UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 6-K

REPORT OF FOREIGN PRIVATE ISSUER PURSUANT TO RULE 13a-16 OR 15d-16 UNDER THE SECURITIES EXCHANGE ACT OF 1934

FOR THE MONTH OF MAY 2022

COMMISSION FILE NUMBER 001-39081

BioNTech SE

(Translation of registrant's name into English)

An der Goldgrube 12
D-55131 Mainz
Germany
+49 6131-9084-0
(Address of principal executive offices)

Indicate by check mark whether the registrant files or will file annual reports under cover Form 20-F or Form 40-F: Form 20-F \boxtimes Form 40-F \square
Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1): \Box
Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7): \Box

DOCUMENTS INCLUDED AS PART OF THIS FORM 6-K				
On May 12, 2022, BioNTech SE (the "Company") issued a corrected	first quarter 2022 financial results conference prese	ntation. The conference call presentation is attached hereto as Exhibit 99.1.		

SIGNATURE

Pursuant to the requirements of the Exchange Act, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

BioNTech SE

By: /s/ Dr. Sierk Poetting

Name: Dr. Sierk Poetting Title: Chief Operating Officer

Date: May 12, 2022

EXHIBIT INDEX

<u>Exhibit</u> <u>Description of Exhibit</u>

99.1 First Quarter 2022: Corporate Update and Financial Results



1st Quarter 2022 Financial Results & Corporate Update

May 9, 2022



Exhibit 99.1

This Slide Presentation Includes Forward-looking Statements

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995, as amended, including, but not limited to, statements concerning: our expected revenues and net profit related to sales of our COVID-19 vaccine, referred to as COMIRNATY® where approved for use under full or conditional marketing authorization, in territories controlled by our collaboration partners, particularly for those figures that are derived from preliminary estimates provided by our partners; our pricing and coverage negotiations with governmental authorities, private health insurers and other third-party payors after our initial asks to national governments; the future commercial demand and medical need for initial or booster doses of a COVID-19 vaccine; competition from other COVID-19 vaccine controlled by our related to our order product and durability of immune responses, the rate and developed market acceptance of our COVID-19 vaccine; competition from other COVID-19 vaccine results, and cost of our research and development programs and our current and future preclinical studies and clinical trials, including statements regarding the triming of initiation and completion of studies or trials and related preparatory work, the period during which the results of the trials will become available and our research and development programs, the triming of collaboration with Pitze to develop and market a COVID-19 vaccine (including a potential booster dose of BNT162b2 and/or a potential booster dose of a variation of BNT162b2 having a modified mRNA sequence); the ability of BNT162b2 and/or a potential booster dose of a variation of BNT162b2 having a modified mRNA sequence); the ability of BNT162b2 and/or and filtral programs, and the continue research and development activities relating to our third-party collaborators to continue research and development activities relating to our third-party collaborators to continue research and development and our third-party collaborators and



Safety Information

COMIRNATY® ▼(the Pfizer-BioNTech COVID-19 vaccine) has been granted conditional marketing authorization (CMA) by the European Commission to prevent coronavirus disease 2019 (COVID-19) in people from 5 years of age. The vaccine is administered as a primary course of 2 doses, 3 weeks apart. In addition, the CAM has been expanded to include a booster dose (third dose) at least 6 months after the second dose in individuals 12 years of age and older, For immunocompromised individuals, a third primary course dose may be given at least 28 days after the second dose. The European Medicines Agency's (EMA's) human medicines committee (CHMP) has completed its rigorous evaluation of COMIRNATY®, concluding by consensus that sufficiently robust data on the quality, safety and efficacy of the vaccine are now available.

- MPORTANT SAFETY INFORMATION:
 Events of anaphylaxis have been reported. Appropriate medical treatment and supervision should always be readily available in case of an anaphylactic reaction following the administration of the vaccine.
 There is an increased risk of myocarditis and pericarditis following vaccination, and more often in younger males. Available data suggest that the course of myocarditis and pericarditis following vaccination, and more often in younger males. Available data suggest that the course of myocarditis and pericarditis following vaccination is not different from myocarditis or general.

 Anxiety-related reactions, including vasovagal reactions (syncope), hyperventilation or stress-related reactions are temporary and resolve on their own. Individuals should be advised to bring symptoms to the attention of the vaccination provider for evaluation. It is important that precautions are in place to avoid injury from fainting.

