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4 **ICH guideline Q12 on technical and regulatory**
5 **considerations for pharmaceutical product lifecycle**
6 **management**
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15 ICH guideline Q12 on technical and regulatory
16 considerations for pharmaceutical product lifecycle
17 management

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58 **1. Introduction**

59 **1.1. Objectives**

60 The concepts outlined in prior ICH Quality Guidelines (ICH Q8, Q9, Q10 and Q11) provide opportunities
61 for science and risk-based approaches for drug development and risk-based regulatory decisions.
62 These guidelines are valuable in the assessment of Chemistry, Manufacturing and Controls (CMC)
63 changes across the product lifecycle. ICH Q8 and Q11 guidelines focus mostly on early stage aspects of
64 the product lifecycle (i.e., product development, registration and launch). Experience with
65 implementation of recent ICH guidelines has revealed technical and regulatory gaps that limit the full
66 realisation of more flexible regulatory approaches to post-approval CMC changes as described in ICH
67 Q8 (R2) and Q10 Annex I. This guideline addresses the commercial phase of the product lifecycle (as
68 described in ICH Q10).

69 A harmonised approach regarding technical and regulatory considerations for lifecycle management will
70 benefit patients, industry, and regulatory authorities by promoting innovation and continual
71 improvement in the biopharmaceutical sector, strengthening quality assurance and improving supply of
72 medicinal products.

73 This guideline provides a framework to facilitate the management of post-approval CMC changes in a
74 more predictable and efficient manner. It is also intended to demonstrate how increased product and
75 process knowledge can contribute to a reduction in the number of regulatory submissions. Effective
76 implementation of the tools and enablers described in this guideline should enhance industry's ability
77 to manage many CMC changes effectively under the firm's Pharmaceutical Quality System (PQS) with
78 less need for extensive regulatory oversight prior to implementation. The extent of operational and
79 regulatory flexibility is subject to product and process understanding (ICH Q8 and Q11), application of
80 risk management principles (ICH Q9), and an effective pharmaceutical quality system (ICH Q10).

81 In certain ICH regions, the current ICH Q12 guideline is not fully compatible with the established legal
82 framework with regard to the use of explicit Established Conditions ('EC') referred to in Chapter 3 and
83 with the Product Lifecycle Management ('PLCM') referred to in Chapter 5 as outlined in this guideline.
84 These concepts will, however, be considered when the legal frameworks will be reviewed and, in the
85 interim, to the extent possible under the existing regulation in these ICH regions.

86 **1.2. Scope**

87 This guideline applies to pharmaceutical drug substances (i.e., active pharmaceutical ingredients) and
88 pharmaceutical drug products, including marketed chemical, and biotechnological/biological products.
89 The guideline also applies to drug-device combination products that meet the definition of a
90 pharmaceutical or biotechnological/biological product. Changes needed to comply with revisions to
91 Pharmacopoeial monographs are not in scope of this guideline.

92 **1.3. ICH Q12 regulatory tools and enablers**

93 Use of the following harmonised regulatory tools and enablers with associated guiding principles, as
94 described in this guideline, will enhance the management of post-approval changes, and transparency
95 between industry and regulatory authorities, leading to innovation and continual improvement.

- 96 • Categorisation of Post-Approval CMC Changes ([Chapter 2](#))

97 Categorisation of Post-Approval CMC Changes is a framework that encompasses a risk-based
98 categorisation for the type of communication expected of the Marketing Authorisation Holder
99 (MAH) with the regulatory authority regarding CMC changes.

100 • Established Conditions (ECs) ([Chapter 3](#))

101 The concept of ECs provides a clear understanding between the MAH and regulatory authorities
102 regarding the necessary elements to assure product quality and identify the elements that require
103 a regulatory submission, if changed. This guideline describes how ECs are identified as well as what
104 information can be designated as supportive information that would not require a regulatory
105 submission, if changed. In addition, guidance is included for managing revisions of the ECs over a
106 product's lifecycle.

107 • Post-Approval Change Management Protocol (PACMP) ([Chapter 4](#))

108 The PACMP is a regulatory tool that provides predictability regarding the information required to
109 support a CMC change and the type of regulatory submission based on prior agreement between
110 the MAH and regulatory authority. Such a mechanism enables planning and implementation of
111 future changes to ECs in an efficient and predictable manner.

112 • Product Lifecycle Management (PLCM) ([Chapter 5](#))

113 The PLCM document serves as a central repository for the ECs and the associated reporting
114 category for changes made to ECs. The document also captures how a product will be managed
115 during the commercial phase of the lifecycle including relevant post-approval CMC commitments
116 and PACMPs.

117 • Pharmaceutical Quality System (PQS) and Change Management ([Chapter 6](#))

118 An effective PQS as described in ICH Q10 and compliance with regional GMPs are necessary for
119 implementation of this guideline. In particular, management of manufacturing changes across the
120 supply chain is an essential part of an effective change management system. This guideline
121 provides recommendations for robust change management across multiple entities involved in the
122 manufacture of a pharmaceutical product.

123 • Relationship between Regulatory Assessment and Inspection ([Chapter 7](#))

124 This guideline outlines the complementary roles of regulatory assessment and inspection, and how
125 communication between assessors and inspectors facilitates the use of the tools included herein.

126 • Post-Approval Changes for Marketed Products ([Chapter 8](#))

127 Approaches to facilitate changes to marketed products are outlined. This guideline provides
128 detailed guidance to enable changes to analytical methods to be made with immediate or other
129 post-implementation notification. Science- and risk-based approaches for stability studies in
130 support of the evaluation of CMC changes are also described.

131 The tools and enablers described above are complementary and are intended to link different phases of
132 the product lifecycle. Pharmaceutical development activities result in an appropriate control strategy,
133 elements of which are considered to be **Established Conditions**. All changes to an approved product
134 are managed through a firm's **Pharmaceutical Quality System**; changes to ECs must also be
135 reported to the regulatory authority. Where the regulatory system provides for **Categorisation of**
136 **Post-approval CMC Changes** for reporting according to risk, the MAH may propose reporting
137 categories for changes to ECs based on risk and knowledge gained through enhanced pharmaceutical

138 development. A system with risk-based reporting categories also facilitates the use of **Post-Approval**
139 **Change Management Protocols**, which provide predictability regarding planning for future changes
140 to ECs. The **Product Lifecycle Management** document is a summary that transparently conveys to
141 the regulatory authority how the MAH plans to manage post-approval CMC changes. The tools and
142 enablers in this guideline do not change the **Relationship between Regulatory Assessment and**
143 **Inspection**; however, collaboration and communication between assessors and inspectors are
144 necessary for the implementation of this guideline. Finally, this guideline proposes approaches to
145 facilitate **Post-Approval Changes to Marketed Products** without the need for regulatory review and
146 approval prior to implementation of certain CMC changes.

147 **2. Categorisation of post-approval CMC changes**

148 Regulatory mechanisms that allow the timely and efficient introduction of CMC changes are important
149 to drug quality, safety, and availability. There is a range of potential CMC changes for which
150 communication between a firm and the regulatory authority is required. CMC changes vary from low to
151 high potential risk with respect to product quality. A well-characterised, risk-based categorisation of
152 regulatory communication requirements is important to the efficient use of industry and regulatory
153 resources.

154 In such a regulatory system, the types of changes in the drug substance, drug product, production
155 process, quality controls, equipment, and facility that invoke communication with regulatory authorities
156 are classified with regard to the potential to have an adverse effect on product quality of the drug
157 product. The regulatory communication category, supporting information/documentation requirements,
158 and associated time frame for evaluation are commensurate with that potential risk.

