SUPPORTING INFECTIOUS DISEASE RESEARCH

#### ML2038/BfrA Recombinant Protein from Mycobacterium leprae

#### Catalog No. NR-19337

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**Product Description:** NR-19337 is a recombinant form of the bacterioferritin protein (ML2038/BfrA) [also known as major membrane protein II (MMP-II) and 22 kDa protein] from *Mycobacterium leprae*. The recombinant His-tagged protein was expressed in *Escherichia coli*, strain BL21(DE3)pLysS and purified using standard chromatographic techniques followed by endotoxin removal procedures.

#### Lot: 61391642

## Manufacturing Date: 310CT2012

QC testing was performed by Colorado State University under the Leprosy Research Support Contract (NIH). The Colorado State University documentation for lot 12.rEC.10.15.coc.MLMMPII is attached.

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# **Recombinant Protein Production and Quality Control Record**

Date Production Started: 9/24/2012

Lot Number: 12.rEC.10.15.coc.MLMMPII bacterioferritin; BEI lot# 61391642

Notebook Number and Page Number: COC TB #1 NOTEBOOK pp. 104-105; COC TB

#2 NOTEBOOK pp. 5-10 & 18-24

Production from Seed Culture/ Clone: no

Production from freshly-transformed Cells: yes

Host Strain used for Gene Expression: E. coli BL21 (DE3) pLysS

Recombinant Plasmid possessing the Recombinant Gene: pET-23b

Culture Type? Shake Flask\_\_\_\_\_ Stationary\_\_\_\_\_ Fermenter X

Culture Size: 5L

Culture Medium: HyperBroth (Athena Enzyme Systems)

Selection (Antibiotic/ Concentration):  $Kan^{50}$ 

Time and Temperature of culture prior to Induction: 2:00, 37.2°C

Final Concentration of IPTG added for Induction: 0.5 mM

Method for Lysis of Cells: Probe Sonication

Protein Purification Procedures: His-bind Resin Purification

Date Production Finished: 10/31/2012

### **NOTES ON PURIFICATION:**

Cells were sonicated on ice with 60 second bursts followed by 90 second intervals.

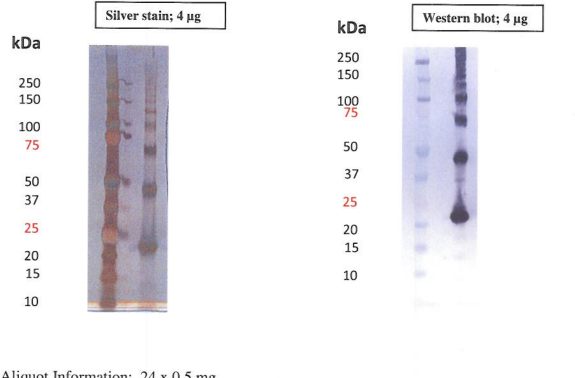
His-bind resin purification per Novagen except for additional Endotoxin (ET) removal steps.

ET removal done by washing column with 10 column volumes (CV) of 10 mM Tris-HCl, followed by 10 CV of 0.5% ASB-14. This was again followed by 10 CV of 10 mM Tris-HCl and eluted with 5 CV of 10 mM Tris-HCl + 1 M Imidazole . All buffers were pH=7.9.

Eluted proteins were exchanged into 10 mM Ammonium Bicarbonate + 4M urea; then into 10 mM Ammonium Bicarbonate + 2M urea; and finally into 10 mM Ammonium Bicarbonate .

## **Quality Control**

Lot Number: 12.rEC.10.15.coc.MLMMPII bacterioferritin Method for Determining Protein Concentration: BCA (Pierce) Final Protein Concentration: 1.454 mg/mL Performed Endotoxin Removal? Yes Endotoxin Contamination: 6.52 ng/mg protein Purity confirmed by SDS-PAGE and Silver Staining (see below) Identity confirmed by Western Blot: \_\_x\_\_\_ or Mass Spectrometry: \_\_\_\_(see below) Antibody used for Western Blot: α-penta-Histidine Polyclonal



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Producer's Name:	Carma O. Cook	_Date: 12/2//2012
Supervisor's Name:	Hand OL	_ Date: 12/27 /2012