Working document on pesticides to be considered for inclusion in the national control programmes to ensure compliance with maximum residue levels of pesticides residues in and on food of plant and animal origin.

This document has been conceived as a working document of the Commission Services. It does not represent the official position of the Commission. It does not intend to produce legally binding effects.

Only the European Court of Justice has jurisdiction to give preliminary rulings concerning the validity and interpretation of acts of the institutions of the EU pursuant to Article 267 of the Treaty.
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1. **Scope**

This document serves the dual purpose of:
- Proposing pesticides to be included in the EU Multi-Annual Control Programme (EU MACP).
- Recommending pesticides to be included in the National Control Programmes (NCPs) of the Member States on a voluntary basis.

The assessment of active substances is based on:
- occurrence data originating from EFSA's annual reporting data
- toxicological reference data found on the EU MRL database and
- analytical capabilities of the EU laboratories which are assessed via an annual survey conducted by the EURL-SRM.

This document is revised each year following the Working Group Meeting of Experts on monitoring of pesticide residues in/on food and is endorsed by the Standing Committee on Plants, Animals, Food and Feed, section pesticides residues (SC PAFF phytopharmaceuticals – section residues) and serves as a preliminary evaluation of the pesticides included on the annual European Commission Regulation.

2. **Introduction**

On 4 October 2013 an Expert Group Meeting on Pesticides Residues Monitoring was held in Brussels. In this meeting it was agreed not to include voluntary analyses in the Regulation concerning the EU MACP for 2015, 2016 and 2017. However, it was deemed necessary to already highlight in advance certain pesticides, which following the assessment detailed in Chapter 3, could be considered for inclusion in the Regulation for the EU MACP. These pesticides are listed in Chapter 4 of this document and can be, on a voluntary basis, taken up in the National Control Programmes of the Member States during the assessment period. After an evaluation of the analytical capability and the monitoring data gathered under the National Control Programmes, their inclusion or non-inclusion in the EU MACP is considered.

The document is completed by a series of Annexes as detailed below:

- **Annex I** includes pesticides for which monitoring data are required for specific risk management questions.
- **Annex II** lists pesticides for which support is needed from the EURel.ks.
- Pesticides that are of interest to EFSA for cumulative risk assessment and which are not taken up in the chapter 4 of this document or the MACP, are included in **Annex III** to this document.
- **Annex IV** includes active substances for which occurrence data indicated very few findings and, thus, can include substances coming from the Chapter 4 assessment or even from the list included in the EU MACP.
- **Annex V** details the assessment methodology of the active substances.
- **Annex VI** includes the form of proposals of pesticides to be assessed by Member States or EURel.ks.
- Substances of interest to be analysed in honey under national control programmes are listed in **annex VII**.
- Commodities of interest to be analysed under the national control programmes are listed in **annex VIII**.
- Substances that have been moved from Chapter 4 of this document into the EU MACP are listed in **Annex IX**.
- **Annex X** provides a brief description of a project regarding the collection of samples of organic products of plant origin for the determination of background levels of dithiocarbamates (CS2), as several false positive analysis results indicate the natural occurrence of CS2 in specific plant products.

**Residue Definitions:**
All pesticides mentioned in this document are recommended to be analysed for their full and legal residue definition according to Reg. (EC) N° 396/2005. In order to avoid that this document would be outdated due to future changes in residue definitions, only the general name of the residue definition is mentioned. For the full details of each residue definition, as well as specific residue definitions for certain commodities, reference is made to the most recent version of Reg. (EC) No 396/2005.

3. Categorisation, prioritisation and assessment

During the SCOFCAH of 12-13 June 2014 the Member States were requested to take a position on the approach for categorisation and prioritisation of the substances that are taken up in this document. A majority of the Member States was in favour of an approach in which the pesticides are divided into specific categories. Based on a limited set of criteria each pesticide is attributed a priority and a time line for evaluation of inclusion or non-inclusion in the MACP.

3.1. Categorisation

The pesticides in Chapter 4 are split up into the following categories:

- Frequent detections, MRL exceedances or RASFF notifications.
  - Based on the occurrence data of the 3 previous years (starting from the year with the latest data available), candidates for inclusion in this WD are substances with (a) MRL exceedances and/or (b) high rate of findings (≥0.5% of samples) for 3 consecutive years (for animal commodities where findings are less, a rate of ≥0.01% can be taken into account).
  - Based on the RASFF notifications of 3 years, the 15 substances with the highest frequency of occurrence in the alerts are examined for findings for 3 years. The above procedure is followed.
- Recently approved substances. Substances approved during the time interval between two consecutive working group meetings.
- Art. 12 priority list.
- High toxicity.

3.2. Prioritisation

The substances included in Chapter 4 of this document are prioritised based on the type of analytical method.

- MRM method: priority 1
- MRM/ SRM or SRM method: priority 2
- In case no standards and/or analytical method are available for substances that qualify to the categories mentioned under chapter 3.1, the substances are not included in chapter 4. They are however taken up in Annex II to this document that lists substances for which support from the EURLs is requested.

A further refinement of the priority is made based on toxicity.

- if ADI ≤ 0.1 mg/kg bw/day or ARfD ≤ 0.1 mg/kg bw, then priority A is assigned.
- if ADI > 0.1 mg/kg bw/day and ARfD > 0.1 mg/kg bw, then priority B is assigned.

Based on the above, prioritization is illustrated in the following table:
Table 1. Prioritization Matrix of Active Substances

<table>
<thead>
<tr>
<th>Toxicity</th>
<th>Analytical Capability</th>
<th>Priority 1</th>
<th>Priority 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Priority A</td>
<td>ADI ≤ 0.1 mg/kg bw/day or ARfD ≤ 0.1 mg/kg bw</td>
<td>1A</td>
<td>2A</td>
</tr>
<tr>
<td>Priority B</td>
<td>ADI &gt; 0.1 mg/kg bw/day and ARfD &gt; 0.1 mg/kg bw or No Toxicological Reference Values Available</td>
<td>1B</td>
<td>2B</td>
</tr>
</tbody>
</table>

- For pesticides with priorities 1A and 1B, the evaluation will be done after 1 year, for categories 2A and 2B after 2 years.
- The sub-priorities A and B, which are linked to the toxicity, don't affect the evaluation timeline and are only for information to the MS, in case they want guidance on which substances should be prioritised.
- In case of RASFF notifications it is possible to accord a higher priority to certain specific substances after discussions in the expert group.

3.3. Assessment

As illustrated in Figure 1, frequently detected substances as defined in 3.1, recently approved substances, substances identified as top-15 in annual RASFF findings, high toxicity substances and Art.12 priority substances can be included in Chapter 4 of this document based on the discussion of the experts during the working group. Based on the datasets of 3 years preceding EFSA’s latest published annual report, in the case a Chapter 4 active substance indicates MRL exceedances and/or findings of more than 0.1% of the analysed samples for 3 years consecutively, and if there is good (≥60%) analytical capability across EU laboratories, then that active substance is eligible for addition on the EU MACP depending on the experts' evaluation. In case analytical capability is <60% then the substance is placed in Annex II for support from the EURLs and is re-evaluated in 1 or 2 years depending on the prioritisation factor of that substance (1yr for 1A/1B, 2yrs for 2A/2B).

Figure 1. Assessment Flow Chart
4. Pesticides to be considered for analysis in products of plant origin

The substances are listed in alphabetical order, separately for commodities of plant origin and of animal origin and per category. Substances newly added to this version of the WD are indicated in white background, while older substances that were evaluated during the 2018 WG are in grey background.

4.1. Pesticides to be considered for analysis in products of plant origin (PO)

4.1.1. Frequent detections\(^1\), MRL exceedances or RASFF notifications

---

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-CPA (4-chlorophenoxyaceticacid) (Not approved)</td>
<td>PO</td>
<td>10/2018</td>
<td>MRM/SRM</td>
<td>2B</td>
<td>after 2 years (10/2020)</td>
<td>no toxicological reference values available</td>
<td>0.03% findings (0.02% MRL exceedances) EFSA 2014</td>
<td>0.03% findings (0.02% MRL exceedances) EFSA 2015</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlordecone (Not approved)</td>
<td>PO</td>
<td>10/2018</td>
<td></td>
<td></td>
<td></td>
<td>no toxicological reference values available</td>
<td>0.74% findings (0.41% MRL exceedances) EFSA 2014</td>
<td>3.53% findings (1.36% MRL exceedances) EFSA 2015</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diafenthiuron (Not Approved)</td>
<td>PO</td>
<td>10/2018</td>
<td>MRM/SRM</td>
<td>1B</td>
<td>after 1 year (10/2019)</td>
<td>no toxicological reference values available</td>
<td>0.03% findings (0.03% MRL exceedances) EFSA 2014</td>
<td>0.00% findings (0.01% MRL exceedances) EFSA 2015</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dinotefuran (Not Approved)</td>
<td>PO</td>
<td>10/2018</td>
<td></td>
<td></td>
<td></td>
<td>no toxicological reference values available</td>
<td>0.07% findings (0.06% MRL exceedances) EFSA 2014</td>
<td>0.01% findings (0.03% MRL exceedances) EFSA 2015</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fenobucarb (Not Approved)</td>
<td>PO</td>
<td>10/2018</td>
<td>MRM</td>
<td>2B</td>
<td>after 1 year (10/2019)</td>
<td>no toxicological reference values available</td>
<td>0.09% findings (0.06% MRL exceedances) EFSA 2014</td>
<td>0.00% findings (0.02% MRL exceedances) EFSA 2015</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>PO</th>
<th>Added</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Toxicity</th>
<th>MRL exceedances</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fosetyl-Al</td>
<td>PO</td>
<td></td>
<td>SRM</td>
<td>2B</td>
<td>after 2 years (10/2017)</td>
<td></td>
<td></td>
<td>1.3% Findings in vegetables, 0.5% in fruits and nuts EFSA 2011</td>
</tr>
</tbody>
</table>

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\(^1\) SRM-compounds are typically analysed on specific commodities so their detection frequencies are typically higher than if they would have been analysed randomly.
4.1 Pesticides to be considered for analysis in products of plant origin

SANCO/12745/2013, Rev.10

Glufosinate ammonium – PO

At request of EFSA, since residues are found in animal origin commodities, interesting to also check soybean which is used both as food and feed.

Method: SRM
Toxicity: ADI = 0.021 mg/kg bw/day, ARfD = 0.021 mg/kg bw
Priority: 2A
Evaluation: after 2 years (10/2017) → 10/2018 → 10/2019

- 0.3% findings in vegetables EFSA 2011
- 0.37% findings in 2011-2013 (EURL priority list)
- 0% findings EFSA 2012
- 0.26% findings EFSA 2013
- 0.03% findings EFSA 2014
- 0.26% findings (0.00% MRL exceedances) EFSA 2015
- 0.82% findings (0.03% MRL exceedances) EFSA 2016

6% labs and 23% MS analysed full RD in 2015
10% labs and 27% MS analysed full RD in 2016
14% labs and 32% MS analysed full RD in 2017

⇒ Analytical capability poor
⇒ Not clear if findings justify inclusion in EU MACP
⇒ Keep extra year in Chapter 4 of WD

Especially relevant for apples, bananas, broccoli, cauliflowers, cereals, cultivated fungi, grapefruit, head cabbage, kiwi, lettuce, melons, onions, oranges, pears, peppers (sweet), potatoes, strawberries, rice, table grapes, tomatoes and wheat. Additionally relevant for several non-MACP commodities such as: celery, cumin, maize and soybeans.

