Investigation of charge transfer complex formation between Fe(III) and 2,6-dihydroxy benzoic acid and its applications for spectrophotometric determination of iron in aqueous media

Mohammad Saeid Hosseini*, Sara Madarshahian
Department of Chemistry, Faculty of Science, Birjand University, Birjand, P.O. Box 414, (IRAN)
E-mail: mshosseini1336@yahoo.com; mshosseinibojd@birjand.ac.ir
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ABSTRACT
On the basis of affinity of Fe(III) ions toward functional groups of 2,6-dihydroxy benzoic acid (2,6-DHB), a charge transfer complex (CT) is formed between Fe(III) ion and 2,6-DHB in acidic media. The stoichiometry and stability constant of the complex was determined using both mole ratio and Job’s methods. It was established a 1:1 complex was formed and the stability constant obtained from both the method, mole ratio: 2.55±0.14×10^5 and Job: 2.70±0.12×10^5 is in agreement with together. This method was used for spectrophotometric determination of Fe(III) in aqueous media. The dynamic range and detection limit (DL) were 1.2×10^-5 to 3.6×10^-4 M and 1.5×10^-6 M, respectively. The relative standard deviation (RSD) was found to be 0.76% in the replicate treatments (n=7) with 1.6×10^-4 M Fe(III). The accuracy of the method was confirmed using a certified reference material (CRM) and it was applied satisfactorily for determination of iron in a multivitamin-multimineral tablet and various water samples.

KEYWORDS
Charge transfer complex; 2,6-Dihydroxybenzoic acid; Fe(III); Spectrophotometry; Mole ratio; Job’s method.

INTRODUCTION
Owing to simplicity, lesser expensive instrumentation and provides high versatility UV–Visible spectrophotometric methods have been developed for the determination of metal ions. It involves using a number of chromogenic reagents for this purpose. Ferric ion is the most stable state of iron in the solutions, hence the determination of iron is generally demonstrated on Fe(III) contents. In recent years, numerous chromogenic reagents, which capable to form high stable complexes with Fe(III) ion, have been widely used for the determination of iron contents in numerous complicated matrices. Among the various reagents that was used, it was found the reagents that capable to form charge transfer (CT) complexes with Fe(III), exhibit more sensitivity for such determination, specially in treatment with the trace amount of Fe(III) ions.

Apparently, presence of Fe(III) in the environmental samples is contributed to construction of high stable complexes with the natural products, such as humic compounds. In order to characterize the structure of such complexes, a number of studies performed on the complexes constructed from Fe(III) and simple humic models, both in solution and in the solid state. In view of this considerations, 2,6-Dihydroxy benzoic acid (2,6-DHBA) is one of the most interesting reagent because of presence two hydroxyl groups adjacent to the carboxyl group in its molecular structure. It is easily soluble in water, in spite of presence of two...
strong intra molecular hydrogen bonds in its structure\textsuperscript{11-13}. It was found from the NMR investigation that there is a fast (in NMR scale) degenerate equilibrium between the two tautomer structures involving the double proton transfer. This phenomenon causes to form stable complexes between 2,6-DHB and some metal ions\textsuperscript{14-24}. Among them, the CT complexes exhibit more stability and absorptivity. In this work, the CT complex of Fe(III)-2,6-DHB was synthesized and characterized by spectrophotometric methods. At the following, it was applied for spectrophotometric determination of trace amount of Fe(III) ions in aqueous solutions.

**EXPERIMENTAL**

**Instrumentation and material**

A Shimadzu model 160A spectrophotometer was used for the absorbance measurements with one pair of 10-mm quartz cell. A Corning model 125 pH meter was used for the pH adjustment. For plotting the curves and obtaining their equations, Grapher version 1.09 program was employed.

