

STABILITY INDICATING RP- HPLC METHOD DEVELOPMENT OF DRUG USED IN MULTIPLE SCLEROSIS

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ABSTRACT:

A basic and specific LC strategy is portrayed for the assurance of DALFAMPRIDINE tablet measurements structures. Chromatographic partition was accomplished on a c18 section utilizing portable stage comprising of a combination of MixedPhosphate cradle (KH₂PO₄+K₂HPO₄) pH:3.5 Acetonitrile (30:70v/v/v), with discovery of 244 nm. Linearity was seen in the reach 35-105 µg/ml for DALFAMPRIDINE (r² =0.998) for drugs assessed by the proposed strategies was in great concurrence with the name guarantee. A few scientific strategies have been proposed for the quantitative assessment of DALFAMPRIDINE independently and in blend with different medications. As far as anyone is concerned straightforward, quick insightful strategy for assurance of DALFAMPRIDINE has not been accounted for up until this point. So endeavor was taken to create and approve a switched stage superior execution fluid chromatographic technique for the quality control of DALFAMPRIDINE in drug arrangements with lower dissolvable utilization alongside the short logical run time that prompts a harmless to the ecosystem chromatographic system and will permit the examination of countless examples in a brief timeframe.

Keywords: Dalfampridine, RP-HPLC, Method development, Validation

INTRODUCTION:

Dalfampridine is a neurofunctional modifier that further develops strolling speed in patients with different sclerosis. In MS, axons are continuously demyelinated which uncovered potassium channels. Subsequently, there is a hole of potassium particles which brings about the repolarization of cells and a lessening in neuronal excitability.¹ The general effect is the hindrance of neuromuscular transmission as setting off an activity potential is more enthusiastically. Dalfampridine represses voltage-gated potassium directs in the CNS to keep up with the transmembrane potential and draw out activity potential.² as such, Dalfampridine attempts to ensure that the current accessible is sufficiently high to animate conduction in demyelinated axons that are uncovered in MS patients. Besides, it works with neuromuscular and synaptic transmission by alleviating conduction blocks in demyelinated axons.³ IUPAC name is 1-[2-(dimethylamino)- 1-(4-methoxyphenyl) ethyl] cyclohexan-1-ol. Sub-atomic equation C₅H₅N₂. Sub-atomic Weight is 94.1.

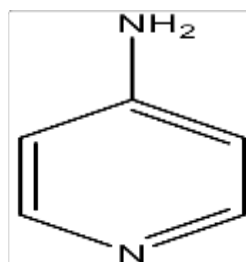


Figure 1: Structure of Dalfampridine

The detailed solvency of Dalfampridine was tracked down in water, methanol, acetonitrile (ACN), CH₃CO, ethyl ether, and exceptionally solvent in ethanol. It was seen as somewhat solvent in ligroin^{4,5}. The security of a medication in definition alludes to the capacity of a specific plan to keep up with its determinations connected with its personality, strength, quality, and immaculateness⁶. Corruption concentrates on over the medication can be determined by uncovering the medication in limits pH conditions (acidic or fundamental), oxidative responses, elated temperature, UV, and dry intensity to a degree of 5-20%^{7,8}. For examination of a medication and its substances, delicate techniques like LC/MS and GC/MS are liked yet are costly. The HPLC is viewed as the most solid and financially savvy^{9,10}. The techniques revealed by the utilization of opposite stage elite execution fluid chromatography (RP-HPLC) for the most part include the inclination method of examination, which makes investigation complex¹¹. Thus, the ongoing work expects to foster an exact, explicit, solidness demonstrating, isocratic technique for the assessment of Dalfampridine in mass and tablet structure.

MATERIALS AND METHODS:

Synthetic substances and Reagents: Dalfampridine Gift tests acquired from Madras drugs, Chennai. NaH₂PO₄ was logical grade provided by Finerchem restricted, Orthophosphoric corrosive (Merck), and Water and Methanol for HPLC (Lichrosolv (Merck)).

Chromatographic Circumstances: The chromatography was performed on a Waters 2695 HPLC framework, furnished with an auto sampler, UV indicator and Engage 2 programming. Investigation was done at 244 nm with segment INERTSIL section, C18(150x4.6 ID) 5µm, aspects at Encompassing temperature. The upgraded portable stage comprises of Methanol: Acetonitrile: water (30:50:20 v/v/v). Stream rate was kept up with at 1 ml/min.

