Modernisation of the spectroscopic General Chapters in the United States Pharmacopeia (USP)

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As part of the USP–NF modernisation process, the General Chapters (Chemical Analysis) Expert Committee has and continues to review collections of chapters in a holistic approach in order to align concepts and content to ensure consistency.\(^{1}\) These revisions are intended to ensure that the scientific expectations defined in the chapters are aligned with current best practices to ensure suitability for use when the appropriate measurements are executed. To these ends, general chapters currently are grouped into two parts: <1000 chapters (i.e., those numbered below 1000), where the focus, as stated in the General Notices, is on content that may be required if called for in a monograph, and >1000 chapters (i.e., those numbered 1000–1999), where these chapters give information, definition or description.

The purpose of spectroscopic General Chapters is to provide the basis for establishing that an instrument or system is "suitable for use" in a USP monograph. This "suitability for use" is usually related to the technical performance specification of the instrument or system and represents a minimum standard. However, on its own, compliance to specification cannot provide overall "suitability for use" and best practice guidance provides the holistic component.

Hence "suitability for use" has two distinct aspects:
- Compliance with technical performance characteristics on the instrument/system itself,
- Operational best practices for an instrument/system when employed as part of a procedure stated in a monograph.

The instrument/system requirements currently described in General Chapter <851>, Spectrophotometry and Light Scattering, covered many spectroscopic techniques and had remained essentially unchanged over many years. Therefore, in the USP Review Cycle 2010 to 2015, it was decided to generate specific pairs of chapters for each of the main spectroscopic types. The below 1000 chapters would define minimum standards for compliance for use in a monograph, the paired above 1000 chapter would give theory, guidance and recommendations for best analytical practices. In general, below 1000 chapters include procedures, instrument qualification and validation/verification sections. Each section involves an assessment of method-specific requirements to ensure the suitability of the system and related measurements. General Chapter <851>, itself would be deleted after all the new chapters were approved.

However, it should be noted that this holistic approach has already been applied to two other spectroscopic chapters, currently in USP 37 2014, Nuclear Magnetic Resonance Spectroscopy <761> and its paired best practices chapter Applications of Nuclear Magnetic Resonance Spectroscopy <1761> and Mass Spectrometry <736> and its paired best practices chapter Applications of Mass Spectrometry <1736>.

The objective was to articulate the basic elements required before and during the execution of a spectroscopic measurement to ensure that suitability for use has been confirmed. As an example, the validation/verification requirements for spectroscopic methods are specific for the procedures in the chapters.

Many of these concepts are not new to USP. The changes to the spectroscopy chapters are consistent with those defined in Chromatography <621>; that is, the user must demonstrate that the system is "suitable for use". It is important that before execution of a measurement the analyst should demonstrate method-specific suitability for the instrument to ensure the integrity of the measurement.

The proposed new below 1000 chapters have a common structure irrespective of the spectroscopic technique namely:
A short introduction to the particular technique and a link to the associated best practices chapter above 1000,

- Qualification requirements for IQ & OQ with recommended procedures and the link to Analytical Instrument Qualification General Chapter <1058>,
- Specific critical attributes of analytical procedures involving the particular technique,
- Validation and verification needs as required by General Chapters <1225> and <1226> for alternative methods being developed for pharmacopeial monographs.

Analytical instrument qualification (AIQ)

Analytical Instrument Qualification <1058> provides a risk-based framework for generating documented evidence over the lifecycle that the instrument or system is, and remains suitable for use in a monograph. <1058> is currently in revision with the intent of extending it to be compatible with GAMP 5² and GAMP Laboratory Good Practice Guide 2nd Edition.³ An article outlining proposals for change has been published.⁴ In addition, the extension proposal has been published.⁵

AIQ is currently defined as the collection of documented evidence that an instrument performs suitably for its intended purpose. Use of a qualified instrument in analyses contributes to confidence in the validity of generated data. Chapter <1058> further characterises the steps in instrument qualification during AIQ:

