HPLC Application

Amino Acids

	us				
Column:	Luna® 5 µm C18(2) 100 Å, LC Column 250 x 4.6 mm, Ea				
Dimensions:	250 x 4.6 mm ID				
Order No:	00G-4252-E0				
Elution Type:	Gradient				
Eluent A:	Water a company and co				
Eluent B:	Acetonitrile				
Gradient	Step No.	Time (min)	Pct A	Pct B	scx cs
Profile:	•	0		РСС В 0	HILL F(2)
Prome:	1	-	100	-	
Flaur Datas	1	20	20	80	Products used in this application:
Flow Rate:	1 mL/min				
Col. Temp.:	ambient				
Detection:	UV-Vis AbsVariable Wave.(UV) @ 225 nm (ambient)				
Analyst Note:	Description of Sample Prep and HPLC: Do not use pipette tips or microcentrifuge tubes that have been autoclaved. The water that may collect from the steam and then dry may contaminate the samples. Collection:				
	- Collect sample (3.5-4 ml unheparizied hematocrit tubes of peripheral blood or minimum of 100 mL culture media)				
	- Centrifuge @ 3000 rpm, 4°C, 10 min				
	- Transfer 75mL* of serum or culture media to a new microcentrifuge tube				
	- Sample can now be stored @ -20 °C until ready to perform extraction				
	Extraction:				
	- Add 1.4 mL (2 x 0.7 mL) HPLC-grade methanol, vortex				
	- Spin @ 15,800 g, 0 °C, 15 min				
	- Transfer 1.2mL (2 x 0.6 mL) of supernatant to a new microcentrifuge tube				
	- Dry completely in Speedvac on low heat (2-3 hr)				
	- Store @ -20 °C until ready to run HPLC or proceed directly to next step				
	HPLC:				
	- Warm samples to room temp; resuspend dry extract in 100 mL distilled-deionized water (ddH2O), let sit at RT for about 30 min then vortex				
	- Transfer 95 mL to HPLC sample vials (we use inserts inside vials); no bubbles				
	- HPLC				
	- run on a Luna C18(2) column (Phenomenex, 250mm x 4.6 mm)				
	- 20mL injection volume with a linear water:acetonitrile gradient from 100:0 to 20:80 over 20 min; 1 mL/min				
	- absorbance is read at 225 nm and integrated peaks are calculated by computer				
	- Beckman System Gold 166 detector				
	Standards: 75mL* of each triple standard is extracted as above. Each standard is run in duplicate with the average AUC used to construct a				
	standard curve, - Prepare standa	re, Use distilled-deionized water (ddH2O) to make up standards. ndard stock solutions of kynuerinine, L-tryptophan and 1-methyl-DL-tryptophan (each = 1mg/ml). Protect from light and store @ -70			
	^{eC} Triple standard is a mix of 50ml each of kynuerinine, L-tryptophan and 1-methyl-DL-tryptophan. When brought to a volume of 1ml, the concentration of each = 50 mg/mL - this dilution is used as the most concentrated standard - From the above dilution, make 4 additional serial dilutions (25, 12.5, 6.25 and 1.25 mg/mL) *A 150 mL sample will give a much more readable result if you suspect low amounts of kynuerinine.				
	Directions for dissolving 1-methyl-DL-tryptophan (1 mg/mlL ~4.6mM)				
	total volume 100mL				
	1-MeTrp	100r	ng		
	1N NaOH	1mL			
	distilled H2O	98ml			
	HCI	~1mL			
	Weigh 1-MeTrp and place in bottle. Add 1N NaOH and mix until majority of powder is dissolved. Add water and stir for ~ 1hr. Titrate to pH=7				
	using 1N HCl. Aliquot and wrap in foil and store @ -70°C.				

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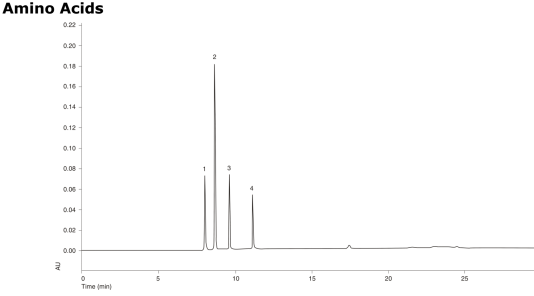


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ANALYTES:

- 5-Hydroxytryptophan 1
- 2 Kynurenine
- 3 Tryptophan
- N-Methyltryptophan 4

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