Quick Method for the Analysis of numerous Highly Polar Pesticides in Foods of Plant Origin via LC-MS/MS involving Simultaneous Extraction with Methanol (QuPPe-Method)

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Note: Changes from V8.1 to 9 are highlighted in yellow

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1. Scope and Short Description

A method is described for the residue analysis of very polar, non-QuEChERS-amenable, pesticides in foods of plant origin such as fruits (including dried fruits), vegetables, cereals and processed products thereof as well as honey.

Residues are extracted from the test portion following Water adjustment and the addition of acidified methanol. The mixture is centrifuged, filtered and directly analyzed by LC-MS/MS. Various options for the simultaneous LC-MS/MS analysis of different combinations of pesticides are provided. Quantification is in most cases performed with the help of isotopically labeled analogues of the target analytes, which are used as internal standards (ILISs). So far available, these ILISs are added directly to the test portion at the beginning of the procedure to compensate for any factors having an influence on the recovery-rates such as volume-deviations, analyte losses during sample preparation as well as matrix-effects during measurement.

2. Apparatus and Consumables

2.1. Powerful sample processing equipment,

e.g. Stephan UM 5 or Retsch Grindomix GM 300.

2.2. 50 mL centrifuge tubes with screw caps,

e.g.: a) reusable 50 mL Teflon® centrifuge tubes with screw caps (e.g. Nalgene/Rochester, USA; Oakridge, article-no. 3114-0050) or b) disposable 50 mL centrifuge tubes (e.g. Sarstedt / Nümbrecht, Germany, 114x28 mm, PP, article-no. 62.548.004).

2.3. Automatic pipettes,

suitable for handling volumes of 10 to 100 µL, 200 to 1000 µL and 1 to 10 mL.

2.4. 10 mL solvent-dispenser,

for the acidified methanol (3.6).

2.5. Centrifuge,

suitable for the centrifuge tubes employed in the procedure (2.2) and capable of achieving > 2500 rpm.

2.6. Syringe filters,

e.g. Cellulose mixed esters filters 0.45 µm pore size, Polyester filters 0.45 µm pore size (both from Macherey-Nagel, Düren, Germany).

Significant levels of Perchlorate and Chlorate were detected in the above mentioned polyester filters. Cellulose mixed esters filters were found to be appropriate for these two compounds. For this suitability test take the worst case scenario into account where the filters are clogged by the extracts, not allowing large volumes (e.g. $200 \mu L$) to pass. Thus elute only small volumes through the filters (e.g. $200 \mu L$). Such clogging was observed using commodities such as industrially milled cereals, pears and pineapples. Furthermore, special attention is required if filters are used to filter diluted extracts as any detected levels in the extracts will have to be multiplied accordingly when calculating the levels in the sample.

2.7. Syringes

e.g. 2 or 5 mL disposable polypropylene syringes suitable for the above mentioned filters 0.

2.8. Autosampler vials,

suitable for LC auto-samplers,

Use plastic vials if pesticides that tend to interact with glass-surfaces are present (e.g. Paraquat, Diquat, Streptomycin and Glyphosate)¹.

2.9. Volumetric flask with stoppers,

for the preparation of stock and working solutions. E.g. 20 mL; 25 mL; 50 mL, 100 mL glass flasks. Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. Paraquat, Diquat, Streptomycin and Glyphosate).

2.10. LC-MS/MS instrumentation,

equipped with ESI source and appropriate columns, see details in chapters 5.7.1 till 5.7.11.

¹The list of compounds requiring plastic vessels might not be comprehensive (this remark applies to the entire document). Such interactions with glass surfaces are typically more pronounced when solutions have low water content and low acidity.



Chemicals

Unless otherwise specified, use reagents of recognized analytical grade. Take every precaution to avoid possible contamination of Water, solvents, sorbents, inorganic salts, etc.

- 3.1. Water (deionized)
- 3.2. Methanol (HPLC quality)
- 3.3. Acetonitrile (HPLC quality)
- 3.4. Formic acid (concentrated; > 95%)
- 3.5. Acetic Acid (concentrated; >98%)
- 3.6. Acidified methanol,

pipette 10 mL Formic acid (3.4) in a 1000 mL volumetric flask and fill up to volume with methanol (3.2).

- 3.7. Citric acid-monohydrate (p.a.)
- 3.8. Dimethylamine,

e.g. 40 % by Fluka (article-no. 38940).

- 3.9. Ammonium formate (p.a.)
- 3.10. Ammonium citrate-tribasic, anhydrous (p.a.)
- 3.11. Sodium hydroxide (p.a.)
- 3.12. Di-Sodiumtetraborate-decahydrate (p.a.)
- 3.13. Dry ice,

technical grade can be used, it should be periodically checked not to contain pesticides at relevant levels.

3.14. Pesticide Standards,

of known purity.

- 3.15. Pesticide stock solutions,
- e.g. 1 mg/mL solutions of pesticide standards (3.14) in a Water miscible solvent (e.g. Water (3.1), methanol (3.2), acidified methanol (3.6), acetonitrile (3.3) or mixtures thereof). See Table 19 for the conversion fac-

tors to be applied between typical purchased standards and analytes and **Table 20** for suggested solvents for the preparation of the stock solutions.

Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. Paraquat, Diquat, Streptomycin and Glyphosate). Keep in mind that some standards are sold as salts or hydrates. Some exemplary conversion factors are shown in **Table 19**.

3.16. Pesticide working solutions / mixtures,

prepared at appropriate concentrations by diluting pesticide stock solutions (3.15) of one or more pesticides with Water-miscible solvents as required for the spiking of samples in recovery experiments (5.4) or for the preparation of calibration standards (5.5). See suggestions in **Table 20** in the Annex.

Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. Paraquat, Diquat, Streptomycin and Glyphosate).

3.17. Internal Standards (ISs),

Exemplary sources are shown in **Table 21**.

3.18. IS Stock solutions,

e.g. 1 mg/mL solutions of ISs (3.17) in a Water miscible solvent (e.g. methanol, acetonitrile, Water or mixtures thereof). For solvent-suggestions see **Table 20** in the Annex.

Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. ILISs of Paraquat, Diquat and Glyphosate as well as Dihydrostreptomycin). Keep in mind that some standards are sold as salts or hydrates. Some exemplary conversion factors are shown in **Table 19**.

Notes:

In general the absolute concentrations of the ILIS-solutions are not important as long as the ILIS-concentration in the final extract is high enough to produce a well measurable signal that is not disturbed by co-eluting matrix components. Important is furthermore that any content of the native analyte within the ILIS-standard (irrespective whether it was present as an impurity of the purchased standard or whether it was generated in the lab during storage of the ILIS-solution or during sample preparation) is low enough to exclude false positive results or significant influence on quantification. For quantification purposes it is of foremost importance that the ratio between the absolute ILIS amount added to the sample prior to extraction (or to the isolated aliquot of the sample extract) and the absolute amount of ILIS added to the calibration standard solutions is known as it is used in calculations.

3.19. IS-working solution I (IS-WS I) for spiking samples prior to extraction,

prepared at appropriate concentrations by diluting IS stock solutions (3.18) of one or more ISs with Water-miscible solvents. Suggestions for solvents are shown in **Table 20** and suggestions for the concentrations in Table 22.

Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. ILIS of Paraquat, Diquat and Glyphosate as well as Dihydrostreptomycin). In presence of Water and especially at high pH levels, Phosphonic acid ¹⁸O₃ will gradually convert to ¹⁸O₂¹⁶O₁, ¹⁸O₁¹⁶O₂ and eventually of ¹⁶O₃ (native) phosphonic acid. The ¹⁶O₃ phosophonic acid standard solution provided by the EURLs should be preferably diluted in acetonitrile, where it was shown to be stable for long periods.

3.20. IS-working solution II (IS-WS II) for preparation of calibration standards,

prepared at appropriate concentrations by diluting IS-WS-I (3.19) with Water-miscible solvents. Suggestions for solvents are shown in **Table 20** and for concentrations in **Table 22**.

Use plastic flasks and stoppers if pesticides that tend to interact with glass-surfaces are present (e.g. ILIS of Paraquat, Diquat and Glyphosate as well as Dihydrostreptomycin). See also sub-note 3 in. For short term usage (e.g. up to one month) the ILIS of Phosphonic acid can be diluted in acidified methanol (3.6).

3.21. LC-MS/MS mobile phases,

see details in chapters 5.7.1 till 0.

4. Disclaimer

This method refers to several trade name of products and instruments which are commercially available and suitable for the described procedure. This information is given for the convenience of the users of this method and does not constitute an endorsement by the EURL of the products named. The application of this method may involve hazardous materials, operations and equipment. It is the responsibility of the users of this method to establish appropriate safety and health practices prior to use. Any consumables and chemicals used in the procedure should be periodically checked, e.g. through reagent blank tests, for any relevant levels of the analytes of interest.



5. Procedure

5.1. Sample preparation

To obtain representative test-portions from the laboratory sample, proceed as required by the respective regulations and guidelines. For fruits and vegetables cryogenic milling (e.g. using dry ice) is to be preferred to minimize degradations, reduce particle size and improve homogeneity and residue accessibility.

For dry commodities (e.g. cereals, pulses) small particle sizes improve the accessibility of residues enclosed in the interior of the materials. Thus fine grinding (e.g. particle size <500µm) is preferable. The larger the particles are the longer extraction times are required to achieve quantitative extraction.

For dried fruits and similar commodities (< 30 % Water content) the following procedure is proposed: Add 850 g of cold Water to 500 g frozen dried fruits and homogenize the mixture using a strong mixer (2.1), if possible with addition of dry ice to prevent or slow down any chemical and enzymatic reactions (3.13). 13.5 g of this homogenate will correspond to 5 g sample.

5.2. Extraction / Centrifugation / Filtration

The extraction procedure is shown at a glance at chapter 5.6.

- 5.2.1.Weigh a representative portion (m_a) of the sample homogenate (5.1) into a 50 mL centrifuge tube (2.2). In case of fresh fruits and vegetables as well as juices take 10 g \pm 0.1 g of the homogenized sample. In case of dried fruits, dried vegetables, dried mushrooms take 5 g \pm 0.05 g or 13.5 g \pm 0.1 g of the re-hydrated and homogenized material according to 5.1 (corresponding to 5 g sample). In case of cereals, dried pulses and honey also take 5 g \pm 0.05 g of the homogenate. Smaller sample portions may have to be used for extract-rich commodities, such as spices or fermented products, or commodities with very high Water absorbing capacity not allowing proper extraction.
- 5.2.2. Add Water (3.1) to a total content of ca. 10 g according to the indications in Table 23.
 - No further Water adjustment is needed where re-hydrated commodities (see **5.1**) are employed. Where no ISs are used or where they are added after extract aliquotation, Water adjustment to 10g is essential. Where the appropriate ISs are employed before any aliquotation has taken place Water adjustment is less critical and may be skipped for commodities containing ≥80% Water (see **Table 23**)

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5.2.3. Add 10 mL acidified methanol (3.6) and 50 μ L of the IS-WS I (3.19) containing isotopically labeled analogues of one or more of the analytes of interest (added IS mass = m_{IS}^{sample}).

The resulting extract volume, taking into account the natural Water content of the sample and the Water added in **5.2.2** sum up ca. 20 mL (corresponds to ca. 0.5 g sample per mL extract if 10 g sample is employed for extraction). Where no ISs are used the aim should be to reach a total volume of the liquid phase that is as close as possible to 20 mL. Keep in mind that the Water volume adjustments in **Table 23** are approximate and that there is a ca. 2.5% volume contraction occurring when methanol is mixed with Water. In any case Water adjustment will help to reduce the bias related to the volume deviation from 20 mL to an acceptable level.

For screening purposes the IS can be alternatively added to a sample extract aliquot (e.g. 1 mL, see 5.2.8), assuming that 1 mL extract corresponds to exactly 0.5 g sample equivalents. This way the added amount of IS per sample can be drastically reduced (e.g. 20-fold if added to 1 mL extract). The IS added at this step will compensate for matrix effects including retention-time shifts. The quantitative result should however be considered as tentative. For more accuracy samples should be re-analyzed with the IS being added in step 5.2.3.

5.2.4.Close the tube and shake vigorously by a mechanical shaker. Shake between 1 min in the case of fresh products and 15 min in the case of dry commodities. In case of dry products the 1 min shaking by hand is to be followed by a soaking period between 15-30 minutes and a subsequent second 1 min vigorous shaking by hand.

In case of dry products (e.g. cereals, pulses) particle size plays an important role as far as extractability is concerned. If a considerable fraction of the particles exceed 500 µm shaking or soaking times may have to be extended.

5.2.5.For **Paraquat and Diquat** the 1 minute shaking is followed by a thermal treatment of 15 minutes at 80 °C in a Water bath. Then shake again for 1 minute and wait for the sample to cool down to room temperature before centrifuging.

1 minute extractions at room temperature with methanol containing 1% Formic acid are well suitable Paraquat and Diquat screening. 15-minute extractions at 80 °C using the same extraction solvent were shown to provide quantitative extraction yields of incurred Diquat and Paraquat residues in wheat and potatoes. In an experiment on **Lentils** containing incurred Diquat residues a stronger extraction solvent was necessary (MeOH/aqueous HCl 0,1M (1:1)) using the same volume, extraction temperature and extraction time as described above².

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² Kolberg DI, Mack D, Anastassiades M, Hetmanski MT, Fussell RJ, Meijer T, Mol HG. Anal Bioanal Chem. 404(8):2465-74 (2012); Development and independent laboratory validation of a simple method for the determination of paraquat and diquat in potato, cereals and pulses.

- 5.2.6. Centrifuge (e.g. for 5 min at 4000 rpm).
- 5.2.7. Filter an aliquot of the extract (e.g. 3 mL) through a syringe filter (0) into a sealable storage vessel.

The extracts of some commodity types (e.g. finely milled cereals) pose difficulties in filtration. To avoid this, place the extraction tubes from (5.2.4) or (5.2.5) for a few hours into the freezer, centrifuge and filter.

Check the filters for any cross-contamination of Perchlorate and Chlorate. Cellulose mixed-ester filters were found to be suitable for the determination of Chlorate and Perchlorate (see also chapter 0. for further information).

5.2.8.Transfer, as required, one or more aliquots (e.g. 1 mL each) of the filtered extract into auto-sampler vials (2.8)

5.3. Blank extracts

Using suitable blank commodities (not containing any detectable residues of the analytes of interest), proceed sample preparation exactly as described in **5.2** but **SKIP THE ADDITION OF ISs.**

5.4. Recovery experiments

Weigh an appropriate portion (see **0**) of a blank commodity homogenate into a 50 mL centrifuge tube (**2.2**) and spike it with a suitable pesticide working solution (**3.16** and **Table 20**).

Spike directly to the matrix, prior to any Water or solvent addition. Use small volumes of pesticide working solutions (e.g. 50-300 μ L), to avoid too strong dilution. Conduct sample preparation exactly as described in **5.2**.

5.5. Preparation of calibration standards

5.5.1. Solvent-based calibration standards

An exemplary pipetting scheme for preparing solvent-based calibration standards is shown in Table 1.

The calculation of the mass-fraction W_R of the pesticide in the sample, when IS is used, is shown in **5.8.1**.

Where solvent-based calibrations are used the use of ILISs for quantification is essential as the IS compensates for any matrix-related signal suppressions / enhancements.



5.5.2. Matrix matched calibration standards

Transfer suitable aliquots of the blank extract (5.3) to auto-sampler vials and proceed as shown in Table 1. The calculation of the mass-fraction W_R of the pesticide in the sample using matrix-matched calibration standards, with and without the use of ILIS, is shown in 5.8.1.1 and 5.8.2.1 respectively.

Table 1: Exemplary pipetting scheme for the preparation of calibration standards

					Calibr	ation stand	dards			
	Solvent based (5.5.1)			Matrix-matched (5.5.2)						
			using IS ⁴		,	without IS ⁵	i		using IS ⁴	
Calibration levels in µg pesticide /mL OR in µg pesticide/ "IS-portion"		0.05 ⁶	0.1	0.25	0.05	0.1	0.25	0.05	0.1	0.25
Blank extract (5.3)		-	-	-	900 μL	900 μL	900 µL	850 µL	850 µL	850 μL
1:1 (v/v) mix of V and acidified Me		900 µL	850 µL	900 µL	50 μL	-	50 μL	50 μL	-	50 μL
Pesticide work-	1 μg/mL	50 μL	100 μL	-	50 μL	100 μL	-	50 μL	100 μL	-
(3.16) ²	5 μg/mL	-	-	50 μL	-	-	50 μL	-	-	50 μL
IS-WS II (3.20) ^{1,3}		50 μL	50 μL	50 μL	-	-	-	50 μL	50 μL	50 μL
Total volume		1000 µL	1000 µL	1000 μL	1000 µL	1000 μL	1000 µL	1000 µL	1000 μL	1000 µL

¹ One IS portion would correspond to the IS mass contained in 50 μL IS-WS II (which in the particular example is added to each calibration standard).

² The concentration of the pesticide working solution(s) should be sufficiently high to avoid excessive dilution of the blank extract which would result in matrix effect deviations.

³For calibration standards of 1 mL it is recommended to prepare the IS-WS II (3.20) by diluting 20-fold the IS-WS I (3.19). The same volume and pipette as in 5.2.3 can then be used for the preparation of the calibration standards.

⁴ When employing IL-ISs matrix-matching and volume adjustments are of less importance as the IS compensates for any matrix-related signal suppressions / enhancements. Also solvent-based calibrations can be used here. Important is that a) the mass ratio of pesticide and IS in the respective calibration standards and b) the ratio between the IS mass added to the sample (5.2.3) and the IS mass added to the calibration standard(s) (5.5.1 and 5.5.2) is known and recorded. For convenience the latter mass ratio should be kept constant throughout all calibration levels (e.g. at 20:1 when preparing calibration standards of 1 mL).

⁴ Where ILISs are <u>not</u> available/employed, matrix-matching via matrix-matched standards Table 1) or via the standard additions approach (**5.5.3**) are particularly important to compensate for matrix effects in measurement. In both cases the total volume of the sample extracts is assumed to be exactly 20 mL, which translates into 0.5 g sample equivalents per mL.

⁶ The calibration level of 0.05 corresponds to 0.1 mg pesticide /kg sample, when using 10 g test portions, or to 0.2 mg/kg sample when using 5 g test portions.



5.5.3. Standard-Additions-Approach

Where no appropriate ISs are available the method of standard additions is a very effective approach for compensating matrix-induced enhancement or suppression phenomena. As this procedure involves a linear extrapolation it is mandatory that pesticide concentrations and detection signals show a linear relationship throughout the relevant concentration range. The procedure furthermore requires knowledge of the approximate (estimated) residue level in the sample ($w_{R(approx)}$). This info is derived from a preliminary analysis.

Prepare 4 equal portions of the final extract and spike 3 of them with increasing amounts of analyte. The amounts to be added should be chosen in such a way to remain within the linear range. It should be avoided that the added levels are too close to the expected analyte level to avoid that measurement variability will influence too much the slope, which is used to calculate the analyte level. In case the concentrations are outside the linear range a dilution of all 4 extracts with the extraction solvent is indicated.

