Optimization of a micro-High-Performance Liquid Chromatography Method for Determination of Metronidazole benzoate in Their standard Powder and in Dosages pharmaceuticals

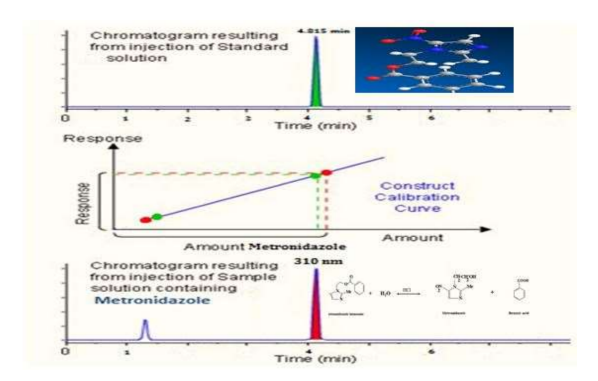
Highlights

- ❖ A new method of estimating Metronidazole benzoate (MET) in pharmaceuticals.
- Use of HPLC-UV technology for LC100 in the estimation of Metronidazole benzoate.
- study the structural synthesis of Metronidazole benzoate in the neutral, acidic and base.
- Studying the relative stability of Metronidazole benzoate during the experimental estimation process.
- ❖ Perform different applications for the purpose of validating the chromatographic method in the estimation of Metronidazole benzoate.

Abstract

context: In this manuscript, a high-performance liquid chromatography method for the determination of metronidazole in pharmaceuticals was described and developed. Method: The RP-HPLC method was developed and the results obtained to determine the form of metronidazole. Chromatographic analysis was performed in HPLC-UV system with Ion Pac column; Arcus EP-C18; 5um, 4.6×250 mm, with [Acetonitrile: triethylamine 30.70 (v/v) + 0.5 M potassiumdihydrogen orthophosphate buffer at pH 4.5 as mobile phase, at a flow rate of 1.0 ml/ min. UV detection in HPLC system was performed at 310 nm. Results: The method was validated for accuracy, precision, specificity, linearity and sensitivity. The retention time for the metronidazole was 9.9 min. Calibration plots was linear over the concentration ranges 1-5 μ g/L for the metronidazole. The Limit of detection (LLOD) was 0.115 µg/ml and the quantification limit (LLOQ) was 0.437 μg/ml. The accuracy of the proposed method was determined by recovery studies and found to be from 93.3% to 100%. Conclusion: Commercial tablet formulation was successfully analyzed using the developed HPLC-UV method that have been validated; accuracy, precision and specificity were found to be within the acceptable limits. Moreover results obtained by the suggested methods showed no significant difference between the results obtained from the suggested method.

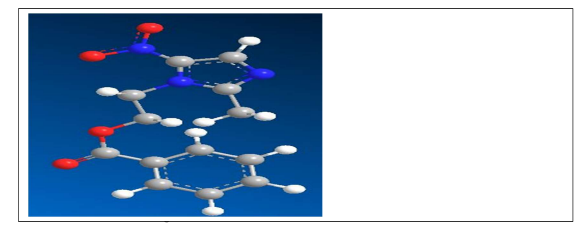
Graphical Abstract



Keywords: *Metronidazole drug, micro-HPLC,* statistical analysis, Detection limit, Quantification limit.

Introduction:

Metronidazole benzoate (MET) is used as an anti-protozoal, , chemical name AUPIC: 2-(2-methyl-5-nitro-1H-imidazol-1-yl)ethyl benzoate ($C_{13}H_{13}N_3O_4$, Mol. Wt. 275.27), Metronidazole benzoate is a white crystalline powder or light yellow odorless, almost soluble in dichloromethane, chloroform, and soluble acetone in ethanol, almost insoluble in water. Its melting point is 99-102, is a benzoic acid derivative used as antiamoebic, antiprotozoal and antibacteria [1-6].



The medical properties of metronidazole are antimicrobial and anti-microbial agents. Nitro-amidazole derivatives are also used in the treatment of anaerobic

bacteria infection. The drug is converted into anaerobic bacteria by the enzyme pyroxene-verdoxine oxidase oxidation. The nitro group in metronidazole is chemically reduced by feridoxin metabolism or by the associated ferredoxin group. Therefore, the new product is responsible for destroying the structure of the DNA spiral chains, thereby inhibiting the synthesis of DNA in microbial and bacterial organisms [7-10].