Nicotine (Not Approved) – PO

Added: 10/2018

Toxicity: ADI = 0.0008 mg/kg bw/day, ARfD = 0.0008 mg/kg bw/day
Method: MRM
Priority: 1A
Evaluation: after 1 year (10/2019)

- 2.66% findings (0.77% MRL exceedances) EFSA 2014
- 0.25% findings (0.13% MRL exceedances) EFSA 2015
- 1.76% findings (0.21% MRL exceedances) EFSA 2016

24% labs and 58% MS analysed full RD in 2016
20% labs and 39% MS analysed full RD in 2017

Also included in Annex I, but listed here as it may be of concern of the EU MACP commodities (e.g. brassica crops).

Novaluron (not approved) – PO

Toxicity: ADI = 0.01 mg/kg bw/day, ARfD NA
Method: MRM
Priority: 1A
Evaluation: after 1 year (10/2018) → 10/2019

- 0.14% findings (0.00% MRL exceedances) EFSA 2013
- 0.12% findings (0.00% MRL exceedances) EFSA 2014
- 0.06% findings (0.00% MRL exceedances) EFSA 2015
- 0.05% findings (0.00% MRL exceedances) EFSA 2016

45% labs and 58% MS analysed full RD in 2016
49% labs and 71% MS analysed full RD in 2017

⇒ Analytical capability medium
⇒ Not clear if findings justify inclusion in EU MACP
⇒ Keep extra year in Chapter 4 of WD

Especially relevant for apples, bananas, broccoli, cauliflowers, cereals, cultivated fungi, grapefruit, head cabbage, kiwi, lettuce, melons, onions, oranges, pears, peppers (sweet), potatoes, strawberries, rice, table grapes, tomatoes and wheat. Additionally relevant for several non-MACP commodities such as: avocado, basil, beans with pods, cherries, Chinese cabbage, clementines, mandarins, fresh herbs (coriander, celery leaves), garlic, lemons, limes, lychee, mangoes, papayas, guavas passion fruits, peas with pods, pineapple, peppers (chili), plums, pomegranates, pomelos, shallots, tea, wild fungi.

Prochloraz – PO

Toxicity: ADI = 0.01 mg/kg bw/day, ARfD = 0.025 mg/kg bw/day
Method: SRM (possible future revision of residue definition that would allow MRM method).
Evaluation: after 1 year (10/2019)

- 1.8% findings EFSA 2012 report
- 1.63% findings EFSA 2013 report
- 1.19% findings (0.01% MRL exceedances) EFSA 2014
- 1.20% findings (0.05% MRL exceedances) EFSA 2015
- 1.17% findings (0.02% MRL exceedances) EFSA 2016

10% labs and 35% MS analysed full RD in 2015
11% labs and 42% MS analysed full RD in 2016
14% labs and 7% MS analysed full RD in 2017

Especially relevant for apples, bananas, broccoli, cauliflowers, cereals, cultivated fungi, grapefruit, head cabbage, kiwi, lettuce, melons, onions, oranges, pears, peppers (sweet), potatoes, strawberries, rice, table grapes, tomatoes and wheat. Additionally relevant for several non-MACP commodities such as: avocado, basil, beans with pods, cherries, Chinese cabbage, clementines, mandarins, fresh herbs (coriander, celery leaves), garlic, lemons, limes, lychee, mangoes, papayas, guavas passion fruits, peas with pods, pineapple, peppers (chili), plums, pomegranates, pomelos, shallots, tea, wild fungi.
<table>
<thead>
<tr>
<th>Substance</th>
<th>Added:</th>
<th>Toxicity:</th>
<th>Method:</th>
<th>Priority:</th>
<th>Evaluation:</th>
<th>Findings</th>
<th>Analytical capability</th>
<th>Findings justify inclusion in EU MACP</th>
<th>Keep extra year in Chapter 4 of WD so that more labs have the time to add this substance to their scope.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyridalil – PO</td>
<td></td>
<td>ADI = 0.03 mg/kg bw/day, ARfD NA</td>
<td>MRM</td>
<td>1A</td>
<td>after 1 year (10/2018) → 10/2019</td>
<td>0.08% findings (0.00% MRL exceedances) EFSA 2013</td>
<td>medium</td>
<td>√ Analysed full RD in 2016</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Pyrethrins – PO</td>
<td></td>
<td>ADI = 0.04 mg/kg bw/day, ARfD = 0.2 mg/kg bw</td>
<td>MRM/SRM</td>
<td>2A</td>
<td>after 2 years (10/2017) → 10/2018 → 10/2019</td>
<td>0.06% findings EFSA 2012</td>
<td>medium</td>
<td>√ Analysed full RD in 2017</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Quinalphos (not approved) – PO</td>
<td>10/2018</td>
<td>no toxicological reference values available</td>
<td>MRM</td>
<td>1B</td>
<td>after 1 year (10/2019)</td>
<td>0.02% findings (0.01% MRL exceedances) EFSA 2014</td>
<td>medium</td>
<td>√ Analysed full RD in 2016</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Spinetoram – PO</td>
<td></td>
<td>ADI = 0.025 mg/kg bw/day, ARfD 0.1 mg/kg bw</td>
<td>MRM</td>
<td>1A</td>
<td>after 1 year (10/2018) → 10/2019</td>
<td>0.12% findings (0.00% MRL exceedances) EFSA 2013</td>
<td>medium</td>
<td>√ Analysed full RD in 2017</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Tolfenpyrad (not approved) – PO</td>
<td>10/2018</td>
<td>no toxicological reference values available</td>
<td>MRM</td>
<td>1B</td>
<td>after 1 year (10/2019)</td>
<td>0.14% findings (0.11% MRL exceedances) EFSA 2014</td>
<td>medium</td>
<td>√ Analysed full RD in 2016</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Trifluralin (not approved) – PO</td>
<td>10/2018</td>
<td>ADI = 0.015 mg/kg bw/day</td>
<td>MRM</td>
<td>2B</td>
<td>after 2 years (10/2020)</td>
<td>0.02% findings (0.01% MRL exceedances) EFSA 2014</td>
<td>medium</td>
<td>√ Analysed full RD in 2016</td>
<td>√ Findings justify inclusion in EU MACP</td>
</tr>
</tbody>
</table>
### 4.1.2. Recently approved substances

<table>
<thead>
<tr>
<th>Substance</th>
<th>Approval Date</th>
<th>Toxicity: ADI</th>
<th>ARfD</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Monitoring Data</th>
<th>Analytical Capability</th>
<th>Findings</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzovindiflpyr – PO</td>
<td>Approved since 03/2016</td>
<td>ADI 0-0.05 mg/kg bw day, ARfD 0.1 mg/kg bw</td>
<td></td>
<td>MRM</td>
<td>1A</td>
<td>after 1 year (10/2017)</td>
<td></td>
<td></td>
<td></td>
<td>Not clear if findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Fenpicoxamid – PO</td>
<td>Approved since 10/2018</td>
<td></td>
<td></td>
<td>MRM</td>
<td>2B</td>
<td>after 2 years (10/2020)</td>
<td></td>
<td></td>
<td></td>
<td>No data on analytical capability.</td>
</tr>
<tr>
<td>Oxathiapipronil – PO</td>
<td>Approved since 02/2014</td>
<td>ADI = 0.15 mg/kg bw/day</td>
<td></td>
<td>MRM</td>
<td>2B</td>
<td>after 1 year (10/2019)</td>
<td>No monitoring data available EFSA 2014, 2015, 2016</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pyriofenone – PO</td>
<td>Approved since 02/2014</td>
<td>ADI = 0.07 mg/kg bw/day, ARfD NA</td>
<td></td>
<td>MRM</td>
<td>1A</td>
<td>after 1 year (10/2018)</td>
<td></td>
<td></td>
<td></td>
<td>Not clear if findings justify inclusion in EU MACP</td>
</tr>
<tr>
<td>Sulfoxaflor – PO</td>
<td>Approved since 08/2015</td>
<td>ADI = 0.04 mg/kg day, ARfD = 0.25 mg/kg</td>
<td></td>
<td>MRM</td>
<td>1B</td>
<td>after 1 year (10/2017)</td>
<td></td>
<td></td>
<td></td>
<td>Not clear if findings justify inclusion in EU MACP</td>
</tr>
</tbody>
</table>
4.1 Pesticides to be considered for analysis in products of plant origin

4.1.3.  **Art. 12 priority list**

No pesticide identified under this category.

4.1.4.  **High toxicity**

No pesticide identified under this category.
### 4.2. Pesticides to be considered for analysis in products of animal origin (AO)