Prepare a working solution (3.16) of the analyte at a concentration level where 50 or 100 μ L of the solution contain the lowest amount of analyte to be added.

Example A: Vial 1) no addition; vial 2) 0.5 x $w_{R(approx)}$, vial 3) 1 x $w_{R(approx)}$, and vial 4) 1.5 x $w_{R(approx)}$, Example B: Vial 1) no addition; vial 2) 1 x $w_{R(approx)}$, vial 3) 2 x $w_{R(approx)}$, and vial 4) 3 x $w_{R(approx)}$.

Adjust the volume within all vials by adding the corresponding solvent amounts.

Table 2 shows an example according to Example B. The calculation of the mass fraction of the pesticide in the sample w_R is shown in **5.8.2.2**.

Table 2 : Exemplary pipetting scheme of a standard additions approach (for a sample extract containing 0.5 g sample equivalents per mL and an estimated residue level ($w_{R(approx)}$) of 0,5 mg/kg = 0.25 μ g/1000 μ l

Additions	Vial 1	Vial 2	Vial 3	Vial 4
Volume of sample extract	1000 μL (= 0.5 g sample)			
IS	none	none	none	none
Added volume of pesticide working solution containing 5 $\mu g/mL$ (3.16)	-	50 μL	100 μL	150 µL
Resulting mass ($m_{\it pest}^{\it std~add}$) of pesticide added to each vial		0.25 µg	0.5 µg	0.75 μg
Volume of solvent	150 µL	100 μL	50 μL	-
Final volume	1150 μL	1150 μL	1150 μL	1150 µL



5.6. QuPPe-PO-Method at a glance

Weigh sample homogenate in 50 mL centrifuge tube

Fresh fruits and vegetables (with high content of water): 10 g \pm 0.1 g, Previously dehydradet dry fruit 13.5 g \pm 0.1 g (containing 5 g sample), Cereals and dried pulses (dried commoditites): 5 g \pm 0.05 g



Adjust water content of sample to 10 mL

e.g. Rye Flour: add 10 g water



Adjust water content of sample to 10 mL

(not mandatory if IL-IS is used) e.g. Potato: add 2 g of water (see Table 23)



Add 100 µL IL-IS (isotopically labeled internal standard) mix



Add 10 mL MeOH containing 1 % formic acid



Shake thoroughly for 1 min/ cereals for 15 min



Centrifuge at 4000 rpm for 5 min



Removal of Lipids

High oil content: C-18 cleanup (e.g. Avocado)
Transfer 4 mL of raw extract to a tube containing
200 mg C18-sorbent (ODS)
shake for 1 min and centrifuge at 4000 rpm for 5 min



Filter 1.5-3 mL supernatant into a plastic vial

(syringe filter 0.45 μm)
(Plastic vials are to be preferred as some compounds – tend to interact with glass)



LC-MS/MS analysis



5.7. LC-MS/MS Measurement

Any suitable LC-MS/MS conditions may be used. Some exemplary instrument measurement conditions are given below. An overview of LC-MS/MS conditions proposed within this document is given in Table 3:

Table 3: Overview and scope of the methods proposed within this document for the QuPPe method:

	M 1.1	M 1.2	M 1.3	M 1.4	M 2	М 3	M 4.1	M 4.2	M 5	M 6	М 7	М8
ESI-mode	Neg.	Neg.	Neg.	Neg.	Neg.	Pos.	Pos.	Pos.	Pos.	Pos.	Pos.	Pos.
Separation principle	Anion Exchange	Anion Exchange	Carbon	Carbon	HILIC	HILIC	HILIC	HILIC	HILIC	HILIC	HILIC	Carbon
Column type	AS 11	AS 11-HC	Hyper- carb	Hyper- carb	Obelisc-R	Obelisc-R	Obelisc-R	BEH- Amide	PFP	Obelisc-R	Trinity P1	Hyper- carb
					NEGATIVE	MODE						
Ethephon	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
HEPA	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
Glufosinate	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
N-Acetyl-glufosinate	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
MPPA	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
Glyphosate	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
AMPA	✓	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
Phosphonic acid	(√)	(√)	✓	✓	NT	NT	NT	NT	NT	NT	-	NT
N-Acetyl-AMPA	NT	✓	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
Fosetyl-Al	-	✓	✓	NT	✓	NT	NT	NT	NT	NT	√ *	NT
Maleic hydrazide	-	-	✓	NT	✓	NT	NT	NT	NT	NT	√ *	NT
Perchlorate	NT	-	✓	✓	✓	NT	NT	NT	NT	NT	√*	NT
Chlorate	NT	-	✓	✓	NT	NT	NT	NT	NT	NT	√*	NT
Bialaphos	NT	NT	✓	NT	NT	NT	NT	NT	NT	NT	-	NT
Cyanuric acid	NT	NT	✓	NT	NT	NT	NT	NT	NT	NT	√ *	NT
Bromide	NT	NT		✓	NT	NT	NT	NT	NT	NT	NT	NT
Bromate	NT	NT	(√)	✓	NT	NT	NT	NT	NT	NT	NT	NT
					POSITIVE							
Amitrole	NT	NT	-	NT	NT	✓	-	✓	NT	NT	NT	NT
ETU	NT	NT	✓	NT	NT	✓	-	✓	✓	NT	NT	NT
PTU	NT	NT	✓	NT	NT	✓	-	✓	✓	NT	NT	NT
Cyromazine	NT	NT	NT	NT	NT	✓	✓	✓	NT	NT	NT	NT
Trimesium	NT	NT	NT	NT	NT	✓	✓	✓	NT	NT	NT	NT
Daminozide	NT	NT	NT	NT	NT	✓	✓	✓	NT	NT	NT	NT
Chlormequat	NT	NT	✓	NT	NT	✓	✓	✓	✓	NT	NT	NT
Mepiquat	NT	NT	✓	NT	NT	✓	✓	✓	✓	NT	NT	NT
Difenzoquat	NT	NT		NT	NT	✓	✓	✓	✓	NT	NT	NT
Propamocarb	NT	NT	NT	NT	NT	✓	✓	✓	NT	NT	NT	NT
Melamine	NT	NT	NT	NT	NT	NT	√	✓	NT	NT	NT	NT
Diquat	NT	NT	-	NT	NT	NT	✓	-	NT	NT	NT	NT
Paraquat	NT	NT	-	NT	NT	NT	✓	-	NT	NT	NT	NT
N,N-Dimethylhydrazine	NT	NT	-	NT	NT	NT	✓	-	NT	NT	NT	NT
Nereistoxin	NT	NT	✓	NT	NT	NT	√	✓	NT	NT	NT	NT
Streptomycin	NT	NT	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT
Kasugamycin	NT	NT	NT	NT	NT	NT	NT	NT	NT	· ✓	NT	NT
Morpholine	NT	NT	NT	NT	NT	NT	(√)	(√)	NT	NT	√	NT
Diethanolamine	NT	NT	NT	NT	NT	NT	(√)	(√)	NT	NT	√	NT
Triethanolamine	NT	NT	NT	NT	NT	NT	(√)	(√)	NT	NT	→	NT
1,2,4-Triazole	NT	NT	NT	NT	NT	NT	(√) (√)	-	NT	NT	NT	- N1
Triazole-alanine	NT	NT	NT	NT	NT	NT	(√) (√)	-	NT	NT	NT	✓
Triazole-aratime Triazole-acetic acid	NT	NT	NT	NT	NT	NT	(√)	-	NT	NT	NT	✓
	NT					NT	NT				NT	→
Triazole-lactic acid		NT	NT	NT	NT			-	NT	NT		
Aminocyclopyrachlor	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT	NT	NT

EU Reference Laboratories for Residues of Pesticides Single Residue Methods

	M 1.1	M 1.2	M 1.3	M 1.4	M 2	М 3	M 4.1	M 4.2	M 5	М 6	М 7	M8
ESI-mode	Neg.	Neg.	Neg.	Neg.	Neg.	Pos.	Pos.	Pos.	Pos.	Pos.	Pos.	Pos.
Separation principle	Anion Exchange	Anion Exchange	Carbon	Carbon	HILIC	HILIC	HILIC	HILIC	HILIC	HILIC	HILIC	Carbon
Column type	AS 11	AS 11-HC	Hyper- carb	Hyper- carb	Obelisc-R	Obelisc-R	Obelisc-R	BEH- Amide	PFP	Obelisc-R	Trinity P1	Hyper- carb
Chloridazon- desphenyl	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT	NT	NT
Mepiquat-4-hydroxy	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT	NT	NT
Propamocarb-N- desmethyl	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT	NT	NT
Propamocarb-N-oxide	NT	NT	NT	NT	NT	NT	NT	✓	NT	NT	NT	NT

 ^{✓ =} satisfactory chomatography and detection sensitivity achieved,
 NT = Not tested under the conditions shown in the respective sections,
 (✓) = possible but compromised due to matrix effects or lacking separation or limited sensitivity or limited information for proper identification,
 "-" analysis was tested and found to be poor under the described conditions,

^{*} Using a gradient (98% B -> 60% B in 5 min, hold 2 min)

EU Reference Laboratories for Residues of Pesticides
Single Residue Methods

Table 4 Practical Information: Mainly used methods used at CVUA Stuttgart

Method	Special remarks on Substances	LC-MS/MS	Comments
Method 1.3 "Glyphosate & Co. Hypercarb" (see 5.7.3)	Glyphosate AMPA N-Acetyl-AMPA Ethephon HEPA Glufosinate N-Acetyl-Glufosinate MPPA Fosetyl-Al Phosphonic acid (first screening) Maleic hydrazide Perchlorate (first screening) Chlorate (first screening) Cyanuric acid Bialaphos	Agilent 1200Sciex QTRAP 5500	Evaluation via solvent calibration and ILISs except for Bialaphos and N-Acetyl-AMPA
Method 1.4 "PerChloPhos" (see 5.7.4)	Perchlorate (quantitative) Chlorate (quantitative) Phosphonic acid (quantitative) Bromide (Screening, quantitative) Bromate (quantitative)	Agilent 1200Sciex QTRAP 5500	 Employed if case screening by 1.3 was positive Dilution 1:5 Evaluation via solvent calibration and ILISs
Method 4.1 "Quats & Co Obelisc R" (see 5.7.7)	Paraquat (for specific commodities) Diquat (for specific commodities)	Waters Acquity UPLC I-ClassSciex QTRAP 5500	Evaluation via matrix-based calibration and ILISs
Method 4.2 "Quats & Co BEH Amide" (see 5.7.8)	Amitrole ETU Chlormequat Mepiquat Daminozide PTU Cyromazine Trimethylsulfonium Nereistoxin Difenzoquat Melamine Propamocarb Morpholine (first screening) Diethanolamine (first screening) Triethanolamine (first screening) Aminocyclopyrachlor Chloridazon-desphenyl Mepiquat-4-hydroxy Propamocarb-N-desmethyl Propamocarb-N-oxide	 Waters Acquity UPLC I-Class Sciex QTRAP 5500 	Evaluation via matrix-based calibration and ILISs (except for Difenzoquat, Amino- cyclopyrachlor, Mepiquat-4-hydroxy, Pro- pamocarb-N-desmethyl, Propamocarb-N- oxide)
Method 6 "Streptomycin and Kasugamycin" (see 5.7.10)	Streptomycin Kasugamycin	Agilent 1200Sciex QTRAP 5500	 Seasonal analyses of selected commodities Evaluation via solvent calibration (using Dihydrostreptomycin as IS for Streptomycin)
Method 7 "Morpholine, Diethanolamine and Trieth- anolamine" (see 5.7.11)	Morpholine (quantitative) Diethanolamine (quantitative) Triethanolamine (quantitative)	Waters Acquity UPLC I-Class Sciex QTRAP 5500	 Employed if case screening by 4.2 was positive Employed if DEA was false negative, by 4.2 e.g. in cereals, dried mushrooms, pomegranates Evaluation via solvent calibration and ILISs
Method 8 "Triazole derivative metabolites (TDMs)" (see 0)	1,2,4-Triazole Triazol-alanine Triazole-acetic acid Triazole-lactic acid	Waters Acquity UPLC I-Class Sciex SelexION Q-Trap® 5500	 Method employed to collect data on TDM-levels in food Evaluation via solvent calibration and ILISs



5.7.1. Method 1.1 "Glyphosate & Co. AS 11"

Table 5: Proposed LC-MS/MS conditions for Ethephon, HEPA (Ethephon metabolite), Glyphosat, AMPA (Glyphosate metabolite), Glufosinate, MPPA (Glufosinate metabolite), N-Acetyl-Glufosinate (Glufosinate metabolite), Phosphonic acid.

Instrument parameters	Conditions	Conditions					
Ionization mode	ESI neg	ESI neg					
Column/temperature (see notes)	Dionex IonPac AS 11 2 x 250 mm (P/N 44077); 40°C						
Pre-column	Dionex IonPac AG 11 2 x	Dionex IonPac AG 11 2 x 50 mm (P/N 44079)					
Pre-filters	e.g. Supelco column saver 2	2.0 µm Filter (optional)					
Eluent A	Water (3.1)						
Eluent B	Note: You will need ca. 0.5	1 mM Citric acid in Water adjusted to pH 11 with Dimethylamine (DMA) Note: You will need ca. 0.5 mL DMA solution for 500 mL 1 mM citric acid in Water Make sure your eluent filters can handle alkaline solvents (see notes)!!					
Gradient	% A	Flow [mL/min]	Time [min]				
	100	0.3	0				
	50	0.3	8				
	50	0.3	15				
	100	0.3	15.1				
	100	0.3	23				
Injection volume	10-20 µL (Note: in case of a on the instrument)	nalyzing only Ethephon 5 μ	uL may be enough -depending				
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS-portion	n* + one level at the report	ing limit				
Acquired mass transitions (m/z)	Compound	Mass Transition	ıs (m/z)				
	Glyphosate: Glyphosate- ¹³ C ₂ , ¹⁵ N ₁ (ILIS):	168/63, 168/124, 171/63	168/63, 168/124, 168/150, 168/81 171/63				
	AMPA: AMPA- ¹³ C ₁ ¹⁵ N ₁ (ILIS):	110/63, 110/79, 112/63	110/63, 110/79, 110/81** 112/63				
	Ethephon: Ethephon-D ₄ (ILIS):	143/107, 143/79, 147/111	145/107				
	HEPA: HEPA-D ₄ (ILIS):	125/79, 125/95, 129/79	125/63				
	Glufosinate: Glufosinate-D ₃ (ILIS):	180/63, 180/136, 183/63	180/85, 180/95				
	N-Acetyl-glufosinate: N-Acetyl-glufosinate-D ₃ (ILIS	222/63, 222/59, 2 3): 225/63	222/136				
	MPPA: MPPA-D ₃ (ILIS):	151/63, 151/107, 154/63	151/63, 151/107, 151/133 154/63				

AMPA: Aminomethylphosphonic acid; MPPA: 3-Methylphosphinicopropionic acid; HEPA: 2-Hydroxyethylphosphonic acid (= hydroxy-ethephon),

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

^{**} See comment 1 under 5.7.1.1 concerning potential interference of AMPA by Fosetyl.

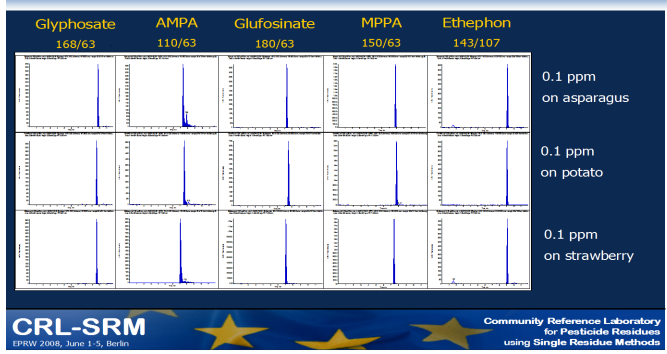


Figure 1: Typical chromatograms of HEPA in real samples

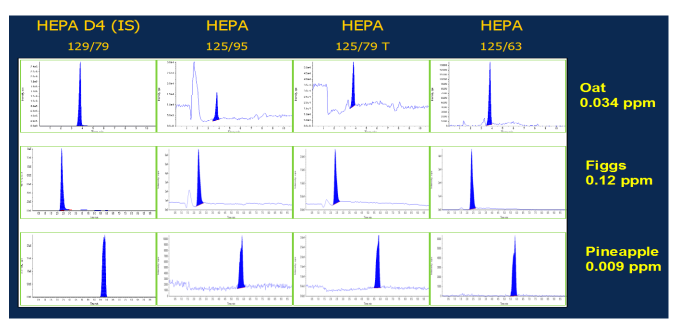


Figure 2: Typical chromatograms of Glyphosate, AMPA, Glufosinate, MPPA and Ethephon spiked on blank-QuPPe extracts of various commodities

5.7.1.1. Hints on Method 1.1

- 1) AMPA and Fosetyl share the mass-transition 110/81. Chromatographic separation is thus needed.
- 2) As the pH of the mobile phase is quite high, it is recommendable to <u>use alkali-compatible components</u>, e.g. metal frits instead of silica frits in the Eluent B reservoir; borosilicate 3.3 bottles instead of glass bottles for eluent B; rotor-seals from alkali-persistent materials, such as PEEK (polyetherketone) or Tefzel, rather than Vespel.
- 3) <u>Priming and reconditioning of column:</u> before first use, after long storage (e.g. >2 weeks), after injection of 50-100 sample extracts for column A or 100-200 extracts for column B):
 - a. Flush column for 30 minutes with **100 mmol aqueous Borax solution** (7,62 g di-sodium tetraborate decahydrate in 200 mL Water) at 0.3 mL/min **OR** Flush for 1 hour with 30 mM NaOH (240 mg NaOH in 200 mL Water) at 0.3 mL/min
 - b. Flush column for 30 minutes with Eluent A (Water) at 0.3 mL/min
 - c. Run system 3-4 times with full gradient (inject standards in matrix)

NOTE: When flushing NaOH or Borax solution through the column make sure that it will go directly into waste and not to the MS ion source!.

- 4) <u>Storage of column</u>: If to be stored for short periods (<2 weeks), columns can be put aside after any normal sequence/run (full gradient). Run system 3-4 times with full gradient to reactivate the column (inject standards in matrix) before starting the sequence. If to be stored for longer periods (e.g. >2 months) recondition the column as described under I.1.a-c
- 5) <u>Pre-filters:</u> If pre-filters are used exchange them as soon as backpressure increases significantly. For practical and convenience reasons it is recommended to exchange pre-filters when performing other maintenance operations such as reconditioning or pre-column exchange.

NOTE: Losses of glyphosate, that could be clearly linked to interactions with a dirty pre-filter, have been once observed.

6) Pre-columns (guard columns):

The pre-column should be exchanged as soon as a clear deterioration of the separation performance (worsening of peak-shape) is noticed. The pre-column of method 1.1. needs to be exchanged more often than that of 1.2 and 1.3. If after pre-filter exchange (see above) the pressure does not come back to normal levels, the frit of the pre-column should be exchanged.

