Research Article

Ion Suppression Study for Tetracyclines in Feed

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Ion suppression in analysis of tetracyclines in feed was studied. The conventional analysis consists of a liquid extraction followed by a clean-up step using solid phase extraction (SPE) technique and analysis of the tetracyclines by liquid chromatography and mass spectrometric detection. Various strategies for extraction and cleanup were tested in the present work, and the effectiveness to decrease the ion suppression on the MS/MS signals was evaluated. Four sample treatment methods were tested with five different feed samples. Extraction solvents tested were McIlvaine buffer and a mixture of McIlvaine buffer dichloromethane (3 : 1). SPE cartridges for cleanup were Oasis HLB, Oasis MCX, and Oasis MAX. The effectiveness of the methods was evaluated in terms of decreasing the ion suppression effect but also of decreasing the variability of ion suppression between samples. The method that provided the most satisfactory results involved a clean-up step based on SPE using mixed-mode cation exchange cartridges (Oasis MCX).

1. Introduction

Tetracyclines are a family of drugs belonging to the group of antibiotics. They are widely used in animal husbandry for therapeutic and prophylactic purposes. Oxytetracycline, tetracycline, chlortetracycline, and doxycycline are by far the most used antibiotics from this family. Their main chemical properties are their amphoteric behaviour due to their several acid-base equilibria and the tendency to act as chelating agents in presence of multivalent ions [1, 2]. Figure 1 shows structures and pKa values of the tetracyclines studied. At pH values below 3 they are positively charged. At pH between pKa1 and pKa2 they are neutral (zwitterionic state), and above pH 8 they are negatively charged.

The use of antibiotics in animal husbandry is strictly regulated to protect consumers, as the presence of antimicrobials residues in food products of animal origin can lead to resistance of bacteria to antibiotics. Therefore, the European Union has developed regulation concerning this issue [3, 4]. Analysis and control of antibiotics in feedstuffs for animals has become an important issue as only authorized feedstuffs can be medicated under specific conditions as stated in Council Directive 90/167/EEC [5]. Use of tetracyclines as feed additives is forbidden in the EU since 2006, as stated in annex II of Commission Recommendation 2005/925/EC [6].

Feed contamination can occur depending on a large number of factors such as human error or handling procedures, but production practices have been identified as the main source [7].

Nowadays, liquid chromatography-tandem mass spectrometry (LC-MS/MS) is the technique of choice for the analysis of veterinary residues in food. The analysis of antibiotics in animal feed, though, has proved to be quite a challenge because of the high complexity and variability of the composition of the matrix. Numerous raw materials and additives are added into the feeds, including grains, seeds, beans, rice, and soy, and thus many interfering components, such as oils, fats, proteins, and salts can occur at very high levels. This complexity causes a strong effect of ion suppression. Ion suppression can be defined as a change in the efficiency of droplet formation or evaporation in the ion source of mass spectrometer, caused mainly by interfering matrix compounds. This affects the amount of charged analyte that reaches the detector and so the signal obtained for it. During the last years a growing concern on this issue has been reported [8–17]. Some factors such
as mobile phase composition [9, 15] or the type of ion source and its geometry [17] have been reported to play a role in ion suppression, but matrix components reaching ion source are the most commonly reported of them. More knowledge on the removal of matrix interferences is needed to overcome ion suppression problems. Many authors have studied this phenomenon by improving sample treatment in residue analysis by HPLC-MS/MS in biological matrices such as whole blood, plasma, serum, or urine [12–14, 16, 18]. In the case of feed samples, it has been proved recently that they are an extreme case, regarding ion suppression, compared to other kind of matrices [19]. Moreover, the changeable composition of each individual feed leads to the obtention of sample extracts with high variation in matrix components, and that leads then to very different extents of ion suppression for each single feed sample. Therefore, not only is it much harder than with other kind of samples (like food) to avoid ion suppression effect, but it is also difficult to obtain at least a homogeneous sample-independent effect. This factor does not allow accurate quantification even when matrix-matched calibration approach is performed. A solution to overcome this effect has been found in the emergence of more isotopically labelled internal standards. The labelled internal standard coelutes with the analyte in question and has similar physicochemical properties. These internal standards, though, still do not ensure correct quantification in all cases [20]. Moreover, their commercial availability is still scarce, and they represent a high cost option. Dilution of the final extract to reduce matrix concentration is also a common option. However, when analyzing samples that may have been contaminated by error or by cross-contamination during production, levels can be very low (in the range of the few parts per billion) and so no great dilution factors are recommended. Presently, the only way to make completely sure that HPLC-MS/MS quantification is fully reliable is to apply standard addition calibration. This ensures a correct quantification of each individual feed. Unfortunately this quantification tool is very time consuming and cumbersome, resulting in only a few feeds being analysed per day, which is hardly affordable for laboratories which have to handle a high number of samples.

