Pediatric Drug Development: Regulatory Considerations — Complying With the Pediatric Research Equity Act and Qualifying for Pediatric Exclusivity Under the Best Pharmaceuticals for Children Act

Guidance for Industry

DRAFT GUIDANCE

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For questions regarding this draft document, contact (CDER) Rosemary Addy at 301-796-1640 or (CBER) the Office of Communication, Outreach, and Development at 800-835-4709 or 240-402-8010.

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U.S. Department of Health and Human Services
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Pediatric Drug Development: Regulatory Considerations —
Complying With the Pediatric Research Equity Act and
Qualifying for Pediatric Exclusivity Under the
Best Pharmaceuticals for Children Act
Guidance for Industry¹

This draft guidance, when finalized, will represent the current thinking of the Food and Drug Administration (FDA or Agency) on this topic. 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. To discuss an alternative approach, contact the FDA staff responsible for this guidance as listed on the title page.

I. INTRODUCTION

This guidance is intended to assist industry developing drug products² to comply with the pediatric study requirements under the Pediatric Research Equity Act (PREA),³ and to describe the process for qualifying for pediatric exclusivity and the protections that pediatric exclusivity offers under the Best Pharmaceuticals for Children Act (BPCA).⁴ In 2010, the Biologics Price Competition and Innovation Act of 2009 extended provisions of the BPCA to biological products.⁵

¹ This guidance has been prepared by the Division of Pediatrics and Maternal Health in the Center for Drug Evaluation and Research in cooperation with the Center for Biologics Evaluation and Research at the Food and Drug Administration.

² For the purposes of this guidance, unless otherwise specified, references to drugs or drug products include drugs approved under section 505 of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 355) and biological products licensed under section 351 of the Public Health Service (PHS) Act (42 U.S.C. 262) that are regulated as drugs.

³ Public Law 108-155 (2003), codified at section 505B of the FD&C Act (21 U.S.C. 355c). Although section 505B has been amended since the passage of PREA, by convention, that section of the FD&C Act is often referred to by the acronym for the Act that created it, PREA. We adopt that convention in this guidance.

⁴ Public Law 107-109 (2002), codified at section 505A of the FD&C Act (21 U.S.C. 355a). Although section 505A has been amended since the passage of the BPCA, by convention, that section of the FD&C Act is often referred to by the acronym for the Act that created it, the BPCA. We adopt that convention in this guidance.

⁵ See section 351(m) of the PHS Act (42 U.S.C. 262(m)).
Note that section 505B(b) of the Federal Food, Drug, and Cosmetic (FD&C) Act on already-marketed drugs and section 409I of the Public Health Service (PHS) Act are only briefly addressed in this guidance. Future guidance may address these issues in greater detail. Furthermore, this guidance only briefly addresses the Food and Drug Administration Reauthorization Act of 2017’s (FDARA’s) amendments to section 505B of the FD&C Act relating to requirements that sponsors of certain adult oncology drugs with molecular targets that are determined to be substantially relevant to the growth or progression of a pediatric cancer submit reports on molecularly targeted pediatric cancer investigations.6,7

The scientific aspects of a pediatric program (e.g., considerations regarding data in pediatric subjects, timing of pediatric studies) are addressed in the draft guidance for industry Pediatric Drug Development Under the Pediatric Research Equity Act and the Best Pharmaceuticals for Children Act: Scientific Considerations (May 2023).8

This guidance, along with the draft guidance for industry Pediatric Drug Development Under the Pediatric Research Equity Act and the Best Pharmaceuticals for Children Act: Scientific Considerations,9 revises and replaces the draft guidance for industry How to Comply With the Pediatric Research Equity Act.10 In addition to addressing the PREA topics covered in the earlier draft guidance (i.e., the pediatric assessment, pediatric plan, waivers and deferrals, compliance issues, and pediatric exclusivity provisions), this guidance addresses statutory changes relating to adverse event reporting, pediatric study plans (PSPs), deferral extensions, and noncompliance.

In general, FDA’s guidance documents do not establish legally enforceable responsibilities. Instead, guidances describe the Agency’s current thinking on a topic and should be viewed only as recommendations, unless specific regulatory or statutory requirements are cited. The use of the word should in Agency guidances means that something is suggested or recommended, but not required.

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6 Certain terms used in this guidance, which appear in bold italics at first mention, are defined for purposes of this guidance in the Glossary.

7 For additional information on FDA’s implementation of these amendments to section 505B of the FD&C Act, see the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act (May 2021). We update guidances periodically. To make sure you have the most recent version of a guidance, check the FDA guidance web page at https://www.fda.gov/regulatory-information/search-fda-guidance-documents.

8 When final, this guidance will represent the FDA’s current thinking on this topic. For the most recent version of a guidance, check the FDA guidance web page at https://www.fda.gov/regulatory-information/search-fda-guidance-documents.

9 When final, this guidance will represent the FDA’s current thinking on this topic.

10 This guidance also addresses certain topics previously addressed in the guidance for industry Qualifying for Pediatric Exclusivity Under Section 505A of the Federal Food, Drug, and Cosmetic Act. That guidance was withdrawn August 7, 2013 (78 FR 48175).
II. OVERVIEW OF REGULATORY STRATEGY FOR PEDIATRIC DRUG DEVELOPMENT

A. General Approach

For purposes of pediatric drug development, FDA generally considers the pediatric population to include those patients from birth to younger than 17 years (i.e., birth through 16 years of age), and to include the subpopulation age groups of neonates, infants, children, and adolescents.\textsuperscript{11} Consistent with International Council for Harmonisation (ICH) guidelines,\textsuperscript{12} FDA considers these subpopulation age groups to be divided as follows:

- Neonates: birth through 27 days (corrected gestational age)
- Infants: 28 days to 23 months
- Children: 2 years to 11 years
- Adolescents: 12 years to younger than 17 years

The BPCA defines \textit{pediatric studies} to mean at least one clinical investigation in “pediatric age groups (including neonates in appropriate cases) in which a drug is anticipated to be used, and, at the discretion of the Secretary, may include preclinical studies.”\textsuperscript{13} For purposes of satisfying the requirements of PREA, assessments of safety and effectiveness must be performed in all relevant pediatric age groups, unless the assessments are waived or deferred.\textsuperscript{14}

The BPCA and PREA are designed to work together to encourage the development of data to inform the safe and effective use of drugs in pediatric populations. Under PREA, pediatric assessments are \textit{required} for drug products with a new active ingredient, new indication, new dosage form, new dosing regimen, or new route of administration unless the drug is for an indication for which orphan designation has been granted.\textsuperscript{15} Also under PREA, molecularly targeted pediatric cancer investigations are \textit{required} for original new drug applications (NDAs) or biologics license applications (BLAs) submitted on or after August 18, 2020, for a new active ingredient, if the drug that is the subject of the application is intended for the treatment of an

\textsuperscript{11} See 21 CFR 201.57(c)(9)(iv)(A) (“the terms \textit{pediatric population(s)} and \textit{pediatric patient(s)} are defined as the pediatric age group, from birth to 16 years, including age groups often called neonates, infants, children, and adolescents”). FDA interprets “birth to 16 years” in 21 CFR 201.57(c)(9)(iv)(A) to mean from birth to younger than 17 years old. See, for example, the guidance for industry \textit{Pediatric Information Incorporated Into Human Prescription Drug and Biological Product Labeling} (March 2019).

\textsuperscript{12} For additional information, see the ICH guidances for industry \textit{E11 Clinical Investigation of Medicinal Products in the Pediatric Population} (December 2000) and \textit{E11(R1) Addendum: Clinical Investigation of Medicinal Products in the Pediatric Population} (April 2018).

\textsuperscript{13} See section 505A(a) of the FD&C Act (21 U.S.C. 355(a)).


\textsuperscript{15} See sections 505B(a)(1)(A) and 505B(k)(1) of the FD&C Act (21 U.S.C. 355c(a)(1)(A) and 21 U.S.C. 355c(k)(1)).
adult cancer, and directed at a molecular target that FDA determines to be substantially relevant to the growth or progression of a pediatric cancer. Studies conducted under the BPCA, on the other hand, are optional because sponsors have the option of declining to undertake them in response to a written request (WR). Nevertheless, it is critical that sponsors consider both laws when planning their pediatric clinical development programs.

PREA requires that any application falling within the requirements of section 505B(a)(1) of the FD&C Act for a new active ingredient, new indication, new dosage form, new dosing regimen, or new route of administration must either include pediatric assessments or reports on the molecularly targeted pediatric cancer investigation (as appropriate); or a request for waiver and/or deferral of the pediatric assessments or reports on the molecularly targeted pediatric cancer investigation. There are certain exceptions; for example, PREA requirements generally do not apply to a drug for an indication for which orphan designation has been granted (see section II.B.4., Orphan Products).

The FD&C Act requires sponsors to submit an initial pediatric study plan (iPSP) during the investigational phase of development, which helps to ensure that sponsors thoroughly consider a pediatric clinical development program earlier in their overall clinical development program. See sections III., Pediatric Research Equity Act, and V., Elements Common to PREA and the BPCA, for additional information about PREA.

While addressing PREA, sponsors should also consider whether to seek a WR under the BPCA. It is important to note that sponsors may qualify for pediatric exclusivity under the BPCA for completed PREA studies when those studies are described in a WR, and the WR is issued by FDA before the sponsor submits any reports about the studies described in the WR. If FDA determines that “information relating to the use of a new drug in the pediatric population may produce health benefits in that population” and issues a WR, a sponsor may qualify for 6 months of exclusivity under the BPCA for conducting studies that are required under PREA. However, as discussed in Section IV. A. 2., Written Request Studies, FDA does not expect to issue WRs solely for studies or planned studies that are required under PREA.

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16 Section 505B(a)(1)(B) of the FD&C Act (21 U.S.C. 355c(a)(1)(B)).

17 For additional information, see the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act.


19 See section 505B(k)(1) of the FD&C Act (21 U.S.C. 355c(k)(1)). Under section 505B(k)(2) of the FD&C Act, this orphan exemption does not apply to products that trigger PREA under section 505B(a)(1)(B) of the FD&C Act.

20 See section 505B(e) of the FD&C Act (21 U.S.C. 355c(e)).

21 See section 505A(h) of the FD&C Act (21 U.S.C. 355a(h)).

22 See section 505A(b)(1) of the FD&C Act (21 U.S.C. 355a(b)(1)).

23 See section 505A(b)(1) of the FD&C Act (21 U.S.C. 355a(b)(1)).
In addition, sponsors should consider other indications for development in the pediatric population based on the current understanding of the effect of the drug (e.g., mechanism of action), and include them as appropriate when seeking a WR. FDA reviews adult safety and effectiveness data and, in some instances, information from postmarketing safety reports, information regarding unapproved uses, and the scientific literature to identify a potential health benefit in pediatric populations when contemplating a WR. This knowledge not only informs FDA’s decision to issue a WR for a given indication, but also whether to include a request to study additional indications in children beyond those already approved in adults. See sections IV., Best Pharmaceuticals for Children Act, and V., Elements Common to PREA and the BPCA, for more information about the BPCA.

B. Developing Drugs for Pediatric Use

During clinical development programs, a sponsor should consider whether the eventual marketing application will trigger the requirements of PREA. FDA encourages sponsors to interact with FDA early, including, when applicable, to discuss studies to meet PREA and other requirements. See section III., Pediatric Research Equity Act, for information about the requirements of PREA.