For further information on the storage and cleanup of column B see: http://www.dionex.com/en-us/webdocs/113497-Man-065463-03-lonPac-AS11-HC-4um-Nov12.pdf



5.7.2. Method 1.2 "Glyphosate & Co. AS 11-HC"

Table 6: Proposed LC-MS/MS conditions for Ethephon, HEPA (Ethephon metabolite), Glyphosat, AMPA (Glyphosate metabolite), Glufosinate, MPPA (Glufosinate metabolite), N-Acetyl-glufosinate (Glufosinate metabolite), Fosetyl-Al, N-Acetyl-AMPA and Phosphonic acid.

Instrument parameters	Conditions	Conditions					
Ionization mode	ESI neg	ESI neg					
Column/temperature (see also notes below)	Dionex IonPac AS 11-HC	Dionex IonPac AS 11- HC 2 x 250 mm (P/N 052961); 40°C					
Pre-column	Dionex IonPac AG11-HC	2 x 50 mm (P/N 052963)					
Pre-filters	e.g. Supelco column saver 2	e.g. Supelco column saver 2.0 µm Filter (optional)					
Eluent A	Water (3.1)						
Eluent B	1 mM tribasic Ammonium ci	trate in Water					
Gradient	%A	Flow [mL/min]	Time [min]				
	100	0.3	0				
	0	0.3	8				
	0	0.3	16				
	100	0.3	16.1				
	100	0.3	23				
Injection volume	10 μL						
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS-portio	n* + one level at the report	ing limit				
Acquired mass transitions (m/z)	Compound	Mass Transition	Mass Transitions (m/z)				
	Glyphosate: Glyphosate- ¹³ C ₂ , ¹⁵ N (ILIS):	168/63, 168/124 171/63	168/63, 168/124, 168/150, 168/81 171/63				
	AMPA: AMPA- ¹³ C, ¹⁵ N (ILIS):	110/63, 110/79, 112/63	110/63, 110/79, 110/81** 112/63				
	N-Acetyl-AMPA:	152/63, 152/79,	152/110				
	Ethephon: Ethephon-D ₄ (ILIS):	143/107, 143/79 147/111	, 145/107				
	HEPA: HEPA-D ₄ (ILIS):	125/79, 125/95, 129/79	125/63				
	Glufosinate: Glufosinate-D ₃ (ILIS):	180/63, 180/136 183/63	, 180/85, 180/95				
	N-Acetyl-glufosinate: N-Acetyl-glufosinate-D ₃ (ILIS	222/63, 222/59, 2 S): 225/63	222/136				
	MPPA: MPPA-D ₃ (ILIS):	151/63, 151/107 154/63	, 151/133				
	Fosetyl-Al: Fosetyl-Al-D ₁₅ (ILIS):	109/81, 109/63 (114/82 (Fosetyl-					
	Phosphonic acid***: Phosphonic acid- ¹⁸ O ₃ (ILIS)	81/79, 81/63 : 87/85					

AMPA: Aminomethylphosphonic acid; MPPA: 3-Methylphosphinicopropionic acid; HEPA: 2-Hydroxyethylphosphonic acid (=hydroxy-ethephon)
* One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

^{**} See comment 1 under **5.7.1.1** concerning potential interference of AMPA by Fosetyl.

^{***} See comment 3 on Phosphonic acid under 5.7.2.1

2.0e5

1.5e5

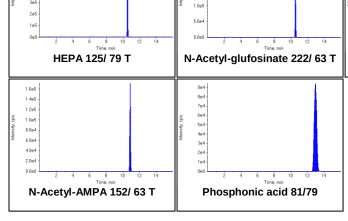
1.0e5

5.0e4

AMPA 110/63 T

3e5

Fosetyl 109/81 T



1.5e5

4e5

3e5

Figure 3: Typical chromatograms of Ethephon, HEPA, Glyphosat, AMPA, Glufosinate, MPPA, N-Acetyl-AMPA, N-Acetyl-glufosinate, Fosetyl-Al and Phosphonic acid at 0.1 mg/L in MeOH with 1% Formic acid.

5.7.2.1. Hints on Method 1.2

- 1) Using this M1.2 some compounds (e.g. Glyphosate) in some commodities tend to give two sharp peaks. The corresponding ILIS typically behaves equally, so that quantification with any of the two peaks remains accurate
- 2) AMPA and Fosetyl share the mass-transition (110/81). Chromatographic separation is thus needed (typically the case).
- 3) Fosetyl and Fosetyl-D₅ tend to degrade to Phosphonic acid both in solutions and in the LC-MS/MS via in-sorce fragmentation. A good chromatographic separation between the two is thus necessary (typically the case).
- 4) When extracts containing high levels of Phosphoric acid (which is naturally contained at high concentrations in many samples) are injected the chromatographic separation of Phosphoric and Phosphonic acid is compromised. This often results in a suppression of the Phosphonic acid signal and in some cases even leads to false negative results. The most important qualifier mass-transition of Phosphonic acid (81/63) also occurs as a minor transition of Phosphoric acid, but as the latter is often present at much higher levels than Phosphonic acid its interference on this mass transition can still be significant, especially if these two elute in close vicinity. The chromatographic separation of Phosphoric and Phosphonic acid considerably improves following dilution of the extracts typically allowing proper detection, identification and quantification of Phosphonic acid next to high levels of phosphoric acid. It is thus beneficial to inject smaller volumes of sample extract (e.g. 1-2 μL) or to dilute QuPPe extracts 5-10-fold before injection. Fortunately both, Phosphoric and Phosphonic acid have at least 1 proper mass-transition (97/63 and 81/79 respectively), which in the case of Phosphonic acid can be used for quantitation and to improve identification certainty. The elution time and peak shape of the Phosphonic acid ILIS can also be used to distinguish it from Phosphoric acid and to avoid false positives. Using signals on the 81/63 mass trace it was calculated that 20 mg/kg Phosphoric acid would simulate 0.1 mg/kg Phosphonic acid if this mass transition was used for quantification.
- 5) <u>Priming and reconditioning of column:</u> before first use, after long storage (e.g. >2 weeks), after injection of 50-100 sample extracts for column A or 100-200 extracts for column B):
 - a. Flush column for 30 minutes with **100 mmol aqueous Borax solution** (7,62 g di-sodium tetraborate decahydrate in 200 mL Water) at 0.3 mL/min <u>OR</u> Flush for 1 hour with 30 mM NaOH (240 mg NaOH in 200 mL Water) at 0.3 mL/min
 - b. Flush column for 30 minutes with Eluent A (Water) at 0.3 mL/min
 - c. Run system 3-4 times with full gradient (inject standards in matrix)

NOTE: When flushing NaOH or Borax solution through the column make sure that it will go directly into waste and not to the MS ion source!.

- 6) <u>Storage of column</u>: If to be stored for short periods (<2 weeks), columns can be put aside after any normal sequence/run (full gradient). Run system 3-4 times with full gradient to reactivate the column (inject standards in matrix) before starting the sequence. If to be stored for longer periods (e.g. >2 months) recondition the column as described under I.1.a-c
- 7) <u>Pre-filters:</u> If pre-filters are used exchange them as soon as backpressure increases significantly. For practical and convenience reasons it is recommended to exchange pre-filters when performing other maintenance operations such as reconditioning or pre-column exchange.

NOTE: Losses of glyphosate, that could be clearly linked to interactions with a dirty pre-filter, have been once observed.

8) Pre-columns (guard columns): The pre-column should be exchanged as soon as a clear deterioration of the separation performance (worsening of peak-shape) is noticed. The pre-column of method 1.2. needs to be exchanged less often than that of 1.1. If after pre-filter exchange (see above) the pressure does not come back to normal levels, the frit of the pre-column should be exchanged



5.7.3. Method 1.3 "Glyphosate & Co. Hypercarb"

Table 7: Proposed LC-MS/MS conditions for Ethephon, HEPA (Ethephon metabolite), Glyphosat, AMPA (Glyphosate metabolite), N-Acetyl-AMPA (Glyphosate metabolite), Glufosinate, MPPA (Glufosinate metabolite), N-Acetyl-glufosinate (Glufosinate metabolite), Fosetyl-Al, Phosphonic acid (Fosetyl metabolite), Maleic hydrazide, Perchlorate, Chlorate, Cyanuric acid and Bialaphos.

Instrument parameters	Conditions	Conditions						
Ionization mode	ESI neg							
Column/temperature	Hypercarb 2.1 x 100 m	Hypercarb 2.1 x 100 mm 5 μm (P/N 35005-102130); 40°C						
Pre-column	Hypercarb Guard 2.1 x	Hypercarb Guard 2.1 x 10 mm 5 μm (P/N 35005-102101)						
Pre-filters	e.g. Supelco column save	e.g. Supelco column saver 2.0 µm Filter (optional)						
Eluent A	1% Acetic acid in Water +	1% Acetic acid in Water + 5% MeOH						
Eluent B	1% Acetic acid in MeOH							
Gradient	% A	Flow [mL/min]	Time [min]					
	100	0.2	0					
	70	0.2	10					
	70	0.4	11					
	70	0.4	18					
	10	0.4	19					
	10	0.4	22					
	100	0.2	22.1					
	100	0.2	30					
Injection volume	5 µL							
Calibration standards and levels		ion* + one level at the reportir	ng limit					
Acquired mass transitions (m/z)	Compound	Mass Transitions	<u> </u>					
Acquired mass transitions (m/2)	Glyphosate: Glyphosate- ¹³ C ₂ , ¹⁵ N (ILIS):	168/63, 168/124, 16	168/63, 168/124, 168/150, 168/81 171/63					
	AMPA**: AMPA- ¹³ C, ¹⁵ N (ILIS):	110/63, 110/79, 110 112/63	//81 **					
	N-Acetyl-AMPA:	152/63, 152/79, 152	152/63, 152/79, 152/110					
	Ethephon: Ethephon-D ₄ (ILIS):	143/107, 143/79, 14 147/111	143/107, 143/79, 145/107 147/111					
	HEPA: HEPA-D ₄ (ILIS):	125/79, 125/95, 125 129/79	6/63					
	Glufosinate: Glufosinate-D ₃ (ILIS):	180/63, 180/136, 18 183/63	180/63, 180/136, 180/85, 180/95 183/63					
	N-Acetyl-glufosinate: N-Acetyl-glufosinate-D ₃ (ILIS		222/63, 222/59, 222/136 225/63					
	MPPA: MPPA-D ₃ (ILIS):	151/63, 151/107, 15 154/63	151/63, 151/107, 151/133 154/63					
	Fosetyl-Al: Fosetyl-Al-D ₁₅ (ILIS):	109/81, 109/63 (dete 114/82 (detected as						
	Phosphonic acid***/****: Phosphonic acid- ¹⁸ O ₃ (ILIS):	81/79, 81/63 (detect 87/85	ted as Phosphonate anion)					
	Maleic hydrazide: Maleic hydrazide-D ₂ (ILIS):	111/82, 111/42, 111 113/42	/55, 111/83					
	Perchlorate: Perchlorate- ¹⁸ O ₄ (ILIS):	99/83, 101/85 107/89						
	Chlorate: Chlorate- ¹⁸ O ₃ (ILIS):	83/67, 85/69 89/71						
	Cyanuric acid: Cyanuric acid- ¹³ C ₃ :	128/42, 128/85 131/43						
	Bialaphos:	322/88, 322/94, 322	2/134					

^{****}For Phosphonic acid, Perchlorate and Chlorate better run Method 1.4 (5.7.4) as these compounds seem to be strongly suppressed by matrix using these LC-conditions

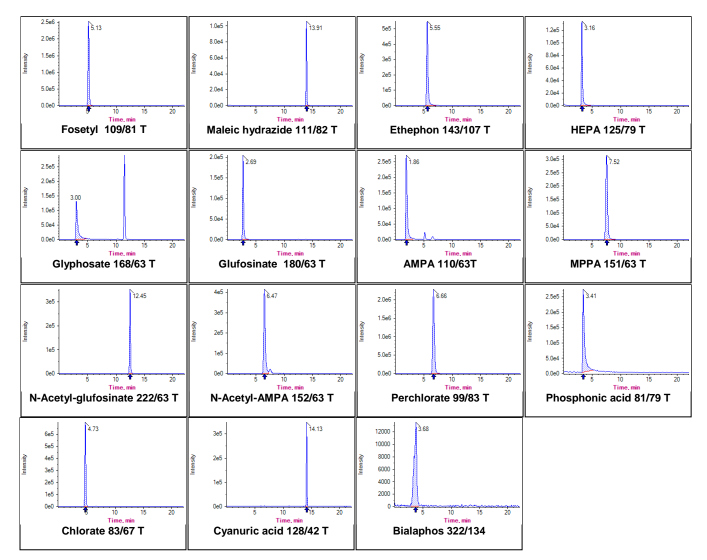


Figure 4: Chromatograms of Ethephon, HEPA, Glyphosat, AMPA, Glufosinate, MPPA, N-Acetyl-AMPA, N-Acetyl-Glufosinate, Fosetyl-Al, Maleic hydrazide, Phosphonic acid, Perchlorate, Chlorate, Cyanuric acid and Bialaphos at 0.1 mg/L in MeOH (with 1% Formic acid).

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

^{**} See comment 1 under 5.7.3.1concerning potential interference of AMPA by Fosetyl.

^{***} See comment 3 on Phosphonic acid under **5.7.3.1**



5.7.3.1. Hints on Method 1.3

1) Priming and reconditioning of the column:

before the first use, the Hypercarb columns and pre-columns have to be thoroughly primed to cover certain active sites on the surface. Priming with solutions containing planar molecules such as chlorophyll and anthocyans accelerates the priming period. A recommendable procedure for priming is the injection of QuPPe extract of spinach (for equilibration of the precolumn inject 10-15 injections spinach extracts, for column and pre-column inject 50 injections spinach extracts, if possible inject 50 µL) or the injection grape skin extract solution, prepared by dissolving 100 mg grape skin extract in 20 mL MeOH + 1% FA-H2O 1:1. This masking of the active sites is temporary and the activity of the column gradually increases with the injection of solvent or diluted extracts. Following a sequence of injections with low or no matrix load will typically raise the need for intermediate conditioning with extracts to restore the column. The impact of priming on the chromatographic properties of the column is exemplary shown in Figures 10, 11 and 12.

Table 8: Proposed LC-MS/MS conditions for priming and reconditioning of the Hypercarb column.

Instrument parameters	Conditions						
Ionisation mode	ESI neg						
Column/temperature	Hypercarb 2.1 x 100 mm 5 μm (P/N 35005-102130); 40°C						
Pre-column	Hypercarb Guard 2.1 x 10 mm 5 μm (P/N 35005-102101)						
Pre-filters	e.g. Supelco column saver 2	2.0 µm Filter (optional)					
Eluent A	1% Acetic acid in Water + 5% MeOH						
Eluent B	1% Acetic acid in MeOH						
Gradient	%A Flow [mL/min] Time [min]						
	100	0.3	0				
	70	0.3	7				
	100	0.3	7.1				
	100	0.3	12				
Injection volume	50 μL						
MS-System	If possible disconnect the M	S-System to prevent contami	nation of the MS.				

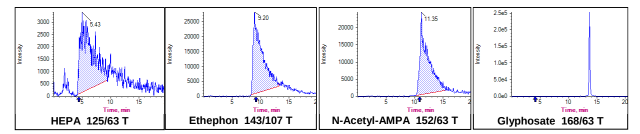


Figure 5: Chromatograms obtained using a new Hypercarb column, poor chromatographic behavior due to strong interactions of analytes with active sites. Same behavior is observed when the pre-column is new.

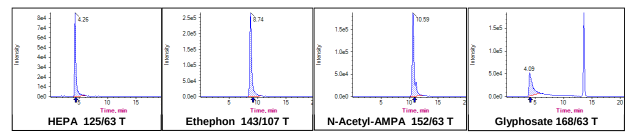


Figure 6: Chromatograms following priming with 10 injections (20µL) of Spinach QuPPe extracts.

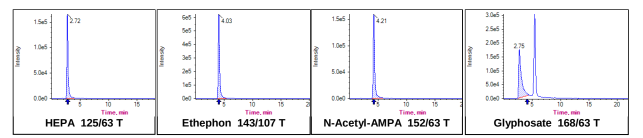


Figure 7: Chromatograms after additional injection of ca. 100 extracts of various fruit and vegetable QuPPe-extracts during normal routine use.

- 2) Pre-columns (guard columns): The pre-column should be exchanged as soon as a clear deterioration of the separation performance (worsening of peak-shape) is noticed. The pre-column of method 1.3 needs to be clearly less often exchanged compared to the pre-columns of methods 1.1 and 1.2. Any exchange of the pre-column requires priming as described above. For this the pre-column does not have to be attached to the column. Connecting several pre-columns in a row and priming them simultaneously is also an option.
- 3) Storage of columns: Following normal operation the column can be stored directly after any normal sequence/run (full gradient). Run system 3-4 times with full gradient to reactivate the column (inject standards in matrix) before starting the sequence. If to be stored for longer periods (e.g. >2 months) it is recommended to recondition the column as described above.
- 4) Pre-filters: If pre-filters are used exchange them as soon as backpressure increases significantly. For practical and convenience reasons it is recommended to exchange pre-filters when performing other maintenance operations such as reconditioning or pre-column exchange. If after pre-filter exchange (see above) the pressure does not come back to normal levels, the frit of the pre-column may need to be exchanged.
 Note: Losses of Glyphosate, that could be clearly linked to interactions with a dirty pre-filter, have been once observed.
- 5) AMPA and Fosetyl share the mass-transition m/z 110/81. Chromatographic separation is thus needed (typically the case).
- 6) Fosetyl and its D5-analogon tend to degrade to Phosphonic acid both in solutions and via in-source fragmentation in LC-MS/MS. A good chromatographic separation between Fosetyl and Phosphonic acid is thus necessary (and is typically the case). Figure 8 shows an example of this in-source fragmentation. Upon injection of 0.1μg/mL Fosetyl a peak showed up on the mass traces of Phosphonic acid at the retention time of Fosetyl. The signal intensity of this peak corresponded to 0.04 μg/mL Phosphonic acid. When injecting Fosetyl-D5 at 0.1 μg/kg the in-source fragmentation was less abundant (corresponding to ca. 0.001 μg/mL Phosphonic acid) but Phosphonic acid as impurity showed up at its proper retention time at a concentration corresponding to ca. 0.007 μg/mL. To be on the safe side Fosetyl-ILIS should thus not be added to calibration solutions or samples or sample extracts intended to be used for the analysis of native phosphonic acid. Furthermore calibration solutions used for the analysis of phosphonic acid should better not contain any native Fosetyl.



