

Feidhmeannacht na Seirbhíse Sláinte, Seirbhís Aisíocaíochta Cúraim Phríomhúil Bealach amach 5, M50, An Bóthar Thuaidh, Fionnghlas Baile Átha Cliath 11, D11 XKF3

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Seán Canney, T.D. Dáil Éireann, Leinster House, Kildare Street, Dublin 2.

30th January 2025

PQ: 1569/25

To ask the Minister for Health when melatonin will be made available under the medical card scheme; and if he will make a statement on the matter. -Seán Canney

Dear Deputy Canney,

The Health Service Executive has been requested to reply directly to you in the context of the above Parliamentary Question (Reference 1569/25), which you submitted to the Minister for Health for response.

Melatonin has never been available through the formal Reimbursement List for reimbursement under Community Drug Schemes. Under the Health (Pricing and Supply of Medical Goods) Act 2013, products that are licensed will undergo a pricing and reimbursement approval process by the National Centre for Pharmacoeconomics (NCPE) on receipt of an application by the HSE from the Marketing Authorisation Holder.

Pharmaceutical companies are required to submit and pricing and reimbursement application should they wish for their product to be added to the Reimbursement List.

Melatonin (Circadin®) underwent a review in 2008 and reimbursement was not recommended by the NCPE as there is currently insufficient evidence to support the reimbursement of this product under the Community Drugs Schemes (available at http://www.ncpe.ie/drugs/melatonin-circadin/).

The HSE Corporate Pharmaceutical Unit received an application for pricing and reimbursement of Melatonin (Slenyto® Prolonged-Release Tablets) in July 2019 from Flynn Pharma for the treatment of insomnia in children and adolescents aged 2-18 with Autism Spectrum Disorder (ASD) and / or Smith-Magenis syndrome, where sleep hygiene measures have been insufficient. The first step in the process is the submission of a rapid review dossier (a clinical and economic dossier) to the National Centre for Phamacoeconomics (NCPE) for assessment. The rapid review was commissioned on 5th July 2022 and

completed by the NCPE on 9th August 2022. The rapid review outcome recommends that a full Health Technology Assessment (HTA) is recommended to assess the clinical effectiveness and cost effectiveness of prolonged release melatonin (Slenyto®) compared to the current standard of care. However, to date the company have not made a HTA submission to the NCPE for assessment.

The position remains that a submission of a completed HTA dossier to the NCPE is required to progress this application, as per the formal processes governing the pricing and reimbursement of medicines.

The HSE Corporate Pharmaceutical Unit received an application for pricing and reimbursement of Melatonin (Voquily® 1mg/ml Oral solution) in September 2024 from Caragen Ltd for sleep onset insomnia in children and adolescents aged 6-17 years with attention-deficit hyperactivity disorder (ADHD) where sleep hygiene measures have been inadequate. The first step in the process is the submission of a rapid review dossier (a clinical and economic dossier) to the National Centre for Phamacoeconomics (NCPE) for assessment. However, to date the company have not made a Rapid Review submission to the NCPE for assessment.

The position remains that a submission of a completed Rapid Review dossier to the NCPE is required to progress this application, as per the formal processes governing the pricing and reimbursement of medicines.

Yours sincerely,

Suzanne Doyle

**Primary Care Reimbursement Service** 

The Health Service Executive operates the General Medical Services Scheme, which includes Medical Cards and GP Visit Cards, under the Health Act 1970, as amended. It has established a dedicated contact service for members of the Oireachtas specifically for queries relating to the status of Medical Cards and GP Visit Cards applications, which the Deputy / Senator may wish to use for an earlier response. Tel: 01-8647180 / email: Oireachtas.pcrs@hse.ie