

DRAFT

MUT/MIN/2016/1

COMMITTEE ON MUTAGENICITY OF CHEMICALS IN FOOD, CONSUMER PRODUCTS AND THE ENVIRONMENT

Minutes of the meeting held at 10.30 am on Thursday 25th February 2016 at the Department of Health in Room 136B Skipton House, Elephant and Castle, London, SE1 6LH.

Present:

Chairman:

Dr D Lovell

Members:

Dr C Beevers
Dr G Clare
Professor S Doak
Professor M O'Donovan
Ms P Hardwick
Professor G Jenkins
Professor D Kirkland
Dr A Lynch
Professor F Martin
Professor D Phillips

Secretariat:

Dr O Sepai (PHE Secretary)
Mr B Maycock (FSA Secretariat)
Dr K Burnett (PHE Tox Unit)
Mr S Robjohns (PHE Secretariat)

Assessors:

Dr L Koshy (HSE) (Item 3 via
teleconference)

In attendance:

Mr Daniel Medlock (PHE)

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1 **ITEM 1: ANNOUNCEMENTS/APOLOGIES FOR ABSENCE**

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3 1. The Chair welcomed Members, the secretariat and assessors. Mr B
4 Maycock was attending for the Food Standards Agency (FSA) and Mr D
5 Medlock was attending as an observer (Public Health England). The Chair
6 also welcomed the assessor Dr L Koshy (Health and Safety Executive)
7 attending via teleconference.

8
9 2. Apologies for absence were received from Dr D Benford (Secretariat
10 FSA), the Member Dr S Dean, and from the assessors Dr H Stemplewski
11 (MHRA), Dr S Fletcher (VMD) and Dr C Ramsay (Health Services Scotland).

12
13 3. Members were reminded of the need to declare any interests before
14 discussion of items and to ensure declarations of interests were kept up to
15 date.

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17 **ITEM 2: MINUTES OF MEETING ON 15th October 2015 (MUT/MIN/2015/2)**

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19 4. Members agreed the minutes subject to minor editorial changes.

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22 **ITEM 3: MATTERS ARISING**

23
24 5. The committee was informed by the HSE assessor that EFSA
25 (European Food Safety Authority) had published its conclusions on
26 glyphosate in November 2015 and concluded that it was unlikely to be
27 carcinogenic. The Joint FAO/WHO Meeting on Pesticides Review (JMPR)
28 would be evaluating glyphosate in May 2016 and the US Environmental
29 Protection Agency was currently conducting a review on glyphosate. One
30 member informed the COM that IARC (the International Agency for Research
31 on Cancer) was about to publish a letter on its opinion of glyphosate.

32
33 6. The secretariat informed the COM that it had received a request from a
34 company regarding the pesticide impurity and metabolite, para-chloroaniline
35 (also a precursor in the dye and pharmaceutical Industry). The COM had
36 previously considered this chemical in 2009 at the request of the UK Advisory
37 Committee on Pesticides (now the Expert Committee on Pesticides (ECP)). It
38 was concluded that para-chloroaniline is an *in vitro* mutagen, but the COM
39 could not conclude on the *in vivo* mutagenicity based on the data provided. A
40 strategy for the conduct of further genotoxicity testing was proposed. The
41 company had conducted the suggested studies and requested that the COM
42 consider the new data. The secretariat liaised with the ECP who confirmed
43 that they support the proposal for the COM to review the new data. It was
44 intended that the company could present the new data to the COM at either
45 the June or October 2016 meetings.

46
47 7. The COM also heard that the Triennial review of the COM was
48 currently going through ministerial approval and that the final report was likely
49 to be approved and published in March 2016. Following recommendations
50 from the review, there would be 'light touch' annual appraisal of members and

1 consideration of further training for the secretariat. Additionally, closer working
2 between various Chairs of advisory committees and with the Directorate of the
3 Public Health England Centre for Radiation, Chemical and Environmental
4 Hazards (CRCE) had been recommended by both the COM Triennial review
5 and the recent CRCE review. Therefore, future meetings would be considered
6 in line with these recommendations.

7 8. The Chair announced that it was the last meeting of Professor David
8 Phillips who had come to the end of his term. It was also the end of his term
9 as Chair of the COC. The Chair thanked Professor Phillips for all his excellent
10 work over the years and the COM provided its best wishes for the future.