FDA has identified a number of scenarios and considerations that may affect a sponsor’s approach to developing a drug product for use in pediatric patients. Some common ones are described below.

1. Drugs for Life-Threatening or Severely Debilitating Conditions and Unmet Medical Needs

Early consultation and discussions are particularly important for drugs intended for life-threatening or severely debilitating conditions. For these drugs, FDA encourages sponsors to discuss the pediatric plan at pre-investigational new drug application (pre-IND) and end-of-phase 1 meetings. In some cases, pediatric studies of drugs for life-threatening or severely debilitating conditions that lack adequate therapies might begin earlier than usual in the drug development process. The need for new therapies might justify early trials despite the relative lack of safety and effectiveness information in humans. Pediatric studies might be considered appropriate when prospects of direct benefit to the enrolled children are sufficient to justify the risks.

24 See the guidance for industry Expedited Programs for Serious Conditions — Drugs and Biologics (May 2014) for more information.


26 See 21 CFR 50.52.
2. Drugs for Diseases or Conditions That Occur Primarily in Pediatric Populations

Sponsors developing drug products for diseases or conditions that primarily or substantially occur in the pediatric population should discuss their plans with FDA as early as possible (e.g., pre-IND meeting). They should consider submitting an iPSP earlier than is required under PREA because the initial clinical studies will likely include pediatric subjects (see section III.B.1., Pediatric Study Plans).

3. Neonates

The complex medical state of neonates makes it critical to evaluate drugs specifically for their use. However, FDA is aware that studies in neonates present special challenges, including the short time period to conduct studies during the neonatal period (e.g., birth through 27 days corrected gestational age), the differences in neonatal physiology that may affect dose and endpoint selection, as well as ethical issues that may be age-specific. Under PREA, studies in neonates may be required. However, it is possible that partial waivers for this and other specific age groups might be appropriate under certain circumstances.27

FDA encourages specific activities regarding neonates. Under PREA, if a sponsor does not plan to study an investigational drug in neonates, the rationale and supporting data explaining why the drug is not appropriate for use in this population should be included in the iPSP.28 Under the BPCA, similar rationale and supporting data explaining why the investigational drug is not appropriate for use in this population should be included in the proposed pediatric study request (PPSR) if a sponsor does not plan to study the drug in neonates. When FDA issues a WR that does not include studies in neonates, the WR must state the rationale for not including neonates.29 See sections IV.A.1., Description of the Written Request, and IV.B., How to Obtain a Written Request, as well as the draft guidance for industry Pediatric Drug Development Under the Pediatric Research Equity Act and the Best Pharmaceuticals for Children Act: Scientific Considerations30 for more information regarding clinical studies in neonates.

4. Orphan Products

PREA requirements generally do not apply “to any drug or biological product for an indication for which orphan designation has been granted;” however, this orphan exemption does not apply to drugs that trigger the PREA requirement for submission of reports on the molecularly targeted

27 See section 505B(a)(5)(B) of the FD&C Act (21 U.S.C. 355(a)(5)(B)).


30 When final, this guidance will represent FDA’s current thinking on this topic.
pediatric cancer investigation. Thus, PREA does not require submission of pediatric assessments for an application (or supplemental application) to market a drug for an indication for which orphan designation has been granted. As FDA has interpreted PREA, if orphan designation is granted after approval of a drug, and postmarketing studies were required under PREA at the time of the drug’s approval, the granting of orphan designation does not alter the already existing requirement for such studies. The PREA orphan exemption is not revisited to retroactively abrogate a PREA requirement that was properly imposed before orphan designation was granted. Additionally, if marketing approval is sought for multiple indications for a drug product, some of which have not been granted orphan designation, the sponsor must submit pediatric assessments for all indications that do not have an orphan designation, unless the assessments are waived or deferred. If the orphan-designated indication(s) involve the pediatric population, we encourage sponsors to conduct studies in the relevant age group(s) whenever appropriate.

Despite this orphan exemption under PREA, a sponsor that submits an application to market a drug for an indication for which orphan designation has been granted may be eligible to qualify for pediatric exclusivity if FDA issues a WR to the sponsor in connection with the application and the sponsor accepts. Sponsors should contact FDA about the feasibility and timing of a WR and about submitting a PPSR, if appropriate. For more information, see sections III.C.2., Submission of Pediatric Assessments or Reports on the Molecularly Targeted Pediatric Cancer Investigation, IV.F.2., Biological Products, and V.B.1., Pediatric Exclusivity Determinations, as well as FDA’s Office of Orphan Products Development web page on developing drug products for rare diseases and conditions, and the guidance for industry Clarification of Orphan Designation of Drugs and Biologics for Pediatric Subpopulations of Common Diseases (July 2018).

31 See sections 505B(a)(1)(B), 505B(a)(3), and 505B(k) of the FD&C Act (21 U.S.C. 355c(a)(1)(B), 21 U.S.C. 355c(a)(3), and 21 U.S.C. 355c(k)). Note that, although section 505B(k) authorizes FDA to issue regulations that would alter the orphan exemption, as of the date of publication of this guidance, FDA has not issued any such regulations.

32 See section 505B(k)(1) of the FD&C Act (21 U.S.C. 355c(k)(1)). For products meeting the criteria in section 505B(a)(1)(B) of the FD&C Act, the requirement to submit reports on the investigation described in section 505B(a)(3) of the FD&C Act applies even if the drug is for an adult indication for which orphan designation has been granted. See section 505B(k)(2) of the FD&C Act (21 U.S.C. 355c(k)(2)). See also the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act.

33 Because the orphan exemption in section 505B(k)(1) of the FD&C Act can affect whether PREA applies to a particular application or supplement submitted to the Agency for review, FDA evaluates whether the orphan exemption is applicable at the time it evaluates whether PREA would otherwise be triggered for any application or supplement under section 505B(a)(1)(A).


35 See the Medical Products for Rare Diseases and Conditions web page at https://www.fda.gov/ForIndustry/DevelopingProductsforRareDiseasesConditions/default.htm.
5. Drug Development in Foreign Countries

Many sponsors conduct their entire clinical programs in other countries and occasionally submit a marketing application with little, if any, prior interaction with FDA. All sponsors who seek to market their drugs in the United States are strongly encouraged to contact FDA as early as possible to avoid any delay in providing any required pediatric information in their applications. Sponsors should include an agreed iPSP in any NDA, BLA, or supplement that is required by PREA to include pediatric assessments or reports on the molecularly targeted pediatric cancer investigation. PREA requirements are described in more detail in section III., Pediatric Research Equity Act.

6. Drugs for Diseases or Conditions That Only Occur in Adults

Sponsors focusing on clinical development programs for drugs for diseases and conditions that only occur in adults must submit an iPSP, assuming the application triggers PREA. A list of diseases and conditions that rarely or never occur in pediatrics can be found on FDA’s website. Generally, applications for drugs for such diseases or conditions that rarely or never occur in pediatrics will qualify for a waiver because the necessary studies would be impossible or highly impracticable. However, sponsors should consider all potential pediatric indications for their drugs. FDA may consider issuance of a WR for other indications that may have health benefits in the pediatric population.

III. PEDIATRIC RESEARCH EQUITY ACT

A. Overview — Requirements of PREA

1. PREA Applicability

With limited exception (for example, the orphan exemption described in section II.B.4., Orphan Products), PREA applies to the following:

- Applications (or supplements to an application) submitted under section 505 of the FD&C Act or section 351 of the PHS Act for a new active ingredient, new indication, new dosage form, new dosing regimen, or new route of administration, and

36 See section 505B(e)(1) of the FD&C Act (21 U.S.C. 355c(e)(1)); see also section 505B(a)(1) of the FD&C Act (21 U.S.C. 355c(a)(1)).

37 See Adult-Related Conditions That Qualify for a Waiver Because They Rarely or Never Occur in Pediatrics, available at https://www.fda.gov/media/101440/download.

38 See section 505B(a)(5) of the FD&C Act (21 U.S.C. 355c(a)(5)). FDA anticipates that there will be additional considerations for applications described in section 505B(a)(1)(B) of the FD&C Act that require submission of reports on the molecularly targeted pediatric cancer investigation described in section 505B(a)(3) of the FD&C Act. For additional information, see the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act.
Contains Nonbinding Recommendations
Draft — Not for Implementation

Original applications for a new active ingredient submitted under section 505 of the FD&C Act or section 351 of the PHS Act on or after August 18, 2020, if the drug that is the subject of the application is intended for the treatment of an adult cancer and is directed at a molecular target that FDA determines to be substantially relevant to the growth or progression of a pediatric cancer.39

PREA also authorizes FDA to require holders of already approved applications to conduct pediatric assessments under certain circumstances.40

2. Scope of Requirements — Generic drugs

Abbreviated new drug applications (ANDAs) submitted under section 505(j) of the FD&C Act most often are applications for drugs containing the same active ingredient(s), strength(s), indication(s), dosage form(s), dosing regimen(s), and route(s) of administration as the listed drugs they reference and are not subject to PREA. ANDA applicants may petition the Agency to request a change from a listed drug per section 505(j)(2)(C) of the FD&C Act and 21 CFR 314.93, a process referred to as a suitability petition. The regulation at 314.93 limits the types of changes that may be permitted to changes in strength, dosage form, route of administration, or of a single active ingredient in a combination drug subject to the restrictions identified in 314.93(d)(1) through (3). ANDAs submitted pursuant to an approved suitability petition for changes in dosage form, route of administration, or for a change in active ingredient in a combination drug do trigger PREA, but they are only eligible for submission as ANDAs if the pediatric assessment or molecularly targeted pediatric cancer investigation requirements are waived.41 If a change proposed in a suitability petition triggers PREA and FDA does not waive the requirement to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation, the suitability petition will be denied, and the proposed drug product will not be eligible for submission as an ANDA.42

B. Development of Drugs for Pediatric Use

1. Pediatric Study Plans

A sponsor planning to submit a marketing application or supplement that is subject to PREA is required to submit an iPSP before submission of pediatric assessments or reports on the molecularly targeted pediatric cancer investigation.43 A sponsor should submit an iPSP to its investigational new drug application (IND) for review by the appropriate review division as early

39 See section 505B(a)(1) of the FD&C Act (21 U.S.C. 355c(a)(1)).

40 See section 505B(b) of the FD&C Act (21 U.S.C. 355c(b)).


42 See, for example, 21 CFR 314.93.

43 See section 505B(e)(1) of the FD&C Act (21 U.S.C. 355c(e)(1)).
as practicable and must submit it no later than 60 calendar days after the date of the end-of-phase 2 meeting or such other time as agreed upon between FDA and the applicant. More information on the timing and contents of iPSPs, the process for reaching agreement with FDA on iPSPs, and the process for amending an agreed iPSP can be found in the guidance for industry Pediatric Study Plans: Content of and Process for Submitting Initial Pediatric Study Plans and Amended Initial Pediatric Study Plans (July 2020). Sponsors are also encouraged to consider the possibility of submitting a PPSR to obtain a WR during the PSP process. An iPSP and a PPSR are different documents and have different considerations for submission; the former is a requirement for compliance with PREA and the latter may be submitted at a sponsor’s discretion to seek a WR under the BPCA. See sections IV.A., Written Requests, and IV.B., How to Obtain a Written Request, for further discussion about PPSRs and WRs.

