PHARMACOVIGILANCE PLAN FOR

BIOLOGIC LICENSE APPLICATION #125742

OF

COMIRNATY® (PFIZER-BIONTECH COVID-19 Vaccine VACCINE, mRNA, BNT162b2, PF-07302048)

Date of Report: 26 August 15 December 2021

Version 1.24

BNT162b2					
1.16 Risk Management Plan	(Non-REMS)	for Biologic	License A	Application #	125742

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LIST OF ABBREVIATIONS

Abbreviation	Definition of Term
AE	adverse event
AESI	adverse event of special interest
A:G	albumin:globulin
ALC-0315	((4-hydroxybutyl)azanediyl)bis(hexane-6,1-diyl)bis(2-
	hexyldecanoate)
ALC-0159	2-[(polyethylene glycol)-2000]-N,N-
	ditetradecylacetamide
ARDS	acute respiratory distress syndrome
BALB/c	bagg albino
BC	Brighton Collaboration
BEST	biologics effectiveness and safety
BLA	biologics license application
BMI	body mass index
BP	blood pressure
CD4, CD8	cluster of differentiation-4, 8
CDC	Centers for Disease Control and Prevention
CI	confidence interval
COPD	chronic obstructive pulmonary disease
COVID-19	coronavirus disease 2019
CSR	clinical study report
СТ	clinical trial
DART	developmental and reproductive toxicology
DCA	data capture aid
DLP	data-lock point
DoD	Department of Defense
ECDC	European Center for Disease Control
EEA	European Economic Area
EMA	European Medicines Agency
eGFR	estimated glomerular filtration rate
EU	European Union
EUA	emergency use authorization
FDA	(US) Food and Drug Administration
GLP	good laboratory practice
HbA1c	glycated hemoglobin
HBV	hepatitis b virus
HCV	hepatitis c virus
HIV	human immunodeficiency virus
IA	interim analysis
ICU	intensive care unit
IFN	interferonInterferon
IL-4	interleukin-4
IM	intramuscular(ly)

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Abbreviation	Definition of Term		
IMD	index of multiple deprivation		
IND	investigational new drug		
LNP	lipid nanoparticle		
LOE	lack of efficacy		
MAAMAH	marketing authorization applicantholder		
MedDRA	Medical Dictionary for Regulatory Activities		
MERS-CoV	Middle East respiratory syndrome–coronavirus		
MHS	Military Health System		
MIS-C	multisystem inflammatory syndrome in children		
MOA	mechanism of action		
modRNA	nucleoside-modified messenger ribonucleic acid		
mRNA	messenger ribonucleic acid		
NCMD	National child mortality database		
NDA	new drug application		
NDS	new drug submission		
NHP	nonhuman primate		
NHS	National Health Service		
NICE	National Institute for Health and Care Excellence		
OCS	oral corticosteroids		
OTIS	Organization of Teratology Information Specialists		
PHN	Pediatric Heart Network		
PK	pharmacokinetic		
PRAC	pharmacovigilance risk assessment committee		
PT	Preferred Term		
PVP	pharmacovigilance plan		
RBC	red blood cell		
RMP	Risk Management Plan		
RNA	ribonucleic acid		
RR	relative risk		
RSV	respiratory syncytial virus		
SAE	serious adverse event		
SARS	severe acute respiratory syndrome		
SARS-CoV-1	severe acute respiratory syndrome coronavirus 1		
SARS-CoV-2	severe acute respiratory syndrome coronavirus 2		
siRNA	small-interfering RNA		
<u>SMQSMSR</u>	standardised MedDRA querysummary monthly safety		
	report		
Tdap	tetanus, diphtheria, and acellular pertussis		
TESSy	The European Surveillance System		
Th1	T helper cell type 1		
Th2	T helper cell type 2		
UK	United Kingdom		
US	United States		
USP	United States pharmacopeia		

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Abbreviation	Definition of Term
V8	variant 8
V9	variant 9
VAED	vaccine-associated enhanced disease
VAERD	vaccine-associated enhanced respiratory disease
WBC	white blood cells
WHO	World Health Organization
WOCBP	women of childbearing potential

1. INTRODUCTION

1.1. Product Details

Table 1.Product Details^a

Product	COVID-19 Vaccine, mRNA, herein after referred to as BNT162b2 is a nucleoside-modified messenger RNA -((modRNA) encoding the viral spike (S) glycoprotein of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2).
Brief description of the product	Chemical class: Nucleoside modRNA formulated in lipid particles. Mechanism of Action: The modRNA in the BNT162b2 is formulated in lipid particles, which enable delivery of the RNAmRNA into host cells to allow expression of the SARS-CoV-2 S antigen. The vaccine elicits an immune response to the S antigen, which protects against COVID-19. Important information about its composition: • • The BNT162b2 is a sterile suspension for injection. • The BNT162b2 is supplied as a frozen suspension in multiple dose vials. • Each vial must be diluted with 1.8 mL of sterile 0.9% Sodium Chloride Injection, USP prior to use to form the vaccine.
	 Each dose of the BNT162b2 contains 30 mcg of a nucleoside-modified messenger RNA (modRNA) encoding the viral spike (S) glycoprotein of SARS-CoV-2. Each <u>0.3 mL</u> dose of the BNT162b2 also includes the following ingredients: lipids (0.43 mg {((4-hydroxybutyl)azanediyl)bis(hexane-6,1-diyl)bis(2-hexyldecanoate), 0.05 mg 2[((polyethylene glycol)-2000]-)_NN-ditetradecylacetamide, 0.09 mg 1,2-distearoyl-sn-glycero-3-phosphocholine, and 0.2 mg cholesterol), 0.01 mg potasium chloride, 0.01 mg monobasic potassium phosphate, 0.36 mg sodium chloride, 0.07 mg dibasic sodium phosphate dihydrate, and 6 mg sucrose. The diluent (0.9% Sodium Chloride Injection, USP) contributes an additional 2.16 mg sodium chloride per dose. The BNT162b2 does not contain preservative. The vial stoppers are not made with natural rubber latex.
Indication	Current: Active immunization to prevent COVID-19 caused by SARS-CoV-2 in individuals 16 years of age and older. Proposed: Active immunization to prevent COVID-19 caused by SARS-CoV- 2 in individuals 12 years of age and older.
Dosage and route of administration	Proposed: A primaryCurrent: BNT162b2 is administered intramuscularly as a series of two doses (0 3 mL each) 3 weeks apart, intramuscularly. A booster dose (a third dose) may be administered intramuscularly A booster dose (a third dose) may be administered intramuscularly
	approximately 6 months after the second dose. There are no data available on the interchangeability of BNT162b2 with other COVID-19 vaccines to complete the vaccination series. Individuals who have received 1 dose of

Table 1. Product Details^a

BNT162b2 sho vaccination set	uld receive a second dose of BNT162b2 to complete the ies.

a. COVID-19 Vaccine, mRNA US Prescribing Information

Data Lock	16 years and older	13 March 2021 (Pfizer Clinical Database)
Point / Data		23 October 2020 (BioNTech Clinical Database)
cut-off:		18 June 2021 (Pfizer Safety Database)
	12 to 15 years older	13 March02 September 2021 (Pfizer Clinical Database)
		18 June 2021 (Pfizer Safety Database)
	Booster dose	17 June 2021 (Pfizer Clinical Database)
		18 June 30 September 2021 (Pfizer Safety Database)*)

a. All safety concerns analysis from the safety database are aligned to this DLP

Merged Cells

2. SAFETY SPECIFICATION

2.1. Elements of the Safety Specification

2.1.1. Non-Clinical

Nonclinical evaluation of BNT162b2 included pharmacology (mouse immunogenicity and NHP immunogenicity and challenge studies), pharmacokinetic (series of biodistribution, metabolism and pharmacokinetic studies), and toxicity (2 GLP rat repeat-dose toxicity and a GLP DART) studies in vitro and in vivo. No additional toxicity studies are planned for BNT162b2.

Nonclinical studies in mice and NHP for BNT162b2 demonstrated both a strong neutralizing antibody response and a Th1-type CD4⁺ and an IFN γ^+ CD8⁺ T-cell response. The Th1 profile is characterized by a strong IFN γ , but not IL-4, response indicating the absence of a potentially deleterious Th2 immune response and is a pattern favored for vaccine safety and efficacy.¹ Rhesus macaques (Study VR-VRT-10671) that had received two IM immunizations with 100 µg BNT162b2 or saline 21 days apart were challenged with 1.05 × 10⁶ plaque forming units of SARS-CoV-2 (strain USA-WA1/2020), split equally between the intranasal and intratracheal routes.² BNT162b2 provided complete protection from the presence of detectable viral RNA in the lungs compared to the saline control with no clinical, radiological or histopathological evidence of vaccine-elicited disease enhancement.

An intravenous rat PK study, using an LNP with the identical lipid composition as BNT162b2, demonstrated that the novel lipid excipients in the LNP formulation, ALC-0315 and ALC-0159, distribute from the plasma to the liver. While there was no detectable excretion of either lipid in the urine, the percent of dose excreted unchanged in feces was ~1% for ALC-0315 and ~50% for ALC-0159. Further studies indicated metabolism played a role in the elimination of ALC-0315. Biodistribution was assessed using luciferase expression as a surrogate reporter formulated like BNT162b2, with the identical lipid composition. After IM injection of the LNP-formulated RNA encoding luciferase in BALB/c mice, luciferase protein expression was demonstrated at the site of injection 6 hours post dose and expression decreased over time to almost reach background levels after 9 days. Luciferase was detected to a lesser extent in the liver; expression was present at 6 hours after injection and was not detected by 48 hours after injection. After IM administration of a radiolabeled LNP-mRNA formulation containing ALC-0315 and ALC-0159 to rats, the percent of administered dose was also greatest at the injection site. Outside of the injection site, total recovery of radioactivity was greatest in the liver and much lower in the spleen, with very little recovery in the adrenal glands and ovaries. The metabolism of ALC-0315 and ALC-0159 was evaluated in blood, liver microsomes, S9 fractions, and hepatocytes from mice, rats, monkeys, and humans. The in vivo metabolism was examined in rat plasma, urine, feces, and liver samples from the PK study. ALC-0315 and ALC-0159 are metabolized by hydrolytic metabolism of the ester and amide functionalities, respectively, and this hydrolytic metabolism is observed across the species evaluated.

In GLP toxicity studies, two variants of the BNT162b2 candidate were tested, designated "variant 8" and "variant 9" (V8 and V9, respectively). The variants differ only in their codon optimization sequences which are designed to improve antigen expression, otherwise the

amino acid sequences of the encoded antigens are identical. BNT162b2 (V9) was evaluated clinically and submitted for application. Two GLP-compliant repeat-dose toxicity studies were performed in Wistar Han rats; one with each variant. Both studies were 17 days in duration with a 3-week recovery period. A GLP-compliant DART study in Wistar Han rats has also been completed. Safety pharmacology, genotoxicity and carcinogenicity studies have not been conducted, in accordance with the 2005 WHO vaccine guideline.³

The IM route of exposure was selected for nonclinical investigations as it is the clinical route of administration. Rats were selected as the toxicology test species as they demonstrated an antigen-specific immune response to the vaccine and are routinely used for regulatory toxicity studies with an extensive historical safety database.

Administration of up to 100 µg BNT162b2 by IM injection to male and female Wistar Han rats once every week, for a total of 3 doses, was tolerated without evidence of systemic toxicity. Expected inflammatory responses to the vaccine were evident such as edema and erythema at the injection sites, transient elevation in body temperature, elevations in WBC count and acute phase reactants, and lower A:G ratios. Injection site reactions were common in all vaccine-administered animals and were greater after boost immunizations. Changes secondary to inflammation included slight and transient reduction in body weights and transient reduction in reticulocytes, platelets and RBC mass parameters. Decreased reticulocvtes were reported in rats treated with the licensed LNP-siRNA pharmaceutical OnpattroTM (NDA # 210922) but have not been observed in humans treated with this biotherapeutic⁴ suggesting this is a species-specific effect. Decreased platelet counts were noted after repeat administration, but were small in magnitude of change, likely related to inflammation-related platelet activation and consumption, and unassociated with other alterations in hemostasis. Elevated levels of gamma-glutamyl transferase were observed in the first repeat-dose toxicity study with BNT162b2 (V8) without evidence of cholestasis or hepatobiliary injury but was not recapitulated in the second repeat dose-toxicity study with BNT162b2 (V9), the final clinical candidate. All changes in clinical pathology parameters and acute phase proteins were reversed at the end of the recovery phase for BNT162b2, with the exception of low magnitude higher red cell distribution width (consistent with a regenerative erythroid response) and lower A:G ratios (resulting from acute phase response) in animals administered BNT162b2. Macroscopic pathology and organ weight changes were also consistent with immune activation and inflammatory response and included increased size and/or weight of draining iliac lymph nodes and spleen. Vaccine-related microscopic findings at the end of the dosing phase consisted of edema and inflammation in injection sites and surrounding tissues, increased cellularity in the draining iliac lymph nodes, bone marrow and spleen and hepatocyte vacuolation in the liver. Vacuolation of periportal hepatocytes, the only test article-related liver microscopic finding, was not associated with any microscopic evidence of hepatic injury or hepatic functional effects (i.e., liver functional enzymes were not elevated) and may be associated with hepatocyte uptake of the LNP lipids.⁵ Microscopic findings at the end of the dosing phase were partially or completely recovered in all animals at the end of the 3-week recovery period for BNT162b2. A robust immune response was elicited to the BNT162b2 antigen.

Administration of BNT162b2 to female rats twice before the start of mating and twice during gestation at the human clinical dose $(30 \ \mu g)$ was associated with non-adverse effects (body

weight, food consumption and effects localized to the injection site) after each dose administration. However, there were no effects of BNT162b2 administration on mating performance, fertility, or any ovarian or uterine parameters in the F0 female rats nor on embryo-fetal or postnatal survival, growth, or development in the F1 offspring. An immune response was confirmed in F0 female rats following administration of each vaccine candidate and these responses were also detectable in the F1 offspring (fetuses and pups).

In summary, the nonclinical safety findings related to BNT162b2 administration primarily represent an expected immune reaction to vaccine administration and are clinically manageable or acceptable risks in the intended population. The key safety findings regarding BNT162b2 from nonclinical studies and their relevance to human usage are presented in Table 2. There was no evidence of vaccine-elicited disease enhancement.

Key Safety findings from Nonclinical Studies ^a	Relevance to Human Usage		
Pharmacology			
 NHP Challenge Model No evidence of vaccine-elicited disease enhancement. Toxicity Injection site reactions: Injection site reactions were common and reversible or showed signs of reversibility at the end of the 3-week recovery period in nonclinical studies. 	 Suggests low risk of vaccine-enhanced disease in humans; being investigated in CTs. In common with other vaccines, BNT162b2 administration has the potential to generate injection site reactions such as edema and erythema at the injection sites. 		
Inflammation and immune activation:			
• Evidence of inflammation or immune activation was common, reversible, and included transiently higher body temperature, higher circulating WBCs, and higher acute phase reactants. Secondarily, transiently lower body weights, reticulocytes, platelets, and RBC mass parameters were observed.	 In common with all vaccines, BNT162b2 administration has the potential to generate inflammation which can lead to increased body temperature, higher circulating WBCs and higher acute phase proteins. Decreased reticulocytes have not been observed in humans treated with the LNP-siRNA pharmaceutical Onpattro⁴, suggesting this finding in rats is a species- specific effect. BNT162b2 administration has the potential to transiently decrease platelets and RBC mass parameters. These slight decreases are not likely to be clinically meaningful due to their small magnitude. 		
 Developmental and Reproductive Toxicity No vaccine-related effects on female fertility or the development of fetuses or offspring were observed in a DART study of BNT162b2 in rats. 	• No effects are anticipated in WOCBP, pregnant women or their offspring.		

Table 2. Ke	y Safety	Findings and	Relevance to	Human	Usage
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a. Safety pharmacology, genotoxicity, and carcinogenicity studies were not conducted, in accordance with 2005 WHO vaccine guideline, as they are generally not considered necessary to support development and licensure of vaccines for infectious diseases.³ In addition, the components of the vaccine construct are lipids and RNA and are not expected to have carcinogenic or genotoxic potential.

2.1.2. Clinical

2.1.2.a. Limitations of the Human Safety Database

The pivotal study was initially planned to enroll approximately 30,000 participants, which would have a probability of 78% of detecting an AE with a frequency of 0.01% (1/1000) and a probability of 95% of detecting an AE with a frequency of 0.02% (1/500). The protocol was amended to enroll approximately 46,000 participants, which would slightly enhance the ability to detect AEs. However, rarer events might not be detected.

Participants in the pivotal study were initially planned to be followed for up to 24 months in order to assess the potential for late-occurring adverse reactions, such as the theoretical risk of VAED. After completing the final efficacy analysis with vaccine efficacy shown to be 95%, and obtaining regulatory authorization to vaccinate in many countries, Pfizer-BioNTech started to unblind all participants to determine those randomized to placebo so that they could be offered vaccine in accordance with local authorization. To date, most placebo subjects have been unblinded to receive active vaccine at or prior to 6 months after the second dose, therefore, a placebo group for comparison of safety data is only available for up to 6 months post Dose 2.

2.1.2.a.1. Clinical Trial Exposure

Brief Overview of Development

Study BNT162-01

BioNTech is conducting a first-in-human dose level–finding Phase 1/2 study (BNT162-01) in Germany to gather safety and immunogenicity data to enable evaluation of 4 vaccines candidates individually to inform the overall clinical development of a BNT162b2.

BNT162-01 is not conducted under the US IND application but is being conducted under a German Clinical Trial Application.

Four vaccine candidates were evaluated in Study BNT162-01. Based on safety and immunogenicity results from this study, 2 vaccine candidates, BNT162b1 and BNT162b2, were selected for evaluation in Study C4591001, which is a Phase 1/2/3 randomized, placebo-controlled, observer-blind, dose-finding, vaccine candidate-selection, and efficacy study in healthy adults (conducted under IND 019736).

Phase 1 of Study C4591001

Phase 1: comprised dose-level-finding evaluations of the 2 selected vaccine candidates; multiple dose levels (some corresponding to those evaluated in Study BNT162-01) were evaluated. Study vaccine was administered using the same 2-dose schedule as in Study BNT162-01 (21 days apart). Dose levels were administered first to an 18- to 55-year age cohort, then to a 65- to 85-year age cohort.

Both vaccine candidate constructs were safe and well tolerated. BNT162b2 at the $30-\mu g$ dose level was selected and advanced to the Phase 2/3 expanded cohort and efficacy evaluation primarily because:

- the reactogenicity profile for BNT162b2 was more favorable than BNT162b1 in both younger and older adults with similar immunogenicity results;
- in the NHP challenge study (VR-VTR-10671, see <u>Section 2.1.1Section 2.1.1</u>), a trend toward earlier clearance of BNT162b2 was observed in the nose.
- Phase 2 of the study (for which (enrollment has completed) comprised the evaluation of safety and immunogenicity data for the first 360 participants (180 from the active vaccine group and 180 from the placebo group, with each group divided between the younger and older age cohorts) entering the study after completion of Phase 1.
- The Phase 3 part of the study (which is ongoing) evaluates the efficacy and safety in all participants (including the first 360 participants from Phase 2). Phase 3 introduced:
 - enrollment of participants 16 to 17 years of age to be evaluated with the 18- to 55year-old cohort,
 - enrollment of a 12- to 15-year-old cohort,
 - immunogenicity data from the 12- to 15-year-old cohort (

Table 3. Exposure to BNT162b2 by Age Group and Dose (C4591001) – Blinded Placebo-Controlled Follow-up Period

- , Table 5, Table 11, Table 13, Table 15, and Table 17), anticipated to bridge to the 16-to 25-year-old cohort.
- Booster groups were subsequently added to evaluate boostability and protection against variant virus strains.

Participants in the pivotal study were initially planned to be followed for up to 24 months in order to assess the potential for late-occurring adverse reactions, such as the theoretical risk of VAED including VAERD. After completing the final efficacy analysis with vaccine efficacy shown to be 95% and obtaining regulatory authorization to vaccinate in many countries, Pfizer-BioNTech started to unblind all participants to determine those participants randomised to placebo so that they could be offered vaccine in accordance with local authorization. To date, most placebo subjects have been unblinded to receive active vaccine at or prior to 6 months after the second dose, therefore, a placebo group for comparison of safety data is only available for up to 6 months post Dose 2.

The initial efficacy analysis on the 16 years and older population was event-driven, with prespecified interim analyses after accrual of at least 62, 92, and 120 cases and a final analysis at 164 cases.

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A further efficacy analysis has been conducted on 12- to \leq 15-year-old cohort participants and on 16 years and older participants cohort participants reported by 13 March 2021.

For evaluation of booster effects and/or protection against emerging SARS CoV 2 variants of concern, approximately 600 existing Phase 3 participants 18 to 55 years of age were randomized 1:1 to receive either receive a booster (Dose 3) at 30 μ g of either BNT162b2 or a prototype based upon the B.1.351 (Beta) variant that originated in South Africa, BNT162b2_{SAT} approximately 6 months after their second dose of BNT162b2.

Other ongoing¹ BNT162b2<u>interventional</u> studies at the cut-off of the clinical database (17 June02 September 2021) also include:

- C4591005: A phase 1/2 study, <u>placebo-controlled</u>, <u>randomized</u>, <u>and observer-blind study</u> to evaluate the safety, tolerability, and immunogenicity of <u>ana SARS-CoV-2</u> RNA vaccine candidate against COVID-19 in healthy Japanese adults.
 One hundred sixty participants were randomly assigned in a 3:1 ratio to study intervention (candidate vaccine: 120, placebo: 40).
- <u>PASS:</u> C4591015: A phase 2/3 study, placebo-controlled, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of SARS-CoV-2 RNA vaccine candidate (BNT162b2) against COVID-19 in healthy pregnant women 18 years of age and older.

Approximately 4000 pregnant women at 24 to 34 weeks gestation are being randomized in a 1:1 ratio to vaccine or placebo.

- C4591017: A phase 3, randomized, observer blind study to evaluate the safety, tolerability, and immunogenicity of multiple production lots and dose levels of BNT162b2 against COVID 19 in healthy participants 12 through 50 years of age and the safety tolerability, and immunogenicity of BNT162b2 RNA based COVID 19 vaccine candidates as a booster dose in healthy participants 18 through 50 years of age..
 - Approximately 340 participants were randomly assigned to each of 3 US lots and to a 20 μ g arm and approximately 170 participants were randomly assigned an EU lot, for a total of approximately 1530 randomized participants in 5 study arms. Protocol amendment 2 has added a booster study in which a subset of the adult participants (18 through 50 years of age) who each received two 30 μ g doses of the designated US lot(s) will be randomly assigned to 1 of 2 arms in a 1:1 ratio (Booster 1: Booster 2), where Booster 1 is BNT162b2 at 30 μ g and Booster 2 is BNT162b2.B.1.351 at 30 μ g. The third dose will be administered approximately 3 months after BNT162b2 Dose 2 of the primary study.
- C4591007: A phase 1, open-label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer-blinded safety, tolerability,

¹ Study C4591017 was completed and therefore is removed from this list.

and immunogenicity study of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy children <12 years of age.

Phase 1 is an open-label dose-finding study that consists of up to 3 different dose levels in each age group, with 16 participants per dose level (total of 144 participants). Phase 2/3 will evaluate the safety, tolerability, and immunogenicity of the selected dose level in each age group from Phase 1, with a total of approximately 4500 participants. Participants will be randomized in a 2:1 ratio to receive active vaccine or placebo.

- C4591020: A phase 3, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of multiple formulations of the vaccine candidate BNT-162B2 against COVID-19 in healthy adults 18 through 55 years of age.
- C4591031 A phase 3 master protocol to evaluate additional dose(s) of BNT162B2 in healthy individuals previously vaccinated with BNT162B2.
- BNT162-01 A multi-site, phase I/II, 2-Part, dose-escalation trial investigating the safety and immunogenicity of four prophylactic SARS-CoV-2 RNA vaccines against COVID-19 using different dosing regimens in healthy and immunocompromised adults.
- BNT162-03² Safety and immunogenicity of SARS-CoV-2 mRNA vaccine (BNT162b1) in Chinese healthy subjects: A phase I, randomized, placebo- controlled, observer-blind study.
- BNT162-04 A multi-site, phase I/II, 2-part, dose escalation trial investigating the safety and immunogenicity of a prophylactic SARS-CoV-2 RNA vaccine (BNT162b3) against COVID-19 using different dosing regimens in healthy adults.
- BNT162-06² Safety and immunogenicity of SARS-CoV-2 mRNA vaccine (BNT162b2) in Chinese healthy population: A phase II, randomized, placebo-controlled, observer-blind <u>study</u>
- BNT162-14 A Phase II, open-label, rollover trial to evaluate the safety and immunogenicity of one or two boosting doses of Comirnaty or one dose of BNT162b2s01 in BNT162-01 trial subjects, or two boosting doses of Comirnaty in BNT162-04 trial subjects
- BNT162-17 A Phase II trial to evaluate the safety and immunogenicity of a SARS-CoV-2
 <u>multivalent RNA vaccine in healthy subjects.</u>

Clinical Trial Exposure

Population for analysis of CTs data in this US Pharmacovigilance Plan includes the following 2 studies:

² This study is conducted by Shanghai Fosun Pharmaceutical Development, Inc. and sponsored by BioNTech SE.

- C4591001: Phase 1/2/3, placebo-controlled, randomized, observer-blind, dose finding, study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.
- BNT162-01: A multi-site, phase I/II, 2-part, dose-escalation trial investigating the safety and immunogenicity of four prophylactic SARS-CoV-2 RNA vaccines against COVID-19 using different dosing regimens in healthy adults.

Participants 16 years of age and older

At the cut-off date of 13 March 2021, a total of 46,505 participants were vaccinated in the BNT162b2 clinical development program:

- 21,745 participants received 2 doses and 360 received 1 dose of BNT162b2 during the blinded follow-up period; 96 participants from study BNT162-01 received 2 doses of the vaccine.
- 19,647 participants, who originally received placebo, then received 1 dose of BNT162b2 in the Open-Label Follow-up period after unblinding. (none from study BNT162-01).

Exposure to BNT162b2 for participants aged 16 years and older in the 2 ongoing studies by number of doses, and demographic characteristics is shown in

Table 3.Exposure to BNT162b2 by Age Group and Dose (C4591001) – Blinded
Placebo-Controlled Follow-up Period

through Table 21.

In addition, exposure in clinical studies in special populations is provided in Table 22 and Table 23.

Participants 12 to 15 years of age

- At the cut-off date of 13 March 2021, a total of 2260 participants <u>12 to 15 years of age</u> were vaccinated in the BNT162b2 clinical development program÷<u>(study</u> <u>C4591001)</u>.
- Clinical At the cut-off date of 02 September 2021, updated clinical study exposure data for the 12- to 15 years of age are provided for the ongoing study C4591001-at the cut off date of 13 March 2021.

In this study

 1124<u>One thousand one hundred twenty-four (1124)</u> participants received 2 doses and 7 received 1 dose of BNT162b2 in the Blinded-Placebo Controlled Followup period.

> • <u>49One thousand and ten (1010)</u> participants who originally received placebo, then received 1 dose of BNT162b2<u>(18) or 2 doses (992)</u> in the Open-Label Follow-up period after unblinding.

Exposure to BNT162b2 for participants aged 12- to 15 years of age by number of doses and demographic characteristics is shown in

Table 3. Exposure to BNT162b2 by Age Group and Dose (C4591001) – Blinded Placebo-Controlled Follow-up Period

,Table 5, Table 11, Table 13, Table 15, Table 17-<u>(at the cut-off date of 13 March 2021)</u> and in Table 24, Table 25, Table 26, <u>Table 27</u>, <u>Table 28</u>, and <u>Table 29 (at the cut-off date of 02 September 2021)</u>. In addition, exposure in clinical studies in special populations is provided in Table 22 and <u>Table 23</u>Table 23, <u>Table 30 and Table 31</u>.

Booster dose

Exposure in participants 1812 years of age and older

At the cut <u>(Studies C4591001 – Cut-off date of 17 June13 March</u> 2021, a total of 306 participants, 18 to 55 years of age, received BNT162b2 30 µg as booster dose (Dose 3); these participant, were originally randomized in the Phase 3 study and completed the BNT162b2 (30 µg) two dose series, and then received a third dose of BNT162b2 (30 µg) approximately 6 months after receipt of Dose 2, with safety and immune response evaluations at 1 month after Dose 3.<u>BNT162-01 – Cut off date 23 October 2020</u>)

Exposure to BNT162b2 for participants who received the booster dose, by number of doses and demographic characteristics, is shown in Table 24 and Table 25. In addition, exposure in special population who received the booster dose is provided in Table 26.

BNT162b2			
1.16 Risk Management Plan (1	Non-REMS) for Biologic	License Applicati	ion # 125742

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
\geq 12 years to \leq 15 years		
Vaccine 30 µg		
1 Dose	7	7
2 Doses	1124	2248
Total	1131	2255
≥ 16 years to ≤ 17 years		
Vaccine 30 µg		
1 Dose	4	4
2 Doses	374	748
Total	378	752
\geq 18 years to \leq 55 years		
Vaccine 10 µg		
2 Doses	12	24
Total	12	24
	12	21
Vaccine 20 µg 2 Doses	12	24
Z Doses Total	12	24 24
	12	24
Vaccine 30 µg	267	267
1 Dose 2 Doses	267 12438	267 24876
2 Doses Total	12438	24876
	12705	25145
>55 years to ≤64 years		
Vaccine 30 µg	67	(7
1 Dose	67	67
2 Doses	4341	8682
Total	4408	8749
\geq 65 years to \leq 74 years		
Vaccine 10 µg	10	
2 Doses	12	24
Total	12	24
Vaccine 20 µg		
2 Doses	9	18
Total	9	18
Vaccine 30 µg		
1 Dose	17	17
2 Doses	3624	7248
Total	3641	7265
≥75 years to ≤84 years		

Table 3. Exposure to BNT162b2 by Age Group and Dose (C4591001) – Blinded

BNT162b2		
1.16 Risk Management Plan	(Non-REMS) for Biologic License Application # 12	25742

Table 3.	Exposure to BNT162b2 by Age Group and Dose (C4591001) – Blinded
	Placebo-Controlled Follow-up Period

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Vaccine 20 µg		
2 Doses	3	6
Total	3	6
Vaccine 30 µg		
1 Dose	3	3
2 Doses	899	1798
Total	902	1801
≥85 years		
Vaccine 30 µg		
1 Dose	2	2
2 Doses	21	42
Total	23	44

Note: 30 µg includes data from phase 1 and phase 2/3.

