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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

## APPLICATION TO MARKET A NEW OR ABBREVIATED NEW DRUG OR BIOLOGIC FOR HUMAN USE

(Title 21, Code of Federal Regulations, Parts 314 & 601)

Form Approved: OMB No. 0910-0338 Expiration Date: March 31, 2020 See PRA Statement on page 3.

1. Date of Submission (mm/dd/yyyy)

Reset Form

08/20/2021

APPLICANT INFORMATION  2. Name of Applicant  Pick Many Sectoring Could I						
BioNTech Manufacturing GmbH  3. Telephone Number (Include country code if applicable and area code) 4. Facsimile (FAX) Number (Include country						
+49 (0) 6131 9084-7593 code if applicable and area code) +49 (0) 6131 9084-390						
5. Applicant Address						
Address 1 (Street address, P.O. box, company name c/o)				Email Address		
An der Goldgrube 12				Ruben.Rizzi@biontech.de		
Address 2 (Apartment, suite, unit, buildir	ng, floor, etc.)			Applicant DUNS		
City	State/Prov	/ince/Region	1	117645848		
Mainz	N/A	<b>.</b>		U.S. License Number if previously issued		
Country	J.	ZIP or Pos	stal Code			
Germany		55131		2229		
6. Authorized U.S. Agent (Required for nor	-U.S. applicants)					
Authorized U.S. Agent Name				Telephone Number (Include area code)		
Elisa Harkins, Global Regulatory Lead,	Pfizer Global Regu	ulatory Affai	irs - Vaccines	215-280-5503		
Address 1 (Street address, P.O. box, cor	mpany name c/o)			FAX Number (Include area code)		
500 Arcola Road						
Address 2 (Apartment, suite, unit, buildir	g, floor, etc.)			845-474-3500		
				Email Address		
City	State			Elisa.HarkinsTull@pfizer.com		
Collegeville	PA			U.S. Agent DUNS		
ZIP Code 19426						
	<u> </u>					
PRODUCT DESCRIPTION	7. NDA, ANDA, c 125742	or BLA Appli	cation Number	8. Supplement Number ( <i>If applicable</i> )		
9. Established Name (e.g., proper name, U	ISP/USAN name)					
[COVID-19 mRNA Vaccine (nucleoside mo	odified)]					
10. Proprietary Name (Trade Name) (If any	<i>'</i> )					
COMIRNATY						
11. Chemical/Biochemical/Blood Product N	, .,					
COVID-19 Vaccine (BNT162, PF-0730204	<del></del>					
12. Dosage Form	13. Strengt	hs		14. Route of Administration		
Liquid	30 mcg			Intramuscular		
15A. Proposed Indication for Use	i	s this indicat	ion for a rare disease (	prevalence <200,000 in U.S.)?		
Active immunization to prevent COVID-19 cause SARS-CoV-2 in individuals ≥16 years of age		oes this pro	duct have an FDA	If yes, provide the Orphan		
SAKS-Cov-2 in individuals \(\geq 10\) years of age	Orphan Designation for this			Designation number for this Continuation		
	indication?			indication: Page for #15		
			Yes No			
15B. SNOMED CT Indication Disease Term	· · ·					
COVID-19; SARS-CoV-2; Disease caused	by severe acute res	spiratory syn	drome coronavirus 2;	SARS-CoV-2 vaccination; COVID-19 vaccination		
APPLICATION INFORMATION	16. Application T (Select one)		New Drug Application			
Abbreviated New Drug Application (ANDA)						
17. If an NDA, identify the type ☐ 505(b)(1) ☐ 505(b)(2) 18. If a BLA, identify the type ☑ 351(a) ☐ 351(k)						
19. If a 351(k), identify the biological refere	nce product that is	s the basis f	or the submission.			
Name of Biologic: Holder of Licensed Application:						
20. If an ANDA, or 505(b)(2), identify the lis	sted drug product t	that is/are th	ne basis for the submis	ssion.		
Name of Drug:			Application Number	of Relied Upon Product:		
Indicate Patent Certification:  P1	Indicate Patent Certification: P1 P2 P3 P4 Section viii - MOU Statement of no relevant patents					

