DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION					Form Approved: OMB No. 0910-0338 Expiration Date: August 31, 2006 See OMB Statement on page 2.	
					FOR FDA USE ONLY	
APPLICATION TO MARKET A NEW DRUG, BI OR AN ANTIBIOTIC DRUG FOR HUMAN					APPLICATION NUMBER	
(Title 21, Code of Fe	deral Regulations, Par	ts 314 & 601	l)			
APPLICANT INFORMATION						
NAME OF APPLICANT			DATE OF SUBMISSION December 9, 2004			
Merck & Co., Inc.						
TELEPHONE NO. (include Area Code) 484-344-4511			FACSIMILE (FAX) Number (include Area Code) 484-344-3682			
APPLICANT ADDRESS (Number, Street, City, State, Country, ZIP Code or Mail Code, and U.S. License number if previously issued): Sumneytown Pike, P.O. Box 4, BLX-29 West Point, PA 19486			AUTHORIZED U.S. AGENT NAME & ADDRESS (Number, Street, City, State, ZIP Code, telephone & FAX number) IF APPLICABLE Florence F. Vickers, Ph.D., F.C.P. Director, Worldwide OTC Regulatory Affairs			
PRODUCT DESCRIPTION						
NEW DRUG OR ANTIBIOTIC APPLICATION N	IUMBER, OR BIOLOGICS L	ICENSE APPL	ICATION NUM	BER (If p	reviously issued) 21-213	
ESTABLISHED NAME (e.g., Proper name, USP Nonprescription lovastatin, MK-08			VACOR D		e name) IF ANY	
CHEMICAL/BIOCHEMICAL/BLOOD PRODUCT 8aβ]]-1,2,3,7,8,8a-hexahydro-3,7-dimethy naphthalenyl 2-methylbutanoate			8β(2S*,4S*), pyran-2-yl)eth	iyl]-1-	CODE NAME (If any)	
DOSAGE FORM: Tablet	STRENGTHS: 20 mg	<u> </u>		ROUTE	OF ADMINISTRATION: Oral	
(PROPOSED) INDICATION(S) FOR USE:	<u> </u>					
To help lower LDL "bad" choleste	rol, which may preve	ent a first h	eart attack	•		
APPLICATION INFORMATION						
APPLICATION TYPE	<u> </u>					
(check one) 🙀 NEW DRUG APPLICA	TION (21 CFR 314.50) DLOGICS LICENSE APPLIC			FED NEW	/ DRUG APPLICATION (ANDA, 21 CFR 314.94)	
IF AN NDA, IDENTIFY THE APPROPRIATE TY	PE 🔯 505 (b)(1)	□ 50	5 (b)(2)			
IF AN ANDA, OR 505(b)(2), IDENTIFY THE RE Name of Drug	FERENCE LISTED DRUG F		AT IS THE BAS er of Approved A			
	ORIGINAL APPLICATION					
		STABLISHMENT	MENT TO A PEND DESCRIPTION S DLS SUPPLEMENT	UPPLEME		
	ALREPORT E E	STABLISHMENT	DESCRIPTION S			
	ALREPORT CHEMISTRY MANUFACTURIN , PROVIDE LETTER DATE	STABLISHMENT	DESCRIPTION S DLS SUPPLEMENT NT TO PARTIA	UPPLEME - L SUBMI		
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LABELING SUPPLEMENT	ALREPORT DE LE CHEMISTRY MANUFACTURIN , PROVIDE LETTER DATE ATE CATEGORY	STABLISHMENT G AND CONTRO OF AGREEME CBE	DESCRIPTION S US SUPPLEMENT NT TO PARTIA		NT CHER SSION: Prior Approval (PA)	
LABELING SUPPLEMENT	ALREPORT E E CHEMISTRY MANUFACTURIN , PROVIDE LETTER DATE ATE CATEGORY	STABLISHMENT G AND CONTRO OF AGREEME CBE	DESCRIPTION S LS SUPPLEMENT NT TO PARTIAL CBE	UPPLEME	Advisory Committee	
LABELING SUPPLEMENT	ALREPORT E E CHEMISTRY MANUFACTURIN , PROVIDE LETTER DATE HATE CATEGORY CATE CATEGORY PRESCRIPTION F THIS A Cablishment information and control sites for drug subst FN), DMF number, and manuf	STABLISHMENT G AND CONTRO OF AGREEME CBE CBE PRODUCT (Rx) PPLICATION IS should be pro ance and drug p acturing steps a	DESCRIPTION S US SUPPLEMENT NT TO PARTIA CBE CBE CBE CBE CBE CBE CBE CBE	UPPLEME -30 -30 -30 -30 	INT CEFFICACY SUPPLEMENT SSION: Prior Approval (PA) Advisory Committee E COUNTER PRODUCT (OTC) PAPER AND ELECTRONIC CELECTRONIC The Application.) ts may be used if necessary). Include name, address,	