Arrangement of arrangements:

Diluent Readiness:

Portable stage is utilized as Diluent.

Standard

weigh precisely 10 mg of DALFAMPRIDINE in 100 ml of volumetric cup and disintegrate in 10ml of versatile stage and make up the volume with portable stage. From above stock arrangement 10 µg/ml of DALFAMPRIDINE is ready by weakening 1ml to 10ml with versatile stage. This arrangement is utilized for recording chromatogram.

Sample

5 TABLETS(each Tab contains 70 mg of DALFAMPRIDINE were gauged and taken into a mortar consistently blended. Test stock arrangements of DALFAMPRIDINE (100µg/ml) and was ready by dissolving weight identical to 100 mg of DALFAMPRIDINE and broken up in adequate portable

stage. After that sifted the arrangement utilizing 0.45-micron needle channel and Sonicated for 5 min and weaken to 100ml with versatile stage. Further weakenings are ready in 5 reproduces of 10 µg/ml of DALFAMPRIDINE was made by adding 1 ml of stock answer for 10 ml of portable stage.

Procedure: 20µL of the norm, test are infused into the chromatographic framework and the regions for tops are estimated and the %Assay are determined by utilizing the formulae.

METHOD:

The created chromatographic strategy was approved for framework appropriateness, linearity exactness, accuracy, toughness and strength according to ICH rules.

System suitability parameters: To assess framework appropriateness boundaries, for example, maintenance time, following variable and USP hypothetical plate count, the versatile stage was permitted to course through the section at a stream pace of 1.0 ml/min for 30 minutes to equilibrate the segment at surrounding temperature. The overlay range of Dalfampridine was gotten and the Dalfampridine showed absorbance's maxima at 267 nm. Chromatographic detachment was accomplished by infusing a volume of 20 µL of standard into INERTSIL segment, C18(150x4.6 ID) 5µm section, the portable period of creation Methanol: Acetonitrile: water (30:50:20 v/v/v) was permitted to move through the segment at a stream pace of 1.0 ml each moment. Maintenance time, following variable and USP hypothetical plate count of the created technique are displayed in table 1.

Assay of pharmaceutical formulation: The proposed approved strategy was effectively applied to decide Dalfampridine in tablet dose structure. The outcome got for was practically identical with the relating marked sums and they were displayed in Table-2.

Approval of Insightful strategy:

Linearity: The linearity study was performed for the convergence of 35 µg/ml to 105 µg/ml level. Each level was infused into chromatographic framework. The region of each level was utilized for estimation of relationship coefficient. Infuse each level into the chromatographic framework and measure the pinnacle region. Plot a diagram of pinnacle region versus fixation (on X-pivot focus and on Y-hub Pinnacle region) and compute the relationship coefficient. The outcomes are displayed in table 3.

Precision studies: The not set in stone by help of recuperation study. The recuperation technique completed at three level 80%, 100%, 120%. Infuse the standard arrangements into chromatographic framework. Work out the Sum found and Sum added for Dalfampridine and compute the singular recuperation and mean recuperation values. The outcomes are displayed in table 4.

Accuracy Studies: accuracy was determined from Coefficient of difference for six reproduce infusions of the norm. The standard arrangement was infused for multiple times and estimated the region for each of the six Infusions in HPLC. The %RSD for the area of six imitate infusions was found. The outcomes are displayed in table 5.

Ruggedness: To assess the middle accuracy of the technique, Accuracy was performed on various day. The standard arrangement was infused for multiple times and estimated the region for each of

the six infusions in HPLC. The %RSD for the area of six recreate infusions was found. The outcomes are displayed in table 6.

Strength: As a feature of the Power, purposeful change in the Stream rate, Portable Stage piece was had to assess the effect on the strategy. The outcomes are displayed in table 7.

LOD and LOQ: The responsiveness of RP not entirely set in stone from LOD and LOQ. Which were determined from the alignment bend involving the accompanying conditions according to ICH rules. The outcomes are displayed in table 8.

