

30 April 2013

Advice to the European Medicines Agency from the Clinical trial Advisory Group on Rules of engagement (CTAG3)

Draft Advice Version 7.0 – With amendments following comments on 6.0

Advice to the European Medicines Agency on rules of engagement for accessing clinical trial data

Draft – ~~0522~~ March 2013 - Version ~~6~~7.0

Preliminary comment: This advisory group discussed the issues and questions listed below and offers the following views and positions for EMA's consideration:

1. Should the marketing authorisation holder be consulted before EMA discloses clinical trial data, in regards of commercial confidential information (CCI)? What elements of the clinical part of the dossier could be considered CCI?

No agreement was reached. The following positions were discussed:

a. EMA should only disclose confidential commercial information from non-clinical and clinical study reports and patient level data when there is an overriding public interest reason for doing so, under conditions which serve that interest. The EMA should always consult with the marketing authorisation holder (MAH) prior to disclosure, to allow the MAH to take any necessary steps to protect against unfair competition and/ or prejudice to regulatory data protection, patent or other IP rights.

Although the situations would be rare (perhaps when working with a new therapeutic class or a rare disease) it is ~~not im~~possible that eCTDs and CSRs would contain competitively valuable information. The sorts of information (with historical examples that are no longer competitively ~~sensitive~~relevant) are:

- Methods to pursue newly validated / devised endpoints that are persuasive to regulators:

e.g., the suite of validated measurements for assessing the effects of migraine on the whole body in support of the first approval of the prototypical 5HT1B/1D agonist sumatriptan p.o. and s.c.

- Identification of investigators that recruit well, especially for rare diseases / difficult patient populations:

e.g., those with sufficient patients to support a clinical trial in cluster headache as a new indication for s.c. sumatriptan

- A novel trial design, streamlining and making more economical the proof of efficacy for an acutely acting compound: e.g., Armitage (adaptive) design that was novel and supported the approval of i.v. dantrolene

- CSRs may contain information on bio-analytical product-characterization methods which are the intellectual property of the MAH - public disclosure could be an infringement of the MAH's IP rights. Furthermore, the use of some specific analytical tests described in the CSR can provide information indicative of the active product substance/molecule that can therefore be identified and used by competitor companies (e.g. tests on molecule-specific epitopes providing information allowing identification of the commercial confidential molecule).

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36 [Commercial sensitivity resides in the effect of EMA's intent to release clinical trial data on products](#)
37 [that rely on data protection laws to prevent generic competition in other territories. In other words,](#)
38 [of particular concern with the proposed proactive broad disclosure of clinical trial data is the](#)
39 [potential for inappropriate use of such data by third parties either to circumvent existing regulatory](#)
40 [data protection \(RDP\) rules, or take advantage of the absence of such rules in the many countries](#)
41 [which do not have robust systems of RDP equivalent to that in the EU. For instance, data](#)
42 [exclusivity in Australia, China and Mexico is directly undermined by publication of the relevant](#)
43 [data, anywhere in the world.](#)

44 [Industry contends that if data are obtained from EMA under its disclosure policy and used lawfully](#)
45 [in a third country then the EU MAH would have no legal redress.](#)

46 [However, even if a CCI was defined \(additional concrete cases must be provided\), open access](#)
47 [should be restricted ONLY for this sensitive part of the CSR. Moreover, EMA consultations to MAH](#)
48 [should not imply long delays in releasing data.](#)

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50 [b. EMA's consultation with the marketing authorisation holder \(MAH\) prior to disclosure may](#)
51 [introduce delays that detract from the concept of "proactive" disclosure. Whether or not a](#)
52 [particular material can be disclosed, and under what terms, should be decided prior to readying](#)
53 [materials for disclosure.](#)

54 [With regards to the examples of CCI listed above: Some of the examples should nowadays not be](#)
55 [legitimate examples of commercial sensitivity. At the time these drugs were being developed, they](#)
56 [may have been thought to be legitimate examples simply because of the way drug development](#)
57 [was done then. Today, these examples should be regarded as being examples that overall make](#)
58 [clinical development more efficient and as such should be shared. Furthermore, if the new method,](#)
59 [endpoint... is an argument for the approval, it should be made publicly available in the EPAR and](#)
60 [properly described in any guideline applying to the evaluation of products in the indication.](#)