2. Developing a Pediatric Formulation

Under PREA, sponsors are required to conduct pediatric studies “using appropriate formulations for each age group” for which the assessment or investigation is required. However, FDA may grant a partial waiver if a sponsor is unable to develop an age-appropriate formulation after reasonable attempts to do so. (See discussion of waivers in section III.D., Waivers and Deferrals Under PREA.) Under PREA, sponsors must submit “a request for approval of a pediatric formulation” used in their pediatric studies, and if a sponsor fails to submit such a request, the drug may be considered misbranded. Accordingly, sponsors should submit an application or supplemental application for any formulation(s) not previously approved that were used during pediatric studies and for which the sponsor has data to assess the safety and effectiveness and to support dosing and administration. To avoid delays in initiation of pediatric clinical studies, sponsors should begin the development of an age-appropriate formulation as early as possible.

C. Pediatric Assessments and Molecularly Targeted Pediatric Cancer Investigations Under PREA

1. Definitions

Pediatric assessments must contain data, gathered using appropriate formulations for each age group for which the assessment is required and that are adequate to:

- Assess the safety and effectiveness of the drug for the claimed indications in all relevant pediatric subpopulations; and

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45 See section 505B(a)(2)(A) and 505B(a)(3)(A) of the FD&C Act (21 U.S.C. 355c(a)(2)(A) and 355c(a)(3)(A)).


47 See section 505B(d) of the FD&C Act (21 U.S.C. 355c(d)).
• Support dosing and administration for each pediatric subpopulation for which the drug is
safe and effective.48

A molecularly targeted pediatric cancer investigation “shall be designed to yield clinically
meaningful pediatric study data, gathered using appropriate formulations for each age group for
which the study is required, regarding dosing, safety, and preliminary efficacy to inform
potential pediatric labeling.”49

2. Submission of Pediatric Assessments or Reports on the Molecularly Targeted
Pediatric Cancer Investigation

Sponsors must submit pediatric assessments or reports on the molecularly targeted pediatric
cancer investigation with any application for which such assessments or reports are required by
PREA, unless FDA defers and/or waives the requirement.50 (For pediatric assessments, if the
drug is for an indication for which orphan designation has been granted, the requirements of
PREA do not apply.)51 See section III.D., Waivers and Deferrals Under PREA, for discussion of
waivers and deferrals. In general, sponsors should include pediatric studies at the time of
submission of an application when there is sufficient knowledge to proceed and it is feasible to
complete studies in children in parallel with adult studies.

Information about the results of pediatric assessments under PREA must be included in product
labeling whether findings are positive, negative, or inconclusive.52 Labeling changes for
approved products must be submitted in accordance with applicable requirements in 21 CFR
601.12 and 21 CFR 314.70. For more information about labeling, see section V.D.,
Considerations for Labeling of Drug Products, and the guidance for industry Pediatric
Information Incorporated Into Human Prescription Drug and Biological Product Labeling
(March 2019).

For information about the specific types of data that may be needed to complete a pediatric
assessment, refer to the draft guidance for industry Pediatric Drug Development Under the
Pediatric Research Equity Act and the Best Pharmaceuticals for Children Act: Scientific
Considerations.53


355c(a)(4), and 21 U.S.C. 355c(a)(5)).

51 See section 505B(k)(1) of the FD&C Act (21 U.S.C. 355c(k)(1)).

52 See section 505B(g)(2) of the FD&C Act (21 U.S.C. 355c(g)(2)).

53 When final, this guidance will represent FDA’s current thinking on this topic.
D. Waivers and Deferrals Under PREA

1. Waivers

PREA authorizes FDA to waive the requirement to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation, based on established criteria, for some or all pediatric age groups.\(^{54}\) FDA can grant a full or partial waiver of the requirements on its own initiative or at the request of an applicant.\(^{55}\) Any applicant requesting a waiver should provide written justification for the waiver and evidence to support the request.

a. Criteria for full waiver

FDA will, as appropriate, grant a full waiver of the requirement to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation if the applicant certifies and FDA finds one or more of the following criteria:

- Necessary studies are impossible or highly impracticable (because, for example, the number of patients is so small or the patients are geographically dispersed).\(^{56}\) For further information, see Section II.B.6., Drugs for Diseases or Conditions That Only Occur in Adults.

- There is evidence strongly suggesting that the drug would be ineffective or unsafe in all pediatric age groups.\(^{57}\)

- The drug (1) does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients; and (2) is not likely to be used in a substantial number of pediatric patients.\(^{58}\)

    Importantly, we note that both criteria must be met for this waiver justification to apply. A drug is considered to represent a meaningful therapeutic benefit over existing therapies if FDA determines that (1) “if approved, the drug or biological product could represent an improvement in the treatment, diagnosis, or prevention of a disease, compared with marketed products adequately labeled for that use in the relevant pediatric population;” or (2) “the drug or biological product is in a class of products or for an indication for which there is a need for additional options.”\(^{59}\) FDA anticipates that improvement over marketed drugs might be demonstrated by

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\(^{54}\) See section 505B(a)(5) of the FD&C Act (21 U.S.C. 355c(a)(5)).

\(^{55}\) See section 505B(a)(5) of the FD&C Act (21 U.S.C. 355c(a)(5)).


\(^{58}\) See section 505B(a)(5)(A)(iii) of the FD&C Act (21 U.S.C. 355c(a)(5)(A)(iii)).

\(^{59}\) See section 505B(c) of the FD&C Act (21 U.S.C. 355c(c)).
showing, for example (1) evidence of increased effectiveness in treatment, prevention, or diagnosis of disease; (2) an improved safety profile; (3) enhancement of compliance (e.g., by virtue of less frequent dosing or mode of delivery); or (4) safety and effectiveness in a new subpopulation for which marketed drugs are not currently labeled.

da. Criteria for partial waiver

FDA will, as appropriate, grant a partial waiver of the requirement to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation with respect to a specific pediatric age group, if the applicant certifies and FDA finds one or more of the following criteria:

- Necessary studies are impossible or highly impracticable (because, for example, the number of patients in that age group is so small or patients in that age group are geographically dispersed);\(^{60}\)
- There is evidence strongly suggesting that the drug would be ineffective or unsafe in that age group;\(^{61}\)
- The drug (1) does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients in that age group; and (2) is not likely to be used in a substantial number of pediatric patients in that age group;\(^{62}\)
- The applicant can demonstrate that reasonable attempts to produce a pediatric formulation necessary for that age group have failed.\(^{63}\)

We note that if a partial waiver is granted on the basis that it is not possible to develop a pediatric formulation, the waiver will cover only the pediatric age groups requiring that formulation.\(^{64}\)

FDA believes that a partial waiver granted based on the inability to develop a pediatric formulation generally should apply to situations in which the applicant can demonstrate that unusually difficult technological problems prevented it from developing a pediatric formulation. In certain cases, FDA may seek appropriate external expert opinion (e.g., from an advisory committee) to help assess whether to grant such a waiver.

\(^{60}\) See section 505B(a)(5)(B)(i) of the FD&C Act (21 U.S.C. 355c(a)(5)(B)(i)).


\(^{64}\) See section 505B(a)(5)(C) of the FD&C Act (21 U.S.C. 355c(a)(5)(C)).
If the sponsor seeks a partial waiver on the grounds that it is not possible to develop a pediatric formulation, the sponsor must submit documentation detailing why a pediatric formulation cannot be developed. This should include a detailed description of the sponsor’s efforts to develop a pediatric formulation. If FDA grants such a waiver, the sponsor’s submission will be made publicly available.

c. Information for requesting a waiver

To request a waiver, sponsors should provide the following:

- The drug name, applicant name, and indication.
- The age group(s) included in the waiver request.
- The statutory reason(s) for requesting a waiver, including reference to the applicable statutory authority.
- Evidence that the request meets the statutory reason(s) for waiver. All relevant scientific/clinical justifications for the waiver request should be included.

d. Waiver decision

FDA grants a waiver at the time of approval of an application that triggers PREA if it determines that the application satisfies the statutory requirements for a waiver. FDA generally includes a preliminary evaluation of the sponsor’s plan to request a waiver in FDA’s comments on the iPSP (see section III.B.1., Pediatric Study Plans). This evaluation reflects FDA’s best judgment at that time.

2. Deferrals

PREA authorizes FDA to defer the requirement to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation, based on established criteria. A deferral acknowledges that pediatric assessments or reports on the molecularly targeted pediatric cancer investigation are required, but permits the applicant to submit the assessments or reports after the

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65 See section 505B(a)(5)(C) of the FD&C Act (21 U.S.C. 355c(a)(5)(C)).

66 A partial waiver on this basis hinges on whether “the applicant can demonstrate” the failure of reasonable attempts to produce a pediatric formulation (section 505B(a)(5)(B)(iv) (21 U.S.C. 355c(a)(5)(B)(iv))).

67 See section 505B(a)(5)(C) of the FD&C Act (21 U.S.C. 355c(a)(5)(C)).


69 See, for example, section 505B(e)(2)(B)(ii) of the FD&C Act (21 U.S.C. 355c(e)(2)(B)(ii)).

70 See section 505B(a)(4) of the FD&C Act (21 U.S.C. 355c(a)(4)).
approval of an NDA, BLA, or supplement. FDA may, on its own initiative or at the request of an applicant, defer the submission of some or all of the pediatric assessments or reports on the molecularly targeted pediatric cancer investigation until a specified date after approval of the drug.71

a. Timeline

Sponsors can discuss a plan for a deferral and the status of a deferred study with FDA as follows:

- Premarketing — It is important to include in the iPSP any plans for a deferral request. In certain cases it may be appropriate to initiate early discussion of a plan for deferral, for example, as part of a pre-IND meeting or during phase 1 of clinical development.

- Application review — FDA grants the deferral, as appropriate, upon approval of the application or supplement.

- Postmarketing — The applicant must submit an annual review of the status of a deferred pediatric study (PREA postmarketing requirement) to FDA until it has submitted the final study report.72 The final due date of a deferred pediatric study may be extended under certain circumstances (see section III.D.2.e., Deferral extensions).

b. Criteria for deferral

FDA may defer the timing of submission of some or all required assessments or reports on the molecularly targeted pediatric cancer investigation if the applicant submits certain required information to FDA, as discussed below, and FDA finds one or more of the following:73

- The drug is ready for approval for use in adults before pediatric studies are complete;

- Pediatric studies should be delayed until additional safety or effectiveness data have been collected; or

- There is another appropriate reason for deferral

An “appropriate reason” for deferral may include, for example, that development of a pediatric formulation is not complete.

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71 See section 505B(a)(4) of the FD&C Act (21 U.S.C. 355c(a)(3)).

72 See section 505B(a)(4)(C) of the FD&C Act (21 U.S.C. 355c(a)(4)(C)).

c. Information for requesting a deferral

To request a deferral, an applicant must provide the following:\(^{74}\)

- A certification of the grounds for deferral;
- A pediatric study plan as described in section 505B(e) of the FD&C Act (see section III.B.1., Pediatric Study Plans);
- Evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time; and
- A timeline for completion of such studies.

d. Deferral review and decision

The decision to defer and the deferral date are determined on a case-by-case basis. FDA may, as appropriate, consider the following in determining whether and how long to defer submission of a pediatric assessment:

- The need for the drug in pediatric patients;
- Availability of sufficient safety data to initiate pediatric clinical studies;
- The nature and extent of pediatric data needed to support pediatric labeling;
- The existence of clearly documented difficulties in enrolling subjects; and/or
- Evidence of technical problems in developing pediatric formulations.

