

# Guide to Product Literature Standard (PLS) for Veterinary Medicinal Products

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## 1 INTRODUCTION

The information in this guide aims to assist applicants with the creation of mock-ups.

Mock-ups should not deviate from the agreed QRD text<sup>1</sup>. Mock-ups should always be submitted for approval via a G.I.15z variation either immediately after the close of a new marketing authorisation procedure (where the product is to be marketed immediately) or a later stage (before marketing). Mock-ups are not routinely required in support of variation applications which result in changes to the product information, and this applies to G.I.18 variations to align the product information with version 9.0 of the QRD templates.

This document includes a list of general labelling requirements, which will assist applicants with the layout and design of their mock-ups and an additional list of national Health Products Regulatory Authority (HPRA) information to be considered for inclusion. This document should be read alongside the HPRA '[FAQs on Processing the Labelling and Package Leaflet for Veterinary Medicinal Products](#)'.

Applicants are advised that the HPRA does not assess shipping packs, datasheets (including Material Safety Data Sheets (MSDSs)), packaging for wholesalers that do not include any labels, display packaging or promotional material.

## 2 MULTI-COUNTRY PACKS

Multi-country packs are medicinal products that are labelled to allow their placing on the market in several Member States with the same packaging.

### 2.1 Joint-labels

Joint-labelling can be achieved between IE and GB, IE and UK (NI), or all three – IE, GB, and UK (NI). For further details, see the HPRA '[Guide to Joint labelling for Veterinary Medicinal Products for use in Ireland and the UK](#)'-and the Veterinary Medicines Directorate's (VMD) '[Joint labelling for veterinary medicines for use in the UK and Ireland](#)' (<https://www.gov.uk/guidance>).

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<sup>1</sup> An exception applies where a shared label/package leaflet with GB is sought and where an interim position has been agreed by the Health Products Regulatory Authority (HPRA) and Veterinary Medicines Directorate (VMD) (cf. [Joint HPRA/VMD Guide to Acceptable Texts for Joint Labelling for Veterinary Medicinal Products for use in Ireland and the UK](#))

## 2.2 Multilingual labels

'Multilingual packaging' refers to the use of two or more languages for at least one component of the packaging material for a medicinal product. The HPRA requires that all labels and packaging be in English, but inclusion of additional languages is permissible if the legibility of the English text is not compromised and the information given is identical in all languages. Multilingual labels require a clear separation between the different languages and all the information provided in each language should be kept together.

## 3 GENERAL LABELLING REQUIREMENTS

### Font type, style, and size

The font size should be as large as possible and should be measured against Times New Roman.

Type of Packaging	Recommended Font Size	Minimum Font Size
Small Immediate Pack Sizes	7 pt	4.75 pt*
Immediate Packaging	7 pt	6 pt
Outer Packaging	7 pt	7 pt
Package Leaflet	9 pt	8 pt

\*Only in exceptional circumstances and on a case-by-case basis. Requires approval via a G.I.15z variation.

### Headings

Use of QRD headings on the immediate and outer packaging is not obligatory, but you must include headings that clearly convey meaning (such as 'withdrawal period').

### Pictograms

Pictograms used should be as per those of the approved QRD guidance on the use of approved pictograms on the packaging of veterinary medicinal products authorised via the centralised (CP), mutual recognition (MR) and decentralised procedures (DCP).

## 4 NATIONAL SPECIFIC INFORMATION

### **Marketing Authorisation (MA) Number**

The marketing authorisation (MA) number is required on the package leaflet and outer package in the format: IE - VPA xxxxx/xxx/xxx.

### **Distribution category**

The distribution category should only be included on the package leaflet as follows with the abbreviation in bold (not in a box) followed by the corresponding explanatory text in brackets:

**POM** (Prescription Only)

**POM(E)** (Prescription Only Exempt)

**CAM** (Companion Animal Medicine)

**LR** (Licensed Retailer)

### **Local representative/ distributor**

The listing of a local representative of the marketing authorisation holder (MAH) or of an entity that functions to physically distribute the product ('a distributor') may be introduced on the package leaflet by the MAH but is not a national requirement.

However, if a local representative is responsible for receiving reports of suspected adverse reactions, then the local representative and their contact details (including telephone number) must be included on the package leaflet, clearly identifying them as performing that task.

To introduce and subsequently amend local representative/ distributor details on the package leaflet, submission of a C.10.a Variation Not Requiring Assessment 'Changes to the labelling or the package leaflet which shall not be connected with the SPC - administrative information concerning the holder's representative' is required. Mock-ups of the labelling and package leaflet are not routinely required.

### **Dedicated dispensing containers**

Dispensing materials intended to be supplied by an MAH to facilitate the dispensing of their product by a registered veterinary practitioner, pharmacist, the holder of veterinary medicinal product retailer's licence (retail responsible person), or a person entered in the 'Companion Animal Medicine Retailers' Register' (registered person), should not include any information other than that on the approved label/package leaflet.

Mock-ups of dispensing materials are not reviewed by the HPRA.

### **QR codes**

Under the provisions of Article 13 of Regulation 2019/6, a QR code may be added, provided the legibility is not affected, and accesses information intended for internal manufacturing,

processing, stock control or anti-counterfeit measures that cannot be accessed by the public or public information, which conforms to the product information approved by the HPRA.

Links to website addresses and/or company websites are considered promotional and cannot be included.

## **APPENDIX I CHECKLIST FOR MOCK-UP PREPARATION**

When preparing mock-ups, please consult the follow checklist:

- Mock-ups contain only the text agreed during the procedure and any additional agreed national-specific information.
- The name of the VMP appears as an integrated unit in the user's field of vision.
- Product specific VPA number is included on the package leaflet and the outer package.
- Font sizes are in line with the requirements as detailed in section 3 above.
- The method of sale and supply is included on the package leaflet denoted by the appropriate abbreviation, as detailed in section 4 above.
- The date the package leaflet was last revised is included (as detailed in the QRD guidance and explanatory texts).
- No company websites have been included.
- Note that the batch number and expiry date will need to be overprinted on the packaging in the format specified in QRD Veterinary Product Information Annotated Template v9.0.
- Only pictograms agreed during the procedure and as listed on the agreed QRD texts are included.
- For joint-labels, all national-specific information should be identified as "IE only" or "UK only".