#### 4.2.1. Frequent detections\(^2\), MRL exceedances or RASFF notifications

<table>
<thead>
<tr>
<th>Pesticide</th>
<th>Added:</th>
<th>Toxicity: ADI</th>
<th>ARfD</th>
<th>Method</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Findings</th>
<th>Analytical capability</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azoxytrobin – AO</td>
<td>10/2018</td>
<td>ADI 0.2 mg/kg bw/day, ARfD NA</td>
<td>NA</td>
<td>MRM</td>
<td>2A</td>
<td>after 2 years (10/2020)</td>
<td>✓ 0.06 % findings (0.00% MRL exceedances) EFSA 2014&lt;br&gt; ✓ 0.46 % findings (0.00% MRL exceedances) EFSA 2015&lt;br&gt; ✓ 0.60 % findings (0.00% MRL exceedances) EFSA 2016</td>
<td>No data on analytical capability.</td>
<td></td>
</tr>
<tr>
<td>Chlorobenzilate (Not approved) – AO</td>
<td>10/2018</td>
<td>ADI 0.02 mg/kg bw/day, ARfD NA</td>
<td>NA</td>
<td>MRM</td>
<td>1B</td>
<td>after 1 year (10/2019)</td>
<td>✓ 0.06 % findings (0.00% MRL exceedances) EFSA 2014&lt;br&gt; ✓ 0.46 % findings (0.00% MRL exceedances) EFSA 2015&lt;br&gt; ✓ 0.60 % findings (0.00% MRL exceedances) EFSA 2016</td>
<td>No data on analytical capability.</td>
<td></td>
</tr>
<tr>
<td>Endrin (Not approved) – AO</td>
<td>10/2018</td>
<td>ADI 0.0002 mg/kg bw/day, ARfD NA</td>
<td>NA</td>
<td>MRM</td>
<td>1A</td>
<td>after 1 year (10/2019)</td>
<td>✓ 0.05 % findings (0.00% MRL exceedances) EFSA 2014&lt;br&gt; ✓ 0.13 % findings (0.00% MRL exceedances) EFSA 2015&lt;br&gt; ✓ 0.14 % findings (0.00% MRL exceedances) EFSA 2016</td>
<td>No data on analytical capability.</td>
<td></td>
</tr>
<tr>
<td>Carbendazim and thiophanate methyl – AO</td>
<td>10/2018</td>
<td>ADI = 0.02 mg/kg bw/day, ARfD = 0.02 mg/kgbw</td>
<td>ADI = 0.02 mg/kg bw/day, ARfD = 0.02 mg/kgbw</td>
<td>MRM/SRM</td>
<td>2A</td>
<td>after 2 years (10/2017)</td>
<td>2.28% findings EFSA 2012&lt;br&gt; 0% findings EFSA 2013 (712 samples)&lt;br&gt; 0.37% findings EFSA 2014 (1350 samples)&lt;br&gt; 1.49% findings (0.00% MRL exceedances) EFSA 2015&lt;br&gt; 0.27% findings (0.00% MRL exceedances) EFSA 2016</td>
<td>51% labs and 68% MS analysed full RD in 2015&lt;br&gt; 42% labs and 72% MS analysed full RD in 2016&lt;br&gt; 38% labs and 64% MS analysed full RD in 2017</td>
<td>Analytical capability medium&lt;br&gt; Findings justify inclusion in EU MACP&lt;br&gt; Keep 1 extra year in chapter 4 of WD so that more labs have the time to add this substance to their scope. Relevant for honey.</td>
</tr>
<tr>
<td>Pendimethalin – AO</td>
<td>10/2018</td>
<td>ADI 0.125 mg/kg bw/day, ARfD 0.3 mg/kg bw</td>
<td>ADI 0.125 mg/kg bw/day, ARfD 0.3 mg/kg bw</td>
<td>MRM</td>
<td>1B</td>
<td>after 1 years (10/2019)</td>
<td>0% findings EFSA 2012 (2 samples)&lt;br&gt; 0% findings EFSA 2013 (100 samples)&lt;br&gt; N.D. EFSA 2014 (93 samples)&lt;br&gt; N.D. EFSA 2015 report (11 samples)&lt;br&gt; N.D. EFSA 2016 preliminary report (91 samples)</td>
<td>21% labs and 56% MS analysed full RD in 2015&lt;br&gt; 26% labs and 43% MS analysed full RD in 2016&lt;br&gt; 18% labs and 25% MS analysed full RD in 2017</td>
<td>Analytical capability poor&lt;br&gt; Not clear if findings justify inclusion in EU MACP&lt;br&gt; Keep 1 extra year in chapter 4 WD. Relevant for muscle, liver, kidney and cow’s milk. It can be used on feed crops.</td>
</tr>
<tr>
<td>Chloromequat – AO</td>
<td>10/2018</td>
<td>ADI 0.04 mg/kg bw/day, ARfD 0.09 mg/kg bw</td>
<td>ADI 0.04 mg/kg bw/day, ARfD 0.09 mg/kg bw</td>
<td>SRM</td>
<td>2A</td>
<td>after 2 years (10/2017)</td>
<td>0% findings EFSA 2012 (2 samples)&lt;br&gt; 0% findings EFSA 2013 (100 samples)&lt;br&gt; N.D. EFSA 2014 (93 samples)&lt;br&gt; N.D. EFSA 2016 report (11 samples)&lt;br&gt; N.D. EFSA 2016 preliminary report (91 samples)</td>
<td>1% labs and 56% MS analysed full RD in 2015&lt;br&gt; 26% labs and 43% MS analysed full RD in 2016&lt;br&gt; 18% labs and 25% MS analysed full RD in 2017</td>
<td>Analytical capability medium&lt;br&gt; Findings justify inclusion in EU MACP&lt;br&gt; Keep 1 extra year in chapter 4 WD. Relevant for muscle, liver, kidney and cow’s milk. It can be used on feed crops.</td>
</tr>
<tr>
<td>Fluazifop-P – AO</td>
<td>10/2018</td>
<td>ADI=0.01 mg/kg bw/day, ARfD=0.017 mg/kgbw</td>
<td>ADI=0.017 mg/kgbw</td>
<td>SRM (hydrolysis required to cover the full residue definition)</td>
<td>2A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glufosinate-ammonium – AO</td>
<td>10/2018</td>
<td>ADI = 0.021 mg/kg bw, ARfD = 0.021 mg/kg bw</td>
<td>ADI = 0.021 mg/kg bw, ARfD = 0.021 mg/kg bw</td>
<td>SRM</td>
<td>2A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^2\) SRM-compounds are typically analysed on specific commodities so their detection frequencies are typically higher than if they would have been analysed randomly.
### 4.2 Pesticides to be considered for analysis in products of animal origin

**Priority: 2A**

Evaluation after 2 years (10/2017) → 10/2018 → 10/2019

- 0% findings EFSA 2012 (148 samples)
- 0% findings EFSA 2013
- 1.03% findings (0.51%MRL exceedances) EFSA 2014
- N.D. EFSA 2015 report (54 samples)
- N.D. EFSA 2016 preliminary report (953 samples)

12% labs and 40% MS analysed full RD in 2015
10% labs and 32% MS analysed full RD in 2016
3% labs and 0% MS analysed full RD in 2017

⇒ **Analytical capability poor**
⇒ **Not clear if findings justify inclusion in EU MACP**
⇒ **Keep 1 extra year in chapter 4 of WD.**

Relevant for animal fat, liver, kidney, eggs, cows’ milk and butter.

### Evaluation after 2 years (10/2017) → 10/2018 → 10/2019

- No monitoring results available in EFSA 2012 report
- No findings EFSA 2013 report (30 samples)
- N.D. EFSA 2014 (31 samples)
- N.D. EFSA 2015 (11 samples)
- N.D. EFSA 2016 (46 samples)

20% labs and 52% MS analysed full RD in 2015
25% labs and 56% MS analysed full RD in 2016
13% labs and 21% MS analysed full RD in 2017

⇒ **Analytical capability poor**
⇒ **Not clear if findings justify inclusion in EU MACP**
⇒ **Keep 1 extra year in chapter 4 WD.**

Relevant for ruminant’s muscle and fat, liver, kidney and cow’s milk.

### Mepiquat – AO

Toxicity: ADI = 0.2 mg/kg bw/day, ARfD = 0.3 mg/kg bw
Method: SRM

**Priority: 2B**

Evaluation after 2 years (10/2017) → 10/2018 → 10/2019

- No monitoring results available in EFSA 2012 report
- 0% findings EFSA 2013 report (30 samples)
- N.D. EFSA 2014 (31 samples)
- N.D. EFSA 2015 (11 samples)
- N.D. EFSA 2016 (46 samples)

20% labs and 52% MS analysed full RD in 2015
25% labs and 56% MS analysed full RD in 2016
13% labs and 21% MS analysed full RD in 2017

⇒ **Analytical capability poor**
⇒ **Not clear if findings justify inclusion in EU MACP**
⇒ **Keep 1 extra year in chapter 4 WD.**

Relevant for ruminant’s muscle and fat, liver, kidney and cow’s milk.
### 4.2 Pesticides to be considered for analysis in products of animal origin

#### 4.2.2. Recently approved

<table>
<thead>
<tr>
<th>Substance</th>
<th>Status</th>
<th>Toxicity: ADI = 0.13 mg/kg bw/day, ARfD = 0.3 mg/kg bw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fenpyrazamine – AO</td>
<td>Approved since 01/2013</td>
<td>Method: MRM</td>
</tr>
<tr>
<td>Priority: 1B</td>
<td>Evaluation: after 1 year (10/2017) → 10/2018 → 10/2019</td>
<td>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.5 mg/kg bw</td>
</tr>
<tr>
<td>14.3% labs and 36% MS analysed full RD in 2015</td>
<td>6% labs and 20% MS analysed full RD in 2015</td>
<td></td>
</tr>
<tr>
<td>17.3% labs and 44% MS analysed full RD in 2016</td>
<td>8.6% labs and 24% MS analysed full RD in 2016</td>
<td></td>
</tr>
<tr>
<td>21% labs and 36% MS analysed full RD in 2017</td>
<td>15% labs and 29% MS analysed full RD in 2017</td>
<td></td>
</tr>
<tr>
<td>⇒ Analytical capability poor</td>
<td>⇒ Analytical capability poor</td>
<td></td>
</tr>
<tr>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td></td>
</tr>
<tr>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td></td>
</tr>
</tbody>
</table>

This substance is not expected to leave significant residues in food of animal origin.

<table>
<thead>
<tr>
<th>Substance</th>
<th>Status</th>
<th>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.25 mg/kgbw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penflufen – AO</td>
<td>Approved since 02/2014</td>
<td>Method: MRM</td>
</tr>
<tr>
<td>Priority: 1A</td>
<td>Evaluation: after 1 year (10/2017) → 10/2018 → 10/2019</td>
<td>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.25 mg/kgbw</td>
</tr>
<tr>
<td>3.6% labs and 12% MS analysed full RD in 2015</td>
<td>3.6% labs and 12% MS analysed full RD in 2015</td>
<td></td>
</tr>
<tr>
<td>3.6% labs and 12% MS analysed full RD in 2016</td>
<td>13% labs and 29% MS analysed full RD in 2017</td>
<td></td>
</tr>
<tr>
<td>⇒ Analytical capability poor</td>
<td>⇒ Analytical capability poor</td>
<td></td>
</tr>
<tr>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td></td>
</tr>
<tr>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance</th>
<th>Status</th>
<th>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.25 mg/kgbw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfoxaflor – AO</td>
<td>Approved since 08/2015</td>
<td>Method: MRM</td>
</tr>
<tr>
<td>Priority: 1B</td>
<td>Evaluation: after 1 year (10/2017) → 10/2018 → 10/2019</td>
<td>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.25 mg/kgbw</td>
</tr>
<tr>
<td>3.6% labs and 12% MS analysed full RD in 2015</td>
<td>3.6% labs and 12% MS analysed full RD in 2015</td>
<td></td>
</tr>
<tr>
<td>3.6% labs and 12% MS analysed full RD in 2016</td>
<td>13% labs and 29% MS analysed full RD in 2017</td>
<td></td>
</tr>
<tr>
<td>⇒ Analytical capability poor</td>
<td>⇒ Analytical capability poor</td>
<td></td>
</tr>
<tr>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
<td></td>
</tr>
<tr>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td>⇒ Keep 1 extra year in chapter 4 WD.</td>
<td></td>
</tr>
</tbody>
</table>
4.3. Evaluation

✓ The evaluation of the chapter 4 substances at the end of the specified evaluation period will be done based on the information listed in Annex V.
✓ The data on the number of labs analysing each substance is collected by the EURLs and stored in the EURL data pool.
✓ The data on the number of MRL exceedances and findings is gathered by EFSA as part of data collection for the National Programmes. These results are then be summarised by COM and added to this document.
✓ In the expert group meeting a decision is taken for moving a substance to the MACP, for deletion from the WD (addition to Annex IV for information for Member States) or for an additional evaluation period in the working document.

5. Proposals for inclusion of new substances in the working document

COM, EFSA, the EURLs and the Member States can put forward substances to be included in the working document by filling out the form in Annex VI. The proposal for inclusion of new substances should be sent to COM by June, prior to the annual expert group meeting on pesticides residues monitoring. During this meeting the submitted proposals will be discussed.

6. Procedure for development of the document

- During the SCOFCAH of 12-13 June 2014 it was decided to develop this document according to an approach in which the pesticides are divided into specific categories. Based on a limited set of criteria each pesticide is attributed a priority and a time line for evaluation of inclusion or non-inclusion in the MACP.
- In Rev.2 of this Working Document this approach was implemented. Details on the substances, criteria, priorities and timelines were discussed in the expert meeting on monitoring on 10 October 2014.
- COM included the decisions taken in the expert group in Rev.3 of this document. In Rev.4 and 5 additional comments from MS experts and the EURLs were taken into account. During the PAFF Committee of 24-25 November 2014 the Member States took note of Rev 5(3).
- Rev 5(3) was applicable to samples analysed in 2015.
- By June 2015 COM, EFSA, the EURLs and Member States could send a proposal to COM for new substances to be included in the working document.
- In October 2015 new substances that were proposed for inclusion in the working document were discussed in the expert group.
- By June 2016 COM, EFSA, the EURLs and Member States could send a proposal to COM for new substances to be included in the working document.
- By August 2016, the EURLs gathered through a survey the information on the % of labs analysing each substance (2015 analyses). By that time the Member States could also submit to EFSA the monitoring data for those substances for which the evaluation timing was set for 10/2016. EFSA summarised these data for the October/November expert group.
In October/November 2016 decisions were taken in the expert group on which chapter 4 substances to move to the MACP 2018, which ones to be deleted from the WD, which ones to be evaluated for an additional period. During this meeting also new substances that were proposed for inclusion in the working document were discussed.