LABELING SUPPLEMENT	ALREPORT E E CHEMISTRY MANUFACTURIN , PROVIDE LETTER DATE ATE CATEGORY CATE CATEGORY C	STABLISHMENT G AND CONTRO OF AGREEME CBE Tefor PRODUCT (Rx) PPLICATION IS should be pro- acturing steps a pady.	DESCRIPTION S US SUPPLEMENT NT TO PARTIAL CBE COLON CBE COLON CBE COLON	-30	INT EFFICACY SUPPLEMENT MOTHER SSION: Prior Approval (PA) Advisory Committee E COUNTER PRODUCT (OTC) PAPER AND ELECTRONIC ELECTRONIC the Application.) ts may be used if necessary). Include name, address, Final dosage form, Stability testing) conducted at the site.	

This a	pplication contains the following items:	(Check all that apply)							
X	1. Index								
	2. Labeling (check 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 CIFIR 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 (e.g., 21 CFR 314.50(d)(4))								
	8. Clinical data section (e.g., 21 CFR 314.50(d)(5); 21 CFR 601.2)								
	9. Safety update report (e.g., 21 CFR 314.50(d)(5)(vi)(b); 21 CFR 601.2)								
	10. Statistical section (e.g., 21 CFR 314.50(d)(6); 21 CFR 601.2)								
	11. Case report tabulations (e.g., 21 CFR 314.50(f)(1); 21 CFR 601.2)								
	12. Case report forms (e.g., 21 CFR 3		<u></u>						
	13. Patent information on any patent		C. 355(b) or (c))						
	14. A patent certification with respect to any patent which claims the drug (21 U.S.C. 355 (b)(2) or 0)(2)(A))								
	15. Establishment description (21 CFI								
	16. Debarment certification (FD&C Ac				· · · · · · · · · · · · · · · · · · ·				
	···		·····						
	17. Field copy certification (21 CFR 314.50 (k)(3))								
	18. User Fee Cover Sheet (Form FDA 3397)								
<u></u>	19. Financial Information (21 CFR Part 54)								
j	X 20. OTHER (Specify) Final Backyround Information for Advisory Committee								
CERTIFICATION									
I agree to update this application with new safety information about the product that may reasonably affect the statement of contra i nd ications, 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.									
	Biological establishment standards in 2								
	Labeling regulations in 21 CFR Parts 20		والمراجع						
	In the case of a prescription drug or biol Regulations on making changes in appl								
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 criminal offense, U.S. Code, title 18, section 1001.									
	TURE OF RESPONSIBLE OFFICIAL OR AG			······	DATE				
SIGNA	TURE OF RESPONSIBLE OFFICIAL OR AS	Florence F. Vic	kers, Ph.D., F.C.P.						
1+		Worldwide OT	C Regulatory Affairs		Dec. 9, 2004				
ADDRI	SS (Street, City, State, and ZIP Code)			Telephone Number	l				
	neytown Pike, P.O. Box 4, BLX-	-29		(484) 344-4511					
	Point, PA 19486	·			····				
Public reporting burden for this collection of information is estimated to average 24 hours 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:									
Food CDER 1401	tment of Health and Human Services and Drug Administration , HFD-99 Rockville Pike ille, MD 20852-1448	Food and Drug Administratic CDER (HFD-94) 12229 Wilkins Avenue Rockville, MD 20852	required to resp		nsor, and a person is not of information unless it rol number.				
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