For additional information on the circumstances in which a deferral may be appropriate for a molecularly targeted pediatric cancer investigation, see the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act.

The iPSP, agreed iPSP, and any subsequent amendments should include the key elements of any planned deferred studies. FDA does not intend to make recommendations on planned deferral requests that are submitted in the absence of an iPSP, except under rare circumstances (e.g., urgent public health need).

FDA grants a deferral, as appropriate, in the approval letter for an NDA, BLA, or supplement.

e. Deferral extensions

The FD&C Act provides a mechanism for FDA to grant an extension of the timeline for a deferral granted by FDA.\(^{75}\) Examples of reasons assessments or investigations may be delayed


\(^{75}\) See section 505B(a)(4)(B) of the FD&C Act (21 U.S.C. 355c(a)(4)(B)).
include, but are not limited to, unexpected difficulties with enrollment, unexpected delays in reaching agreement with FDA on protocols for the pediatric clinical studies, or an unanticipated need for additional safety or effectiveness data before proceeding with studies in children. During consideration of the deferral extension request, in general, FDA considers whether the applicant could have prevented or foreseen the delay. FDA also generally considers the likelihood that studies can be completed given the circumstances.

To request a deferral extension, an applicant must submit a new timeline for the completion of pediatric studies along with any significant updates to the following information from the original deferral request:

- A certification of the grounds for deferral;
- PSP; and
- Evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time.

Applicants should submit information to support the need for an extension of the timeline for deferred studies. For example, if an applicant needs additional time to complete the deferred pediatric studies because of difficulty recruiting subjects, the applicant should provide information outlining the reasons for the difficulties, evidence supporting the reasons outlined (including information on the incidence of the condition and global geographic distribution, if applicable), and information outlining its efforts to increase enrollment such as the number of clinical sites contacted and the number of subjects screened and enrolled.

An applicant must submit a request for deferral extension, along with the required information, at least 90 days before the date that the studies are due. FDA will respond to such request within 45 days of receipt of the request. If FDA grants the deferral extension, the specified date will be the new due date for submission of the deferred assessments or deferred reports on the molecularly targeted pediatric cancer investigation.

3. Annual Review

Pediatric assessments deferred under PREA are required postmarketing studies subject to annual status reporting requirements under PREA and FDA regulations.

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An applicant’s annual status report under PREA must contain the following:\(^1\)

- Information detailing the progress made in conducting pediatric studies
- If no progress has been made in conducting such studies, evidence and documentation that such studies will be conducted with due diligence and at the earliest possible time
- The projected completion date for pediatric studies
- The reason(s) that a deferral or deferral extension continues to be necessary

### E. Compliance With PREA

If an applicant submits an application or supplement subject to PREA and fails to comply with applicable PREA requirements, FDA may, as appropriate, refuse to file the application or issue a complete response letter after reviewing the application.\(^2\) If an applicant fails to fulfill required deferred pediatric studies under PREA that were established in the approval letter, FDA will issue to the applicant a noncompliance letter that informs it of such failure.\(^3\) The applicant must respond to this letter in writing within 45 days and may request a deferral extension as part of that response.\(^4\)

FDA will post the noncompliance letter and the applicant’s response on the FDA public website after redacting any information protected by applicable law.\(^5\) If FDA grants a deferral extension before the initial study due date, FDA does not intend to issue a noncompliance letter unless and until the newly established due date has passed.

After FDA issues a noncompliance letter, it may take additional steps to ensure compliance if needed. The drug may be considered misbranded solely because of the applicant’s failure to comply with PREA and subject to relevant enforcement action.\(^6\) For an approved drug, the failure to submit pediatric assessments or reports on the molecularly targeted pediatric cancer investigation, or a request for waiver or deferral of those studies, will not be the basis for

\(^1\) See section 505B(a)(4)(C)(i) of the FD&C Act (21 U.S.C. 355c(a)(4)(C)(i)).


\(^3\) See section 505B(d)(1) of the FD&C Act (21 U.S.C. 355c(d)(1)).

\(^4\) See section 505B(d)(1) of the FD&C Act (21 U.S.C. 355c(d)(1)).


\(^6\) See section 505B(d)(2) of the FD&C Act (21 U.S.C. 355c(d)(2)).
withdrawing approval of a drug or revoking a license for a biological product. However, if FDA finds a drug to be misbranded, the drug could be subject to an injunction or seizure proceedings.

IV. BEST PHARMACEUTICALS FOR CHILDREN ACT

A. Written Requests

1. Description of the Written Request

A WR is a document issued by FDA requesting submission of a study or studies intended to provide meaningful health benefits in the pediatric population. The WR specifies the elements of the study or studies that the sponsor or application holder must complete to qualify for pediatric exclusivity. FDA can issue a WR at the request of an interested party or on its own initiative. Completion of studies described in a WR is voluntary. FDA does not limit issuance of a WR to a specific drug product, and the WR can result in only one 6-month period of pediatric exclusivity for that sponsor, as described in section IV.F., Attaching the Period of Pediatric Exclusivity After a Determination That a Drug Qualifies for Pediatric Exclusivity. FDA’s authority to issue a WR extends to use of an active moiety for indications that may produce health benefits in the pediatric population, regardless of whether the indications have been previously approved in adults.

Generally, a WR seeks all applicable information necessary to establish safety and effectiveness of a drug for use in all relevant pediatric populations, including study information (e.g., type, timing, endpoints), drug-specific safety concerns to be monitored, statistical analysis plan, and timeline for completing the studies.

FDA can use a PPSR to develop a WR or use alternative information (see section IV.B., How to Obtain a Written Request). As a greater understanding of the indication or of the mechanism of action of a particular drug or drug class develops, WRs, including elements within a study or studies necessary to qualify for pediatric exclusivity, may evolve.

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87 See section 505B(d)(2) of the FD&C Act (21 U.S.C. 355c(d)(2)).


89 See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).

90 Section 505A of the FD&C Act does not require the sponsor or application holder to conduct pediatric studies; instead, it creates an exclusivity incentive to encourage such studies. However, the sponsor or application holder may be required to conduct pediatric studies of certain new and marketed drugs under section 505B of the FD&C Act.

In addition, the BPCA requires that, in issuing a WR, FDA take into account adequate representation of children of ethnic and racial minorities.\(^\text{92}\) If a WR does not request studies in neonates, the request will include a statement describing the rationale for not requesting such studies.\(^\text{93}\)

A sponsor will not be eligible for pediatric exclusivity based on requirements or requests to conduct postmarketing studies (e.g., studies required under PREA) or other communications about pediatric studies unless it is in receipt of a WR.\(^\text{94}\)

### 2. Written Request Studies

In general, FDA decides what studies to include in a WR by determining what information is needed to use the drug appropriately in the pediatric population. When making this determination, in general, FDA obtains the following from the sponsor:

- Information on any other indications for this product that may have health benefits in children. For example, the sponsor should provide information on any indications for which there are ongoing clinical studies in adults and/or children or for which the sponsor has opened an IND.

- Information that exists in the literature on the drug or on pharmacologically related drugs.

In some instances, FDA may ask a sponsor to submit information to an IND before issuing a WR. Similarly, in some cases, a sponsor may wish to submit pediatric study data to its IND in support of an amendment to its WR. Pediatric studies previously submitted to an IND can be used as the basis of a PPSR or can be submitted to an NDA or BLA to qualify for pediatric exclusivity in response to a WR; however, FDA does not consider pediatric studies a sponsor submits to an NDA or BLA (either in an original application, amendment, or supplement) before FDA issues a WR as being responsive to that WR.

In certain situations, FDA may determine that a WR for additional pediatric studies will not be issued. Such situations may include the following:

- Sufficient pediatric information has already been submitted to the NDA or BLA, even if the pediatric information is not yet included in the labeling;

- Study of the drug for the specified indication(s) in the pediatric population would not offer a health benefit in that population;

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\(^\text{92}\) See section 505A(d)(1)(A) of the FD&C Act (21 U.S.C. 355a(d)(1)(A)).

\(^\text{93}\) See section 505A(d)(1)(A) of the FD&C Act (21 U.S.C. 355a(d)(1)(A)).

\(^\text{94}\) See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).


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- It is not possible to conduct a study or studies of the drug for the specified indications in the pediatric population in a manner that would provide useful information (e.g., the study population is too small to yield interpretable results); and/or

- Outstanding safety concerns from studies or significant theoretical concerns need to be clarified with additional studies to support conducting studies in the pediatric population.

Historically, FDA has at times issued WRs solely for studies required under PREA, even if there were no other indications that may produce health benefits in the pediatric population. However, over time, data on pediatric labeling changes pursuant to BPCA and/or PREA have been collected. Between 2002 and 2019, there were 768 products with pediatric labeling changes under BPCA and/or PREA. Sixty-three percent of these labeling changes were based on studies conducted under PREA/Pediatric Rule alone; 21 percent were based on studies conducted under BPCA alone; 16 percent were based on studies conducted under both the BPCA and PREA. These data suggest that studies required under PREA are successfully completed, and that PREA requirements have resulted in an increase in pediatric labeling, even without the added incentive of the BPCA.

The BPCA provides FDA with discretion to determine whether to issue, and the appropriate scope of, WRs based on the information that “may produce health benefits” in the pediatric population.95 In light of the data on pediatric labeling changes pursuant to the BPCA and/or PREA, FDA believes WRs should be reserved for those sponsors who conduct additional pediatric studies — beyond what is required under PREA — that may produce health benefits in children. Thus, upon finalization of this guidance, FDA does not expect to issue WRs solely for studies or planned studies that are required under PREA. In general, FDA expects that a WR that includes studies or planned studies required under PREA will also include additional indications or populations. If there are no additional studies for indications or populations that may produce health benefits in the pediatric population beyond the studies or planned studies required under PREA, then FDA does not expect to issue a WR for that drug. For example, if a sponsor has an iPSP that includes a plan for deferred studies of a drug for pediatric juvenile idiopathic arthritis (pJIA), FDA does not expect to issue a WR solely for studies of pJIA in the same pediatric population. However, if FDA determines that this drug may produce health benefits in pediatric systemic juvenile idiopathic arthritis (sJIA), and there are no studies or planned studies required under PREA for this indication, then it may be appropriate for FDA to issue a WR for pediatric studies for both pJIA and sJIA.

In general, when considering issuance of a WR, FDA evaluates the need for studies for all pediatric subpopulations and for all indications for which the drug is being used or could be used in the pediatric population. In general, FDA considers the indications already approved for adults, indications pending for adults, and unapproved uses including uses that might be specific to the pediatric population. A single WR may address multiple indications and uses that are both approved and unapproved.96

95 See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).

96 See section 505A(d)(1)(B) of the FD&C Act (21 U.S.C. 355a(d)(1)(B)).
FDA can issue a WR that includes nonclinical studies. Section 505A(a) of the FD&C Act defines the term pediatric studies to mean “at least one clinical investigation (that, at the Secretary’s discretion, may include pharmacokinetic studies) in pediatric age groups (including neonates in appropriate cases) in which a drug is anticipated to be used, and, at the discretion of the Secretary, may include preclinical studies.” FDA may need certain toxicology studies in immature animals to evaluate the safety of drugs for use in pediatric populations. Accordingly, FDA may request that a sponsor or holder of an approved application conduct nonclinical studies before completing pediatric studies in humans. FDA may need to review such studies before it can determine whether information relating to use of the drug could produce health benefits in the pediatric population; thus, FDA may need these studies to determine if it will issue a WR.

