AUTHORISATION FOR A PLANT PROTECTION PRODUCT

PLANT PROTECTION PRODUCTS REGULATION (EC) No 1107/2009

Extent of Authorisation: Great Britain and Northern Ireland

Product name: Soar

Formulation: a suspension concentrate formulation containing 250 g / l azoxystrobin details of which are specified in the ‘Confidential Conditions’ section (Appendix 3) of this authorisation.

MAPP number: 19525

Authorisation holder: Agform Limited, Maidenstone Heath, Blundell Lane, Bursledon, Southampton, SO31 1AA. (Registered company number: 4046444)

This authorisation ends:

(a) 31 December 2025 except as set out in (b) and (c) below:

(b) 30 June 2026 for sale and distribution of existing stocks

(c) 30 June 2027 for the disposal, storage and use of existing stocks

This authorisation will be withdrawn or amended before the end dates above if any of the active substances contained in this product are withdrawn from the Approvals Register or list of approved active substances included in Regulation (EU) No 540/2011, or if a decision is taken to withdraw or amend this authorisation under Regulation (EC) No 1107/2009 on any other grounds.
This and the attached Appendices 1 to 3 are signed by the Health and Safety Executive for and on behalf of the Secretary of State, the Welsh Ministers, the Scottish Ministers and the Department of Agriculture, Environment and Rural Affairs in Northern Ireland.

Date of issue 16 February 2023.

EXPLANATORY NOTES

1. This product has been authorised in accordance with the Uniform Principles for product evaluation and agreed end points for the active substance.

2. This is Authorisation Number 0447 of 2023 and replaces Notice(s) of Authorisation Number(s) 1524 of 2020 which will begin a staged withdrawal from the end of the month of issue of this notice.

3. This authorisation, other than Appendix 3, will be published on HSE’s website.


5. In this notice Regulation (EC) No 1107/2009 means:
   In relation to Great Britain, Regulation (EC) No 1107/2009 as it has effect in Great Britain
   In relation to Northern Ireland, Regulation (EC) No 1107/2009 as it has effect by virtue of the Protocol on Ireland/Northern Ireland in the EU withdrawal agreement

6. In this notice Regulation (EU) No 540/2011 means:
   In relation to Northern Ireland, Regulation (EU) No 540/2011 as it has effect by virtue of the Protocol on Ireland/Northern Ireland in the EU withdrawal agreement

7. In this notice Regulation (EC) No 1272/2008 means:
   In relation to Great Britain, Regulation (EC) No 1272/2008 as it has effect in Great Britain
   In relation to Northern Ireland, Regulation (EC) No 1272/2008 as it has effect by virtue of the Protocol on Ireland/Northern Ireland in the EU withdrawal agreement
APPENDIX 1: CONDITIONS OF AUTHORISATION

Failure to comply with the following conditions will result in the withdrawal or amendment of the authorisation under Regulation (EC) No 1107/2009 and may result in other enforcement action, including prosecution.

Placing on the market:

Packaging: The authorisation holder must only place this product on the market in the following containers:

i) 1 to 20 litre high density polyethylene container.

ii) 1 to 20 litre polyethylene terephthalate container.

Label: The authorisation holder must only sell and supply the product with the agreed label (Co. ref.: SOAR Label - 04D (002)), which is the label submitted with the application on 7 December 2021 (HSE ref.: W002036186), label amendments as specified in Annex A to HSE’s letter dated 16 February 2023 and labelling requirements according to the Regulation on Classification, Labelling and Packaging of Substances and Mixtures (EC) No. 1272/2008 (see Appendix 3 of this authorisation).

Classification:

Labels must comply with the Regulation (EC) No 1272/2008. Based on the information held by HSE at the time of issue of this authorisation the classification requirements include:

<table>
<thead>
<tr>
<th>Hazard Class and Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute toxicity hazard category 4</td>
<td></td>
</tr>
<tr>
<td>Hazardous to the aquatic environment - Acute 1</td>
<td></td>
</tr>
<tr>
<td>Hazardous to the aquatic environment - Chronic 1</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hazard Statement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>H332 Harmful if inhaled</td>
<td></td>
</tr>
<tr>
<td>H400 Very toxic to aquatic life</td>
<td></td>
</tr>
<tr>
<td>H410 Very toxic to aquatic life with long lasting effects</td>
<td></td>
</tr>
</tbody>
</table>

Authorisation holders are reminded that they are responsible for ensuring that the classification complies with the Regulation (EC) No 1272/2008 and other information affecting the classification of the product.