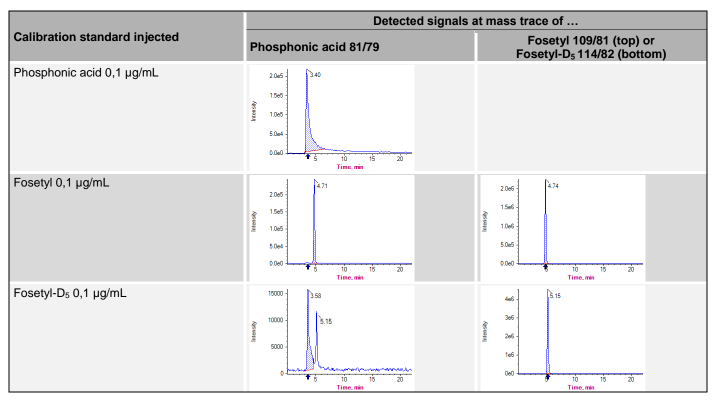


Figure 8: Chromatograms of Phosphonic acid, Fosetyl and Fosetyl-D₅ (each at 1,0 μg/mL). In addition to the proper mass-traces of Fosetyl and Fosetyl-D₅ the mass trace of Phosphonic acid is also shown to demonstrate the occurrence of in-source fragmentation of Fosetyl and Fosetyl-D₅ towards Phosphonic acid as well as the presence of Phosphonic acid as an impurity of the Fosetyl-D₅ standard solution.

7) A degradation of Ethephon to Phosphonic acid in solution is also observed. Figure 9 shows a small peak of Phosphonic acid (corresponding to 0.002 μg/mL) that showed up when Ethephon standard at 1 μg/ mL was injected. This contamination is considered negligible. However Figure 6 also shows chromatograms of an unsuitable Ethephon-D4 standard containing only ca. 0.08 μg/mL Ethephon-D4 and ca. 0.8 μg/mL Phosphonic acid instead of the expected 1 μg/mL Phosphonic acid. The use of such an ILIS would contaminate the sample with Phosphonic acid leading to false positive results. To be on the safe side Ethephon-ILIS should thus not be added to calibration solutions, samples or sample extracts intended for the analysis of native phosphonic acid. Furthermore calibration solutions used to analyse phosphonic acid should better not contain any native Ethephon.

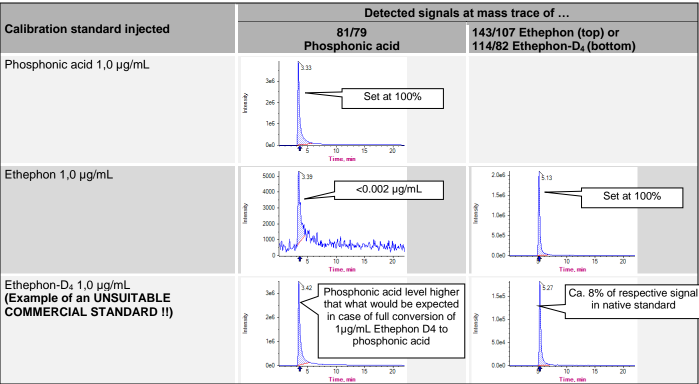


Figure 9: Chromatograms of Phosphonic acid, Ethephon and an unsuitable Ethephon- D_4 standard (each at 1,0 μ g/mL). Whereas Phosphonic acid is only contained at very low concentrations in the Ethephon standard the content of Phosphonic acid in the Ethephon-D4 standard is unacceptably high with the Phosphonic acid having already been present at high contents in the purchased standard.

8) In case of the determination of Fosetyl and Phosphonic acid on the Hypercarb-column, we refer to the patent of D. Rosati and C. Venet from Bayer CropScience (Patent-No. WO 2006079566 A1).



5.7.4.Method 1.4 "PerChloPhos"

Table 9: Proposed LC-MS/MS conditions for Phosphonic acid (Fosetyl metabolite), Perchlorate, Chlorate, Bromide and Bromate.

Instrument parameters	Conditions						
Ionisation mode	ESI neg						
Column/temperature	Hypercarb 2.1 x 100 mm 5 μm (P/N 35005-102130); 40°C						
Pre-column	Hypercarb Guard 2.1 x 1	Hypercarb Guard 2.1 x 10 mm 5 μm (P/N 35005-102101)					
Pre-filters	e.g. Supelco column saver 2	e.g. Supelco column saver 2.0 µm Filter (optional)					
Eluent A	1% Acetic acid in Water + 5	% MeOH					
Eluent B	1% Acetic acid in MeOH						
Gradient	% A	Flow [m	nL/min]	Time [min]			
	100	0.	4	0			
	70	0.	4	10			
	100	0.4		10.1			
	100	0.	4	15			
Injection volume	5 μL						
Dilution	1:5 dilution MeOH + 1% Fo (1 µL sample extract + 5 µl		Formic acid)				
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion	n* + one level a	t the reporting	limit			
Acquired mass transitions	Compound		Mass Transi	tions (m/z)			
	Bromate: Bromate ¹⁸ O ₃ (ILIS):		127/95, 129/113, 127/111, 129/97 135/117				
	Bromide*:		81/81, 79/79				
	Chlorate: Chlorate- ¹⁸ O ₃ (ILIS):		83/67, 85/69 89/71				
	Perchlorate: 99/83, 101/85 Perchlorate- ¹⁸ O ₄ (ILIS): 107/89						
* The 1.5 dilution is used for Bromide ser	Phosphonic acid: Phosphonic acid ¹⁸ O ₃ (ILIS):		81/79, 81/63 87/85				

^{*} The 1:5 dilution is used for Bromide screening. For quantification purposes where Bromide exceeds ca. 1 mg/kg, the sample extracts should be diluted e.g. 1:250 (1:50 manually and 1:5 by the HPLC).

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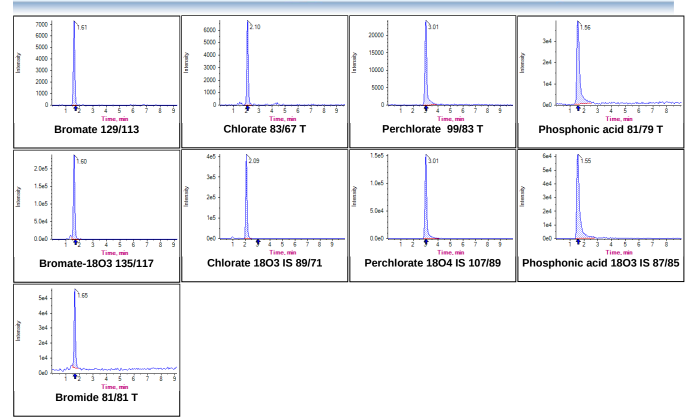


Figure 10: Chromatograms of Bromate (0.02 mg/kg in Currant), Bromide (1 mg/kg in Currant), Phosphonic acid (0.05 mg/kg in Currant), Perchlorate (0.01 mg/kg in Currant) and Chlorate (0.01 mg/kg in Currant).

5.7.4.1. Hints on Method 1.4

- 1) The hypercarb column and its pre-column should be thoroughly primed before usage, see hint on Method 1.3.
- 2) Check the filters for any cross-contamination of Perchlorate and Chlorate. See comments under **0** Cellulose mixed ester filters were found to be suitable for this application!
- 3) Fosetyl and Ethephon as well as their respective ILIS's degrade to Phosphonic acid. To be on the safe side Fosetyl, Ethephon and their respective ILIS's should thus not be added to calibration solutions or samples or sample extracts intended to be used for the analysis of native phosphonic acid. Furthermore calibration solutions used for the analysis of phosphonic acid should better not contain any native Fosetyl. See also hints on method 1.3.
- 4) When extracts containing high levels of Phosphoric acid (which is naturally contained at high concentrations in many samples) are injected, the chromatographic separation of Phosphoric and Phosphonic acid is compromised. This often results in a suppression of the Phosphonic acid signal and in some cases even leads to false negative results. The most important qualifier mass-transition of Phosphonic acid (81/63) also occurs as a minor transition of the in-source fragment of Phosphoric acid, but as the latter is often present at much higher levels than Phosphonic acid the interference on this mass transition can still be significant, especially if these two elute in close vicinity (exemplarily shown at the chromatograms in Figure 11). The chromatographic separation of Phosphoric and Phosphonic acid considerably improves following dilution of the extracts typically allowing proper detection, identification and quantification of Phosphonic acid next to high levels of phosphoric acid. It is thus beneficial to inject smaller volumes of sample extract (e.g. 1-2 µL) or to dilute QuPPe extracts 5-10-fold before injection. Fortunately both, Phosphoric and Phosphonic acid have at least one proper mass-transition (m/z 97/63 and 81/79 respectively, shown in Figure 11 which in the case of Phosphonic acid can be used for quantitation and to improve identification certainty. The elution time and peak shape of the Phosphonic acid ILIS can also be used to distinguish it from Phosphoric acid and to avoid false positives. Using signals on the m/z 81/63 mass trace it was calculated that approx. 200 mg/kg Phosphoric acid would fake 0.1 mg/kg Phosphonic acid if this mass transition was used for quantifica-



tion. In an experiment using Differential Mobility Separation (DMS) technique (see Figure 8 and Figure 9) a separation of Phosphoric acid and Phosphonic acid at the mass trace m/z 81/63 was achieved.

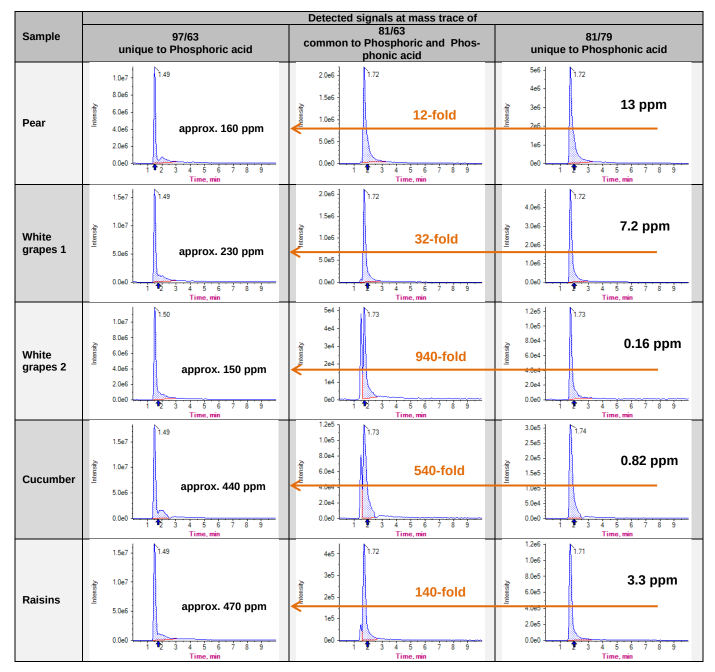


Figure 11: Chromatographic and mass-spectrometric separation of Phosphoric and Phosphonic acid.

5) High levels of Phosphoric acid (which is naturally contained at high concentrations in many samples) or Phosphonic acid (that is used as insecticide) could affect the determination of bromide. Depending on the condition of the column, the separation of these three compounds could be insufficient, resulting in compromised identification and quantification. Bromine is mainly composed of two naturally occurring stable isotopes, that are almost equally frequent (⁷⁹Br and ⁸¹Br). Being an element, no MS/MS fragmentation is possible so that MS/MS analysis has to rely on "parent/parent" analysis. The mass trace m/z 81/81 is recommended for quantifications whereas m/z 79/79 can be used as a qualifier.

The mass trace m/z 81/81 is interfered by Phosphonic acid (m/z of [H2PO3]-=81) whereas m/z 79/79 is highly affected by Phosphoric acid due to in-source fragmentation (Figure 12, the two columns declared as "CE -5 V"), the two left columns). At the mass trace m/z 81/81, 10 ppm Phosphonic acid simulate 7 ppm Bromide. At the mass trace m/z 79/79, 10 ppm Phosphoric acid simulate aprox. 2,5 ppm bromide. In practice the interference by Phosphoric acid is more critical as it is naturally contained at high levels (e.g. 100-2000 mg/kg) in various samples. A 50-fold dilution of QuPPe extracts typically allows better identification and quantification of bromide next to high levels of Phosphoric and Phosphonic acid as this improves chromatographic separation and reduces matrix-effects.

To improve selectivity and increase quantification accuracy and identification certainty, the interferences caused by Phosphoric and Phosphonic acid can be further reduced by increasing the Collision Energy (CE) for the m/z 81 and 79 (Figure 12, the two columns declared as "CE -70 V"). While Bromide cannot be fragmented, the interfering quasi-molecular ion of Phosphonic acid (m/z 81) as well as the interfering in-source fragments of Phosphoric and Phosphonic acid (m/z 79) are largely destroyed by increased collision induced dissociation. While losing up to a 100-fold of absolute sensitivity, the interferences were largely decreased resulting in a better signal-to-noise ratio.

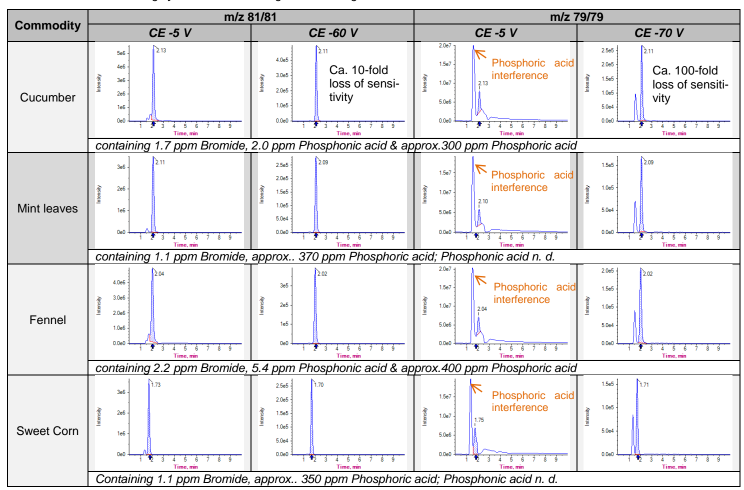


Figure 12: Chromatograms of Bromide using non-optimized collision energies (CE -5 V) showing the interference by Phosphoric acid and Phosphonic acid as well as optimized collision energies (CE -60 V and -70 V, the) showing reduced interferences.



6) Chlorate can be a minor contaminant of Perchlorate solutions and is also a minor in-source fragment of Perchlorate. In the experiment shown below Perchlorate standard at 0.2 μg/mL was injected resulting in two peaks on the mass traces of Chlorate (see Table 5). One originating from Chlorate contained as impurity in the Perchlorate solution (at ca. 0.35%) and one originating from in-source fragmentation at the retention time of Perchlorate, corresponding to a Chlorate amount of 0,001 μg/mL. This means that calibration solutions containing both chlorate and perchlorate at the same level the chlorate signal will be overestimated by ca. 0.5% which is negligible. Also samples containing perchlorate may fake the presence of chlorate at very low levels normally well below the reporting level of chlorate. When chlorate ILIS is co-injected misidentification is highly unlikely as the two compounds typically separate well chromatographically.

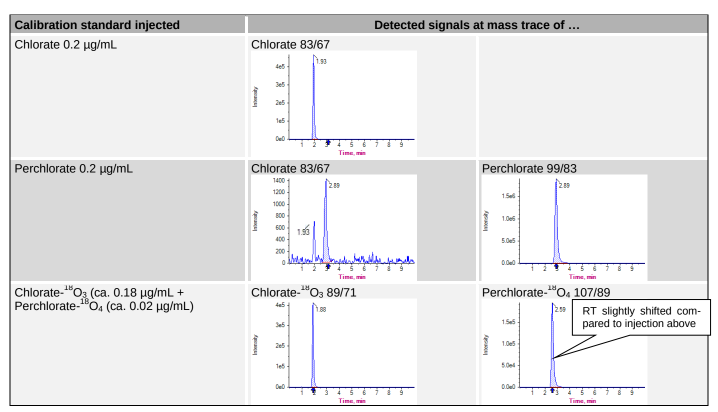


Figure 13: Chromatograms of Chlorate and Perchlorate at 0.2 μ g/mL and of a mixture of Chlorate-¹⁸O₃ and Perchlorate-¹⁸O₄, containing ca. 0.2 μ g/mL Chlorate ¹⁸O₃ and ca. 0.02 μ g/mL Perchlorate-¹⁸O₄.



5.7.5. Method 2 "Fosetyl and Maleic Hydrazide"

Table 10: Proposed LC-MS/MS conditions for Fosetyl-Al, Maleic hydrazide and Perchlorate

Instrument parameters								
Ionization mode	ESI neg	ESI neg						
Column/temperature	Obelisc R 2.1 x 150 mm 5 µm 100 Å; (SIELC; OR-21.150.0510)							
Pre-filters	e.g. Supelco column saver 2	e.g. Supelco column saver 2.0 µm Filter						
Pre-column	Obelisc R 2.1 x 10mm 5 µm (SIELC; OR-21.G.0510)	Obelisc R 2.1 x 10mm 5 µm (SIELC; OR-21.G.0510)						
Eluent A	50 mmol NH ₄ -formate in Wa use brown glass bottles	ter + 0.1 % Formic acid						
Eluent B	Acetonitrile							
Gradient	%A	Flow [mL/min]	Time [min]					
	3	0.3	0					
	10	0.3	6					
	70	0.5	15					
	70	0.5	18					
	3	0.5	18.1					
	3	0.5	28					
Injection volume	5 μL							
Calibration standards and levels	For Maleic hydrazide (MH) a	one level at the reporting limit an additional level at 1 or 2 μ onsider that MH is typically on ons etc.)	g/mL may be useful as well,					
Acquired mass transitions	Compound	Mass Transition	ns (m/z)					
	Fosetyl-Al: Fosetyl-Al-D ₁₅ (ILIS): Maleic hydrazide:	109/81, 109/63 (114/82 (detected 111/82, 111/42,	• •					
	Maleic hydrazide-D ₂ (ILIS):	113/42	111/33, 111/03					
	Perchlorate: Perchlorate- ¹⁸ O ₄ (ILIS):	99/83, 101/85 107/89						

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

Note: It should be kept in mind that standards of isotopically labeled pesticides may contain small amounts of native (unlabelled) compounds as impurities. Typically these impurities are at low levels, so that the added amounts of native-pesticides, resulting from the addition of ISs, are insignificant. In the case of Maleic hydrazide (MH), however, the amount of IS added is comparably high due to the low detection sensitivity achieved for this compound. Assuming native MH being contained as impurity in D2-MH at 0.25 % the resulting concentration of native MH following the addition of 20 µg D2-MH to 10 g sample will be at 0.005 mg /kg sample. This aspect is to be considered when setting the Reporting Limits of MH as well as when judging residue levels in samples having low MRLs (e.g. baby food) or organic food.