In response to a WR, sponsors may, as appropriate, submit studies conducted by a third party. However, the sponsor should submit reports of studies that it has not conducted only if (1) the data from the studies appear to provide useful pediatric information that fairly responds to the WR issued by FDA; and (2) the sponsor obtains a right of reference to submit the reports of studies along with the underlying data.

A sponsor can use data it collects before or after FDA issues a WR to respond to the WR. Although FDA can request literature reviews as part of a larger WR, reviews of published literature alone are not pediatric studies that will qualify a drug for pediatric exclusivity.

Although a sponsor may, as appropriate, use studies it conducts to meet PREA requirements to qualify also for pediatric exclusivity, as mentioned, FDA does not consider pediatric studies a sponsor submits to an NDA or BLA (either in an original application, amendment, or supplement) before FDA issues a WR as being responsive to that WR. To qualify for pediatric exclusivity, sponsors and holders of an approved application should obtain a WR or an amendment to an existing WR before submitting the pediatric studies to an application.

3. Amended Written Requests

Each WR states that the WR may be amended. A sponsor may request an amendment to the WR, or FDA may issue an amendment on its own initiative. However, FDA does not anticipate amending WRs in the absence of scientific, medical, or regulatory justification.

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97 See section 505A(a) of the FD&C Act (21 U.S.C. 355a(a)).

98 We support the principles of the 3Rs (reduce/refine/replace) for animal use in testing when feasible. The FDA encourages sponsors to consult with review divisions when considering a nonanimal testing method believed to be suitable, adequate, validated, and feasible. The FDA will consider if the alternative method could be assessed for equivalency to an animal test method.

99 See sections 505A(a), 505A(b), and 505A(c) of the FD&C Act (21 U.S.C. 355a(a), 21 U.S.C. 355a(b), and 21 U.S.C 355a(c)).

100 As noted earlier in this section, going forward, FDA does not expect to issue WRs solely for studies or planned studies that are required under PREA.
Sponsors can request a change in the studies’ due date in the WR by submitting a proposed amendment to the WR. If a sponsor believes it will be unable to meet the time frames in a WR, it should contact FDA to request such a change as soon as possible. If FDA agrees to the change, it intends to notify the sponsor in writing regarding the extension time. Any request for an extension of time should take into consideration that completed reports of all studies should be submitted for filing at least 15 months before the expiration of the patent or exclusivity to which the pediatric exclusivity would attach. Sponsors should be aware that if they choose to submit within less than 15 months, FDA may not be able to complete the review and make a determination in time to meet the 9-month deadline.\footnote{See sections 505A(b)(2), 505A(c)(2), and 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(b)(2), 21 U.S.C. 355a(c)(2), and 21 U.S.C. 355a(d)(4)).}

FDA intends to issue all amendments to a WR in writing. Sponsors should also request any amendments to a WR in writing. Discussion of a proposed amendment at a meeting with FDA does not constitute a request to amend a WR nor does it constitute FDA’s amendment of the WR. In addition, if a sponsor has submitted a protocol that is inconsistent with the WR, and FDA has not commented on the protocol, the sponsor should not assume FDA agrees that the protocol is consistent with the WR. Even a minor change to a study, such as a change in the number of subjects, the age groups enrolled, or the elimination of certain testing requirements, may warrant a change in the protocol and a revision of the WR if it relates to a specified term of the WR.

Sponsors can submit preliminary data to an IND in support of a request for amendment. Sponsors that believe that their studies may not fairly respond to the WR as issued but nonetheless provide valuable pediatric information should (1) seek to obtain an amended WR \textit{before} submitting any pediatric study reports to their NDAs or BLAs; and (2) submit proposed amendments to their WRs early enough to ensure enough time for FDA to issue an amended WR and to ensure the sponsor has enough time to submit its studies at least 15 months before the expiration of any patent or exclusivity to which pediatric exclusivity would attach.

Sponsors should not submit the requested study reports to their NDAs or BLAs until \textit{after} they have received FDA’s response to requested amendments in writing. Reports of studies that do not fairly respond to the existing WR will \textit{not} qualify for pediatric exclusivity (see sections IV.C., How to Submit Study Reports in Response to a Written Request, IV.D., Qualifying for Pediatric Exclusivity, and IV.E., Determining Eligibility For Pediatric Exclusivity).\footnote{See sections 505A(b)(1), 505A(c)(1), 505A(d)(4), and 505A(h) of the FD&C Act (21 U.S.C. 355a(b)(1), 21 U.S.C. 355a(c)(1), 21 U.S.C. 355a(d)(4), and 21 U.S.C. 355a(h))).}

**B. How to Obtain a Written Request**

Historically, WRs have generally been issued after a drug is approved for use in adults. However, there may be situations in which it is appropriate to issue a WR before such an action. See sections II., Overview of Regulatory Strategy for Pediatric Drug Development, and III.B.1., Pediatric Study Plans, and previous subsections of this section for additional considerations relating to timing of a WR. Additionally, the appropriate timing of the submission of a specific
PPSR can be discussed with the relevant review division(s). FDA may issue a WR either in response to a PPSR or on its own initiative (see sections IV.A., Written Requests).

1. Submitting a PPSR

If a sponsor has exclusivity or patent protection for a drug or exclusivity for a biological product, or anticipates that it will have such exclusivity or patent protection, the sponsor can submit a PPSR to the appropriate review division (such a proposal can help to expedite FDA’s issuance of a WR). Sponsors should mark PPSRs with the header PROPOSED PEDIATRIC STUDY REQUEST and submit it to the appropriate IND. Sponsors that seek to qualify for pediatric exclusivity to attach to existing patents and/or exclusivities should plan to submit their PPSRs with sufficient time to:

- Permit FDA to review the PPSR, confer with the party submitting the PPSR as necessary, and issue the WR (including review by the FDA’s internal pediatric review committee (the PeRC) as required before issuance103);
- Allow time for the sponsor, after the WR is issued, to initiate the studies, complete the studies, and submit the reports for filing; and
- Provide FDA 180 days to review the studies and make an exclusivity determination, with a remaining, nonoverlapping 9 months before expiration of the patent or exclusivity period.104

The PPSR should describe the studies the sponsor or application holder proposes to conduct to qualify for pediatric exclusivity. The PPSR should include (1) a background section, (2) nonclinical studies, (3) drug information, (4) clinical studies, (5) known drug safety concerns and monitoring, (6) statistical information, including power of studies and statistical assessments, and (7) time frame for submitting reports of the study or studies.

It is important to note that a PPSR is not a substitute for an iPSP (section III.B.1., Pediatric Study Plans). Although these submissions may have some similarities, each one is submitted under a different statutory scheme and serves a distinct purpose. See sections II., Overview of Regulatory Strategy for Pediatric Drug Development, III., Pediatric Research Equity Act, and V.C., PREA and Pediatric Exclusivity, for additional information.

FDA intends to consider PPSRs that include requests to study multiple pediatric age groups in the same study, as appropriate. FDA recognizes that studies defined by age may be inappropriate when it is reasonable to define subgroups using methods other than age, such as development stage. If the sponsor submits data as part of a PPSR to indicate that a drug should

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103 See section 505A(f) of the FD&C Act (21 U.S.C. 355a(f)).

104 See sections 505A(b)(2), 505A(c)(2), and 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(b)(2), 21 U.S.C. 355a(c)(2), and 21 U.S.C. 355a(d)(4)).
be studied in pediatric groups identified by characteristics other than age, in general, FDA intends to consider that data when developing the WR.

FDA has 120 days after a sponsor submits a PPSR to review and act on the submission. Our response to the PPSR is either a WR or a PPSR inadequate letter, in which we inform the sponsor of the reasons we will not issue a WR at this time. In general, FDA also makes suggestions as to what the sponsor should include in a resubmitted PPSR that might support our issuance of a WR.

2. Issuance and Acceptance of a WR

The sponsor or application holder must respond to FDA within 180 days after receiving the WR indicating whether it will conduct the studies and, if so, indicate when it will initiate the studies.

The procedure for qualifying for exclusivity and the protections that exclusivity will confer are described in more detail in sections IV.E., Determining Eligibility For Pediatric Exclusivity, and IV.F., Attaching the Period of Pediatric Exclusivity After a Determination That a Drug Qualifies for Pediatric Exclusivity.

If a sponsor declines a WR that is issued by FDA, the sponsor must provide the reasons it declined the request. If the sponsor declines the WR because it is not possible to develop an appropriate pediatric formulation, the sponsor must submit to FDA the reasons such pediatric formulation cannot be developed. If a sponsor declines a WR, the sponsor is not eligible to qualify for pediatric exclusivity for the studies under that WR.

C. How to Submit Study Reports in Response to a Written Request

To qualify for pediatric exclusivity, sponsors or application holders must submit study reports in accordance with FDA requirements for filing. Studies submitted in an application or supplement that does not meet requirements for filing of an NDA, BLA, or supplement (i.e., FDA refuses to file the application or supplement) are not considered submitted to FDA.

In general, sponsors should also submit study reports in accordance with the guidance for industry Guideline for the Format and Content of the Clinical and Statistical Sections of an Application (July 1988) and the ICH guidance for industry E3 Structure and Content of Clinical Study Reports (July 1996).

105 See section 505A(d)(3) of the FD&C Act (21 U.S.C. 355a(d)(3)).


109 See section 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(d)(4)). For additional information on filing requirements and refusal to file, see, for example, 21 CFR 314.50, 21 CFR 314.101, and 21 CFR 601.2.
To help ensure that pediatric study reports are evaluated for eligibility for pediatric exclusivity in a timely manner, sponsors should include the following with the application or supplement:

- A header that states SUBMISSION OF PEDIATRIC STUDY REPORTS — PEDIATRIC EXCLUSIVITY DETERMINATION REQUESTED;
- A copy of the WR and any amendments;
- A summary of the pediatric studies conducted in response to the WR;
- An annotated WR indicating how and where in the submitted study reports each term of the WR has been addressed; and
- Proposed labeling that includes information regarding the results of the study or studies.

If there is information that is adequate to support the evaluation of dosing, safety, and efficacy in a subpopulation of the population included in the WR, FDA encourages the sponsor to submit an NDA, BLA, or supplement to incorporate that information into labeling for the drug before the determination of exclusivity. If making multiple submissions in response to a single WR, the sponsor should, in the final submission, reference prior submissions (including relevant submission dates) and mark only the last submission as described above.

D. Qualifying for Pediatric Exclusivity

We note at the outset that a commitment to complete a study at some future date is not sufficient to qualify a drug for pediatric exclusivity. Rather, to qualify for an initial period of pediatric exclusivity, a sponsor must submit study reports that fairly respond to an issued WR, were conducted in accordance with commonly accepted scientific principles and protocols, and have been reported in accordance with filing requirements. It is not necessary for the uses studied under the WR to be approved.