All solutions were prepared using deionized and double distilled water. An approximately 0.1 M solutions of 2,6-DHBA, obtained from Fluka (Switzerland), and Fe(NO\textsubscript{3})\textsubscript{3}, obtained from Merck (Germany), were prepared and standardized against KOH Titrisol and Titriplex\textsuperscript{®} III solutions, respectively. The working solutions were made by suitable dilution of these standard solutions. Both the standard and working solutions were stored in PTFE bottles. The buffer solution required to adjust the pH of the solutions were prepared by mixing 1 M formic acid/sodium formate and acetic acid/sodium acetate (all the reagents obtained from Merck) for the pH range of 2.5-3.5 and 3.5-5.5, respectively. The other reagents were of analytical grade, obtained from Merck, and were used without further purification.

**Spectrophotometric determination of Fe(III) ions**

To a solution containing up to 0.5mg of Fe(III) ions, 0.5mL of 0.05 M 2,6-DHB solution was added and its pH was adjusted to 3.00±0.05 using formic acid/sodium formate solutions. The solution was diluted to the mark using a 25-mL volumetric flask. It was shaken for a few minutes and finally the absorbance was measured at 560 nm against a reagent blank prepared by the same manner.

**RESULTS AND DISCUSSION**

**Characterization of the complex formation**

The proposed method involves formation a soluble bluish CT complex. As shown in figure 1, the product exhibits a maximum absorption in the visible region on 560nm against the reagent blank, whereas both the reactants show negligible absorption at this wavelength interval, solitarily. Consequently, the absorbance is directly related to the complex construction, which is made greater by increasing both the concentration and pH of the solution. Obviously, there is a risk of precipitate the Fe(III) content as hydroxide, when the pH is so increased. As it is shown in figure 1, this phenomenon causes to decrease the absorbance at the pHs more than 3. Hence, the future examinations were carried out at the buffering pH of 3±0.05 as the optimum pH for such determinations.

The effect of 2,6-DHB concentration on the complex formation was examined in the range of 0 -6×10\textsuperscript{-4}M using the solutions in which the concentrations of Fe(III) were fixed on 1.2×10\textsuperscript{-4}M. As shown in figure 2, the absorbance measurements established that the requested ligand concentration to complete the complex formation should be at least three times of Fe(III) concentration. Furthermore, it should be considered that the reaction between Fe(III) and 2,6-DHB tends to complete more rapidly at the presence of excess amount of the ligand.

In order to investigate the effect of non aqueous
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medium on formation of the complex, a series of mixed methanol-water solutions containing 20%, 40% and 60% methanol were prepared in which the concentrations of Fe(III) and 2,6-DHB were fixed on 1.2×10^{-4}M and 3.6×10^{-4}M, respectively. It was found that the absorbance values from 0.203 respects to aqueous solution were increased up to 0.245 for 60% methanol. With regards to this observation, occurrence of the charge transfer phenomenon, which is enhanced by decreasing the polarity of the solvent, was confirmed.

**Determination of the complex formation by mole ratio method**

In order to calculate formation constant of the complex by using the mole ratio method\(^{[25]}\), the solutions that were prepared to investigate the effect of 2,6-DHB concentration on the complex formation was used for this purpose. As it is shown in figure 2, the break point arising from the extrapolated tangent lines of the curve confirms that 2,6-DHB/Fe(III) ratio in the CT complex is 1:1. By considering to the equation curve fitted to the spectrophotometric data, the absorbance value at 1:1 ratio in which the CT complex is in equilibrium with the metal/ligand constituents, is directly related to the complex concentration. By using the molar absorbivity of the complex that was obtained from the linear calibration curve, the complex concentration is calculated and consequently, the metal and ligand concentrations were determined by subtracting from the initial ligand quantity (1.2×10^{-4}M). In this manner, the formation constant was found to be 2.55±0.14×10^{5}.

