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AOP DEVELOPERS' HANDBOOK v. 2.8: SUPPLEMENT TO THE GUIDANCE DOCUMENT FOR DEVELOPING AND ASSESSING AOPs

FOREWORD

This document is the AOP Developers' Handbook supplement to the Guidance Document for developing and assessing Adverse Outcome Pathways (AOPs) [ENV/JM/MONO(2013)6, Second Edition]. The Guidance Document provides a historical background for the AOP development programme, and outlines the elements required to construct an AOP as well as the principles of the AOP framework.

The AOP Developers' Handbook (previously "Users' Handbook") supplement was prepared initially in June 2014 by a subgroup of the Extended Advisory Group on Molecular Screening and Toxicogenomics (EAGMST). At that time it was acknowledged that the Handbook should be revised as expert groups and member countries acquire experience in developing, assessing, and applying AOPs. The present version of the AOP Developers' Handbook reflects the most recent principles, practices, and recommendations pertaining to AOP development as implemented and supported via Release 2.8 of the adverse outcome pathway Wiki (AOP-Wiki; aopwiki.org) and overseen by the Emerging Science for Chemical Assessment (ESCA) advisory group.

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117 **AOP DEVELOPERS’ HANDBOOK: SUPPLEMENT TO THE GUIDANCE**
118 **DOCUMENT FOR DEVELOPING AND ASSESSING ADVERSE OUTCOME**
119 **PATHWAYS (AOPs)**

120
121 **ABOUT THIS DOCUMENT**

122
123 This document, the OECD AOP Developers’ Handbook, is a supplement to the Guidance Document
124 for developing and assessing Adverse Outcome Pathways (AOPs) [ENV/JM/MONO(2013)6,
125 Second Edition] (AOP guidance hereafter).

126
127 The AOP Guidance, originally published in 2013 and revised in 2017, provides an introduction to
128 the terminology and concepts of AOP development, including the identification and use of relevant
129 scientific data and resulting knowledge. The Guidance also briefly outlines some potential
130 applications of AOPs.

131
132 While the AOP Guidance document provides a
133 set of definitions and the conceptual background
134 behind AOP development, this AOP Developers’
135 Handbook is designed to provide focused, in-
136 depth, and practical instructions concerning
137 development and review of AOP descriptions in
138 the **AOP knowledgebase (AOP-KB)**, generally
139 accessed via the **AOP-Wiki (aopwiki.org)**. The
140 AOP Developers’ Handbook can be thought of as
141 being analogous to the “instructions for authors”
142 used in preparing a journal article. However,
143 rather than describing the preparation of a
144 technical manuscript, this Handbook (organized
145 into sections) details how to develop, structure,
146 and document an AOP description in the AOP-
147 Wiki. Each section corresponds to “pages” in the
148 AOP-Wiki which are presented as standardized
149 template forms to be filled in during developer
150 AOP construction within the AOP-Wiki
151 environment. The guidance provided in each
152 section of this Handbook includes descriptions of
153 documentation strategies for AOP development
154 i.e., AOP component descriptions, and
155 organisation of that information into each section
156 of the template Wiki AOP pages. This Handbook also provides more explicit guidance on
157 documentation of the information and the factors considered during collection of the evidence
158 relevant to the AOP and evaluating overall weight of evidence (WoE) considerations that inform
159 both the potential fit-for-purpose applications of the AOP and its relevance to different life stages,
160 sex, taxa, susceptible populations etc.

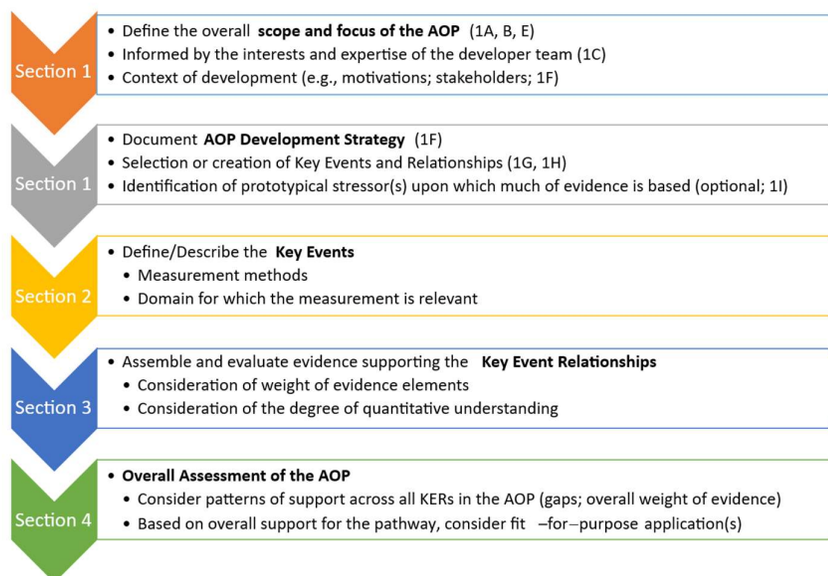
AOP Knowledgebase (AOP-KB) refers to the accumulated machine-readable text and data organized and stored in a MySQL database in accordance with the current AOP Data Model and compiled in the AOP XML.

AOP-Wiki (aopwiki.org) is a web-based interface that provides read/write access to the AOP-KB and serves as the official and primary tool for entering new AOP information in accordance with OECD guidance.

A variety of other tools have read access to the AOP-KB via the XML downloads and can make use of the information contained therein for a variety of purposes. At present, the AOP-Wiki is the only portal for entry of new information into the AOP-KB.

161
162 Although there is no one-size-fits-all approach to AOP development, the sections of the handbook
163 are organized according to a generalized workflow that applies to many AOP development projects
164 (Figure 1). As with the AOP Guidance itself, this Handbook is not intended to provide a review or
165 summary of the literature informing the AOP concept. It focuses on practical aspects of AOP
166 development and assessment and is intended to promote consistency and ensure all AOP developers
167 and contributors understand the approach for AOP development and contribution within the AOP-
168 Wiki. The template and practices outlined herein, to the extent feasible, are intended to support
169 efficient assembly of information pertinent to an AOP and its components (the focus of Handbook
170 Sections 1-3), as well as transparent documentation of information considered during evaluation of
171 evidence confidence and the overall assessment, including WoE, of the AOP (the focus of Section

172 4) along with critical gaps and uncertainties that are relevant to decisions regarding appropriate
173 regulatory applications.
174



175
176

177 **Figure 1.** A generalized workflow for AOP development that has informed the organization of the
178 Developer’s Handbook.

179

180 Developers are encouraged to consult **Annex 1** which outlines a set of guiding questions for
181 evaluating the evidence considered in the overall support for an AOP. Familiarity with these
182 questions before starting an AOP development project can guide the initial scoping including expert
183 solicitation and review of existing literature and/or the design of novel studies toward the data that
184 best inform and support AOPs. Review of the guiding questions and weight of evidence
185 considerations are intended to cue developers on the types of studies that are most influential in
186 providing support for regulatory applications. AOPs are generally best supported by studies that
187 consider multiple key events where comparisons of the concentration, time, or incidence of biological
188 effect in the sample population is not confounded by variations in experimental design. Essentiality
189 of any given key event along the pathway is best evaluated by examining the effects of its prevention
190 or modulation on all downstream events. Searching for or designing studies that best address the
191 guiding questions in **Annex 1** can be expected to lead to both efficient, and high quality AOP
192 development.

193

194 AOP descriptions developed as part of the OECD AOP Development Programme are peer-reviewed
195 according to procedures outlined by the OECD [[Guidance Document for the Scientific Review of](#)
196 [AOPs; ENV/CBC/MONO\(2021\)22](#)]. Because AOP descriptions within the AOP-Wiki are viewed as
197 living documents, they are expected to continue to evolve over time, as new evidence may increase
198 or decrease the overall confidence and certainty in an AOP or its component(s). Consequently, AOPs
199 that are reviewed and endorsed by the OECD will have multiple versions, namely, a static pdf version
200 created at the time of the review or endorsement (termed a “snapshot”), and the current version in
201 the AOP-Wiki, which can continue to change over time. Reviews are performed on these static
202 versions which are permanently stored in the AOP-KB. In this way, users can distinguish content
203 that has been peer-reviewed and endorsed from that which may have been added or modified
204 afterward. The time-stamped, static versions corresponding to the endorsed version of the AOP are
205 also published in the [OECD series on Adverse Outcome Pathways](#) (through 2025) or as an OECD
206 Monograph (2026 and beyond).

207 INTRODUCTION TO ADVERSE OUTCOME PATHWAYS (AOPs)

208

209 An AOP describes a sequence of events commencing with initial interaction(s) of a stressor with a
210 biomolecule within an organism that causes a perturbation in its biology (i.e., molecular initiating
211 event, MIE), which can progress through a dependent series of intermediate key events (KEs) and
212 culminate in an adverse outcome (AO) considered relevant to risk assessment or regulatory
213 decision-making (Table 1). AOPs are composed of a causal sequence of upstream to downstream
214 KEs, representing a cascading series of measurable biological changes that can be expected to
215 occur if the perturbation is sufficiently severe (i.e., in terms of potency, duration, frequency) to
216 drive the pathway all the way to the AO. **Importantly, AOPs do not describe every detail of the
217 biology but instead focus on describing critical steps or check-points along the path to
218 adversity, which are both measurable and have potential predictive value for regulatory
219 application.** While the focus of AOP development is to capture and organise what is known, the
220 process of AOP development may also identify current knowledge gaps which, if filled, could
221 further improve predictive utility.

222

223 **Table 1:** Definitions of key terms and abbreviations used in this Handbook (see AOP guidance for
224 additional terminology relevant to the AOP framework and its application).

225

Molecular initiating event	MIE	A specialised type of key event that represents the initial point of chemical/stressor interaction at the molecular level within the organism that results in a perturbation that starts the AOP.
Key event	KE	A change in biological or physiological state that is both measurable and essential to the progression of a defined biological perturbation leading to a specific adverse outcome.
Key event relationship	KER	A scientifically-based relationship that connects one key event to another, defines a causal and predictive relationship between the upstream and downstream event, and thereby facilitates inference or extrapolation of the state of the downstream key event from the known, measured, or predicted state of the upstream key event.
Adverse Outcome	AO	A specialised type of key event that is generally accepted as being of regulatory significance on the basis of correspondence to an established protection goal or equivalence to an apical endpoint in an accepted regulatory guideline toxicity test.

226

227 KEs are measurable biological changes that are essential to the progression along an AOP.
228 Essentiality indicates that the KEs play a causal role in the pathway, such that if a given KE is
229 prevented or fails to occur, progression to subsequent KEs in the pathway will not occur. While
230 KEs are essential to progression along the AOP, they are not necessarily sufficient. The extent of
231 triggering of the pathway (influenced by intensity and duration of exposure to a stressor)
232 determines whether it will progress all the way to the AO. The conditions under which progression
233 can be expected are described as quantitatively as possible, in the KERs that link an upstream to a
234 downstream KE.

235

236 The suitability of a given AOP for application in different regulatory contexts is influenced by (1)
237 the confidence and precision with which the KEs can be measured, (2) the level of confidence in
238 the relationships between the KEs linked in an AOP (KERs) based on biological plausibility and
239 empirical support for the KERs; and (3) WoE for the overall hypothesised pathway, taking into
240 account additional considerations including any uncertainties and inconsistencies. Therefore,
241 overall assessment of AOPs is best supported by providing thorough descriptions of the KEs
242 [Section 2], relationships between those KEs [i.e., KERs, Section 3], and by final consideration of
243 the overall patterns of support including plausibility and other direct and indirect empirical
244 evidence of causal relationships across the key events defined for the pathway that increase or
245 decrease overall confidence in the AOP [Section 4]. The overall patterns of support, ultimately

246 inform the suitability (i.e., fit-for-purpose) for various types of applications. Consequently, both
247 the Handbook and AOP-Wiki are structured in a manner that include structured pages and prompts
248 for AOP developers to provide relevant types of supporting documentation.

249

250 *Principles of AOP Development and their Implications for AOP Description*

251

252 As a pragmatic convention, AOPs are conceptualised as a single sequence of events proceeding
253 from the MIE to the AO via a series of intermediate KEs (Villeneuve et al. 2014a). That is, they
254 describe how one particular molecular perturbation may cause one AO, not every possible AO that
255 perturbation may cause, nor every perturbation leading to a particular AO. MIEs, KEs, and AOs
256 may be shared by more than one AOP to form an AOP network. Consequently, KEs should be
257 constructed as discrete (modular) units without reference to a specific MIE, AO, or other KEs.
258 Likewise, it is important that KERs describing relationships between discrete pairs of KEs are
259 independent of other elements of the AOP. This facilitates generation of self-contained KE and
260 KER descriptions that can be linked to multiple other AOPs. Such an approach both fosters
261 consistency and increases efficiencies in the AOP development process, by eliminating the need
262 for AOP developers to completely re-describe biological measurements (KEs) or evidence
263 supporting the relationship between two KEs (KERs) that another developer may have already
264 detailed. Maintaining KE and KER descriptions as discrete units that avoid reference to other
265 elements of the AOP also facilitates the updating of KE and KER descriptions as new methods for
266 measuring KEs or new evidence supporting KERs are developed. Finally, it facilitates the
267 construction and conceptualisation of AOP networks.

268

269 An AOP network is defined as an assembly of two or more AOPs that share one or more KEs
270 (Knapen et al. 2018). Because the components of an AOP (KEs and KERs) are described in the
271 AOP-Wiki, in a modular fashion, AOP networks emerge from the description of individual AOPs
272 that share KEs. AOP networks capture broader knowledge concerning the range of possible AOs
273 which a perturbation may cause, or the variety of upstream KEs which can lead to a given AO.
274 AOP networks are also suited to address exposures to multiple stressors that lead to the same AO
275 or individual stressors that activate multiple MIEs (Knapen et al., 2015; Villeneuve et al., 2014a,
276 b; Knapen et al. 2018).

277

278 In describing the KEs and KERs of an AOP, the content of each information field of the KE or
279 KER description should be completed where possible and supported by citation of primary
280 literature and other relevant sources. Nevertheless, AOP descriptions reflect current knowledge
281 and will evolve as additional information becomes available, so AOP descriptions should be
282 regarded as “living documents” that reflect the state of knowledge at the time they were last
283 updated. It is expected that, as “living documents”, AOPs may have gaps that may be addressed
284 over time as the science progresses or as other researchers contribute. This also encourages
285 collaboration and contributions between experts in various areas of research and the regulatory risk
286 assessment community.

287

288 AOPs thus provide a relevant construct to promote collaboration and better coordinate and tailor
289 research to practical application, such as the development of KE-based testing strategies. The
290 AOP-Wiki facilitates this by providing a tool to organise and share the relevant data and
291 information. Consequently, it is recommended that descriptions are structured using presentation
292 of bullets or tables and organised into topical subsections rather than as extensive narrative text.

293

294 In this Handbook, particular emphasis is placed on sections related to the description of the MIE,
295 KEs and AO in an AOP (Section 2), the assembly of available scientific evidence supporting the
296 KERs (Section 3) and the overall support for the AOP as a whole (Section 4) and may additionally
297 consider its potential application (Figure 1).

298

299 AOP descriptions should be supported with well documented and transparent citation of the
300 appropriate peer-reviewed literature and/or other relevant sources. Authors are encouraged to

301 provide references formatted according to the OECD Style Guide
302 (<https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-Edition.pdf>).

