## Procedure for Endotoxin removal with EndoBind-R<sup>TM</sup>

- Remove the top cap.
- Remove the bottom cap.
- Allow the 0.02% sodium azide storage solution to drain from the column.
- Wash the column with 10.0 ml endotoxin-free water to remove sodium azide
- Equilibrate the column with 1.0-2.0 ml of sample buffer.
- umn. Collect the flow-through in an endotoxin-free tube. Add 0.5-1.0 ml of sample to EndoBind-RTM and allow it to penetrate the col-
- this step until a total of five 1.0 ml fractions have been collected Add 1.0 ml of sample buffer to the column and collect flow-through. Repeat

fractions. Check for endotoxin removal using an appropriate assay. The majority of protein and DNA generally pass through the column in the first two sorbance of the flow-through fractions at OD<sub>280</sub> for proteins and OD<sub>260</sub> for DNA each fraction for endotoxin-free product. This can be done by measuring the ab-Substances pass through the column at different rates, so it is important to check

Refer to EndoBind-RTM Application Notes for guidance on proper sample buffer

## Procedure for EndoBind-RTM Storage and Maintenance

After each use the EndoBind-RTM column should be washed before storage:

- Rinse the column with 10.0 ml endotoxin-free water.
- Wash the column with 2.0 ml of 2.0 M sodium chloride
- crucial to remove all traces of salt before treatment with detergent Rinse the column with 10.0 ml endotoxin-free water. This step is
- Wash the column with 2.0 ml of 1% sodium deoxycholate.
- Rinse the column with 10.0 ml endotoxin-free water.
- 4.00 Add 4.0 ml of 0.02% sodium azide and store upright at 4°C

The column is shipped in 0.02% sodium azide. each column is dedicated to the purification of a specific protein or DNA solution In addition to routine cleaning, EndoBind-RTM shows optimal performance when

> EndoBind-R™ EndoBind-R™ EndoBind-R™ Endotoxin Removal Products: Soluble Biotech, Inc. offers superior products for detection, removal and neutralization of bacterial toxins • For Research Use Only • 1 ml column 5 ml column Bulk resin EBR-3001.01 EBR-3005.01 Inquire



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removal of endotoxin. It can be used to remove media. Under optimized conditions it can also samples with high specificity and minor product loss. It can be used over a broad range of endotoxin from water, buffer and cell culture EndoBind-R<sup>TM</sup> is an agarose-based affinity chromatography media specifically for the remove endotoxin from protein and DNA conditions.

#### **Product Description**

Hydrophilicity minimizes non-specific binding

pH range (Buffer)

2,000,000 EU/ml resin

 $10^{-7} - 10^{-8} \text{ M}$ 

pH 2.0-9.0

**Product Characteristics** 

EndoBind-RTM

Purity

>98% Factor C Sushi Peptide

Regular use between 4°C and room temperature

Temperature Stability

Flow Rate

Gravity

Binding Affinity **Binding Capacity** 

- Large pore size
- No special buffers required for binding or washing
- Chemically-synthesized compound minimizes lot-to-lot variability
- Non-cytotoxic; non-hemolytic
- High binding affinity and capacity

#### Advantages

## Predictive Oncology

### EndoBind-R<sup>TM</sup>

Catalog No. EBR-3001

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thelium and arterial smooth muscle cells. Recentlevels of endotoxin can trigger immune responses cells including monocytes, neutrophils, dendritic cell culture media and protein and DNA preparaendotoxin. On average, a single E. coli cell conly, a 34 amino acid Sushi domain was identified endotoxin using EndoBind-R" is fast, easy and tains 2,000,000 LPS molecules. Sub-nanogram in the Factor C enzyme of the LAL cascade that outer membrane of gram-negative bacteria concells, hepatocytes, vascular and respiratory epiand cell culture media. It has also been used to remove endotoxin from protein and DNA solu-The removal of endotoxin from water, buffers, tions is a priority. The majority of lipid in the used to remove endotoxin from water, buffers, and alter the phenotype and function of many shows very high affinity for LPS. It has been tions with minimal product loss. Removal of sists of lipopolysaccharide (LPS), also called inexpensive.