- Design Qualification (DQ): The documented collection of activities that define the functional and operational specifications of the instrument and criteria for vendor selection based on the instrument’s intended purpose.
- Installation Qualification (IQ): The documented collection of activities necessary to establish that an instrument is delivered as designed and specified, that it is properly installed in the selected environment and that this environment is suitable for the instrument.
- Operational Qualification (OQ): The documented collection of activities necessary to demonstrate that an instrument will function according to its operational specification in the selected environment. The extent of OQ testing that an instrument undergoes depends on its intended applications. Therefore, no specific OQ tests for any instrument or application are offered in <1058>. It should be noted that routine analytical tests do not constitute OQ testing.
- Performance Qualification (PQ): The documented collection of activities necessary to demonstrate that an instrument consistently performs according to the specifications defined by the user and is appropriate for the intended use. PQ tests may resemble those performed during OQ, but the specifications for their results may be set differently if required. Procedures for OQ and PQ testing that comply with regulatory expectations are not well defined and are subject to different interpretations.

The qualification tests and acceptance criteria are included in the below 1000 chapters to establish minimum pharmacopeial requirements for instrument performance. Ultimately, the use of a qualified instrument contributes to confidence in the validity of analytical data used to guarantee patient safety. The chapters will stipulate that manufacturer-recommended OQ and PQ tests and acceptance criteria may be used if deemed scientifically equivalent. This will prevent the AIQ tests in the <1000 chapters from becoming excessively prescriptive.

Validation and verification

As part of the overall revision process for spectroscopy general chapters, validation requirements also are being aligned with current standards. The validation of spectroscopic procedures
is covered under both the ICH guidelines and Validation of Compendial Procedures <1225> even though the predominant theme for each of these documents historically has been validation of chromatographic procedures. The objective of the current modernisation effort is to include validation sections in the various spectroscopy chapters thus providing both general direction for the validation of procedures based on these methods and specific validation requirements that are important for specific spectroscopic approaches.

 mass spectrometry, <1736> and applications of nuclear magnetic resonance spectroscopy, <1773>.

Revision of General Chapter <851>
As noted earlier, the current General Chapter <851> covers multiple spectroscopic types, and has not been subject to a major revision for many years. Whilst the general spectroscopic principles detailed in the current <851> describe the fundamental requirements, it was felt that they are too vague to meet the extensive and increasing use of instrumental spectroscopic techniques, and the associated current (and future) regulatory requirements. In addition the proliferation of the application of new spectroscopic techniques makes revision of the current <851> essential.

The following new spectroscopic chapter pairs have been developed in accordance with the approaches outlined previously:
- Atomic Absorption Spectroscopy, <852> & <1852>
- Fluorescence Spectroscopy, <853> & <1853>
- Mid-Infrared Spectroscopy, <854> & <1854>
- Turbidimetry and Nephelometry, <855>
- Near Infrared (NIR) Spectroscopy, <856> and <1856>
- Ultraviolet-Visible Spectroscopy, <857> & <1857>
- Raman Spectroscopy, <858> & <1858>

These together with other spectroscopic chapters are shown in relationship to the current <851> in Figure 1, “Revision Paths for Spectroscopy General Chapters.” Note that neither NIR nor Raman spectroscopies are adequately covered in <851> and had previously only above 1000 information chapters. This situation has now been rationalised.

Revision history during the 2010–2015 cycle
The drafts and revisions for these Spectroscopy General Chapters were
After extensive drafting and review, the new Chemometrics Chapter will be very important. In addition CD/ORD, XRF and XRD may be included in the cycle. At present there are no definite plans for a companion chapter <1855> for <1855> Nephelometry, Turbidimetry and Visual Comparison, although it may well happen especially as there is growing interest in these techniques from a bioanalytical perspective. In addition there is likely to be a revision of Analytical Instrument Qualification General Chapter <1058>.

We hope that practicing spectroscopists will make the time and effort to keep in touch with these developments and comment. Over the years we have heard many practitioners complaining about the appropriateness of monographs and General Chapters. You now have the chance to assist with the change process. We strongly encourage you to use it!

References