**Determination of the complex formation by Job’s method**

In order to confirm the determination of the complex formation constant obtained from the mole ratio method, the calculation was repeated by using the Job’s method\(^{[26]}\). Subsequently, a series of 25-mL solutions (n=21) were prepared in which the Fe(III) and 2,6-DHB contents were just inversely varied over the concentration range of 0 to 4.0×10^{-4}M so that the total mole fractions were fixed to 1. The pH of the solutions was adjusted to 3.00±0.05 by addition of adequate quantities of formic acid/sodium formate solutions. The solutions were then transferred to 25-mL volumetric flasks and diluted to the mark. After shaking them for a few minutes, the absorbance values were measured at 560nm against the buffering blank containing formic acid/sodium formate solutions. Figure 3 represents variation of absorbance (A) vs. mole fraction of 2,6-DHB reagent. As it is shown, the maximum absorbance was obtained within the range of 0.45-0.50, which denotes that Fe(III)/2,6-DHB ratio in the CT complex is 1:1. According to the discussion present in the literature\(^{[27]}\), a little deviation observed between the experimental and theoretical values is predictable, which is attributed to...
the dissociation phenomenon of 2,6-DHB. Curve 1 in figure 3, represents extrapolation of the linear sections of curve 2. By intersecting the extrapolated lines equations, the coordination of the break point was obtained as $A_1=0.354$ and $X_c=0.50$. This break point locates situation in which dissociation of the complex is negligible. The normal maximum absorbance with coordination of $A_1=0.309 \pm 0.001$ and $X_c=0.50$, was considered as the maximum point of curve 2. Essentially, the $A_1$ and $A_2$ should be obtained at an equal mole fraction value of 2,6-DHB. Hence, the mole fraction respected to $A_1$ was inserted in equation of curve 2 and the resulted absorbance was considered as $A_2$ value. The complex formation constant, $K_r$, was then calculated from the following equation:

$$K_r = \frac{A_2/A_1}{C_{M}(1-A_2/A_1)^2}$$

(1)

where $C_M$ is the initial analytical concentration of Fe(III) at the maximum points of $A_1$ and $A_2$. In this manner, the $K_r$ was obtained as $2.70 \pm 0.12 \times 10^5$. No significance difference was observed using the F and t tests at 95% confidence level between the two values obtained from both the above techniques.

**Analytical figures of merit**

Under the optimum conditions, a linear calibration curve was constructed with equation of $A=1966.7C +0.0053$, ($r^2=0.9987$), for the determination of Fe(III) over the concentration range of $1.2 \times 10^{-5}$ to $3.6 \times 10^{-4}$ M (0.28-20 μg mL$^{-1}$). The RSD obtained in the replicate treatments (n=7) with $1.6 \times 10^{-4}$M Fe(III) was found to be 0.76%. In treatment with the blank solutions, the DL was obtained as 1.5×10$^{-6}$M.

**Effect of foreign ions**

The effects of several foreign ions were studied by introducing each one to 25-mL aliquots of the solutions containing Fe(II) with the concentration of $5.0 \times 10^{-8}$M. Each ion was considered as interfering agent, when the absorbance value exhibited a deviation more than ±5%. It was found that 200-fold excess of the alkaline and alkaline earth metal ions as chloride or nitrate compounds did not interfere in the determination of Fe(III) content. As shown in TABLE 1, some of the other metal ions were tolerated at moderately high foreign ion/Fe(III) ratios.

**Application**

In order to test accuracy of the method, a labora-

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**TABLE 1**: Tolerance ratio of the foreign ions on the determination of $5 \times 10^{-4}$ M of Fe(III) in 25ml of the aqueous phase

| Mole ratio of interfering ion to Ag(I) | Ions |
|--------------------------------------|------|
| 100                                  | Zn(II), Co(II), La(III), Cd(II), Cu(II), Ca(II), Hg(II), As(III), Pb(II), Ce(III), Ni(II) |
| 50                                   | Ag(I), Ce(III) |
| 10                                    | Mn(II), Al(III), Cr(III), V(III), Saragasso |