61 [It is emphasised that "competitively valuable information" is not necessarily CCI. For example, a](#)
62 [negative study result is obviously competitively valuable information, but this should not make it](#)
63 [CCI.](#)

64 [Study methods and study results are never CCI. The information is essential for the interpretation](#)
65 [of the study results and should be available for the public. EMA's policy will ensure that this will be](#)
66 [done only after a decision about marketing authorisation has been made.](#)

67 [Third-party requestors may need some of this "competitively sensitive" information to carry out](#)
68 [proper re-analysis and verification of results, such as trial protocols, but may not necessarily need](#)
69 [all of them \(e.g. identification of investigators that recruit well\). Most of the information on 'good](#)
70 [investigators' in CTD and CRS will also be available in publications.](#)

71 [Identity of investigators should always be public in order to make clear any conflicts of interest](#)
72 [between MAH and professionals.](#)

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74 [Note from EMA: stakeholders are invited to present additional concrete \(historic?\) examples and](#)
75 [case scenarios how confidential commercial information from CSRs could be used for unfair](#)
76 [competition and/ or prejudice to regulatory data protection, patent or other IP rights and what](#)
77 ['necessary steps' might be required. \(See also comment under section3\) –stakeholders are invited](#)

78 to specifically comment on the question: What elements of the clinical part of the dossier could be
79 considered CCI?

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81 The questions listed below addressed the issue: ~~What-what~~ steps will a requester have to go
82 through before being able to access clinical trial data from the EMA website? After accessing the
83 dedicated domain of the EMA website:

84 **42. Should requesters have to identify themselves?**

85 It is useful to distinguish between access to (1) aggregate data (e.g. lists of studies conducted, ICH
86 compliant clinical study reports including the study protocol, statistical analysis plan and other
87 appendices, but excluding patient level data) and (2) patient-level data (e.g. individual case record
88 forms, SAS files with line listings).

89 1. Aggregate data: No agreement was reached. The following positions were discussed:

90 a. There is no convincing rationale that identification of requesters could or should be
91 required. Such data should be accessible freely (similar to EPAR information today).
92 It is assumed that aggregate data contains no or few personal data (any personally
93 identifiable information must be removed prior to release unless justified to
94 remain). It is pointed out that the aim of transparency shouldn't be only to allow a
95 potential reanalysis. For example, drug independent bulletins need full information
96 of clinical trials not for research purposes but for education purposes in health
97 areas. A watchdog activity is high useful to citizens and also for drug regulatory
98 bodies. So in many cases there won't be a "legitimate scientific question" to be
99 considered. Transparency goes beyond reanalysis purposes.

100 b. In the interest of transparency, requesters should be identified, logged and their
101 identity made public, primarily to ensure patient confidentiality is not compromised
102 and to avoid the misuse of patient level data by third parties with commercial
103 interests that are not related to healthcare research. It is technically possible to
104 accurately identify requestors; one could perhaps use an ORCID ID to identify
105 requestors. Requestors of clinical trial data should also have sufficient qualifications
106 and experience for any subsequent analysis of data obtained from clinical trials, as
107 aligned with ICH-E9 and 'statistical principles for clinical trials'. Also, in order for
108 any analysis of data obtained from clinical trials, there should be a legitimate
109 scientific question being proposed in order for the request for data access to be
110 considered. Requesters should not only identify themselves, but they should also
111 provide details of their qualifications and experience which supports they are
112 sufficiently educated and trained to implement any subsequent analysis of the data
113 being requested. This information should be made transparent by the requester at
114 the time of seeking access to data.

115 NOTE from EMA: such proposals may not be compatible with the legal framework under
116 which EMA operates as a public body; to be discussed at upcoming CTAG3 meeting

117 2. Patient-level data: No agreement was reached. The following positions were discussed:

118 a. These data should be freely accessible without the need for identification.
119 Arguments in favour of this position include (not in order of importance):