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:42)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

./nda2 unblinded/C4591001 PVP BLA/adsl s912

Table 4.Exposure to BNT162b2 by Age Group and Dose (C4591001) – Open-Label
Follow-up Period – Subjects Who Originally Received BNT162b2

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
≥16 years to ≤17 years		
Vaccine 30 µg		
1 Dose	3	3
≥18 years to ≤55 years		
Vaccine 30 µg		
1 Dose	58	58
>55 years to ≤64 years		
Vaccine 30 µg		
1 Dose	17	17
≥65 years to ≤74 years		
Vaccine 30 µg		
1 Dose	8	8

Table 4.Exposure to BNT162b2 by Age Group and Dose (C4591001) – Open-Label
Follow-up Period – Subjects Who Originally Received BNT162b2

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
≥75 years to ≤84 years		
Vaccine 30 µg		
1 Dose	1	1
≥85 years		
Vaccine 30 µg		
1 Dose	2	2

27MAR2021 (12:46)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/adsl s9123

Table 5.Exposure to BNT162b2 by Age Group and Dose (C4591001) – Open-Label
Follow-up Period – Subjects Who Originally Received Placebo and Then
Received BNT162b2 After Unblinding

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
≥12 years to ≤15 years ^a		
Vaccine 30 µg		
1 Dose	30	30
2 Doses	19	38
Total	49	68
≥ 16 years to ≤ 17 years		
Vaccine 30 µg		
1 Dose	107	107
2 Doses	186	372
Total	293	479
≥18 years to ≤55 years		
Vaccine 30 µg		
1 Dose	2713	2713
2 Doses	8419	16838
Total	11132	19551
>55 years to ≤64 years		
Vaccine 30 µg		
1 Dose	655	655
2 Doses	3330	6660

BNT162b2
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742

Exposure to BNT162b2 by Age Group and Dose (C4591001) - Open-Label Table 5. Follow-up Period - Subjects Who Originally Received Placebo and Then **Received BNT162b2 After Unblinding**

Age Group Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Total	3985	7315
≥65 years to ≤74 years		
Vaccine 30 µg		
1 Dose	128	128
2 Doses	3286	6572
Total	3414	6700
≥75 years to ≤84 years		
Vaccine 30 µg		
1 Dose	23	23
2 Doses	783	1566
Total	806	1589
≥85 years		
Vaccine 30 µg		
1 Dose	1	1
2 Doses	16	32
Total	17	33

Note: 30 μ g includes data from phase 1 and phase 2/3.

a. Includes subjects who became eligible for unblinding at 16 years of age, confirmed to have received

placebo originally and then received BNT162b2 post unblinding. PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/ads1 s9122

Table 6. Exposure to BNT162b2 by Age Group and Dose (BNT162-01)

Age Group Dose Exposure (Number of Doses Received)	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
≥18 years to ≤64 years		
Vaccine 1 µg		
1 Dose	1	1
2 Doses	11	22
Total	12	23

BNT162b2
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742

Age Group Dose Exposure (Number of Doses Received)	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
1 Dose	0	0
2 Doses	12	24
Total	12	24
Vaccine 10 µg		
1 Dose	1	1
2 Doses	11	22
Total	12	23
Vaccine 20 µg		
1 Dose	0	0
2 Doses	17	34
Total	17	34
Vaccine 30 µg		
1 Dose	0	0
2 Doses	18	36
Total	18	36
≥65 years to ≤74 years		
Vaccine 1 µg		
1 Dose	0	0
2 Doses	0	0
Total	0	0
Vaccine 3 µg		
1 Dose	0	0
2 Doses	0	0
Total	0	0
Vaccine 10 µg		
1 Dose	0	0
2 Doses	5	10

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BNT162b2	
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age Group Dose Exposure (Number of Doses Received)	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
Total	5	10
Vaccine 20 µg		
1 Dose	0	0
2 Doses	6	12
Total	6	12
Vaccine 30 µg		
1 Dose	0	0
2 Doses	6	12
Total	6	12
75 years to ≤84 years		
Vaccine 1 µg		
Dose	0	0
2 Doses	0	0
Fotal	0	0
Vaccine 3 µg		
Dose	0	0
2 Doses	0	0
Fotal	0	0
Vaccine 10 µg		
1 Dose	0	0
2 Doses	1	2
Total	1	2
/accine 20 μg		
1 Dose	0	0
2 Doses	1	2
Total	1	2
Vaccine 30 µg		
1 Dose	0	0
2 Doses	0	0
Total	0	0

PFIZER CONFIDENTIAL SDTM Creation: 24NOV2020 (15:06) Source Data: adsl Table Generation: 10MAR2021 (11:32) (Cutoff date:23OCT2020, Snapshot Date: 23OCT2020) Output File: ex_b2_age_dose2 rtf

Table 7.	Exposure to BNT162b2 by Dose (Totals) (C4591001) – Blinded Placebo-
	Controlled Follow-up Period

Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Vaccine 10 µg		
2 Doses	24	48
Total	24	48
Vaccine 20 µg		
2 Doses	24	48
Total	24	48
Vaccine 30 µg		
1 Dose	367	367
2 Doses	22821	45642
Total	23188	46009

Note: 30 µg includes data from phase 1 and phase 2/3.

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/ads1 s922

Table 8. Exposure to BNT162b2 by Dose (Totals) (C4591001) – Open-Label Followup Period – Subjects Who Originally Received BNT162b2

Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Vaccine 30 µg			
1 Dose	89	89	
Note: 30 μg includes data from phase 1 and phase 2 Note: Subjects who received 2 nd Dose of BNT162b/ PFIZER CONFIDENTIAL SDTM Creation: 25MA 27MAR2021 (12:46) (Cutoff Date: 13MAR2021, Snapshot Date: 25MAF ./nda2 unblinded/C4591001 PVP BLA/adsl s922;	2 after unblinding. R2021 (23:24) Source Data: adsl Ta R2021) Output File:	able Generation:	

Table 9. Exposure to BNT162b2 by Dose (Totals) (C4591001) – Open-Label Followup Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Vaccine 30 µg			
1 Dose	3657	3657	
2 Doses	16039	32078	
Total	19696	35735	

BNT162b2					
1.16 Risk Management Plan	(Non-REMS)	for Biologic	License Ap	plication #	125742

Exposure to BNT162b2 by Dose (Totals) (C4591001) - Open-Label Follow-Table 9. up Period - Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

Dose	Number of Subjects	Total Number of
Exposure (Number of Doses Received)	Exposed to BNT162b2	Vaccine Doses

Note: 30 µg includes data from phase 1 and phase 2/3.

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46) (Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/ads1 s9222

Dose Exposure (Number of Doses Received)	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
Vaccine 1 µg		
1 Dose	1	1
2 Doses	11	22
Total	12	23
Vaccine 3 µg		
1 Dose	0	0
2 Doses	12	24
Total	12	24
Vaccine 10 µg		
1 Dose	1	1
2 Doses	23	46
Total	24	47
Vaccine 20 µg		
1 Dose	0	0
2 Doses	24	48
Total	24	48
Vaccine 30 µg		
1 Dose	0	0
2 Doses	24	48
Total	24	48

Table 10. Exposure to BNT162b2 by Dose (Totals) (BNT162-01)

BNT162b2	
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Table 10. Exposure to BNT162b2 by Dose (Totals) (BNT162-01)

Dose Exposure (Number of Doses Received)	No. of Subjects Exposed to	Total No. of Vaccine Doses
	BNT162b2	

PFIZER CONFIDENTIAL SDTM Creation: 24NOV2020 (15:06) Source Data: adsl Table Generation: 10MAR2021 (11:49) (Cutoff date:23OCT2020, Snapshot Date: 23OCT2020)

Output File: ex_b2_dose rtf

		Number of Subjects Exposed to BNT162b2		ber of Vaccine loses
Dose	Male	Female	Male	Female
Age Group				
Vaccine 10 µg				
≥ 18 years to ≤ 55 years	5	7	10	14
\geq 65 years to \leq 74 years	2	10	4	20
Total	7	17	14	34
Vaccine 20 µg				
≥ 18 years to ≤ 55 years	6	6	12	12
≥65 years to ≤74 years	4	5	8	10
≥75 years to ≤84 years	1	2	2	4
Total	11	13	22	26
Vaccine 30 µg				
≥ 12 years to ≤ 15 years	567	564	1128	1127
≥ 16 years to ≤ 17 years	187	191	373	379
≥ 18 years to ≤ 55 years	6456	6249	12770	12373
>55 years to ≤64 years	2231	2177	4421	4328
\geq 65 years to \leq 74 years	1934	1707	3858	3407
\geq 75 years to \leq 84 years	511	391	1020	781
≥85 years	12	11	23	21
Total	11898	11290	23593	22416

Note: 30 μg includes data from phase 1 and phase 2/3. PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File: ./nda2 unblinded/C4591001 PVP BLA/adsl s932

Table 12. Exposure to BNT162b2 by Dose, Age Group, and Gender (C4591001) -**Open-Label Follow-up Period – Subjects Who Originally Received** BNT162b2

		Number of Subjects Exposed to BNT162b2		ber of Vaccine Doses
Dose Age Group	Male	Female	Male	Female
Vaccine 30 µg				
≥ 16 years to ≤ 17 years	0	3	0	3
≥ 18 years to ≤ 55 years	24	34	24	34
>55 years to ≤64 years	12	5	12	5
\geq 65 years to \leq 74 years	4	4	4	4
≥75 years to ≤84 years	0	1	0	1
≥85 years	1	1	1	1
Total	41	48	41	48

Note: 30 μg includes data from phase 1 and phase 2/3. Note: Subjects who received 2nd Dose of BNT162b2 after unblinding. PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46) (Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

./nda2 unblinded/C4591001 PVP BLA/adsl s9323

Table 13. Exposure to BNT162b2 by Dose, Age Group, and Gender (C4591001) -**Open-Label Follow-up Period – Subjects Who Originally Received Placebo** and Then Received BNT162b2 After Unblinding

		Number of Subjects Exposed to BNT162b2		ber of Vaccine oses
Dose Age Group	Male	Female	Male	Female
Vaccine 30 µg				
≥ 12 years to ≤ 15 years ^a	26	23	36	32
≥ 16 years to ≤ 17 years	152	141	250	229
≥ 18 years to ≤ 55 years	5424	5708	9450	10101
>55 years to ≤64 years	1973	2012	3602	3713
≥65 years to ≤74 years	1801	1613	3530	3170
\geq 75 years to \leq 84 years	495	311	976	613
≥85 years	13	4	25	8
Total	9884	9812	17869	17866

Note: 30 µg includes data from phase 1 and phase 2/3.

a. Includes subjects who became eligible for unblinding at 16 years of age, confirmed to have received placebo originally and then received BNT162b2 post unblinding.

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46)

BNT162b2	
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 12	5742

Table 13. Exposure to BNT162b2 by Dose, Age Group, and Gender (C4591001) – Open-Label Follow-up Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

		Number of Subjects Exposed to BNT162b2		Total Number of Vaccine Doses	
Dose Age Group	Male	Female	Male	Female	

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File: ./nda2_unblinded/C4591001_PVP_BLA/adsl_s932_open

	No. of Subjects Ex BNT162b2				
Dose Age Group	Male	Female	Male	Female	
Vaccine 1 µg			I		
≥ 18 years to ≤ 64 years	7	5	14	9	
≥65 years to ≤74 years	0	0	0	0	
≥75 years to ≤84 years	0	0	0	0	
Total	7	5	14	9	
Vaccine 3 µg					
≥18 years to ≤64 years	5	7	10	14	
≥65 years to ≤74 years	0	0	0	0	
≥75 years to ≤84 years	0	0	0	0	
Total	5	7	10	14	
Vaccine 10 µg					
≥18 years to ≤64 years	8	10	16	19	
≥65 years to ≤74 years	3	2	6	4	
≥75 years to ≤84 years	1	0	2	0	
Total	12	12	24	23	
Vaccine 20 µg					
≥18 years to ≤64 years	7	10	14	20	
≥65 years to ≤74 years	1	5	2	10	
≥75 years to ≤84 years	0	1	0	2	
Total	8	16	16	32	
Vaccine 30 µg					
≥18 years to ≤64 years	10	8	20	16	
≥ 65 years to ≤ 74 years	2	4	4	8	
\geq 75 years to \leq 84 years	0	0	0	0	

 Table 14.
 Exposure to BNT162b2 by Dose, Age Group, and Gender (BNT162-01)

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BNT162b2					
1.16 Risk Management Plan	Non-REMS) for Biologic L	license Ap	plication #	125742

Table 14. Exposure to BNT162b2 by Dose, Age Group, and Gender (BNT162-01)

	No. of Subjects Exposed to BNT162b2 Total No. of Va		No. of Vaccine Doses	
Dose Age Group	Male	Female	Male	Female
Total	12	12	24	24

PFIZER CONFIDENTIAL SDTM Creation: 24NOV2020 (15:06) Source Data: adsl Table Generation: 10MAR2021 (11:53) (Cutoff date:23OCT2020, Snapshot Date: 23OCT2020)

Output File: ex_b2_age_dose_sex rtf

Table 15.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) – Blinded Placebo-Controlled Follow-up Period

Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
\geq 12 years to \leq 15 years		
Vaccine 30 µg		
Racial origin		
White	971	1937
Black or African American	52	103
Asian	72	143
American Indian or Alaska Native	4	8
Native Hawaiian or other Pacific Islander	3	6
Multiracial	23	46
Not reported	6	12
Total	1131	2255
Ethnic origin		
Hispanic/Latino	132	263
Non-Hispanic/non-Latino	997	1988
Not reported	2	4
Total	1131	2255
≥ 16 years to ≤ 17 years		
Vaccine 30 µg		
Racial origin		
White	309	614
Black or African American	30	60
Asian	22	44
American Indian or Alaska Native	4	8
Native Hawaiian or other Pacific Islander	3	6
Multiracial	10	20
Total	378	752
Ethnic origin		
Hispanic/Latino	49	98

1 16 Risk Management Plan (Non-PEMS) for Biologic License Application # 125742	BNT162b2
1.10 Kisk Management I fan (Non-KEWIS) for Biologie Electise Application # 125742	1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742

Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Non-Hispanic/non-Latino	329	654
Total	378	752
\geq 18 years to \leq 55 years		
Vaccine 10 µg		
Racial origin		
White	11	22
Asian	1	2
Total	12	24
Ethnic origin		
Hispanic/Latino	1	2
Non-Hispanic/non-Latino	11	22
Total	12	24
Vaccine 20 µg		
Racial origin		
White	10	20
Black or African American	2	4
Total	12	24
Ethnic origin		
Hispanic/Latino	1	2
Non-Hispanic/non-Latino	11	22
Total	12	24
Vaccine 30 µg		
Racial origin		
White	9923	19637
Black or African American	1400	2764
Asian	683	1358
American Indian or Alaska Native	161	311
Native Hawaiian or other Pacific Islander	40	80
Multiracial	427	851
Not reported	71	142
Total	12705	25143
Ethnic origin		
Hispanic/Latino	4000	7874
Non-Hispanic/non-Latino	8650	17160
Not reported	55	109
Total	12705	25143
>55 years to ≤64 years		
Vaccine 30 µg		
Racial origin		

BNT162b2	
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742	!

Table 15.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) - Blinded Placebo-Controlled Follow-up Period

Age Group Dose	Number of Subjects	Total Number of Vaccine Doses
Race/Ethnic Origin	Exposed to BNT162b2	vaccine Doses
White	3719	7388
Black or African American	430	849
Asian	135	267
American Indian or Alaska Native	30	58
Native Hawaiian or other Pacific Islander	8	15
Multiracial	76	152
Not reported	10	20
Total	4408	8749
Ethnic origin		
Hispanic/Latino	965	1903
Non-Hispanic/non-Latino	3413	6786
Not reported	30	60
Total	4408	8749
≥65 years to ≤74 years		
Vaccine 10 µg		
Racial origin		
White	12	24
Total	12	24
Ethnic origin		
Non-Hispanic/non-Latino	12	24
Total	12	24
Vaccine 20 µg		
Racial origin		
White	9	18
Total	9	18
Ethnic origin		
Non-Hispanic/non-Latino	9	18
Total	9	18
Vaccine 30 µg		
Racial origin		
White	3272	6528
Black or African American	219	437
Asian	82	164
American Indian or Alaska Native	22	44
Native Hawaiian or other Pacific Islander	6	12
Multiracial	30	60
Not reported	10	20
Total	3641	7265
Ethnic origin		

BNT162b2		
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Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Hispanic/Latino	583	1158
Non-Hispanic/non-Latino	3038	6067
Not reported	20	40
Total	3641	7265
≥75 years to ≤84 years		
Vaccine 20 µg		
Racial origin		
White	3	6
Total	3	6
Ethnic origin		
Non-Hispanic/non-Latino	3	6
Total	3	6
Vaccine 30 µg		
Racial origin		
White	838	1673
Black or African American	22	44
Asian	31	62
American Indian or Alaska Native	3	6
Native Hawaiian or other Pacific Islander	1	2
Multiracial	7	14
Total	902	1801
Ethnic origin		
Hispanic/Latino	107	213
Non-Hispanic/non-Latino	789	1576
Not reported	6	12
Total	902	1801
≥85 years		
Vaccine 30 µg		
Racial origin		
White	20	38
Asian	1	2
American Indian or Alaska Native	1	2
Multiracial	1	2
Total	23	44
Ethnic origin		
Hispanic/Latino	2	4
Non-Hispanic/non-Latino	21	40
Total	23	44

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Table 15. Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin (C4591001) – Blinded Placebo-Controlled Follow-up Period

Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
27MAR2021 (12:46)	ation: 25MAR2021 (23:24) Source Data: adsl	Table Generation:
(Cutoff Date: 13MAR2021, Snapshot L./nda2 unblinded/C4591001 PVP BL		

Table 16. Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin (C4591001) – Open-Label Follow-up Period – Subjects Who Originally Received BNT162b2

Age Group	Number of Subjects	Total Number of
Dose Race/Ethnic Origin	Exposed to BNT162b2	Vaccine Doses
≥ 16 years to ≤ 17 years		
Vaccine 30 µg		
Racial origin		
White	3	3
Total	3	3
Ethnic origin		
Non-Hispanic/non-Latino	3	3
Total	3	3
≥ 18 years to ≤ 55 years		
Vaccine 30 µg		
Racial origin		
White	46	46
Black or African American	2	2
Asian	2	2
American Indian or Alaska Native	8	8
Total	58	58
Ethnic origin		
Hispanic/Latino	31	31
Non-Hispanic/non-Latino	27	27
Total	58	58
>55 years to ≤64 years		
Vaccine 30 µg		
Racial origin		
White	14	14
Asian	1	1
American Indian or Alaska Native	2	2
Total	17	17

Table 16.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) – Open-Label Follow-up Period – Subjects Who Originally
	Received BNT162b2

Age Group Dose	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Race/Ethnic Origin			
Ethnic origin			
Hispanic/Latino	10	10	
Non-Hispanic/non-Latino	7	7	
Total	17	17	
≥65 years to ≤74 years			
Vaccine 30 µg			
Racial origin			
White	8	8	
Total	8	8	
Ethnic origin			
Hispanic/Latino	5	5	
Non-Hispanic/non-Latino	3	3	
Total	8	8	
≥75 years to ≤84 years			
Vaccine 30 µg			
Racial origin			
White	1	1	
Total	1	1	
Ethnic origin			
Non-Hispanic/non-Latino	1	1	
Total	1	1	
≥85 years			
Vaccine 30 µg			
Racial origin			
White	2	2	
Total	2	2	
Ethnic origin			
Non-Hispanic/non-Latino	2	2	
Total	2	2	

Note: Subjects who received 2nd Dose of BNT162b2 after unblinding. PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File: ./nda2 unblinded/C4591001 PVP BLA/adsl s9423

BNT162b2		
1.16 Risk Management Plan (Non-F	REMS) for Biologic License	Application # 125742

Table 17.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) – Open-Label Follow-up Period – Subjects Who Originally
	Received Placebo and Then Received BNT162b2 After Unblinding

Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
\geq 12 years to \leq 15 years ^a			
Vaccine 30 µg			
Racial origin			
White	45	62	
Asian	3	5	
Multiracial	1	1	
Total	49	68	
Ethnic origin			
Hispanic/Latino	2	4	
Non-Hispanic/non-Latino	47	64	
Total	49	68	
≥ 16 years to ≤ 17 years			
Vaccine 30 µg			
Racial origin			
White	251	410	
Black or African American	11	19	
Asian	14	25	
American Indian or Alaska Native	2	4	
Native Hawaiian or other Pacific Islander	1	2	
Multiracial	12	16	
Not reported	2	3	
Total	293	479	
Ethnic origin			
Hispanic/Latino	26	43	
Non-Hispanic/non-Latino	266	434	
Not reported	1	2	
Total	293	479	
≥ 18 years to ≤ 55 years			
Vaccine 30 µg			
Racial origin			
White	8806	15340	
Black or African American	1087	1899	
Asian	619	1136	
American Indian or Alaska Native	128	236	
Native Hawaiian or other Pacific Islander	17	32	
Multiracial	405	781	
Not reported	70	127	
Total	11132	19551	
Ethnic origin			

BNT162b2					
1.16 Risk Management Plan	Non-REMS) for Biologic L	icense App	plication #	125742

Table 17.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) – Open-Label Follow-up Period – Subjects Who Originally
	Received Placebo and Then Received BNT162b2 After Unblinding

Age Group Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Hispanic/Latino	3441	5300	
Non-Hispanic/non-Latino	7635	14157	
Not reported	56	94	
Total	11132	19551	
>55 years to ≤64 years			
Vaccine 30 µg			
Racial origin			
White	3416	6271	
Black or African American	331	592	
Asian	120	227	
American Indian or Alaska Native	35	67	
Native Hawaiian or other Pacific Islander	4	7	
Multiracial	63	120	
Not reported	16	31	
Total	3985	7315	
Ethnic origin			
Hispanic/Latino	901	1560	
Non-Hispanic/non-Latino	3067	5724	
Not reported	17	31	
Total	3985	7315	
≥65 years to ≤74 years			
Vaccine 30 µg			
Racial origin			
White	3093	6076	
Black or African American	187	360	
Asian	78	154	
American Indian or Alaska Native	20	39	
Native Hawaiian or other Pacific Islander	6	12	
Multiracial	22	43	
Not reported	8	16	
Total	3414	6700	
Ethnic origin			
Hispanic/Latino	547	1060	
Non-Hispanic/non-Latino	2842	5590	
Not reported	25	50	
Total	3414	6700	
≥75 years to ≤84 years Vaccine 30 µg			

BNT162b2					
1.16 Risk Management Plan	Non-REMS) for Biologic L	license Ap	plication #	125742

Table 17.	Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnic Origin
	(C4591001) – Open-Label Follow-up Period – Subjects Who Originally
	Received Placebo and Then Received BNT162b2 After Unblinding

Age Group	Number of Subjects	Total Number of Vaccine Doses
Dose Race/Ethnic Origin	Exposed to BNT162b2 Vacc	
Racial origin		
White	752	1483
Black or African American	22	42
Asian	17	34
American Indian or Alaska Native	4	8
Multiracial	6	12
Not reported	5	10
Total	806	1589
Ethnic origin		
Hispanic/Latino	89	174
Non-Hispanic/non-Latino	706	1393
Not reported	11	22
Total	806	1589
≥85 years		
Vaccine 30 µg		
Racial origin		
White	15	29
Asian	1	2
Multiracial	1	2
Total	17	33
Ethnic origin		
Non-Hispanic/non-Latino	17	33
Total	17	33

Note: 30 μ g includes data from phase 1 and phase 2/3.

a. Includes subjects who became eligible for unblinding at 16 years of age, confirmed to have received placebo originally and then received BNT162b2 post unblinding.
 PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation:

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:46) (Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

./nda2 unblinded/C4591001 PVP BLA/ads1 s942 open

Table 18. Exposure to BNT162b2 by Dose and Race/Ethnic Origin (C4591001) – Blinded Placebo-Controlled Follow-up Period

Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses
Vaccine 10 µg		
Racial origin		
White	23	46

BNT162b2					
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Table 18.	Exposure to BNT162b2 by Dose and Race/Ethnic Origin (C4591001) –
	Blinded Placebo-Controlled Follow-up Period

Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Asian	1	2	
Total	24	48	
Ethnic origin			
Hispanic/Latino	1	2	
Non-Hispanic/non-Latino	23	46	
Total	24	48	
Vaccine 20 µg			
Racial origin			
White	22	44	
Black or African American	2	4	
Total	24	48	
Ethnic origin			
Hispanic/Latino	1	2	
Non-Hispanic/non-Latino	23	46	
Total	24	48	
Vaccine 30 µg			
Racial origin			
White	19052	37815	
Black or African American	2153	4257	
Asian	1026	2040	
American Indian or Alaska Native	225	437	
Native Hawaiian or other Pacific Islander	61	121	
Multiracial	574	1145	
Not reported	97	194	
Total	23188	46009	
Ethnic origin			
Hispanic/Latino	5838	11513	
Non-Hispanic/non-Latino	17237	34271	
Not reported	113	225	
Total	23188	46009	

Note: 30 µg includes data from phase 1 and phase 2/3.

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:47)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

./nda2_unblinded/C4591001_PVP_BLA/ads1_s952

BNT162b2				
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Table 19. Exposure to BNT162b2 by Dose and Race/Ethnic Origin (C4591001) -Open-Label Follow-up Period - Subjects Who Originally Received BNT162b2

Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Vaccine 30 µg			
Racial origin			
White	74	74	
Black or African American	2	2	
Asian	3	3	
American Indian or Alaska Native	10	10	
Total	89	89	
Ethnic origin			
Hispanic/Latino	46	46	
Non-Hispanic/non-Latino	43	43	
Total	89	89	

Note: 30 μg includes data from phase 1 and phase 2/3. Note: Subjects who received 2nd Dose of BNT162b2 after unblinding. PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (23:24) Source Data: adsl Table Generation: 27MAR2021 (12:47)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File: /nda2 unblinded/C4591001 PVP BLA/adsl s9523

Table 20. Exposure to BNT162b2 by Dose and Race/Ethnic Origin (C4591001) -**Open-Label Follow-up Period – Subjects Who Originally Received Placebo** and Then Received BNT162b2 After Unblinding

Dose Race/Ethnic Origin	Number of Subjects Exposed to BNT162b2	Total Number of Vaccine Doses	
Vaccine 30 µg			
Racial origin			
White	16378	29671	
Black or African American	1638	2912	
Asian	852	1583	
American Indian or Alaska Native	189	354	
Native Hawaiian or other Pacific Islander	28	53	
Multiracial	510	975	
Not reported	101	187	
Total	19696	35735	
Ethnic origin			
Hispanic/Latino	5006	8141	
Non-Hispanic/non-Latino	14580	27395	
Not reported	110	199	
Total	19696	35735	

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Table 20. Exposure to BNT162b2 by Dose and Race/Ethnic Origin (C4591001) – Open-Label Follow-up Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

Dose	Number of Subjects	Total Number of
Race/Ethnic Origin	Exposed to BNT162b2	Vaccine Doses
Note: 30 µg includes data from phase 1 and phase PFIZER CONFIDENTIAL SDTM Creation: 25.		able Generation:

27MAR2021 (12:47)

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/adsl s952 open

Table 21.	Exposure to	BNT162b2 by D	ose and Race/Ethnic	Crigin (BNT162-01)
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Dose Race/Ethnic Origin	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
Vaccine 1 µg		
Desist Ostata		
White	12	23
Total	12	23
Ethnic Origin		
Non-Hispanic/non-Latino	12	23
Total		23
Vaccine 3 µg		
Racial Origin		
White	12	24
Total	10	24
Ethnic Origin		
Non-Hispanic/non-Latino	12	24
Total	12	24
Vaccine 10 µg		
Racial Origin		
White		47
Total	24	47
Ethnic Origin		
Non-Hispanic/non-Latino	24	47
Total	24	47
Vaccine 20 µg		
Racial Origin		
White	24	48
Total	24	48
Ethnic Origin		

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Table 21.	Exposure to BN	T162b2 by Dose and	Race/Ethnic Origin (BN	T162-01)

Dose Race/Ethnic Origin	No. of Subjects Exposed to BNT162b2	Total No. of Vaccine Doses
Non-Hispanic/non-Latino	24	48
Total	24	48
Vaccine 30 µg Racial Origin		
White	24	48
Total	24	48
Ethnic Origin		
Non-Hispanic/non-Latino	24	48
Total	24	48

Only race, ethnic origins collected on the case report form with a count of at least one in either column are displayed PFIZER CONFIDENTIAL SDTM Creation: 24NOV2020 (15:06) Source Data: adsl Table Generation: 10MAR2021 (12:27) (Cutoff date:23OCT2020, Snapshot Date: 23OCT2020)

Output File: ex_b2_dose_race rtf

Table 22. Exposure to BNT162b2 (30 µg) by Special Population (C4591001) – Blinded Placebo-Controlled Follow-up Period

Population	Number of Subjects Exposed to BNT162b2 (30 µg) (N ^a =23188) n ^b	of
Subjects with any baseline comorbidity	10371	26487
AIDS/HIV	100	196
Any Malignancy + Metastatic Solid Tumor + Leukemia + Lymphoma	852	1696
Chronic Pulmonary Disease	1901	3774
Renal Disease	140	279
Rheumatic Disease	75	147
Mild Liver Disease + Moderate or Severe Liver Disease	154	302
Cerebrovascular Disease + Peripheral Vascular Disease + Myocardial Infarction + Congestive Heart Failure	651	1298
Dementia	7	14
Diabetes With/Without Chronic Complication	1706	3385
Hemiplegia or Paraplegia	4	8
Peptic Ulcer Disease	63	126
Obese	7689	15262

Note: Comorbidity is based on Charlson Comorbidity Index categories. Participants identified as belonging to these categories were identified by medical history data collected during the study.

Note: 30 μ g includes data from phase 1 and phase 2/3.

Note: Hemiplegia or Paraplegia only includes preferred terms Hemiplegia and Paraplegia.

a. N = number of subjects in the specified group.

(Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

./nda2 unblinded/C4591001 PVP BLA/admh s953

Table 22. Exposure to BNT162b2 (30 µg) by Special Population (C4591001) – Blinded Placebo-Controlled Follow-up Period

Population	Number of Subjects Exposed to BNT162b2 (30 µg) (N ^a =23188) n ^b	of
b. n = Number of subjects reporting at least 1 occurrence of any Years of age] or BMI ≥95 th percentile [12-15 Years of age]). PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (19:25) 27MAR2021 (12:47)	• · ·	0 1

Table 23. Exposure to BNT162b2 (30 μg) by Special Population (C4591001) – Open-

Fable 23. Exposure to BN1162b2 (30 μg) by Special Population (C4591001) – Open-Label Follow-up Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

Population	Number of Subjects Exposed to BNT162b2 (30 µg) (N ^a =19696) n ^b	Total Number of Vaccine Doses
Subjects with any baseline comorbidity	8981	21590
AIDS/HIV	86	161
Any Malignancy + Metastatic Solid Tumor + Leukemia + Lymphoma	734	1406
Chronic Pulmonary Disease	1590	2953
Renal Disease	139	262
Rheumatic Disease	66	122
Mild Liver Disease + Moderate or Severe Liver Disease	102	193
Cerebrovascular Disease + Peripheral Vascular Disease + Myocardial Infarction + Congestive Heart Failure	567	1075
Dementia	9	17
Diabetes With/Without Chronic Complication	1555	2928
Hemiplegia or Paraplegia	4	8
Peptic Ulcer Disease	76	145
Obese	6760	12320

Note: Comorbidity is based on Charlson Comorbidity Index categories. Participants identified as belonging to these categories were identified by medical history data collected during the study. Note: 30 µg includes data from phase 1 and phase 2/3.

Note: Hemiplegia or Paraplegia only includes preferred terms Hemiplegia and Paraplegia.

a. N = number of subjects in the specified group.

b. n = Number of subjects reporting at least 1 occurrence of any comorbidity or obese (BMI \geq 30 kg/m² [\geq 16 Years of age] or BMI \geq 95th percentile [12-15 Years of age]).

PFIZER CONFIDENTIAL SDTM Creation: 25MAR2021 (19:25) Source Data: admh Table Generation: 27MAR2021 (12:47)

 Table 23. Exposure to BNT162b2 (30 μg) by Special Population (C4591001) – Open-Label Follow-up Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

Population Number of Subjects Total Number Exposed of to BNT162b2 (30 µg) Vaccine Doses (N²=19696) n^b (Cutoff Date: 13MAR2021, Snapshot Date: 25MAR2021) Output File:

/nda2 unblinded/C4591001 PVP BLA/admh s953 open

Exposure in participants 12-15 years of age – (Study C4591001-6-month follow-up period – Cut-off date 02 September 2021)

Table 24. Exposure to BNT162b2 by Dose, Age Group, and Gender (Totals) (C4591001) -BNT162b2 Experienced Subjects Who Were Rerandomized to Receive 1 Booster Dose of BNT162b2 (30 µg) Booster Safety Population 12-15 Years - Blinded Placebo-Controlled Follow-up Period

Dose Exposure (Number of Doses Received)	Number of Subjects Exposed to BNT162b2	<u>Total Number</u> <u>of</u> Vaccine Doses
Dose — Age Group®	Male	Female
Vaccine 30 µg		
Vaccine 30 µg 1 Dose	<u>7</u>	<u>7</u>
<u>≥18 years to ≤55 years2 Doses</u>	140<u>1124</u>	166 2248
Total	140<u>1131</u>	166 <u>2255</u>

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Note: Only phase 3 subjects who already received two doses of BNT162b2 in the original phase of the study and consented to participate the booster evaluation phase were rerandomized to receive booster dose.

a. Based on age at booster dose.

PFIZER CONFIDENTIAL SDTM Creation: 29JUL2021 (23:0130SEP2021 (11:35) Source Data: adsl Table Generation: 17AUG2021 (09:15) 04NOV2021 (12:52)

(Data Cutoff Date: 17JUN202102SEP2021, Database Snapshot Date: 27JUL202127SEP2021) Output File: /nda2 unblinded/C4591001 G1_PVP adl6mpd2/ads1 boost 6932.5922

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 Table 25. Exposure to BNT162b2 by Age Group, Dose, and Race/Ethnie Origin (Totals) (C4591001) -BNT162b2 Experienced 12-15 Years - Open-Label Follow-up Period - Subjects Who Were Rerandomized to Receive 1 Booster Dose of BNT162b2 (30 μg) - Booster Safety PopulationOriginally Received Placebo and Then Received BNT162b2 After Unblinding

Age Group [®] —Dose — Race/Ethnic Origin Exposure (Number of Do <u>Received)</u>	Number of Subjects Exposed to BNT162b2 ISES	<u>Total Number of</u> <u>Vaccine Doses</u>		Inserted Cells
Subjects ≥18 years to ≤55 years	-		1	
Vaccine 30 µg				Inserted Cells
-Racial origin	-			
	249			
-Black or African American	28			
-Asian	16			
	2			
Native Hawaiian or other Pacific Islander 1 Do	<u>.18</u>	18		Deleted Cells
	4			Inserted Cells
Not reported	6			Inserted Cells
	306			
-Ethnie origin	-			
	85			
<u>Non Hispanic/non Latino</u>	219			
Not reported 2 Do	<u>992</u>	<u>,1984</u>		Deleted Cells
Total	306 1010	2002	$\langle \ \rangle$	Inserted Cells
Note: Only phase 3 Includes subjects who already beca	me eligible for unblinding at 16 ve	ars of age confirmed		Inserted Cells

Note: Only phase 3<u>Includes</u> subjects who alreadybecame eligible for unblinding at 16 years of age, confirmed to have received two doses of placebo originally and then received BNT162b2 in the original phase of the study and consented to participate the booster evaluation phase were rerandomized to receive booster dose. a. Based on age at booster dose.post unblinding.

