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6	IDENTIFICATION OF METABOLITES AND THERMAL TRANSFORMATION
7	PRODUCTS OF QUINOLONES IN RAW COW MILK BY LIQUID
8	CHROMATOGRAPHY COUPLED TO HIGH RESOLUTION MASS
9	SPECTROMETRY
10 11 12 13 14 15 16	A. Junza ¹ , S. Barbosa ² , R. Codony ³ , A. Jubert ³ , J. Barbosa ^{1,2} , D. Barrón ¹ *
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19	¹ Universitat de Barcelona, Departament de Química Analítica, Campus de l'Alimentació
20	de Torribera.
21	Avda. Prat de la Riba, 171, 08921 Sta. Coloma de Gramanet, Barcelona, Spain
22	² Universitat de Bracelona, Departament de Química Analítica,
23	Martí i Franquès, 1-11, 08028 Barcelona, Spain
24	³ Laboratori Interprofessional Lleter de Catalunya (ALLIC)
25	Ctra. Vilassar a Cabrils s/n, 08348 Cabrils, Barcelona, Spain
26 27 28	* To whom correspondence should be sent. Phone: +34-934033797
29	E-mail: dolores.barron@ub.edu

30 ABSTRACT

The presence of residues of antibiotics, metabolites and thermal transformation products
(TPs), produced during thermal treatment to eliminate pathogenic microorganisms in milk,
could be represent a risk for people.

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35 Cow milk samples spiked with enrofloxacin (ENR), ciprofloxacin (CIP), difloxacin (DIF) 36 and sarafloxacin (SAR) and milk samples from cows medicated with ENR were submitted 37 to several thermal treatments. The milk samples were analysed by liquid chromatography 38 mass spectrometry (LC-MS) to find and identify TPs and metabolites. In this work, 27 TPs 39 of four quinolones and 24 metabolites of ENR were found. Some of these compounds had 40 been reported previously, but others were characterized for the first time, including lactose-41 conjugated CIP, the formamidation reaction for CIP and SAR, and hydroxylation or ketone 42 formation to produce three different isomers for all quinolones studied.

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Keywords: antibiotics, milk, quinolones, thermal treatments, transformation products,
metabolites, elucidation, ToF, LTQ-Orbitrap

49 **1. INTRODUCTION**

Quinolones are one of the classes of antibiotics that are most widely used in veterinary practice to treat bacterial infections of animals in livestock farming and bovine milk production. Antibiotics are also used as growth promoters, although this practice has been forbidden by the European Union (EU) since 2006¹.

The presence of antibiotic residues in food is a potential risk to consumers because of direct toxic effects such as allergic reactions or the induction of resistant strains of bacteria^{2,3}. To ensure human health, the EU has established maximum residue limits (MRL) for some antibiotics in foodstuffs of animal origin (37/2010/EC)⁴. Active substances are usually included in this regulation. However, their metabolites such as ciprofloxacin, the main metabolite of enrofloxacin, are only considered in a few cases.

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61 Milk is one of the most commonly consumed foods in the world. Drug residues and metabolites can be found in milk if the time between drug administration and milking is 62 63 too short. Before consumption, cow milk has to be subjected to thermal treatment to eliminate pathogenic microorganisms. Two common thermal processes are pasteurization, 64 65 which consists of heating at 60°C for 30 min or at 72°C for 15 s, and sterilization, by which milk is heated at 120°C for 20 min^{5,6}. During thermal processes, the antibiotic residues and 66 67 their metabolites could undergo transformations, depending on the temperature and the duration of the process. 68

The study of metabolites and their thermal degradation products is of interest because these new compounds may be a health risk to consumers and could be more persistent and harmful than the drug administered⁷.

72 Numerous papers have been published on analytical methods to determine target substances and their main metabolites in several matrices⁸⁻²¹. However, there are few 73 74 studies on the identification and determination of metabolites, degradation, and 75 transformation products (TPs). The analytical methods consist of chromatographic 76 separation prior to detection with a mass spectrometer. Several strategies are used to identify non-targets, including accurate mass measurement to determine the elemental 77 composition and distinguish isobaric molecular ions, multiple mass fragmentations (MSⁿ), 78 and complementary techniques such as NMR to confirm structure^{7,22-34}. Some metabolites 79 of quinolones are described in the bibliography^{25,35-41}, but to our knowledge no study has 80 81 been carried out on metabolites in milk which has been exposed to thermal treatment.

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The aim of this study was to characterize the metabolites and TPs of four quinolones ENR, CIP, DIF and SAR (Figure 1) in cow's milk subjected to thermal treatment. In order to test the new metabolites and TPs found in spiked milk samples, milk samples from animals medicated with ENR were submitted to the same thermal treatment. Samples were analysed by liquid chromatography (LC) coupled to high resolution mass spectrometry (ToF and LTQ-Orbitrap).

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90 2. EXPERIMENTAL PROCEDURES

91 **2.1. Chemicals and apparatus**

92 The standards were purchased from several pharmaceutical firms: ENR from Cenavisa
93 (Reus, Spain), CIP from Ipsen Pharma (Barcelona, Spain), DIF from Abbott (Madrid,
94 Spain) and SAR from AK Scientific Inc. (CA,USA).

All reagents were of analytical grade unless indicated. Formic acid (HOCHO), acetic acid
(HOAc), acetonitrile (MeCN), methanol (MeOH), sodium dihydrogenphosphate and
sodium hydroxide (NaOH) were supplied by Merck (Darmstadt, Germany). Water was
ultrapurified by a Milli-Q system (Millipore, MA, USA).

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101 The quinolone individual stock solutions (ENR, CIP, DIF and SAR) were prepared at a 102 concentration of 500 μ g·mL⁻¹ in 50 mM HOAc aqueous solution. The individual working 103 solutions that were used to spike the milk samples were prepared by dilution of the 104 individual stock solution at a concentration of 20 μ g·mL⁻¹ in water.

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Sodium dihydrogenphosphate solution 0.1 M adjusted with NaOH 5 M to pH 10 was addedto milk samples prior to the clean-up step.

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109 The SPE cartridges used in this study were Oasis HLB (3 cm³/60 mg), obtained from
110 Waters (Milford, MA, USA).

111

112 To simulate the thermal treatment of milk, a laboratory oven (Memmert) was used to warm 113 the samples. A Rotanta 460RS (Hettich Zentrifuguen, Germany) centrifuge was used to 114 separate precipitated proteins and fat of supernatant after the extraction. The SPE step was 115 carried out in a Supelco vacuum manifold with disposable liners for 24 cartridges 116 (Bellefonte, PA, USA), connected to a Supelco vacuum tank. Evaporation to dryness was 117 performed in a TurboVap LV (Caliper Life Science, Hopkinton, MA, USA). A Crison 118 2002 potentiometer (±0.1 mV) (Crison, Barcelona, Spain) with a Crison 5203 combined pH electrode from Orion Research (Boston, MA, USA) was used to measure the pH of the 119 120 phosphate solution and the mobile phase.