#### Introduction to Endotoxin

# Removing Endotoxin from Protein Solutions

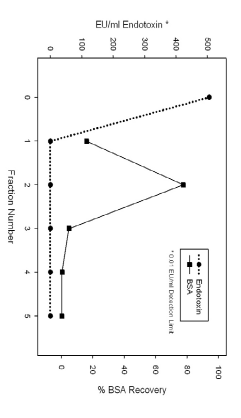


Figure 1. Endotoxin Removal from BSA. BSA samples at 1 mg/ml were prepared in 20 mM sodium acetate at pH 5.0 with 150 mM sodium chloride and 50 ng/ml *E. coll* Co55: *B5* endotoxin and applied to EndoBind-R<sup>78</sup>. The protein was recovered in four subsequent 1 ml washes. Protein recovery was determined by absorbance and endotoxin levels were determined by PyroGene (Lonza) assay.

a rather low value of about 0.05 EU/mg. E. coli O55:B5 endotoxin at a concentration of 50 ng/ml (500 EU/ml) was added to the protein solution and purified experiments showed that a 20 mM sodium acetate buffer at pH 5.0 containing moved from the starting material doBind-R<sup>TM</sup> column. Even the 0.05 0.01 EU/ml in all five column fractions. This represents more than 99.998 % tained less than 6% of the initial BSA load. The LPS content in the sample load initial with EndoBind-R<sup>TM</sup>. pared in 20 mM sodium acetate at pH 5.0 containing 150 mM sodium chloride BSA at these conditions was measured. A 1 mg/ml solution of BSA was prehigher than the isoelectric point of BSA (4.6). Next, endotoxin removal from tric point. As an example, bovine serum albumin (BSA) was purified. ionic strength of the buffer should be optimized in regard to the protein isoelec-For optimal protein purification and recovery using EndoBind-R<sup>TM</sup>, the pH and (fraction 0) measured 506 EU/ml and was reduced to below the detection limit of The low endotoxin BSA was tested for contaminating endotoxin and found to be 150 mM sodium chloride gave the best product recovery. This pH is slightly removal protein (Figure 1). fraction 2, as a 77% protein peak. Fractions 3 through 5 combined conand over 99% protein recovery after purification with the En-The flow-through, fraction 1, contained about 16% of the However, the majority of BSA eluted into the first EU/ml contaminating endotoxin was re-First,

For a more detailed explanation of buffer optimization and purification of protein solutions using **EndoBind-R<sup>TM</sup>**, refer to the **Bi-Objech**, **Inc. EndoBind-R<sup>TM</sup>** protein Purification Application Notes. This document outlines both salt and pH optimization protocols and their application to purify proteins such as bovine serum albumin, human transferrin, bovine liver catalase, hemoglobin from bovine erythrocytes, and rabbit IgG.

## Removing Endotoxin from DNA Solutions

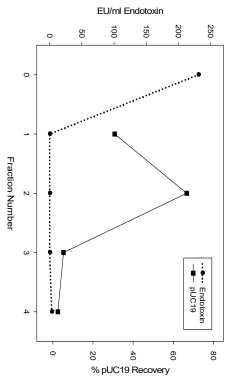


Figure 2. Endotoxin Removal from pUC19. pUC19 samples at 30 µg/ml were prepared in TE (10 mM Tris, 1 mM EDTA) at pH 8.0 with 1 M sodium chloride and 25 ng/ml 18. coli 1053/58 endotoxin and applied to EndoBind-R<sup>TM</sup>. The DNA was recovered in three subsequent 1 ml washes. DNA recovery was determined by absorbance and endotoxin levels were determined by PyroGene (Lonza) assay.

cloning vector pUC19. Previous experiments showed that a TE buffer at pH 8.0 lar experiments with small linear DNA fragments gave nearly identical results. moval of over 99.99% of endotoxin with near complete product recovery. Simi-EU/ml (fraction 0) and was reduced to below the level of detection (0.01 EU/ml) 2). In addition, endotoxin removal was nearly complete. eluting in the flow-through and a peak value of nearly 67% in fraction 2 (Figure was rinsed with three 1 ml washes of TE pH 8.0 with 1 M sodium chloride R<sup>TM</sup> column. The flow-through was collected as fraction 1. Next, the column concentration of 25 ng/ml (250 EU/ml) (fraction 0) and added to the EndoBindendotoxin removal, a 30 µg/ml pUC19 solution was prepared in TE pH 8.0 with containing 1 M sodium chloride was sufficient for high DNA recovery. To test DNA purification using EndoBind-RTM was investigated using the common in all samples collected from the  ${f EndoBind ext{-}R^{TM}}$  column. This represents re-1 M sodium chloride. E. coli O55:B5 endotoxin was added to the solution at a (fractions 2-4). DNA recovery was very high with about 30% of the initial load The load contained 23

For a more detailed explanation of buffer optimization and purification of DNA solutions using **EndoBind-R<sup>TM</sup>**, refer to the **BioDtech**, **Inc. EndoBind-R<sup>TM</sup>** DNA Purification Application Notes. This document outlines both salt and pH optimization protocols and their application to purify small, linear DNA fragments as well as plasmid samples with both high and low levels of endotox in contamination.