

# **Resolution on the draft Commission implementing decision authorising the placing on the market of products containing, consisting of or produced from genetically modified soybean DAS-81419-2 pursuant to Regulation (EC) No 1829/2003 of the European Parliament and of the Council**

2021/2759(RSP) - 07/07/2021 - Text adopted by Parliament, single reading

The European Parliament adopted by 470 votes to 199, with 23 abstentions, a resolution **objecting** to the draft Commission implementing decision authorising the placing on the market of products containing, consisting of or produced from genetically modified soybean DAS-81419-2 pursuant to Regulation (EC) No 1829/2003 of the European Parliament and of the Council.

On 9 February 2012, Dow Agro Sciences Ltd submitted an application to the national competent authority of the Netherlands for the placing on the market of foods, food ingredients and feed containing, consisting of or produced from genetically modified soybean DAS-81419-2 ('the GM soybean'). The application also concerned the placing on the market of products containing or consisting of the GM soybean for uses other than food and feed, with the exception of cultivation. On 26 October 2016, the European Food Safety Authority (EFSA) adopted a favourable opinion in relation to that application.

## ***Main concerns from Member States***

Members pointed out that Member States have submitted many critical comments to the EFSA, including that:

- data concerning pest and disease pressure are insufficient for a detailed analysis of ecological interaction of the GM soybean with the environment;
- the applicant only refers to substantial unintended losses of the GM soybean during loading and unloading as a route for environmental exposure and that other routes of exposure of the environment by waste materials from processing or use of the soybean (e.g. manure, faeces from animals fed the GM soybean) were not specifically assessed;
- the proposed monitoring plan does not address relevant questions for the general surveillance of human and animal health and cannot be regarded as sufficiently elaborated.

## ***Lack of assessment of the complementary herbicides and outstanding questions concerning Bt toxins***

A number of studies show that side effects have been observed that may affect the immune system following exposure to Bt toxins and that some Bt toxins may have adjuvant properties, meaning that they can increase the allergenicity of other proteins

with which they come into contact.

Members pointed out that assessment of herbicide residues, and herbicide break-down products, found on GM plants is considered outside the remit of the EFSA Panel on Genetically Modified Organisms and is therefore not undertaken as part of the risk assessment. This is problematic since residues from spraying with glufosinate are known to disturb the microbiome which, for example, may enhance immune reactions in combination with Bt toxins.

### ***Undemocratic decision-making***

Members recalled that the vote on 17 May 2021 of the Standing Committee on the Food Chain and Animal Health delivered no opinion, meaning that the authorisation was not supported by a qualified majority of Member States.

The Commission recognised that the fact that GMO authorisation decisions continue to be adopted by the Commission without a qualified majority of Member States in favour, which is very much the exception for product authorisations as a whole but which has become the norm for decision-making on GM food and feed authorisations, is problematic. It should be noted that in its ninth term, the European Parliament has already adopted 18 objections to placing GMOs on the market.

### ***Recommendations***

On the basis of these considerations, Parliament considered that the Commission's implementing decision exceeds the implementing powers provided for in Regulation (EC) No 1829/2003 and that it is not consistent EU law. Consequently, it called on the Commission to:

- withdraw its draft implementing decision;
- move forward with the utmost urgency concerning the development of sustainability criteria, with full involvement of Parliament;
- not to authorise herbicide-tolerant GM crops until the health risks associated with the residues have been comprehensively investigated on a case-by-case basis, which requires a full assessment of the residues from spraying such GM crops with complementary herbicides, an assessment of the herbicide break-down products and any combinatorial effects, including with the GM plant itself;
- fully integrate the risk assessment of the application of complementary herbicides and their residues into the risk assessment of herbicide-tolerant GM plants, regardless of whether the GM plant concerned is to be cultivated in the Union or is for import into the Union for food and feed uses;
- not to authorise the import for food or feed uses of any GM plant which has been made tolerant to a herbicide-active substance that is not authorised for use in the Union;
- immediately suspend the import of GM soybeans cultivated in Brazil and Argentina until effective legally binding mechanisms have been put in place to prevent the placing on the Union market of products associated with deforestation and related human rights violations.

Lastly, Parliament reiterated its call for the implementation of a European vegetable protein production and supply strategy, which would enable the Union to become less dependent on GM soybean imports and to create shorter food chains and regional markets.