121 **2.2. Milk samples**

Milk samples used in the work have been supplied by the Laboratori Interprofessional Lleter de Catalunya (ALLIC), control laboratory of milk in Catalonia. The samples were analysed previously in ALLIC using a screening method to determine if the milk contained residues of antibiotics. The milk samples that were negative in antibiotics were used as blank samples in this work. Four positive samples in ENR were used to study the metabolites.

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129 **2.3. LC-MS instrumentation and working conditions**

130 **2.3.1. LC conditions**

Quinolones, their metabolites and degradation products were separated in a Zorbax Eclipse XDB-C8 column (5 μ m, 4.6 \times 150 mm) from Agilent Technologies (Santa Clara, CA, USA), using a pre-column Kromasil C8 (5 μ m, 4.6 \times 15 mm) supplied by Akady (Barcelona, Spain) when samples were analysed by LC-ToF. A Pursuit UPS C18 column (2.4 μ m, 2 \times 50 mm) from Varian (Harbor City, CA, USA) was used when samples were injected into the LC-LTQ-Orbitrap.

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For both columns, the mobile phase was composed of an aqueous solution of 0.1% HOCHO (solvent A) and MeCN with 0.1% HOCHO (solvent B) at a constant flow rate of 1 mL·min⁻¹ and 0.3 mL·min⁻¹, respectively. The injection volume was 20 μ L in LC-ToF and 10 μ L in LC-LTQ-Orbitrap. Separations were carried out in a Zorbax Eclipse XDB-C8 column for 11 min under the following gradients: from 0 to 1 min, 15% B; 4 min, 45% B; 7 min, 56% B; 8.5 min, 15% B; and 11 min, 15% B. Separations were carried out in a Pursuit UPS C18 column under the following conditions: from 0 to 3.5 min, 1% B; 4.5 145 min, 25% B; 5 min, 50% B; 6.5 min, 50% B; 7.5 min, 25% B; 8.5 min, 10% B; and 11
146 min, 1% B.

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148 **2.3.2. LC-MS (ToF)**

A HP Agilent Technologies 1100 LC system was equipped with an autosampler and coupled to a 6220 oa-ToF LC/MS mass spectrometer with an electrospray ionisation source (ESI) (Agilent Technologies, Santa Clara, CA, USA). The system was controlled by Mass Hunter workstation software (Agilent Technologies, Santa Clara, CA, USA) for the acquisition and processing of data from the ToF mass spectrometer.

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The optimum parameters of ToF in positive mode were as follows: capillary voltage 4000 V, drying gas (N₂) temperature 300°C, drying gas (N₂) flow rate 9 L·min⁻¹, nebulizer gas (N₂) 40 psi, fragmentor voltage 150 V, skimmer voltage 60 V, and OCT 1 RF voltage 250 V. The ToF-MS mass resolving power was approximately 10,000 FWHM (Full width at half maximum) at m/z 922. Spectra were acquired over the m/z 50-1100 range. Data storage was in profile and centroid modes.

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163 **2.3.3. LC-MS (LTQ-Orbitrap)**

An Accela LC system (Thermo Scientific, Hemel Hempstead, UK) was equipped with a thermostatic autosampler and coupled to an LTQ Orbitrap Velos mass spectrometer (Thermo Scientific, Hemel Hempstead, UK), with an ESI source. XCalibur software was used for the data analyses.

169 The operation parameters used were source voltage, 3.5 kV; sheath gas (N₂): 40 (arbitrary 170 units); auxiliary gas (N_2) : 10 (arbitrary units); sweep gas (N_2) : 10 (arbitrary units); and 171 capillary temperature, 275°C. Default values were used for most other acquisition parameters (Fourier transform (FT) automatic gain control (AGC) target 1.10⁶ for MS 172 mode and $5 \cdot 10^4$ for MSⁿ mode). Milk samples were first analysed in full MS mode with the 173 174 Orbitrap mass resolving power set at 30,000 FWHM at m/z 400. The following analyses were carried out in MSⁿ mode with the Orbitrap mass resolving power set at 15,000 175 176 FWHM at m/z 400. The maximum injection time was set to 100 ms with one micro scan for MS mode and to 500 ms with one micro scan for MSⁿ mode. TPs were fragmented in 177 178 the HCD Collision Cell because some metabolites were not fragmented in the trap. The 179 selected HCD voltage was 45 V, although in some cases it was raised to 50 and 60 V. The 180 mass range was from m/z 100 to 1000.

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182 **2.4. Procedures**

183 **2.4.1 Thermal study**

In this study, individual standards of quinolones (ENR, CIP, DIF and SAR) at 1 µg·mL⁻¹ in 184 185 water were warmed to degrade the antibiotics. All quinolones were heat-treated under 186 conditions of temperature and time corresponding to three thermal processes of milk: 187 pasteurization at 60°C for 30 min (T1) or 72°C for 15 s (T2) and sterilization at 120°C for 188 20 min (T3). Subsequently, individual milk samples spiked with ENR, CIP, DIF and SAR 189 were warmed at 120°C for 20 and 60 min (T3.20 and T3.60). The samples heated at 190 different temperatures were compared with spiked non-heated samples (T0) and with blank 191 milk samples at same thermal condition, ie, for example samples at T3.20 were compared 192 with blank at T3.20. Milk samples from animals medicated with ENR were also thermally treated at T3.20 and T3.60 and compared with the same sample but non-heated. Allexperiments were made by triplicate.

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196 **2.4.2. Sample treatment and clean-up (SPE)**

197 The method for analyzing the milk sample was used previously to determine the antibiotics 198 in the milk⁹⁻¹². The extraction method consist of the addition of 0.5 mL of phosphate 199 solution 0.1 M at pH 10 and 2 mL of water to 2 g of milk, centrifugation of samples, and an 200 SPE process. The SPE cartridges used were Oasis HLB and these were preconditioned 201 with 1 mL of MeOH, 1 mL of water and 1 mL of 0.1 M phosphate solution at pH 10. After 202 sample loading, the cartridge was washed with 3 mL of water and the analytes were eluted 203 with 2 mL of MeOH.

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205 The methanolic fraction eluted from the SPE was evaporated to dryness under a nitrogen 206 stream by TurboVap LV and a water temperature of 35°C. The extract was then 207 reconstituted with 200 μ L of water and filtered through membrane filters (Ultra free 208 Durapore PVDF 0.22 μ m from Millipore), before injection into the LC-LTQ-Orbitrap.

