







Certificate of Analysis

Product Name:	Crystel GOLD CRY 101	
Reference:	CRY/GOL/019	
Batch Number:	P813701	
Date of Manufacture:	2018-03-23	
Expiry Date:	2020-03-23	

This is to certify that Crystel Gold solution above was tested on 2018-03-23 and found to comply with all requirements as set out in the specifications and method of analysis.

PROPERTIES	SPECIFICATION	RESULTS
APPEARANCE	Clear and free from suspended matter	Conforms
ODOUR	Alcohol like	Conforms
SPECIFIC GRAVITY	0.870 - 0.890	0.875
STERILITY	No growth	Conforms
IRRADIATION	Achieves minimum dose	Conforms
ENDOTOXIN	<0.25EU/ml	<0.1
LINDOTOMIN	Recovery 50-200%	137%

Completed By:	Reviewed By:
Quality Department: Carareus	<u> Hidinska</u>
Date: 2015-07-05	Date: 2018 - 07 - 05

DRF1574 CRY/GOL/006/Issue4/Sept2012





Sterility Testing Analytical Report

Please note the results below are not generated at Associates of Cape Cod Int. Inc., but reported from a subcontractor's final report.

Sample ID: CRY101 Gold

Lot No: P813701 ICN: 0618015D

Subcontractor's Report Number: 2018110008

Lab Book Reference: ST03-176 Date Received: 18 JUN 2018 Date Completed: 03 JUL 2018

No. of samples: 10

Sterility Test - Membrane Filtration

Medium	Result
Tryptone Soy Broth	No Growth
Fluid Thioglycollate Medium	No Growth

Conclusions:

The samples submitted comply with the test for sterility in accordance with the client's methodology.

Results Reported by:	
RELLEY	Date: 03 5UL2018
Laboratory Technician/Designee: L Ellis	
Reviewed by:	
Quality Assurance Manager	Date:
☐UK General Manager ☐ Technical Manager	
Laboratory Manager	
QA Approved by:	
I harrier .	Date: C4 Jul 2018
☑ Quality Assurance ☑ UK General Manager ☑ Technical Manager	

Revision History:

DCCF No:	Revision:	Description:	Originator:	Effective Date:
N/A	1	Revision 1, no changes	L Ellis	03 JUL 2018

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ST03-176

ICN: 0618015D

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REVISION NO: 1

DATE: 03 JUL 2018

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LIMULUS AMEBOCYTE LYSATE (LAL) ANALYTICAL REPORT Release Test - Kinetic Turbidimetric Method (SOPCTS012)

Sample ID: CRY101 Gold

Lot No:P813701 ICN No:0618015D Lab Book Reference: EL194-134 Date Received: 18 JUN 2018 Date Completed: 21 JUN 2018

Reagents

Pyrotell®-T: Lot Number 517-10-848-T. Pyrotell®-T was reconstituted with Pyrosol® Buffer and used to Assay all dilutions.

Pyrosol® Buffer: Lot Number 226-414.

LAL Reagent Water (LRW): Lot Number AC10244546. LRW was used to reconstitute endotoxin, prepare dilutions of endotoxin and sample(s), and serve as the negative control.

Instrumentation: Pyros Kinetix Incubating Kinetic Tube Reader; S/N; 1E-1055.

Control Standard Endotoxin (CSE): Lot Number 159, The potency of the CSE is 14 EU/ng when measured against Reference Standard Endotoxin (RSE), lot number H0K354 (Food and Drug Administration, CBER) with Pyrotell®-T lot number 517-10-848-T.

The sensitivity of the assay is the lowest concentration of CSE used to construct the calibration curve. A series of 10-fold dilutions between 0.001 and 1.0 EU/mL was tested in duplicate and the results were used to construct the standard curve.

Preparation and Testing of Sample: The samples were diluted with LRW and analysed at a 1:100 dilution, in accordance with validation EL168-03. Positive product controls were tested, in parallel, on sample dilutions spiked with an Endotoxin concentration of 0.1 EU/mi. The sample to lysate ratio was 4:1.

Results: The Release Test is performed according to a validated method and is used to release finished products. The test can also be performed for release of raw/in-process materials. The absolute value for the correlation coefficient for the regression of log onset time versus log endotoxin concentration was greater than or equal to 0.980. The negative control (LRW) was tested in duplicate and contained less than 0.001 EU/mL. Inhibition or enhancement of the bacterial Endotoxin test by the sample is absent if the concentration of the Endotoxin assayed in the Positive Product Controls are within 50% to 200% of the nominal concentration.

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EL194-134 REVISION NO:1

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ICN: 0618015D DATE:21 JUN 2018

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TSCTS015-3.1 Rev 18

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Sample Results:

Sample Identification	% Spike Recovery	EU/ml
P813701	137	<0.1

All Samples analysed passed the specification of < 0.25 EU/ml.

proval:	
Techncian: Laboratory Technician: L Ellis	Date: 21 JUN 2018
Reviewed By: Review Board Laboratory Manager	Date: 25500 2019
QA Approval:	Date: 267UN7018

References:

Associates of Cape Cod SOPCTS012, General Kinetic LAL Turbidimetric Assay
United States Pharmacopoeia <85> Bacterial Endotoxins Test
European Pharmacopoeia, Section 2.6.14.

Revision History:

DCCF No:	Revision:	Description:	Originator	Effective
ALA				Date:
NA	1	Revision 1 No Changes	L Ellis	21JUN2018

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http://www.steris-ast.com

Certificate of Irradiation

Date Issued: 11-Jun-2018

UK33S12118562-1-1

This is to certify that AST Daventry Synergy Health PLC, a STERIS Company has where appropriate delivered an irradiation process in accordance with the current certified standards:

EN ISO 11137-1 Sterilisation of Health Care Products EN ISO 13485 Quality System - Medical Devices

The Tristel Company Ltd
Unit 4C Lynx Bus Park, Fordham Rd
Newmarket
Suffolk CB8 7NY
UNITED KINGDOM

Order Information		
Account Number:	100708	
Synergy Health Sales Part Reference:	1108717	
Customer Reference Number:	707323	
Product Description:	ALCOHOL 1 LTR TRIGGER SPRAY DV4778	
	20-45kGy	
Validation Reference: 4778		
Quantity Received:	41	
Customer Minimum Specification kGy:	20.0	
Customer Maximum Specification kGy:	45.0	
Customer Unit Lot/Batch Number:	BN: P813701, 1 PLT	
I	rradiation Data	
Date and Time of Irradiation:	11-Jun-2018 04:20	
Reference Dose Range kGy:	34.0 - 34.5	
Calculated Minimum Dose kGy:	27.5	
Calculated Maximum Dose kGy:	39.8	

Irradiation Release Authorised By Synergy Health Sterilisation UK, a STERIS Company