Synthesis, Magnetic, Spectral and Antibacterial Properties of Some Metal(II) Complexes of Mixed Drugs, Aspirin and Vitamin B₂

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Introduction

Metal complexes are now the focus of drug designs due to the success of metallo drugs like Auranofin, Flamazine and Cis-platin[1]. This has expanded the data base on chemo-therapeutic agents which are less toxic, with improved anticancer, antioxidant and antimicrobial activities. Thus, metal complexes could serve as lead compounds for the replacement of existing ones in the combat against toxicity and resistance[2]. Aspirin is an analgesic with anti-inflammatory and antipyretic actions[3,4]. On the other hand, riboflavin is a water-soluble vitamin that is used in the treatment of neonatal jaundice, migraine and together with UV light is effective for inactivation of pathogens in body fluids. Its deficiency results in sore throat, edema and anemia[5,6]. Literature is available on metal complexes of individual analgesics and vitamins including Aspirin and Riboflavin[5-10]. However, no information is available yet on the metal(II) complexes of the mixed drugs-Aspirin and Riboflavin[11-16]. Since metallo drugs contain a metal and drug(s) as active ingredients, we thought of using the metals, Mn, Fe, Co, Ni, Cu & Zn, and Riboflavin (vitamin B₂) being essential in humans, to act as mineral and vitamin supplements. The combination of the various metals, Aspirin and Riboflavin are also expected to improve antibacterial activities[17,18]. Our group, has over a decade focused on the search for novel metallo-analgesic and metallo-vitamin complexes as lead compounds in drug research for the management of pain and vitamin deficiency[13-16]. Thus, our aim was to synthesize the metal(II) complexes of the mixed drugs-Aspirin and Riboflavin{M = Mn, Fe, Co, Ni, Cu & Zn} and investigate their magnetic properties for cooperative phenomena (spin crossover, ferro- and anti-ferromagnetisms). In addition their antibacterial properties against some pathogenic bacteria were also verified for their suitability as broad-spectrum antibacterial agents.
Materials and Methods

[Fe(HL)(HL²)SO₄]·H₂O was prepared by the addition of 1.08 g (3.89 x 10⁻³ mol) of Fe⁺² to a stirring homogeneous solution of 0.70 g (3.89 x 10⁻³ mol) of Aspirin, HL) and 1.46 g (3.89 x 10⁻³ mol of Riboflavin, HL²) in 40 mL of hot methanol. The resulting coloured solution was then refluxed for six hours, after which the product formed on cooling to room temperature. The product was isolated by filtration, washed with methanol and dried over anhydrous CaCl₂. To prepare the complexes, similar procedures were used for the preparation and isolation of the Mn(II), Co(II), Ni(II), Cu(II), and Zn(II) complexes from their acetate, chloride and sulphate salts respectively. A typical equation is shown below;

\[ \text{HL} + \text{HL}^2 + \text{FeSO}_4 \cdot 7\text{H}_2\text{O} \rightarrow [\text{Fe(HL)(HL}^2\text{SO}_4] \cdot \text{H}_2\text{O} + 6\text{H}_2\text{O} \]

Physical measurements

IR (as KBr disc) and electronic (solid reflectance) spectra were recorded using a Perkin-Elmer FT-IR spectrum BX and a Perkin-Elmer λ25 spectrometer respectively. The melting points of the ligands and the complexes were determined using a Gallenkamp melting point apparatus, while percentage metal was determined by complexometric titration using EDTA. An electrochemical analyzer consort C933 was used to determine molar conductance measurements of 1mM solutions in dimethyl sulphoxide. Single temperature magnetic moment susceptibility measurements were carried out using a Sherwood susceptibility balance MSB Mark 1 at a temperature of 30°C.

Antibacterial studies

Laboratory strains of E. coli, S. epidermidis, S. aureus, Enterobacteria sp., Klebsiella sp., Bacillus sp., and Pseudomomas sp. were used for the screening. The antimicrobial experiments were carried out at the Department of Microbiology, University of Ibadan, Ibadan, Nigeria.

Preparation of agar plates/samples

Antibacterial susceptibility tests were performed using the agar diffusion technique. 10 cm² of Muller Hinton’s agar was poured into each of the Petri dishes to set; afterwards their surfaces were uniformly inoculated with 0.3 cm³ of 18 h old test bacteria cultures. This was followed by the addition of 1 mM solution of each compound in 3 cm³ of dimethyl sulphoxide to 6 mm wells bored into the agar. The plates were then left on the bench for 30 minutes to allow diffusion of the compounds into the agar, after which they were incubated at 37°C for 24 hours. The inhibitory zones diameters in mm were taken as an indicator of antimicrobial activities.

Results and Discussion

Analytical data

The metal complexes all showed expected coloration due to d-d and M→L CT transitions. The melting points of the ligands and their metal(II) complexes were distinctively different, that is, Aspirin decomposed at 136°C, Vitamin B₂ was melted at 280°C, and the metal complexes mostly decomposed above or below the melting point and decomposition temperature of their respective parent drugs due to coordination. The percentage metal and experimental values, of the complexes were well within the range of theoretical values (Table 1) and the covalent nature of the metal complexes in dimethyl sulphoxide was confirmed by molar conductance values in the range 19.68 – 46.80 Ω⁻¹cm² mol⁻¹[19].