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210 **2.4.2. Data treatment**

The LC-ToF data were treated using two procedures to find metabolites and TPs in milk samples from medicated cow with ENR. The first corresponds to the manual comparison of mass spectra of blanks and samples. This procedure was used initially to detect the TPs in standards and spiked cow milk. The other option consists of the use of Mzmine2^{42,43} free software, which provides a list of all m/z on mass spectra. The list can be shortened by removing all the ions that only appear in the blank spectra and those that have a similar or higher intensity in the blanks than in the samples. Only the ions which appeared between 1 to 9 min and from 150 to 800 Da were considered. Also, a mass defect filter (MDF) of 120
mDa was applied.

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222 **3. RESULTS AND DISCUSSION**

223 **3.1. Determination of thermal TPs by LC-ToF-MS**

Blank and spiked water samples were subjected to the thermal conditions explained in Section 2.3.1 and analysed by LC-ToF-MS. The total ion current chromatograms (TIC) of spiked water samples were compared with the TIC of blank water samples to examine for possible TPs. The two chromatograms were nearly identical, except for the spiked antibiotic peak.

229

The mass spectra of blank and spiked water samples were compared by overlapping. The only difference between the spiked sample and the blanks was the peak of parent compound. However, a decrease in the area of the protonated molecule of quinolones was noticed when the temperature rose. The highest decrease was seen when the samples were heated at 120°C for 20 min (T3.20) where in some cases up to 20%. A new condition of 120°C for 60 min (T3.60) was also applied to ensure that TPs were obtained in sufficient high concentrations to be detected.

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Raw cow milk samples free of quinolones were spiked and heated under the selected conditions. To assess whether any TPs had formed, the same procedure as that used for the water standards was followed. In this case, some ions were identified in the mass spectra of the spiked milk samples that were not observed in the blank milk samples. Some ions were identified as doubly charged protonated molecules, with characteristic losses of water and

243 carbon dioxide. These ions were also detected in the water samples. The doubly protonated identity was confirmed by the isotopic pattern. The difference between $[M+2H]^{2+}$ and 244 $[M+1+2H]^{2+}$ was m/z 0.5. In addition, the same retention time (t_R) of the compounds in the 245 extracted ion chromatogram (EIC) was observed⁴⁴. The losses of H₂O (18.0105) and CO₂ 246 (43.9898) are not considered as TPs because these were also observed in the source by 247 CID-in-source fragmentation^{40,45-48}. The ions m/z 332.1405 and 386.1311 were detected in 248 samples that were spiked with ENR and DIF respectively. These m/z values are consistent 249 with the m/z of CIP and SAR. The EIC were obtained for blank and spiked samples to 250 251 check the presence of TPs. Table 1 shows all the ions found in milk and their identification, sorted by the original antibiotic. As can be observed, the errors between 252 253 experimental mass and assigned structure were lower than 3.0 ppm.

254

To assign a structure to the remaining of the ions that were observed, an in-house database was created using a list of 106 metabolic reactions⁴⁹ that can take place in milk, and another list of milk components (including carbohydrates, amino acids, vitamins and lipids; a total of 188 components)⁵⁰. The exact mass of the starting antibiotic or a known fragment is introduced into the database and a series of combinations of reactions and/or possible additions are obtained that give rise to a molecule with an exact mass similar to that observed in the mass spectra of ToF.

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- 263

264 **3.2. Identification of thermal TPs by LC-LTQ-Orbitrap-MS**

As a ToF spectrometer cannot fragment molecules, we used an LTQ-Orbitrap to elucidate the TPs, as it has a higher mass resolving power than ToF and can fragment the compound to obtain structural information. For each quinolone, Table 2 shows the retention time (t_R), m/z values, the gains and losses of mass compared to the parent quinolone, the m/z of the TPs observed, and the fragments observed in mass spectra. Table 2 also shows the proposed reactions, with the assigned molecular formula, the double bond equivalents (DBE) of neutral molecules, the theoretical mass, and the error in ppm calculated for the formula. The information is provided as common reactions that occurred in all the studied quinolones, and other reactions that were only observed in some quinolones.

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277 *Common reactions of the four quinolones*

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279 Several gains and losses of mass were common among the four quinolones studied. Our database shows that the gain of 15.9949 m/z (Table 2) could correspond to two different 280 281 reactions: hydroxylation and N-oxide formation. An evaluation of MS/MS spectra could be 282 used to differentiate between both structures. As an example, Figure 2A shows the MS/MS 283 spectrum of ions m/z 416.1416 in samples of DIF and discussion about elucidation is given 284 in supporting information. The position of OH could not be assigned with certainty, but our 285 proposal is hydroxylation in position 3 of the piperazine ring, as in oxo-ciprofloxacin, the metabolite described in the bibliography^{36,37,40}. For the four quinolones, the hydroxylated-286 287 TPs were observed, but the concentration of TPs from ENR and DIF were higher than CIP and SAR ones. N-oxide formation only occurred in ENR and DIF in N⁴-alkylpiperazinyl. 288 This metabolite has been described previously^{38,51}. For this structure, a loss of OH is 289 290 characteristic in fragmentation and was experimentally observed in Figure 2B of the 291 MS/MS spectrum of ions m/z 416.1416 for DIF. In this case, the DBEs of TPs undergo no 292 alteration of values in comparison to the DBE of parent quinolones. Different fragmentation patterns were observed for the hydroxylated and N-oxide compounds that were found, as can be observed in Figure 2A and B. In Figure 2B, corresponding to the Noxide TP, the peak of the precursor ion disappeared almost completely when the same voltage was applied (HCD: 45). This finding suggests that hydroxylated compounds are more stable than N-Oxide TP. The structures assigned to the rest of the ions are shown in Figure 2. Table 2 shows the fragmentation ions, the molecular formula, the DBE, and the calculated error in the assignation of the different structures.

300

301 A gain of 29.9740 m/z was also observed for the four quinolones, as shown in Table 2. 302 Three peaks were found for ENR and DIF, whereas only one peak was observed for their 303 main metabolites (CIP and SAR), at a concentration too low to be isolated and fragmented. 304 The mass spectra of all peaks of ENR and DIF were studied. Figure 3 shows the mass 305 spectra of ions with m/z 390.1460 of ENR. In Figure 3A, the separation by LC-LTQ-306 Orbitrap is shown with peaks at 5.71, 7.57 and 7.99 min. Although the retention time of 307 ENR matches peak 1, the mass spectra are different, as can be observed in Figures 3B and 308 C. A comparison of both mass spectra revealed only two ions (m/z 245.1077 and 72.0805) 309 in common. The m/z 245.1077 indicates that the differences between the TP of peak 1 and 310 ENR are in the loss fragment (part of the piperazinyl ring). The accurate mass of the 311 fragment m/z 128.0706 corresponds to molecular formula C₆H₁₀NO₂ with a DBE value of 312 3 and is the key to the elucidation as is shown in Figure 3C and in the supporting 313 information⁵².

