

ICH – International Council for Harmonisation

Your Objectives:

At the end of the lesson, you should be able to summarise the role of ICH.

The International Council for Harmonisation of Technical Requirements for Pharmaceuticals

for Human Use (hereafter ICH) is unique in that it invites the
authorities and industry to discuss scientific and technical
 of pharmaceuticals and develop ICH guidelines.

Since its inception in 1990, the ICH has gradually evolved to respond to increasingly global
 in the pharmaceutical sector, and these ICH guidelines are
applied by a growing number of regulatory authorities, the aim being to achieve greater
 worldwide so as to ensure that safe and
, high-quality are developed and
that these be registered and maintained in the most vigilant yet resource-efficient way. Since
its organisational changes, in October of 2015, the ICH has grown as an organisation and
currently includes at least 17 members* and 32 observers*.

The ICH develops, under consensus, standardized, commendable
guidelines for the evaluation of the , efficacy and safety of
medicines in a multi-stage process. These include Good Clinical Practice (GCP) guidelines for
 trials of medicinal products, Good Manufacturing Practice (GMP)
guidelines for a flawless manufacturing practice. Guidelines for the standardization of
 terminology (MedDRA) and information transfer (ESTRI) in the
regulatory area are just two examples.

Also vital is a standardized for the submission of documents for approval, known as the **Common Technical Document** (CTD) - see Image 2. Within the EU, the ICH guidelines are adopted by the Committee for Medicinal Products for Human Use (CHMP) at the European Medicines Agency (EMA).

Note that ICH guidelines, although they are not mandatory as such, a commitment to them and due implementation on a national or regional does strengthen the ICH process. The guidelines, nevertheless, should only be deviated from in justifiable cases. The ICH topics fall into four (4) categories of guidelines, and **topic codes** are assigned accordingly:

- Safety
- Quality
- Efficacy
- Multidisciplinary

Process of Harmonisation

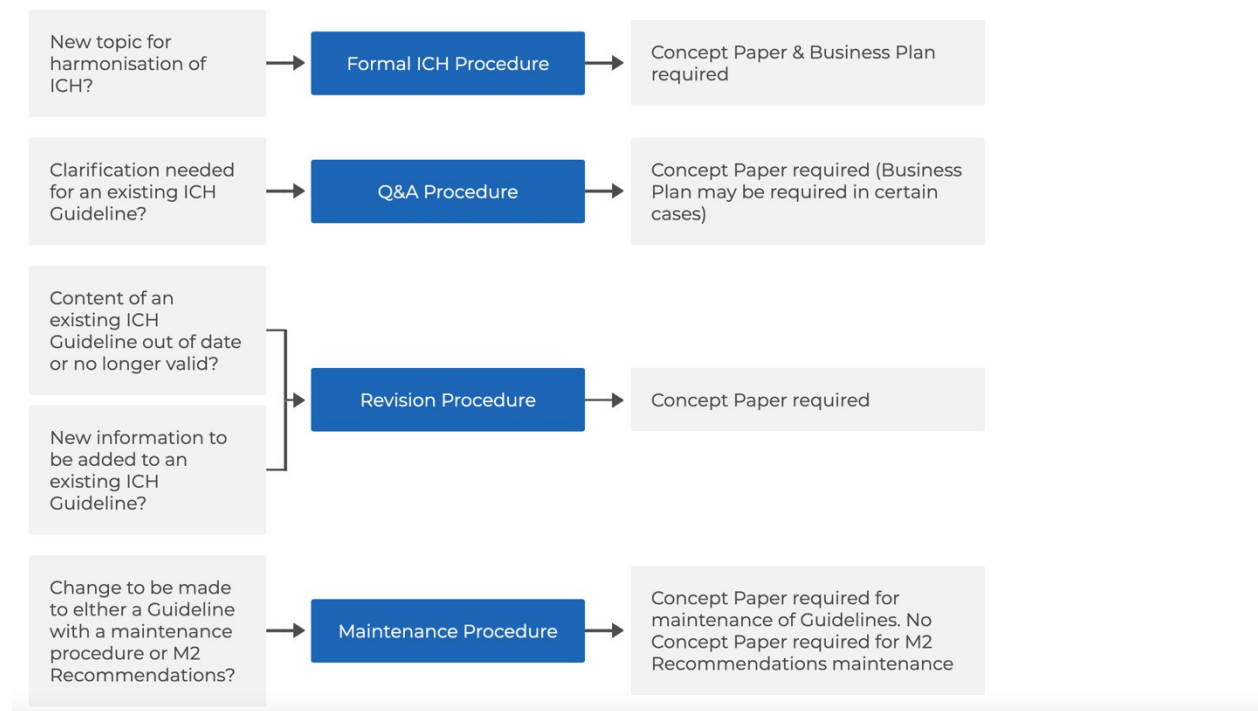
ICH harmonisation activities, too, fall into four (4) categories of procedures:

- Formal ICH,
- Q&A, (question & answer)
- Revision, and
- Maintenance, depending on the activity to be undertaken -see Image 1 on Page 2.

Each harmonisation activity is initiated by a **concept paper**, which is a short of the proposal. Depending on the category of harmonisation activity, a **business** may also be required. This would outline the costs and benefits of harmonising the topic proposed by the concept paper.

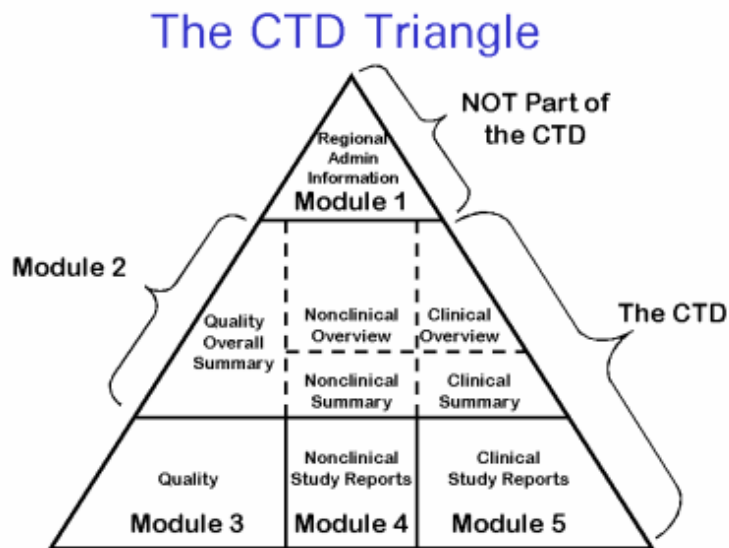
* For the list of founding and regulatory members and legislative and administrative observers, go here: <https://www.ich.org/page/members-observers>

Image 1:



source: <https://www.ich.org/page/process-harmonisation>

Image 2:



source: Wikipedia: https://en.wikipedia.org/wiki/Common_Technical_Document

Aufgabe Lückentext:

Folgende Wörter bitte in den Lückentext einfüllen.

Jedes Wort kommt einmal vor.

Bitte Gross- und Kleinbuchstaben beachten.

aspects, clinical, developments, effective, format, harmonisation, level, medicines, medical, pharmaceutical, plan, quality, regulatory, scientific, summary