	Previous Page N	lext Page							
21. §   	Submission (See instructions) Product Correspondence Request for Proprietary Na		Labeling Supple Supplement  Other (Speci	Postm	CMC Supplement carketing Requirements or 0	☐ Efficacy Supplement ☐ Annual Report Commitments ☐ Periodic Safety Report			
	Submission Presubmis Sub-Type Initial Subr	_	Amendment Resubmission		23. If a supplement, identified the appropriate categ	' I LUDE I PHOLADOROVALIPA)			
	For Originals and all Supplem combination product (21 CFR				bination Product (See instructions)	Request for Designation (RFD) Number			
	Does the submission contain:  Only Pediatric data?  Yes	✓ No	ıman factors inforn Î Yes <b>√</b> No	nation?	26. Proposed Marketing  Prescription Prod	Status (Select one) duct (Rx) Over-The-Counter Product (OTC)			
	Reasons for Submission oonse to FDA 20 August 2021 Sev	enth Round of Co	omments on the Draf	t Packag	ge Insert for COMIRNATY				
E	Establishment Information <i>(Fu</i> Establishment Name Pharmacia and Upjohn Company I		nt information shou	ıld be p	provided in the body of the	application.)			
7	Address 1 (Street address, P.O 7000 Portage Road Address 2 (Apartment, suite, ur		,			Registration (FEI) Number 1810189			
 	Dity		State/Province/F	Region		MF Number			
C	Kalamazoo Country USA		MI ZIP 490		tal Code	Establishment DUNS Number 618054084			
Is	s the establishment new to the		✓ Yes  No		What is the status of the e	establishment?  Active Inactive Withdrawn			
E	Establishment Contact Informa	ation at the site.	/facility						
	lame of Contact for the Establ	shment				Telephone Number (Include area code)			
(b) (d	, , ,					(b) (6)			
						FAX Number (Include area code) (b) (6)			
						Email Address			
						(b) (6)			
I	Manufacturing Steps and/or Type of Testing  LNP production and bulk drug product formulation, Fill and finish, Primary packaging, Secondary packaging,  Drug product testing  Is the site ready for inspection?  If No, when will site be ready? (mm/dd/yyyy)								
						Continuation Page for #28			
	•		,	` , .	, ,	MFs referenced in the current application.)			
IND	IND 19736, DMF 012683, DMF 9543, DMF 15209, DMF 011793, DMF 011820, DMF 011321, DMF 10953,  Contin. Page for #29								
30.	This application contains the f	ollowing items	(Select all that ap	ply)					
	1. Index 2. Labeling (Select one): Draft Labeling Final Printed Labeling 3. Summary (21 CFR 314.50 (c))								
	4. Chemistry Section  A. Chemistry, manufacturing, and controls information (e.g., 21 CFR 314.50(d)(1); 21 CFR 601.2)  B. Samples (21 CFR 314.50 (e)(1); 21 CFR 601.2 (a)) (Submit only upon FDA's request)  C. Methods validation package (e.g., 21 CFR 314.50(e)(2)(i); 21 CFR 601.2)								
	5. Nonclinical pharmacology and toxicology section (e.g., 21 CFR 314.50(d)(2); 21 CFR 601.2)  6. Human pharmacokinetics and bioavailability section (e.g., 21 CFR 314.50(d)(3); 21 CFR 601.2)								
	7. Clinical microbiology section (e.g., 21 CFR 314.50(d)(4))  8. Clinical data section (e.g., 21 CFR 314.50(d)(5); 21 CFR 601.2)  Item 30 continued on page 3								

