tobacco products removed into tobacco product class' volume of assessment to be allocated among specified classes of tobacco products. In the Federal Register of May 31, 2013 (78 FR 32581), FDA issued a proposed rule to add part 1150 (21 CFR part 1150) to require domestic tobacco product manufacturers and importers to submit to FDA information needed to calculate the amount of user fees to assess each domestic manufacturer and importer under the FD&C Act. In the Federal Register of July 10, 2014 (79 FR 39302), FDA finalized portions of the User Fee proposed rule related to cigarettes, snuff, chewing tobacco, and roll-your-own tobacco, which is codified at part 1150. In the Federal Register of May 10, 2016 (81 FR 28707), FDA finalized a rule that requires domestic manufacturers and importers of cigars and pipe tobacco to submit information needed to calculate the amount of user fees assessed under the FD&C Act.

FDA is issuing this draft guidance consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the current thinking of FDA on the responses to the frequently asked questions set forth in the guidance. It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

I. Background

FDA is announcing the availability of a draft guidance for industry entitled “Tobacco Product User Fees: Responses to Frequently Asked Questions.” This draft guidance provides information in response to frequently asked questions related to tobacco product user fees assessed and collected under section 919 of the FD&C Act (21 U.S.C. 387s). In particular, this draft guidance provides information regarding the submission of information needed to assess user fees owed by each domestic manufacturer or importer of tobacco products and how FDA determines whether a company owes user fees in each quarterly assessment. The current Form FDA 3852, “Report of Tobacco Product Removals Subject to Tax for Tobacco Product User Fee Assessments,” discussed in this draft guidance, is available at https://www.fda.gov/media/88957/download.

The Family Smoking Prevention and Tobacco Control Act (Pub. L. 111–31) (Tobacco Control Act) was enacted on May 20, 2010. The Tobacco Control Act requires FDA, in accordance with that act, to regulate tobacco products. Included in the Tobacco Control Act is the requirement that FDA assess and collect user fees.

Section 919(a) of the FD&C Act requires FDA, in accordance with that section, to “assess user fees on, and collect such fees from, each manufacturer and importer of tobacco products subject to” the tobacco product provisions of the FD&C Act (chapter IX of the FD&C Act). Under the calculations required by section 919 of the FD&C Act, the tobacco products that are subject to user fee assessments are cigarettes, snuff, chewing tobacco, roll-your-own tobacco, cigars, and pipe tobacco. The total amount of user fees for each fiscal year is specified in section 919(b)(1) of the FD&C Act, and, under section 919(a), FDA is to assess and collect one-fourth of that total each quarter of the fiscal year. The FD&C Act provides for the total quarterly assessment to be allocated among specified classes of tobacco products. The class allocation is based on each tobacco product class’ volume of tobacco products removed into commerce. Within each class of tobacco products, an individual domestic manufacturer or importer is assessed a user fee based on its market share for that tobacco product class.

II. Paperwork Reduction Act

While this guidance contains no collection of information, it does refer to previously approved FDA collections of information. Therefore, clearance by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501–3521) is not required for this guidance. The previously approved collections of information are subject to review by OMB under the PRA. The collections of information in part 1150 have been approved under 0910–0749.

III. Electronic Access

exception applies. A biological product (defined in section 351(i) of the Public Health Service Act (PHS Act) (42 U.S.C. 262(i)) with an approved license under section 351 of the PHS Act is not required to have an approved application under section 505 of the FD&C Act.

For decades, FDA has interpreted the word “drug” in the term “new drug” to refer to the entire drug product and not just its active ingredient. This interpretation has significant implications for public health. An active ingredient can have different effects on the body depending on the formulation of the drug and its route of administration (e.g., topical vs. intravenous), among other things. That is why when it reviews an application, FDA carefully evaluates, for each drug product, not only the active ingredient but also information about the drug’s formulation, route of administration, labeling, inactive ingredients, bioavailability, and manufacturing processes. In accordance with this approach, FDA has consistently argued in the courts that the term “drug” in “new drug” means the entire drug product and not only an active ingredient, and courts, including the U.S. Supreme Court, have agreed with FDA’s interpretation. See U.S. v. Generix Drug Corp., 460 U.S. 453, 458–59 (1983) (the FD&C Act’s definition of “new drug” applies to the entire drug product rather than the active ingredient); see also U.S. v. Premo Pharmaceutical Lab., 629 F.2d 785 (2d Cir. 1980). FDA regulations incorporate FDA’s interpretation of “new drug” (see 21 CFR 314.200), and a product-specific interpretation of “new drug” underpins FDA’s drug regulatory system.

