

# STUDY OF COMPLEX FORMATION BETWEEN COPPER (II) METAL ION AND DRUG TAMSULOSIN WITH BIOLOGICAL IMPORTANT AMINO ACID LIKE PHENYLALANINE AND GLUTAMIC ACID IN 80% ETHANOL –WATER MIXTURE P<sup>H</sup> METRICALLY.

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**Abstract :** The stability constant of complexes of Copper (II) ion with new Drug Tamsulosin ( FLOMAX) as Primary ligand and biological molecules such as Phenyl alanine and glutamic acid as a secondary ligands have been determined P<sup>H</sup> metrically in 80 % (v/v) ethanol-water medium at 27°C and fixed ionic strength 0.1M NaClO<sub>4</sub> by computational program SCOG

**Key Words** - Tamsulosin, ΔlogK, Copper metal ion, Ternary Complexes.

## I. INTRODUCTION:

The metal ligand complexes are very important as per as the biological phenomenon's are concerned, every biological system involves the complex mechanism consisting of different components like metal ions and biological important amino acids etc. and hence any drug delivered in the biological system is expected to form the complexes with already present moieties to form the chelates thus study of complexation of the drug with metal ion is purposeful. The present study aims understanding the nature of drug complex by studying the equilibriums leading to determination of stability constant values of ternary complexes formed between the transition metal ion Cu<sup>2+</sup>, Drug Tamsulosin and the Amino acids Phenyl alanine and the glutamic acid.

The Drug Tamsulosin is available in market by name Flomax used for the treatment of symptomatic Benign Prostatic Hyperplasia, helps with Passage of Kidney stone <sup>[1], [2]</sup> and for the Urinary retentions also used for the treatment of acute urinary retention. Though the drug preliminarily used to treat BHP however appears to be useful for the kidney stone of the order 4 mm to 10 mm size <sup>[3]</sup>. The structure of the Drug Tamsulosin is illustrated in fig.1

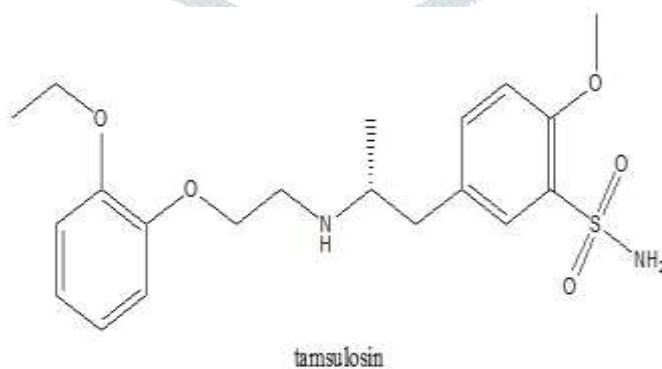


Fig. 1 Chemical Structure of Ligand Tamsulosin

This drug is alpha blocker and bladder neck muscle fibers are relaxed by its use and make prostate easier to urinate. Tamsulosin is used to treat men who have symptoms of an enlarged prostate gland, which is also known as benign enlargement of the prostate

(Benign Prostatic Hyperplasia or BPH). Benign enlargement of the prostate is a problem that can occur in men as they get older. The prostate gland is located below the bladder. As the prostate gland becomes bigger some muscles in that gland may become hard and get in the way of the tube that take urine from the bladder, which can cause problems in urinating, such as a need to urinate often, a weak stream when urinating, or a feeling of not being able to empty the bladder completely. as It is an Alpha adrenergic receptor antagonist. Therefore it is clear that this drug excretes to level good and thus it may become possible along with that it may take out certain excess amount of the metals from the body<sup>[4]</sup>.

The metal ions are integral parts of enzymes and play an important role in the biological system, such as to trigger a reaction, control reaction mechanism, stabilize protein structure, maintain structure of cell walls etc. Latest information indicates regulation of metabolism and growth of animal cell is dependent upon the mobilization of divalent and trivalent metal ions.