 Vaccination should be postponed in individuals suffering from acute severe febrile illness or acute infection. The presence of a minor infection and/or low-grade fever should not delay vaccination. As with other intramuscular injections, the vaccine should be given with caution in individuals receiving anticoaquant therapy or those with thrombocytopenia or any coaquilation disorder (such as haemophilia) because bleeding or bruising may occur following an intramuscular administration in these individuals.

 The efficacy and safety of the vaccine has not been assessed in immunocompromised individuals. As with their the vaccine has not been assessed in immunocompromised individuals. As with any vaccine, vaccination with COMRNATY® may not protect all vaccine recipients, inclined tables, adverse reactions in participants 16 years of age and older were injection site pain (>80.96), highigue of the vaccine has 7.090, highigue and the 162 of 3.090, highigue and
- For complete information on the safety of COMIRNATY® always make reference to the approved Summary of Product Characteristics and Package Leaflet available in all the languages of the European Union on the EMA website.

The black equilateral triangle ▼ denotes that additional monitoring is required to capture any adverse reactions. This will allow quick identification of new safety information. Individuals can help by reporting any side effects they may get. Side effects can be reported to EudraVigilance or directly to BioNTech using email medinto@biontech.de, telephone +49 6131 9084 0, or via the website www.biontech.de



Safety Information

AUTHORIZED USE IN THE U.S.
COMIRNATY* (COVID-19 Vaccine, mRNA) is an FDA-approved COVID-19 vaccine for active immunization to prevent coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-COV.2) in individuals 15 years of age and older. It is also authorized under EUA to provide a 2-dose primary series to individuals 5. years of age and older. It is also authorized under EUA to provide a 2-dose primary series to a part and older. It is also authorized under EUA to provide a 2-dose primary series to the law of the primary series dose to individuals 5 years of age and older who have completed primary series with Pfleze-BioNTech COVID-19 Vaccine or COMIRNATY*, a single booster dose to individuals 18 years of age and older who have completed primary series with Pfleze-BioNTech COVID-19 Vaccine or COMIRNATY*, a single booster dose to individuals 12 years of age and older who have received a first booster dose of any authorized COVID-19 vaccine; and a second booster dose to individuals 12 years of age and older who have received a first booster dose of any authorized COVID-19 vaccine.

The booster schedule is based on the labeling information of the vaccine used for the primary series.

- IMPORTANT SAFETY INFORMATION
 Individuals should not get the vaccine if they:

 had a severe allergic reaction after a previous dose of this vaccine

 had a severe allergic reaction to any ingredient of this vaccine

Individuals should tell the vaccination provider about all of their medical conditions, including if they: Individuals should tell the vaccination provider about all of their medical conditions, including if they:

have any allegrige.

have had myocarditis (inflammation of the heart muscle) or pericarditis (inflammation of the lining outside the heart)

have a lever.

have a bleeding disorder or are on a blood thinner

are immunocompromised or are on a medicine that affects the immune system

are pregnant, plan to become pregnant, or are breastfeeding

have received another COVID-19 vaccine

have received another COVID-19 vaccine

have ever fainted in association with an injection



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BIONTECH

Our Vision: Harnessing The Power Of The Immune System To Fight Human Diseases



BIONTECH

Highlights in Q1 2022



- Reported Q1 total revenues of €6.4 bn¹
- Signed pandemic preparedness contract with Federal Republic of Germany through 2027
- Multiple new deals signed:
 - Matinas lipid nanocrystal collaboration
 - Regeneron collaboration to advance BNT116 in combination with Libtayo in NSCLC
- Invoiced ~750 m doses globally in Q1
- FDA authorized 4th dose in adults 50 years+ and in immunocompromised individuals 12 years+
- Continued label expansion in multiple regions for booster dose in 12+ years
- BNT211 (CLDN6 CAR-T cell therapy) Phase 1/2 data presented at AACR showed manageable safety profile and signs of clinical activity
- First RiboMab BNT141 (CLDN18.2 antibody) entered Phase 1 clinical study in solid tumors

1 BioNTech's profit share is estimated based on preliminary data shared between Pfizer and BioNTech as further described in the Annual Report on Form 20-F for the year ended December 31, 202 as well as the Quarterly Report as of and for the three months ended March 31, 2022, filed as an exhibit to BioNTech's Current Report on Form 6-K. Any changes in the estimated share of the collaboration anderly stores conflict with the reconsized cross-cribed.



