Exploiting the diazotization reaction of 4- minoacetophenone for Methyldopa determination.

Jwan A.AbdulSattar*

Received 8, January, 2013 Accepted 4, March, 2013

Abstract:

Based on the diazotization reaction of 4-aminoacetophenone with sodium nitrite in acid medium to form diazonium salt, which was coupled with Methyldopa to form a violet reddish soluble azo dye with maximum absorbance at 560 nm,a batch procedure had been developed for the estamination of Methyldopa. Under optimum experimental parameters affecting on the development and stability of the colored product, Beer's law obeyed in the range $(0.5-45) \mu g.ml^{-1}$ with a correlation coefficient (0.9979).The proposed method was successfully applied to the determination of Methyldopa in either pure form and in commercial brands of pharmaceuticals, no interference was observed from common excipients in the formulations. The analytical results obtained by applying this method were in good agreement with labeled values.

Key words: Methyldopa, determination, diazotization, pharmaceutical preparation.

Introduction:

Due to the magnitude use of Methyldopa (MTD). chemically known α -methyl-3,4-dihydroxy as alanine which is phenyl а catecholamine derivative widely used as an antihypertensive agent. The Methyldopa is a centrally acting α -2adrenoreceptor agonist, which reduces sympathetic tone and produces a fall in blood pressure [1].Besides, it is necessary for routine quality control in the analysis of produced medicines, thus there is an important demand for rapid and simple methods for the determination of methyldopa in pharmaceutical preparations. The official method reported in USP[2]describes non aqueous a the titration for assay of MTD, although this later one is kind of simple and being used in routine laboratories, it is analysis time consuming and very tedious. Several researches have been devoted to the

development of new high performance alternative procedures for determinationof Methyldopa in medicines and/or biological specimens using different techniques Electrochemical [3-6], such as Chromatography[7-10], Nanotechnology [11].How-ever these methods either requires sophisticated equitement:or involves procedures rigorous of the with control experimental conditions.

Molecular absorption spectrophotometry is by far the instrumental technique of choice in industrial laboratories. Owning mainly to its simplicity, often demanding low cost equipment and lending itself to easy automation of trace analysis procedures, therefore a various number of UV-Visible spectrophotophotometric methods for

Methyldopa determination have been reported which involves the

use of diverse chromogenic reagents such as: O-chloranil(O-CIN), Chloranilic acid (CIA) and Dichlorodicyanobenzoqu-inone (DDQ) [12], Vanadium[13],2,6-dichloroquinone -4chlorimide(DCQ[14],p-Chloranil[15]

tris(1,10phenanthroline[16],Molybdate [17,18],Salicylic acid[19].4-amino benzoic acid[20]and 3-amino pyridine[21].Nevertheless, most of the mentioned methods present some disadvantage and drawbacks such as long time for the color development[13],complex

procedure[17] and limited Beer's law range[14,18].

A vast number of aromatic such medicines amines as: Sulfamethoxazole[22], Pramipexole Dihyochloride[23],Metronidazole[24], Ceftazidime[25] and Metoclopramide Hydrochloride and Dapsone[26] have been determined by the diazotization reaction, which is based on the conversion of free primary aryl amine into a diazonium salt by the reaction with nitrous acid; the salt then rapidly form an azo dye with a chromogenic reagent. The procedure requires the removal of excess nitrous acid by sulfamic acid, the stabilization of intermediary diazonium salt at low temperature and the expulsion of nitrogen bubbles [22].

The present paper describes the application of 4-amino acetophenone as an inexpensive new diazotization agent for the determination of MTD in medication. The method is based on the diazotization reaction of 4-amino acetophenone with sodium nitrite in hydrochloric acid medium;the formed diazonium salt is then coupled with MTD in sodium hydroxide medium to form a water soluble mono azo dye. This method does not need to get rid of excess sodium nitrite (by addition sulfamic acid or ammonium sulfamate) because of the low concentration of sodium nitrite used by adding

equimolar solution of 4-amino acetophenone and sodium nitrite. The reaction product has been spectrophotometrically measured at 560 nm.

Material and Methods: Equipment:

Shimadzu UV-VIS double beam spectrophotometer (VARIAN UV-Visible) with 1cm matched quartz cells was used for all spectral measurements of the resulting solutions.

Reagents and Chemicals:

All chemicals were of analytical reagents grade.

