ICH – International Council for Harmonisation

Your	Obje	ctives:

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

The International	Council for	Harmonisation	of Tech	nical Requ	irements fo	or Pharm	aceutica	ıls
for Human Use	(hereafter I	CH) is unique ir	that it	invites th	e			
authorities and			indu	stry to d	iscuss scie	ntific ar	nd techr	nical
		of pharmaceuti	cals and	l develop I	CH guidelin	es.		
Since its inceptio	n in 1990,	the ICH has gra	dually 6	evolved to	respond t	o increas	singly glo	obal
		in the pharm	aceutica	al sector,	and these	ICH gu	idelines	are
applied by a gro	wing numb	er of regulatory	y autho	rities, the	aim being	to achi	ieve gre	ater
		worldwide	so	as to	ensure	that	safe	and
		, high-quality				are de	veloped	and
that these be reg its organisational currently includes	changes, i	n October of 20	015, the	e ICH has			•	
The ICH develops	, under			consens	us, standar	dized, co	ommend	able
guidelines for th	e evaluatio	n of the			, ef	ficacy ar	nd safet	y of
medicines in a m	ulti-stage p	rocess. These in	clude G	ood Clinic	al Practice	(GCP) gı	uidelines	for
		trials of medici	nal prod	lucts, Goo	d Manufact	uring Pra	actice (G	MP)
guidelines for a	flawless r	_ nanufacturing p	ractice.	Guidelin	es for the	standa	rdizatior	າ of
		terminology (N	∕ledDRA	and info	rmation tr	ansfer (E	STRI) in	the
regulatory area as	re iust two e	z zamnles						

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 ($\rm CHMP$) at the European Medicines Agency ($\rm EMA$).						
Note that ICH guidelines, although they are not mandato	ory 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) catego	ries of procedures:					
 Formal ICH, Q&A, (question & answer) Revision, and Maintenance, depending on the activity to be under 	dertaken -see Image 1 on Page 2.					

activity, a business

Each harmonisation activity is initiated by a concept paper, which is a short

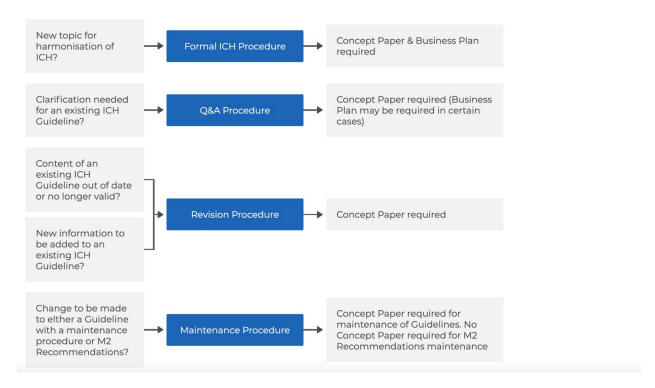
of the proposal. Depending on the category of harmonisation

may also be required. This would outline the

costs and benefits of harmonising the topic proposed by the concept paper.

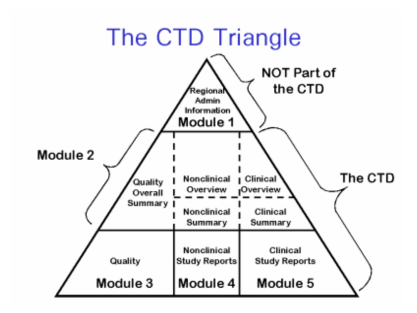
^{*} For the list of founding and regulatory members and legislative and administrative observers, go here: https://www.ich.org/page/members-observers

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