A number of different analytical methods for testing and identifying metronidazole benzoate in pharmaceutical preparations. Quantification of metronidazole is measured in pharmaceuticals using high performance liquid chromatography [11-14].

In this study, high-performance liquid chromatography (RP-HPLC) was developed using ultraviolet detector, a simple, fast and sensitive method for quantitative determination of metronidazole in pharmaceuticals. Stability of samples was determined in different laboratory conditions. It is very important to develop an appropriate analytical method to estimate the content of metronidazole in its pharmaceutical forms. In the HPLC method, an eleunt solution consisting of a mixture of solvents such as Acetonitrile, Methanol and Potassium Hydrogen Phosphate is used. A chromatographic separation column (Ion Pac Arcus EP-C18; $5\mu m$, 4.5×250 mm)is selected with a qualitative and quantitative estimate of this type of pharmaceutical and appropriate separation conditions are applied. This method was validated in accordance with the Food and Drug Administration (FDA) Guidance Document, entitled "Dynamic Verification Method" (May 2001). [15-20] The RP-HPLC method was also validated in accordance with *ICH* guidelines.

metronidazole Synthesis

metronidazole is produced by amidazole synthesis or ethylenediamine and acetic acid, followed by treatment with lime, then nickel as in the steps of the following equation [21,22].

The objective of the study

The objective of the study was to develop and verify the RP-HPLC method with an UV detector for quantitative determination of metronidazole benzoate in pharmaceuticals.

1.Experimental

1.1.Instrumentation:

LC-100 series S-HPLC features fully automatic digital computer control. Its electronic circuit design, internal mechanical structure design, processing technology, functions of cinematography workstation and the technical criteria make it a leading instruments with excellent stability and reliability, apparatus consisted of USA HPLC class LC series guipped with double beam UV-Visible spectrophotometer (Angstrom Advanced Inc., USA), model UV-100 PC with 1 cm path length quartz cell is used and it is connected to IBM compatible computer, The software was UVPC personal spectroscopy software version Matlab, R2003b was used for the proposed chemometric methods, the PLS was performed with PLS_Tool box for use with Matlab R2003b,VP pumps and variable wavelength programmable UV detector. Peak areas were integrated using a Angstrom Advanced Inc. LC solution software program. The chromatographic separation and quantification were performed on Ion Pac column; Arcus EP-C18; (250 mm × 4.6 mm; particle size 5 μm) analytical column maintained at room temperature. The mobile phase, drug standard solutions, tablet sample solutions was filtered through a millipore membrane filter before injection into the HPLC system[23-28].

2-Chemicals and reagents

2.1.Pure Standard:

Standard MET with claimed purity of 98%, according to manufacturer certificate and were kindly donated by AARTI drug Industries Pharma-India; for medical devices and pharmaceuticals).

2.2.Market sample:

Flagyl Espagne-France® tablets batch No. 75014-Paris France, were labeled to contain 500 mg MET per tablet were manufactured by Sanofi Pharma for Pharmaceuticals and Medical Appliances Flagyl-France, the other drug METROSULE-500 tablets batch No. MBO/12120686, were manufactured by (Limited Ajanta).

2.3.Configure the Samples for measurement

-HPLC grade (Sigma-Aldrich® Chemie GmbH, Germany). Solutions

-Stock standard solutions of MET were prepared in Acetonitrile: triethylamine 30:70 (v/v) + 0.5 M potassium dihydrogen orthophosphate buffer at pH 4.5 to prepare concentration of 1 mg/ml from MET [29-32].

-Working standard solutions of MET was prepared in Acetonitrile: triethylamine 30:70 (v/v)] + 0.5 M potassium dihydrogen orthophosphate buffer at pH 4.5 to prepare the concentration of $(1.0,2.0,3.0,4.0 \text{ and } 5.0) \mu g/ml$.

