







Update on our COVID-19 vaccine

December 22, 2020

This slide presentation includes forward-looking statements

Forward-Looking Statements

Various statements in this slide presentation concerning the future expectations of BioNTech, its plans and prospects, including the Company's views with respect to its efforts to combat COVID-19; the collaboration between BioNTech and Pfizer to develop a potential COVID-19 vaccine; its expectations regarding the potential characteristics of BNT162b2 in its Phase 2/3 trial and/or in commercial use based on data observations to date; the expected time point for additional readouts on efficacy data of BNT162b2 in our Phase 2/3 trial; the nature of the clinical data, which is subject to ongoing peer review, regulatory review and market interpretation; the timing for submission of data for, or receipt of, any marketing approval or Emergency Use Authorization; its contemplated shipping and storage plan, including its estimated product shelf life at various temperatures; and the ability of BioNTech to manufacture and supply the quantities of BNT162 to support clinical development and, if approved, market demand, including our production estimates for 2020 and 2021, are forward-looking statements reflecting the current beliefs and expectations of management made pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995, as amended. Words such as "expects," "plans," "potential," "target," "continue" and variations of these words or similar expressions are intended to identify forward-looking statements. Such statements are based on the current beliefs and assumptions of the management team of BioNTech and on the information currently available to the management team of BioNTech, and are subject to change. The Company will not necessarily inform you of such changes. These forward looking statements are subject to known and unknown risks, uncertainties, assumptions and other factors that could cause the Company's actual results, performance or achievements to be materially different than any future results, performance or achievements expressed or implied by the forward-looking statements. Actual results may differ materially from those indicated by these forward-looking statements as a result of various important factors, including but are not limited to: our ability to meet the pre-defined endpoints in clinical trials; competition to create a vaccine for COVID-19; the ability to produce comparable clinical or other results, including our stated rate of vaccine effectiveness and safety and tolerability profile observed to date, in the remainder of the trial or in larger, more diverse populations upon commercialization; the ability to effectively scale our productions capabilities; and other potential difficulties. Any forward-looking statements represent the Company's views only as of today and should not be relied upon as representing its views as of any subsequent date. The Company explicitly disclaims any obligation to update any forward-looking statements. The mRNA vaccine discussed in this slide presentation is an investigational product being developed by BioNTech and its collaborators and are not currently approved by the FDA, EMA or any other regulatory authority.



Safety information

Authorized use in the U.S.:

The Pfizer-BioNTech COVID-19 Vaccine is authorized for use under an Emergency Use Authorization (EUA) for active immunization to prevent coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) in individuals 16 years of age and older.

Important safety information from U.S. FDA emergency use authorization prescribing information:

- Do not administer Pfizer-BioNTech COVID-19 Vaccine to individuals with known history of a severe allergic reaction (e.g., anaphylaxis) to any component of the Pfizer-BioNTech COVID-19 Vaccine.
- Appropriate medical treatment used to manage immediate allergic reactions must be immediately available in the event an acute anaphylactic reaction occurs following administration of Pfizer-BioNTech COVID-19 Vaccine.
- Immunocompromised persons, including individuals receiving immunosuppressant therapy, may have a diminished immune response to the Pfizer-BioNTech COVID-19 Vaccine.
- The Pfizer-BioNTech COVID-19 Vaccine may not protect all vaccine recipients.
- In clinical studies, adverse reactions in participants 16 years of age and older included pain at the injection site (84.1%), fatigue (62.9%), headache (55.1%), muscle pain (38.3%), chills (31.9%), joint pain (23.6%), fever (14.2%), injection site swelling (10.5%), injection site redness (9.5%), nausea (1.1%), malaise (0.5%), and lymphadenopathy (0.3%).
- Severe allergic reactions have been reported following the Pfizer-BioNTech COVID-19 Vaccine during mass vaccination outside of clinical trials. Additional adverse reactions, some of which may be serious, may become apparent with more widespread use of the Pfizer-BioNTech COVID-19 Vaccine.
- Available data on Pfizer-BioNTech COVID-19 Vaccine administered to pregnant women are insufficient to inform vaccine-associated risks in pregnancy.
- Data are not available to assess the effects of Pfizer-BioNTech COVID-19 Vaccine on the breastfed infant or on milk production/excretion.
- There are no data available on the interchangeability of the Pfizer-BioNTech COVID-19 Vaccine with other COVID-19 vaccines to complete the vaccination series. Individuals who have received one dose of Pfizer-BioNTech COVID-19 Vaccine should receive a second dose of Pfizer-BioNTech COVID-19 Vaccine to complete the vaccination series.
- Vaccination providers must report Adverse Events in accordance with the Fact Sheet to VAERS at https://vaers.hhs.gov/reportevent.html or by calling 1-800-822-7967. The reports should include the words "Pfizer-BioNTech COVID-19 Vaccine EUA" in the description section of the report.
- Vaccination providers should review the Fact Sheet for mandatory requirements and Information to Provide to Vaccine Recipients/Caregivers and the Full EUA Prescribing Information for Requirements and Instructions for Reporting Adverse Events and Vaccine Administration Errors.