$LOD = 3.3\sigma/S$ and

$LOQ = 10 \sigma/S$, where

σ = Standard deviation of y block of relapse line,

S = Incline of the alignment bend

Forced degradation studies:

The constrained debasement study is viewed as a fundamental insightful part of the medication improvement program for little atoms. Constrained debasement, normally known as pressure testing, The ICH meaning of pressure testing for the medication item is "review embraced to survey the impact to serious circumstances on the medication item. Such investigations incorporate photograph strength testing and explicit testing on specific items like metered portion inhalers, creams, emulsions and so on. According to FDA rule "Solidness is characterized as the limit of a medication substance or medication item to stay inside laid out details to keep up with its personality, strength, quality, and immaculateness all through the retest or termination dating periods". The outcomes are displayed in table 9.

THERMAL DEGRADATION:

Stress testing is probably going to be completed on single bunch of the medication substance (Programming interface). Thermolytic corruption might prompt hydrolysis/lack of hydration/isomerization/epimerization/decarboxylation/adjustments and a few sorts of polymerization responses. ICH rules propose that thermolytic debasement study ought to be completed at temperatures (in 10 augmentations for example 50oC, 60oC, and so on) over that for sped up testing and pull out the example at various time stretches during response condition. On the off chance that sensible corruption (for example 5-20%) has seen, testing can be halted as of now.

PHOTOLYTIC DEGRADATION

The photochemical soundness of the medication was concentrated by uncovering the 100µg/ml answer for UV light by keeping the receptacle in UV chamber for 24 hours. For HPLC study, the resultant arrangement was infused into the framework and the chromatogram were recorded to evaluate the security of test.

ACIDIC DEGRADATION: Test arrangement (100µg/ml) ready and moved into a 50ml volumetric flagon and disintegrate in versatile stage up to 75% then sonicate it for 10 minutes then add 1 ml of 0.1N HCl then kept in broiler at 60°C for 1 hour then cool and add 1 ml of 0.1N NaOH it then make up the volume up to 50ml with portable stage, then, at that point, place the example in the vial and measure the chromatogram.

BASE DEGRADATION:

Test arrangement (100µg/ml) ready and moved into a 50ml volumetric flagon and disintegrate in portable stage up to 75% then sonicate it for 10 minutes then, at that point, add 1 ml of 0.1N NaOH then kept in broiler at 60°C for 1 hour then cool it and add 1 ml of 0.1N HCl then make up the volume up to 50ml with versatile. stage, then place the example in the vial and measure the chromatogram

PEROXIDE DEGRADATION:

Test arrangement of Ribociclib (100µg/ml) and 1 ml of 20% hydrogen peroxide (H₂O₂) was blended. For HPLC study, 100µg/ml were infused into the framework and the chromatogram was recorded to survey the solidness of test.

