AGENDA

Section A  Information and/or discussion

A.01 Summary Report of previous meetings.

A.02 New active substances:
   1. New admissible dossiers to be noted:
      (No new dossiers.)
   2. Exchange of views on new European Food Safety Authority (EFSA) conclusions
      (No specific conclusions identified.)

Commission Draft Review Report and Regulation concerning the (non-) approval of:
   Beta-cypermethrin  (No detailed discussion; Member States are requested to send in comments after the meeting)
   Bacillus amyloliquefaciens FZB24
   Beauveria bassiana strain 147
   Beauveria bassiana NPP 11B005

A.03 Renewal of approval:
   1. Annex I Renewal Projects: State of play
   2. Exchange of view on EFSA conclusions
      (No specific conclusions identified)
   3. Draft Review/Renewal Reports and Regulations for discussion:
i. Flupyrdsulfuron-methyl
ii. Pymetrozine
iii. Imazamox
iv. Maleic hydrazide
v. Flazasulfuron
vi. Coniothyrium minitans strain CON/M/91-08
vii. Mesosulfuron-methyl (No detailed discussion; Member States are requested to send in comments after the meeting)
viii. Mesotrione
ix. Pendimethalin
x. 2,4-DB
xi. Carfentrazone-ethyl
xii. Acetamiprid
xiii. Propyzamide
xiv. Propoxycarbazone-sodium (No detailed discussion; Member States are requested to send in comments after the meeting)
xv. Benzoic acid (No detailed discussion; Member States are requested to send in comments after the meeting)
xvi. Diquat

A.04 Confirmatory Data:
1. Bifenthrin
2. Thiamethoxam
3. Clothianidin
4. Imidacloprid
5. Tetraconazole
6. Diclofop (revised Review Report to be noted)
7. Cyflumetofen
8. 8-hydroxyquinoline (revised Review Report to be noted)
9. AOB

A.05 Article 21 Reviews:
i. Diflubenzuron (Draft Review Report and draft Implementing Regulation for discussion)
ii. Thiametoxam, other uses than seed treatments and granules (revised Review Report to be noted)
iii. Clothianidin, other uses than seed treatments and granules (revised Review Report to be noted)
iv. Imidacloprid, other uses than seed treatments and granules (revised Review Report to be noted)

A.06 Amendment of the conditions of approval:
1. New admissible dossiers to be noted
i. Paraffin oils

2. 8-Hydroxyquinoline

3. Penflufen

A.07 Basic substances:

1. Pilot projects: state of play

2. New dossiers received (only for information):
   i. Fructose (extension of use)
   ii. Propolis

3. Exchange of view on EFSA Technical Reports

(No specific report identified)

4. Draft Review Reports for discussion:

Capsicum spice (No detailed discussion; Member States are requested to send in comments after the meeting.)
Millefolii herba (No detailed discussion; Member States are requested to send in comments after the meeting.)

A.08 Exchange of views and possible taking note of the following Guidance Documents:

1. Draft Guidance Document on the assessment of exposure of operators, workers, residents and bystanders in risk assessments for plant protection products (Doc. SANTE/10832/2015) (to be noted)
2. Guidance Document on the renewal of approval of active substances to be assessed in compliance with Regulation (EU) No 844/2012 (for discussion - changes of specification)

A.09 Notifications under Article 44(4) of Regulation (EC) No 1107/2009 (to be noted).

A.10 Notifications under Article 36(3) of Regulation (EC) No 1107/2009 (to be noted).

A.11 Notifications under Article 53 of Regulation (EC) No 1107/2009 (to be noted).

A.12 News from European Food Safety Authority (EFSA).
   1. Follow up workshop formulation laboratories
   3. Article 68 Enforcement Working group

A.14 Report from working groups:
   1. Plant Protection Products Application Management System (PPPAMS)
   2. Post Approvals Issues group (PAI) (no meeting since December)
   3. Sustainable plant protection experts group Dutch proposal

A.15 OECD.

A.16 Bees:
   1. Review of Fipronil – state of play
   4. AOB

A.17 Court cases:
   - Cases C-442/14 and C-673/13 - Judgements announced for 23/11/2016

A.18 Endocrine disruptors.

A.19 Minor Uses.

