

United Kingdom Public Assessment Reports (UKPARs)

Nelio 20 mg Tablet for Dogs

This product has been authorised via the Decentralised procedure or via the Mutual Recognition Procedure (MRP). It is expected that a Public Assessment Report (PuAR) for this product will be available on the Heads of Medicines Agencies (HMA(v)) website by approximately 3 months from its date of authorisation.

HMA(v) represents the veterinary regulatory agencies of the EU.

There is a section in the HMA(v) website called 'VMRI': Veterinary Mutual Recognition Index. This contains a list of the veterinary medicinal products approved via the Decentralised Procedure or via the MRP.

To go to the PuAR for this veterinary product, click on the link to the HMA(v) website then click on VMRI. Whilst there are various ways to search for a product, clicking on 'ABC' and then finding the product alphabetically is probably the easiest.

In this list, the first column gives the Reference Member State, i.e. the Member State who first evaluated the product. If it says UK, the assessment report was written by the Veterinary Medicines Directorate. If it gives another Member State, the assessment report was written by the regulatory agency of that Member State.

As the EU legislation requiring a PAR came into force on 30 October 2005, it only applies to products authorised on or after 30 October 2005 (i.e. granted a Marketing Authorisation on or after 30 October 2005). As the Scientific Discussion (the largest part of the PuAR) has to be specially written for the PuAR, this document may not be available for products authorised before 30 October 2005.

[Link to HMA\(v\) Website](#)