Proactive Approach to Managing COVID-19 at a Global Scale

Strong global position to tackle COVID-19 pandemic

Delivered nearly 3.4 bn¹ doses cumulatively to >175 countries and regions

On track to achieve pledge to deliver a total of 2 bn doses to low- and middle-income countries by end of 2022

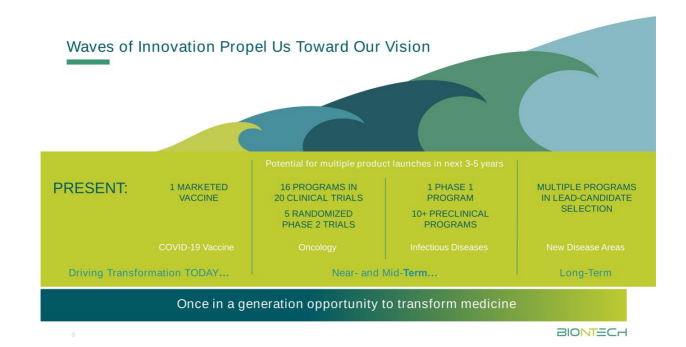
Innovation to stay ahead of COVID-19

- Optimized formulation
- Pediatric label expansion
 - Submission for boosters in children 5 to <12 yrs
 - Evaluating 3-dose primary regimen in children
 6 months to <5 yrs; data expected in coming weeks
- Future pandemic preparedness
 - Monitoring of emerging variants
 - Rapid data-guided vaccine adaptation
- Pre-emptive approach to variants
 - Comprehensive variant-adapted and next-gen vaccine development program
 - Broad research program to study anti-SARS-CoV-2 immune profile after vaccinations, boosters, breakthrough infections to inform strategy

1 As of end-April 2022

his slide has been updated to remove an incorrect graphic regarding the countries to which the Pfizer-BioNTech COVID-19 vaccine has been delivere





Agenda

01 First Quarter 2022 Highlights Ugur Sahin, CEO

Pipeline Update Özlem Türeci, CMO

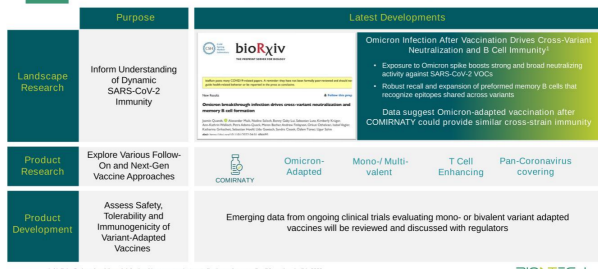
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1.0

COVID-19 Vaccine R&D Strategy to Drive Pandemic Preparedness

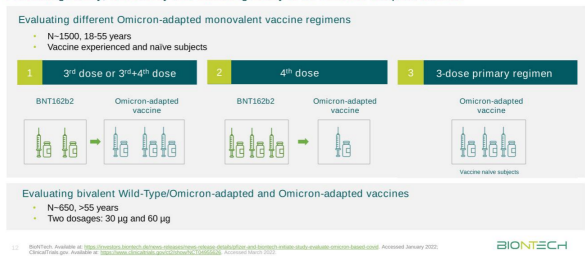


bioRxiv. Omicron breakthrough infection drives cross-variant neutralization and memory B cell formation; April 1, 2022.
 Available at. https://www.biorxv.org/content/10.1101/2022.04.01.486695y1.full.pdf
 VOC., variants of concern

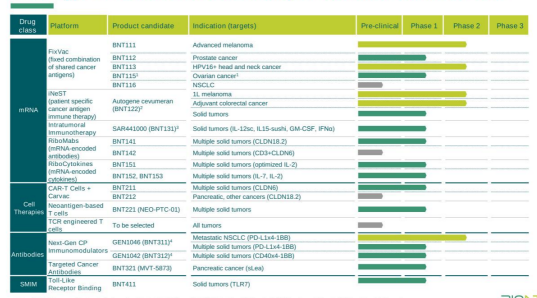


Comprehensive Clinical Response Strategy to Omicron Variant

Assessing Safety, Tolerability and Immunogenicity of an Omicron-Adapted Vaccine



Oncology: Advancement Across Multiple Modalities and Indications



¹BNT115 is currently being studied in an investigator-initiated Phase 1 trial. ²Collaboration with Genentech ³Collaboration with Sanofi, ⁴Collaboration with Genma SMIM, Small Molecule Immunomodulators