Copper is a transition metal ion are integral parts of enzymes and play an important role in the biological system, such as to trigger a reaction, control reaction mechanism, stabilize protein structure, maintain structure of cell walls etc. Latest information indicates regulation of metabolism and growth of animal cell is dependent upon the mobilization of divalent and trivalent metal ions. It is widely distributed throughout the body.<sup>[5]</sup> The identification of mammalian homologues of these proteins reveals a remarkable structural and functional conservation of copper metabolism between bacteria, yeast and humans. Furthermore, studies on the function and localization of the products of the Menkes and Wilson's disease genes, which are defective in patients afflicted with these diseases, have provided valuable insight into the mechanisms of copper balance and their role in maintaining appropriate copper distribution in mammals<sup>[6]</sup>

Copper (Cu) is an essential trace element required for survival by all organisms from bacterial cells to humans. All amino acids are polymer and regarded as building block of protein. Some amino acids are studied in this research.<sup>[7]</sup>

## II. MATERIALS AND METHODS:

Present investigation deals with the P<sup>H</sup> metric studies on copper (II) metal complexes with Benign Prostatic Hyperplasia Tamsulosin and amino acids Phenyl alanine and Glutamic acid in 80% (v/v) ethanol-water medium at 27°C and fixed ionic strength of 0.1 M NaClO<sub>4</sub>.

The nitrates of copper, of A.R. grade were obtained from Doodle (India). Metal ion was used in the form of their perchlorates to avoid the possibility of complex formation with anions. The perchlorates were prepared from the corresponding nitrates.<sup>[8]</sup> The concentration of metal ions was estimated by the standard procedures.<sup>[9-11]</sup> Sodium perchlorate (Merck) was dissolved in carbon dioxide free distilled water.

The solution of sodium hydroxide was also prepared in carbonate free distilled water by allowing the solution to stand for a long time till any carbonate if present precipitated. The solution was used as titrant for the potentiometric titration. As a routine, the solution was standardized at least once every day by titrating with standard oxalic acid solution. Per chloric acid of Reidal (Germany) was used for the preparation of the stock solution. Its exact normality was obtained by titrating it conductometrically using standard sodium hydroxide solution. Amino acids from Merck (Germany) and Fluka (Germany) were prepared by dissolving A.R. grade sample in 80% (v/v) ethanol – water medium.

Solution of the Drug Tamsulosin was prepared by dissolving sample as received in 80% (v/v) ethanol-water medium. Drugs samples in pure form were obtained from pharmacy industries.

The Methodology were used in the study of ternary metal complexes by the potentiometric titration technique, involves the titrations of carbonate free solution of against standard sodium hydroxide, where drug Tamsulosin (D) and amino acids (R) are the ligands.

The ionic strength of the solutions was maintained constant i.e. 0.1 M by adding appropriate amount of 1M sodium perchlorate solution. The titrations were carried out at 27°C in an inert atmosphere by bubbling oxygen free nitrogen gas through an assembly containing the electrode to expel out CO<sub>2</sub>. The experimental procedure, in the study of ternary metal complexes by the potentiometric titration technique, involves the titration of carbonate free solution of in 80 % ( v/v) ethanol-water, were corrected by method of Vansittart and Hass. The formation constant of ternary complexes were determined by computational programmed SCOGS to minimize the standard derivation.

The systems taken for the titration are set as follows:

I	Free HClO <sub>4</sub> ( A )
II	Free HClO <sub>4</sub> ( A ) + Tamsulosin ( D )
III	Free HClO <sub>4</sub> ( A ) + Tamsulosin ( D ) + Copper ion ( M )
IV	Free HClO <sub>4</sub> ( A ) + Amino acids ( R )
V	Free HClO <sub>4</sub> ( A ) + Amino acids ( R ) + Copper ion ( M )
VI	Free HClO <sub>4</sub> ( A ) + Tamsulosin ( D ) + Amino acids ( R ) + Copper ion ( M )

### III. RESULT AND DISCUSSIONS:

#### 3.1 Binary metal complexes

The  $P^K$  and the  $\log K$  values of the amino acids are important for the determination of the Stability constant of the studied ternary complexes and hence are taken as it is investigated [12-18]

