Commissioner Kyriakides explained the epidemiological situation, the increases of COVID-19 cases across the EU and the concern with the new variant.

EFPIA, MfE and Business Europe raised concern in relation to the export authorisation scheme. EFPIA acknowledged the importance of transparency but considered that this particular measure impacts the supply chain and undermines the cooperation with international partners. This increases the risk of retaliation with the US and China as it is perceived as an export ban. Moreover, they referred to a hiccup in handling the requests. They referred to an incident occurring with an export authorisation in Germany. The associations called for reducing the scope of the Regulation to exempt samples and small quantities and the duration of the measure.

Commissioner clarified that we have not banned any export – apart from one case associations are aware of, where we followed the opinion of the national authority. This concerned one particular company with which we have particular challenges on the performance of our agreement. Commissioner called on industry to ensure that the production capacity is in place and can guarantee delivery of vaccines to EU citizens according to our agreements.

Commissioner confirmed that we are committed to implementing a smooth procedure and the measure is temporary. As far as samples are concerned, EC confirmed that they are covered and the customs cannot check if the shipments are for analytical tests, clinical trials or commercial use. We want to avoid applicants to circumvent the rules by sending small shipments to third countries.