314

315 Unlike the first molecule, the other two ions of m/z 390.1460 (peaks 2 and 3) had low 316 fragmentation as can be observed in Figure 3. Nevertheless, their structures were 317 elucidated. The retention time of these peaks are different, but their mass spectra are 318 identical, which indicates that they are isomers (Figure 3D). The m/z 245.1085 observed in 319 the mass spectrum of ENR (Figure 3B) could not be found in the spectra of peaks 2 and 3. 320 The molecules were broken up from the piperazinyl ring, resulting in the ions 100.0756 321 and 291.0774. The fragment 100.0756 is the same as that observed in Figure 3C, which 322 indicates that there was only a ketone in the ring. The fragmentation of 291.0774 makes us 323 to conclude that OH is on the aromatic ring. Two different vacant positions in the aromatic 324 ring led to the two different isomers. For DIF, the fragmentations were similar to those 325 shown in ENR, as can be observed in Table 2.

326

327 TPs corresponding to net loss of 26.0146 were observed for all compounds (ENR, CIP, DIF and SAR). For the TPs of ENR and DIF the losses of N⁴-alkylamine from piperazinyl 328 ring were observed. However, as CIP and SAR do not have an alkyl group in N^4 , the losses 329 330 of ammonia (17.0275) were seen. Moreover, the TPs had DBE one unit lower than the 331 parent compound, due to the break in the piperazinyl ring. Therefore, the peaks at m/z332 334.1560, 306.1246, 374.1307 and 360.1156 have been assigned to desethylenequinolone TPs³⁴⁻³⁷. As Table 2 shows, when Orbitrap was used to elucidate the structures, the 333 334 calculation errors (ppm) were lower than 2 ppm. Slightly higher error values were obtained 335 when ToF was used.

336

337 Other reactions

The major metabolites, CIP and SAR, underwent other transformations, as can be observed in Table 2. Due to the absence of the alkyl chain in the N⁴-piperazinyl ring, two reactions acetylation (Acetyl-CIP (m/z 374.1509) and Acetyl-SAR (m/z 428.1416)) and formamidation (Formamide-CIP (m/z 375.1463) and Formamide-SAR (m/z 429.1416))

took place. The corresponding explanation of elucidation is in supplementary material.

344 In addition to the above-mentioned reactions, CIP gave three more TPs (m/z: 274.0986, 345 573.1429 and 656.2457). The ion 573.1429 was observed at a very low concentration and 346 could not be fragmented. For the mass 274.0986, the ion 292.1091 could also be observed, 347 at less intensity, corresponding to the gain of a water molecule. The main ion obtained by MS^2 of the ion 274.0986 was 231.0564, and this was isolated and fragmented again. In the 348 349 MS^3 spectrum, the ion m/z 249.0668 was also observed with m/z 231.0564 which 350 corresponds to gain of a water molecule. This gain of water was only observed in the ions 351 that kept the carbonyl group of acid, which is capable of hydrating the initial carboxylic 352 acid again. The rest of the ions were mainly due to fragmentation of the piperazine ring. 353 The structure assigned to m/z 292.1091 was CIP, with loss of the cyclopropyl group.

354

The last TP (656.2457) was obtained by heating the spiked milk sample with CIP at T3.60. When a HCD voltage of 45 V was used, only fragments with low m/z were found. To obtain structural information, the voltage was reduced to 35 V. The CIP ion (m/z 332.1400) could be identified among all the fragments as can be seen in Figure 4. As can be followed in supporting information, the molecule CIP (332.1405) is combined with the lactose (galactose + glucose) present in milk at high concentration. This new product was favoured when samples were heated at a high temperature for a long time (60 min).

362

363 3.3. Effect of temperature on the stability of TPs

For each TP ion, the areas obtained from milk samples were plotted *vs*. the heating conditions (T0, T3.20 and T3.60) (Figure 5). For ENR and CIP (Figure 5A and 5B), all the TPs were observed at T0 except for hydroxylated quinolones (m/z 376.1656_1 and 348.1334, respectively). The concentration of these ions was raised by heating at T3.20, and disappeared at T3.60. In the case of ENR (Figure 5A), the concentration of some TPs, including CIP (332.1402), desethylene-ENR (334.1569) and two isomers of hydroxylation and ketone formation (390.1461_1 and 390.1461_3), increased slightly by warming at T3.20, and decreased at T3.60. For CIP, T3.60 was required for conjugation with lactose (656.2477) to occur (Figure 5B). The acetylated (374.1535) and formamidated (375.1463) were present at T0 and their concentrations increased by heating. The 362.1148 ion (corresponding to gain $[M+H+29.974]^+$) was initially present (T0), but was destroyed by heating.

376

377 In contrast, the concentration of most of the ions in DIF samples increased slightly when 378 the temperature was increased (Figure 5C). A general downward trend of DIF ions was 379 seen at T3.60. Completely different behaviour was observed for the 430.1185_3 ion, which 380 underwent sharp decrease at T3.20 and a sharp rise in formation when the warming time 381 was increased, possibly due to the ion coming from different sources, and for the 382 430.1185 1 ion, which disappeared completely at T3.20. The acetylated (428.1407) and 383 formamidated (429.1391) TPs of SAR (Figure 5D) behaved in the same way as those of 384 CIP, *i.e.*, their concentration increased with temperature.

385

386 3.4 Determination of metabolites and TPs in medicated cow milk

Four raw milk samples from cows medicated with ENR were submitted to the same thermal treatments as the spiked milk samples. The extracts were analysed by LC-ToF and the metabolites and TPs were fragmented by LC-LTQ-Orbitrap.

390

In this case, metabolites and TPs were found using the two data treatments (Section 2.3.3).
When exact mass, retention time and MS/MS spectra were compared, some of the
compounds in milk spiked with ENR and CIP were also present in medicated cow milk.

Table 3 shows the retention time (t_R) of metabolites in samples analysed by LC-ToF, the accurate m/z of metabolites determined by ToF and LTQ-Orbitrap, the assigned molecular formula, the theoretical mass, the error in ppm calculated for the assigned formula, the proposed structures, and the double bond equivalents (DBE) of the neutral molecule.

399

400 CIP (m/z 332.1405), the main metabolite of ENR, was one of the common compounds. In 401 addition, desethylene metabolites of ENR (m/z 334.1561)(ENR-6) and CIP (m/z402 306.1248)(ENR-3) were observed. The TPs of ENR corresponded to hydroxylation (ENR-403 16) and N-Oxide formation (ENR-17)(m/z 376.1667) and two isomers of compound m/z404 390.1460 (ENR-18 and ENR-19) (hydroxylation and ketone formation, Figure 3D) were 405 also found. Four TPs of CIP were observed in medicated cow milk. At 3.6 min was found 406 the hydroxy-CIP (m/z 348.1354)(ENR-8) and at 6.1 min the m/z 362.1147(ENR-11), although this time the MS^2 spectrum could not be obtained due to its low concentration. 407 408 The other metabolites were from acetylation (m/z 374.1511)(ENR-14) and formamidation 409 (*m*/*z* 375.1463)(ENR-15) of CIP.