303

304 **REVIEW AND ENDORSEMENT of AOP-WIKI CONTENT**

305 AOPs developed and evaluated according to the guidance in the Handbook may be submitted for
306 technical review via the OECD AOP Development Programme, submitted for potential
307 publication in a partner journal¹, or have a review managed by an approved third party
308 organization, provided the reviews are managed as described in the [Guidance Document for the
309 scientific review of Adverse Outcome Pathways](#). AOPs that are accepted after review and
310 revision according to the guidance are then eligible to be added to the OECD AOP Development
311 Workplan and considered for endorsement by the OECD Working Party on Hazard Assessment
312 (WPHA) and/or Working Group of the National Coordinators for the Test Guidelines Program
313 (WNT). See also [AOP Review and Endorsement Info page](#).

314

315

316 **OBTAINING AUTHOR ACCESS TO THE AOP-Wiki**

317

318 **Read-access** to all contents of the AOP-KB is publicly available via the AOP-Wiki (aopwiki.org)
319 without need to create a user profile, login ID, or password.

320

321 **Commentor access:** A self-created user account, with a verified email address, grants the user the
322 ability to comment on all pages in the AOP-Wiki including AOPs, KEs, and KERs. Users can
323 create an account on the AOP-Wiki by clicking the “Register” button on the AOP-Wiki home page.

324

325 **Author Access:** In order to create or edit AOPs, KEs, or KERs, the user must request author access
326 to the AOP-Wiki by following the instructions [here](#).

327

328

329 **A NOTE ON AOP DESCRIPTIONS IN THE AOP-Wiki**

330

331 AOP descriptions in the AOP-Wiki consist of both structured information and free text.

332

333 **Structured information** fields in the AOP-Wiki employ standardised ontologies or controlled
334 vocabularies available through look-up tables or by making selections from a drop-down list.
335 Structured information fields within the AOP-Wiki populate a back-end database and can be
336 exported in a machine-readable format (i.e., XML) that can be used in a variety of computational
337 analyses, and more complex querying, and searching of the AOP-KB. For example, construction
338 of AOP networks from the modular units of individual AOP descriptions relies on these structured
339 annotation fields.

340

341 **Free text** sections in the AOP-Wiki provide AOP developers with much greater descriptive
342 flexibility than structured information fields. While free text is searchable, it is not standardised
343 and machine-readable and is not part of the XML download, thus limiting its use from a
344 computational standpoint.

345

346 **CONTENT LICENSING**

347 By default, all content in the AOP-Wiki is licensed under a Creative Commons, Attribution,
348 Share Alike ([CC BY-SA](#)) license. This license stipulates the following:

- 349 • Users must not **restrict access** to the work using technical measures, or otherwise
350 attempt to impose limitations on the freedoms to use, study, apply, redistribute, or
351 distribute derivative works.
- 352 • Users must **give proper attribution to the author and retain the license notice**.
- 353 • Users must **release derivative works under identical license terms**.

354

355 Any reuse of AOP-Wiki content or derivative of AOP-Wiki content requires appropriate attribution
356 including a link to the license and indication of any changes made. AOPs are, however, represented
357 by pages within the AOP-Wiki that have page-specific accessibility properties. AOP page licensing
358 options (Table 2) are described below.

359

360 Key **Event** and Key Event **Relationship** pages in the AOP-Wiki are shared pages that any
361 author can edit. Consequently, at present, only a BY-SA license can be applied. Authors
362 wishing to protect unpublished content on an Event or Relationship page, are encouraged to
363 develop their content on an external pre-print server, and then cite the appropriate DOI on the
364 relevant Event or Relationship pages in the AOP-Wiki. To facilitate attribution, authors may
365 also want to “tag” content they have added to these shared pages with their name or initials.

366

367 **AOP Pages** have restricted author access in the AOP-Wiki. They can only be edited by authors
368 listed as contributors. Consequently, there is an option to directly protect content of an AOP
369 page, if desired. At the time an AOP page is first created in the AOP-Wiki (**and only at that**
370 **time**), authors have the option to override the default CC BY-SA license and instead select a
371 “©; Copyright, All Rights Reserved” license. A © license indicates that the author retains all
372 rights provided by copyright law, and prohibits others from reproducing, distributing, and/or
373 adapting any part of the work without the copyright holder’s permission. Conceptually, this
374 allows AOP-pages on the AOP-Wiki to function as a pre-print server. While the content under
375 development is visible to other authors and potential users, the content is restricted and
376 protected by law. This option is provided to encourage transparent AOP development on the
377 AOP-Wiki, while protecting the intellectual property of the authors and the effort they have put
378 into developing the AOP.

379

380 To ensure the ultimate accessibility and usability of information in the AOP-Wiki, All Rights
381 Reserved licenses in the AOP-Wiki automatically revert to CC BY-SA after 12 months from
382 the AOP page creation date, unless the authors take action to extend the All Rights Reserved
383 license, prior to its expiration. The All Rights Reserved License can be extended at any time,
384 prior to its expiration by clicking the “Edit” button on the AOP page and then clicking the
385 “Extend current All Rights Reserved License” button from the Editing page. An active All
386 Rights Reserved License can be extended multiple times. However, it is the authors
387 responsibility to monitor the All Rights Reserved expiration date and take action to extend the
388 term before the expiration date. The current expiration date for an All Rights Reserved License
389 can be found on the Editing page, in blue highlighted text positioned directly above the “Extend
390 current All Rights Reserved License” button.

391

392 Once the All Rights Reserved license expires, the AOP page defaults automatically to a CC BY
393 SA license. The authors can also switch to a CC BY-SA license at any time by clicking the
394 “Edit” button on the AOP page, then making a new license selection. Note, **any switch to a CC**
395 **BY-SA license is irreversible**. Once an AOP page defaults or is switched by the authors to a
396 CC BY-SA license, it cannot be changed back to an All Rights Reserved license.

397

398 In addition to the default CC BY-SA license, authors also have the option to select a CC BY-
399 SA License with an “Open for Adoption” tag. This option applies the same license terms as the
400 CC BY-SA license, however, it is used to signal that the original authors are no longer actively

401 developing the AOP and invite new authors to take over development. New authors wishing to
 402 take over development of the AOP can do so by contacting the AOP-Wiki gardening team at
 403 aopwiki@googlegroups.com. Note, an All Rights Reserved License cannot be applied to an
 404 AOP page that has been opened for adoption.

405
 406
 407

Table 2: AOP page License Options Overview¹

License Option	Terms	Implementation Notes
All Rights Reserved	Re-use of the content of the AOP page, in any form, requires advanced, written permission from the authors.	Must be selected at the time the AOP page is first created. Expires after 12 months unless extended by the authors. Once an All Rights Reserved license expires or a different license type is selected, it is not possible to revert back to an All Rights Reserved license.
BY-SA	This license allows users to distribute, remix, adapt and build upon the material in any medium or format so long as attribution is given to the creator(s). The license allows for commercial use. However, if you remix, adapt, or build upon the material all derivative works must be licensed under identical terms.	This is the default license applied at the time of AOP page creation, unless an All Rights Reserved license was selected at that time. Authors can switch from All Rights Reserved (if applicable) to BY-SA at any time. However, it is not possible to revert back to All Rights reserved once a BY-SA selection has been made.
BY-SA Open for Adoption	This license allows users to distribute, remix, adapt and build upon the material in any medium or format so long as attribution is given to the creator(s). The license allows for commercial use. However, if you remix, adapt, or build upon the material all derivative works must be licensed under identical terms.	This option is available on the Editing Page, accessed by clicking the “Edit” button on the AOP page. This selection is used to signal that the original authors are no longer developing the page and invite other developers to take over. An All Rights Reserved license cannot be applied to an AOP page that was opened for adoption.

408
 409
 410

¹ License options described apply only to AOP pages in the AOP-Wiki. Key event and key event relationship pages are BY-SA only.

411
 412

SECTION 1 – AOP DESCRIPTION

413
 414
 415

This section is for information on the AOP to be entered on the upper portion of an AOP page within the AOP-Wiki. Here the overall structure of the AOP is introduced, the motivation and strategy for its development described and the component KEs and KERs are listed.

416
 417

1A. AOP Identifier and Title

418
 419

This subsection provides guidance for naming the AOP.

420
 421
 422

i. AOP Identifier

Each AOP is automatically given a numerical AOP identifier by the AOP-Wiki when it is created (e.g., AOP: ###).

423
 424

ii. (AOP) Title

425
 426

Each AOP should be given a descriptive title that takes the form “MIE leading to AO via distinctive KE”. For example, “Aromatase inhibition [MIE] leading to reproductive dysfunction

427 [AO] via reduced vitellogenin production” or “Thyroxine production inhibition [MIE] leading to
428 decreased cognitive function [AO] via decreased circulating thyroid hormone concentrations”.
429 While each AOP is distinguished in the AOP-KB and AOP-Wiki by their AOP page ID numbers
430 and unique URL, in a growing number of cases where AOPs linking the same MIE to the same
431 AO are being entered into the AOP-Wiki, the “via distinctive KE” descriptor makes it easier to
432 distinguish different AOPs within a network of closely related AOPs.
433

434 In cases where the MIE is unknown or undefined, the earliest known KE in the sequence (i.e.,
435 furthest upstream) should be used in lieu of the MIE and it should be made clear that the stated
436 event is a KE and not the MIE.

437 *iii. Short Name*

438 A short name should also be provided that succinctly summarises the information from the title.
439 This name should not exceed 90 characters.
440

441 **1B. Graphical Representation of the AOP**

442 A graphical summary of the AOP listing all the KEs in sequence, including the MIE (if
443 known) and AO, and the pair-wise relationships (links or KERs) between those KEs should
444 be provided. This is easily achieved using the standard box and arrow AOP diagram (Figure
445 2).
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451 **Figure 2.** Generic AOP diagram, where boxes represent KEs and arrows represent KERs.

452

Development tip 1 – Graphical Representation: The graphical representation (AOP diagram) serves as a useful road-map to guide AOP development in the AOP-Wiki. For this reason, it is recommended that an AOP diagram be developed prior to creating an AOP description in the AOP-Wiki. Starting with the graphical summary provides a useful overview of the KE and KER pages that will need to be included. Ideally, development of a graphical overview of the AOP should be followed by a search of existing content to determine whether analogous AOPs and/or KEs or KERs already exist in the knowledgebase. This prevents duplicated effort and help to ensure that KEs and KERs are shared among AOPs, allowing for de facto creation of AOP networks. Once existing KE and KER pages relevant to the AOP have been identified, the developer then knows which pages in the AOP-KB will need to be edited or created de novo.

470

The graphical summary is prepared and uploaded by the user (and is often included) as part of the proposal when AOP development projects are submitted to the OECD AOP development workplan.

The graphical representation, or AOP diagram, provides a useful and concise overview of the KEs that are included in the AOP, and the sequence in which they are linked together. This can aid both the process of development, as well as review and use of the AOP.

471

Development tip 2 – Number of KEs to include: Determining the number of KEs to include in an AOP and the specificity with which they are defined is one of the more challenging aspects of AOP development. In describing KEs within an AOP, it is important to recognise their distinction from “mechanism of action”. AOPs provide a description of a limited number of essential, measurable events (check-points or nodes of convergence of mechanistic pathways most relevant to informing application) leading to induction of the relevant toxicity endpoint. They do not necessarily provide a comprehensive molecular description of every aspect of the biology involved. With that in mind, the following “rules of thumb” can help guide the process of KE definition (Villeneuve et al. 2014a, b):

Where possible and appropriate for application, try to include at least one KE at each major level of biological organisation (molecular, cellular, tissue, organ, individual).

Where feasible/appropriate, focus on KEs that can be measured in a relatively routine manner over those that require highly specialised expertise, equipment, or supplies to measure. These will tend to be the KEs for which empirical evidence to support KERs is more likely to be available to support the WoE evaluation.

Select a limited number of KEs that are measurable and for which evidence supports plausibility and potential predictive utility. Where relevant, more detailed description of the underlying biology involved can be incorporated into the descriptions of the biological plausibility linking two KEs (see section 3 – KER descriptions).

472

Development tip 3 – Branching of AOPs captured on a single AOP page

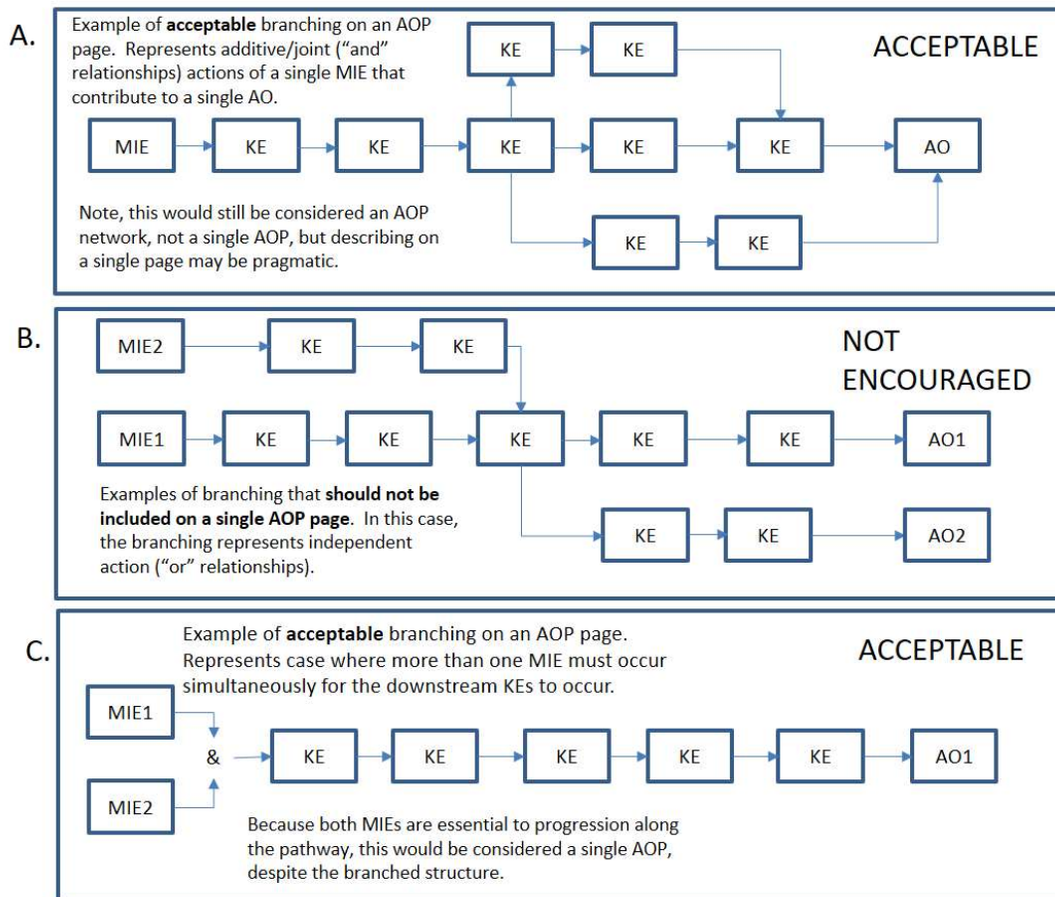
In principle, an individual AOP is defined as a single, non-branching sequence of KEs, linked by KERs that connect a single MIE to an AO (Villeneuve et al. 2014a). Consequently, most AOPs pages should define a single, non-branching, sequence of KEs linked by KERs. However, it is recognized that in some cases there may be exceptions for which representation of a simple AOP network on an AOP page is a more pragmatic unit of development and evaluation (see Leist et al. 2017 for examples and further explanation). In such cases, representation of a branched structure on an AOP page is acceptable, so long as the principles of modularity of the KEs and KERs and overall coherence to the framework is maintained.

For example, representation of branching on an AOP is acceptable when there are multiple KEs, causally linked to the MIE and AO that are occurring concurrently and acting in concert to drive the downstream effects. In such cases, the various KEs cannot be placed neatly into a single temporal sequence because they are effectively occurring simultaneously. Likewise it cannot be determined which of the concurrent KEs is most essential or critical, because there are multiple KEs contributing jointly such that it cannot be effectively determined whether one could cause the pathway to progress without the other. This is contrasted with cases where KEs act independently such that one event or the other, alone, would allow progression toward the outcome.

