“Anti-microbial Study and Synthesis of Schiff bases of 3-actyl 4-hydroxy Quinolin-2-one”

SHIVRAJ S. ANJANIKA1 and SANTOSH S. CHANDOLE2*

1Department of Chemistry, Sharadchandra College, Naigaon, District-Nanded MS-431709, India.
2Department of Chemistry, S. G. B. College, Purna Jn., MS-431511, India.
*Corresponding author E-mail: schandole@rediffmail.com

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ABSTRACT

Quinoline based new Schiff bases were synthesized from 3-Acetyl 4-Hydroxy Quinolin-2-(1H)-one and screened their antibacterial and antifungal activity. The Schiff bases 4-hydroxy-3-(1-((4-picolin-2-yl)imino)ethyl)quinolin-2-(1H)-one(L1), 4-hydroxy-3-(1-((5-picolin-2-yl)imino)ethyl)quinolin-2(1H)-one(L2), 4-hydroxy-3-(1-((6-picolin-2-yl)imino)ethyl)quinolin-2(1H)-one(L3) and 4-hydroxy-3-(1-((3-nitro-4-picolin-2-yl)imino)ethyl)quinolin-2(1H)-one(L4) were prepared from 3-Acetyl 4-Hydroxy Quinolin-2-One with 2-amino picolines. The structures of Schiff bases were confirmed by Infrared, mass, proton-NMR and 13CNMR spectral analysis. In vitro studies of these Schiff bases were carried out for their antibacterial activity by Agar contact method and antifungal activity by the poison plate method. The bacterial species used were B. subtilis, E. coli, S. typhi and S. aureus. Fungal species used were F. monelliforme, A. niger, A. flavus, and P. chrysogenum.

Keywords: 3-Actyl 4-Hydroxy Quinolin-2-One, Amino picoline, Spectral study, Schiff bases, Biological study.

INTRODUCTION

Study of nitrogen containing heterocyclic moiety is being very popular area of interest for researchers as these possess pharmacological properties. Quinoline is an important fused heterocyclic aromatic compound. 4-hydroxyquinolin-2(1H)-one, the essential moiety of major interest of the many research as it has importance due to its synthetic, medicinal values. The importance of this structural moiety is due to its presence in many naturally occurring organic heterocyclic compounds. Bucharidine (I) and foliosidine (II) is quinoline alkaloid, extracted from Haplophyllum bucharicum and Haplophyllum foliosum respectively. Both have estrogenic action. Viral-RNA polymerase Inhibitory action of few 4-hydroxyquinoline-2-one derivatives such as Compounds (III) and (IV) strongly prevent the replication of the Hepacivirus C.

In addition to this 4-Hydroxy-2(1H)-quinolinones along with their derivatives are among the group of valuable heterocyclic compounds associated with many pharmacological various medicinal values such as analgesic, anti-
inflammatory,15 diuretic,16 antiallergen,17 orally active antagonists,18 cardiovascular agents,19 anticonvulsant,20 antimicrobial (antibacterial and antifungal),21-23 antitubercular,24 dye-stuffs.25

Heating mantle, the reaction mixture in the alcohol is heated for three hours at refluxing temperature. The mixture is cooled three hours later. The solid Schiff base is vacuum filtered after being washed with ethanol. The Schiff base is dried and recrystallized from ethanol. The purity of the Schiff bases was checked by m.p. and TLC.

**Biological activity**

**Antibacterial activity**

Anti-bacterial activity was performed by agar contact method.30 B. subtilis and S. typhi were gram+ve bacteria that were utilized as test organisms, whereas S. aureus and E. coli were gram-ve microorganisms. Mueller Hinton Agar for bacteria was used for all tests for antibacterial activity. Ampicillin was used as positive control for bacteria. The solvent and positive control used was DMSO. Antibiotics and dehydrated media powder were brought from Hi-Media, India. Using sterile wire-loop, test organisms were aseptically added to sterile MH broth before being incubated at 37°C for 18 hours. This suspension was utilized as an inoculant. Wells in the media plates with a 10mm diameter were made using a sterile cork borer for the addition of compound solutions and controls. With the aid of a micropipette, 100 µL of the compound solution was aseptically poured to the wells to reach a ultimate strength of 10 g of compound in each well. As controls, the same quantity of DMSO and ampicillin solution were introduced. The plates were cooled for 30 min to allow solutions to diffuse through the agar substrate. Further, Plates were incubated at 37°C for a period of 24 hours. The zone margin should be regarded as the region that does not clearly display any expansion that the unaided eye can see. With a measuring scale in millimetres, the clean zone was measured.