Use:

Field of use: **ONLY AS A FUNGICIDE**
User: Professional

<table>
<thead>
<tr>
<th>Crops/situations:</th>
<th>Maximum individual dose: (l product / ha)</th>
<th>Maximum total dose:</th>
<th>Maximum number of treatments: (per crop)</th>
<th>Latest time of application:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Barley (spring), barley (winter), durum wheat, rye (spring), rye (winter), spelt, triticale, wheat (spring), wheat (winter)</td>
<td>0.5</td>
<td>-</td>
<td>2</td>
<td>35 days before harvest (BBCH 60)</td>
</tr>
</tbody>
</table>

Environmental protection:

(1) Crops/situations with 5m buffer zone:

Since there is a risk to aquatic life from use, users not applying the statutory buffer zone must either themselves carry out or ensure that someone else has carried out a Local Environment Risk Assessment for Pesticides (LERAP) on their behalf before each spraying operation from a horizontal boom sprayer. Users must not allow direct spray from horizontal boom sprayers to fall within 5m of the top of the bank of any static or flowing waterbody or within 1m of a ditch which is dry at the time of application (these distances to be measured as set out in the guidance documents available from HSE Chemical Regulation Division’s website and any amendments that are made to it) unless:

(a) The LERAP indicates that a narrower buffer zone will be sufficient; and

(b) Any measures indicated by the LERAP as justifying the narrower buffer zone are complied with in full and in accordance with any conditions applicable to them.

Spray must be aimed away from water.

Spray from hand-held sprayers must not be allowed to fall within 1m of the top of the bank of a static or flowing waterbody. Spray must be aimed away from water.

(2) The results of the LERAP must be recorded in written form and must be available for a period of...
three years for inspection to any person entitled to exercise enforcement powers under or in connection with the Plant Protection Products Regulations 2011 or the Plant Protection Products (Sustainable Use) Regulations 2012. (An electronic record will satisfy the requirement for a written record, providing it is similarly available for inspection and can be copied).

(3) Detailed guidance on LERAPs and how to conduct a LERAP are contained in the guidance documents available from HSE Chemicals Regulation Division’s website. All LERAPs must be carried out in accordance with this Guidance and any amendments that are made to it.

Other specific restrictions:

To reduce the risk of resistance developing in target diseases the total number of applications of product containing QoI fungicides made to any cereal crop must not exceed two.
APPENDIX 2: GENERAL CONDITIONS OF AUTHORISATION

Failure to comply with the following conditions will result in the withdrawal or amendment of the authorisation under Regulation (EC) No 1107/2009 and may result in other enforcement action, including prosecution.

Label:
The authorisation holder must follow the detailed requirements of all relevant Parts of ‘The Labelling Handbook’ available on HSE’s website at www.hse.gov.uk/pesticides. The authorisation holder must not make any amendments to the authorised label that are not consistent with the existing authorised use(s) and authorised label text. Classification is based on the information held by HSE.

Authorisation holders shall classify the product or update the label according to Regulation (EC) No 1272/2008 without undue delay following

- any change to the classification and labelling requirements required in the relevant legislation or;
- other information becoming available affecting the classification and labelling of the product.

In these cases HSE must be informed of the changes (refer to HSE’s website at www.hse.gov.uk/pesticides for details).

Packaging:
The authorisation holder must design and construct the packaging and fastenings to make sure they are strong and solid throughout so they will not come apart and will safely withstand normal handling; they can be repeatedly refastened and the contents cannot accidentally escape; the contents cannot attack either the packaging or the fastenings or form harmful or dangerous compounds with them and they are unlikely to attract children or arouse their curiosity.

Mixing:
The authorisation holder must ensure that any plant protection or adjuvant product(s) recommended for mixing with this product are authorised or have an official List Entry for use in the UK (refer to HSE’s website). The authorisation holder must ensure that any tank mix recommendations comprising this product and any other plant protection or adjuvant product(s) are supported by data as referenced in the compatibility assurance statement for this product.

Adverse effects:
The authorisation holder must immediately notify the Secretary of State, the Welsh Ministers, the Scottish Ministers and the Department of Agriculture, Environment and Rural Affairs in Northern Ireland (care of the Health and Safety Executive), if they have any new information on the potentially adverse effects of the authorised product, or of residues of an active substance in that product when used in accordance with the conditions of this authorisation. For those products authorised under Regulation (EC) No 1107/2009 as it has effect by virtue of the Protocol on Ireland / Northern Ireland in the EU withdrawal agreement, authorisation holders must also tell the relevant competent authorities of the EC Member States (a list of which is available from the Health and Safety Executive) and the EC Commission. Failure to comply with this requirement is an offence.
Provision of information:
The authorisation holder must comply with all requests for information required by, or on behalf of, the Secretary of State, the Welsh Ministers, the Scottish Ministers or the Department of Agriculture, Environment and Rural Affairs in Northern Ireland in accordance with Regulation (EC) No 1107/2009.
APPENDIX 3: CONFIDENTIAL CONDITIONS OF AUTHORISATION

This section is confidential to the authorisation / permit holder. Therefore, it is not available to those other than the data owner and has been removed.