**Table No. 1.** The proton ligand constant and metal ligand stability constant of drug Tamsulosin and amino acids with Copper (II) determined in 80 % (v/v) ethanol-water mixture at 27°C and ionic strength  $\mu = 0.1$  M  $\text{NaClO}_4$  are shown in the

Ligands	$pK_1$	$pK_2$	Copper	
			$\text{Log}K_1$	$\text{Log}K_2$
Tamsulosin	2.5444	5.9636	6.6043	-
Phenyl alanine	13.14	9.300	8.990	7.670
Glutamic acid	3.136	5.899	1098	8.640

The  $pK$  and  $\log K$  value of drug here is important for the explanation of stability constant of Metal ligand ternary complexes. The figure 2 and the figure3 illustrates the scheme of complex formation between the Metal ion and the Primary Ligand

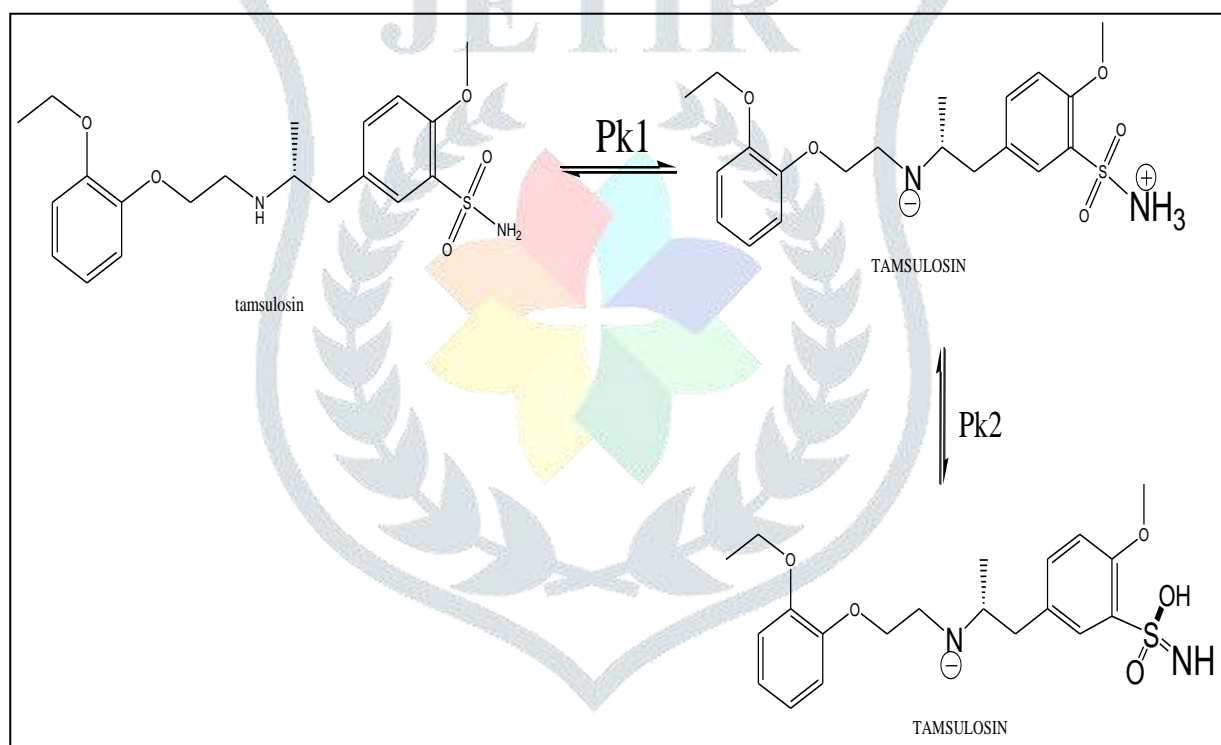


Fig. 2. Proton Dissociation Scheme for free Ligand Tamsulosin

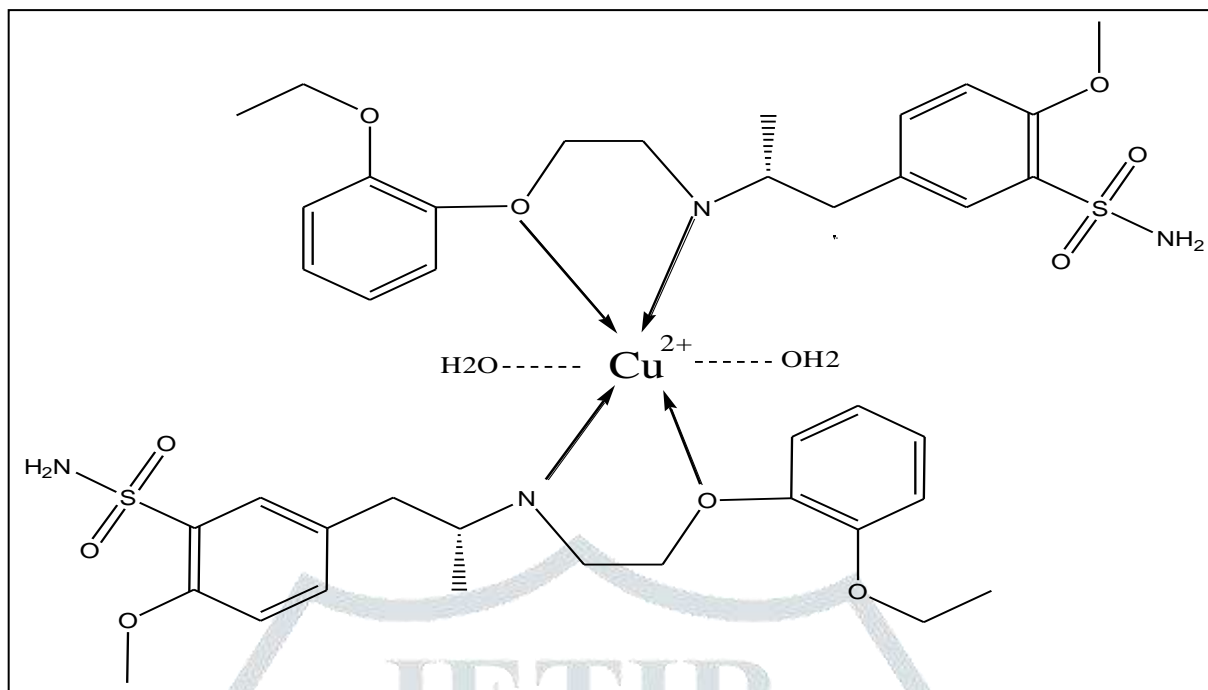


Fig.3. Proposed Structure of Cu (II) Tamsulosin Complex:

### 3.2 Ternary Metal Complexes:

The potentiometric titration, ternary systems shows that the mixed ligand curve coincide with A+D complex curve up to the pH ~ 2.8 and after this pH, it deviates. Theoretical composite curve remains toward left of the mixed ligand complex curve. After pH ~ 2.9, the mixed ligand curve drifts towards X-axis, indicating the formation of hydroxide species. Since the mixed ligand curve coincide with individual metal complex titration curves, the formation of 1:1:1 complex by involving stepwise equilibrium.

The primary ligand Tamsulosin (D) form 1:1 and secondary ligand (R) form 1:1 and 1:2 complexes with Cu (II). It is evident from the figure of percentage concentration species of Cu (II) - Tamsulosin - Phenyl alanine and glutamic acid systems that the percentage distribution curves of free metal decreases sharply with increasing pH, indicates involvement of metal ion in the complex formation process. Percentage concentration of free ligands Tamsulosin and glycine increases and this increase may be due to the dissociation of ligand present in the system, as a function of pH.

### 3.3 Species distribution studies:

To explain the equilibrium and evaluate the calculated stability constant of ternary complexes Cu (II) - Tamsulosin - phenyl alanine and Cu (II) - Tamsulosin - glutamic acid, species distribution curves have been plotted as a function of pH at temperature 27°C and  $\mu = 0.1$  M NaClO<sub>4</sub> by using SCOG programmed.