In cases where an additive (“and”) relationship must be assumed, representation of a simple AOP network on a single AOP page within the AOP-KB may be more practical from both a development and use stand-point than breaking those multiple highly related pathways into separate AOP descriptions. As long as KEs and associated KERs are each represented as separate modular pages in the AOP-KB (as described below), capturing such networks on single AOP pages does not create problems for modular AOP network building. Indeed, it can strengthen the overall AOP by capturing the evidence for pleiotropic effects of the same MIE that ultimately contribute to the same outcome.

Note, such branched AOP structures should only be included on a single AOP page when all the branches diverge from a common MIE (or MIEs in the case that two or more MIEs MUST occur to drive the pathway) and converge to a common AO (Figure 3A) and two or more of the KEs contributing causally to the AO occur concurrently such that it is experimentally intractable to isolate and identify which is playing the dominant causal role and all KEs have predictive value.

Branched structures should not be included on a single AOP page when they diverge to independent adverse outcomes (e.g., Figure 3B) and/or are operating largely independent of one another and can be experimentally resolved from one another in space or time. Following this logic, two or more MIEs may occur on an AOP page, when two or more MIEs MUST occur simultaneously in order for the pathway to be triggered (Figure 3C).



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Figure 3. Illustration of general guidance regarding inclusion of simple AOP networks or branched AOP structures (A) on a single AOP page. Branching representing independent actions leading to more than AO should not be included in an AOP description (B). Branching indicating multiple KEs (including MIEs) that MUST occur for the pathway to progress downstream should be included in an AOP description. In case multiple MIEs are essential, branching of MIEs are acceptable (C).

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482

1C. Authors of the AOP

This section provides guidance on author identification.

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485

i. Authors and Affiliations

List the name and affiliation information of the individual(s)/organisation(s) that created/developed the AOP. In the context of the OECD AOP Development Workplan, this would typically be the individuals and organisation that submitted an AOP development proposal to ESCA and further considered under an OECD working party (e.g., WPHA, WNT). Significant contributors to the AOP should also be listed. A corresponding author with contact information may be provided here. This author does not need an account on the AOP-Wiki and can be distinct from the point of contact below. The list of authors will be included in any snapshot made from an AOP.

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ii. Point of Contact

Indicate the point of contact for the AOP-Wiki entry itself. This person is responsible for managing the AOP entry in the AOP-Wiki and controls write access to the page by defining the contributors as described below. Clicking on the name will allow any wiki user to correspond with the point of contact via the email address associated with their user profile in the AOP-

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501 Wiki. This person can be the same or vary from the corresponding author listed in the authors
502 section. In cases where the individuals are different, the corresponding author would be the
503 appropriate person to contact for scientific issues whereas the point of contact would be the
504 appropriate person to contact about technical issues with the AOP-Wiki entry itself.

505

506 Corresponding authors and the point of contact are encouraged to monitor comments on their
507 AOPs and develop or coordinate responses as appropriate. Selecting the “Watch” ()
508)option on the AOP page will allow an e-mail alert to be sent whenever changes to the AOP
509 page or linked KE or KER pages are made.

510

511 *iii. AOP-Wiki Contributors*

512 List user names of all authors contributing to or revising pages in the AOP-Wiki that are linked
513 to the AOP description. Identification of contributors in this section controls write access to the
514 AOP page. Only contributors listed here, with author rights in the AOP-Wiki, can edit the AOP
515 page.

516

517 *iv. Coach(es)*

518 This field is used to identify coaches who supported the development of the AOP.

519 Coaches are experienced AOP developers that are familiar with the guidance document, AOP
520 development principles, and navigation within the AOP-Wiki. They assist AOP developers by
521 answering questions about the framework, the organization of information in the AOP-Wiki and
522 facilitate compliance with the guidance document and best practices. Upon acceptance of the
523 AOP development project under the OECD workplan, a coach will be assigned. AOP
524 developers without an OECD workplan – related project can request a coach from the SAAOP
525 (Society for the Advancement of AOPs) via aopwiki@googlegroups.com.

526 Identification of coaches in this section provides acknowledgement of the volunteer
527 contributions made by the coach(es) and professional recognition.

528

529 **1D. Handbook Versioning and OECD Status**

530

531 *i. Handbook Version*

532 ● As the AOP framework evolves and information fields, features, or functions are added
533 or modified in the AOP-Wiki, the AOP Developers’ Handbook (this document) is
534 updated to reflect the current state of the AOP-Wiki. In many cases, the AOP-Wiki and
535 Handbook may undergo several updates over the duration of an AOP development
536 project. Newly added AOPs are required to comply with the version of the Handbook
537 that was current on the date the AOP was created, or newer. Where feasible, authors
538 are encouraged to update their AOPs for consistency with the current Handbook
539 version. However, this is not always possible or practical. Consequently, the
540 “Handbook Version” column of the “Status” table is used to indicate the version of the
541 Handbook that the authors used to guide their development.

542 ● When a developer creates an AOP, the current version of the Handbook, on the date of
543 creation, will be automatically populated into the “Handbook Version” column of the
544 “Status” table, along with a link to that version of the Handbook. This information will
545 also display in the “Title” section of the AOP page, right under the “Short name”. As
546 newer versions are released, the authors have the option to switch to a newer Handbook
547 version by selecting from a drop down menu on the Edit page. However, they cannot
548 select versions that pre-date the creation date of their AOP. Both archived handbook
549 versions and release notes summarizing the major changes can be found on the
550 Developers’ Handbooks archive page (<https://aopwiki.org/handbooks>).

551

552 *ii. OECD Status*

553 For AOPs that are included in a project that has been accepted into the OECD AOP Development
554 Workplan (see [http://www.oecd.org/chemicalsafety/testing/projects-adverse-outcome-
555 pathways.htm](http://www.oecd.org/chemicalsafety/testing/projects-adverse-outcome-pathways.htm)), the status with regard to progress through OECD review and endorsement

556 processes is indicated. ‘OECD status’ tracks the level of review/endorsement of the AOP . This
557 designation is managed and updated by the OECD. It cannot be changed by the AOP author(s).
558 AOPs in the AOP-Wiki can be filtered by their OECD status either using the table heading filters
559 on the AOP listing page, or by clicking the “With OECD status” button on the listing page,
560 which toggles to the OECD view and contains only those AOP development projects that are
561 part of the OECD workplan. The OECD status designations for filtering purposes are the
562 following:

- 563 ● WPHA/WNT Endorsed
- 564 ● ESCA Approved
- 565 ● Under Review
- 566 ● Under Development

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568

569 **iii. OECD Project Number**

570 The OECD project number is assigned upon acceptance into the OECD AOP development
571 workplan and indicated along with the current OECD status of the AOP. This designation is
572 managed and updated by the OECD. It cannot be changed by the AOP author(s). OECD project
573 numbers are listed in the all AOPs listing table (blank for AOP development projects not on the
574 OECD workplan), and are displayed on the “OECD View” page, which is accessed by clicking
575 the “With OECD status” button on the AOP listing page.

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578 **iv. Date Modified**

579 The date the AOP was last modified is automatically tracked by the AOP-Wiki. The date
580 modified field can be used to evaluate how actively the page is under development and how
581 recently the version within the AOP-Wiki has been updated compared to any snapshots that
582 were generated.

583
584

585 **1E. ABSTRACT**

586 In the abstract section, authors should provide a concise and informative summation of the AOP
587 under development. Abstracts should typically be 200-400 words in length (similar to an abstract
588 for a journal article). Suggested content for the abstract includes the following: (1) the
589 background/purpose for initiation of the AOP’s development (if there was a specific intent); (2) a
590 brief description of the MIE, AO, and/or major KEs that define the pathway; (3) a short summation
591 of the overall WoE supporting the AOP and identification of major knowledge gaps (if any); (4) a
592 brief statement about how the AOP may be applied (optional). The aim is an "executive summary"
593 to capture the highlights of the AOP and its potential scientific and regulatory relevance.

594

595 **1F. AOP Development Strategy**

596 This subsection describes key elements of “Why” (Context) and “How” (Strategy) the AOP was
597 developed. The content informs other developers, reviewers and users about the strategy and focus
598 for identification and assimilation of the relevant evidence base for KEs and KERs in the AOP.

599

600 Context:

601 This subsection describes key elements of why the AOP was developed and for whom (e.g.,
602 funding sources; stakeholders; etc.).

603 Below are examples of the types of information to include:

- 604 ● Key research question(s) or regulatory needs being addressed
- 605 ● Scope and basis for the evidence gathering/literature search scope
 - 606 ○ e.g., focused on a specific taxonomic group?
 - 607 ○ adding new branches to an existing AOP?
 - 608 ○ development of an additional KE/KER?
- 609 ● Acknowledgement of the source of funding (if applicable)
- 610 ● The overall objective/envisaged use of the AOP that informed its development, e.g., to

- 611 ○ document biology based on specialized expertise,
- 612 ○ establish the relevance and utility of an assay,
- 613 ○ develop an organizing construct in stressor specific (quantitative) hazard
- 614 characterization,
- 615 ○ contribute to development of an integrated approach to testing and assessment, etc.
- 616 ○ indication of interesting biology encompassed by the AOP that is not necessarily
- 617 evident from the KE and KER descriptions;
- 618 ○ as part of a network-guided approach to AOP development, noting other AOP(s)
- 619 developed as part of the effort
- 620 ● Other information that may be useful to the AOP developer and/or user that facilitates
- 621 understanding of motivation/objective/scope for AOP development.

622

623 Strategy

624 This subsection describes *how* the AOP was developed to address the context indicated in the

625 background and acknowledgements above. Specifically, what was the strategy, focus and

626 workflow for identification and assembly of relevant evidence to meet the objective/envisaged

627 application? This information is critical to facilitate the reuse of components and expansion of

628 AOPs. Transparency of the rationale for identification and selection of supporting data also

629 contributes to confidence for regulatory application of AOPs and/or their components.

630

631 Developers should tailor the contents of this section to their particular AOP context and approach,

632 depending e.g., on the scope, nature of prior documentation of the pathway, the starting point for

633 development (e.g., the molecular initiating event or adverse outcome), complexity, and/or

634 envisaged application(s). For example, it may build on previously well-documented and accepted

635 pathways, with focus on particular aspects of uncertainty or particular components of the pathway.

636

637 Content may include:

- 638 ● **Level of resolution / detail in terms of the KEs and KERs represented** in the pathway.
- 639 The goal is to identify notable milestones or checkpoints in the progression of and adverse
- 640 biological response that are both measurable and have predictive utility relevant to
- 641 regulatory application, rather than detailed elements of biology. It is important, then, to
- 642 specify the basis for selection of which KEs and KERs are explicitly, versus implicitly,
- 643 represented in the AOP.
- 644
- 645 ● **Overall data search and identification strategy/ies**, including general strategies (i.e.,
- 646 workflow) for information search, retrieval, and screening (and possibly assessment).
- 647 Example content includes:
 - 648 - reliance on prior knowledge and/or documentation of the pathway, e.g.,
 - 649 ○ expert knowledge
 - 650 ○ previously conducted stressor specific (systematic) reviews documenting key
 - 651 events
 - 652 ○ previous AOP descriptions
 - 653 - overview of data identification and search strategies, including initial and refined
 - 654 approaches, e.g.,
 - 655 ○ search terms, search strings, etc. and databases searched, the time period of
 - 656 searching, and returned results,
 - 657 - novel data – describe the type(s) of experiments that were conducted, specialized
 - 658 software and tools used for assimilation, screening and assessment of information
 - 659 for relevance to the AOP,

660

661 The description in this section provides an *overview* of the search strategy relevant to inclusion of

662 the KEs and KERs in the AOP. Considerations for documentation of more detailed information on

663 search and assimilation strategies for individual KERs is presented in Section 3.

664

665

666 1G. KE and KER Tables

667 Tables listing each KE and KER are automatically created in the AOP-KB as KE pages to
668 link to the AOP are selected or created and as KERs are defined.

- 669 • **KE Table:** This table summarises all of the KEs of the AOP, including the MIE and
670 AO. This table is populated in the AOP-Wiki as KEs are added to the AOP. Each
671 table entry acts as a link to the individual KE description page. For guidance on
672 completing the KE descriptions see Section 2.
- 673 • **Relationship Table:** This table summarises all of the KERs of the AOP and is
674 populated in the AOP-Wiki as KERs are added to the AOP. Each table entry acts as a
675 link to the individual KER description page. For guidance on completing the KER
676 descriptions see Section 3.

677

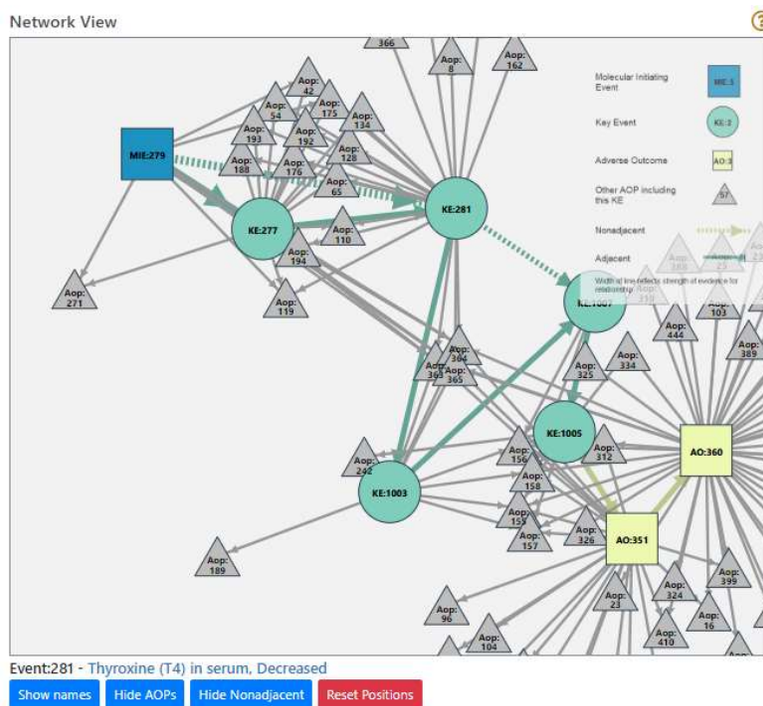
678 1H. Network View

679 The AOP-Wiki automatically generates a network view of the AOP (Figure 4). This network
680 graphic is based on the information provided in the MIE, KEs, AO, KERs and WoE summary
681 tables. The width of the arrows (representing the KERs) is determined by its WoE confidence level,
682 with thicker lines representing higher degrees of confidence. This network view also shows which
683 KEs are shared with other AOPs. Visibility of non-adjacent relationships and/or other AOPs that
684 share KEs with the AOP in question can be toggled on and off, as can the names of KEs. Users can
685 customize the layout of network representation of the viewer. If logged in, that customized view is
686 retained when returning to the AOP-Wiki.

687

688 With AOP-Wiki release 2.6 there is also an option to display the AOP in third party tools that allow
689 for alternative visualization of the AOP in an AOP network context. These third party options can
690 be accessed via the “Explore in a Third Party Tool” button.

691



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693

694 **Figure 4.** Example of the default network view in the AOP-Wiki. Note the option to hide or show
695 AOPs that share one of more or the same KEs, non-adjacent relationships, and event names.