RESULTS AND DISCUSSION

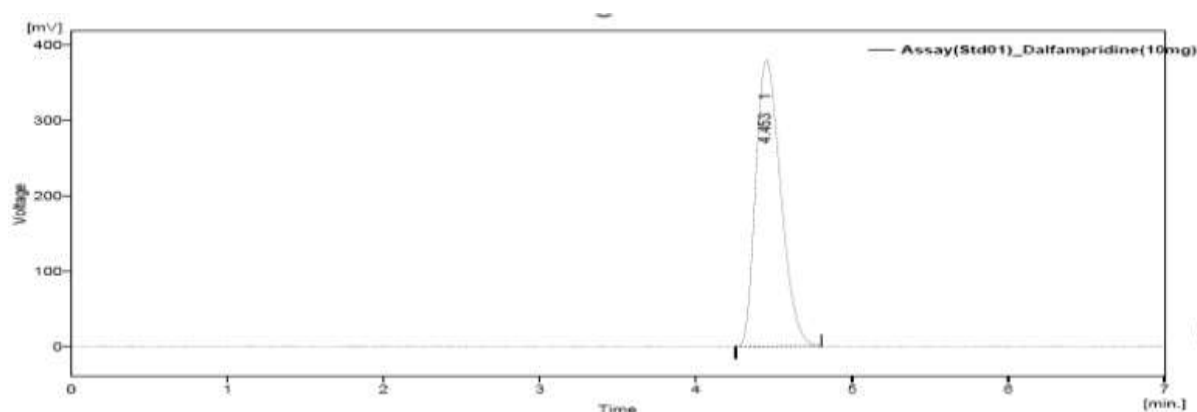


Figure 2: Standard chromatogram

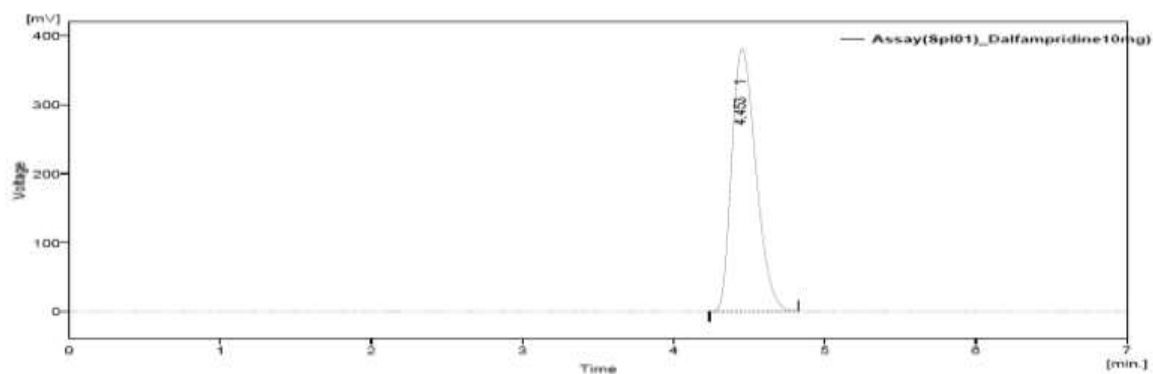


Figure 3: Sample chromatogram

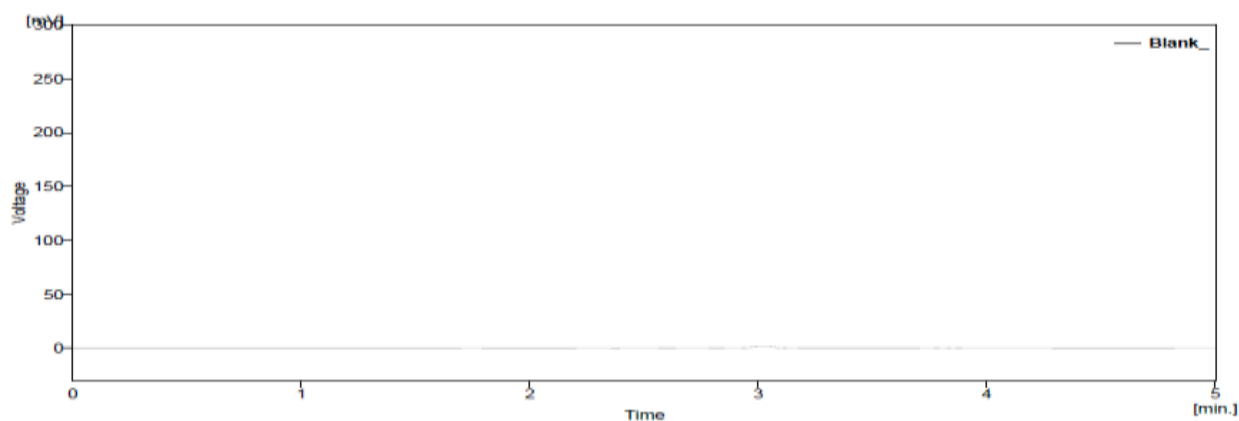


Figure 4: Blank chromatogram

Table 1: System suitability parameters

Injection	Retention time (min)	Peak area
1	4.389	4256.908
2	4.387	4254.980
3	4.387	4250.876
4	4.387	4256.980
5	4.433	4256.083
6	4.438	4250.980
Mean	4.4035	4254.468
SD	0.0248	2.836
%RSD	0.14	0.27

Table 2: Assay results for Dalfampridine

DALFAMPRIDINE		
	Standard Area	Sample Area
Injection-1	4236.32	4260.493
Injection-2	4256.047	4236.32
Injection-3	4250.49	4256.047
Injection-4	4236.32	4262.647
Injection-5	4260.403	4252.55
Average Area	4247.916	4253.611
Tablet average weight	81	
Standard weight	20	
Sample weight	700	
Label amount	70	
std. purity	99.87	
Amount found in mg	2.03	
Assay(%purity)	101.25	