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- i. Lowering the hurdle for patients who wish to access data related to their own disease. [Asking requesters to publicly share their personal details, education and training before getting access would violate data protection regulations and induce a hurdle for non-professional user groups. Also, the rules of engagement should not include any pre-selection or pre-identification and publication of the requester name for a simple reason: a patient can ask for the data about a product he has to take for his/her disease. If specific qualifications are requested, one will easily know who are the requesters with a personal interest in the product \(those without clear qualifications\).](#)+
 - ii. Proper verification of identity of the requester is near-impossible;
 - iii. If the data are used for illegal actions such as illegitimate commercial use, there are legal actions which can be taken against the firm/country benefiting from the illegal action. Thus, this point should not be an argument to force requester-identification. Furthermore, if someone wishes the data for illegal action, he will surely and easily use a wrong identification or could only ask others to also request data in order to increase the number of suspects;
 - iv. Any patient-level data that EMA makes available will be de-identified/anonymised, therefore the risk of retro-active patient identification is considered acceptably low, and the patient data protection is not an issue (it is argued that there is even no need to distinguish between aggregate data and patient level data). Therefore, there is no need to verify the identity of the requester (*Note: reference is made to CTAG1, which is discussing standards for de-identification/anonymisation to ensure patient data protection*);
 - v. [There are cases of harassment by pharmaceutical industry when a physician declared an adverse event to an agency \(example: Dr Chiche in Marseilles about the Mediator story\). If the name of the requesters is given to EMA, how will EMA make sure that the name of the requester will not be known by the Marketing Authorisation Holder? In case of harassment linked to a data request, what would be EMA's responsibility?](#)
 - ~~vi.~~ [Any suggestion that requestors of clinical trial data should also have sufficient qualifications and experience for any subsequent analysis of data is neither practical nor desirable for either aggregate data or patient-level data. It would entail subjective and arbitrary judgements about what qualifications and experience are "sufficient".](#)
 - ~~vi.~~vii. [The privacy of study participants is important and their privacy should be warranted. On the other hand, the privacy should also be warranted for study participants, patients or other \(EU\) citizens who like to access patient-level data for their own private use. Namely, publication of their name on the internet involves the risk of unintended use of the personal data of this person, especially if this information can be detected by search engines such as Google. For example, the information \(name + type of medication\) may be detected during a background search performed for a](#)

165 job application; the information can be used by insurance companies; or
166 the information can be used for direct marketing for registered or falsified
167 medicines, including spamming. This is an argument to carefully consider
168 whether the benefits of publication of the names of private persons
169 outweigh the risks of unintended use and breach of privacy of those who
170 access data. Thus, benefits of publication of the names of those who access
171 patient level data may not outweigh the risks, because publication of
172 personal data in combination with (type of) medicines for which data have
173 been accessed creates the possibility for unintended and undesirable use of
174 personal data;

175 [viii.](#) As data would be anonymous there is no sensitive data. Retrospective
176 patient identification cannot be prevented by verifying the identity of the
177 requester, nor can any violator necessarily be identified through such
178 knowledge as there will usually be no conclusive link between the violation
179 and the requester. We should keep in mind article 6.1. b and c. in directive
180 95/46/EC of the European Parliament and of the Council of 24 October
181 1995 on the protection of individuals with regard to the processing of
182 personal data and on the free movement of such data. Pursuant to this
183 article collection of data must be adequate, relevant and not excessive in
184 relation to the purposes. Registering the requester is also processing of
185 personal data and should only be done for legitimate reasons and should
186 not be excessive in relation to the purpose.

187 [vii-ix.](#) [Concerns about inappropriate analyses are misplaced, since the scientific](#)
188 [community will or will not give their support to these analysis based on its](#)
189 [scientific value.](#)

190 b. These data should be freely accessible only after verification of the identity of the
191 requester. Arguments in favour of this position include (not in order of
192 importance):

193 i. Patient-level data is too sensitive to allow anonymous requesters to access
194 because the risk of retrospective patient identification is never zero. The
195 legal liability associated with the release of the patient data from a data
196 privacy perspective needs to be considered. There is reference to the risk of
197 retro-active patient identification being “acceptably low”, yet that still
198 presents a risk to patient identification. Legal accountability needs to be
199 addressed if a patient is in fact identified and this is used improperly
200 against an individual patient;

201 ii. The level of de-identification required to render patient-level data suitable
202 for open public access is likely to seriously compromise the utility of that
203 data for the purpose of research in the interest of public health. Much of
204 the value of analysis of patient-level data over aggregate data is the ability
205 to link and take account of patient characteristics in analyses. For example,
206 if age and gender were to be removed from the dataset, it would not be
207 possible to investigate possible treatment interactions with these
208 characteristics or with these in combination with other characteristics that
209 remain in the dataset. If dates are removed this reduces scope for scrutiny
210 and (unless replaced with a series of derived times from event to event)