Focused Execution in 2022 Across 5 Phase 2 Programs in Various Solid Tumor Types

Platform	FixVac Off-the-shelf mRNA vaccine				Bispecific Next-generation immunotherapy
Program	BNT111 R/R Melanoma	BNT113 HPV16+ HNSCC	Autogene cevumeran BNT122 ¹ 1L Melanoma	Autogene cevumeran BNT122 ¹ Adjuvant colorectal cancer	BNT311 ² R/R NSCLC
How	Encodes 4 tumor- associated antigens covering >90% of cutaneous melanoma patients U.S. Fast Track	• Encodes HPV16 oncoproteins E6 & E7	Targets 20 neo-antigens unique to each patient Data update expected 2H 2022	Targets 20 neo-antigens unique to each patient	Conditional 4-1BB co- stimulation while blocking PD(L)1 axis
	Designation and Orphan Drug Designation				
Why	Potential to improve outcomes in combo with anti-PD1	 Potential for synergistic anti-tumor effect in combination with anti- PD1 	Trial success may unlock 1L use of iNeST as combination therapy with anti-PD(L)1 in anti-PD1- naive advanced cancers	Potential to address residual cancer cells that remain – focus on recurrence free survival	 Enhances T cell and NK cell function and targets them to tumor lesions

R/R. refractory/relapsed: HPV16+, human papilloma virus type 16 positive; HNSCC, head and neck squamous cell carcinoma; NK cell, Natural killer cell, CPI, checkpoint inhibitor 1 Collaboration with Generatech, 2 Collaboration with Generatech, 2 Collaboration with Generatech.



BNT211: CAR-T Cell Program with Potential Targeting Multiple High-Need Solid Tumors

BNT211 CAR Structure CLDN6 not present in healthy tissues 2nd generation CAR Directed against CLDN6 Cancer specific carcino-embryonic αCLDN6 scFv antigen Expressed in multiple solid cancers with CD8 hinge CLDN6 expressed in multiple cancers high medical need 4-1BB CARVac drives in vivo expansion, persistence and efficacy of CAR-T cells СD3ζ Part 1: monotherapy CLDN6 CAR-T dose escalation: 1x10⁷ CAR-T (DL1) 1x10⁸ CAR-T (DL2) 1x10⁹ CAR-T (DL3) Part 3 Expansion Cohorts Patients with CLDN6+ Ovarian Cancer High CLDN6 relapsed/ refractory Testicular Cancer Endometrial Cancer Lung Cancer solid tumors t 2: combination thera CLDN6 CAR-T dose alation + CLDN6 CARVa 1x10⁷ CAR-T (DL1) 1x10⁸ CAR-T (DL2) 1x10⁹ CAR-T (DL3) (up to 36 patients) Gastric Cancer Tumors NOS

CLDN6, Claudin-6; CAR-T cells, chimeric antigen receptor engineered T cells; scFv, single chain variable fragment; RP2D, recommended Phase 2 dose; NOS, not otherwise specific Reinhard K. et al. Science 2002: 367-446-453



BNT211: CAR-T in Solid Tumors Encouraging Efficacy and Safety Profiles Presented at AACR



CLDN6 CAR-T cells as monotherapy or combined with CARVac well tolerated at dose levels evaluated to date (1x10 7 and 1x10 8 CAR-T)

- Grade 1-2 CRS seen in 70% of patients at 1x108 CAR-T dose, manageable by administration of tocilizumab
- 2 DLTs observed, both patients fully recovered and showed clinical benefit
- MTD not reached yet



- Robust CAR-T engraftment achieved in all patients translating into clinical activity: ORR 43%, DCR of 86% in evaluable patients (n=14; 1x10⁷ and 1x10⁸ CAR-T)
 - 6 PR, 5 SD+, 1 SD (Testicular, ovarian and other tumors, 6 weeks post-infusion)
 - 5 testicular cancer patients show promising responses at 1x10⁸ CAR-T: ORR 80%, DCR 100%; 1 CR, 3 PR, 1 SD
- CARVac supports CAR-T engraftment and mediates physiologic expansion plus upregulation of survival pathways
- Some patients show continuing CAR-T persistence (>150 days post infusion)
- Patients with initial PR showed further deepening of responses

