GUIDELINES ON SUBMISSION OF DOCUMENTATION FOR A MULTISOURCE ( GENERIC) FINISHED PHARMACEUTICAL PRODUCT: QUALITY PART

DRAFT FOR COMMENT

Should you have any comments on the attached text, please send these to Dr Sabine Kopp, Manager, Medicines Quality Assurance Programme, Quality Assurance and Safety: Medicines, World Health Organization, 1211 Geneva 27, Switzerland; e-mail: kopps@who.int; fax: (+41 22) 791 4730 (kopps@who.int) and to Ms Marie Gaspard (gaspardm@who.int), by 25 April 2013.

Working documents are sent out electronically and they will also be placed on the Medicines web site for comment. If you do not already receive directly our draft guidelines please let us have your e-mail address (to bonnyw@who.int) and we will add it to our electronic mailing list.

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1. INTRODUCTION

1.1 Background

Through the International Conference on Harmonisation (ICH) process, considerable harmonization has been achieved on the organization for the Quality Module of registration documents with the issuance of the Common Technical Document (CTD) - Quality (ICH M4Q) guideline. This recommended format in the M4Q guideline for the quality information of registration applications has become widely accepted by regulatory authorities both within and beyond the ICH Regions.

This document, Guidelines on submission of documentation for a multisource (generic) finished pharmaceutical product: quality part, provides recommendations on the quality information for active pharmaceutical ingredients (APIs) and finished pharmaceutical products (FPPs) that should be submitted to national medicines regulatory authorities (NMRAs) to support product dossiers (PDs).

Alternate approaches to the principles and practices described in this document may be acceptable provided they are supported by adequate scientific justification. It is also important to note that the NMRAs may request information or material, or define conditions not specifically described in this guidance, in order to adequately assess the quality of a pharmaceutical product.

1.2 Objectives

These guidelines are intended to:

- assist applicants on the preparation of the Quality Module of PDs for multisource products by providing clear general guidance on the format of these dossiers;
- adopt the modular format of the Common Technical Document - Quality (M4Q) as developed by ICH;
- provide guidance on the technical and other general data requirements.

These measures are intended to promote effective and efficient processes for the development of these PDs by applicants and the subsequent assessment procedures by NMRAs.

1.3 Scope

These guidelines apply to PDs for multisource pharmaceutical products containing existing APIs of synthetic or semi-synthetic origin. For the purposes of these guidelines an existing API is one that has been previously approved through a finished product by a stringent regulatory authority\(^1\) or WHO. Fermentation, biological, biotechnological and herbal APIs are covered by other guidelines.

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\(^1\) Stringent regulatory authority (SRA): a regulatory authority which is:

- a member of the International Conference on Harmonisation (ICH) (as specified on www.ich.org);
- or

an ICH observer, being the European Free Trade Association (EFTA), as represented by Swiss Medic, and Health Canada (as may be updated from time to time).
1.4 General principles

To facilitate the preparation of the PD these guidelines are organized in accordance with the structure of the Common Technical Document – Quality (M4Q) guideline, as developed by ICH.

The text of the M4Q (CTD-Q) guideline has been restated in these guidelines in **bold text**, verbatim, with minor modifications to accommodate WHO terminology and include certain text that would be appropriate for multisource pharmaceutical products, notably:

- “Drug substance” is replaced with “active pharmaceutical ingredient” or “API”;
- “Drug product” is replaced with “finished pharmaceutical product” or “FPP”;
- “application” is replaced with “product dossier” or “PD”;
- “combination product” is replaced with “fixed-dose combination” or “FDC”;
- “clinical batches” is replaced with “comparative bioavailability or biowaiver batches”.

Following the **bold** text of the M4Q (CTD-Q) guideline additional guidance is provided in plain text to easily distinguish from the ICH text and is included to provide further clarity on general expectations for the content of PDs. This approach is intended to facilitate the identification and origin of the text in the guidelines (i.e. from ICH or additional information).

The content of these guidelines should be read in conjunction with relevant information described in other existing NMRA guidelines, WHO guidelines or ICH reference documents and guidelines. The quality of existing APIs and corresponding multisource products should not be inferior to new APIs and innovator FPPs. Therefore, the principles of the ICH guidelines that are referenced throughout these and other guidelines may also equally apply to existing APIs and multisource products.

Scientific literature may be appropriate to fulfil the requirements for some of the information or parameters outlined in these guidelines (e.g. qualification of specified identified impurities). Furthermore the requirements outlined in certain sections may not be applicable for the proposed API or FPP. In these situations a summary and the full reference to the scientific literature should be provided or the non-applicability of the requested information should be clearly indicated as such with an accompanying explanatory note.

1.5 Guidance on format

For the format and presentation of the PD recommendations outlined in the WHO general filing guidelines: *Guidelines on submission of documentation for a multisource (generic) finished pharmaceutical product: preparation of product dossiers in common technical document format* (TRS 961, Annex 15) may be followed, with the understanding that module 1 contains regionally required information and therefore the required contents will vary depending on the NMRA to which the PD is filed.

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*time*; or

*a regulatory authority associated with an ICH member through a legally-binding, mutual recognition agreement including Australia, Iceland, Liechtenstein and Norway (as may be updated from time to time).*
There may be a number of instances where repeated sections can be considered appropriate. Whenever a section is repeated it should be made clear what the section refers to by creating a distinguishing title in parentheses following the M4Q (CTD-Q) guideline heading, e.g. 3.2.S Drug substance (or API) (name, Manufacturer A).

Following are recommendations for the presentation of the information in the Quality Module for different scenarios that may be encountered:

- The Open part (non-proprietary information) of each DMF should always be included in its entirety in the PD, as an annex to 3.2.S:

- For an FPP containing more than one API: one complete “3.2.S” section should be provided for one API, followed by other complete “3.2.S” sections for each other API;

- For an API from multiple manufacturers: one complete “3.2.S” section should be provided for the API from one manufacturer, followed by other complete “3.2.S” sections for each other API manufacturer;

- For an FPP with multiple strengths (e.g. 10, 50, 100 mg): one complete “3.2.P” section should be provided with the information for the different strengths provided within the subsections. One complete copy of the PD should be provided for each FPP strength;

- For an FPP with multiple container closure systems (e.g. bottles and unit dose blisters): one complete “3.2.P” section should be provided with the information for the different presentations provided within the subsections;

- For multiple FPPs (e.g. tablets and a parenteral product): a separate dossier is required for each FPP;

- For an FPP supplied with reconstitution diluent(s), one complete “3.2.P” section should be provided for the FPP, followed by the information on the diluent(s) in a separate part “3.2.P”, as appropriate;

- For a co-blistered FPP, one complete “3.2.P” section should be provided for each product.
2. GLOSSARY

The definitions provided below apply to the words and phrases used in these guidelines.
Although an effort has been made to use standard definitions as far as possible, they may have different meanings in other contexts and documents. The following definitions are provided to facilitate interpretation of the guidelines.

active pharmaceutical ingredient (API)
Any substance or mixture of substances intended to be used in the manufacture of a pharmaceutical dosage form, and that, when so used, becomes an active ingredient of that pharmaceutical dosage form. Such substances are intended to furnish pharmacological activity or other direct effect in the diagnosis, cure, mitigation, treatment or prevention of disease, or to affect the structure and function of the body (WHO Technical Report Series, No. 961, Annex 10, 2011).

API starting material
A raw material, intermediate, or an API that is used in the production of an API and that is incorporated as a significant structural fragment into the structure of the API. An API starting material can be an article of commerce, a material purchased from one or more suppliers under contract or commercial agreement, or produced in-house (ICH Q7). See also starting materials for synthesis.

applicant
The person or company who submits an application for marketing authorization of a new pharmaceutical product, an update to an existing marketing authorization or a variation to an existing market authorization. (WHO Technical Report Series, No. 929, Annex 5, 2005).

BCS highly soluble
An API for which the highest dose recommended by WHO (if the API appears on the WHO Model List of Essential Medicines) or highest dose strength available on the market as an oral solid dosage form (if the API does not appear on the WHO Model List of Essential Medicines) is soluble in 250 ml or less of aqueous media over the pH range of 1.2–6.8 at 37°C (WHO Technical Report Series, No. 937, Annex 7, 2006).

commitment batches
Production batches of an API or FPP for which the stability studies are initiated or completed post-approval through a commitment made in a regulatory application (WHO Technical Report Series, No. 953, Annex 2, 2009).

comparator product (reference product)
A pharmaceutical product with which the generic product is intended to be interchangeable in clinical practice. The comparator or reference product will normally be the innovator product for which efficacy, safety and quality have been established (WHO Technical Report Series, No. 937, Annex 7, 2006).

existing API
An API that is not considered a new active substance, that has been previously approved through a finished product by a stringent regulatory authority or WHO, but requires the filing of a dossier. This would include, for example, new PDs and variations to multisource products.

finished pharmaceutical product (FPP)
A finished dosage form of a pharmaceutical product, which has undergone all stages of manufacture, including packaging in its final container and labelling (WHO Technical Report Series, No. 961, Annex 10, 2011).

**innovator pharmaceutical product**
Generally the pharmaceutical product that was first authorized for marketing (normally as a patented product) on the basis of documentation of efficacy, safety and quality (WHO Technical Report Series, No. 937, Annex 7, 2006).

**manufacturer**
A company that carries out operations such as production, packaging, repackaging, labelling and relabelling of pharmaceuticals. (WHO Technical Report Series, No. 961, Annex 3, 2011).

**multisource (generic) pharmaceutical products**
Pharmacologically equivalent or pharmaceutically alternative products that may or may not be therapeutically equivalent. Multisource pharmaceutical products that are therapeutically equivalent are interchangeable (WHO Technical Report Series, No. 937, Annex 7, 2006).

**officially recognized pharmacopoeia (or compendia)**
Those pharmacopoeias whose standards are officially recognized by an NMRA. These may be national, regional or international pharmacopoeia, at the discretion of the NMRA.

**ongoing stability study**
The study carried out by the manufacturer on production batches according to a predetermined schedule in order to monitor, confirm and extend the projected retest period (or shelf-life) of the API, or confirm or extend the shelf-life of the FPP (WHO Technical Report Series, No. 953, Annex 2, 2009).

**pilot-scale batch**
A batch of an API or FPP manufactured by a procedure fully representative of and simulating that to be applied to a full production-scale batch. For example, for solid oral dosage forms, a pilot scale is generally, at a minimum, one-tenth that of a full production scale or 100,000 tablets or capsules, whichever is the larger; unless otherwise adequately justified (WHO Technical Report Series, No. 953, Annex 2, 2009).

**primary batch**
A batch of an API or FPP used in a stability study, from which stability data are submitted in a registration application for the purpose of establishing a re-test period or shelf-life (WHO Technical Report Series, No. 953, Annex 2, 2009).

**production batch**
A batch of an API or FPP manufactured at production scale by using production equipment in a production facility as specified in the application (WHO Technical Report Series, No. 953, Annex 2, 2009).

**starting materials for synthesis**
Materials that mark the beginning of the manufacturing process as described in an application or in a DMF. A starting material for a synthetic API is a chemical compound of defined molecular structure that contributes to the structure of the API. See also API starting material.
3. QUALITY SUMMARIES

3.1 Module 2.3: Quality overall summary – product dossiers (QOS-PD)

The Quality Overall Summary (QOS) is a summary that follows the scope and the outline of the Body of Data in Module 3. The QOS should not include information, data or justification that was not already included in Module 3 or in other parts of the CTD.

The QOS should include sufficient information from each section to provide the Quality assessor with an overview of Module 3. The QOS should also emphasise critical key parameters of the product and provide, for instance, justification in cases where guidelines were not followed. The QOS should include a discussion of key issues that integrates information from sections in the Quality Module and supporting information from other Modules (e.g., qualification of impurities via toxicological studies), including cross-referencing to volume and page number in other Modules.

The WHO Quality overall summary – product dossiers (QOS-PD) template or the QOS template associated with the intended NMRA, if available, should be completed for multisource pharmaceutical products containing APIs of synthetic or semi-synthetic origin (see 1.3 Scope for further clarification) and their corresponding FPPs. For simplicity, these guidelines will refer to the QOS-PD, which can be downloaded from the WHO web site.

All sections and fields in the QOS-PD template that would be applicable should be completed. It is understood that certain sections and fields may not apply and should be indicated as such by reporting “not applicable” in the appropriate area with an accompanying explanatory note.

The use of tables to summarize the information is encouraged, where possible. The tables included in the template may need to be expanded or duplicated (e.g. for multiple strengths), as necessary. These tables are included as illustrative examples of how to summarize information. Other approaches to summarize the information can be used if they fulfil the same purpose.

4. MODULE 3: QUALITY

4.1 Table of contents of Module 3

A Table of contents for the filed product dossier should be provided.

4.2 Body of data

3.2.S Drug substance (or active pharmaceutical ingredient (API))

The NMRA, at its discretion, may accept API information in one or more of the following three options:

- Option 1: Certificate of suitability of the European Pharmacopoeia (CEP); or
- Option 2: Drug master file (DMF) procedure;
- Option 3: Full details in the PD.

The applicant should clearly indicate at the beginning of the API section (in the PD and in the QOS-PD) how the information on the API for each API manufacturer is being submitted. The
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API information submitted by the applicant/FPP manufacturer should include the following for each of the options used.

- **Option 1: Certificate of Suitability of the European Pharmacopoeia (CEP)**

  A complete copy of the CEP (including any annexes) should be provided in Module 1. The declaration of access for the CEP should be duly filled out by the CEP holder on behalf of the FPP manufacturer or applicant to the NMRA who refers to the CEP.

  In addition, a written commitment should be included that the applicant will inform the NMRA in the event that the CEP is withdrawn. It should also be acknowledged by the applicant that withdrawal of the CEP would require additional consideration of the API data requirements to support the PD. The written commitment should accompany the copy of the CEP in Module 1.

Along with the CEP, the applicant should supply the following information in the dossier, with data summarized in the QOS-PD.

- 3.2.S.1.3 General properties – discussions on any additional applicable physicochemical and other relevant API properties that are not controlled by the CEP and Ph.Eur. monograph, e.g. solubilities and polymorphs as per guidance in this section.
- 3.2.S.3.1 Elucidation of structure and other characteristics – studies to identify polymorphs (exception: where the CEP specifies a polymorphic form) and particle size distribution, where applicable, as per guidance in this section.
- 3.2.S.4.1 Specification – the specifications of the FPP manufacturer including all tests and limits of the CEP and Ph.Eur. monograph and any additional tests and acceptance criteria that are not controlled in the CEP and Ph.Eur. monograph, such as polymorphs and/or particle size distribution.
- 3.2.S.4.2 / 3.2.S.4.3 Analytical procedures and validation – for any methods used by the FPP manufacturer in addition to those in the CEP and Ph.Eur. monograph.
- 3.2.S.4.4 Batch analysis – results from two batches of at least pilot scale, demonstrating compliance with the FPP manufacturer’s API specifications.
- 3.2.S.5 Reference standards or materials – information on the FPP manufacturer’s reference standards.
- 3.2.S.6 Container closure system – specifications including descriptions and identification of primary packaging components. Exception: where the CEP specifies a container closure system and the applicant declares to use the same container closure system.
- 3.2.S.7 Stability – exception: where the CEP specifies a re-test period that is the same as or of longer duration, and storage conditions which are the same or higher temperature and humidity as proposed by the applicant.

