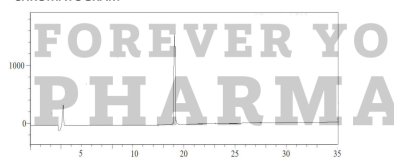


CERTIFICATE OF ANALYSIS

SAMPLE INFORMATION

Product Name	Retatrutide 5 mg	
Client Name/Lot No.	Forever Young Pharmacy / Lot# 25RET51	
Sequence	Tyr-{Aib}-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Ile-{α-Me-Leu}-Leu-Asp-Lys-{diacid-C20-gamma-Glu-(AEEA)-Lys}-Ala-Gln-{Aib}- Ala-Phe-Ile-Glu-Tyr-Leu-Leu-Glu-Gly-Gly-Pro-Ser-Ser-Gly-Ala- Pro-Pro-Pro-Ser-NH2 (sodium salt)	
Dissolution condition	100% H2O	
Length	39AA	
Molecular Weight	4813.6 g/mol	

CHROMATOGRAM



Peak#	Ret. Time	Area %
1	18.462	0.085
2	19.992	99.711
3	20.201	0.174
4	21.558	0.030

TEST RESULTS

	Specifications	Results
Strength	5.00 mg	5.26 mg
Appearance	White to off white lyophilized powder	Conforms
Purity	≥98.0%	99.7%
pH value	6.0-8.0	7.0
Impurity	Single Impurity ≤1.0%	0.2%
	Total Impurity ≤2.0%	0.3%

TEST PARAMETERS

Pump A	0.1% trifluoroacetic in 100% water
Pump B	0.1% trifluoroacetic in 100% acetonitrile
Total Flow	1.0ml/min
Wavelength	220nm
Analytical Column Type	Agilent ZORBAX StableBond 5μm C18 (4.6*250mm*5 μm)
Dissolution Method	100% H2O
Injection Volume	30uL

CONCLUSION

One 3ml vial contained a white lyophilized powder and has a black cap with silver crimp and is Lot# 25RET51.

The sample was analysed using Reverse Phase High Performance Liquid Chromatography (RP-HPLC) and determined to contain 99.7% retatrutide (5.26 mg), and the rest are impurities of minor significance.

CERTIFIED BY:

Date D.

Dane Fredericksen Analytical Chemist