Previous Page Next Pag	e							
30. This application contains the following	items (Continued; s	select all that a	apply)					
9. Safety update report (e.g., 21 CFR 314.50(d)(5)(vi)(b); 21 CFR 601.2)								
11. Case report tabulations (e.g., 2 21 CFR 601.2)	; [	12. Case report forms (e.g., 21 CFR 314.50 (f)(2); 21 CFR 601.2)				CFR 601.2)		
13. Patent information on any patent that claims the drug/ biologic (21 U.S.C. 355(b) or (c))  14. A patent certification with respect to any patent that claims drug/biologic (21 U.S.C. 355 (b)(2) or (j)(2)(A))							at claims the	
15. Establishment description (21	CFR Part 600, if app	plicable)	16. Deba	arment cert	tification <i>(FD&amp;C A</i>	act 306 (k)(1))		
17. Field copy certification (21 CFF	R 314.50 (I)(3))	[				Form FDA 3397, G 92, or MDUFA Forr		
19. Financial Disclosure Informatio	n <i>(21 CFR Part 54)</i>	-						
20. Other (Specify):	,							
CERTIFICATION I agree to update this application with new safety information about the product that may reasonably affect the statement of contraindications, warnings, precautions, or adverse reactions in the draft labeling. I agree to submit safety update reports as provided for by regulation or as requested by FDA. If this application is approved, I agree to comply with all applicable laws and regulations that apply to approved applications, including, but not limited to, the following:  1. Good manufacturing practice regulations in 21 CFR Parts 210, 211 or applicable regulations, Parts 606, and/or 820.  2. Biological establishment standards in 21 CFR Part 600.  3. Labeling regulations in 21 CFR Parts 201, 606, 610, 660, and/or 809.  4. In the case of a prescription drug or biological product, prescription drug advertising regulations in 21 CFR Part 202.  5. Regulations on making changes in application in FD&C Act section 506A, 21 CFR 314.71, 314.72, 314.97, 314.99, and 601.12.  6. Regulations on Reports in 21 CFR 314.80, 314.81, 600.80, and 600.81.  7. Local, state, and Federal environmental impact laws.  If this application applies to a drug product that FDA has proposed for scheduling under the Controlled Substances Act, I agree not to market the product until the Drug Enforcement Administration makes a final scheduling decision.  The data and information in this submission have been reviewed and, to the best of my knowledge, are certified to be true and accurate.								
Warning: A willfully false statement is a cri 31. Typed Name and Title of Applicant's Re			·			32. Date (mm/c	ld/yyyy)	
Elisa Harkins, Global Regulatory Lead, Global F	<del>,                                     </del>					08/20/2021		
33. Telephone Number (Include country code if applicable and area code) 215-280-5503	34. FAX Number applicable an 845-474-3500		ntry code if		il Address kinsTull@pfizer.co			
36. Address of Applicant's Responsible Off				Енѕа.паі	T unaprizer.com	III		
Address 1 (Street address, P.O. box, cor 500 Arcola Road Address 2 (Apartment, suite, unit, buildir	mpany name c/o)				_			
City Collegeville	State/Provi	ince/Region						
Country	111	ZIP or Postal	I Code					
United States of America		19426						
37. Signature of Applicant's Responsible C Other Authorized Official Elisa Harkins Digitally signed by Eli DN: o-Pfizer Inc, cn=	sa Harkins Tull	Sign 38	38. Countersi	gnature of	Authorized U.S.	Agent	Sign	
Tull  DN: o=Pfizer Inc, cn=Elisa Harkins Tull Reason: I attest to the accuracy and integrity of this document Date: 2021 08 20 16:32:33 -04'00'								
The information below applies only to requirements of the Paperwork Reduction Act of 1995.								
The burden time for this collection of informat including the time to review instructions, sea data needed and complete and review the countries burden estimate or any other aspect of for reducing this burden to the address to the	ion is estimated to av rch existing data sou ollection of informatic this information coll	rerage 24 hours rces, gather and on. Send comme	s per response nd maintain the nents regardin	e, E g C s F	Department of Heal Food and Drug Adn Office of Operations	Ith and Human Sei ninistration s on Act (PRA) Staff		
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_	T CONTINUESTION DACE FOR ITEM 20. Fate						

FIRST CONTINUATION PAGE FOR ITEM 28 – Establishment Information	Provide information for additional establishments below, as needed.		
Establishment Name Pfizer Manufacturing Belgium NV			
Address 1 (Street address, P.O. box, company name c/o)	Registration (FEI) Number		
Rijksweg 12	1000654629		
Address 2 (Apartment, suite, unit, building, floor, etc.)	MF Number		
City State/Province/Region			
Puurs N/A	E ( I I I I I I I I I I I I I I I I I I		
Country ZIP or Postal Code	Establishment DUNS Number		
Belgium 2870	370156507		
Is the establishment new to the application?  What is the status of the			
✓ Yes No Pending	Active Inactive Withdrawn		
Establishment Contact Information at the site/facility			
Name of Contact for the Establishment	Telephone Number (Include area code)		
(b) (6) (b) (6)	(b) (6)		
	FAX Number (Include area code)		
	(b) (6)		
	Email Address		
	(b) (6)		
Manufacturing Steps and/or Type of Testing	Is the site ready Yes No N/A		
LNP production and bulk drug product formulation, Fill and finish, Primary packaging, Secondary	for inspection?		
packaging, Drug product testing	If No, when will site be ready? (mm/dd/yyyy)		
	ready: (//////dd/yyyyy)		
Establishment Name			
Wyeth BioPharma Division of Wyeth Pharmaceuticals LLC	Registration (FEI) Number		
Address 1 (Street address, P.O. box, company name c/o)  1 Burtt Road			
Address 2 (Apartment, suite, unit, building, floor, etc.)	1222181		
	MF Number		
City State/Province/Region			
Andover	Establishment DUNS Number 174350868		
Country ZIP or Postal Code United States 01810			
Is the establishment new to the application?  What is the status of the	 establishment?		
✓ Yes □ No ✓ Pending	Active Inactive Withdrawn		
Establishment Contact Information at the site/facility			
Name of Contact for the Establishment (b) (6)	Telephone Number (Include area code)		
(b) (6)	(b) (6)		
	FAX Number (Include area code)		
	, ,		
	(b) (6)		
	Email Address		
	(b) (6)		
Manufacturing Stone and/or Type of Tacting	Is the site ready.		
Manufacturing Steps and/or Type of Testing	Is the site ready Yes No N/A N/A		
Manufacturing Steps and/or Type of Testing  Manufacture of drug substance, Drug substance testing, Drug product testing	for inspection?  If No, when will site be		
I	for inspection?		