PFIZER CONFIDENTIAL SDTM Creation: 29JUL2021 (23:0130SEP2021 (11:35) Source Data: adsl Table Generation: 17AUG2021 (09:16) 04NOV2021 (12:52)

(Data Cutoff Date: 17JUN202102SEP2021, Database Snapshot Date: 27JUL202127SEP2021) Output File: ./nda2_unblinded/C4591001_G1_PVP_adl6mpd2/ads1_boost_6942_s9222_

Population	BNT162	bjects Exposed to 2b2 -(30-µg) =306) # ^b	<u>,Total Number (</u>	<u>of Vaccine Doses</u>
Age Group ^a	Male	<u>Female</u>	Male	<u>Female</u>
$\frac{2}{2} \frac{2}{2} \frac{30 \mu \text{g}}{2}$	<u>567</u>	- <u>564</u>	- <u>1128</u>	- <u>1127</u>

Label 27. Exposure to BNT162b2 by Gender (C4591001) – 12-15 Years – Open Label Follow-up Period – Subjects Who Originally Received Placebo and Then Received BNT162b2 After Unblinding

-	-	jects Exposed to 162b2	Total Number	of Vaccine Doses
Dose Age Group ^a	Male	<u>Female</u>	<u>Male</u>	<u>Female</u>
- Vaccine 30 μg				
<u>≥12 years to ≤15</u> years	<u>518</u>	<u>492</u>	<u>1027</u>	<u>975</u>
a. Based on age at vi confirmed to have rece PFIZER CONFIDENT 04NOV2021 (12:52)	ived placebo origina		BNT162b2 post unblu	nding.

(Data Cutoff Date: 02SEP2021, Database Snapshot Date: 27SEP2021) Output File: /nda2_unblinded/C4591001_PVP_adl6mpd2/adsl_1215_s932_plac

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BNT162b2	
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 1257	742

<u>Age Group^a Dose</u> Race/Ethnic Origin	<u>Number of Subjects</u> <u>Exposed to BNT162b2</u>	<u>Total Number of</u> <u>Vaccine Doses</u>
12 years to ≤15 years		
/accine 30 µg		
Racial origin		
White	<u>970</u>	<u>1935</u>
Black or African American	<u>52</u>	<u>103</u>
Asian	<u>72</u>	<u>143</u>
American Indian or Alaska Native	<u>4</u>	<u>8</u>
Native Hawaiian or other Pacific slander	<u>3</u>	<u>6</u>
Multiracial	<u>24</u>	<u>48</u>
Not reported	<u>6</u>	<u>12</u>
Total	<u>1131</u>	2255
Ethnic origin		
Hispanic/Latino	<u>132</u>	<u>263</u>
Non-Hispanic/non-Latino	<u>997</u>	<u>1988</u>
Not reported	<u>2</u>	<u>4</u>
Total	<u>1131</u>	2255

04NOV2021 (12:52) (Data Cutoff Date: 02SEP2021, Database Snapshot Date: 27SEP2021) Output File: /nda2 unblinded/C4591001 PVP adl6mpd2/adsl 1215 s942 blind

BNT162b2
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742

Table 29.	Exposure to BNT162b2 by Race/Ethnic Origin (C4591001) – 12-15 Years
	- Open-Label Follow-up Period - Subjects Who Originally Received
	Placebo and Then Received BNT162b2 After Unblinding

<u>Age Groupª</u> <u>Dose</u> Race/Ethnic Origin	<u>Number of Subjects</u> Exposed to BNT162b2	<u>Total Number of</u> <u>Vaccine Doses</u>
-		
≥ 12 years to ≤ 15 years		
Vaccine 30 µg		
Racial origin		
White	<u>866</u>	<u>1718</u>
Black or African American	<u>48</u>	<u>96</u>
Asian	<u>62</u>	<u>123</u>
American Indian or Alaska	<u>2</u>	<u>3</u>
Native		
Multiracial	<u>26</u>	<u>52</u>
Not reported	<u>6</u>	<u>10</u>
Total	<u>1010</u>	<u>2002</u>
Ethnic origin		
Hispanic/Latino	<u>115</u>	222
Non-Hispanic/non-Latino	<u>892</u>	<u>1774</u>
Not reported	<u>3</u>	<u>6</u>
Total	<u>1010</u>	<u>2002</u>

a. Based on age at vaccination. Includes subjects who became eligible for unblinding at 16 years of age, confirmed to have received placebo originally and then received BNT162b2 post unblinding. PFIZER CONFIDENTIAL SDTM Creation: 30SEP2021 (11:35) Source Data: adsl Table Generation: 04NOV2021 (12:52)

04NOV2021 (12:52) (Data Cutoff Date: 02SEP2021, Database Snapshot Date: 27SEP2021) Output File: /nda2_unblinded/C4591001_PVP_adl6mpd2/adsl_1215_s942_plac_

Table 30. Exposure to BNT162b2 (30 μg) by Special Population (C4591001) – 12-15 Years – Blinded Placebo-Controlled Follow-up Period

Population	<u>Number of Subjects</u> <u>Exposed</u> <u>to BNT162b2 (30 μg)</u> <u>(N^a=1131)</u> <u>n^b</u>	<u>Total Number of</u> <u>Vaccine Doses</u>
Subjects with any baseline comorbidity	155 249	<u>494</u>
Any Malignancy + Metastatic Solid Tumor + Leukemia + Lymphoma	8	
Chronic Pulmonary Disease	4 <u>0119</u>	235
Mild Liver Disease + Moderate or Severe Liver Disease	5 2	<u>4</u>
C erebrovascular Disease + Peripheral Vascular Disease + Myocardial Infarction + Congestive Heart Failure	1	
Diabetes With/Without Chronic Complication	6 2	4
Obese	122<u>143</u>	<u>284</u>

Note: Includes subjects who became eligible for unblinding at 16 years of age.

Note: Comorbidity is based<u>on</u> Charlson Comorbidity Index categories. Participants identified as belonging to these categories were identified by medical history data collected during the study.

Note: Only phase 3 subjects who already received two doses of BNT162b2 in the original phase of the study and consented to participate the booster evaluation phase were rerandomized to receive booster dose. Note: Hemiplegia or Paraplegia only includes preferred terms Hemiplegia and Paraplegia. <u>No participants</u> were identified.

N = number of subjects in the specified group.

b. n = Number of subjects reporting at least 1 occurrence of any comorbidity or obese (BMI $\geq \frac{30 \text{ kg/m}^2}{1695 \text{ th percentile [12-15 Years of age]-]}}$.

PFIZER CONFIDENTIAL SDTM Creation: 29JUL2021 (13:4605OCT2021 (18:33) Source Data: admh Table Generation: 17AUG2021 (09:16) 04NOV2021 (12:52)

(Data Cutoff Date: 17JUN202102SEP2021, Database Snapshot Date: 27JUL202127SEP2021) Output File: /nda2_unblinded/C4591001_G1_PVP_adl6mpd2/admh_boost1215_s953_blind

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BNT162b2
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742

Table 31.Exposure to BNT162b2 (30 µg) by Special Population (C4591001) – 12-15Years – Open-Label Follow-up Period – Subjects Who Originally
Received Placebo and Then Received BNT162b2 After Unblinding

<u>Population</u>	<u>Number of Subjects Exposed</u> <u>to BNT162b2 (30 µg)</u> <u>(N^a=1010)</u> n ^b	<u>Total Number of</u> <u>Vaccine Doses</u>
- Subjects with any baseline comorbidity	<u>214</u>	<u>425</u>
Chronic Pulmonary Disease	<u>114</u>	<u>226</u>
Rheumatic Disease	<u>2</u>	<u>4</u>
Diabetes With/Without Chronic Complication	<u>2</u>	<u>4</u>
Obese	<u>116</u>	<u>229</u>

Note: Includes subjects who became eligible for unblinding at 16 years of age.

Note: Includes subjects confirmed to have received placebo originally and then received BNT162b2 post unblinding.

Note: Comorbidity is based on Charlson Comorbidity Index categories. Participants identified as belonging to these categories were identified by medical history data collected during the study.

Note: Hemiplegia or Paraplegia only includes preferred terms Hemiplegia and Paraplegia. No participants were identified.

a. N = number of subjects in the specified group.

b. n = Number of subjects reporting at least 1 occurrence of any comorbidity or obese (BMI >95th percentile [12-15 Years of age]).

PFIZER CONFIDENTIAL SDTM Creation: 05OCT2021 (18:33) Source Data: admh Table Generation: 04NOV2021 (12:52)

(Data Cutoff Date: 02SEP2021, Database Snapshot Date: 27SEP2021) Output File:

/nda2 unblinded/C4591001 PVP adl6mpd2/admh 1215 s953 plac

2.1.2.a.2. Inclusion and Exclusion Criteria

Detailed descriptions of all inclusion and exclusion criteria for clinical studies are provided in the individual CSRs which were filed to IND 019736.

Inclusion criteria

- Healthy participants who are determined by medical history, physical examination (if required), and clinical judgment of the investigator to be eligible for inclusion in the study.
- Healthy participants with pre-existing stable disease, defined as disease not requiring significant change in therapy or hospitalization for worsening disease during the 6 weeks before enrollment, can be included. In order for the overall Phase 3 study population to be as representative and diverse as possible, the inclusion of participants with known chronic stable infection with HIV, HCV, or HBV was permitted as the study progressed. Specific criteria for these Phase 3 participants can be found in the C4591001 protocol, Section 10.8.
- Phase 2/3 only: Participants who, in the judgment of the investigator, are at higher risk for acquiring COVID-19 (including, but not limited to, use of mass transportation, relevant demographics, front-line essential workers, and others).
- The participants enrolled were 12 years of age and older; the 12- to 15-year-old cohort was included in the protocol in October 2020.

Exclusion criteria

Phase 1 exclusion criteria were stricter than criteria in Phases 2 and 3 of the study. Participants were excluded from the studies according to the general criteria listed below:

• Previous vaccination with any coronavirus vaccine

<u>Reason for exclusion</u>: To avoid confounding the assessment of serological or clinical immune response in the study population.

Is it considered to be included as missing information? No.

Rationale: Minimal potential clinical impact on the target population.

Previous clinical or microbiological diagnosis of COVID-19

<u>Reason for exclusion</u>: Phase 1 excluded participants with a previous clinical or microbiological diagnosis of COVID-19 because these participants may have some degree of protection from subsequent infection by SARS-CoV-2 and therefore would confound the pivotal efficacy endpoint. During Phase 2/3, participants with prior undiagnosed infection were allowed to be enrolled. Screening for SARS-CoV-2 with nucleic acid amplification test by nasal swab or antibodies to non-vaccine SARS-CoV-2

antigen by serology was not conducted before vaccine administration in Phase 2/3, but samples were taken to run these assays after vaccination, thus identifying participants with unidentified prior infection. This group will be assessed to identify whether prior infection affects safety.

Is it considered to be included as missing information? No.

<u>Rationale</u>: Safety in study participants with prior infection will be assessed in the pivotal study.

• Immunocompromised individuals with known or suspected immunodeficiency, as determined by history and/or laboratory/physical examination.

<u>Reason for exclusion</u>: Immunocompromised participants may have impaired immune responses to vaccines and would therefore limit the ability to demonstrate efficacy, which is the primary pivotal endpoint.

Is it considered to be included as missing information? No.

<u>Rationale</u>: Participants with potential immunodeficient status were not specifically included in the study population. However, since the study population is intended to be as representative as possible of the vulnerable population to COVID-19 illness, sub-analyses of immunogenicity data in future studies may provide further understanding of immune responses in this population.

• Receipt of blood/plasma products or immunoglobulin, from 60 days before study intervention administration or planned receipt throughout the study

<u>Reason for exclusion</u>: To avoid confounding the assessment of serological or clinical immune response in the study population.

Is it considered to be included as missing information? No.

Rationale: No impact on the safety of the target population.

• Women who are pregnant or breastfeeding

Reason for exclusion: To avoid use in a vulnerable population.

Is it considered to be included as missing information? Yes.

<u>Rationale</u>: It is not known if maternal Maternal vaccination with <u>BNT162b2 would</u> <u>haveCOVID-19 mRNA vaccine is being studies in C4591015 to explore</u> unexpected negative consequences to the embryo or <u>fetusfoetus</u>.

• Other medical or psychiatric condition including recent (within the past year) or active suicidal ideation/behavior or laboratory abnormality that may increase the risk of study participation or, in the investigator's judgment, make the participant inappropriate for the study

<u>Reason for exclusion</u>: To avoid misleading results deriving from non-compliance to study procedures.

Is it considered to be included as missing information? No.

Rationale: Safety profile of BNT162b2 is not expected to differ in these subjects when properly administered.

2.1.2.a.2.1. Non-Study Post-Authorization Exposure

Cumulatively, through <u>18 June30 September</u> 2021, approximately <u>774,478,4401,709,812,866</u> doses of BNT162b2 were shipped worldwide, corresponding to <u>642,817,105approximately</u> <u>1,402,241,841</u> estimated administered doses.

The worldwide number of shipped doses may serve as a reasonable indicator of subject exposure, considering that approximately 8382% of the shipped doses were administered.

Cumulative worldwide estimated exposure³ by dose, and region based on or extrapolated from internal data (number of shipped doses) and published data (number of doses administered) is displayed in Table 27.

The estimated cumulative number of shipped and administered doses of BNT162b2 by region based on data provided in the shipment tracker (Order Book),⁴ from the receipt of the first temporary authorization for emergency supply on 01 December 2020 through 30 September 2021, are summarized in Table 32.

Table 32. Cumulative Estimated Shipped-and-/Administered Doses of BNT162b2 by Region Worldwide, through 18 June 2021

Region/Country/Ot her	% of Doses	Total Number of Shipped Doses	Total Number of Administered Doses
Europe	41. <u>81%</u>	323502270 <u>7032671</u> 10	268506884 <u>5714739</u> 11
European Union ^{ag} (27)	33.3 <u>30.0</u> <u>%</u>	2576283455135057 85	2 <u>138315264159396</u> <u>86</u>
Additional EEA Countries ^{#a} (3)	0. <u>54%</u>	3559335<u>7006155</u>	2954248<u>5674986</u>
Switzerlanda	<u>0.3%</u>	<u>4500990</u>	<u>3690812</u>

³ Including data from license partners.

⁴ The Order Book is the most accurate tracker of shipment used as data source for all the Regions and Countries; US shipment data not available in the Order Book were taken from the Order Management Dashboard and data for Fosun License Partner territories, Hong Kong and Macau, were provided by BioNTech.

Region/Country/Ot her	% of Doses	Total Number of Shipped Doses	Total Number of Administered Doses
$\overline{\rm UK^b}$	<u>3.6%</u>	<u>61213230</u>	50194849
Other Countries ^{bc}	<u>8.06.3%</u>	<u>6231459010721704</u> <u>5</u>	<u>5172111087917977</u>
Commonwealth of Independent States ^d	<u>0.6%</u>	<u>9823905</u>	8055602
North America [®]	29.8 <u>18.5</u> %	230593605 <u>3160936</u> 95	191392692<u>2645972</u> 40
US	26.6 <u>15.8</u> %	205645305 2700205 05	170685603<u>2268172</u> 24
Canada	3. 2.7%	2494830046073190	2070708937780016
Central and South America ^{d<u>f</u>}	7.4 <u>12.5</u> %	5764473021308568 0	47845126 <u>17473025</u> <u>8</u>
Asia	19.5<u>23.6</u> %	15073 9485 <mark>4040775</mark> 81	1251137733313436 16
Japan ^a Japan ^a	12.2<u>10.7</u> %	94169790<u>18349812</u> 0	78160926 <u>15046845</u> 8
Other Countries^eOther Countries ^g	7.3<u>12.9</u> <u>%</u>	<u>5656969522057946</u> <u>1</u>	4 6952847<u>18087515</u> <u>8</u>
Oceania	0.71.4%	568152023158980	471566218990364
Australia/New Zealand ⁻ Zealand ^a	0.7 <u>1.4%</u>	5681520 <u>23158980</u>	471566218990364
Other Countries	0.0%	0	0
Africa ^f Africa ^h	0.82.9%	6316830 <u>50129820</u>	524296941106452
Total	100.0%	774478440 <u>1709812</u> 866	642817105 <u>1402241</u> 841

Table 32. Cumulative Estimated Shipped-and-/Administered Doses of BNT162b2 by Region Worldwide, through 18 June 2021

a. Conditional approval. In this Region BNT162b2 was conditionally approved;

b. Includes:

<u>In the UK, with both authorisation the authorization</u> for emergency supply under regulation 174 and the conditional marketing authorisationauthorization approval, are currently active for BNT162b2. c. Includes Albania, Kosovo, and North Macedonia and Switzerland with conditional approval, Georgiawhere BNT162b2 was conditionally approved. Serbia and Ukraine withwhere it received

authorization for emergency supply,

Azerbaijan, Bosnia and Moldova where BNT162b2it was shipped for COVAX, Turkey where it was shipped according to a pharmacovigilance agreement in place by the MAH and the Turkish government-: c. Authorization for emergency supply.

d. Includes: Georgia and Ukraine where BNT162b2 received authorization for emergency supply and Moldova where it was conditionally approved; in Azerbaijan BNT162b2 was shipped for COVAX, and Tajikistan and Uzbekistan are part of US government donations;

Brazil and Peru with conditional approval,

e. In this Region BNT162b2 initially received authorization for emergency supply; in the US, a full approval (BLA) was also granted on 23 August 2021 and in Canada a full approval (NDS) replacing the pre-existing authorization for emergency supply was granted during the current reporting period on 16 September 2021;

<u>F___Includes</u> Chile, Colombia, Costa Rica, Dominican Republic, Ecuador, El Salvador, Honduras, Mexico, Panama, Paraguay and Uruguay with authorisationwhere BNT162b2 received authorization for emergency supply, Argentina, Brazil and Peru where BNT162b2 was conditionally approved; Bolivia where BNT162b2 was shipped for COVAX and Antigua&Barbuda, Bahamas, Barbados, Belize, Dominica, Grenada, Guyana, Jamaica, St. Kitts&Nevis, St. Lucia, StVin&Grenadine, Suriname and Trinidad&Tobago that are part of US Government donations;

Table 32. Cumulative Estimated Shipped-and /Administered Doses of BNT162b2 by Region Worldwide, through 18 June 2021

Region/Country/Ot	% of Doses	Total Number of Shipped	Total Number of
her		Doses	Administered Doses
g. Includes :			
Hong Kong, Malaysia	and South Korea with	conditional approval,	
Bahrain, Bhutan, Indon	iesia, Iraq, Israel, Jord	lan, Kuwait, Lebanon, Macau, <u>Ma</u>	<u>lldives, Mongolia, O</u> man,
Pakistan, Palestine, Pal	cistan<u>Philippines</u>, Qat	ar, Saudi Arabia, Singapore, Sri L	anka-and, United Arab Emirates
withand Vietnam where	e BNT162b2 received	authorization for emergency sup	ply ,
; Hong Kong, Malaysia	a, South Korea and Th	ailand where BNT162b2 was con	ditionally approved and
Bangladesh, Bhutan, L	aos , Maldives, Monge	olia, Philippines and West Bank &	c Gaza where BNT162b2 was
shipped for COVAX;			
h. Includes f.Includes	s:		
Rwanda, Tunisia and S	outh Africa where BN	JT162b2 received authorisation for	or emergency supply,
Angola, Botswana, Cape Verde, Chad, Ivory Coast, LibyaLybia and Togo where BNT162b2 was shipped for			
COVAX; Benin, Cong	o, Gabon, Namibia, S	eychelles, Sierra Leone and Ugan	da that are parts of US
Government donations	; Botswana, Egypt, Es	watini, Kenya, Mauritius, Moroco	co, Rwanda, South Africa and
Tunisia where BNT162b2 received authorization for emergency supply.			

Out of the total shipped and administered doses, 213,475,665 and 177,184,802 respectively, were shipped to Rest Of World (Non EEA countries, Canada, Central and South America, Asian countries [excluding Japan], Oceania and Africa).

Method Used to Calculate Exposure

Not applicable.

Exposure

Not applicable.

2.1.2.a.3. Regulatory Actions Related to Safety

There were no withdrawals for safety reasons up to 28 February 30 September 2021.

2.1.2.b. Populations Not Studied in the Pre-Approval Phase

There has been limited exposure to BNT162b2 in some special populations and no epidemiologic studies have been conducted in pregnant/lactating women, pediatric participants (<12 years of age), and specific subpopulations that were initially excluded from the BNT162b2 program.

Table 33. Exposure of Special Populations Included or not in Clinical Trial Development Programs

Type of special population	Exposure
Pregnant women	Available data on BNT162b2 administered to pregnant women are
-	insufficient to inform on vaccine-associated risks in pregnancy. In a
	reproductive and developmental toxicity study, no vaccine-related

Table 33. Exposure of Special Populations Included or not in Clinical Trial Development Programs

Type of special population	Exposure
	adverse effects on female fertility, fetal development, or postnatal development were reported.
	Participants 16 years of age and older
	Through the cut-off date of 13 March 2021, there were 50 cases (52 events) originating from Study C4591001 in participant 16 years of age and older, and all were unique pregnancies.
	Participants in Booster group12-15 years of age
	Through the cut-off date of 17 June02 September 2021, there were no <u>CT</u> cases originating f pregnancies from Studystudy C4591001 in participants enrolled in the booster group. <u>12-15 years of age.</u>
Breastfeeding women	Breastfeeding women were not initially included in the BNT162b2 clinical development program.
	Data are not available to assess the effects of BNT162b2 on the breastfed infant or on milk production/excretion.
	The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for BNT162b2 and any potential adverse effects on the breastfed child from BNT162b2 or from the underlying maternal condition. For preventive vaccines, the underlying maternal condition is susceptible to disease prevented by the vaccine.
	Participants 16 years of age and older
	Through the cut-off date of 13 March 2021, there were no CT cases indicative of exposure during breastfeeding from study C4591001 in participants 16 years of age and older.
	Participants in Booster group12-15 years of age
	Through the cut-off date of 17 June02 September 2021, there were no <u>CT</u> cases indicative of exposure during breastfeeding originating from <u>Studystudy</u> C4591001 in participants enrolled in the booster group12-15 years of age.
Participants with relevant comorbidities:	Healthy participants with pre-existing stable disease, defined as disease not requiring significant change in therapy or hospitalization for worsening disease during the 6 weeks before enrollment, were
 Participants with hepatic impairment Participants with renal impairment Participants with cardiovascular disease 	included. This allowed enrollment of a proportion of participants with common comorbidities such as cardiovascular diseases including hypertension, chronic pulmonary diseases, asthma, chronic liver disease, BMI >30 kg/m ² , participants with stage 3 or worse chronic kidney disease, and participants with varying disease severity.
 Immunocompromised participants Participants with a disease severity different from inclusion criteria in CTs 	Participants with potential immunodeficient status were not specifically included in the study population. Please refer to Table 22, Table 23, <u>Table 30</u> and <u>Table 26Table 31</u> for the exposure of special populations.

Table 33. Exposure of Special Populations Included or not in Clinical Trial Development Programs

Type of special population	Exposure
Participants of different racial	Please refer to Table 21 Table 15, Table 16, Table 17, Table 18,
and/or ethnic origin	Table 19, Table 20, Table 21, Table 28 and Table 25 Table 29 for
	exposure information by ethnic origin from the studies.
Subpopulations carrying known and relevant polymorphisms	No data available.
Pediatric participants	The safety and effectiveness of BNT162b2 in individuals younger than 16.5 years of age have not been established.
	Participants 16 to 17 years of age and older A total of 671 pediatric participants 16 to 17 years of age received
	BNT162b2 through the DLP of 13 March 2021:
	 378 participants in the blinded-placebo controlled follow-up period (Error! Not a valid result for table.).
	• 293 participants in the open-label follow-up period after the unblinding (Table 5).
	Participants 12 to 15 years of age One thousand andone hundred eighty (1180thirty-one (1131) pediatric participants 12 to 15 years of age received in the blinded controlled follow-up period; 1010 participants, who originally received placebo, then received BNT162b2 in the Open-Label Follow-up period after unblinding through the cut-off date of 13 March02 September 2021 (Table 3 Table 24 and Table 5 Table 25).
Elderly (≥65 years old)	The safety and effectiveness of BNT162b2 in elderly participants was consistent with that seen in younger adult participants.
	Clinical studies of BNT162b2 included a total of 8846 participants 65 years of age and over; of these, 8827 were from study C4591001, through the cut-off date of 13 March 2021:
	 4590 participants in the blinded-placebo controlled follow- up period (Error! Not a valid result for table.)
	• 4237 participants in the open-label follow-up period after unblinding (Table 5).
	Nineteen (19) participants 65 years of age and over were from study BNT162-01 study through the cut-off date of 23 October 2020 (Table 6).

Abbreviations: EUA = emergency use authorization; BMI = body mass index; COVID-19 = coronavirus disease 2019; CT = clinical trial

2.1.2.c. Adverse Events / Adverse Reactions

2.1.2.c.1. Identification of Safety Concern in the Initial PVP Submission

2.1.2.c.1.1. Risks not Considered Important for Inclusion in the List of Safety Concerns in the PVP

Not all potential or identified risks for the vaccine are considered to meet the level of importance necessitating inclusion in the list of safety concerns in the PVP:

- Risks with minimal and temporary clinical impact on patients (in relation to the severity of the disease prevented).
- The following reactogenicity events are identified risks not included in the list of safety concerns in the PVP: Injection site pain, Fever, Chills, Fatigue, Headache, Muscle pain, and Joint pain.
- Very rare potential risks for any medicinal treatment, including vaccines, which are well known to healthcare professionals are not included in the list of safety concerns.

2.1.2.c.2. Important Identified and Potential Risks and Missing Information

2.1.2.c.2.1. Presentation of Important Identified Risks and Important Potential Risks

Important Identified Risks

Table 34. Myocarditis and Pericarditis

Potential	A mechanism of action (MOA) by which the vaccine could cause myocarditis and
mechanisms,	pericarditis has not been established. Nonclinical studies, protein sequence
evidence source and	analyses and animal studies in rats and non-human primates have not identified a
strength of evidence	MOA. Hypotheses for MOA include an immune stimulated response (including
	the possibility of molecular mimicry), a general systemic inflammatory response
	from vaccination or a hypersensitivity response.
Characterisation of	Participants 16 years of age and older
the risk	Data from the CT dataset ^a (cut-off date: 18 June 2021)
	Two cases were retrieved with the myocarditis and pericarditis search strategy ^b in the clinical trial dataset through the cut-off date of 18 June 2021. These cases originated from Phase 3 clinical study C4591001 and are summarized below:
	Myocarditis:
	There were no cases reporting myocarditis as SAE.
	Pericarditis (2 cases):
	Two (2) serious adverse events [PT Pericarditis] were reported, both deemed not related to study treatment by the Investigator.
	Data from the safety database: (cut-off date: 18 June 2021):
	Since the first temporary authorization for emergency supply under Regulation 174 in the UK (01 December 2020) and through 18 June 2021, 823 potentially relevant cases (0.3% of the total post-authorization dataset) were retrieved from the Myocarditis and Pericarditis search strategy: ^b 490 cases reported events related to myocarditis and 371 cases reported events related to pericarditis (in 38 of these 823 cases, the subjects developed both myocarditis and pericarditis related events).
	Myocarditis (490 cases):
	These 490 cases were individually reviewed and assessed according to Brighton Collaboration (BC) Myocarditis Case Definition and Level of Certainty Classification (version 1.4.2, 30 May 2021), as shown in the Table below:
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Table 34. Myocarditis and Pericarditis

	Number of cases
BC 1	41
BC 2	44
BC 3	42
BC 4	337
BC 5	26
Total	490
evidence to meet the case definition" and Lev Fhere were 464 cases meeting BC Level 1 to 4 Country of incidence: Israel (135), US (78), G (taly, Japan (13 each), Austria (10), Greece, Sp Norway (6 each), Ireland (5); the remaining 23 countries. Gender: Females (133), Males (325), Unknow Age (n=443) ranged from 16 to 97 years (mear Reported relevant PTs: Myocarditis (463) and	which are presented below: ermany (76), UK (55), France (21) ain (8 each), Sweden (7), Canada, cases originated from 17 different n (6). = 37.2 years, median = 32.0 years
	Total Events
<u> </u>	N = 464 (%)
Serious events	459 (98.9)
Events with Criterion of Hospitalization	
Events with Criterion of Hospitalization Distribution of events by Outcome	459 (98.9) 337 (72.6)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death	459 (98.9) 337 (72.6) 14 (3.0)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving	459 (98.9) 337 (72.6) 14 (3.0) 149 (32.1)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	459 (98.9) 337 (72.6) 14 (3.0) 149 (32.1) 106 (22.8)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved Outcome: Resolved with sequelae	459 (98.9) 337 (72.6) 14 (3.0) 149 (32.1) 106 (22.8) 10 (2.2)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	459 (98.9) 337 (72.6) 14 (3.0) 149 (32.1) 106 (22.8)
Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved Outcome: Resolved with sequelae	459 (98.9) 337 (72.6) 14 (3.0) 149 (32.1) 106 (22.8) 10 (2.2) 185 (39.9) rael (50), UK (38), Italy (33), ia (9), Greece (7), Germany (6), 5 each); the remaining 20 cases n (5).

Table 34. Myocarditis and Pericarditis

	Total Events N = 372 (%)
Serious events	370 (99.5)
Events with Criterion of Hospitalization	206 (55.4)
Distribution of events by Outcome	
Outcome: Death	3 (0.8)
Outcome: Resolved/Resolving	213 (57.3)
Outcome: Not resolved	63 (16 9)
Outcome: Resolved with sequelae	7 (1.9)
Outcome: Unknown/No data	86 (23 1)

Participants 12 to 15 years of age

Data from the CT dataset:* database (cut-off date 02 September 2021): No cases were One (1) case was retrieved reporting with the Myocarditis and Pericarditis as SAEsearch strategy^b in the clinical trial datasetCT database through the cut-off date of 18 June 02 September 2021. This case originated from the clinical study C4591001. Myocarditis (1 case): One (1) SAE (PT Myocarditis) was reported 3 days after the administration of the second dose of BNT162b2; the participant recovered the following day. The SAE was deemed not related to study treatment by the investigator. Pericarditis: There were no cases reporting pericarditis as SAE. Data from the safety database: (cut-off date 30 September 2021): Through 18 June 30 September 2021, 15180 potentially relevant cases (0.03% of the total post-authorization dataset) were retrieved from the Myocarditis and Pericarditis search strategy: b 13154 cases reported myocarditis and 461 cases reported pericarditis (in 235 of these 15180 cases, the subjects developed both myocarditis and pericarditis). Myocarditis (13154 cases) These 13154 cases were individually reviewed and assessed according to Brighton Collaboration (BC) Myocarditis Case Definition and Level of Certainty Classification, as shown in the Table below: Brighton Collaboration Level Number of cases BC 1 θ 14 BC 2 09 BC 3 0 BC 4 ++130BC 5 <u>21</u> 13154 Total Level 1 indicates a definitive case with the highest level of diagnostic certainty of myocarditis, level 2 indicates a probable case, and level 3 indicates a possible case. Level 4 is defined as "reported event of myocarditis with insufficient evidence to meet the case definition" and Level 5 as not a case of myocarditis.