In the case of sterile APIs, data on the sterilization process of the API, including validation data, should be included in the PD.

- **Option 2: Drug master file (DMF) procedure**

  Full details of the chemistry, manufacturing process, quality controls during manufacturing and process validation for the API may be submitted to the NMRA as a
DMF by the API manufacturer, for example as outlined in WHO’s *Guidelines on active pharmaceutical ingredient master file procedure* (Technical Report Series, No. 948, Annex 4, 2008).

In such cases, the *Open part* (non-proprietary information) needs to be included *in its entirety* in the PD as an annex to 3.2.S. In addition, the applicant/FPP manufacturer should complete the following sections in the PD and QOS-PD *in full* according to the guidance provided unless otherwise indicated in the respective sections:

General information S.1.1 through S.1.3  
Manufacture S.2  
  Manufacturer(s) S.2.1  
  Description of manufacturing process and process controls S.2.2  
  Controls of critical steps and intermediates S.2.4  
Elucidation of structure and other characteristics S.3.1  
Impurities S.3.2  
Control of the API S.4.1 through S.4.5  
Reference standards or materials S.5  
Container closure system S.6  
Stability S.7.1 through S.7.3  

It is the responsibility of the applicant to ensure that the complete DMF (i.e. both the applicant’s *Open part* and the API manufacturer’s *Restricted part*) is supplied to the NMRA directly by the API manufacturer and that the applicant has access to the relevant information in the DMF concerning the current manufacture of the API.

A copy of the letter of access should be provided in the PD *Module 1*.

DMF holders can use the guidance provided for the option “Full details in the PD” for preparation of the relevant sections of the Open and Restricted parts of their DMFs. Reference can also be made to the DMF guidelines in WHO Technical Report Series, No. 948, Annex 4.

- **Option 3: Full details in the PD**

  Information on the 3.2.S *Active pharmaceutical ingredient* sections, including full details of chemistry, manufacturing process, quality controls during manufacturing and process validation for the API, should be submitted in the PD as outlined in the subsequent sections of these guidelines. The QOS-PD should be completed as per Section 3.1 of these guidelines.

3.2.S.1 General Information (name, manufacturer)  

3.2.S.1.1 Nomenclature (name, manufacturer)  

Information on the nomenclature of the API should be provided. For example:

- (Recommended) International Non-proprietary Name (INN);  
- Compendial name, if relevant;  
- Chemical name(s);
Company or laboratory code;
• Other non-proprietary name(s) (e.g., national name, United States Adopted Name (USAN), British Approved Name (BAN)); and
• Chemical Abstracts Service (CAS) registry number.

The listed chemical names should be consistent with those appearing in scientific literature and those appearing on the product labelling information (e.g. summary of product characteristics, package leaflet (also known as patient information leaflet or PIL), labelling). Where several names exist, the preferred name should be indicated.

3.2.S.1.2 Structure (name, manufacturer)

The structural formula, including relative and absolute stereochemistry, the molecular formula, and the relative molecular mass should be provided.

This information should be consistent with that provided in Section 3.2.S.1.1. For APIs existing as salts, the molecular mass of the free base or acid should also be provided.

3.2.S.1.3 General properties (name, manufacturer)

A list should be provided of physicochemical and other relevant properties of the API.

This information can be used in developing the specifications, in formulating FPPs and in the testing for release and stability purposes.

The physical and chemical properties of the API should be discussed including the physical description, solubilities in common solvents (e.g. water, alcohols, dichloromethane, acetone), quantitative aqueous pH solubility profile (e.g. pH 1.2 to 6.8, dose/solubility volume), polymorphism, pH and pKa values, UV absorption maxima and molar absorptivity, melting point, refractive index (for a liquid), hygroscopicity, partition coefficient, etc (see table in the QOS-PD). This list is not intended to be exhaustive, but provides an indication as to the type of information that could be included.

Some of the more relevant properties to be considered for APIs are discussed below in greater detail.

Physical description

The description should include appearance, colour and physical state. Solid forms should be identified as being crystalline or amorphous (see 3.2.S.3.1 for further information on API solid forms).

Solubilities/quantitative aqueous pH solubility profile

The following should be provided for all options for the submission of API data.

The solubilities in a number of common solvents should be provided (e.g. water, alcohols, dichloromethane, acetone).
The solubilities over the physiological pH range (pH 1.2 to 6.8) in several buffered media should be provided in mg/ml. If this information is not readily available (e.g. literature references), it should be generated in-house.

For solid oral dosage forms, the dose/solubility volume should be provided as determined by:

\[
\text{dose/solubility volume} = \frac{\text{largest dosage strength (mg)}}{\text{the minimum concentration of the drug (mg/ml)}}
\]

* corresponding to the lowest solubility determined over the physiological pH range (pH 1.2 to 6.8) and temperature (37 ± 0.5°C).

As per the Biopharmaceutics Classification System (BCS), highly soluble (or highly water soluble) APIs are those with a dose/solubility volume of less than or equal to 250 ml.

For example, compound A has as its lowest solubility at 37 ± 0.5°C, 1.0 mg/ml at pH 6.8 and is available in 100 mg, 200 mg and 400 mg strengths. This API would not be considered a BCS highly soluble API as its dose/solubility volume is greater than 250 ml (400 mg/1.0 mg/ml = 400 ml).

Polymorphism

As recommended in ICH’s CTD-Q Questions and answers/location issues document the following refers to where specific data should be located in the PD:

• the polymorphic form(s) present in the proposed API should be listed in Section 3.2.S.1.3;

• the description of manufacturing process and process controls (3.2.S.2.2) should indicate which polymorphic form is manufactured, where relevant;

• the literature references or studies performed to identify the potential polymorphic forms of the API, including the study results, should be provided in Section 3.2.S.3.1;

• if a polymorphic form is to be defined or limited (e.g. for APIs that are not BCS highly soluble and/or where polymorphism has been identified as an issue), details should be included in 3.2.S.4.1 through 3.2.S.4.5.

Additional information is included in the referenced sections of these guidelines.

Particle size distribution

As recommended in ICH’s CTD-Q Questions and Answers/Location Issues document, the studies performed to identify the particle size distribution of the API should be provided in Section 3.2.S.3.1 (refer to this section of these guidelines for additional information).

Information from literature

Supportive data and results from specific studies or published literature can be included within or attached to this section.
Reference documents: ICH Q6A

3.2.S.2 Manufacture (name, manufacturer)

3.2.S.2.1 Manufacturer(s) (name, manufacturer)

The name, address, and responsibility of each manufacturer, including contractors, and each proposed production site or facility involved in manufacturing and testing should be provided.

The facilities involved in the manufacturing, packaging, labelling, testing and storage of the API should be listed. If certain companies are responsible only for specific steps (e.g. milling of the API), this should be clearly indicated.

The list of manufacturers/companies should specify the actual addresses of production or manufacturing site(s) involved (including block(s) and units(s)), rather than the administrative offices. Telephone number(s), fax number(s) and e-mail address(es) should be provided.

A valid manufacturing authorization should be provided for the production of APIs. If available, a certificate of GMP compliance should be provided in the PD in Module 1.

3.2.S.2.2 Description of Manufacturing Process and Process Controls (name, manufacturer)

The description of the API manufacturing process represents the applicant’s commitment for the manufacture of the API. Information should be provided to adequately describe the manufacturing process and process controls. For example:

A flow diagram of the synthetic process(es) should be provided that includes molecular formulae, weights, yield ranges, chemical structures of starting materials, intermediates, reagents and API reflecting stereochemistry, and identifies operating conditions and solvents.

A sequential procedural narrative of the manufacturing process should be submitted. The narrative should include, for example, quantities of raw materials, solvents, catalysts and reagents reflecting the representative batch scale for commercial manufacture, identification of critical steps, process controls, equipment and operating conditions (e.g., temperature, pressure, pH, time).

Alternate processes should be explained and described with the same level of detail as the primary process. Reprocessing steps should be identified and justified. Any data to support this justification should be either referenced or filed in 3.2.S.2.5.

Where the DMF procedure is used, a cross-reference to the Restricted part of the DMF may be indicated for confidential information. In this case, if detailed information is presented in the Restricted part, the information to be provided for this section of the PD includes a flow chart (including molecular structures and all reagents and solvents) and a brief outline of the manufacturing process, with special emphasis on the final steps including purification procedures. However, for sterile APIs full validation data on the sterilization process should
be provided in the Open part (in cases where there is no further sterilization of the final product).

The following requirements apply to the third option for submission of API information, where full details are provided in the dossier.

As discussed in ICH Q7 and WHO Technical Report Series, No. 957 Annex 2, the point at which the API starting material is introduced into the manufacturing process is the starting point of the application of GMP requirements. The API starting material itself needs to be proposed and its choice justified by the manufacturer and accepted as such by assessors. The API starting material should be proposed taking into account the complexity of the molecule, the proximity of the API starting material to the final API, the availability of the API starting material as a commercial chemical and the quality controls placed upon the API starting material. This justification should be documented in the dossier and be available for review by WHO GMP inspectors.

In situations where the API starting material is a complex molecule and only a minimal number of synthetic steps from the final API, a further molecule called the starting material for synthesis should be proposed and its choice justified by the applicant. The starting material for synthesis defines the starting point in the manufacturing process for an API to be described in an application. The applicant should propose and justify which substances should be considered as starting materials for synthesis. See section 3.2.S.2.3 for further guidance. In the case where the precursor to the API is obtained from fermentation, or is from plant or animal origin, such a molecule can be considered the API starting material regardless of complexity.

A one step synthesis may be accepted in exceptional cases, for example where the API starting material is covered by a CEP, or where the API starting material is an API accepted through the DMF procedure, or when the structure of the API is so simple that a one step synthesis can be justified, e.g. ethambutol or ethionamide.

In addition to the detailed description of the manufacturing process as per ICH M4Q, the recovery of materials, if any, should be described in detail with the step in which they are introduced into the process. Recovery operations should be adequately controlled such that impurity levels do not increase over time. For recovery of solvents, any processing to improve the quality of the recovered solvent should be described. Regarding recycling of filtrates (mother liquors) to obtain second crops, information should be available on maximum holding times of mother liquors and maximum number of times the material can be recycled. Data on impurity levels should be provided to justify recycling of filtrates.

Where there are multiple manufacturing sites for one API manufacturer, a comprehensive list in tabular form should be provided comparing the processes at each site and highlighting any differences.

All solvents used in the manufacture (including purification and/or crystallization step(s)) should be clearly identified. Solvents used in the final steps should be of high purity. Use of recovered solvents in the final steps of purification and/or crystallization is not recommended, however their use can be justified on presentation of sufficient data demonstrating that recovered solvents meet appropriate standards as outlined in ICH Q7.
Where polymorphic/amorphous forms have been identified, the form resulting from the synthesis should be stated.

Where particle size is considered a critical attribute (see 3.2.S.3.1 for details), the particle size reduction method(s) (milling, micronization) should be described.

Justification should be provided for alternate manufacturing processes. Alternate processes should be explained with the same level of detail as the primary process. It should be demonstrated that batches obtained by the alternate processes have the same impurity profile as the principal process. If the obtained impurity profile is different it should be demonstrated to be acceptable according to the requirements described under S.3.2.

It is acceptable to provide information on pilot scale manufacture, provided it is representative of production scale and scale-up is reported immediately to the NMRA according to the requirements of the associated variation guidelines (e.g. WHO Technical Report Series, No. X, Annex Y, 2013).

3.2.S.2.3 Control of Materials (name, manufacturer)

Materials used in the manufacture of the API (e.g., raw materials, starting materials, solvents, reagents, catalysts) should be listed identifying where each material is used in the process. Information on the quality and control of these materials should be provided. Information demonstrating that materials meet standards appropriate for their intended use should be provided, as appropriate. (Details in 3.2.A.2)

Where the DMF procedure is used, a cross-reference to the Restricted part of the DMF is considered sufficient for this section.

The following requirements apply to the third option for submission of API information, where full details are provided in the dossier.

The API starting material should be fully characterized and suitable specifications proposed and justified, including at a minimum control for identity, assay, impurity content and any other critical attribute of the material. For each API starting material, the name and manufacturing site address of the manufacturer(s) should be indicated. A brief description of the preparation of the API starting material should be provided for each manufacturer, including the solvents, catalysts and reagents used. A single set of specifications should be proposed for the starting material that applies to material from all sources. Any future changes to the API starting material manufacturers, mode of preparation or specifications should be notified.

As indicated in section 3.2.S.2 there are occasions where a starting material for synthesis may also need to be defined. In general, the starting material for synthesis described in the PD should:

- be a synthetic precursor of one or more synthesis steps prior to the final API intermediate. Acids, bases, salts, esters and similar derivatives of the API, as well as the racemate of a single enantiomer API, are not considered final intermediates;
- be a well characterized, isolated and purified substance with its structure fully elucidated including its stereochemistry (when applicable);
• have well defined specifications that include among others one or more specific identity tests and tests and limits for assay and specified, unspecified and total impurities; and
• be incorporated as a significant structural fragment into the structure of the API.

Copies of the specifications for the materials used in the synthesis, extraction, isolation and purification steps should be provided in the PD, including starting materials, reagents, solvents, catalysts and recovered materials. Confirmation should be provided that the specifications apply to materials used at each manufacturing site. A certificate of analysis of the starting material for synthesis should be provided. A summary of the information on starting materials should be provided in the QOS-PD.

The carry-over of impurities of the starting materials for synthesis into the final API should be considered and discussed.

A letter of attestation should be provided confirming that the API and the starting materials and reagents used to manufacture the API are without risk of transmitting agents of animal spongiform encephalopathies.

When available, a CEP demonstrating TSE-compliance should be provided. A complete copy of the CEP (including any annexes) should be provided in Module 1.

Reference documents: ICH Q6A

3.2.S.2.4 Controls of Critical Steps and Intermediates (name, manufacturer)

Critical Steps: Tests and acceptance criteria (with justification including experimental data) performed at critical steps identified in 3.2.S.2.2 of the manufacturing process to ensure that the process is controlled should be provided.

Intermediates: Information on the quality and control of intermediates isolated during the process should be provided.

Where the DMF procedure is used a cross-reference to the Restricted part of the DMF is considered sufficient for this section of the PD, with the exception of information that is also relevant for the applicant (ref: APIMF guidelines in WHO Technical Report Series, No. 948, Annex 4).

The following requirements apply to the third option for submission of API information, where full details are provided in the dossier.

The critical steps should be identified. These can be among others: steps where significant impurities are removed or introduced, steps introducing an essential molecular structural element such as a chiral centre or resulting in a major chemical transformation, steps having an impact on solid-state properties and homogeneity of the API that may be relevant for use in solid dosage forms.

Specifications for isolated intermediates should be provided and should include tests and acceptance criteria for identity, purity and assay, where applicable.

Reference documents: ICH Q6A
3.2.S.2.5 Process Validation and/or Evaluation (name, manufacturer)

Process validation and/or evaluation studies for aseptic processing and sterilisation should be included.

Where the DMF procedure is used, a cross-reference to the Restricted part of the DMF is considered sufficient for this section of the PD.