211 precludes time to event analyses. This would mean, for example, that
212 survival analyses in cancer trials would not be possible. This is an important
213 consideration for individual participant data systematic (IPD) reviews and
214 meta-analyses. Re-consider whether tiered access is feasible. Open public
215 access for all documentation including clinical study reports, results, and
216 aggregate data. Access to IPD restricted to being for the purpose of
217 research in the interest of public health - as demonstrated by provision of a
218 protocol or research plan, disclosure of investigator name and affiliation
219 and declaration of any potential conflict of interest (preferably at the point
220 of release of data, but delayed if necessary);

221 iii. Strict assurances about the specific use of personal data are given as part
222 of the consent process to trial entry; they do not include release except
223 under strict rules. Release of individual patient data, even anonymised,
224 contravenes the information provided as part of the consent process, and
225 thereby infringes human rights.

226 iv. It is possible (and will be even easier in the future) to combine anonymised
227 data sets with other data that is readily available publically to identify
228 individuals. This is important for privacy particularly as the data contains
229 health information that can be sensitive and assumed to be private by the
230 clinical trial participant. For example please see :
231 [http://online.wsj.com/article/SB1000142412788732378370457824784249](http://online.wsj.com/article/SB10001424127887323783704578247842499724794.html)
232 [9724794.html](http://online.wsj.com/article/SB10001424127887323783704578247842499724794.html) and the original article 'Identifying Personal Genomes by
233 Surname Inference. Melissa Gymrek et al. Science 339:321, 2013'.

234 v. Requesters of patient-level clinical trial data should also have sufficient
235 qualifications and experience for any subsequent analysis of data obtained
236 from clinical trials, as aligned with ICH-E9 and 'statistical principles for
237 clinical trials'. Also, in order for any analysis of data obtained from clinical
238 trials, there should be a legitimate scientific question being proposed in
239 order for the request for data access to be considered. Requesters should
240 not only identify themselves, but they should also provide details of their
241 qualifications and experience which supports they are sufficiently educated
242 and trained to implement any subsequent analysis of the data being
243 requested. This information should be made transparent by the requester
244 at the time of seeking access to data.

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246 ~~iii-vi.~~ There is a risk of illegitimate commercial use of patient-level data (please
247 refer to **point 3**). To mitigate this risk the identity of the requester must be
248 verified;

249 ~~iv-vii.~~ The identity of the requester should be available and public. It is widely
250 accepted in science that people have to disclose their financial interest. This
251 principle should be applied here as well;

252 viii. The objective is clearly to restore trust in the system, not to create an all-
253 purpose research tool. Patient data is not to be diverted to research
254 purposes for which it was never intended or to "data mining", be it
255 academic or commercial. Such misuse could otherwise lead to false claims

256 of efficacy and safety of medicines. The EMA has previously stated the
257 objective is to "(...) enable the independent re-analysis of the evidence
258 used by the Agency's committees to determine their benefits and risks and
259 is expected to lead to public-health benefits." The access process should be
260 developed with this public health principle in mind;

261 ix. It is not clear how providing patients access to data relating to their own
262 disease is aligned with the remit of access to data which is being able to
263 independently re-analyse the benefit-risks. Anyone wishing to re-analyse
264 data should have minimal qualifications and expertise and it should not be
265 suggested that individuals who are not equipped with the relevant skills
266 should attempt to re-analyse data.

267 ~~v-x.~~ It should be recognised that clinical trial participants are providing sensitive
268 health information while those who are accessing anonymised data would
269 not be required to provide sensitive health information. For example they
270 would only be required to provide their name, address and research
271 institution. It is also difficult to understand why the name of a
272 researcher/requester who accessed data for a particular disease would
273 result in insurance or any other consequence. Merely accessing the data
274 does not indicate or suggest that the individual has that disease or
275 condition. In addition if an email address is not made public (and there is
276 no reason to do so) there is little or no risk of spamming.