By June 2017 COM, EFSA, the EURs and Member States could send a proposal to COM for new substances to be included in the working document.

By August 2017, the EURs gathered, through a survey, the information on % of labs analysing each substance (2016 analyses). By that time the Member States could also submit to EFSA the monitoring data for those substances for which the evaluation timing was set for 10/2017. EFSA summarised these data for the October/November expert group.

During the Standing Committee on Plants, Animals, Food and Feed (PAFF) – section Residues of 21-22 November 2017, the Member States took note of the Rev 9(1) of the document.

By June 2018 COM, EFSA, the EURs and Member States could have sent a proposal to COM for new substances to be included in the working document.

By October 2018, the EURs will gather through a survey the information on % of labs analysing each substance (2017 analyses). By that time the Member States will also submit to EFSA the monitoring data for those substances for which the evaluation timing was set for 10/2018. EFSA will summarise these data for the October expert group.

In October 2018, decisions were taken in the expert group on which chapter 4 substances to move to the MACP 2020, which ones to be deleted from the WD and which ones to be evaluated for an additional period. During this meeting also new substances were proposed for inclusion in the working document.
Annex I: Substances for which information on residues is needed for specific risk management questions.

Monitoring data for these substances could be used for answering specific risk management questions. These substances are for the time being no candidates for uptake in the MACP.

- Anthraquinone, especially relevant for tea, dried herbs and dried spices.
- Benzalkonium chloride\(^3\)
- Chlorates\(^4\)
- Didecyldimethylammonium chloride\(^5\)
- Glyphosate in soybean
- Nicotine, especially relevant in goji berries (small fruits and berries), mushrooms, tea, chives, brassica crops, moringa. ARfD exceedances reported.

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\(^3\) The results should be reported as mixture of alkylbenzyldimethylammonium chlorides with alkyl chain lengths of C8, C10, C12, C14, C16 and C18.

\(^4\) The results for chlorates (including Mg, Na and K chlorates), should be expressed as chlorate.

\(^5\) The results should be reported as mixture of alkyl-quaternary ammonium salts with alkyl chain lengths of C8, C10 and C12.
Annex II: Substances for which analytical support is requested from the EURLs

For the substances listed in this Annex, support is needed from the EURLs because no validated analytical method and/or no standards are available. To be checked and updated with EURLs.

Substances relevant for plant origin commodities.

(a) Support required due to residue definition

<table>
<thead>
<tr>
<th>Substance</th>
<th>Support Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bifenazate – PO</td>
<td>Toxicity: ADI = 0.01 mg/kg bw/day, ARfD NA No validated method available for the full residue definition (applicable from 19/08/2014). Method: MRM/SRM ✓ 0.3% findings in vegetables (EFSA 2011 report) ✓ 0.24% findings EFSA 2012 report (parent) ✓ 0.29% findings EFSA 2013 report (parent) ✓ 0.30% findings EFSA 2014 report ✓ 0.17% findings EFSA 2015 report ✓ 0.24% findings EFSA 2016 report 7% labs and 23% MS analysed full RD in 2016 54% labs and 71% MS analysed full RD in 2017</td>
</tr>
</tbody>
</table>

| Cyflufenamid – PO | Standard is now available. Support needed to improve analytical capability. Toxicological, occurrence and laboratory capability data in §4.1.1 |

Desmethyl-chlorpyrifos-methyl – PO

EFSA investigated the metabolism of chlorpyrifos-methyl in post-harvest treatment in cereals. Desmethyl-chlorpyrifos-methyl was observed as a significant metabolite as a result of degradation of the parent compound under standard hydrolytic conditions. Toxicological data for desmethyl-chlorpyrifos-methyl are missing and should be provided. EFSA proposed an enforcement residue definition (specific to chlorpyrifos-methyl) which includes the parent compound (in all crops) and its desmethyl metabolite (in cereals and processed commodities only); chlorpyrifos-methyl can be enforced in plant commodities with a limit of quantification (LOQ) of 0.01 mg/kg, while analytical methods are not commercially available for its desmethyl metabolite and should be developed.

Fosetyl-Al – PO

The EURL-SRM has published a method for fosetyl and phosphonic acid (QuPPe). The method is available on-line. Standards are available. The EURL-SRM also provides OfLs isotope labelled standard of phosphonic acid, synthesized at the EURL-SRM. An interlaboratory validation study is planned. Toxicological, occurrence and laboratory capability data in §4.1.1

<table>
<thead>
<tr>
<th>Glufosinate ammonium – PO</th>
<th>Guazatine (not approved) – PO</th>
</tr>
</thead>
<tbody>
<tr>
<td>At request of EFSA, residues are found in animal origin commodities, interesting to also check soybean which is used both as food and feed. The EURL-SRM has published a method for glufosinate, MPPA and N-acetyl glufosinate (QuPPe). The method is available on-line and several labs use it for these compounds. Standards are available. An interlaboratory validation study is planned. Toxicological, occurrence and laboratory capability data in §4.1.1</td>
<td></td>
</tr>
<tr>
<td>The toxicity of guazatine is ADI = 0.0048 mg/kg bw/day, ARfD = 0.04 mg/kg bw Especially relevant for citrus fruits and cereals based on use pattern ✓ No monitoring data EFSA 2012, 2013, 2014, 2015 or 2016</td>
<td></td>
</tr>
</tbody>
</table>
Glyphosate (future residue definition 'sum of glyphosate, AMPA and N-acetylglucosinolate) – PO

In the upcoming Art. 12 review the residue definition for glyphosate will be changed. The EURL-SRM has published a method for glyphosate, N-acetylglucosinolate and AMPA (QuPPe). The method available on-line and many labs use it. An interlaboratory validation is planned. Standards are available

Meptyldinocap (approved since 01/04/2015) – PO

No method available for full residue definition, 2,4-DNOP and 2,4-DNOC standards are available. The EURL-SRM is working on a method for this compound which should be published next year (2018).

Toxicity: ADI = 0.016 mg/kg bw/day, ARfD = 0.12 mg/kg bw

0.04% findings EFSA 2012 report
0.0% findings EFSA 2013 report
0.04% findings EFSA 2014 report
0.00% findings EFSA 2015 report
0.13% findings EFSA 2016 report
9% labs and 29% MS analysed full RD in 2017
Especially relevant for melons, strawberries and table grapes.

Prochloraz – PO

Current SRM method does not cover full RD (possible future revision of residue definition that would allow MRM method).

Nicotine (Not approved) – PO

Toxicological, occurrence and laboratory capability data in §4.1.1

Triclopyr – PO

This substance shares the same metabolites as chlorpyriphos and chlorpyriphos-methyl. For these substances new toxicological studies are available requiring the review of certain MRLs. As these metabolites are not taken up in the current residue definition, method development should only start once the Art. 12 Regulation is voted.

Toxicity: ADI = 0.03 mg/kg bw/day, ARfD = 0.3 mg/kg bw
Method: MRM/SRM, method was developed by the EURL-SRM, the report will be published in the near future.
Relevant for oranges, mandarins, apples, pears
✓ 0.07% findings EFSA 2012 report (parent)
✓ 0.03% findings EFSA 2013 report (parent)
✓ 0.02% findings EFSA 2014 report
✓ 0.06% findings EFSA 2015 report (19604 samples)
✓ 0.03% findings EFSA 2016 report (22614 samples)
42% labs and 77% MS analysed full RD in 2017
43% labs and 79% MS analysed full RD in 2017
Especially relevant for bananas, kiwi, pears, oranges, strawberries and table grapes. Additionally relevant for some non-MACP commodities such as: apricots, mandarins/clementines, lemons, limes and plums.

Tritosulfuron – PO

New residue definition after Art. 12 review: separate MRLs are set for tritosulfuron and 2-amino-4-methoxy-6-(trifluormethyl)-1,3,5-triazine (AMTT).

A method for AMTT has been developed by the EURL-SRM and it is now available on-line. AMTT standard is available.
Toxicity parent: ADI = 0.06 mg/kg bw/day, ARfD NA
Toxicity AMTT: ADI and ARfD 0.0001 mg/kg bw/day
Method: MRM/SRM method for AMTT available
Standard for AMTT is not commercially available.
Especially relevant for rice, wheat, rye and oats
✓ 0% findings EFSA 2012 report
✓ 0% findings EFSA 2013 report
✓ 0% findings EFSA 2014 report (7447 samples)
✓ 0% findings EFSA 2015 report (4160 samples)
✓ 0% findings EFSA 2016 report (7002 samples)
25% labs and 50% MS analysed full RD in 2016
25% labs and 46% MS analysed full RD in 2017

(b) Support required due to other reasons

Benzovindiflupyr – PO
Toxicological, occurrence and laboratory capability data in §4.1.2

Fenpyrazamine – PO
Toxicological, occurrence and laboratory capability data in §4.1.2

Fluensulfone – PO
Not approved in EU, recently approved outside EU
No method available
ADI 0-0.01 mg/kg bw day, ARfD 0.1 mg/kg bw
Relevant commodities: fruiting vegetables

Lambda-cyhalothrin, Gamma-cyhalothrin – PO
Cyhalothrin is not approved in the EU since 1994, hence the default MRL of 0.01* mg/kg applies. It is constituted by four isomers (2 diastereomeric pairs): R.R; R.S; S.R and S.S, as follows:
Lambda-cyhalothrin is a 1:1 mixture of two of the four cyhalothrin components, the R,R and S,R isomers (numbers 1 and 3) and its approval was renewed by Regulation (EU) 2016/146 of 4 February 2016. Gamma-cyhalothrin is constituted by only the most toxic of the four components, the S,R isomer (the third one), which is also contained in lambda-cyhalothrin. As a result, gamma cyhalothrin is twice as toxic as lambda-cyhalothrin and four times more toxic than cyhalothrin. It is an approved active substance under Regulation (EU) 1334/2014 of 16 December 2014.

Following a Commission investigation in September 2016, it was found that most authorisations of gamma-cyhalothrin PPPs in MSs are based on reference to lambda-cyhalothrin, i.e to a less toxic compound of isomers than the actual substance used in the PPPs. Currently no validated analytical methods are available to distinguish between the more toxic residues of gamma-cyhalothrin and the residues of lambda-cyhalothrin, while both share the same residue definition.

As part of the outcome of the discussion held during the SC PAFF of 21-22 September 2017 it was requested that the EURLs would continue their effort to develop a routine method which can discriminate between the two substances.

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**Novaluron (Not approved) – PO**
Toxicological, occurrence and laboratory capability data in §4.1.1. Support needed to improve analytical capability.

**Paraquat – PO**
For the analysis of paraquat in soybean (high fat matrix) it is challenging to enforce the MRL set at the LOQ of 0.02* mg/kg. A method was developed but it does not show the robustness needed. The EURLs are requested to validate a method and to circulate it to the labs. The analysis of paraquat in soyabean is no candidate for the EU MACP. It can be considered for the national programmes.

