

## Cetrotide

Procedural steps taken and scientific information after the authorisation\*

\*Due to the Agency`s update of its procedure management systems, an additional document, reflecting the historical lifecycle may be available in the 'Assessment history' section. For the complete product lifecycle procedures, you may need to also refer to EPAR - Procedural steps taken and scientific information after authorisation (archive).

Application number	Scope	Notification	Product Information affected <sup>3</sup>	Summary
Article 61(3) /	- Notification acc. Article 61(3) - Accepted	24/11/2025	Labelling and	

<sup>&</sup>lt;sup>1</sup> Notifications are issued for type I variations and Article 61(3) notifications (unless part of a group including a type II variation or extension application or a worksharing application). Opinions are issued for all other procedures.



<sup>&</sup>lt;sup>2</sup> A Commission decision (CD) is issued for procedures that affect the terms of the marketing authorisation (e.g. summary of product characteristics, annex II, labelling, package leaflet). The CD is issued within two months of the opinion for variations falling under the scope of Article 23.1a(a) of Regulation (EU) No. 712/2012, or within one year for other procedures.

<sup>&</sup>lt;sup>3</sup> SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).

EMA/N/0000308409	Update of the Labelling to align instruction for Use (IFU) concerning the injection angle and site. The update provides clearer guidance on the recommended injection site in the lower abdominal area and on maintaining an appropriate distance from the navel, as well as specifying a preferred injection angle to support correct subcutaneous administration. The MAH is also taking the opportunity to introduce some minor editorial changes to the labelling and package leaflet.			PL	
Variation type IB / EMA/VR/0000307159	This was an application for a group of variations.  B.I.a.1 Change in the manufacturer of a starting material/reagent/intermediate used in the manufacturing process of the active substance or change in the manufacturer (including where relevant quality control testing sites) of the active substance, where no Ph. Eur. Certificate of Suitability is part of the approved dossier - B.I.a.1.z Other variation - Accepted  B.I.d.1.a Re-test period/storage period - B.I.d.1.a.4 Extension or introduction of a retest period/storage period supported by real time data - Accepted	10/11/2025	N/A		