696

697

698 1I. Prototypical Stressor(s)

699 The Prototypical Stressor field is a structured data field that can be used to identify one or more
700 “prototypical” stressors that act through this AOP. However, please recall that an AOP should
701 not be stressor-specific. Prototypical stressors are stressors for which responses at multiple key
702 events in addition to the MIE have been well documented. Experiments with the prototypical
703 stressor(s) may have provided much of the empirical support for the AOP and/or quantitative
704 understanding of the key event relationships. Thus, prototypical stressors identified may serve as
705 useful “positive controls” for evaluating responses of other stressors that may act on this pathway
706 and/or provide insights into the types of structures or properties that may be relevant to the
707 stressor domain that is relevant to this AOP. The relative potency of various other stressors,
708 compared to the prototypical stressor(s) may also be informative relative to quantitative
709 understanding of the KERs and associated applications of the AOP.

710 Please note:

- 711 ● This field is NOT intended to provide a comprehensive listing of all stressors known to
712 act through this AOP.
- 713 ● It is NOT intended that AOPs will be searchable by prototypical stressor(s)
- 714 ● Identification of a prototypical stressor does NOT indicate the AOP is stressor specific.

715 In the case of prototypical stressors that are chemicals, chemical names can be selected from
716 established chemical ontologies. However, non-chemical stressors such as radiation, genetic
717 or environmental factors, disease vectors or viruses, etc. may also be identified. Authors are
718 encouraged to utilize appropriate ontologies wherever possible.

719

720

721 **1J. Life Stage/Taxonomic/and Sex Applicability**

722 See Section 4 on Overall Assessment of the AOP

723

724 **1K. Overall Assessment of the AOP**

725 See Section 4

726

727

Development tip 4 – Sharing of KEs:

Use existing KEs when possible - when adding KEs to an AOP it is strongly recommended to use KEs that already exist in the AOP-Wiki as much as possible. When adding a new KE in the AOP-Wiki, the system will identify events using related terms to aid in reviewing whether suitable KEs already exist.

Existing KE requires modification - If an existing KE requires modification to make it suitable, changes to the content on that page should be coordinated with the point(s) of contact for other AOPs sharing the KE to ensure that the original meaning is not altered.

AOP-KB Etiquette – When using an existing KE, it is the responsibility of the person making changes to ensure that KEs used in multiple AOPs are not altered in such a way as to diminish the applicability of that KE for the existing AOPs. Please be courteous to your fellow AOP developers.

Creating new KEs - If no suitable KEs are available in the AOP-Wiki, or if the revisions needed to make an existing KE description suitable for the AOP under-development would make it unsuitable for use in AOPs it is already linked to, then a new KE should be created.

729

730 **2A. Event ID**

731 When a KE is created, an ID number is automatically assigned to it (Event: ###). This number is
732 used for tracking the KE in the AOP-KB and corresponds with a unique URL of the form
733 <https://aopwiki.org/events/###>.

734

735 **2B. KE Title**

736 The KE title should describe a discrete biological change that can be measured. It should generally
737 define the biological object or process being measured and whether it is increased, decreased, or
738 otherwise definably altered relative to a control state. For example “enzyme activity, decreased”,
739 “hormone concentration, increased”, or “growth rate, decreased”, where the specific enzyme or
740 hormone being measured is defined.

741

742 **2C. Short Name**

743 The KE short name should be a reasonable abbreviation of the KE title and is used in labelling this
744 object throughout the AOP-Wiki. The short name should be less than 80 characters in length.

745

746 **2D. Level of Biological Organisation**

747 Structured terms, selected from a drop-down menu, are used to identify the level of biological
748 organisation for each KE (e.g. molecular, cellular, organ). Note that KEs should be defined within
749 a particular level of biological organisation. Only KERs should be used to transition from one level
750 of organisation to another. Selection of the level of biological organisation defines which structured
751 terms will be available to select when defining the Event Components (below).

752

753 **2E. KE Components and Biological Context**

754

755 Because one of the aims of the AOP-Wiki is to facilitate generation of AOP networks through the
756 use of shared KE and KER elements, authors are strongly encouraged to define their KEs using a
757 set of structured ontology terms (Event Components); in the absence of structured terms, the same
758 KE could have a variety of titles. In order to make synonymous KEs more machine-readable, they
759 should be defined by one or more “event components” consisting of a **biological process**, **object**,
760 and **action** with each term originating from one of 22 biological ontologies (Ives, et al., 2017).
761 **Biological process** describes dynamics of the underlying biological system (e.g., receptor
762 signalling). The biological **object** is the subject of the perturbation (e.g., a specific biological

763 receptor that is activated or inhibited). **Action** represents the direction of perturbation of this system
764 (generally increased or decreased; e.g., ‘decreased’ in the case of a receptor that is inhibited to
765 indicate a decrease in the signalling by that receptor).

766

767 AOP-Wiki release 2.8 integrated new features to assist with Event Component selection.

768

- **Action** terms (required) are selected from a drop-down list.

769

- For **Biological Process** and **Biological Object** terms, entering a term into the search bar will search available ontologies and populate a list of potential terms to choose from. The appropriate term can then be selected using the “Add” radio button at the end of each row. A single Event Component can only have one Process and one Object term, but multiple Event Components can be added to a single Key Event .

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- A warning will display if the user attempts to add a combination of Event Component terms that have already been applied to the event.

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- Authors are strongly encouraged to make use of these features to make their AOP information more machine readable and to facilitate interoperability with other systems that rely on ontologies and controlled vocabularies.

777

778

779

Development tip 5– How specifically should my KE be defined: The following are some general recommendations and “rules of thumb” concerning how specifically to define a KE (see also Villeneuve et al. 2014a, b):

Define the KE with enough specificity that it is clear what to measure to determine the state of the KE. For example “histological changes” is too broad; “oocyte atresia” or “hyperplasia” would be better.

KEs should refer to/focus on a single measurable event within a specific biological level of organisation, rather than compounding events together. For example, it would be better to define a KE as “enzyme activity, increased” (if that can be measured), rather than “transcription and translation leading to enzyme activity, increased”.

The biological context of the KE (e.g., the tissue type/taxa/life stage/sex etc.) should only be restricted (e.g., “enzyme activity in liver, decreased” or “hormone concentration in females, increased”) to the extent that function changes with context. If the function is equivalent in both sexes, do not restrict the context by sex. If the function is equivalent in all cell types, do not restrict to a specific cell type.

780

781 **2F. Other AOPs that use this KE**

782

All of the AOPs that are linked to this KE will automatically be listed in this subsection. This table
783 can be particularly useful for identifying AOP networks which include the KE.

784

785

2G. KE Description

786

A description of the biological state being observed or measured, the biological compartment in
787 which it is measured, and its general role in the biology should be provided. For example, the
788 biological state being measured could be the activity of an enzyme, the expression of a gene or
789 abundance of an mRNA transcript, the concentration of a hormone or protein, neuronal activity,
790 heart rate, etc. The biological compartment may be a particular cell type, tissue, organ, fluid (e.g.,
791 plasma, cerebrospinal fluid), etc. The “role in the biology” could describe the reaction that an
792 enzyme catalyses and the role of that reaction within a given metabolic pathway; the protein that a
793 gene or mRNA transcript codes for and the function of that protein; the function of a hormone in
794 a given target tissue, physiological function of an organ, etc. Care should be taken to avoid
795 reference to other KEs, KERs or AOPs. Only describe this KE as a single isolated measurable
796 event/state. This will ensure that the KE is modular and can be used in other AOPs, thereby
797 facilitating construction of AOP networks. Additionally, avoid the use of semi-quantitative terms
798 that suggest an undefined threshold (e.g., insufficient, inadequate, sustained). Quantitative
799 understanding of the magnitude or duration of change in the KE required to impact a downstream
800 event should be defined in the KER (see Section 3G), not in the KE description or title.

801

802 **2H. How it is Measured or Detected**

803 One of the primary considerations in evaluating AOPs is reliability and relevance of the methods
804 used to measure the KEs. The aim of this section of the KE description is not to provide detailed
805 protocols, but rather to capture, in a sentence or two, per method, the type(s) of measurements that
806 can be employed to evaluate the KE and the relative level of scientific confidence in those
807 measurements. These can range from citation of specific validated test guidelines, to citation of
808 specific methods published in the peer reviewed literature, to outlines of a general protocol or
809 approach (e.g., a protein may be measured by ELISA).

810
811 Key considerations regarding scientific confidence in the measurement approach include whether
812 the assay is fit for purpose, whether it provides a direct or indirect measure of the biological state
813 in question, evidence that it is reproducible, and the extent to which it is accepted in the scientific
814 and/or regulatory community. Information can be obtained from the [OECD test guidelines website](#)
815 and the EURL ECVAM Database Service on Alternative Methods to Animal Experimentation
816 ([DB-ALM](#)).

817
818 **2I. Biological Domain of Applicability**

819 The biological domain(s) of applicability of the KE in terms of sex, life-stage, taxa, and other
820 aspects of biological context are defined in this section. In essence, the taxa/life-stage/sex
821 applicability is defined based on the species or groups of organisms for which the measurements
822 represented by the KEs can be made based on direct evidence from the literature (i.e., empirical
823 domain of applicability) or based on one or more lines of scientific reasoning (i.e., biologically
824 plausible domain of applicability) [see Development tip 6]. Defining the taxonomic, life stage and
825 sex relevance of each KE helps to bound the domain of applicability of the AOP as a whole and
826 provides an understanding of how broadly data represented by a KE measurement may be applied.
827

Development tip 6 – Domain of applicability: When defining domain of applicability, it is useful to think about it in two ways

Empirical domain of applicability: Species, sexes, life stages, for which there is already demonstrable evidence that the measurement can be made (KEs), the relationship applies (KERs) or the AOP in its entirety is relevant (AOPs).

Biologically plausible domain of applicability: The broad range of species, sexes, life stages for which the measurement (KE), relationship (KER), or AOP is likely to apply based on scientific reasoning (i.e., molecular conservation of targets/pathways; phylogenetic relatedness; similarity in life history; analogy).

Authors are encouraged to present both, and to clearly distinguish between the two based on the “evidence calls” made in the structured table and/or the explanatory text provided in the free text field.

828
829 As a general guide, whether defining the domain of applicability empirically or based on biological
830 plausibility, there are two primary considerations for a KE:

- 831
832 1. **Structure:** Is there evidence that the biological object being measured/observed is
833 present/conserved in the taxa/sex/life-stage of interest? Here biological object may refer
834 to a protein, a cell type, an organ, etc.
835 2. **Function:** Is there evidence that the function of that biological object and the process being
836 measured via the KE are conserved and relevant in the taxa/sex/life-stage of interest. Does
837 it play the same role?

838
839 For example, if the KE involves binding to the estrogen receptor, but invertebrates lack a functional
840 homolog of the estrogen receptor, one could reasonably conclude that the AOP is not relevant to
841 invertebrates on the basis of a lack of conserved structure. Evidence supporting this biologically
842 plausible taxonomic domain of applicability could be collected from bioinformatics approaches
843 and existing toxicity data across species to support this broad extrapolation to all invertebrates.

844 Depending on the evidence supporting the taxonomic domain of applicability, the specific
845 (common or Latin) species name or taxonomic group (e.g., class, order, family) may be reported
846 with the appropriate NCBI taxonomy ID in the “Taxonomic Applicability” table of the AOP-Wiki.
847 Likewise, if the KE involves a measurement in ovarian tissue, its applicability domain in terms of
848 sex would be restricted to females. Such information would be captured in the “Sex Applicability”
849 table of the AOP-Wiki using predefined terms like: male, female, mixed, asexual, hermaphrodite,
850 or unspecified. If a KE involved altered organogenesis (e.g., heart formation), the KE would only
851 be relevant to the life-stage during which the heart is actually formed, not adult life stages in which
852 organ development has already completed. Life-stage can be described in the “Life Stage” table of
853 the AOP-Wiki by selecting from structured ontology terms. If an applicable life-stage term cannot
854 be found, new terms may be added on request by the AOP-Wiki administrators.

855
856 Biological domain of applicability is defined in the AOP-KB using a combination of structured
857 fields and free text. Selection of structured terms to describe the applicability domain can aid AOP
858 network construction as well as facilitating other types of computational processing and searching
859 of information captured in the AOP-KB.

860
861 When the developer selects structured ontology terms to help define the domain of applicability of
862 the KE, there is also an option to make evidence calls related to applicability of the specific KE for
863 that category term. These calls should be based on expert knowledge of the biology and the extent
864 of supporting evidence. Recommendations for these calls are:

- 865
866
- 867 • Low: With the understanding that by definition a KE must be measurable in the
868 species/taxonomic group/lifestage/sex defined, no such measurements have been
869 reported or shown experimentally *in vitro* or *in vivo* to date; however, there are one or
870 more scientifically-based lines of evidence suggesting that measurement could plausibly
871 be made (e.g., in silico or bioinformatic evidence of protein or pathway conservation).
 - 872 • Moderate: The measurement associated with the KE can plausibly be made for the
873 species/taxonomic group/lifestage/sex, and there is at least some supporting *in vitro* or *in*
874 *vivo* experimental evidence, although though it may not involve direct measurement of
875 the KE.
 - 876 • High: The measurement associated with the KE has been made repeatedly *in vitro* or *in*
877 *vivo* and/or with multiple orthogonal methods for the species/taxonomic
878 group/lifestage/sex.

879 ***i. Taxonomic Applicability***

880 Latin or common names of a species or broader taxonomic grouping (e.g., class, order, family)
881 can be selected from an ontology. In many cases, individual species identified in these structured
882 fields will be those for which the evidence used in constructing the AOP was strongest in
883 relation to this KE.

884
885 ***ii. Life Stage Applicability***

886 The structured ontology terms for life-stage are more comprehensive than those for taxa, but
887 may still require further description/development and explanation in the free text section.

888
889 ***iii. Sex Applicability***

890 The authors must select from one of the following: Male, female, mixed, asexual, third gender,
891 hermaphrodite, or unspecified.

892
893 ***iv. Evidence for Biological Domain of Applicability***

894 This free text section should be used to elaborate on the scientific basis for the indicated
895 domains of applicability and the WoE calls (if provided). While structured terms may be
896 selected to define the taxonomic, life stage and sex applicability (see structured applicability
897 terms, above) of the KE, the structured terms may not adequately reflect or capture the
898 overall biological applicability domain (particularly with regard to taxa). Likewise, the

899 structured terms do not provide an explanation or rationale for the selection. The free-text
900 section on evidence for taxonomic, life stage, and sex applicability can be used to elaborate
901 on why the specific structured terms were selected, and provide supporting evidence,
902 references and background information. This information should also indicate the type of data
903 used as evidence (e.g., in silico, in vitro, in vivo).

904

905 **2J. AO-Specific Content**

906 An AO is a specialised KE that represents an adverse outcome of regulatory significance, (“apical
907 endpoint”). For KEs that are designated as an AO, one additional field of information (regulatory
908 significance of the AO) should be completed, to the extent feasible. If the KE is being described is
909 not an AO, simply indicate “not an AO” in this section.

910

911

Regulatory Significance of the AO

912 A key criterion for defining an AO is its relevance for regulatory decision-making (i.e., it
913 corresponds to an accepted protection goal or common apical endpoint in an established
914 regulatory guideline study). For example, in humans this may constitute increased risk of
915 disease-related pathology in a particular organ or organ system in an individual or in either the
916 entire or a specified subset of the population. In wildlife, this will most often be an outcome of
917 demographic significance, e.g., population sustainability. In addition to describing the biological
918 state associated with the AO, how it can be measured, and its taxonomic, life stage, and sex
919 applicability, it is useful to describe regulatory examples using this AO.

920

921

922 **2K. References**

923 List of the literature that was cited for this KE description. References should either be numbered
924 [#], and cited by number, or cited in (Author, Year) style at locations on the Event page
925 corresponding to the statement(s) they support. Ideally, the list of references should conform with
926 the OECD Style Guide ([https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-
927 Edition.pdf](https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-Edition.pdf)) (OECD, 2015).