159 Regulatory authorities are encouraged to utilise a system that incorporates risk-based regulatory
160 processes for (a) requesting approval from the regulatory authority, (b) notifying the regulatory
161 authority, or (c) simply recording CMC changes, with associated information requirements and, where
162 applicable, timeframes for decision. Such a system would include the following categories for
163 regulatory communications with one or more levels in each case:

- 164 • **Prior-approval:** Certain changes are considered to have sufficient risk to require regulatory
165 authority review and approval prior to implementation and are requested by the MAH in a suitably
166 detailed regulatory submission. An inspection may be associated with such changes.
- 167 • **Notification:** Certain moderate- to low-risk changes are judged to not require prior approval and
168 generally require less information to support the change. These changes are communicated to the
169 regulatory authority as a formal notification that takes place within a defined period of time before
170 or after implementation, according to regional requirements. A mechanism for immediate
171 notification is useful when prior approval is not required, but timely awareness of the change by
172 the regulator is considered necessary.

173 In addition, the lowest risk changes are only managed and documented within the PQS and not
174 reported to regulators, but may be verified on routine inspection.

175 Harmonisation or convergence toward a system of risk-based categorisation of post-approval changes
176 is encouraged as an important step toward achieving the objectives of this guideline. Such a system
177 provides inherent, valuable flexibility in regulatory approach and a framework that can support
178 additional regulatory opportunities such as:

- 179 • Facilitating the use of tools and enablers described in this guideline by providing a range of request
180 and notification categories available as a target for a lowering of regulatory submission
181 requirements.
- 182 • The use of a lower category for request/notification if certain criteria/conditions are met and the
183 relevant supporting documentation is provided as described in regional regulatory guidance; the
184 need for regulatory inspection associated with the change may preclude the ability to use a lower
185 category.
- 186 • Options for possible regulatory convergence regarding the association of a certain type of change
187 with a particular category when reasons for being different from other regulatory authorities are
188 not clearly established.

189 A risk-based categorisation system may be accomplished by having the principles captured in
190 regulations with further details in guidance, which can provide additional flexibility to modify
191 expectations as science and technology evolve. For examples of risk-based categorisation systems,
192 refer to existing regulations and guidance of ICH members, and WHO guidelines and guidance on
193 changes to approved products.

194 **3. Established conditions (ECs)**

195 **3.1. Introduction**

196 Although the Common Technical Document (CTD) format has been defined for a marketing application,
197 there are no previously harmonised approaches to defining which elements in an application are
198 considered necessary to assure product quality and therefore would require a regulatory submission if
199 changed post-approval. These elements are being defined in this guideline as “Established Conditions
200 for Manufacturing and Control” (referred to as ECs throughout this guideline).

201 **3.2. Definition of ECs and their role in the regulatory submission**

202 **3.2.1. ECs definition**

203 ECs are legally binding information (or approved matters) considered necessary to assure product
204 quality. As a consequence, any change to ECs necessitates a submission to the regulatory authority.

205 **3.2.2. ECs in a regulatory submission**

206 All regulatory submissions contain a combination of ECs and supportive information (refer to [Appendix](#)
207 [1](#)). Supportive information is not considered to be ECs, but is provided to share with regulators the
208 development and manufacturing information at an appropriate level of detail, and to justify the initial
209 selection of ECs and their reporting category.

210 ECs should not be confused with CMC regulatory commitments (e.g., stability and other commitments)
211 made by a MAH to provide data or information to the regulatory agency in a marketing authorisation
212 application (MAA). Such information, in the context of this guideline, is considered supportive
213 information. Changes to CMC regulatory commitments are not addressed in this guideline, but are
214 managed according to existing regional regulations and guidance.

215 ECs in a submission are either implicit or explicit:

216 • Implicit ECs are elements that are not specifically proposed by the MAH but are derived from and
217 revised according to regional regulation or guidance related to post-approval changes.

218 • Explicit ECs are specifically identified and proposed by the MAH together with their proposed
219 reporting category as part of a regulatory submission (see [Chapter 3.2.3](#)). This guideline provides
220 the opportunity to identify explicit ECs and associated reporting categories. Unless otherwise
221 specified by regional requirement, identifying explicit ECs for a given product is not mandatory.

222 An MAH may use one or both approaches as described above to define ECs and their associated
223 reporting categories. If the MAH wishes to propose a different reporting category than provided in
224 regional regulation and guidance for an implicit EC, the explicit EC approach should be used.

225 The MAH should provide rationales for the ECs and associated reporting categories in the appropriate
226 CTD sections in Module 3.

227 See [Appendix 1](#) for more information regarding sections of the marketing application that may contain
228 ECs and supportive information.

229 **3.2.3. Identification of ECs**

230 This chapter outlines approaches to define ECs for manufacturing processes and analytical methods. A
231 similar approach can be used to define other types of ECs (e.g., performance of the container closure
232 system) and should be justified by the applicant and approved by the regulatory agency.

233 The extent of ECs may vary based on the firm's development approach and potential risk to product
234 quality.

235 **3.2.3.1. Identification of ECs for the manufacturing processes**

236 In addition to the unit operation and the sequence of steps, and in considering the overall control
237 strategy, ECs proposed and justified in a manufacturing process description should be those inputs
238 (e.g., process parameters, material attributes) and outputs (that may include in-process controls) that
239 are necessary to assure product quality. These should include critical process parameters (CPPs, as
240 defined in ICH Q8(R2)), as well as key process parameters (KPPs), which are parameters of the
241 manufacturing process that may not be directly linked to critical product quality attributes, but need to
242 be tightly controlled to assure process consistency as it relates to product quality.

243 The details of ECs and the associated reporting category will depend on the extent to which the firm
244 can apply knowledge from product and process understanding (i.e., their development approach) to
245 manage the risks to product quality. Appropriate justification should be provided to support the
246 identification of ECs and proposed reporting categories. Different approaches can be used alone, or in
247 combination, to identify ECs for manufacturing processes; these include, but are not limited to the
248 following:

249 • A **parameter based approach**, in which product development prior to regulatory submission
250 provides a limited understanding of the relationship between inputs and resulting quality
251 attributes, will include a large number of inputs (e.g., process parameters and material attributes)
252 along with outputs (including in-process controls).

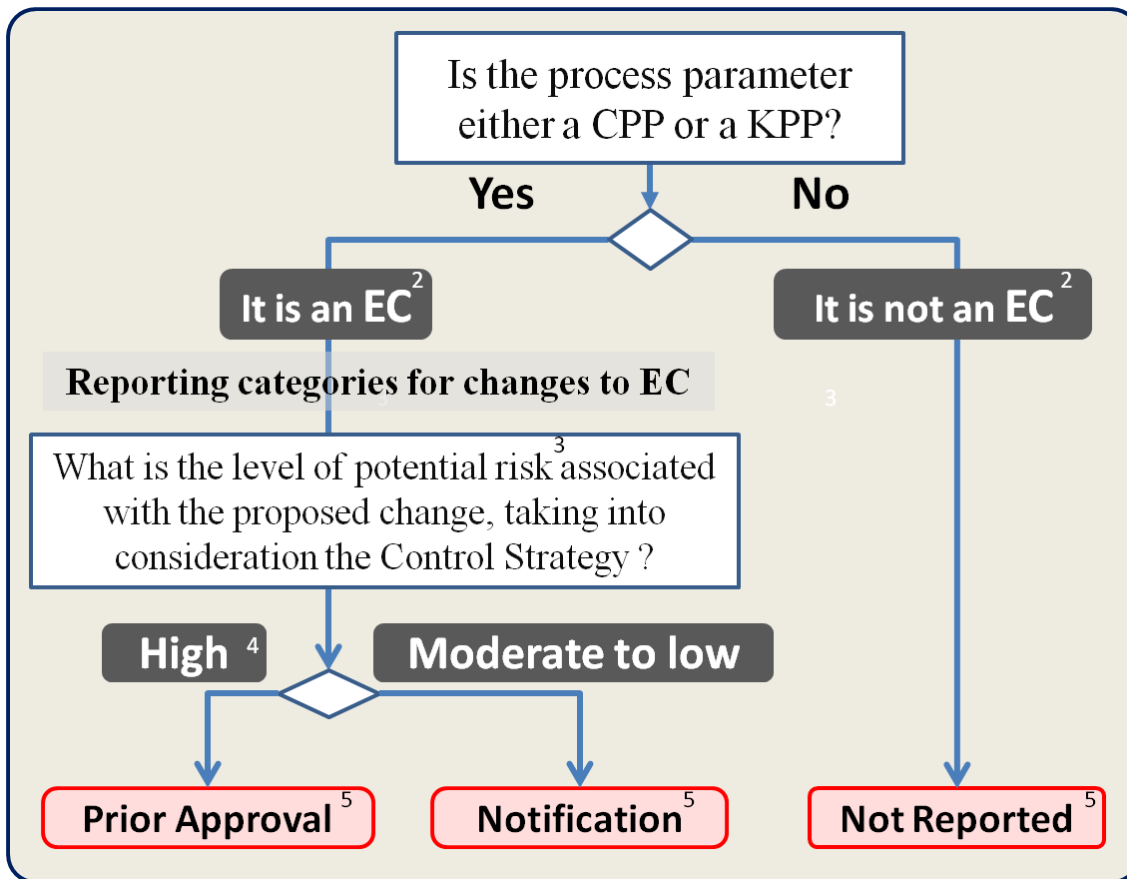
253 • An **enhanced approach** with increased understanding of interaction between inputs and product
254 quality attributes together with a corresponding control strategy can lead to identification of ECs
255 that are focused on the most important input parameters along with outputs, as appropriate.