Development of analytical methodologies for analysis of veterinary drugs in feed by HPLC-MS/MS has started to increase in number for the last recent years. Although almost all authors are aware about ion suppression/enhancement phenomena in feed analysis, only a few of them have developed their methods including standard addition calibration [21, 22]. Some others decided to perform this calibration technique by building calibration curve with spiked aliquots of the processed sample prior to HPLC-MS/MS analysis [23, 24], assuming that extraction recovery is a factor of much less impact in the final results than matrix effects. Others assume that their extraction and clean-up techniques are good enough to compensate this effect [25–27] or do not even mention it [28–30].

The aim of this work was to investigate some clean-up methodologies and to evaluate their effectiveness to reduce ion suppression in the analysis of antibiotic residues in feed by LC-MS. For this evaluation, some strategies mentioned in the literature [9, 11–13, 18] were used. Tetracyclines were chosen as a model group to perform the experiments. No papers have been found reporting ion suppression concern in LC-MS/MS tetracycline analysis, and only one in LC-MS analysis (single quadrupole and ion trap) in soils [31]. Several sample treatment procedures were tested and compared using different kinds of feed samples.
2. Experimental

2.1. Chemicals and Reagents. Oxytetracycline (OTC), tetracycline (TC), chlortetracycline (CTC), doxycycline (DC) and demeclocycline (DMC) were purchased from Sigma (St. Louis, MO, USA). Methanol (MeOH) and acetonitrile (ACN) were obtained from Biosolve (Valkenswaard, The Netherlands) and acetic acid (AA), ammonia solution 25% v/v (NH₃), formic acid (FA), and dichloromethane (DCM) from Merck (Darmstadt, Germany). Solid reagents were purchased from Merck and included citric acid, potassium dihydrogen phosphate, ethylenediaminetetraacetic acid disodium salt (EDTA), and sodium hydroxide (NaOH). All reagents were analytical-reagent grade.

Figure 3: Ion suppression profiles of a blank, a dry, and a wet feed sample obtained with sample treatment method 1.
Table 1: Sample pretreatment methods tested for the cleanup of tetracyclines from animal feeds.

| Method | 1 | 2 | 3 | 4 |
|--------|---|---|---|---|
| Extraction | 40 mL McIlvaine buffer-EDTA 0.1 M | 40 mL DCM McIlvaine buffer-EDTA 0.1 M (1:3) | 40 mL McIlvaine buffer-EDTA 0.1 M | 40 mL McIlvaine buffer-EDTA 0.1 M |
| SPE Loading pH | 4.2 | 4.2 | 2.5 | 10 |
| Cartridge | Oasis HLB (60 mg) | Oasis HLB (60 mg) | Oasis MCX (60 mg) | Oasis MAX (60 mg) |
| Wash (2 mL) | H₂O | H₂O | (1) FA 2% (v/v) | (1) NH₃ |
| Elution (2 mL) | MeOH | MeOH | MeOH : NH₃ (95:5) | MeOH : FA (95:5) |
| Final step | Evaporation with N₂ and reconstitution with mobile phase | Evaporation with N₂ and reconstitution with mobile phase | Dilution with 8 mL acetic acid 10% (v/v) | Dilution with 8 mL H₂O (v/v) |

Table 2: LC-MS/MS precursor/product ion combinations (quantifier bold) monitored in MRM ESI positive mode.

| Tetracycline | Retention Time (min) | Precursor ion (m/z) | Product ions (m/z) |
|--------------|----------------------|--------------------|-------------------|
| TC           | 9.2                  | 445.2              | 410.1, 154.1      |
| CTC          | 10.2                 | 479.1              | 444.1, 154.1      |
| DC           | 10.4                 | 445.2              | 428.1, 154.1      |
| OTC          | 8.9                  | 461.2              | 337.1, 201.1      |
| DMC          | 9.7                  | 465.1              | 154.1             |

The pH of the extract was adjusted when necessary, and filtration through glass fiber filters was performed before SPE step. Final extracts obtained were filtered through 0.45μm nylon syringe filters before injection into the LC-MS/MS system.

