

To: Dienstpostbus GMT-secretariaat[5.1.2e]@minvws.nl
Cc: [5.1.2e] [5.1.2e]@minvws.nl
From: [5.1.2e]
Sent: Wed 12/2/2020 9:43:34 AM
Subject: graag webex inplannen FW: new survey: new SC to start discussions on monoclonal antibodies - please participate by Thu 3 Dec
Received: Wed 12/2/2020 9:43:34 AM

Ha dames

Zouden jullie voor morgen een webex van 15 min willen inplannen met

[5.1.2e] (VWS)

[5.1.2e] (RIVM)

[5.1.2e] (RIVM)

Mij zelf

Gaat over: JP nieuwe covid behandelingen

Dank!

[5.1.2e]

Van: [5.1.2e]

Verzonden: woensdag 2 december 2020 10:41

Aan: [5.1.2e] < [5.1.2e]@rivm.nl>; [5.1.2e] < [5.1.2e]@rivm.nl>

CC: [5.1.2e] < [5.1.2e]@minvws.nl>

Onderwerp: FW: new survey: new SC to start discussions on monoclonal antibodies - please participate by Thu 3 Dec

Ha [5.1.2e] [5.1.2e]

Hierbij het verzoek vanuit de commissie waar we zojuist even over spraken.

We zullen een kort overleg inplannen voor morgen (met jou, [5.1.2e] [5.1.2e] en ik)

Groet [5.1.2e]

Van: [5.1.2e] < [5.1.2e]@minvws.nl>

Verzonden: dinsdag 1 december 2020 12:39

Aan: [5.1.2e] < [5.1.2e]@minvws.nl>

Onderwerp: FW: new survey: new SC to start discussions on monoclonal antibodies - please participate by Thu 3 Dec

Hoi [5.1.2e]

Ik heb onderstaande gisteren uitgezet bij GMT, maar wellicht hoort dit ook bij jullie. Vanuit de JP steering committee. De vraag is of wij kunnen aangeven of er in NL behoefte is aan VIR-7831 en Otilimab (wanneer dit geautoriseerd wordt in de EU) en wanneer dit het geval is, of we een national focal point willen doorgeven die hier namens NL aanspreekpunt voor wordt.

Zou jij dit verzoek verder willen oppakken en mij kunnen laten weten wat we aan de JPA kunnen laten weten?

Groet, [5.1.2e]

Van: [5.1.2e]

Verzonden: maandag 30 november 2020 17:45

Aan: [5.1.2e] < [5.1.2e]@minvws.nl>

Onderwerp: FW: new survey: new SC to start discussions on monoclonal antibodies - please participate by Thu 3 Dec

Hoi [5.1.2e]

Zie onderstaand verzoek dat wij ontvingen van de Joint Procurement steering committee. Zou jij dit verzoek binnen GMT willen uitzetten?

Groet, 5.1.2e

Van: 5.1.2e @ec.europa.eu <5.1.2e@ec.europa.eu>

Verzonden: maandag 30 november 2020 17:39

Aan: 5.1.2e @ec.europa.eu

Onderwerp: new survey: new SC to start discussions on monoclonal antibodies - please participate by Thu 3 Dec

Dear members of the joint procurement agreement steering committee,

You may be aware that the US Food and Drug Administration has authorized the use of two therapeutics to treat COVID-19 patients that are monoclonal antibodies. These are: **Bamlanivimab** (Eli Lilly): authorized for COVID-19 patients who are 12 years of age and older weighing at least 40 kilograms, and who are at high risk for progressing to severe COVID-19 and/or hospitalization (<https://www.fda.gov/media/143602/download>) and the **"cocktail" of casirivimab and imdevimab** (Regeneron/Roche (in Europe)) to be administered for the treatment of mild to moderate COVID-19 in adults and pediatric patients (12 years of age or older weighing at least 40 kilograms and who are at high risk for progressing to severe COVID-19 (<https://www.fda.gov/media/143891/download>)). The **European Medicines Agency** plans to start a "rolling review" for both therapeutics soon.

In addition, the pharmaceutical company GSK is working on two possible therapeutics, namely **VIR-7831 and Otilimab**. The latter is in the clinical phase and the European Medicines Agency provided advice.

Therefore, we would like to invite you to nominate your representatives for a specific steering committee to start the discussions and know about your possible interest to purchase these therapeutics in case they get authorized in Europe.

We would like to thank you for your support in completing the requested data for your country. The needs for assessment and nominated member for the Specific Procurement Procedure Steering Committee should be completed by **THURSDAY 3 DECEMBER** through this survey:

5.1.2h

Kind regards,

JPA Team