FDA has long employed a risk-based enforcement approach with respect to new drugs marketed without an approved application. In October 2003, the Agency published a draft guidance, entitled “Marketed Unapproved Drugs—Compliance Policy Guide,” to clarify how FDA generally intended to exercise its enforcement discretion regarding illegally marketed unapproved new drugs (October 23, 2003, 68 FR 60702). In June 2006, FDA finalized the 2003 draft guidance in a final guidance entitled “Marketed Unapproved Drugs—Compliance Policy Guide Sec. 440.100, Marketed New Drugs Without Approved NDAs or ANDAs” (CPG 440.100 guidance) (June 9, 2006, 71 FR 33466). The CPG 440.100 guidance described how FDA intended to prioritize regulatory action under its existing enforcement authority regarding currently marketed unapproved new drugs, including that FDA generally intended to apply a risk-based approach.

In 2011, FDA updated the CPG 440.100 guidance to clarify that unapproved new drugs introduced onto the market after September 19, 2011, were subject to enforcement action at any time without regard to the enforcement priorities set out in CPG 440.100 (September 21, 2011, 76 FR 58398). As described in the updated version of the CPG 440.100 guidance, FDA generally intended to encourage manufacturers of unapproved new drugs to submit applications for their products, while continuing to apply a risk-based approach to removing unapproved new drugs from the market and preserving access to medically necessary drugs.

The CPG 440.100 guidance was part of FDA’s UDI, which focuses on addressing the continued illegal marketing in the United States of drug products that lack the required FDA review and approval for safety and efficacy. To address this problem, FDA’s UDI adopts a risk-based approach for removing from the market unapproved new drugs, particularly those that pose serious risks to patients, with the goal of also preserving patient access to medically necessary drugs and encouraging manufacturers of unapproved new drugs to submit applications for their products. The UDI has a two-pronged approach to help assure patient safety. First, the Agency encourages manufacturers of unapproved new drugs to obtain approval to be legally marketed in the United States. Second, FDA works to remove unapproved new drugs from the...
market consistent with risk-based enforcement priorities and existing enforcement authorities.

As a result of the UDI, FDA has initiated 45 actions since 2006 (some affecting multiple unapproved new drugs) that have led to hundreds of potentially unsafe drugs being voluntarily removed from the market, including several drugs with significant safety concerns. These drugs were removed from the market in response to FDA Federal Register notices announcing that FDA intended to take enforcement action (13 of the actions), warning letters (15 of the actions), or at FDA’s informal request through communications such as a teleconference (17 of the actions). In all 45 actions, safety concerns supported removal of the unapproved new drug products from the market, such as serious adverse events, labeling that did not adequately warn healthcare professionals of risks, or potential risks of harm resulting from adulterated drugs produced by facilities with current good manufacturing practice violations.

The following are well-documented examples of significant adverse events associated with unapproved new drugs that resulted in compliance actions to remove an entire class of unapproved new drugs from the market. As noted below, these compliance actions have also spurred manufacturers to seek and obtain FDA approval of safe and effective versions of these drugs:

- Carbinoxamine-containing products
  - Between 1969 and 2006, FDA received 665 adverse events reports, including 93 deaths, associated with unapproved quinine sulfate use. Among the more common types of events with serious outcomes reported to the Agency were cardiac events, renal failure, and events related to overdose. FDA approved its first quinine sulfate product in August 2005, and the approved labeling for quinine sulfate provides extensive warnings to ensure its safe use. After a safe and effective FDA-approved quinine sulfate product became available, in December 2006, FDA issued a Federal Register notice announcing that it intended to take enforcement action against unapproved drug products containing quinine (including quinine sulfate and other salts of quinine) and persons who cause the manufacture of such products or their shipment in interstate commerce because these products presented serious safety risks that the unapproved drug labeling did not comprehensively describe. As of February 2021, there are five FDA-approved quinine sulfate capsules, including four generic drug products, available in the marketplace. As noted above, these compliance actions have resulted in potentially unsafe unapproved new drugs being removed from the market as well as FDA approval of safe and effective versions of drug products previously marketed without approval. Approval of formerly unapproved new drugs helps reduce concerns about a potential market disruption or shortage of these drugs, because the manufacturers of approved drugs have invested in a manufacturing process that helps to ensure the drug is produced reliably and consistently. This lowers the risk of quality problems, which are one of the main causes of shortages. In addition, the approval of previously unapproved new drugs assures the American public that the approved versions of those drugs are safe and effective for their intended uses, manufactured in accordance with Federal quality standards, and bear accurate and complete labeling regarding risks, benefits, and safe use.

  - Quinine

  2 Between 1983 and 2006, FDA became aware of 21 deaths in children under 2 years of age associated with the use of carbinoxamine-containing drugs, including unapproved drugs. FDA had concerns about the risks associated with these products because, although their safety and effectiveness had not been studied in infants and young children, they were promoted for use in this vulnerable age group. As a result, in June 2006, FDA issued a Federal Register notice announcing that it intended to take enforcement action against unapproved drug products containing carbinoxamine and those who cause the manufacture of such products. As of February 2021, six FDA-approved carboxinamine-containing drug products, including five generic versions, are available on the marketplace and are labeled as contraindicated in children under 2 years old.


