



Leukotoxin (LtxA) Detection Report

Protocol No.: DET-PRO-017

Name of the product: Lumineux Oral Essentials clean and fresh

Approved by: *Nour Abochama* Date: 2019-03-01
Nour Abochama
Director QA/QC, American Testing Lab Inc.

Approved by: *Kourosh Maddahi DDS* Date: 3/1/2019
Kourosh Maddahi DDS
Founder, Oral Essentials Inc.

Table of Contents

1	Objective:	3
2	Scope:	3
3	Responsibility:	3
4	Materials, Equipments and documents:	3
4.1	List of Equipment:	3
4.2	Injection Run:	4
4.3	System Suitability:	4
4.4	Calculations:	4
5	Protocol	5
7	Results:	5
8	Conclusion:	5

1 Objective:

To determine the effect of Lumineux Oral Essentials clean and fresh of Leukotoxin (LtxA) Fats concentrations.

2 Scope:

A 3T3 clone A31 Cell Line from mouse was exposed to a 5000 µg/ml solution of test product for 24 hours. A determination of Leukotoxin (LtxA) Fats concentration was performed using HPLC method.

3 Responsibility:

The persons responsible for the process will perform the and record the information. The responsible person will supervise the study, verify the completion of the records and write the report. Quality Assurance will review and approve the Report.

Name of the Personnel	Designation	Signature
___Nour Abochama_____	___Director QA/QC___	<u><i>Nour Abochama</i></u>
___Reem Samman_____	___Chemist_____	<u><i>ras</i></u>

4 Materials, Equipments and documents:

List with Equipment No. and all Standard Operation Procedures (SOPs) for the normal processes under test.

4.1 List of Equipment:

Name of Equipment	Equipment No.
GC with MS detector	LAB-2442
Balance	LAB-2233
Pipette	LAB-2422

Complied By: ras Date: 2019-02-28
Reviewed By: Nour Abochama Date: 2019-02-28

4.2 Injection Run:

- 4.2.1 1 inj Mobile Phase BLK;
- 4.2.2 5 inj working standard STD;
- 4.2.3 1 inj samples;
- 4.2.4 1 inj control (working standard) QSTD

4.3 System Suitability:

- 4.3.1 From the 6 injections of standard, area RSD should not be more than 5.0%.
- 4.3.2 Recovery between standard control and mean area of standard should be within 95.0%-105.0%.

4.4 Calculations:

- 4.4.1 Calculate against the mean area of first 5 injection of standard.

4.4.1.1 Recovery standard

$$\% \text{ Recovery} = (\text{Area}_{\text{qstd}} / \text{Mean Area}_{\text{std}}) \times 100$$

Where:

$\text{Area}_{\text{qstd}}$ = Area of peak of interest in standard or Control.
 $\text{Mean area}_{\text{std}}$ = Mean area of peak of interest in standard

4.4.1.2 Sample Calculation

$$\% \text{ of Leukotoxin (LtxA) Fats in Products} = (\text{Area}_{\text{spl}} / \text{Mean Area}_{\text{std}}) \times (\text{C}_{\text{std}} / \text{W}_{\text{spl}}) \times 100 \text{ mL} \times 100$$

Where:

Area_{spl} = Area of of Leukotoxin (LtxA) Fats in Sample preparation
 $\text{Mean Area}_{\text{std}}$ = Mean Area of Fats in working standard preparation
 C_{std} = Concentration of standard in working standard (mg/mL)
 W_{spl} = Weight of sample (mg)

Complied By: ras Date: 2019-02-28

Reviewed By: Nour Abochama Date: 2019-02-28

5 Protocol

- 5.1 The test item is prepared y adding 10% (v/v) FCS and 1% antibiotics (v/v) to the artificial Sylvia
- 5.2 Prepare 5000 µg/mL solution of Lumineux Oral Essentials clean and fresh. Expose the test
- 5.3 item to the product solution for 24, 36, 48, 60, 72, 84 and 96 hours then prepare as described in section 5.4.

6 Approval:

- 6.1 Submit the Document to QA for review and approval.

7 Results:

Exposure Time	Blank (µg/ml)	Test Product (µg/ml)	Water (µg/ml)	Ref. Product (µg/ml)
0 hours	0.00	0.19	0.19	0.19
24 hours	0.00	0.15	0.19	0.19
36 hours	0.00	0.10	0.19	0.19
48 hours	0.00	0.08	0.18	0.19
60 hours	0.00	0.07	0.18	0.18
72 hours	0.00	0.04	0.19	0.18
84 hours	0.00	0.03	0.19	0.05
96 hours	0.00	0.03	0.19	0.05

8 Conclusion:

The toxin concentration have been 84% decreased in test item after 96 hours of exposure. The solution at 5000 µg/mL is prepared form Lumineux Oral Essentials clean and fresh Lot# 17K13B2

Approved By: Nour Abochama Date: 2019-03-01
Nour Abochama, M. Sc.,