2.4.sample updating:

To perform model updating, the optimized Partial least squares (PLS) calibration set was augmented with different samples of Flagyl Espagne-France® tablets containing known amounts from standard MET and METROSULE-500 tablets were manufactured by (Limited Ajanta). One known concentration to three unknown concentrations of samples containing different concentrations of each were added purpose for done the initial calibration and the predictive ability of the updated sample was checked using external validation samples, then calculate the perform sample updating for each component using the developed method RP-HPLC with three concentrations of the added updating samples [33-36].

3.Procedure:

3.1.Standard drug solution

The mobile phase was used as solvent for the preparation of standard solutions. Standard stock solution of metronidazole (500 $\mu g/mL$) was prepared by dissolving an accurately weighed amount of metronidazole (50 mg) in 100 mL of mobile phase in 250 mL volumetric flask. The flask was then made up to the mark with mobile phase. The stock solution was diluted aptly with mobile phase to prepare the working standard solutions of metronidazole (1, 2, 3, 4 and 5 $\mu g/mL$).

3.2.Chromatographic conditions:

Table 1 shows the values of the basic parameters obtained using the reverse phase chromatography system (RP-HPLC).

Mobile phase	Acetonitrile: triethylamine $30:70 (v/v)$] + 0.5
	M potassium dihydrogen orthophosphate
	buffer at pH 4.5
Flow rate	1.0 mL/min
Detection wavelength	310 nm
Column temperature	Room temperature
Injection volume	20 μL
Run time	15 min

Table 1. parameters of RP-HPLC method

4.The results:

4.1. The calibration curve:

Calibration curves of the proposed method were prepared over concentration range of 1-5 μ g/ml for metronidazole benzoate. solution was prepared in triplicate and 20 μ l of each solution was injected onto the column. The peaks were determined at 310 nm. The calibration curve of metronidazole was constructed by plotting the peak area vs concentration.

4.2.Stress degradation studies:

Stress degradation studies was carried out using different ICH prescribed stress conditions such as acidic, basic, oxidative, thermal and photolytic stresses [37-39].

4.2.1.Acid degradation:

60 mg from tablet powder of metronidazole was taken in 100 ml volumetric flask. 5 ml of 0.1 N HCl was added to the flask and kept at 70-80°C reflux condition for 2-3 h. After completion of the stress, the solution was neutralized by using 0.1 N NaOH and completed up to the mark with mobile phase. Hydrolysis of metronidazole benzoate may be hydrochloric acid:

One such reaction is hydrolysis, "splitting with water." Hydrolysis of esters is stimulated by any acid or base

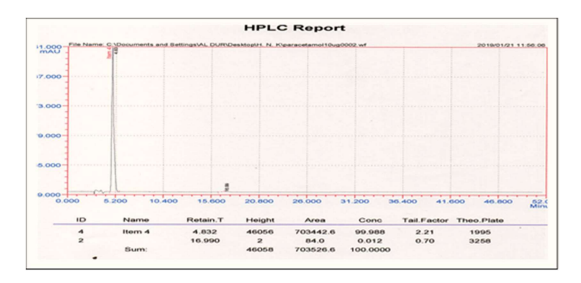


Figure .2. Chromatogram of acid degradation

4.2.2. The base degradation:

When using base such as NaOH or Potassium Hydroxide to suppress ester, products of Carboxylic Salt and Alcohol. 60 mg from tablet powder of metronidazole was taken in 100 ml volumetric flask. 5ml of 0.1 N NaOH was added in the flask and kept at 70-80°C reflux condition for 2-3 h. After completion of the stress, the solution was neutralized by using 0.1N HCl and completed up to the mark with mobile phase.

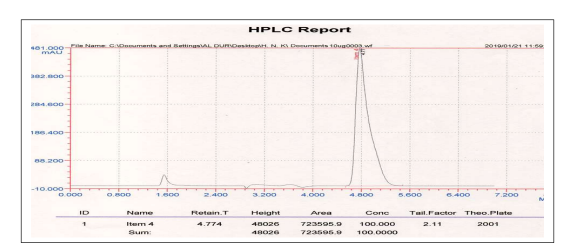
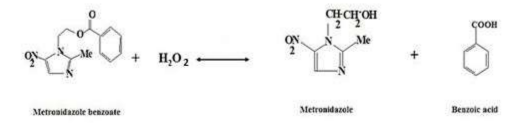


Figure .3. Chromatogram of Base degradation

4.2.3. Oxidative degradation:

60~mg from tablet powder of metronidazole benzoate and 5~ml of $20\%~H_2O_2$ were added in 100~ml volumetric flask. The flask was kept at $70\text{-}80^\circ\text{C}$ reflux condition for 2-3 h. After completion of the stress, the flask was completed up to the mark with mobile phase.