2.3. LC-MS/MS Conditions. A Waters 2690 separations module HPLC system (Waters Corporation, USA) coupled to a Quattro Ultima tandem mass detector (Micromass/Waters, Manchester, UK), both operating under MassLynx software, was used for sample analysis. The mass spectrometer was operated in electrospray positive mode, and data acquisition was in multiple reaction monitoring mode (MRM). The precursor/product ions monitored are listed in Table 2. The source settings were as follows: capillary voltage 2.7 kV, source temperature 120°C, desolvation temperature 300°C, cone nitrogen gas flow 180 Lh⁻¹, and desolvation gas flow 580 Lh⁻¹. Argon (3.2 × 10⁻³ mbar) was used as the collision gas, and the multiplier was operated at 750 V. The cone voltage was set at 20 V, and collision energy changed during analysis depending on the analyte (25 eV for TC and DC, 26 eV for OTC, and 30 eV for CTC and DMC). The HPLC system was equipped with a Symmetry C₁₈ (5 μm, 3.0 × 150 mm column, Waters) at 10°C. A binary gradient mobile phase was used at a flow rate of 0.4 mL min⁻¹ with solvent A (ammonium acetate 1 mM, pH 2.6) and solvent B (ammonium acetate 10 mM: ACN, 10:90). The gradient started isocratic for 1 min at 0% B, followed by a linear gradient.

Figure 4: Chromatogram from a standard injection of TC, CTC, OTC, DC, and DMC (1 mg L⁻¹).

Standard solutions (1000 mg L⁻¹) were prepared in methanol monthly and stored at 4°C. Mixtures of OTC, TC, CTC, and DC (10 and 100 mg L⁻¹) were prepared by dilution of the concentrated solutions and stored at 4°C for a week. An internal standard (IS) stock solution (DMC, 100 mg L⁻¹) in MeOH was prepared monthly. Working standard solutions were prepared daily by mobile phase dilution of the 10 and 100 mg L⁻¹ mixtures.

SPE materials were obtained from Waters (Micromass/Waters, Manchester, UK).

2.2. Sample Treatment Procedures. Five porcine feed samples were used for the study: one premix sample, two dry feed samples, and two slurry feed samples. Feeds were chosen that were representative for the range of different feeds available. Four different sample treatment procedures, which are summarized in Table 1, were tested. Sample weight was 2 grams, and extraction solution volume was 40 mL. Extraction was carried out by means of a head-over-head shaker for 20 minutes in all cases, and samples were subsequently centrifuged at 3000 rpm for 15 minutes.
increase to 50% B in 9 min. The gradient remained isocratic at 50% B for 3 min. Subsequently the gradient linearly increased to 100% B in 1 min. The gradient remained at this % B for a further 3 min. Afterwards the gradient returned to 0% B for equilibration of the column. Sample injection volume was 10 µL.

2.4. Qualitative Assessment of Ion Suppression. The experiments for qualitative assessment of ion suppression were carried out using a postcolumn infusion setup coupled to the chromatographic system described in Section 2.3 through a T piece. The setup is shown in Figure 2. The infusion pump flushes a constant flow at 10 µL min\(^{-1}\) of a 5 mg L\(^{-1}\) standard solution of all tetracyclines in mobile phase. The quantification transition for each tetracycline is monitored in the MS/MS system. When mobile phase is injected into the system, a reference baseline for each transition is obtained due to the constant infusion of the standard solution of analytes. When feed extracts free from tetracyclines are injected, ion suppression profiles for each transition are obtained, and the influence of ion suppression in the tetracyclines infusion baseline due to the eluted matrix components can be evaluated. In fact, these profiles show the effect of compounds eluting from the chromatographic system on the analytes MS/MS signals. The signal intensity of the baseline decreases when matrix components causing ion suppression elute, and it increases when substances enhancing ionization elute.

Observing the signal variation at the time window where every analyte elutes, a good qualitative prediction can be made, whether suppression or enhancement are expected for that analyte.