1. For a Drug Product That Is the Subject of a New Drug Application or Biologics License Application

A drug product qualifies for pediatric exclusivity when all of the following have occurred:

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110 See sections 505A(b) and 505A(c) of the FD&C Act (21 U.S.C. 355a(b) and 21 U.S.C. 355A(c)).

111 See sections 505A(b), 505A(c), and 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(b), 21 U.S.C. 355a(c), and 21 U.S.C. 355a(d)(4)).

112 Approval is required for a sponsor to qualify for a second 6-month period of pediatric exclusivity under section 505A(g) of the FD&C Act (21 U.S.C. 355a(g)). See section IV.F.1.b., A second 6-month period of pediatric exclusivity, for further discussion of that process.

113 See sections 505A(b), 505A(c), and 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(b), 21 U.S.C. 355a(c), and 21 U.S.C. 355a(d)(4)).
• FDA issued a WR for pediatric studies, and the sponsor or holder of an approved application agreed to the request.

• The sponsor or the holder of an approved application has submitted reports of the requested studies. Such reports should be submitted to the NDA or BLA after FDA issues the WR.

• The studies were completed using appropriate formulations for each age group and within the requested time frame.

• FDA has determined the studies fairly respond to the WR, have been conducted in accordance with commonly accepted scientific principles and protocols, and have been reported in accordance with filing requirements.

• FDA makes an exclusivity determination at least 9 months before the expiration date of the patent and/or exclusivity protection to which the pediatric exclusivity will attach.\footnote{See section 505A(b)(2) and 505A(c)(2) of the FD&C Act (21 U.S.C. 355a(b)(2) and 21 U.S.C. 355a(c)(2)) and section 351(m)(4) of the PHS Act (42 U.S.C. 262(m)(4)).}

2. Nonprescription Drugs

Nonprescription drugs marketed under an approved application may be eligible for pediatric exclusivity, and the recommendations in this guidance apply to those nonprescription drugs.

Nonprescription drugs that are marketed pursuant to a monograph developed under the over-the-counter drug review are not eligible for exclusivity under section 505A of the FD&C Act.

E. Determining Eligibility For Pediatric Exclusivity

For a drug to be considered eligible for pediatric exclusivity, FDA recommends that the sponsor submit a complete report of all studies to FDA at least 15 months before the expiration of any existing patent or exclusivity it wishes to protect (i.e., the 9-month time period\footnote{See section 505A(b)(2) and 505A(c)(2) of the FD&C Act; 21 U.S.C. 355a(b)(2) and 21 U.S.C. 355a(c)(2).} plus the 180-day exclusivity determination review period\footnote{See section 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(d)(4)).}). If sponsors choose to submit within less than 15 months, FDA may not be able to complete its review of such studies and make a determination about the drug’s eligibility for pediatric exclusivity in time to meet the 9-month deadline.

In making an eligibility determination, FDA will evaluate whether the studies fairly respond to the WR, were conducted in accordance with commonly accepted scientific principles and
Contains Nonbinding Recommendations
Draft — Not for Implementation

protocols, and have been reported in accordance with filing requirements.\textsuperscript{117} FDA’s pediatric exclusivity boards (Boards), which are comprised of representatives from the Center for Drug Evaluation and Research or the Center for Biologics Evaluation and Research, including representatives with pediatric expertise, make this determination. In general, the Boards consider the following when assessing, as required by statute, whether the studies \textit{fairly respond} to a WR:

- The purpose of the pediatric exclusivity provision as described in the statute, with reference to the legislative history. The statute makes clear that its purpose is to generate meaningful clinical information on the use of drug products in children that will result in a health benefit to pediatric populations.\textsuperscript{118}

- Whether the sponsor met the terms of a WR.

- The information sought in the WR and the objectives stated in the WR.

In general, the Boards ask whether the studies were designed and carried out by the sponsor in a way likely to meet those objectives specified in the WR and underlying the exclusivity provision as a whole. When a sponsor meets the terms of a WR, the resulting studies fairly respond to that WR because studies that are carried out in accordance with the trial’s plans and objectives, as expressed in the WR, generally satisfy the statutory goal of obtaining meaningful pediatric use information. Sometimes, a sponsor fails to produce meaningful pediatric information despite conducting the studies in the manner requested. Under such circumstances, FDA nevertheless considers the sponsor to have fairly responded to the WR. FDA understands that the failure to generate meaningful information in such cases is at least partially attributable to study design, and FDA and the sponsor generally design studies described in a WR jointly.

Where the sponsor has not met the terms of the WR, FDA evaluates whether the information generated by the studies is nevertheless sufficient to meet the objectives of the WR in light of the information sought in the WR. If FDA determines that the objectives of the WR were met, then FDA concludes that the sponsor has fairly responded, even if it did not meet the terms of the WR. FDA considers studies that do not meet the terms of the written request to have fairly responded if, considering the data provided by the sponsor as a whole (i.e., by considering all relevant data, and not just data generated by those studies), the sponsor meets the objectives of the WR by generating clinically meaningful information of the general type (quality and quantity) the WR contemplates.

\textsuperscript{117} See section 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(d)(4)).

\textsuperscript{118} See, for example, section 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)) requiring FDA to determine “that information relating to the use of a new drug in the pediatric population may produce health benefits,” section 505A(f)(2) of the FD&C Act (21 U.S.C. 355a(f)(2)) requiring review by the PeRC before FDA issues a WR, section 505A(f)(3) of the FD&C Act, providing that the same committee may review the reports of studies conducted in response to a WR before FDA makes a determination regarding pediatric exclusivity, section 505A(f)(6)(E) of the FD&C Act (21 U.S.C. 355a(f)(6)(E)) requiring FDA to publicly report, among other things, labeling changes made as a result of studies conducted in response to a WR, and section 505A(k)(2) of the FD&C Act (21 U.S.C. 355a(k)(2)) requiring sponsors to distribute the same to physicians and other health care providers.
For example, if a specific number of subjects is requested or a specific study duration or endpoint is specified to ensure that the study will generate adequate data to provide a health benefit, failure to comply with these elements of the WR may result in a denial of exclusivity. Denial is likely if, in the absence of compliance with the terms of the WR, the studies are not expected to be interpretable or will not provide information that otherwise yields a health benefit to the pediatric populations addressed in the WR.

Where a WR is capable of more than one interpretation, the Boards generally consider a fair response to be one that interprets the WR in a manner likely to generate information that will provide a health benefit (including meaningful pediatric labeling) in the relevant populations that the WR asked the sponsor to study. If the studies submitted fairly respond to the WR, the Boards will recommend that the drug is eligible for pediatric exclusivity (assuming the other statutory requirements for pediatric exclusivity are met). If, on the other hand, the sponsor responds to the WR in such a way that the possibility of a health benefit (including meaningful pediatric labeling in relevant age groups) from the studies conducted is not likely, the Boards are likely to conclude that the submission does not fairly respond to the WR.

Generally, FDA expects to notify sponsors or holders of an approved application within the 180-day period after the study reports are submitted whether the study reports fairly responded to the WR and the drug qualifies for pediatric exclusivity.

F. Attaching the Period of Pediatric Exclusivity After a Determination That a Drug Qualifies for Pediatric Exclusivity

1. Drug Products

a. An initial 6-month period of pediatric exclusivity

Pediatric exclusivity will attach to all unexpired exclusivities and patents listed in the Approved Drug Products With Therapeutic Equivalence Evaluations publication (the Orange Book).

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119 See sections 505A(b)(1), 505A(c)(1), and 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(b)(1), 21 U.S.C. 355a(c)(1), and 21 U.S.C. 355a(d)(4)).

120 See section 505A(d)(4) of the FD&C Act (21 U.S.C. 355a(d)(4)).

121 For the purposes of this section IV.F.1., references to drugs or drug products do not include biological products licensed under section 351 of the PHS Act (42 U.S.C. 262). Although the considerations in this section are generally relevant to such biological products, pediatric exclusivity for biological products differs in certain ways from pediatric exclusivity for other drug products. For more information see section IV.F.2., Biological Products, below.

122 Pediatric exclusivity that has attached to the end of the patent term will block for an additional 6 months after the patent expires approval of an ANDA or 505(b)(2) application if (1) the ANDA or 505(b)(2) sponsor did not seek approval until the end of the patent term; or (2) the ANDA or 505(b)(2) sponsor’s patent challenge has been unsuccessful. See, for example, sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)). In addition, if an ANDA or 505(b)(2) sponsor files a paragraph IV certification challenging a listed patent, and the patent litigation is ongoing when the patent expires, the pediatric exclusivity will attach at the
For studies a sponsor conducts on a previously unapproved drug, pediatric exclusivity will attach to any exclusivities or patents that will be listed in the Orange Book upon approval of that drug and to certain later listed patents or exclusivities. For studies a sponsor conducts on a previously approved drug, pediatric exclusivity will attach to protections listed for the previously approved drug at the time the drug qualifies for pediatric exclusivity and to certain later listed patents and exclusivities, as described further in section IV.F.1.c., Later-filed applications containing the same drug. Pediatric exclusivity for combination drugs may raise additional considerations that are not addressed in this guidance.

b. A second 6-month period of pediatric exclusivity

Each WR may result in only one 6-month period of pediatric exclusivity. However, after a drug has qualified for an initial period of pediatric exclusivity, a sponsor submitting a supplement to an application can submit additional pediatric studies meeting the relevant statutory requirements in response to a second WR. The second 6-month period of pediatric exclusivity will attach only to any new 3-year exclusivity period for which the supplemental application qualifies. In addition, several other considerations regarding a second period of pediatric exclusivity are presented as follows:

- A second WR can result in a 6-month period of exclusivity only if the response to the WR results in an approved supplemental application for a new use.

- A new use is a use not included in the approved labeling of an approved drug. For example, expansion of the labeling to include a new pediatric population constitutes a new use.

end of the patent term to block approval of that ANDA or 505(b)(2) application for an additional 6 months after the patent expires (Ranbaxy Labs. v. FDA).

123 The Orange Book is available at https://www.accessdata.fda.gov/scripts/cder/ob/index.cfm.

124 See section 505A(c) of the FD&C Act (21 U.S.C. 355a(c)).

125 See section 505A(b) of the FD&C Act (21 U.S.C. 355a(b)).

126 See section 505A(c) of the FD&C Act (21 U.S.C. 355a(c)).

127 See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).

128 See section 505A(g) of the FD&C Act (21 U.S.C. 355a(g)).

129 See section 505A(g) of the FD&C Act (21 U.S.C. 355a(g)).

130 See section 505A(g) of the FD&C Act (21 U.S.C. 355a(g)).

131 See 21 CFR 99.3(g).
The supplement for a new use submitted in response to the second WR must qualify for 3-year exclusivity\(^{132}\) or no 6-month period of pediatric exclusivity will attach.

The second 6-month period of pediatric exclusivity attaches *only* to the 3-year exclusivity applied to the supplement for a new use containing the studies submitted in response to the second WR and not to any other exclusivity or patent protections applicable to the drug.