277 xi. There is also a risk of other unintended consequences: Some requesters
278 may present out-of-context results that would lead to false impressions of
279 drug safety issues and lead to unfounded health scares (e.g.
280 <http://www.biomedcentral.com/1471-2458/2/6>). This risk is of high
281 importance to the ultimate decision of whether patient level data should
282 have open access and the long term consequences should be discussed.
283 However, sometimes it's in fact the opposite. Some requesters use data
284 from drug regulatory agencies to minimize unfounded health scares with
285 potential harms in other senses: for example, the PPI-Clopidogrel
286 interaction case:
287 <http://www.nature.com/ajg/journal/v106/n7/full/ajg2011126a.html>

288 xii. -If a requestor uses data for an illegitimate use, is the EMA liable for failing
289 to protect patient confidentiality? There is no secure path forward when
290 granting control to anyone to secure patient confidentiality. Industry can do
291 certain measures to ensure that data confidentiality is given within a
292 dataset. But there is no measure available to secure this when a requester
293 has access to the clinical trial data for the purpose to re-analyse it, as they
294 would then have the potential to merge the clinical trial data with other
295 available data. The only way to secure patient confidentiality is to have a
296 step that checks the request for access is scientific (good intent) and clear
297 rules noting that data cannot be further disseminated. If the rules require
298 the uploading of a protocol or analysis plan then this using a restrictive
299 access approach increases the protection against unintended use of the
300 data. The policy will need to clarify who is liable for any illegitimate use of
301 data.

302 [vi-xiii. Although the identity of the requester indeed should be known to the](#)
303 [database owner, it is not conclusive to request publication of these names](#)
304 [and addresses.](#)

305 c. [Several types of compromises could be envisaged:](#) For access, a hierarchy for
306 different user groups should be foreseen with access to different types of data. For
307 the EMA pharmacovigilance database, such an access policy already exists.
308 (EMA/759287/2009 corr., EudraVigilance access policy for medicines for human
309 use) This paper is adopted after consultation with the Patients' and Consumers'
310 Working Party and consultation with the Health Care Professional Working Group.
311 The paper defines 4 types of stakeholder groups:

- 312 • Medicines Regulatory Authorities, the European Commission and the
313 Agency (hereafter referred to as Stakeholder Group I)
- 314 • Healthcare Professionals and the General Public (hereafter referred to as
315 Stakeholder Group II)
- 316 • Marketing Authorisation Holders and Sponsors of Clinical Trials
317 (hereafter referred to as Stakeholder Group III)
- 318 • Research Organisations (hereafter referred to as Stakeholder Group IV)

319 There is a need to modify the categories according to an optional user identification
320 process, granting access to e.g. patient level after authorisation. [If hierarchy for](#)
321 [different user groups were finally considered, healthcare professionals should have](#)
322 [access to the higher possible level of information.](#) This would also allow for the
323 processes discussed under topics 3, 4 and 6, setting reminders or making
324 registered users aware of possible consequences after misuse.

325 [Those specific trials should be identified where retroactive patient identification is a](#)
326 [risk, and alternatives should be provided for these cases to harmonize patient and](#)
327 [health professional rights. For example, access to data on clinical studies conducted](#)
328 [in patients with rare diseases should be restricted and treated under different](#)
329 [provisions, such as mandatory registration and identity verification of the](#)
330 [requestor, and contractual agreements covering the consequences of misuse and/or](#)
331 [inadvertent identification.](#)

332 [Alternatively, open access could be granted for aggregate anonymised data and](#)
333 [restricted access for patient level data where access is controlled by EMA.](#)

334 [Consider differentiating between requests for data to "independently re-analyse](#)
335 [trial data" and requests for data to be used in "secondary analysis to address new](#)
336 [clinical questions" and how this could determine the level of data access required.](#)
337 [The complexity of taking patient level data and all the associated meta-data should](#)
338 [be noted, and this complexity could lead to incorrect analyses being generated](#)
339 [unless appropriate checks are put in place to deal with such situations.](#)

340 [Note whether it would be feasible for the EMA themselves to re-analyse patient-](#)
341 [level trial data to address the "independent re-analysis" of trial data. If this](#)
342 [approach was possible, this could lead to granting open access to aggregate](#)
343 [anonymised data, and EMA and other nominated stakeholders considered](#)
344 ["independent" to access to patient level data.](#)

