Form Approved: (OMB No. 0910 - (297 Expiration Date:	December 31, 2006	See instructions for OMB Statement.	

DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION

PRESCRIPTION DRUG USER FEE COVERSHEET

A completed form must be signed and accompany each new drug or biologic product application and each new supplement. See exceptions on the reverse side. If payment is sent by U.S. mail or courier, please include a copy of this completed form with payment. Payment instructions and fee rates can be found on CDER's website: http://www.fda.gov/cder/pdufa/default.htm

1. APPLICANT'S NAME AND ADDRESS	4. BLA SUBMISSION TRACKING NUMBER (STN) / NDA NUMBER				
	5. DOES THIS APPLICATION REQUIRE CLINICAL DATA				
2. TELEPHONE NUMBER	FOR APPROVAL?				
	[]YES []NO				
	IF YOUR RESPONSE IS "NO" AND THIS IS FOR A				
	SUPPLEMENT, STOP HERE AND SIGN THIS FORM				
	IF RESPONSE IS "YES", CHECK THE APPROPRIATE				
	RESPONSE BELOW:				
	[] THE REQUIRED CLINICAL DATA ARE CONTAINED IN THE APPLICATION				
	[] THE REQUIRED CLINICAL DATA ARE SUBMITTED BY REFERENCE TO:				
3. PRODUCT NAME	6. USER-FEE I.D. NUMBER				
7. IS THIS APPLICATION COVERED BY ANY OF THE FOLLOWING USER FEE EXCLUSIONS? IF SO, CHECK THE					
APPLICABLE EXCLUSION/					
[] A LARGE VOLUME PARENTERAL DRUG PRODUCT [] A 505(b)(2) APPLICATION THAT DOES NOT REQUIRE A					
APPROVED-UNDER SECTION 505 OF THE FEDERAL FOOD, DRUG, AND COSMETIC ACT BEFORE 9/1/92 (Self Explanatory					
[] THE APPLICATION QUALIFIES FOR THE ORPHAN	/ [] THE APPLICATION IS SUBMITTED BY A STATE OR				
EXCEPTION UNDER SECTION 736(a)(1)(E) of the Federal	FEDERAL GOVERNMENT ENTITY FOR A DRUG THAT IS NOT				
Food,Drug, and Cosmetic Act	DISTRIBUTED COMMERCIALLY				
8. HAS A WAIVER OF AN APPLICATION FEE BEEN GRANTED	FOR THIS APPLICATION? [] YES [] NO				
	stimated to average 30 minutes per response, including the time for				
reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing					
the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:					
	rug Administration An agency may not conduct or				
Food and Drug Administration CDER, HFD	0-94 sponsor, and a person is not				
	lawn Drive, Room 3046 required to respond to, a collection				
1401 Rockville Pike Rockville, M Rockville, MD 20852-1448	ID 20852 of information unless it displays a currently valid OMB control				
	number.				
SIGNATURE OF AUTHORIZED COMPANY REPRESENTATIVE	TITLE DATE				
9. USER FEE PAYMENT AMOUNT FOR THIS APPLICATION					
Form FDA 3397 (12/03)					