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No. S 1078

HEALTH PRODUCTS ACT (CHAPTER 122D)

HEALTH PRODUCTS (CLINICAL RESEARCH MATERIALS) (AMENDMENT NO. 3) REGULATIONS 2021

In exercise of the powers conferred by section 72(1) of the Health Products Act, the Health Sciences Authority, with the approval of the Minister for Health, makes the following Regulations:

Citation and commencement

1. These Regulations are the Health Products (Clinical Research Materials) (Amendment No. 3) Regulations 2021 and come into operation on 3 January 2022.

Amendment of regulation 2

2. Regulation 2(1) of the Health Products (Clinical Research Materials) Regulations 2016 (G.N. No. S 332/2016) is amended by deleting the definition of “licensed healthcare institution” and substituting the following definition:

““licensed healthcare institution” means —

- (a) any premises or conveyance specified in a licence granted under the Healthcare Services Act 2020 for the provision of any licensable healthcare service; or
- (b) a healthcare institution that is licensed under the Private Hospitals and Medical Clinics Act 1980;”.

[G.N. Nos. S 94/2019; S 108/2021; S 730/2021]

Made on 30 December 2021.

BENJAMIN ONG
Chairman,
Health Sciences Authority,
Singapore.

[401:04/01-000; AG/LEGIS/SL/122D/2020/10 Vol. 1]

(To be presented to Parliament under section 72(5) of the Health Products Act).