Table 34. Myocarditis and Pericarditis

NoThe details of 153 cases met BC levels(excluding 1 to 3. There were 11 cases meeting BC Level 4, which 5 case) are presented below: Country of incidence: US (10) and Bahrain (1). Country of incidence: Hong Kong (39), US (25), Germany (18), France (17), Italy (8), Israel (7), Austria and Spain (6 each), Denmark and Japan (5 cases); the remaining 17 cases originated from 13 different countries. Gender: Females (20Female (1), Males (10130), and not reported (3). Age (n= $\frac{11153}{1}$) ranged from 12 to 15 years (mean = 13.8-9 years, median = 14.0 years). Reported relevant PT: Myocarditis (11153). Overall event seriousness and outcome of these 44153 cases are summarized below. **Total Events** N = 11153 (%) 10153* (100.0) Serious events Events with Criterion of Hospitalization <u>110 (71 9)</u> **Distribution of events by Outcome** Outcome: Death θ Outcome: Resolved/Resolving 3 Outcome: Not resolved 4 θ Outcome: Resolved with sequelae Outcome: Unknown/No data Pericarditis (4 cases) Country of incidence: US (4). Gender: Males (4). Age (n=4) ranged from 12 to 15 years (mean = 13.5 years, median = 13.5 years). Reported relevant PT: Pericarditis (4). Overall event seriousness and outcome of these-4 cases are summarized below. **Total Events** -4 Serious events 3 Events with Criterion of Hospitalization 4 Distribution of events by Outcome 0 Outcome: Death Outcome: Resolved/Resolving 479 (51.6) Outcome: Not resolved 17 (11.1) Outcome: Resolved with sequelae 0 Outcome: Unknown/No data 257 (37.3) *Includes 1 case where myocarditis was captured as non-serious and upgraded to serious after the DLP. Pericarditis (61 cases)

Table 34. Myocarditis and Pericarditis

These 61 cases were individually reviewed	
Colloboration (BC) Pericarditis Case Def Classification, as shown in the Table belo	
Classification, as shown in the Table belo	<u>Sw</u>
Brighton Collaboration Level	Number of cases
BC 1	1
BC 2	4
BC 3	0
BC 4	56
BC 5	0
<u>Total</u>	<u>61</u>
Level 1 indicates a definitive case with	the highest level of diagnostic certaint
of pericarditis, level 2 indicates a proba	
possible case. Level 4 is defined as "re	
insufficient evidence to meet the case of	lefinition" and Level 5 as not a case of
pericarditis.	
The details of 61 cases are presented below	<u>DW:</u>
Country of incidence: Hong Kong (29), I	taly (7), France (6), US (4), Canada (3)
Australia, Belgium, Germany, and Japan	
from 4 different countries.	,
Gender: Males (48) and Females (13).	
Age (n=61) ranged from 12 to 15 years (n	mean = 14.0 years, $median = 14.0$ year
Reported relevant PT: Pericarditis (61).	
Overall event seriousness and outcome o	f these Booster Group
Data from the CT database:	
Through DLP 17 June 2021, no cases we	re retrieved reporting myocarditis and
pericarditis in the participants who receive	
perieducidas in the participants who recert	
Data from the safety database:	
Through DLP 18 June 2021, no cases we	re retrieved reporting myocarditic and
pericarditis in the subjects who received	
below.	<u></u>
[Total Events
	$\frac{\text{Total Events}}{N = 61 (\%)}$
Serious events	$\frac{N = 01 (70)}{61 (100.0)}$
Events with Criterion of Hospitalization	
Events with Criterion of Hospitalization Distribution of events by Outcome	<u>1/(2/9)</u>
Outcome: Death	0
Outcome: Deam Outcome: Resolved/Resolving	18 (29 5)
Outcome: Not resolved	9 (14.8)
Outcome: Resolved with sequelae	<u>9 (14.8)</u> 1 (1.6)
Outcome: Unknown/No data	33 (54 1)

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Table 34. Mv	ocarditis and	Pericarditis
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Risk factors and risk groups	Post-authorization reports have been received for more males than females, over a wide age range and following dose 1 and dose 2 of the vaccine. Evaluation by the US CDC has found reports to be most frequent in adolescent and young adult male patients following the second dose of vaccine.
Preventability	Due to an unknown MOA, preventative measures are not clear for individuals with or without a personal history of myocarditis or pericarditis.
Impact on the risk- benefit balance of the biologic product	The vaccine continues to have a favorable risk benefit balance
Public health impact	Considering the low rates of myocarditis and pericarditis reported following vaccination, balanced with the risk of death and illness (including myocarditis) caused by SARS-CoV-2, the public health impact of post-vaccination myocarditis and pericarditis is minimal.

a. Please note that CT dataset from the safety database includes only cases reporting SAEs.

b. Search criteria: the following PTs were used to retrieve cases of Myocarditis and Pericarditis: Autoimmune myocarditis; Eosinophilic myocarditis; Giant cell myocarditis; Hypersensitivity myocarditis; Immune-mediated myocarditis; Myocarditis; Autoimmune pericarditis, Pericarditis; Pericarditis adhesive; Pericarditis constrictive; Pleuropericarditis.

Table 35. Anaphylaxis

mechanisms,histaevidence source andcontastrength of evidencedysp	raction of an allergen with IgE on basophils and mast cells triggers release of amine, leukotrienes and other mediators that cause diffuse smooth muscle raction and vasodilation with plasma leakage. This can manifest clinically with onea, hypotension, swelling (sometimes leading to airway compromise), and (including hives).
the risk Data Three clim part Inve Data Three data Ana Ana Ana Ana	ticipants 16 years of age and older a from the CT dataset ^a (cut-off date 18 June 2021) bugh 18 June 2021, ^b there was 1 case from the CT dataset (from Phase 3 icical study C4591001) of serious Anaphylactoid reaction in a 17-year-old icipant reported as resolved and deemed related to study treatment by the stigator: a from the safety database: (cut-off date 18 June 2021): bugh 18 June 2021, ^b there were 3822 cases (1.2% of the total post authorization set) reporting a total of 3914 events in individuals 16 years and older including: phylactic reaction (3414) phylactic shock (420) phylactoid rection (75) phylactoid shock (5) rall event seriousness and outcome are summarized below:

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Table 35. Anaphylaxis

Serious events Events with Criterion of Hospitalization Distribution of events by Outcome* Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	N = 3914 (%)
Events with Criterion of Hospitalization Distribution of events by Outcome* Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	3868 (98.8)
Distribution of events by Outcome* Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	1231 (31.5)
Outcome: Death Outcome: Resolved/Resolving Outcome: Not resolved	
Outcome: Not resolved	28 (0.7)
Outcome: Not resolved	2958 (75.6)
Out	171 (4.4)
Outcome: Resolved with sequelae	56 (1.4)
Outcome: Unknown	704 (18)
*For the outcome count, the multiple Lowest Level within a case are counted and presented individually total count of the event outcome may exceed the tot	y Therefore, for selected PTs th
urticipants 12 to 15 years of age ata from the CT dataset; *database (cut-off date (nrough 18 June02 September 2021, there were n porting Anaphylactic reaction/shock, Anaphylac om the CT datasetdatabase.	to cases ^b there were no cases
ata from the safety database: (cut-off date 30 Se	ptember 2021):
nrough 18 June 2021, ^b30 September 2021, there <u>1 Anaphylactic</u> reaction, <u>4 Anaphylactic shock</u> , <u>ockAnaphylactoid reaction</u>) in individuals 12 to <u>ost-authorization dataset</u>); overall event seriousn clow:	and 1 anaphylactic o 15 years of age; (0.01% of t ess and outcome are summar
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there <u>1 Anaphylactic</u> reaction, <u>4 Anaphylactic shock</u> , <u>ockAnaphylactoid reaction</u>) in individuals 12 to <u>ost-authorization dataset</u>); overall event seriousn	and 1 anaphylactic 15 years of age; (0.01% of t ess and outcome are summar Total Events
nrough <u>18 June 2021</u> , ^b 30 September 2021, there <u>1 Anaphylactic reaction</u> , <u>4 Anaphylactic shock</u> , <u>oekAnaphylactoid reaction</u>) in individuals 12 to <u>sst-authorization dataset</u> ; overall event seriousn clow:	and 1 anaphylactic 0 15 years of age; (0.01% of the set of the s
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there <u>1 Anaphylactic reaction</u> , <u>4 Anaphylactic shock</u> , <u>oekAnaphylactoid reaction</u>) in individuals 12 to <u>isst-authorization dataset</u> ; overall event seriousn clow:	and 1 anaphylactic 15 years of age; (0.01% of the set
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there <u>1 Anaphylactic</u> reaction, <u>4 Anaphylactic shock</u> , <u>oekAnaphylactoid reaction</u>) in individuals 12 to <u>ost-authorization dataset</u>); overall event seriousn clow: <u>Serious events</u> Events with Criterion of Hospitalization	and 1 anaphylactic 0 15 years of age; (0.01% of the set of the s
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there <u>1 Anaphylactic</u> reaction, <u>4 Anaphylactic shock</u> , <u>oekAnaphylactoid reaction</u>) in individuals 12 to <u>oest-authorization dataset</u>); overall event seriousn clow: <u>Serious events</u> <u>Events with Criterion of Hospitalization</u> Distribution of events by Outcome	and 1 anaphylactic 0 15 years of age; (0.01% of t ess and outcome are summar Total Events N = 546 (%) 546 (100) ↓15 (32.6)
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there 1 Anaphylactic reaction, 4 Anaphylactic shock, oekAnaphylactoid reaction) in individuals 12 to ist-authorization dataset); overall event seriousn clow: Serious events Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death	and 1 anaphylactic 15 years of age; (0.01% of the sess and outcome are summaries and outcome a
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there 1 Anaphylactic reaction, 4 Anaphylactic shock, oekAnaphylactoid reaction) in individuals 12 to ist-authorization dataset); overall event seriousn elow: Serious events Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death Outcome: Resolved/Resolving	and 1 anaphylactic 0 15 years of age; (0.01% of the sess and outcome are summarian and outcome are summarian and the s
nrough <u>18 June 2021</u> , ^b <u>30 September 2021</u> , there 1 Anaphylactic reaction, 4 Anaphylactic shock, oekAnaphylactoid reaction) in individuals 12 to ist-authorization dataset); overall event seriousn clow: Serious events Events with Criterion of Hospitalization Distribution of events by Outcome Outcome: Death	and 1 anaphylactic 15 years of age; (0.01% of the sess and outcome are summaries and outcome a

Table 35. Anaphylaxis

	Conclusion: Evaluation of Anaphylactic reaction/shock, Anaphylactoid reaction/shock cases through <u>18 Jun30 September</u> 2021 did not reveal any significant new safety information. Anaphylaxis is appropriately described in the product labeling as are non-anaphylactic hypersensitivity events. Surveillance will continue.
Risk factors and risk groups	Known hypersensitivity to any components of the vaccine.
Preventability	Prevention of anaphylaxis may not be possible, particularly with the 1 st dose of a vaccine; therefore, healthcare professionals administering the vaccine must be vigilant for early signs and symptoms.
Impact on the risk- benefit balance of the biologic product	Anaphylactic reaction in an individual can be impactful (medically important) because it is a potentially life-threatening event requiring medical intervention.
Public health impact	Minimal due to rarity of the event. Although the potential clinical consequences of an anaphylactic reaction are severe, this is a known risk of vaccines to healthcare professionals with negligible public health impact.

a. Please note that CT dataset from the safety database includes only cases reporting SAEs.

b. Updated search criteria starting from the 6th SMSR (see 5th Monthly Safety Update preliminary PRAC Assessment Report; EMEA/H/C/005735/MEA/002.4): PTs Anaphylactic reaction, Anaphylactic shock, Anaphylactoid reaction, Anaphylactoid shock, without Brighton Collaboration criteria applied.

Important Potential Risks

Table 36. Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)

Potential	This potential risk is theoretical because it has not been described in association with
mechanisms,	the BNT162b2 or it has not been reported from any other late phase clinical trial of
evidence source	other human vaccine. Animal models of SARS-CoV-2 infection have not shown
and strength of	evidence of VAED after immunization, whereas cellular immunopathology has been
evidence	demonstrated after viral challenge in some animal models administered SARS-CoV-1
	(murine, ferret and non-human primate models) or MERS-CoV (mice model)
	vaccines. ^{1,6} This potential risk has been included based on these animal data with
	these related betacoronaviruses. Historically, disease enhancement in vaccinated
	children following infection with natural virus has been observed with an inactivated
	respiratory syncytial virus vaccine. ⁷
	Potential mechanisms of enhanced disease may include both T cell-mediated [an
	immunopathological response favoring T helper cell type 2 (T _H 2) over T helper cell
	type 1 (T _H 1)] and antibody-mediated activity (antibody responses with insufficient
	neutralizing activity leading to formation of immune complexes and activation of
	complement or allowing for Fc-mediated increase in viral entry to cells).8
Characterization	Participant 16 years and older
of the risk	Data from the CT database (set off data 12 Marsh 2021)
	Data from the CT database (cut-off date 13 March 2021)

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Table 36.	Vaccine-Associated Enhanced Disease (VAED), including Vaccine-
Associated Enhanced Respiratory Disease (VAERD)	

		BNT162b2 (30 μg) (N ^a =23164)		lacebo =23155)
Timing	n^{b} (%)	(95% CI ^c)	n ^b (%)	(95% CIC
PD1 Before Dose 2	0	(0.0, 0.0)	6 (0.0)	(0.0, 0.1)
Within 7 days PD1	0	(0.0, 0.0)	0	(0.0, 0.0)
PD2	1 (0.0)	(0.0, 0.0)	25 (0 1)	(0.1, 0.2)
Within 7 days PD2	0	(0.0, 0.0)	2 (0.0)	(0.0, 0.0)
Total ^d	1 (0.0)	(0.0, 0.0)	31 (0 1)	(0.1, 0.2)
Output File: /nda2 u If VAED/VAERD modified and/or mo upon subsequent na lower risk for sever risk for severe COV fatal outcomes, or f cases in vaccinated challenging to asses specific clinical or theoretical risk is b above shows a favc BNT162b2 versus t risk of VAED/VAE Data from the CT d	were to occur i ore severe clini tural infection te COVID-19 f /ID-19 (e.g. ol or observation individuals wh ss for VAED/V laboratory mar est performed a trable balance of chose receiving ERD at this tim	in vaccinated indi cal presentation of . This may result having more seven der or immunoco of an unfavorable nen compared to t /AERD on an indi kers at this time, ra at a population lev of severe COVID g placebo, providin e.	viduals, it may of SARS-CoV-2 in individuals a re disease, for in mpromised) hav e imbalance in s hose not vaccin ividual case bas rather surveillar vel, ⁹ as noted at -19 cases in par ng reassurance a	viral infectior assumed to be a adividuals at kn ving higher rate severe COVID ated. It is sis, given the la coce for this poove. The table ticipants receive
Data from the C1 d There were no case the DLP of 18 June Data from the safet No post-authorized	s indicative of 2021 .^a. y database (cur AE reports ha	VAED/VAERD : t-off date 18 June ve been identified	as SAEs in the (ED/VAERD,

Table 36. Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)

The search criteria utilised to identify potential cases of VAED for this report includes PTs indicating a lack of effect of the vaccine and PTs potentially indicative of severe or atypical COVID-19^a. Since the first temporary authorization for emergency supply under Regulation 174 in

the UK (01 December 2020) and through the DLP 18 June 2021, there were 584 cases (0.2% of the total post-authorization dataset), reporting 1427 potentially relevant events.

Seriousness criteria for the total 584 cases: Medically significant (452, of which 10 also serious for disability), Hospitalization required (non-fatal/non-life threatening) (115, of which 3 also serious for disability), Life threatening (34, of which 22 were also serious for hospitalization), Death (160).

Gender: Females (298), Males (268), Unknown (18);

Age (n=553) ranged from 17 to 103 years (mean = 70.3 years, median = 77.0);

Overall event seriousness and outcome are summarized below:

	Total Events N = 1427 (%)
Serious events	1261 (88.4)
Events with Criterion of	612 (42.9)
Hospitalization	
Distribution of events by Outcome	*
Outcome: Death	311 (21.8)
Outcome: Resolved/Resolving	375 (26.3)
Outcome: Not resolved	246 (17.2)
Outcome: Resolved with sequelae	14 (1.0)
Outcome: Unknown/No data	484 (33.9)

* For the outcome count, the multiple Lowest Level Terms that code to the same PT within a case are counted and presented individually. Therefore, for selected PTs the total count of the event outcome may exceed the total number of events.

The most frequently reported relevant PTs ($\geq 2\%$) were: Drug ineffective (390), Vaccination failure (194), Dyspnoea (180), COVID-19 pneumonia (179), Diarrhoea (111), Respiratory failure (52), Vomiting (50), Pulmonary embolism (33).

Conclusion: VAED may present as severe or unusual clinical manifestations of COVID-19. Overall, there were 425 subjects with confirmed COVID 19 following one or both doses of the vaccine; 288 of the 425 cases were severe, resulting in hospitalization, disability, life threatening consequences or death. None of the 288 cases could be definitively considered as VAED/VAERD.

In this review of subjects with COVID-19 following vaccination, based on the current evidence, VAED/VAERVAED remains a theoretical risk for the vaccine. Surveillance will continue.

Participants 12 to 15 years of age

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Table 36. Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)

Data from the CT dataset database (cut-of	f date 02 September 2021):			
There were no cases reporting VAED/VAI through the DLP of 18 June 02 September				
Data from the safety database: ^b (cut-off date 30 September 2021):				
Through the DLP <u>18 June30 September</u> 2021, there were <u>no cases2 cases (reporting 6 potentially relevant events)</u> indicative of VAED or VAERD in the safety database involving individuals 12 to 15 years of age. [#] .				
Booster Group				
Data from the CT database:				
No AEs wereSeriousness criteria for the 2 cases: Life threatening (1 also serious for hospitalization) and Hospitalization required (non-fatal/non-life threatening) (1).				
Gender: Males (2);				
Age (n=2): 12 years and 15 years (1 case e	each);			
Overall event seriousness and outcome are	summarized below:			
	$\frac{\text{Total Events}}{N=6}$			
Serious events	<u>6</u>			
Events with Criterion of	<u>6</u>			
Hospitalization Distribution of events by Outcome				
	0			
Outcome: Death	0			
Outcome: Resolved/Resolving Outcome: Unknown/No data	<u>4</u> 2			
<u>The relevant PTs</u> reported that suggested a among participants in the Phase 3 BNT162 cut-off date (17 June 2021).				
Data from the safety database:				
Through DLP 18 June 2021, nothese 2 cases were retrieved reporting severe COVID- 19: Diarrhoea, Drug ineffective, Multisystem inflammatory syndrome in the subjects who received booster dose. ⁻ children, Seizure, Vaccination failure, and Vomiting (1 each).				
Conclusion: VAED may present as severe COVID-19. In both cases, the subjects had the vaccine. Upon review, these 2 cases un course was not descriptive of an unusual c infection.	confirmed COVID-19 following 2 doses of likely represent VAED as the clinical			

Table 36. Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)

Risk factors and risk groups	In this review of subjects with COVID-19 following vaccination, based on the current evidence, VAED/VAERD remains a theoretical risk for the vaccine. Surveillance will continue. It is postulated that the potential risk may be increased in individuals producing lower neutralizing antibody titers or in those demonstrating waning immunity. ^{8,9}
Preventability	An effective vaccine against COVID-19 that produces high neutralizing titers and a $T_{\rm H}1$ predominant CD4 ⁺ T cell response and strong CD8 ⁺ T cell response, is expected to mitigate the risk of VAED/VAERD; ^{1,8} that immune profile is elicited by BNT162b2 in clinical and preclinical studies. ^{10,11}
Impact on the risk-benefit balance of the biologic product	If there were an unfavorable balance in COVID-19 cases, including severe cases, in the pivotal clinical study between the vaccine and placebo groups, that may signal VAED/VAERD.
Public health impact	The potential risk of VAED/VAERD could have a public health impact if large populations of individuals are affected.

a. Please note that CT dataset from the safety database includes only cases reporting SAEs.a.
 b. Search criteria updated to include new PTs introduced in the MedDRA version 24.0. The updated search criteria is: PTs Vaccine associated enhanced disease OR Vaccine associated enhanced respiratory disease OR Standard Decreased Therapeutic Response Search AND at least 1 of the following PTs Dyspnoea; Tachypnoea; Hypoxia; COVID 19 pneumonia; Respiratory Failure; Acute Respiratory Distress Syndrome; Cardiac Failure; Cardiogenic shock; Acute myocardial infarction; Arrhythmia; Myocarditis; Vomiting; Diarrhoea; Abdominal pain; Jaundice; Acute hepatic failure; Deep vein thrombosis; Pulmonary embolism; Peripheral Ischaemia; Vasculitis; Shock; Acute kidney injury; Renal failure; Altered state of consciousness; Seizure; Encephalopathy; Meningitis; Cerebrovascular accident; Thrombocytopenia; Disseminated intravascular coagulation; Chillblains; Erythema multiforme; Multiple organ dysfunction syndrome; Multisystem inflammatory syndrome in children. Note: the "Standard Decreased Therapeutic Response" search includes the Lack of efficacy PTs (Drug ineffective/Vaccination failure).

2.1.2.c.2.2. Presentation of Missing Information

Table 37. Use in Pregnancy and Lactation

Evidence source:

The safety profile of the vaccine is not known in pregnant or lactating women due to their exclusion from the pivotal clinical study. There may be pregnant women who choose to be vaccinated despite the lack of safety data. It will be important to follow these women for pregnancy and birth outcomes. The timing of vaccination in a pregnant woman and the subsequent immune response may have varying favorable or unfavorable impacts on the embryo/fetus. The clinical consequences of SARS-CoV-2 infection to the woman and fetus during pregnancy is not yet fully understood and the pregnant woman's baseline health status may affect both the clinical course of her pregnancy and the severity of COVID-19 disease. These factors and the extent to which the pregnant woman may be at risk of exposure to SARS-CoV-2 will influence the benefit risk considerations for use of the vaccine.

Population in need of further characterization:

The lack of data <u>will beis</u> communicated in product labeling; one clinical study of the safety and immunogenicity of the BNT162b2 in pregnant <u>and lactating</u> women is ongoing (C4591015); 42 non-interventional studies (C4591009, <u>and C4591011,) are planned and 2 non-interventional studies</u> (C4591021, and C4591022) <u>are ongoing</u> to assess whether sub-cohortsuse of interest, such as pregnant women,

Table 37. Use in Pregnancy and Lactation

experience increased risk of safety events of interest following receipt of the BNT162b2 are planned/ongoingin pregnancy (see 3 1.3 – Action plan for safety issues).

Data from the Safety Database^a (Cut-off date 18 June 2021)

Since the first temporary authorization for emergency supply under Regulation 174 in the UK (01 December 2020) and through 18 June 2021, there were 2636 cases (0.8 % of the total Post-authorization dataset) reporting use during pregnancy or lactation.

Overall event seriousness and outcome are summarized below:

	Total Events N = 6215 (%)
Serious events	2464 (39.6)
Events with Criterion of Hospitalization	314 (5 1)
Distribution of events by Outcome*	
Outcome: Death	61 (1)
Outcome: Resolved/Resolving	1657 (26.7)
Outcome: Not resolved	602 (9.7)
Outcome: Resolved with sequelae	65 (1)
Outcome: Unknown/No data	3864 (62.2)

* For the outcome count, the multiple Lowest Level Terms that code to the same PT within a case are counted and presented individually Therefore, for selected PTs the total count of the event outcome may exceed the total number of events

The most frequently reported relevant PTs ($\geq 2\%$) were: Maternal exposure during pregnancy (867), Exposure via breast milk (791), Exposure during pregnancy (402), Off label use (296), Abortion spontaneous, Product use issue (277 each), Headache (184), Maternal exposure during breast feeding (161), Fatigue (155), Pyrexia (134), Pain in extremity (119), Vaccination site pain (91), Myalgia (79), Chills (75), Maternal exposure timing unspecified (73), Nausea, Pain (72 each), and Dizziness (56).

Participants 12 to 15 years of age

Data from the safety database: (Cut-off date 30 September 2021)

Through 30 September 2021, there was 1 case reporting use of BNT162b2 during pregnancy in the safety database. The serious case involved a 12-year-old female who received first dose of BNT162b2 during pregnancy (trimester of exposure unknown) and had miscarriage after 2 weeks of vaccine administration (PTs Maternal exposure during pregnancy, Fatigue, and Abortion spontaneous). Patient outcome was reported as recovered with sequelae. Through 30 September 2021, there were no cases reporting use of BNT162b2 during lactation.

a. Cumulative RMP tables on Missing information are provided as per previous FDA's request to include a cumulative analysis, from post-authorization experience, of the Important Missing Information identified in the Pharmacovigilance Plan.

Table 38. Vaccine Effectiveness

Evidence source:

Although vaccine efficacy in a controlled clinical study is the objective of the pivotal study, real-world vaccine effectiveness when the BNT162b2 is used in a large and more diverse population is unknown.

Anticipated risk/consequence of missing information:

Efficacy information obtained from clinical study data <u>will beis</u> communicated in the product labeling. Four post-authorization effectiveness studies in real-world use are <u>planned/ongoing</u>: 1 interventional study (BNT162-01 cohort 13), 1 non-interventional study (C4591014) and 2 low-interventional studies (WI235284 and WI255886) to determine the effectiveness of BNT162b2 when administered outside of the clinical setting (see 3.1.3 – *Action plan for safety issues*).

Data from the Safety Database^a (Cut-off date: 18 June 2021)

Since the first temporary authorization for emergency supply under Regulation 174 in the UK (01 December 2020) and through 18 June 2021, there were 6373 cases (1.9% of the total Post-authorization dataset) reporting lack of efficacy.

Overall event seriousness and outcome are summarized below:

	Total Events N = 6373 (%)
Serious events	6373 (100)*
Events with Criterion of Hospitalization	616 (9.7)
Distribution of events by Outcome	
Outcome: Death	334 (5 2)
Outcome: Resolved/Resolving	1253 (19.7)
Outcome: Not resolved	721 (11.3)
Outcome: Resolved with sequelae	21 (0.3)
Outcome: Unknown/No data	4044 (63.5)

*Includes 26 cases where LOE was captured as non-serious and upgraded to serious after the DLP.

The PT Drug ineffective was reported in 4765 cases, Vaccination failure was reported in 1608 cases; the most frequently co-reported PTs ($\geq 2\%$) were: COVID-19 (5022), Asymptomatic COVID-19 (502), Pyrexia (412), Suspected COVID-19 (379), SARS-CoV-2 test positive (359), Headache (327), Fatigue (262), Cough (227), Dyspnoea (180), COVID-19 pneumonia (179), Myalgia (162), Asthenia (156), Malaise (152), and Chills (133).

Participants 12 to 15 years of age

Data from the safety database: (Cut-off date 30 September 2021)

Through 30 September 2021, there were 29 cases retrieved reporting lack of efficacy. Upon review, 1 case was not considered to be true lack of efficacy because the subject developed SARS-CoV-2 infection during the early days from the first dose (days 1-13); the development of a vaccine preventable disease during this time is not considered a lack of effect of the vaccine. Therefore, there were 28 relevant cases reporting lack of efficacy in individuals 12 to 15 years of age; overall event seriousness and outcome are summarized below:

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Table 38. Vaccine Effectiveness

	$\frac{\text{Total Events}}{N = 28 (\%)}$
berious events	28 (100)
Events with Criterion of Hospitalization	<u>2 (7.1)</u>
Distribution of events by Outcome	
Outcome: Death	<u>0</u>
Outcome: Resolved/Resolving	4 (14.3)
Outcome: Not resolved	<u>1 (3.6)</u>
Outcome: Unknown/No data	23 (82 1)

The PTs Drug ineffective and Vaccination failure were reported in 19 and 9 cases, respectively; the co-reported events reported more than once were coded to the PTs: COVID-19 (24), Pyrexia (5), Headache (4), Suspected COVID-19 (3), and Fatigue (2).

a. Cumulative RMP tables on Missing information are provided as per previous FDA's request to include a cumulative analysis, from post-authorization experience, of the Important Missing Information identified in the Pharmacovigilance Plan.

Table 39. Use in Paediatric Individuals <125 Years of Age[§]

Evidence source:

BNT162b2 has not been initially studied in pediatric individuals younger than 12 years of age due to their exclusion from the pivotal clinical study.

Paediatric individuals may display different reactogenicity and safety profiles compared to adults, due to lower body mass and differently matured immunological responses.

Population in need of further characterization:

The are no data in individuals less than $\frac{125}{2}$ years of age; ain the pediatric population:

- 3 clinical studies [C4591001 (≥12 to ≤15 years of age), C4591007 (<12 years of age) and C4591007 substudy - Troponin group (5 to <12 years of age and 12 to <16 years of age) are ongoing;
- <u>1</u> clinical study of the safety, tolerability, immunogenicity and efficacy of BNT162b2 in individuals younger than 12 years [C4591007 (< 12 years of age)]^a is ongoing (see 3.1.3 — Action plan for safety issues); a[C4591023 (<6 months; >5 to <12 years of age) is planned;
- <u>2</u> non-interventional study (studies [C4591009) is planned to assess the occurrence of safety events of interest in a general US population (< 12 and ≥ 12 to ≤15 years of age) (see 3.1.3 — Action plan for safety issues); a non-interventional study (and C4591038 (former C4591021) to assess potential increased risk of AESI, in a general EU population substudy) (<12 years of age)] are planned;
- 1 low interventional study is planned [C4591036 (<21 years of age, including <12 years is ongoing (of age)]

For details on these studies, see 3.1.3 – Action plan for safety issues); a non-interventional study (C4591036) to characterize the clinical course, risk factors, long term sequelae, and quality of life in children and young adults <21 years with acute post vaccine myocarditis/pericarditis is planned (see 3.1.3 – Action plan for safety issues).

Data from the Safety Database^bDatabase^a (cut-off date 30 September 2021)

Since the first temporary authorization for emergency supply under Regulation 174 in the UK (01 December 2020) and through <u>18 June30 September</u> 2021, there were <u>132 cases56 cases</u>^b (0.0401% of the total Post-authorization dataset) involving individuals below <u>125</u> years of age. Overall event seriousness and outcome are summarized below:

	Total Events N = 343172 (%)
Serious events	34 (9.9<u>30 (17.4</u>)
Events with Criterion of Hospitalization	<u>3 (</u> 1 (0.3 .7)
Distribution of events by Outcome*	
Outcome: Death	2 (0.6<u>1</u> 2)
Outcome: Resolved/Resolving	101 (29.4<u>61 (35.5</u>)
Outcome: Not resolved	53 (15.5<u>46</u> (26.7)
Outcome: Resolved with sequelae	0
Outcome: Unknown/No data	192 (56.0<u>63</u> (36.6)

* For the outcome count, the multiple Lowest Level Terms that code to the same PT within a case are counted and presented individually. Therefore, for selected PTs the total count of the event outcome may exceed the total number of events.

The most frequently reported PTs (22%)(>3 occurrences) were: Product administered to patient of inappropriate age (4321), Off label use (3717), Product use issue (2615), Pyrexia (1311), Fatigue, Headache (11 each), Pain in extremity (10), Nausea (8), Malaise, Myalgia (7 each), Arthralgia, Dizziness, Pain (6 each), Chills, Swelling (5 each), Diarthoea, Pruritus, Rash (, and Nausea (4 each), Abdominal pain upper, Circumstance or information capable of leading to medication error, Cough, Injection site pain, Nasopharyngitis, Peripheral swelling, Vaccination site pain, Vaccination site swelling, and Vomiting (3 each).

a. Phase 1 open label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer blinded safety, tolerability, and immunogenicity, study of a SARS-CoV-2 RNA vaccine candidate against COVID 19 in healthy children <12 years of age.</p>
b & Microir information has been available to the first the available to the safety.

b-§ Missing information has been reworded to reflect the current state.

a. Cumulative RMP tables on Missing information are provided as per previous FDA's request to include a cumulative analysis, from post-authorization experience, of the Important Missing Information identified in the Pharmacovigilance $Plan_{\overline{\tau}_2}$

b. Please note that at the DLP of 18 June 2021 there were 29 additional pediatric cases under 5 years of age; at the DLP of 30 September 2021, follow-up information was received for these cases and they were identified to refer to adult subjects, rather than to pediatric subjects under 5 years of age.

2.1.2.d. Identified and Potential Interactions, Including Food-Biologic Product and Drug-Biologic Product Interactions

As noted in the WHO Guidelines on Nonclinical Evaluation of Vaccines,³ pharmacokinetics testing is not required for final formulation. No interaction linked to metabolism is expected with vaccines. The only potential for interaction is with other vaccines administered concomitantly and with immunosuppressive drugs.

Co-administration studies with BNT162b2 have not been done, therefore there is not sufficient data to understand the effect on vaccine effectiveness of BNT162b2 or co-administered vaccines. A co-administration study with seasonal influenza vaccine is planned. If BNT162b2 is given at the same time as other injectable vaccine(s), the vaccine(s) should be administered at different injection sites.

2.1.2.e. Epidemiology of Indication and Target Population

Indication

Active immunization against to prevent COVID-19 disease caused by SARS-CoV-2 virus, in individuals ≥ 1612 years of ageand older.