345 [It is also noted that in order to allow for public access to patient-level data in the future, they](#)
346 [would have to be a mandatory part of the clinical submission documents, and reflected in the](#)
347 [relevant CHMP guideline documents such as CHMP/EWP/2998/03. Furthermore, the potential use of](#)
348 [patient-level data outside of the clinical study scope should be covered in the study informed](#)
349 [consent form such that the subject agrees to the future "secondary use" of patient-level data](#)
350 [outside of the study scope.](#)

351 **23. Should requesters be required to 'Agree' to respect personal data protection?**

352 It is agreed that this point is only relevant for patient-level data.

353 It is agreed that any requirement for the requester to actively agree to respect personal data
354 protection would depend on whether the identity of the requester can be/has been verified. (No
355 agreement was reached on that point, see above)

356 If the identity of the requester has not been verified (two positions):

357 a) Without requester identification, such `agreement` to respect personal data protection is
358 only for information, but cannot be legally binding. As far as CTAG1 rules for patient data
359 anonymisation are applied and effective, respect of personal data protection mainly forbids
360 linking the data obtained from EMA with other databases/information.

361 b) Even if the identity of a requester cannot be verified, a disclaimer about the need for
362 personal data protection should be "read and accepted" by the requester.

363 If the identity of the requester has been verified:

364 Should it be/have been possible to verify the identity of the requester, and the requester actively
365 agrees to respect personal data protection, any violation of this agreement should be legally
366 enforceable.

367 Requesters have to be made aware of EU and local data protection regulations. Ticking a box
368 implies a contractual relationship between the requester and the database owner/holder of the
369 data. However, in that case both contractual parties need to be fully identifiable. A contractual but
370 not necessarily public "digital" agreement appears to be preferable compared to a purely
371 anonymous process.

372 Details of a contractual agreement should clarify that if any individuals are provided access to
373 clinical trial data, then the holders of the data cannot be held accountable in any way for what the
374 requesters subsequently do with the data; any re-analysis of the data is at the responsibility of the
375 requester. If subsequent issues are found with respect to an incorrect re-analysis, misuse of the
376 data for purposes outside of the research proposal originally specified, or any potential fraudulent
377 behaviour, the original owner of the source data cannot be held accountable in any way.

378 **34. Should the requester be required to 'Agree' to refrain from unintended**
379 **commercial uses of information retrieved?**

380 There is general agreement that EMA's policy on Access to clinical trial data should further the
381 interest of public health, but should not abet usage of data for unintended commercial uses
382 (~~e.g. such as~~ obtaining a marketing authorisation in a third, non-EU, jurisdiction). EMA's policy
383 should attempt to mitigate this risk without compromising transparency. The option of requiring
384 anonymous data requesters to tick a 'read and accepted' tick box is considered ineffectual.

385 No agreement was reached on the following point (two positions):

- 386 a) The requester should be required to sign a legally binding agreement affirming that the
387 information and data will only be used for the agreed public health research purpose and
388 not for any commercial use. Requests for patient level data from requesters to the EMA
389 must be handled on a case-by-case basis, and follow consistent criteria to establish if and
390 how the information provided will be used for valid scientific purposes and to benefit
391 patients. [\(Please refer to discussion of CCI under Question 1\)](#)
- 392 b) It is unclear which situations we are talking about and "unintended commercial uses" may
393 be used as a "killer argument". For example, if industry fears that one cannot exclude that
394 a full CSR may be used for obtaining a marketing authorisation in a non-EU jurisdiction,
395 this may prevent full transparency. [The relationship between knowledge and profit-making
396 is too complex to have it be contractually bound during the data release process; there is
397 no simple distinction between using data for public health research and commercial use.
398 The party suggesting a legally binding contract requiring the requestor to guarantee to use
399 the data for public health purposes and not commercial purposes, should be clarified as to
400 how commercial purposes and public health purposes will be defined and disentangled in
401 practice. Some real-life examples of "unintended commercial uses" should be given during
402 the next CTAG3 session.](#)

403 **45. Should the requester be made aware of quality standards for additional /**
404 **secondary analyses?**

405 No agreement was reached on this point (two positions):

- 406 a) It is emphasised that advising requesters of quality standards for additional secondary analyses
407 should not and cannot impose any obligations on the requester. [However, it would be
408 appropriate to ask EMA to communicate their quality standards when a public statement is
409 issued.](#) (Note: Reference is made to the work of CTAG4).