11 12 13 **ITEM 4: ASSAYS USED TO EVALUATE GERM CELL DNA INTEGRITY IN** 14 **HUMAN FERTILITY INVESTIGATIONS (MUT2016/01)**

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16 9. At the previous October 2015 meeting members considered a paper on
17 germ cell mutagenesis and aging. A paper on radiation and transgenerational
18 effects was also considered. Recent developments in germ cell mutagenicity
19 in general had also been discussed, including the possibility that a chemical
20 could be a germ cell mutagen, but not a somatic cell mutagen, and that air
21 pollution should be classified as a human germ cell mutagen.

22
23 10. During a literature review and preliminary investigations for a further
24 paper on whether air pollution was a germ cell mutagen, it was noted that
25 assays for DNA integrity utilised in assisted reproductive technologies (ART)
26 were often used as a marker of DNA damage in human sperm. It was
27 therefore considered appropriate for the COM to evaluate these assays
28 before evaluating in more detail the suggestion that air pollution is a germ cell
29 mutagen. It was not clear what value the sperm chromatin structure assay
30 (SCSA) and the TUNEL (terminal deoxynucleotidyl transferase dUTP nick end
31 labelling) assays may have in investigating potential germ cell mutagenesis in
32 humans.

33
34 11. Some information on germ cell DNA damage in humans has been
35 obtained from studies investigating infertility or those aiming to improve the
36 outcome of assisted reproduction technologies (ART). A report by Yauk et al.,
37 2015 on an IWGT workshop on germ cell assays described these two assays
38 (i.e. the SCCA and the TUNEL) and noted that while they generally correlate
39 well with each other they measure different aspects of DNA integrity and
40 therefore will differ in sensitivity. Paper MUT 2016/01 provided an overview of
41 both the SCSA and the TUNEL assays for consideration by the COM.

42
43 12. Members noted that both the SCCA and the TUNEL were primarily
44 assays for the detection of DNA strand breaks. They should be considered
45 only as indicator assays and not informative on the consequences of the DNA
46 strand breaks or downstream events. For example, they were not informative
47 on whether the DNA strand breaks lead to a mutation, to apoptosis, or
48 whether they would be repaired. The papers referred to in MUT/2016/01
49 mainly looked at effects on fertility. A key question raised was whether the
50 observed reduced fertility was due to a genotoxic effect or a toxic effect.

Members also agreed that although both assays measured DNA strand breaks, they measured different types of DNA stand breaks. The significance and cause of the detected DNA strand breaks was also unclear. The observed DNA fragmentation could have arisen due to different reasons, such as chemical induced oxidative stress, apoptosis, or from another process not involving genotoxicity. There also appeared to be a relatively high background level and range of DNA strand breaks present in sperm, which would make it difficult to detect a chemically induced increase in DNA fragmentation. Furthermore, it was not clear at what point in spermatogenesis the DNA damage occurs. Members considered that it may be useful for the COM to have some input from an expert in this field, such as reproductive biologist with some knowledge of genotoxicity.

13. The COM considered that that there was some lack of consistency and conflicting results reported in the data and papers provided (e.g. there were conflicting reports on the correlation between the SCSA and the TUNEL assays).

14. Overall, members considered that there were a number of reasons why the results of both the SCSA and the TUNEL assays would be difficult to interpret in terms of germ cell mutagenicity e.g. they were indirect methods for evaluating potential germ cell mutagenicity; there was a lack of consistency between some of the data and in the test methods used; uncertainty over the underlying biology leading to the formation of DNA strand breaks and downstream effects; a large variation in background levels and range of effects; and a lack of validation of the test methods.

15. The COM considered that these assays may be able to indicate or contribute to lines of evidence for potential DNA damage caused by genotoxic chemicals, but there were a number of uncertainties as outlined above. It would be useful to harmonise these methodologies and for the validation of these assays to be undertaken. The COM would wait for the outcome of an IARC working group and further scientific developments on germ cell mutagenicity and the suggested germ cell mutagenicity of air pollution before deciding whether to conduct its own detailed review.