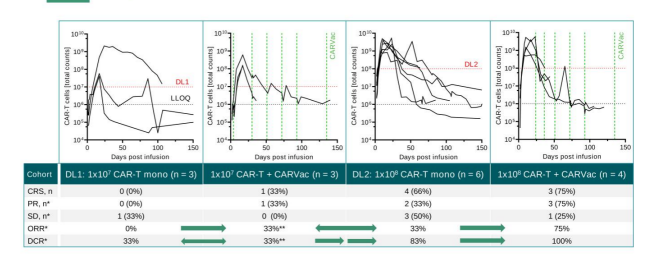
Data cut-off: MAR 10, 2022

DL1: 1x107 CAR-T; DL2: 1x108 CAR-T

CLDN6, Claudin-6; DLT, dose-limiting toxicity; MTD, maximum tolerated dose; CRS, cytokine release syndrome; CR, complete response; DCR, disease control rate; DL, dose level; ORR, overall response; DCR and isease



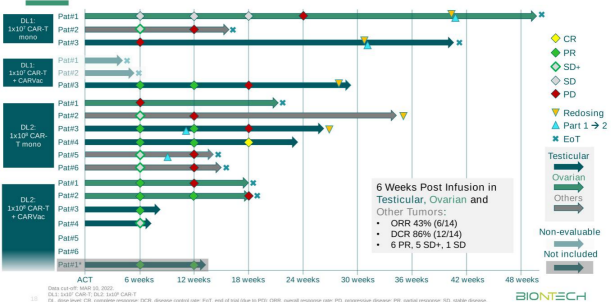
Robust CAR-T Engraftment Seen in all Patients and Persisting CAR-T in Responding Patients



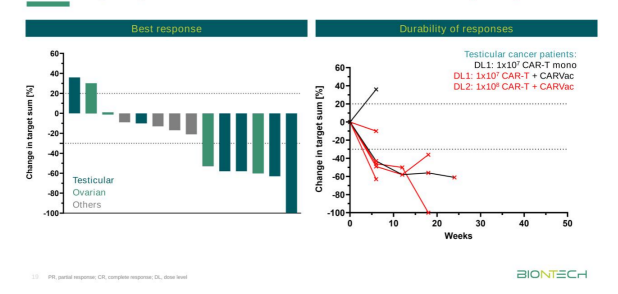
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DL1: Ixt0* CAR-T; DL2: Ixt0* CAR-T; CR5, cytokine release syndrome; DCR, disease control rate; DL, dose level; DLT, dose-limiting toxicity; ORR, overall response rate; PR, partial response; SD, stable disease; *Af first tumor assessment

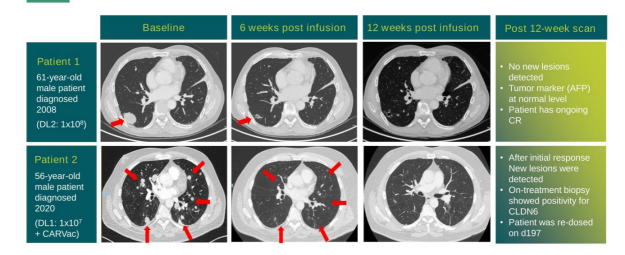
Efficacy Observed at 6 Weeks Post Infusion



Continuing Responses in Testicular Cancer with One PR Deepening to CR



Responses in Two Testicular Cancer Patients with Relapse After Prior Treatment



Data cut-off: MAR 10, 2022
AFP, alpha-fetoprotein; CR, complete response; CLDN6, Claudin-6.



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Key Highlights For Q1 2022

Total Revenues ¹	Operating Result
€6.4 bn	© €4.8 bn
Diluted EPS	Cash and Trade Receivables
<u>.ff.</u> €14.24	€6.2 bn + €12.7 bn

1. BioNTech's profit share is estimated based on preliminary data shared between Pfizer and BioNTech as further described in the Annual Report on Form 20-F for the year ended December 31, 202, field as an exhibit to BioNTech's Current Report on Form 8-K. Any changes in the estimated share of the collaboration partner's gross profit will be recognized prospectively.



First Quarter 2022 COVID-19 Vaccine Commercial Revenues



Strong Q1 2022 - BioNTech reiterates 2022 financial year guidance

1. BioNTech's profit share is estimated based on preliminary data shared between Pfizer and BioNTech as further described in the Annual Report on Form 20-F for the year ended December 31, 2021 as well as the Quarterly Report as of and for the three months ended March 31, 2022, filed as an exhibit to BioNTech's Current Report on Form 6-K. Any changes in the estimated share of the collaboration partner's gross profit will be recognized prospectively.



