

Brussels, 26 October 2018 F.2/Art. 4.1(b) (2018)6176672

# NOTE FOR THE ATTENTION OF MARC VANHEUKELEN HEAD OF THE EU MISSION TO THE WTO

Subject: Instructions for the regular meeting of the WTO Dispute Settlement Body on 29 October 2018

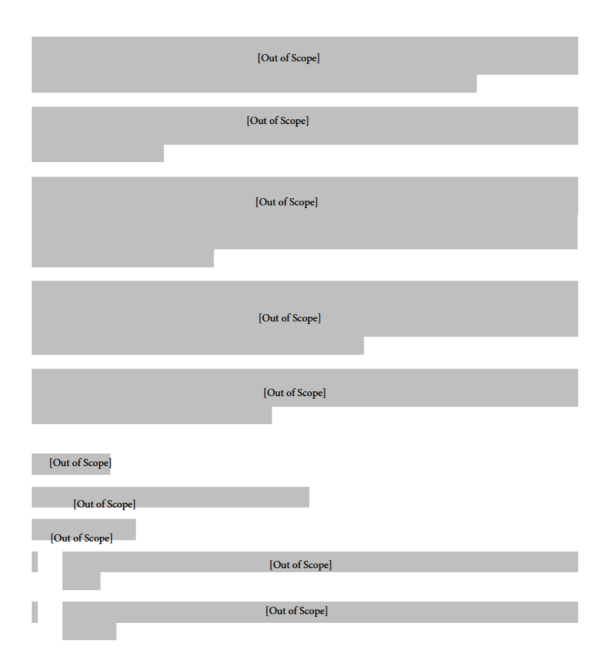
### Summary:

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Eight returning disputes are listed under implementation surveillance:			
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, and in EC-Biotech (DS 291) brought by the US.			
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C. EUROPEAN COMMUNITIES – MEASURES AFFECTING THE APPROVAL AND MARKETING OF BIOTECH PRODUCTS: STATUS REPORT BY THE EUROPEAN UNION (WT/DS291/37/ADD.126)

#### **Background**

In 2003, the US brought WTO proceedings regarding certain EU level and certain Member State level measures relating to the approval and marketing of biotech products claiming that these affected imports of agricultural and food imports from the United States.

Similar WTO proceedings were brought by Canada and Argentina (the three WTO proceedings were later merged into one single dispute, DS 291). The DSB adopted a panel report in November 2006, which found that the EU violated the SPS Agreement on three grounds:

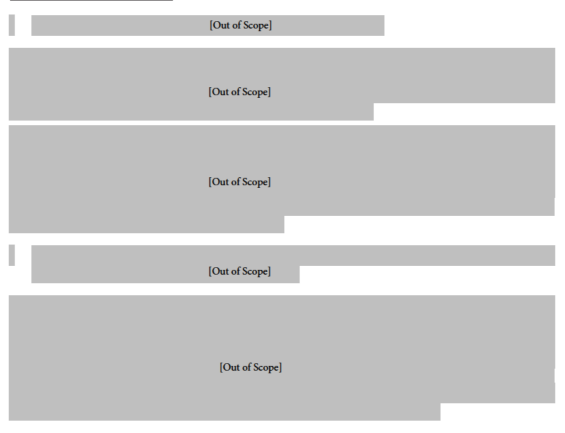
- a) the EU applied a **general** *de facto* **moratorium** on approval of GM products between June 1999 and August 2003;
- b) **undue delays** in the completion of 21 product-specific approval procedures brought forward by the US (out of 25 cases considered by the Panel);
- c) **national safeguard measures** taken by 6 Member States, which were found not to be based on appropriate risk assessments.

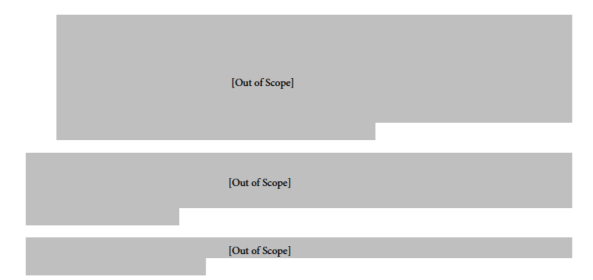
On 19 December 2006, the EU informed the DSB of its intention to implement the Panel's recommendations and findings. The EU signed final settlements with CAN and with Argentina in 2010. The mutually agreed solutions provide for the establishment of regular bilateral dialogues on agricultural biotechnology market access issues of mutual interest.