Table 1: Analytical data of Aspirin, Riboflavin and their metal(II) complexes.

| Complexes | Formula Mass | Color | D.T (°C) | %Yield | %M(Exp) | 1m | μ_m (B.M) |
|-----------|--------------|-------|----------|---------|---------|-----|-----------|
| HL        | 180.16       | White | #136     | -       | -       | -   | -         |
| HL²       | 376.36       | Yellow| 280      | -       | -       | -   | -         |
| [(Mn(HL)(HL²)Cl].4H₂O | 1365.94 | Yellow | 270 | 50 | 8.05(7.46) | 19.68 | 1.42 |
| [Fe(HL)(HL²)SO₄].H₂O | 726.31 | Brown | 275 | 60 | 7.67(7.67) | 26.80 | 3.92 |
| [Co(HL)(HL²)Cl₂].1/2H₂O | 638.38 | Yellow | 265 | 70 | 8.62(8.67) | 46.80 | 4.63 |
| [Ni(HL)(HL²)Cl₂].H₂O | 722.22 | Yellow | 272 | 60 | 8.08(8.13) | 22.70 | 2.06 |
| [(Cu(HL)(HL²)Cl₁] | 1311.94 | Yellow | 290 | 60 | 9.76(9.20) | 19.76 | 1.45 |
| [Zn(HL)(HL²)SO₄] | 717.89 | Yellow | 277 | 70 | 9.10(9.12) | 31.10 | 0.54 |

Keywords: D.T- Decomposition temperature; *Melting point; M- Metal, Exp- Experimental, HL- Aspirin, HL²-Riboflavin, A₁ₗ= Ω cm⁻¹mol⁻¹.

Electronic spectra and magnetic moment measurements

The solid reflectance spectra of the drugs showed two bands at 33.00 kK and 41.49 kK due to π → π* and charge transfer(CT) respectively[19]. The spectrum of the Mn(II) complex indicated 2T₂g → 2A₁g and 2T₂g → 2B₁g transitions at 12.73 kK and 22.22 kK respectively, typical of low spin octahedral geometry. This complex had a moment of 1.42 B.M which was lower than the reported value of 1.73 - 2.2 B.M for low spin octahedral Mn(II) complexes due to antiferromagnetism, operating through a Mn-Mn bond in a dimeric structure [Figure 1].

The Fe(II) complex exhibited transitions of both high spin (T₂g → 3E) and low spin (A₁g → 3T₁g) octahedral geometry at 12.12 kK and 20.83 kK respectively. A moment of 5.0 - 5.6 B.M is usually expected for high spin Fe(II) complexes while low spin octahedral Fe(II) complexes are expected to be diamagnetic. In this study, an observed moment of 3.92 B.M was complementary of equilibrium between high spin and low spin octahedral forms[20].

Similarly, the Ni(II) complex exhibited transitions 1A₁g → 3T₂g and 1A₁g → 3B₁g, typical of spin crossover at 12.77 kK and 18.62 kK respectively. Room temperature magnetic moments in the range 2.8 - 3.3 B.M have been reported for high spin octahedral Ni(II) complexes, while low spin octahedral Ni(II) complexes had moments in the range 0.78 - 1.68 B.M. However, an observed moment of 2.06 B.M for this complex corroborated spin cross over, i.e. equilibrium between the high spin octahedral
Complexes of Mixed Drugs Aspirin and Vitamin B₂

The antibacterial activities of the drugs and its metal complexes against Bacillus sp., Pseudomonas sp., Klebsiella sp., Enterobacteria sp., S. aureus, S. epidermidis and E. coli are presented in Table 3. Riboflavin, Aspirin and all the metal(II) complexes showed no activity against all the bacteria used. Exceptions were [Co(HL)(HL²)Cl₂]Cl₂, [Ni(HL)(HL²)Cl₂]Cl₂, H₂O and [Cu(HL)(HL²)Cl₂]Cl₂, H₂O which showed activities of 12.0 mm, 9.0 mm, 6.0 mm and 11.0 mm against Klebsiella sp., Bacillus sp., S. epidermidis and S. aureus respectively. Generally, metal(II) complexes are expected to be more effective than the metal-free ligand, due to chelation, which reduces the polarity of the metal atom and increases its lipophilic character, thus favoring its permeation through the lipid layers of the bacterial membrane. Hence, the general inactivity of these metal complexes may be attributed to their probable lipophobic nature[12], and the fact that Aspirin and Riboflavin are bacteriostatic and growth supplements respectively[6,7].