928

929

930

931

932

933 **SECTION 3 – KER DESCRIPTIONS**

934

935 The utility of AOPs for regulatory application is defined, to a large extent, by the confidence and
936 precision with which they facilitate extrapolation of data measured at low levels of biological
937 organisation to predicted outcomes at higher levels of organisation and the extent to which they can
938 link biological effect measurements to their specific causes. Within the AOP framework, the
939 predictive relationships that facilitate extrapolation are represented by the KERs. Consequently, the
940 overall WoE for an AOP is a reflection in part, of the level of confidence in the underlying series of
941 KERs it encompasses. Evidence related to determination of confidence in the supporting data for the
942 KER as part of the AOP is included here. The confidence in the overall AOP pathway is considered
943 in Section 4, taking into account the KER specific evidence and patterns of support across all levels
944 of biological organization in the AOP.

945

946 Describing the KERs in an AOP involves assembling and organising the types of information and
947 evidence that defines the scientific basis for inferring the probable change in, or state of, a
948 downstream KE from the known or measured state of an upstream KE. Before describing a KER,
949 developers should carefully consider the following:

950

951 KERs are always described in the form of a directed relationship (one-way arrow) linking an
952 upstream “causing” event to a downstream “responding” event. The pair of KEs linked via a KER
953 may either be adjacent to one another in the sequence of KEs that define a given AOP, or non-

954 adjacent (Figure 5). Regardless of adjacency, one event is always positioned upstream of the other.
 955 By convention (and for clarity), KERs linking adjacent KEs in an AOP are represented using solid
 956 arrows, while KERs that link KEs that are not adjacent to one another in sequence are linked via
 957 dashed arrows (e.g., Figure 5). This is a graphical convention only which has no bearing on the type
 958 of content to include in the KER description.

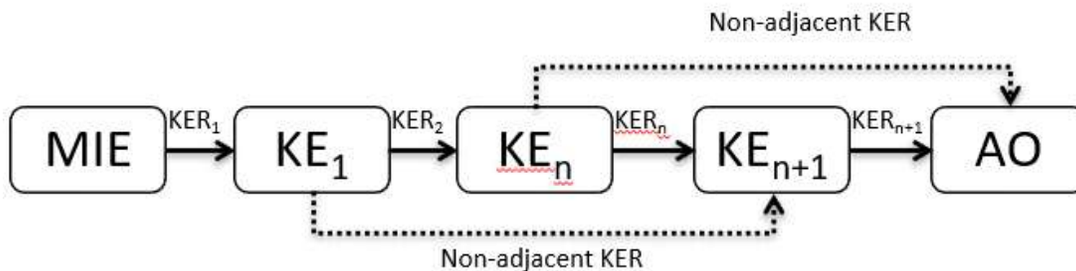
959

960 A KER description must be created for each adjacent upstream-downstream pair of KEs in the
 961 pathway. Graphically speaking, there should always be at least one solid arrow path connecting each
 962 KE in the pathway into a sequence. There should be no KEs that are unconnected or are only
 963 connected via a non-adjacent path (represented as a dashed arrow) only.

964

965 Inclusion and description of non-adjacent KERs within an AOP can be particularly useful for
 966 assembling evidence supporting the AOP and in the consideration of the overall support across the
 967 entire AOP (section 4). For example, some KE measurements may be fairly difficult to make, such
 968 that they are rarely made in routine studies. While there may be sufficient data or plausibility to
 969 establish an intermediate KE as part of the AOP, much of the available WoE may ignore or “leap
 970 over” that particular KE. Including KER descriptions for non-adjacent KE pairs allows the WoE for
 971 these relationships to be readily described and linked to other AOPs without compromising the
 972 principle of modularity with regard to the KER descriptions. With this in mind, the upstream-
 973 downstream pair of KEs linked via a KER may be adjacent in one AOP and non-adjacent in another
 974 (Figure 6).

975

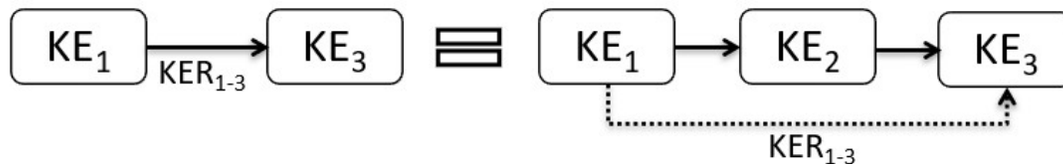


976

977

978 **Figure 5.** Generic AOP diagram illustrating the graphical convention for depicting KERs linking
 979 adjacent (solid arrow) versus non-adjacent (dashed arrow) upstream-downstream KE pairs within an
 980 AOP. Regardless of adjacency, each KER represents a predictive relationship between a pair of KEs
 981 and can be supported by WoE.

982



983

984

985 **Figure 6.** Graphical depiction of the modular functionality of KERs connecting KE1 to KE3. The
 986 content of KER1-3 is identical despite the fact that the KE1 and KE3 are adjacent in one AOP and
 987 non-adjacent in the other.

988

989 Overall, the subsections of the KER descriptions are intended to aid the user in collecting relevant
 990 information that will support evaluation of the level of confidence in each KER, which in turn
 991 contributes to the assessment of the WoE of the AOP overall (section 4).

992

993

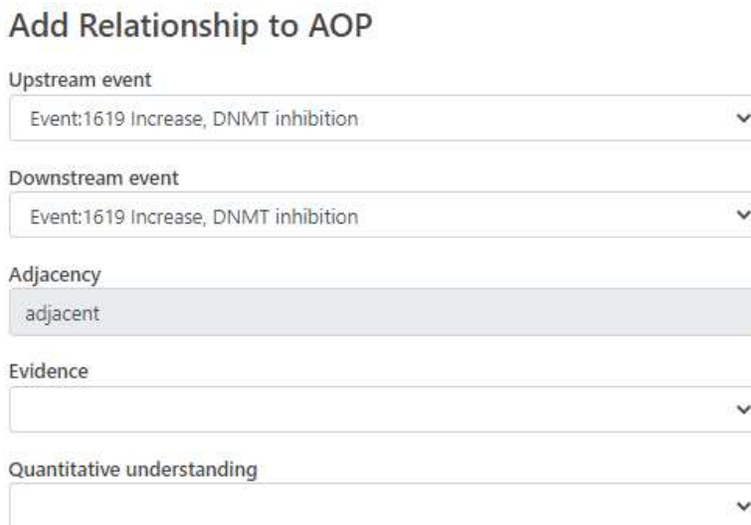
994

3A. Relationship ID

995 When a KER is created, an ID number is automatically assigned to it (Relationship: ###). This
996 number is used for tracking the KER in the AOP-KB and corresponds with a unique URL of the
997 form <https://aopwiki.org/relationships/###>.
998

999 3B. KER Title

1000 All KER titles take the form “upstream KE leads to downstream KE”. KER titles are generated
1001 automatically by selecting an upstream KE and downstream KE to link in the AOP-Wiki (Figure 7).
1002



The image shows a web form titled "Add Relationship to AOP". It consists of five vertically stacked dropdown menus. The first two are labeled "Upstream event" and "Downstream event", both showing the selected option "Event:1619 Increase, DNMT inhibition". The third is labeled "Adjacency" and shows "adjacent". The fourth is labeled "Evidence" and is currently empty. The fifth is labeled "Quantitative understanding" and is also empty. Each dropdown menu has a small downward-pointing arrow on its right side.

1003
1004

1005 **Figure 7.** Add Relationship dialog from AOP-Wiki. Note, user will select KEs from a drop-down
1006 menu of options, therefore the KER title is created automatically. This also means that the KEs
1007 must be created before a KER can be defined.
1008

1009 3C. AOPs Referencing Relationship

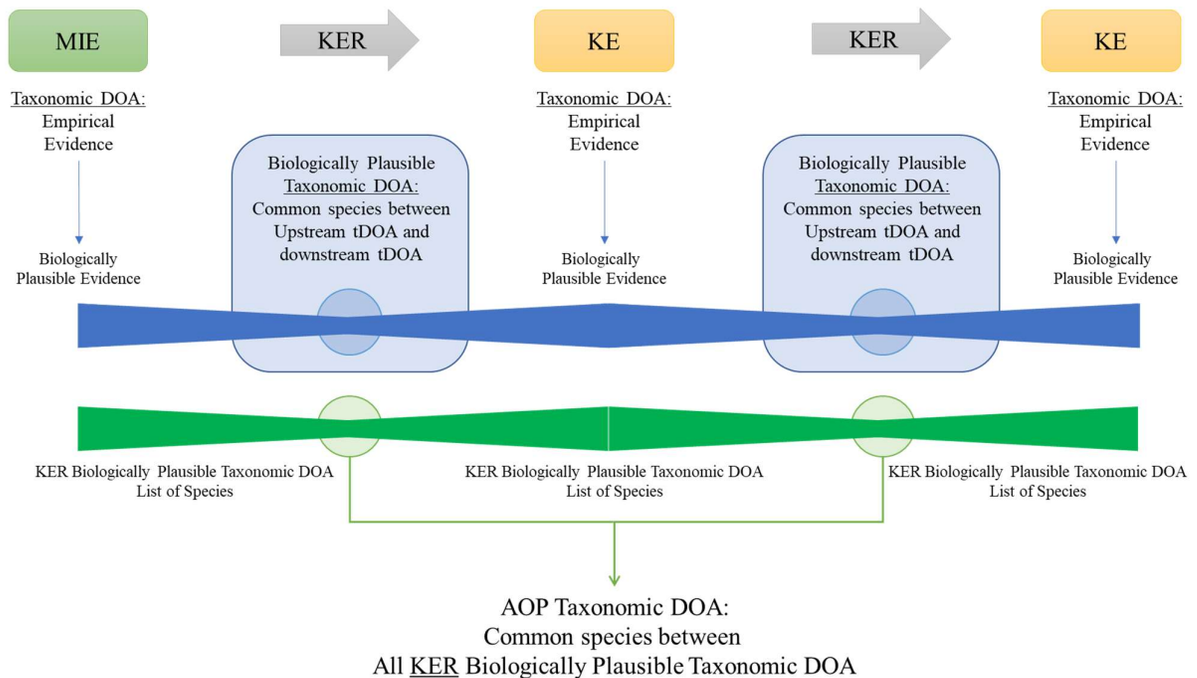
1010 All of the AOPs that are linked to this KER will automatically be listed in this subsection.
1011

1012 3D. Biological Domain of Applicability

1013 Developers have the option to select one or more structured terms that help to define the biological
1014 applicability domain of the KER. As a rule, the biological domain of applicability of a KER can
1015 never be broader than the more restrictive of the two KEs it links together. For example, if the
1016 upstream KE is relevant to all vertebrates but the downstream KE is relevant only to sexually
1017 mature, egg-laying female vertebrates, the KER would be relevant to sexually mature egg-laying
1018 female vertebrates. This concept applies whether considering the empirical domain of
1019 applicability, or the biologically plausible domain of applicability and once again authors should
1020 clearly indicate both.
1021

1022 Thus, the biological applicability domains of the two KEs being linked is a strong determinant of
1023 the biological domain of applicability of a KER (Figure 8).

1024
1025



1026 Figure 8. Example for determining the taxonomic domain of applicability (tDOA) considering both
1027 the empirical evidence and biologically plausible evidence and combining upstream KE and
1028 downstream KE tDOA to determine KER tDOA. Further, considering the KER tDOAs across the
1029 AOP the most restrictive tDOA across all KERs defines the tDOA for the AOP. The blue horizontal
1030 line considers each KE to define the biologically plausible tDOA of the KER, whereas the green
1031 horizontal line considers each KER to define the biologically plausible tDOA for the entire AOP.
1032 Figure modified from Jensen et al. 2022.

1033

1034 However, in some cases, the biological applicability domain of the KER may be even more
1035 restrictive. This is because in addition to structural and functional conservation, the KER also
1036 considers the conservation of a biological relationship between two KEs. The three considerations
1037 that generally guide definition of the biological domain of applicability are thus:

1038

- 1039 1. Structure: Is there evidence that the biological object(s) being measured/observed in the
1040 context of the two KEs being linked present/conserved in the taxa/sex/life-stage of
1041 interest?
- 1042 2. Function: Is there evidence that the functions of those biological objects and the
1043 processes being measured in the two KEs are conserved and relevant in the taxa/sex/life-
1044 stage of interest? Does the object/process play the same role in both KEs?
- 1045 3. Regulation: Is there evidence that the regulation of the KEdownstream by KEupstream
1046 is conserved and relevant in the taxa/sex/life-stage of interest?

1047

1048 Selection of structured terms to describe the biological domain of applicability can aid AOP network
1049 construction as well as facilitating other types of computational processing and searching of
1050 information captured in the AOP-Wiki.

1051

1052 Upon selection of structured biological applicability domain terms, developers have the option to
1053 classify the extent of the supporting evidence for the terms they have selected:

1054

- 1055 • Low the relationship is biologically plausible, but has not been shown experimentally *in*

1056

1057 *in vitro* or *in vivo* in this species/taxonomic group/lifestage/sex; evidence may be
1058 computationally derived by models or other available tools for evaluating structural and
1059 functional conservation (e.g., *in silico* or bioinformatic evidence of protein or pathway
1060 conservation).

- 1061 • **Moderate** the relationship is biologically plausible, and there is some limited supporting *in*
1062 *vitro* and/or *in vivo* experimental evidence in the species/taxonomic group/lifestage/sex of
1063 interest; computationally derived data to support the biologically plausible domain of
1064 applicability could be included as evidence toward structural conservation and used for
1065 extrapolation.
- 1066 • **High** the relationship is biologically plausible, and there is considerable supporting evidence
1067 in the species/taxonomic group/lifestage/sex, including evidence of temporal, dose-
1068 response, and/or incidence concordance between the two KEs for the group in question.

1070
1071 **i. Taxonomic Applicability**

1072 Authors can indicate the relevant taxa for this KER in this subsection. The process is similar to
1073 that described for KEs (Section 2).

1074
1075 **ii. Life Stage Applicability**

1076 Authors can indicate the relevant life stage for this KER in this subsection. The process is similar
1077 to that described for KEs (Section 2).

1078
1079 **iii. Sex Applicability**

1080 Authors can indicate the relevant sex for this KER in this subsection. The process is similar to
1081 that described for KEs (Section 2).

1082
1083 **iv. Evidence Supporting the Biological Domain of Applicability**

1084 As for the KEs, there is also a free-text section of the KER description that the developer can use
1085 to explain his/her rationale for the structured terms selected with regard to taxonomic, life stage, or
1086 sex applicability, or provide a more exact description of the applicability domain than may be
1087 feasible using standardised terms. Developers are also encouraged to distinguish the empirical
1088 domain of applicability from the more expansive biologically plausible domain of applicability
1089 (see *Development tip 5*). Here developers can indicate what type(s) of evidence were used to
1090 support the domain of applicability (e.g., *in silico*, *in vitro*, *in vivo*) and cite the methods if
1091 relevant.

1092
1093
1094 **3E. KER Description**

1095 Provide a brief, descriptive summation of the KER. While the title itself is fairly descriptive, this
1096 section can provide details that are not inherent in the description of the KEs themselves (see Section
1097 2, recommendations regarding number of KEs to include). For example, if the upstream KE was
1098 antagonism of a specific receptor, the description could stipulate that “persistent antagonism of the
1099 receptor for a period of days” will trigger the downstream KE. Shorter term antagonism of the same
1100 receptor (i.e., same upstream KE) may trigger a different downstream KE, and thus would be
1101 described in a different KER. This description section can be viewed as providing the increased
1102 specificity in the nature of upstream perturbation (KE_{upstream}) that leads to a particular downstream
1103 perturbation (KE_{downstream}), while allowing the KE descriptions to remain generalised so they can
1104 be linked to different AOPs. The description is also intended to provide a concise overview for
1105 readers who may want a brief summation, without needing to read through the detailed support for
1106 the relationship (covered below). Care should be taken to avoid reference to other KEs that are not
1107 part of this KER, other KERs or other AOPs. This will ensure that the KER is modular and can be
1108 used by other AOPs.