A.20 Interpretation issues:
      i. Larvex
      ii. Siltac
      iii. Colour spray
2. Questions and answers

A.21 Classifications under Regulation (EC) No 1272/2008 / REACH:
   1. Status of harmonised classifications
   2. Preparation of Harmonised Classification and Labelling dossiers (CLH dossiers) by Member States
   3. Report from the WG on AR template (merging CLH and xAR templates)

A.22 Glyphosate:
   • State of the dossier

A.23 Exchange of information from the Pesticide Residues section of the Committee: possible impact on authorisations.

A.24 Proposal on amendment of criteria for the approval of low risk active substances (SANTE/12376/2015).


A.26 Commission Communications amending Commission Communications (2013/C 95/01-95/02) – General update.

Section B Draft(s) presented for an opinion


(B.01_SANTE_11619_2016 )

Legal Basis: Article 13(2)(c) of Regulation (EC) No 1107/2009
Procedure: Examination procedure
B.02 Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation confirming the conditions of approval of the active substance acrinathrin, as set out in Implementing Regulation (EU) No 540/2011. (Draft Review Report SANTE/11357/2011 Rev. 7)

(B.02_SANTE_11038_2016 Rev. 2)

Legal Basis: Article 13(2)(c) of Regulation (EC) No 1107/2009

Procedure: Examination procedure


(B.03_SANTE_11977_2016 Rev. 0)


Procedure: Examination procedure


(B.04_SANTE_11979_2016 Rev. 0)


Procedure: Examination procedure


(B.05_SANTE_11266_2016 Rev. 0)
**Legal Basis:** Article 13(2)(a) of Regulation (EC) No 1107/2009  
**Procedure:** Examination procedure


(B.06_SANTE_11806_2016 Rev. 0)

**Legal Basis:** Article 13(2)(a) of Regulation (EC) No 1107/2009  
** Procedure:** Examination procedure


(B.07_SANTE_11899_2016 Rev. 0)

**Legal Basis:** Article 13(2)(a) of Regulation (EC) No 1107/2009  
** Procedure:** Examination procedure


(B.08_SANTE_10680_2015 Rev. 4)

**Legal Basis:** Article 20(1) of Regulation (EC) No 1107/2009  
** Procedure:** Examination procedure

**B.09** Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation concerning the non-approval of the active substance cyclaniliprole, in accordance with Regulation (EC) No 1107/2009 of the European Parliament and of the Council concerning the placing of plant protection products on the market
(B.09_SANTE_11597_2016 Rev. 0)

**Legal Basis:** Article 13(2)(b) of Regulation (EC) No 1107/2009  
**Procedure:** Examination procedure

**B.10** Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation concerning the non-approval of the active substance *Pseudozyma flocculosa* ATTC 64874 in accordance with Regulation (EC) No 1107/2009 of the European Parliament and of the Council concerning the placing of plant protection products on the market. (Draft Review Report SANTE/10615/2016 Rev. 1)

(B.10_SANTE_10614_2016 Rev. 0)

**Legal Basis:** Article 13(2)(b) of Regulation (EC) No 1107/2009  
**Procedure:** Examination procedure

**B.11** Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation amending Implementing Regulation (EU) No 540/2011 as regards the conditions of approval of the active substance buprofezin (Draft Review Report SANCO/12256/2010 Final).

(B.11_SANTE_10311_2016 Rev. 1)

**Legal Basis:** Article 21(3) and Article 78(2) of Regulation (EC) No 1107/2009  
**Procedure:** Examination procedure

**B.12** Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation amending Implementing Regulation (EU) No 540/2011 as regards the conditions of approval of the active substance oxyfluorfen (Draft Review Report SANCO/11136/2011 Rev. 3).

(B.12_SANTE_10984_2016 Rev. 1)

**Legal Basis:** Article 21(3) and Article 78(2) of Regulation (EC) No 1107/2009  
**Procedure:** Examination procedure

**B.13** Exchange of views and possible opinion of the Committee on a draft Commission Implementing Regulation amending Implementing Regulation (EU) No 540/2011 as regards the extension of the approval periods of several active substances part of the AIR IV renewal programme and listed in Part B of the Annex to Implementing Regulation (EU) No 686/2012
Legal Basis: First paragraph Article 17 and Article 78(2) of Regulation (EC) No 1107/2009
Procedure: Examination procedure

Miscellaneous

M.01 New Scientific publications and information submitted by stakeholders.

M.02 Antibiotics – Yearly reporting by Member States

M.03 Date of next meeting.