No more than two 6-month periods of exclusivity under the BPCA are possible for any specific drug product.

c. Later-filed applications containing the same drug

In situations where a sponsor submits an application or supplement containing an active moiety for which the sponsor previously qualified for pediatric exclusivity, pediatric exclusivity does not attach to new (not previously listed) patents or exclusivity covering the later filed applications or supplements unless the subsequent drug product could not be labeled without the data that qualified the previously approved drug product for the prior pediatric exclusivity.\(^{133}\)

FDA notes that if pediatric exclusivity for which a drug product previously qualified has attached to a listed patent or exclusivity protecting the previously approved application that also protects the new application or new supplement held by the same sponsor, the pediatric exclusivity also attaches to that patent in conjunction with the new application or supplement.\(^{134}\) For example, if a sponsor qualifies for a 6-month pediatric exclusivity that attaches to a 5-year exclusivity, that exclusivity attaches to each of the sponsor’s NDAs protected by that 5-year exclusivity, regardless of when the new application or supplement is filed or what it contains. The following examples are provided:

- **Example 1** — Drug 1 (D1) qualifies for pediatric exclusivity. The sponsor or holder of an approved application for D1 later files a different application for a drug product containing D1 or a supplement to an existing application for a drug product containing D1. FDA does not need any of the data the sponsor or holder of an approved application submitted for pediatric exclusivity to approve the new application or new supplement. The pediatric exclusivity does not attach to any exclusivities or patents that apply solely to the new application or the new supplement.

- **Example 2** — D1 qualifies for pediatric exclusivity. The sponsor or holder of the approved application for D1 later files a different application for a drug product containing D1 or a supplement to an existing application for a drug product containing D1. The drug product could not be labeled with the data submitted in the later-filed applications or supplements without the data the sponsor or holder of an approved

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\(^{133}\) See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).

\(^{134}\) See sections 505A(b)(1) and 505A(c)(1) of the FD&C Act (21 U.S.C. 355a(b)(1) and 21 U.S.C. 355a(c)(1)).
application previously submitted for pediatric exclusivity. The pediatric exclusivity
attaches to any exclusivities or patents that apply to the new application or the new
supplement. In addition, if the pediatric exclusivity attaches to a patent or exclusivity
that also protects the new application or new supplement, the pediatric exclusivity applies
to the new application or supplement to the same extent it applies to the previously
approved application.

2. Biological Products

Pediatric exclusivity for biological products differs in some ways from provisions applicable to
other drug products. First, as described below, pediatric exclusivity only attaches to reference
product and orphan drug exclusivity periods. Second, unlike other drug products, pediatric
exclusivity does not attach to patents for biological products.135

Under section 351(k)(7)(A) of the PHS Act, approval of an application for a biosimilar or
interchangeable biological product submitted under section 351(k) of the PHS Act may not be
made effective until 12 years after the date on which the reference product was first licensed
under section 351(a) of the PHS Act. Moreover, under section 351(k)(7)(B) of the PHS Act, an
application for a biosimilar or interchangeable biological product submitted under section 351(k)
of the PHS Act may not be submitted for review until 4 years after the date on which the
reference product was first licensed under section 351(a) of the PHS Act. An additional 6-month
period of pediatric exclusivity will attach to the 12- and 4-year periods if the sponsor meets the
requirements for pediatric exclusivity pursuant to section 505A of the FD&C Act.136
Furthermore, an additional 6-month period of pediatric exclusivity will also attach to the 7 years
of orphan drug exclusivity for a biological product designated under section 526 of the FD&C
Act for a rare disease or condition.137

V. ELEMENTS COMMON TO PREA AND THE BPCA

A. The Pediatric Review Committee

Section 505C of the FD&C Act directed FDA to establish the PeRC that must review all WRs
and all requests for deferrals, deferral extensions, and waivers.138 The PeRC also provides
consultation on pediatric assessments and on iPSPs, agreed iPSPs, and any significant
amendments to such plans.139

135 See section 351(m) of the PHS Act (42 U.S.C. 262(m)).

136 See section 351(m) of the PHS Act (42 U.S.C. 262(m)).

137 See section 527(a) of the FD&C Act and section 351(m) of the PHS Act (21 U.S.C. 360cc(a); 42 U.S.C. 262(m)).

138 See sections 505C, 505A(f), and 505B(f) of the FD&C Act (21 U.S.C. 355d, 21 U.S.C. 355a(f), and 21 U.S.C.
355c(f)).

139 See section 505B(f) of the FD&C Act (21 U.S.C. 355c(f)).
As a general matter, the PeRC also reviews significant amendments to WRs and PPSR inadequate letters to ensure consistency. Members of the PeRC include FDA employees with expertise in pediatrics (including representation from the Office of Pediatric Therapeutics (OPT)), neonatology, biopharmacology (i.e., pharmacology/toxicology), statistics, chemistry, legal issues, pediatric ethics, the appropriate expertise pertaining to the pediatric drug under review, and other individuals as needed. In addition to the responsibilities above, the PeRC also provides consultation on tracking information regarding pediatric assessments and labeling changes. As a general matter, members of the relevant drug review division provide background information to the PeRC and are present during the discussion of an application.

B. Publishing Information About Pediatric Studies

1. Pediatric Exclusivity Determinations

FDA posts on its website a list of exclusivity determinations on approved drugs that have qualified for pediatric exclusivity. FDA also publishes pediatric exclusivity information for drugs in the Patent and Exclusivity Information section of the Orange Book and its supplements in the same manner as FDA publishes information regarding 5-year exclusivity, 3-year exclusivity, patent listings, and orphan drug exclusivity.

2. Medical, Statistical, and Clinical Pharmacology Reviews

Sections 505A(k) and 505B(h) of the FD&C Act require that FDA publish the medical, statistical, and clinical pharmacology reviews of pediatric studies conducted under the BPCA and of pediatric assessments under PREA. For studies submitted in response to a WR, FDA must do so within 210 days after the submission. For most pediatric assessments submitted under PREA, FDA must do so within 330 days after the submission. FDA makes such

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141 See sections 505A(f)(6) and 505B(f)(6) of the FD&C Act (21 U.S.C. 355a(f)(6) and 21 U.S.C. 355c(f)(6)).

142 See the FDA’s Pediatric Exclusivity Granted web page at https://www.fda.gov/drugs/development-resources/pediatric-exclusivity-granted.


144 See section 505A(k)(1) of the FD&C Act (21 U.S.C. 355a(k)(1)). For drug products for which exclusivity determinations were made before September 27, 2007, we have posted summaries of medical and clinical pharmacology reviews of studies conducted under section 505A of the FD&C Act, consistent with applicable BPCA requirements at that time.

145 See section 505B(h)(1) of the FD&C Act (21 U.S.C. 355c(h)(1)). This provision of the law first went into effect on September 27, 2007.
information publicly available consistent with section 301(j) of the FD&C Act, the Freedom of Information Act, and the Trade Secrets Act.146

Before disclosure, the medical, statistical, and clinical pharmacology reviews are redacted, as appropriate, for trade secret information and for confidential commercial information.147

As FDA interprets section 505A(k) of the FD&C Act, the 210-day BPCA disclosure requirement is triggered when a sponsor or holder of an approved application submits an application or a supplement in response to a WR that FDA determines meets filing requirements (see section IV.C., How to Submit Study Reports in Response to a Written Request). FDA interprets the disclosure requirement to apply to such applications and supplements submitted in response to a WR under the BPCA, regardless of whether (1) the application process is completed or it is later withdrawn; (2) the drug qualifies for pediatric exclusivity; or (3) the application or supplement is approved or the sponsor receives a complete response letter. In addition, FDA interprets the disclosure requirement to apply to partial responses to a WR under the BPCA that meet the filing requirements. (See section IV.C., How to Submit Study Reports in Response to a Written Request, for a discussion of partial responses.)

3. Other Pediatric Information

FDA maintains a web page148 containing extensive information about pediatric studies conducted under sections 505A and 505B of the FD&C Act. The web page includes, among other things, statistics and other information regarding the relevant studies, drugs, labeling changes, and reports. Statistics we post include the number of waivers, deferrals, and deferral extensions granted; the number of pediatric formulations developed; and the number of formulations not developed (including the reason they were not developed).149 The web page also identifies drugs approved for use in a pediatric population for which a pediatric formulation

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147 21 CFR 314.430 is FDA’s regulation regarding the public disclosure of information in a drug application or abbreviated application. 21 CFR 601.51 is FDA’s regulation regarding the public disclosure of information in a biological product file. Under 21 CFR 314.430(b), we will not publicly disclose the existence of an application or abbreviated application before an approval or tentative approval letter is sent, unless the existence of the application or abbreviated application has been previously publicly disclosed or acknowledged. Under 21 CFR 601.51(b), we will not disclose the existence of a biological product file before the application has been approved if the existence of the file has not been previously publicly disclosed or acknowledged. Under 21 CFR 314.430(d)(1) and 601.51(d)(1), if the existence of the application or biological product file has been publicly disclosed or acknowledged, as a general matter, no data or information contained in the application, abbreviated application, or biological product file is available for public disclosure before we send an approval letter or before a license is issued. We note that 21 CFR 314.430 and 601.51 were promulgated before the passage of PREA and the BPCA and do not specifically discuss the disclosure of medical, statistical, and clinical pharmacology reviews of pediatric studies conducted under the BPCA and of pediatric assessments under PREA. As a general matter, FDA discloses such reviews only to the extent the drug product studied has already been approved. Additionally, the protected status of particular information in the reviews is determined on a case-by-case basis.


149 See section 505B(f)(6) of the FD&C Act (21 U.S.C. 355c(f)(6)).
was developed and that qualified for pediatric exclusivity, but the formulation was not marketed within 1 year of the exclusivity determination.\textsuperscript{150} FDA also maintains a web page containing information about section 409I of the PHS Act.\textsuperscript{151}

A list of approved drugs for which WRs have been issued is published on the FDA website.\textsuperscript{152} In addition, WRs, including any amendment(s) if not otherwise incorporated into one document, are posted on the FDA website within 30 days of the determination that the requirements for exclusivity have been met.\textsuperscript{153}

Information from a required annual review following the granting of a deferral will be posted publicly within 90 days of submission.\textsuperscript{154} The posting will include the information submitted through the annual review, the name of the applicant, the date on which the drug was approved, and the date of each deferral or deferral extension.\textsuperscript{155}

Finally, FDA posts guidances, relevant regulations, relevant presentations from conferences, press releases, and reports, among other information. Transcripts from Pediatric Advisory Committee meetings, beginning September 2004, are posted as well, as are transcripts from past meetings of the Pediatric Advisory Subcommittee of the Anti-Infective Drugs Advisory Committee, held between April 1999 and June 2004.\textsuperscript{156} FDA intends to update these sites regularly.

C. PREA and Pediatric Exclusivity

To qualify for pediatric exclusivity, the pediatric studies conducted to satisfy the requirements of PREA must be the subject of a WR and satisfy all other requirements for pediatric exclusivity.

\textsuperscript{150} See section 505A(e)(2) of the FD&C Act (21 U.S.C. 355a(e)(2)).

\textsuperscript{151} See the Off-Patent Studies Under BPCA web page at https://www.fda.gov/drugs/development-resources/patent-studies-under-bpca.

\textsuperscript{152} See the Written Requests Issued web page at https://www.fda.gov/drugs/development-resources/written-requests-issued. On occasion, information obtained by FDA subsequent to issuance of a WR causes FDA to rescind the WR. This list is not updated to indicate when a WR has been rescinded.

\textsuperscript{153} See section 505A(e)(1) of the FD&C Act (21 U.S.C. 355a(e)(1)). See also the FDA’s Pediatric Exclusivity Granted web page at https://www.fda.gov/drugs/development-resources/pediatric-exclusivity-granted.