The following requirements apply to the third option for submission of API information, where full details are provided in the dossier.

It is expected that the manufacturing processes for all APIs are properly controlled. If the API is prepared as sterile, a complete description should be provided for aseptic processing and/or sterilization methods. The controls used to maintain the sterility of the API during storage and transportation should also be provided. Alternate processes should be justified and described (see guidance in 3.2.S.2.2 for the level of detail expected).

3.2.S.2.6 Manufacturing Process Development (name, manufacturer)

A description and discussion should be provided of the significant changes made to the manufacturing process and/or manufacturing site of the API used in producing comparative bioavailability or biowaiver, scale-up, pilot, and, if available, production scale batches.

Reference should be made to the API data provided in Section 3.2.S.4.4.

Where the DMF procedure is used, a cross-reference to the Restricted part of the DMF is considered sufficient for this section of the PD.

3.2.S.3 Characterisation (name, manufacturer)

3.2.S.3.1 Elucidation of Structure and other Characteristics (name, manufacturer)

Confirmation of structure based on e.g., synthetic route and spectral analyses should be provided. Information such as the potential for isomerism, the identification of stereochemistry, or the potential for forming polymorphs should also be included.

Elucidation of structure

The PD should include quality assurance (QA) certified copies of the spectra, peak assignments and a detailed interpretation of the data of the studies performed to elucidate and/or confirm the structure of the API. The QOS-PD should include a list of the studies performed and a conclusion from the studies (e.g. if the results support the proposed structure).

For APIs that are not described in an officially recognized pharmacopoeia, the studies carried out to elucidate and/or confirm the chemical structure normally include elemental analysis, infrared (IR), ultraviolet (UV), nuclear magnetic resonance (NMR) and mass spectra (MS) studies. Other tests could include X-ray powder diffraction (XRPD) and differential scanning calorimetry (DSC).
For APIs that are described in an officially recognized pharmacopoeia, it is generally sufficient to provide copies of the IR spectrum of the API from each of the proposed manufacturer(s) run concomitantly with an officially recognized pharmacopoeial reference standard. See Section 3.2.S.5 for details on acceptable reference standards or materials.

Isomerism/Stereochemistry

When an API is chiral, it should be specified whether specific stereoisomers or a mixture of stereoisomers have been used in the comparative biostudies, and information should be given as to the stereoisomer of the API that is to be used in the FPP.

Where the potential for stereoisomerism exists, a discussion should be included of the possible isomers that can result from the manufacturing process and the steps where chirality was introduced. The identicality of the isomeric composition of the API to that of the API in the comparator product should be established. Information on the physical and chemical properties of the isomeric mixture or single enantiomer should be provided, as appropriate. The API specification should include a test to ensure isomeric identity and purity.

The potential for interconversion of the isomers in the isomeric mixture, or racemisation of the single enantiomer should be discussed.

When a single enantiomer of the API is claimed for non-pharmacopoeial APIs, unequivocal proof of absolute configuration of asymmetric centres should be provided such as determined by X-ray of a single crystal.

If, based on the structure of the API, there is not a potential for stereoisomerism, it is sufficient to include a statement to this effect.

Polymorphism

Many APIs can exist in different physical forms in the solid state. Polymorphism is characterized as the ability of an API to exist as two or more crystalline phases that have different arrangements and/or conformations of the molecules in the crystal lattice. Amorphous solids consist of disordered arrangements of molecules and do not possess a distinguishable crystal lattice. Solvates are crystal forms containing either stoichiometric or nonstoichiometric amounts of a solvent. If the incorporated solvent is water, the solvates are also commonly known as hydrates.

Polymorphic forms of the same chemical compound differ in internal solid-state structure and, therefore, may possess different chemical and physical properties, including packing, thermodynamic, spectroscopic, kinetic, interfacial and mechanical properties. These properties can have a direct impact on API processability, pharmaceutical product manufacturability and product quality/performance, including stability, dissolution and bioavailability. Unexpected appearance or disappearance of a polymorphic form may lead to serious pharmaceutical consequences.

Applicants and API manufacturers are expected to have adequate knowledge about the polymorphism of the APIs used and/or produced. Information on polymorphism can come from the scientific literature, patents, compendia or other references to determine if polymorphism is a concern, e.g. for APIs that are not BCS highly soluble. In the absence of
published data for APIs that are not *BCS highly soluble*, polymorphic screening will be
necessary to determine if the API can exist in more than one crystalline form. Polymorphic
screening is generally accomplished via crystallization studies using different solvents and
conditions.

There are a number of methods that can be used to characterize the polymorphic forms of an
API. Demonstration of a nonequivalent structure by single crystal X-ray diffraction is
currently regarded as the definitive evidence of polymorphism. XRPD can also be used to
provide unequivocal proof of polymorphism. Other methods, including microscopy, thermal
analysis (e.g. DSC, thermal gravimetric analysis and hot-stage microscopy) and spectroscopy
(e.g. IR, Raman, solid-state nuclear magnetic resonance (ssNMR)) are helpful to further
characterize polymorphic forms. Where polymorphism is a concern, the
applicants/manufacturers of APIs should demonstrate that a suitable method, capable of
distinguishing different polymorphs, is available to them.

Decision tree 4(1) of ICH Q6A can be used where screening is necessary and 4(2) can be used
to investigate if different polymorphic forms have different properties that may affect
performance, bioavailability and stability of the FPP and to decide whether a preferred
polymorph should be monitored at release and on storage of the API. Where there is a
preferred polymorph, acceptance criteria should be incorporated into the API specification to
ensure polymorphic equivalence of the commercial material and that of the API batches used
in the comparative bioavailability or biowaiver studies. The polymorphic characterization of
the API batches used in comparative bioavailability or biowaiver studies by the above
mentioned methods should be provided. The method used to control polymorphic form
should be demonstrated to be specific for the preferred form.

Polymorphism can also include solvation or hydration products (also known as
pseudopolymorphs). If the API is used in a solvated form, the following information should
be provided:

- specifications for the solvent-free API in 3.2.S.2.4, if that compound is a synthetic
  precursor;
- specifications for the solvated API including appropriate limits on the weight ratio of
  API to solvent (with data to support the proposed limits);
- a description of the method used to prepare the solvate in 3.2.S.2.2.

*Particle size distribution*

For APIs that are not *BCS highly soluble* contained in solid FPPs, or liquid FPPs containing
undissolved API, the particle size distribution of the material can have an effect on the in vitro
and/or in vivo behaviour of the FPP. Particle size distribution can also be important in dosage
form performance (e.g. delivery of inhalation products), achieving uniformity of content in
low-dose tablets (e.g. 2 mg or less), desired smoothness in ophthalmic preparations and
stability of suspensions.

If particle size distribution is an important parameter (e.g. as in the above cases), results from
an investigation of several batches of the API should be provided, including characterization
of the batch(es) used in the comparative bioavailability or biowaiver studies. API
specifications should include controls on the particle size distribution to ensure consistency
with the material in the batch(es) used in the comparative bioavailability and biowaiver
studies (e.g. limits for d10, d50 and d90). The criteria should be established statistically based
on the standard deviation of the test results from the previously mentioned studies. The following is provided for illustrative purposes as possible acceptance criteria for particle size distribution limits:

- \(d_{10}\) not more than (NMT) 10% of total volume less than \(X\) \(\mu\)m
- \(d_{50}\) \(XX\) \(\mu\)m – \(XXX\) \(\mu\)m
- \(d_{90}\) not less than (NLT) 90% of total volume less than \(XXXX\) \(\mu\)m.

Other controls on particle size distribution can be considered acceptable, if scientifically justified.

Reference documents: ICH Q6A

### 3.2.S.3.2 Impurities (name, manufacturer)

**Information on impurities should be provided.**

Details on the principles for the control of impurities (e.g. reporting, identification and qualification) are outlined in the ICH Q3A, Q3B and Q3C impurity guidelines. Additional information to provide further guidance on some of the elements discussed in the ICH guidelines is outlined below.

Regardless of whether a pharmacopoeial standard is claimed, a discussion should be provided of the potential and actual impurities arising from the synthesis, manufacture, or degradation of the API. This should cover starting materials, by-products, intermediates, chiral impurities and degradation products and should include the chemical names, structures and origins. The discussion of pharmacopoeial APIs should not be limited to the impurities specified in the API monograph.

The tables in the QOS-PD template should be used to summarize the information on the API-related and process-related impurities. In the QOS-PD, the term *origin* refers to how and where the impurity was introduced (e.g. “Synthetic intermediate from Step 4 of the synthesis”, “Potential by-product due to rearrangement from Step 6 of the synthesis”). It should also be indicated if the impurity is a metabolite of the API.

The ICH thresholds for reporting, identification (used to set the limit for individual unknown impurities) and qualification are determined on the basis of potential exposure to the impurity, e.g. by the maximum daily dose (MDD) of the API. For APIs available in multiple dosage forms and strengths having different MDD values, it is imperative that the thresholds and corresponding controls for each of the presentations be considered to ensure that the risks posed by impurities have been addressed. This is normally achieved by using the highest potential daily MDD, rather than the maintenance dose. For parenteral products, the maximum hourly dose of the API should also be included.

It is acknowledged that APIs of semi-synthetic origin do not fall within the scope of the ICH impurity guidelines. However, depending on the nature of the API and the extent of the chemical modification steps, the principles on the control of impurities (e.g. reporting, identification and qualification) could also be extended to APIs of semi-synthetic origin. As an illustrative example, an API whose precursor molecule was derived from a fermentation process, or a natural product of plant or animal origin that has subsequently undergone several chemical modification reactions generally would fall within this scope, whereas an API whose
sole chemical step was the formation of a salt from a fermentation product generally would not fall within this scope. It is understood that there is some latitude for these types of APIs.

Identification of impurities

It is recognized by the pharmacopoeias that APIs can be obtained from various sources and thus can contain impurities not considered during the development of the monograph. Furthermore, a change in the production or source may give rise to additional impurities that are not adequately controlled by the official compendial monograph. As a result, each PD is assessed independently to consider the potential impurities that may arise from the proposed route(s) of synthesis. For these reasons, the ICH limits for unspecified impurities (e.g., NMT 0.10% or 1.0 mg per day intake (whichever is lower) for APIs having a maximum daily dose $\leq 2$ g/day) are generally recommended, rather than the general limits for unspecified impurities that may appear in the official compendial monograph that could potentially be higher than the applicable ICH limit.

Qualification of impurities

The ICH impurity guidelines should be consulted for options on the qualification of impurities. The limit specified for an identified impurity in an officially recognized pharmacopoeia is generally considered to be qualified. The following is an additional option for qualification of impurities in existing APIs:

The limit for an impurity present in an existing API can be accepted by comparing the impurity results found in the existing API with those observed in an innovator product using the same validated, stability-indicating analytical procedure (e.g., comparative HPLC studies). If samples of the innovator product are not available, the impurity profile may also be compared to a different approved FPP with the same route of administration and similar characteristics (e.g., tablet versus capsule). It is recommended that the studies be conducted on comparable samples (e.g., age of samples) to obtain a meaningful comparison of the impurity profiles.

Levels of impurities generated from studies under accelerated or stressed storage conditions of the innovator or approved FPP are not considered acceptable/qualified.

A specified impurity present in the existing API is considered qualified if the amount of the impurity in the existing API reflects the levels observed in the innovator or approved FPP.

Basis for setting the acceptance criteria

The basis for setting the acceptance criteria for the impurities should be provided. This is established by considering the identification and qualification thresholds for API-related impurities (e.g., starting materials, by-products, intermediates, chiral impurities or degradation products) and the concentration limits for process-related impurities (e.g., residual solvents) as per the applicable ICH guidelines (e.g., Q3A, Q3C).

The qualified level should be considered as the maximum allowable limit. However, limits which are considerably wider than the actual manufacturing process capability are generally discouraged. For this reason, the acceptance criteria are also set taking into consideration the actual levels of impurities found in several batches of the API from each
manufacturer, including the levels found in the batches used for the comparative
bioavailability or biowaiver studies. When reporting the results of quantitative tests, the actual
numerical results should be provided rather than vague statements such as “within limits” or
“conforms”. In the cases where a large number of batches have been tested it is acceptable to
summarize the results of the total number of batches tested with a range of analytical results.

If there are identified impurities specified in an official compendial monograph that are not
controlled by the proposed routine in-house analytical procedure, a justification for their
exclusion from routine analyses should be provided (e.g. “Impurities D, E and F listed in the
Ph.Int. monograph are not potential impurities from the proposed route of synthesis used by
manufacturer X”). If acceptable justification cannot be provided it should be demonstrated
that the routine in-house method is capable of separating and detecting the impurities
specified in the official compendial monograph at an acceptable level (e.g. 0.10%). If such a
demonstration cannot be performed, a one-time study should be conducted applying the
pharmacopoeial method to several recent batches to demonstrate the absence of the
pharmacopoeial listed impurities.

ICH class II solvent(s) used prior to the last step of the manufacturing process may be
exempted from routine control in API specifications if suitable justification is provided.
Submission of results demonstrating less than 10% of the ICH Q3C limit (option I) of the
solvent(s) in three consecutive production-scale batches or six consecutive pilot-scale batches
of the API or a suitable intermediate would be considered acceptable justification. The last
step solvents used in the process should always be routinely controlled in the final API.

For guidance on acceptable residual solvent limits, refer to ICH Q3C. The limit for residues
of triethylamine (TEA) is either 320 ppm on the basis of ICH Q3C option I or 3.2 mg/day on
the basis of permitted daily exposure (PDE).

The absence of known established highly toxic impurities (genotoxic) used in the process or
formed as a by-product should be discussed and suitable limits should be proposed. The limits
should be justified by appropriate reference to available guidances (e.g.
EMEA/CHMP/QWP/251344/2006 or USFDA Guidance for Industry: Genotoxic and
carcinogenic impurities in drug substances and products, recommended approaches,
December 2008) or by providing experimental safety data or published data in peer-reviewed
journals.

Residues of metal catalysts used in the manufacturing process and determined to be present in
batches of API are to be controlled in specifications. This requirement does not apply to
metals that are deliberate components of the pharmaceutical substance (such as a counter ion
of a salt) or metals that are used as a pharmaceutical excipient in the FPP (e.g. an iron oxide
pigment). The guidelines on the specification limits for residues of metal catalysts or metal
reagents EMEA/CHMP/SWP/4446/2000 or any equivalent approaches can be used to address
this issue. The requirement normally does not apply to extraneous metal contaminants that are
more appropriately addressed by GMP, GDP or any other relevant quality provision such as
the heavy metal test in monographs of recognized pharmacopoeias that cover metal
contamination originating from manufacturing equipment and the environment.

Reference documents: ICH Q3A, Q3C, Q6A

3.2.S.4 Control of the API (name, manufacturer)
3.2.S.4.1 Specification (name, manufacturer)

The specification for the API should be provided.

As defined in ICH’s Q6A guideline, a specification is:

“a list of tests, references to analytical procedures and appropriate acceptance criteria, which are numerical limits, ranges, or other criteria for the tests described. It establishes the set of criteria to which an API or FPP should conform to be considered acceptable for its intended use. “Conformance to specifications” means that the API and/or FPP, when tested according to the listed analytical procedures, will meet the listed acceptance criteria. Specifications are critical quality standards that are proposed and justified by the manufacturer and approved by regulatory authorities.”