2.5. Quantitative Assessment of Ion Suppression. 1 mg L\(^{-1}\) tetracyclines standard solutions and matrix-matched recovery standards (MMRSs) at the same concentration were injected into the LC-MS/MS system. MMRs are extracts from blank feed samples that have been spiked with the analytes at the end of the sample treatment process. Ion suppression or enhancement percentages were determined for each tetracycline as the peak area ratio of the MMRs to the standard in solution multiplied by 100. Values lower than 100% were an indicative of ion suppression whereas values higher than 100% indicated ion enhancement.

For each analyte, the response factors (\(\text{Area}_{\text{analyte}}/\text{Area}_{\text{IS}}\)) in the 5 studied feed samples were determined in MMRs for the set of samples. Each MMRs was analysed by triplicate, and the average was calculated. The RSD (%) between the averages of the five tested samples (\(n = 5\)) was used to quantify the variation in ion suppression due to differences between feed samples for each sample treatment method.

3. Results and Discussion

Four different sample treatments, summarized in Table 1, were tested in this study. All of them are based in an extraction step using McIlvaine buffer-EDTA 0.1 M (pH 4.2) and a further clean-up step of solid phase extraction. McIlvaine buffer has been extensively reported to be efficient for tetracycline extraction in a large number of matrices, as stated in some reviews [1, 2].

Method 1 is currently in use at this laboratory for routine analysis. In this method, the cleanup of the extracts is performed with the reversed phase Oasis HLB cartridges. The loading of the extract into the cartridge does not require any pH adjustment since the maximum interaction of TCs with the sorbent occurs when the neutral form of the analytes is prevalent, like at pH 4.2 (Figure 1). Finally the elution of TCs is achieved with methanol.

The effect of addition of dichloromethane (DCM) in the extraction step was investigated in method 2, as DCM might assist to the removal of some nonpolar matrix compounds
and thus provide cleaner extracts. After extraction, the aqueous layer was processed throughout an Oasis HLB cartridge as in method 1.

The pH of the McIlvaine buffer extract was modified after extraction in methods 3 and 4 to reach a suitable pH for SPE. Mixed-mode cation exchange (Oasis MCX, method 3) or mixed-mode anion exchange (Oasis MAX, method 4) cartridges were used. These cartridges base their performance in a combination of ion exchange and reverse phase mechanisms. Therefore, they are expected to be more selective for targeted analytes and so to provide a more efficient cleanup [18, 32–34].

**Figure 6:** Ion suppression profiles of a blank, a dry, and a wet feed sample obtained with sample treatment method 3.
For method 3, before loading into the cartridge, pH of the extract was decreased to 2.5 in order to have the analytes positively charged. Theoretically SPE cartridge performance may be compromised, as the pH is only slightly lower than $pK_{a1}$ (Table 1). However decreasing pH more is not recommended, as it has been reported to induce epimerization of tetracyclines [1, 2]. At these conditions good SPE recoveries for all analytes were obtained for the complete optimized method (74–100%).

For method 4, pH of the extract was brought to 10 to ensure all analytes were in anionic form. At these conditions, SPE recoveries were good for the analytes (97–100%) except for CTC (ca. 30%). This is possibly due to partial degradation of this analyte to iso-CTC, as this analyte is particularly prone to form this derivative at high pH values [2].

Ion suppression profiles of feed extracts obtained for the four sample pretreatment methods were recorded. These profiles were studied separately for each analyte in every feed extract and compared. Focusing in the behaviour of the profile at the retention time at which tetracycline elutes, and comparing it with the reference “blank” signal (which corresponds to the injection of mobile phase), a good qualitative assessment can be made, whether suppression or enhancement are expected in a significant extent for each tetracycline in each studied feed extract.

Ion suppression profiles of a blank, a dry and a slurry (wet) feed sample extract obtained with sample treatment method 1 are shown in Figure 3. Both feeds exhibit ion suppression in the chromatogram time window where tetracyclines elute (8–12 min, Figure 4).

A similar trend was observed when using method 2. The addition of DCM to the extraction solution was therefore not an improvement. Numeric results from the ion suppression quantification experiment agreed with these results, and the suppression factors (see Section 2.5) for all the analytes in the set of feed samples were clearly below 100% for both methods (Figure 5).