**Pyridalil – PO**
Toxicological, occurrence and laboratory capability data in §4.1.1

**Pyrethrins – PO**
Toxicological, occurrence and laboratory capability data in §4.1.1

**Pyriofenone – PO**
Approved since 2/2014
Method and standard available in the meanwhile.
Toxicological, occurrence and laboratory capability data in §4.1.2

**Sulfoxaflor – PO**
Toxicological, occurrence and laboratory capability data in §4.1.1. Support needed to improve analytical capability.

**Spinetoram – PO**
Toxicological, occurrence and laboratory capability data in §4.1.1
### Substances relevant for animal origin commodities

**(a) Support required due to residue definition**

<table>
<thead>
<tr>
<th>Substance</th>
<th>AO</th>
<th>Toxicity</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Boscalid</strong> – AO</td>
<td>No method available for the full AO residue definition, standard M510F01 is commercially available, but the development of an analytical method is pending.</td>
<td>Toxicity: ADI = 0.04 mg/kg bw/day, ARfD NA</td>
<td>✓ 0% findings EFSA 2012 report</td>
<td>Relevant for ruminant's and poultry liver, ruminant's kidney</td>
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<td>✓ 0% findings EFSA 2013 report</td>
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<td>✓ 0.30% findings EFSA 2014 report</td>
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<td>✓ 0.39% findings EFSA 2015 report</td>
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<td>✓ 0.14% findings EFSA 2016 report</td>
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<tr>
<td><strong>Chlorpropham</strong> – AO</td>
<td>No method available for the full AO residue definition; a method for 4-HAS and its validation are pending (not needed for the analysis of code 1016000 (poultry) and 1030000 (eggs).</td>
<td>Toxicity: ADI = 0.05 mg/kg bw/day, ARfD = 0.5 mg/kg bw</td>
<td>✓ 0.19% findings EFSA 2012 report</td>
<td>Relevant for ruminant's and swine kidney</td>
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<td>✓ 0% findings EFSA 2013 report</td>
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<td>✓ 0% findings EFSA 2014 report (866 samples)</td>
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<td>✓ 0% findings EFSA 2015 report (502 samples)</td>
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<td></td>
<td>✓ 0% findings EFSA 2016 report (1818 samples)</td>
<td></td>
</tr>
<tr>
<td><strong>Fenpropidin</strong> – AO</td>
<td>No method available for full AO residue definition, standards of 2-methyl-2-[4-(2-methyl-3- piperidin-1-yl-propyl)-phenyl]propionic acid commercially not available</td>
<td>Toxicity: ADI = 0.02 mg/kg bw/day, ARfD = 0.02 mg/kg bw</td>
<td>✓ 0% findings EFSA 2012 report</td>
<td>Relevant for ruminant's and swine liver and kidney</td>
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<td></td>
<td>✓ 0% findings EFSA 2013 report</td>
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<td>✓ 0% finding EFSA 2014 report (356 samples)</td>
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<td></td>
<td>✓ 0% findings EFSA 2015 report (294 samples)</td>
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<td></td>
<td></td>
<td>✓ 0% findings EFSA 2016 report (1016 samples)</td>
<td></td>
</tr>
<tr>
<td><strong>Fenpropimorph</strong> – AO</td>
<td>No validated method available for the full AO residue definition</td>
<td>Method MRM/ SRM. The standard for metabolite fenpropimorph carboxylic acid is now commercially available. Successfull validation at 0.01 mg/kg by EURL-SRM using QuEChERS without PSA cleanup in milk and swine meat. Data publication pending.</td>
<td>Toxicity: ADI = 0.003 mg/kg bw/day, ARfD = 0.03 mg/kg bw</td>
<td>Relevant for ruminant's fat, swine and ruminant's muscle, liver and kidney and cow's milk</td>
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<td>✓ 0% findings EFSA 2012 report (396 sample)</td>
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<td></td>
<td>✓ 0% findings EFSA 2013 report (453 samples)</td>
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<td>✓ 0% findings EFSA 2014 report (238 samples)</td>
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<td></td>
<td>✓ 0% findings EFSA 2015 report (154 samples)</td>
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<td></td>
<td>✓ 0% findings EFSA 2016 report (2064 samples)</td>
<td></td>
</tr>
<tr>
<td><strong>Fluazifop-P</strong> – AO</td>
<td>Method: SRM (hydrolysis required to cover the full residue definition). Toxicalogical, occurrence and laboratory capability data in §4.2.1</td>
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<td></td>
<td>✓ 0% findings EFSA 2012 report</td>
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<td></td>
<td>✓ 0% findings EFSA 2013 report (83 samples)</td>
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<td>✓ 0% findings EFSA 2014 report (173 samples)</td>
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<td>✓ 0% findings EFSA 2015 report (107 samples)</td>
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<td></td>
<td></td>
<td></td>
<td>✓ 0% findings EFSA 2016 report (1138 samples)</td>
<td></td>
</tr>
<tr>
<td><strong>Fluopyram</strong> – AO</td>
<td>No method available for the full AO residue definition.</td>
<td>Toxicity: ADI = 0.012 mg/kg bw/day, ARfD = 0.5 mg/kg bw</td>
<td>✓ 0% findings EFSA 2012 report</td>
<td></td>
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<td>✓ 0% findings EFSA 2013 report</td>
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<td></td>
<td></td>
<td>✓ 0% findings EFSA 2014 report (173 samples)</td>
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<td>✓ 0% findings EFSA 2015 report</td>
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<td></td>
<td>✓ 0% findings EFSA 2016 report</td>
<td></td>
</tr>
</tbody>
</table>
#### Annex II

<table>
<thead>
<tr>
<th>Chemical Name</th>
<th>Method</th>
<th>Toxicological, occurrence and laboratory capability data in §4.2.1</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Glufosinate-ammonium – AO</strong></td>
<td>SRM, but validation is needed for products of animal origin.</td>
<td></td>
<td>Glyphosate (future residue definition 'sum of glyphosate, AMPA and N-acetylglglyphosate) – AO</td>
</tr>
<tr>
<td><strong>Haloxypin – AO</strong></td>
<td>SRM (hydrolysis required to cover conjugates). Method for food of animal origin (including conjugates) is pending.</td>
<td></td>
<td>Ioxynil – AO</td>
</tr>
<tr>
<td><strong>Spioxamine – AO</strong></td>
<td>No method available for full AO residue definition, standard spioxamine carboxylic acid is commercially not available. Toxicity: ADI = 0.025 mg/kg bw/day, ARfD = 0.1 mg/kgbw</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Aminocyclopyrachlor – AO</strong></td>
<td>Not approved in EU, recently approved outside EU ADI 0-3 mg/kg bw day, ARfD N/A Standard commercially available. Successfully validated by EUROL-SRM using QuPPe in food of plant origin. Validations in products of animal origin are pending. Relevant commodities animal fat, milk, liver and kidney.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Carbendazim and Thiophanate methyl – AO</strong></td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.1.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fenpyrazamine – AO</strong></td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.2.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Maleic hydrazide – AO</strong></td>
<td>Method: SRM. QuPPe amenable but validation is needed for products of animal origin. Toxicity: ADI = 0.25 mg/kg bw/day, ARfD NA Priority: 2B Evaluation after 2 years (10/2017)</td>
<td>10/2018</td>
<td>10% labs and 28% MS analysed full RD in 2015 12% labs and 36% MS analysed full RD in 2016 6% labs and 14% MS analysed full RD in 2017 Relevant for all commodities of animal origin.</td>
</tr>
<tr>
<td><strong>Benzovindiflupyr – AO</strong></td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.2.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Chloromequat – AO</strong></td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.1.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chemical</td>
<td>AO</td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.1.</td>
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</tr>
<tr>
<td>Mepiquat</td>
<td>AO</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Penflufen</td>
<td>AO</td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.2.</td>
<td></td>
</tr>
<tr>
<td>Penthiopyrad</td>
<td>AO</td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.2.</td>
<td></td>
</tr>
<tr>
<td>Sulfoxaflor</td>
<td>AO</td>
<td>Toxicological, occurrence and laboratory capability data in §4.2.2.</td>
<td></td>
</tr>
</tbody>
</table>
Annex III: Substances that are of interest for cumulative risk assessment

EFSA is currently establishing common assessment groups for cumulative risk assessment. In order to have sufficient data to calculate the background exposure, monitoring results would be needed for compounds from the acute neurotoxicity group, the chronic neurotoxicity group and the thyroid group. Some of these pesticides are not taken up in the MACP or in chapter 4 of this document that lists pesticides that could be considered for future uptake in the MACP. However, since monitoring data for these substances would be of interest for the further development of the CRA methodology, they are listed in this annex, for information only.

- 2,4-DB (especially relevant for citrus fruits and pome fruits. Additionally relevant for the non-MACP commodity: chamomile)
- Amitrole
- Cyhalofop-butyl (especially relevant for rice)
- Dazomet
- Flufenacet (especially relevant for beans with pods, grapes, potatoes, rye, oats, strawberries, leek, lettuce, wheat, cucumber and rice. Additionally relevant for several non-MACP commodities such as: celeriac, chives, currants, dill, fennel, raspberries, parsley, strawberries)
- Glufosinate ammonium (especially relevant for potatoes, strawberries and rice. Additionally relevant for several non-MACP commodities such as: berries, tea)
- Ioxynil (especially relevant for cereals, leek, lettuce, tomatoes. Additionally relevant for the non-MACP commodity: chives and dill)
- Isoxaflutole
- MCPA and MCPB (especially relevant for aubergines, cultivated fungi, head cabbage, table grapes, lettuce, peaches, wheat, rye and strawberries. Additionally relevant for several non-MACP commodities such as: Chamomile, berries, cherries, mint, thyme, lentils, tea)
- Milbemectin (this substance has two isomers A3 and A4 of 1920 £ each, relevant for strawberries)
- Metconazole
- Molinate
- Oxadiargyl
- Oxasulfuron
- Oxyfluorfen
- Picolinafen
- Propaquizafop
- Pyridate (especially relevant for grapefruit, oranges, sweet pepper. Additionally relevant for several non-MACP commodities such as: avocado, Brussel's sprouts, celery, dill, leek, mandarins and tea) (SRM method, support EURLs needed)
- Quinoclamidine
- Quizalofop, including quizalfop-P (especially relevant for carrots, head cabbage, spinach, broccoli, spinach and potatoes. Additionally relevant for several non-MACP commodities such as: celeriac, parsley, coriander, caraway, fennel, dill, herbs (balm, basil, mint, thyme); beet, chard, artichoke, chicory)
- Sulfuryl fluoride (especially relevant for nuts, oilseeds and dried fruit)
- Tri-allate
Annex IV: Substances with a low level of findings

This annex contains substances for which few residues were detected during their evaluation under chapter 4. They were moved to this annex for information of the Member States that are interested of keeping them in their National Programmes as most of them are analysed by a large fraction of laboratories and Member States.