Copies of the API specifications, dated and signed by authorized personnel (e.g. the person in charge of the quality control or quality assurance department) should be provided in the PD, including specifications from each API manufacturer as well as those of the FPP manufacturer.

The FPP manufacturer’s API specification should be summarized according to the table in the QOS-PD template under the headings tests, acceptance criteria and analytical procedures (including types, sources and versions for the methods).

- The standard declared by the applicant could be an officially recognized compendial standard (e.g. Ph.Int., Ph.Eur., BP, USP, JP) or an in-house (manufacturer’s) standard.
- The specification reference number and version (e.g. revision number and/or date) should be provided for version control purposes.
- For the analytical procedures, the type should indicate the kind of analytical procedure used (e.g. visual, IR, UV, HPLC, laser diffraction), the source refers to the origin of the analytical procedure (e.g. Ph.Int., Ph.Eur., BP, USP, JP, in-house) and the version (e.g. code number/version/date) should be provided for version control purposes.

In cases where there is more than one API manufacturer, the FPP manufacturer’s API specifications should be one single compiled set of specifications that is identical for each manufacturer. It is acceptable to lay down in the specification more than one acceptance criterion and/or analytical method for a single parameter with the statement “for API from manufacturer A” (e.g. in the case of residual solvents).

Any non routine testing should be clearly identified as such and justified along with the proposal on the frequency of non routine testing.

The ICH Q6A guideline outlines recommendations for a number of universal and specific tests and criteria for APIs.

Reference documents: ICH Q3A, Q3C, Q6A, officially recognized pharmacopoeia

3.2.S.4.2 Analytical Procedures (name, manufacturer)

The analytical procedures used for testing the API should be provided.
Copies of the in-house analytical procedures used to generate testing results provided in the
PD, as well as those proposed for routine testing of the API by the FPP manufacturer, should
be provided. Unless modified, it is not necessary to provide copies of officially recognized
compendial analytical procedures.

Tables for summarizing a number of the different analytical procedures and validation
information (e.g. HPLC assay/impurity methods, GC methods) can be found in the 2.3.R
Regional information section of the QOS-PD (i.e. 2.3.R.2). These tables may be used to
summarize the in-house analytical procedures of the FPP manufacturer for determination of
the residual solvents, assay and purity of the API, in section 2.3.S.4.2 of the QOS-PD. Other
methods used to generate assay and purity data in the PD can be summarized in 2.3.S.4.4 (c)
or 2.3.S.7.3 (b) of the QOS-PD. Officially recognized compendial methods need not be
summarized unless modifications have been made.

Although HPLC is normally considered the method of choice for determining API-related
impurities, other chromatographic methods such as GC and TLC can also be used, if
appropriately validated. For determination of related substances, reference standards should
normally be available for each of the identified impurities, particularly those known to be
toxic and the concentration of the impurities should be quantitated against their own reference
standards. Impurity standards may be obtained from pharmacopoeias (individual impurities
or resolution mixtures), from commercial sources or prepared in-house. It is considered
acceptable to use the API as an external standard to estimate the levels of impurities, provided
the response factors of those impurities are sufficiently close to that of the API, i.e. between
80 and 120%. In cases where the response factor is outside this range, it may still be
acceptable to use the API, provided a correction factor is applied. Data to support calculation
of the correction factor should be provided for an in-house method. Unspecified impurities
may be quantitated using a solution of the API as the reference standard at a concentration
corresponding to the limit established for individual unspecified impurities (e.g. 0.10%). The
test for related substances in the Ph.Int. monograph for lamivudine serves as a typical
example.

The system suitability tests (SSTs) represent an integral part of the method and are used to
ensure the adequate performance of the chosen chromatographic system. As a minimum,
HPLC and GC purity methods should include SSTs for resolution and repeatability. For
HPLC methods to control API-related impurities, this is typically done using a solution of the
API with a concentration corresponding to the limit for unspecified impurities. Resolution of
the two closest eluting peaks is generally recommended. However, the choice of alternate
peaks can be used if justified (e.g. choice of a toxic impurity). In accordance with the Ph.Int.
section on Methods of Analysis, the repeatability test should include an acceptable number of
replicate injections. HPLC assay methods should include SSTs for repeatability and in
addition either peak asymmetry, theoretical plates or resolution. For TLC methods, the SSTs
should verify the ability of the system to separate and detect the analyte(s) (e.g. by applying a
spot corresponding to the API at a concentration corresponding to the limit of unspecified
impurities).


3.2.S.4.3 Validation of Analytical Procedures (name, manufacturer)

Analytical validation information, including experimental data for the analytical
procedures used for testing the API, should be provided.
Copies of the validation reports for the analytical procedures used to generate testing results provided in the PD, as well as those proposed for routine testing of the API by the FPP manufacturer, should be provided.

Tables for summarizing a number of the different analytical procedures and validation information (e.g. HPLC assay/impurity methods, GC methods) can be found in the 2.3.R Regional information section of the QOS-PD (i.e. 2.3.R.2). These tables may be used to summarize the validation information of the analytical procedures of the FPP manufacturer for determination of residual solvents, assay and purity of the API, in section 2.3.S.4.3 of the QOS-PD. The validation data for other methods used to generate assay and purity data in the PD can be summarized in 2.3.S.4.4 (c) or 2.3.S.7.3 (b) of the QOS-PD.

As recognized by regulatory authorities and pharmacopoeias themselves, verification of compendial methods can be necessary. The compendial methods as published are typically validated based on an API or an FPP originating from a specific manufacturer. Different sources of the same API or FPP can contain impurities and/or degradation products that were not considered during the development of the monograph. Therefore the monograph and compendial method should be demonstrated suitable to control the impurity profile of the API from the intended source(s).

In general verification is not necessary for compendial API assay methods. However, specificity of a specific compendial assay method should be demonstrated if there are any potential impurities that are not specified in the compendial monograph. If an officially recognized compendial method is used to control API-related impurities that are not specified in the monograph, full validation of the method is expected with respect to those impurities.

If an officially recognized compendial standard is claimed and an in-house method is used in lieu of the compendial method (e.g. for assay or for specified impurities), equivalency of the in-house and compendial methods should be demonstrated. This could be accomplished by performing duplicate analyses of one sample by both methods and providing the results from the study. For impurity methods, the sample analyzed should be the API spiked with impurities at concentrations equivalent to their specification limits.

Reference documents: ICH Q2

3.2.S.4.4 Batch Analyses (name, manufacturer)

Description of batches and results of batch analyses should be provided.

The information provided should include batch number, batch size, date and production site of relevant API batches used in comparative bioavailability or biowaiver studies, preclinical and clinical data (if relevant), stability, pilot, scale-up and, if available, production-scale batches. This data is used to establish the specifications and evaluate consistency in API quality.

Analytical results should be provided from at least two batches of at least pilot scale from each proposed manufacturing site of the API and should include the batch(es) used in the comparative bioavailability or biowaiver studies. A pilot-scale batch should be manufactured by a procedure fully representative of and simulating that to be applied to a full production-scale batch.
Copies of the certificates of analysis, both from the API manufacturer(s) and the FPP manufacturer, should be provided for the profiled batches and any company responsible for generating the test results should be identified. The FPP manufacturer’s test results should be summarized in the QOS-PD.

The discussion of results should focus on observations noted for the various tests, rather than reporting comments such as “all tests meet specifications”. For quantitative tests (e.g. individual and total impurity tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms”.

A discussion and justification should be provided for any incomplete analyses (e.g. results not tested according to the proposed specification).

Reference documents: ICH Q3A, Q3C, Q6A

3.2.S.4.5 Justification of Specification (name, manufacturer)

Justification for the API specification should be provided.

A discussion should be provided on the inclusion of certain tests, evolution of tests, analytical procedures and acceptance criteria, differences from the officially recognized compendial standard(s), etc. If the officially recognized compendial methods have been modified or replaced, a discussion should be included.

The justification for certain tests, analytical procedures and acceptance criteria may have been discussed in other sections of the PD (e.g. impurities, particle size distribution) and does not need to be repeated here, although a cross-reference to their location should be provided.

Reference documents: ICH Q3A, Q3C, Q6A, officially recognized pharmacopoeia

3.2.S.5 Reference Standards or Materials (name, manufacturer)

Information on the reference standards or reference materials used for testing of the API should be provided.

Information should be provided on the reference standard(s) used to generate data in the PD, as well as those to be used by the FPP manufacturer in routine API and FPP testing.

The source(s) of the reference standards or materials used in the testing of the API should be provided (e.g. those used for the identification, purity, assay tests). These could be classified as primary or secondary reference standards.

A suitable primary reference standard should be obtained from an officially recognized pharmacopoeial source (e.g. Ph.Int., Ph.Eur., BP, USP, JP) where one exists and the lot number should be provided. Where a pharmacopoeial standard is claimed for the API and/or the FPP, the primary reference standard should be obtained from that pharmacopoeia when available. Primary reference standards from officially recognized pharmacopoeial sources do not need further structural elucidation.

Otherwise, a primary standard may be a batch of the API that has been fully characterized (e.g. by IR, UV, NMR, MS analyses). Further purification techniques may be needed to
render the material acceptable for use as a chemical reference standard. The purity requirements for a chemical reference substance depend upon its intended use. A chemical reference substance proposed for an identification test does not require meticulous purification, since the presence of a small percentage of impurities in the substance often has no noticeable effect on the test. On the other hand, chemical reference substances that are to be used in assays should possess a high degree of purity (such as 99.5% on the dried or water/solvent free basis). Absolute content of the primary reference standard must be declared and should follow the scheme: 100% minus organic impurities (quantitated by an assay procedure, e.g. HPLC, DSC, etc.) minus inorganic impurities minus volatile impurities by loss on drying (or water content minus residual solvents).

A secondary (or in-house) reference standard can be used by establishing it against a suitable primary reference standard, e.g. by providing legible copies of the IR of the primary and secondary reference standards run concomitantly and by providing its certificate of analysis, including assay determined against the primary reference standard. A secondary reference standard is often characterized and evaluated for its intended purpose with additional procedures other than those used in routine testing (e.g. if additional solvents are used during the additional purification process that are not used for routine purposes).

Reference standards should normally be established for specified impurities. Refer to 3.2.S.4.2 for additional guidance.


3.2.S.6 Container Closure System (name, manufacturer)

A description of the container closure system(s) should be provided, including the identity of materials of construction of each primary packaging component, and their specifications. The specifications should include description and identification (and critical dimensions with drawings, where appropriate). Non-compendial methods (with validation) should be included, where appropriate.

For non-functional secondary packaging components (e.g., those that do not provide additional protection), only a brief description should be provided. For functional secondary packaging components, additional information should be provided.

The suitability should be discussed with respect to, for example, choice of materials, protection from moisture and light, compatibility of the materials of construction with the API, including sorption to container and leaching, and/or safety of materials of construction.

The WHO Guidelines on packaging for pharmaceutical products (WHO Technical Report Series, No. 902, Annex 9, 2002) and the officially recognized pharmacopoeias should be consulted for recommendations on the packaging information for APIs.

Primary packaging components are those that are in direct contact with the API or FPP. The specifications for the primary packaging components should be provided and should include a specific test for identification (e.g. IR).

Copies of the labels applied on the secondary packaging of the API should be provided and should include the conditions of storage. In addition, the name and address of the
manufacturer of the API should be stated on the container, regardless of whether relabeling is conducted at any stage during the API distribution process.

3.2.7 Stability (name, manufacturer)

3.2.7.1 Stability Summary and Conclusions (name, manufacturer)

The types of studies conducted, protocols used, and the results of the studies should be summarised. The summary should include results, for example, from forced degradation studies and stress conditions, as well as conclusions with respect to storage conditions and re-test date or shelf-life, as appropriate.

The WHO guidelines on Stability testing of active pharmaceutical ingredients and finished pharmaceutical products (WHO Technical Report Series, No. 953, Annex 2) should be consulted for recommendations on the core stability data package.

As outlined in the WHO stability guidelines, the purpose of stability testing is to:

“provide evidence of how the quality of an API or FPP varies with time under the influence of a variety of environmental factors such as temperature, humidity and light.”

The tables in the QOS-PD template should be used to summarize the results from the stability studies and related information (e.g. conditions, testing parameters, conclusions, commitments).

Stress testing

As outlined in the ICH Q1A guidance document, stress testing of the API can help identify the likely degradation products, which can in turn help establish the degradation pathways and the intrinsic stability of the molecule and validate the stability indicating power of the analytical procedures used. The nature of the stress testing will depend on the individual API and the type of FPP involved.

Stress testing may be carried out on a single batch of the API. For examples of typical stress conditions refer to WHO Technical Report Series, No. 953, Annex 2, Section 2.1.2, as well as, “A typical set of studies of the degradation paths of an active pharmaceutical ingredient” in WHO Technical Report Series, No. 929, Annex 5, Table A.1.

The objective of stress testing is not to completely degrade the API, but to cause degradation to occur to a small extent, typically 10-30% loss of active by assay when compared with non-degraded API. This target is chosen so that some degradation occurs, but not enough to generate secondary products. For this reason, the conditions and duration may need to be varied when the API is especially susceptible to a particular stress factor. In the total absence of degradation products after 10 days, the API is considered stable under the particular stress condition.

The tables in the QOS-PD template should be used to summarize the results of the stress testing and should include the treatment conditions (e.g. temperatures, relative humidities, concentrations of solutions, durations) and the observations for the various test parameters.
The discussion of results should highlight whether mass balance was observed.

Photostability testing should be an integral part of stress testing. The standard conditions are described in ICH Q1B. If “protect from light” is stated in one of the officially recognized pharmacopoeia for the API, it is sufficient to state “protect from light” on labelling, in lieu of photostability studies, when the container closure system is shown to be light protective.

When available, it is acceptable to provide the relevant data published in the scientific literature (inter alia WHOPARs, EPARs) to support the identified degradation products and pathways.

**Accelerated and long-term testing**

Available information on the stability of the API under accelerated and long-term conditions should be provided, including information in the public domain or obtained from scientific literature. The source of the information should be identified.

The storage conditions and the lengths of studies chosen should be sufficient to cover storage and shipment. Refer to the WHO stability guidelines WHO Technical Report Series, No. 953 Annex 2.

To establish the re-test period, data should normally be provided on not less than three batches of at least pilot scale. The batches should be manufactured by the same synthesis route as production batches and using a method of manufacture and procedure that simulates the final process to be used for production batches. The stability testing programme should be summarized and the results of stability testing should be summarized in the dossier and in the tables in the QOS-PD.

The information on the stability studies should include details such as storage conditions, batch number, batch size, container closure system and completed (and proposed) test intervals. The discussion of results should focus on observations noted for the various tests, rather than reporting comments such as “all tests meet specifications”. Ranges of analytical results where relevant and any trends that were observed should be included. For quantitative tests (e.g. individual and total degradation product tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms”. Where different from the methods described in S.4.2, descriptions and validation of the methodology used in stability studies should be provided.

Refer to WHO Technical Report Series, No. 953, Annex 2 for further information regarding the minimum data required at the time of submitting the dossier, storage conditions, container closure system, test specifications and testing frequency.