410

411 Other compounds were found in milk from medicated cows, but not in spiked milk. These 412 compounds were produced by metabolism of the antibiotic in the cow, and some of them 413 were identified for the first time.

414

The metabolite with the highest intensity after CIP (332.1405) and desethylene-ENR (334.1561)(ENR-6) had m/z 263.0834 (ENR-1). The m/z of the metabolite was lower than the m/z of ENR. Therefore, the MS/MS spectrum was analysed to find which part of the parent compound had been modified. Figure 6A shows MS/MS spectrum of ENR-1. It

419 was concluded that the metabolite structure corresponded to CIP without a piperazine ring420 (Supporting information). Only the amino group was maintained, as shown in Table 3.

421

422 In the MS/MS spectrum of metabolite ENR-2 with m/z 291.0783 (Figure 6B), the ion m/z423 263.0834 (ENR-1), discussed above in supporting information and in Figure 6A, was 424 observed. The difference between both ions corresponded to a carbonyl group (27.9949). 425 The assigned structure for this metabolite is given in Table 3.

426

427 Another metabolite elucidated was m/z 307.1095 (ENR-4). Some explanations of the 428 fragments are shown in Figure 6C and supporting information. Its structure can be 429 observed in Table 3.

430

The ion m/z 334.1189 (ENR-5) was identified as formyldesethylene-CIP²⁵. Its structure was checked by its mass spectrum. The molecule was broken up from the piperazinyl ring, resulting in the ions m/z 72.0445 and 263.0834 cited before. Other important fragments were m/z 245.0730 [263.0834-H₂O], 230.0489 [334.1189-H₂O-Cyclopropyl-Formamide] and 217.0411 [230.0489-CH].

436

437 A compound (ENR-9) with the same m/z as hydroxy-CIP (m/z 348.1354)(ENR-8) was also 438 studied, because the retention time differed by about 4 minutes from the peak observed in 439 spiked samples. The MS/MS spectrum (Figure 6D) was very similar to the spectrum of 440 formyldesethylene-CIP (m/z 334.1189)(ENR-5). The most striking difference between both 441 spectra was that instead of fragment m/z 72.0444 was observed a fragment of m/z 86.0601. 442 This fragment corresponds to the addition of a methyl group to fragment m/z 72.0444. In 443 conclusion, a structure of acetyldesethylene-CIP (ENR-9) is suggested, which could be 444 formed from the oxidation of the side chain of desethylene-ENR (m/z 334.1561)(ENR-6).

445

446 The EIC of ion m/z 362.1137 (ENR-12 and ENR-13) from a full-scan of ToF was very 447 similar to that obtained for the m/z 390.1460 (ENR-18 and ENR-19) in spiked samples 448 with ENR (Figure 3). Three peaks were observed in the EIC. The first one was at 6.1 min, 449 which coincides with the peak observed in samples fortified with CIP. Also were seen two 450 peaks at 6.8 and 7.1 min, with the same shape as the two peaks at 7.6 and 7.99 in Figure 3. 451 Unfortunately, when the sample was analysed by LC-LTQ-Orbitrap, it was degraded and 452 only one peak was observed and fragmented. In the MS/MS spectrum, the ions m/z453 291.0776 and 259.0820 were present, as elucidated previously in Figure 3D. Another 454 fragment that helped to determine the structure corresponded to m/z 72.0444, which is like 455 fragment 100.0757 of metabolite 390.1460 without the alkyl chain of ENR. Due to the 456 resemblances between the EICs and mass spectra of ions m/z 362.1137 (ENR-12 and ENR-457 13) and 390.1460 (ENR-18 and ENR-19), can be assume that these compounds are the 458 same, but one is from CIP and the other from ENR, respectively.

459

460 The last two elucidated metabolites corresponded to oxo-CIP (m/z 346.1198)(ENR-7) and 461 formyl-CIP (360.1354)(ENR-10), which have been described previously in literature^{36,40}. 462 Their structures were confirmed by their MS/MS spectra.

463

464 The rest of the metabolites found in medicated cow milk could not be elucidated. In any 465 case, Table 3 shows their m/z, retention time (t_R), hypothetical molecular formula, double 466 bond equivalents (DBE) and errors in ppm.

468 **AUTHOR INFORMATION**

469 **Funding source**

470 The authors are grateful to the Ministerio de Ciencia e Innovación (MICINN) (Project

- 471 CTQ2010-19044/BQU) for financial support. A.J. received support from MICINN (BES-
- 472 2011-045849) for the training of researchers.
- 473

474 Supporting Information Available

- 475 Explanation about MS elucidation of some TPs and metabolites of quinolones (ENR, CIP,
- 476 DIF and SAR) has been included. This material is available, free of charge, via the Internet
- 477 at http://pubs.acs.org.
- 478

480

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637	
638	FIGURE CAPTIONS
639	
640	Fig 1
641	Structures of the four quinolones studied (ENR, CIP, DIF and SAR).
642	
643	Fig 2
644	A) MS ² spectrum for the hydroxylated TP of DIF. B) MS ² spectrum for the N-oxide TP of
645	DIF.
646	
647	Fig 3
648	A) EIC of ion m/z 390.1460 in the milk sample spiked with ENR. B) MS ² spectrum for
649	ENR. C) MS^2 spectrum for the first peak (t _R 5.71 min). D) MS^2 spectrum for the remaining
650	peaks (t _R 7.57 and 7.99 min).
651	
652	Fig 4
653	Product ion scan (MS ²) of CIP lactose conjugate (precursor ion m/z 656.2457). Top left,
654	extended mass spectrum where losses of water molecules are shown.
655	
656	Fig 5
657	Influence of the different warming conditions in areas of different TPs of ENR, CIP, DIF
658	and SAR.
659	
660	Fig 6
661	MS^2 spectra for the metabolites of ENR. A) m/z 263.0828 (ENR-1). B) m/z 291.0776
662	(ENR-2). C) <i>m/z</i> 307.1089 (ENR-4). D) <i>m/z</i> 348.1354 (ENR-9).
663	

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Table 1. Accurate masses of ions observed in milk spiked samples but not in blank milk samples by LC-ToF. The errors (ppm) are calculated from the theoretical and proposed structures.