Pesticides relevant to products of plant origin

Previously listed in Chapter 4.1.1 (Frequent detections, MRL exceedances or RASFF notifications)

<table>
<thead>
<tr>
<th>Substance</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Found in EFSA Reports</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amitraz (Not approved) – PO</td>
<td>2A</td>
<td>10/2018</td>
<td>0.03% findings 2012 EFSA report, 0.27% findings EFSA 2013 report, 0.09% findings (0.01% MRL exceedances) EFSA 2014, 0.06% findings (0.04% MRL exceedances) EFSA 2015, 0.05% findings (0.03% MRL exceedances) EFSA 2016</td>
</tr>
<tr>
<td>Benalaxyl including other mixtures of constituent isomers including benalaxyl-M – PO</td>
<td>1A</td>
<td>10/2017</td>
<td>0.1% findings in vegetables EFSA 2011 report, 0.05% findings EFSA 2012 report, 0.02% findings EFSA 2013 report, 0.02% findings EFSA 2014 report, 0.04% findings, 0.00% MRL exceedances EFSA 2015</td>
</tr>
<tr>
<td>Chlorfluazuron (Not approved) – PO</td>
<td>1B</td>
<td>10/2016</td>
<td>0.01% findings (0.01% MRL exceedances) EFSA 2013, 0.09% findings (0.09% MRL exceedances) EFSA 2014, 0.01% findings (0.02% MRL exceedances) EFSA 2015, 0.00% findings (0.02% MRL exceedances) EFSA 2016</td>
</tr>
<tr>
<td>Clomazone – PO</td>
<td>1B</td>
<td>10/2016</td>
<td>0.01% findings (0.01% MRL exceedances) EFSA 2013, 0.09% findings (0.09% MRL exceedances) EFSA 2014, 0.01% findings (0.02% MRL exceedances) EFSA 2015, 0.00% findings (0.02% MRL exceedances) EFSA 2016</td>
</tr>
<tr>
<td>Heptachlor (Not approved) – PO</td>
<td>1A</td>
<td>10/2016</td>
<td>0.3% findings in animal commodities, 0.1% in vegetables (EFSA 2011 report), 0.06% findings EFSA 2012 report, 0.05% findings EFSA 2013 report, 0.02% findings EFSA 2014 report, 0.01% findings, 0.00% MRL exceedances EFSA 2015</td>
</tr>
<tr>
<td>Quintozene (Not approved) – PO</td>
<td>1A</td>
<td>10/2016</td>
<td>0.01% findings, 0.00% MRL exceedances</td>
</tr>
</tbody>
</table>
67% labs and 92% MS analysed full RD in 2015  
⇒ Analytical capability good  
⇒ Few findings

Tetramethrin (Not approved) – PO

Toxicity: no toxicological reference values available
Method: MRM
Priority: 1B
Evaluation after 1 year (10/2016) ⇒ 10/2018
✓ 0.02% findings EFSA 2012 report
✓ 0.02% findings EFSA 2013 report
✓ 0.04% findings (0.01% MRL exceedances) EFSA 2014
✓ 0.00% findings (0.01% MRL exceedances) EFSA 2015
✓ 0.01% findings (0.01% MRL exceedances) EFSA 2016
68% labs and 92% MS analysed full RD in 2015
⇒ Low findings
⇒ Good analytical capability

Previously listed in Chapter 4.1.2 (Recently Approved)

Fluxapyroxad – PO

Approved since 1/2013
Method: MRM
Toxicity: ADI = 0.02 mg/kg bw/day, ARfD = 0.25 mg/kg bw
Priority: 1A
Evaluation: after 1 year (10/2016, extended to 10/2017)
✓ 0% findings EFSA 2012 report
✓ 0.12% findings EFSA 2013 report
✓ 0.01% findings EFSA 2014 report
✓ 0.04% findings (0.01% MRL exceedances) EFSA 2015 report (19016 samples)
✓ 0.01% findings (0.00% MRL exceedances) EFSA 2016 report (21906 samples)
42% labs and 85% MS analysed full RD in 2015
45% labs and 81% MS analysed full RD in 2016
⇒ Findings don’t justify inclusion in EU MACP
⇒ Medium analytical capability

Isopyrazam – PO

Approved since 4/2013
Method: MRM
Toxicity: ADI = 0.03 mg/kg bw/day, ARfD = 0.2 mg/kg bw
Priority: 1A
Evaluation: after 1 year (10/2016) extended with an extra year (10/2017)
✓ No monitoring results EFSA 2012 report
✓ 0% findings EFSA 2013 report (473 samples)
✓ 0% findings EFSA 2014 report
✓ 0.04% findings (0.00% MRL exceedances) EFSA 2015 report (2668 samples)
✓ 0.05% findings (0.00% MRL exceedances) EFSA 2016 report (6568 samples)
27% labs and 69% MS analysed full RD in 2015
42% labs and 73% MS analysed full RD in 2016
⇒ Analytical capability medium
⇒ Findings don’t justify inclusion in EU MACP

Penflufen – PO

Approved since 02/2014
Toxicity: ADI = 0.04 mg/kg bw/day, ARfD = 0.5 mg/kg bw
Method: MRM
Priority: 1A
Evaluation: after 1 year (10/2017) ⇒ 10/2018
✓ No monitoring data available EFSA 2012, 2013 or 2014
✓ N.D. EFSA 2015, 2016 (4161 samples)
14% labs and 46% MS analysed full RD in 2015
26% labs and 65% MS analysed full RD in 2016
33% labs and 57% MS analysed full RD in 2017
⇒ Analytical capability poor
⇒ Low findings

Penthiopyrad – PO

Approved since 5/2014
Method: MRM
Toxicity: ADI = 0.1 mg/kg bw/day, ARfD = 0.75 mg/kg bw
Priority: 1B
Evaluation: after 1 year (10/2017)
✓ No monitoring data available EFSA 2012 report
✓ No monitoring data available EFSA 2013 report
✓ 0.08% findings EFSA 2014 report
✓ 0.04% findings (0.00% MRL exceedances) EFSA 2015 report (2595 samples)
✓ 0.06% findings (0.00% MRL exceedances) EFSA 2016 report (8298 samples)
19% labs and 50% MS analysed full RD in 2015
40% labs and 77% MS analysed full RD in 2016
⇒ Analytical capability medium
⇒ Findings don’t justify inclusion in EU MACP
### Previously listed in Chapter 4.1.4 (High toxicity)

**Ethoprophos – PO**

| Toxicity: ADI = 0.0004 mg/kg bw/day, ARfD = 0.01 mg/kgbw | Method: MRM |
| Priority: 1A | |
| Evaluation: after 1 year (10/2016) | 0.01% findings EFSA 2012 report |
| 0.02% findings EFSA 2013 report | 0.01% findings EFSA 2014 report |
| 0.01% findings, 0.00% MRL exceedances 2015 EFSA | 83% labs and 100% MS analysed full RD in 2015 |
| EURL comment: a lot of laboratories use this as an internal standard. If there are significant findings then this practice is called into question. Also this compound is unstable in protic solvents and therefore is unlikely to be found | ⇒ Analytical capability good |
| ⇒ Few findings |

### Previously listed in Chapter 4.1.5 (Voluntary in Reg. (EU) N° 788/2012)

**Phenthoate (Not approved) – PO**

| Footnote i) in Reg. (EC) N° 788/2012 |
| Method: MRM |
| Toxicity: ADI = 0.003 mg/kg bw/day, ARfD NA |
| Priority: 1A |
| Evaluation after 1 year (10/2016) | 0.01% findings EFSA 2012 report |
| 0% findings EFSA 2013 report | 0.03% findings EFSA 2014 report |
| 0.01% findings, 0.00% MRL exceedances 2015 EFSA | 78% labs and 100% MS analysed full RD in 2015 |
| ⇒ Analytical capability good |
| ⇒ Few findings |

**Prothiofos (Not approved) – PO**

| Footnote g) in Reg. (EC) N° 788/2012 |
| Method: MRM |
| Toxicity: no ADI or ARfD available in database |
| Priority: 1B |
| Evaluation after 1 year (10/2016) | 0.01% findings EFSA 2012 report |
| 0.01% findings EFSA 2013 report | 0.01% findings EFSA 2014 report |
| 0.01% findings, 0.00% MRL exceedances 2015 EFSA | 66% labs and 96% MS analysed full RD in 2015 |
| ⇒ Low findings |
| ⇒ Substance mainly of interest for imported commodities |
| ⇒ Good analytical capability |

**Rotenone (Not approved) – PO**

| Footnote g) in Reg. (EC) N° 788/2012 |
| Method: MRM |
| Toxicity: no ADI or ARfD in database |
| Priority: 1B |
| Evaluation after 1 year (10/2016) | 0% findings EFSA 2012 report |
| 0% findings EFSA 2013 report | 0.01% findings EFSA 2014 report |
| 0.01% findings, 0.00% MRL exceedances 2015 EFSA | 50% labs and 89% MS analysed full RD in 2015 |
| ⇒ Low findings |
| ⇒ Medium analytical capability |

**Triticonazole – PO**

| Footnote i) in Reg. (EC) N° 788/2012 |
| Method: MRM |
| Toxicity ADI = 0.025 mg/kg bw/day, ARfD = 0.05 mg/kg bw |
| Priority: 1A |
| Evaluation after 1 year (10/2016) | 0% findings EFSA 2012 report |
| 0% findings EFSA 2013 report | 0.02% findings EFSA 2014 report |
| 0.01% findings, 0.01% MRL exceedances 2015 EFSA | 77% labs and 100% MS analysed full RD in 2015 |
| ⇒ Low findings |
| ⇒ Good analytical capability |
Pesticides for analysis in products of animal origin

Previously listed in Chapter 4.2.1 (Frequent detections, MRL exceedances or RASFF notification)

<table>
<thead>
<tr>
<th>Pesticide</th>
<th>Method</th>
<th>Toxicity</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azinphos ethyl (Not approved) – AO</td>
<td>MRM</td>
<td>no toxicological information available</td>
<td>1B</td>
<td>after 1 year (10/2017)</td>
<td>0% findings EFSA 2012 report</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td>0.12% findings EFSA 2013 report</td>
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<td>0% findings EFSA 2014 report</td>
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<td></td>
<td>0.00% findings (0.00% MRL exceedances) EFSA 2015 report (73 samples)</td>
</tr>
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<td>0.00% findings (0.00% MRL exceedances) EFSA report (2092 samples)</td>
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<td></td>
<td>62% labs and 92% MS analysed full RD in 2015</td>
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<td></td>
<td>⇒ Analytical capability good</td>
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<td></td>
<td></td>
<td>⇒ No findings</td>
</tr>
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<td></td>
<td>Relevant for animal muscle and fat.</td>
</tr>
<tr>
<td>Haloxyfop – AO</td>
<td>SRM (hydrolysis</td>
<td>ADI=0.00065 mg/kg bw/day, ARfD=0.075</td>
<td>2A</td>
<td>after 2 years (10/2017) → 10/2018</td>
<td>0% findings EFSA 2012 report</td>
</tr>
<tr>
<td></td>
<td>required to</td>
<td>mg/kg bw</td>
<td></td>
<td></td>
<td>0% findings EFSA 2013 report (171 samples)</td>
</tr>
<tr>
<td></td>
<td>cover conjugates</td>
<td></td>
<td></td>
<td></td>
<td>0% findings EFSA 2014 report (258 samples)</td>
</tr>
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<td>N.D EFSA 2015 (16 samples)</td>
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<td>N.D EFSA 2016 (708 samples)</td>
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<td>14% labs and 40% MS analysed full RD in 2015</td>
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<td>9% labs and 24% MS analysed full RD in 2016</td>
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<td>4% labs and 0% MS analysed full RD in 2017</td>
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<td>⇒ Analytical capability poor</td>
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<td></td>
<td>⇒ No findings</td>
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<td>Relevant for cows’ milk, kidney, liver, butter and poultry fat.</td>
</tr>
<tr>
<td>Ioxynil – AO</td>
<td>SRM</td>
<td>ADI = 0.005 mg/kg bw/day, ARfD 0.04 mg/kg bw</td>
<td>2A</td>
<td>after 2 years (10/2017) → 10/2018</td>
<td>No monitoring results available in EFSA 2012 report</td>
</tr>
<tr>
<td></td>
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<td>0% findings EFSA 2013 report (177 samples)</td>
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<td>0% findings EFSA 2014 report (563 samples)</td>
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<td>N.D EFSA 2015 report (21 samples)</td>
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<td>N.D EFSA 2015 report (44 samples)</td>
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<td>4% labs and 12% MS analysed full RD in 2015</td>
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<td>6% labs and 16% MS analysed full RD in 2016</td>
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<td>3% labs and 7% MS analysed full RD in 2017</td>
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<td></td>
<td>⇒ Analytical capability poor</td>
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<td>⇒ No findings</td>
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<td>Relevant for ruminant fat, muscle, kidney and liver.</td>
</tr>
<tr>
<td>Benzovindiflupyr – AO</td>
<td>MRM</td>
<td>ADI 0-0.05 mg/kg bw day, ARfD 0.1 mg/kg bw</td>
<td>1A</td>
<td>after 1 year (10/2017) → 10/2018</td>
<td>No EFSA monitoring data for 2012, 2013, 2014, 2015, 2016.</td>
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<td></td>
<td>0% labs and 0% MS analysed full RD in 2015</td>
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<td>4.9% labs and 16% MS analysed full RD in 2016</td>
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<td>13% labs and 29% MS analysed full RD in 2017</td>
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<td>⇒ Analytical capability poor</td>
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<td>⇒ Not clear if findings justify inclusion in EU MACP</td>
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<td>⇒ Already kept in chapter 4 of WD for an extra year.</td>
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<td>Relevant for animal fat and liver.</td>
</tr>
<tr>
<td>Bixafen – AO</td>
<td>MRM</td>
<td>ADI = 0.02 mg/kg bw/day, ARfD = 0.2 mg/kg bw</td>
<td>1A</td>
<td>after 1 year (10/2017)</td>
<td>0% findings EFSA 2012 report</td>
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<tr>
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<td></td>
<td></td>
<td>0% findings EFSA 2013 report (527 samples)</td>
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<td>0% findings EFSA 2014 report (480 samples)</td>
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<td>0.00% findings (0.00% MRL exceedances) EFSA 2015 report (22854 samples)</td>
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<td>0.00% findings (0.00% MRL exceedances) EFSA 2016 report (104 samples)</td>
</tr>
<tr>
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<td>0% labs and 0% MS analysed full RD in 2015</td>
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<td>1% labs and 4% MS analysed full RD in 2016</td>
</tr>
</tbody>
</table>