For Perchlorate better run Method 1.3 or 1.4

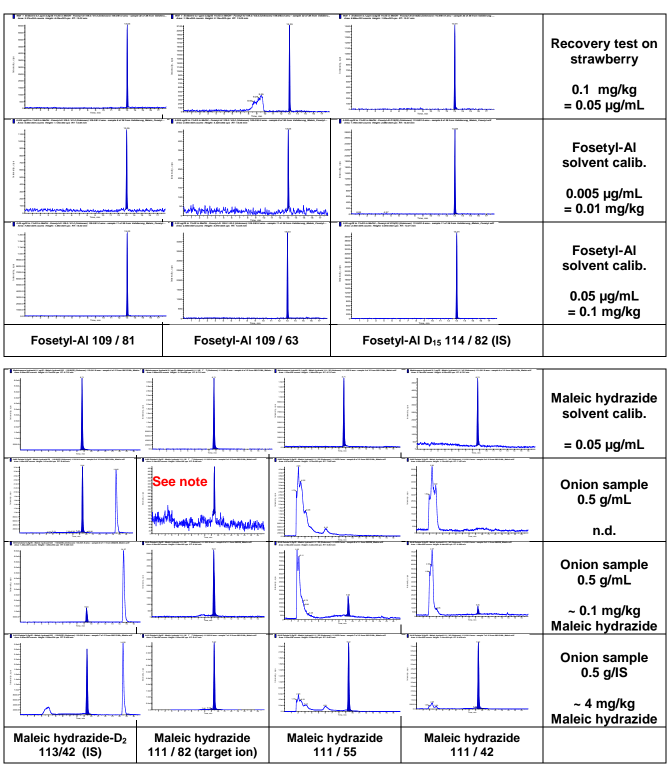


Figure 14: Typical chromatograms of Fosetyl-Al and Maleic hydrazide in various types of extracts and in pure solvent



5.7.6. Method 3 "Amitrole & Co"

Table 11: Proposed LC-MS/MS conditions for Amitrole, Chlormequat, Mepiquat, Daminozide, ETU, PTU, Trimesium, Difenzoquat and Cyromazine.

Instrument parameters	Conditions			
Ionisation mode	ESI pos			
Column/temperature	Obelisc R 2.1 x 150 mm 5 μm 100 Å (SIELC; OR-21.150.0510); 40°C			
Pre-column	Obelisc R 2.1 x 10 mm 5 μm (SIELC; OR-21.G.0510)			
Pre-filters	e.g. Supelco column saver 2.0 µm Filter			
Eluent A	5 mmol NH ₄ -formate in Water Use brown glass bottles			
Eluent B	5 mmol NH ₄ -formate Acetonitrile/Water 95 :5 (v/v)			
Gradient	% A	Flow [mL/min]	Time [min]	
	2	0.4	0	
	2	0.4	2.5	
	80	0.4	5	
	80	0.4	11	
	2	0.4	11.1	
	2	0.4	18	
Injection volume	5 μL			
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion* +	e.g. 0.05 or 0.1 µg/IS portion* + one level at the reporting limit		
Acquired mass transitions	Compound	Mass Tra	Mass Transitions (m/z)	
	Amitrole: Amitrole- ¹⁵ N (ILIS): Amitrole- ¹⁵ N ₂ , ¹³ C ₂ (ILIS):	85/43, 85/ 86/43 89/44		
	Chlormequat: Chlormequat-D ₄ (ILIS):	122/58, 12 126/58	122/58, 122/63, 124/58 126/58	
	Mepiquat: Mepiquat-D ₃ (ILIS):	114/98, 1 ⁻ 117/101	114/98, 114/58 117/101	
	Daminozide: Daminozide- ¹³ C ₄ (ILIS): Daminozide-D ₆ (ILIS):	161/143, 165/147 167/149		
	Cyromazine: Cyromazine-D ₄ (ILIS):	167/68, 10 171/86	167/68, 167/125, 167/85, 167/108, 171/86	
	ETU (Ethylenethiourea): ETU-D ₄ (ILIS):	103/44, 10 107/48	103/44, 103/60, 103/86 107/48	
	PTU (4-Methyl-2-imidazolidinet PTU-D6 (N,N'-Propylenethioure		117/100, 117/58, 117/60, 117/72 <i>(123/64)</i>	
	Trimethylsulfonium: Trimethylsulfonium-D ₉ (ILIS):	77/62, 77/ 86/68	77/62, 77/47 86/68	
	Difenzoquat: No ILIS currently available	249/77, 24 -	49/130, 249/193	

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

Note: For Paraquat, Diquat, Trimethylsulfonium and N,N-Dimethylhydrazine better run Method 4 (5.7.7)

^{**} The acronym PTU, commonly used for the propineb degradant 4-Methyl-2-imidazolidinethione (N,N'-iso-propylenethiourea), is also used for N,N'-propylenethiourea (= N,N'-Trimethylenethiourea). The IS tested corresponds to N,N'-propylenethiourea D6.

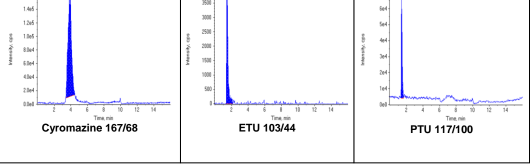


Figure 15: Typical chromatograms of Amitrole, Chlormequat, Mepiquat, Daminozide, ETU, PTU and Cyromazine in apple extract at 0.01 mg/kg



5.7.7. Method 4.1 "Quats & Co Obelisc R"

Table 12: Proposed LC-MS/MS conditions Diquat, Paraquat, Chlormequat, Mepiquat, Daminozide N,N-Dimethylhydrazine, Cyromazine, Trimethylsulfonium, Nereistoxin, Difenzoquat, Melamine and Propamocarb.

Instrument parameters	Conditions					
Ionisation mode	ESI pos	ESI pos				
Column/temperature	Obelisc R 2.1 x 150 mm 5 µm	Obelisc R 2.1 x 150 mm 5 µm 100 Å (SIELC; OR-21.150.0510); 40°C				
Pre-filters	e.g. Supelco column saver 2.0	e.g. Supelco column saver 2.0 µm Filter				
Pre-column	Obelisc R 2.1 x 10 mm 5 µm (SIELC; OR-21.G.0510)					
Eluent A		ter (adjust to pH 3 with Formi 20 mmol NH ₄ -formate in Wate				
Eluent B	Acetonitrile					
Gradient	% A	Flow [mL/min]	Time [min]			
	20	0.4	0			
	80	0.4	4			
	80	0.4	12			
	20	0.4	12.1			
	20	0.4	20			
Injection volume	10 μL					
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion* + one level at the reporting limit (use plastic vials if Paraguat and Diquat are within your scope!)					
Acquired mass transitions	Compound	Mass Transitions (n	Mass Transitions (m/z)			
104411041114111411	Diquat**: Diquat-D ₄ (ILIS):	184/128, 183/157, 184/ 188/160	184/128, 183/157, 184/156 188/160			
	Paraquat**: Paraquat-D ₆ (ILIS):	186/171, 171/77, 171/1 192/174	186/171, 171/77, 171/155 192/174			
	Chlormequat: Chlormequat-D ₄ (ILIS):	122/58, 122/63, 124/58 126/58	122/58, 122/63, 124/58 126/58			
	Mepiquat: Mepiquat-D₃ (ILIS):	114/98, 114/58 117/101	·			
	Daminozide: Daminozide- ¹³ C ₄ (ILIS): Daminozide-D ₆ (ILIS):	161/143, 161/61, 161/1 165/147 167/149				
	N,N-Dimethylhydrazine: N,N-Dimethylhydrazine-D ₆ (ILIS):	61/44, 61/45 67/49				
	Cyromazine: Cyromazine-D ₄ (ILIS):	167/68, 167/125, 167/8 171/86	167/68, 167/125, 167/85, 167/108, 171/86			
	Trimethylsulfonium: Trimethylsulfonium-D ₉ (ILIS):	77/62, 77/47 86/68				
	Nereistoxin: Nereistoxin-D ₆ (ILIS):	150/105, 150/61, 150/7 156/105	150/105, 150/61, 150/71 156/105			
	Difenzoquat: No ILIS currently available	249/77, 249/130, 249/1	93			
	Melamine: Melamine- ¹⁵ N ₃ (ILIS):	127/85, 127/68, (127/60 130/87	0)			
	Propamocarb: Propamocarb-D ₇ (ILIS):	189/144, 189/102, 189/ 196/103				

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

Note: For Morpholin, Diethanolamine (DEA) and Triethanolamine (TEA) better run Method 7 (5.6.9). As DEA converts to Morpholine in the ion source, chromatographic separation of these two is paramount. With Method 4.1 these two peaks do not sufficiently separate.

^{**} Diquat and Paraquat require special extraction conditions (see 5.2.5)

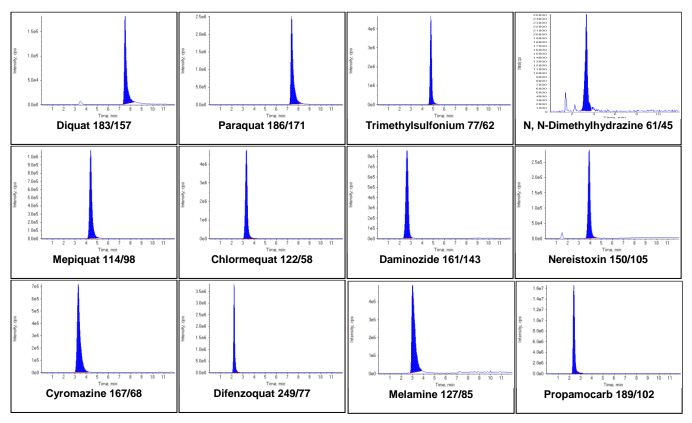


Figure 16: Typical chromatograms of Diquat, Paraquat, Chlormequat, Mepiquat, Daminozide, N,N-Dimethylhydrazine, Trimethylsulfonium, Cyromazine, Nereistoxin, Difenzoquat, Melamine and Propamocarb in apple extract at 0.1 mg/kg



5.7.8. Method 4.2 "Quats & Co BEH Amide"

Table 13: Proposed LC-MS/MS conditions for Aminocyclopyrachlor, Amitrole, Chlormequat, Chloridazon-desphenyl, Cyromazine, Daminozide, Diethanolamine, Difenzoquat, ETU, Melamine, Mepiquat, Mepiquat-4-hydroxy, Morpholine, Nereistoxin, Propamocarb, Propamocarb-N-desmethyl, Propamocarb-N-oxide, PTU, Triethanolamine, Trimesium (Trimethylsulfonium).

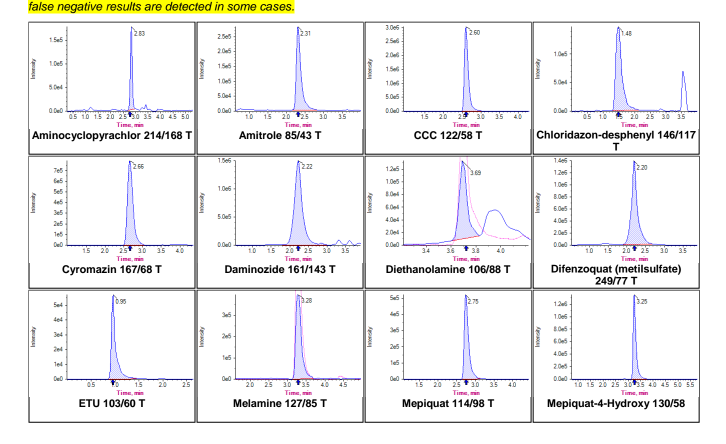
Instrument parameters	Conditions					
Ionisation mode	ESI pos.					
Column/temperature	BEH Amide 2.1 x 100mm 1.7 μm (P/N: 186004801); 40°C					
Pre-filters	e.g. Supelco column saver 2.0 µm Filter					
Pre-column	BEH Amide 1.7 μm (P/N: 186004799)					
Eluent A	50 mmol NH ₄ -formate in Wat	er (adjust to pH	3 with Formic	acid) Use brown glass!		
Eluent B	Acetonitrile					
Gradient	% A	Flow [ml	_/min]	Time [min]		
	3	0.5		0		
	3	0.5		0.5		
	30	0.5		4.0		
	60	0.5		5.0		
	60	0.5		6.0		
	3	0.5		6.1		
	3	0.5		10		
Injection volume	2 μL					
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion	* + one level at	the reporting	limit		
Acquired mass transitions	Compound		Mass Transi			
	Aminocyclopyrachlor:		214/170, 214/168, 214/101			
	Amitrole: Amitrole- ¹⁵ N (ILIS): Amitrole- ¹⁵ N ₂ ¹³ C ₂ (ILIS):		85/43, 85/57, 85/58 86/43 89/44			
	Chlormequat: Chlormequat-D ₄ (ILIS):		122/58, 124/58, 122/63 126/58			
	Chloridazon-desphenyl: Chloridazon-desphenyl- ¹⁵ N ₂ (ILIS):		146/117, 146/101, 146/66 148/117			
	Cyromazine: Cyromazine-D ₄ (ILIS):		167/68, 167/125, 167/108, 167/85 171/86			
	Daminozide: Daminozide- ¹³ C ₄ (ILIS); Daminozide-D ₆		161/143, 161/61, 161/101 , 161/115, 161/44			
	Diethanolamine*** (DEA): Diethanolamine-D4 (ILIS):		106/88, 106/70, 106/45 110/92			
	Difenzoquat: No ILIS currently available		249/130, 249/77, 249/193, -			
	ETU (Ethylenethiourea): ETU-D ₄ (IS):		103/60, 103/44, 103/86 107/48			
	Melamine: Melamine- ¹⁵ N ₃ (ILIS):		127/85, 127/68, (127/60) 130/87			
	Mepiquat: Mepiquat-D ₃ (ILIS):		114/98, 114/58 117/101			
	Mepiquat-4-hydroxy:		130/58, 130/5	9 <mark>6, 130/114</mark>		
	Morpholine***: Morpholine-D ₈ (ILIS):		88/70, 88/45, 88/44 96/78			
	Nereistoxin: Nereistoxin-D ₆ (ILIS):		150/105, 150/61, 150/71 156/105			

Propamocarb: Propamocarb-D ₇ (ILIS):	189/144, 189/74, 189/102 196/103
Propamocarb-N-desmethyl:	175/102, 175/144, 175/74
Propamocarb-N-oxide:	205/102, 205/144, 205/74
PTU (4-Methyl-2-imidazolidinethione)**: PTU-D6 (N,N'-Propylenethiourea-D ₆):	117/100, 117/58, 117/60, 117/72 123/64
Triethanolamine*** (TEA): Triethanolamine-D ₁₂ (ILIS):	150/132, 150/70, 150/88 162/144
Trimethylsulfonium: Trimethylsulfonium-D ₉ (ILIS):	77/62, 77/47 86/68

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

**The acronym PTU, commonly used for the propineb degradant 4-Methyl-2-imidazolidinethione (N,N'-iso-propylenethiourea), is also used for N,N'-propylenethiourea (= N,N'-Trimethylenethiourea). The IS tested corresponds to N,N'-propylenethiourea D₆.

***For Morpholin, Diethanolamine and Triethanolamine better run Method 7 (5.6.9) as these compounds seem to be strongly suppressed by matrix using these LC-conditions. This effect is reduced if the extract is diluted e.g. 5/10 fold. For Diethanolamin even



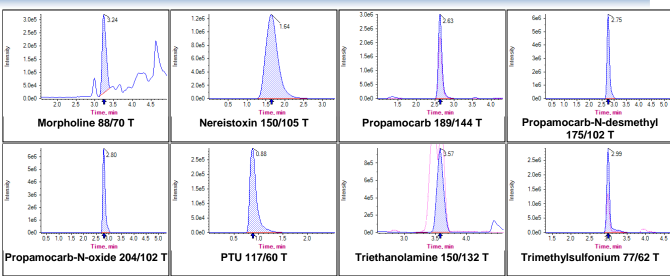


Figure 17: Typical chromatograms of Aminocyclopyrachlor, Amitrole, Chlormequat, Chloridazon-desphenyl, Cyromazine, Daminozide, Diethanolamine, Difenzoquat, ETU, Melamine, Mepiquat, Mepiquat-4-hydroxy, Morpholine, Nereistoxin, Propamocarb, Propamocarb-N-desmethyl, Propamocarb-N-oxide, PTU, Triethanolamine, Trimesium (Trimethylsulfonium) in tomato extracts spiked at 0.05 mg/kg.



5.7.9. Method 5 "Quats & Co. MonoChrom MS"

Table 14: Proposed alternative LC-MS/MS conditions for Chlormequat and Mepiquat

Instrument parameters	Conditions				
Ionisation mode	ESI pos				
Column/temperature	MonoChrom MS 100x2 mm; 5	µm (Varian); at 40°C			
Eluent A	5 mmol/L NH ₄ -acetate + 0.1% A	Acetic acid in Water			
Eluent B	Acetonitrile				
Gradient	% A	Flow [mL/min]	Time [min]	
	5	0.4		0	
	95	0.4		2	
	95	0.4		5	
	5	0.4		5.1	
	5 0.4 15				
Injection volume	5 μL				
Calibration standards and levels	e.g. 0.05 or 0.1 μg/IS portion*+	one level at the repor	ting limit		
Acquired mass transitions	Compound		Mass Tra	ansitions (m/z)	
	Chlormequat: Chlormequat-D ₄ (ILIS):		122/58, 122/63, 124/58 126/58		
	Mepiquat: Mepiquat-D ₃ (ILIS):		114/98, 114/58 117/101		
	Difenzoquat: No IS currently available		249/77, 249/130, 249/193 -		
	ETU (Ethylenethiourea): ETU-D ₄ (ILIS):		103/44, 103/60, 103/86 107/48		
	PTU (4-Methyl-2-imidazolidinethione)**: PTU-D6 (N,N'-Propylenethiourea-D ₆):		117/100, 117/58, 117/60, 117/72 123/64		

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

For more information on method 5 please refer to the following document within the EURL homepage: http://www.crl-pesticides.eu/library/docs/srm/meth_ChlormequatMepiquat_CrlSrm.pdf

^{**}The acronym PTU, commonly used for the propineb degradant 4-Methyl-2-imidazolidinethione (N,N'-iso-propylenethiourea), is also used for N,N'-propylenethiourea (= N,N'-Trimethylenethiourea). The IS tested corresponds to N,N'-propylenethiourea D_6 .



5.7.10. Method 6 "Streptomycin and Kasugamycin"

Table 15: Proposed LC-MS/MS conditions Streptomycin and Kasugamycin

Instrument parameters	Conditions	Conditions				
Ionisation mode	ESI pos	ESI pos				
Column	Obelisc R 2.1 x 150 mm 5µm (SIELC; OR-21.150.0510); 4					
Pre-filters	e.g. Supelco column saver 2	.0 μm Filter				
Pre-column	Obelisc R 2.1 x 10 mm 5 μm (SIELC; OR-21.G.0510)					
Eluent A	0.1% Formic acid in Water					
Eluent B	0.1% Formic acid in acetonitrile					
Gradient	%A Flow [mL/min] Time [min]					
	20	0.3	0			
	20	0.3	8			
	<mark>20</mark>	0.3	<mark>13</mark>			
	80	<mark>0.5</mark>	<mark>18</mark>			
	<mark>80</mark>	<mark>0.5</mark>	<mark>23</mark>			
Injection volume	20 μL; dwell time increased t	o 200 ms				
Calibration standards and levels		e.g. 0.05 or 0.1 µg/IS portion* one level at the reporting limit (use plastic vials if Streptomycin is within your scope)				
Acquired mass transitions	Compound	Compound Mass Transitions (m/z)				
	Streptomycin: Dihydrostreptomycin (IS):					
	Kasugamycin: No IS currently available	380/112, 380/200 -				

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

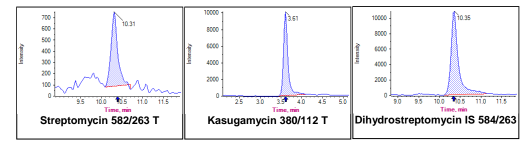


Figure 18: Typical chromatograms of Streptomycin and Kasugamycin in apple extracts spiked at 0.01 mg/kg.