Incidence:

The COVID-19 is caused by a novel coronavirus labeled as SARS-CoV-2. The disease first emerged in December 2019, when a cluster of patients with pneumonia of unknown cause was recognized in Wuhan City, Hubei Province, China.¹² The number of infected cases rapidly increased and spread beyond China throughout the world. On 30 January 2020, the WHO declared COVID-19 a Public Health Emergency of International Concern and thus a pandemic.¹³

Estimates of SARS-CoV-2 incidence change rapidly. We<u>The MAH</u> obtained incidence and prevalence estimates using data from Worldometer, a trusted independent organization that collects COVID-19 data from official reports and publishes current global and country-_specific statistics online.¹⁴

As of 03 March15 August 2021, the overall number of people who had been infected with SARS-CoV-2 was over 115207 million worldwide,¹⁵ an increase of nearly 100-92 million in the 75 months since 28 July 202003 March 2021.¹⁶ Table 35 Table 40 shows the incidence and prevalence as of 03 March15 August 2021 for the US, UK, and EU-27 countries. In the CONFIDENTIAL

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EU and the UK, by <u>03 March15 August</u> 2021 the total number of confirmed cases had accumulated to <u>almost 2741</u> million people, or <u>5,2268.074</u> per 100,000 people (from <u>1.727</u> million, or <u>3375,226</u> per 100,000 by <u>28 July 202003 March 2021</u>). Across countries in the EU, the number of confirmed cases ranged from <u>1,0722.118</u> to <u>11,83615,620</u> cases per 100,000 people. Finland and <u>GreeceGermany</u> reported the lowest incidence rates while Czech Republic, Slovenia, and Luxembourg reported the highest. ¹⁵

In the US, the number of confirmed cases had reached over $\frac{2937}{2000}$ million (8,864cases (11.236) per 100,000 people) by $\frac{03 \text{ March } 2021.^{15} \text{ 15 August } 2021.^{15}}{15 \text{ August } 2021.^{15}}$ This is an increase from 4.529 million (1,3578.864 per 100,000) by $\frac{28 \text{ July } 2020.^{17} \text{ 03 March } 2021.^{16}}{16}$

	Total Cases	Incidence: Total Cases/ 100,000	Active Cases[®]Cases	Prevalence: Active Cases/ 100,000	Total Deaths	Mortality: Deaths / 100,000	Population
Global	115,760,943 207.731.370	-1,485 2.665	21,707,680 17.141.537	- <u>278-220</u>	2,571,518 4.371.692	-33-<u>56</u>	7,794, 824,793<u>798,124</u>
EU-27	22,642,536 35.243.565	-5,083 7.910	6,113,464 2.000.178	-1,462-<u>149</u>	553,363 747.450	<u>-124-168</u>	445,4 <u>24,167-541,383</u>
UK	4,194,785 6,241,011	-6,157 <u>9,140</u>	1,065,282 313,343	1, 564-<u>923</u>	123,783 130,894	<u>-182-192</u>	68,125,249<u>284,715</u>
EU-27 + UK	26,837,321 <u>41,484,576</u>	-5,226 <u>8,074</u>	7,178,746 <u>3,313,521</u>	-1,398-<u>645</u>	677,146 <u>878,344</u>	- <u>132-171</u>	513, 549,416-<u>826,098</u>
US	29,456,377 <u>37,435,835</u>	<u>-8,864</u> <u>11,236</u>	<u>8,921,400</u> <u>6,653,787</u>	-2,685 <u>1,997</u>	531,652 637,439	<u>-160-191</u>	332,304,437 333,172,543
EU-27 Countr	ies						
Austria	4 65,322-<u>668,732</u>	5,147 <u>7,378</u>	21,028 8,559	<u>-233-94</u>	<u>- 8,625</u> 10,756	95<u>119</u>	9, 040,866 . <u>063,848</u>
Belgium	.774,344.<u>1,149,869</u>	<u> </u>	<u>699,566</u> 52,835	- <u>6,019-454</u>	22,141 25,287	191<u>217</u>	11, 623,476-<u>646,025</u>
Bulgaria	-253,183-<u>4</u>32,962	3,662 <u>6,284</u>	33,770 <u>14,645</u>	<u>-488-213</u>	10,413 <u>18,339</u>	151<u>266</u>	6, 913,156.<u>889,852</u>
Croatia	- <u>244,205-367,022</u>	-5,973 <u>9,002</u>	3,322 <u>1,903</u>	<u>-81-47</u>	-5,555 <u>8,283</u>	136 203	4, 088,197-<u>076,913</u>
Cyprus	- 35,620 - <u>108,707</u>	-2,936 <u>8,931</u>	33,331 <u>17,496</u>	2,747 <u>1,437</u>	232 <u>456</u>	19<u>38</u>	1, 213,250-<u>217,182</u>
Czech Republic	1, 269,058-<u>676,222</u>	-11,836 <u>15,620</u>	-154,580 <u>2,441</u>	1,442-<u>23</u>	-20,941 <u>30,373</u>	195<u>283</u>	10, 722,330 _ <u>731,206</u>
Denmark	- <u>212,798-330,777</u>	- 3,665 <u>5,688</u>	- 6,995 <u>12,854</u>	120 - <u>221</u>	2, 370-<u>560</u>	41 <u>44</u>	5, 805,897-<u>815,014</u>
Estonia	-69,193 - <u>136,992</u>	<u>5,214</u> <u>10,319</u>	-17,938 5,131	1,352-<u>387</u>	-615 <u>1,279</u>	4 <u>696</u>	1,327, 135-<u>533</u>
Finland	-59,442-<u>117,531</u>	1,072 2.118	-12,683 <u>70.536</u>	229 <u>1,271</u>	759-<u>995</u>	1 4 <u>18</u>	5, 546,504-<u>550,349</u>
France	3,<u>6,449,863</u>810,216	<u>5,829</u> <u>9,857</u>	3,461,485 <u>455,926</u>	5,295-<u>697</u>	87,542 <u>112,612</u>	134<u>172</u>	65, 370,546 <u>4</u>35,079
Germany	2,472,896 <u>3,825,039</u>	2,945 <u>4,549</u>	-126,785 53,169	<u>-151-63</u>	_71,711 <u>92,370</u>	85<u>110</u>	-83,963,843 <u>84,083,573</u>
Greece	197,279-<u>535,237</u>	-1,899 <u>5,163</u>	<u>21,157</u> <u>37,611</u>	- <u>204-363</u>	-6,597 <u>13,174</u>	64<u>127</u>	10, 388,744-<u>366,043</u>

Table 40. Incidence, Prevalence, and Mortality of COVID-19 as of 03 March-15 August 2021-1515

	Total Cases	Incidence: Total Cases/ 100,000	Active Cases [®] <u>Cases</u>	Prevalence: Active Cases/ 100,000	Total Deaths	Mortality: Deaths / 100,000	Population
Hungary	<u>810,316</u> 439,900	- 4,561 8,412	-98,361 14,326	1,020-<u>149</u>	-15,324 30,038	159<u>312</u>	9, 643,837<u>6</u>32,892
Ireland	- <u>221,189-322,989</u>	4,44 6 6,461	-193,468 42,205	3,889-<u>844</u>	- 4,357 5,059	88<u>101</u>	4, 974,683 - <u>999,386</u>
Italy	2,976,274 4.435.008	4 <u>,92</u> 7 7.347	<u>437,421</u> 126.466	724-<u>210</u>	<u>98,635</u> 128,413	163 213	60, 401,999- <u>362,319</u>
Latvia	- <u>88,022-140,122</u>	4,702 7,522	<u>9,233</u> 1,218	- <u>493-65</u>	-1,654 2,561	<u>88138</u>	1, 872,109-<u>862,827</u>
Lithuania	- <u>200,349-289,810</u>	7,430 <u>10,815</u>	<u>10,859</u> 12,355	4 <u>03461</u>	3,281 <u>4,451</u>	122<u>166</u>	2, 696,596-<u>679,705</u>
Luxembourg	- <u>55,902-74,595</u>	8,834 <u>11,704</u>	-3,074-<u>705</u>	- <u>486-111</u>	<u>-643-828</u>	102<u>130</u>	632,773-<u>637,340</u>
Malta	- <u>23,226-35,337</u>	5,251 7,979	_3,000 <u>1,043</u>	<u>-678-236</u>	<u>-321-430</u>	73 97	442, 333-<u>858</u>
Netherlands	1, 101,430 <u>901,900</u>	-6,418 <u>11,072</u>	- <u>124,498</u>	- <u>725</u>	- 15,697 <u>17,909</u>	92<u>104</u>	17, 160,343 - <u>177,282</u>
Poland	1,735,406 2,885,333	4,589 7,633	249,567 <u>154,721</u>	- 660-<u>409</u>	<u>44,360</u> 75,299	117<u>199</u>	37, 818,722- 800,220
Portugal	806,626-<u>1,003,335</u>	7,926 9,872	64,797 45,367	637-<u>446</u>	-16,430 17,562	161<u>173</u>	10, 176,690-<u>163</u>,426
Romania	812,318 <u>1,087,223</u>	4,242 5,694	44,953 2,982	235-<u>16</u>	-20,586 34,348	108<u>180</u>	19 , 151,141_<u>093,951</u>
Slovakia	314,359-<u>393,529</u>	-5,756 7,204	51,570-<u>825</u>	944-<u>15</u>	7,489 12,544	137<u>230</u>	5,4 61,420<u>462,601</u>
Slovenia	192,266-<u>261,428</u>	<u>9,247</u> 12,573	10,751 <u>2,150</u>	517-<u>103</u>	3,874 4,433	186<u>213</u>	2,079, 130-<u>258</u>
Spain	3,136,321 4,693,540	6,706 10,034	343,770 722,353	- 735 - <u>1.544</u>	-70,247 82,470	150<u>176</u>	46, 766,95 4- <u>775.041</u>
Sweden	- <u>675,292-1,110,147</u>	6,659 10,916	- <u>15,858</u>	- <u>156</u>	12,964 14,621	128<u>144</u>	10, 141,493<u>169,660</u>

Table 40. Incidence, Prevalence, and Mortality of COVID-19 as of 03 March-<u>15 August</u> 2021-¹⁶¹⁵

a Active case counts were not available for Netherlands and Sweden; therefore, those two countries are excluded from the overall prevalence calculations for EU 27 and EU 27. LIK.

The reported numbers refer only to cases that have been tested and confirmed to be carrying the virus and sometimes, depending upon the country, also presumptive, suspect, or probable cases of detected infection. There are large geographic variations in the proportion of the population tested as well as in the quality of reporting across countries. People who carry the virus but remain asymptomatic are less likely to be tested and therefore mild cases are likely underreported. The numbers should therefore be interpreted with caution. ¹⁸

Prevalence:

The prevalence of SARS-CoV-2 infection is defined as active cases per 100,000 people including confirmed cases in people who have not recovered or died. On <u>03 March15</u> <u>August</u> 2021, the overall prevalence <u>estimates</u> for the EU and UK (though not available for <u>Swedenwere 449</u> and the Netherlands) was 1,398923 active cases per 100,000,¹⁵ respectively¹⁵ compared to <u>51approximately 1,500</u> per 100,000 on <u>28 July 2020.¹⁶ for both</u>

<u>the EU and UK on 03 March 2021.</u>¹⁶ The range of reported prevalence was <u>8115</u> to <u>6,0191,544</u> per 100,000: <u>Croatia, DenmarkSlovakia, Romania</u>, and <u>GermanyCzech Republic</u> reported the lowest prevalence while <u>Belgium, FranceSpain, Cyprus</u>, and <u>IrelandFinland</u> reported the highest (Table 40).

In the US, the prevalence on 03 March15 August 2021 was nearly twice as high assimilar to the combined EU+UK estimates, with 2,685-1,997 active cases per 100,000.⁴⁵¹⁵ -The prevalence in the US was 653This is a decrease of approximately 700 per 100,000 on 28 July 2020since 03 March 2021, when the prevalence was 2,685 per 100,000.⁴⁶¹⁶

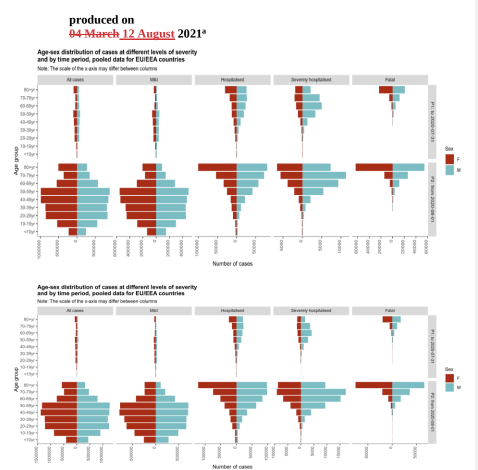
Demographics of the population in the proposed indication and risk factors for the disease:

Since the beginning of the pandemic, the ECDC has continuously collected COVID-19 information from all countries who are members of <u>the</u> EU/EEA-and the UK. In the ECDC's TESSy database, COVID-19 case-based data, including age and gender, are available for over 80% of the official number of cases reported by ECDC epidemic intelligence,¹⁹ enabling estimates of age and gender distribution representative of the European population. TESSy data on age and sex distributions by severity of symptoms as posted on <u>04-March12 August</u> 2021 are shown in Figure 1.²⁰

The top half of the figure represents data ending on 31 July 2020 and the bottom half presents data from 01 August 2020 to 04 March08 August 2021 (Figure 1). In general, the age-sex patterns before 01 August 2020 have remained the same since then. The gender distribution of persons testing positive for SARS-CoV-2 in the European population is similar for most age groups. Cases reported in TESSy have been older than the general population throughout the pandemic, with few cases observed in people aged younger than 20 years. This likely reflects the age distribution of people who met the requirements for being tested and is unlikely to reflect the actual distribution of infections in the population. Those with severe outcomes (hospitalized, severely hospitalized, <u>[admitted to intensive care and/or required respiratory support]</u>, or fatal) have been disproportionately older and male compared to COVID-19 cases overall. While age-sex patterns have remained consistent throughout the pandemic, a notable difference between the periods before and since 01 August 2020 is that the absolute numbers of cases have increased dramatically in the latter period compared to the earlier one.

Figure 1. Age-Sex distribution of COVID-19 Cases as Different Levels of Severity, <u>Pooled Data for EU/EEA and UKCountries</u>. Case-based Data from TESSy

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Note: <u>""</u><u>inild"="</u> = a case that has not been reported as hospitalized or a case that resulted in death a Data from ECDC COVID-19 Surveillance report Week <u>831</u>, 2021 <u>4 March12 August</u> 2021 "2 2 Age-sex pyramids"

Accessed 6 March 15 August 2021²⁰

US distributions of COVID cases and deaths by age, sex, and race, as well as the cross--tabulation of age and sex, are shown in Table 36.²¹ Table 41 as of 14 August 2021.²² At that time, the CDC reported that the US had recorded a total of 36,556,516 cases of COVID and 618,591 deaths attributable to the disease. However, because demographic data were not available for all US COVID cases and deaths, the numbers in Table 41 and Table 42 are drawn, respectively, from 29,346,352 cases and 513,204 deaths. Those under age 50 account for 65roughly 67% of cases but less thanapproximately 5% of deaths. For ages 18-74, males account for less than half of cases but over 60% of deaths. Among the pediatric population, there is close to a 50-50 case distribution between males and females across ages 0-17. However, the pediatric mortality distribution is highly irregular between the sexes,

with males being 51.5% of COVID deaths among 0-4 year olds, 55.9% among 5-11 year olds, 46.7% among 12-15 year olds, and 68.7% among 16-17 year olds.

Table 41. Distributions Distribution of Cases (n=21,895,936) and Deaths (n=382,00929,346,352) by Age, Sex, Race, and Cross-Tabulated Age and Sex --- United States as of 08 March14 August 2021^{21,aa} 22

								Age x Sex %		
Event	Age Grou p	Age %	Sex	Sex %	Race ^b Race ^b	Race %	Age Grou p	Males <u>%</u>	Females <u>%</u>	
Cases	0-4	<u>2.</u> 2	Males	47. 8 <u>7</u>	H/L	<u>20.728.</u> <u>3</u>	0-4	51.7	48.3	
	5- 17<u>11</u>	<u>9.54.</u>	Female s	52. 2 3	AI/AN	1 .2	5- 17 <u>11</u>	49 <u>50</u> .8	50.2 <u>49.</u>	
	12-15	<u>2</u> 3.8	5	<u> </u>	Asian	3.2	12-15	49.6	<u>1</u> 50.4	
	16-17	2.6			Black	11.6	16-17	48.3	51.7	
	18-29	22. <u>47</u>			<u>NH/PI</u> Asian	<u>0.</u> 3 .6	18-29	47.1 <u>46.</u> 9	52.9 <u>53.</u> 1	
	30-39	16. <mark>3</mark> 6			<u>White</u> Black	<u>12.250.</u> <u>3</u>	30-39	<u>48.2</u> 47. 9	<u>51.8</u> <u>52.</u> 1	
	40-49	14. <mark>9</mark> 8			M/ONH/PI	0.4 5.3	40-49	47.7	52.3	
	50-64	20.5			White	56	50-64	48. 5 6	51. <mark>5</mark> 4	
	65-74	7. <mark>8</mark> 3			<u>M/O</u>	6	65-74	49 48.7	51 <u>.3</u>	
	75-84	4 <u>.1</u> <u>3.</u> 7					75-84	45.7	54.3	
	85+	2.4 <u>1</u>					85+	33.9<u>34.</u> 4	<u>66.165.</u> 6	
Death s	0-4	<0.1	Males	54.3	H/L	12.2	0-4	47.6	<u>52.4</u>	
	5 17	0.1	Female s	45.7	AI/AN	4	5 17	57.7	42.3	
	18-29	0.5			Asian	4 .3	18-29	63	37	
	30-39	1.1			Black	14.7	30-39	66	34	
	4 0 -49	2.8		ļ	NH/PI	0.2	4 0 -49	66-5	33.5	
	50-64	14.5			White	<u>63.1</u>	50-64	65	35	
	<u>65-74</u>	<u>21.3</u>			M/O	4.4	<u>65-74</u>	<u>61.4</u>	<u>38.6</u>	
	75-84	27.7					75-84	55.8	44.2	
	85+	32.1					85+	4 1.8	58.2	

a. Percentage of missing demographic data varied by types of event and demographic.

b. Except for Hispanics/Latinos, all categories refer to non-Hispanics

Abbreviations: Al/AN=American Indian/Alaska Native, H/L=Hispanic/Latino, M/O=Multiple/Other, NH/PI=Native Hawaiian/Other Pacific Islander

Table 42. Distribution of Deaths (n=513,204) by Age, Sex, Race, and Cross-Tabulated Age and Sex -- United States as of 14 August 2021^a

Event	<u>Age</u> Group	<u>Age %</u>	<u>Sex</u>	<u>Sex %</u>	Race ^b	<u>Race</u> <u>%</u>	<u>Age</u> Group	<u>Males</u> <u>%</u>	<u>Females</u> <u>%</u>
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Deaths	<u>0-4</u>	<u><01</u>	Males	<u>54.2</u>	H/L	<u>18.5</u>	<u>0-4</u>	<u>51.5</u>	48.5
	5-11	<u><01</u>	Females	<u>45.8</u>	AI/AN	<u>12</u>	5-11	<u>55.9</u>	44.1
	12-15	<u><01</u>			Asian	<u>3.8</u>	12-15	<u>46.7</u>	<u>53.3</u>
	<u>16-17</u>	<u><01</u>			Black	13.8	<u>16-17</u>	68.7	<u>31.3</u>
	<u>18-29</u>	0.6			NH/PI	<u>02</u>	<u>18-29</u>	<u>64</u>	<u>36</u>
	<u>30-39</u>	<u>1.3</u>			White	<u>58.7</u>	<u>30-39</u>	<u>65.1</u>	<u>34.9</u>
	<u>40-49</u>	3.1			<u>M/O</u>	3.8	<u>40-49</u>	65.3	<u>34.7</u>
	<u>50-64</u>	15.4					<u>50-64</u>	<u>64</u>	<u>36</u>
	<u>65-74</u>	21.6					<u>65-74</u>	<u>60.6</u>	<u>39.4</u>
	<u>75-84</u>	27.3					<u>75-84</u>	55.5	44.5
	85+	30.7					85+	41.8	58.2

a. Percentage of missing demographic data varied by types of event and demographic

b. Except for Hispanics/Latinos, all categories refer to non-Hispanics

Abbreviations: AI/AN=American Indian/Alaska Native, H/L=Hispanic/Latino, M/O=Multiple/Other, NH/PI=Native Hawaiian/Other Pacific Islander

In general, disease has been much less severe among ages 0-24 compared to ages \geq 25 years, with 2.5% hospitalized, 0.8% admitted to an intensive care unit, and <0.1% dying among ages 0-24, versus 16.6% hospitalized, 8.6% intensive care, and 5% dying among ages \geq 25 years.²³ Among hospitalized cases with COVID-19 in the US, approximately 90% are over 40 years old, and between 58% to 66% are at least 60 years old.²⁴ The majority (approximately 60%) of COVID-19 patients admitted to hospitals in the US have been male.^{24,25,26,27,28}

African American COVID-19 patients have been reported to have an increased risk of hospitalization^{25,29} and mortality,³⁰ compared to white patients in the United States. A CDC report examined demographic trends among US COVID-19 deaths from May to August of 2020.³¹ During the observation period, the percentage of US COVID-19 deaths that were Hispanic increased from 16.3% in May to 26.4% in August, the only racial or ethnic group among whom the percentage of deaths increased during that time. In terms of setting, 64.3% of deaths occurred in inpatient hospitals and 21.5% in nursing homes or long-term care facilities.

As of 08 March 2021, the CDC estimated that The most recent CDC estimate of the total number of excess deaths (as opposed to overall deaths in the preceding paragraph) across the US from 0126 January 2020 to 27 February 2020 to the present2021 from all causes (COVID-19 and otherwise) ranged from 509,890 624,307.³² A CDC report examining US 545,600-660,200, with an estimated 75-88% of excess deaths being associated with race and age, restricted to the period 26 January 2020 to 03 October 2020, estimated that 66% of USCOVID-19.³³ An earlier CDC report on excess deaths covering 26 January 2020 through 3 October 2020 broke down excess deaths by demographics³³: by age during that period-were attributable to COVID 19.³⁴ By age, the largest increase in deaths compared to average expected deaths occurred among adults aged 25-44 (26.5% increase). By race, increases in deaths compared to expectation were largest among Hispanics (53.6% increase), Asian Americans (36.6% increase), African Americans (32.9% increase), and Native Americans and Native Alaskans (28.9% increase), all compared to an excess 11.9% deaths among non-Hispanic whites.

While research earlier in the pandemic tended to focus on adults, more recent data have given greater attention to children and adolescents. For the period January 1-March 31 2021 across 14 states (the most recently available data), the CDC's COVID-NET database recorded 204 adolescents aged 12-17 who were hospitalized for likely primarily COVID-19-related reasons.³⁵ The 204 adolescents were 47.5% male—consistent with the COVID case sex distribution across all ages—and disproportionately from minorities, with 31.4% Hispanic and 35.8% non-Hispanic African Americans.³⁵

Another recent CDC report described demographic trends in US COVID-19 incidence among 15,068 cases aged 0-24 years across 16 jurisdictions during the period 01 January 2020 through 31 December 2020.³⁶ The report broke down incidence by age groups and 2020 sub-periods that are presented in Table 43. The table shows that early in 2020, 5-9 year olds were experiencing less COVID-19 than 0-4 year olds, but by the end of the year this pattern had reversed. Compared to 5-9 year olds, the age categories 10-14, 15-19, and 20-24 years old showed progressively greater incidence rates, a pattern that held throughout 2020.

<25 years across three periods of 2020 in 16 U.S. jurisdictions ³⁰										
<u>2020</u>	Age Group	<u>Number</u>	Cases per 100,000 population	Rate Ratio						
Sub-Period	(years)	of Cases	<u>(95% CI)</u>	<u>(95% CI)</u>						
<u>Jan 1-Apr 30</u>	<u>0-4</u>	<u>956</u>	<u>21 (20-23)</u>	<u>1.28 (1.17-1.41)</u>						
	<u>5-9</u>	772	<u>17 (16-18)</u>	<u>Reference</u> ^a						
	<u>10-14</u>	<u>1,184</u>	<u>25 (23-26)</u>	<u>1.49 (1.36-1.63)</u>						
	<u>15-19</u>	<u>3,267</u>	<u>67 (65-70)</u>	<u>4.03 (3.72-4 36)</u>						
	<u>20-24</u>	<u>8,889</u>	<u>175 (171-178)</u>	<u>10.47 (9.72-11.26)</u>						
<u>May 1-Aug 31</u>	<u>0-4</u>	<u>14,017</u>	<u>314 (309-319)</u>	<u>1.01 (0.98–1.03)</u>						
	<u>5-9</u>	14,406	<u>312 (307-317)</u>	Reference ^a						
	<u>10-14</u>	20,490	<u>430 (424-436)</u>	<u>1.38 (1.35–1.41)</u>						
	<u>15-19</u>	50,210	<u>1,034 (1,025-1,043)</u>	3.32 (3.26–3.38)						
	<u>20-24</u>	78,655	<u>1,547 (1,536-1,557)</u>	<u>4.96 (4.88–5.05)</u>						
Sep 1-Dec 31	<u>0-4</u>	33,595	<u>752 (744–760)</u>	0.71 (0.70-0.72)						
	<u>5-9</u>	48,824	<u>1,056 (1,047–1,066)</u>	<u>Reference</u> ^a						
	<u>10-14</u>	76,922	<u>1.615 (1.604–1.627)</u>	<u>1.53 (1.51–1.55)</u>						
	<u>15-19</u>	149,660	<u>3,083 (3,067–3,098)</u>	2.92 (2.89–2.95)						
	<u>20-24</u>	187,825	<u>3,693 (3,677–3,710)</u>	<u>3.50 (3.46–3.53)</u>						

Table 43. COVID-19 incidence and rate ratios, by age group among persons aged <25 years across three periods of 2020 in 16 U.S. jurisdictions³⁶

a Reference to imply that incidence rate in 5-9 year-old age group is used as comparison to calculate rate ratios for other age groups

Other US pediatric data are generally consistent with the CDC findings. Table 44 summarizes demographic results for a retrospective cohort of 135,794 individuals under the age of 25 who were tested for COVID-19 by 08 September 2020 within the PEDSnet network of US pediatric health systems.³⁷ The Table 44 shows that, among the pediatric population, children age 12-17 were more frequently infected than those under age 12. African Americans and Hispanics had elevated frequencies of testing positive relative to their proportion of the cohort.

A study of 1,945,831 individuals aged 0-18 recorded in the Premier Healthcare Database between March and October 2020 included 20,714 pediatric cases of COVID-19; the authors reported similar patterns to what is shown in Table **43**, with the additional observation that

COVID-19 cases aged 0-1 and 12-18 years were more likely to develop serious illness than those aged 2-11.³⁸

Table 44. Demographics of 135,794 US individuals under age 25 tested for COVID-19 by 08 September 2020³⁷

Characteristic	Patients, n (%)				
	COVID-19 negative	COVID-19 positive,	COVID-19 positive,		
	(n=130,420)	Asymptomatic or mild	Severe illness (n=359)		
		<u>illness (n=5,015)</u>			
Age, years					
<1	17,431 (13)	494 (10)	<u>72 (20)</u>		
1-4	32,619 (25)	808 (16)	<u>40 (11)</u>		
5-11	35,617 (27)	<u>1,029 (21)</u>	<u>72 (20)</u>		
12-17	32,362 (25)	1,521 (30)	117 (33)		
18-24	12,391 (10)	<u>1,163 (23)</u>	<u>58 (16)</u>		
Sex					
Female	<u>61,637 (47)</u>	2,527 (50)	<u>172 (48)</u>		
Male	68,701 (53)	2,485 (50)	<u>187 (52)</u>		
Other or Unknown	<u>82 (0.06)</u>	<u>3 (0.06)</u>	<u>0</u>		
Race/ethnicity					
Hispanic	<u>14,156 (11)</u>	<u>918 (18)</u>	<u>108 (30)</u>		
API	<u>4,471 (3)</u>	<u>151 (3)</u>	<u>9 (3)</u>		
Black or AA	18,646 (14)	1,424 (28)	<u>119 (33)</u>		
White	77,540 (60)	1,988 (40)	<u>97 (27)</u>		
Multiple	3,883 (3)	126 (3)	<u>5 (1)</u>		
Other or Unknown	<u>11,724 (9)</u>	<u>408 (8)</u>	<u>21 (6)</u>		

AA=African American, API=Asian or Pacific Islander

Risk Factors

While anyone can become infected with SARS-CoV-2, symptoms of COVID-19 disease can range from very mild (or no symptoms) to severe or fatal. A person's risk of initial infection increases through spending time in close physical proximity to others, especially in indoor spaces with poor ventilation.³⁹ People living in long-term care facilities or high-density apartment homes, or working in occupations with close proximity to others (e.g. healthcare, transportation), have a higher risk of infection.^{39,40, 39,41} Among children, the primary source of infection is an infected adult living in the same household.⁴² According to the CDC, people ages 18 29some ethnic minority groups have the highesta higher risk of infection, but age is not associated with risk of initial infection, while children age 4 among people aged 5 and under have the lowest rateolder (Table 37).⁴³ Risk of infection is also higher among some ethnic minority groups.^{-, 5}Table 45).^{46, 47}

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Table 45. Risk for COVID-19 Infection, Hospitalizationinfection, Hospitalisation, and Death by Age Group ³⁶-and by Race/Ethnicity-³⁷⁴⁷

	Rate ratios ^a				
Age Group (years)	CasesCases ^c	Hospitalization Hospitalisation ^d	DeathDeathe		
0-4	<1	<u>2<1</u>	<u>2<1</u>		
<u>-5-17 - 5-17</u>	1	<u>≤</u> 1	<u>≤</u> 1		
<u>-18-29 18-29</u> ^b	<u>31</u>	7 <u>1</u>	<u> 451</u>		
30-39	<u>21</u>	<u>+++2</u>	4 <u>54</u>		
40-49	<u>21</u>	<u>152</u>	<u>13010</u>		
50-64	<u>21</u>	<u>254</u>	400 <u>35</u>		
65-74	<u>21</u>	<u>356</u>	<u>110095</u>		
75-84	<u>21</u>	<u>559</u>	2800 <u>230</u>		
85+	<u>21</u>	<u>8015</u>	7900<u>600</u>		
Race/Ethnicity					
Non-Hispanic White bf	1	1	1		
American Indian or Alaska Native,	1. 9 7	3.7 <u>4</u>	2.4		
non-Hispanic					
Asian, non-Hispanic	0.7	1.4 <u>0</u>	1.0		
Black or African American, non-	11	2. <u>98</u>	1.9 2.0		
Hispanic					
Hispanic or Latino	1. 3 9	3. 2.8	2.3		

na. Rates are expressed as whole numbers, with values less than 10 rounded to the nearest integer, two-digit numbers rounded to nearest multiple of five, and numbers greater than 100 rounded to two significant digits.

b. Rate ratios for each age group are relative to the <u>5-1718-29</u>-year age category. <u>This group was selected</u> as the reference group because it has accounted for the largest cumulative number of COVID-19 cases compared to other age groups.

bc. Includes all cases reported by state and territorial jurisdictions (accessed on July 12, 2021). The denominators used to calculate rates were based on the 2019 Vintage population (https://www.census.gov/newsroom/press-releases/2019/popest-nation.html).

d. Includes all hospitalizations reported through COVID-NET (from March 1, 2020 through July 3, 2021, accessed on July 12, 2021). Rates were standardized to the 2020 US standard COVID-NET catchment population (https://www.cdc.gov/coronavirus/2019-ncov/covid-data/covid-net/purpose-methods.html).

e. Includes all deaths in National Center for Health Statistics (NCHS) provisional death counts (accessed on July 12, 2021). The denominators used to calculate rates were based on the 2019 Vintage population (https://data.cdc.gov/NCHS/Provisional-COVID-19-Deaths-by-Sex-and-Age/9bhg-hcku).

f. Rate ratios for each race/ethnicity group are relative to the Non-Hispanic White category.

