

Attachment B

Please return this form along with the updated lists of products in Attachment B by **August 15, 2017**.

For instructions on how to complete Attachment B, please refer to the attached Dear Colleague letter.

Product Checklist	
1.	☐ Reviewed all products in Attachment B and compared it to the three publicly available lists, i.e. Prescription Drug Product List , CDER Billable Biologic Product List & CBER Billable Biologic Product List
2.	☐ Added/Deleted products, as appropriate - Notified appropriate Agency point of contact per section II and III of DCL letter
3.	☐ Contacted Orange Book Staff to discontinue CDER prescription products as needed
4.	☐ Contacted CDER User Fee Staff to discontinue CDER biologic products as needed
5.	☐ Contacted CBER User Fee Staff to discontinue CBER biologics products as needed

See examples on next page

Attachment B Example 1 - Edit Existing Product List

CDER PRODUCTS

Billing Firm: Firm Name			
Owner of Products: Product Owner Name		Notes for PDUFA User Fee staff	
NDA #/Prod	Trade Name/ Ingredient		Dosage Form/ Strength
12345 1	Product 1 Active Ingredient	e.g. the NDA was transferred to firm B on 07/02/2017	Injectable; subcutaneous 2,500IU/0.2ml (12,500IU/ML)
12345 2	Product 2 Active Ingredient	Gained TE code, should not be billed	Injectable; subcutaneous 5,000IU/0.2ml (25,000IU/ML)
56789 1	Product 1 Active Ingredient	Cross out discontinued product	Tablet, Extended Release; Oral EQ 4 MG BASE
56789 2	Product 2 Active Ingredient	Notes	Tablet, Extended Release; Oral EQ 4 MG BASE

Attachment B Example 2 - Missing PDUFA Eligible Products

CDER PRODUCTS / BIOLOGIC PRODUCTS

Billing Firm: Firm Name			
Owner of Products: Product Owner Name			
NDA #/Prod	Trade Name/ Ingredient	Dosage Form/ Strength	Notes for PDUFA User Fee Staff
NDA 082101 / 1	New NDA Product New Product Active Ingredient	New product dosage form Strength	New Approval on 06/28/2017
BLA 163590 / 0	New BLA Product New Product Active Ingredient	New product dosage form Strength 1	New Approval on 07/15/2017
BLA 163590 / 0	New BLA Product New Product Active Ingredient	New product dosage form Strength 2	New Approval on 07/15/2017
NDA 222536 / 2	NDA Product Product Active Ingredient	Product dosage form Strength	Transferred from firm xxx on 04/15/2017

Attachment B - Missing PDUFA Eligible Products

CDER PRODUCTS / BIOLOGIC PRODUCTS

Billing Firm: Firm Name

Owner of Products: Product Owner Name

NDA #/Prod	Trade Name/ Ingredient	Dosage Form/ Strength	Notes for PDUFA User Fee Staff
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