410 The use of such advice is questioned. This may discourage non-professional users from
411 downloading and using such data. There is no benefit from such advice but it may mean a
412 subjective additional hurdle to lay groups/patients.

- 413 b) The requester should be advised of quality standards for additional secondary analyses.
414 The same standards must be applied equally to the requester as would be applied to the
415 MAH. It is emphasised that such advice should imply clear obligations on the requester.

416 **56. Should the requester have to declare whether they wish to upload a protocol /**
417 **analysis plan?**

418 There is agreement that good scientific practise requires those who wish to engage in secondary
419 data analysis to complete and submit a study protocol before accessing the data. Therefore, the
420 opportunity (but not obligation) to upload a protocol on an EMA managed repository is welcomed.
421 There was no consensus as to the time of publication of such uploaded protocols. Options discussed
422 were:

- 423 a) Immediately after uploading the protocol
- 424 b) After a fixed time span (e.g. 1 month, 1 year?)
- 425 c) Around the time of publication of the results of secondary analysis

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426 d) Timing of publication decided by requester

427 Several comments/views along the following lines were expressed:

428 A requester should have to submit a protocol or analysis plan before being granted access to the
429 data as this enables full transparency of the purpose and intention for requesting access to the
430 data and this helps to minimise any misuse by third parties. In order to ensure there is a legitimate
431 research question(s) being proposed, pre-specifying the clinical hypotheses to be investigated
432 ensures the scientific credibility of the research to be undertaken.

433 [The process to be followed could be tailored to the remit for the request for access to data -
434 independent re-analysis versus secondary analyses of existing data.](#)

435 [A protocol could be either uploaded or provided as link to a " trial register". An \(ethics committee\)
436 review of the protocol should be provided by the requester.](#)

437 Provision of a protocol demonstrating good research methods, fair use of data and the purpose to
438 which it will be put seems an entirely reasonable exchange for access to data. There seems to be a
439 danger of introducing double standards with requirement for access to clinical trial protocols and
440 clinical trial data, but not to protocols for subsequent use. For IPD, make provision of a protocol
441 (with delayed public access if necessary) a prerequisite for access to or release of data. A link to a
442 formally published protocol would be acceptable. [Protocols should be given a unique identifier,
443 which is also quoted in each publication that arises from the analyses.](#)

444 ~~Therefore t~~he protocol must be reviewed before the patient level data is provided.

445 ~~NOTE from EMA: such proposals may not be compatible with the legal framework under which EMA
446 operates as a public body; to be discussed at upcoming CTAG3 meeting~~

447 **67. Should requesters be allowed to share accessed data?**

448 It was agreed that this would become uncontrollable in case identification of the requester is not
449 verifiable.

450 No agreement was reached on the following point of sharing data (two positions):

451 a) Should it be/have been possible to verify the identity of the requester, EMA may consider
452 restricting data sharing. However, in such case any third party would have to be given
453 access to the same data as the first requester directly from the EMA. [If a collaboration
454 between 2 requesters is necessary \(e.g. Academia + industry or data management
455 company\), EMA should be informed and give approval. This can be anticipated in the
456 analysis plan.](#)

457 b) Requesters should not be allowed to share accessed data because that way the validity of
458 the dataset cannot be controlled. [Requestors will be responsible for the security of the data
459 they gain access to. Without this accountability, the sharing of data could quickly become
460 widespread; this can be avoided if requesters have restricted access to data sets in a
461 controlled system.](#) Requesters should need to explicitly confirm that they will not forward
462 the downloaded original dataset to third parties. It is acknowledged that others must be
463 able to repeat research findings; that is a basic principle of research. However, such groups
464 would then have to identify themselves separately before accessing the same data.

465 ~~b)c)~~ [The validity of the dataset cannot be controlled in any way; everybody can alter the](#)
466 [original dataset once it is released by the drug agency. So the ban of sharing data is](#)
467 [useless.](#)

468 **78. How should EMA's policy be rolled out (timelines)?**

469 There was brief discussion as to whether the policy should be rolled out in a staggered way,
470 starting with high-level (aggregated) data, followed by more granular (patient-level) data sets. No
471 conclusion was reached (three positions).