1109
1110 **3F. Evidence Collection Strategy**

1111 Include a description of the approach for identification and assembly of the evidence base for the

- 1112 KER. For the literature searches and surveys, include, for example:
1113
1114 i. Sources and dates of information consulted including expert knowledge, databases searched and
1115 associated search terms/strings,
1116 ii. Study screening criteria and methodology (e.g., inclusion/exclusion criteria, specialized software
1117 tools, number of reviewers); any constraints on the search.
1118 iii. Study quality assessment considerations including links to existing resources (e.g., existing tools
1119 applied)
1120 iii. Data extraction strategy, specialized software tools and/or data management strategy, and
1121 iv. Links to any repositories/databases of relevant references

1122
1123 Tabular summaries and links to relevant supporting documentation are encouraged, wherever
1124 possible.

1125
1126 Alternatives to literature search-based approaches include, but are not limited to, novel
1127 experimentation, application of biologically-based models, identification of sources of canonical
1128 knowledge, etc.

1129
1130 **3G. Evidence Supporting this KER**

1131 Assembly and description of the scientific evidence supporting KERs in an AOP is an important
1132 step in the AOP development process that sets the stage for overall assessment of the AOP relevant
1133 to regulatory application (Section 4). To do this, biological plausibility, empirical support, and the
1134 current quantitative understanding of the KER are evaluated with regard to the predictive
1135 relationships/associations between defined pairs of KEs as a basis for considering WoE (Section 4).
1136 In addition, uncertainties and inconsistencies are considered.

1137
1138 ***i. Biological Plausibility***

1139 Define, in free text, the biological rationale for a connection between KEupstream and
1140 KEdownstream. What are the structural or functional relationships between the Kes (see Annex
1141 1)? For example, there is a functional relationship between an enzyme's activity and the product
1142 of a reaction it catalyses.

1143
1144 Contextual citation of supporting references should be included. However, it is recognised that
1145 there may be cases where the biological relationship between two KEs is very well established,
1146 to the extent that it is widely accepted and consistently supported by so much literature that it is
1147 unnecessary and impractical to cite the relevant primary literature (i.e., canonical knowledge).
1148 Citation of review articles or other secondary sources, like text books, may be reasonable in such
1149 cases. The primary intent is to provide scientifically credible support for the structural and/or
1150 functional relationship between the pair of KEs if one is known.

1151
1152 In general, the structural and/or functional relationship supporting biological plausibility is based
1153 on understanding of "normal" biological function, rather than response to a specific stressor. The
1154 description of biological plausibility can also incorporate additional mechanistic detail that helps
1155 inform the relationship between KEs, but is not practical/pragmatic to represent as separate KEs
1156 due to the difficulty or relative infrequency with which it is likely to be measured. For example,
1157 in the case of G protein coupled receptor activation (KEupstream) leading to increased activity of
1158 a specific enzyme (KEdownstream), there may be numerous mechanistic steps between these KEs
1159 (e.g., alterations in signal transduction pathways, transcriptional regulation, post-translational
1160 modifications, etc.). These underlying details, if known, can be captured in the description of
1161 biological plausibility (if desired) rather than represented as independent KEs. The KER
1162 descriptions are the appropriate place for "embedding" this type of biological detail without
1163 compromising the reusability of the KE descriptions within the AOP-Wiki. However, it should
1164 be kept in mind that added detail should only be included to the extent that it enhances the
1165 predictive utility of the AOP for regulatory application. Detail may be particularly useful in
1166 considering the differences across taxonomic groups or species that may dictate the broad utility

1167 of the AOP (i.e., the taxonomic domain of applicability). In part, the AOP is intended to filter
 1168 through much of the mechanistic detail to focus on what important causal events for the adverse
 1169 outcome have predictive value for regulatory application. Thus, efforts should be made to keep
 1170 the descriptions focused and concise.

1171

1172 **Empirical Evidence**

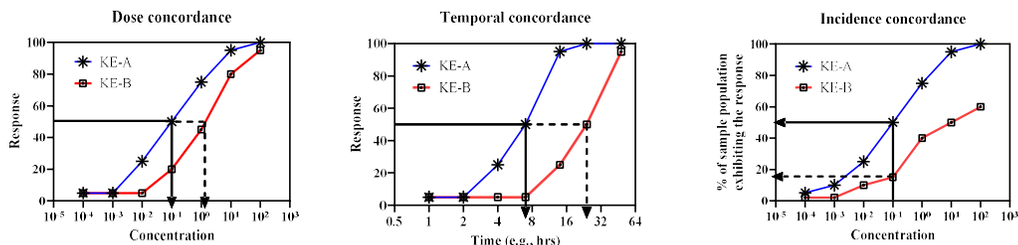
1173 In this section authors are encouraged to cite specific evidence relevant to assessment of changes
 1174 in the upstream KE (KE_{upstream}) leading to, or being associated with, a predictable subsequent
 1175 change in the downstream KE (KE_{downstream}).

1176

1177 In particular, it is useful to cite direct evidence showing that stressors that perturb KE_{upstream}
 1178 also perturb KE_{downstream}. Because this section of the KER description cites evidence from
 1179 specific studies, it is also helpful to provide as much detail as possible about the toxicological and
 1180 biological context in which the measurements were made. While the KER itself is not intended to
 1181 be stressor-specific, this information addresses whether supporting data on quantitative patterns
 1182 of relationships between key events is consistent with what's expected, if the KER is operative.
 1183 Expected patterns are that the upstream KE is impacted at doses/concentrations of the stressor that
 1184 are equal to or lower than those that impact the downstream KE (dose concordance; Figure 9),
 1185 that at any given dose of stressor, the upstream is impacted earlier in the time-course of exposure
 1186 than the downstream event (temporal concordance; Figure 9), and likewise for any given dose and
 1187 duration of exposure to the stressor, the upstream event is observed in an equal to or greater
 1188 proportion of the sample population than the downstream event (incidence concordance; Figure
 1189 9). Deviations from these expected patterns may be due to factors such as experimental design,
 1190 the relative sensitivity of methods for measuring KEs, and other factors; experimental details that
 1191 could influence apparent concordance or lack thereof, should therefore be considered when
 1192 assembling and presenting evidence.

1193

1194



1195

1196 **Figure 9.** Examples of dose concordance, temporal concordance, and incidence concordance.
 1197 Note that dose concordance and temporal concordance are comparing the relative dose or time at
 1198 which a defined level of response is observed for KE_A compared to KE_B. Incidence concordance
 1199 compared the fraction of the population impacted at the same dose and time point for KE_A versus
 1200 KE_B.

1201

1202

1203 The consideration of empirical support in the form of bulleted lists or tables that include a short
 1204 description of the nature of the observed empirical support along with the corresponding
 1205 reference(s) is preferred as a basis to consider whether available data consistently supports
 1206 expected patterns. An example is provided below (Table 3). However, authors are free to modify
 1207 the format to best suit their approach to support the consideration of weight of evidence for the
 1208 pathway. To the extent possible, entries in the table should be based on benchmark doses to
 1209 facilitate comparative assessment of effect, thus normalizing for groupsizes and dose spacing.
 1210

1211

Table 3. Example of an empirical evidence table assembled for a KER¹.

Species,	Stressor(s)	Upstream	Downstream	Effect on	Effect on	Citation
----------	-------------	----------	------------	-----------	-----------	----------

life-stage, sex tested		Effect (Y/N)	Effect (Y/N)	Upstream Event (descriptive)	Downstream Event (descriptive)	
Adult, female, rainbow trout	Gemfibrozil	Y	Y	Benchmark dose (BMD) 15 µg/L	BMD 45 µg/L	Smith et al. 1978
Adult, F, Sprague Dawley rat	Low fat diet	Y	N	Significant decrease at 100 mg/kg/day, after 3 days	No effect at concentrations up to 2 g/kg/d, fed up to 10 days	Zonk 2018
Juvenile, M, mouse	Clofibrac acid	N	Y	BMD 45 mg/kg/d, measured 5 d post-injection	BMD 5 mg/kg/d, measured 5 d post-injection	Doe et al. 2012
Larval zebrafish	UV radiation @ UV index = 90	Y	Y	Significant decrease in 80% of sampled population after 48 h	Significant increase in 22% of sampled population after 96 h	Lee et al. 1994

¹ Entries in this table are for illustrative purposes only. They do not refer to results from real studies. Any resemblance to existing scientific results or authors is coincidental.

Dose Concordance

In the case of dose-response concordance, the aim is not to consider dose-dependence of a single KE in the pair, but rather to assess the extent of the evidence that KE upstream is generally impacted at doses (or stressor severities) equal to or less than those at which KE downstream is impacted (data row 2 of Table 3 shows an example of dose concordance; row 3 does not follow the expected pattern for dose concordance).

Temporal Concordance

In the case of temporal concordance, it is desirable to assemble evidence relevant to assessing whether effects on KE upstream are observed earlier in a time-course than effects on the downstream KE (data row 3 of Table 3 shows an example of temporal concordance, as well as dose concordance).

Incidence Concordance

In the case of incidence concordance, evidence should be assembled that addresses whether, at an equivalent dose or stressor severity, KEupstream occurs more frequently than KEdownstream (data row 4 of Table 3 shows an example of incidence concordance, as well as temporal concordance).

Other Evidence (optional)

Although evidence that demonstrates dose, temporal or incidence concordance is preferred, other evidence that empirically supports the relations that a sufficient change in KEupstream will lead to a change in KEdownstream, but do not fall into the above three categories, can be cited in this subsection.

Uncertainties and Inconsistencies

In addition to outlining the evidence supporting a particular linkage, it is also important to identify inconsistencies or uncertainties in the relationship. This could include, for example, empirical

1243 evidence showing changes in KEupstream that did not elicit alterations in KEdownstream. It could
1244 also include descriptions of gaps in biological understanding that lend to uncertainties in
1245 understanding of the exact nature of the structural or functional relationship between the two KEs.
1246 Additionally, while there are expected patterns of concordance that support a causal linkage
1247 between the KEs in the pair, it is also helpful to identify experimental details that may explain
1248 apparent deviations from the expected patterns of concordance. An example of this would be a
1249 case where methods for measuring the upstream KE are relatively insensitive compared to those
1250 for measuring the downstream KE, leading to the appearance of dose-response or incidence
1251 discordance that is simply an artefact of the measurement techniques employed. In this regard,
1252 when assembling information from multiple disparate studies, it is important to capture variables
1253 that directly influence how well concordance can be assessed (i.e., information regarding the
1254 doses tested in various experiments and the time-points at which various KE measurements were
1255 made). Identification of uncertainties and inconsistencies contributes to evaluation of the overall
1256 WoE supporting the AOPs that contain a given KER (see Section 4), and to the identification of
1257 research gaps that warrant investigation.

1258

1259 Given that AOPs are intended to support regulatory applications, AOP developers should focus
1260 on those inconsistencies or gaps that would have a direct bearing or impact on the confidence in
1261 the KER and its use as part of an AOP for inference or extrapolation in a regulatory setting.
1262 Uncertainties that would have little impact on regulatory application do not need to be described.
1263 In general, this section details evidence that may raise questions regarding the overall validity and
1264 predictive utility of the KER (including consideration of both biological plausibility and empirical
1265 support). It also contributes, along with other elements, to the overall evaluation of the WoE for
1266 the KER (see, Section 4).

1267

1268 **3H. Known Modulating Factors**

1269 This section presents information regarding modulating factors/variables known to alter
1270 quantitative aspects of the response-response function that describes the relationship between
1271 the two KEs (for example, an iodine deficient diet causes a significant increase in the
1272 sensitivity of the downstream event to changes in the upstream event [alters the slope of the
1273 relationship]; a particular genotype doubles the sensitivity of KEdownstream to changes in
1274 KEupstream). Information on these known modulating factors should be listed in this
1275 subsection, along with relevant information regarding the manner in which the modulating
1276 factor alters the relationship (if known). Note: this section should focus on those modulating
1277 factors for which solid evidence supported by relevant data and literature are available. It
1278 should NOT list all possible/plausible modulating factors. In this regard, it is useful to bear in
1279 mind that many risk assessments conducted through conventional apical guideline testing-
1280 based approaches generally consider few if any modulating factors.

1281

1282 It is recommended that information regarding known modulating factors be captured in a
1283 tabular format (Table 4), providing the following information about each:

- 1284 • What it is – the modulating factor for which there is solid evidence that it influences
1285 this KER.
- 1286 • Details of the modulating factor – specify which features (classes or subsets?) of this
1287 modulating factor are relevant for this KER.
- 1288 • Describe the known effect(s) of the modulating factor on the KER.
 - 1289 ○ E.g., increases magnitude of effect on downstream KE by two-fold
 - 1290 ○ E.g., reduces the probability of effect on the downstream event by 40%
 - 1291 ○ E.g., delays onset of the downstream event by 12-18 h
 - 1292 ○ E.g., increases sensitivity to the upstream event by a factor of four
- 1293 • Reference(s) – provide one or more references that provide supporting scientific
1294 evidence that establishes the effect of the modulating factor on the KER.

1295

1296 **Table 4.** Recommended tabular format for capturing information regarding known modulating
1297 factors¹.

Modulating Factors	MF details	Effects on the KER	References
Age	>55 years old (human)	Sensitivity of downstream event to change in upstream event increased by factor of 4	Smith et al. 1978
Genotype	BRCA1 truncation mutation in nucleotides 2401-4109)	Probability of downstream event increased by 40%	Zonk 2018
Diet	Iodine deficient	Delays onset of downstream effect by 5-10 d	Doe et al. 2012
Disease state	Type 2 diabetes	Increases risk of downstream event by 10 fold	Lee et al. 1994
Previous exposure	Within 3 years of Covid 19 infection	Magnitude of effect on downstream event increased 2-fold Delay	Walla Walla and Grant, 2022

1299 ¹ Entries in this table are for illustrative purposes only. They do not refer to results from real
1300 studies. Any resemblance to existing scientific results or authors is coincidental.

1301

1302

1303 3I. Quantitative Understanding

1304 The quantitative understanding section of the KER description is intended to capture information
1305 that helps to define how much change in the upstream KE, and/or for how long, is needed to elicit
1306 a detectable and defined change in the downstream KE. While empirical support (see previous
1307 section 3G Evidence Supporting this KER) addresses whether data on the relationship between the
1308 two KEs are consistent with the patterns that are expected if the upstream event is causing the
1309 downstream event, the quantitative understanding section helps to define the precision with which
1310 the state of the downstream KE can be predicted from knowledge of the state of the upstream KE.
1311 The higher the confidence in empirical support for a KER, the greater the likelihood that the
1312 response-response relationship can be quantified. These quantitative relationships may be defined
1313 in terms of correlations, response-response relationships, dose-dependent transitions or points of
1314 departure (i.e., a threshold of change in KE_{upstream} needed to elicit a change in KE_{downstream}),
1315 etc. They may take the form of simple mathematical equations or sophisticated biologically-based
1316 computational models that consider other modulating factors such as compensatory responses, or
1317 interactions with other biological or environmental variables. Regardless of form, the idea is to
1318 briefly describe what is known regarding the quantitative relationship between the KEs and cite
1319 appropriate literature that defines those relationships and/or provides support for them.

1320

1321 Data that confer quantitative understanding of a KER are not necessarily independent of those
1322 addressing other weight of evidence considerations. Rather, the quantitative understanding section
1323 collects additional detail about the nature of the quantitative relationship generally from the same
1324 studies used to establish empirical support. These further details are intended to support
1325 quantitative prediction of the probability or magnitude of change in KE_{downstream} based on a
1326 known state of KE_{upstream}. For transparency, the toxicological and biological context in which

1327 the quantitative relationships were defined should be indicated within the description. The ultimate
1328 goal is to identify quantitative relationships that generalise across the entire applicability domain of
1329 the two KEs being linked via the KER.

