Neutron™ Needlefree Catheter Patency Device
Microbial Ingress Test
Report of a study commissioned by ICU Medical Inc. and conducted by Alcami

OBJECTIVE
A protocol was developed and executed by Alcami (Durham, North Carolina), a world-class contract development and manufacturing organization (CDMO), to evaluate microbial ingress on the Neutron needlefree catheter patency device over a 7-day period.

METHODS
Devices were subjected to a repeated sequence of multiple disinfections and multiple activations. This was followed by inoculation with a microbe known to cause central line-associated bloodstream infections. Devices were then disinfected, and a flush fluid was collected for assay.

MICROBIAL SELECTION AND CULTURE PREPARATION
Staphylococcus aureus was selected for this performance test, as it is considered one of the most common infectious pathogens of concern in the clinical setting.

Test organisms from the American Type Culture Collection (ATCC), a nationally recognized culture organization, were inoculated from stock cultures and incubated at a temperature of 30–35°C to create the $1-5 \times 10^3$ colony-forming units (CFU) minimum inoculation solution.

MICROBIAL PERFORMANCE TEST
Testing was performed in a controlled environment. The test procedure consisted of the following sequential steps: the clear septum of each of the 20 test devices was inoculated with $1-5 \times 10^3$ CFU of Staphylococcus aureus, followed by 1 minute of drying in ambient conditions, then the top of the surface of the connector was disinfected using a 70% isopropyl alcohol pad for 3 seconds and allowed to dry. The device was activated using a syringe with 10 mL of sterile 0.9% saline. The fluid was then collected, filtered, and plated to determine growth at 30–35 °C for 2–3 days. Each of the 20 test devices was activated 20 times per day for 7 days, totaling 140 activations per device.

CONTROLS
Two positive controls and two negative controls followed the same test procedure as the devices for each type of test organism. The positive controls demonstrated the viability of the challenge organisms throughout the test period. The positive controls were not disinfected after inoculation with the test organism, and the negative controls were not inoculated.

Positive monitor recovery control titers demonstrated that the minimum challenge level meets the United States FDA 2008 intravascular administration set guidance for microbial ingress.

ACCEPTANCE CRITERIA
Negative controls must be negative for the challenge organisms. Positive controls must be representative of the test setup. All study acceptance criteria were met.
RESULTS
Analysis of growth on plated fluid samples found that:
› All devices tested negative for growth (0 CFU), meaning there was no microbial contamination of the device fluid path.
› Negative controls were negative for growth, and positive controls tested positive for growth.

Activations per test organism:
› Final total activations/test device: (20 activations) x (7 days) = 140

<table>
<thead>
<tr>
<th></th>
<th>Test Devices 1–20</th>
<th>Positive Controls</th>
<th>Negative Controls</th>
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<tbody>
<tr>
<td>Day 1</td>
<td>0 CFU</td>
<td>++</td>
<td>0 CFU</td>
</tr>
<tr>
<td>Day 2</td>
<td>0 CFU</td>
<td>++</td>
<td>0 CFU</td>
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<tr>
<td>Day 3</td>
<td>0 CFU</td>
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<td>Day 4</td>
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<td>0 CFU</td>
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<tr>
<td>Day 7</td>
<td>0 CFU</td>
<td>++</td>
<td>0 CFU</td>
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CONCLUSIONS
The neutron needlefree catheter patency device passed the microbial ingress test, and no contamination was found after repeated use, inoculation, and disinfection with 70% isopropyl alcohol. Therefore, the Neutron needlefree catheter patency device is suitable for use for 7 days on the appropriate vascular access or infusion device.