**TABLE 2**: Analytical values for NIES certified reference material no. 9 “Sargasso”

| Element | Content (Wt.%) | Trac contents |
|---------|----------------|--------------|
| Calcium | 1.34±0.05      | Lead         |
| Sodium  | 1.70±0.08      | Vanadium     |
| Potassium| 6.10±0.20     | Silver       |
| Strontium| 0.100±0.003   | Cadmium      |
| Magnesium| 0.65±0.03     | Cobalt       |

**TABLE 3**: Comparison of the result obtained from this work with that of FAAS for the determination of Fe(III) in various environmental samples. The results are reported as the average values from five sample measurements with the related standard deviations

| Samples     | Spiked (μg) | Found (μg) | FAAS | Crit./Exp. | Crit./Exp. |
|-------------|-------------|------------|------|------------|------------|
| Multi.-multi. tablet | 56          | 174±1.7    | 187±2.3 | 6.39/1.83 | 3.31/2.02 |
| Tap water   | 56          | 58.3±0.6   | 57.2±0.4 | 6.39/2.25 | 3.31/0.5  |
| Well water  | 56          | 59.7±0.8   | 58.4±0.4 | 6.39/4.00 | 3.31/0.6  |
| Waste water | 28          | 21.3±0.4   | 22.8±0.6 | 6.39/2.25 | 3.31/0.5  |

a: A multivitamin-multimineral tablet formula: VA, 5000U; VD, 400 U; VE, 30U; VBI, 1.5 mg; VB2, 1.5mg; VB6, 2mg; VC, 60mg; VB12, 6μg; VK1, 25 μg; biotin, 30μg; folic acid, 400μg; niacinamide, 20 mg; pantothenic acid, 10mg; Ca, 162 mg; P, 125mg; K, 40mg; Cl, 36.3 mg; Fe, 18mg; Cu, 2mg; Zn, 15mg; Mn, 25.8mg; I, 150μg; Mo, 25μg; Se, 25μg; Ni, 5μg; Si, 10μg; Sn, 10μg; V, 10μg.

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A multi. multimineral tablet formula: VA, 5000 U; VD, 400 U; VE, 30 U; VBI, 1.5 mg; VB2, 1.5 mg; VB6, 2 mg; VC, 60 mg; VB12, 6 μg; VK1, 25 μg; biotin, 30 μg; folic acid, 400 μg; niacinamide, 20 mg; pantothenic acid, 10 mg; Ca, 162 mg; P, 125 mg; K, 40 mg; Cl, 36.3 mg; Fe, 18 mg; Cu, 2 mg; Zn, 15 mg; Mn, 25.8 mg; I, 150 μg; Mo, 25 μg; Se, 25 μg; Ni, 5 μg; Si, 10 μg; Sn, 10 μg; V, 10 μg.
applied to the determination process. In treatment with the water sample, they were initially filtered through a membrane filter with pore size of 0.45μm. To confirm the accuracy of the result obtained by this method, the measurements were also repeated by a well-known method, such as FAAS\(^{[29]}\). The results for the determination and recovery of Fe(III) contents are shown in TABLE 3. As it is shown, the results obtained from both the methods were agreed with together according to F and t tests. These results are denoted with the applicability of this method for properly determination of Fe(III) in various water samples and similar sample solutions.

CONCLUSION

A new type of complex as a CT complex of Fe(III)-2,6-DHP was developed for determination of iron with spectrophotometric method. The Fe(III)-2,6-DHP complex benefits from a moderately high formation constant in acidic media. The proposed method provides an appropriate selectivity and sensitivity for easy determination of Fe(III) ions. One of the advantages of this method is directly measuring of Fe(III) in various aqueous media, such as pharmaceutical samples. A number of metal ions, which may be found either in water or pharmaceutical samples, did not interfere in the determination. The RSD and DL of the proposed method were respectively 0.76% and 1.5×10^{-6}M, which are less than the respected values for iron determination by traditional direct spectrophotometric method, such as Fe(II)-1,10-phenanthroline. Furthermore, the method benefits form a widespread dynamic range in comparison with the above method and FAAS.