Conditional market authorization in 27 European States on 21 December for a COVID-19 vaccine









A concerted and large-scale global effort

Conditional Marketing Authorization in the EU and Switzerland¹

Approved Emergency Use Authorization / Temporary Use Approval

Vaccination with our COVID-19 vaccine already underway under Emergency Use Authorization/Temporary Use Approval

Rolling application for emergency use authorization to further countries underway.



Commitment to equitable supply of vaccine globally



EU's dose allocation is based on the member states' population as agreed between the European Commission and its member states

- All countries across the EU that have requested doses will receive them in the next 5 days
- Parallel vaccine shipments to multiple EU countries planned immediately following completion of final paperwork (manufacturing batch release)





Vaccine transportation: minimal changes to pre-existing cold chain



Bespoke vaccine freezer boxes; each freezer box can host between approx. 1000 to 5000 doses



Vaccine storage: administered like many other vaccines



Administration to vaccinees at room temperature

Injected intramuscular (arm); no additional equipment needed for administration at mass vaccination center



Once removed from the freezer, the unopened vaccine can be stored for up to 5 days at 2 °C to 8 °C

And up to 2 hours at temperatures up to 30 °C, prior to use



-70 °C Freezers Long-term storage not necessary at vaccination centers

Unless vaccination centers want to store for up to 6 months Special warehouses already identified





Example: decentralized distribution in Germany: BioNTech delivers to 25 distribution centers, run by federal states



Accountability of the federal states



Example: comprehensive information provided to professionals and vaccination centers in Germany





Medical Information Center (HCP)



BNT162

Handlungshilfe

für die Apotheke

Virtual Service Agent (Al chatbot)



Vaccination Center Starter Kit

BIONTE



COMIRNATY®:

A journey from scientific discovery to approval



Project Lightspeed – a 10-month journey to an effective and safe vaccine









Data from Phase 3 study shows 95% efficacy



- Analysis indicates efficacy rate of 95% in participants with and without prior SARS-CoV-2 infection
- Final analysis of unblinded data by independent data monitoring committee conducted on Nov 18, 2020
- Vaccinated participants will continue to be monitored for efficacy and safety for up to 2 years





Phase 3 trial data suggest rapid onset of protection against COVID-19

People are vaccinated twice with 30 µg of mRNA, 21 days apart



protection against COVID-19 7 days after 2nd dose²



12 days after the 1st dose

¹ https://www.nejm.org/doi/full/10.1056/NEJMoa2034577?query=featured_home 15 ² Individuals may not be fully protected until 7 days after their second dose of vaccine.

Clinical data indicate COVID-19 Occurrence From 7 Days After Dose 2 by Comorbidity Status¹

	BNT162b2 (30 μg) N=18,198		Placebo N=18,325			
	n	Surveillance Time (n)	n	Surveillance Time (n)	VE (%)	(95% CI)
Overall	8	2.214 (17,411)	162	2.222 (17,511)	95.0	(90.0, 97.9)
Comorbidity						
No comorbidity	4		76		94.7	(85.9, 98.6)
Any comorbidity	4		86		95.3	(87.7, 98.8)
Any malignancy	1		4		75.7	(-145.8, 99.5)
Cardiovascular	0		5		100.0	(-0.8, 100.0)
Chronic pulmonary disease	1		14		93.0	(54.1, 99.8)
Diabetes	1		19		94.7	(66.8, 99.9)
Obese (≥30.0 kg/m²)	3		67		95.4	(86.0, 99.1)
Hypertension	2		44		95.4	(82.6, 99.5)
Diabetes (including gestational diabetes)	1		20		95.0	(68.7, 99.9)



Clinical trial data indicates vaccine is highly efficacious with a favorable safety profile

Gold standard of clinical research – randomized large-scale clinical trial – to ensure safety and efficacy. We took important steps in parallel to accelerate the process together with the authorities – without shortcuts