1-Methyl dopa (MTD) stock standard solution (1000 μ g.ml⁻¹) was prepared by dissolving 0.0500 g of pure MTD (SDI) and made up to 50 ml volumetric flask with distilled water. Working standard solutions were prepared by suitable dilution of the stock standard solution.

2-Hydrochloric acid solution (0.8 M) was prepared by diluting 6.68 ml of 11.64 M of concentrated hydrochloric acid (BDH) with distilled water in 100 ml volumetric flask.

3-Sodium hydroxide solution (0.5 M) was prepared by dissolving 1.0000 g of sodium hydroxide (BDH) in distilled water and diluting to the mark in 50 ml volumetric flask.

4-4-amino acetophenone(3mM) solution was prepared by dissolving 0.0405 of 4-amino g acetophenon(BDH) in 5 ml ethanol ,adding 20 ml distilled water and finally the acidity was adjusted with 1 ml of 0.8M hydrochloric acid. This solution was frozen to zero degree using ice bath for 5 min. To this solution equimolar of sodium nitrite was added with shaking for 10 min., after that the azo solution was transferred to 100 ml volumetric flask,

Procedures:

1-Assay of Pure Methyl dopa (MTD):

Into series of 25 ml а volumetric falsk.transfer increasing volumes of standard stock $\mu g.ml^{-1}$) solution(1000 containing (0.01-0.9) ml of Methyldopa to cover the range of calibration curve(0.5- $45)\mu g.ml^{-1}$ in a final volume of 25 ml.to this solution add 4ml of azo 4amino acetophenon, the solution was shacked thoroughly and 2ml of 0.5M NaOH was added. The contents were diluted to the mark with distilled water. after 15 min. the absorbance of the azo dye was measured at 560 nm against the corresponding reagent blank.

2-Assay of MTD in tablets dosage form:

Ten tablets were finely weight, ground and powdered. A quality corresponding to the weight of one tablet (250 mg) was carefully weighted and made up to 100 ml with distilled water. The resultant solution was filtered and diluted to 3 different concentrations which were analyzed in five replicate as described under the assay of pure Methyldopa (MTD0.

Result and Discussion: Preliminary Studies

The proposed method involves diazotization of the 4-amino acetophenone with sodium nitrite in hydrochloric acid medium to form diazonium salt, which on coupling with Methyldopa in sodium hydroxide medium yielding a water-soluble azo dye. The visible spectrum (Fig.1) of product vielded reaction the demonstrates that the best analytical wavelength is located at 560 nm, which has a negligible absorbance at reagent blank at the corresponding $\lambda_{max}.$



Fig (1): Absorption spectra of azo dye against reagent blank (A) and agent blank against distilled water(B).

Optimization conditions:

The optimum reaction conditions have been established by varying the factors one at a time and keeping the other parameters fixed and observing the effects of the product on the adsorbance.These factors include the NaOH, 4-amino acetophenone, HCl volumes, time and addition sequence.

The effect of different volumes of (0.5 M) (0.5-4) ml Sodium hvdroxide. (1-7)ml 4-amino acetophenone (3 mM) and (0.5-3) ml Hydrochloric acid (0.8 M) were used for color development. Fig (2) depict the obtained results showing that (2ml) of NaOH (0.5 M), (4 ml) 4-amino acetophenone (3 mM) and (1 ml) of HCl (0.8 M) were efficient for accurate reproducible volumes use.

141



Fig (2): Optimum conditions for determination of Methyldopa

Different diazotization reaction times were tried in the proposed experiments. It was shown from Fig (3) that the absorbance was maximum and stable within (15 min.) min.Thus 15 min. was adopted as the diazotization reaction time.



Fig (3): Effect of time.

Acceleration of color intensity was applied by varying the addition sequence of the drug (MTD), base (NaOH) and the reagent(4-amino acetophenone).Best absorption values was achieved by adopting the following sequence (Drug+ Reagent+ Base).Fig(4).



Fig (4): Sequence effect. A=Drug+Azo reagent+Base B=Azo reagent+Base+Drug C=Base+Drug+azo reagent

Study of the dye:

The composition of the formed complex had been established using Mole ratio method which was based on the measurement of series of solution in which increased volumes(0.5-4) ml of (3 mM) diazonium reagent were added to a fixed volume(1) ml of(3 mM) Methyldopa, under optimum conditions mentioned in the analytical procedure, the results obtained in Fig(5) indicate that 1:1 azo dye was formed between Methyldopa(D) and diazonium reagent(R).The azo probable reaction path might be written as follow:



Fig (5): mole ratio plot.