1330

1331 Based on recommendations from workshops held in September 2015 (Wittwehr et al. 2016) and
1332 April 2017 (LaLone et al. 2017), description of the quantitative understanding of the KER has been
1333 organised into subsections in order to more consistently capture information useful for both
1334 quantitative AOP and AOP network applications. As with other areas of the AOP descriptions,
1335 authors are encouraged to complete the subsections to the extent feasible, but it is recognized that
1336 supporting information may not be adequate to address all.

1337

1338 ***i. Response-response relationship***

1339 This subsection should be used to define sources of data that define the response-response
1340 relationships between the KEs. A response-response relationship is a mathematical
1341 function that describes the magnitude, probability, or severity of change in the
1342 downstream KE (B) as a function of the measured (or predicted) state of the upstream
1343 KE (A). Information regarding the general form of the relationship (e.g., linear, exponential,
1344 sigmoidal, threshold, etc.) should be captured if possible. If there are specific mathematical
1345 functions or computational models relevant to the KER in question that have been defined,
1346 those should also be cited and/or described where possible, along with information concerning
1347 the approximate range of certainty with which the state of the KE_{downstream} can be predicted
1348 based on the measured state of the KE_{upstream} (i.e., can it be predicted within a factor of two,
1349 or within three orders of magnitude?). For example, a regression equation may reasonably
1350 describe the response-response relationship between the two KEs, but that relationship may
1351 have only been validated/tested in a single species under steady state exposure conditions. It is
1352 important to note such uncertainties.

1353

1354 ***ii. Time-scale***

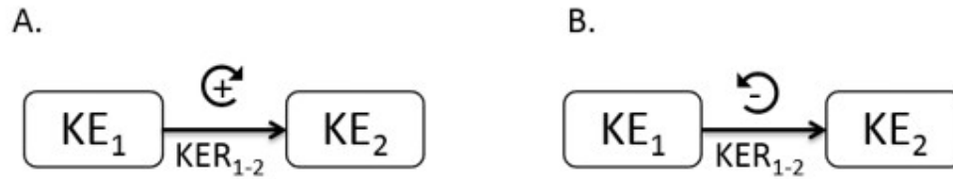
1355 This sub-section should be used to provide information regarding the approximate time-scale
1356 of the changes in KE_{downstream} relative to changes in KE_{upstream} (i.e., do effects on
1357 KE_{downstream} lag those on KE_{upstream} by seconds, minutes, hours, or days?). This can be
1358 useful information both in terms of modelling the KER, as well as for analysing the critical or
1359 dominant paths through an AOP network (e.g., identification of an AO that could kill an
1360 organism in a matter of hours will generally be of higher priority than other potential AOs that
1361 take weeks or months to develop). Identification of time-scale can also aid the assessment of
1362 temporal concordance. For example, for a KER that operates on a time-scale of days,
1363 measurement of both KEs after just hours of exposure in a short-term experiment could lead to
1364 incorrect conclusions regarding dose-response or temporal concordance if the time-scale of the
1365 upstream to downstream transition was not considered.

1366

1367

1368 ***iii. Known Feedforward/Feedback loops influencing this KER***

1369 KERs are depicted in a manner that suggests that the upstream event is independent of the
1370 downstream event. However, in biological systems, feedback relationships are common. This
1371 subsection should define whether there are known positive or negative feedback loops involved
1372 and what is understood about their time-course and homeostatic limits. In some cases where
1373 feedback processes are measurable and causally linked to the outcome, they may be represented
1374 as KEs (see development tip 5). However, in most cases these features are expected to
1375 predominantly influence the shape of the response-response and time-course, behaviours
1376 between selected KEs (i.e., the KER). For example, if a feedback loop acts as an auto-regulatory
1377 loop designed to maintain a homeostatic range of concentrations between some upper and lower
1378 limit, the feedback loop will directly shape the response-response relationship between the KEs.
1379 It is recommended that an annotation indicating a positive or negative feedback loop (Figure
1380 10) in a KER be added to the graphical representation, and that details be provided in this
1381 subsection of the KER description.



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Figure 10. Recommended graphical annotation to indicate that a known (A) positive feedback (i.e., feedforward) or (B) negative feedback loop is involved in the transition from one KE to the next in the AOP. Note, this is an optional annotation. See Development tip 7 for more information on describing positive and negative feedback processes using the AOP framework.

Development tip 7 – Capturing information on positive or negative feedback loops.

Ways to capture/represent known positive or negative feedback loops have emerged as a frequently asked question in relation to use of the AOP framework. Thus, a few general guidelines are provided here.

In cases where feedback loops play a direct causal role in the progression of a biological perturbation leading to an AO, they can be included as KEs as long as they are measurable. For example, for an AOP in which a negative feedback process results in decreased hormone signalling that leads to the AO, a measurable event indicative of or involved in the activation of the negative feedback could be included as a KE.

In cases where a feedback loop may act as a key compensatory or adaptive mechanism that dictates how severely the KEupstream needs to be impacted in order to affect the KEdownstream, but does not play a direct causal role in the AOP (other than defining the relevant point of departure), the feedback should not be included as a separate KE. Rather it should be detailed as part of the quantitative understanding section of the KER description. In the user supplied graphical representation, a forward or backward looping symbol could be added above the arrow linking the two KEs to indicate that a known positive or negative feedback loop is involved in the transition (Figure 10B).

In cases where two measurable KEs in an AOP are part of a positive feedback loop, it can be challenging to define which should be upstream and which downstream, as they are amplifying or altering one another in a cycle. A two headed arrow is undesirable as it can incorrectly suggest that the AOP is reversible. However, in practice an AOP with a positive feedback loop could be accurately represented as two different AOPs in the AOP-Wiki, in which the KEs involved in the positive feedback are presented in either order. This effectively creates a bi-directional arrow when the AOP network is assembled. Rather than creating two nearly identical AOP pages with the KE order reversed for each, the current recommendation is to select either order for the KEs and connect them with a unidirectional arrow, but add a forward looping symbol above the arrow in the user-supplied graphical representation to indicate that a known feedforward loop is involved. (Figure 10A).

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iv. Classification of quantitative understanding

To aid in overall assessment of the AOP and whether it is fit-for-purpose for various applications, developers are also asked to classify the extent of quantitative understanding of the KER as low, moderate, or high, taking into account the extent of data and resulting confidence in empirical support, but also the extent to which quantitative impact of relevant modulating factors is understood. General guidance for classification of the level of quantitative understanding of a KER as low, moderate, or high (Annex 2) is based on several key considerations:

- The accuracy and precision with which a change in KEdownstream can be predicted based on KEupstream.

- 1400 • The precision with which uncertainty in the prediction of KEdownstream can be quantified.
1401 • The extent to which known modulating factors or feedback mechanisms are accounted for.
1402 • The extent to which the relationships described can be reliably generalised across the biological
1403 applicability domain of the KER.
1404

1405 **3J. References**

1406 List of the literature that was cited for this KER description using the appropriate format. Ideally,
1407 the list of references, should conform, with the OECD Style Guide
1408 (<https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-Edition.pdf>) (OECD, 2015).
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SECTION 4 – OVERALL ASSESSMENT OF THE AOP

This section addresses the relevant biological domain of applicability of the AOP as a whole (i.e., in terms of taxa, sex, life stage, etc.) and WoE for the overall AOP. Both are critical for determining the AOP's fit-for-purpose for various applications. This overall assessment is captured on the lower portion of the AOP pages within the AOP-Wiki. **The goal of the overall assessment is not to reproduce or reiterate all the content assembled as part of sections 1-3, but rather to provide a high level synthesis and overview of the relative confidence in the AOP and any significant gaps or weaknesses.** While description and evaluation of modular components facilitate development through sharing, regulatory applications, such as integrated approaches to testing and assessment and stressor specific mode of action, require integrated, pathway-level, analyses. Assimilation and assessment of the extent to which experimental data support expected patterns across all the KERs for the AOP informs relative confidence relevant to consideration of its suitability for specific regulatory applications. For example, the confidence required for prioritizing testing is normally less than that for screening assessment or full assessment to inform risk management.

Determination of confidence in the overall AOP is based on the biological plausibility, empirical support, and extent of quantitative understanding for the KERs (Section 3) and the evidence supporting essentiality of the KEs.

Assessment of the AOP is organised into a number of steps. Guiding questions that inform evaluation at each step are included in Annex 1. The questions are designed to facilitate assignment of categories of high, moderate, or low confidence for each consideration. While it is not necessary to repeat lengthy text that appears elsewhere in the AOP description (or related KE and KER descriptions), a brief explanation or rationale for the selection of high, moderate, or low confidence should be made, based on the guiding questions detailed below.

4A. Define the Biological Domain of Applicability of the AOP

The relevant biological domain(s) of applicability in terms of sex, life-stage, taxa, and other aspects of biological context are defined in this section. Biological domain of applicability is informed by the “Description” and “Biological Domain of Applicability” sections of each KE and KER description (see sections 2G and 3E for details). In essence the taxa/life-stage/sex applicability is defined based on the groups of organisms for which the measurements represented by the KEs are relevant and the structural, functional, and regulatory relationships represented by the KERs are operative.

The relevant biological domain of applicability, including the biologically plausible domain of applicability of the AOP as a whole will nearly always be defined based on the most narrowly restricted of its KEs and KERs. For example, if most of the KEs apply to either sex, but one is relevant to females only, the biological domain of applicability of the AOP as a whole would be limited to females. While much of the detail defining the domain of applicability may be found in the individual KE and KER descriptions, the rationale for defining the relevant biological domain of applicability of the overall AOP should be briefly summarised on the AOP page.

4B. Assess the Essentiality of All KEs

An important aspect of assessing an AOP is evaluating the essentiality of its KEs. This normally entails assessment of the impact of manipulation of a given KE (e.g., experimentally blocking or exacerbating the event) on the downstream sequence of KEs defined for the AOP. Consequently, evidence supporting essentiality is collated on the AOP page, rather than on the independent KE pages that are as stand-alone modular units that do not reference other KEs in the sequence. That said, such evidence can also be captured through the description of adjacent and non-adjacent KERs.

The nature of experimental evidence that is relevant to assessing essentiality relates to the impact on

- 1467 downstream KEs and the AO if upstream KEs are prevented or modified. This includes:
- 1468 ● Direct evidence: directly measured experimental support that blocking or preventing a KE
- 1469 prevents or impacts downstream KEs in the pathway in the expected fashion. Depending on
- 1470 the nature of the KE, could also be evidence that overexpression of the object of the KE
- 1471 prevents or impacts the downstream KEs in a manner consistent with its causal, and
- 1472 essential, role in the pathway.
- 1473 ● Indirect evidence: evidence that modulation or attenuation in the magnitude of impact on a
- 1474 specific KE (increased effect or decreased effect) is associated with corresponding changes
- 1475 (increases or decreases) in the magnitude or frequency of one or more downstream KEs.
- 1476

1477 When evaluating the overall support for essentiality of the KEs, authors may want to summarize

1478 their evaluation of relative levels of support in a tabular format (e.g., Table 5). The objective is to

1479 summarise briefly investigations in which the essentiality of KEs has been experimentally explored

1480 either directly or indirectly. In some cases, the impact of blocking or modifying an early KE on all

1481 downstream KEs in the pathway has been determined; in other cases, the impact only on a single

1482 adjacent or non-adjacent downstream KE has been measured.

1483

1484 When assembling support for essentiality of the KEs, it is not necessary to repeat lengthy text on the

1485 design or results of relevant investigations that may appear in other parts of the AOP description

1486 (e.g., as biological plausibility or empirical support for a KER). Rather, the entries should briefly

1487 address the extent of the supporting and contradictory data through a short description of the nature

1488 of the direct or indirect evidence addressing essentiality, along with relevant references. The

1489 objective is to provide an overview of the extent and nature of supporting and inconsistent data on

1490 essentiality of the KEs in a format that will facilitate a “call” on the overall degree of support for

1491 essentiality across the AOP. Some examples of brief narratives addressing support for essentiality

1492 are included here. The specific nature of these narratives necessarily vary, depending on the nature

1493 of key events in the AOP. See https://aopwiki.org/info_pages/2/info_linked_pages/6 for additional

1494 examples:

1495

- 1496 For direct evidence:
- 1497 ● Knock-out of KE1 or early KEs leads to blockage of all downstream KEs
- 1498 ● Overexpression or underexpression of KE1 leads to effect on all downstream KEs
- 1499 ● One or more downstream KEs is blocked or reversed by inhibiting (or allowing recovery of)
- 1500 upstream KEs
- 1501 ● Overexpression or underexpression in repair enzyme for early KEs leads to decreased or
- 1502 increased incidence of downstream KEs
- 1503 ● Antagonism or agonism of upstream KE leads to expected pattern of effects on downstream
- 1504 KEs
- 1505

- 1506 For indirect evidence:
- 1507 ● Impact of a known modulating factor for early KEs leads to expected pattern of effects on
- 1508 later KEs
- 1509

1510 **Table 5:** Example of a Table Format for summarizing the relative evidence supporting the

1511 Essentiality of KEs in the pathway.

1512

Event	Direct Evidence	Indirect Evidence	No experimental evidence	Contradictory experimental evidence
MIE	****	**		
KE1	*	****		
KE2			****	
KE3.....	**			*
KE _n				

1513

1514

1515 ***Uncertainties or Inconsistencies:***

1516 In addition to outlining the evidence supporting essentiality, it is also important to identify
1517 inconsistencies or uncertainties. This could include, for example, evidence in specific studies that
1518 did not support that blockage or attenuation of an early KE impacted later KEs in the AOP.
1519 Discordance with the results of other studies should be considered based on evaluation of the
1520 adequacy of study design, taking into account, for example, the sensitivity of the detection of impact.
1521 It could also include, for example, gaps in knowledge concerning the essentiality of the MIE or
1522 particular KEs where there are data on essentiality only for one or a few. To the extent possible,
1523 inconsistencies and uncertainties should focus on data gaps important for potential envisaged
1524 regulatory applications as a basis for indicating priorities for further research.

1525

1526 Based on the assembled evidence on essentiality for the KEs, confidence in the supporting data on
1527 essentiality is considered for the entire AOP, including KERs and KEs. This is commonly based on
1528 the extent of direct and/or indirect evidence for one, several or all of the KEs.

1529

1530 Confidence in the supporting data for essentiality of KEs within the AOP is considered:

- 1531 • **High** if there is direct evidence from specifically designed experimental studies illustrating
1532 prevention or corresponding impact on downstream KEs and/or the AO if upstream KEs are
1533 blocked or modified [e.g., via stop exposure/reversibility studies, antagonism, knock out
1534 models, etc.];
- 1535 • **Moderate** if there is indirect evidence that modification of one or more upstream KEs is
1536 associated with a corresponding (increase or decrease) in the magnitude or frequency of
1537 downstream KEs [e.g., augmentation of proliferative response (KE_{upstream}) leading to
1538 increase in tumour formation (KE_{downstream} or AO)];
- 1539 • **Low** if there is no or contradictory experimental evidence that blocking or
1540 modulating/attenuating any of the KEs influences the KEs downstream or AO (Annex 1).