Risk for severe or fatal COVID-19 disease has been shown to increase with older age, male sex, or ethnic minority status.³⁶.__46.37.38.48.49.50_Risks^{47,5152,53,54} Children aged 5-17 typically experience a milder disease course and have lower risk of hospitalization andor death.^{46,55,56}Among adults, these risks increase dramatically for every 10-year age group above age 1739 (Table 37Table 42).^{36,41,46,57} Table 37Table 45 also gives estimated rate ratios for COVID-19 hospitalizationhospitalisation and death by race/ethnicity relative to white, non-Hispanic persons in the US. The highest risks of hospitalizationhospitalisation and death were observed among American Indian or Alaska native persons (RR = 3.74 for hospitalizationhospitalisation and 2.4 for death) and Hispanic or Latino persons (RR = 3.2.8 for hospitalizationhospitalisation and 2.3 for death). These differences in risk among ethnic groups may be attributed to differences in underlying factors that are correlated with

race/ethnicity including socioeconomic status, access to health care, and occupation-related virus exposure.³⁷⁴⁷

Risk of severe or fatal COVID-19 disease is higher among persons who are current or former smokers, have lower socioeconomic status, have no or public insurance, or live in neighborhoods with higher rates of limited English proficiency.^{38,40,41,58} ^{51,53,57} The CDC has also recognized other socio-demographic groups who may need to take extra precautions against COVID-19 due to increased risk for severe illness: pregnant women; breastfeeding mothers; people with disabilities-or developmental/behavioral disorders; people living in rural communities, nursing homes, long term care facilities, or prisons; people experiencing homelessness; people with developmental, behavioural, or substance abuse disorders; and newly resettled refugee populations.⁵⁹

RiskAmong adults, risk for severe or fatal COVID-19 disease also-increases with the presence of chronic medical conditions, including obesity, respiratorychronic lung diseases (e.g., COPD or asthma), cardiovascular disease, diabetes, cancer, liver disease, neurological diseases (e.g., stroke or dementia), chronic kidney disease, sickle cell disease, autoimmune conditions and immunosuppression, or<u>HIV</u>, higher scores on the WHO Clinical Progression Scale and Charlson Comorbidity Index.^{38,39,40,41,4252,57,60,51,53}. Table 38 Table 46 shows the estimated hazard ratios of COVID-19 mortality associated with these chronic conditions and socio-demographics from a cohort study of 17 million adults in England.⁴¹(with 17,000 COVID-19-related deaths) in England.⁵⁷

The presence of one or more underlying medical conditions also increases risk of severe or fatal disease among children aged 5-17.^{61,62,63,64} In particular, childhood obesity has been consistently associated with two to three times the risk of severe disease or hospitalization.^{61, 64,65,66,} For many other individual comorbid conditions, pediatric sample sizes are very small and different studies produce conflicting results, so it is difficult to estimate precise risk ratios based on current literature.^{42,63}

Table 46.	Hazard Ratios and 95% Confidence Intervals for COVID-19related
	Death ⁴¹⁵⁷

Characteristic	Category	COVID-19 death	Hazard Ratio
		Adjusted for	Fully adjusted
		age <u>, sex</u> , and <u>sexNHS</u> administrative region	
Age	18-39	0 05 (0 04-0 0706)	0 06 (0 04-0 08<u>07</u>)
-	40-49	0 <u>2832</u> (0 <u>2328</u> -0 <u>3338</u>)	0 30 <u>34</u> (0 25 <u>29</u> -
			0 36 <u>39</u>)
	50-59	1 00 (ref)	1 00 (ref)
	60-69	2 79 93 (2 52<u>69</u>-3 <u>1020</u>)	2 4 <u>057</u> (2 16<u>35</u>-
			2 66<u>80</u>)
	70-79	<u>9 17 (</u> 8 62 (7.84 <u>48</u> -	6 07 (5.51-<u>74</u>
		9 4 <u>693</u>)	<u>(6 6921-7 31</u>)
	80+	38 29 (35.02-	20.60 (18.70-<u>24</u> 10
		41.87)43 16 (40 03-	<u>(</u> 22 68 <u>23-26 13</u>)
		<u>46 53)</u>	

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Characteristic	Category	COVID-19 death Hazard Ratio		
		Adjusted for	Fully adjusted	
		age <u>, sex,</u> and <u>sexNHS</u>		
		administrative region		
Sex	Female	1 00 (ref)	1 00 (ref)	
	Male	1 78<u>73</u> (1 71<u>68</u>-1 8578)	1 59<u>55</u> (1 <u>5350</u>-	
			1 65 <u>60</u>)	
BMI (kg/m ²)	Not obese	1 00 (ref)	1 00 (ref)	
	30-34.9 (obese class I)	1 23 (1 17 <u>18-</u>1 <u>3028</u>)	1 05<u>07</u> (1 00 <u>0</u>3. 1 112)	
	35-39.9 (obese class II)	1 81<u>79</u> (1 68–<u>1</u> 95<u>90</u>)	1 40 <u>44</u> (1 30 <u>36</u> 1 <u>5254</u>)	
	40+ (obese class III)	2 66 76 (2 39 2 95<u>54-</u> 3 00)	<u>2 11 (1 92 (1.72</u> 93-2 13 29)	
Smoking	Never	1 00 (ref)	1 00 (ref)	
Smoning	Former	1 4 <u>344</u> (1 <u>37–40-</u> 1 49)	1 <u>+926</u> (1 <u>+4-22-</u> 1 24 30)	
	Current	1 14<u>17</u> (1 05 <u>10-</u> 1 23<u>25</u>)	0 89<u>97</u> (0 82 0.97 91-1 04)	
Ethnicity-	White	1 00 (ref)	1 00 (ref)	
	Mixed	1 62 59 (1 26 2.08 28-	1 43 (1 11-15-	
	Mixed	1 02 <u>55</u> (1 20 2.00 <u>28-</u> 1 97)	1 4 3 (1 11 <u>15</u> 1 84 78)	
	South Asian	1 <u>6997</u> (1 <u>54</u> <u>1.8482</u> 2 <u>14</u>)	1 4 <u>570</u> (1 <u>32–55-</u>	
	D11.		1 <u>5885</u>)	
	Black	1 <u>8882</u> (1 <u>65 61-</u> 2 <u>1405</u>)	1 48 <u>44</u> (1 29–<u>27-</u> 1 69<u>63</u>)	
	Other	1 37<u>38</u> (1 <u>13–17-</u> 1 6563)	1 33<u>38</u> (1 10–<u>16-</u> 1 6163)	
IMD quintile ^{ea}	1 (least deprived)	1 00 (ref)	1 00 (ref)	
1	2	1 16<u>17</u> (1 08<u>11</u>-1 23)	1 <u>1213</u> (1 05 <u>07-</u> 1 19)	
	3	1 31<u>37</u> (1 23 _30-	1 22<u>25</u> (1 15–<u>19-</u>	
		1 4 <u>044</u>)	1 30 <u>32</u>)	
	4	1 69<u>77</u> (1 59 <u>68-</u> 1 7986)	1 51<u>53</u> (1 42<u>46-</u> 1 61)	
	5 (most deprived)	2 11 (1.98 2 2501 2 22)	1 79<u>71</u> (1 <u>68 <u>62</u> <u>1 0180</u>)</u>	
D1 1	NT- march		1 <u>9180</u>)	
Blood pressure	Normal High BP or diagnosed	1 00 (ref)	1 00 (ref)	
	hypertension	1 09 (1 05 <u>06-</u>1 <u>14<u>13</u>)</u>	0 89<u>90</u> (0 85–<u>87-</u> 0 9394)	
Respiratory disease exc	21	1.95 (1.86–2.04)	1 63 <u>66</u> (1 55 – <u>59-</u>	
h d h d			1 71 <u>73</u>)	
Asthma ^b (vs. none)	With no recent OCS use	1 <u>1315</u> (1 07 – <u>10-</u>	<u>1 00 (</u> 0 99 (0 93)	
	W.1 0.000	1 <u>2021</u>)	<u>95-</u> 1 05)	
	With recent OCS use	1 <u>5561</u> (1 <u>39–47-</u>	1 <u>1315</u> (1 01 <u>05-</u>	
		1 73 75)	1 26)	
Chronic heart disease		1.57 (1.51–1.64)	1.17 (1.12 1.22)	
Diabetes ^{eb} (vs. none)	With HbA1c < 58 mmol/mol	1 58<u>53</u> (1 51 47. 1 66<u>59</u>)	1 <u>3120</u> (1 <u>24–16-</u> 1 <u>3725</u>)	
	With HbA1c \geq 58 mmol/mol	2 61 57 (2 46 -45-	1 95 83 (1 83	
		2 7770)	2.08 74-1 93)	

Table 46. Hazard Ratios and 95% Confidence Intervals for COVID-19-related Death-157

Table 46.	Hazard Ratios and 95% Confidence Intervals for COVID-19related
	Death ⁻¹⁵⁷

Characteristic Category		COVID-19 death	Hazard Ratio
		Adjusted for	Fully adjusted
		age <u>, sex,</u> and <u>sexNHS</u>	
		administrative region	
	With no recent HbA1c	2 27<u>19</u> (2 06 _02_	1 90<u>71</u> (1 72–
	measure	2 50<u>37</u>)	2.09<u>58-1</u>86)
Cancer (non-	Diagnosed <1 year ago	1 81<u>47</u> (1 58 2.07<u>31-</u>	1 72<u>44</u> (1 50 _28_
hematological, vs none)		<u>1 65</u>)	1 96 62)
	Diagnosed 1-4.9 years ago	1 20<u>13</u> (1 10 _04 -	1 15<u>11</u> (1 05 <u>0</u>3-
		1 <u>3222</u>)	1 27<u>20</u>)
	Diagnosed \geq 5 years ago	0 99 (0 93–<u>95-</u>1 <u>0604</u>)	0.96 (0.91–<u>2</u> 41
			<u>(1 0386-3 13</u>)
Hematological	Diagnosed <1 year ago	<u>2 54 (1 96-3 02 (2.24</u>	2 80 (2 08-3 78)
malignancy (vs. none)		4.0829)	
	Diagnosed 1-4.9 years ago	2 56 (28 (1 95-2 14 -	2 4 6 (25 (1 92-
		3.06<u>66</u>)	2 06 2.95<u>62</u>)
	Diagnosed \geq 5 years ago	1 70 71 (1 4 6 51 -	1 61 65 (1 39 46-
		1 98 93)	1 87)
Reduced kidney	eGFR 30-60	1 5650 (1 49-45-	1 33 30 (1 28 -25-
function ^{dc} (vs. none)		1 63 55)	1 40 <u>35</u>)
	eGFR <u><15-</u> < 30	3.48 (3.23 3.75)274	2 52 (2 33–2 72)
		<u>(2 56-2 93)</u>	
	eGFR <15 or dialysis	<u>6 40 (5 75-7 12)</u>	<u>4 42 (3 93-4 98)</u>
Liver disease		2 39 <u>27</u> (2 06 – <u>01–</u>	1 75 (1 51 2.03<u>54-</u>
		2 77<u>57</u>)	<u>1 98</u>)
Dementia		4 59 (4 33-4 87)	<u>3 62 (3 41-3 84)</u>
Stroke-or dementia		2 57 (<u>03</u> (1 95- 2 46 -	2.16 (2.06 -
		2.70<u>12</u>)	2 27)<u>1 53 (1 46-</u>
			<u>1 59)</u>
Other neurological disea	se	3 08<u>15</u> (2 85–<u>96-</u>	2 58 <u>72</u> (2 38 <u>55-</u>
		3 33 36)	2 79<u>90</u>)
Organ transplant		6.00<u>5 54</u> (4 73 -	3.53 (<u>1 61 (1 28-</u>
		7.61<u>51-681</u>)	2 77 4.49<u>02</u>)
Asplenia		1 62<u>50</u> (1 19 - 2 - 21<u>16-</u>	1 34<u>26</u> (0 98–<u>97-</u>
		<u>1 95</u>)	1 83<u>64</u>)
Rheumatoid arthritis, lug	ous, or psoriasis	1.30 (1.21–1.38)	1 19<u>23</u> (1 11–<u>17-</u>
			1 27<u>30</u>)
Other immunosuppressive condition		2.75 (2.10-3.62)	2 21<u>00</u> (1 68 <u>57-</u>
			2 90<u>54</u>)

a. Ethnicity hazard ratios were estimated from a model restricted to those with recorded ethnicity.

b. For OCS use, 'recent' refers to during the year before baseline.

ea. Classification by HbA1c is based on measurements the most recent measurement within 15 months of baseline

db. eGFR is measured in <u>mlmL</u> min⁻¹ per 1.73 m² and <u>takenderived</u> from the most recent serum creatinine measurement.

ec. Index of Multiple Deprivation (derived from the patient's postcode)

Models were adjusted for age using a four-knot cubic spline for age, except for estimation of age-group hazard ratios Ref, reference group; 95% CI, 95% confidence interval

The main existing treatment options:

Through 28 February 30 September 2021, other COVID-19 vaccines were authorized and recommended for use in the United States including vaccines from Moderna (NCT04470427), and Johnson & Johnson/Janssen (NCT04505722). Others may subsequently be approved.

Natural history of the indicated condition in the untreated population, including mortality and morbidity:

Symptoms of COVID-19

The clinical manifestations of COVID-19 vary widely, from asymptomatic infection in 17-20%, 67. 45% across age groups, 68, 69 70, 71 to critical illness and death. The rate of asymptomatic infection decreases with increasing age and long-term care facilities are associated with a lower rate of asymptomatic infection when compared to household transmission or other healthcare facilities.⁷¹ A recent meta-analysis has estimated that 46.7% of infections in children are asymptomatic.⁷¹ The most common symptoms of COVID-19 are fever, cough, and shortness of breath (Table 39).⁷²-for both children and adults (Table 47).⁷³ 74

Table 47. Signs and symptoms among 291 pediatric (age <18 years) and 10,944 adult (age 18-64 years) patients^a with laboratory confirmed COVID-19 -United States, 12 February- 2April 2020⁴⁶.73

	No. (%) with sign/symptom			
Sign/Symptom	Pediatric	Adult		
Fever, cough, or shortness of breath ^b	213 (73)	10,167 (93)		
Fever ^d	163 (56)	7,794 (71)		
Cough	158 (54)	8,775 (80)		
Shortness of breath	39 (13)	4,674 (43)		
Myalgia	66 (23)	6,713 (61)		
Runny nose ^c	21 (7.2)	757 (6 9)		
Sore throat	71 (24)	3,795 (35)		
Headache	81 (28)	6,335 (58)		
Nausea/Vomiting	31 (11)	1,746 (16)		
Abdominal pain ^e pain ^c	17 (5.8)	1,329 (12)		
Diarrhea	37 (13)	3,353 (31)		

Cases were included in the denominator if they had a known symptom status for fever, cough, shortness of breath, nausea/vomiting, and diarrhea. Total number of patients by age group: <18 years (N = 2,572), 18-64 years (N = 113,985).

Includes all cases with one or more of these symptoms. b.

Runny nose and abdominal pain were less frequently completed than other symptoms; therefore,

percentages with these symptoms are likely underestimates.

d. Patients were included if they had information for either measured or subjective fever variables and were considered to have a fever if "yes" was indicated for either variable

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Progression and Timeline of Mild to Moderate Disease

Mild to moderate disease is defined as the absence of viral pneumonia and hypoxia. For those who develop symptoms, the incubation period is usually 4 to 5 days, with 97.5% experiencing symptoms within 11 days of exposure.^{75,76} Those with mild COVID-19 recover at home with supportive care and guidance to self-isolate. Those with moderate disease are monitored at home and are sometimes recommended to be hospitalized if conditions worsen-⁴⁸.⁷⁶ Data on rates of re-infection are limited but variants that are not neutralized by immune antisera, such as the recent <u>beta (South African)</u> variant, may lead to increased risk of re-infection in the future-⁴⁷.⁷⁵

Progression and Timeline of Severe Disease Requiring Hospitalization

Those with severe disease will require hospitalization to manage their illness. Based on data that have been systematically collected for the US by the CDC between 01 August 2020 and 02 March05 September 2021, there were 1,814,6062,816,280 new hospital admissions for patients with confirmed COVID-19 in the US.⁷⁷ For the week ending 28 February22 August 2021, 103.5 patients per 100,000 population were hospitalizedhospitalised due to COVID-19 in 2221 countries of the EU/EEA with available data.⁷⁸ Based on data from 23 states and New York City, as of August 19, 2021, 1.6%-3.6% of children with COVID-19 have been hospitalised and 0.0-0.03% of children with COVID-19 have died.⁷⁹

The most common symptoms in patients are fever (42-80%), shortness of breath (35-71%), fatigue (33-62%), cough (77-84%), chills (63%), myalgias (63%), headache (59%), and diarrhea (33%).^{80,81,82,83} COVID-19 patients also commonly experience gustatory disorders (44%) and olfactory disorders (53%).⁸⁴ Among unhospitalised children < 18 years of age, 89% experienced one or more typical symptoms of COVID, including fever, cough, shortness of breath, and 22% experienced all three.⁸¹Approximately 17% to 40% of those hospitalizedhospitalised with COVID-19 experience severe symptoms necessitating intensive care, 23 , 24,28,51 , 29,80 with 31% of children hospitalised experiencing severe COVID-19 that necessitates intensive care or invasive ventilation or ends in death. Risk factors for severe COVID-19 in hospitalised children include presence of a comorbid condition, younger age, and male sex.³⁸ More than 75% of patients hospitalizedhospitalised with COVID-19 require supplemental- oxygen.⁸⁵

Studies early in the pandemic demonstrated that time from onset of illness to ARDS was 8-12 –days and time from onset of illness to ICU admission was 9.5–12 days.⁴⁷.⁷⁵ In 17 countries of the EU/EEA with available data, 1.8 patients per 100,000 population were in the ICU due to COVID-19 for the week ending 28 February 2021.⁵⁹.⁸⁶ A recent meta-analysis found that, of patients <19 years of age, 11% went to the ICU, non-invasive ventilation was administered among 12%, and 4% required mechanical ventilation.⁴⁵...⁶⁹

Mortality

As of $\frac{07 \text{ March} 17 \text{ August}}{2021}$, there were $\frac{522,973620,493}{22,973620,493}$ deaths reported in the US for all age groups among $\frac{28,771,74936,951,181}{2021}$ cases $(1.\frac{87}{7}\% \text{ of cases})$.⁴⁹).⁸⁷ As of $\frac{28 \text{ February} 17}{2021}$ August 2021 there were $\frac{547,267746,566}{247,267746,566}$ deaths reported for all age groups in the EU/EEA

among 22,527,37035,381,520 cases (2.41% of cases).⁸⁸ As of 7 March17 August 2021, the UK has seen 124,736131,466 deaths from COVID-19 in all age groups among 4,231,1666,352,224 cases (2.91% of cases).⁸⁹ According to a recent meta-analysis of pediatricpaediatric studies published through October 2020, the mortality for paediatric patients <19 is 0.1-2%.^{69,23} In a study from January through June 2020 using the National Child Mortality Database (NCMD) in England, 5.7% of 437 children 0-17 years of age is 2%.⁴⁵ who died were SARS-CoV-2 PCR-positive and those who died of COVID-19 were older and were more likely to be non-White ethnicity.⁹⁰

Mortality data are also presented from Worldometer, an independent organizationorganisation that publishes current, reliable COVID-19 statistics online.⁴⁷¹⁴ The mortality of SARS-CoV-2 infection is defined as the cumulative number of deaths among detected cases.

As of 03 March15 August 2021, the overall SARS-CoV-2 mortality for the EU + UK was 677,146-878,344 deaths, or 132171 per 100,000 people. Reported mortality among EU countries and the UK ranged from 1418 to 195312 deaths per 100,000 (Table 35). Table 40). Finland and Cyprus reported the lowest mortality; <u>Hungary</u>, Czech Republic, <u>Belgium</u> and <u>SloveniaBulgaria</u> reported the highest.⁴⁵¹⁵

Overall reported mortality among hospitalized hospitalised COVID-19 patients varies from 12.8% to 26% in the EU-and UK.²⁸, UK, and US.^{29,30,91,92}_31,93,94. Mortality rates are declining over time, presumably due to an improved understanding of COVID-19 and its management.^{58,95}.96

Complications of COVID-19 and Long-COVID

Complications of COVID-19 include impaired function of the heart, brain, lung, liver, kidney, and coagulation system. $\frac{24,27,97}{24,27,97}$ Based on a meta-analysis of 42 studies, the risk of thromboembolism was 21% overall and 31% in the ICU, with the pooled odds of mortality being 74% higher among those who experienced thromboembolism compared to those who did not.⁹⁸

COVID-19 symptoms can persist weeks or months beyond the acute infection.^{99,100} The NICE guideline scope published on 30 October 2020 defined "Long COVID" signs and symptoms that continue or develop after acute COVID-19. It includes both ongoing symptomatic COVID-19 (from 4 to 12 weeks) and post-COVID-19 syndrome (12 weeks or more and for which signs and symptoms are not explained by an alternative diagnosis).¹⁰¹

A meta analysis of 31 studies among patients between 18 to 49 years of age found that COVID-19 symptoms were experienced for 14 days to 3 months post-infection, including persistent fatigue (39–73%), breathlessness (39–74%), decrease in quality of life (44–69%), impaired pulmonary function, abnormal CT findings including pulmonary fibrosis (39–83%), evidence of peri-/perimyo-/myocarditis (3–26%), changes in microstructural and functional CONFIDENTIAL

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brain integrity with persistent neurological symptoms (55%), increased incidence of psychiatric diagnoses (5.8% versus 2.5–3.4% in controls), and incomplete recovery of olfactory and gustatory dysfunction (33–36%).¹⁰² Post-acute COVID symptoms in children with asymptomatic or mild disease appear to be less severe than in adults, with the most common symptoms being a post-viral cough (4%), fatigue (2%), or both symptoms (1%) with the duration of symptoms lasting 3 to 8 weeks.¹⁰³

Children who are infected with COVID-19 are at risk of subsequent multisystem inflammatory syndrome (MIS-C) and often develop a rash following resolution of COVID-19, ^{101,105,45} 6^{9,106,107} As of August 19, 2021 there were 4,403 cases of MIS-C reported to health departments in the US. ¹⁰⁸ Additional symptoms of MIS-C include abdominal pain, bloodshot eyes, chest tightness or pain, diarrhea, lethargy, headache, low blood pressure, neck pain, and vomiting. ¹⁰⁹

Important co-morbidities:

Important comorbidities in hospitalized COVID-19 patients include hypertension, diabetes, obesity, cardiovascular disease, chronic pulmonary disease or asthma, chronic kidney disease, cancer, and chronic liver disease.^{24,25,26,51,54}.^{25,26,27,80,83}. Prevalence of these conditions have been reported to be lower in mild cases and higher among fatal cases, as shown as shown for EuropeanEU/EEA countries in Table 40 Table 48 below-using TESSy data posted on 12 August 2021.¹¹⁰

Table 48. Preconditions among COVID-19 Patients in EU/EEA-and-UK, by Severity of Disease. Case-based Data from TESSy Produced 04 MarchReported 12 August 2021¹¹⁰

	EU/EEA, producedreported on 04 March12 August 2021			
	Mild	HospHospitalised	Severe	Fatal
Total N	1, 155,969 948,252	214,784<u>356,472</u>	35,46852,365	67,011<u>109,878</u>
Asplenia (%)	0	0	0	0
Asthma (%)	0 5 6	1. <u>62</u>	1. <u>73</u>	1. <u>62</u>
Cancer, malignancy (%)	2<u>3</u>.1	7.2<u>9.1</u>	9.7<u>10</u>	<u>9.311.1</u>
Cardiac disorder, excluding	6.2 9.1	18.4 23.7	20.7 22.8	24.7 29.4
hypertension (%)				
Chronic lung disease, excluding	1.8	<u>4.73.6</u>	53<u>4.4</u>	5. 3 <u>.6</u>
asthma (%)				
Current smoking (%)	0.9	0. <u>31</u>	0.4 <u>2</u>	0.1
Diabetes (%)	3 3<u>5</u>	13.9<u>17.1</u>	18 9 20.5	15.6 19.2
Haematological disorders (%)	0	0. <u>32</u>	0.1	0. <u>21</u>
HIV/other immune deficiency (%)	0 <u>+2</u>	0. <mark>97</mark>	<u>+0.7</u>	0. <mark>8<u>5</u></mark>
Hypertension (%)	0. <u>78</u>	<u>32</u> 9	4.4 <u>3.2</u>	6. 3 <u>.8</u>
Kidney-related condition, renal	0.3	2.3<u>1.8</u>	2-2<u>1.9</u>	<u>32</u> .7
disease (%)				
Liver-related condition, liver disease	0 2 3	0.7	0.7	0.6
(%)				
Neuromuscular disorder, chronic	0. 6 7	2.4<u>1.8</u>	1. <u>64</u>	<u>2.</u> 4 .2
neurological (%)				
Obesity (%)	0 2 1	0 2	0.4 <u>5</u>	0.2

Table 48. Preconditions among COVID-19 Patients in EU/EEA-and UK, by Severity of Disease. Case-based Data from TESSy Produced 04 MarchReported 12 August 2021¹¹⁰

EU/EEA, produced reported on 04 March12 August 2021			
Mild	HospHospitalised	Severe	Fatal
0.4 <u>3</u>	0 2	0.1	0.1
0	0	0	0
0	0	0	0
<u>82.576.7</u>	<u>42.8</u> 36.7	<u>32.73</u>	27.3 25
	Mild 0.4 <u>3</u> 0 0	Mild HospHospitalised 0.43 0.2 0 0 0 0 0 0	Mild HospHospitalised Severe 0.43 0 2 0.1 0 0 0 0 0 0

Abbreviation: nosp = nospita

Table 41

Table 49 below summarizes comorbidities among US COVID-19 patients in a retrospective cohort study conducted among 629,953 individuals tested for COVID-19 in a large health system in the US Northwest between 01 March and 31 December 2020.³⁸⁵¹ The most common comorbidities were similar in the full cohort and among those who tested positive: obesity, hypertension, diabetes, and asthma. Among those hospitalized for COVID-19, a large number of comorbidities had elevated prevalence compared to the full cohort and those who tested positive: obesity, hypertension, diabetes, kidney disease, congestive heart failure, coronary artery disease, and chronic obstructive pulmonary disease.

Table 49.Comorbidities in individuals tested for COVID-19 in the Providence St.
Joseph Health System – States of California, Oregon, and Washington,
01 March–31 December 202038 51

	Tested (N= 629,953)	Positive (N= 54,645)	Hospitalized (N= 8,536)
Comorbidity	%	%	%
Hypertension	23.3	19.8	40.2
Diabetes	9.4	10.9	28.3
Weight			
Underweight	2.1	1.7	3.1
Normal	29.0	23.9	24.3
Overweight	31.7	32.6	30.3
Class 1 Obesity	19.8	22.3	21.2
Class 2 Obesity	9.6	11.1	10.9
Class 3 Obesity	7.7	8.6	10.3
Asthma	6.5	5.3	6.7
Chronic Obstructive Pulmonary Disease	4.0	2.6	8.3
Coronary Artery Disease	5.5	3.6	9.7
Myocardial Infarction	2.2	1.6	5.5
Congestive Heart Failure	5.3	3.9	13.2
Kidney Disease	5.6	5.3	17.2
Liver Disease	3.1	2.5	4.0
Cancer	6.1	3.0	6.3

In a retrospective cohort of 135,794 individuals under the age of 25 who were tested for COVID-19 by 08 September 2020 within the PEDSnet network of US pediatric health systems, the proportion of obese individuals was similar among those who tested negative (18%) and among mild or asymptomatic COVID-19 cases (19%), but clearly elevated among severe COVID-19 cases (37%).³⁷ Those with severe cases of COVID-19 more commonly had chronic conditions in at least two body systems, with 25% of COVID-19 negative individuals, 17% mild or asymptomatic cases, and 38% of severe cases having multiple chronic conditions.

More recent data provide insight into comorbidities among the pediatric population. For the period January 1-March 31 2021 across 14 states, the CDC's COVID-NET database recorded 204 adolescents aged 12-17 who were hospitalized for likely primarily COVID-related reasons.³⁵ Among the 204 adolescents, 70.6% had at least one major underlying medical condition, the most common conditions being obesity (35.8%), chronic lung diseases including asthma (30.9%), and neurologic disorders (14.2%).³⁵

2.1.2.f. Pharmacological Class Effects

There are 2 vaccines (including BNT162b2) with a mRNA platform authorized for emergency use in multiple US jurisdictions since 11 December 2020. Theoretical concerns in mRNA vaccines have included the risk of the presence of naked extracellular RNA in the body which may lead to edema or coagulation and concerns about aberrant immune responses to the RNA or lipid particles. The immunogenicity and efficacy data from study C4591001 are indicative of the vaccine delivery system's success in transfecting the RNA into the appropriate target cells to stimulate an immune response. The RNA itself cannot integrate into the DNA genome.^{111,112} The probability of any sequences from the vaccine RNA being integrated into the human genome by a reverse transcription mediated mechanism is considered remote, no higher than the probability of host RNA sequences being re-inserted into the genome, especially given the small quantity of RNA in the vaccine, the barriers to transfected RNA reaching the nucleus, the non-replicating nature of the vaccine RNA, the limited stability of RNA in a cellular context, and the expected targeting of transfected cells for elimination by T cells elicited by the vaccine antigen expressed from the RNA.

3. PHARMACOVIGILANCE PLAN

3.1. Structure of the Pharmacovigilance Plan

3.1.1. Summary of Ongoing Safety Concerns

Table 50. Ongoing Safety Concerns

Important Identified Risks	Anaphylaxis
	Myocarditis and Pericarditis
Important Potential Risks	Vaccine-associated enhanced disease (VAED) including Vaccine- associated enhanced respiratory disease (VAERD)
Missing Information	Use in pregnancy and lactation
	Vaccine effectiveness
	Use in pediatric individuals $<\frac{125}{2}$ years of age [§]

Table 50. Ongoing Safety Concerns

Important Identified Risks	Anaphylaxis	
	Myocarditis and Pericarditis	
§ Missing information has been reworded to reflect the current state.		

3.1.2. Routine Pharmacovigilance Practices

- Routine pharmacovigilance activities <u>isare</u> a critical component of activities relating to the detection, assessment, understanding and prevention of risks. The objective of routine pharmacovigilance is to have processes in place to assure the ongoing and timely collection, processing, follow-up, and analysis of individual AE reports globally, following global safety Standard Operating Procedures and regulatory guidance.
- Pfizer, on behalf of the marketing authorization applicant (MAAholder(MAH), monitors the safety profile of its products, evaluates issues potentially impacting product benefitrisk profiles in a timely manner, and ensures that appropriate communication of relevant information is conveyed in a timely manner to regulatory authorities and other interested parties as appropriate and in accordance with international principles and prevailing regulations.
- Pfizer, on behalf of the MAAMAH, conducts scientific data gathering activities for the detection and evaluation of AEs in order to ensure safety monitoring, which is commensurate with product characteristics.
- Signal detection activities include periodic literature review for the life cycle of the product. This includes reviewing the medical literature for individual case reports that should be entered into the safety database as well as periodic aggregate literature review for broader signal detection.
- Safety signal evaluation requires the collection, analysis, and assessment of information to evaluate whether there is a potential causal association between an event and the administration of the product and includes subsequent qualitative or quantitative characterization of the relevant safety risk to determine appropriate pharmacovigilance and risk mitigation actions.
- Routine pharmacovigilance activities will include the use of DCAs. They are intended to facilitate the capture of clinical details about:
 - the nature and severity of COVID-19 illness in individuals who have received the COVID-19 vaccine and is anticipated to provide insight into potential cases of vaccine lack of effect or VAED.
 - potential anaphylactic reactions in individuals who have received the COVID-19 vaccine.
- A web-based AE reporting portal will be available for vaccine providers and recipients, to assist with anticipated high volume of reports (based on expected large target

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population). The portal will capture key adverse event data in the initial interaction and will provide automated intake into the Pfizer safety database via E2B for safety review.

• At the country level, the Drug Safety Unit performs routine pharmacovigilance activities including the collection of AEs from various sources and the reporting of AEs to the regulatory authority as per local regulatory guidelines.