Table 3: Linearity results of Dalfampridine

S. No.	Conc.(µg/ml)	Area
1	35	2490.485
2	52.5	3351.245
3	70	4207.305
4	87.5	5059.046
5	105	5933.427

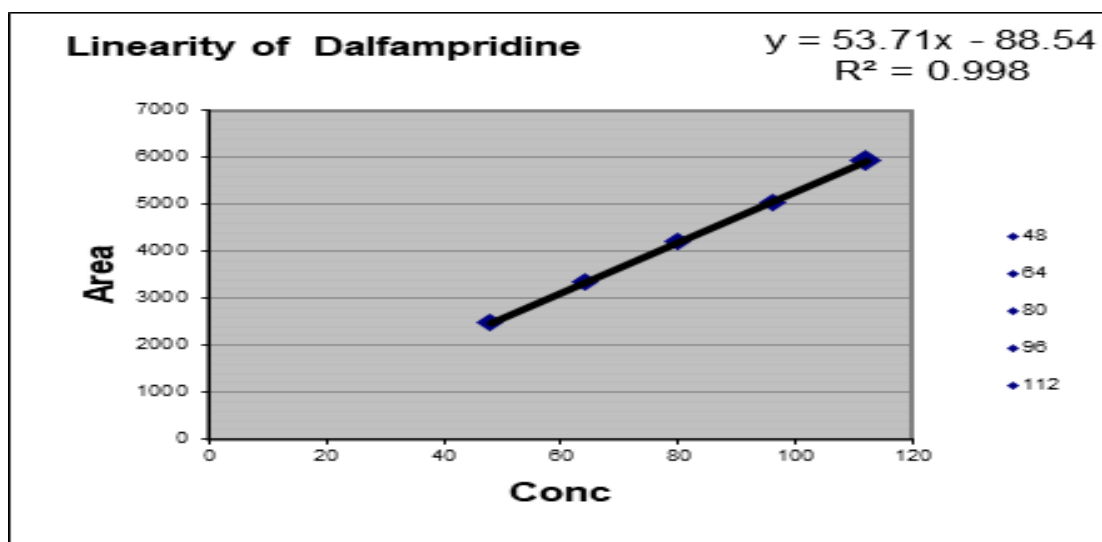


Figure 5: Linearity graph for Dalfampridine

Table 4: Showing accuracy results for Dalfampridine

Recovery level	Accuracy					Average % Recovery
	Amount taken(mcg/ml)	Area	Average area	Amount recovered(mcg/ml)	%Recovery	
80%	70	4266.47	4255.722	71.27	101.59	100.45%
	70	4240.819				
	70	4259.877				
100%	87.5	5166.633	5091.955	87.82	100.86	
	87.5	5054.273				
	87.5	5354.958				
120%	105	5833.427	5838.054	105.78	98.91	
	105	5822.224				
	105	5858.512				

Table 5: Precision results for Dalfampridine

DALFAMPRIDINE		
S.No.	Rt	Area
1	4.38	4255.073
2	4.367	4184.798
3	4.437	4259.877
4	4.397	4262.854
5	4.437	4286.183
6	4.437	4259.877
avg	4.4092	4251.444
St dev	0.0319	34.450
%RSD	0.72	0.81

Table 6. Ruggedness results of Dalfampridine

DALFAMPRIDINE	%Assay
Analyst 01	99.97
Analyst 02	100.11
%RSD	0.13

Table 7: Robustness results for Dalfampridine

Parameter	DALFAMPRIDINE	
	Retention	Tailing factor

	time(min)	
Flow		
0.8ml/min	5.383	1.600
1.2ml/min	3.847	1.471
Wavelength		
331nm	4.463	1.487
335nm	4.450	1.526

Table 8: LOD, LOQ of Dalfampridine

Drug	LOD	LOQ
Dalfampridine	61.74	4.73

Table 9: Forced degradation study of Dalfampridine

Name of the Degradation	Condition	Peak Purity	Peak Purity Value	%Assay
Thermal Degradation	60°C/7Days	PASS	+	99.07
Photolytic Degradation	1.2mill/LUX Hours	PASS	+	99.01
Acid Degradation	5mL of 3N HCl/4Hrs at 80°C	PASS	+	98.94
Base Degradation	5mL of 3N NaOH Solution/4Hrs at 80°C	PASS	+	98.97
Peroxide Degradation	5mL of 10% H ₂ O ₂ /4Hrs at Bench top	PASS	+	99.00

CONCLUSION:

The Developed HPLC method was validated and it was found to be simple, precise, accurate and sensitive for the estimation of Dalfampridine in its pure form and in its pharmaceutical

dosage forms. Hence, this method can easily and conveniently adopt for routine quality control analysis of Dalfampridine in pure and its pharmaceutical dosage forms.

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