**Proposed storage statement and re-test period**

A storage statement should be established for display on the label based on the stability evaluation of the API. The WHO stability guidelines include a number of recommended storage statements that should be used, when supported by the stability studies.

A re-test period should be derived from the stability information and should be displayed on the container label.
After this re-test period, a batch of API destined for use in the manufacture of an FPP could be re-tested and then, if in compliance with the specification, could be used immediately (e.g. within 30 days). If re-tested and found compliant, the batch does not receive an additional period corresponding to the time established for the re-test period. However, an API batch can be re-tested multiple times and a different portion of the batch used after each re-test, as long as it continues to comply with the specification. For APIs known to be labile (e.g. certain antibiotics), it is more appropriate to establish a shelf-life rather than a re-test period (reference: ICH Q1A).

Limited extrapolation of the real time data from the long-term storage condition beyond the observed range to extend the re-test period can be undertaken at the time of assessment of the PD, if justified. Applicants should consult the ICH Q1E guideline for further details on the evaluation and extrapolation of results from stability data (e.g. if significant change was not observed within 6 months at accelerated condition and the data show little or no variability, the proposed re-test period could be up to two times the period covered by the long-term data, but should not exceed the long-term data by 12 months).

Reference documents: ICH Q1A, Q1B, Q1D, Q1E, WHO Technical Report Series, No. 953, Annex 2

3.2.5.7.2 Post-approval Stability Protocol and Stability Commitment (name, manufacturer)

The post-approval stability protocol and stability commitment should be provided.

Primary stability study commitment

When available long-term stability data on primary batches do not cover the proposed re-test period granted at the time of assessment of the PD, a commitment should be made to continue the stability studies in order to firmly establish the re-test period. A written commitment (signed and dated) to continue long-term testing over the re-test period should be included in the dossier when relevant.

Commitment stability studies

The long-term stability studies for the commitment batches should be conducted through the proposed re-test period on at least three production batches. Where stability data was not provided for three production batches, a written commitment (signed and dated) should be included in the dossier.

The stability protocol for the commitment batches should be provided and should include, but not be limited to, the following parameters:

- number of batch(es) and different batch sizes, if applicable;
- relevant physical, chemical, microbiological and biological test methods;
- acceptance criteria;
- reference to test methods;
- description of the container closure system(s);
- testing frequency;
description of the conditions of storage (standardized conditions for long-term testing as described in these guidelines and consistent with the API labelling, should be used);
and
other applicable parameters specific to the API.

Ongoing stability studies

The stability of the API should be monitored according to a continuous and appropriate programme that will permit the detection of any stability issue (e.g. changes in levels of degradation products). The purpose of the ongoing stability programme is to monitor the API and to determine that the API remains and can be expected to remain within the re-test period in all future batches.

At least one production batch per year of API (unless none is produced during that year) should be added to the stability monitoring programme and tested at least annually to confirm the stability. In certain situations, additional batches should be included. A written commitment (signed and dated) for ongoing stability studies should be included in the dossier.

Refer to WHO Technical Report Series, No. 953, Annex 2, Section 2.1.11 for further information on ongoing stability studies.

Any differences in the stability protocols used for the primary batches and those proposed for the commitment batches or ongoing batches should be scientifically justified.

Reference documents: ICH Q1A, Q1B, Q1D, Q1E, WHO Technical Report Series, No. 953, Annex 2

3.2.S.7.3 Stability Data (name, manufacturer)

Results of the stability studies (e.g., forced degradation studies and stress conditions) should be presented in an appropriate format such as tabular, graphical, or narrative.

Information on the analytical procedures used to generate the data and validation of these procedures should be included.

The actual stability results used to support the proposed re-test period should be included in the dossier. For quantitative tests (e.g. individual and total degradation product tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms”.

Reference documents: ICH Q1A, Q1B, Q1D, Q1E, Q2, WHO Technical Report Series, No. 953, Annex 2

3.2.P Drug product (or finished pharmaceutical product (FPP)) (name, dosage form)

3.2.P.1 Description and Composition of the FPP (name, dosage form)

A description of the FPP and its composition should be provided. The information provided should include, for example:

- Description of the dosage form
The description of the FPP should include the physical description, release mechanism (e.g. immediate, modified (delayed or extended)), as well as any other distinguishable characteristics, e.g.

“The proposed XYZ 50mg Tablets are available as white, oval, film-coated tablets, debossed with ‘50’ on one side and a break-line on the other side.

The proposed XYZ 100mg Tablets are available as yellow, round, film-coated tablets, debossed with ‘100’ on one side and plain on the other side.”

- **Composition, i.e., list of all components of the dosage form, and their amount on a per unit basis (including overages, if any), the function of the components, and a reference to their quality standards (e.g., compendial monographs or manufacturer’s specifications)**

The tables in the QOS-PD template should be used to summarize the composition of the FPP and express the quantity of each component on a per unit basis (e.g. mg per tablet, mg per ml, mg per vial) and percentage basis, including a statement of the total weight or measure of the dosage unit. The individual components for mixtures prepared in-house (e.g. coatings) should be included in the tables, where applicable.

*All components* used in the manufacturing process should be included, including those that may not be added to every batch (e.g. acid and alkali), those that may be removed during processing (e.g. solvents) and any others (e.g. nitrogen, silicon for stoppers). If the FPP is formulated using an active moiety, then the composition for the active ingredient should be clearly indicated (e.g. “1 mg of active ingredient base = 1.075 mg active ingredient hydrochloride”). All overages should be clearly indicated (e.g. “contains 2% overage of the API to compensate for manufacturing losses”).

The components should be declared by their proper or common names, quality standards (e.g. Ph.Int., Ph.Eur., BP, USP, JP, in-house) and, if applicable, their grades (e.g. “Microcrystalline Cellulose NF (PH 102)”) and special technical characteristics (e.g. lyophilized, micronized, solubilised, emulsified).

The function of each component (e.g. diluent/filler, binder, disintegrant, lubricant, glidant, granulating solvent, coating agent, antimicrobial preservative) should be stated. If an excipient performs multiple functions, the predominant function should be indicated.

The qualitative composition, including solvents, should be provided for all proprietary components or blends (e.g. capsule shells, colouring blends, imprinting inks). This information (excluding the solvents) is to be listed in the product information (e.g. summary of product characteristics, labelling, package leaflet).

- **Description of accompanying reconstitution diluent(s)**

For FPPs supplied with reconstitution diluent(s) that are commercially available or have been assessed and considered acceptable in connection with another approved PD, a brief description of the reconstitution diluents(s) should be provided.
For FPPs supplied with reconstitution diluent(s) that are not commercially available or have not been assessed and considered acceptable in connection with another approved PD, information on the diluent(s) should be provided in a separate FPP portion (“3.2.P”), as appropriate.

- Type of container and closure used for the dosage form and accompanying reconstitution diluent, if applicable

The container closure used for the FPP (and accompanying reconstitution diluent, if applicable) should be briefly described, with further details provided under 3.2.P.7 Container closure system, e.g.

“The product is available in HDPE bottles with polypropylene caps (in sizes of 100’s, 500’s and 1000’s) and in PVC/Aluminum foil unit dose blisters (in packages of 100’s (cards of 5x2, 10 cards per package).”

Reference documents: ICH Q6A

3.2.P.2 Pharmaceutical Development (name, dosage form)

The Pharmaceutical Development section should contain information on the development studies conducted to establish that the dosage form, the formulation, manufacturing process, container closure system, microbiological attributes and usage instructions are appropriate for the purpose specified in the product dossier. The studies described here are distinguished from routine control tests conducted according to specifications. Additionally, this section should identify and describe the formulation and process attributes (critical parameters) that can influence batch reproducibility, product performance and FPP quality. Supportive data and results from specific studies or published literature can be included within or attached to the Pharmaceutical Development section. Additional supportive data can be referenced to the relevant nonclinical or clinical sections of the product dossier.

Pharmaceutical development information should include, at a minimum:

- the definition of the quality target product profile (QTPP) as it relates to quality, safety and efficacy, considering for example the route of administration, dosage form, bioavailability, strength and stability;
- identification of the potential critical quality attributes (CQAs) of the FPP so as to adequately control the product characteristics that could have an impact on quality;
- discussion of the potential CQAs of the API(s), excipients and container closure system(s) including the selection of the type, grade and amount to deliver drug product of the desired quality;
- discussion of the selection criteria for the manufacturing process and the control strategy required to manufacture commercial lots meeting the QTPP in a consistent manner.

These features should be discussed as part of the product development using the principles of risk management over the entire lifecycle of the product (ref: ICH Q8).
For a discussion of additional pharmaceutical development issues specific to the development of FDCs, reference should be made to WHO Technical Report Series, No. 929, Annex 5, Section 6.3.2.

Reference documents: ICH Q6A, Q8, Q9, Q10

3.2.P.2.1 Components of the FPP (name, dosage form)

3.2.P.2.1.1 Active Pharmaceutical Ingredient (name, dosage form)

The compatibility of the API with excipients listed in 3.2.P.1 should be discussed. Additionally, key physicochemical characteristics (e.g., water content, solubility, particle size distribution, polymorphic or solid state form) of the API that can influence the performance of the FPP should be discussed. For fixed-dose combinations, the compatibility of APIs with each other should be discussed.

Physicochemical characteristics of the API may influence both the manufacturing capability and the performance of the FPP.

Guidance on compatibility studies is provided in Appendix 3 of the WHO Guidelines for registration of fixed-dose combination medicinal products (WHO Technical Report Series, No. 929, Annex 5, 2005). In addition to visual examination, chromatographic results (assay, purity) are required to demonstrate API-API and API-excipient compatibility. In general, API-excipient compatibility is not required to be established for specific excipients when evidence is provided (e.g. SmPC or product leaflet) that the excipients are present in the comparator product.

3.2.P.2.1.2 Excipients (name, dosage form)

The choice of excipients listed in 3.2.P.1, their concentration, their characteristics that can influence the FPP performance should be discussed relative to their respective functions.

When choosing excipients, those with a compendial monograph are generally preferred and may be required in certain jurisdictions. Other resources are available for information on acceptable excipients and their concentrations, such as the US-FDA IIG list and the Handbook of Pharmaceutical Excipients. Use of excipients in concentrations outside of established ranges is discouraged and generally requires justification. In addition, available guidelines should be referenced which address particular excipients to be avoided, for example azo colorants as listed in the EMA Guideline CPMP/463/00. Other guidance such as the WHO Guidelines on development of Paediatric Medicines (TRS 970, Annex 5) may provide useful general guidance in this regard.

Ranges or alternates for excipients are normally not accepted, unless supported by appropriate process validation data. Where relevant, compatibility study results (e.g. compatibility of a primary or secondary amine API with lactose) should be included to justify the choice of excipients. Specific details should be provided where necessary (e.g. use of potato or corn starch).
Where antioxidants are included in the formulation, the effectiveness of the proposed concentration of the antioxidant should be justified and verified by appropriate studies.

Antimicrobial preservatives are discussed in 3.2.P.2.5.

3.2.P.2.2 Finished Pharmaceutical Product (name, dosage form)

3.2.P.2.2.1 Formulation Development (name, dosage form)

A brief summary describing the development of the FPP should be provided, taking into consideration the proposed route of administration and usage. The differences between the comparative bioavailability or bio waiver formulations and the formulation (i.e., composition) described in 3.2.P.1 should be discussed. Results from comparative in vitro studies (e.g., dissolution) or comparative in vivo studies (e.g., bio equivalence) should be discussed, when appropriate.

The requirements for bioequivalence studies should be taken into consideration for example when formulating multiple strengths and/or when the product(s) may be eligible for a biow aiver. WHO reference documents (e.g., WHO Technical Report Series, No. 937, Annex 7) may be consulted.

Product scoring may be recommended or required, for example when division into fractional doses may be necessary according to approved posology.

If the proposed FPP is a functionally scored tablet, a study should be undertaken to ensure the uniformity of dose in the tablet fragments. The data provided in the PD should include a description of the test method, individual values, mean and relative standard deviation (RSD) of the results. Uniformity testing (i.e. content uniformity for split portions containing less than 5 mg or less than 5% of the weight of the dosage unit portion, or mass uniformity for other situations) should be performed on each split portion from a minimum of 10 randomly selected whole tablets. As an illustrative example, the number of units (i.e. the splits) would be 10 halves for bisected tablets (one half of each tablet is retained for the test) or 10 quarters for quadrisection tablets (one quarter of each tablet is retained for the test). At least one batch of each strength should be tested. Ideally, the study should cover a range of the hardness values. The splitting of the tablets should be performed in a manner that would be representative of that used by the consumer (e.g. manually split by hand).

The uniformity test on split portions can be demonstrated on a one-time basis and does not need to be added to the FPP specification(s). The tablet description in the FPP specification and in the product information (e.g. summary of product characteristics, labelling, package leaflet) should reflect the presence of a score.

If splitting of a tablet is intended for a paediatric dose, a demonstration of content uniformity of tablet fragments may be required.

Where relevant, labelling should state that the score line is only to facilitate breaking for ease of swallowing and not to divide into equal doses.

In vitro dissolution or drug release
A discussion should be included as to how the development of the formulation relates to development of the dissolution method(s) and the generation of the dissolution profile.

The results of studies justifying the choice of in vitro dissolution or drug release conditions (e.g. apparatus, rotation speed, medium) should be provided. Data should also be submitted to demonstrate whether the method is sensitive to changes in manufacturing processes and/or changes in grades and/or amounts of critical excipients and particle size where relevant. The dissolution method should be sensitive to any changes in the product that would result in a change in one or more of the pharmacokinetic parameters. Use of a single point test or a dissolution range should be justified based on the solubility and/or biopharmaceutical classification of the API.

For slower dissolving immediate-release products (e.g. Q=80% in 90 minutes), a second time point may be warranted (e.g. Q=60% in 45 minutes).

Modified-release FPPs should have a meaningful in vitro release rate (dissolution) test that is used for routine quality control. Preferably this test should possess in vitro-in vivo correlation. Results demonstrating the effect of pH on the dissolution profile should be submitted if appropriate for the type of dosage form.

For extended-release FPPs, the testing conditions should be set to cover the entire time period of expected release (e.g. at least three test intervals chosen for a 12-hour release and additional test intervals for longer duration of release). One of the test points should be at the early stage of drug release (e.g. within the first hour) to demonstrate absence of dose dumping. At each test period, upper and lower limits should be set for individual units. Generally, the acceptance range at each intermediate test point should not exceed 25% or ±12.5% of the targeted value. Dissolution results should be submitted for several lots, including those lots used for pharmacokinetic and bioavailability or bio waiver studies.

Recommendations for conducting and assessing comparative dissolution profiles can be found in Appendix 1.

3.2.P.2.2.2 Overtages (name, dosage form)

Any overages in the formulation(s) described in 3.2.P.1 should be justified.

Justification of an overage to compensate for loss during manufacture should be provided, including the step(s) where the loss occurs, the reasons for the loss and batch analysis release data (assay results).

Overages for the sole purpose of extending the shelf-life of the FPP are generally not acceptable.