Administrative Procedure Act (APA; 5 U.S.C. 551 et seq.) and FDA’s regulations on good guidance practices (§ 10.115 (21 CFR 10.115)). Under the APA, FDA may use guidance documents to “advise the public prospectively of the manner in which the agency proposes to exercise a discretionary power.” Accordingly, FDA’s good guidance practice regulations define “guidance documents” to include “documents that relate to . . . enforcement policies.” (§ 10.115(b)(2)). Additionally, the HHS Notice is supported by flawed facts. It cites, for the proposition that the UDI and CPG 440.100 guidance resulted in price increases for certain new drugs, only a single observational study of 26 products, which included pricing estimates that were not inflation-adjusted over the 4-year observational period, which could lead to an overestimation of real price changes. The HHS Notice also erroneously ties the 2015 price increase for the drug DARAPRIM to the UDI. DARAPRIM was approved as a new drug under the FD&C Act in 1953. Following the 1962 FD&C Act amendments, which required drugs to demonstrate not only safety but efficacy, DARAPRIM was found to be effective, in 1971, as part of FDA’s review of all new drugs that had been approved only for safety before 1962. DARAPRIM was then fully approved by FDA as a safe and effective drug. For years after its approval, DARAPRIM was an off-patent, off-exclusivity drug eligible for generic competition, but no drug manufacturer sought and obtained approval of a generic version during this period. It was during this period, in 2015, that the holder of the approved application for DARAPRIM significantly raised the price of the drug. FDA recently approved a generic version of this product on February 28, 2020.

Due to the HHS Notice’s legal and factual inaccuracies, including those described above, HHS and FDA believe it is appropriate to withdraw the HHS Notice at this time. The HHS Notice does not accurately reflect the Department’s or FDA’s thinking because it is inconsistent with the FD&C Act, FDA regulations, and judicial precedent, among other legal authorities, and is not supported by the facts. In addition, the HHS Notice could result in significant harm to public health by suggesting that unsafe or ineffective drugs could circumvent the drug approval process. Although the withdrawal of FDA’s CPG 440.100 guidance does not change the legal obligations that apply to new drugs, or FDA’s existing enforcement authority over unapproved new drugs, we recognize that the withdrawal of the CPG may have created confusion for the public, including regulated industry, as to how FDA intends to prioritize its enforcement resources in this area. FDA therefore plans to issue guidance on this topic consistent with good guidance practices. The guidance will provide appropriate updates regarding FDA’s enforcement priorities for marketed unapproved new drugs. In the interim, before such guidance is issued, FDA will continue to exercise its existing general approach to prioritizing regulatory and enforcement action, which involves risk-based prioritization in light of all the facts of a given circumstance. Risk-based enforcement best supports FDA’s public health priorities.

FDA’s longstanding interpretation of the statutory terms “new drug,” “grandfathered,” and “GRASÉ” are unchanged and the HHS Notice did not affect the requirements that apply to new drugs under the statutes FDA administers. The HHS Notice did not, and legally could not, provide a new pathway for the legal marketing of unapproved new drugs. Neither HHS nor FDA has the authority to exempt a product or class of products that are new drugs under the FD&C Act from the new drug approval requirements of the FD&C Act. See Cutler v. Kennedy, 475 F. Supp. 838, 856 (D.D.C. 1979); Hoffman-LaRoche v. Weinberger, 425 F. Supp. 890, 892–894 (D.D.C. 1975).

Janet Woodcock,
Acting Commissioner of Food and Drugs.

Xavier Becerra,
Secretary, Department of Health and Human Services.

[FR Doc. 2021–11257 Filed 5–26–21; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2021–N–0335]

Authorizations of Emergency Use of Certain Biological Products During the COVID–19 Pandemic; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the issuance of two Emergency Use Authorizations (EUAs) (the Authorizations) under the Federal Food, Drug, and Cosmetic Act (FD&C Act) for biological products for use during the COVID–19 pandemic. FDA has issued one Authorization for biological products as requested by Eli Lilly and Company and one Authorization for a biological product as requested by Janssen Biotech, Inc. The Authorizations contain, among other things, conditions on the emergency use of the authorized products. The Authorizations follow the February 4, 2020, determination by the Secretary of Health and Human Services (HHS) that there is a public health emergency that has a significant potential to affect national security or the health and security of U.S. citizens living abroad and that involves a novel (new) coronavirus. The virus, now named SARS–CoV–2, causes the illness COVID–19. On the basis of such determination, the Secretary of HHS declared on March 27, 2020, that circumstances exist justifying the authorization of emergency use of drugs and biological products during the COVID–19 pandemic, pursuant to the FD&C Act, subject to the terms of any authorization issued under that section. The Authorizations, which include an explanation of the reasons for issuance, are reprinted in this document.


ADDRESSES: Submit written requests for single copies of the EUAs to the Office of Counterterrorism and Emerging Threats, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 4338, Silver Spring, MD 20993–0002. Send one self-addressed adhesive label to assist that office in processing your request or include a Fax number to which the Authorizations may be sent. See the SUPPLEMENTARY INFORMATION.