



Quality and Stability Report for Tesamorelin

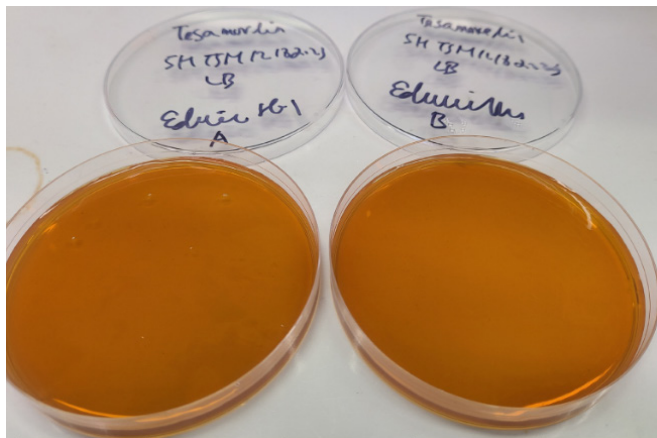
Product Name	Tesamorelin	Manufacture Date	April 18th, 2025
Batch No#	5HTSMIZ182025	Analysis Date	April 25th, 2025
Quantity	2 vial	Expire Date	One Year

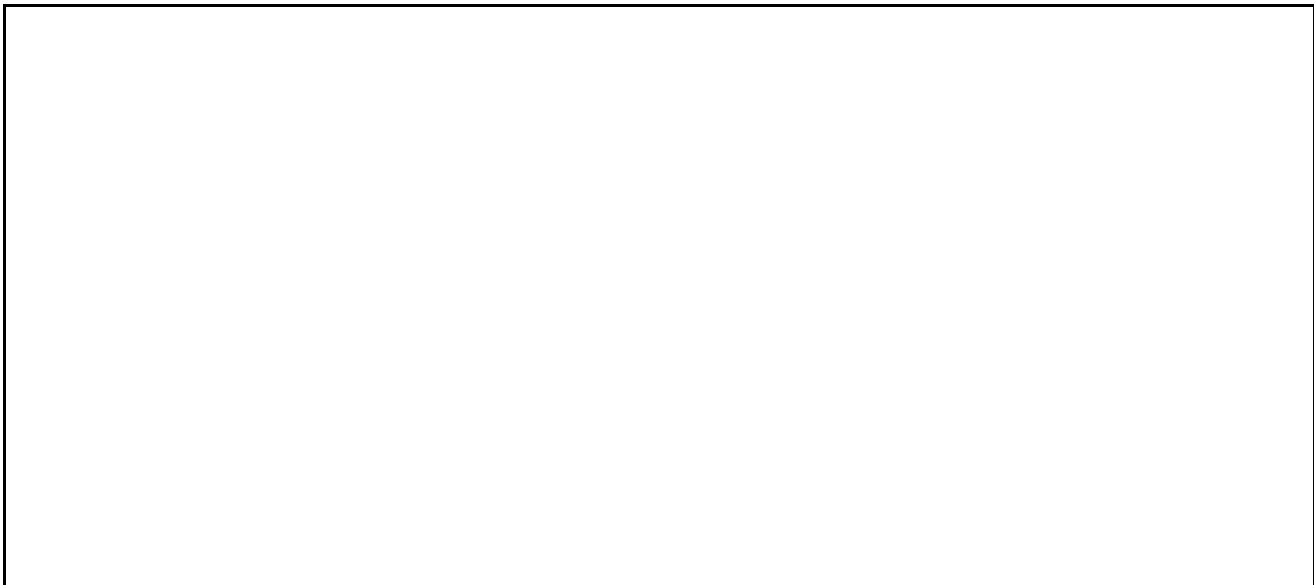
Tested Item	Specification	Results
Appearance	White Powder	Complies
Odor	Odorless	Complies
Fluid Clarity	Solution with precipitates	Complies
pH value	pH=7.0 after dissolved	Complies
Heavy Metals	Test	Complies
Arsenic	Test	Complies
Lead	Test	Complies
Mercury	Test	Complies

Microbie Test		
Total Plate Count	2 plate for 1ml of sample each plate	Complies
Yeast & Mold	Negative (Potate LB Plates)	Complies
Bacteria	Negative (Potate LB Plates)	Complies
Endotoxin Level	<0.05 EU/ml	Complies

Conclusion	The QC data shows no bacterial/fungal contamination present. This product is qualified for each specification for release.
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Note: LB agar and enriched LB culture media were used to test microbie in the samples. 48 hours culture time was used for mold and bacterial growth. Endotoxin was below 0.05 EU/ml, kit from Genscripts Inc.





QC Scientist	Edwin Hu, PhD	Report Date	25-Apr-25
Signature	<i>Edwin Hu</i>	Date	<i>25-Apr-25</i>