Remove Continuation Page	Return to Form	
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SECOND CONTINUATION PAGE FOR ITEM 28 – Establishment Information						Provide information for additional establishments below, as needed.			
	Establishment Name Pfizer Inc								
	Address 1 (Street address, P.O. box, company i	F	Registration (FEI) Number						
	875 Chesterfield Parkway West		1940118						
	Address 2 (Apartment, suite, unit, building, floor, etc.)						MF Number		
	City	State/Province/Region							
	Chesterfield	MO	MO				t DUNS Nun	ahor	
	Country		ZIP or Pos	tal Code			it DONS Null	ibei	
	United States		63017			004954111			
	Is the establishment new to the application?	Yes	No	What is the status of the Pending	e est	ablishment? Active	Inactive	Withdrawn	
	Establishment Contact Information at the site/f	acility							
	Name of Contact for the Establishment	<b>y</b>			П	elephone Nu	ımber (Includ	de area code)	
	(b) (6)					•	,	,	
(b)	(6)					(b) (6)			
, ,					F	AX Number	(Include are	a code)	
						(b) (6)			
						Email Addres	s		
						b) (6)			
	Manufacturing Steps and/or Type of Testing		I.			le the site r	oady 🗔		
						Is the site ready for inspection?			
	Drug substance testing, Drug product testing					If No, when	will site be		
						ready? (mr	n/dd/yyyy)		
	Establishment Name								
	Pfizer Ireland Pharmaceuticals								
	Address 1 (Street address, P.O. box, company i	name c/o)			F	Registration (	FEI) Numbe	r	
	Grange Castle Business Park Clondalkin				_   3	3004145594			
	Address 2 (Apartment, suite, unit, building, floor	; etc.)				//F Number			
		S			-  '	ii Number			
	City	State/Provi	nce/Region						
	Dublin 22	N/A	ZIP or Pos	tal Cada	E	Establishmen	t DUNS Nun	nber	
	Country Ireland		N/A	ital Code	9	985586408			
	Is the establishment new to the application?		1071	What is the status of the	e est	ablishment?			
		Yes [	No	✓ Pending		Active	Inactive	Withdrawn	
	Establishment Contact Information at the site/f	acility							
	Name of Contact for the Establishment (b) (6)				Т	elephone Nu	ımber ( <i>Includ</i>	de area code)	
(b)	(6)					(b) (6)			
				-	F	AX Number	(Include are	a code)	
					_  (	b) (6)			
					E	mail Addres	s		
					(	b) (6)			
	Manufacturing Steps and/or Type of Testing					Is the site r	eady 🗔		
	Drug product testing					for inspecti		es No N/A	
	Drug product testing					If No, when ready? (mr			
								tion Page for #28	
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**Remove Continuation Page** Return to Form Provide information for additional THIRD CONTINUATION PAGE FOR ITEM 28 – Establishment Information establishments below, as needed. Establishment Name Hospira Zagrab Ltd.