3.2.P.2.2.3 Physicochemical and Biological Properties (name, dosage form)

Parameters relevant to the performance of the FPP, such as pH, ionic strength, dissolution, redispersion, reconstitution, particle size distribution, aggregation, polymorphism, rheological properties, biological activity or potency, and/or immunological activity, should be addressed.
3.2.P.2.3 Manufacturing Process Development (name, dosage form)

The selection and optimisation of the manufacturing process described in 3.2.P.3.3, in particular its critical aspects, should be explained. Where relevant, the method of sterilisation should be explained and justified.

Where relevant, justification for the selection of aseptic processing or other sterilization methods over terminal sterilization should be provided.

Differences between the manufacturing process(es) used to produce comparative bioavailability or biowaiver batches and the process described in 3.2.P.3.3 that can influence the performance of the product should be discussed.

The rationale for choosing the particular pharmaceutical product (e.g. dosage form, delivery system) should be provided. The scientific rationale for the choice of the manufacturing, filling and packaging processes that can influence FPP quality and performance should be explained (e.g. wet granulation using high shear granulator). API stress study results may be included in the rationale. Any developmental work undertaken to protect the FPP from deterioration should also be included (e.g. protection from light or moisture).

The scientific rationale for the selection, optimization and scale-up of the manufacturing process described in 3.2.P.3.3 should be explained, in particular the critical aspects (e.g. rate of addition of granulating fluid, massing time, granulation end-point). A discussion of the critical process parameters (CPP), controls and robustness with respect to the QTPP and CQA of the product should be included (ref: ICH Q8).

3.2.P.2.4 Container Closure System (name, dosage form)

The suitability of the container closure system (described in 3.2.P.7) used for the storage, transportation (shipping) and use of the FPP should be discussed. This discussion should consider, e.g., choice of materials, protection from moisture and light, compatibility of the materials of construction with the dosage form (including sorption to container and leaching) safety of materials of construction, and performance (such as reproducibility of the dose delivery from the device when presented as part of the FPP).

Testing requirements to verify the suitability of the container closure system contact material(s) depend on the dosage form and route of administration. The pharmacopoeias provide standards that are required for packaging materials, including for example the following:

<table>
<thead>
<tr>
<th>Material</th>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass containers</td>
<td>USP &lt;660&gt;</td>
</tr>
<tr>
<td></td>
<td>Ph. Eur 3.2.1</td>
</tr>
<tr>
<td>Plastic containers</td>
<td>Ph. Eur 3.2.2, 3.2.2.1</td>
</tr>
<tr>
<td></td>
<td>USP &lt;661&gt;, &lt;671&gt;</td>
</tr>
<tr>
<td>Rubber/Elastomeric closures</td>
<td>USP &lt;381&gt;</td>
</tr>
<tr>
<td></td>
<td>Ph. Eur 3.2.9</td>
</tr>
</tbody>
</table>
The following table outlines the general recommendations for the various dosage forms for one-time studies to establish the suitability of the container closure system contact materials.

<table>
<thead>
<tr>
<th>Solid oral products</th>
<th>Oral liquid and topical products</th>
<th>Sterile products (including ophthalmics)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description of any additional treatments*</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Extraction studies</td>
<td>---</td>
<td>X</td>
</tr>
<tr>
<td>Interaction studies (Migration/Sorption)</td>
<td>---</td>
<td>X</td>
</tr>
<tr>
<td>Moisture permeability</td>
<td>X (uptake)</td>
<td>X (usually loss)</td>
</tr>
<tr>
<td>Light transmission</td>
<td>X**</td>
<td>X</td>
</tr>
</tbody>
</table>

* X = information should be submitted
** --- = information does not need to be submitted
* X = information should be submitted
** --- = information does not need to be submitted
* X = information should be submitted
** --- = information does not need to be submitted

For solid oral dosage forms and solid APIs, compliance with regulations on food-contact plastic materials, (for example (EU) No. 10/2011) can be considered acceptable in lieu of extraction studies.

The suitability of the container closure system used for the storage, transportation (shipping) and use of any intermediate/in-process products (e.g. premixes, bulk FPP) should also be discussed.

For a device accompanying a multidose container, the results of a study should be provided demonstrating the reproducibility of the device (e.g. consistent delivery of the intended volume), generally at the lowest intended dose.

In accordance with the Ph.Int. general chapter Liquid Preparations for Oral Use:

“Each dose from a multidose container is administered by means of a device suitable for measuring the prescribed volume. The device is usually a spoon or a cup for volumes of 5 ml or multiples thereof, or an oral syringe for other volumes or, for oral drops, a suitable dropper.”

For a device accompanying a multidose container, the results of a study should be provided demonstrating the reproducibility of the device (e.g. consistent delivery of the intended volume), generally at the lowest intended dose.

A sample of the device should be provided with Module 1.

### 3.2.P.2.5 Microbiological Attributes (name, dosage form)

Where appropriate, the microbiological attributes of the dosage form should be discussed, including, for example, the rationale for not performing microbial limits testing for non-sterile products and the selection and effectiveness of preservative
systems in products containing antimicrobial preservatives. For sterile products, the integrity of the container closure system to prevent microbial contamination should be addressed.

Where an antimicrobial preservative is included in the formulation, the amount used should be justified by submission of results of the product formulated with different concentrations of the preservative(s) to demonstrate the least necessary but still effective concentration. The effectiveness of the agent should be justified and verified by appropriate studies (e.g. USP or Ph.Eur. general chapters on antimicrobial preservatives) using a batch of the FPP. If the lower limit for the proposed acceptance criterion for the assay of the preservative is less than 90.0%, the effectiveness of the agent should be established with a batch of the FPP containing a concentration of the antimicrobial preservative corresponding to the lower proposed acceptance criteria.

As outlined in the WHO stability guidelines (WHO Technical Report Series, No. 953, Annex 2, 2009), a single primary stability batch of the FPP should be tested for effectiveness of the antimicrobial preservative (in addition to preservative content) at the proposed shelf-life for verification purposes, regardless of whether there is a difference between the release and shelf-life acceptance criteria for preservative content.

3.2.P.2.6 Compatibility (name, dosage form)

The compatibility of the FPP with reconstitution diluent(s) or dosage devices (e.g., precipitation of API in solution, sorption on injection vessels, stability) should be addressed to provide appropriate and supportive information for the labeling.

Where a device is required for oral liquids or solids (e.g. solutions, emulsions, suspensions and powders/granules for such reconstitution) that are intended to be administered immediately after being added to the device, the compatibility studies mentioned in the following paragraphs are not required.

Where sterile, reconstituted products are to be further diluted, compatibility should be demonstrated with all diluents over the range of dilution proposed in the labelling. These studies should preferably be conducted on aged samples. Where the labelling does not specify the type of containers, compatibility (with respect to parameters such as appearance, pH, assay, levels of individual and total degradation products, subvisible particulate matter and extractables from the packaging components) should be demonstrated in glass, PVC and polyolefin containers. However, if one or more containers are identified in the labelling, compatibility of admixtures needs to be demonstrated only in the specified containers.

Studies should cover the duration of storage reported in the labelling (e.g. 24 hours under controlled room temperature and 72 hours under refrigeration). Where the labelling specifies co-administration with other FPPs, compatibility should be demonstrated with respect to the principal FPP as well as the co-administered FPP (i.e. in addition to other aforementioned parameters for the mixture, the assay and degradation levels of each co-administered FPP should be reported).

3.2.P.3 Manufacture (name, dosage form)

3.2.P.3.1 Manufacturer(s) (name, dosage form)
The name, address, and responsibility of each manufacturer, including contractors, and each proposed production site or facility involved in manufacturing and testing should be provided.

The facilities involved in the manufacturing, packaging, labelling and testing should be listed. If certain companies are responsible only for specific steps (e.g. manufacturing of an intermediate), this should be clearly indicated. (Ref: WHO good distribution practices for pharmaceutical products, WHO Technical Report Series, No. 957, Annex 5.)

The list of manufacturers/companies should specify the actual addresses of production or manufacturing site(s) involved (including block(s) and unit(s)), rather than the administrative offices.

For a mixture of an API with an excipient, the blending of the API with the excipient is considered to be the first step in the manufacture of the final product and therefore the mixture does not fall under the definition of an API. The only exceptions are in the cases where the API cannot exist on its own. Similarly, for a mixture of APIs, the blending of the APIs is considered to be the first step in the manufacture of the final product. Sites for such manufacturing steps should be included in this section.

A valid manufacturing authorization for pharmaceutical production is generally required and a marketing authorization may be required to demonstrate that the product is registered or licensed in accordance with national requirements (Module 1, 1.2.2).

For each site where the major production step(s) are carried out, when applicable, a WHO-type certificate of GMP may be required, issued by the competent authority in terms of the WHO Certification Scheme on the Quality of Pharmaceutical Products Moving in International Commerce (Module 1, 1.2.2).

Justification for any differences to the product in the country or countries issuing the WHO-type certificate(s)

When there are differences between the product for which this application is submitted and that marketed in the country/countries which provided the WHO-type certificate(s), data to support the applicability of the certificate(s) despite the differences should be provided. Depending on the case, it may be necessary to provide validation data for differences in site of manufacture, specifications, formulation, etc. Note that only minor differences are likely to be acceptable. Differences in container labelling need not normally be justified.

Regulatory situation in other countries

The countries should be listed in which this product has been granted a marketing authorization, this product has been withdrawn from the market and/or this application for marketing has been rejected, deferred or withdrawn (Module 1, 1.2.2).


3.2.P.3.2 Batch Formula (name, dosage form)
A batch formula should be provided that includes a list of all components of the dosage form to be used in the manufacturing process, their amounts on a per batch basis, including overages, and a reference to their quality standards.

The tables in the QOS-PD template should be used to summarize the batch formula of the FPP for each proposed commercial batch size and express the quantity of each component on a per batch basis, including a statement of the total weight or measure of the batch.

All components used in the manufacturing process should be included, including those that may not be added to every batch (e.g., acid and alkali), those that may be removed during processing (e.g., solvents) and any others (e.g., nitrogen, silicon for stoppers). If the FPP is formulated using an active moiety, then the composition for the active ingredient should be clearly indicated (e.g. “1 kg of active ingredient base = 1.075 kg active ingredient hydrochloride”). All overages should be clearly indicated (e.g. “Contains 5 kg (corresponding to 2%) overage of the API to compensate for manufacturing losses”).

The components should be declared by their proper or common names, quality standards (e.g. Ph.Int., Ph.Eur., BP, USP, JP, in-house) and, if applicable, their grades (e.g. “Microcrystalline Cellulose NF (PH 102)”) and special technical characteristics (e.g. lyophilized, micronized, solubilised, emulsified).

3.2.P.3.3 Description of Manufacturing Process and Process Controls (name, dosage form)

A flow diagram should be presented giving the steps of the process and showing where materials enter the process. The critical steps and points at which process controls, intermediate tests or final product controls are conducted should be identified.

A narrative description of the manufacturing process, including packaging, that represents the sequence of steps undertaken and the scale of production should also be provided. Novel processes or technologies and packaging operations that directly affect product quality should be described with a greater level of detail. Equipment should, at least, be identified by type (e.g., tumble blender, in-line homogeniser) and working capacity, where relevant.

Steps in the process should have the appropriate process parameters identified, such as time, temperature, or pH. Associated numeric values can be presented as an expected range. Numeric ranges for critical steps should be justified in Section 3.2.P.3.4. In certain cases, environmental conditions (e.g., low humidity for an effervescent product) should be stated.

The maximum holding time for bulk FPP prior to final packaging should be stated. The holding time should be supported by the submission of stability data, if longer than 30 days. For an aseptically processed FPP, sterile filtration of the bulk and filling into final containers should preferably be continuous; any holding time should be justified.

Proposals for the reprocessing of materials should be justified. Any data to support this justification should be either referenced or filed in this section (3.2.P.3.3).

The information above should be summarized in the QOS-PD template and should reflect the production of the proposed commercial batches.
For the manufacture of sterile products, the class (e.g. A, B, C etc.) of the areas should be stated for each activity (e.g. compounding, filling, sealing etc), as well as the sterilization parameters for equipment, container/closure, terminal sterilization etc.

Reference documents: ICH Q8, Q9, Q10

### 3.2.P.3.4 Controls of Critical Steps and Intermediates (name, dosage form)

**Critical Steps:** Tests and acceptance criteria should be provided (with justification, including experimental data) performed at the critical steps identified in 3.2.P.3.3 of the manufacturing process, to ensure that the process is controlled.

**Intermediates:** Information on the quality and control of intermediates isolated during the process should be provided.

Examples of applicable in-process controls include:

- granulations: moisture (limits expressed as a range), blend uniformity (e.g. low dose tablets), bulk and tapped densities, particle size distribution;
- solid oral products: average weight, weight variation, hardness, thickness, friability, and disintegration checked periodically throughout compression, weight gain during coating;
- semi-solids: viscosity, homogeneity, pH;
- transdermal dosage forms: assay of API-adhesive mixture, weight per area of coated patch without backing;
- metered dose inhalers: fill weight/volume, leak testing, valve delivery;
- dry powder inhalers: assay of API-excipient blend, moisture, weight variation of individually contained doses such as capsules or blisters;
- liquids: pH, specific gravity, clarity of solutions;
- parenterals: appearance, clarity, fill volume/weight, pH, filter integrity tests, particulate matter, leak testing of ampoules, pre-filtration and/or pre-sterilization bioburden testing.

Reference documents: ICH Q2, Q6A, Q8, Q9, Q10, WHO Technical Report Series, No. 929, Annex 5

### 3.2.P.3.5 Process Validation and/or Evaluation (name, dosage form)

Description, documentation, and results of the validation and/or evaluation studies should be provided for critical steps or critical assays used in the manufacturing process (e.g., validation of the sterilisation process or aseptic processing or filling). Viral safety evaluation should be provided in 3.2A.2, if necessary.

The following information should be provided for all products:

a) a copy of the process validation protocol, specific to this FPP, described below;

b) a commitment that three consecutive, production-scale batches of this FPP will be subjected to prospective validation in accordance with the above protocol;
applicant should submit a written commitment that information from these studies will be available for verification after approval by the NMRA inspection team;

c) if the process validation studies have already been conducted (e.g. for sterile products), a copy of the process validation report should be provided in the PD in lieu of (a) and (b) above.

One of the most practical forms of process validation, mainly for non-sterile products, is the final testing of the product to an extent greater than that required in routine quality control. It may involve extensive sampling, far beyond that called for in routine quality control and testing to normal quality control specifications and often for certain parameters only. Thus, for instance, several hundred tablets per batch may be weighed to determine unit dose uniformity. The results are then treated statistically to verify the "normality" of the distribution and to determine the standard deviation from the average weight. Confidence limits for individual results and for batch homogeneity are also estimated. Strong assurance is provided that samples taken at random will meet regulatory requirements if the confidence limits are well within compendial specifications.

Similarly, extensive sampling and testing may be performed with regard to any quality requirements. In addition, intermediate stages may be validated in the same way, e.g. dozens of samples may be assayed individually to validate mixing or granulation stages of low-dose tablet production by using the content uniformity test. Certain product characteristics may occasionally be skip tested. Thus, subvisual particulate matter in parenteral preparations may be determined by means of electronic devices, or tablets/capsules tested for dissolution profile if such tests are not performed on every batch.