256 • In certain cases, applying knowledge from a data-rich environment enables a **performance based**
257 **approach** in which ECs could be primarily focused on control of unit operation outputs rather than
258 process inputs (e.g., process parameters and material attributes). For example, a performance-
259 based approach could be considered for manufacturing process steps with in-line continuous
260 monitoring (e.g., using appropriate process analytical technologies such as NIR for the control of a
261 blending process).

262 When considering this approach, it is important to ensure that all relevant parameters and material
263 attributes that have a potential to impact product quality are monitored and equipment used remains
264 qualified in order to assure a stable process. In certain cases, such as a path-dependent process where
265 a specific outcome cannot be defined (e.g., fluid bed granulation and drying), select parameters or
266 attributes may need to be specified as ECs (e.g., differences in granular properties can affect the final
267 product quality).

268 A suitably detailed description of the manufacturing process is important to provide a clear
269 understanding of what is and is not necessary to assure product quality. Use of this guidance should
270 not lead to a less detailed description of the manufacturing process in Module 3 of the CTD.

271 A decision tree to identify ECs and associated reporting categories for manufacturing process
272 parameters is shown in Figure 1. This decision tree is intended to guide the identification of ECs based
273 on an assessment of criticality (i.e., CPPs) or impact on the process consistency as it relates to product
274 quality (i.e., KPPs). The corresponding reporting category is dependent on the potential risk to quality.
275 Risk assessment activities should follow approaches described in ICH Q9. In assessing the risk and
276 subsequent reporting category, an MAH should consider the overall control strategy and any possible
277 concurrent changes. Appropriate justification should be provided in support of the identification of ECs
278 and those aspects that are not ECs.



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281 **Figure 1.** Figure 1. Decision Tree for Identification of ECs and Associated Reporting Categories for
 282 Manufacturing Process Parameters⁵

283 Information regarding product-specific post-approval change activities, such as post-change
 284 monitoring, may be provided as supporting information to aid in the determination of ECs and
 285 associated reporting categories.

286 Criticality and risk should be evaluated periodically during the lifecycle of the product and, using the
 287 decision tree, the ECs should be updated based on acquired knowledge.

288 Additionally, an MAH should consider the impact of concurrent changes when assessing the appropriate
 289 reporting category.

290 **3.2.3.2. Identification of ECs for analytical procedures**

291 ECs related to analytical procedures should include elements which assure performance of the
 292 procedure. Appropriate justification should be provided to support the identification of ECs for
 293 analytical procedures. The extent of ECs could vary based on the method complexity, development and
 294 control approaches.

¹ This diagram does not apply as is for the performance-based approach.

² Appropriate justification is expected for ECs and non-ECs

³ Assessment of risk to quality using tools and concepts found in ICH Q9

⁴ In some cases, moderate risk changes may require prior approval.

⁵ See [Chapter 2](#) for further guidance on reporting categories and see [Chapter 3.3](#), regarding roles and responsibilities related to managing changes and maintaining an approved application.

295 • Where the relationship between method parameters and method performance has not been fully
296 studied at the time of submission, ECs will incorporate the details of operational parameters
297 including system suitability.

298 • When there is an increased understanding of the relationship between method parameters and
299 method performance defined by a systematic development approach including robustness studies,
300 ECs are focused on method-specific performance criteria (e.g., specificity, accuracy, precision)
301 rather than a detailed description of the analytical procedure.

302 A suitably detailed description of the analytical procedures in Module 3 is expected to provide a clear
303 understanding regardless of the approach used to identify ECs for analytical procedures. Use of this
304 guideline should not lead to providing a less detailed description of analytical procedures in the MAA.

305 **3.2.4. Revision of ECs**

306 It may be necessary to change approved ECs as a result of knowledge gained during the product
307 lifecycle (e.g., manufacturing experience, introduction of new technologies or changes in the control
308 strategy).

309 Options available for the MAH to change approved ECs, and to revise the associated reporting category
310 for approved ECs include:

311 • Submission of an appropriate post-approval regulatory submission describing and justifying the
312 proposed revision to the approved ECs. Justification may include information such as validation
313 data and batch analyses.

314 • Submitting a PACMP, in the original marketing application or as part of a post-approval submission,
315 describing a revision to ECs or reporting categories, and how the change will be justified and
316 reported.

317 • Revisions to ECs could also be made utilising an approved post-approval regulatory commitment,
318 as appropriate.

319 **3.3. Roles and responsibilities**

320 The management of all changes to and maintenance of the approved marketing application is the
321 responsibility of the MAH. There is a joint responsibility to share and utilise information between the
322 MAH and any manufacturing organisations to assure the marketing application is maintained, reflects
323 current operations, and that changes are implemented appropriately across relevant sites. Maintenance
324 of the marketing application (including aspects that are not identified as ECs) should follow regional
325 expectations. See [Chapter 6](#) for information related to interactions between an MAH and any
326 manufacturing organisations.

327 For any referenced submission (e.g., Type II Drug Master File, Active Substance Master File, etc.) in a
328 marketing application, the holder of the referenced submission has a responsibility to report changes
329 to their ECs to the MAH referencing their submission, so that the MAH can assess the impact of the
330 change and report any related change to the ECs found in the approved MAA, as necessary and per
331 regional requirements.

332 The approval of ECs and subsequent changes to ECs is the responsibility of the regulatory authorities.

333 **4. Post-approval change management protocol (PACMP)**

334 **4.1. Definition of a PACMP**

335 A PACMP is a regulatory tool that provides predictability and transparency in terms of the requirements
336 and studies needed to implement a change as the approved protocol provides an agreement between
337 the MAH and the regulatory authority. A protocol describes the CMC change an MAH intends to
338 implement during the commercial phase of a product, how the change would be prepared and verified,
339 including assessment of the impact of the proposed change, and the suggested reporting category in
340 line with regional requirements, i.e., a lower reporting category and/or shortened review period as
341 compared to similar change procedure without an approved PACMP. The PACMP also identifies specific
342 conditions and acceptance criteria to be met. A PACMP can address one or more changes for a single
343 product, or may address one or more changes to be applied to multiple products (see [Chapter 4.5](#)).
344 The PACMP may be submitted with the original MAA or subsequently as a stand-alone submission. The
345 PACMP requires approval by the regulatory authority, and the conditions and acceptance criteria
346 outlined in the protocol must be met in order to implement the change(s).