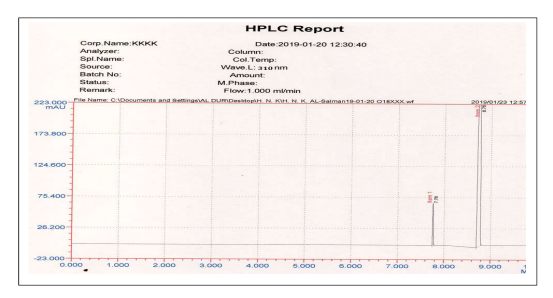


Figure .4. Chromatogram of Oxidative degradation

4.2.4Photolytic degradation:

For photolytic degradation study, 60 mg from tablet powder of metronidazole benzoate was transferred into a glass petri dish and placed in the direct sunlight for 2-3 h. After completion of the stress, the tablet powder was transferred to a 100 ml volumetric flask and made up to the mark with mobile phase, The infrared spectrum of the solution is then analyzed. The process of decomposition in this way leads to the partial disintegration of the metronidazole compound and the uncontrolled interference with pharmaceutical additives and this is evident in Figure 5, where the peaks of HPLC-UV appear irregular and sometimes overlapping.

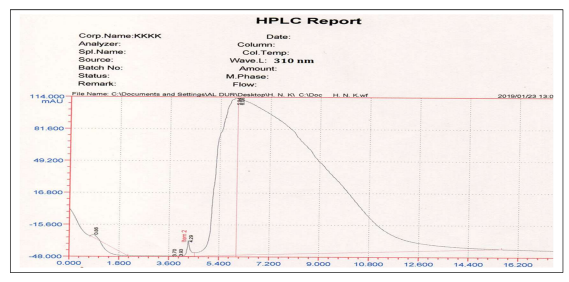


Figure .5. Chromatogram of Photolytic degradation

4.2.5. Thermal degradation:

For this, 60 mg from tablet powder of metronidazole benzoate was taken in glass petri dish and placed in hot air oven at 105°C for 2-3 h. After specified time, the tablet powder was transferred to a 100 ml volumetric flask and made up to the mark with mobile phase. Increasing the temperature of the metronbdazole solution above 100°C indicates that it is difficult to control the synthetic structure of the metronidazole and thus obtain complete thermal dissolution of the compound, this is shown in Figure 6.

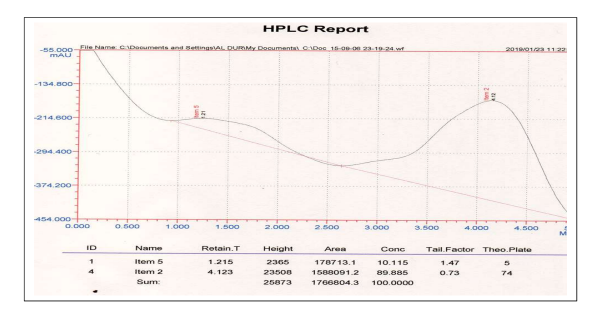


Figure .6. Chromatogram of Thermal degradation

5.Infrared spectrum of metronidazole:

5.1.For pure metronidazole powder

In the infrared spectrum of metronidazole (Fig. 7, Table 2) stretching vibrations of –OH-associated group were indicated by two absorption bands at 3220.85 cm⁻¹ and 3100.79 cm⁻¹. Bonds –C=C– and –C=N– of imidazole cycle are characterized by fluctuations in the frequency range from 1700 cm⁻¹ up to 1500 cm⁻¹, but in the spectrum of pure metronidazole they are pronounced not strongly [41].

