Final Agenda

Tenth meeting of the Committee for Risk Assessment

16 – 18 March 2010
Helsinki, Finland
16 March: starts at 9:00
18 March: ends at 15:00

Item 1 – Welcome & Apologies

Item 2 – Adoption of the Agenda

RAC/A/10/2010
For adoption

Item 3 – Declarations of conflicts of interest to the Agenda

Item 4 – Adoption of the draft minutes of RAC-9

- Adoption of the draft minutes

RAC/M/09/2010 draft final
For adoption

Item 5 – Administrative issues and information items

a. Status report on the RAC - 9 action points
b. Outcome of written procedures
c. Report from other ECHA bodies and activities

RAC/10/2010/18
ROOM DOCUMENT
For information
d. Feedback on the annual survey of members

RAC/10/2010/18
ROOM DOCUMENT
For information
e. Update on the financial arrangements for (co-) rapporteurs for restriction dossiers
Item 6 – Feedback from the MB decision on approval of RAC Rules of procedure

- Handling minority positions

For discussion and decision

Item 7 – Requests according to Art 77(3)(c) of REACH

a. Final draft opinion on boric acid and its compounds in photographic applications
   
   For adoption

b. Framework for dealing with requests according to Art 77(3)(c) of REACH
   
   RAC/10/2010/12
   
   For agreement

Item 8 – CLH

8.1 CLH Dossiers

a. Epoxiconazole
   
   For adoption

b. Abamectin/Avermectin B1a
   
   For discussion and possible adoption

c. Gallium arsenide
   
   For discussion and possible adoption

d. Tetrahydrofuran
   
   For discussion

e. TDCP
   
   For discussion

f. Leucomalachite Green – accordance check
   
   For discussion

8.2 Appointment of RAC (co-) rapporteurs for CLH dossiers

- Appointment of RAC (co-) rapporteurs for CLH dossiers
   
   RAC/10/2010/20
   
   ROOM DOCUMENT
   
   For agreement

8.3 General CLH issues

a. Templates for the CLH opinion and BD and Commission’s feedback on RAC request

b. Substances already agreed at TC C&L

c. Note H, hazard statements on reprotoxicity, justification for non-CMR&RS proposals
d. State of play of the submitted CLH dossiers

RAC/10/2010/22
ROOM DOCUMENT
For discussion

e. Feedback from the Ad Hoc meeting for exchanging experience on accordance check for CLH dossiers

RAC/10/2010/13
For discussion

f. Handling a group of substances

For discussion

Item 9 – Restrictions

9.1 Report from the meeting of RAC and SEAC (co-)rapporteurs and ECHA Secretariat

For information

9.2 General restriction issues

• Update on intended restriction dossiers

For information

Item 10 – RAC manual of conclusions and recommendations

• Revised RAC manual of conclusions and recommendations

RAC/10/2010/14 & RAC/10/2010/15
For discussion and possible outline approval

Item 11 – Authorisation

a. Working procedure for the appointment of rapporteurs for applications for authorisations

RAC/10/2010/16 & RAC/10/2010/17
For agreement

b. RAC role in the authorisation process

For information

Item 12 – Guidance issues

a. Feedback from the guidance update on the DNEL/DMEL derivation from human data

b. Feedback from the RAC consultation on the CLH guidance document

RAC/10/2010/23
ROOM DOCUMENT
For information
c. Report on other guidance activities

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<th>Item 14 – Main conclusions and Action Points of RAC-10</th>
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