Previously listed in Chapter 4.2.3 (Voluntary in Reg. (EU) N° 788/2012)
### Chlorobenzilate (not approved) – AO

Footnotes g) and i) in Reg. (EC) N° 788/2012.

- **Method:** MRM
- **Toxicity:** ADI = 0.02 mg/kg bw/day, ARfD NA
- **Priority:** 1A
- **Evaluation after 1 year (10/2016)**
  - ✔ 0.96% findings EFSA 2012 report
  - ✔ 0.03% findings EFSA 2013 report
  - ✔ 0.05% findings EFSA 2014 report
  - ✔ 0.00% findings, 0.00% MRL exceedances 2015 EFSA
  - 55% labs and 84% MS analysed full RD in 2015

*Relevant for animal fat, milk and eggs.*

- **⇒ Analytical capability medium**
- **⇒ Findings don’t justify inclusion in EU MACP**

### Cyfluthrin – AO

Footnote i) in Reg. (EC) N° 788/2012

- **Method:** MRM
- **Toxicity:** ADI = 0.003 mg/kg bw/day, ARfD = 0.02 mg/kg bw
- **Priority:** 1A
- **Evaluation after 1 year (10/2016)**
  - ✔ 0% findings EFSA 2012 report
  - ✔ 0% findings EFSA 2013 report (3531 samples)
  - ✔ 0% findings EFSA 2014 report (4189 samples)
  - ✔ 0.00% findings, 0.00% MRL exceedances 2015 EFSA
  - 82% labs and 96% MS analysed full RD in 2015

*Relevant for animal fat.*

- **⇒ Analytical capability good**
- **⇒ No findings**

### Cyproconazole – AO

No footnote, remark in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in liver (2014), it does not need to be analysed in poultry meat (2014). Not relevant for commodities listed in 2013/2015.’

- **Method:** MRM
- **Toxicity:** ADI = 0.02 mg/kg bw/day, ARfD = 0.02 mg/kg bw
- **Priority:** 1A
- **Evaluation after 1 year (10/2016)**
  - ✔ 0% findings EFSA 2012 report
  - ✔ 0% findings EFSA 2013 report (902 samples)
  - ✔ 0% findings EFSA 2014 report (2164 samples)
  - ✔ 0.00% findings, 0.00% MRL exceedances 2015 EFSA data
  - 46% labs and 76% MS analysed full RD in 2015

*Relevant for liver.*

- **⇒ Analytical capability medium**
- **⇒ No findings**

### Dichlorprop (Not approved) – AO

No footnote, remark in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in liver (2014), it does not need to be analysed in poultry meat (2014). Not relevant for commodities listed in 2013/2015.’

- **Method:** SRM (hydrolysis required to cover conjugates)
- **Toxicity:** no ADI or ARfD in COM database, non-approved substance
- **Priority:** 2B
- **Evaluation after 2 years (10/2017)**
  - ✔ 0% findings EFSA 2012 report (124 samples)
  - ✔ 0% findings EFSA 2013 report (234 samples)
  - ✔ 0% findings EFSA 2014 report (531 samples)
  - ✔ 0.00% findings (0.00% MRL exceedances) EFSA 2015 report (53 samples)
  - ✔ 0.00% findings (0.00% MRL exceedances) EFSA 2016 report (111 samples)
  - 16% labs and 40% MS analysed full RD in 2015
  - 27% labs and 44% MS analysed full RD in 2016

- **⇒ Analytical capability poor**
- **⇒ No findings**
  - Relevant for liver and kidney.

### Epoxiconazole – AO

No footnote, remark in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in liver (2014), it does not need to be analysed in poultry meat (2014). Not relevant for commodities listed in 2013/2015.’

- **Method:** MRM
- **Toxicity:** ADI = 0.008 mg/kg bw/day, ARfD = 0.023 mg/kg bw
- **Priority:** 1A
- **Evaluation after 1 year (10/2016)**
  - ✔ 0% findings EFSA 2012 report
  - ✔ 0% findings EFSA 2013 report
  - ✔ 0% findings EFSA 2014 report

*⇒ Analytical capability medium*  
*⇒ No findings*  

### Etofenprox – AO


- **Method:** MRM
- **Toxicity:** ADI = 0.03 mg/kg bw/day, ARfD = 1 mg/kg bw
- **Priority:** 1A
- **Evaluation after 1 year (10/2016)**
  - ✔ 0% findings EFSA 2012 report
Annex IV

- 0 % findings EFSA 2012 report
- 0 % findings EFSA 2013 report
- 0 % findings EFSA 2014 report
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA data

Relevant for liver
⇒ Analytical capability medium
⇒ No findings

Fenthion (Not approved) – AO

Footnote i) in Reg. (EC) N° 788/2012
Method: MRM
Toxicity: ADI = 0.007 mg/kg bw/day, ARfD = 0.01 mg/kg bw
Priority: 1A
Evaluation after 1 year (10/2016)
- 0 % findings EFSA 2012 report
- 0 % findings EFSA 2013 report (2260 samples)
- 0 % findings EFSA 2014 report (3598 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
31% labs and 76% MS analysed full RD in 2015
Relevant for animal fat, cows’ milk and butter.
⇒ Analytical capability medium
⇒ No findings

Fluquinconazole – AO

No footnote, remark h) in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in milk (2013), liver (2014) and butter (2015), it does not need to be analysed in swine meat (2013), poultry meat (2014) and egg (2015).’
Method: MRM
Toxicity: ADI = 0.002 mg/kg bw/day, ARfD = 0.02 mg/kg bw
Priority: 1A
Evaluation after 1 year (10/2016)
- 0.35 % findings EFSA 2012 report
- 0% findings EFSA 2013 report (1280 samples)
- 0% findings EFSA 2014 report (2703 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
48% labs and 76% MS analysed full RD in 2015
Relevant for cows’ milk, liver and butter.
⇒ Analytical capability medium
⇒ No findings

Flusilazole (not approved) – AO

No footnote, remark in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in swine meat (2013) and liver (2014), it does not need to be analysed in milk (2013) and poultry meat (2014). Not relevant for commodities listed in 2015.’
Method: MRM
Toxicity: ADI = 0.002 mg/kg bw/day, ARfD = 0.005 mg/kg bw
Priority: 1A
Evaluation after 1 year (10/2016)
- 0 % findings EFSA 2012 report
- 0% findings EFSA 2013 report (222 samples)
- 0% findings EFSA 2014 report (1027 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
1% labs and 4% MS analysed full RD in 2015
Relevant for animal fat, kidney and liver.
⇒ Analytical capability low
⇒ No findings

Metaflumizone – AO

No footnote, remark in Reg. (EC) N° 788/2012: ‘To be analysed on voluntary basis in swine meat (2013), poultry meat (2014) and egg (2015), it does not need to be analysed in milk (2013), liver (2014) and butter (2015).’
Method: MRM
Toxicity: ADI = 0.01 mg/kg bw/day, ARfD = 0.13 mg/kg bw
Priority: 1A
Evaluation after 1 year (10/2016),
- 0 % findings EFSA 2012 report
- 0% findings EFSA 2013 report (669 samples)
- 0% findings EFSA 2014 report (1074 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
31% labs and 72% MS analysed full RD in 2015
Relevant for swine muscle, poultry muscle and eggs.
⇒ Analytical capability low
⇒ No findings

Metazachlor – AO

Footnote h) in Reg. (EC) N° 788/2012 and remark: ‘To be analysed on voluntary basis in liver (2014), it does not need to be analysed in poultry meat (2014). Not relevant for commodities listed in 2013/2015.’
Method: SRM
Toxicity: ADI = 0.08 mg/kg bw/day, ARfD = 0.5 mg/kg bw
Priority: 2A
Evaluation after 2 years (10/2017)
- 0 % findings EFSA 2012 report
- 0% findings EFSA 2013 report (1366 samples)
- 0% findings EFSA 2014 report (1959 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
44% labs and 80% MS analysed full RD in 2015
Relevant for animal fat, cows’ milk and butter.
⇒ Analytical capability medium
⇒ No findings

