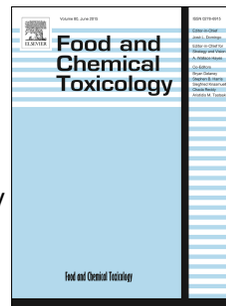


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Preface to the special issue of food and chemical toxicology on the outcomes of the MARLON project on veterinary epidemiology of potential health impacts of genetically modified feeds in livestock

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1 **Preface to the special issue of Food and Chemical Toxicology on the**
2 **outcomes of the MARLON project on veterinary epidemiology of**
3 **potential health impacts of genetically modified feeds in livestock**

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7
8 This special issue is a collection of articles dedicated to the impact of genetically
9 modified (GM) animal feeds on the health of livestock. It features the outcomes
10 of the research project titled "Monitoring of Animals for Feed-related Risks in the
11 Long Term (MARLON)". MARLON was a three-year research project funded by
12 the European Union's (EU) Seventh Framework Program for research and
13 technology development (MARLON, 2015). It focused on monitoring livestock in
14 commercial production scenarios for potential health impacts associated with
15 feeds containing GM ingredients. There has been a rapid rate of adoption of GM
16 feed ingredients especially GM-soybean, maize, oilseed rape and cottonseed in
17 non-EU countries, since their commercial large-scale introduction in the mid-90s
18 (James, 2017). They are now common ingredients in livestock feed. In addition
19 to addressing GM feeds, the methodology and insights gained from the project
20 can be broadly applied to the surveillance of health effects of livestock feeds and
21 hazards in general.

22
23 MARLON operated at the interface of science, legislation and policy. It brought
24 together scientists from 11 European institutions with complementary expertise
25 including veterinary health and epidemiology, livestock production, biosafety of
26 GM crops, food and feed safety. MARLON's activities included stakeholder
27 consultations, a training course and a final conference. The main outcomes of
28 MARLON were new knowledge, guidelines and methodologies for monitoring GM
29 feed ingredient-related health impacts for key stakeholders including European
30 regulators. MARLON's progressive approach relates to the possibility that
31 monitoring of GM feeds and foods may become necessary because they are
32 regulated and can only be marketed after regulatory approval. In fact, EU
33 legislation on GM foods and feeds has a provision that a post-market monitoring
34 *may* be required based on the outcome of the pre-market risk assessment (EU,
35 2003). Thus, a key outcome of MARLON was providing European policy makers
36 with pertinent new knowledge on several safety topics regarding GM crops which
37 added to the outcomes from other EU projects including GMSAFOOD, GRACE and
38 G-TwYST projects and the French government-funded GMO90+ projects that
39 focussed on *in vivo* and *in vitro* biomarkers, methods for safety testing of GM
40 crops and collation of the existing evidence on the potential health,
41 environmental and economic impacts of GM crops.

42
43 When the MARLON project began, there were a few challenges to address. For
44 example, a requirement for *case-specific monitoring* to verify potential risks
45 identified or assumptions made during the pre-market safety assessment was
46 only necessary for foods derived from GM oilseed crops with a modified fatty
47 acid composition, which might have potential to impact human nutrition and
48 health. In addition, there was a requirement for general, passive surveillance of

49 animal health impacts as part of the *general surveillance* carried out for all GM
50 crops that had received approval for “environmental release” (e.g., field
51 cultivation, import and processing of seeds and other viable GM crop products)
52 (EU, 2001). Therefore, MARLON activities had to be anticipatory because it was
53 not possible to rely on past case-specific monitoring experience. This was
54 especially challenging because negative health effects due to approved GM
55 products were not known to exist. Another challenge was that the approach to
56 developing monitoring methods should be universally applicable because it was
57 not possible to foresee for which combination of GM feed, livestock species and
58 health parameters monitoring could be required. Additionally, it was necessary
59 to use data from existing animal health surveillance activities, which were either
60 generic or focused on the impact of non-feed-related animal health.

61

62 This special issue addresses the following questions:

- 63 • Are there known health impacts related to GM-crop-derived livestock feeds?
64 (De Vos and Swanenburg, 2018)
- 65 • Would it be possible to measure the exposure of animals to specific GM feed
66 ingredients? (Nadal et al., 2018)
- 67 • Can health indicators be specified by regulators for the monitoring of
68 livestock for possible health effects of GM feeds? (De Santis et al., 2018)
- 69 • How are different feed and livestock production chains organized and
70 characterised (e.g., husbandry practices, regulations, traceability, and health
71 checks)? (Kleter et al., 2018)
- 72 • Is it possible to design monitoring programs to exploit data from existing
73 surveillance for the detection of unusual trends in animal health parameters
74 which is collated in IPAFEED and relate this to the consumption of GM feeds?
75 (Vince et al., 2018)

76

77 De Vos and Swanenburg (De Vos and Swanenburg, 2018) in their review of the
78 literature addressed whether there are there known health impacts related to
79 GM-crop-derived livestock feeds. They focussed on controlled livestock feeding
80 studies with the aim to collate state-of-the-art knowledge on health parameter
81 (e.g., serum chemistry, pathology, etc.) perturbations associated with feeding of
82 GM ingredients. They explored relevant previously reported health parameter
83 changes and provided a systematic annotation of all health parameters
84 measured in a newly established database called IPAFEED (IPAFEED, 2015;
85 Vince et al., 2018), which is available to regulators to assess the design of
86 monitoring schemes based on previously reported health parameters. Notably,
87 there were no adverse effects linked to GM feeds on any health parameters.

