Synthesis, Characterization and Antibacterial Activity of Some Penicillin Derivatives

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Abstract

This study illustrated the synthesis of two new di amidine compounds ([c] and [d]) by reaction of 6-amino penicillanic acid (6-APA) with di α-amino nitrile compounds ([a] and [b]). [a] and [b] di α-amino nitrile compounds synthesized from the condensation reaction on aldehyde and di amine in the presence of potassium cyanid as one pot three components reaction. The new di amidine compounds ([c] and [d]) have been proven their efficiency by inhibiting some types of bacteria (Staphylococcus aurous, Streptococcus, Escherichia coli and klebsiella). Amidine compound [d] showed better effect than [c] against the selected bacteria. The synthesized compounds were characterized by conventional techniques using infrared spectrophotometer (IR) and proton nuclear magnetic resonance (¹H-NMR).

Keywords:
Amidine
6-amino penicillanic acid
Alpha-amino nitrile
Penicillin derivatives
E.coli

1. Introduction

Amidine is an organic compounds have form N=C–N and considered to be analogue to carboxylic acid, ester or amide [1]. Amidine was first synthesized by Gerhardt by reacting N-phenylbenzimidyl chloride and aniline in the nineteenth century [2]. Amidines enter various organic reactions to prepare different compounds, especially cyclic compounds [3], just like Schiff bases [4]. Amidines and bis amidine compounds are play an important role in treating many diseases and almonds, such as inflammations [5] cancer [6] alzheimer’s disease [7] parasites [8], and the most common application in the past was as anti-bacterial agent [9-12].

Amidines can be prepared in different ways using multiple chemicals [13-15], especially from nitrile compounds [16,17]. Penicillin is anti-bacterial agent, useful in treating some human diseases but is limited by the ability of some bacteria to acquire resistance to the drug. Therefore new derivatives are required, some researchers synthesize new organic compound or to control the resistance of bacteria. The interpretation of this work is to synthesize derivatives of penicillin with two beta-lactam groups to achieve the role of structure activity-relationship property [18].

2. Experimental

Chemicals and Instruments

All materials used in this study were provided and purchased from the original manufacturing places Aldrich, Fluka and B.D.H. Infra-red spectra were recorded with FT-IR BRUKER spectrophotometer and SHIMADZU spectrophotometer 8300 in wave number range of (400-4000) cm⁻¹ and ¹H-NMR spectra were recorded on VARIAN 500 MHz spectrometers using deuterated dimethyl sulfoxid (DMSO-d6) as solvent.

Organic Syntheses

Synthesis of α-amino nitrile compounds [a] and [b] [19]:
2 mmole of aldehyde (verataldehyde or vanillin) was dissolved in 15 mL glacial acetic acid followed by the addition of 1 mmole of di-amine compound (benzidine). Small amount of p-toluene sulfonic acid was added to the reaction as catalyst with stirring to produce di imine compound, 4 mmole of KCN (excess) was added to the reaction after half hour and stirred for two days. The reaction mixture poured onto crashed ice and made slightly alkaline with ammonia solution addition, then left over night. The precipitate was filtered by Buchner funnel and washed with water to yield di-α-amino nitrile compounds [a] and [b] respectively. Scheme 1 shows the reaction of aldehydes with amine compound in the presence of potassium cyanide. To diagnose the presence of the nitrile group chemically, small amount of [a] or [b] was heated with sodium hydroxide solution (10%), the released ammonia gas was detected by wet red litmus paper that indicated the formation α-amino nitrile compound. Table 1 included physical data of [a] and [b] α-amino nitrile compounds.

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Synthesis of amidine compounds [c] and [d] [20]:

1 mmole of α-amino nitrile [a] or [b] was dissolved in 15 mL absolute methanol and mixed with 5 mL sodium methoxide (23 mg sodium metal dissolved in 3 mL methanol). The mixture stirred for two hours, and then 2 mmole of 6-aminopenicillanic acid was added. The reaction mixture was refluxed for 8 hours. Amidine compounds [c] and [d] were produced respectively. Scheme 1 shows the reaction of α-amino nitrile compound with amine. Compounds [c] and [d] filtered and washed with water, then purified and crystallized by ethyl acetate/methanol solvents. Table 1 included physical data of [c] and [d] amidine compounds.

![Scheme 1](image)

Table 1. The physical properties of organic compounds.

| Symbol | Chemical formula | Molecular weight (g mole⁻¹) | Color | M.p °C | Yield % |
|--------|------------------|-----------------------------|-------|--------|--------|
| [a]    | C₃₂H₃₀N₄O₈       | 534.23                      | Yellow| 200-205| 88     |
| [b]    | C₃₀H₂₆N₄O₄       | 506.2                       | Red   | 220-225| 80     |
| [c]    | C₄₈H₅₄N₈O₁₀S₂    | 966.34                      | Orange| 213-215| 77     |
| [d]    | C₄₆H₄₆O₁₀S₂      | 954.34                      | Brawn | 218-220| 70     |

Antimicrobial activity

In this study, the desired products amidine compounds [c] and [d] were prepared and biological efficacy was tested against four types of bacteria, two are gram positive (Staphylococcus aureus and Streptococcus) and two are gram negative (Escherichia coli and Klebsiella) using the lowest concentration of the substances 10⁻³ M.

The antibacterial activities were evaluated using well diffusion method on Mueller-Hinton agar by working zones in millimeter (mm) which filled with 100 μl of the test samples and incubated at 37 °C for one day. After incubation, the diameter of the inhibition zones measured using a ruler. The area of inhibition is clear and surrounding the place of injection of the sample and bacterial not growth in it.

