Prior to Use Storage and Patient Administration Periods for CSPs

USP - NF 27 <797> Pharmaceutical Compounding - Sterile Preparations

The subject of “beyond use dating” for compounded sterile preparations published in USP-NF 27 <797> has been a continuing source of questions and concern for pharmacies. Questions regarding storage times, sterility test requirements and administration times are commonplace in the industry. <797> has specific requirements that must be met:

- Appropriateness of containers to preserve sterility and strength
- Before-administration storage periods
- Patient administration times

This technical bulletin seeks to clarify the USP issues and provide the pharmacist with a clear understanding of and a rationale for I-Flow*’s position on the subject as well as a list of available I-Flow* references for chemical and microbiological studies performed in support of this position.

USP-NF 27 <797> is concerned with two issues, risk of microbiological contamination and chemical stability. In the section titled RESPONSIBILITY OF COMPOUNDING PERSONNEL, item number 8, the USP states “Packaging selected for CSPs is appropriate to preserve the sterility and strength until the beyond-use date”.

To satisfy this requirement I-Flow* has performed protocol Microbial Ingress Testing of the I-Flow* ON-Q* Device (PSI-04073). In this study, ON-Q* pumps were filled with a microbial growth supporting medium and then immersed for 24 hours in a circulating bath containing Brevundimonas diminuta. Brevundimonas is the bacteria of choice for filter bacteria retention studies and packaging ingress studies due to its extremely small size and motility. Following the exposure to the bacterial challenge solution, the filled devices were incubated at 20-25°C for 7 days. At the end of the seven days the pumps were emptied and the growth medium inspected for bacterial contamination. No bacterial contamination was present. This study clearly demonstrates that the ON-Q* pump meets the USP requirements for an appropriate container to maintain product integrity and content sterility. Item number 11 of the same USP section states, “Beyond-use dates are assigned based on direct testing or extrapolation from reliable literature sources and other documentation”.

The referenced testing performed by I-Flow*, at an independent contract laboratory, meets the requirement of “other documentation” as stated in <797>.

The ON-Q* Pain Relief System is intended to provide continuous delivery of medication (such as local anesthetics) to or around surgical wound sites and/or in close proximity to nerves for preoperative, perioperative, and postoperative regional anesthesia and/or pain management. In USP-NF 27 <797> under Examples of Medium-Risk Compounding, item number 3 provides for a description of a medium risk device such as the ON-Q* pump, and states “Filling of reservoirs of injection and infusion devices with volumes of sterile drug solutions that will be administered over several days at ambient temperatures between 25° and 40°”.

Devices filled in the surgical suite are classified as high-risk is because the air quality in the surgical suite does not meet the standard of ISO Class 5. USP <797> specifically states that if a device is filled in an area with an environmental classification of greater than ISO Class 5, that the device must be classified as a high risk device. Surgical suites are typically operated with a controlled environment greater than ISO Class 5. The USP has provided for classification of CSPs that will be administered over several days.

Since USP has chosen to include devices such as the ON-Q* pump as medium-risk, the storage conditions before administration must meet the USP guidelines (see General Notices and Requirements). “For a medium-risk preparation, in absence of passing a sterility test, the storage periods cannot exceed the following time periods:

- Before administration, the CSPs are properly stored and are exposed for not more than 30 hours at controlled room temperature,
- For not more than 9 days at cold temperatures,
- And for 45 days in solid frozen state at -20° or colder.”

The General Notices Section of USP lists requirements for temperature monitoring, etc. Cold temperatures are defined by the USP as 2-8°C. For a high-risk CSP the room temperature storage period drops to < 24 hours and the cold temperature time to < 3 days.
These sections of the USP are clearly discussing the risk of microbial contamination and the ability of the CSP to remain sterile for both the storage period and the administration period. To this end, I-Flow* has performed protocol Microbial Storage Stability of the I-Flow* ON-Q* Device (PSI-04063). In this study, 10 ON-Q* pumps were filled with a microbial growth supporting medium and then stored at 2-8 °C for seven days. At the completion of the seven days, the pumps were transferred to room temperature storage (20-25 °C) for an additional 14 days. Another group of 10 pumps was stored at room temperature only for 14 days. Appropriate media growth promotion studies and bacteriostasis/fungistasis studies were performed with this stability study. At the completion of the prescribed storage periods the pumps were examined for microbial growth. In all cases all pumps were sterile. This study demonstrates that the ON-Q* pump, when filled under aseptic conditions, by appropriately trained personnel, maintains its contents sterile for time periods that exceed the USP pre-administration storage periods. The USP states in the section on DETERMINING BEYOND-USE DATES that: “Compounding personnel may consult the manufacturer of particular products for advice on assigning beyond-use dates based on chemical and physical stability parameters.”

I-Flow* has also performed the following protocols:

- Medium Risk Media Fill Protocol of the I-Flow* ON-Q* Device using the BAXA Repeater Pump (PSI-04062)
- Medium Risk Media Fill Protocol of the I-Flow* ON-Q* Device (PSI-04059)
- High Risk Media Fill Protocol of the I-Flow* ON-Q* Device (PSI-04061)

In these three studies the media fill qualifications recommendations of USP-NF 27 <797> were followed. The final container filled and incubated for 14 days was the ON-Q* pump. These three studies provide additional documentation as to the suitability of the ON-Q* pump as a final container for CSPs.

In addition, I-Flow* has performed chemical stability studies for the most commonly used local anesthetics. These studies demonstrate that the anesthetics are stable for periods that exceed the USP prior to administration storage periods. Chemical stability of the pump has been verified with the following drugs:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Concentration</th>
<th>Storage Time at RT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bupivacaine</td>
<td>0.25 – 0.5%</td>
<td>30 days</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>1%</td>
<td>30 days</td>
</tr>
<tr>
<td>Ropivacaine</td>
<td>0.2%</td>
<td>30 days</td>
</tr>
</tbody>
</table>

Summary

- Microbial ingress studies and media fill challenges performed by I-Flow* have demonstrated the ON-Q* pump is an appropriate container for CSP storage.
- The USP makes a clear difference between “before-administration” storage periods and patient administration periods.
- Studies performed by I-Flow* have demonstrated both microbiological and chemical stability of the filled ON-Q* pump for time periods that exceed the USP requirements for before-administration storage.

There are inherent risks in all medical devices. Please refer to the product labeling for Indications, Cautions, Warnings and Contraindications. Failure to follow the product labeling could directly impact patient safety. Physician is responsible for prescribing and administering medications per instructions provided by the drug manufacturer. Refer to www.iflo.com for additional product safety Technical Bulletins.

Please contact the Clinical Services Department at 800-444-2728 or 949-923-2400 if you have any questions regarding this information.