472 a) If the name of the requester is not needed for aggregated data, then most points do not
473 need further discussion. A staggered roll-out should not delay implementation of the rules
474 to make data publicly available.

475 There is no obvious benefit and no reason to use a staggered way other than limited
476 capacity. Hence, there is no reason to postpone access to patient-level data

477 b) A staggered roll-out would be preferable as there are already many challenges to opening
478 up access to aggregated data which need to be solved. Aligning with the roll-out of the
479 EudraCT version 9 and access to results for many clinical trials could be an important step
480 forward. [Aggregated data, after consultation with the MAH for removal of CCI and PPD, is](#)
481 [more likely to have value to a wider audience and therefore should be of initial focus. A](#)
482 [staggered roll-out should be done by running several pilots to evaluate potential issues.](#)

483 c) A staggered approach would be pragmatic and could achieve much almost immediately.
484 There are many issues around the release of IPD, particularly around open public access
485 versus some model of conditional access. If this could be set aside for now with focus on
486 release of aggregate data and results of all statistical analyses as set out in the trial
487 protocol, rapid progress could be made. Access to IPD could follow after sufficient time for
488 discussion and enquiry. For example, potential impact of public release of IPD on
489 participant consent needs to be investigated. Therefore, separate the issues of (1) release
490 and access to trial information, results and aggregate data from (2) release and access to
491 IPD, and move ahead immediately with 1. Do not delay implementation of 1 while 2 is
492 addressed (it is much more complex and requires careful consideration). Extend the time
493 period to allow proper consideration and investigation of issues pertaining to 2. [However,](#)
494 [the delay of the access to IPD should only be delayed for a short time - one year.](#)

495 **89. Should requesters be encouraged to provide feedback?**

496 There is agreement that users of data should be encouraged to link back the results of their
497 analyses to the accessed data in order to ensure two-way transparency.

498 While a link back of results of individual analyses is desirable, it should be located on a separate
499 database in order to not increase subjective hurdles to lay people. This database should/could be
500 linked to the database of analysis plans/protocols.

501 It may also be useful to add a user/log-in concept to the repository to allow requesters to build
502 project websites. These project websites would give requesters the opportunity to publish
503 timelines, the protocol and the results of their project (or links to such documents).

504 Several comments/views along the following lines were expressed:

30 April 2013

Advice to the European Medicines Agency from the Clinical trial Advisory Group on Rules of engagement (CTAG3)

Draft Advice Version 7.0 – With amendments following comments on 6.0

- 505 | • Just encouraging requesters to link their analyses back to the data accessed is not
506 | sufficient. Further discussion is needed on how any resulting publications arising from
507 | secondary analyses are linked back to data access requests. Principles should be included
508 | on minimal expectations of requesters and what should be fed back having been granted
509 | access to data. For example, should the requester have to summarise their key findings of
510 | their analyses as a minimum? Publishing has to be accepted not only in the form of articles
511 | in journals but also as other documents with open access from the internet.
- 512 | • EMA should be committed to comment / answer in some way whatever new evidence
513 | brought up by requesters after its analysis.
- 514 | • —
- 515 | • On the assumption that access to anonymised patient level data is granted for a defined
516 | research project, access to a secure area should be granted for a defined duration (the
517 | duration necessary to complete the project). An open-ended access (beyond the research
518 | project) would undermine the benefits of identification and declaration of research
519 | purposes. NOTE from EMA: such proposals may not be compatible with the legal framework
520 | under which EMA operates as a public body; to be discussed at upcoming CTAG3 meeting
- 521 | • Requesters should be given a time frame within which they are obliged to publish/make
522 | public any outcomes and conclusions resulting from their analyses.
- 523 | • Requestors should be required to make publications derived from this work open access
524 | either via a journal or via deposition in a publicly available repository within 12 months of
525 | the completion of the work and a copy of the work supplied to EMA.
- 526 | — There should be no requirement for a time frame within which requestors are obliged to
527 | publish/make public the results of their analysis. However, if the EMA is constructing a
528 | database that will showcase the requests that have come in, also indicating which parties
529 | accessed what data, it would be nice to also include space for requestors to not only say
530 | what outcomes have resulted from their analysis (e.g. publications) but also encourage
531 | requestors who did not publish any resulting analyses to explain the reasons for no
532 | publication.
- 533 | •