It can be seen that, the concentration of Cu (II) - Tamsulosin-glycine increases from pH~2.6 whereas the concentration for the formation of D (Tamsulosin) and HR (Phenyl alanine) show continuous decrease with increasing pH which indicates the formation of Cu(II)-Tamsulosin - Phenyl alanine . The concentration of DRH species continuously increases; confirm the formation of ternary complexes.

From the SCOG distribution curve it is concluded that the formation of ternary complex started only after the metal primary ligand complex has attained its maximum concentration. This indicate that metal primary ligand complex Cu (II)- Tamsulosin is formed first then the secondary ligands such as Phenyl alanine and glutamic acid coordinated to it, resulting the formation of ternary complex

### 3.4 The Stability Constants of Ternary Complexes.

The relative stabilities of the binary and ternary complexes are quantitatively expressed in term of  $\beta_{11}$ ,  $\beta_{20}$ ,  $\beta_{02}$ ,  $K_D$ ,  $K_R$ ,  $K_T$  and  $\Delta \log K$  value which are represented in table II.

For the system ligand which form both 1: 1 and 1:2 binary complexes. The magnitude of the constant is the measure of stability of mixed ligand complexes. Water and  $K_a$  calculated statistically expected value 0.6 log units by considering with probabilities for a variety of reason discussed by Sigel.  $\Delta \log K$  value can be calculated by using first or second approach. The calculated  $\Delta \log K$  values for all systems are given in table II.

The Comparison of  $\beta_{11}$  with  $\beta_{20}$  and  $\beta_{02}$  of this system show that preferential formation of ternary complexes over binary complex of primary as well as secondary ligand. The considerably positive value of  $K_D$  &  $K_R$  indicates high stability of ternary

complexes with respect to that of primary as well as secondary ligands. The  $K_f$  value of this complex is positive but the magnitude is smaller which indicates lower stability of ternary complexes.

Result of the present investigations show that the stability constant of ternary complexes formed are less stable. The negative  $\Delta \log K$  value of this system in case of ternary system of glutamic acid indicates that the ternary complex is less stable than the binary 1:1 copper -Tamsulosin & metal-amino acids complex. This is in accordance with statistical considerations. The negative value of  $\Delta \log K$  does not mean that the complex is not formed. The negative value may be due to the higher stability of its binary complexes, reduced number of coordination sites, steric hindrance. [19-22] Electronic consideration [23-24] difference in bond type, geometrical structure etc.

**Table No. II.**

Parameters based on some relationship between the formation of ternary complexes of Copper (II) metal ion with Tamsulosin in the presence of amino acids (1:1:1) system at temp = 27°C I = 0.1 M NaClO<sub>4</sub> Medium = 80% (V/V) Ethanol-Water.

Amino acids	$\beta_{11}$	B <sub>20</sub>	B <sub>02</sub>	K <sub>D</sub>	K <sub>R</sub>	K <sub>F</sub>	$\Delta \log K$
Phenyl alanine	16.4443	6.6043	16.660	9.8400	7.4543	2.4577	0.8500
Glutamic acid	17.5800	6.6043	19.6200	10.975	6.600	2.3851	-0.0043

#### IV CONCLUSION:

The  $\Delta \log K$  value of this system is higher than the statistically expected value showing the stabilized nature of the ternary complex. The primary ligand Tamsulosin having smaller size.

Therefore its  $\Delta \log K$  value is less negative. Negative  $\log K$  value of ternary complexes is due to the electrostatic repulsion between the negative charges on Tamsulosin & amino acids. Steric hindrance consideration is the most important factor because in the present studies of ternary complex, primary ligand Tamsulosin coordinates with the metal ion in the lower pH range and form 1:1 complex. In solution, ternary complex forms as the titration curve run below the Cu (II)-Tamsulosin titration curve. So, it is evident that the entry of the secondary ligand amino acids faces steric hindrance due to bigger size of the Cu (II)-Tamsulosin complex as compared to aqua ion, which tries to restrict the entry of the secondary ligand in the coordination sphere of the Cu (II) metal ion & thus reduces the stability of ternary complexes. The order of stability of ternary complexes of Cu (II) with respect of secondary ligand for respective primary ligands is Tamsulosin = phenyl alanine > glutamic acid

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