5.7.11. Method 7 "Morpholine, Diethanolamine and Triethanolamine"

Table 16: Proposed LC-MS/MS conditions Morpholine, Diethanolamine and Triethanolamine

Instrument parameters	Conditions				
Ionisation mode	ESI pos				
Column	Dionex Acclaim Trinity P1	2.1 x 10	00 mm (3 μm) (P/N 0	71389); 40°C	
Pre-filters	e.g. Supelco column saver 2	2.0 µm F	Filter		
Pre-column	Dionex Acclaim Trinity P1	2.1 x 10	0 mm (3 μm) (P/N 07	1391)	
Eluent A	50 mmol NH ₄ -formate in Wa Use brown glass bottles!	iter (adji	ust to pH 3 with Form	ic acid)	
Eluent B	Acetonitrile				
Gradient	%A	F	Flow [mL/min]	Time [min]	
	10		0.4	0	
	10		0.4	10	
Injection volume	5 μL				
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion	n+ one I	level at the reporting I	imit	
	Compound		Mass Transitions (r	n/z)	
Acquired mass transitions	Morpholine: Morpholine-D ₈ (IS):		88/70, 88/45, 88/44 96/78		
Acquired made transitions	Diethanolamine (DEA): Diethanolamine-D ₄ (IS).		106/88, 106/70, 106/45 110/92		
	Triethanolamine (TEA). Triethanolamine-D ₁₂ (IS):		150/132, 150/70, 150/88 162/144		

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

Morpholin, DEA and TEA are not pesticides, they are additive of waxes used to coat crops (citrus, apples and mangoes etc). They are included in this method for the sake of convenience and synergy. As these three compounds can be analyzed very sensitively 5-10-fold dilution of the extracts before injection is recommendable where possible, especially in absence of an IS requiring standard additions approach (5.5.3)

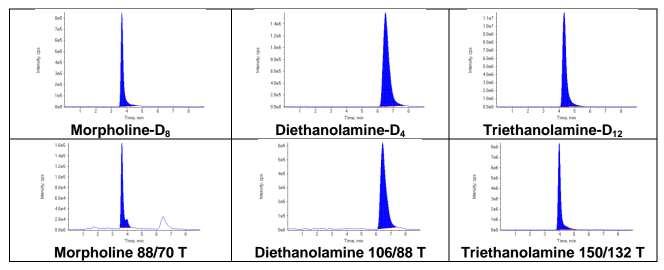


Figure 19: Typical chromatograms of Morpholine, Diethanolamine and Triethanolamine in apple extracts at 0.05 mg/kg (extract were diluted 10-fold before injection)



5.7.12. Method 8 "Triazole derivative metabolites (TDMs)"

Table 17: Proposed LC-MS/MS conditions 1,2,4-Triazole, Triazole-alanine, Triazole-acetic acid, Triazole-lactic acid and 1,2,3-Triazole

Instrument parameters	Conditions					
Ionisation mode	ESI pos					
Column	Hypercarb 2.1 x 100 mm 5 μm (P/N 35005-102130); 40°C					
Pre-column	Hypercarb Guard 2.1 x 10 mm	n 5 μm (P/N 35005-102101)				
Pre-filter	e.g. Supelco column saver 2.0 µr	m Filter (optional)				
Eluent A	1% Acetic acid in Water + 5% Me	eOH				
Eluent B	1% Acetic acid in MeOH					
	%A	Flow [mL/min]	Time	[min]		
	100	0.6		0		
Gradient	10	0.6		5		
	10	0.6		6		
	100 0.6 6.1					
	100	0.6	•	10		
Injection volume	2 µL					
Calibration standards and levels	e.g. 0.05 or 0.1 µg/IS portion* one	e level at the reporting limit				
	Compound**	Mass Transitions (m/z)	Selexion Q-Tr DMS-Condition (DMS temperation)	ons		
		,	COV (V)	SV (V)		
	1,2,4-Triazole ^{#:} 1,2,4-Triazole- ¹³ C ₂ , ¹⁵ N ₃ (IS).	70/43, 70/70 75/46	-10 -13.75	2600 3000		
Acquired mass transitions	Triazole-alanine: Triazole-alanine- ¹³ C ₂ , ¹⁵ N ₃ (IS):	157/70, 157/88, 157/42 162/75	-2.0 -1.75	3000 3100		
	Triazole-acetic acid: 128/70, 128/43, 128/73 Triazole-acetic acid- ¹³ C ₂ , ¹⁵ N ₃ (IS): 133/75		-6.0 -6.0	3100 3500		
	Triazole-lactic acid: Triazole-lactic acid- ¹³ C ₂ , ¹⁵ N ₃ (IS):	158/70, 158/43, 158/112 : 163/75	-3.0 -2.25	3300 3500		
	1,2,3-Triazole [#] : No IS currently available	70/43 -	<mark>-12</mark> -	<mark>3000</mark> -		

^{*} One IS portion is the absolute IS-mass contained in the prepared calibration standard solution (see also Table 1).

^{# 1,2,4-}Triazole and 1,2,3-Triazole are used as nitrification inhibitors in fertilizers

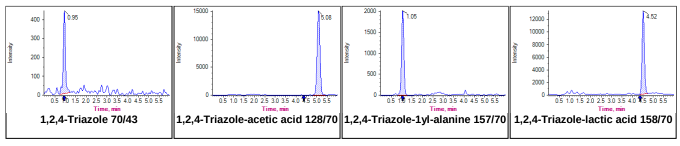


Figure 20: Typical chromatograms of TDMs in avocado extracts spiked at 0.01 mg/kg.

^{**} All ILISs were a friendly donation and are at the time not commercially available.

^{***} DMS condition differ to some extent from instrument to instrument (further parameters: CUR 20, GS1 60, GS2 70, DMO -3.0)



5.8. Calibration and Calculations

5.8.1. **Using IS**

5.8.1.1. Where IS is added to the sample before any aliquotation:

The following calculation approach requires that the ratio of the IS masses added to the test portions (5.2.3) and to the calibration standard(s) (5.5) (m_{IS}^{sample} / $m_{IS}^{cal\ mix}$) is known and constant. By keeping the IS constant throughout the calibration levels the peak ratio $PR^{cal\ mix}$ ($A_{pest}^{cal\ mix}$ / $A_{IS}^{cal\ mix}$) of each calibration level can be plotted against the absolute mass of the pesticide $m_{pest}^{cal\ mix}$ rather than the ratio $m_{pest}^{cal\ mix}$ / $m_{IS}^{cal\ mix}$ (the $m_{IS}^{cal\ mix}$ is set as 1).

The calibration graph (to be plotted for each pesticide separately) is described by the following formula:

$$PR^{calmix} = a_{cal} \times m_{pest}^{calmix} + b_{cal}$$
 (1)

The mass fraction (w_R) of a given pesticide in a given sample can be calculated as follows using the respective peak ratio of pesticide and internal standard obtained from the sample extract ($PR^{sample} = A_{pest}^{Sample} / A_{IS}^{Sample}$), the correction factor ($m_{IS}^{sample} / m_{IS}^{cal mix}$) as well as the weight of the test portion (m_a).

$$w_{R} = \frac{(PR^{Sample} - b_{cal})}{a_{cal}} \times \frac{1}{m_{a}} \times \frac{m_{ISTD}^{Sample}}{m_{ISTD}^{cal mix}} \left(\frac{\text{mg}}{\text{kg}}\right)$$
(2)

Reasonably (but not necessarily) the calibration standards should be prepared in such a way that the ratio $m_{IS}^{sample} / m_{IS}^{cal \ mix}$ equals 20 (the assumed volume ratio of sample extract versus calibration standard). The absolute masses of the IS-WS I and II do not need to be necessarily known (see also the notes of Table 1).

5.8.1.2. Where IS is added to an aliquot of the extract

When adding the IS to an aliquot of the extract (e.g. 1 mL) the knowledge of the exact total volume of the sample extract becomes important. Water adjustment is thus essential and if it is done as described in **5.2.2** and **Table18**, the total volume can be assumed to be exactly 20 mL. In this case 1 mL sample extract will correspond to $1/20^{th}$ of the test portion (m_a). The mass of the IS to be added to an aliquot (m_{IS} aliquot) should be scaled according to the aliquot volume used (V_{aliquot}) with the IS mass ratio (m_{IS} aliquot / m_{IS} cal mix) being important for the calculation. Reasonably (but not necessarily) m_{IS} aliquot should be derived using the following formula m_{IS} aliquot = m_{IS} sample x V_{aliquot}/20 , with m_{IS} sample being the IS mass that would have been added to the entire sample portion according to **5.2.2** and **Table 18**.

Following the above, the mass fraction (w_R) of a given pesticide in a given sample can be calculated as follows using the respective peak ratio of pesticide and internal standard obtained from the sample extract ($PR^{sample} = A_{pest}^{sample} / A_{IS}^{sample}$), the correction factor ($m_{IS}^{aliquot} / m_{IS}^{cal mix}$) as well as the weight of the sample equivalents in the aliquot ($m_{aliquot} = m_a \times V_{aliquot}/20$).



$$w_R = \frac{(PR^{sample} - b_{cal})}{a_{cal}} \times \frac{1}{m_{aliquot}} \times \frac{m_{ISTD}^{aliquot}}{m_{ISTD}^{cal mix}} \left(\frac{\text{mg}}{\text{kg}}\right)$$
 (3)

Variables used

Mass of pesticide in calibration mixture	$m_{\it pest}^{\it calmix}$	μg
Mass of pesticide in final extract	$m_{\it pest}^{\it sample}$	μg
Mass of internal standard in calibration mixture	m ^{cal mix} ISTD	μg
Mass of internal standard added to test portion (sample)	m_{ISTD}^{sample}	μg
Mass of internal standard added to aliquot of sample extract	$m_{ISTD}^{aliquot}$	μg
Volume of sample extract aliquot used (5.8.1.2 and 5.5.3) to spike the IS or for standard additions	$V^{\it aliquot}$	mL
Mass of test portion	m_a	g
Mass of test portion represented in an aliquot	m _{aliquot}	g
Mass fraction of pesticide in the sample	W_R	mg/kg
Peak area of pesticide obtained from calibration standard (mixture)	$A_{pest}^{cal\ mix}$	(counts)
Peak area of IS obtained from calibration standard (mixture)	A_{ISTD}^{calmix}	(counts)
Peak area of pesticide obtained from the injected extract	A_{pest}^{sample}	(counts)
Peak area of IS obtained from the injected extract	$A_{ISTD}^{ sample}$	(counts)
Peak ratio of pesticide vs. IS obtained from calibration mixture	PR cal mix	(dimensionless)
Peak ratio of pesticide vs. IS obtained from injected extract	PR sample	(dimensionless)
Slope of calibration graph	a cal	(dimensionless)
Bias of calibration graph (intercept)	$b_{\it cal}$	(dimensionless)



5.8.2. Not using IS

If no appropriate ISs are used it is of high importance to properly compensate for matrix effects. For the compensation of matrix effects matrix-matched calibrations (5.5.2) and the standard additions approach (5.5.3) are recommended. In both cases the assumption is made that the total volume of the sample extract is exactly 20 mL. Adjustment of the Water content (and extract volume) in the sample is thus paramount.

5.8.2.1. Calculations when employing matrix-matched calibration without IS

The calibration graph (to be plotted for each pesticide separately) is described by the following formula:

$$A_{pest}^{cal\;mix} = a_{cal} \times C_{pest}^{cal\;mix} + b_{cal}$$
 (1)

The mass fraction (w_R) of a given pesticide in a given sample can be calculated as follows using the respective peak area of the pesticide obtained from the sample extract (A_{pest}^{sample}) and a correction factor (V) as well as the weight of the test portion (m_a).

$$w_{R} = \frac{(A_{pest}^{Sample} - b_{cal})}{a_{cal}} \times \frac{1}{m_{a}} \times V_{end} \left(\frac{\text{mg}}{\text{kg}}\right)$$
 (2)

where V_{end} is the total volume of the sample extract (20 mL).

All other variables are listed in 5.8.1.2.

5.8.2.2. Calculations when employing the standard additions approach

The standard additions approach is the method of choice where no appropriate IL-IS is available. This approach typically compensates matrix effect better than the matrix-matched calibrations (**5.5.2**). The mass fraction of the pesticide in the sample (w_R) is calculated via linear regression using a graphical presentation as shown in Figure 1. The Y-intercept of the calibration graph will indicate the pesticide mass contained in the non-fortified aliquot of the sample extract.

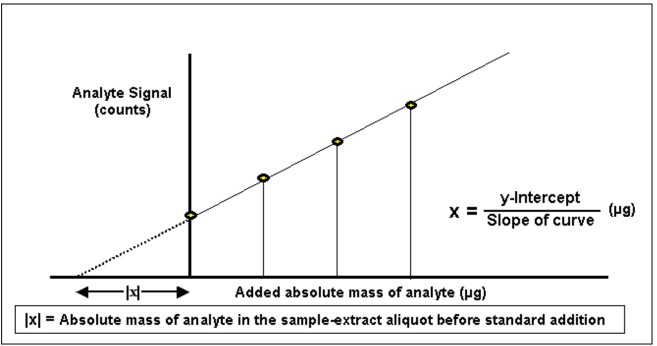


Figure 21: Internal calibration using the procedure of standard additions, schematically

Key:

Y Peak area of analyte

X Added absolute mass of analyte $m_{pest}^{std\ add}$ in μg

absolute amount of analyte in the sample extract (in μ g) before standard addition (y = 0)

With
$$x = \frac{y - \text{int } iercept (b)}{slope \ of \ the \ curve (a)}$$
 (µg)

The calculation is performed as follows using the regression graph shown in

$$w_R = \frac{b}{a} \times \frac{V_{end}}{V_{al} \times m_a} \left(\frac{\text{mg}}{\text{kg}} \right)$$

where:

b Y-intercept of the calibration graph of the analyte in question;

a Slope of the calibration graph of the analyte in question $(1/\mu g)$;

V_{end} Volume of sample extract (mL) (should be 20 mL)

Volume of aliquots used for the standard additions approach (mL)

 m_a Weight of initial sample portion (g)



6. Performance Data

Validation data of the presented methods according to SANTE/11945/2015 guidance document are shown at the EURL validation database at www.eurl-pesticides-datapool.eu. Exemplary LOQs of the presented methods are listed in **Table 18.**

Table 18: Overview of approximate limits of quantification (LOQs)^a

Method	Pesticide	Most fruits and Veg-	Citrus	Cereals
		etables (tested on	(tested on Orange)	(tested on Barley,
		Tomato, Cucumber,	[mg/kg]	Oat, Rice) [mg/kg]
		Apples) [mg/kg]		
M1.1/M1.2/M1.3	Ethephon	0.01/0.01/0.01	0.01/0.01/0.01	0.02/0.02/0.02
M1.1/M1.2/M1.3	HEPA	0.01/0.01/0.02	0.01/0.01/0.02	0.02/0.02/0.02
M1.1/M1.2/M1.3	Glyphosate	0.01/0.01/0.02	0.02/0.01/0.02	0.02/0.02/0.02
M1.1/M1.2/M1.3	AMPA	0.01/0.01/0.02	0.02/0.01/0.02	0.02/0.02/0.02
M1.1/M1.2/M1.3	Glufosinate	0.01/0.01/0.02	0.02/0.02/0.02	0.02/0.02/0.02
M1.1/M1.2/M1.3	MPPA	0.01/0.01/0.01	0.02/0.02/0.01	0.02/0.02/0.02
M1.1/M1.2/M1.3	N-Acetyl-glufosinate	0.02/0.02/0.01	0.02/0.02/0.01	0.02/0.02/0.02
M1.2/M1.3	N-Acetyl-AMPA	0.01/0.01	0.01/0.01	0.02/0.02
M1.3/ <mark>M1.4</mark> /M2	Perchlorate ^b	0.01/ <mark>0.005</mark> /0.01	0.01/ <mark>0.01</mark> /0.01	0.01/ <mark>0.01</mark> /0.01
M1.2/M1.3 ^b /M1.4 ^b	Phosphonic acid	0.1/0.1/ <mark>0.05</mark>	0.1/0.1/ <mark>0.05</mark>	0.1/0.1/ n.a.
M1.3/ <mark>M1.4</mark>	Chlorate ^b	0.01/ <mark>0.005</mark>	n.a./ <mark>0.005</mark>	n.a./ <mark>n.a.</mark>
M1.4	Bromide ^c	1	1	n.a.
M1.4	Bromate ^b	0.02	0.02	<mark>n.a.</mark>
M1.1/M1.3/M2	Fosetyl	0.1/0.005/0.005	n.a. / 0.005/0.005	n.a. / 0.005/ 0.005
M2/M1.3	Maleic hydrazide	0.01/0.01	0.01/0.01	0.02/0.02
M3/M4.2	Amitrole	0.01/ <mark>0.01</mark>	0.01/ <mark>0.01</mark>	0.02/ <mark>0.02</mark>
M3/ <mark>M4.2</mark> /M5	ETU	0.01/ <mark>0.05</mark> /0.01	0.02/ <mark>0.05</mark> /n.a.	0.02/ <mark>0.1</mark> /n.a.
M3/M4.2/M5	PTU	0.01 <mark>/0.05</mark> /0.01	0.02/ <mark>0.05</mark> /n.a.	0.02/ <mark>0.1</mark> / n.a.
M3/M4.1/ <mark>M4.2</mark> / M5	Chlormequat	0.005/0.005/ <mark>0.01</mark> /0.01	0.005/0.005/ <mark>n.a.</mark> /0.01	0.01/0.01/ <mark>0.02</mark> /0.01
M3/M4.1/ <mark>M4.2</mark> / M5	Mepiquat	0.005/0.01/ <mark>0.01</mark> /0.01	0.005/0.01/ <mark>n.a.</mark> /0.01	0.001/0.02/ <mark>0.02</mark> /0.02
M3/M4.1/ <mark>M4.2</mark>	Cyromazine	0.01/0.01/ <mark>0.01</mark>	0.01/0.01/ <mark>n.a.</mark>	0.02/0.02/ <mark>0.02</mark>

