



Príomhoifigeach Cliniciúil
Oifig an Phríomhoifigigh Cliniciúil

Ospidéal Dr Steevens, Lána Steevens
Baile Átha Cliath 8 , D08 W2A8

Chief Clinical Officer
Office of the Chief Clinical Officer

Dr Steevens Hospital, Steevens Lane
Dublin 8, D08 W2A8

www.hse.ie
[@hselive](https://twitter.com/hselive)

t 01 635 2000
e cco@hse.ie

BY EMAIL ONLY

Deputy Jennifer Whitmore
Dáil Éireann
Leinster House
Kildare Street
Dublin 2

24th March 2023

PQ10869/23-Deputy Jennifer Whitmore- To ask the Minister for Health if he will outline what the review of the HSE policy on provision of Cariban will entail; when it will be completed; and if he will make a statement on the matter.

Dear Deputy Whitmore,

Thank you for your representation.

An exceptional arrangement was put in place effective 1st January 2023 to support the reimbursement of the Exempt Medicinal Product (EMP), Cariban® (doxylamine/pyridoxine), under the Community Drug Schemes for the treatment of nausea and vomiting of pregnancy (NVP) against the dedicated funding of €1.3m as part of Budget 2023. Only licensed products are added to the formal GMS Reimbursement List in line with the Health (Pricing and Supply of Medical Goods) Act 2013

The National Centre for Pharmacoeconomics (NCPE) completed a rapid review for Xonvea® on 15th August 2019. The NCPE noted that doxylamine/pyridoxine is associated with a greater benefit than placebo but the size of the effect is small and the quality of the evidence is low. The NCPE outlined that there is no evidence presented for an increased clinical benefit for doxylamine/pyridoxine compared to other therapeutic options such as promethazine or cyclizine. The NCPE recommended that Xonvea® not be considered for reimbursement at the submitted price. To date the MA Holder of Xonvea® has not progressed the pricing and reimbursement application further with the HSE. The MA Holders for Exeltis® and Navalem® have not submitted pricing and reimbursement applications to the HSE to date. The HSE therefore encourages clinicians, along with the Institute of Obstetricians and Gynaecologists and the National Clinical Programme for Obstetrics and Gynaecology, and other healthcare professionals to encourage the MA Holders of the licensed medicinal products (Xonvea®, Exeltis® and Navalem®) to progress with the formal pricing and reimbursement process in Ireland.

Following a review of the available evidence and in light of the exceptional circumstances of unmet clinical need for this patient cohort, the HSE considered the potential for a time-limited, quantity-dependent reimbursement arrangement for women with NVP, to access Cariban® ULM, when prescribed by a consultant obstetrician and in line with clear clinical criteria, as set



out by the HSE. This recommendation was made with the understanding that the HSE has no agreement in place with the manufacturers of Exempt Medicinal Products (EMPs) (i.e. unlicensed medicines), in terms of pricing and supply under the Health (Pricing and Supply of Medical Goods) Act 2013. Manufacturers can cease supply and increase the price of these medicines without consultation and notification to the HSE. Due to the uncertainty around the cost of EMPs, the HSE retains discretion as to the continued reimbursement support for any EMP. If a licensed product was subsequently approved for reimbursement this arrangement could then be reviewed/revised.

Since 1st January 2023, applications for this medicine are being processed by PCRs and are being turned around in a short timeframe.

Yours sincerely

A handwritten signature in black ink, appearing to read 'Sharon Hayden', written over a thin horizontal line.

Sharon Hayden
General Manager
Office of the Chief Clinical Officer