Clinical Efficacy		~44,000	No serious safety		
95% >94%		participants in phase 3 trials	CONCERNS reported by the independent Data Monitoring Committee (DMC) to date		
in all subjects	bjects in subjects >65 y/o in U.S., Germany, Turkey, South Afr Brazil and Argentina More than 40% between 65-85 years				
Generally well tolerated Observed side-effects are common reactions to vaccination and transient. ¹			The only Grade 3 adverse events greater than 2% in frequency following dose 2 were:		
Adverse events were usually mild to moderate in intensity and resolved within a few days after vaccination.			Headache 2.0%	Fatigue 3.8%	
Most frequently chills, joint pain	observed adverse eve and fever.	ents were injection site pain and swellin	g, fatigue, headac	che, muscle pain,	

17 ¹Full safety assessment has been completed for ~38,000 study participants; BioNTech is also collecting safety data from adolescents and planning a pediatric study and a study on any effects on pregnancy



How mRNA works:

A deep dive into the technology



What is messenger RNA?



- The first molecule of life, involved in almost all aspects of cell biology
- Can be synthesized and engineered to resemble mRNA molecules as they occur naturally in the cytoplasm of human cells and transiently deliver proteins of interest
- mRNA has a transient messenger function and is rapidly degraded in the body



mRNA vaccines are a natural solution that avoid the use of viruses

Natural molecule studied for > 50 years	Does not require addition of adjuvants or use of a viral vector for administration	Highly scalable production		
with well-characterized bio-safety properties	High purity and animal material free	Precision vaccine Virus-free Non-integrating into DNA Non-infectious		
S' Cap 5' UTR VIRUS ANTIGEN 3'				
Genetic information Vaccine SARS-CoV-2 mRNA	mRNA Clinical LNP testing	Phase 3EUA /Vaccinationtrialsapproval		



How mRNA vaccines work – training the immune system for a real infection



Multiple levers of immune response: Strong antibody and robust T cell responses observed



BioNTech Publications:

- 1. Holtkamp et al. Modification of antigen-encoding RNA increases stability, translational efficacy, and T-cell stimulatory capacity of dendritic cells. Blood 2006.
- 2. Orlandini von Niessen et al. Improving mRNA-based therapeutic gene delivery by expression-augmenting 3' UTRs identified by cellular library screening. Molecular Therapy, 2019.
- 3. Vogel et al. A prefusion SARS-CoV-2-spike RNA vaccine is highly immunogenic and prevents lung infection in non-human primates. Nature 2020.
- 4. Walsh et al. Safety and immunogenicity of two RNA-based Covid-19 vaccine candidates: New England Journal of Medicine, 2020
- 5. Sahin et al. Concurrent human antibody and TH1-type cell responses elicited by a Covid-19 RNA vaccine. Nature, 2020.
- 6. Polack et al. Safety and efficacy of the BNT162b2 mRNA covid-19 vaccine: New England Journal of Medicine, 2020
- 7. Sahin et al. BNT162b2 induces SARS-CoV-2-neutralising antibodies and T cells in humans. Medrxiv, 2020.





BNT162b2 induced Antibodies cross-neutralize mutant SARS-COV-2 variants in pVNT assay



BION

What does COMIRNATY® contain – and why?

mRNA*

Active ingredient

This encodes the viral spike glycoprotein of the SARS-CoV-2 virus.

Salt

4 different salts

These buffer the vaccines to stabilize the pH, so that it matches the pH in our bodies.

* Each dose: of 0.3 mL with 30 micrograms mRNA



Water for injection

Sugar

Sucrose

This is a cryoprotectant. It ensures the lipids don't get too sticky at cold storage temperatures.

Lipids

4 different molecules

They form a protective capsule around the RNA, aiding in the delivery of the RNA, and protect the RNA from degradation.





What COMIRNATY[®] contains in detail – from the prescribers information



- mRNA: Active Ingredient
 - mRNA

Salt: 4 different salts

- potassium chloride
- potassium dihydrogen phosphate
- sodium chloride
- disodium phosphate dihydrate

- Sugar: Sucrose
 - Sucrose

Lipids: 4 different molecules

- ((4-hydroxybutyl)azanediyl)bis(hexane-6,1-diyl)bis(2-hexyldecanoate) (ALC-0315)
- 2-[(polyethylene glycol)-2000]-N,Nditetradecylacetamide (ALC-0159)
- 1,2-Distearoyl-sn-glycero-3phosphocholine (DSPC)
- cholesterol





Q&A







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Thank you for your participation!

The press conference has now ended.