ITEM 5: GERM CELL ADVERSE OUTCOME PATHWAYS (MUT/2016/02)

16. As part of the COM's ongoing review of germ cell mutagenesis, the secretariat were made aware of recent papers by a group from Health Canada (Yauk et al., 2015 and Marchetti et al., 2015) regarding adverse outcome pathways (AOP) for germ cell endpoints. An individual AOP is specific to a molecular initiating event (MIE) and is not chemical specific. Key events (KE) are identified for the toxicological effect, which should be measurable. The connection between KE's is referred to as a KE relationship (KER). Modified Bradford Hill criteria are used to evaluate the empirical evidence and biological knowledge and this evaluation establishes the KERs. The DNA alkylation AOP (Yauk et al., 2015) focused on premeiotic germ cell DNA alkylation using ethylnitrosourea as a model alkylating agent. Unique features of germ cells suggest that they should be considered separately from

1 somatic cells. The AOP makes the assumption that the processes of DNA
2 repair and damage are conserved across eukaryotic cells. The tubulin binding
3 AOP (Marchetti et al., 2015) uses colchicine as a model example and says
4 that the majority of evidence is generated from rodents. It was noted that
5 benzimidazoles induce this AOP. Members were asked for their views on the
6 AOP approach and whether this would have any impact on the COM's 2007
7 statement on benzimidazoles (COM/07/S3).

8
9 17. Members considered that AOPs were useful for capturing and clarifying
10 information obtained from systems biology approaches and to provide
11 frameworks to aid in the communication of mode of actions, but there was
12 some way to go before they could be used to evaluate the safety of
13 chemicals. They had the potential to help communicate and explain expert
14 evaluations and scientific reports (i.e. to make them more accessible to lay
15 individuals). The two examples provided were very specific and more
16 qualitative than quantitative. The AOPs would likely build over time and could
17 aid predictive toxicology. It was noted that they were intended to be
18 chemically agnostic. The two papers provided were considered to illustrate
19 how to develop AOPs with two already well understood mechanisms (e.g.
20 tubulin binding leading to microtubule depolymerisation and alkylation).
21 Currently, one of the main difficulties was the different terminology used by
22 different specialist areas of toxicology. There was no consensus on
23 terminology, which would need to be addressed. It was noted that systems
24 biology may help with this, as it already had a number of agreed terms (e.g.
25 for receptor binding, antagonism, agonist etc.).

26
27 18. There was some discussion of whether AOPs could be used to
28 evaluate mixtures of chemicals or in risk assessment. The COM considered
29 that they could not currently be used for either as there were too many
30 uncertainties, alternative pathways and potential chemical interactions (e.g.
31 each chemical could have more than one AOP). Regarding the COM 2007
32 statement on benzimidazoles, where a 'common mechanism' of toxicity had
33 been identified (aneugenicity via inhibition of tubulin polymerisation),
34 members considered that there was some similarity between the flowchart in
35 the statement illustrating the benzimidazole 'common mechanism group' and
36 the more detailed AOP. Although the terminology used was a little different to
37 that used currently, the COM agreed that the statement still remained valid
38 and did not need to be changed.

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40 19. Overall, members noted that there was a lot of work and interest in the
41 area of AOPs (e.g. ECVAM were interested due to the potential to reduce
42 animal testing). There was likely some way to go before they could be used
43 by the COM. However, in the longer term, AOPs could aid communication of
44 expert opinions (e.g. in explaining the difference between hazard and risk).
45 The committee agreed to keep a watching brief on the development of AOPs
46 for mutagenicity.

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48 **ITEM 6: ANNUAL REPORT 2015 (MUT/2016/03)**
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1 20. A draft COM annual report for 2015 had been prepared. The committee
2 was asked to provide any comments. Comments such as typographical
3 amendments could also be sent to the secretariat. A revised version would be
4 circulated following members comments and this would be cleared by Chair's
5 action before publication.

6 7 **ITEM 7: ANY OTHER BUSINESS**

8 9 **1) Dominant lethal assay**

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11 21. The dominant lethal test was discussed due to an email from the
12 National Toxicology Program in the USA requesting that the OECD Test
13 Guideline 478 not be deleted because it was still used (e.g. in the USA) and
14 the assay was still considered useful for detecting germ cell mutagens.

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16 22. The COM considered that it would support the current UK view to
17 delete the OECD Test Guideline 478 for the dominant lethal test.

18 19 **2) COC statement on alcohol and cancer**

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21 23. Professor Phillips, the Chair of the COC, thanked the COM for its
22 advice on alcohol that had been used in the COC's detailed review and
23 statement on the carcinogenicity of alcohol. The COC published its statement
24 on alcohol and cancer in January 2016 at the same time as the Chief Medical
25 Officer's (CMO) report on alcohol. They were both broadly compatible in
26 terms of cancer.

27 28 **ITEM 8: DATE OF NEXT MEETING**

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30 24. 16th June 2016.
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