Infrared spectroscopy

The IR spectra of the drugs and their metal complexes are recorded in Table 2. The IR spectra of Aspirin and Riboflavin showed bands at 3457 cm⁻¹ and 3441 cm⁻¹ respectively assigned to ν(O-H) stretching vibration[8,9]. This band in Aspirin was the same in the metal complexes, corroborating non coordination of the hydroxyl oxygen in aspirin. Meanwhile, this band shifted in Riboflavin due to the coordination of the hydroxy oxygen atom[10]. The strong bands at 1754 cm⁻¹, 1683 cm⁻¹, 1606 cm⁻¹, 1732 cm⁻¹, 1648 cm⁻¹ and 1622 cm⁻¹, in Aspirin (HL) and Riboflavin (HL²) respectively were assigned as ν(C=O). The former three bands shifted to the range 1732 – 1621 cm⁻¹, corroborating coordination of the carbonyl oxygen atoms of the Aspirin in the metal complexes. However, the latter three bands remained un-shifted in Riboflavin due to non coordination of the carbonyl oxygen atom of Riboflavin[11].

![Figure 1: Proposed structure for the Mn(II){x=4} and Cu(II){x=0} complexes.](Image)

Table 2: Infrared (cm⁻¹) and Electronic spectral data of Aspirin, Vitamin B₂ and their mixed metal.

| Compound        | νN-H/ νO-H | νC=O | νC=N | νM-O | Absorption Band (kK) |
|-----------------|------------|------|------|------|----------------------|
| HL              | 3457       | 1754| 1683| 1606| 51.02 41.49 33.00    |
| HL²             | 3441 3377  | 1732| 1648| 1622| 51.02 46.51 44.05    |
| [{Mn(HL)(HL²)Cl₂}]Cl₂·4H₂O | 3496 3376| 1733| 1649| 1621| 51.02 46.51 44.05    |
| [Fe(HL)(HL²)SO₄] | 3438      | 1732| 1647| 1622| 22.22 12.73          |
| [Co(HL)(HL²)Cl₂]Cl₂·H₂O | 3421      | 1732| 1647| 1581 1546 1505| 572 533 518| 595 502 486 474| 22.22 12.73 |
| [Ni(HL)(HL²)Cl₂]H₂O | 3496 3388| 1732 | 1648| 1621| 595 519 501 486| 18.62 12.77 |
| [{Cu(HL)(HL²)Cl₂}]Cl₂ | 3440      | 1732| 1648| 1622| 595 533 518 501| 48.31 13.70 |
| [Zn(HL)(HL²)SO₄] | 3453      | 1732| 1647| 1580 1547 1502| 571 422| 50.76 46.51 |

Furthermore, the bands at 1581 cm⁻¹, 1548 cm⁻¹ and 1504 cm⁻¹ were assigned to ν(C≡N) in Riboflavin. These bands remained un-shifted in the metal complexes thus confirming non coordination through its imine nitrogen atom[10]. The new bands in the range 422 - 595 cm⁻¹ were assigned to ν(M-O) due to the coordination of the hydroxy oxygen atoms of the Riboflavin, oxygen atoms of the carbonyl in Aspirin and sulphate to metal ions[9,10]. On the contrary, these bands were absent in the spectra of the Riboflavin and Aspirin thus confirming coordination in the metal complexes.

Antibacterial activities

The antibacterial activities of the drugs and its metal complexes against Bacillus sp., Pseudomonas sp., Klebsiella sp., Enterobacteria sp., S. aureus, S. epidermidis and E. coli are
Table 3: Antibacterial activities of Aspirin, Riboflavin and their metal(II) complexes.

| Bacteria    | Complex                  | S. aureus | Enterobacteria sp | Pseudomonas sp | S. epidermidis | Klebsiella sp | Bacillus sp | E. coli |
|------------|--------------------------|-----------|-------------------|---------------|----------------|--------------|------------|---------|
| Streptomycin | HL                        | 30        | 20                | 25            | 20             | 32           | 25         | 13      |
|             | HL^2                     | R         | R                 | R             | R              | R            | R          | R       |
|             | [{(Mn(HL)(HL^2))}_2Cl_2]H_2O | R         | R                 | R             | R              | R            | R          | R       |
|             | [Fe(HL)(HL^2)SO_4]        | R         | R                 | R             | R              | R            | R          | R       |
|             | [{Co(HL)(HL^2)Cl}_2]H_2O  | R         | R                 | 12            | 9              | R            | R          | R       |
|             | [{Ni(HL)(HL^2)Cl}_2]H_2O  | 6         | R                 | R             | R              | R            | R          | R       |
|             | [{Cu(HL)(HL^2)Cl}_2]H_2O  | R         | R                 | R             | R              | R            | R          | R       |

HL- Aspirin; HL^2–Riboflavin; R-Resistance

Conclusion

Some Mn(II), Fe(II), Co(II), Ni(II), Cu(II) and Zn(II) complexes of the mixed drugs, Aspirin and Riboflavin (Vitamin B3) were synthesized and characterized. Room temperature magnetic moment measurements indicated that the Mn(II) and Cu(II) complexes were antiferromagnetic, while Fe(II) and Ni(II) complexes exhibited spin crossover. In addition the metal complexes assumed a probable 6–coordinate octahedral geometry and were non-electrolytes in DMSO. The in-vitro antibacterial activities studies showed that most of the metal(II) complexes and the drugs had no activity against Bacillus sp, E. coli, Pseudomonas sp, Klebsiella sp, Staphylococcus spp and Enterobacteria sp.

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