## **DEPARTMENT OF HEALTH AND HUMAN SERVICES**

Food and Drug Administration

Form Approved: OMB No. 0910-0001 Expiration Date: March 31, 2024 See PRA Statement on last page.

## TRANSMITTAL OF ADVERTISEMENTS AND PROMOTIONAL LABELING FOR DRUGS AND BIOLOGICS FOR HUMAN USE

1. Date Submitted 10/01/2021

2. Application Information		imes Single product	Multiple products	For multiple products, submit completed form and		
Application Type: BLA				specimen of advertising/promotional materials to one application of choice, and attach separate sheet addressing items 3-5 for remainder of products. Refer to No. 3 on instruction sheet.		
Application Number: 12	5742 /					
NOTE: Form FDA 2253 is	required by law	v. Reports are required t	or approved NDAs,	ANDAs (21 CFR 314.81), and BLAs (601.1	12(f)(4))	
3. Proprietary Name			4. Established Name			
COMIRNATY			[COVID-19 mRNA Vaccine (nucleoside modified)]			
			Product Code No			
5. Package Insert Date an (Latest final printed lab			6. Manufacturer Name			
08/21 LAB-1448-1.0	<b>- - - - - - - - - </b>		License No. (Biologics):			
7.		Advertisement / Prom				
a. Please check only one	: X Profession	nal Consumer				
Material Type (use FDA codes)	Dissemination/ Publication	Material ID Code	Material Description			
b.	Date c.	d.		e.		
www-website	10/01/2021	PP-CVV-USA-0429	US HCP Website - 1 Update	6+ BLA Approval - Checklist/LDV Resource	Delete Row	
f. Comments  PP-CVV-USA-0429 is USA-0357 filed 08/26/ Code PP-CVV-USA-04	2021.	rsion of PP-CVV-USA	0422 filed 09/24/	2021 and previous version PP-CVV-goes live.		

8.	Applicant's (or Agent's) Return Address	9. Responsible Official's (or Agent's)						
	Address 1 (Street address, P.O. box, company	a. Telephone Number (Include area code)						
	An der Goldgrube 12	(484) 865-5035						
	Address 2 (Apartment, suite, unit, building, floor,	b. FAX Number (Include area code)						
		(845) 474-3500						
	City	State/Pro	vince/Region	c. Email Address  Donna.Boyce@pfizer.com				
	Mainz	N/A						
	Country		ZIP or Postal Code					
	Germany 5		55131					
10	). Typed Name and Title of Responsible Offici	nt 11. Signature of Responsible Officia	al or Agent	12. Date				
Donna Boyce M.S., Senior Vice President, Global Regulatory Affairs, Global Product Development			(b) (6)  D p is by signed by (b) (6) The Conf. (c)		10/01/2021			
13. For CBER Products Only (Check one)								
Draft ★ Final								

This section applies only to requirements of the Paperwork Reduction Act of 1995.

## \*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.\*

The burden time for this collection of information is estimated to average 2 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."