

AUTHORISATION NUMBER: IE/BPA 70796

EUROPEAN COMMUNITIES (AUTHORISATION, PLACING ON THE MARKET, USE AND CONTROL OF BIOCIDAL PRODUCTS) REGULATIONS

CERTIFICATE OF AUTHORISATION

The Competent Authority for Biocides in Ireland, pursuant to the provisions of Regulation (EU) No 528/2012 of the European Parliament and of the Council concerning the making available on the market and use of biocidal products, as amended by Regulation (EU) No 334/2014, and European Union (Biocidal Products) Regulations, 2013, (S.I. 427 of 2013), grants authorisation to make available on the market in Ireland, the biocidal product:

Biocidal Product Family Name:	ARVO XY PE	
Name and address of the	Name	STOCKMEIER FRANCE SAS
authorisation holder	Address	3 rue de la Buhotière 35091 Rennes France
Authorisation number	IE/BPA 7	70796
Authorisation type	Mutual recognition in sequence (NA-MRS)	
Date of the authorisation	26 th May 2021	
Expiry date of the authorisation	05 th June 2029	

subject to the conditions detailed in the Annexes to this certificate.

Authorisation granted on behalf of the Competent Authority for Biocides in Ireland by

Pesticide Control Division (PCD)

Official Stamp:



Version: 1.2

$\underline{\textbf{ANNEX I}}$ Product Family Summary and Conditions of Authorisation

Biocidal Product Name	ARVO XY PE IE/BPA 70796				
Additional Trade names					
(with suffixes to the	02 SAFE 7.4	IE/BPA 70796-001			
Authorisation number)	SPRAY OXY PAE	IE/BPA 70796-002			
R4BP asset number	IE-0026285-0000				
Marketing Company, Address	To be confirmed				

Active Substance(s) (% w/w):	Hydrogen peroxide (7.44 % w/w)			
Product-Type:	PT 02 – Disinfectants and algaecides not intended for direct application to humans or animals (Disinfectants) PT 04 – Food and feed area (Disinfectants)			
Product Composition:	See Confidential PAR on R4BP3			
Substance(s) of Concern:	N/A			
Comparative Assessment	NO			
Formulation Type:	AL – Any other liquid			
Area of Use:	Indoor use			
Statement of use:	ARVO XY PE is a disinfectant containing hydrogen peroxide (7.44 % w/w) to be used by professional users for surface disinfection (sanitary, surfaces, equipment, and furniture). Claimed uses are: -for PT2, the disinfection in the medical sector, and institutional buildings (hotels, sport halls), -for PT4, the disinfection in the food industry, kitchens and canteens. The product is applied either with a trigger spray or via airbone diffusion (fogging).			
User Category:	Professional			
Special labelling provisions for Ireland:	In addition to the details recorded on the SPC, the following details shall be recorded on the product label(s). Use Biocides Safely and Sustainably It is illegal to use this product for uses or in a manner other than that prescribed on this label. Poison Information: For information or to report a poisoning incident contact The National Poisons Information Centre, Beaumont Hospital, Dublin (01-8092166), retain the label for reference.			

This authorisation may be subject to review in accordance with Regulation (EU) No 528/2012, as amended by Regulation (EU) No 334/2014, or the European Union (Biocidal Products) Regulations, 2013, (S.I. 427 of 2013). The outcome of such a review may lead to amendments to or the revocation of this authorisation.

The following conditions and restrictions apply:

- 1. Product may **not** be made available on the market or used in the Republic of Ireland unless it complies with the Annexes of this authorisation.
- 2. The requirements and conditions, specified in the Annexes, of this authorisation may **not** be altered without prior approval of modifications by the Irish Competent Authority for Biocides in Ireland. Where any amendments are made to the original authorisation in another Member State, the Irish Competent Authority for Biocides in Ireland must be informed by the Authorisation Holder.
- 3. The holder of this certificate for authorisation must inform or provide the Irish Competent Authority for Biocides with any new or requested information/data, respectively, that shows this biocidal product and/or any of its active substances cause or may cause an adverse effect on human or animal health, ground water or the environment.
- 4. All product made available on the market in Ireland must comply with the classification, labelling and packaging requirements established in: Article 69 of Regulation (EU) No 528/2012; the Chemicals Act 2008 (as amended) transposing Regulation (EC) No 1272/2008; and the classification, labelling and Safety Data Sheet information detailed in the Annex II to this certificate.
- 5. All biocidal products advertised must comply with Article 72 of Regulation (EU) No 528/2012.
- 6. A printed copy of the Irish label in accordance with the Annexes of this authorisation must be submitted to the Irish Competent Authority for Biocides prior to any product being made available on the market in Ireland. All product labels must carry the authorisation number of the form: IE/BPA 70796.
- 7. Safety Data Sheets (SDS) for the biocidal product(s) shall be prepared and made available in accordance with Article 70 of the Biocidal Products Regulation 528/2012 (as amended). Relevant sections of the SDS must be updated post-authorisation in accordance with Annex II of the authorisation certificate. In particular, Section 15 of the SDS should be updated to contain the authorisation number IE/BPA 70796. The SDS must be submitted to the Irish Competent Authority for Biocides and the National Poisons Information Centre of Ireland http://www.poisons.ie/manufacturers.asp before the product is made available on the market for sale or use.
- 8. On an annual basis, details of the quantities of this product (by pack size) manufactured in Ireland, imported into Ireland and/or exported from Ireland must be submitted to the Irish Competent Authority for Biocides by 31 January of the following year.
- 9. Fees are payable for the maintenance of the product on the Register of Biocidal Products and shall be paid by the 31st December of the following year and each year thereafter.

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(b) Amendments to Authorisation

The following amendments apply to the conditions of authorisation for the biocidal product family:

Issue	Re-issue	Version	Modifications applied ²
26/05/2021	-	1.0	Original certificate
-	19/01/2022	1.1	Additional tradename
-	03/07/2023	1.2	Amendments to product formulator NA-ADC BC-GC083028-53

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ANNEX II

Summary of Product Characteristics (SPC) for a biocidal product family

The following conditions, outlined in the summary of product characteristics (SPC), apply to the authorisation for the biocidal product family as provided for in Article 22 of Regulation (EU) No 528/2012 as amended. The authorised biocidal product family SPC file is referenced below:

Issue	Re-issue	Version	File Name
26/05/2021	-	1.0	spc_ARVO XY PE_IE_cn_202003191250
-	19/01/2022	1.1	spc_ARVO_XY_PE_IE_en_202108051424
-	03/07/2023	1.2	spc_ARVO_XY_PE_IE_en_202301021630