zeylanoides). Ta študija bi lahko pomagala pri načrtovanju novih bolj učinkovitih spoj in kvasom iz rodu Candida, ki bi bile v prihodnosti celo uporabne v terapevtske namene.