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REFERENCES

[1] B.Stearns, K.A.Losee; J.Bernstein, Med.Pharm. Chem., 6, 201 (1963).
[2] B.Morelli; Analyst, 108, 870 (1983).
[3] N.Meddoure, T.Douadi, S.Chafaa, M.Khan, G.Bouet; Comptes Rendus Chimie, 7, 1113 (2004).
[4] L.Vladescu, R.Lerch-Gurguta; Talanta, 40, 1127 (1993).
[5] M.C.Chattopadhyaya; J.Inorg.Nucl.Chem., 38, 569 (1976).
[6] M.S. Abu-Bakr, H.Sedaira, E.Y.Hashem; Talanta, 41, 1669 (1994).
[7] R.C.Hider, D.Bickar, I.E.G.Morrison, J.Silver; J.Am.Chem.Soc., 106, 6981 (1984).
[8] A.C.Dash, G.M.Harris; Inorg.Chem., 21, 2336 (1982).
[9] R.B.Jordan; Inorg.Chem., 22, 4160 (1983).
[10] J.Xu, R.B.Jordan; Inorg.Chem., 27, 1502 (1988).
[11] N.S.Goluber, G.S.Denisov; J.Md.Struct., 270, 263 (1992).
[12] N.S.Goluber, G.S.Denisov, L.A.Kuzina S.N. Smirnov; J.Gen.Chem.(Rass), 64, 1162 (1994).
[13] G.S.Denisov, N.S.Goluber, V.M.Schreiber, S.S. Shajakhmedov A.V.Shurukhina; J.Mol.Struct., 381, 73 (1996).
[14] S.F.M.Ali, V.R.Rao; J.Inorg.Nucl.Chem., 37, 1041 (1975).
[15] W.Brzyska, A.Kula, Z.Rzacynska, M.Jarioniec; Pol.J.Chem., 72, 2087 (1998).
[16] W.Brzyska, A.Kula, Z.Rzacynska, M.Jarioniec; Pol.J.Chem., 72, 2524 (1998).
[17] T.Glowiak, W.Brzyska, A.Kula, Z.Rzacynska, M.Jarioniec; JCOORD.Chem., 48, 477 (1999).
[18] W.P.Griffith, H.I.S.Nogueira, B.C.Parkin, R.N. Sheppard, J.P.White, D.J.Williams; J.Chem.Soc., Dalton Trans., 1775 (1995).
[19] G.Bandoli, A.Dolmella, T.I.A.Gerber, J.Perils, J.GH.Preez; Inorg.Chim.Acta, 294, 114 (1999).
[20] F.Carita, L.erre, G.emicra, A.Panzanelli; Inorg. Chim.Acta, 80, 57 (1983).
[21] G.Smith, C.H.L.Kennard, T.C.W.Z.Mak; Kristallogr., 184, 275 (1988).
[22] P.C.R.Soares-Santos, H.I.S.Nogueira, F.A.Almeida Paz, R.A.Sa Ferreira, L.D.Carlos, J.Klinowski, T. Trindade; Eur.J.Inorg.Chem., 3609 (2003).
[23] F.Carititi, E.Strinna, G.Micera, A.Panzanelli, G.Cianni; Inorg.Chim.Acta, 80, 57, (1983).
[24] T.C.W.Mak, G.Smith C.H.L.Z.Kennard; Kristallogr., 270, 33 (1993).
[25] A.E.Harvey, D.L.Manning; J.Am.Chem.Soc., 72, 4488 (1950).
[26] W.Likussar, D.F.Boltz; Anal.Chem., 43, 1265 (1971).
[27] A.R.Das, V.S.K.Nair; J.Inorg.Nucl.Chem., 34, 1271 (1972).
[28] M.J.Benolie, P.Quevauviller; Analyst, 123, 977 (1998).
[29] ‘Standard Methods for Examination of Water and Wastewater’, 16th ed., APHA-AWWA.WPCF, Washington, D.C. (1985).