

# Rare melanocortin-4 receptor pathway diseases: Clinical features and genetic confirmation

Rare melanocortin-4 receptor (MC4R) pathway diseases can be caused by genetic variants within the pathway, which impair signalling that controls hunger.<sup>1</sup>

**Hyperphagia** (pathological, insatiable hunger) and **early-onset, severe obesity** are clinical features of a rare MC4R pathway disease.<sup>1</sup> If you see these features in your patients, **they may be living with a rare MC4R pathway disease.**<sup>1</sup>



## Hyperphagia<sup>2</sup>

Also known as an abnormally strong sensation of hunger or desire to eat.

Characteristics and behaviours include:



Heightened and prolonged hunger



Longer time to reach satiety



Shorter duration of satiety



Severe preoccupation with food (hyperphagic drive)



Food-seeking behaviours (sneaking or stealing food)

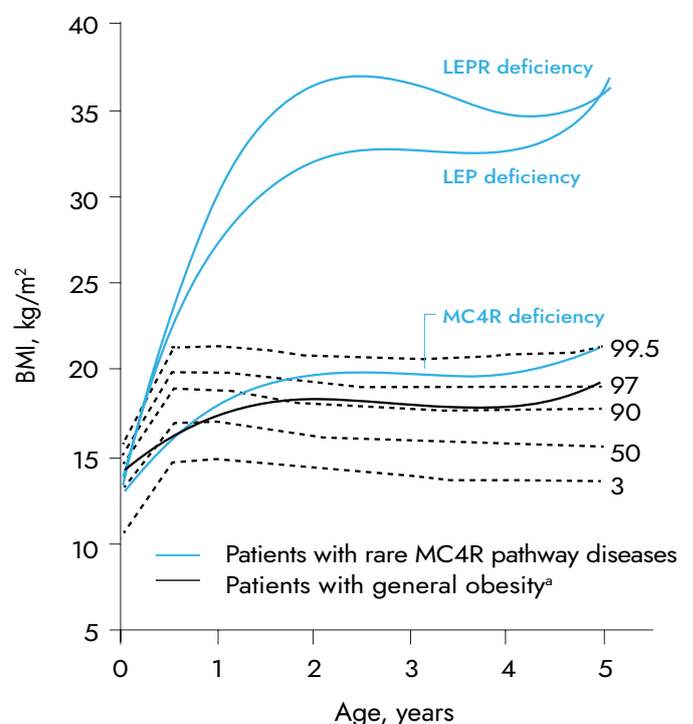


Distress and functional impairment if denied food



## Early-onset, severe obesity<sup>3</sup>

Defined as having a BMI  $\geq 120\%$  of the 95<sup>th</sup> percentile and onset before the age of 5.<sup>4</sup>



a) Patients with general obesity have a BMI  $>30$  kg/m<sup>2</sup> by age 14 to 16 years and do not have a variant in *LEP*, *LEPR*, or *MC4R*.

Reprinted with permissions from Springer Nature from Kohlsdorf K, et al. *Int J Obes (Lond)*. 2018;42(9):1602–1609.

Proactive identification of clinical features, and appropriate referral for genetic confirmation using correct gene panels can help move children living with a rare MC4R pathway disease onto their most appropriate care path.<sup>5</sup>



Access to appropriate tools means genetic variants that cause rare MC4R pathway diseases can be diagnosed early.<sup>5</sup>

## The diagnosis pathway<sup>5</sup>



Patient visits HCP to discuss symptoms



HCP identifies clinical features



HCP refers patient for genetic testing



Genetic testing results confirm if the patient has a rare MC4R pathway disease



Through a correct referral, children with genetic variants that cause rare MC4R pathway diseases can be screened, and appropriately cared for



Current genetic screening allows many more rare diseases to be identified – rare MC4R pathway diseases can now be part of this



If you need more information on genetic confirmation or locating expert centres in your country, please visit our website [Path4hcps.com](https://Path4hcps.com), accessible via the QR code:



## References:

1. Loos, RJF and Yeo, GSH. *Nat Rev Genet*. 2022;23:120–133.
2. Heymsfield SB, et al. *Obesity (Silver Spring)*. 2014;22(suppl 1):S1–S17
3. Kohlsdorf K, et al. *Int J Obes (Lond)*. 2018;42(9):1602–1609
4. Hampl SE, et al. *Pediatrics*. 2023;151(2):e2022060640
5. Styne DM, et al. *J Clin Endocrinol Metab*. 2017;102(3):709–757