## Attachment B

Please return this form along with the updated lists of products in Attachment B by June 3, 2024.

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. <u>Prescription Drug Product List</u> , <u>CDER Billable Biologic Product List</u> & <u>CBER Billable Biologic Product List</u>
2.	Added/Deleted products, as appropriate - Notified appropriate Agency point of contacts per section III and IV 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 <u>CBER User Fee Staff</u> 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								
Trade Name: Ingredient:	Trade Name Active Ingre							
NDA/BLA #/F	Prod Approval Date		Strength / Dosage Form	Notes for PDUFA User Fee staff				
123456 / 1	03/	10/2018	Injectable; subcutaneous 2,500IU/0.2ml (12,500IU/ML)	e.g. the NDA was transferred to firm B on 03/12/2021				
123456 / 2	05/	15/2007	Injectable; subcutaneous 5,000IU/0.2ml (25,000IU/ML)	Gained TE code, should not be billed				
567890 / 1		01/2014	Tablet, Extended Release; Oral EQ 4 MG BASE	Cross out discontinued product				
567890 / 2	02/	01/2014	Tablet, Extended Release; Oral EQ 4 MG BASE	Notes				

# **Attachment B Example 2 - Missing PDUFA Eligible Products**

### **CDER PRODUCTS / BIOLOGIC PRODUCTS**

Billing Firm: Firm Name								
Owner of Products: Product Owner Name								
NDA/BLA #/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 11/28/2020					
BLA 163590/0	<i>New BLA Product</i> <i>New Product Active Ingredient</i>	New product dosage form Strength 1	New Approval on 02/18/2021					
BLA 163590 / 0	New BLA Product New Product Active Ingredient	New product dosage form Strength 2	New Approval on 02/18/2021					
NDA 222536 / 2	NDA Product Product Active Ingredient	Product dosage form Strength	Transferred from firm xxx on 03/20/202					