Some improvements were clearly observed in the ion suppression profiles of the extracts obtained according to methods 3 and 4. No ion suppression or enhancement was observed in the profiles between 8 and 12 min As a matter of example Figure 6 shows the ion suppression profiles of a blank, a dry and a slurry (wet) feed sample extract obtained with method 3. Similar profiles were obtained when using method 4. The results of the ion suppression quantification experiment are consistent with these qualitative results. The percentage of signal obtained in 1 mg L$^{-1}$ MMRs in the five feeds studied compared to a standard solution was close to 100% (Figure 5). Only in the case of CTC in one of the two slurry feeds, significant enhancement (>150%) was found, what seems to show that this sample contains some particular substances that enhance the ionization of this particular analyte under the mentioned sample treatment conditions. The overall clear improvement provided by methods 3 and 4 has been achieved by the combination of the use of ion exchange cartridges (more selective) and the dilution of the extracts instead of evaporation and reconstitution (less introduction of contaminants and human error).

### Table 3: Variation (RSD %) between the five different studied feed samples processed with the four different sample pretreatment methods at 1 mg L$^{-1}$.

| Method | 1 RSD (%) | 2 RSD (%) | 3 RSD (%) | 4 RSD (%) |
|--------|-----------|-----------|-----------|-----------|
| TC     | 12.3      | 19.4      | 11.1      | 2.6       |
| CTC    | 17.9      | 25.8      | 12.3      | 21.1      |
| DC     | 30.8      | 39.7      | 4.9       | 6.5       |
| OTC    | 14.7      | 24.3      | 9.4       | 8.6       |

As expected, a high variation between the different feed samples was found for all tetracyclines (Table 3). Methods 3 and 4 provide the best results regarding the variability on the response factors (Area$_{analyte}$/Area$_{IS}$) in MMS due to the feed sample. CTC is an exception (method 4) due to degradation at pH 10. That indicates that the extracts obtained for the five feeds with these two methods are more uniform than the ones obtained by methods 1 and 2.

### 4. Conclusions

Ion suppression in LC-MS/MS analysis of tetracyclines in feed was studied. The study of this phenomenon in feed samples has proven to be of a high level of complexity. Four sample pretreatment methods were tested with five different feed samples in terms of ion suppression profiles, ion suppression quantification, and variation. The method that seemed to provide less ion suppression and more uniform extracts without significant degradation of any analyte was method 3, which involved SPE with Oasis MCX cartridges. Although these results are still not sufficient to replace the current protocol (method 1) which relies heavily on standard addition, they provided valuable information for future research, which should include studies with larger numbers of different feed samples, different concentration levels, and different levels of extract dilution.

### References

[1] H. Oka, Y. Ito, and H. Matsumoto, “Chromatographic analysis of tetracycline antibiotics in foods,” Journal of Chromatography A, vol. 882, no. 1-2, pp. 109–133, 2000.

[2] C. R. Anderson, H. S. Rupp, and W. H. Wu, “Complexities in tetracycline analysis—chemistry, matrix extraction, cleanup, and liquid chromatography,” Journal of Chromatography A, vol. 1075, no. 1-2, pp. 23–32, 2005.

[3] Commission Decision. Regulation (EC) No 882/2004 of the European Parliament and of the Council of 29 April 2004 on official controls performed to ensure the verification of compliance with feed and food law, animal health and animal welfare rules, 2004.

[4] Regulation. Regulation (EC) No 1831/2003 of the European Parliament and of the Council of 22 September 2003 on additives for use in animal nutrition, 2003.

[5] Council Directive 90/167/EEC. Council Directive 90/167/EEC of 26 March 1990 laying down the conditions governing the...
preparation, placing on the market and use of medicated
feedingstuffs in the Community, 1990.

[6] Commission Recommendation 2005/925/EC of 14 December
2005 on the coordinated inspection programme in the field of
animal nutrition for the year 2006 in accordance with Council
Directive 95/53/EC.

[7] J. D. G. McEvoy, “Contamination of animal feedingstuffs as
a cause of residues in food: a review of regulatory aspects,
iccidence and control,” Analytica Chimica Acta, vol. 473, no.
1-2, pp. 3–26, 2002.

[8] L. L. Jessome and D. A. Volmer, “Ion suppression: a major
concern in mass spectrometry,” LC-GC North America, vol. 24, no.
5, pp. 498–510, 2006.

[9] T. M. Annlesley, “Ion suppression in mass spectrometry,” Clinical
Chemistry, vol. 49, no. 7, pp. 1041–1044, 2003.