ENR				CIP			
<i>m/</i> 2 (exp)	<i>m/z</i> (theorical)	Structure	Error (ppm)	<i>m/</i> 2 (exp)	<i>m/z</i> (theorical)	Structure	Error (ppm)
157.5685	157.5686	$[CIP+2H-H_2O]^{2+}$	0.6	157.5690	157.5686	$[CIP+2H-H_2O]^{2+}$	2.5
166.5740	166.5739	$[CIP+2H]^{2+}$	0.6	166.5742	166.5739	$[CIP+2H]^{2+}$	1.8
167.5820	n.a *	$[334.1569]^{2+}$	I	274.0989	n.a	n.a	I
171.5845	171.5843	$[ENR+2H-H_2O]^{2+}$	1.2	288.1511	288.1507	$[CIP+H-CO_2]^+$	1.4
180.5897	180.5897	$[ENR+2H]^{2+}$	0.1	314.1293	314.1288	$[CIP+H-H_2O]^+$	1.6
316.1817	316.1820	[ENR+H-CO ₂] ⁺	0.9	332.1400	332.1405	[CIP+H] ⁺	1.5
332.1402	332.1405	[CIP+H] ⁺	0.9	348.1334	n.a	n.a	I
334.1569	n.a	n.a	I	362.1148 ^b	n.a	n.a	I
342.1603	342.1612	$[ENR+H-H_2O]^+$	2.6	374.1509	n.a	n.a	I
360.1719	360.1718	[ENR+H] ⁺	0.3	375.1563	n.a	n.a	ı
376.1656 ^a	n.a	n.a	I	573.1431	n.a	n.a	I
390.1461 ^b	n.a	n.a	1	656.2477	n.a	n.a	I
719.3349	719.3363	[2ENR+H] ⁺	1.9				
DIF				SAR			
2/ M	2/ W	Striicture	Error	2/ M	2/ W	Structure	Error
(exp)	(theorical)	211111111	(mdd)	(exp)	(theorical)		(mdd)
191.5714	191.5717	$[DIF+2H-H_2O]^{2+}$	0.5	184,5637	184.5639	$[SAR+2H-H_2O]^{2+}$	1.1
200.5761	200.5770	$[DIF+2H]^{2+}$	3.0	193,5692	193.5692	[SAR+2H] ²⁺	0.1
299.0980	n.a	n.a	I	342,1422	342.1412	$[SAR+H-CO_2]^+$	2.9
374.1284	n.a	n.a	I	360,1181	n.a	n.a	I
382.1367	382.1362	$[DIF+H-H_2O]^+$	1.3	368,1193	368.1205	[SAR+H-H ₂ O] ⁺	3.3
386.2209	386.1311	[SAR+H] ⁺	0.3	386.1311	386.1311	$[SAR+H]^+$	0.1
400.1446	400.1467	[DIF+H] ⁺	2.0	402.1241	n.a	n.a	I
416.1396^{a}	n.a	n.a	I	416,1069	n.a	n.a	I
430.1185 ^b	n.a	n.a	I	428.1410	n.a	n.a	I
				429.1395	n.a	n.a	ı

(*) n.a: Non-assigned.
(a) 2 chromatographic peaks with the same m/z (at different retention time).
(b) 3 chromatographic peaks with the same m/z (at different retention time).

Table 2. Accurate masses of parent quinolones, given reactions, all TPs observed and the fragments obtained by LC-LTQ-Orbitrap. The DBE calculated with the deprotonated molecular formula and the calculated error (ppm) of the assignation of structure.

	t. ^a		Gains or	Reactions			Molecular		Theorical	Fror
	(mim)	m/z	losses	(*)	m/z TPs	Fragmentation	formula	DBE	mass	(mdd)
Commo	n reacti	ons of 4 quin	olones							
ENR	4.7	360.1718	+15.9949	ΧН	376.1666	358.1560, 340.1121, 286.1018, 244.0912, 243.0877	$\mathrm{C}_{19}\mathrm{H}_{23}\mathrm{FN}_{3}\mathrm{O}_{4}$	10	376.1667	-0.3
ENR	5.8	360.1718	+15.9949	O-N	376.1665	359.1639, 330,1611, 315.1741, 300.1506, 257.1085, 231.0928, 84.0807	$C_{19}H_{23}FN_{3}O_{4}$	10	376.1667	-0.5
CIP	3.6	332.1405	+15.9949	НΧ	348.1355	330.1248, 310.1184, 136.0617	$C_{17}H_{19}FN_3O_4$	10	348.1354	0.3
DIF	6.0	400.1467	+15.9949	НХ	416.1416	398.1307, 343.0880, 329.0725, 316.0647, 256.0801	$C_{21}H_{20}F_2N_3O_4$	13	416.1416	0.0
DIF	6.3	400.1467	+15.9949	0-N	416.1418	399.1388, 370.1359, 355.1489, 299.09899	$C_{21}H_{20}F_2N_3O_4$	13	416.1416	0.5
SAR	5.8	386.1311	+15.9949	ХН	402.1261	384.1156, 382.1197, 364.1089, 343.0882, 316.0653, 136.0758	$C_{20}H_{18}F_2N_3O_4\\$	13	402.1260	0.2
ENR	5.8	360.1718	+ 29.9740	HX + K	390.1458	372.1353, 362.1509, 346.1560, 318.1246, 301.0982, 289.09813, 245.1083, 128.0706, 100.0756, 72.0808	$C_{19}H_{21}FN_3O_5$	11	390.1460	-0.5
ENR	6.7	360.1718	+ 29.9740	HX + K	390.1462	372.1356, 362.1510, 291.07754, 273.0670, 259.0514, 100.0757, 72.0808	$C_{19}H_{21}FN_3O_5$	11	390.1460	0.5
ENR	6.8	360.1718	+ 29.9740	HX + K	390.1461	372.1355, 362.1509, 291.0774, 273.0669, 259.0513, 100.0756, 72.0809	$C_{19}H_{21}FN_3O_5$	11	390.1460	0.3
CIP	6.1	332.1405	+29.9740	HX + K	362.1141 ^b		$C_{17}H_{17}FN_3O_5$	11	362.1147	-1.7
DIF	6.0	400.1467	+ 29.9740	HX + K	430.1209	412.1102, 402.1259, 386.1309, 368.1204, 329,0730, 299.0988, 197.1029, 114.0549, 70.0650	$C_{21}H_{18}F_2N_3O_5$	14	430.1209	0.0
DIF	6.9	400.1467	+ 29.9740	HX + K	430.1207	412.1101, 402.1258, 384.1151, 345.0678, 327.0572, 86.0599	$C_{21}H_{18}F_2N_3O_5$	14	430.1209	-0.5
DIF	7.1	400.1467	+ 29.9740	HX + K	430.1207	412.1101, 402.1258, 384.1151, 345.0678, 327.0572, 86.0599	$C_{21}H_{18}F_2N_3O_5$	14	430.1209	-0.5
SAR	6.7	386.1311	+29.9740	HX + K	416.1053 ^b		$C_{20}H_{16}F_2N_3O_5$	14	416.1053	0.0
ENR	4.4	360.1718	- 26.0146	DE + H	334.1560	316.1464, 311.3906, 296.1393, 289.098, 263.0825, 219.0927, 164.1069, 86.0600, 72.0807	$C_{17}H_{21}FN_3O_3$	6	334.1561	-0.3
CIP	3.6	332.1405	- 26.0146	DE + H	306.1246	288.1141, 268.108, 263.0824, 245.1084	$C_{15}H_{17}FN_3O_3$	6	306.1248	-0.7
DIF	5.8	400.1467	- 26.0146	DE + H	374.1307	354.1246, 343.0884, 336.1138, 317.0728, 299.0988, 280.1242, 273.0830	$C_{19}H_{18}F_2N_3O_3$	12	374.1311	-1.1
SAR	5.9	386.1311	- 26.0146	DE + H	360.1156	343.0883, 342.1048, 340.1089, 322.0986, 317.07310, 299.0991	$C_{18}H_{16}F_2N_3O_3$	12	360.1154	0.6

