Review of USP Chapter <1223> Validation of Alternative Microbiological Methods Presented at the 2007 Global Conference on Pharmaceutical Microbiology

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Introduction

This article is based on a presentation given by the author on USP chapter <1223> Validation of Alternative Microbiological Methods during the 2007 PDA Global Conference on Pharmaceutical Microbiology, held in Bethesda, Md, October 30 – November 1, 2007. The author has had a few further thoughts on the subject that will be given in the conclusion of this article.

Several regulatory documents have a bearing on our understanding of USP chapter <1223>. The first of these is the draft guidance document on Comparability Protocols [1]. Comparability Protocols are used in part to address modifications to methods used in previous filings to the FDA. This method has been identified by the FDA as the preferred method for introducing alternate microbiological methods into an approved product submission.

GMPs for the 21st Century – A Risk-based Approach (Sept. 29, 2004) is also relevant to this discussion [2]. This document draws together many regulatory guidance documents relevant to GMP, and provides web-based links to access the information. Two guidance documents referred to in this 2004 report deserve special attention: “Guidance for Industry: Sterile Drug Products Produced by Aseptic Processing — Current Good Manufacturing Practice” and “Guidance for Industry PAT — A Framework for Innovative Pharmaceutical Development, Manufacturing and Quality Assurance.” The potential use of PAT (process analytical technology) exists for alternate microbiological methods, particularly rapid ones as microbiological analyses account for the majority of the manufacturing timeline [3-6].

PDA Technical Report 33 (PDA TR #33) “Evaluation, Validation and Implementation of New Microbiological Testing Methods” is considered to be, in a sense, the predecessor of chapter <1223>. It considered the validation of microbiological methods to be necessarily approached in a different manner from the validation of chemical methods. Sections from TR 33 delineated were: Vendor and Method Requirements, Designing the Validation Plan, Installation Qualification (IQ), Operational Qualification (OQ) and Performance Qualification (PQ).

Finally, any discussion of validation must address the statistics of the system. This is a particularly difficult subject with microbiology as the numbers obtained generally do not follow a “normal” distribution. Given the importance of the appropriate application of statistical methods well-suited to the data and ana-
Analytical needs, microbiologists should feel free to consult any of the many available references, including the three references that were provided [7-9]. USP Chapter <1010> Analytical Data—Interpretation and Treatment [10] is also a good reference.

**Development of Compendial Chapters**

There are currently two official compendial chapters covering the validation of alternative microbiological methods. These are the European Pharmacopeia's "5.1.6 Alternative Methods for Control of Microbiological Quality" [11] and the USP's "<1223> Validation of Alternative Microbiological Methods" [12]. The publication history of chapter <1223> is enlightening. The first version of the chapter appeared in Pharmacopeial Forum (PF) in 2002 [13]. Many comments received concerned the chapter being confusing and difficult to use. The USP expert committee (Analytical Microbiology during the 2000-2005 USP cycle of revision) in charge of the chapter responded to those comments by preparing another version that was published one year later [14]. The nature of the comments was virtually unchanged. Therefore, the experts decided a major rewrite was in order, and the next version, published in 2005 [15], had the entire section pertaining to validation of microbiological identification removed. This was done to simplify the topic and focus the chapter [16]. It was this version that became official in the 2nd Supplement.

**<1223> Validation of Alternative Microbiological Methods**

There are two critical components contained in the introduction of the chapter. The first pertains to the comparative nature of the validation of an alternative method. This differs substantially from the validation methods described in USP chapter <1225> Validation of Compendial Procedures. In the case of <1225>, the validation is not comparative in nature. That is to say, one is not validating a method as an alternative to a compendial method. You are rather validating a critical aspect of the alternate method against that used in the compendial method (method of enumeration, demonstration of viable cells on a membrane, etc.). The critical importance of this is the question to be addressed in the validation of an alternative method: does the alternate method yield results equivalent to or better than the results generated by the compendial method [17]?

The second critical component discussed in the introduction pertains to method variability. Microbiological methods typically have much higher levels of variability associated with them. Relative standard deviations (RSD) of 15 to 35 percent are common, whereas for analytical chemistry methods, typical RSDs range between one and three percent. The RSD is defined as the standard deviation divided by the mean.
The currently official version of the chapter addresses the validation of two types of microbiological methods: Qualitative tests and quantitative tests. A qualitative test is concerned with only one thing: are any viable microorganisms present. This implies that the result from such a test is binary, as in something is there or isn't. The second type of method covered is quantitative. In such a method, one is clearly interested in measuring a range of microorganisms (enumerating them). The list of validation elements to address differs depending upon the type of test.