Scheme (1): Reaction path.

The product formed was soluble in water. The apparent stability constant was calculated by comparing the absorbance of a solution containing stoichiometric amount of MTD (3)mM (A_s) that of a solution containing a five-fold excess of Diazionium $reagent(A_m)$ and according to analytical procedure. The stability A_s)/ A_m ; A_m =0.11, A_s =0.16,C=3×10⁻³M] of the product in water under the described experimental of conditions was 2.0×10^{3} L.mol⁻¹.

Figures of merits:

For proposed method, the calibration graph was obtained by the procedure described previous and a series of standard solutions was analyzed to test the linearity. The molar absorptivity, the sandell's sensitivity, the slope, the intercept and correction coefficient were evaluated by a least squares regression analysis and were included in Table (1).

Statistical evaluation of regression line gave the values of standard deviations for residuals(Sy/x), slop (Sa) and intercept(Sb) at 95% confidence demonstrated are in Table(1)These small figures point out to the high precision of the proposed method. limit also the of detection(LOD) limit of and

quantitative(LOQ) were calculated and shown in the same table.

Simplifier in the second			
Parameters	Values		
Correlation coefficient, r ²	0.9979		
Test for a significant correlation (at confidence level 95%).	2.262		
Regression equation	Y=0.0077x+0.0233		
Slope, b(ml. µg ⁻¹)	7.7×10^{-3}		
Intercept, a	2.33×10 ⁻²		
Conf. limit for slope $b\pm t_{sb}$	0.0077 ± 0.000259		
Conf. Limit for intercept $a \pm t_{sa}$	0.0233±0.017721		
Standard deviation of the residuals, $S_{y/x}$	5.6929× 10 ⁻³		
Standard deviation of the slop, S_b	1.1472×10^{-4}		
Standard deviation of the intercept, S _a	7.8346× 10 ⁻³		
Conf. limit conc. μg.ml ⁻¹ 95% C.I.	14.96±0.141261		
Conf. limit Abs. 95% C.I.	0.138±0.001496		
Linearity range (µg.ml ⁻¹)	0.5-45		
Molar absorptivity, ϵ (l.mol ⁻¹ . cm ⁻¹)	1.0434×10^4		
Sandell's sensitivity, S (µg. ml ⁻¹)	2.0241×10^{-2}		
Limit of detection, LOD (µg. ml ⁻¹)	1.1364× 10 ⁻¹		
Limit of quantification, LOQ (µg. ml ⁻¹)	$7.3933 imes 10^{-1}$		

Table (1): Data for calibration graph for MTD.

Accuracy and Precision of the proposed method.

The accuracy and precision of the proposed method were tested by analyzing four replicate of MTD by proposed spectrophotometric method for three different concentrations of MTD.The low values of Erel.% which are summarized in Table(2) indicate to the high accuracy and precision of the proposed method.

Table (2): Accuracy and precision ofthe proposed method.

Conc.of M	ΓD μg.ml ⁻¹	E 0/	Dag %	
Present	Found	E %	Rec. %	
15.000	14.960	-0.266	99.733	
35.000	34.875	-0.357	99.642	
45.000	44.922	-0.172	99.827	

Pharmaceutical Applications:

In order to demonstrate the applicability of the proposed method for the determination of Methyldopa. the method was successfully applied to the analysis of different pharmaceutical preparation containing MTD and results are summarized in Table (3).For all preparations examined, assay the results of proposed method were in good agreement with the labeled content.

Table (3): Application of the
proposed method for the
determination of MTD in
pharmaceutical preparations.

Pharmaceuti-cal preparation	Conc.of MTD ,µg.ml ⁻¹		Е %	Rec.%
ALDOSAM (250 mg)	15.000	14.922	-0.519	99.480
	35.000	34.854	-0.421	99.57
	45.000	45.025	+0.057	100.057
ALDOMETHYL(250 mg)	15.000	14.896	-0.692	99.307
	35.000	34.805	-0.556	99.443
	45.000	44.935	-0.144	99.855

Conclusion:

The developed methodology is very adequate for the determination of methyldopa in aqueous solution and in pharmaceutical preparation samples at a concentration level of traces(ppm) and without requiring any previous separation step nor a temperature or pH control.Morever the proposed methods are very economical when compared to other methods such as those based on the use of HPLC.