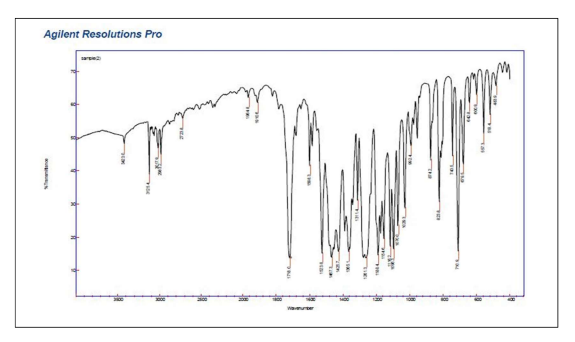


Figure: 7. IR spectrum of metronidazole

5.2. metronidazole and its crud:

The characteristic absorption bands at 1535.59 cm-1 and 1368.90 cm-1 characterize stretching vibrations of nitro group, as is evidenced by the data in comparative Table 2, the changes in significant fluctuations do not occur, so we can conclude that it is not involved in the formation of the bonds with the other materials in crud. The characteristic absorption bands at 1265.53 cm-1 at 1187.45 cm-1, which correspond to -C=C- and -C=N- bonds in the structure of metronidazole, based on the data in Table 2, are stored in the test samples.

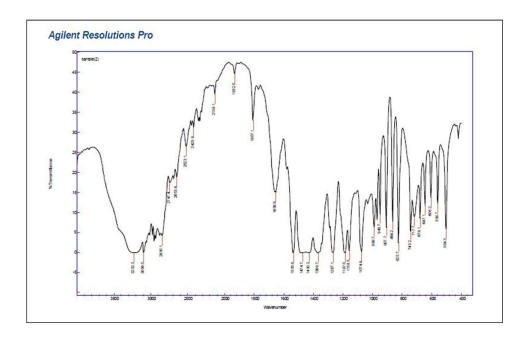


Figure. 8. IR-spectrum of metronidazole and its crud

Table .2. Characteristic absorption bands in the infrared spectra of metronidazole and its crud

	MET	MET-Crud
ν (O-H bound)	3710.70	-
ν(0-H associated)	3220.85 3100.79	3220.52 3100.88
ν(C=O)	2342.02	-
ν(C=C) ν(C=N)	-	-
ν(N=0)	1535.59 1368.90	1535.71 1368.91
v(C=C)	1265.53	1265.55
ν (C-N)	1187.45	1187.51
ν(C-O)	1074.51	1074.47
o (C-H)	825.72	825.72
M-OH	-	-

6.Discussion of the results:

6.1.The Optimization of HPLC conditions:

The chromatographic conditions were developed to separate all the degradation products from the peaks of metronidazole. During the process of HPLC method optimization, several trials were taken using Ion Pac Arcus EP-C18; 5 μ m, 4.5×250 mm, with use suitable organic phase, Acetonitrile: triethylamine 30:70 (v/v)] + 0.5 M potassium dihydrogen orthophosphate buffer at pH 4.5 and 1 ml/min flow rate. The wavelength were monitored at 310 nm. The retention time for metronidazole was 4.815 min. Good peak shape was observed of the a new analytical method .(Figure 9).

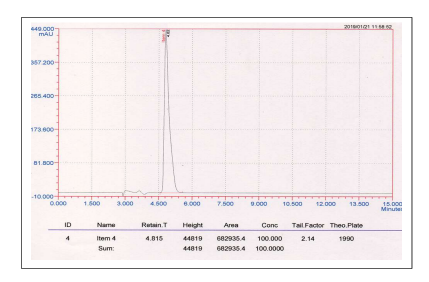


Figure 9: Chromatogram obtained after method optimization

6.2.The System suitability:

Studies were carried out for the purpose of adapting the HPLC-UV system. The standard metronidazole (3 μg / mL) was used through three replicas of the same concentration that was replicated using the optimal method. Table 3 shows the System suitability. These results meet the requirements of separation method and metronidazole estimates in various pharmaceuticals.

Parameters	Value of Metronidazole	Recommended limits
Retention time	4.815(%RSD 0.532)	RSD ≈1
Peak area	682935.4 (%RSD 0.447)	RSD ≤ 2
USP plate count	1990	≥2000-2500
USP tailing factor	2.14	≤ 2-2.5
Resolution	1.5 min	≥ 3

Table. 3. System suitability.