Contrary to the other two complainants, the US did not consider the establishment of a dialogue sufficient in terms of compliance of the EU and thus, it was not ready to settle the dispute. To the contrary, it made a retaliation request on 17 January 2008, to which the EU objected. On 15 February 2008, and according to the sequencing agreement concluded with the US, both parties requested the suspension of the sanctions arbitration under Article 22.6 DSU. Those sanction arbitration procedures can only be resumed after conclusion of a compliance procedure based on Article 21.5 DSU.

An EU-US **technical dialogue on plant biotechnology** has been taking place on a regular basis since 2008. The last meeting took place on 17 July 2018.

#### Recent EU developments

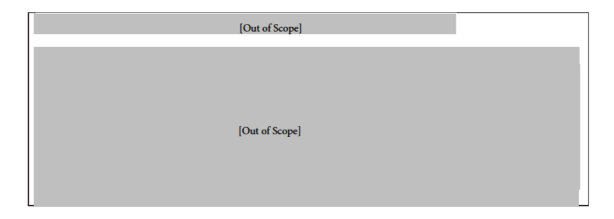




• The ruling of the Court of Justice of the European Union concerning new mutagenesis techniques

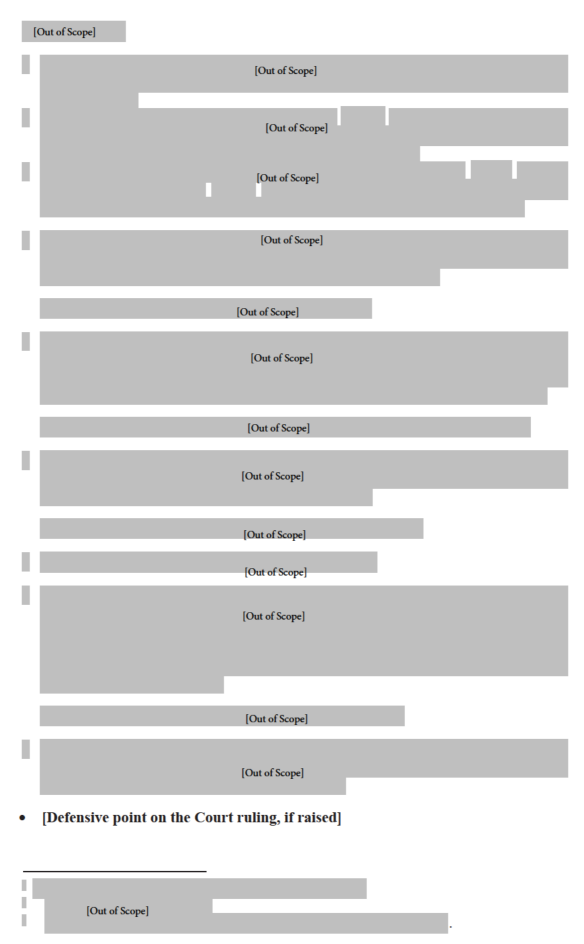
The Court of Justice of the European Union was requested to give a preliminary ruling regarding the regulatory status of organisms produced by means of certain biotechnological techniques known as "new mutagenesis techniques", and in particular on whether such techniques are exempted from the GMO-legislation. The Directive 2001/18/EC, on the deliberate release of GMOs, contains an exemption applicable to "mutagenesis techniques", which already existed before the adoption of the Directive. The ruling was rendered on 25 July 2018.

The outcome of CJEU ruling is that the only organisms obtained by means of techniques or methods of mutagenesis, which have conventionally been used in a number of applications and have a long safety record, are exempted. Therefore, the GMO legislation is applicable to organisms obtained by mutagenesis techniques, which have emerged since the adoption of Directive 2001/18/EC.

