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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: February 28, 2023 See PRA Statement on page 3.

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

APPLICANT INFORMATION 2	2. Name of Applic	cant					
3. Telephone Number (Include country code i	if applicable and a	area code)	4. Facsimile (FAX) N code if applicable a	umber (Include country and area code)			
5. Applicant Address							
Address 1 (Street address, P.O. box, comp	pany name c/o)			Email Address			
Address 2 (Apartment, suite, unit, building	, floor, etc.)			Applicant DUNS			
City	State/Province/Region						
Country	ZIP or Postal Code			U.S. License Number if previously issued			
6. Authorized U.S. Agent (Required for non-l	U.S. applicants)			II			
Authorized U.S. Agent Name	Telephone Number (Include area code)						
Address 1 (Street address, P.O. box, comp	Address 1 (Street address, P.O. box, company name c/o)						
Address 2 (Apartment, suite, unit, building	Address 2 (Apartment, suite, unit, building, floor, etc.)						
City	State			U.S. Agent DUNS			
ZIP Code	U.S. Agent DONS						
PRODUCT DESCRIPTION 7	7. NDA, ANDA, o	r BLA Appli	cation Number	8. Supplement Number (If applicable)			
I Reges Bestur Herr	, , , ,			(),,,			
9. Established Name (e.g., proper name, US	P/USAN name)						
10. Proprietary Name (Trade Name) (If any)							
11. Chemical/Biochemical/Blood Product Na	me (If any)						
12. Dosage Form	13. Strength	ns		14. Route of Administration			
15A. Proposed Indication for Use	Is	this indicat	on for a rare disease (prevalence <200,000 in U.S.)?			
	О		duct have an FDA gnation for this	If yes, provide the Orphan Designation number for this indication: Continuation Page for #15			
15B. SNOMED CT Indication Disease Term ((Use continuation	n page for e		on and respective coded disease term)			
APPLICATION INFORMATION	16. Application Ty	ne \Box	Name Davis A I' I'	(AIDA) Distance III A. F. F. (BLA)			
AFFEIGATION IN ORMATION	(Select one)		New Drug Application Abbreviated New Drug				
17. If an NDA, identify the type 505(b			18. If a BLA, identify	the type 351(a) 351(k)			
19. If a 351(k), identify the biological reference	ce product that is	the basis f					
Name of Biologic:			Holder of Licensed A				
20. If an ANDA, or 505(b)(2), identify the lists	ed drug product t	hat is/are th					
Name of Drug: Application Number of Relied Upon Product:							
Indicate Patent Certification: P1	☐ P2 ☐ F	P3 []	P4 Section viii	- MOU Statement of no relevant patents			

	Previous Page Next Page									
21.	Submission (See									
22.	2. Submission Sub-Type Presubmission Amendment Sub-Type Submission Resubmission Resubmission									
24.	. For Originals and all Supplements, is the product a Combination Product combination product (21 CFR 3.2(e))?					Request for Designation (RFD) Number				
	Does the submission contain: Only Pediatric data? Yes No Yes No 26. Proposed Marketing Status (Select one) Prescription Product (Rx) Over-The-Counter Product (OTC)									
	27. Reasons for Submission									
28.	Establishment Information (Full establishment Establishment Name	t information	should be p	provided in the body of t	he applica	ation.)				
	Establishment Name									
	Address 1 (Street address, P.O. box, company name c/o)					Registration (FEI) Number				
	Address 2 (Apartment, suite, unit, building, floor	Address 2 (Apartment, suite, unit, building, floor, etc.)				MF Number				
	City	State/Provi	nce/Region			12 1	1 DUNG N			
-	Country	ZIP or Pos	tal Code	Estab	Establishment DUNS Number					
	Is the establishment new to the application?	No	What is the status of the		establishment? Active Inactive Withdrawn					
	Establishment Contact Information at the site/	facility								
	Name of Contact for the Establishment Telephone Number (Include area code)							area code)		
-	Address 1 (Street address, P.O. box, company name c/o)									
	Address 2 (Apartment, suite, unit, building, floor, etc.)					FAX Number (Include area code)				
	City	State/Provi	State/Province/Region ZIP or Postal Code			Email Address				
	Country									
	Manufacturing Steps and/or Type of Testing						Is the site ready Yes No N/A for inspection? If No, when will site be ready? (mm/dd/yyyy)			
							Continuation Pa	age for #28		
29.	Cross References (List related BLAs, INDs, N	IDAs, PMAs,	, 510(k)s, IC	Es, BMFs, MAFs, and	DMFs refe	erenced	d in the current	application.)		
								Contin. Page for #29		
30.	30. This application contains the following items (Select all that apply)									
	1. Index 2. Labeling (Select one): Draft Labeling Final Printed Labeling 3. Summary (21 CFR 314.50 (c))							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 6. Human pharmac				cokinetics	okinetics and bioavailability section 4.50(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)						; 21 CFR 601.2)			
							Item 30 co	ntinued on page 3		

Previous Page Next Page	е								
30. This application contains the following it	tems (Continued; s	select all tha	at apply)						
9. Safety update report (e.g., 21 CF 21 CFR 601.2)	(b);	10. Statistical section (e.g., 21 CFR 314.50(d)(6); 21 CFR 601							
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)							
	☐ 13. Patent information on any patent that claims the drug/ ☐ 14. A patent certification						tion with respect to any patent that claims the $U.S.C.\ 355\ (b)(2)\ or\ (j)(2)(A))$		
15. Establishment description (21 C	CFR Part 600, if app	olicable)	16. Deb	arment cert	tification (FD&C A	ct 306 (k)(1))			
17. Field copy certification (21 CFR		18. User Fee Cover Sheet (PDUFA Form FDA 3397, GDUFA Form FDA 3794, BsUFA Form FDA 3792, or MDUFA Form FDA 3601)							
19. Financial Disclosure Information	n (21 CFR Part 54)								
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 crin		Code, title	18, section 10	01. ———		<u> </u>			
31. Typed Name and Title of Applicant's Responsible Official						32. Date (mm/dd/yyyy)			
33. Telephone Number (Include country code if applicable and area code) 34. FAX Number (Include applicable and area code)				35. Ema	il Address				
36. Address of Applicant's Responsible Offi	cial								
Address 1 (Street address, P.O. box, con	npany name c/o)								
Address 2 (Apartment, suite, unit, buildin	Address 2 (Apartment, suite, unit, building, floor, etc.)								
City	State/Provi	noo/Pogion							
City State/Province/Region									
Country ZIP or Po			ostal Code						
37. Signature of Applicant's Responsible Official or Other Authorized Official Sign 38. Countersignature of Authorized U.S. Agent Sign Sign									
The information below applies only to requirements of the Paperwork Reduction Act of 1995.									
The burden time for this collection of information is estimated to average 24 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 the address to the right: Department of Health and Human Services Food and Drug Administration Office of Operations Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov									
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