Where ranges of batch sizes are proposed, it should be shown that variations in batch size would not adversely alter the characteristics of the finished product. It is envisaged that those parameters listed in the following validation scheme will need to be re-validated once further scale-up is proposed after approval.

The process validation protocol should include inter alia the following:

- a reference to the current master production document;
- a discussion of the critical equipment;
- the process parameters that can affect the quality of the FPP (critical process parameters (CPPs)) including challenge experiments and failure mode operation;
- details of the sampling: sampling points, stages of sampling, methods of sampling and the sampling plans (including schematics of blender/storage bins for uniformity testing of the final blend);
- the testing parameters/acceptance criteria including in-process and release specifications and including comparative dissolution profiles of validation batches against the batch(es) used in the bioavailability or biowaiver studies;
- the analytical procedures or a reference to appropriate section(s) of the dossier;
- the methods for recording/evaluating results;
- the proposed timeframe for completion of the protocol.

The manufacture of sterile FPPs needs a well-controlled manufacturing area (e.g. a strictly controlled environment, highly reliable procedures and appropriate in-process controls).
detailed description of these conditions, procedures and controls should be provided, together with actual copies of the following standard operating procedures:

a) washing, treatment, sterilization and depyrogenation of containers, closures and equipment;
b) filtration of solutions;
c) lyophilization process;
d) leaker test of filled and sealed ampoules;
e) final inspection of the product;
f) sterilization cycle.

The sterilization process used to destroy or remove microorganisms is probably the single most important process in the manufacture of parenteral FPPs. The process can make use of moist heat (e.g. steam), dry heat, filtration, gaseous sterilization (e.g. ethylene oxide), or radiation. It should be noted that terminal steam sterilization, when practical, is considered to be the method of choice to ensure sterility of the final FPP. Therefore, scientific justification for selecting any other method of sterilization should be provided.

The sterilization process should be described in detail and evidence should be provided to confirm that it will produce a sterile product with a high degree of reliability and that the physical and chemical properties as well as the safety of the FPP will not be affected. Details such as Fo range, temperature range and peak dwell time for an FPP and the container closure should be provided. Although standard autoclaving cycles of 121°C for 15 minutes or more would not need a detailed rationale, such justifications should be provided for reduced temperature cycles or elevated temperature cycles with shortened exposure times. If ethylene oxide is used, studies and acceptance criteria should control the levels of residual ethylene oxide and related compounds.

Filters used should be validated with respect to pore size, compatibility with the product, absence of extractables and lack of adsorption of the API or any of the components.

For the validation of aseptic processing of parenteral products that cannot be terminally sterilized, simulation process trials should be conducted. This involves filling containers with culture media under normal conditions, followed by incubation. Refer to current WHO GMP guidelines for details.

Reference documents: ICH Q8, Q9, Q10, WHO Technical Report Series, No. 961, Annex 3

3.2.P.4 Control of Excipients (name, dosage form)

3.2.P.4.1 Specifications (name, dosage form)

The specifications for excipients should be provided.

The specifications from the applicant or the FPP manufacturer should be provided for all excipients, including those that may not be added to every batch (e.g. acid and alkali), those that do not appear in the final FPP (e.g. solvents) and any others used in the manufacturing process (e.g. nitrogen, silicon for stoppers).

If the standard claimed for an excipient is an officially recognized compendial standard, it is sufficient to state that the excipient is tested according to the requirements of that standard,
rather than reproducing the specifications found in the officially recognized compendial monograph.

If the standard claimed for an excipient is a non-compendial standard (e.g. in-house standard) or includes tests that are supplementary to those appearing in the officially recognized compendial monograph, a copy of the specification for the excipient should be provided.

In general, excipients with an officially recognized pharmacopoeial monograph should be used. Exceptions should be justified.

For excipients of natural origin, microbial limit testing should be included in the specifications. Skip testing is acceptable if justified (submission of acceptable results of five production batches).

For oils of plant origin (e.g. soy bean oil, peanut oil) the absence of aflatoxins or biocides should be demonstrated.

The colours permitted for use should be limited to those listed in suitable guidelines such as the “Japanese pharmaceutical excipients”, the EU “List of permitted food colours”, and the FDA “Inactive ingredient guide”. For proprietary mixtures, the supplier’s product sheet with the qualitative formulation should be submitted, in addition to the FPP manufacturer’s specifications for the product including identification testing.

For flavours the qualitative composition should be submitted, as well as a declaration that the excipients comply with foodstuff regulations (e.g. USA or EU).

Information that is considered confidential may be submitted directly to the NMRA by the supplier with reference to the specific related product.

Other certifications of at-risk components may be required on a case-by-case basis.

If additional purification is undertaken on commercially available excipients details of the process of purification and modified specifications should be submitted.

Reference documents: ICH Q6A

3.2.P.4.2 Analytical Procedures (name, dosage form)

The analytical procedures used for testing the excipients should be provided, where appropriate.

Copies of analytical procedures from officially recognized compendial monographs do not need to be submitted.

Reference documents: ICH Q2

3.2.P.4.3 Validation of Analytical Procedures (name, dosage form)

Analytical validation information, including experimental data, for the analytical procedures used for testing the excipients should be provided, where appropriate.
Copies of analytical validation information are generally not submitted for the testing of excipients, with the exception of the validation of in-house methods where appropriate.

Reference documents: ICH Q2

3.2.P.4.4 Justification of Specifications (name, dosage form)

Justification for the proposed excipient specifications should be provided, where appropriate.

A discussion of the tests that are supplementary to those appearing in the officially recognized compendial monograph should be provided.

3.2.P.4.5 Excipients of Human or Animal Origin (name, dosage form)

For excipients of human or animal origin, information should be provided regarding adventitious agents (e.g., sources, specifications, description of the testing performed, viral safety data). (Details in 3.2.A.2).

The excipients to be addressed in this section may include gelatin, phosphates, stearic acid, magnesium stearate and other stearates. If from plant origin a declaration to this effect will suffice.

For these excipients from animal origin, a letter of attestation should be provided confirming that the excipients used to manufacture the FPP are without risk of transmitting agents of animal spongiform encephalopathies.

Materials of animal origin should be avoided whenever possible.

When available, a CEP demonstrating TSE-compliance should be provided. A complete copy of the CEP (including any annexes) should be provided in Module 1.

Reference documents: ICH Q5A, Q5D, Q6B, WHO Technical Report Series, No. 908, Annex 1

3.2.P.4.6 Novel Excipients (name, dosage form)

For excipient(s) used for the first time in an FPP or by a new route of administration, full details of manufacture, characterisation, and controls, with cross references to supporting safety data (nonclinical and/or clinical) should be provided according to the API and/or FPP format. (Details in 3.2.A.3).

At their discretion, an NMRA may choose not to accept the use of novel excipients in submitted PDs. For the purpose of these guidelines, a novel excipient is one that has not been used (at a similar level and by the same route of administration) in a product approved by an SRA or WHO. If novel excipients are accepted, full information should be provided in 3.2.A.3.

3.2.P.5 Control of FPP (name, dosage form)

3.2.P.5.1 Specification(s) (name, dosage form)

The specification(s) for the FPP should be provided.
As defined in ICH’s Q6A guideline, a specification is:

“a list of tests, references to analytical procedures and appropriate acceptance criteria, which are numerical limits, ranges, or other criteria for the tests described. It establishes the set of criteria to which an API or FPP should conform to be considered acceptable for its intended use. “Conformance to specifications” means that the API and/or FPP, when tested according to the listed analytical procedures, will meet the listed acceptance criteria. Specifications are critical quality standards that are proposed and justified by the manufacturer and approved by regulatory authorities.”

A copy of the FPP specification(s) from the applicant (as well as the company responsible for the batch release of the FPP, if different from the applicant), dated and signed by authorized personnel (i.e. the person in charge of the quality control or quality assurance department) should be provided in the PD. Two separate sets of specifications may be set out: after packaging of the FPP (release) and at the end of shelf-life.

The specifications should be summarized according to the tables in the QOS-PD template including the tests, acceptance criteria and analytical procedures (including types, sources and versions for the methods):

- the standard declared by the applicant could be an officially recognized compendial standard (e.g. Ph.Int., BP, USP, JP) or an in-house (manufacturer’s) standard;
- the specification reference number and version (e.g. revision number and/or date) should be provided for version control purposes;
- for the analytical procedures, the type should indicate the kind of analytical procedure used (e.g. visual, IR, UV, HPLC), the source refers to the origin of the analytical procedure (e.g. Ph.Int., Ph.Eur., BP, USP, JP, in-house) and the version (e.g. code number/version/date) should be provided for version control purposes.

ICH’s Q6A guideline outlines recommendations for a number of universal and specific tests and criteria for FPPs. Specifications should include, at minimum, tests for appearance, identification, assay, purity, performance tests (e.g. dissolution), physical tests (e.g. loss on drying, hardness, friability, particle size), uniformity of dosage units, and as applicable, identification and assay of antimicrobial or chemical preservatives (e.g. antioxidants) and microbial limit tests.

The following information provides guidance for specific tests that are not addressed by ICH’s Q6A guideline:

- fixed-dose combination FPPs (FDC-FPPs):
  - analytical methods that can distinguish each API in the presence of the other API(s) should be developed and validated,
  - acceptance criteria for degradation products should be established with reference to the API they are derived from. If an impurity results from a chemical reaction between two or more APIs, its acceptance limits should in general be calculated with reference to the worst case (the API with the smaller area under the curve). Alternatively the content of such impurities could be calculated in relation to their reference standards,
o a test and limit for content uniformity is required for each API present in the FPP at less than 5 mg or less than 5% of the weight of the dosage unit,
o for the API(s) present at equal or greater than 5 mg and equal or greater than 5% of the weight of the dosage unit, a test and limit for weight variation may be established in lieu of content uniformity testing;

- modified-release products: a meaningful API release method;
- inhalation and nasal products: consistency of delivered dose (throughout the use of the product), particle or droplet size distribution profiles (comparable to the product used in in vivo studies, where applicable) and if applicable for the dosage form, moisture content, leak rate, microbial limits, preservative assay, sterility and weight loss;
- suppositories: uniformity of dosage units, melting point;
- transdermal dosage forms: peel or shear force, mean weight per unit area, dissolution.

Unless there is appropriate justification, the generally accepted limit for the API content of the FPP in the release specifications is ± 5% of the label claim (i.e. 95.0-105.0%).

For products such as tablets, capsules and suppositories where a test for uniformity of single dose preparations is required, a test and limit for content uniformity is required when the API is present in the FPP at less than 5 mg or less than 5% of the weight of the dosage unit.

Otherwise, the test for mass uniformity may be applied.

Skip testing is generally acceptable for parameters such as identification of colouring materials and microbial limits, when justified by the submission of acceptable supportive results for five production batches. When skip testing justification has been accepted, the specifications should include a footnote, stating at minimum the following skip testing requirements: at minimum every tenth batch and at least one batch annually is tested. In addition, for stability-indicating parameters such as microbial limits, testing will be performed at release and shelf-life during stability studies.

Any differences between release and shelf-life tests and acceptance criteria should be clearly indicated and justified. Note that such differences for parameters such as dissolution are normally not accepted.

Reference documents: ICH Q3B, Q3C, Q6A

**3.2.P.5.2 Analytical Procedures (name, dosage form)**

The analytical procedures used for testing the FPP should be provided.

Copies of the in-house analytical procedures used during pharmaceutical development (if used to generate testing results provided in the PD) as well as those proposed for routine testing should be provided. Unless modified, it is not necessary to provide copies of officially recognized compendial analytical procedures.

Tables for summarizing a number of the different analytical procedures and validation information (e.g. HPLC assay/impurity methods) can be found in the 2.3.R Regional information section of the QOS-PD (i.e. 2.3.R.2). These tables may be used to summarize the analytical procedures used for determination of the assay, related substances and dissolution of the FPP.
Refer to section 3.2.S.4.2 of these guidelines for additional guidance on analytical procedures.

Reference documents: ICH Q2

3.2.P.5.3 Validation of Analytical Procedures (name, dosage form)

Analytical validation information, including experimental data, for the analytical procedures used for testing the FPP, should be provided.

Copies of the validation reports for the in-house analytical procedures used during pharmaceutical development (if used to support testing results provided in the PD) as well as those proposed for routine testing should be provided.

Tables for summarizing a number of the different analytical procedures and validation information (e.g. HPLC assay/impurity methods, GC methods) can be found in the 2.3.R Regional information section of the QOS-PD (i.e. 2.3.R.2). These tables may be used to summarize the validation information of the analytical procedures used for determination of the assay, related substances and dissolution of the FPP.

As recognized by regulatory authorities and pharmacopoeias themselves, verification of compendial methods may be necessary. The compendial methods, as published, are typically validated based on an API or an FPP originating from a specific manufacturer. Different sources of the same API or FPP can contain impurities and/or degradation products or excipients that were not considered during the development of the monograph. Therefore the monograph and compendial method(s) should be demonstrated suitable for the control of the proposed FPP.

For officially recognized compendial FPP assay methods, verification should include a demonstration of specificity, accuracy and repeatability (method precision). If an officially recognized compendial method is used to control related substances that are not specified in the monograph, full validation of the method is expected with respect to those related substances.

If an officially recognized compendial standard is claimed and an in-house method is used in lieu of the compendial method (e.g. for assay or for related compounds), equivalency of the in-house and compendial methods should be demonstrated. This could be accomplished by performing duplicate analyses of one sample by both methods and providing the results from the study. For related compound methods, the sample analyzed should be the placebo spiked with related compounds at concentrations equivalent to their specification limits.

Reference documents: ICH Q2

3.2.P.5.4 Batch Analyses (name, dosage form)

A description of batches and results of batch analyses should be provided.

Information should include strength and batch number, batch size, date and site of production and use (e.g. used in comparative bioavailability or biowaiver studies, preclinical and clinical studies (if relevant), stability, pilot, scale-up and, if available, production-scale batches) on relevant FPP batches used to establish the specification(s) and evaluate consistency in manufacturing.
Analytical results tested by the company responsible for the batch release of the FPP (generally, the applicant or the FPP manufacturer, if different from the applicant) should be provided for a minimum of three batches, or in the case of conventional dosage forms with APIs that are demonstrated to be stable, at least two batches of at least pilot-scale. The third batch can be smaller, if justified. These batches should be manufactured by a procedure fully representative of and simulating that to be applied to a full production-scale batch.

The testing results should include the batch(es) used in the comparative bioavailability or biowaiver studies. Copies of the certificates of analysis for these batches should be provided in the PD and the company responsible for generating the testing results should be identified.

The discussion of results should focus on observations noted for the various tests, rather than reporting comments such as “all tests meet specifications”. This should include ranges of analytical results, where relevant. For quantitative tests (e.g. individual and total impurity tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms” (e.g. “levels of degradation product A ranged from 0.2 to 0.4%”). Dissolution results should be expressed at minimum as both the average and range of individual results. Recommendations for conducting and assessing comparative dissolution profiles can be found in Appendix 1.

A discussion and justification should be provided for any incomplete analyses (e.g. results not tested according to the proposed specification).

Reference documents: ICH Q3B, Q3C, Q6A

3.2.P.5.5 Characterisation of Impurities (name, dosage form)

Information on the characterisation of impurities should be provided, if not previously provided in “3.2.S.3.2 Impurities”.