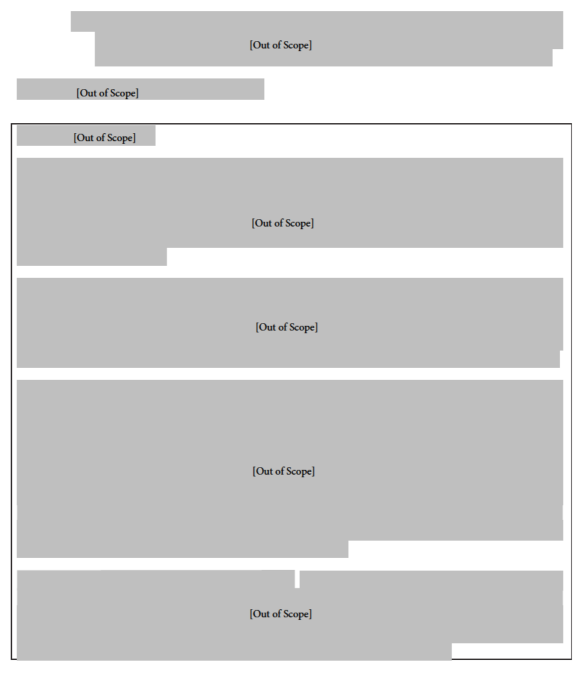


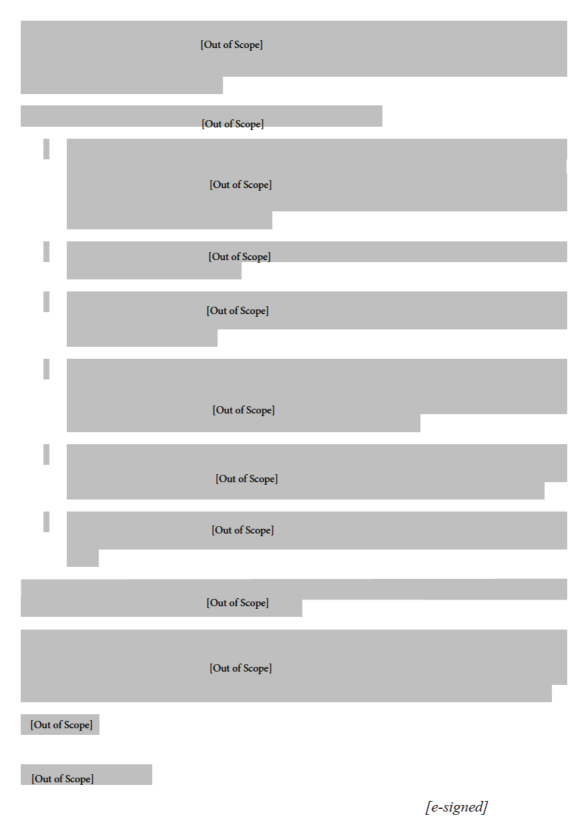
### Line to take

The EU continues to be committed to acting in line with its WTO obligations.

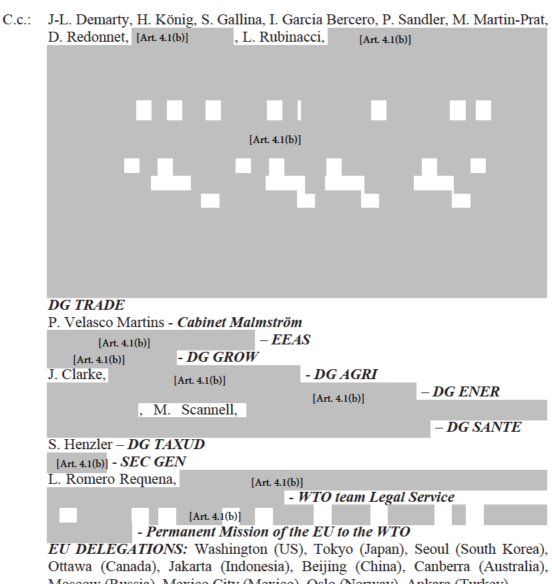


- In July, the Court of Justice of the European Union provided important clarification on the scope of application of the GMO legislation in relation to organisms obtained by mutagenesis techniques.
- The CJEU ruled that organisms obtained by means of new techniques/methods of mutagenesis, which have appeared or have been mostly developed since the adoption of Directive 2001/18, fall within the scope of the Directive.
- The Commission is now analysing the ruling to ensure its proper implementation.
- Operators in and outside the EU remain responsible for ensuring that products which are placed on the market are safe and comply with all relevant regulatory requirements.





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