347 A PACMP should describe changes with a level of detail commensurate with the complexity of the
348 change. Once approved, in cases where implementation (see “step 2” below) is pending, there is an
349 assumption that the proposed approach is re-evaluated by the MAH on a regular basis and its validity
350 reconfirmed prior to implementation of the change(s). Specifically, before implementing the change(s),
351 the risk assessment provided in the initial PACMP submission should be reviewed by the MAH to ensure
352 that the outcomes of that risk assessment as they pertain to the planned change(s) are still valid. If
353 the review of the initial risk assessment indicates an increased level of risk associated with execution of
354 the change, the previously approved reporting category should no longer be considered appropriate. In
355 such cases, existing guidance should be followed or a consultation with the relevant regulatory
356 authority should be sought. In addition, the MAH should confirm that the control strategy continues to
357 ensure that the product will be produced consistently following implementation of the change(s).

358 Finally, the use of a PACMP is enabled through an effective PQS that incorporates quality risk
359 management principles (ICH Q9) and an effective change management system (ICH Q10, Appendix 2).
360 The MAH is responsible for ensuring that whenever a CMC change is to be introduced under a PACMP,
361 the facility meets the regulatory requirements of the regulatory jurisdiction where the PACMP was
362 approved with respect to GMP compliance, and inspection or licensing status.

363 **4.2. Application of a PACMP**

364 Step 1: Submission of a written protocol that describes the proposed change(s), its rationale(s), risk
365 management activities, proposed studies and acceptance criteria to assess the impact of the
366 change(s), other conditions to be met (e.g., confirmation that there is no change to the approved
367 specification), the proposed reporting category for the change(s), and any other supportive information
368 (see also below). This protocol is reviewed and approved by the regulatory authority in advance of
369 execution of the protocol.

370 Step 2: The tests and studies outlined in the protocol are performed. If the results/data generated
371 meet the acceptance criteria in the protocol and any other conditions are met, the MAH submits this
372 information to the regulatory authority according to the categorisation (classification) in the approved
373 protocol for review by the regulatory authority as appropriate. Depending on the reporting category,
374 approval by the regulatory authority may or may not be required prior to implementation of the
375 change. If the acceptance criteria and/or other conditions in the protocol (see step 1) are not met, the

376 change cannot be implemented using this approach and should follow existing regulation or guidance
377 instead.

378 Significant changes to the manufacturing process or controls that were not anticipated in the PACMP
379 step 1 (e.g., change of order of unit operations) cannot be implemented as part of step 2 and should
380 be the subject of a regulatory submission as governed by regional regulation or guidance. However,
381 minor unanticipated modifications of the process or controls related to the intended change and not
382 affecting the technical principles of the protocol are normally considered within scope, if appropriately
383 justified.

384 No change outlined in a PACMP should introduce any additional risks to patient safety, product quality
385 or efficacy. A CMC change that would require supportive efficacy, safety (clinical or non-clinical), or
386 human PK/PD data to evaluate the effect of the change (e.g., certain formulation changes, clinical or
387 non-clinical studies to evaluate new impurities, assessment of immunogenicity/antigenicity) is
388 generally not suitable for inclusion in a PACMP.

389 **4.3. Elements of a PACMP**

390 The development of the PACMP is informed by the application of process and product understanding
391 gained from product development and/or manufacturing experience. A PACMP includes some, if not all,
392 of the following elements:

- 393 • A detailed description of the proposed change(s), including a rationale. The differences before and
394 after the proposed change(s) should be clearly highlighted (e.g., in a tabular format).
- 395 • Based on an initial risk assessment, a list of specific tests and studies to be performed to evaluate
396 the potential impact of the proposed change(s), such as: characterisation, batch release, stability
397 (as appropriate, see [Chapter 8.2.1](#)), in-process controls. The PACMP should include an appropriate
398 description of the analytical procedures and proposed acceptance criteria for each test or study.
- 399 • Discussion regarding the suitability of the approved control strategy or any changes needed to the
400 control strategy associated with the planned change(s).
- 401 • Any other conditions to be met, such as confirmation that certain process qualification steps will be
402 completed before implementation.
- 403 • Where applicable, supportive data from previous experience with the same or similar products
404 related to: development, manufacturing, characterisation, batch release, and stability to allow for
405 risk mitigation.
- 406 • Proposed reporting category for the implementation of step 2 of the PACMP.
- 407 • Confirmation that ongoing verification will be performed under the PQS to continue to evaluate and
408 ensure that there is no adverse effect of the change(s) on product quality. In cases where
409 monitoring of the impact on product quality following implementation of the change(s) is required,
410 a summary of the quality risk management activities should be provided to support the proposed
411 PACMP. If multiple changes are to be implemented, these activities should address the potential
412 risk from the cumulative effect of multiple changes and how they are linked.

413 The MAH should demonstrate in the PACMP suitable scientific knowledge and understanding of aspects
414 impacted by the proposed change in order to conduct an appropriate risk assessment of the proposed
415 change(s). Typically, more complex changes would require enhanced product/process understanding.

416 **4.4. Modification to an approved PACMP**

417 A modification to an already approved PACMP such as replacement or revision of a test, study or
418 acceptance criterion should provide the same or greater capability to assess the effect of the proposed
419 change on the product quality. Such changes would normally require a notification type of
420 communication with the regulatory authority. A modification that more significantly alters the content
421 of the protocol may require either prior approval of a protocol amendment or submission of a new
422 protocol, as agreed upon with the regulatory authority.

423 **4.5. Types of PACMPs**

424 There are different types of PACMPs:

- 425 • One or more change(s) to a single product – see above and Annex IIA, for content and
426 implementation. A PACMP can also be designed to be used repeatedly to make a specified type of
427 CMC change over the lifecycle of a product, applying the same principles.

428 If the protocol describes several changes for a particular product, a justification should be added
429 showing how the changes are related and that inclusion in a single protocol is appropriate.

- 430 • Broader protocols – the general principles outlined above apply. The risk of the proposed change(s)
431 should be similar across products; additional considerations should be taken into account
432 depending on the approach, for example:
 - 433 a. One or more changes to be implemented across multiple products (e.g., change in stopper
434 across multiple products that use the same container closure system): the same risk mitigation
435 strategy should be applicable across all impacted products;
 - 436 b. One or more changes to be implemented across multiple products and at multiple sites (e.g.,
437 change in analytical method across multiple sites, change in manufacturing site(s) across
438 multiple products): the same risk mitigation strategy should be applicable across all impacted
439 products and/or sites (see Annex IIB).

440 **5. Product lifecycle management (PLCM)**

441 The PLCM document outlines the specific plan for product lifecycle management that is proposed by the
442 MAH, includes key elements of the control strategy, the ECs, proposed reporting categories for changes
443 to ECs, PACMPs (if used) and any post-approval CMC commitments. This will encourage prospective
444 lifecycle management planning by the MAH and facilitate regulatory assessment and inspection. The
445 PLCM document should be updated throughout the product lifecycle as needed.

446 **5.1. PLCM document: scope**

447 The PLCM document serves as a central repository in the MAA for ECs and reporting categories for
448 making changes to ECs. It includes the key elements described in [Chapter 5.2](#) below and references to
449 the related information located elsewhere in the MAA (see Annex III). Submission of the PLCM
450 document is encouraged; however, the document is expected when the MAH proposes explicit ECs.