	3	5	5	0	0	0	0		9
	-0-	ġ	°,	0.(0.0	0.(0.0		-0-
	332.1405	386.1311	374.1511	428,1416	375.1463	429.1369	274.0986		656.2461
	10	13	11	14	11	14	10		12
	$C_{17}H_{19}FN_3O_3$	$C_{20}H_{18}F_2N_3O_3$	$C_{19}H_{21}FN_3O_4$	$C_{22}H_{20}F_2N_3O_4$	$C_{18}H_{20}FN_4O_4$	$C_{21}H_{19}F_2N_4O_4\\$	$C_{14}H_{13}FN_{3}O_{2}$		$C_{29}H_{39}FN_3O_{13}$
Other reactions			$\begin{array}{c} 356.1402, \ 314.1297, \ 300.0777, \ 272.0827, \ 249.0669, \\ 243.0563, \ 231.0562, \ 215.0249 \end{array}$	410.1310, 382.1361, 368.1203	$\begin{array}{c} 357.1356,\ 340.1096,\ 332.1402,\ 314.1297,\ 270.0671,\\ 243.0563,\ 231.0563,\ 216.0692 \end{array}$	411.1264, 386.1312, 368.1205, 348.1141	$\begin{array}{ccccccc} 231.0562^{\circ}, & 217.0404^{\circ}, & 203.0611, & 187.0663, \\ 175.0664, 163.0665, 148.0556, 136.557^{\circ} \end{array}$		638.2450, 620.2246, 494.1929, 476.1825, 404.1616, 344.1400, 332.1401
	332.1404	386.1309	374.1509	428.1416	375.1463	429.1369	274.0986 ^c	573.1429 ^b	656.2457
	DE	DM	Α	А	н	ц	DY +DCP		LC
	- 28.0303	- 14.0157	+ 42.0103	+ 42.0103	+ 43.0056	+43.0056	- 58.0419	+241.0024	+ 324.1056
	360.1718	400.1467	332.1405	386.1311	332.1405	386.1311	332.1405	332.1405	332.1405
	5.0	6.0	7.0	8.0	6.6	7.2	5.0	5.0	4.0
	ENR	DIF	CIP	SAR	CIP	SAR	CIP	CIP	CIP

^a Retention time when LC-ToF is used.

^b Concentration too low to be fragmented; ^c These masses were also observed with a molecule of water (+ 18.0105) but at low concentration.

(*) HX: hydroxylation; N-O: N-Oxide; K: ketone formation; DE: deethylation; H: Hydrogenation; CC: cysteine conjugation; DC: Decarboxylation; DF: Reductive defluorination; DM: Demethylation; A: Acetylation; F: Formamidation; DY: Dehydration; DCP: Decyclopropylation; LC: Lactose conjugation.

and their theorical mass assigned to molecular formula are given with calculated error (ppm) for the assignation of suggested structure for the Table 3. Metabolites and TPs found in medicated cow milk with ENR sorted by accurate mass. Accurate masses analysed by ToF and Orbitrap

two analy:	sers.			-		-			
2/ m	<i>m/z</i>	<i>z/m</i>	$t_{ m R}{}^{ m a}$	Molecular	Error	Error	Summestad Structura	DRF	
(ToF)	(Orbitrap)	theorical	(min)	formula ^b	(ToF)	(Drbitrap)	ouggesteu ou ucunte	DDE	
263.0828	263.0834	263.0826	6.85	$C_{13}H_{12}FN_2O_3^+$	0.8	3.0	H H H H H H	6	ENR-1
288.1349	288.1345	288.1343	2.9	$C_{15}H_{18}O_3N_3^+$	2.1	0.7	n.a		
291.0770	291.0783	291.0776	6.7	$C_{14}H_{12}FN_{2}O_{4}^{+}$	-2.1	2.4		10	ENR-2
302.1501	302.1506	302.1499	5.7	$C_{16}H_{20}N_{3}O_{3}^{+}$	0.7	2.3	n.a		
306.1239	306.1256	306.1248	3.6	$C_{15}H_{17F}N_{3}O_{3}^{+}$	-2.9	2.6	The second secon	6	ENR-3
307.1083	307.1097	307.1089	6.5	$C_{15}H_{16}FN_2O_4^+$	-1.9	1.9		6	ENR-4
329.1485	329.1498	329.1496	3.4	$C_{18}H_{21}N_2O_4^+$	-3.3	0.6	n.a		
329.1485	329.1498	329.1496	4.9	$C_{18}H_{21}N_2O_4^+$	-3.3	0.6	n.a		

CIP	ENR-5	ENR-6	ENR-7	ENR-8
10	10	6	11	10
		HO H	HO +D +D +D +D +D +D +D +D +D +D +D +D +D	HO H
0	2.7	1.5	1.7	-1.1
0.9	-2.7	-6.5	-2.3	-3.1
C ₁₇ H ₁₉ FN ₃ O ₃ +	$C_{16}H_{17}FN_{3}O_{4}^{+}$	$C_{17}H_{21}FN_3O_3^+$	$C_{17}H_{17}FN_{3}O_{4}^{+}$	$C_{17}H_{19}FN_{3}O_{4}^{+}$
5.0	6.4	4.35	6.5	3.6
332.1405	334.1198	334.1561	346.1198	348.1354
332.1405	334.1207	334.1566	346.1204	348.1350
332.1408	334.1189	334.1539	346.1190	348.1343