A discussion should be provided of all impurities that are potential degradation products (including those among the impurities identified in 3.2.S.3.2 as well as potential degradation products resulting from interaction of the API with other APIs (FDCs), excipients or the container closure system) and FPP process-related impurities (e.g. residual solvents in the manufacturing process for the FPP).

Reference documents: ICH Q3B, Q3C, Q6A

3.2.P.5.6 Justification of Specification(s) (name, dosage form)

Justification for the proposed FPP specification(s) should be provided.

A discussion should be provided on the omission or inclusion of certain tests, evolution of tests, analytical procedures and acceptance criteria, differences from the officially recognized compendial standard(s), etc. If the officially recognized compendial methods have been modified or replaced, a discussion should be included.

The justification for certain tests, analytical procedures and acceptance criteria (e.g. degradation products, dissolution method development) may have been discussed in other
sections of the PD and does not need to be repeated here, although a cross-reference to their location should be provided.

ICH Q6A should be consulted for the development of specifications for FPPs.

3.2.P.6 Reference Standards or Materials (name, dosage form)

Information on the reference standards or reference materials used for testing of the FPP should be provided, if not previously provided in “3.2.S.5 Reference Standards or Materials”.

See Section 3.2.S.5 for information that should be provided on reference standards or materials. Information should be provided on reference materials of FPP degradation products, where not included in 3.2.S.5.


3.2.P.7 Container Closure System (name, dosage form)

A description of the container closure systems should be provided, including the identity of materials of construction of each primary packaging component and its specification. The specifications should include description and identification (and critical dimensions, with drawings where appropriate). Non-compendial methods (with validation) should be included, where appropriate.

For non-functional secondary packaging components (e.g., those that neither provide additional protection nor serve to deliver the product), only a brief description should be provided. For functional secondary packaging components, additional information should be provided.

Suitability information should be located in 3.2.P.2.

The WHO Guidelines on packaging for pharmaceutical products (WHO Technical Report Series, No. 902, Annex 9, 2002) and the officially recognized pharmacopoeias should be consulted for recommendations on the packaging information for FPPs.

Descriptions, materials of construction and specifications (of the company responsible for packaging the FPP, generally the FPP manufacturer) should be provided for the packaging components that are:

- in direct contact with the dosage form (e.g. container, closure, liner, desiccant, filler);
- used for drug delivery (including the device(s) for multi-dose solutions, emulsions, suspensions and powders/granules for such);
- used as a protective barrier to help ensure stability or sterility;
- necessary to ensure FPP quality during storage and shipping.

Primary packaging components are those that are in direct contact with the API or FPP.
The specifications for the primary packaging components should include a specific test for identification (e.g. IR). Specifications for film and foil materials should include limits for thickness or area weight.

Information to establish the suitability (e.g. qualification) of the container closure system should be discussed in Section 3.2.P.2.4. Comparative studies may be warranted for certain changes in packaging components (e.g. comparative delivery study (droplet size) for a change in manufacturer of dropper tips).

3.2.P.8 Stability (name, dosage form)

3.2.P.8.1 Stability Summary and Conclusions (name, dosage form)

The types of studies conducted, protocols used, and the results of the studies should be summarised. The summary should include, for example, conclusions with respect to storage conditions and shelf-life, and, if applicable, in-use storage conditions and shelf-life.


As outlined in the WHO stability guidelines, the purpose of stability testing is to provide evidence of how the quality of an API or FPP varies with time under the influence of a variety of environmental factors such as temperature, humidity and light. The stability programme also includes the study of product-related factors that influence its quality, for example, interaction of API with excipients, container closure systems and packaging materials.

**Stress testing**

As outlined in the WHO stability guidelines, photostability testing should be conducted on at least one primary batch of the FPP if appropriate. If “protect from light” is stated in one of the officially recognised pharmacopoeia for the API or FPP, it is sufficient to state “protect from light” on labelling, in lieu of photostability studies, when the container closure system is shown to be light protective. Additional stress testing of specific types of dosage forms may be appropriate (e.g. cyclic studies for semi-solid products, freeze-thaw studies for liquid products).

**Accelerated, intermediate (if necessary) and long-term testing**

Stability data must demonstrate stability of the medicinal product throughout its intended shelf-life under the climatic conditions prevalent in the target countries. Merely applying the same requirements applicable to other markets could potentially lead to substandard products, e.g. stability studies conducted for countries in Climatic Zone I/II when the products are supplied in Climatic Zones III and IV countries. Refer to WHO Technical Report Series, No. 953, Annex 2, Appendix 1 for information on climatic zones.

Refer to WHO Technical Report Series, No. 953, Annex 2 for further information regarding the storage conditions, including the minimum data required at the time of submitting the dossier.
To establish the shelf-life, data from stability studies should be provided on a minimum of three batches, or in the case of conventional dosage forms with APIs that are demonstrated to be stable, at least two batches of at least pilot-scale. The third batch can be smaller, if justified. The batches should be of the same formulation and packaged in the same container closure system as proposed for marketing. The manufacturing process used should simulate that to be applied to production batches and should provide product of the same quality and meeting the same specification as that intended for marketing. Where possible, batches of the FPP should be manufactured using different batches of the API(s). Stability studies should be performed on each individual strength, dosage form and container type and size of the FPP unless bracketing or matrixing is applied.

The stability testing programme should be summarized and the results of stability testing should be reported in the dossier and summarized in the tables in the QOS-PD. Bracketing and matrixing of proportional strengths can be applied, if scientifically justified.

For sterile products sterility should be reported at the beginning and end of shelf-life. For parenteral products, subvisible particulate matter should be reported frequently, but not necessarily at every test interval. Bacterial endotoxins need only be reported at the initial test interval. Weight loss from plastic containers should be reported over the shelf-life.

Any in-use period and associated storage conditions should be justified with experimental data, for example after opening, reconstitution and/or dilution of any sterile and/or multidose products or after first opening of FPPs packed in bulk multidose containers (e.g. bottles of 1000s). If applicable, the in-use period and storage conditions should be stated in the product information.

The information on the stability studies should include details such as

- storage conditions;
- strength;
- batch number, including the API batch number(s) and manufacturer(s);
- batch size;
- container closure system including orientation (e.g. erect, inverted, on-side) where applicable (e.g. semi-solids and liquids in plastic containers);
- completed (and proposed) test intervals.

The discussion of results should focus on observations noted for the various tests, rather than reporting comments such as “all tests meet specifications”. This should include ranges of analytical results and any trends that were observed. For quantitative tests (e.g. individual and total degradation product tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms”.

Dissolution results should be expressed at minimum as both the average and range of individual results.

Applicants should consult ICH’s Q1E guideline for details on the evaluation and extrapolation of results from stability data (e.g. if significant change was not observed within 6 months at accelerated condition and the data show little or no variability, the proposed shelf-life could be up to two times the period covered by the long-term data, but should not exceed the long-term data by 12 months).
Proposed storage statement and shelf-life

The proposed storage statement and shelf-life (and in-use storage conditions and in-use period, if applicable) for the FPP should be provided.

The recommended labelling statements for use, based on the stability studies, are provided in the WHO stability guidelines.

Reference documents: WHO Technical Report Series, No. 953, Annex 2, ICH Q1A, Q1B, Q1C, Q1D, Q1E, Q3B, Q6A

3.2.P.8.2 Post-approval Stability Protocol and Stability Commitment (name, dosage form)

The post-approval stability protocol and stability commitment should be provided.

Primary stability study commitment

When available long-term stability data on primary batches do not cover the proposed shelf-life granted at the time of assessment of the PD, a commitment should be made to continue the stability studies in order to firmly establish the shelf-life. A written commitment (signed and dated) to continue long-term testing over the shelf-life period should be included in the dossier.

Commitment stability studies

The long-term stability studies for the Commitment batches should be conducted through the proposed shelf-life on at least three production batches of each strength in each container closure system. Where stability data was not provided for three production batches of each strength, a written commitment (signed and dated) should be included in the dossier.

Ongoing stability studies

As described in the WHO stability guidelines, an ongoing stability programme is established to monitor the product over its shelf-life and to determine that the product remains and can be expected to remain within specifications under the storage conditions on the label. Unless otherwise justified, at least one batch per year of product manufactured in every strength and every container closure system, if relevant, should be included in the stability programme (unless none is produced during that year). Bracketing and matrixing may be applicable. A written commitment (signed and dated) to this effect should be included in the dossier.

Any differences in the stability protocols used for the primary batches and those proposed for the commitment batches or ongoing batches should be scientifically justified.

Reference documents: ICH Q1A

3.2.P.8.3 Stability Data (name, dosage form)

Results of the stability studies should be presented in an appropriate format (e.g., tabular, graphical, narrative). Information on the analytical procedures used to generate the data and validation of these procedures should be included.
Information on characterisation of impurities is located in 3.2.P.5.5.

The actual stability results/reports used to support the proposed shelf-life should be provided in the PD. For quantitative tests (e.g. individual and total degradation product tests and assay tests), it should be ensured that actual numerical results are provided rather than vague statements such as “within limits” or “conforms”. Dissolution results should be expressed at minimum as both the average and range of individual results.

Reference documents: ICH Q1A, Q1B, Q1C, Q1D, Q1E, Q2

3.2.A Appendices

3.2.A.1 Facilities and equipment

Not applicable (i.e. not a biotech product).

3.2.A.2 Adventitious agents safety evaluation

3.2.A.3 Novel excipients

At their discretion, an NMRA may choose not to accept the use of novel excipients in submitted PDs. If novel excipients are accepted, full information should be provided in the format of the sections in 3.2.P.

3.2.R Regional Information

3.2.R.1 Production documentation

3.2.R.1.1 Executed production documents

A minimum of three batches, or in the case of conventional dosage forms with APIs that are demonstrated to be stable, at least two batches of at least pilot-scale, should be manufactured for each strength. The third batch can be smaller, if justified. These batches should be manufactured by a procedure fully representative of and simulating that to be applied to a full production-scale batch.

For solid oral dosage forms, pilot scale is generally, at a minimum, one-tenth that of full production scale or 100 000 tablets or capsules, whichever is the larger.

Copies of the executed production documents should be provided for the batches used in the comparative bioavailability or biowaiver studies. Any notations made by operators on the executed production documents should be clearly legible.

If not included in the executed batch records through sufficient in-process testing, data should be provided for the batch used in comparative bioavailability or biowaiver studies that demonstrates the uniformity of this batch. The data to establish the uniformity of the biobatch should involve testing to an extent greater than that required in routine quality control.

English translations of executed records should be provided, where relevant.

3.2.R.1.2 Master production documents
Copies of the FPP master production documents should be provided for each proposed strength, commercial batch size and manufacturing site.

The details in the master production documents should include, but not be limited to, the following:

a) master formula;

b) dispensing, processing and packaging sections with relevant material and operational details;

c) relevant calculations (e.g. if the amount of API is adjusted based on the assay results or on the anhydrous basis);

d) identification of all equipment by, at minimum, type and working capacity (including make, model and equipment number, where possible);

e) process parameters (e.g. mixing time, mixing speed, milling screen size, processing temperature range, granulation end-point, tablet machine speed (expressed as target and range));

f) list of in-process tests (e.g. appearance, pH, assay, blend uniformity, viscosity, particle size distribution, LOD, weight variation, hardness, disintegration time, weight gain during coating, leaker test, minimum fill, clarity, filter integrity checks) and specifications;

g) sampling plan with regard to the:

i. steps where sampling should be done (e.g. drying, lubrication, compression),

ii. number of samples that should be tested (e.g. for blend uniformity testing of low dose FPPs, blend drawn using a sampling thief from x positions in the blender),

iii. frequency of testing (e.g. weight variation every x minutes during compression or capsule filling);

h) precautions necessary to ensure product quality (e.g. temperature and humidity control, maximum holding times);

i) for sterile products, reference to SOPs in appropriate sections and a list of all relevant SOPs at the end of the document;

j) theoretical and actual yield;

k) compliance with the GMP requirements.


3.2.R.2 Analytical procedures and validation information
The tables presented in section 2.3.R.2 in the QOS-PD template may be used to summarize the analytical procedures and validation information from sections 3.2.S.4.2, 3.2.S.4.3, 3.2.S.4.4 (c), 2.3.S.7.3 (b), 3.2.P.5.2 and 3.2.P.5.3, where relevant.

4.3 Literature references

References to the scientific literature relating to both the API and FPP should be included in this section of the PD when appropriate.

5. REFERENCES


APPENDIX 1
RECOMMENDATIONS FOR CONDUCTING AND ASSESSING COMPARATIVE DISSOLUTION PROFILES

The dissolution measurements of the two FPPs (e.g. test and reference (comparator), or two different strengths) should be made under the same test conditions. A minimum of three time points (zero excluded) should be included, the time points for both reference and test product being the same. The sampling intervals should be short for a scientifically sound comparison of the profiles (e.g. 5, 10, 15, 20, 30, 45 (60, 90, 120) minutes). Inclusion of the 15 minute time point in the schedule is of strategic importance for profile similarity determinations (very rapidly dissolving scenario). For extended-release FPPs, the time points should be set to cover the entire time period of expected release, e.g. 1, 2, 3, 5 and 8 hours for a 12-hour release and additional test intervals for longer duration of release.

Studies should be performed in at least three (3) media covering the physiological range, including pH 1.2 hydrochloric acid, pH 4.5 buffer and pH 6.8 buffer. International Pharmacopoeia buffers are recommended; alternative compendia buffers with the same pH and buffer capacity are also accepted. Water may be considered as an additional medium, especially when the API is unstable in the buffered media to the extent that the data is unusable.

If both the test and reference products show more than 85% dissolution in 15 minutes, the profiles are considered similar (no calculations required). Otherwise:

- **Similarity** of the resulting comparative dissolution profiles should be calculated using the following equation that defines a similarity factor ($f_2$):

\[
f_2 = 50 \log \left\{\frac{1}{n} \sum_{t=1}^{n} (R_t - T_t)^2 \right\}^{0.5} \times 100
\]

where $R_t$ and $T_t$ are the mean percent API dissolved in reference and test product, respectively, at each time point. An $f_2$ value between 50 and 100 suggests the two dissolution profiles are similar;

- a maximum of one time-point should be considered after 85% dissolution of the reference product has been reached. In the case where 85% dissolution cannot be reached due to poor solubility of the API, the dissolution should be conducted until an asymptote (plateau) has been reached;

- at least 12 units should be used for each profile determination. Mean dissolution values can be used to estimate the similarity factor, $f_2$. To use mean data, the % coefficient of variation at the first time point should be not more than 20% and at other time points should be not more than 10%;

- when delayed-release products (e.g. enteric coated) are being compared, the recommended conditions are acid medium (pH 1.2) for 2 hours and buffer pH 6.8 medium;
• when comparing extended-release beaded capsules, where different strengths have been achieved solely by means of adjusting the number of beads containing the API, one condition (normally the release condition) will suffice;

• surfactants should be avoided in comparative dissolution testing. A statement that the API is not soluble in any of the media is not sufficient and profiles in absence of surfactant should be provided. The rationale for the choice and concentration of surfactant should be provided. The concentration of the surfactant should be such that the discriminatory power of the test will not be compromised.

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