451 The elements of the PLCM document are summarised below:

- 452 • **Summary of Product Control Strategy:** A high level summary of the product control strategy
453 should be included in the PLCM document to clarify and highlight which elements of the control
454 strategy should be considered ECs.
- 455 • **ECs** (refer to [Chapter 3](#)): The proposed ECs for the product should be listed in the PLCM document.
456 The identification and justification of ECs are located in the relevant sections of the CTD.
- 457 • **Reporting category for making changes to approved ECs** (refer to [Chapter 3](#)): The proposed
458 reporting categories when making a change to an EC should be listed in the PLCM document. The
459 detailed justification of the reporting categories is located in the relevant sections of the CTD. The
460 reporting category may be based on regional regulations or guidance, or MAH justification.
- 461 • **PACMPs** (refer to [Chapter 4](#)): PACMPs that are submitted to prospectively manage and implement
462 one or more post-approval changes should be listed along with the corresponding ECs to be
463 changed. The approval date of the PACMP should be noted in subsequent submissions. If the
464 PACMP is submitted and approved after approval of the original MAA, an updated PLCM document
465 should accompany the PACMP
- 466 • **Post-approval CMC commitments:** CMC commitments (e.g., specific process monitoring,
467 revisions to ECs) that will be implemented during the commercial phase should be listed in the
468 PLCM document

469 **5.2. Submitting the PLCM document**

470 The initial PLCM document is submitted with the original MAA or with a supplement/variation for
471 marketed products where defining ECs ([Chapter 3.2.3](#)) may facilitate regulatory change management.
472 Following regulatory review and approval of the MAA, the PLCM document will contain ECs and
473 associated reporting categories.

474 **5.3. Maintenance of the PLCM document**

475 An updated PLCM document should be included in post-approval submissions for CMC changes. The
476 updated PLCM document will capture the change in ECs and other associated elements (reporting
477 category, commitments, PACMP). The MAH should follow regional expectations for maintaining a
478 revision history for the PLCM document.

479 **5.4. Format and location of PLCM document**

480 A tabular format is recommended to capture certain elements of PLCM described in [Chapter 5.2](#), but
481 other appropriate formats can be used. See Annex III for an example PLCM table.

482 The PLCM document can be located in either the CTD Module 1, 2, or 3 based on regional
483 recommendations.

484 **6. Pharmaceutical quality system (PQS) and change** 485 **management**

486 **6.1. General considerations**

487 An effective PQS as established in ICH Q10 and in compliance with regional GMPs is the responsibility
488 of a firm (manufacturing sites and MAH where relevant) and it is not the intent of this guideline to

489 require a specific inspection assessing the state of the PQS before the firm can use the principles in
490 this guideline. The conduct of routine inspections in connection with submitted marketing applications
491 and surveillance will nevertheless continue as foreseen by regional regulatory requirements.

492 In the event that the PQS is found not to be compliant, it may result in restrictions on the ability to
493 utilise flexibility in this guideline.

494 Consistent with the basic requirements of ICH Q10, an effective change management system is
495 necessary for implementation of this guideline and is summarised in [Appendix 2](#).

496 **6.2. Management of manufacturing changes in the supply chain**

497 In many cases, a firm has to manage communication of information and interactions of PQSs across
498 multiple entities (internal and external). Therefore, the implementation of robust change management
499 across multiple sites (outsourced or not) is necessary. In conjunction with change control principles in
500 [Appendix 2](#), the following change management activities should be considered to support the
501 approaches defined in this guideline:

- 502 • Changes to ECs should be communicated in a timely fashion between the MAH and the regulators,
503 and between the MAH and the manufacturing chain (and vice versa).
- 504 • The timeliness of communication is driven by the impact of any change related to ECs and should
505 be targeted to those entities in the chain that need to be aware of or to implement the change over
506 the lifecycle of the product.
- 507 • Process knowledge and continual improvement are drivers for change. For example, a Contract
508 Manufacturing Organisation (CMO) may be in a position to propose process improvements which
509 significantly improve control and product consistency. These data can be utilised to revise the ECs
510 and associated PLCM document. The organisation responsible for batch release should be aware of
511 all relevant changes and where applicable, be involved in the decision making.
- 512 • The communication mechanisms regarding MAA changes and GMP issues should be defined in
513 relevant documentation, including contracts with CMOs.

514 **7. Relationship between regulatory assessment and** 515 **inspection**

516 Regulatory assessment and inspection are complementary activities and their fundamental roles
517 remain unchanged by this guideline. Facility-related information obtained on inspection should be
518 available to assessors and the most recent PLCM document, when applicable, should be available to
519 inspectors.

520 Communication between assessors and inspectors can facilitate regulatory review of a specific product
521 submission. When required, information relating to GMP and marketing authorisation compliance may
522 be communicated from inspectors to assessors, and vice-versa, via established mechanisms. The
523 communications can also occur between regulators across regions in accordance with appropriate
524 bilateral/multilateral arrangements.

525 **8. Post-approval changes for marketed products**

526 Marketed products can benefit from the application of ECs and PACMPs as described in this guideline.
527 Specifically, ECs and reporting categories can be proposed for a marketed product via a post-approval
528 regulatory submission; a PACMP can also be proposed for planned change(s) to a marketed product. In
529 addition, such products would also benefit from additional approaches to facilitate changes. This
530 chapter describes a strategy for a structured approach for frequent CMC changes (e.g., analytical
531 methods) and data requirements for CMC changes (e.g., stability).

532 **8.1. Structured approach to analytical procedure changes**

533 Marketed products have existing analytical procedures that may benefit from advances made in
534 analytical sciences. The intent of this chapter is to incentivize structured implementation of equivalent
535 analytical procedures that are fit for purpose. An approach wherein specific criteria are defined for
536 changes to analytical procedures used to test marketed products is described below. If this approach is
537 followed and all criteria are met, the analytical procedure change can be made with immediate or other
538 post-implementation notification, as appropriate, to the relevant regulatory authorities.

539 The following situations are out of scope of this chapter:

- 540 • Procedures where the specification does not adequately reflect the complex information provided
541 by the method. In particular, procedures for which only a subset of the peaks are identified and
542 specified (e.g., assay for identity by peptide map, assay for complex drug substances), or where
543 the specification acceptance criteria include a general comparison to a reference standard beyond
544 specified peaks (e.g., “comparable to reference standard” such as for naturally derived products,
545 biotechnology products made in living systems).
- 546 • Change(s) to a test method based on a biological/immunological/immunochemical principle or a
547 method using a biological reagent (e.g., bioassay, binding assay, ELISA, testing for viral
548 adventitious agents).
- 549 • Changes to predictive models used with multivariate methods.

550 It is important to note that with the exception of the above exclusion criteria, all other methods are in
551 scope including those used for biotechnological/biological products.

552 Making use of Chapter 8.1 is dependent on the regional implementation of ICH guidelines (e.g., ICH
553 Q2, Q9 and Q10) and routine application of these guidelines by industry. The flexibility provided in
554 Chapter 8.1 may not be available in all regions and in all situations; some specific changes may require
555 prior approval as defined in regional guidance.

556 **8.1.1. Principles**

557 In order for this approach to be used, the following should be met:

- 558 • The high-level description of the original method and the revised method should be the same (e.g.,
559 chromatography with spectroscopic detection)
- 560 • Validation results should demonstrate that the revised method is equivalent to or better than the
561 original method
- 562 • Test results obtained using the original method and revised method should be equivalent to each
563 other. This should be assessed in two ways: First, the revised method should give an equivalent

564 outcome, i.e., the same quality decision will be made regardless of whether the data was obtained
565 by the original or the revised method. Second, the validation protocol should contain explicit
566 criteria that compare results obtained using the new and revised method. See step 2 below for
567 further details.

568 • System suitability requirements should be established for the revised method. System suitability
569 ensures the day-to-day performance of the method during routine use.

570 • Specification changes (e.g., total impurities, potency) cannot be introduced using this mechanism
571 unless allowed by existing regional regulations.

572 • This approach may not be used if toxicological or clinical data are required as a result of the
573 method change.

574 If these criteria are met, the methods are equivalent and changes can be made with immediate or
575 other post-implementation notification, as appropriate, to regulatory authorities.