Method	Pesticide	Most fruits and Veg-	Citrus	Cereals
		etables (tested on	(tested on Orange)	(tested on Barley,
		Tomato, Cucumber, Apples) [mg/kg]	[mg/kg]	Oat, Rice) [mg/kg]
M3/M4.1/ <mark>M4.2</mark>	Daminozide	0.01/0.02/ <mark>0.01</mark>	0.01/0.02/ <mark>n.a.</mark>	0.02 /0.04/ <mark>0.02</mark>
M3/M4.1/ <mark>M4.2</mark>	Trimethylsulfonium-Cation	0.01/0.005/ <mark>0.01</mark>	0.01/0.005/ <mark>n.a.</mark>	0.02/0.01/ <mark>0.02</mark>
M3/M4.1/ <mark>M4.2</mark>	Nereistoxin	0.01/0.01/ <mark>0.01</mark>	n.a./n.a./ <mark>n.a.</mark>	n.a./n.a./ <mark>0.02</mark>
M3/M4.1/ <mark>M4.2</mark>	Propamocarb	n.a./n.a./ <mark>0.01</mark>	n.a./n.a./ <mark>n.a.</mark>	n.a./n.a./ <mark>0.02</mark>
M4.1	N,N-Dimethylhydrazine	0.005	0.005	0.01
M4.1	Diquat	0.005	0.005	0.005
M4.1	Paraquat	0.005	0.005	0.005
M4.1/ <mark>M4.2</mark>	Melamine	n.a./ <mark>0.01</mark>	n.a./ <mark>n.a.</mark>	n.a./ <mark>0.02</mark>
M4.2	Aminocyclopyrachlor	0.01	<mark>n.a.</mark>	0.02
M4.2	Chloridazon-desphenyl	0.01	<mark>n.a.</mark>	0.02
M4.2	Difenzoquat	0.01	<mark>n.a.</mark>	0.02
M4.2	Mepiquat-4-hydroxy	0.01	<mark>n.a.</mark>	0.02
M4.2	Propamocarb-N- desmethyl	0.01	<mark>n.a.</mark>	0.02
M4.2	Propamocarb-N-oxide	0.01	n.a.	0.02
M6	Streptomycin	0.005	n.a.	n.a.
M6	Kasugamycin	0.01	n.a.	n.a.
M7	Morpholine ^d	0.01	0.01	n.a.
M7	Diethanolamine ^d	0.01	0.01	n.a.
M7	Triethanolamine ^d	0.01	0.01	n.a.
M8	1,2,4-Triazole	0.01	0.01	0.02
M8	Triazole-alanine	0.01	0.01	0.02
M8	Triazole-acetic acid	0.01	0.01	0.02
M8	Triazole-lactic acid	0.01	0.01	0.02
M8	1,2,3-Triazole	n.a.	<mark>n.a.</mark>	<mark>n.a.</mark>

using Q-Trap Sciex 5500 instrument;

b value derived from 5-fold diluted extract (0.1 g sample equivalents/mL)

value derived from 250-fold diluted extract (0.002 g sample equivalents/mL) d value derived from 10-fold diluted extract (0.05g sample equivalents/mL)

References

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Kolberg DI, Mack D, Anastassiades M, Hetmanski MT, Fussell RJ, Meijer T, Mol HG. Anal Bioanal Chem. 404(8):2465-74 (2012); Development and independent laboratory validation of a simple method for the determination of paraguat and diguat in potato, cereals and pulses

Alder L. and Startin J. R. (2005); Determination of Chlormequat and Mepiquat in Foods by Liquid Chromatography/Mass Spectrometry or Liquid Chromatography/Tandem Mass Spectrometry: Interlaboratory Study; Journal of AOAC International Vol. 88, No. 6: 1762-1776

Vahl, M. et al. (1998); Analysis of Chlormequat residues in grain using liquid chromatography-mass spectrometry (LC-MS/MS); Fresenius J Anal Chem 361:817-820



8. ANNEX

Table 19: Conversion factors between typical purchased standards and target analytes (3.15):

Compound	MW [g/mol]	Compound as sold	MW [g/mol]	Conversion factor
Bialaphos	323.3	Bialaphos-sodium	345.3	1.07
Bromate	<mark>127.9</mark>	Bromate-potassium	<mark>167.0</mark>	1.31
Bromide	<mark>79.9</mark>	Bromide-potassium	119.0	1.49
Chlorate (anion)	83.6	Chlorate-sodium	106.1	1.27
Chlormequat (cation)	117.6	Chlormequat-chloride	158.1	1.34
Chlormequat-D ₄ (cation)	121.6	Chlormequat-D ₄ -chloride	162.1	1.33
Difenzoquat (cation)	249.3	Difenzoquat-methylsulfate	360.4	1.45
Diquat (dication)	184.2	Diquat-dibromide-monohydrate	362.1	1.97
Diquat-D ₄ (dication)	188.2	Diquat-D ₄ -dibromide-monohydrate	366.1	1.95
Fosetyl	110.0	Fosetyl-Al	118.0	1.07
Foodbyl D	115.0	Fosetyl-D ₅ -1/3aluminium	123.0	1.07
Fosetyl-D₅	115.0	Fosetyl-D ₅ -sodium	137.0	1.19
Glufosinate	182.2	Glufosinate-ammonium	198.2	1.09
Glufosinate-D ₃	185.1	Glufosinate-D ₃ -hydrochloride	220.6	1.19
Kasugamycin	379.4	Kasugamycin-hydrochloride- monohydrate	433.8	1.14
Mepiquat (cation)	114.2	Mepiquat-chloride	149.7	1.31
Mepiquat-D ₃ (cation)	117.2	Mepiquat-D ₃ -iodide	244.1	2.08
Mepiquat-4-hydroxy	130.2	Mepiquat-4-hydroxy-chloride	<mark>165.7</mark>	1.27
N. N-Dimethylhydrazine-D ₆	66.1	Dimethylhydrazine-D ₆ hydrochloride	102.6	1.55
N-Acetyl-glufosinate	223.2	N-Acetyl-glufosinate-disodium	267.2	1.20
N-Acetyl-glufosinate-D ₃	226.2	N-Acetyl-glufosinate-D ₃ -disodium	270.2	1.19
Nereistoxin	149.3	Nereistoxin-oxalate	239.3	1.60
Nereistoxin-D ₆	155.3	Nereistoxin-D ₆ -oxalate	245.3	1.58
Paraquat (dication)	186.3	Paraquat-dichloride	257.2	1.38
Paraquat-D ₆ (dication)	192.3	Paraquat-D ₆ -diiodide	446.0	2.32
Propamocarb-N-oxide	205.2	Propamocarb-N-oxide x HCl	240.7	1.17
Streptomycin	581.6	Streptomycin-sesquisulfate	728.7	1.25
Trimethylsulfonium	77.2	Trimethylsulfonium-iodide	204.1	2.64
Trimethylsulfonium-D ₉	86.2	Trimethylsulfonium-D ₉ -iodide	213.1	2.47



Table 20: Exemplary concentrations of pesticide stock and working solutions (3.15 and 3.16), solvent proposals also apply to ILISs (see 3.18, 3.19 and 3.20).

Compound	Stock Solution (exemplary)		Working Solutions including mixtur (exemplary)	res
	Solvent used to prepare	[mg/mL]	Solvent used to prepare	[µg/mL]
Aminocyclopyrachlor	Methanol	1	Methanol	5/1/0.1
Amitrole	Methanol	1	Methanol	5/1/0.1
AMPA	Water/Methanol (3:1)	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
Bromate	Water/Methanol (50:50)	1	Methanol	5/1/0.1
Bromide	Methanol	1	Methanol	5/1/0.1
Chlorate	Methanol	1	Methanol + 1% Formic acid	5/1/0.1
Chloridazon-desphenyl	Methanol	1	Methanol	5/1/0.1
Chlormequat	Methanol	1	Methanol	5/1/0.1
Cyromazine	Methanol	1	Methanol	5/1/0.1
Daminozide	Methanol	1	Methanol	5/1/0.1
Diethanolamine	Acetonitrile	1	Methanol	5/1/0.1
Difenzoquat	Acetonitrile	1	Methanol	5/1/0.1
Diquat*	Methanol + 1% Formic acid	1	Methanol + 1% Formic acid	5/1/0.1
Ethephon	Methanol + 1% Formic acid	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
ETU	Methanol	1	Methanol	5/1/0.1
Fosetyl	Water / methanol (3:1)	0.1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
Glufosinate	Water / methanol (10:1)	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
Glyphosate*	Water / methanol (3:1)	0.2	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
HEPA	Methanol	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
Kasugamycin	Methanol	1	Methanol	5/1/0.1
Maleic hydrazide	Methanol	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
Mepiquat	Methanol	1	Methanol	5/1/0.1
Mepiquat-4-hydroxy	Methanol	1	Methanol	5/1/0.1
Morpholine	Methanol	1	Methanol	5/1/0.1
MPPA	Water	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1
N,N-Dimethylhydrazine	Methanol	1	Methanol	5/1/0.1
N-Acetyl- AMPA	Methanol Methanol	1	Water/Methanol+1% Formic acid (50:50)	5/1/0.1

Compound	Stock Solution (exemplary)		Working Solutions including mixtures (exemplary)	Working Solutions including mixtures (exemplary)		
·	Solvent used to prepare	[mg/mL]	Solvent used to prepare	[µg/mL]		
N-Acetyl-glufosinate	Methanol	1	Methanol + 1% Formic acid	5/1/0.1		
Nereistoxin	Methanol / Water (3:1)	1	Methanol	5/1/0.1		
Paraquat**	Methanol	1	Methanol	5/1/0.1		
Perchlorate	Methanol	1	Methanol	5/1/0.1		
Phosphonic acid	Water (Acetonitrile for the ILIS)	1	Methanol	5/1/0.1		
Propamocarb	Acetonitrile	1	Methanol			
Propamocarb-N-desmethyl	Acetonitrile:Aceton (1 mL Aceton to initially dissolve)	1	Methanol	5/1/0.1		
Propamocarb-N-oxid	Methanol	1	Methanol Methanol			
PTU	Methanol	1	Methanol	5/1/0.1		
Streptomycin*	Water / methanol (1:1)	1	Methanol	5/1/0.1		
Triethanolamine	Methanol	1	Methanol	5/1/0.1		
Trimethylsulfonium (tri- mesium)	Methanol	1	Methanol	5/1/0.1		

^{*} Use plastic vessels and stoppers for compounds that tend to interact with glass surfaces



Table 21: Exemplary providers of isotopically labeled internal standards 3.17.

.,			A (1 1 N)	Conc.	Amount		Prices in €-cent		
Name		Source	Article-No.	[µg/mL]	per unit	1 unit	2 μg* 0.1 μg**		
	¹⁵ N	1	XA10240100ME	100	1.1 mL	165 €	300 c	15 c	
A '. I	¹⁵ N ¹³ C	1	XA10240110AL	100	1.1 mL	332€	604 c	30 c	
Amitrole	$^{15}N_2$ $^{13}C_2$	7	A633382		10 mg	1496€	30 c	1.5 c	
	$^{15}N_4$ / $^{13}C_2$	8	C4313		10 mg				
	130 151	1	CIL-CDNLM-6786-1.2	100	1.2 mL	464 €	773 c	39 c	
	¹³ C ₂ . ¹⁵ N ₂	5	CDNLM-6786-1.2	100	1.2 mL	464 €	773 c	39 c	
AMPA	¹³ C. ¹⁵ N	7	A617342		10 mg	1687€	34 c	1.7 c	
	C. N	1	XA10205100WA	100	1.1 mL	332€	604 c	30 c	
	¹³ C. ¹⁵ N.D ₂	10	CDNLM-6786-1.2	100	1.2 mL	465€	775 c	39 c	
Bromate- ¹⁸ O₃		1	CIL-OLM-8283-18O-1.2	100	1.2 mL	<mark>406.35</mark>	<mark>6.77 c</mark>	0.34 c	
Chlorate-18O3***		12		200	5 mL	250 €	50 c	2.5 c	
		1	X 11340100DO	100	10 mL	286 €	57 c	2.9 c	
		1	XA11340100DO	100	1.1 mL	73 €	133 c	6.6 c	
Chlormequat-chloride	1.1.2.2-D ₄	6	D3386		10 mg	756 €	15 c	0.8 c	
		1	CA11340100		5 mg	389 €	16 c	0.8 c	
	D ₉	3	673151		5 mg	310 €	12 c	0.6 c	
0	¹³ C ₃	9	32679		10 mg	408	8.2 c	0.4 c	
Cyanuric acid	¹⁸ O ₃	3	673141		10 mg	299€	6.0 c	0.3 c	
Cyromazine-D ₄			XA11920010EA	100	1.1 mL	118€	215 c	11 c	
Cyromazine-D ₄	romazine-D ₄		C989302		10 mg	1047 €	21 c	1.1 c	
Daminozide-D ₆		1	XA11960100AL	100	1.1 mL	87€	158 c	7.9 c	
Diethanolamine	D ₄	4	D-5307		100 mg	432€	0.9 c	0.04 c	
Dietrianolamine	D ₈	7	D441902		100 mg	1100€	2.2 c	0.1 c	
Dibudraatrantamusia	sesquisulfate-hydrate	1	C 12635300		100 mg	29€	0.1 c	0.003 c	
Dihydrostreptomycin	sulfate	1	EPD1954000		25 mg	120 €	1.0 c	0.048	
		1	XA12960010DO	100	1.1 mL	82€	149 c	7.5 c	
		4	D-3932		10 mg	144 €	2.9 c	0.1 c	
Diquat-D₄-dibromide (e	ethylene-D ₄)	6	D17071		50 mg	840 €	3.4 c	0.2 c	
(mostly as monohydrat		7	D492902		5 mg	117€	4.7 c	0.2 c	
		10	B130022-10		10 mg	1109€	22 c	1.1 c	
		11	sc-218246		5 mg	234 €	9.4 c	0.5 c	
			XA13230100AC	100	1.1 mL	127 €	231 c	12 c	
	5	1	DRE-C13230100		10 mg	1197€		1.2 c	
Ethephon	D_4	6	D8328		5 mg	1387 €	56 c	2.8 c	
		7	C366177		10 mg	1122€	22 c	1.1 c	
	¹³ C ₂	7	C366178		2.5 mg	1650 €	132 c	6.6 c	
		1	C 13330100		50 mg	316€	1.3 c	0.06 c	
Education and the Section	TU D4)		XA13330100AC	100	1.1 mL	127 €	231 c	12 c	
Ethylenethiourea-D4 (E	= I U-D4)	6	D1965		100 mg	733 €	1.5 c	0.07 c	
		7	1367002		10 mg	98€	2.0 c	0.1 c	
	D ₁₅ (Aluminium)	1	CA13940010		10 mg	380 €	7.6 c	0.4 c	
Fosetyl	D ₅ (Sodium)	8	C5607		10 mg	825 €	17 c	0.8 c	
Glufosinate-D ₃		2	-	friendly o					



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	Source Article-No. Conc.	Audala Na	Conc.	Amount		Prices in €-cent		
Name		Source	Article-No.	[µg/mL]	per unit	1 unit	2 μg*	0.1 μg**
		7	G596952		10 mg	1870 €	37 c	1.9 c
		1	XA14050100WA	100	1.1 mL	304 €	553 c	28 c
		_	CNLM-4666-1.2	100	1.2 mL	361 €	602 c	30 c
		5	CNLM-4666-10X-1.2	1000	1.2 mL	1173€	196 c	9.8 c
01 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		1	CIL-CNLM-4666-1.2	100	1.2 mL	344 €	573 c	29 c
Glyphosate-1,2- ¹³ C ₂ , ¹⁵	N	6	CN10570		5 mg	1991 €	80 c	4.0 c
		7	G765002		10 mg	1048€	21 c	1.0 c
		9	608629-SPEC		10 mg	247 €	4.9 c	0.25 c
		11	sc-280758		1 mg	262€	52 c	2.6 c
		1	CA13230200		10 mg	256 €	5.1 c	0.3 c
HEPA (Hydroxy-Ethep	hon)-D.	7	H939652		25 mg	1125€	9.0 c	0.5 c
TILI A (Flydroxy-Ethep	110117-04	2	-		- 3	friendly o		
		3	676639	100	1 mL	99 €	2.0 c	0.1 c
		1	C 14730100		10 mg	235 €	4.7 c	0.2 c
Maleic hydrazide-D ₂ (N	MH-D ₂)	3	673799		10 mg	199 €	20 c (10μg)	1 c (0.5 µg
42 45		3	673055		10 mg	289 €	5.8 c	0.3 c
Melamine- ¹³ C ₃ , ¹⁵ N ₃		1	CIL-CNLM-8150-10X-1.2	1000	1.2 mL	1145€	229 c	12 c
	D ₁₆ -chloride-	6	D14539		50 mg	1350€	5.4 c	0.3 c
Mepiquat-	_ ,	1	X 14880100DO	100	10 mL	378 €	76 c	3.8 c
	D ₃ (methyl-D ₃) -iodide	1	XA14880100DO	100	1.1 mL	68€	124 c	6.2 c
Morpholine-D8		4	D-1895/0.5		500 mg	468 €	0.94 c (10μg)	0.05 (0.5μg)
	D ₃ (methyl-D ₃)	2	-	friendly o	friendly donation			
N-Acetyl-glufosinate	D ₃ . (Acetyl amino-D ₃ disodium salt	7	A178237		5 mg	141 €	5.6 c	0.3 c
Nereistoxin-oxalate-D ₆	- }	1	C 15502010		10 mg	245€	5 c	0.3 c
		2	-	friendly o	donation			
MPPA-D₃		7	M326162		10 mg	1825€	37 c	1.8 c
	D ₆ -diiodide	1	C 15870200		50 mg	256 €	1.0 c	0.05 c
Paraquat-	D ₈ -dichloride	7	P191902		25 mg	1125€	9.0 c	0.5 c
	- 0	5	OLM-7310-1.2	100	1.2 mL	326 €	272 c	14 c
Perchlorate- ¹⁸ O₄		12***		40	5 mL	250 €	125 c	6.3 c
Phosphonic acid- ¹⁸ O ₃		12		2000	1 mL	125	6.3 c	0.3 c
Propamocarb-D ₇		4	DER-XA16390100AC	100	1.1 mL	82€	149 c	7.5 c
PTU-D ₆ = N,N' -(1,2-Propylene = (4-Methyl-2-imidazol		6	D535 (not available)	100	100 mg	756 €	1.5 c	0.1 c
PTU-D ₆ (1,3-Propylene-d6 Thick, (not exactly co-eluting		7	P836802		10 mg	1100€	22 c	1.1 c
(not exactly co-eluting with target analyte) 1, 2, 4-Triazole- ¹³ C ₂ , ¹⁵ N ₃		2	-	friendly o	donation			
1, 2, 4-Triazole- C ₂ , TN ₃ 1, 2, 4-Triazole-acetic acid- ¹³ C ₂ , ¹⁵ N ₃		2	-	friendly o				
1, 2, 4-Triazole-alanine- ¹³ C ₂ , ¹⁵ N ₃		2	-	friendly o				
1, 2, 4-Triazole-lactic a		2	-	friendly o				
	"D ₁₅ " (in reality D ₁₂)	1	CIL-DLM-7663	,	1 mg	153 €	31 c	1.5 c
Triethanolamine								0.45
Trietnanoiamine	D ₁₂	7	T775582		10mg	141 €	2.8 c	0.15 c

			Conc.	Amount		Prices in €-	cent
Name	Source	Article-No.	[µg/mL]	per unit	1 unit	2 μg*	0.1 μg**

Providers of compounds::

- 1: LGC Standards
- 2: Bayer Crop Science
- 3: HPC (High Purity Compounds)
- 4: CDN Isotopes (via Dr. Ehrenstorfer)
- 5: Cambridge Isotope Lab. Inc.
- 6: Medical isotopes
- 7: Toronto Research Chemicals
- 8: ALSACHIM
- 9: Sigma-Aldrich
- 10. Cerilliant (by Sigma Aldrich)
- 11. Santa Cruz biotechnology. inc.
- 12. EURL-SRM (hosted at CVUA Stuttgart)

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^{* 2} µg IS are typically employed to samples (typically 10 g) at the beginning of the procedure

^{** 0.1} µg are typically added to 1 mL aliquots of sample extracts (typically corresponding to 0.5 g sample), in this case only matrix-effects are compensated

pensated
*** Due to manufacturing process the stock solution of ¹⁸O₃-Chlorate is accompanied by ¹⁸O₄-Perchlorate (ca. 40 µg/mL). As perchlorate has typically a 5-fold higher sensitivity compared to chlorate the signal intensities of the two are typically within the same range.



