

Beat: Health

UE AUTHORIZE MARKETING OF BIONTECH PFIZER VACCINE AGAINST COVID19 TO BE LAUNCHED

ON 27DEC SAID PDT URSULA VON DER LYEN

Paris, Washington DC, Brussels, 25.12.2020, 18:30 Time

USPA NEWS - The European Union countries shall be starting the collective on December 27, as announced by the President of European Commission , Ursula Von Der Lyen. « As we have promised, this vaccine will be available for all EU countries, at the same time, on the same conditions. » she declared. « We granted conditional marketing authorisation to the vaccine produced by BioNTech and Pfizer. The European Medicines Agency assessed this vaccine thoroughly. And it concluded that it is safe and effective against COVID-19. Based on this scientific assessment, we proceeded to authorise it for the European Union market. » "EMA will issue its opinion on the second vaccine, that of Moderna, on 6 January. "continued President of European Commission , Ursula Von Der Lyen. We publish her full statement.

STATEMENT BY PRESIDENT OF EUROPEAN COMMISSION URSULA VON DER LYEN OVER BIONTECH PFIZER VACCINE-----

« Good evening.Today we add an important chapter in our fight against COVID-19. We took the decision to make available for European citizens the first COVID-19 vaccine. We granted conditional marketing authorisation to the vaccine produced by BioNTech and Pfizer. The European Medicines Agency assessed this vaccine thoroughly. And it concluded that it is safe and effective against COVID-19. Based on this scientific assessment, we proceeded to authorise it for the European Union market. As we have promised, this vaccine will be available for all EU countries, at the same time, on the same conditions. The first batches of this vaccine will be shipped from Pfizer's manufacturing site here in Belgium within the next days.

I have always said, during this pandemic, that we are in this together. So vaccination can start at the same time, during the European Union vaccination days, on 27, 28 and 29 December. This is a very good way to end this difficult year, and to finally start turning the page on COVID-19."-----

This is our first vaccine. More will be approved soon, if they prove to be safe and effective.

EMA WILL ISSUE ITS OPINION ON THE 2ND VACCIN MODERNA ON 6 JANUARY-----"EMA will issue its opinion on the second vaccine, that of Moderna, on 6 January. And while we deliver vaccines for Europeans, we also help secure them for the rest of the world. Team Europe is the largest contributor to the COVAX Facility, with over EUR 800 million for procuring vaccines for the low- and middle-income countries. Finally, allow me to say how proud I am that the first COVID-19 vaccine available in Europe is a true product of European innovation. BioNTech has received more than EUR 9 million of EU research funding over the past decade, to develop these really ground-breaking technologies. And it secured a EUR 100 million loan from the European Investment Bank, that is backed by the European Union, in June. This helped expand its manufacturing capacities and supply the vaccine quickly worldwide. And this is a true European success story. Thank you so much. » President of European Commission, Ursula Von Der Lyen Stated. Source: European Commission,

Article online:

<https://www.uspa24.com/bericht-17956/ue-authorize-marketing-of-biontech-pfizer-vaccine-against-covid19-to-be-launched.html>

Editorial office and responsibility:

V.i.S.d.P. & Sect. 6 MDSIV (German Interstate Media Services Agreement): Jedi Foster P/O Rahma Sophia Rachdi

Exemption from liability:

The publisher shall assume no liability for the accuracy or completeness of the published report and is merely providing space for the submission of and access to third-party content. Liability for the content of a report lies solely with the author of such report. Jedi

Foster P/O Rahma Sophia Rachdi

Editorial program service of General News Agency:

UPA United Press Agency LTD

483 Green Lanes

UK, London N13NV 4BS

contact (at) unitedpressagency.com

Official Federal Reg. No. 7442619