Reference

- 1-Hoffman, B.B,Lefkowitz, R.S,in: A.G. Gilman, J.G.Hardman, L.E. Limbird, P. B. Molinoff, R.W.Ruddon (Eds.).1996."The pharmacological Basis of Therapeutics",MacGrow-Hill,New York,9th ed.,Chapter 12.
- 2-The United states Pharmacopeia, 2000.The United States Pharmacopoeial Convention: Rockville, MD, 24th ed.
- 3-Saeed, S., Reyhaneh-Sadat, S. and Zahra, K. 2011.Sensitive Electrochemical for Sensor Determination of Methyldopa Based Polypyrrole/Carbon on Nanoparticle Composite Thin Film Made Situ by In Electropolymerization, Electr. analysis.23(9):2248-2254.
- 4-Mohammad, B. G. and Masoud, A.2009. Preparation of Polypyrrole/Nuclear Fast Red Films on Gold Electrode and Its application on the Electrocatalytic Determination of Methyl-dopa and Ascorbic Acid ,Electranalysis. 21(22):2461-2467.
- 5-Norouzi, P. Ganjali, M.R. hahtaheri, S.J. Dinarvand, R. and amzehpoor, 2009. Monitoring A. of Methyldopa by Fast Fourier Cyclic Transform Continuous Voltammetrv Gold at Microelectrode .Chin. J. Chem.27(4):732-738.

- 6-Azhar Ali, S. and Sami, M. A.2005. Cyclic Voltammetric Study Of α-Methyldopa At Carbon Paste Electrode, Pak. J. Pharm. Sci.18(1):6-17.
- 7-Valizadeh, H.,Nemati, M.,Hallaj-Nezhadi, S.,Ansarin, M. and Zakeri-Milani, P.2010. Single dose bioequivalence study of alphmethyldopa tablet formulations using a modified HPLC method, Arzneimittelforschung. 60(10):607-11.
- M..Erica. B..Lucia. 8-Chiara. T.,Brunetta, P.,Iiaria, C.,Roberto, P. and Roberto. G.2008. Simultaneous determination of serum concentrations of dopamine, 3-O-Levodopa, methyldopa and α -methyldopa by HPLC, Bio. and Pharm .62(4):253-258.
- 9-Gholamreza, B., Amir, Κ. and Shahla, M.2006.A rapid high performance liquid chromatographic determination of methyldopa in human serum with fluorescence detection and alumina extraction: Application to bioequivalence а study, J. Chromatog.B.832(2):197-201.
- 10-Wang, C., Wang, Z., Han, D., Hu, Y., Zhao, J., Yang, X. and Song, S.2006.Simultaneous determination of levodopa and methyldopa in human serum by capillary electrophoresis, Chin. J. Chromatog.24(4):389-91.
- 11-Ali, M.,Abdolmajid, B.,Rassoul, D.,Jalil, B.,Fatemeh, A. and Ali,A.2008. Bioelectrocatalysis of Methyldopa by Adsobed Tyrosine on the surface of Modified Glassy Carbon with Carbon Nanotubes, Int.J. Electrochem. Sci. 3:1248-1257,
- 12-Shama, K.,Sharma, SP and Lahiri, SC.2012.Spectrophotometric,Fouri er transform infrared spectroscopic and theoretical studies of the

charge-transfer complexes between methyldopa[(S)-2 amino-3-(3,4-dihydroxyphenyl)-2-methyl propanoic acid] and the acceptors (chloranilic acid,o-chloranil and dichlorodicyanoben-zoquinone) in acetonitrile and their thermodynamic properties, Spectro chem Acta A Mol Biomol Spectrosc. 15(92):212-224.