**Antifungal activity**

The poison plate approach was used to provide antifungal activity.31 For the evaluation of antifungal activity, Potato Dextrose Agar (PDA) media was utilized as a culture. The sterilization of the medium was archived by autoclaving at 120-125°C for 25-30 min under 15 psi of pressure. 20 mL of sterilized, melted PDA was added to sterilized petri plates with 2 mL of each component, and the mixture was then gently stirred in a circular motion to get homogenized. With positive Neomycin and negative DMSO controls, the identical process
was followed. *A. niger, A. flavus, F. moneliforme,* and *P. chrysogenum* were chosen to assess the antifungal activities. The fungal spores from the slant culture were transferred to a test tube containing sterile saline and thoroughly mixed with a sterile wire loop. As an inoculant, this spore solution was employed. The plates were kept for incubation for 100 h at room temperature. Further, the growth of the infected fungi was monitored on the plates. The outcomes were noted.

**RESULT AND DISCUSSION**

All reactions were conducted using standard procedures. In the presence of sodium ethoxide, methyl anthranilate and ethylacetoacetate were refluxed to produce the intermediate 3-acetyl-4-hydroxy-quinolin-2(1H)-one (I) needed for the synthesis of Schiff bases. The purity of the intermediate product (I) was assessed by TLC after it was recrystallized in ethanol. Various substituted 4-hydroxy-3-(1-(heteroarylimino)ethyl)quinolin-2-one (L₁-L₄) were prepared carrying out reaction in ethanol for 4 hours.

In the analytical results as detailed above, the significance of the peaks identified in the IR, ¹H NMR, and ¹³CNMR spectra of the compounds (L₁-L₄) is clarified. The compound (L₃-L₄) IR spectra have shown a prominent band at 3507-3498 cm⁻¹ and is given to the (-OH) vibration, confirming the presence of enolic -OH group present in Schiff bases. The two bands at 1570-1504 cm⁻¹ and 1470-1422 cm⁻¹ are designated to the aromatic ring. Strong band between 1668 and 1658 cm⁻¹ is assigned for lactam carbonyl.

Each of the (L₁-L₄) ¹H NMR spectra showed a singlet(3H) in the range 2.23-2.38 ppm that was attributed to an methyl hydrogen bonded to imine group. A singlet (3H) in the region 2.22-2.65 ppm is given to picoline’s methyl substituent . The peaks observed in the region 8.2 and 7.0 ppm were ascribed for aromatic Hydrogen atoms. The existence of the 4-hydroxyl group is confirmed by a wide singlet at 15.63–15.92 ppm. The peak observed between 10.50 – 10.64 ppm reveals the presence of secondary amino group. The lactam carbon revealed peaks in the range of 165–161 ppm, while imine carbon showed peaks in the range of 176–175 ppm. The explanation provided for other peaks found in ¹H NMR, ¹³CNMR, and mass spectra, as well as molecular ion peaks, supports the structures of compounds (L₁-L₄).

The synthesized Schiff’s bases were investigated for anti-bacterial with *Bacillus subtilis* and *Salmonella typhi* (Gram-positive bacteria) while *Staphylococcus aureus* and *Escherichia coli* (Gram-negative bacteria). The results are reported in Table 1. All compounds have displayed good antibacterial activity with all bacterial species in the range of 10-14mm diameter of zone of inhibition but lesser, except L₄ which shown maximum zone of inhibition within range of 16-18mm of diameter than reference used. The enhanced activity observed in L₄ might be due to presence of nitro group in the moiety. The screening test for antifungal activity against *F. moneliforme, A. niger, A. flavus,* and *P. chrysogenum.* fungi revealed that (L₁-L₄) exhibit significant activity, especially L₄ have shown minimum growth of all fungi.