[10] J. P. Antignac, K. De Wasch, F. Monteau, H. De Brabander, F.
Andre, and B. Le Bizec, “The ion suppression phenomenon
in liquid chromatography-mass spectrometry and its conse-
fuences in the field of residue analysis,” Analytica Chimica Acta,
vol. 529, no. 1-2, pp. 129–136, 2005.

[11] Y. Hsieh, M. Chintala, H. Mei et al., “Quantitative screening
and matrix effect studies of drug discovery compounds in
monkey plasma using fastgradient liquid chromatogra-
y/tandem mass spectrometry,” Rapid Communications in
Mass Spectrometry, vol. 15, no. 24, pp. 2481–2487, 2001.

[12] B. K. Matuszewski, M. L. Constanzer, and C. M. Chavez-Eng,
“Strategies for the assessment of matrix effect in quantitative
bioanalytical methods based on HPLC-MS/MS,” Analytical
Chemistry, vol. 75, no. 13, pp. 3019–3030, 2003.

[13] C. Müller, P. Schäfer, M. Störtzel, S. Vogt, and W. Wein-
mann, “Ion suppression effects in liquid chromatography-
electrospray-ionisation transport-region collision induced
dissociation mass spectrometry with different serum extrac-
tion methods for systematic toxicological analysis with mass
spectra libraries,” Journal of Chromatography B, vol. 773, no. 1,
pp. 47–52, 2002.

[14] R. Bonfiglio, R. C. King, T. V. Olah, and K. Merkle, “The
effects of sample preparation methods on the variability of the
electrospray ionization response for model drug compounds,” Rapid Communications in Mass Spectrometry, vol. 13, no. 12,
pp. 1175–1185, 1999.

[15] C. R. Mallet, Z. Lu, and J. R. Mazzeo, “A study of ion sup-
pression effects in electrospray ionization from mobile phase
additives and solid-phase extracts,” Rapid Communications in
Mass Spectrometry, vol. 18, no. 1, pp. 49–58, 2004.

[16] J. X. Shen, R. J. Motyka, J. P. Roach, and R. N. Hayes,
“Minimization of ion suppression in LC/MS/MS analysis
through the application of strong cation exchange solid-
phase extraction (SCX-SPE),” Journal of Pharmaceutical and Biomedical Analysis, vol. 37, no. 2, pp. 359–367, 2005.

[17] M. Hočapček, K. Volná, P. Janda et al., “Effects of ion-pairing
reagents on the electrospray signal suppression of sulphonated
dyes and intermediates,” Journal of Mass Spectrometry, vol. 39,
no. 1, pp. 43–50, 2004.

[18] E. Chambers, D. M. Wagrowski-Diehl, Z. Lu, and J. R. Mazzeo,
“Systematic and comprehensive strategy for reducing matrix
effects in LC/MS/MS analyses,” Journal of Chromatography B,
vol. 852, no. 1-2, pp. 22–34, 2007.

[19] H. G. J. Mol, P. Plaza-Bolaños, P. Zomer, T. C. De Rijk, A. A.
M. Stolker, and P. P. J. Mulder, “Toward a generic extraction
method for simultaneous determination of pesticides, myco-
toxins, plant toxins, and veterinary drugs in feed and food
matrixes,” Analytical Chemistry, vol. 80, no. 24, pp. 9450–9459,
2008.

[20] N. Lindegardh, A. Annerberg, N. J. White, and N. P. J. Day,
“Development and validation of a liquid chromatographic-
tandem mass spectrometric method for determination of
pipeazine in plasma. Stable isotope labeled internal stand-
ard does not always compensate for matrix effects,” Journal of
Chromatography B, vol. 862, no. 1-2, pp. 227–236, 2008.

[21] L. Kantiani, M. Farré, J. M. Grases I Freixiedas, and D. Barceló,
“Determination of antibacterials in animal feed by pressurized
liquid extraction followed by online purification and liquid
chromatography- electro spray tandem mass spectrometry,” Analytical and Bioanalytical Chemistry, vol. 398, no. 3, pp.
1195–1205, 2010.

[22] F. Van Holthoon, P. P. J. Mulder, E. O. Van Bennekom, H.
Heskamp, T. Zuidema, and H. J. A. Van Rhijn, “Quantitative
analysis of penicillins in porcine tissues, milk and animal
feed using derivatisation with piperidine and stable iso-
dilution liquid chromatography tandem mass spectrometry,” Analytical and Bioanalytical Chemistry, vol. 396, no. 8, pp.
3027–3040, 2010.