3.1.3. Action Plan for Safety Issues

Action Plan for Important Identified Risks

Table 51. Ac	ction Plan for II	nportant Identified	Risk "Myocardi	tis and Pericarditis"
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Actions proposed	• Communication of this important identified risk via label (Section 5.2 - <i>Myocarditis and Pericarditis</i> , Section 6.2 - <i>Post Marketing Experience</i>).
	 C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-vaccine in the United States.
	C4591011: Active safety surveillance of the Pfizer-BioNTech COVID-19 vaccine in the US Department of Defense population following Emergency Use Authorization.
	C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer-BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
	C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
	• <u>C4591038 (former, C4591021 substudy: Post Conditional approval active</u> surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine. Substudy to investigate natural history of post-vaccination myocarditis and pericarditis.
	 C4591036: Pediatric Heart Network Study: Working title: MyocarditisLow interventional cohort study of myocarditis/pericarditis follow up study within the Pediatric Heart Networkassociated with Comirnaty in persons less than 21 years of age.
	 C4591031 substudy <u>B</u>: A <u>Phase 3</u>randomized, placebo-controlled, observer- blind, cross-over substudy of 1000 participants with documented receipt of 2 prior-to evaluate the safety and tolerability of a booster (third) dose of BNT162b2. Participants ≥12 years of age to ≤30 µgyears of age who have completed a 2-dose primary series of BNT162b2 (30 µg doses of BNT162b2 (the second dose received) at least 6 months ago), 16 to 30 years of age (randomized 1:1 in a crossover design to receive 30 µg BNT162b2 or placebo at baseline and the alternative 4 weeks later),(≥12 months for those 12-17 years of age) prior to randomization will be enrolled.
	 C4591007 substudy <u>— Troponin group</u>: A Phase 3 substudy of 750 participants 5 to <12 years of age (randomized 2:1 to receive BNT162b2 10 µg or placebo) and 500 participants 12<u>-15_<16</u> years of age (open label receipt of BNT162b2 30 µg).
Objective of proposed actions	 Labelling communicates the risk of myocarditis and pericarditis. C4591009: To assess the occurrence of safety events of interest, including myocarditis and pericarditis, in the general US population of all ages, pregnant women, the immunocompromised and persons with a prior history of COVID-19 within selected data sources participating in the US Sentinel System.
	 C4591011: To assess whether individuals in the US DoD Military Health System (MHS) experience increased risk of safety events of interest, including myocarditis and pericarditis, following receipt of the Pfizer-BioNTech COVID-19 Vaccine.
	 C4591012: To assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, including

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	myocarditis and pericarditis, following receipt of the Pfizer-BioNTech COVID-19 Vaccine.
	 C4591021: To assess the potential increased risk of adverse events of special interest (AESI), including myocarditis/pericarditis after being vaccinated with COVID-19 mRNA vaccine.
	 <u>C4591038 (former, C4591021 substudy÷)</u>: To describe the natural history of post-vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes within 1 year of myocarditis/pericarditis diagnosis among individuals vaccinated with BNT162b2 as well as individuals not vaccinated with a COVID-19 vaccine.
	 C4591036: Pediatric Heart Network Study: To characterize the clinical course, risk factors, resolution, long-term sequelae, and quality of life in children and young adults <21 years with acute post-vaccine myocarditis/pericarditis.
	 C4591031 substudy <u>B</u>: To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of subclinical myocarditis amongst individuals <u>1612</u> to 30 years of age.
	 C4591007 substudy <u>– Troponin group</u>: To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of subclinical myocarditis amongst individuals 5 to <u>45<16</u> years of age.
Rationale for proposed actions	 Labeling communicates to health care provider the risk of myocarditis and pericarditis.
	 C4591009 and C4591021: Robust surveillance is needed to ensure comprehensive understanding of real-world safety of the Pfizer-BioNTech COVID-19 Vaccine in large samples of general US (C4591009) and EU (C4591021) populations and in subcohorts of interest, including pregnant women, immunocompromised individuals and persons with a prior history of COVID-19 infection.
	 C4591011 and C4591012: Robust surveillance is needed to ensure comprehensive understanding of real-world safety. This surveillance strategy consists of complementary approaches to ensure timely signal identification and evaluation in populations who have received the Pfizer-BioNTech COVID-19 Vaccine under an Emergency Use Authorization (EUA).
	 <u>C4591038 (former, C4591021 substudy:)</u>: Study is needed to describe natural history of myocarditis and pericarditis in persons after vaccination with Pfizer-BioNTech COVID-19 vaccine and in unvaccinated persons. <u>Use in persons <12 will be captured as age is not a criterion for study eligibility.</u>
	 C4591036: Pediatric Heart Network Study: Need to collect evidence of safety of patients <21 years presenting to PHN sites after receiving a first or second dose of a COVID-19 vaccine and who were diagnosed with myocarditis/pericarditis
	• C4591031 substudy <u>B</u> : The first ~4 days post-vaccination is the time period when symptomatic myocarditis cases have most frequently been reported. Elevated Troponin I may be an indicator of subclinical myocarditis.
	 C4591007 substudy <u>— Troponin group</u>: The first ~4 days post-vaccination is the time period when symptomatic myocarditis cases have most frequently

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	been reported. Elevated Troponin I may be an indicator of subclinical myocarditis.
Monitoring by the sponsor for safety issue and proposed actions	 C4591009: Post-approval observational studies using real-world data are needed to assess the association between Pfizer-BioNTech COVID-19 Vaccine and safety events of interest, among persons administered the vaccine in both the overall US population and in populations of interest (e.g., pregnant women, the immunocompromised and persons with a prior history of COVID-19 infection). This observational study will capture safety events (based on AESI) including myocarditis and pericarditis, in individuals of any age who received the Pfizer-BioNTech COVID-19 Vaccine since its availability under an EUA using electronic health records and claims data from data partners participating in the Sentinel System. This study, will capture hospitalizations, deaths and serious safety events of interest, including myocarditis and pericarditis, as well as selected pregnancy-related and birth outcomes.
	• C4591011 and C4591012:
	 The collection of safety data in vaccine recipients is critical to our understanding of the vaccine safety profile and to enable safety signal detection and, if needed, further risk mitigation during the EUA. In addition to the collection and monitoring of AEs reported voluntarily by healthcare professionals providing the vaccine and by individuals receiving the vaccine, active surveillance studies of the Pfizer- BioNTech COVID-19 Vaccine under EUA are also planned. Active surveillance of large numbers of individuals vaccinated with the Pfizer-BioNTech COVID-19 Vaccine is necessary to confirm the safety profile demonstrated in the clinical study in a broader population under real-world conditions. Pfizer-BioNTech is conducting active surveillance studies of individuals vaccinated with the Pfizer—BioNTech COVID-19 Vaccine under an EUA in populations prioritized in the early stages of the EUA, e.g., active military and elderly, as described in the study protocols C4591011 (study planned) and C4591012 (study ongoing) submitted to FDA on 29 January 2021. The study period is/will be approximately 30 months following availability of vaccine under EUA. The studies capture hospitalizationss deaths and serious safety events of interest, including myocarditis and pericarditis.
	 C4591021/<u>C4591038 (former, C4591021 substudy=):</u> Post-approval observational studies using real-world data are needed to assess the association between Pfizer-BioNTech COVID-19 Vaccine and safety events of interest, among large numbers of persons administered the vaccine in both the overall population and in populations of interest (e.g., pregnant women, the immunocompromised) and to assess the natural history of post-vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes within 1 year of myocarditis/pericarditis diagnosis among individuals vaccinated with BNT162b2 as well as individuals not vaccinated with a COVID-19 vaccine.
	 C4591036: Pediatric Heart Network Study :: Prospective cohort study on patients <21 years presenting to PHN sites after receiving a first or second dose of a COVID-19 vaccine and who were diagnosed with myocarditis/pericarditis. This study characterizes the clinical course, risk

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Table 31. Action 1	Tan for important fuentineu Risk Astyocarulus and renearulus
	factors, long-term sequelae, and quality of life in children and young adults <21 years with acute post-vaccine myocarditis/pericarditis.
	 C4591031 substudy B: If testing of Troponin I levels in individuals who did not receive BNT162b2 (currently in preparation) indicates that Troponin I could be a reliable indicator of potential subclinical myocarditis, obtaining serum samples for potential Troponin I testing during the period of increased risk of clinical myocarditis may help characterize the absence/presence and frequency of subclinical myocarditis amongst individuals <u>1612</u> to 30 years of age. C4591007 substudy <u>- Troponin group</u>: If testing of Troponin I levels in individuals who did not receive BNT162b2 (currently in preparation) indicates that Troponin I could be a reliable indicator of potential subclinical myocarditis, obtaining serum samples for potential Troponin I testing during the period of increased risk of clinical myocarditis may help characterize the absence/presence and frequency of subclinical myocarditis may help characterize the absence/presence and frequency of subclinical myocarditis may help characterize the absence/presence and frequency of subclinical myocarditis amongst individuals 5 to <u>15<16</u> years of age.
Milestones for	• C4591009:
evaluation and	Protocol submission: 31 August 2021
reporting	Monitoring report submission: 31 October 2022
	Interim Analysis submission: 31 October 2023
	Study completion: 30 June 2025
	• Final study report submission: 31 October 2025.
	• C4591011:
	 Interim study reportsNot applicable^a will be submitted on the following dates based on data collected post EUA in target populations:
	 31 December 2021 30 June 2022 31 December 2022 Final study reports submission: 31 December 2023.
	• C4591012
	Protocol amendment submission: 31 August 2021
	 Interim study reports will be submitted on the following dates based on data collected post-EUA in target populations:
	• 31 December 2021
	 30 June 2022 31 December 2022
	Study completion: 30 June 2023
	• Final study reports submission: 31 December 2023.
	• C4591021
	Protocol submission: 11 August 2021
	Progress report submission: 30 September 2021
	• Interim study reports will be submitted on the following dates:
	 31 March 2022 30 September 2022 31 March 2023 30 September 2023 31 March 2024 Study completion: 31 March 2024
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	Final study report submission: 30 September 2024
	• <u>C4591038 (former, C4591021 substudy)</u>
	 Final protocol submission: 31 January 2022
	Study completion: 31 March 2024
	• Final study report submission: 30 September 2024.
	• C4591036
	 Protocol submission: 30 November 2021
	Study completion: 31 December 2026
	Final study report submission: 31 May 2027
	• C4591031 substudy <u>B</u>
	Final protocol submission: 30 November 2021
	Study completion: 30 June 2022
	Final study report submission: 31 December 2022
	C4591007 substudy <u>– Troponin group</u>
	Final protocol submission: 30 September 2021
	Study completion: 30 November 2023
	Final study report submission: 31 May 2024
	as this is a voluntary sponsor study (as per FDA was informed (Response to F
May 2021 Information	Pagnest Pagarding Active Surveillance Studies) that received to PLA 12574

a. <u>Milestones deleted as this is a voluntary sponsor study (as per FDA was informed (Response to FDA 12 May 2021</u>—Information Request Regarding Active Surveillance Studies) that<u>received to BLA 125742/0</u>, <u>dated 13 August 2021</u>, where the first milestone (Interim Report submission due 30 June 2021) is delayed due to a change in study collaboration;FDA characterized both studies as "voluntary" and therefore it has been removed from this table.no longer commitment).

Table 52. Action Plan for Important Identified Risk "Anaphylaxis"

Actions proposed	Communication of this important identified risk via label (Sections 4 - Contraindications, 5.1 - Management of Acute Allergic Reactions, Section 6 - Adverse reactions - and 6.2 - Post Authorization Experience).
	• C4591001: Phase 1/2/3, placebo-controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.
	C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19-mRNA vaccine in the United States.
	C4591011: Active safety surveillance of the Pfizer-BioNTech COVID-19 vaccine in the US Department of Defense population following Emergency Use Authorization.
	 C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer-BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
	 C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
Objective of proposed actions	Labelling communicates the risk of anaphylaxis.

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Table 52. Action Plan for Important Identified Risk "Anaphylaxis"

Table 52. Action I	ian for important identified Risk "Anaphylaxis"
	 C4591001: To evaluate the safety, tolerability, immunogenicity, and efficacy of BNT162b2. Further, an unfavorable imbalance between the vaccine and control groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, may indicate the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2. C4591009: To assess the occurrence of safety events of interest in the general US population of all ages, pregnant women, the immunocompromised and persons with a prior history of COVID-19 within selected data sources participating in the US Sentinel System. C4591011: To assess whether individuals in the US DoD Military Health System (MHS) experience increased risk of safety events of interest,
	following receipt of the BNT162b2.
	 C4591012: To assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.
	 C4591021: To assess the potential increased risk of adverse events of special interest (AESI), including anaphylaxis after being vaccinated with COVID 19 mRNA-vaccine.
Rationale for proposed	Labeling communicates to health care provider the risk of anaphylaxis.
actions	• C4591001: Long-term monitoring throughout the clinical study for up to 2 years to assess the risk for vaccine-associated enhanced disease.
	 C4591009: Robust surveillance is needed to ensure comprehensive understanding of real-world safety of the BNT162b2 in the in large samples of general US population and in subcohorts of interest, including pregnant women, immunocompromised individuals and persons with a prior history of COVID-19 infection.
	 C4591011 and C4591012: Robust surveillance is needed to ensure comprehensive understanding of real-world safety. This surveillance strategy consists of complementary approaches to ensure timely signal identification and evaluation in populations expected to receive the BNT162b2 under an Emergency Use Authorization (EUA).
	 C4591021: Robust surveillance is needed to ensure comprehensive understanding of real-world safety of the Pfizer-BioNTech COVID-19 Vaccine in the general EU population.
Monitoring by the sponsor for safety issue and proposed actions	 C4591001: Safety evaluations will include AESI, including anaphylaxis; these will be collected systemically and monitored throughout the Phase 3 study.
	 C4591009 and C4591021: Post-approval observational studies using real-world data are needed to assess the association between BNT162b2 and safety events of interest, among persons administered the vaccine in both the overall US (C4591009) and EU (C4591021) population and in populations of interest (e.g., pregnant women, the immunocompromised and persons with a prior history of COVID-19 infection). This observational study will capture safety events (based on AESI) including anaphylaxis, in individuals of any age who received the BNT162b2 since its availability under an EUA using electronic health records and claims data from data partners participating in the Sentinel System. This study, will capture hospitalizations, deaths and serious safety events of interest, including anaphylaxis, as well as selected pregnancy-related and birth outcomes. C4591011 and C4591012:
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Table 52. Action Plan for Important Identified Risk "Anaphylaxis"

	 The collection of safety data in vaccine recipients is critical to our understanding of the vaccine safety profile and to enable safety signal detection and, if needed, further risk mitigation during the EUA. In addition to the collection and monitoring of AEs reported voluntarily by healthcare professionals providing the vaccine and by individuals receiving the vaccine, active surveillance studies of the BNT162b2 under EUA are also planned.
	2. Active surveillance of large numbers of individuals vaccinated with the BNT162b2 is necessary to confirm the safety profile demonstrated in the clinical study in a broader population under real-world conditions. Pfizer-BioNTech plans to conduct active surveillance studies of individuals vaccinated with the BNT162b2 under an EUA in populations prioritized in the early stages of the EUA, e.g., active military and elderly, as described in the study protocols C4591011 and C4591012 submitted to FDA on 29 January 2021. The study period will be approximately 30 months following availability of vaccine under EUA. The studies will capture hospitalizations, deaths and serious safety events of interest, including anaphylaxis.
	• C4591021: The collection of safety data in vaccine recipients, including pregnant women, is critical to our understanding of the vaccine safety profile and to enable robust safety signal detection and evaluation and, if needed, further risk mitigation under BLA.
Milestones for evaluation and reporting	 C4591001 (ongoing Study): CSR submission upon regulatory request: at any time Final CSR submission with supplemental follow-up: 31 August 2023.
	 C4591009: Protocol submission: 31 August 2021 Monitoring report submission: 31 October 2022 Interim Analysis submission: 31 October 2023 Study completion: 30 June 2025 Final study report submission: 31 October 2025. C4591011
	 Interim study reports.<u>Not applicable</u>^a will be submitted on the following dates based on data collected post EUA in target populations:
	 C4591012 <u>Protocol amendment submission: 31 August 2021</u> Interim study reports will be submitted on the following dates based on data collected post-EUA in target populations: 31 December 2021 30 June 2022 31 December 2021 31 December 2022 Study completion: 30 June 2023
	Final study reports submission: 31 December 2023.C4591021



Protocol submission: 11 August 202	21
Progress report submission: 30 Sep	tember 2021
Interim study reports will be submit	tted on the following dates:
o 31 March 2022	
o 30 September 2022	
 31 March 2023 	
o 30 September 2023	
o 31 March 2024	
Study completion: 31 March 2024	
Final study report submission: 30 S	September 2024
a. Milestones deleted as this is a voluntary sponsor study (as per FD.	A was informed (Response to FDA 12
May 2021—Information Request Regarding Active Surveillance Studie	es) that received to BLA 125742/0,
date d 12 Assesset 2021 subara the first milestone (Interim Depart subm	1.1.1.1.1.20 L

dated 13 August 2021, where the first milestone (Interim Report submission due 30 June 2021) is delayed due to a change in study collaboration; FDA characterized both studies as "voluntary" and therefore, it has been removed from in this table. no longer commitment). Action Plan for Important Potential Risks

 Table 53.
 Action Plan for Important Potential Risk "Vaccine-associated enhanced disease (VAED) including Vaccine-associated enhanced respiratory disease (VAERD)"

() The	,
Actions proposed	 C4591001: Phase 1/2/3, placebo-controlled, randomized, observer-blind, dose- finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.
	• C4591008: HERO Together: A post-Emergency Use Authorization observational cohort study to evaluate the safety of the Pfizer-BioNTech COVID-19 Vaccine in US healthcare workers, their families, and their communities.
	 C4591009: A non-interventional post-approval safety study of the Pfizer— BioNTech COVID-19-mRNA vaccine in the United States.
	• C4591011: Active safety surveillance of the Pfizer—BioNTech COVID-19 vaccine in the US Department of Defense population following Emergency Use Authorization.
	C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer—BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
	 C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine.
Objective of proposed actions	 C4591001: to evaluate the safety, tolerability, immunogenicity, and efficacy of BNT162b2. An unfavorable imbalance between the vaccine and control groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, may indicate the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2.
	 C4591008, C4591009, C4591011, C4591012, and C4591021: to characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among individuals vaccinated with the BNT162b2 since EUA.

Table 53. Action Plan for Important Potential Risk "Vaccine-associated enhanced disease (VAED) including Vaccine-associated enhanced respiratory disease (VAERD)"

(VALA	D)
Rationale for	• C4591001: Robust and long-term monitoring throughout the clinical study for up
proposed actions	to 2 years to assess the risk for vaccine-associated enhanced disease.
	 C4591008, C4591009, C4591011, C4591012 and C4591021: Robust surveillance is needed to ensure comprehensive understanding of real-world safety. This surveillance strategy consists of complementary approaches to ensure timely signal identification and evaluation in populations expected to receive the vaccine in the early stages of an EUA as well as with broader vaccination roll-out.
Monitoring by the sponsor for safety issue and proposed actions	• C4591001: Protocol prespecified stopping and alert rules were set for detecting enhanced COVID-19.
	Participants in all stages of the study will be monitored for COVID-19 illness including severe COVID-19 from Visit 1 onward. Cases will undergo blinded review to identify whether any features of each case appear unusual, in particular greater severity. Indicators of severity may include accelerated deterioration, need for hospitalization, need for ventilation, or death. The Data Monitoring Committee, supported by an unblinded medical monitor, will look for adverse imbalances between vaccine and control groups in COVID-19 disease outcomes, in particular for cases of severe COVID-19, that may be a signal for vaccine-associated enhanced disease on an ongoing basis and at interim analyses. Stopping rules were set so that enrollment could be paused in the event of an adverse imbalance.
	Additional safety evaluations will include AESI that could represent symptoms of severe COVID-19 disease; these will be collected systemically and monitored throughout the Phase 3 study.
	C4591008, C4591011, C4591012: The collection of safety data in vaccine recipients is critical to our understanding of the vaccine safety profile and to enable efficient safety signal detection and, if needed, further risk mitigation during the EUA. In addition to the collection and monitoring of AEs reported voluntarily by healthcare professionals providing the vaccine and by individuals receiving the vaccine, active surveillance studies of the BNT162b2 under EUA are also planned. Active surveillance of large numbers of individuals vaccinated with the BNT162b2 is necessary to confirm the safety profile demonstrated in the clinical study in a broader population under real-world conditions. Pfizer-BioNTech plans to conduct active surveillance studies of vaccinated individuals in populations prioritized in the early stages of the EUA, e.g., healthcare workers, active military, and elderly, as described in C4591008 protocol submitted to FDA on 28 January 2021; C4591011 protocol submitted to FDA on 29 January 2021 and C4591012 protocol submitted to FDA on 29 January 2021. The study period will be approximately 30 months following availability of vaccine under EUA. The studies will capture hospitalizations, deaths and serious safety events of interest, including severe COVID-19 (which, if associated with vaccination, may indicate VAED/VAERD).
	 C4591009 and C4591021: Surveillance of large numbers of individuals vaccinated with the BNT162b2 is necessary to confirm the safety profile demonstrated in the clinical study in a broader population under real-world conditions. This study is intended to capture a broader sample of vaccinated individuals of any age in the general US (C4591009) and EU (C4591021) population using large scale data sources.

Table 53. Action Plan for Important Potential Risk "Vaccine-associated enhanced disease (VAED) including Vaccine-associated enhanced respiratory disease (VAERD)"

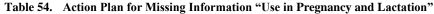
Milestones for	• C4501001 (an anima Stude))
evaluation and	C4591001 (ongoing Study):
reporting	CSR submission upon regulatory request: at any time
	• Final CSR submission with supplemental follow-up: 31 August 2023.
	 Three<u>Two</u> observational post-authorization safety studies for EUA (C4591008, C4591011, and C4591012) and 1 voluntary study (C4591008):
	 C4591008: Interim study reports will be submitted on the following dates based on data collected post_EUA in target populations:
	○ 31 December 2021
	○ 30 June 2022
	• <u>Final study reportsC4591008</u> ª
	• Not applicable
	• <u>C4591012:</u>
	• <u>Protocol amendment</u> submission: 31 <u>December 2023</u> . <u>August 2021</u>
	 C4591012: Interim study reports will be submitted on the following dates based on data collected post-EUA in target populations:
	• 31 December 2021
	o 30 June 2022
	 31 December 2022 Study completion: 30 June 2023
	 Final study reports submission: 31 December 2023.
	 C4591011: Interim study reports* will be submitted on the following dates based
	on data collected post-EUA in target populations:
	0 31 December 2021
	 → 30 June 2022 → 31 December 2022
	 S1 December 2022 Final study reports submission: 31 December 2023.
	 C4591011:
	• Not applicable ^a
	• C4591009:
	 Protocol submission: 31 August 2021
	 Monitoring report submission: 31 August 2021 Monitoring report submission: 31 October 2022
	 Interim Analysis submission: 31 October 2023
	 Study completion: 30 June 2025 Final study report submission: 31 October 2025
	• Final study report submission: 31 October 2025.
	• C4591021
	<u>Protocol submission: 11 August 2021</u>
	Progress report submission: 30 September 2021
	• Interim study reports will be submitted on the following dates:
	 31 March 2022 30 September 2022
	 30 September 2022 31 March 2023
	 30 September 2023
	• 31 March 2024
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Table 53. Action Plan for Important Potential Risk "Vaccine-associated enhanced disease (VAED) including Vaccine-associated enhanced respiratory disease (VAERD)"

Study completion: 31 March 2024Final study report submission: 30 September 2024

a. FDA was informed (Response to FDA 12 May 2021 – Information Request Regarding Active Surveillance Studies) that the first milestone (Interim Report submission due 30 June 2021) is delayed due to a change in study collaboration;, therefore, it has been removed from in this table. a.Milestones deleted as this is a voluntary sponsor study (as per FDA Information Request received to BLA 125742/0, dated 13 August 2021, where the FDA characterized both studies as "voluntary" and therefore no longer commitment).

Action Plan for Missing Information



Actions proposed	 C4591015:C4591015^a: A phase 2/3, placebo-controlled, randomized, observer blind study to evaluate the safety, tolerability, and immunogenicity of a SARS- CoV-2 RNA vaccine candidate (BNT162b2) against COVID-19 in healthy pregnant women 18 years of age and older.
	 C4591009: A non-interventional post-approval safety study of the Pfizer— BioNTech COVID-19-mRNA vaccine in the United States.
	 C4591011: Active safety surveillance of the Pfizer-BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization.
	 C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaceineC4591022vaccine
	 <u>C4591022</u>: Pfizer-BioNTech COVID-19 Vaccine exposure during pregnancy: A non-interventional post-approval safety study of pregnancy and infant outcomes in the Organization of Teratology Information Specialists (OTIS)/MotherToBaby Pregnancy Registry.
Objective of proposed actions	 C4591015:C4591015*: To assess safety and immunogenicity of BNT162b2 in pregnant women. In addition, exploratory objectives include: To describe the immune response in infants born to breastfeeding maternal participants vaccinated with prophylactic BNT162b2 during pregnancy. To describe the safety of maternal immunization in infants born to breastfeeding maternal participants vaccinated with prophylactic BNT162b2 during pregnancy. C4591009^{ab}: To assess whether pregnant women experience increased risk of safety events of interest following receipt of the BNT162b2. C4591011^{ab}: To assess whether sub-cohorts of interest, such as pregnant women, in the MHS experience increased risk of safety events of interest following receipt of the BNT162b2. C4591021^{ab}: To assess whether pregnant women experience increased risk of safety events of interest following receipt of the BNT162b2.
	 C4591022^{ab}: To assess whether pregnant women receiving BNT162b2 experience increased risk of pregnancy and infant safety outcomes, including major

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Table 54.	Action Plan for Mis	sing Information	"Use in Pregnancy	and Lactation"

	congenital malformations, spontaneous abortion, stillbirth, preterm delivery, small for gestational age, and small for age postnatal growth to one year of age.
Rationale for proposed actions	Acquisition of data in an unstudied population with potentially different safety considerations from the time vaccine is available.
	 considerations from the time vaccine is available. C4591015: Monitoring via ongoing clinical study. C4591009: The collection of safety data in vaccine recipients, including pregnant women, is critical to our understanding of the vaccine safety profile and to enable robust safety signal detection and evaluation and, if needed, further risk mitigation under BLA. C4591009:Post-approval observational studies using real-world data are needed to assess the association between Prizer-BioNTech COVID-19 Vaccine and safety events of interest, among persons administered the vaccine in both the overall US population and in populations of interest (e.g., pregnant women, the immunocompromised and persons with a prior history of COVID-19 infection). This observational study will capture safety events (based on AESI) including myocarditis and pericarditis, in individuals of any age who received the Pfizer-BioNTech COVID-19 Vaccine since its availability under an EUA using electronic health records and claims data from data partners participating in the Sentinel System. This study will capture hospitalizations, deaths and serious safety events of interest, including myocarditis and pericarditis, as well as selected pregnancy-related and birth outcomes. C4591011: The collection of safety data in vaccine recipients is critical to our understanding of the vaccine safety profile and to enable efficient safety signal detection and, if needed, further risk mitigation. Active surveillance studies of the BNT162b2 under EUA are also planned. Active surveillance of large numbers of individuals vaccinated with the BNT162b2 under teus verveillance studies of individuals vaccinated with the BNT162b2 under an EUA in populations. Prizer-BioNTech plans to conduct active surveillance studies of individuals vaccine under EUA. The study will capture hospitalizations, deaths and serious safety events of interest, including anaphylaxis. C4591021: The collection of s
	be conducted to evaluate if the pregnant women receiving the vaccine during pregnancy experience increased risk of pregnancy and infant outcomes compared with 1) pregnant women who are unvaccinated and 2) pregnant women who have received an influenza or tetanus, diphtheria, and acellular pertussis (Tdap) vaccine
	during pregnancy C4591021: The collection of safety data in vaccine recipients, including pregnant women, is critical to our understanding of the vaccine safety



	profile and to enable robust safety signal detection and evaluation and, if needed,
	further risk mitigation under BLA.
Milestones for evaluation and	•C4591015:= <u>a.</u>
reporting	Study completion: 31 OctoberAugust 2022
	 Final study report submission: 31 <u>MayMarch</u> 2023.
	• C4591009:
	Protocol submission: 31 August 2021
	Monitoring report submission: 31 October 2022
	Interim Analysis submission: 31 October 2023
	Study completion: 30 June 2025
	• Final study report submission: 31 October 2025.
	• C4591011:
	 Interim study reports^b will be submitted on the following dates based on data collected post EUA in target populations:
	\sim 31 December 2021
	0 30 June 2022
	<u>↔ 31 December 2022</u>
	 Final study report submission: 31 December 2023.
	• Not applicable ^c
	• C4591021
	Protocol submission: 11 August 2021
	 Progress report submission: 30 September 2021
	 Interim study reports will be submitted on the following dates:
	o 31 March 2022
	 30 September 2022 31 March 2023
	 30 September 2023
	o 31 March 2024
	Study completion: 31 March 2024
	 Final study report submission: 30 September 2024
	• C4591022:
	Interim reports submission:
	o 31 January 2022
	o 31 January 2023
	o 31 January 2024
	31 January 2025Study completion: 30 June 2025
an Enrolmont of	 Final study report submission: 31 December 2025 participants into study C4591015 was stopped due to recruitment challenges as a result of
	participants into study C4591015 was stopped due to recruitment challenges as a result of dations for COVID 19 vaccination in pregnant women and the increased availability of
	es. Enrolment of new participants was stopped on 25 October 2021. Participants already
	inue follow up evaluations until study end as planned. For this reason, study completion
	from 31 October 2022 to 31 August 2022 and final study report submission date was
changed from 31 M	May 2023 to 31 March 2023".

b. Study assesses pregnancy only.

b. FDA was informed (Response to FDA 12 May 2021 - Information Request Regarding Active

Surveillance Studies) that the first milestone (Interim Report submission due 30 June 2021) is delayed due to

Table 54. Action Plan for Missing Information "Use in Pregnancy and Lactation"

a change in study collaboration; therefore, it has been removed from in this table.c.Milestones deleted as this is a voluntary sponsor study (as per FDA Information Request received to BLA 125742/0, dated 13 August 2021, where the FDA characterized both studies as "voluntary" and therefore no longer commitment).

Table 55. Action Plan for Missing Information "Vaccine Effectiveness"

Action proposed	C4591014: Pfizer-BioNTech COVID-19 BNT162b2 Vaccine Effectiveness Study Kaiser Permanente Southern California.
	 WI235284: Determining RSV Burden and Outcomes in Pregnant Women and Older Adults Requiring Hospitalization. COVID-19 Amendment for COVID VE/ Sub-study 6.
	• WI255886: Avon Community Acquired Pneumonia Surveillance Study: A Pan- pandemic Acute Lower Respiratory Tract Disease Surveillance Study.
	 BNT162-01 cohort 13: Immunogenicity of Pfizer-BioNTech COVID-19 Vaccine in immunocompromised subjects, including assessment of antibody responses and cell-mediated responses.
	 C4591007 substudy <u>– Lower dose evaluation group</u>: To study lower dose levels of BNT162b2 in individuals 12 through <<u>3018</u> years of age.
Objective of proposed actions	 C4591014: To estimate the effectiveness of 2 doses of BNT162b2 against hospitalization and emergency department admission for acute respiratory illness due to SARS-CoV-2 infection.
	 WI235284: To estimate the effectiveness of 2 doses of BNT162b2 against hospitalization for acute respiratory illness due to SARS-CoV-2 infection.
	 WI255886: To estimate the effectiveness of 2 doses of BNT162b2 against hospitalization for acute respiratory illness due to SARS-CoV-2 infection.
	 BNT-162-01 cohort 13: To assess potentially protective immune responses in immunocompromised adults.
	 C4591007 substudy <u>– Lower dose evaluation group</u>: To evaluate the immunogenicity and safety of lower dose levels of BNT162b2 in individuals 12 through <<u>3018</u> years of age.
Rationale for proposed actions	 C4591014: To determine the effectiveness of BNT162b2 when administered outside of the clinical setting.
	 WI235284: To determine the effectiveness of BNT162b2 when administered outside of the clinical setting.
	 WI255886: To determine the effectiveness of BNT162b2 when administered outside of the clinical setting.
	 BNT-162-01 cohort 13: To determine whether the BNT162b2 has potential to protect immunocompromised adults.
	 C4591007 substudy <u>– Lower dose evaluation group</u>: To assess whether lower dose levels of BNT162b2 could provide an adequate immune response with improved tolerability.

