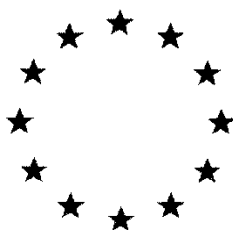


# *European Commission*



**Combined Draft Renewal Assessment Report prepared according to  
Regulation (EC) N° 1107/2009  
and  
Proposal for Harmonised Classification and Labelling (CLH Report)  
according to Regulation (EC) N° 1272/2008**

**HEPTAMALOXYLOGLUCAN**

**Volume 3 – B.1 (PPP) – PEL101GV**

Rapporteur Member State: France  
Co-Rapporteur Member State: Spain

## Version History

| When    | What        |
|---------|-------------|
| 2020-09 | Initial RAR |
|         |             |
|         |             |
|         |             |

### Introduction

The applicant Elicityl prepared a draft renewal assessment report to support the renewal of inclusion of the active substance Heptamaloxyloglucan.

Heptamaloxyloglucan was included in Annex I of Directive 91/414/EEC under Commission Directive 2010/14/EU, which entered into force on 01 June 2010. According to Regulation (EU) No 540/2011, heptamaloxyloglucan is deemed to have been approved under Regulation (EC) No 1107/2009. An extension of approval has been granted by Regulation (EU) 2017/1527 until 31/05/2021.

The Annex I Inclusion Directive for Heptamaloxyloglucan (2010/14/EU) provides specific provisions under Part B which need to be considered by the applicant in the preparation of their submission and by the MS prior to granting an authorisation:

For the implementation of the uniform principles of Annex VI, the conclusions of the review report on heptamaloxyloglucan (SANCO/10502/09 – final, 27/11/2009), and in particular Appendices I and II thereof, as finalised in the Standing Committee on the Food Chain and Animal Health on 27/11/2009 shall be taken into account.

Heptamaloxyloglucan is included in AIR 4 program (SANTE-2016-10616–rev 9, June 2018). The rapporteur Member State is France and the co-rapporteur Member State is Spain (Commission Implementing Regulation (EU) 2016/183 of 11 February 2016).

The draft renewal assessment report has been prepared according to Commission Regulation No.844/2012.

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## B.1. IDENTITY

### B.1.1. IDENTITY OF THE PLANT PROTECTION PRODUCT

|  |   |
|--|---|
| <b>B.1.1.1. Applicant</b>  | ELICITYL SA<br>746 avenue Ambroise Croizat<br>F-38920 Crolles, France   |
| <b>B.1.1.2. Producer of the plant protection product</b>   | ELICITYL SA<br>746 avenue Ambroise Croizat<br>F-38920 Crolles, France   |
| <b>B.1.1.3. Trade name or proposed trade name and producer's development code number of the plant protection product</b> | PEL101GV  |
| <b>B.1.1.4. Detailed quantitative and qualitative information on the composition of the plant protection product</b>     |   |
| <b><i>B.1.1.4.1. Composition of the plant protection product</i></b>   | Technical Heptamaloxyloglucan: 1000 g/kg  |
| <b><i>B.1.1.4.2. Information on the active substances</i></b>  | Heptamaloxyloglucan, min. 780g/kg   |
| <b><i>B.1.1.4.3. Information on safeners, synergists and co-formulants</i></b>   | None  |
| <b>B.1.1.5. Type and code of the plant protection product</b>  | The formulation code which are the closest to the preparation are SP, SG or ST, however the applicant claims that PEL101GV is neither a powder nor a granule nor a tablet. It is a solid to be used after dissolution in water.<br><br>Considering the properties of the product, RMS considers it should be considered as a water soluble powder (SP). |
| <b>B.1.1.6. Function</b>   | Protection against frost damage   |
| <b>B.1.1.7. Field of use envisaged</b>   | Grape   |
| <b>B.1.1.8. Effects on harmful organisms</b>   | No claimed effect on harmful organisms  |

### B.1.2. REFERENCES RELIED ON

| Data Point | Author(s) | Year | Title Report No. Document No. Source (where different from company) GLP/ Officially recognised testing facilities <sup>2,3</sup> | Vertebrate study Y/N | Data protection claimed Y/N | Justification if data protection is claimed | Owner | Previously used <sup>1</sup><br>Y/N<br><br>If yes, for which data point? |
|------------|-----------|------|--|----------------------|-----------------------------|---|-------|--|
|------------|-----------|------|--|----------------------|-----------------------------|---|-------|--|

|  |  |  | Published or not                   |  |  |  |  |  |
|--|--|--|------------------------------------|--|--|--|--|--|
|  |  |  | No data submitted for this section |  |  |  |  |  |
|  |  |  |                                    |  |  |  |  |  |

<sup>1</sup> In order to facilitate the compilation of the final list of the tests and studies relied upon and the corresponding data protection, indicate whether the study was used in the previous DAR/RAR or, when the information is available, whether the study was already submitted in the framework of national authorisations.

<sup>2</sup> See Art.3 of Annex of Regulation No 283/2013 and 284/2013

<sup>3</sup> The RMS shall check that the GLP statement has been properly signed in the study report, that the study results are properly reported in accordance with GLP standards and following the relevant guidance by OECD on the review of the GLP status of non-clinical safety data (currently under development).