<table>
<thead>
<tr>
<th>Synthesized Schiff base</th>
<th>Zone of Inhibition (diameter measured in mm)</th>
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<tbody>
<tr>
<td></td>
<td>Gram-positive</td>
</tr>
<tr>
<td></td>
<td>S. typhi</td>
</tr>
<tr>
<td>Ampicillin (Reference)</td>
<td>19</td>
</tr>
<tr>
<td>L₁</td>
<td>13</td>
</tr>
<tr>
<td>L₂</td>
<td>14</td>
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<tr>
<td>L₃</td>
<td>13</td>
</tr>
<tr>
<td>L₄</td>
<td>17</td>
</tr>
</tbody>
</table>

Spectral data for the synthesized compound is given as below

L₁: 4-hydroxy-3-(1-(4-picolin-2-yl)imino)ethyl quinolin-2(1H)-one

Yield: 72%; Colour: Yellow; Melting Point: 234-236°C IR (KBr, cm⁻¹): 3500 Broad –O–H and enolic, 3402 Broad and weak >N–H, 1668(>C=O)

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**Fig. 2. Synthesis of Schiff bases L₁ to L₄**

**Table 1: Anti-bacterial activity**

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**Synthesized Schiff base**

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**Yield:** 72%; **Colour:** Yellow; **Melting Point:** 234-236°C **IR (KBr, cm⁻¹):** 3500 Broad –O–H and enolic, 3402 Broad and weak >N–H, 1668(>C=O)
of lactam, 1614 (>C=N) of imine, 1600 (>C=N) of picoline, 1570 and 1505 and 1450 aromatic (>C=C), 1242 (enolic –OH interaction), 750 (>N-H). 1HNMR (CDCl3) (300 MHz): 2.23 (S, 3H, imine –CH3), 2.40 (S, 3H, –CH3 of substituted at Py-moiety), 7.23 S (1H) of Py-moiety, 8.23 (d, 2H of Py-moiety ), 7.23(d, 8.09-7.28 (Ar─H of Quinoline moiety), 81 for C3, 114-140 for aromatic carbons, 161 for C4, 162 for lactam carbon, and 175 for imine carbon. Mass Spectra[M+1]+ : 294.27.

L2: 4-hydroxy-3-(1-((5-picolin-2-yl)imino)ethyl) quinolin-2(1H)-one
Yield: 75%; Colour: Yellow; Melting Point: 220-222°C. IR (KBr, cm⁻¹): 3506Broad –OH and enolic, 3292Broad and weak >N-H, 1658(>C=O) of lactam, 1611 (>C=N) of imine, 1599 (>C=N) of picoline, 1566 and 1445 aromatic (>C=C), 1252 (enolic –O-H) interaction, 773 (>N─H). 1HNMR (CDCl3) (300 MHz): 2.23 (S, 3H, imine –CH3), 2.49 (S, 3H, –CH3 of substituted at Py-moiety), 8.36 S (1H) of Py-moiety, two doublets at 8.27-8.26 and 6.96-6.88 of 3C & 4C hydrogens of Py-moiety), 8.06-7.21 (Ar─H of quinolone moiety), 16.75 (bs, S, 1H, O-H), 10.50 (bs, S,1H, N-H). 13CNMR (CDCl3) (300 MHz): 19 and 20 (imine –CH3 carbon and –CH3 of substituted at picoline ring), 81 for C3, 114-140 for aromatic carbons, 160 for C4, 163 for lactam carbon, and 175 for imine carbon. Mass Spectra[M+1]+:294.27.

L3: 4-hydroxy-3-(1-((6-picolin-2-yl)imino)ethyl) quinolin-2(1H)-one

L4: 4-hydroxy-3-(1-((3-nitropicolin-2-yl)imino)ethyl) quinolin-2(1H)-one
Yield: 70%; Colour: Green; Melting Point: 239-241°C. IR (KBr, cm⁻¹): 3507 Broad –O–H and

**CONCLUSION**

New Schiff bases were synthesized by condensation of 3-acetyl-4-hydroxy-quinolin-2(1H)-one with substituted amino picoline. All the compounds were characterized by spectral study which favors the structure of targeted molecules. These synthesized Schiff’s bases were further subjected to their antimicrobial activity and revealed that they possess good antibacterial and antifungal activity.

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REFERENCES