Methidathion (Not approved) – AO

Footnote i) in Reg. (EC) N° 788/2012
Method: MRM
Toxicity: ADI = 0.001 mg/kg bw/day, ARfD = 0.01 mg/kg bw
Priority: 1A
Evaluation after 1 year (10/2016)
- 0 % findings EFSA 2012 report
- 0% findings EFSA 2013 report (3707 samples)
- 0% findings EFSA 2014 report (4804 samples)
- 0.00% findings, 0.00% MRL exceedances 2015 EFSA
<table>
<thead>
<tr>
<th>Substance</th>
<th>Method</th>
<th>Toxicity</th>
<th>Priority</th>
<th>Evaluation</th>
<th>Findings of EFSA Reports</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parathion-methyl (Not approved) – AO</strong></td>
<td></td>
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<tr>
<td>Footnote i) in Reg. (EC) N° 788/2012</td>
<td>MRM</td>
<td>ADI = 0.003 mg/kg bw/day, ARfD = 0.03 mg/kg bw</td>
<td>1A</td>
<td>After 1 year (10/2016)</td>
<td>0% findings EFSA 2012 report, 0% findings EFSA 2013 report, 0.00% findings EFSA 2015 report, 0.00% findings EFSA 2016 report</td>
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<tr>
<td><strong>Profenofos (Not approved) – AO</strong></td>
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<tr>
<td>Footnote i) in Reg. (EC) N° 788/2012</td>
<td>MRM</td>
<td>ADI = 0.03 mg/kg bw/day, ARfD = 1 mg/kg bw</td>
<td>1A</td>
<td>After 1 year (10/2016)</td>
<td>0% findings EFSA 2012 report, 0% findings EFSA 2013 report, 0.06% findings EFSA 2014 report</td>
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<tr>
<td><strong>Prothioconazole – AO</strong></td>
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<tr>
<td>No footnote, remark in Reg. (EC) N° 788/2012</td>
<td>MRM/ SRM</td>
<td>ADI = 0.01 mg/kg bw/day, ARfD = 0.01 mg/kg bw</td>
<td>2A</td>
<td>After 2 years (10/2017)</td>
<td>0% findings EFSA 2012 report, 0% findings EFSA 2013 report, 0.00% findings EFSA 2014 report</td>
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<tr>
<td><strong>Resmethrin (Not approved) – AO</strong></td>
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<tr>
<td>Footnote i) in Reg. (EC) N° 788/2012</td>
<td>MRM</td>
<td>ADI = 0.03 mg/kg bw/day, ARfD = NA</td>
<td>1A</td>
<td>After 1 year (10/2016)</td>
<td>0% findings EFSA 2012 report, 0% findings EFSA 2013 report, 0.06% findings EFSA 2014 report</td>
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<tr>
<td><strong>Tetraconazole – AO</strong></td>
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<tr>
<td>No footnote, remark in Reg. (EC) N° 788/2012</td>
<td>MRM</td>
<td>ADI = 0.004 mg/kg bw/day, ARfD = 0.05 mg/kg bw</td>
<td>1A</td>
<td>After 1 year (10/2016)</td>
<td>0% findings EFSA 2012 report, 0% findings EFSA 2013 report, 0.00% findings EFSA 2014 report</td>
</tr>
</tbody>
</table>

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**Annex IV**

- 0% findings EFSA 2013 report (701 samples)
- 0% findings EFSA 2014 report (1650 samples)
- 0.00% findings (0.00% MRL exceedances) EFSA 2015 report (821 samples)
- 0.00% findings (0.00% MRL exceedances) EFSA 2016 report (51 samples)

1% labs and 4% MS analysed full RD in 2015
6% labs and 16% MS analysed full RD in 2016

⇒ Analytical capability poor
⇒ No findings

Relevant for liver and kidney of swine and ruminants.

- 70% labs and 92% MS analysed full RD in 2015

⇒ Analytical capability medium
⇒ No findings

Relevant for animal muscle, fat, milk and eggs.

- 0.00% findings, 0.00% MRL exceedances EFSA 2015 report (821 samples)
- 0.00% findings (0.00% MRL exceedances) EFSA 2016 report (51 samples)

⇒ Analytical capability good
⇒ No findings

Relevant for animal fat, milk and eggs.

- 70% labs and 92% MS analysed full RD in 2015

⇒ Analytical capability low
⇒ Few findings

Relevant for animal fat, muscle, liver, kidney, cow's milk and eggs.

- 0% findings EFSA 2013 report (3342 samples)
- 0% findings EFSA 2014 report (4097 samples)

⇒ Analytical capability medium
⇒ No findings

Relevant for animal muscle, fat, milk and eggs.

- 70% labs and 92% MS analysed full RD in 2015

⇒ Analytical capability poor
⇒ No findings

Relevant for ruminant's and swine liver and kidney.

- 0% findings EFSA 2013 report (3342 samples)
- 0.06% findings EFSA 2014 report (3372 samples)

⇒ Analytical capability medium
⇒ No findings

Relevant for animal muscle, fat, milk and eggs.

- 70% labs and 92% MS analysed full RD in 2015

⇒ Analytical capability poor
⇒ No findings

Relevant for liver and kidney of swine and ruminants.
<table>
<thead>
<tr>
<th>Substance</th>
<th>Method</th>
<th>Toxicity: ADI</th>
<th>ARfD</th>
<th>Priority</th>
<th>Evaluation after 1 year</th>
<th>Monitoring Results</th>
<th>Lims Analysis</th>
<th>Relevant for</th>
<th>Analytical Capability</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiacloprid – AO</td>
<td>MRM</td>
<td>ADI = 0.01 mg/kg bw/day</td>
<td>ARfD = 0.03 mg/kg bw</td>
<td>1A</td>
<td>2015 preliminary EFSA data 26.6% findings, 0.5% MRL exceedances in honey. Not tested on other AO commodities.</td>
<td>8% labs and 24% MS analysed full RD in 2015</td>
<td>Relevant for ruminant’s liver and kidney.</td>
<td>Analytical capability low</td>
<td>No findings</td>
<td></td>
</tr>
<tr>
<td>Triazophos (Not approved) – AO</td>
<td>MRM</td>
<td>ADI = 0.001 mg/kg bw/day</td>
<td>ARfD = 0.001 mg/kg bw</td>
<td>1A</td>
<td>2015 preliminary EFSA data 26.6% findings, 0.5% MRL exceedances in honey. Not tested on other AO commodities.</td>
<td>8% labs and 24% MS analysed full RD in 2015</td>
<td>Relevant for animal fat, eggs and milk.</td>
<td>Analytical capability good</td>
<td>No findings</td>
<td></td>
</tr>
<tr>
<td>Topramezone (Approval pending) – AO</td>
<td>MRM</td>
<td>ADI = 0.001 mg/kg bw/day</td>
<td>ARfD = 0.001 mg/kg bw</td>
<td>1A</td>
<td>2015 preliminary EFSA data 26.6% findings, 0.5% MRL exceedances in honey. Not tested on other AO commodities.</td>
<td>8% labs and 24% MS analysed full RD in 2015</td>
<td>Relevant for ruminant's liver and kidney.</td>
<td>Analytical capability low</td>
<td>No findings</td>
<td></td>
</tr>
</tbody>
</table>
Annex V: Evaluation at the end of the evaluation period

Information to be gathered for evaluation at the end of the evaluation period

_Pesticide X_

- Analytical capability (data collection via EURLs)
  - % of labs that took part in the survey
  - % of Member States that took part in the survey
  - % of the labs that is able to analyse the full residue definition
  - % of the labs that analyses part of the residue definition
  - % of the Member States that is able to analyse the full residue definition
  - % of the Member States that analyses part of the residue definition

- MRL exceedances/ findings (data collection by EFSA as part of the data collection for the National Programmes)
  - N° of samples analysed
  - % of samples with findings > LOQ
  - % of samples numerically exceeding the MRL
  - % of samples analysed according to full residue definition (SSD code P005)
  - % of samples analysed for part of the residue definition (SSD code P004)
  - N° of RASFF notifications
  - N° of ARfD exceedances (not systematically calculated by EFSA, only mentioned if specific MS information is available)

Evaluation summarised by COM in Working Document

_Pesticide X_

- % of labs that is able to analyse the full residue definition
- % of samples with residues > MRL
- % of findings
- N° of RASFF notifications
Annex VI: Proposals for uptake of new substances in the Working Document

Proposal sheet to be filled out by COM, EFSA, EURLs or Member States

- Proposal made by:
- Substance:
- Proposed category or annex:
- Findings and/or MRL exceedances:
- Method:
- Toxicity:
- Proposed priority:
- Proposed evaluation period:
- Relevant commodities:
- Additional information:
Annex VII: Substances of interest to be analysed in honey under the national control programmes

EFSA recommended in its 2014 annual report to analyse honey samples for the substances that are listed in the EU MACP in commodities of plant origin, in order to allow estimating the exposure of bees and adapting certain MRLs for honey. Member States are encouraged to conduct these analyses under their national programmes and to clearly report to EFSA which MRL (pesticides MRL or veterinary medicinal product MRL) was used for the evaluation. For honey the residue definition for plant products applies. Next to residue information for the residue definition for plant products, also information on residues in line with the residue definition for animal origin can be useful to get a view on other specific metabolites that might occur in bees.

Substances for which residues frequently occur in honey:

- Acetamiprid
- Amitraz (veterinary medicinal product)
- Azoxystrobin
- Benzalkonium chloride
- Boscalid
- Carbendazim and thiophanate methyl
- Chlorates
- Chlordane
- Clothianidin
- Chlorfenvinphos
- Coumaphos (veterinary medicinal product)
- Didecyldimethylammonium chloride
- Dimoxystrobin
- Dimethoate
- Glyphosate
- Iprodione
- Imidacloprid
- Lambda-cyhalothrin
- Orthophenylphenol
- Thiacloprid

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6 The results should be reported as mixture of alkyl-quaternary ammonium salts with alkyl chain lengths of C8, C10 and C12.
Annex VIII: Commodities of interest to be analysed under the national programmes

EFSA recommended focusing monitoring activities on commodities that frequently contain pesticides residues or that have the potential to result in a significant short-term intake:

- Small fruits and berries
- Grapefruits
- Rucola
- Apricots
- Celeriacs
- Brussels sprouts
- Cherries
- Tea

As currently little monitoring data are available for pesticides residues in feed, EFSA recommended to include animal feed commodities in the monitoring programmes in order to get a view on the animal exposure. On the basis of residue data for feed EFSA is able to estimate the exposure of humans to the pesticides residues:

- Rapeseed
- Soybean
Annex IX: Substances moved from the working document to the EU MACP

- Ametocradin (2019 EU MACP)
- Cyazofamid (2019 EU MACP)
- Cyflufenamid (2020 EU MACP)
- Fenpyrazamine (2020 EU MACP)
- Emamectin benzoate B1a, expressed as emamectin (2019 EU MACP)
- Etoxazole (2019 EU MACP)
- Fluopicolide (2018 EU MACP)
- Glyphosate (2019 EU MACP)
- Metrafenone (2019 EU MACP)
- Proquinazid (2020 EU MACP)
- Prothiocarbazole (2018 EU MACP)
- Prosulfocarb (2018 EU MACP)
- Spirotetramat (2019 EU MACP)
- Tricyclazole (2020 EU MACP)

7 Introduced for Products of Animal Origin. Analytical capability of full RD:
2015 (survey on 84 labs/25MSs): 23% of labs, 48% of MSs
2016 (survey on 81 labs/25MSs): 24% of labs, 48% of MSs
3.74% findings (2.04% MRL exceedances) EFSA 2016 report (294 samples)
Relevant for ruminant kidney, liver and honey. To be checked whether relevant for cows’ milk, animal muscle and fat.
Annex X: Special Project on dithiocarbamates (CS2) in organic samples

The existence of naturally occurring dithiocarbamates (CS2) in specific plant commodities can lead to false positive results of MRL exceedances. An effort from the Commission, EFSA and the EURLs has been initiated to examine the background levels of dithiocarbamates in certain plant products.

In order to better understand this issue and in view of the preparation of Art.12 reviews, data on dithiocarbamates background levels in organic products should be made available to EFSA by December 2019. As such MSs should include sampling of organic products for the analysis of dithiocarbamates their National Control Programs in 2018 and deliver the results to EFSA.

Further details on the project can be found following the path below on CIRCA BC:
CIRCABC > SANTE > EURLs for Pesticides
Then in the Library section follow the path:
Library > eurl-pesticides-srm > Project on Phytogenic Levels of Carbon Disulfide (Dithiocarbamates)