DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Individual Patient Expanded Access Investigational New Drug Application (IND)

(Title 21, Code of Federal Regulations (CFR) Part 312)

Form Approved: OMB No. 0910-0814 Expiration Date: May 31, 2022 See PRA Statement on last page.

1. Patient's Initials	2. Date of Submission (mm/dd/yyyy)		
3.a. Initial Submission Select this box if this form is an initial submission for an individual patient expanded access IND,	3.b. Follow-Up Submission Select this box if this form accome a follow-up submission to an existence individual patient expanded access.	sting	
and complete only fields 4 through 8, and fields 10 and 11.	and complete the items to the rig section, and fields 8 through 11.		
4. Clinical Information			
Indication			
Brief Clinical History (Patient's age, genderequest, including an explanation of why		rapy, response to prior therapy, reason for ns)	
5. Treatment Information			
Investigational Drug Name			
Name of the entity that will supply the dru	g (generally the manufacturer)		
FDA Review Division (if known)			
Treatment Plan (Including the dose, route modifications to the treatment plan in the		d duration, and monitoring procedures. Also include	
6. Letter of Authorization (LOA), if app	icable (generally obtained from the mai	nufacturer of the drug)	
I have attached the LOA. (Attach the	e LOA; if electronic, use normal PDF func	tions for file attachments.)	
Note: If there is no LOA, consult the Fo	rm Instructions.		
license number, current employment, a	and job title. Alternatively, attach the first	ar of graduation, medical specialty, state medical few pages of physician's curriculum vitae (CV), ormal PDF functions for file attachments.)	
8. Physician Name, Address, and Cont	act Information		
Physician Name (Sponsor)	Email Address of Physician		
Address 1 (Street address, No P.O. boxes)			
Address 2 (Apartment, suite, unit, building,	Telephone Number of Physician		
City	State	Facsimile (FAX) Number of Physician	
ZIP Code	Physician's IND number, if known		

9. Contents of Submission									
This submission contains the following n follow-up communications, use Form FE	· · · · · · · · · · · · · · · · · · ·	The state of the s	all that app	oly). If none of th	he following	apply to the			
☐ Initial Written IND Safety Report			nange in Treatment Plan						
Follow-up to a Written IND Safety I	Genera	eneral Correspondence							
Annual Report	Annual Report			Response to FDA Request for Information					
Summary of Expanded Access Use	e (treatment completed)	Respon	nse to Clinical Hold						
10.a. Request for Authorization to Use	e Form FDA 3926								
I request authorization to submit this Form FDA 3926 to comply with FDA's requirements for an individual patient expanded access IND.									
10.b. Request for Authorization to Use Alternative IRB Review Procedures									
 I request authorization to obtain co the treatment use begins, in order review and approval at a convened 	to comply with FDA's require	ements for IRB review a	and approva	al. This concurr					
required materials unless I receive continue clinical investigations cov- informed consent, and that an Inst approval of this treatment use, cor request, treatment may begin with working days of treatment. I agree	vered by the IND if those itutional Review Board (Insistent with applicable Fout prior IRB approval, perto conduct the investigation.	studies are placed o RB) will be responsit DA requirements. I u rovided the IRB is no tion in accordance w	n clinical hole for initial inderstand otified of the ith all other	nold. I also cert al and continui I that in the cas se emergency t er applicable re	tify that I wi ing review a se of an em treatment w	ll obtain and nergency vithin 5			
WARNING: A willfully false sta	tement is a criminal of	fense (U.S.C. Title	18, Sec.	1001).					
Signature of Physician									
To enable the signature field, please fill out all prior required fields. For a list of required fields which have not yet been filled out, please click here.									
For FDA Use Only									
Date of FDA Receipt	Is this an emergency indiv	vidual patient IND?	Is this indication for a rare disease (prevalence < 200,000 in the U.S.)?						
IND Number	☐ Yes	No	☐ Yes ☐ No		☐ No				
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 45 minutes 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 Operations 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."