Table 22: Exemplary concentrations of IS working solutions (3.19)

	IS -Addition to s	samples (5.2.3)	· ·	calibration stand-	Expected approx. IS-		
IS*	Suggested concentration of IS-WS I (3.19)	Absolute mass of IS spiked to sample (50 µL IS-WS I) (mis sample)	Suggested concentration of IS- WS II (3.20) **	Absolute mass of IS spiked to calibration standard (50 µL IS-WS II) (mis cal mix)	concentration in sample		
	μg/mL	μg	μg/mL	μg	μg/mL		
Amitrole-(¹⁵ N)/ (¹⁵ N ₂ , ¹³ C ₂)	40	2	2	0,1	0,1		
AMPA- ¹³ C, ¹⁵ N	40	2	2	0,1	0,1		
Bromate- ¹⁸ O ₃	<mark>60</mark>	3	3	0.15	0.15		
Chlorate- ¹⁸ O ₃	40	2	2	0,1	0,1		
Chloridazon-desphenyl-	40	2	2	0.1	0.1		
Chlormequat-D ₄	40	2	2	0,1	0,1		
Cyromazine-D ₄	40	2	2	0,1	0,1		
Daminozid-D ₆	40	2	2	0,1	0,1		
Diethanolamine-D ₆	40	2	2	0,1	0,1		
Dihydrostreptomycin****	40	2	2	0,1	0,1		
Diquat-D ₄	40	2	2	0,1	0,1		
Ethephon-D ₄	40	2	2	0,1	0,1		
ETU-D₄	40	2	2	0,1	0,1		
Fosetyl-D ₅ (from fosetyl-aluminium-D ₁₅)	40	2	2	0,1	0,1		
Glufosinat-D ₃	40	2	2	0,1	0,1		
Glyphosat-13C2,15N	40	2	2	0,1	0,1		
HEPA-D₄	40	2	2	0,1	0,1		
Maleic hydrazide-D ₂	200	10	10	0,5	0,5		
Mepiquat-D₃	40	2	2	0,1	0,1		
Morpholine-D ₈	40	2	2	0,1	0,1		
MPPA-D ₃	40	2	2	0,1	0,1		
N-Acetyl-AMPA*	40	2	2	0,1	0,1		
N-Acetyl-glufosinate-D ₃	40	2	2	0,1	0,1		
Nereistoxin-D ₄	40	2	2	0,1	0,1		
Paraquat-D ₆	40	2	2	0,1	0,1		
Perchlorate-18O ₄	40	2	2	0,1	0,1		
Phosphonic acid	40	2	2	0,1	0,1		
PTU-D ₆	40	2	2	0,1	0,1		
Triethanolamine-D ₁₂	40	2	2	0,1	0,1		

^{*} The concentration of the IS should be high enough to ensure good detection with little influence of the noise on the signal (e.g. S/N>20 is fine). It should be kept in mind, however, that isotopically labeled ISs (IL-ISs) typically contain small amounts of the non-labeled analogues. To minimize the risk of false positives the amount of IL-IS added to the samples should thus not be higher than necessary. Quantification of the parent is typically not affected to a great extend as the cross-contamination is typically at low levels and as similar concentrations of the native pesticide originating from the IL-IS will also be present in the calibration standards and thus subtracted via the intercept. In the case of Maleic hydrazide, where the IL-IS is added at higher concentrations to the samples special attention is necessary (see also comments under 5.7.2).

NOTE: If detections of a compound are rather seldom and the IS expensive it is advisable to add the IS to the 1 mL aliquot transferred to the autosampler vial (see **5.2.7**). Alternatively, it can be even skipped entirely in the first screening analysis and only added in a second analysis in case the

^{**} a 20-fold dilution of the IS working solution used to spike samples in step 5.2.3 .

^{***} Dihydrostreptomycin is not isotopically labeled but still suitable for compensation of matrix effects on Streptomycin if LC conditions are adjusted in such a way to ensure exact co-elution and thus equivalent matrix-effects

first one was positive. The first approach is to be preferred especially where the retention times of a compound tends to shift. By comparing the retention time between the IS and the suspected peak as well as the peak shape the certainty of identification significantly improves.

Table 23: Water content of selected foods and Water amount to be added to test portions prior to extraction (5.2.2) depending on the analytical approach

	е апагушсаг аррго				
		Typical	mL of Water to be add tions [g] (where Water- ent sample weights this	addition refers to differ-	
Commodity group	Commodity	Water content g/100 g	When quantifying with IS that was added at the beginning of the procedure (5.2.3)	When no IS is used or when IS is added after aliquotation (5.8.1.2)	Remarks
Fruits					
Citrus fruit	Citrus juices	90	-	1	
	Grapefruit	90	-	1	
	Lemon/lime	85	-	1.5	
	Orange	85	-	1.5	
	Tangerine	90	-	1	
Pome fruit	Apple	85	-	1.5	
	Apple (dried)	30	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
	Apple sauce	80	-	2	
	Apple juice	90	-	1	
	Pear	85	-	1.5	
	Quince	85	-	1.5	
Stone fruit	Apricot	85	-	1.5	
	Apricot (dried)	30	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
	Apricot nectar	85	-	1.5	
	Cherry	85	-	1.5	
	Mirabelle	80	-	2	
	Nectarine	85	-	1.5	
	Peach	90	-	1	
	Peach (dried)	20	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
	Plum	85	-	1.5	
	Plum (dried)	20	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
Soft and small	Blackberry	85	-	1.5	
fruit	Blueberry	85	-	1.5	



			mL of Water to be add tions [g] (where Water-		
		Typical	ent sample weights this		
Commodity group	Commodity	Water con- tent g/100 g	When quantifying with IS that was added at the beginning of the procedure (5.2.3)	When no IS is used or when IS is added after aliquotation (5.8.1.2)	Remarks
	Currant	85	-	1.5	
	Elderberry	80	-	2	
	Gooseberry	90	-	1	
	Grapes	80	-	2	
	Raspberry	85	-	1.5	
	Raisins	20	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
	Strawberry	90	-	1	
	Pineapple	85	-	1.5	
Other fruits	Banana	75	2.5	2.5	
	Fig	80	-	2	
	Fig (dired)	20	8.5 to 5 g sample (see 5.2.2)	8.5 to 5 g sample (see 5.2.2)	Weigh 13.5 g rehydratized homogenate
	Kiwi	85	-	1.5	
	Mango	80	-	2	
	Papaya	90	-	1	
Vegetables					
Root and tu- ber vegetables	Beetroot	90	-	1	
bei vegetables	Carrot	90	-	1	
	Celeriac	90	-	1	
	Horseradish	75	2.5	2.5	
	Parsley root	90	-	1	
	Radish	95	-	0.5	
	Black salsify	80	-	2	
	Potato	80	-	2	
	Garlic	60	7 to 5 g sample	7 to 5 g sample	
Leek plants	Onion	90	-	1	
	Leek	85	-	1.5	
	Shallot	80	-	2	

			ml of Water to be add	ad to 10 a test war	
		Typical	mL of Water to be add tions [g] (where Water- ent sample weights this	addition refers to differ-	
Commodity group	Commodity	Water content g/100 g	When quantifying with IS that was added at the beginning of the procedure (5.2.3)	When no IS is used or when IS is added after aliquotation (5.8.1.2)	Remarks
	Chive	85		1.5	
Fruiting vege-	Aubergine	90	-	1	
tables	Cucumber	95	-	0.5	
	Melon	90	-	1	
	Pepper, sweet	90	-	1	
	Pumpkin	95	-	0.5	
	Tomato	95	-	0.5	
	Zucchini	95	-	0.5	
	Broccoli	90	-	1	
Cabbage	Brussel sprouts	85	-	1.5	
	Cauliflower	90	-	1	
	Chinese cabbage	95	-	0.5	
	Kale	90	-	1	
	Kohlrabi	90	-	1	
	Red cabbage	90	-	1	
	Savoy cabbage	90	-	1	
	White cabbage	90	-	1	
	Lettuce varieties	95	-	0.5	
	Endive	95	-	0.5	
Leafy vegeta- bles and herbs	Cress	90	-	1	
bies and nerbs	Lamb's lettuce	85	-	1.5	
	Parsley	80	-	2	
	Rucola	85	-	1.5	
	Spinach	90	-	1	
Stem	Asparagus	95	-	0.5	
vegetables	Celery	95	-	0.5	
	Leek	85	-	1.5	
	Rhubarb	95	-	0.5	

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		Typical	mL of Water to be add tions [g] (where Water- ent sample weights this		
Commodity group	Commodity	Water con- tent g/100 g	When quantifying with IS that was added at the beginning of the procedure (5.2.3)	When no IS is used or when IS is added after aliquotation (5.8.1.2)	Remarks
	Artichokes	85	-	1.5	
Legumes	Beans, peas, lentils (dried)	<10	10 to 5 g sample	10 to 5 g sample	
	Beans, peas	75	2.5	2.5	
Miscellaneo	us				
Cereals	Grain, flour etc.	10	10 to 5 g sample	10 to 5 g sample	Different sample amounts may be used depending on Water-absorbing properties of material
Extract-rich	Coffee beans	<10	10 to 2 g sample	10 to 2 g sample	Different sample
("difficult") commodities	Tea	<10	10 to 2 g sample	10 to 2 g sample	amounts may be
	Dry herbs and spices	<10	10 to 2 g sample	10 to 2 g sample	used depending on extract-richness
Other	Mushrooms	90	-	1	
	Wine	90	-	1	
	Honey	20	9 to 5 g sample	9 to 5 g sample	

Table 24: Exemplary LC-MS/MS parameters for Sciex QTrap 5500

	Methods	Method	Method	Method	Method	Method	Method	Method	Method
	1.1 / 1.2	1.3	1.4	2	3 + 4.1 + 5	4.2	6	7	8
Ion source (ESI, Turbo Ion Spray) Mode	negative	negative	negative	negative	positive	positive	positive	positive	positive with Sele- xlon TM
Curtain gas (Nitrogen)	30 psi (2.07 bar)	40 psi (2.76 bar)	40 psi (2.76 bar)	30 psi (2.07 bar)	30 psi (2.07 bar)	30 psi (2.07 bar)	30 psi (2.07 bar)	40 psi (2.76 bar)	20 psi (1.38 bar)
Collision gas					medium				
lon spray voltage	-4500	-4500	-4500	-4500	1500	5000	5500	1500	5500
Gas 1 (Zero Grade Air or Nitro- gen)	50 psi (3,45 bar)	60 psi (4,14 bar)	60 psi (4,14 bar)	50 psi (3,45 bar)	50 psi (3,45 bar)	60 psi (4,14 bar)	50 psi (3,45 bar)	60 psi (4,14 bar)	60 psi (4,14 bar)
Gas 2 (Zero Grade Air or Nitro- gen)	60 psi (4,14 bar)	60 psi (4,14 bar)	70 psi (4,83 bar)	60 psi (4,14 bar)	60 psi (4,14 bar)	50 psi (3,45 bar)	60 psi (4,14 bar)	70 psi (4,83 bar)	70 psi (4,83 bar)
Temperatur Gas 2	600°C	550°C	550°C	500°C	500°C	500°C	550°C	500°C	550°C
Resolution MS 1		unit (ca. 0.7 amu FWHM*)							
Resolution MS 2		unit (ca. 0.7 amu FWHM)							
Dwell time	20	20	20	50	20	10	50	20	20

^{*}FWHM = full width at half maximum



Table 25: Document History

Action	When?	Version
Development of Method by the CRL-SRM	2006-2008	
Presentation of method at the EPRW in Berlin (oral presentation plus poster)	June 2008	-
Drafting of V1	NovDec. 2008	V/4
Placing of V1 in CRL-Website	Jan. 2009	V1
Update of Table 1, Expected concentrations of ISs were calculated with a wrong dilution factor in previous version. Arithmetical errors were corrected. Introduction of measurement conditions for HEPA within the "Glyphosate & Co." method	Aug. 2009	V2
Introduction of measurement conditions for the screening of diquat and paraquat within the "Quats & Co. method"		
Introduction of measurement conditions for Amitrole, chlormequat, mepiquat and daminozide "Amitrole & Co." method	Nov 2009	V3
Extensive text revisions		
Introduction of measurement conditions for Streptomycin Kasugamycin		
Introduction of measurement conditions for the screening of Perchlorate ion	May 2010	V4
Extensive text revisions		
Extensive text revisions and restructuring of document		
Introduction of measurement conditions for ETU, ETU D_4 , PTU, PTU D_6 , Cyromazine, Cyromazine D_4 , N-Acetyl-Glufosinate, N-Acetyl-Glufosinate D_3 , MPPA D_3 , Morpholin, Morpholin D_8	Nov 2010	V5
Introduction of an acronym for the method (QuPPe)		
Advice to use plastic vessels and stoppers for Glyphosate		
Minor modification and additional instructions in Method 1 (M1)		
Modification of mobile phase of M3 to improve analysis of ETU and PTU		
Introd. of measurement cond. for Amitrole 15N13C and Amitrole 15N in M3		
Introd. of measurement cond. for Nereistoxin and Nereistoxin D6 in M4	July 2011	V6
New method (M7) for the analysis of Morpholin/Morpholin D_8 ; Diethanonamine/diethanolanmine D_6 ; Triethanolamine/Triethanolamine D_{12} (M7) Removal of Morpholin from M4 as it does not separate from the interfering diethanolamine		
Introduction of ETU and PTU and their corresponding ILISs in Method 5		
Correction of dimension of stock solutions conc. in Table 12 (to mg/mL)		
Text and Table revisions		

Action	When?	Version
Extensive revision of table concerning possible sources of purchase of ISs		
Some additions in "Apparatus and Consumables" chapter		V7
Clarifications in chapter concerning standard additions		
Overview table concerning the scope of the methods 1.1, 1.2, 1.3 and 2		
Addition of Phosphonic acid in Method 1.1 ("Glyphosate & Co.")		
New LC-method (Method 1.2) for "Glyphosate & Co." using a Dionex ionPac AS11-HC column and an Eluent with near to neutral pH; additionallycovering Fosetyl		
New LC-method (Method 1.3) for "Glyphosate & Co." using a Hypercarb column and an acidic Eluent covering all analytes covered by Method 1.1, Method 1.2 and Method 2 (including perchlorate).		
Update of practical considerations for methods 1.1-1.3		
Update of table with performance data		
Table with exemplary recovery data was deleted (recovery figures can be obtained in the EURL-DataPool		
Update of table with LOQs		
Update of table with providers of ILISs		
Elimination of errors in text		
Addition of Chlorate in Method 1.3	Nov. 2013	V7.1
Update of practical considerations for methods 1.1-1.3 (Column C)		
Update of table with performance data		
Update of table with LOQs		
Introduction of Trimethylsulfonium-D9 and N,N-Dimethylhydrazine-D6 in Method 4		
Thorough revision of text and elimination of errors		V8
Practical advices on the choice of filter materials		
New Table 15: Conversion factors between standard materials and analytes		
Advices as regards the use of ILISs		
Update of Table 5.6 LC-MS/MS measurement conditions		
New chapters "Hints on Method $1.1-1.4$ " and replacement of the section "Practical care and use considerations concerning the columns of methods $1.1-1.3$. This includes information on various potential sources of errors such as in-source fragmentations of Fosetyl and Ethephon to Phosphonic acid and of Perchlorate to Chlorate as well as degradation of compounds in solution.		
Introduction of Cyanuric acid and Bialaphos in M1.3		

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Action	When?	Version
Correction of a typing error concerning the mass-transitions of Phosphonic acid (81/79 instead of 81/81)		
Introduction of the ILIs of Phosphonic acid and chlorate in M1.3 and 1.4		
New LC Method (1.4) for "PerChloPhos" using a Hypercarb column and an acidic Eluent optimized for chlorate, perchlorate, phosphonic acid compared to Method 1.3		
Change of name of former M4 to M4.1		
Introduction of Melamine and Propamocarb as well as the corresponding ILISs in M4.1		
New LC Method (M4.2) employing a Hilic-Type BEH Amide column allowing the simultaneous analysis of many polar pesticides		
Reduction of injection volume and increase of dwell-time in method M6		
New LC-method (M8) for the analysis of triazole derivative metabolite (TDMs) and their corresponsing ILISs		
Update of Table 17: Providers of isotopically labeled internal standards		
5.1 Sample preparation: note to importance of having small particle sizes		V8.1
5.2.4 notes to extraction time for dry products and the influence of particle size	May 2015	
5.6 information on the methods currently routinely used at CVUA Stuttgart		
Update Table 20 Exemplary LC-MS/MS parameters for Sciex QTRAP 5500		
Update of chapter 5 Procedure including the extraction procedure at a glance	Mar. 2016	V9
Update of Table 3 Overview and scope of the methods proposed within this document for the QuPPe method:		
Update of Table 4 Practical Information: Mainly used methods used at CVUA Stuttgart		
Update of chapter 5.7.3.1.Hints on Method 1.3		
Update of method 1.4 Introduction of measurement conditions for the measurement of Bromate and Bromide ion		
Update of chapter 5.7.4.1.Hints on Method 1.4		
Update of Method 4.2 "Quats & Co BEH Amide" including Aminocyclopyrachlor, Chloridazon-desphenyl, Mepiquat-4-hydroxy, Propamocarb-N-desmethyl, Propamocarb-N-oxide		
Update of Method 6 "Streptomycin and Kasugamycin", change of gradient and new chromatograms		
Update of		
Method 8 "Triazole derivative metabolites (TDMs)" new DMS parameters		
Update of Table 18: Overview of approximate limits of quantification (LOQs)*		

Action	When?	Version
Update of Table 19: Conversion factors between typical purchased standards and target analytes (3.15):		
Update of Table 20: Exemplary concentrations of pesticide stock and working solutions		
Update of Table 21: Providers of isotopically labeled internal standards		
Update of Table 22: Exemplary concentrations of IS working solutions		