Quantitative microbiological tests require that the following validation parameters be addressed:
1. Accuracy: The closeness of the test results obtained by the alternate test method to the value obtained by the traditional method.
2. Precision: The degree of agreement among individual test results when the procedure is applied repeatedly to multiple samplings of suspensions of laboratory microorganisms across the range of the test.
3. Specificity: The ability to detect a panel of microorganisms suitable to demonstrate that the method is fit for its intended purpose.
4. Limit of Quantification: The lowest number of microorganisms that can be accurately counted.
5. Linearity: The ability to produce results that are proportional to the concentration of microorganisms present in the sample within a given range.
6. Range: The interval between the upper and lower levels of microorganisms that have been demonstrated to be determined with precision, accuracy and linearity.
7. Limit of Detection, Ruggenedness and Robustness, with definitions equivalent to those for qualitative tests.

Qualitative microbiological methods require considerably fewer data parameters be addressed. These parameters are:
1. Specificity: The ability to detect a range of microorganisms.
2. Limit of Detection: Determination of the lowest amount detectable, that is to say, the minimum number that will generate a signal above the background noise.
3. Ruggenedness: The resistance to influences of operational and environmental variables (often considered random effects).
4. Robustness: The capacity to remain unaffected by small but deliberate (nonrandom) variations in method parameters.

There are a couple of general notes about the compendial methods. First, the microorganisms that should be included in the method validation should represent species of importance to one's specific requirements. If one finds a particular species during environmental monitoring that is not included in pharmacopeial lists, its exclusion from the pharmacopeia does not obviate the need for tracking this particular species. Second, validation of an alternative method to a compendial method is comparative in nature. Therefore, all data elements listed above should involve comparisons between the results of the alternative and compendial methods. The requirement is that the alternative method results be equivalent or better than the compendial method results.

Differences Between the EP and the USP Chapters

The two compendial chapters are similar in many respects. The differences are substantial in other respects. For example, the USP chapter does not address the validation of microbial identification methods, whereas the EP chapter does. The USP chapter has dropped the parameters of accuracy and precision from the list to be addressed for qualitative methods. This is because the USP considers these parameters to be operationally no different from the limit of detection. There are other differ-
ences of a more statistical nature.

Conclusion

My impression from talking to many people about this chapter is that it does serve a useful function. Recently, I have heard some interesting concerns about this chapter. One concern is that perhaps this chapter is slanted too much towards rapid microbiological methods. This may be reflected in portions of the chapter text pertaining to robustness and ruggedness. Here, it is suggested that much of the responsibility for the validation of these parameters rests with the vendor of the method. One could reasonably infer the use of rapid methods here, since most of these are instrument-dependent. From a compendial perspective, this underplays the importance of having potentially affected parties of a new method try the method using their own test articles, and then reporting problems to the USP. Another concern pertains to the importance of distinguishing between validation and verification (demonstration of method suitability). It is considered of utmost importance to be able to state prior to any validation exercise just what it is that you are attempting to validate. That is to say, what is the intended purpose of the method for which you are undertaking validation? One final concern pertains to the importance of using appropriate statistics. Many statistical methods make certain assumptions about underlying data distributions that should be verified. Also, some methods are better suited to smaller sample sizes than others. Always remember that there are always several ways to skin the microbiological cat statistically.

References

11. Ph. Eur. 2006. 5.1.6 Alternative Methods for Control of Microbiological Quality. Pharm Eur. 5.5:4131-4142
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He began his industrial career at Bausch & Lomb, where he managed the microbiology and toxicology areas. He also managed the biostatistics area in the clinical affairs group. His next industrial stop was as the leader of the biological science section of the Advanced Concepts area at the cosmetics company, Revlon. Numerous new analytical methods were developed in the pursuit of these actives. He was also a coinventor on two patents.

Porter completed his doctoral work in Zoology at the University of California, Berkeley, with an emphasis in comparative endocrinology. His postdoctoral work included studies of the molecular biology of avian gonadotropic hormones and mammalian parathyroid hormone and its receptors.

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