	Registration (FEI) Number							
	Address 1 (Street address, P.O. box, company name c/o)							
Prudnicka cesta 60								
Address 2 (Apartment, suite, unit, building, flo	or, etc.)			MF Number				
				-   Will Multiper				
City	State/Prov	rince/Regio	n					
Prigorje	Brdovecko	1		Establishment DUNS Number				
Country		ZIP or Po	ostal Code	500625201				
Croatia		10291		300023201				
Is the establishment new to the application?	✓ Yes	] No	What is the status of the Pending	establishment? Active Inactive Withdrawn				
Establishment Contact Information at the site	e/facility							
Name of Contact for the Establishment				Telephone Number (Include area code)				
(b) (6)				Totophone Humber (molado drea code)				
(b) (6)				(b) (6)				
				FAX Number (Include area code)				
				(b) (6)				
				Email Address				
				-				
				(b) (6)				
Manufacturing Steps and/or Type of Testing				Is the site ready  Yes  No N/A				
Drug Product Release Testing (Sterility)				for inspection?				
				If No, when will site be				
				ready? (mm/dd/yyyy)				
Establishment Name								
SGS Lab Simon SA				11				
Address 1 (Street address, P.O. box, compan	y name c/o)			Registration (FEI) Number				
Vieux Chemin du Poete 10				3004186644				
Address 2 (Apartment, suite, unit, building, flo	or, etc.)			MF Number				
				- I Will Marine				
City	State/Prov	ince/Regio	n					
Wavre	N/A			Establishment DUNS Number				
Country			ostal Code	283063907				
Belgium		1301						
Is the establishment new to the application?		_	What is the status of the					
	✓ Yes	No	✓ Pending	Active Inactive Withdrawn				
Establishment Contact Information at the site	e/facility							
Name of Contact for the Establishment				Telephone Number (Include area code)				
(b) (6)								
(b) (6)				(b) (6)				
				FAX Number (Include area code)				
				AA Nullibel (Iliciude alea code)				
				(b) (6)				
				Email Address				
				†				
				(b) (6)				
Manufacturing Steps and/or Type of Testing				Is the site ready Yes No N/A				
Drug Product Release Testing (Sterility)				for inspection? Yes I NO I N/A				
Drug Froduct Release Testing (Steffilly)	If No, when will site be							
	ready? (mm/dd/yyyy)							
				Add Fourth Continuation Page for #28				
FORM FDA 356h (08/18 - PREVIOUS EDITION	NS OBSOL	ETE\	Page 6 of 7	_FDA-CBER-2021-568β-1078764 _				

Remove Continuation Page Return to Form Provide information for additional FOURTH CONTINUATION PAGE FOR ITEM 28 – Establishment Information establishments below, as needed. Establishment Name Fresenius Kabi USA LLC (b) (4) mber Number

			ve Withdrawn
	Establishment Contact Information at the site/facility		
	Name of Contact for the Establishment	Telephone Number (	Include area code)
	Anthony Giessert		
b)	(4), (b) (6)	(b) (4), (b) (6)	
		FAX Number (Include	e area code)
		N/A	
		Email Address	
		(b) (6)	
	Manufacturing Steps and/or Type of Testing	Is the site ready	7 Vee
	manufacture, testing and release of diluent (0.9% Sodium chloride Injection, USP)	for inspection?	✓ Yes ☐ No ☐ N/A
	manufacture, testing and release of and in (0.770 bounding emoritie injection, Obt.)	If No, when will site	
		ready? (mm/dd/yy)	(y)
	Establishment Name		
	Hospira Inc.		
(b)	Address 1 (Street address PO hov company name c/o)	Registration (FEI) Ni	ımber
(U)	(7)		
			S Number
		t	ive   Withdrawn
	Establishment Contact Information at the site/facility		
	Name of Contact for the Establishment	Telephone Number (	Include area code)
/I \	Paul Lucas	(h) (d) -(l) -(0)	
(p)	(4), (b) (6)	(b) (4), (b) (6)	
		FAX Number (Include	e area code)
		(b) (4), (b) (6)	
		Email Address	
		(b) (6)	
	Manufacturian Ctans and/ou Turns of Tasting	1	
	Manufacturing Steps and/or Type of Testing	Is the site ready for inspection?	✓ Yes □ No □ N/A
	manufacture, testing and release of diluent (0.9% Sodium chloride Injection, USP)	If No, when will site	
		ready? (mm/dd/yy	
		Add Fifth Con	tinuation Page for #28
-01	RM FDA 356h (08/18 - PREVIOUS EDITIONS OBSOLETE) Page7 of 7	FDA-CBER-202	1-568 <b>β</b> -1 <u>0</u> 78765
JI	Rem	ove Continuation Page	1-5688-1078765 Return to Form