88

89 The review by Nadal and co-workers (Nadal et al., 2018) focussed on whether it
90 would be possible to measure the exposure of animals to specific GM feed
91 ingredients with the notion that it might link exposure to ingredients with
92 adverse health effects in an individual or cohort of animals. They also addressed
93 whether ingredients could be traced and detected throughout the feed and
94 livestock production chains, and whether there are markers for exposure in
95 livestock animals, (e.g., via uptake of GM material from feed into tissues, fluids,
96 and excreta). The review concludes that although it is possible to measure the
97 presence of specific ingredients in feeds based on GM-crop-related proteins or
98 DNA, there are no markers for animal exposure. Therefore, traceability
99 measures have not yet traced and quantified an animal's exposure to a specific
100 GM feed ingredient.

101
102 De Santis and co-workers addressed whether health indicators can be specified
103 by regulators for the monitoring of livestock for possible health effects of GM
104 feeds in their review (De Santis et al., 2018). They explored four scenarios: 1)
105 potential allergic reactions induced by GM feeds as compared to non-GM feeds in
106 livestock animals; 2) horizontal gene transfer from consumed GM crops to
107 intestinal microorganisms and the host animal; 3) reduced incidence of
108 mycotoxins in GM feeds; and 4) nutritionally altered GM crops. The authors
109 concluded that there were no adverse effects to GM feeding in the literature for
110 these scenarios. There is evidence, however, of beneficial effects related to a
111 reduction of the mycotoxin fumonisin in insect-resistant GM crop varieties and
112 through nutritional enhancement of crops.

113
114 The Kleter and co-workers' (Kleter et al., 2018) review approached the
115 organization and characterization of different feed and livestock production
116 chains (e.g., husbandry practices, regulations, traceability, and health checks)
117 with the aim to determine how the movement of feeds and animals through the
118 production chain are recorded and how this could be used for monitoring of
119 animal health effects. The authors concluded that for livestock, traceability at
120 herd or individual levels are possible depending on the livestock species.
121 Although the existing animal traceability systems are designed for infectious
122 disease reporting or animal identification, it is not possible to trace GM
123 ingredients (even if it is labelled correctly) back to the farms of origin or to verify
124 the distribution and inclusion level in the feeds.

125
126 The last review by Vince and co-workers addresses designing monitoring
127 programs to exploit data from existing surveillance for the detection of unusual
128 trends in animal health parameters, which is collated in IPAFEED, and relate this
129 to the consumption of GM feeds. The review reveals a broad methodology to
130 assist the monitoring of GM feed-related health effects (Vince et al., 2018). This
131 was a key challenge for MARLON because there were no recorded GM feed-
132 related health effects and no methods for quantifying prior exposure of the
133 animals to GM feed. The authors reviewed various types of existing veterinary
134 surveillance, which aim at providing evidence for freedom from specific diseases,
135 early detection of outbreaks, and monitoring of endemic diseases including
136 syndromic surveillance (e.g., non-specific health parameters), risk-based
137 surveillance, collection-point-based surveillance, checkpoint surveillance,
138 sentinel herd surveillance and representative surveys. In addition, they reported
139 on a probabilistic approach towards syndromic surveillance that estimates the
140 likelihood that an effect will be detected above background variability through
141 existing reporting systems. Importantly, this approach considers the production
142 chain, from farmer via veterinarian and clinical laboratory, to post-mortem
143 abattoir inspection. The authors concluded that observed effects or deviating
144 trends would need follow-up with confirmatory research under controlled
145 conditions and suggested that managers should consider cost-effectiveness
146 compared with alternative options when setting up a surveillance scheme.

147
148 In summary, the reviews in this section provide comprehensive overviews of
149 published controlled feeding livestock studies with GM crops checking for health
150 impacts beyond solely performance, the possibilities to detect GM feed
151 ingredients and the animals' exposure to them, the possibility to employ health
152 indicators for four hypothetical scenarios including allergenicity, horizontal gene

153 transfer, changed mycotoxin levels, and nutritional improvement, the
154 organization of feed and livestock production chains in Europe, and the
155 possibilities for applying veterinary epidemiology to surveillance of possible GM
156 feed-related health impacts. The generic, statistical approach developed within
157 MARLON will assist in the design of monitoring programs based on syndromic
158 surveillance for GM and non-GM feeds which would then rely on existing data of
159 health syndromes commonly measured in livestock production. This also concurs
160 with recent trends in syndromic surveillance, mining of big data and domestic
161 animal toxicology.

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