[23] M. J. G. de la Huebra, U. Vincent, and C. von Holst, “Determi-
nation of semduramycin in poultry feed at authorized level by
liquid chromatography single quadrupole mass spectrometry,” Journal of Pharmaceutical and Biomedical Analysis, vol. 53, no. 4,
pp. 860–868, 2010.

[24] U. Vincent, Z. Ezerskis, M. Chedin, and C. von Holst, “Deter-
nmination of ionophore coccidiostats in feeding stuffs by
liquid chromatography-tandem mass spectrometry. Part II.
Application to cross-contamination levels and non-targeted
feed,” Journal of Pharmaceutical and Biomedical Analysis, vol.
54, no. 3, pp. 526–534, 2011.

[25] M. Cronly, P. Behan, B. Foley, E. Malone, P. Shearan, and L.
Regan, “Determination of eleven coccidiostats in animal feed
by liquid chromatography-tandem mass spectrometry at cross
contamination levels,” Analytica Chimica Acta, vol. 700, no. 1-
2, pp. 26–33, 2011.

[26] R. Liu, W. Hei, P. He, and Z. Li, “Simultaneous determination
of fifteen illegal dyes in animal feeds and poultry products by
ultra-high performance liquid chromatography tandem mass
spectrometry,” Journal of Chromatography B, vol. 879, no. 24,
pp. 2416–2422, 2011.

[27] W. Li, T. J. Herrman, and S. Y. Dai, “Determination of afla-
toxins in animal feeds by liquid chromatography/tandem mass
spectrometry with isotope dilution,” Rapid Communications in
Mass Spectrometry, vol. 25, no. 9, pp. 1222–1230, 2011.

[28] P. Delahaut, G. Pierret, N. Ralet, M. Dubois, and N. Gillard,
“Multi-residue method for detecting coccidiostats at carry-
over level in feed by HPLC-MS/MS,” Food Additives and
Contaminants—Part A, vol. 27, no. 6, pp. 801–809, 2010.

[29] C. Van Poucke, F. Dumoulin, and C. Van Peteghem, “Deter-
cination of banned antibacterial growth promoters in animal feed
by liquid chromatography-tandem mass spectrometry: opti-
misation of the extraction solvent by experimental design,” Analytica Chimica Acta, vol. 529, no. 1-2, pp. 211–220, 2005.

[30] C. Van Poucke, K. De Keyser, A. Baltusmièkine, J. D. G.
McEvoy, and C. Van Peteghem, “Liquid chromatographic-
tandem mass spectrometric detection of banned antibacterial
growth promoters in animal feed,” Analytica Chimica Acta,
vol. 483, no. 1-2, pp. 99–109, 2003.

[31] S. O’Connor, J. Locke, and D. S. Aga, “Addressing the chal-
leges of tetracycline analysis in soil: extraction, clean-up, and
matrix effects in LC-MS,” Journal of Environmental Monitor-
ing, vol. 9, no. 11, pp. 1234–1262, 2007.

[32] M. Lavin, T. Alsberg, Y. Yu, M. Adolfsson-Erici, and H. Sun,
“Serial mixed-mode cation- and anion-exchange solid-phase
extraction for separation of basic, neutral and acidic pharmaceuticals in wastewater and analysis by high-performance liquid chromatography-quadrupole time-of-flight mass spectrometry,” *Journal of Chromatography A*, vol. 1216, no. 1, pp. 49–62, 2009.

[33] A. Tölgyesi, L. Tölgyesi, V. K. Sharma, M. Sohn, and J. Fekete, “Quantitative determination of corticosteroids in bovine milk using mixed-mode polymeric strong cation exchange solid-phase extraction and liquid chromatography-tandem mass spectrometry,” *Journal of Pharmaceutical and Biomedical Analysis*, vol. 53, no. 4, pp. 919–928, 2010.

[34] D. R. Baker and B. Kasprzyk-Hordern, ”Multi-residue analysis of drugs of abuse in wastewater and surface water by solid-phase extraction and liquid chromatography-positive electrospray ionisation tandem mass spectrometry,” *Journal of Chromatography A*, vol. 1218, no. 12, pp. 1620–1631, 2011.
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