ENR-9	ENR-10	ENR-11	ENR-12	ENR-13	ENR-14
10	11	11	11	11	11
					H H H H H H H H H H H H H H H H H H H
2.9		1.4	1.4	1.4	2.7
-3.1	0.8	-2.8	-2.8	-2.8	-0.3
$C_{17}H_{19}FN_{3}O_{4}^{+}$	C ₁₇ H ₁₉ FN ₃ O ₄		$\mathrm{C}_{17}\mathrm{H}_{17}\mathrm{FN}_{3}\mathrm{O}_{5}^{+}$	$C_{17}H_{17}FN_3O_5^+$	$C_{19}H_{21}FN_{3}O_{4}^{+}$
6.5	6.9		6.8	7.1	7.0
348.1354	348.1354		362.1147	362.1147	374.1511
348.1364	360.1360	362.1152	362.1152	362.1152	374.1523
348.1343	360.1357	362.1137	362.1137	362.1137	374.1510

ENR-15	ENR-16	ENR-17	ENR-18	ENR-19
11	10	10	11	11
	To			
0.3	1.1	1.6	0.3	0.3
-0.3	-2.9	-2.9	-0.3	-0.3
$C_{18}H_{20}FN_4O_4^+$	$C_{19}H_{23}FN_3O_3^+$	$C_{19}H_{23}FN_3O_3^+$	$C_{19}H_{21}FN_{3}O_{5}^{+}$	$C_{19}H_{21}FN_{3}O_{5}^{+}$
6.6	4.6	5.8	6.65	6.85
375.1463	376.1667	376.1667	390.1460	390.1460
375.1464	376.1671	376.1673	390.1461	390.1461
375.1462	376.1656	376.1656	390.1459	390.1459

^a Retention time when LC-ToF is used. ^b These molecular formulas are hypothetical in case of metabolites without assigned structure. n.a : Non-assigned





Fig 1







Fig 4





Fig 6

Electronic Supporting Information for the research article:

IDENTIFICATION OF METABOLITES AND THERMAL TRANSFORMATION PRODUCTS OF QUINOLONES IN RAW COW MILK BY LIQUID CHROMATOGRAPHY COUPLED TO HIGH RESOLUTION MASS SPECTROMETRY

A. Junza¹, S. Barbosa², R. Codony³, A. Jubert³, J. Barbosa^{1,2}, D. Barrón¹*

¹Departament de Química Analítica, Campus de l'Alimentació de Torribera Avda. Prat de la Riba, 171, 08921 Sta. Coloma de Gramanet, Barcelona, Spain

²Departament de Química Analítica

Martí Franquès, 1-11, 08028 Barcelona, Spain

³ Laboratori Interprofessional Lleter de Catalunya (ALLIC)

Ctra. Vilassar a Cabrils s/n, 08348 Cabrils, Barcelona,

* Email: dolores.barron@ub.edu

Content:

• Explanations about MS elucidation of some TPs and metabolites of quinolones (ENR, CIP, DIF and SAR).

Section 3.2: Common reactions of the four quinolones

1. TPs of DIF (*m*/*z* 416.1416)

In the mass spectrum of the ion m/z 416.1416 (Figure 2A), two losses of water (18.0105 m/z) were observed. The first was due to carboxylic acid and the same loss was observed for the rest of the quinolones. The second loss of water observed in the spectra is only possible if an OH group is in position 2 or 3 of the piperazine ring.

2. TPs of ENR (*m/z* 390.1460) Peak 1

The accurate mass of the fragment m/z 128.0706 corresponds to molecular formula $C_6H_{10}NO_2$ with a DBE value of 3 and is the key to the elucidation (Figure 3C). This was followed by two consecutive fragmentations, corresponding to losses of 27.9949 (CO) [128.0706 - CO \rightarrow 100.0756; 100.0756 - CO \rightarrow 72.0808] and the reduction of one unit of DBE. These fragments indicate that a ketone and an alcohol are present in the piperazinyl ring, as observed in the suggested fragmentation shown in Figure 3C. The rest of the fragments are due to losses of CO₂, H₂O and pieces of the piperazine ring. It should be noted that the DBE of some fragments such as m/z 301.0982 and 328.1454 are 10.5. The structures represented in the mass spectrum (Figure 3C) have 11 unsaturations (double bonds and rings). The DBE calculated is the average of the resonance structures of drawn structure (DBE: 11) and the carbocation (DBE:10)⁵².

Section 3.2: Other reactions

1. TPs of CIP and SAR (acetylation and formamidation)

Two different reactions took place on CIP and SAR. The first was acetylation, which could be justified by a loss of acetyl group (42.0103) (Acetyl-CIP(-H₂O): 356.1402 \rightarrow 314.1297 and Acetyl-SAR(-H₂O): 410.1310 \rightarrow 368.1203), as observed in the MS spectra (not shown). The second reaction corresponds to formamidation. For CIP and SAR, the loss of formamide and subsequent gain of 2H occurred in one step (Formamide-CIP(-H₂O): 357.1356 \rightarrow 314.1296 and Formamide-SAR(-H₂O): 411.1264 \rightarrow 368.1205). Moreover, in the case of CIP, the loss of NH₃ (357.1356 \rightarrow 340.1096), followed by the loss of a carbonyl and gain of 2H (340.1096 \rightarrow 314.1296), were also observed. Due to the absence of the alkyl chain in the N⁴-piperazinyl ring, both reactions (acetylation and formamidation) took place.

2. TPs of CIP (*m*/*z* 656.2457)

Figure 4 shows the mass spectrum of this TP with its fragmentations. The two main characteristic losses were the loss of -180.0633 (656.2457 $\rightarrow 476.1825$), which corresponds to the loss of a glucose, and the loss of a galactose without a water molecule (476.1825 $\rightarrow 344.1400$). In the spectrum, steady losses of six water molecules were observed due to the OH present on the sugars, as can be seen in the smaller mass spectrum in Figure 4. All these losses could be explained if the molecule CIP (332.1405) is combined with the lactose (galactose + glucose) present in milk at high concentration. This new product was favoured when samples were heated at a high temperature for a long time (60 min).

Section 3.4: Determination of metabolites and TPs in medicated cow milk

1. Metabolite ENR-1 (*m/z* 263.0834)

In the MS/MS spectrum of ENR-1 (Figure 6A), the loss of a water molecule (263.0834 \rightarrow 245.0729) and CO (245.0729 \rightarrow 217.0778) from carboxylic acid can be observed. The presence of cyclopropyl can also be justified by ion *m/z* 222.0441, which is the result of the loss of cyclopropyl from the parent metabolite (263.0834 – 41.0391 \rightarrow 222.0441).

2. Metabolite ENR-4 (*m/z* 307.1095)

In this case, characteristic losses of the carboxylic group and cyclopropyl could be observed in the spectrum of ENR-4 (Figure 6C). The modifications of structure with respect to ENR, were again found over the piperazine ring. As can be seen, the ion m/z 248.0596 loses the methyl group, which results in the ion m/z 235.0520 that later loses a molecule of water (235.0520 \rightarrow 217.0411).