- 13-Padmarajaiah, N., Ashwinee, K.,Anantharaman, S.,Naef, G.and Avinask, K.2011. Spectrophotometric Determination of Catecholamine Using Vanadium And Eriochrome Cyanine R, Quim.Nova.34(3):1-4.
- 14-E.A., Gadkariem,K.E.E., Ibrahim, N.A.A., Kamil,M.E.M. ,Haga and H.A., El-Obeid.2009. A new spectrophotometric method for the determination of methyldopa, Saudi Pharm. J. 17(4): 289-293.
- 15- Gotardo, M.A.,Lima,L.S., R. Sequinel, Rufino J.L., Pezza,L. and Pezza, H.R. 2008. A simple spectrophotometric method for the determination of methyldopa using p-chloranil in the presence of hydrogen peroxide, Ecl. Quim. 33(3):7-12.
- 16-M., Chamsaz,A., Safavi and J., Fadaee.2007.Simultaneous kinetic spectrophotometric determination of carbidopa,levodopa and methyldopa in the presence of citrate with the aid of multivariate calibration and artificial neural networks, Analytica Chimica Acta.603:140-146.
- 17-Paulo R.,Leonardo, P.and Helena, R.P.2006.Determination of Methyldopa in Pharmaceutical Formulations by combined Spot Test-Diffuse Reflectance spectroscopy, J. Braz. Chem. Soc.17(4):674-679.
- 18-P.R.S., Riberiro,L., Pezza and H.R., Pezza.2005.Spectrophotometric determination of methyldopa in

pharmaceutical formulations, Ecl.Quim.30(3):23-28.

- 19-Tubino, M.,Batista, D. and Rodrigues, J.A.2006.Kinetic Method for the Determination of α -Methyldopa in Pharmaceutical Preparations:analytical Proceudure and Reaction Mechanism Considerations,Analytic. Lett.39(2):327-339.
- 20-Tayyebeh, M.,Abbas, A.,Lida, K.and Massoumeh, M.2006. Spectrophotomrtic Determination of Catecholamines based on their Oxidation Reaction Followed with 4-Aminobenzoic Acid,J. Braz. Chem. Soc.17(7):1259-1265.
- 21-Mouyed, Q.Al-Abachi and Muneer, A.Al-Daamy.2005.Determination of catechol Amine Drug in Pharmaceutical preparation via Oxidative Coupling Reaction with 3-Amino Pyridine and Sodium Periodate, National J. Chem. 18:226-234.
- 22-Raghad, S. and Wasan, A. Al-Uzri.2011.Spectrophotometric Method for Determination of Sulfamethoxazole in Pharmaceutical Preparations by Diazotization-Coupling Reaction, J. Al-Nahrain Univ..14(3):9-16.

- 23-B.M., Gurupadayya,V., Vishwajith and N., Srujana. 2009. Spectrophotometric Methods for the Estimation of Pramipexole Dihydrochloride in Pharmaceutical Formulations, sWorld J. Chem. 4(2):157-160.
- 24-P. Thulasamma and P. Venkateswarlu.2009.Spectrophoto metric Method for the Determination of Metronidazole in Pharmaceutical Pure and Dosage Forms,Rasayan J. Chem. 2(4),865-868.
- 25-Basavaraj, H.,Bennikallu, H.and Mruthyun, J.2008.Development and validation of spectrophotometric methods for determination of ceftazidine in pharmaceutical dosage form, Acta Pharm. 58:275-285.
- 26-Hosakere, D. R.and Malligere, A. V.
 2006. Sensitive Spectrophotometric determination of Metoclopramide Hydrocloride and Dapsone in Bulk Sample and Dosage Forms, Sci. Asia. 32:319-321
- 27-M.Q.,Al-Abachi and T.S.,Al-Ghabsha.1983.Fundamentals of Analytical Chemistry.Press of Mousl University, Mousl, , pp 346.

استكشاف تفاعل الازوتة ل4-امينو اسيتوفينون لتقدير عقار المثيل دوبا

*قسم الكيمياء/كلية العلوم/الجامعة المستنصرية.

الخلاصة:

اعتمادا على تفاعل الازونة ل 4-امينو اسيتوفينون مع كمية مكافئة من نتريت الصوديوم في وسط حامض الهيدروكلوريك لتكوين ملح الدايازونيوم الذي يزدوج مع كاشف مثيل الدوبا لتكوين صبغة ارجوانية محمرة اللون التي اعطت اعلى امتصاصية عند الطول الموجي الاعظم(560) نانومتر, تم تطوير طريقة طيفية جديدة لتقدير المثيل دوبا بصورته النقبة وفي المستحضرات الصيدلانية .عند دراسة الظروف المثلى المؤثرة على استقرارية المركب الناتج وجد بأن مدى الخطية الذي يطيع قانون بير كان ضمن المدى(6-0.5) مايكرو غرام. ملى-1 ومعامل الارتباط(0.9979) . وقد دلت النتائج المستحصلة على نجاح تطبيق الطريقة المذكورة انفا للتقدير الطيفى الكمى لدواء المثيل دوبا في المستحضرات الصيدلانية.