BNT162b2	
1.16 Risk Management Plan (Non-REMS) for Biologic License Application # 125742	

Table 55.	Action Plan	for Missing	Information	"Vaccine	Effectiveness"
Table 55.	Action 1 Ian	ior missing	mor mation	vacune	Encenveness

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Monitoring by the sponsor for safety	C4591014: Use of primary and secondary data sources to monitor COVID-19 infection in vaccinated individuals.
issue and proposed actions	• WI235284: Use of primary and secondary data sources to monitor COVID-19 infection in vaccinated individuals.
	• WI255886: Use of primary and secondary data sources to monitor COVID-19 infection in vaccinated individuals.
	• BNT-162-01 cohort 13: Reactogenicity, AE and SAE assessment.
	C4591007 substudy <u>– Lower dose evaluation group</u> : Reactogenicity, AE and SAE and immunogenicity assessment.
Milestones for	• C4591014:
evaluation and	Study completion: 31 December 2022
reporting	• Final CSR submission: 30 June 2023.
	• WI235284: Final CSR submission: 30 June 2023.
	• WI255886: Final CSR submission: 30 June 2023.
	• BNT-162-01 cohort 13: First IA submission: 30 September 2021.
	• C4591007 substudy (to study lower_Lower dose levels) evaluation group:
	Final protocol submission: 30 September 2021
	Study completion: 30 November 2023
	Final report submission: 31 May 2024

Actions proposed	 C4591001 ≥12 to ≤15 years of age: Phase 1/2/3, placebo-controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals^a. Randomised placebo-controlled study in 2000 participants (1000 active recipients) of 2 doses of BNT162b2 at a 21-day interval.
	C4591007 <12 years of age: - Lower dose evaluation group:
	 Phase 1 (low dose evaluation): open label dose finding study tothat will evaluate the safety, tolerability, and immunogenicity of 10 µg BNT162b2 from 2 schedule in 2 age groups (participants 12 to <16 and phase16 to <18 years of age).
	 <u>Phase 2/3 placebo controlled, observer blinded lower dose evaluation: Is the portion of the study that will evaluate the</u> safety, tolerability, and immunogenicity study of a SARS CoV 2 RNA vaccine candidate against COVID-19 in healthy children <in (12="" 16="" <16="" age="" and="" each="" group="" li="" to="" to<="" years=""> </in>
	<18 years of age-
).at the selected dose schedule from the Phase 1: lower-dose evaluation. In
	this open-label dose finding portion up to 3 age groups (study, all participants ≥ 5 to \leq will have blood drawn at baseline prior to Dose 1 and at 1, 6, and 12
	$\frac{1}{2}$ years, ≥ 2 to <5 years, and ≥ 6 -months to $<\underline{a}$ fter Dose 2-years of age) with 16
	participants per dose level. Dose finding is being initiated in this study in.
	Immunobinding to comparator participants ≥ 5 to < 12 years of age based on
	the acceptable blinded safety assessment of the 30-µg dose in 12-to 15-year-
	olds in the C4591001 study. The purpose of Phase 1 is to identify preferred
	dose level(s) of BNT162b2 from up to 3 different dose levels in each age
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	 group. Phase 2/3: Children ≥5 to <12 years of age are randomized 2:1 at selected dose level of BNT162b2 at a 21 day interval (4500 total subjects; 3000 active vaccine). Children 2 to <5 years and 6 to 23 months of age randomized 2:1 placebo controlled at selected dose level of BNT162b2 at a 21 day interval (1125 total subjects per age group; 750 active vaccine per will be based on immunogenicity data collected at baseline and 1 month after Dose 2. The persistence of the immune response will be based on immunogenicity data collected at baseline and 1 month after Dose 2. The persistence of the immune response will be based on immunogenicity data collected in participants at baseline and 1, 6, and 12 months after Dose 2. (300 participants per age group). C4591007 substudy: Subelinical myocarditis evaluation. – Troponin group: A Phase 3 substudy of 750 participants 5 to <12 years of age (randomized 2:1 to receive BNT162b2 10 µg or placebo) and 500 participants 12–15–<16 years of age (open label receipt of BNT162b2 30 µg).
	 C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA vaccine in the United States.
	 C4591021/<u>C4591038 (former.</u> C4591021 substudy: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer_BioNTech Coronavirus Disease 2019 (COVID-19) vaccine. Substudy to investigate natural history of post-vaccination myocarditis and pericarditis.
	 C4591023: Phase 1 open label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer- blinded safety, tolerability, and immunogenicity study of a SARS-CoV-2 RNAmRNA vaccine candidate against COVID-19 in healthy infants <6 months of age. Phase 1: open-label dose finding portion up to 3 dose levels with 16 participants per dose level. Phase 2/3: Children ≥5 to <12 years of age are randomized 2:1 at selected dose level of BNT162b2 at a 21-day interval. Design and subject numbers to be confirmed.
	 C4591036: Pediatric Heart Network Study: Working title: <u>Myocarditis/pericarditis follow up study within the Pediatric Heart Network Low</u> <u>Interventional Cohort Study of Myocarditis/Pericarditis Associated with</u> <u>Comirnaty in persons less than 21 years of age.</u>
Objective of proposed actions	 C4591001 ≥12 to ≤15 years of age: Safety compared to placebo and immune-non- inferiority of neutralizing antibody immune response compared to subjects 16-25 years of age.
	 C4591007 <<u>12 years of age_Lower dose evaluation group</u>: Immunobridging analysis of immune responses in participants within each age group (participants ≥5 to <12 years, ≥2 to <5 years, and ≥6 months to <2 years of age) to those in participants 16 to 25 years of age in the Phase 3 C4591001 efficacy study. Efficacy if sufficient cases accrue.
	 C4591007 substudy (<u>Subclinical myocarditis evaluation) – Troponin group</u>: To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of subclinical myocarditis amongst individuals 5 to <u>15<16</u> years of age.
	 C4591009-and C4591021: To assess the use of vaccine in persons <12 for whom vaccine is not yet approved and and occurrence of safety events of interest, including myocarditis and pericarditis, in the general US and EU populations

TED Tears of Age				
		within selected broad population based data sources participating in the US sentinel system.		
	•	C4591021: To assess the potential increased risk of Adverse Events of Special		
		Interest (AESI), including myocarditis/pericarditis after being vaccinated with COVID-19 vaccine.		
	•	C4591038 (former, C4591021 substudy: To describe the natural history of post- vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes within 1 year of myocarditis/pericarditis diagnosis among individuals, including any individuals <12 years, vaccinated with BNT162b2 as well as individuals not vaccinated with a COVID-19 vaccine.		
	•	C4591023: To evaluate safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy infants <6 months of age.		
	•	C4591036: Pediatric Heart Network Study: To characterize the clinical course, risk factors, resolution, long-term sequelae, and quality of life in children and young adults <21 years, including any individuals <12 years, with acute post-vaccine myocarditis/pericarditis.		
Rationale for proposed actions	•	C4591001 \geq 12 to \leq 15 years of age: Need to collect evidence of safety and effectiveness to support immunization in this age group.		
	•	C4591007 <12 years of age_Lower dose evaluation group: Need to collect evidence of safety and effectiveness to support immunization in this age group.		
	•	C4591007 substudy <u>– Troponin group</u> : The first ~4 days post-vaccination is the time period when symptomatic myocarditis cases have most frequently been reported. Elevated Troponin I may be an indicator of subclinical myocarditis.		
	•	C4591009 and C4591021: Long-term surveillance of large numbers of individuals vaccinated with the BNT162b2 is necessary to confirm the safety profile demonstrated in the clinical study in a broader population under real-world conditions. Use in persons <12 will be captured as age is not a criterion for study eligibility		
	•	C4591038 (former, C4591021 substudy:): Study is needed to describe natural history of myocarditis and pericarditis in persons after vaccination with Pfizer-BioNTech COVID-19 vaccine and in unvaccinated persons. Use in persons <12 will be captured as age is not a criterion for study eligibility		
	•	C4591023: Need to collect evidence of safety and effectiveness to support immunization in this age group.		
	•	C4591036: Pediatric Heart Network Study: Study is necessary to characterize the clinical course, risk factors, resolution, long-term sequelae, and quality of life in children and young adults <21 years with acute post-vaccine myocarditis/pericarditis. Use in children <12 will be captured as age is not a criterion for study eligibility		
Monitoring by the	•	C4591001 \geq 12 to \leq 15 years of age:		
sponsor for safety		• Electronic diary for reactogenicity 7 days following each dose of vaccine.		
issue and proposed actions		Adverse events for one month after second dose.		
actions		• Serious Adverse Events for 6 months after the second dose.		
		Related SAEs and related deaths for 24 months after the second dose.		

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 Collection of COVID-19 and MIS-C cases up to 24 months after the second dose.
 C4591007 <12 years of age_Lower dose group/C4591007 substudy – Troponin group:
• Electronic diary for reactogenicity 7 days following each dose of vaccine.
Adverse events for one month after second dose.
• Serious Adverse Events for 6 months after the second dose.
 Related SAEs and related deaths for 24 months after the second dose for < 12 years.
 Collection of COVID-19 and MIS-C cases up to 24 months after the second dose < 12 years.
• C4591009:
 Longitudinal medical care information on outpatient medication dispensing, vaccine administrations, and inpatient and outpatient diagnoses and procedures in addition to adjudication of select events via medical records.
 Incidence rates and comparative incidence rate ratios of safety events of interest (AESIs from FDA's BEST System¹¹³ and CDC's Vaccine Safety Datalink¹¹⁴ in addition to vaccine-associated enhanced respirator disease).
 Study period to start on date that BNT162b2 became available under EUA (December 11, 2020) and will end a minimum of 3 years after this date.
• Risk windows will be defined for safety events of interest that have a hypothesized increased risk during specific time periods following vaccination. For other safety events of interest, patients will be followed for a maximum of 1 year.
• C4591021:
Longitudinal medical care and vaccination information.
 Incidence rates and comparative incidence rate ratios of safety events of interest.
 Study period to start on date that BNT162b2 became available in EU and will end after 2-3 years depending on outcome.
 Risk windows will be defined for safety events of interest that have a hypothesized increased risk during specific time periods following vaccination.
 <u>C4591038 (former, C4591021 substudy=):</u> Describe natural history of post-vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes for 1 year Study population includes individuals vaccinated with BNT162b2 as well as individuals not vaccinated with a COVID-19 vaccine C4591023 < 6 months of age
• Electronic diary for reactogenicity 7 days following each dose of vaccine.
Adverse events for one month after second dose.
• Serious Adverse Events for 6 months after the second dose.
• Related SAEs and related deaths for 12 months after the second dose for < 12 years.
 Collection of COVID-19 and MIS-C cases up to 12 months after the second dose < 12 years.

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Tais of Age			
	 C4591036: Pediatric Heart Network Study: Prospective cohort study on patients <21 years presenting to PHN sites after receiving a first or second dose of a COVID-19 vaccine and who were diagnosed with myocarditis. This study characterizes the clinical course, risk factors, long-term sequelae, resolution and quality of life in children and young adults <21 years with acute post-vaccine myocarditis/pericarditis. 		
Milestones for	• C4591001 ≥12 to ≤15 years of age:		
evaluation and	Reports:		
reporting	• 6-month post dose 2 (safety): 31 OctoberDecember 2021 ^b		
	 24-month post dose 2 (safety): 30 April 2023^c. 		
	Study completion: 31 May 2023		
	Final report submission: 31 October 2023		
	• C4591007 < 12 years of age: _ Lower dose group:		
	 First report with up to 1-month post dose 2 in ≥5 to <12 years of age (safety): 30 September 2021 		
	• Further reports:		
	 6-month post dose 2 (safety): 31 March 2022 		
	 24-month post dose 2 (safety): 30 September 2023. 		
	Study Completion: 30 November 2023		
	Final Report Submission: 31 May 2024		
	C4591007 substudy (Subclinical myocarditis evaluation):- Troponin group:		
	Final protocol submission: 30 September 2021		
	Study completion: 30 November 2023		
	Final report submission: 31 May 2024		
	• C4591009:		
	Protocol submission: 31 August 2021		
	Monitoring report submission: 31 October 2022		
	Interim Analysis submission: 31 October 2023		
	Study completion: 30 June 2025		
	• Final study report submission: 31 October 2025.		
	• C4591021:		
	Protocol submission: 11 August 2021		
	Progress report submission: 30 September 2021		
	Interim study reports will be submitted on the following dates:		
	• 31 March 2022		
	• 30 September 2022		
	o 31 March 2023		
	o 30 September 2023		
	• 31 March 2024		
	Study completion: 31 March 2024 Final study ranget submission: 30 Sentember 2024		
	Final study report submission: 30 September 2024		
	<u>C4591038 (former,</u> C4591021 substudy Final protocol submission: 31 January 2022		
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	Study completion: 31 March 2024
	• Final study report submission: 30 September 2024.
•	C4591023
	Final protocol submission: 31 January 2022
	Study completion: 31 July 2024
	• Final study report submission: 31 October 2024.
•	C4591036:
	 Protocol submission: 30 November 2021
	Study completion: 31 December 2026
	• Final study report submission: 31 May 2027

a. Study originally included in the PVP to address the Missing Information "Use in pediatric individuals \leq 16 years of age".

b. Due date updated from 31 July 2021 because the last subject visit for this group will not be until September 2021.October 2021 to 31 December 2021 to implement additional CBER requests (received on 24 September 2021) for the 12 to ≤15 years sBLA (FDA already informed of this change on 22 October 2021).

c. Due date updated from 31 January 2023 for the same reason above.

3.1.4. Summary of Actions to be Completed, Including Milestones

Table 57.	Summary of Safety Concerns and Action Plans
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Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Myocarditis and Pericarditis	C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-vaccine in the United States. <i>Planned</i>	To assess the occurrence of safety events of interest, including myocarditis and pericarditis, in the general US population of all ages, pregnant women, the immunocompromised and persons with a prior history of COVID-19 within selected data sources participating in the US Sentinel System	 Protocol submission: Monitoring report submission: Interim analysis submission: Study completion: Final study report submission: 	 31 August 2021 31 October 2022 31 October 2023 30 June 2025 31 October 2025
	C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization <i>Planned</i>	To assess whether individuals in the US DoD Military Health System (MHS) experience increased risk of safety events of interest, including myocarditis and pericarditis, following receipt of the Pfizer-BioNTech COVID-19 Vaccine.	 Interim reports submission:⁻ Final study report submission:<u>Not applicable*</u> 	 31 December 2021 30 June 2022 31 December 2022 31 December 2023<u>Not applicable*</u>
	C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer-BioNTech <u>Coronavirus</u> <u>Disease 2019 (COVID-19)</u> Vaccine. Ongoing	To assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, including myocarditis and pericarditis, following receipt of the Pfizer-BioNTech COVID-19 Vaccine	 Protocol amendment submission: Interim reports submission: Study completion: Final study report submission: 	 31 August 2021 31 December 2021 30 June 2022 31 December 2022 30 June 2023 30 June 2023 31 December 2023
	C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer	To assess the potential increased risk of adverse events of special interest (AESI), including myocarditis/pericarditis after	 <u>Protocol submission</u> Progress report submission: Interim analysis submission: 	 11 August 2021 30 September 2021 31 March 2022 30 September 2022

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Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
	BioNTech Coronavirus Disease 2019 (COVID-19) vaccine Ongoing	being vaccinated with COVID-19 mRNA-vaccine.	Study completion:Final study report submission:	31 March 2023 30 September 2023 31 March 2024 31 March 2024 30 September 2024
	C4591038 (former, C4591021 substudy=): Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine. Substudy to investigate natural history of post-vaccination myocarditis and pericarditis. <i>Planned</i>	To describe the natural history of post-vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes within 1 year of myocarditis/pericarditis diagnosis among individuals vaccinated with BNT162b2 as well as individuals not vaccinated with a COVID-19 vaccine.	 Final protocol submission: Study completion: Final study report submission 	 30 September 2024 31 January 2022 31 March 2024 30 September 2024.
	C4591036: Pediatric Heart Network Study: Working title: <u>Myocarditis</u> Low interventional cohort study of <u>myocarditis</u> /pericarditis follow- up study within the Pediatric <u>Heart Network</u> , associated with Comirnaty in persons less than 21 years of age	To characterize the clinical course, risk factors, resolution, long-term sequelae, and quality of life in children and young adults <21 years with acute post- vaccine myocarditis/pericarditis.	 Protocol submission: Study completion: Final study report submission: 	 30 November 2021 31 December 2026 31 May 2027
	Planned C4591031 substudy:- <u>B:</u> A Phase 3randomized, placebo- controlled, observer-blind, cross- over substudy to evaluate the safety and tolerability of 1000	To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of	Final protocol submission:Study completion:Final study report submission:	 30 November 2021 30 June 2022 31 December 2022

Table 57.	Summary of Safety Concerns and Action Plans
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Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Safety Concerns	Ongoing/Planned Action participants with documented receipt of 2 prior 30 µg doses_a booster (third) dose_of BNT162b2-(the second dose received_Participants ≥12 years of age to <30 years of age who have completed a 2-dose primary series of BNT162b2 (30 µg doses) at least 6 months ago), 16 to 30 years of age (randomized 1:1 in a crossover design to receive 30 µg BNT162b2 or placebo at baseline and the alternative 4 weeks later).(≥12 months for those 12-17 years of age) prior to randomization will be enrolled. PlannedOrgoing C4591007 substudyTroponin group: A Phase 3 substudy of 750 participants 5 to <12 years of age (randomized 2:1 to receive BNT162b2 10 µg or placebo) and 500 participants 12-15 16 years of age (open label receipt of BNT162b2 30 µg).	Summary of Objectives subclinical myocarditis amongst individuals <u>1612</u> to 30 years of age. To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of subclinical myocarditis amongst individuals 5 to <u>15≤16</u> years of age.	 Milestones Final protocol submission: Study completion: Final study report submission: 	• 30 September 2021 • 30 November 2023 • 31 May 2024
Anaphylaxis	Ongoing C4591001: Phase 1/2/3, placebo- controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity,	To evaluate the safety, tolerability, immunogenicity, and efficacy of BNT162b2. An unfavorable imbalance between the vaccine and control	 CSR submission upon regulatory request: Final CSR submission with supplemental follow-up: 	At any time31 August 2023

Table 57.	Summary of Safety Concerns a	and Action Plans
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Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals. <i>Ongoing</i>	groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, may indicate the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2.		
C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-Vaccine in the United States. <i>Planned</i>	To assess the occurrence of safety events of interest in the general US population of all ages, pregnant women, the immunocompromised and persons with a prior history of COVID-19 within selected data sources participating in the US Sentinel System.	 Protocol submission: Monitoring report submission: Interim analysis submission: Study completion: Final study report submission: 	 31 August 2021 31 October 2022 31 October 2023 30 June 2025 31 October 2025
C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization.	To assess whether individuals in the US DoD Military Health System (MHS) experience increased risk of safety events of interest, following receipt of the BNT162b2.	 Interim reports submission^e: Final study report submission: Not applicable* 	 31 December 2021 30 June 2022 31 December 2022 31 December 2023 31 December 2023 Not applicable*
Planned C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer-BioNTech <u>Coronavirus</u> <u>Disease 2019 (COVID-19)</u> Vaccine.	To assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.	 Protocol amendment submission: Interim reports submission: Study completion: Final study report submission: 	 31 August 2021 31 December 2021 30 June 2022 31 December 2022 30 June 2023 30 June 2023 31 December 2023
	and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals. <i>Ongoing</i> C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-Vaccine in the United States. <i>Planned</i> C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization. <i>Planned</i> C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer-BioNTech <u>Coronavirus</u> <u>Disease 2019 (COVID-19)</u>	and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, may indicate the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2.C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-Vaccine in the United States.To assess the occurrence of safety events of interest in the general US population of all ages, pregnant women, the immunocompromised and persons with a prior history of COVID-19 within selected data sources participating in the US Sentinel System.C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization.To assess whether individuals in the US DoD Military Health System (MHS) experience increased risk of safety events of interest, following receipt of the BNT162b2.PlannedTo assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.	and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, in particular for severe COVID-19 disease, in particular for severe COVID-19 disease, may indicate the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2.Protocol submission: • Monitoring report submission: • Interim analysis submission: • Interim analysis submission: • Study completion: • Final study report submission: • Final study report submission: • Final study report submission: • Final study report submission: • Not applicable*C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization.To assess whether individuals in the US DoD Military Health System (MHS) experience interest, following receipt of the BNT162b2.• Interim reports submission: • Not applicable*PlannedTo assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.• Interim reports submission: • Not applicable*PlannedTo assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.• Protocol amendment submission: • Not applicable*Piraer-BioNTech COVID-19)To assess whether individuals in the US Veteran's Affairs Health System experience increased risk of safety events of interest, following receipt of the BNT162b2.• Study completion:Piraese 2019 (COVID-19)To assess w

Table 57. Summary of Safety Concerns and Action Plans

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
	C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine Ongoing	To assess the potential increased risk of adverse events of special interest (AESI), including anaphylaxis after being vaccinated with COVID 19 mRNA-vaccine.	 Protocol submission: Progress report submission: Interim analysis submission: 	 11 August 2021 30 September 2021 31 March 2022 30 September 2022 31 March 2023 30 September 2023 30 September 2023 31 March 2024
Vaccine- associated enhanced disease (VAED) including	C4591001: Phase 1/2/3, placebo- controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity,	To evaluate the safety, tolerability, immunogenicity, and efficacy of BNT162b2. An unfavorable imbalance	 Study completion: Final study report submission: CSR submission upon regulatory request: Final CSR submission with supplemental follow-up: 	 31 March 2024 30 September 2024 Any time 31 August 2023
vaccine- associated enhanced respiratory disease (VAERD)	and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals. Ongoing	between the vaccine and control groups in the frequency of COVID-19 disease, in particular for severe COVID-19 disease, may suggest the occurrence of VAED/VAERD. Surveillance is planned for 2 years following Dose 2		

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Table 57.	Summary of Safety	Concerns and Action Plans
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Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
	C4591008 HERO Together: A post- Emergency Use Authorization observational cohort study to evaluate the safety of the Pfizer- BioNTech COVID-19 Vaccine in US healthcare workers, their families, and their communities. C4591008: <i>Ongoing</i>	To characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among individuals vaccinated with the BNT162b2 since EUA	Interim reports submission: Final study report submission:Not applicable*	 31 December 2021 30 June 2022 31 December 2022 31 December 2023Not applicable*
	C4591012: Post-emergency use authorization active safety surveillance study among individuals in the Veteran's Affairs Health System receiving Pfizer- BioNTech Coronavirus Disease 2019 (COVID-19) vaccine. Ongoing	To characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among individuals vaccinated with the BNT162b2 since EUA	 Protocol amendment <u>submission:</u> Interim reports submission: Study completion: Final study report submission: 	 31 August 2021 31 December 2021 30 June 2022 31 December 2022 30 June 2023 30 June 2023 31 December 2023
	C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the US Department of Defense population following Emergency Use Authorization	To characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among individuals vaccinated with the BNT162b2 since EUA	 Interim reports submission* Final study report submissionNot applicable* 	 31 December 2021 30 June 2022 31 December 2022 31 December 2023Not applicable*
	C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-Vaccine in the United States	To characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among	 Protocol submission: Monitoring report submission: Interim analysis submission: Study completion Final study report submission: 	 31 August 2021 31 October 2022 31 October 2023 30 June 2025 31 October 2025

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Safety Concerns	Ongoing/Planned Action Planned C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine Ongoing C4591015: A phase 2/3, placebo-controlled, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate (BNT162b2) against COVID-19 in healthy pregnant women 18	Summary of Objectives individuals vaccinated with the BNT162b2_since_EUA To characterize the real-world incidence of safety events of interest, including events indicative of severe or atypical COVID-19 disease, among individuals vaccinated with the BNT162b2 since EUA. To assess safety and immunogenicity of BNT162b2 in pregnant women. Exploratory objectives include: To describe the immune response in infants born to breastfeeding maternal participants vaccinated with prophylactic BNT162b2	Milestones • Protocol submission • Progress report submission: • Interim analysis submission: • Study completion • Final study report submission: • Study completion: • Study completion: • Final study report submission:	Due dates • 11 August 2021 • 30 September 2021 • 31 March 2022 30 September 2022 31 March 2023 30 September 2023 31 March 2023 31 March 2024 • 31 March 2024 • 31 March 2024 • 31 October August 2022 • 31 MayMarch 2023
	years of age and older. Ongoing C4591011: Active safety surveillance of the Pfizer- BioNTech COVID-19 Vaccine in the United States Department of Defense population following	during pregnancy. To describe the safety of maternal immunization in infants born to breastfeeding maternal participants vaccinated with prophylactic BNT162b2 during pregnancy. To assess whether sub-cohorts of interest, such as pregnant women, in the MHS experience increased risk of safety events of interest following receipt of the	Interim reports submission: Final study report submission: Not and lock lock	• <u>31 December 2021</u> <u>30 June 2022</u> <u>31 December 2022</u> • <u>31 December</u> 2022Net en jischle*
	Emergency Use Authorization	BNT162b2.	submission: <u>Not applicable*</u>	2023Not applicable*

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Table 57. Summary of Safety Concerns and Action Plans

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Safety Concerns	Planned	Summary or Objectives	winestones	Duc uaits
	C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-Vaccine in the United States. <i>Planned</i>	To assess whether pregnant women, experience increased risk of safety events of interest following receipt of the BNT162b2.	 Protocol submission: Monitoring report submission: Interim analysis submission: Study completion: Final study report submission: 	 31 August 2021 31 October 2022 31 October 2023 30 June 2025 31 October 2025
	C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine	To assess whether pregnant women experience increased risk of safety events of interest following receipt of the BNT162b2.	 Protocol submission: Progress report submission: Interim analysis submission: 	 11 August 2021 30 September 2021 31 March 2022 30 September 2022 31 March 2023 30 September 2023 31 March 2024
	Ongoing		Study completion:Final study report submission:	31 March 202430 September 2024
	C4591022: Pfizer-BioNTech COVID-19 Vaccine exposure during pregnancy: A non- interventional post-approval safety study of pregnancy and infant outcomes in the Organization of Teratology Information Specialists (OTIS)/MotherToBaby Pregnancy Registry <u>Ongoing</u>	To assess whether pregnant women receiving BNT162b2 experience increased risk of pregnancy and infant safety outcomes, including major congenital malformations, spontaneous abortion, stillbirth, preterm delivery, small for gestational age, and small for age postnatal growth to one year of age.	 Interim reports submission: Study completion: Final study report submission: 	 31 January 2022 31 January 2023 31 January 2024 31 January 2025 30 June 2025 31 December 2025

Table 57.	Summary of Safety Concerns and Action Plans
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Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Vaccine effectiveness	C4591014: Pfizer-BioNTech COVID-19 BNT162b2 Vaccine Effectiveness Study - Kaiser Permanente Southern California. <u>Ongoing</u> Planned	To estimate the effectiveness of 2 doses of BNT162b2 against hospitalization and emergency department admission for acute respiratory illness due to SARS- CoV-2 infection.	Study completion: Final CSR submission:	 31 December 2022 30 June 2023
Vaccine effectiveness (Cont'd)	WI235284: Determining RSV Burden and Outcomes in Pregnant Women and Older Adults Requiring Hospitalization. Amendment for COVID VE/ Sub-study 6. <u>Ongoing</u> Planned	To estimate the effectiveness of 2 dosed of BNT162b2 against hospitalization for acute respiratory illness due to SARS- CoV-2 infection.	Final CSR submission:	• 30 June 2023

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
	WI255886: Avon Community Acquired Pneumonia Surveillance Study: A Pan- pandemic Acute Lower Respiratory Tract Disease Surveillance <u>Study</u> . <i>PlannedOngoing</i>	To estimate the effectiveness of 2 doses of BNT162b2 against hospitalization for acute respiratory illness due to SARS- CoV-2 infection.	Final CSR submission:	• 30 June 2023
	BNT162-01 cohort 13: Immunogenicity of Pfizer- BioNTech COVID-19 Vaccine in immunocompromised subjects, including assessment of antibody responses and cell- mediated responses. <i>Ongoing</i>	To assess potentially protective immune responses in immunocompromised adults.	First IA submission:	• 30 September 2021
	C4591007 substudy <u>– Lower dose</u> evaluation group: To study lower dose levels of BNT162b2 in individuals 12 through < <u>3018</u> years of age. <i>Planned</i>	To evaluate the immunogenicity and safety of lower dose levels of BNT162b2 in individuals 12 through < <u>3018</u> years of age	Final protocol submission:Study completion:Final report submission:	 30 September 2021 30 November 2023 31 May 2024
Use in pediatric individuals < 125 years of age	C4591001 ≥12 to ≤15 years of age: Phase 1/2/3, placebo-controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine	Safety compared to placebo and immune-non-inferiority of neutralizing antibody immune response compared to subjects 16-25 years of age.	 Report 6-month post dose 2 (safety): Report 24-month post dose 2 (safety): Study completion: Final report submission: 	 31 October 2021^e 31 December 2021^c 30 April 2023^{dd} 31 May 2023 31 October 2023

 Table 57.
 Summary of Safety Concerns and Action Plans

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Table 57. Summary of Safety Concerns and Action Plans

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates	
	candidates against COVID-19 in healthy individuals ^b . Ongoing				Field Code Changed
	C4591007	Immunobridging analysis of immune responses in participants within each age group (participants ≥5 to <12 years, ≥2 to <5 years, and ≥6 months to <2 years of age) to those in participants 16 to 25 years of age	 First report with up to 1-month post dose 2 (safety) in ≥5 to <12 years of age: Report 6-month post dose 2 (safety) in ≥5 to <12 years of age: Report 24-month post dose 2 	 30 September 2021 31 March 2022 30 September 2023 	
	safety, tolerability, and immunogenicity, study of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy children <12 years of age. <u>PlannedOugoing (started in</u> <u>March)</u>	in the Phase 3 C4591001 efficacy study. Efficacy if sufficient cases accrue.	 (safety) in ≥5 to <12 years of age: Study completion date: Final Report Submission: 	 30 November 2023 31 May 2024 	

Table 57. Summary of Safety Concerns and Action Plans

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
	C4591007 substudy <u>– Troponin</u> group: A Phase 3 substudy to evaluate subclinical myocarditis in participants 5 to <12 years of age (randomized 2:1 to receive BNT162b2 10 μg or placebo) and 12 <u>-15</u> <16 years of age (open label receipt of BNT162b2 30 μg).	To obtain serum samples within the first ~4 days after vaccination for potential Troponin I testing, in order to evaluate the frequency of subclinical myocarditis amongst individuals 5 to $\frac{15 \le 16}{9}$ years of age.	Final protocol submission:Study completion:Final report submission:	 30 September 2021 30 November 2023 31 May 2024
	C4591009: A non-interventional post-approval safety study of the Pfizer-BioNTech COVID-19 mRNA-vaccine in the United States. <i>Planned</i>	To assess the occurrence of safety events of interest, including myocarditis and pericarditis in athe general US population (including use in persons <12 years administered vaccine) within selected broad population-based data sources-participating in the US sentinel system.	 Protocol submission: Monitoring report submission: Interim analysis submission: Study completion: Final study report submission: 	 31 August 2021 31 October 2022 31 October 2023 30 June 2025 31 October 2025
	C4591021: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer BioNTech Coronavirus Disease 2019 (COVID-19) vaccine Ongoing	To assess the occurrencepotential increased risk of safety eventsAdverse Events of interest in a general EU population ,Special Interest (AESI), including use in <12 receivingmyocarditis/pericarditis after being vaccinated with COVID-19 vaccine, within selected broad population based data sources.	 Protocol submission Progress report submission: Interim analysis submission: Study completion: Final study report submission: 	 11 August 2021 30 September 2021 31 March 2022 30 September 2022 31 March 2023 30 September 2023 31 March 2024 31 March 2024 30 September 2024

Safaty Concome	Orgaing/Donnad Action	Summory of Objectives	Milestones	Due dates
Safety Concerns	Ongoing/Planned Action C4591038 (former, C4591021 substudy: Post Conditional approval active surveillance study among individuals in Europe receiving the Pfizer-BioNTech Coronavirus Disease 2019 (COVID-19) vaccine. Substudy to investigate natural history of post- vaccination myocarditis and pericarditis. Planned	Summary of Objectives To describe the natural history of post-vaccination myocarditis/pericarditis, including recovery status, risk factors, and/or identification of serious cardiovascular outcomes within 1 year of myocarditis/pericarditis diagnosis among individuals vaccinated with BNT162b2 including any use in <12 year-old as well as individuals not vaccinated with a COVID-19 vaccine.	 Final protocol submission: Study completion: Final study report submission: 	 31 January 2022 31 March 2024 30 September 2024
	C4591023: Phase 1 open label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer- blinded safety, tolerability, and immunogenicity study of a SARS-CoV-2 RNAmRNA vaccine candidate against COVID-19 in healthy infants <6 months of age. Phase 1: open-label dose finding portion up to 3 dose levels with 16 participants per dose level. Phase 2/3: Children ≥5 to <12 years of age are randomized 2:1 at selected dose level of BNT162b2 at a 21-day interval. Design and subject numbers to be confirmed.	To evaluate safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy infants < 6 months of age	 Final protocol submission: Study completion: Final study report submission: 	 31 January 2022 31 July 2024 31 October 2024

Table 57. Summary of Safety Concerns and Action Plans

Safety Concerns	Ongoing/Planned Action	Summary of Objectives	Milestones	Due dates
Safety Concerns	Oligoling/Thanked Tectori	Summary of Objectives	intrestones	Duc unics
	Planned			
	C4591036: Pediatric Heart Network Study: Working title:	To characterize the clinical course, risk factors, resolution,	Protocol submission:	• 30 November 2021
	MyocarditisLow interventional	long-term sequelae, and quality	Study completion:	• 31 December 2026
	<u>cohort study of</u> <u>myocarditis</u> /pericarditis follow-	of life in children and young adults <21 years, including any	• Final study report submission:	• 31 May 2027
	up study within the Pediatric	use in < 12 -year oldsold, with		
	Heart Networkassociated with Comirnaty in persons less than	acute post-vaccine myocarditis/pericarditis.		
	21 years of age.	inyocardins/pericardins.		
	Planned			
a. FDA was informed (Response to FDA 12 May 2021 Information Request Regarding Active Surveillance Studies) that the first milestone (Interim				
Report submission due 30 June 2021) is delayed due to a change in study collaboration; therefore, it has been removed from this table. ba. Enrolment of participants into study C4591015 was stopped due to recruitment challenges as a result of global recommendations for COVID-19				
vaccination in pregnant women and the increased availability of COVID-19 vaccines. Enrolment of new participants was stopped on 25 October 2021.				
Participants already enrolled will continue follow up evaluations until study end as planned. For this reason, study completion date was changed from 31				
October 2022 to 31 August 2022 and final study report submission date was changed from 31 May 2023 to 31 March 2023".				
b. Study originally included in the PVP to address the Missing Information "Use in pediatric individuals < 16 years of age".				
e.c. Due date updated from 31 JulyOctober 2021 because the last subject visit for this group will not be until to 31 December 2021 to implement additional				
CBER requests (received on 24 September 2021-) for the 12 to \leq 15 years sBLA (FDA already informed of this change on 22 October 2021).				
d-d. Due date updated from 31 January 2023 for the same reason above.				

* Milestones deleted as this a voluntary sponsor study (as per FDA Information Request received to BLA 125742/0, dated 13 August 2021, where the FDA characterized both studies as "voluntary" and therefore no longer commitment).

ANNEX

3.2. Pharmacovigilance Methods

- BNT162b2 Vaccine: BNT162b2 Data Capture Aids:
 - o Pfizer-BioNTech COVID-19 Vaccine VAED Data Capture Aid.
 - Pfizer-BioNTech COVID-19 Vaccine Anaphylactic Reaction Data Capture Aid.

3.2.1. List of Studies Included in the Pharmacovigilance Plan

C4591001

C4591007

C4591007 substudy (Subclinical myocarditis evaluation Troponin group)

C4591007 substudy (To study lowerLower dose levels of BNT162b2evaluation group)

C4591008

C4591009

C4591011

C4591012

C4591014

C4591015

C4591021

C4591038 (former, C4591021 substudy

C4591022

C4591023

C4591031 substudy B

BNT162-01 cohort 13 WI235284

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WI255886

C4591036

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 https://www.worldometers.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronavirus/#countrieshttps://www.worldoweters.info/coronaviru
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