1541

1542 **4C. Evidence Assessment.**

1543 The biological plausibility, empirical support, and quantitative understanding from each KER
1544 in an AOP are assessed together:

1545

1546 ***i. Review the Biological Plausibility of Each KER***

1547 Biological plausibility of each of the KERs in the AOP is the most influential consideration in
1548 assessing WoE or degree of confidence in an overall hypothesised AOP for potential regulatory
1549 application (Meek et al., 2014; 2014a). The defining question for biological plausibility (Annex
1550 1) is: Is there a mechanistic (i.e., structural or functional) relationship between KE_{upstream} and
1551 KE_{downstream} consistent with established biological knowledge? Confidence in the WoE for the
1552 biological plausibility of the KERs would be considered:

- 1553 • **High** if it is well understood based on extensive previous documentation and has an
1554 established mechanistic basis and broad acceptance (canonical knowledge; e.g., increased
1555 follicle stimulating hormone signalling leading to increased estrogen synthesis, increased
1556 incidence of alkylated DNA leading to increased incidence of mutations)
- 1557 • **Moderate** if the KER is plausible based on analogy to accepted biological relationships
1558 but scientific understanding is not completely established
- 1559 • **Low** if there is empirical support for a statistical association between KEs but structural
1560 or functional relationship between them is not understood.

1561

1562 ***ii. Review the Empirical Support for Each KER***

1563 Empirical support entails consideration of experimental data in terms of the associations between
1564 KEs – namely dose-response concordance and temporal relationships between and across multiple
1565 KEs. It is examined most often in studies of dose-response/incidence and temporal relationships
1566 for stressors that impact the pathway at multiple levels of biological organization. These patterns
1567 are most evident when considered across all KERs of the AOP with experimental protocols
1568 optimally designed to address incidence and severity of key events in the AOP at multiple or all

1569 levels of biological organization. While less influential than biological plausibility and essentiality
1570 (Meek et al., 2014; 2014a), empirical support contributes to the assessment of confidence in an
1571 AOP for regulatory application.

1572
1573 It is important to recognise that empirical support relates to the “concordance” of dose response,
1574 temporal and incidence relationships for KERs; the defining question is not whether or not there
1575 is a dose response relationship for a specific KE but rather, whether there is expected concordance
1576 with the dose-response relationships for KERs – i.e., between KEs (Figure 9).

1577
1578 The defining questions for empirical support (Annex 1) are: Does KEupstream occur at lower
1579 doses and earlier time points than KEdownstream; is the incidence or frequency of KEupstream
1580 greater than that for KEdownstream for the same dose of tested stressor? Inconsistencies in
1581 empirical support across taxa, species and stressors that don’t align with the expected pattern for
1582 the hypothesised AOP as described in Section 3 should be identified and their basis considered.

1583
1584 Empirical support for each of the KERs would be considered:

- 1585
- 1586 • **High** if there is dependent change in both events following exposure to a wide range of
1587 specific stressors (extensive evidence for temporal, dose-response and incidence
1588 concordance) and no or few data gaps or conflicting data.
- 1589 • **Moderate** if there is demonstrated dependent change in both events following exposure
1590 to a small number of specific stressors and some evidence inconsistent with the expected
1591 pattern that can be explained by factors such as experimental design, technical
1592 considerations, differences among laboratories, etc.;
- 1593 • **Low** if there are limited or no studies reporting dependent change in both events following
1594 exposure to a specific stressor (i.e., endpoints never measured in the same study or not at
1595 all), and/or lacking evidence of temporal or dose-response concordance, or identification
1596 of significant inconsistencies in empirical support across taxa and species that don’t align
1597 with the expected pattern for the hypothesised AOP.
- 1598

1599 Although developers should evaluate the support for each KER, most critically for the Overall
1600 Assessment of the AOP is to consider the overall level of support across all of the KERs. It may
1601 not be uncommon that the degree of supporting evidence for some KERs in the pathway are quite
1602 limited. However, when there is strong plausibility for the pathway as a whole, and there are well
1603 supported non-adjacent relationships that bridge across some of the weaker intermediate KERs,
1604 the support for the pathway as a whole may still be quite strong. While evidence assembly may
1605 be done in a highly modular fashion, the Overall Assessment of the AOP should once again step
1606 back and evaluate the evidence supporting the pathway as a whole. It is that more integrated and
1607 holistic view that really informs application.

1608
1609 Tables summarising the relevant experimental data for tested stressors across all the KEs may be
1610 helpful in considering the extent of empirical support and to the extent possible should be based
1611 on benchmark doses. For example, benchmark doses (BMDs) for specified similar increases in
1612 each of the KEs are entered in the cells of the table. If the hypothesised linkages in the AOP are
1613 supported by empirical data, there is a pattern of increasing BMDs from the top lefthand corner
1614 to the bottom right hand corner for each of the tested stressors. Presentation in this manner readily
1615 identifies any exceptions to the expected patterns that are considered as inconsistencies and
1616 diminish from the overall weight of empirical support (see Table 6).

1617
1618 **Table 6.** Generic example of a concordance table for evaluating overall empirical support for an

1619 AOP.
1620

Benchmark Dose (mg/kg/d)	KE 1	KE 2	KE 3	KE 5	KE 6	KE 7
0.01	----	----	----	----	----	----
0.05	+++	++	---	++	----	----
0.1		+	+++	+++	----	----
0.5					++	----
1.0					+	++++

1621 a. Benchmark dose at which a specified level of change in the KE relative to controls was inferred, based on the empirical results.
1622 (Note, where concentrations tested are inadequate to determine a BMD, LOEC or NOEC could also be considered, but
1623 concentrations tested in different studies must be taken into account).
1624
1625

1626 **4D. Known Modulating Factors**

1627 The evidence supporting the influence of various modulating factors is assembled within the
1628 individual KERs. As part of the Overall Assessment of the AOP, authors should list the known
1629 modulating factors that have been identified, briefly note their expected influence on the outcome,
1630 and list the specific KER(s) involved. This can be captured in a simple table (e.g., Table 7).
1631 Additional details or notes can be supplied as free text below the table.
1632

1633 **Table 7.** Example of suggested tabular format for identifying critical information concerning known
1634 modulating factors that may be expected to influence the AOP.

Modulating Factor	Influence on Outcome	KER(s) Involved

1635
1636 **4E. Review the Quantitative Understanding of the KERs**

1637 The extent of quantitative understanding of the KERs in an AOP is critical with regard to potential
1638 regulatory application. For some applications (e.g., dose- response analysis in an in-depth risk
1639 assessment), quantitative characterization of downstream KERs may be essential, while for others
1640 quantitative understanding of upstream KERs may be most important (e.g., QSAR modelling for
1641 category formation for testing). Because evidence that contributes to quantitative understanding of
1642 the KER is generally not mutually exclusive with the empirical support for the KER (i.e., expected
1643 patterns of quantitative relationships), evidence that contributes to quantitative understanding will
1644 generally be considered to some extent as part of the evaluation of the WoE supporting the KER (see
1645 Section 3.E. and Annex 1, footnote b). However, specific attention is also given to how precisely
1646 and accurately one can potentially predict an impact on KEdownstream based on some measurement
1647 of KEupstream. This is captured in the form of quantitative understanding calls for each KER, i.e.,
1648 as low, moderate, or high (Annex 2). As noted in section 3, general guidance for characterising the
1649 level of quantitative understanding of a KER is based on several key considerations:

- 1650 ● The extent to which a change in KEdownstream can be precisely predicted based on
1651 KEupstream.
- 1652 ● The precision with which uncertainty in the prediction of KEdownstream can be quantified.
- 1653 ● The extent to which known modulating factors or feedback mechanisms are accounted for.
- 1654 ● The extent to which the relationships described can be reliably generalized across the
1655 applicability domain of the KER.

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1657 As with the other parts of the overall assessment of the AOP, it is not necessary to repeat all the
1658 details provided in the KER descriptions. The overall evaluation of the quantitative understanding
1659 should briefly explain the rationale for the assigned level of quantitative understanding of each
1660 KER. It should then consider the overall pattern of quantitative understanding across all KERs to
1661 indicate how precisely outcomes along the entire pathway may be predicted for a given exposure
1662 scenario. If certain parts of the pathway can be predicted with quantitative precision, while others

1663 cannot, the potential implications for application may be discussed.

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1666 **4F. Considerations for Potential Applications of the AOP (optional)**

1667 The Overall Assessment of the AOP is intended to help inform decisions about an AOP's fit-for-
1668 purpose for different types of applications. Consequently, at their discretion, following their
1669 assessment of the AOP, the developers may want to discuss the type(s) of application(s) they feel
1670 the AOP would be suited for, based on their evaluation. This may include, for example, possible
1671 utility for test guideline development or refinement, development of integrated testing and
1672 assessment approaches, development of (Q)SARs / or chemical profilers to facilitate the grouping
1673 of chemicals for subsequent read-across, screening-level hazard assessments or even risk
1674 assessment. This section can consider whether the AOP assembled can support the intended
1675 application that was outlined previously in the "AOP Development Strategy" section. It may also be
1676 that new potential applications or limitations which become apparent when developing the AOP and
1677 assessing the evidence could also be noted in this section.

1678 It is further recognized, that developers may not be aware of all the potential applications for any
1679 given AOP. Consequently, users of the AOP-Wiki are encouraged to leave comments on the
1680 discussion pages, or via the [AOP Forum](#) if they identify suitable applications for a given AOP.
1681 Listing these applications can aid others in using the AOP.

1682

1683 **4G. References**

1684 References cited elsewhere on the AOP page should be listed here. This is not a compilation of all
1685 references cited on the linked KE and KER pages. Ideally, the list of references, should conform
1686 with the OECD Style Guide ([https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-
1687 Edition.pdf](https://www.oecd.org/about/publishing/OECD-Style-Guide-Third-Edition.pdf)) (OECD, 2015).

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ANNEX 1: Guidance for Assessing Relative Level of Confidence in the Overall AOP

Examples of complete tables for selected AOPs are available:

AOP	Assessment Summary File
https://aopwiki.org/aops/15	https://aopwiki.org/system/dragonfly/production/2017/05/19/7s1ibrunwt_RevisedAssessmentSummaryAop_15.pdf
https://aopwiki.org/aops/23	https://aopwiki.org/system/dragonfly/production/2017/03/20/3usvv7naq8_Annex1_for_AOP_23_AR_reproductive_dys_2017_03_20.pdf
https://aopwiki.org/aops/38	https://aopwiki.org/aops/38#evidence
https://aopwiki.org/aops/42	https://aopwiki.org/system/dragonfly/production/2017/03/24/6u60jhkjp8_TPO_AOP_Summary_Tables.pdf

1. Support for Biological Plausibility of KERs ¹	Defining Question	High ^{2,3}	Moderate	Low
	Is there a mechanistic (i.e., structural or functional) relationship between KEup and KEdown consistent with established biological knowledge?	Extensive understanding based on extensive previous documentation and broad acceptance -Established mechanistic basis	The KER is plausible based on analogy to accepted biological relationships but scientific understanding is not completely established.	There is empirical support for a statistical association between KEs (See 3.), but the structural or functional relationship between them is not understood.
⁴ MIE => KE1: (copy and paste the KER description into this cell)	Biological Plausibility of the MIE => KE1 is xxx. Rationale:			
KE1 => KE2: (copy and paste the KER description into this cell)	Biological Plausibility of KE1 => KE2 is xxx Rationale:			
KE2 => KE3 (copy and paste the KER description into this cell)	Biological Plausibility of KE2 => KE3 is xxx. Rationale:			

¹Rank ordered Bradford Hill considerations adapted from Meek et al. (2014b)

²The guidance for “high”, “moderate” and “low” draws on limited current experience. Additional delineation of the nature of relevant evidence in these broadly defined categories requires more experience with larger numbers of documented AOPs.

³“Direct evidence” implies specifically designed experiments to consider the relevant element. “Indirect evidence” may overlap with other elements.

⁴To the extent possible, each of the relevant Bradford Hill considerations is addressed for each of the KERs (biological plausibility and empirical support) and KEs (essentiality) and separate rationales provided.

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2. Support for Essentiality of KEs ⁵	Defining Question	High	Moderate	Low
	What is the impact on downstream KEs and/or the AO if an upstream KE is modified or prevented?	Direct evidence from specifically designed experimental studies illustrating prevention or impact on downstream KEs and/or the AO if upstream KEs are blocked or modified	Indirect evidence that modification of one or more upstream KEs is associated with a corresponding (increase or decrease) in the magnitude or frequency of downstream KEs	No or contradictory experimental evidence of the essentiality of any of the KEs.
AOP	Rationale for Essentiality of KEs in the AOP is xxx:			

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⁵While the extent of the *supporting* data on the essentiality of each of the KEs is addressed separately (Table 3), delineation of the degree of confidence is based on consideration of evidence for all of the KEs within the AOP and therefore, only one rationale is required. This call is normally based on the extent of the available evidence for a range of KEs in the AOP.

3. Empirical Support for KERs				
Defining Questions	High	Moderate	Low	
Does KEup occur at lower doses and earlier time points than KE down and at the same dose of stressor, is the incidence of KEup > than that for KEdown? ^{6,7} . Are there inconsistencies in empirical support across taxa, species and stressors that don't align with expected pattern for hypothesised AOP?	Multiple studies showing dependent change in both events following exposure to a wide range of specific stressors. (Extensive evidence for temporal, dose-response and incidence concordance) and no or few critical data gaps or conflicting data	Demonstrated dependent change in both events following exposure to a small number of specific stressors and some evidence inconsistent with expected pattern that can be explained by factors such as experimental design, technical considerations, differences among laboratories, etc.	Limited or no studies reporting dependent change in both events following exposure to a specific stressor (i.e., endpoints never measured in the same study or not at all); and/or significant inconsistencies in empirical support across taxa and species that don't align with expected pattern for hypothesised AOP	
MIE => KE1: (copy and paste the KER description into this cell) ^b	Empirical Support of the MIE => KE1 is xxx. Rationale:			
KE1 => KE2: (copy and paste the KER description into this cell)	Empirical Support of the KE1 => KE2 is xxx. Rationale:			
KE2 => KE3 (copy and paste the KER description into this cell)	Empirical Support of the KE2 => KE3 is xxx. Rationale:			
<p>^b In many cases, evidence that contributes to quantitative understanding (Section 4 of a KER description) will also provide empirical support for the relationship. Consequently, relevant information from the "Quantitative Understanding" section of the KER description should be considered as part of the overall weight of evidence evaluation of the concordance of empirical observations and consistency for the KER.</p>				

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⁶This is normally considered on the basis of tabular presentation of available data on temporal and dose-response aspects, in a template that documents the extent of support. See, for example, Table 4.

⁷Note that this relates to concordance of dose response, temporal and incidence relationships for KERs rather than the KEs; the defining question is not whether or not there is a dose response relationship for the KE but whether there is concordance with that for earlier and later KEs. This is normally demonstrated in studies with different types of stressors.

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ANNEX 2: General guidance for characterizing the level of quantitative understanding of a KER as low, moderate, or high.

Extent of Quantitative Understanding	Characteristics
High	<p>Change in KEdownstream can be precisely predicted based on a relevant measure of KEupstream.</p> <p>Uncertainty in the quantitative prediction can be precisely estimated from the variability in the relevant measure of KEupstream.</p> <p>Known modulating factors and feedback/feedforward mechanisms are accounted for in the quantitative description.</p> <p>There is evidence that the quantitative relationship between the KEs generalizes across the relevant applicability domain of the KER.</p>
Moderate	<p>Change in KEdownstream can be precisely predicted based on a relevant measure of KEupstream.</p> <p>Uncertainty in the quantitative prediction is influenced by factors other than the variability in the relevant measure of KEupstream.</p> <p>Quantitative description does not account for all known modulating factors and/or known feedback/feedforward mechanisms.</p> <p>The quantitative relationship has only been demonstrated for a subset of the overall applicability domain of the KER (e.g., based on a single species).</p>
Low	<p>Only a qualitative or semi-quantitative prediction of the change in KEdownstream can be determined from a measure of KEupstream.</p> <p>Known modulating factors and/or known feedback/feedforward mechanisms are not accounted for.</p> <p>The quantitative relationship has only been demonstrated for a narrow subset of the overall applicability domain of the KER (e.g., based on a single species).</p>

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