6.3. The validation of method and assay:

In accordance with (ICH) guidelines, the new chromatographic method HPLC-UV and parameters such as Specificity, Linearity range and Sensitivity, Regression, Precision, accuracy and rigidity were used to validate the method used [40]. In order to assess the method validity, the effect experimental conditions on the peak areas of the analytes was examined. The validity of the method was checked at concentration 3 μ g/mL for metronidazole. Table 4 summarized all The results. The results revealed that the peak areas for the drugs were unaffected small changes in flow rate, composition of mobile phase, temperature and detection wavelength indicating significant validity of the method.

Table .4. Results of method robustness

Parameter	Metronidazole (3 μg/mL)				
	Found (µg/mL)	%RSD			
Analyst	3.0	100.0	0.301		
Column	2.9	96.66	0.184		
System	2.8	93.33	0.115		

6.4.The Specificity: [41]

The specificity of the proposed method was studied using the study of forced degradation. The analysis was performed to ensure that the proposed method was able to separate metronidazole from the potential degradation products generated during the study of forced degradation. Studies were performed using acid, base, oxidation, photolysis and heat for the tablet sample at a concentration of 3 μg / ml of metronidazole. Table 5 shows the results of forced decomposition. Chromatograms shapes are shown in the Figures: 2-6. The highest percentage of deterioration occurred under the alkaline conditions of the drug. The lowest percentage of degradation of metronidazole occurred in the case of thermal and in the case of photosynthesis. One peak degradation was observed in decomposition products. Other degradation products due to stress do not interfere with the detection of metronidazole, so the method can be considered as an indicator of stability.

Table. 5. Results of forced degradation studies

Type of degradation	Metronidazo	Metronidazole (60 ug/mL)		
	% Recovery	% Degradation		
Undegraded	100.02	0.000		
Acid	98.459	1.541		
Base	94.372	5.628		
Oxidative	95.179	4.821		
Photolytic	98.114	1.886		
Thermal	98.882	1.118		

6.5. The Linearity range and Sensitivity: [42,43]

Under the optimum experimental conditions, a linear relationship was established by plotting the peaks areas for drug against the drug concentration ($\mu g/mL$). The concentration range was found to be 1 $\mu g/mL$ to 5 $\mu g/mL$ for metronidazole. the Linear regression analysis of the data gave from the following equations:

$$y = 120130x + 933.5$$
 ($R^2 = 0.9998$) for metronidazole

On the assumption that: y = peak area, x = concentration of the drug ($\mu g/mL$) and $R^2 = Regression$ coefficient. The high values of regression coefficients with small

intercept indicate the good linearity of the calibration curve that shows in Figure 10.

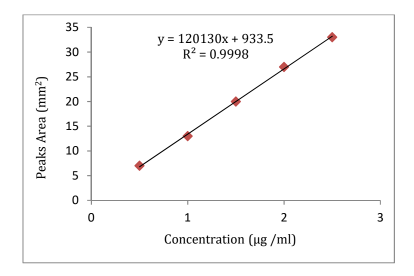


Figure . 10. linearity of the calibration curve

6.6.The Regression:[44]

The sensitivity of the proposed method was assessed by calculating limit of quantitation (LLOQ) and limit of detection (LLOD). The LOD and LLOQ were calculated as follows:

LLOQ=10×SD/S; LLOD=3.3×SD/S

Where SD = standard deviation of the drug response and S = Slope of the calibration curve. LLOD values were found to be 0.115 μ g/ml while LLOQ values were found to be 0.437 μ g/ml. These values demonstrate the satisfactory sensitivity of the proposed method for the analysis of selected drug, table 6 shows the results of Regression statistics of the proposed method.

R ²	0.9998
Standard Error	0.3830
Standard Error Estimate	0.3651
Intercept	120130X
Slope	933.5
LLOD (μg/ml)	0.115
LLOQ (μg/ml)	0.437

Table .6. Regression statistics of the proposed method

6.7.The Accuracy:[45]

for the pre analysis tablet sample solutions, a known amount of standard solution was added at three different levels, 10%, 20% and 30%. The solutions were reanalyzed by the proposed method. The results of recovery studies. The % recovery was between 98% and 100% with % RSD<0.4%. The results indicate

good accuracy of the method. The selectivity of the method was demonstrated by the noninterference of the excipients with the analysis of the analytes, The results are summarized in Table 7.