576 **8.1.2. Structured approach**

577 • Step 1: Evaluate the high-level method description. Examples include:

578 – Gravimetric analysis

579 – Volumetric analysis

580 – Atomic absorption

581 – Microscopy

582 – Thermal analysis

583 – Electrochemical analysis

584 – Column chromatography (e.g., HPLC, UPLC)

585 – Plate chromatography (e.g., TLC); if used as an ID test or limit test a change to another type
586 of method description may be made if the criteria in this chapter are met

587 – Electrophoresis

588 – Changes to spectroscopic procedures should remain within same specific technology, e.g., UV
589 to UV, NMR to NMR

590 When two techniques are used together (e.g., HPLC with UV detection), both would be part of the
591 method description (i.e., column chromatography with spectroscopic detection).

592 • Step 2: A prospective analytical validation protocol should be prepared and approved internally by
593 the firm. It should be based on a comparison of the current and proposed method and knowledge
594 of the original validation protocol. The validation should assure that the revised method will be fit
595 for its intended purpose and should contain at least the following:

596 – The principles of ICH Q2 should be followed to validate the change. All validation
597 characteristics relevant to the type of method being validated should be executed as described
598 in ICH Q2.

599 – The validation protocol should include, at minimum, the tests used to validate the existing
600 method and all other relevant tests in ICH Q2. For example, if specificity, linearity, precision

- 601 and accuracy were assessed during validation of the original method, then specificity, linearity,
602 precision and accuracy should also be included in the validation of the revised method. The
603 protocol acceptance criteria should reflect appropriate expectations for method performance
604 and be justified scientifically. They should also be developed in the context of the validation
605 acceptance criteria for the original method to assure that the revised method is fit for purpose.
- 606 – The validation should assess equivalency of the results of the revised method to those of the
607 original method using parallel testing of an adequate number of samples of appropriate
608 concentration based on the intended use of the method. The assessment of equivalency should
609 include the requirement that the new method does not lose any meaningful information
610 provided by the old method. Also the same quality decision should result when assessing data
611 from the same samples tested using the original and revised methods.
 - 612 – If there is a switch from manual to automated methods, the validation should also assess the
613 impact of any related changes in critical reagents, reference standards or software.
 - 614 – The protocol should also contain the detailed operating conditions of both the original method
615 and the revised method to assure the changes being made are clear. The description of the
616 method may be included by attachment.
- 617 • Step 3: Consider the system suitability criteria that exist in the current method, if any, and
618 determine, based on method development data and any additional knowledge gained from
619 commercial production, the system suitability criteria aspects that should be part of the new
620 method. System suitability in this context includes all criteria used to evaluate the day-to-day
621 performance of the method when used for routine testing.
 - 622 • Step 4: Execute the validation protocol and compare the results to the predetermined acceptance
623 criteria. If any criterion is not met, an assessment should be performed to evaluate the impact of
624 the failure to meet the criterion on the validity of the method. If all criteria are met, the method is
625 considered acceptable for its intended use.
 - 626 • Step 5: Consider new product information, if any, identified as a result of a change in the context
627 of the current regulatory filing. If new or revised specifications (e.g., total impurities, potency) are
628 required based on results obtained during method validation, this structured approach may not be
629 used unless allowed by existing regional regulations. In addition, this approach may not be used if
630 toxicological or clinical data are required as a result of the method change. Thus, the method
631 change should have no impact on safety, efficacy, purity, strength, identity, or potency of the
632 product.
 - 633 • Step 6: Prepare a written summary report documenting the outcome of the validation versus the
634 protocol criteria.
 - 635 • Step 7: Follow the internal change process as defined within the firm's PQS to implement the
636 change.
 - 637 • Step 8: Unless new information is identified as a result of this process (see step 5), provide a post-
638 implementation notification of the method change to the regulatory authority after the change is
639 implemented as per regional reporting requirements. This may include the updated method
640 description, the protocol, and the summary report of the validation.
 - 641 • Step 9: Complete post-change monitoring. The firm's change control system (refer to Appendix 2)
642 should explicitly identify and document a mechanism to assure the change was effective with no

643 unintended consequences. The outcome of the assessment should be documented with a
644 conclusion indicating the acceptability of the change.

- 645 • Step 10: All information related to the method change should be available for verification during
646 routine regulatory inspection

647 **8.2. Data requirements to support CMC changes**

648 The data needed for submission to the regulatory authority in support of a post-approval change is
649 established by regional regulations and guidance. This guideline provides science- and risk-based
650 approaches that can be used to develop strategies for confirmatory stability studies supporting post-
651 approval changes to enable more timely filing, approval, and implementation of the changes. Such
652 approaches could be proposed in a PACMP (see Annex IIB).

653 **8.2.1. Stability data approaches to support the evaluation of CMC change**

654 Unlike the formal stability studies recommended in ICH Q1A(R2), whose objective is to establish a
655 useful shelf-life and storage conditions for a new, never-marketed drug substance/drug product, the
656 purpose of stability studies, if needed, to support a post-approval CMC change is to confirm the
657 previously approved shelf-life and storage conditions. The scope and design of such stability studies
658 are informed by the knowledge and experience acquired for the drug product and drug substance.
659 Approaches to the design of such studies should be appropriately justified and may include:

- 660 • Identifying the stability-related quality attributes and shelf-life limiting attributes
- 661 • Stability risk assessments to determine what factors can affect stability relative to the proposed
662 CMC changes
- 663 • Use of appropriate tools to evaluate the impact of the proposed change. These may include:
 - 664 – Drug substance and/or drug product accelerated and/or stress studies on representative
665 material (which may be pilot or laboratory scale rather than full scale)
 - 666 – Pre-and post-change comparability studies on representative material
 - 667 – Statistical evaluation of informal and formal stability studies or other relevant data
 - 668 – Predictive degradation and other empirical or first-principles kinetic modelling
 - 669 – Application of relevant institutional knowledge and knowledge from the scientific literature
 - 670 – Use of confirmatory studies post-change instead of submission of data as part of a regulatory
671 change submission

672 Where applicable, a commitment to initiate or complete ongoing, long-term stability testing on post-
673 change batches can assure that the approved shelf life and storage conditions continue to be applicable
674 after implementing the CMC change.

675 **9. Glossary**

Term	Definition
CAPA	Corrective Action and Preventive Action – System that focuses on investigating,

Term	Definition
	understanding, and correcting discrepancies while attempting to prevent their occurrence
CMO(s)	Contract Manufacturing Organisation(s)
CPP	Critical Process Parameter – process parameter whose variability has an impact on a critical quality attribute and therefore should be monitored or controlled to assure the process produces the desired product quality. (Q8R2)
CQA	Critical Quality Attribute – a physical, chemical, biological or microbiological property or characteristic that should be within an appropriate limit, range, or distribution to assure the desired product quality. (Q8R2)
CTD	Common Technical Document
ECs	Established Conditions
Firm	Manufacturing sites and MAH where relevant
KPP	Key Process Parameter - parameters of the manufacturing process that may not be directly linked to critical product quality attributes, but need to be tightly controlled to assure process consistency as it relates to product quality
MAA	Marketing Authorisation Application
MAH	Marketing Authorisation Holder
Notification	The submission of a change in ECs that does not require approval prior to implementation.
PACMP	Post-Approval Change Management Protocol
PLCM	Product Lifecycle Management
Post-approval CMC commitments	Commitment by the MAH to undertake specific CMC activities to be implemented during the commercial phase.
Prior-approval	Change to an approved established condition that requires regulatory review and approval prior to implementation
PQR	Periodic Quality Review – regular periodic review of API or drug products with the

Term	Definition
	objective to verify process consistency, to highlight any trends and to identify product and process improvements
PQS	Pharmaceutical Quality System
QRM	Quality Risk Management

676 **10. References**

677 ICH M4: *The CTD – Quality*

678 ICH Q1A(R2) *Stability Testing of New Drug Substances and Products*

679 ICH Q2(R1) *Validation of Analytical Procedures: Text and Methodology*

680 ICH Q5E *Comparability of Biotechnological/Biological Products Subject to Changes in Their*
681 *Manufacturing Process*