3. Results and Discussion

This study aims to synthesize new penicillin compounds of two symmetrical terminals possessing amidine and beta-lactam groups and study the behavior against different types of bacteria. The organic compounds that have been prepared have proven good efficacy against the types of bacteria used in order to contain two effective groups of beta-lactams with addition to the final formulation of amidine (N=C=N), which has previously proven effective as an anti-bacterial so, expected that the synthesized compounds are promising as antibiotic compounds in the pharmaceutical field against different kinds of bacteria and especially for the resistant type.
Infra-red spectroscopy
The absorption bands of infrared spectra of [a] and [b] showed peaks at 2228 cm\(^{-1}\) and 2235 cm\(^{-1}\) respectively which could be attributed to nitrile group that generally appeared some times as a weak peak at range (2250-2200) cm\(^{-1}\) [21,22]. This supported by chemical test mentioned previously in the experimental. The reaction of [a] or [b] with 6-APA to produce [c] and [d] must involve disappearing of nitrile group and forming amidine group instead. The absorption bands of infrared spectrum of [c] and [d] showed new peaks at 1633 cm\(^{-1}\) and 1614 respectively which represented (N=C–N) starching that normally presented near this area [20,23,24]. The attached 6-APA part showed characteristic peaks of carboxylic carbonyl (COOH) and beta–lactam carbonyl (C=O) that occurred at 1670 cm\(^{-1}\) and 1730 cm\(^{-1}\) for [c] and [d] respectively. Hydroxyl group in [c] and [d] appeared as broad peak between 3300-3500 cm\(^{-1}\) [22]. Table 2 shows the characteristic absorption bands of [a], [b], [c] and [d] compounds.

| Compound | O–H | νN–H | νC≡N | νC=O carboxylic | νC=O β-lactam | νN=C–N Amidine | δ N–H Binding | C=C Aromatic |
|----------|-----|------|------|-----------------|----------------|----------------|----------------|--------------|
| [a]      | –   | 3355 | 2228 | –               | –              | –              | 1610           | 1503,1430    |
| [c]      | –   | 3355 | 1672 | 1733            | 1630           | 1609           | 1585           |
| [b]      | 3364| 3220 | 2235 | –              | –              | –              | 1609           | 1510,1447    |
| [d]      | 3504| 3352 | –    | 1670            | 1730           | 1614           | –              | 1597         |

Proton nuclear magnetic resonance
Proton nuclear magnetic resonance \(^1\)HNMR was used for the characterization of the newly synthesized amidine compound. \(^1\)HNMR spectra were used to confirm the final structure of compounds.

\(^1\)H-NMR amidine compound
The \(^1\)H-NMR spectrum of amidine compound [c], Figures 1 showed singlet signal near (1.4) ppm which could be attributed to methyl groups which appeared at range (1-2.5) ppm rang [25,26]. Dihydrothiazine ring (–SH–C–) signal appeared at (3.2) ppm which generally noticed at range (3-3.6) ppm [26]. Amine group (N–H) which occurred at rang (3-5) ppm normally [22].

Three peaks appeared as doublet for β-lactam ring were observed around (4.2-4.4) ppm could be attributed to (CH–CH), (CO–CH) and (N–CH) which appeared at range (4.4.9) [26]. New peaks appeared at (10.5-10.6) ppm which attributed to hydroxyl (O–H) of carboxylic acid which generally fixed at range (10.5-12) ppm rang [22]. Amidine group (–HN–CH=NH) appeared as singlet signal in two different locations this is because the double bond was in delocalization forms, so one peak attributed to (–NH–C–) form, which appeared at range (1.3-2.8) ppm, while the other peak could be attributed to (NH=–C–) form, which appeared at range (4-7) ppm [24].
Biological study
In our study, compounds [c] and [d] were evaluated in vitro against four types of bacteria, two of grams positive (Staphylococcus aurous and Streptococcus) and two of grams negative (Escherichia coli and Klebsiella), using well diffusion method and follow the amount of inhibition of these ligands through the use of the least concentration of the inhibitor. The compounds appeared good results, due to the fact that compounds contained two antibacterial groups which are amidine [27] and beta-lactam groups [28] postulated in two terminals. Figure 2 shows the inhibition zones of [c] and [d] on agar culture. Table 3 shows the inhibition zones of the prepared compounds against bacteria.
Table 3. The inhibition zone of [c] and [d] compounds against bacteria.

| Compounds | Escherichia coli | Klebsiella | Staphylococcus aureus | Streptococcus |
|-----------|-----------------|------------|----------------------|---------------|
| [c]       | 11              | 10         | 10                   | 12            |
| [d]       | 18              | 20         | 11                   | 20            |

4. Conclusion

Compounds [c] and [d] were evaluated in vitro against four types of bacteria, and showed good efficacy by inhibiting bacteria diffusion, due to the presence of two well known beta-lactam groups as anti bacterial, in addition to the presences of two amidine groups. Eventually, novel compound structures achieved, having powerful structure-activity relationship property. This is in the interest of fighting drug-resistance bacteria. Amidine compound [d] showed better effect than [c] against the selected bacteria. the presence of hydroxyl group (i.e. [d]) instead of methoxy group (i.e. [c]) could play role in the structure-activity relationship, that it form hydrogen bonding which could assisted in the interaction toward bacteria.

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