Q1 2022 Financial Results – Profit or Loss

(in millions, except per share data)¹	Q1 2022	Q1 202
Research & development revenues	€12.4	€20.
Commercial revenues ²	6,362.2	2,027.
Total revenues	€6,374.6	€2,048.
Cost of sales	(1,294.1)	(233.1
Research and development expenses	(285.8)	(216.2
Sales and marketing expenses	(14.3)	(8.7
General and administrative expenses	(90.8)	(38.9
Other operating income less expenses	63.1	110.
Operating income	€4,752.7	€1,662.
Finance income less expenses	265.4	(19.9
Income taxes	(1,319.3)	(514.2
Profit for the period	€3,698.8	€1,128.
Earnings per share		
Basic profit for the period per share	€15.13	€4.6
Diluted profit for the period per share	€14.24	€4.3

Numbers have been rounded, numbers presented may not add up precisely to the totals and may have been adjusted in the table context. Presentation of the consolidated statements of profit or loshas been condensed.

² BioNTech's profit share is estimated based on preliminary data shared between Pfizer and BioNTech as further described in the Annual Report on Form 20-F for the year ended December 31, 2021 at well as the Quarterly Report as of and for the three months ended March 31, 2022, filed as an exhibit to BioNTech's Current Report on Form 6-K. Any changes in the estimated share of the collaboration partner's gross profit will be recognized prospectively.



2022 Financial Year Guidance Reiterated

COVID-19 Vaccine Revenues for FY 2022 ¹	
Estimated BioNTech COVID-19 vaccine revenues	€ 13 – 17 bn
Planned FY 2022 Expenses and Capex ¹	
R&D expenses	€ 1,400 - 1,500 m
SG&A expenses	€ 450 - 550 m
Capital expenditure	€ 450 - 550 m
Estimated FY 2022 Tax Assumptions	
BioNTech Group estimated annual effective income tax rate	~28%²





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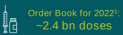
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Significant Pipeline Expansion and Maturation Expected in 2022

Continue COVID-19 2 · Label & geographic First randomized Initiate 4 FIH vaccine trials Autoimmune disease expansion Phase 2 readout • 10+ additional mRNA · Regenerative medicine Next-generation vaccines Prepare for vaccine programs Cardiovascular disease registrational trials Innovations for pandemic · Precision antibacterials POC data for CAR-T preparedness cell therapy Invest in Foundation to Enable Accelerated Innovation and Expansion Digital & Al Capabilities | Technologies | Development Team | Manufacturing | Global Footprint

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COVID-19 Vaccine Outlook 2022





Upcoming Data

BNT162b2	Timing
 Data for 4th dose in adults, aged 16 to 65 years² 	ongoing
 Data for 3-dose regimen in children, aged 6 months to <5 years 	coming weeks

Follow-on and next generation vaccines	
Omicron-adapted vaccine: monovalent, bivalent - 3rd and/or 4th dose	coming weeks
Multiple updates: Follow on and next-gen vaccines	2H

28 1 As of mid-April 2022; In combination with contracts entered into by Pfizer



Further Expected Pipeline Milestones in 2022

4 Infectious Disease First-In-Human	Trial Starts
• Shingles vaccine ¹	2H
Tuberculosis vaccine ²	2H
HSV 2 vaccine	2H
Malaria vaccine	2Н
3 Oncology First-in-Human Trial	Starts
BNT141 – RiboMab, solid tumors	FPD in January
BNT142 – RiboMab, solid tumors	1H
BNT116 - FixVac in combination w/Libtayo, NSCLC	2Н
3 Data Updates	
BNT161 – Influenza mRNA vaccine¹	2022
BNT122 ³ Phase 2 – iNeST in combination w/Pembrolizumab, 1L Melanoma	2Н
BNT211 Phase 1/2 – CAR-T/CLDN6+, multiple solid tumors	2Н
HSV 2, Herpes simplex virus type 2; FPD, first patient dosed; CLDN, Claudin; NSCLC, non-small cell lung cancer 1 Partnered with Briser 2 Collaboration with BMCE 3 Partnered with Concented.	BIONTE



Annual General Meeting
June 1, 2022

Virtual Capital Markets Day June 29, 2022