682 ICH Q8(R2) *Pharmaceutical Development*

683 ICH Q9 *Quality Risk Management*

684 ICH Q10 *Pharmaceutical Quality System*

685 ICH Q11 *Development and Manufacture of Drug Substances*

686 ICH Q8, Q9, and Q10 *Questions and Answers*

687 ICH Q8, Q9, & Q10 *Questions and Answers -- Appendix: Q&As from Training Sessions (Q8, Q9, & Q10*
688 *Points to Consider)*

689

690 **Appendix 1: CTD sections that contain ECs**

691 Notes:

- 692 • This table does not contain a complete list of ECs for a product. The intention of the table is to provide general guidance about the elements of
693 manufacture and control that constitute ECs and their location within the CTD structure.
- 694 • White rows indicate CTD sections where ECs are generally located. Grey rows indicate CTD sections where supportive information is generally located.
- 695 • CTD sections containing ECs may contain elements of supportive information.
- 696 • B = applicable to biotechnological/biological products
- 697 • For delivery system information, the location or the relevant content within the CTD structure may vary depending on the design of the particular product
698 and region

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.S	DRUG SUBSTANCE	
3.2.S.1	General Information	
3.2.S.1.1	Nomenclature	Drug Substance Name, Structure.
3.2.S.1.2	Structure	
3.2.S.1.3	General properties	Supportive information
3.2.S.2	Manufacture	

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.S.2.1	Manufacturer(s)	Drug Substance Manufacturing Site(s) (including testing)
3.2.S.2.2	Description of manufacturing process and process controls	<p>Individual unit operations and their sequence in the manufacturing process</p> <p>For levels/details of ECs for inputs (process parameters and material attributes) and outputs of individual unit operations, reference is made to Chapter 3.2.3.1 – <i>Identification of ECs for the Manufacturing Processes</i></p>
3.2.S.2.3	Control of Materials	<p>Starting material specifications (test, elements of analytical procedure and acceptance criteria)</p> <p>Raw material/reagent/solvent critical controls</p> <p>Source of materials (e.g., cell and seed source, raw materials) and control of critical materials of biological origin</p> <p>Generation and control of Master - Working Cell Bank / Master, - Working Seed Lot, etc. (B)</p>
3.2.S.2.4	Control of critical steps and intermediates	Specifications (e.g., test, elements of analytical procedure and acceptance criteria) for critical steps and intermediates including storage conditions of critical intermediates
3.2.S.2.5	Process validation and/or evaluation	Supportive information

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.S.2.6	Manufacturing process development	Supportive information
3.2.S.3	Characterisation	Supportive information
3.2.S.3.1	Elucidation of structure and other characteristics	Supportive information
3.2.S.3.2	Impurities	
3.2.S.4	Control of Drug Substance	
3.2.S.4.1	Specification	Drug Substance Specification For each Quality Attribute on the specification <ul style="list-style-type: none"> • Test Method • Acceptance Criteria
3.2.S.4.2	Analytical Procedures	Reference is made to Chapter 3.2.3.2 . – <i>Identification of ECs for Analytical Procedures</i>
3.2.S.4.3	Validation of analytical procedure	Supportive information
3.2.S.4.4	Batch analyses	Supportive information
3.2.S.4.5	Justification of specification	Supportive information

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.S.5	Reference Material	Reference Material qualification (e.g., test, elements of analytical procedure, where appropriate, and acceptance criteria)
3.2.S.6	Container Closure	Material of construction and specification
3.2.S.7	Stability	
3.2.S.7.1	Stability Summary and Conclusions	Drug Substance storage conditions and shelf-life (or Retest period for chemicals)
3.2.S.7.2	Post-approval stability protocol and stability commitments	Supportive information (also see Chapter 3.2.2.)
3.2.S.7.3	Stability data	Supportive information
3.2.P	DRUG PRODUCT	
3.2.P.1	Description and Composition of Drug Product	Drug Product qualitative and quantitative composition
3.2.P.2	Pharmaceutical development	
3.2.P.2.1	Components of the drug product	

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.P.2.2	Drug product	Supportive information
3.2.P.2.3	Manufacturing process development	
3.2.P.2.4	Container closure system	
3.2.P.2.5	Microbiological attributes	
3.3.P.2.6	Compatibility	
3.2.P.3	Manufacture	
3.2.P.3.1	Manufacturer(s)	Drug Product Manufacturing (including: testing, primary packaging, device assembly for drug product-device combination products) sites
3.2.P.3.2	Batch Formula	Drug Product Batch Formula (Qualitative and Quantitative)
3.2.P.3.3	Description of manufacturing process and process controls	Individual unit operations and their sequence in the manufacturing process For levels/details of ECs for inputs (process parameters and material attributes) and outputs of individual unit operations, reference is made to Chapter 3.2.3.1 – Identification of ECs for the Manufacturing Processes
3.2.P.3.4	Controls of Critical Steps and Intermediates	Specifications (e.g., test, elements of analytical procedure and acceptance criteria) for critical steps and intermediates including storage conditions of critical intermediates
3.2.P.3.5	Process validation and/or	

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
	evaluation	Supportive information
3.2.P.4	Control of Excipients	
3.2.P.4.1	Specifications	<p>Excipient Specification</p> <p>For each Quality Attribute on the specification</p> <ul style="list-style-type: none"> • Test Method • Acceptance Criteria <p>Or, if applicable,</p> <p>Reference to pharmacopoeial monograph</p>
3.2.P.4.2	Analytical Procedures	Reference to pharmacopoeial monograph and if none exists, refer to Chapter 3.2.3.2 – <i>Identification of ECs for Analytical Procedures</i>
3.3.P.4.3	Validation of analytical procedures	Supportive information

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.3.P.4.4	Justification of specifications	Supportive information
3.2.P.4.5	Excipients of Human or Animal Origin	Excipient source and controls should be specified (for human- or animal-derived excipients only)
3.2.P.4.6	Novel excipients	(If Novel excipient specification is not described in 3.2.P.4.1) Novel Excipient Specification For each Quality Attribute on the specification <ul style="list-style-type: none"> • Test Method • Acceptance Criteria
3.2.P.5	Control of Drug Product	
3.2.P.5.1	Specification(s)	Drug product specification For each Quality Attribute on the specification <ul style="list-style-type: none"> • Test Method • Acceptance Criteria

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.P.5.2	Analytical Procedures	Reference is made to Chapter 3.2.3.2 – <i>Identification of Established Conditions for Analytical Procedures</i>
3.2.P.5.3	Validation of analytical procedures	Supportive information
3.3.P.5.4	Batch analyses	Supportive information
3.2.P.5.5	Characterisation of impurities	
3.2.P.5.6	Justification of specification(s)	
3.2.P.6	Reference Materials	Reference material qualification (e.g., test, elements of analytical procedure, where appropriate, and acceptance criteria)
3.2.P.7	Container Closure System	Supplier/manufacturer of container closure Material of construction and specification

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
3.2.P.8	Stability	
3.2.P.8.1	Stability Summary and Conclusion	Drug product storage conditions and shelf-life (or retest period for chemicals) Where applicable, in-use storage conditions and shelf-life
3.2.P.8.2	Post-approval stability protocol and stability commitment	Supportive information (also see Chapter 3.2.2.)
3.3 P.8.3	Stability data	Supportive information
3.2.A	APPENDICES	
3.2.A.1	Facilities and equipment	Regional regulation and guidance apply
3.2.A.2	Adventitious agents safety evaluation	Supportive information
3.2.A.3	Excipients	Supportive information
3.2.R	REGIONAL INFORMATION	

CTD SECTION	SECTION TITLE	ESTABLISHED CONDITIONS – General List with notes
	Not Applicable	Regional regulation and guidance apply. For EU, Medical Device information or CE mark confirmation

699 **Appendix 2: principles of change management**

700 Consistent with the basic requirements of ICH Q10, an effective change management system supports
701 the principles of this guideline and is described below:

- 702 1. Captures stimuli for change including those that can improve product performance or process
703 robustness;
- 704 2. Ensures full understanding of the scope of the change and its implications for all aspects of the
705 process and control strategy including the impact on ECs and aspects that are not ECs in affected
706 marketing authorisations;
- 707 3. Leverages existing process performance and product quality knowledge;
- 708 4. Requires a science and data based risk assessment and risk-categorisation of the proposed change
709 including the management of risk in the event the proposed change is not implemented;
- 710 5. Determines data (existing and/or to be newly generated) needed to support the change and
711 accordingly develops study protocols describing the methods, prospective acceptance criteria as
712 well as additional post-implementation process performance and/or product quality monitoring as
713 necessary;
- 714 6. When required, ensures that a regulatory submission is developed (e.g., supplement/variation,
715 PACMP) and submitted;
- 716 7. Uses a defined change control process to approve or reject the change and involve appropriate
717 stakeholders, including but not restricted to Manufacturing, Quality, and Regulatory personnel;
- 718 8. Ensures implementation of the change is based on:
 - 719 a. Review that the change as implemented remains aligned with the relevant protocols, any PLCM
720 document and/or any PACMP;
 - 721 b. Assessment of data generated to demonstrate that the change objective and acceptance
722 criteria were met;
- 723 9. Ensures that risk-mitigating steps are developed in case of deviations from acceptance criteria, or
724 identification of unanticipated risks;
- 725 10. Captures new product/process knowledge gained during implementation of the change;
- 726 11. Verifies, post-implementation, that changes have been effective in achieving the desired outcome
727 with no unintended consequences;
 - 728 a. If deviations associated with post-approval changes are detected, ensures that the issue is
729 managed via the firm's deviation management process and appropriate corrective and/or
730 preventive actions are identified and undertaken via the firm's corrective and preventive action
731 (CAPA) system
 - 732 b. Where applicable, ensures that regulatory filings are updated and an assessment is made as to
733 whether updates to the PLCM document are needed
 - 734 c. Requires a post-implementation lessons-learned exercise to build on the product and process
735 knowledge gained with a view to continual improvement, including improvement of the PQS
 - 736 d. Ensures that the change is included and assessed as part of the Product Quality Review (PQR)

737 12. The change management system should be organised and available for review during
738 audit/inspection.

739 *Management Review*

740 Details of Management Review are extensively described in ICH Q10 including the use of appropriate
741 performance indicators as a means to assess the effectiveness of a PQS. These should be meaningful,
742 simple and data-driven. In addition to the requirements of ICH Q10 in the context of ensuring an
743 effective change management system, the following could be considered in the Management Review:

- 744 • Monitoring the timeliness of the change management system to assure that changes are
745 implemented in a timely manner commensurate with the urgency identified for the change. When
746 implementation is delayed, an assessment and mitigation of any risks associated with the delay
747 should be made;
- 748 • Monitoring the performance of the change management system, such as assessing the frequency
749 of proposed changes that are not approved for implementation upon first submission;
- 750 • Ensuring that post-implementation verification occurs and reviewing the results of that verification
751 as a measure of change management effectiveness (e.g., to identify improvements to the change
752 management system);

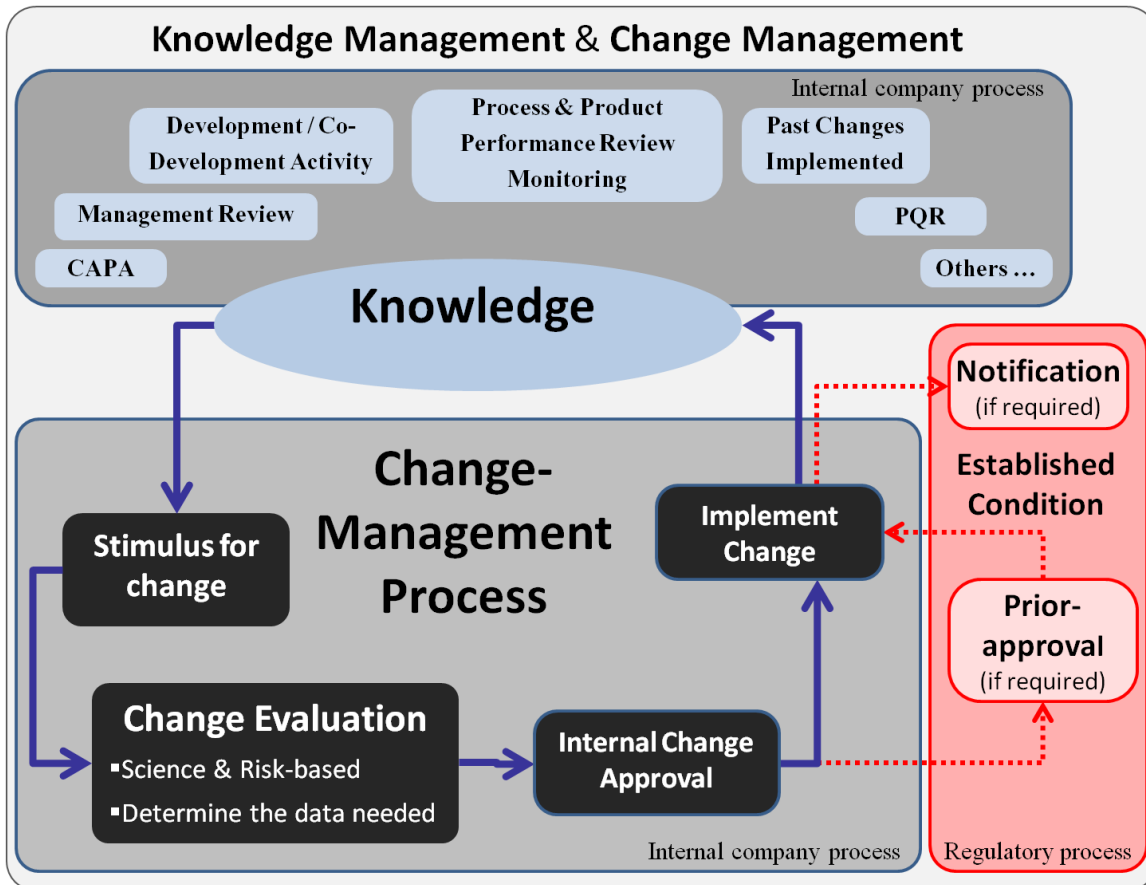
753 *Use of Knowledge in Change Management*

754 An effective change management system includes active knowledge management, in which information
755 from multiple sources is integrated to identify stimuli for changes needed to improve product and/or
756 process robustness. The connection between knowledge management and change management is
757 illustrated in Figure A1.

758 As indicated in ICH Q10 and shown in Figure A1, these sources can include, but are not limited to,
759 developmental studies, process understanding documents, product or process trending, and product-
760 specific CAPA outcomes. They should be comprehensive across the product lifecycle, including all
761 relevant stakeholders (R&D, manufacturing, CMOs, suppliers, etc.). With respect to sharing knowledge
762 between the firm and suppliers, and between the firm and CMOs, considerations for sharing knowledge
763 that relates to product and process robustness or otherwise informs changes should be built into
764 quality agreements and/or contracts.

765 In addition to individual sources of information, there should be a mechanism to provide a holistic view
766 of quality performance for a specific product or product family on a regular basis, as captured in the
767 PQR and shown in Figure A1. This should include steps taken to identify and manage variability
768 introduced from raw materials and the manufacturing process that could impact on product quality
769 during its lifecycle. This allows for the identification of further need for change not apparent when the
770 data are viewed in isolation.

771 Use of knowledge is the responsibility of the firm and should be described in the PQS (for more
772 detailed information reference is made to ICH Q8, Q9, Q10, Q11, Q/IWG Q&A). As described in ICH
773 Q10, there is no added regulatory requirement for a formal knowledge management system.



774

775 **Figure A1** Connection between knowledge management and change management process

776