

**Instructions For Use**

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**Access System Check Solution****REF** 81910**FOR PROFESSIONAL USE ONLY**

Rx Only

**PRINCIPLE****INTENDED USE**

For use with the Access Immunoassay Systems in the weekly maintenance System Check Procedure.

**REAGENTS****PRODUCT INFORMATION**

- Provided ready to use.
- Stable until the expiration date stated on the label when stored at 2 to 8°C.

**R4****Access System Check Solution: 6 x 4.0 mL.**

Alkaline phosphatase, 1% bovine serum albumin (BSA), 0.25% ProClin\* 300, &lt; 0.1% sodium azide.

\*ProClin is a trademark of LANXESS Corp.

**WARNING AND PRECAUTIONS**

- For *in vitro* diagnostic use.
- For hazards presented by the product refer to the following sections: REACTIVE INGREDIENTS and GHS HAZARD CLASSIFICATION.

**REACTIVE INGREDIENTS** **CAUTION****Sodium azide preservative may form explosive compounds in metal drain lines. See NIOSH Bulletin: Explosive Azide Hazard (8/16/76).****To avoid the possible build-up of azide compounds, flush wastepipes with water after the disposal of undiluted reagent. Sodium azide disposal must be in accordance with appropriate local regulations.****GHS HAZARD CLASSIFICATION**

## SYSTEM CHECK SOLUTION WARNING



H317	May cause an allergic skin reaction.
H412	Harmful to aquatic life with long lasting effects.
P273	Avoid release to the environment.
P280	Wear protective gloves, protective clothing and eye/face protection.
P333+P313	If skin irritation or rash occurs: Get medical advice/attention.
P362+P364	Take off contaminated clothing and wash it before use.

reaction mass of: 5-chloro-2-methyl-4-isothiazolin-3-one [EC# 247-500-7] and 2-methyl-4-isothiazolin-3-one [EC# 220-239-6](3:1) < 0.05%

SDS

Safety Data Sheet is available at [beckmancoulter.com/techdocs](http://beckmancoulter.com/techdocs)

## TESTING PROCEDURE(S)

### DIRECTIONS FOR USE

Refer to the appropriate system Manuals and/or Help system for detailed instructions.

### ADDITIONAL INFORMATION

For a patient/user/third party in the European Union and in countries with identical regulatory regime (Regulation 2017/746/EU on In vitro Diagnostic Medical Devices); if, during the use of this device or as a result of its use, a serious incident has occurred, please report it to the manufacturer and/or its authorized representative and to your national authority.

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May be covered by one or more pat. -see [www.beckmancoulter.com/patents](http://www.beckmancoulter.com/patents).

### REVISION HISTORY

#### Revision L

IFU updated to add Dutch, Finnish, Macedonian, Traditional Chinese, and Estonian

#### Revision M

New release of IVDR compliant IFU.

#### Revision N

Added Translations.

## Revision P

Updated ProClin trademark statement.

## Revision R


Added Translations.

## SYMBOLS KEY

Glossary of Symbols is available at [beckmancoulter.com/techdocs](http://beckmancoulter.com/techdocs) (document number C02724).

EC	REP
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