Table .7. summarized results of accuracy

Claimed Conc. (µg mL ⁻¹)	Found conc. (µg mL ⁻¹)	Recovery ± RSD
١,٠	1.0	100 ± 0.378
2.0	2.0	100 ±0.317
٣,٠	2.84	94.6 ± 0.322
3.0 μg mL ⁻¹ for Drugs (Flagyl Espagne-France® tablets)	2.98	99.3 ± 0.359

6.8.The Precision:[46]

The precision was established by analyzing metronidazole at a concentration of 3 μ g/ml. The system precision was tested by applying the developed method for the determination of metronidazole in the pure standard metronidazole for three successive times (n=3). The method precision was tested by repeated analysis of metronidazole in tablet sample for three successive times(n=3). The results are summarized in Table 8. The %RSD values for system precision and method precision were $\leq 0.4\%$, indicating that the proposed method has good precision in the analysis of metronidazole.

Table. 8. Results of precision studies

	Intra-day Inter-day			
Claimed conc. (μg mL-1)	Found (μg mL·1)	±Recovery % RSD	Found (µg /ml)	Recovery ± RSD%
١,٠	1.0	۱۰۰ ± 0.۳78	1.0	1 · · ± · ,٣٣٣
۲,۰	2.0	۱۰۰ ±0.۳17	2.1	11. ± .,٣
3.0	۲,۸٤	9 £ , 7 ± • , 77 7	2.75	91.6 ± •,٣٩٩
4.0	٣,٩٨	99.5 ± • , ٣٩٨	4.0	1 · · ± · , ٣٢ ·
5.0	٤,٩٠	98.0 ± 0. * 59	4.87	9 V.4 ± • , ٣ ٨ V
3.0 μg/ml Drug (Flagyl Espagne- France® tablets)	۲,۹۸	99.3 ± 0. * 59	2.90	97,7 ± •,٣٣٨

7. The applications of method:

The analytical method of metronidazole and METROSULE-500 were assessed by examining commercially available tablets (Negazole tablets, Gulf pharmaceutical Industries Limited U.A.E.; that claiming to contain 500 mg of metronidazole). The percentage of metronidazole was found where the values were 99.3% $\pm 0.359\%$, while the ratio of metronidazole in METROSULE-500 (Limited Ajanta) was found where the values were 102.5% $\pm 0.167\%$. this results indicating the values of % Recovery and RSD% that the proposed method was accurate and Precision in metronidazole analysis in dosages forms, table 9 summarized the applications results.

Analyte	Labeled claim	Found	Mean	%Recovery	%RSD
	(mg)	(mg)	(mg)		
Negazole tablets	900	850	880	99.3	± 0.359
	900	900			
	900	890			
METROSULE-500	650	650	651.66	102.5	±0.167
	650	645			
	650	660			

Table. 9. Assay of metronidazole in tablets

Conclusion:

This work described HPLC System (LC100 Angstrom advanced)equipped with a UV detector for metronidazole determination in two commercial pharmaceutical drugs. This developed method considered as simple, inexpensive and needs only a very small volume of the sample as well as used it's an ultraviolet detector makes this system very specific because of one peak in the chromatogram. In this application, there is no need for high sensitivity since the pharmaceutical drugs have a very low concentration. The method was validated as per the HPLC-UV guidelines and the developed method obeys Beer's law over the concentration range of $1.0-5.0~\mu g/mL$ for drugs.

Based on the results, this study divulges with important analytical method used to determine the presence of metronidazole in the dosage form. The developed and validated stability indicating HPLC-UV method for the quantification of metronidazole is simple, accurate, precise, sensitive, specific, rugged and robust. The proposed method can thus be applied for routine analysis of metronidazole in tablet dosage form.

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CONFLICTS OF INTEREST: There are no conflicts of interest for anyone.

DEDICATION: Give my work humility to the faculty of pharmacy and professors specialized in the fields of chemistry and pharmaceuticals.

FUNDING OF RESEARCH: The research was funded by the researchers himself.

AUTHOR'S CONTRIBUTIONS: This research was done individually in the laboratories of the College of pharmacy, University of Basrah. This research was completed over a period of 3 months with serious and continuous work, and therefore, excellent results were obtained in finding an easy and sensitive method to estimate the metronidazole.

Ethics: We undertake to address any ethical or health issues that may arise after the publication of this manuscript.

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