\textsuperscript{156} See https://www.fda.gov/advisory-committees/pediatric-advisory-committee/past-meeting-materials-pediatric-advisory-committee.
under the BPCA.\textsuperscript{157} As discussed in section IV.A.2., Written Request Studies, FDA does not expect to issue WRs solely for studies or planned studies that are required under PREA.

For already marketed drugs, FDA may require pediatric assessments under PREA if it finds, for example, that the drug is used for a substantial number of pediatric patients for the labeled indications and adequate pediatric labeling could confer a benefit on pediatric patients.\textsuperscript{158} In some cases, FDA may first issue a WR under section 505A of the FD&C Act before requiring studies under section 505B(b). In those cases, if the sponsor declines the WR for the labeled indication(s), FDA may still require those studies under PREA.\textsuperscript{159}

It is important to note the distinction between the scope of the studies FDA requests under the BPCA and those required under PREA. The scope of studies described in a WR may be broader than those required under PREA. FDA’s authority to issue a WR extends to the use of an active moiety for indications that may produce health benefits in the pediatric population, regardless of whether it has previously approved the indications in adults.\textsuperscript{160} Under PREA, pediatric assessments are required only for those indications the sponsor has included in the pending application.\textsuperscript{161} To learn more about eligibility for pediatric exclusivity, see section IV., Best Pharmaceuticals for Children Act, or contact the relevant review division.

### D. Considerations for Labeling of Drug Products

#### 1. Labeling Study Results

Study results submitted in response to PREA or a WR must be described in labeling regardless of whether these findings support safety and/or effectiveness, do not support safety and/or

\textsuperscript{157} See sections 505A(b)(1), 505A(c)(1), 505A(d) and 505A(h) of the FD&C Act (21 U.S.C. 355a(b)(1), 21 U.S.C. 355a(c)(1), 21 U.S.C. 355a(d), and 21 U.S.C. 355a(h)).

\textsuperscript{158} See section 505B(b)(1) of the FD&C Act (21 U.S.C. 355c(b)(1)).

\textsuperscript{159} See section 505B(b) of the FD&C Act (21 U.S.C. 355c(b)). This section states that the Secretary may require a sponsor or holder of an approved application to submit pediatric assessments as described under section 505B(a)(2) of the FD&C Act if the Secretary finds that “(A) (i) the drug or biological product is used for a substantial number of pediatric patients for the labeled indications; and (ii) adequate pediatric labeling could confer a benefit on pediatric patients; (B) there is reason to believe that the drug or biological product would represent a meaningful therapeutic benefit over existing therapies for pediatric patients for 1 or more of the claimed indications; or (C) the absence of adequate pediatric labeling could pose a risk to pediatric patients.”

\textsuperscript{160} See sections 505A(a), 505A(b)(1), 505A(c)(1), and 505A(d)(1)(B) of the FD&C Act (21 U.S.C. 355a(a); 21 U.S.C. 355a(a), 21 U.S.C. 355a(b)(1), 21 U.S.C. 355a(c)(1), and 21 U.S.C. 355a(d)(1)(B)).

\textsuperscript{161} See sections 505B(a)(1)(A) and 505B(a)(2) of the FD&C Act (21 U.S.C. 355c(a)(1)(A) and 21 U.S.C. 355c(a)(2)). Note, however, that molecularly targeted pediatric cancer investigations are based on molecular mechanism of action rather than clinical indication. See sections 505B(a)(1)(B) and 505B(a)(3) of the FD&C Act (21 U.S.C. 355c(a)(1)(B) and 21 U.S.C. 355c(a)(3)). For additional information, see the guidance for industry FDARA Implementation Guidance for Pediatric Studies of Molecularly Targeted Oncology Drugs: Amendments to Sec. 505B of the FD&C Act.
effectiveness, or are inconclusive.\textsuperscript{162} If a full or partial waiver is granted because there is evidence that a drug would be unsafe or ineffective in pediatric populations, applicants must include this information in labeling.\textsuperscript{163}

Applicants must distribute information to health care providers describing any labeling changes that are approved as a result of these studies as required by FDA.\textsuperscript{164}

2. Dispute Resolution

The BPCA and PREA provide for a dispute resolution process when FDA and the applicant fail to agree on appropriate labeling changes.\textsuperscript{165} If the applicant does not agree within the specified time period after FDA’s request to make labeling changes, FDA must refer the matter to the Pediatric Advisory Committee (PAC).\textsuperscript{166} The PAC then has 90 days after receiving the referral to review the pediatric study reports and make a recommendation to FDA.\textsuperscript{167} FDA will consider the recommendation, and, if appropriate, within 30 days after receiving the recommendation, make a request to the applicant to make the labeling changes FDA determines to be appropriate.\textsuperscript{168} If the applicant fails to agree to make the labeling changes within 30 days after receiving such a request, the drug may be deemed misbranded.\textsuperscript{169}

3. Priority Review of Applications and Labeling Supplements

Any application or supplement to an application that proposes a labeling change as a result of pediatric studies a sponsor conducts under section 505A of the FD&C Act will be considered a priority application or supplement.\textsuperscript{170} This priority status applies even if the studies submitted

\textsuperscript{162} See sections 505A(j) and 505B(g)(2) of the FD&C Act (21 U.S.C. 355a(j) and 21 U.S.C. 355c(g)(2)). See also the guidance for industry Pediatric Information Incorporated Into Human Prescription Drug and Biological Product Labeling.

\textsuperscript{163} See section 505B(a)(5)(D) of the FD&C Act (21 U.S.C. 355c(a)(5)(D)).

\textsuperscript{164} See sections 505A(k)(2) and 505B(h)(2) of the FD&C Act (21 U.S.C. 355a(k)(2) and 355c(h)(2)).

\textsuperscript{165} See sections 505A(i)(2) and 505B(g)(1) of the FD&C Act (21 U.S.C. 355a(i)(2) and 355c(g)(1)).

\textsuperscript{166} See sections 505A(i)(2)(A) and 505B(g)(1)(A) of the FD&C Act (21 U.S.C. 355a(i)(2)(A) and 355c(g)(1)(A)).

\textsuperscript{167} See sections 505A(i)(2)(B) and 505B(g)(1)(B) of the FD&C Act (21 U.S.C. 355a(i)(2)(B) and 355c(g)(1)(B)).

\textsuperscript{168} See sections 505A(i)(2)(C) and 505B(g)(1)(C) of the FD&C Act (21 U.S.C. 355a(i)(2)(C) and 355c(g)(1)(C)).

\textsuperscript{169} See sections 505A(i)(2)(D) and 505B(g)(1)(D) of the FD&C Act (21 U.S.C. 355a(i)(2)(D) and 355c(g)(1)(D)).

\textsuperscript{170} See section 505A(i)(1) of the FD&C Act (21 U.S.C. 355a(i)(1)). This priority review provision applies only to applications and supplements containing studies conducted under section 505A of the FD&C Act; it does not apply to an application or supplement solely because it contains pediatric information. Note that NDAs and BLAs may be otherwise eligible for priority review. For information on Prescription Drug User Fee Act VI performance goals and procedures, see https://www.fda.gov/industry/fda-user-fee-programs/prescription-drug-user-fee-amendments.
did not respond completely to the WR or did not otherwise qualify for pediatric exclusivity.\textsuperscript{171} Applications that include a pediatric assessment submitted with the sole intention of responding to PREA requirements do not necessarily receive priority review.

For more information about priority review, see FDA’s Prescription Drug User Fee Act reauthorization performance goals and procedures document.\textsuperscript{172}

E. Adverse Event Reporting for Drug Products Subject to the BPCA and PREA

At the same time that a sponsor submits reports of studies responding to a WR, the sponsor must also provide FDA with all available postmarketing adverse event reports regarding the studied drug.\textsuperscript{173} The format of the postmarketing adverse event report should follow the model for a periodic safety update report described in the ICH guidance for industry \textit{E2C(R2) Periodic Benefit-Risk Evaluation Report (PBRER)} (July 2016). In addition, the sponsor may contact the review division for further information.

Eighteen months after the date of a labeling change made to reflect studies conducted under PREA or the BPCA, the applicable center refers to OPT a report of all adverse events received by FDA for the drug product.\textsuperscript{174} As a general matter, OPT presents a report and analysis to the PAC, and the PAC reviews this analysis and recommends whether additional monitoring (other than the usual surveillance) is necessary. When the PAC considers additional monitoring necessary after the 18-month period, the center generally continues to refer adverse event reports to OPT.

VI. ADDITIONAL INFORMATION

The Division of Pediatrics and Maternal Health can provide general information about complying with PREA and the BPCA. Additional pediatric information and contact information is available on the Pediatric Product Development web page.\textsuperscript{175}

\textsuperscript{171} See section 505A(i)(1) of the FD&C Act (21 U.S.C. 355a(i)(1)).

\textsuperscript{172} See PDUFA Reauthorization Performance Goals and Procedures Fiscal Years 2013 Through 2017 at \url{https://www.fda.gov/media/81306/download}.

\textsuperscript{173} See section 505A(d)(2)(B) of the FD&C Act (21 U.S.C. 355a(d)(2)(B)).

\textsuperscript{174} See sections 505A(l) and 505B(i) of the FD&C Act (21 U.S.C. 355a(l) and 355e(i)).

\textsuperscript{175} Available at \url{https://www.fda.gov/drugs/development-resources/pediatric-and-maternal-health-product-development}. 
Assessment — Contains data, gathered using appropriate formulations for each age group for which the assessment is required, that are adequate to assess the safety and effectiveness as well as support the dosing and administration of a drug product for each relevant pediatric age group (see section 505B(a)(2) of the FD&C Act (21 U.S.C. 355c(a)(2))).

Certify/Certification — In general, a certification is a statement from the applicant that the data provided to support a deferral¹ and/or waiver² request are accurate and complete.

Deferral — Defers submission of some or all of the required assessments or reports on the molecularly targeted pediatric cancer investigation until a specified date (see section 505B(a)(4) of the FD&C Act (21 U.S.C. 355c(a)(4))).

Pediatric Study Plan (PSP) — An outline of planned pediatric studies, along with any deferral and/or waiver requests, that is submitted by a sponsor for an application subject to PREA before the submission of assessments or reports on the molecularly targeted pediatric cancer investigation (see section 505B(e) of FD&C Act (21 U.S.C. 355c(e))). For more information, see the guidance for industry Pediatric Study Plans: Content of and Process for Submitting Initial Pediatric Study Plans and Amended Initial Pediatric Study Plans (July 2020).³

Proposed Pediatric Study Request (PPSR) — In general, a PPSR is a submission describing what pediatric studies a sponsor or application holder believes will yield information about the use of a drug that may produce health benefits in the pediatric population.

Molecularly Targeted Pediatric Cancer Investigation — An investigation of a drug described in section 505B(a)(1)(B) of the FD&C Act that must be designed to yield clinically meaningful pediatric study data, gathered using appropriate formulations for each age group for which the study is required, regarding dosing, safety, and preliminary efficacy to inform potential pediatric labeling (see section 505B(a)(3) of the FD&C Act (21 U.S.C. 355c(a)(3))).

Waiver — Waives the requirement to submit assessments or reports on the molecularly targeted pediatric cancer investigation for the entire pediatric population or specific age group(s) (see section 505B(a)(5) of the FD&C Act (21 U.S.C. 355c(a)(5))).

Written Request (WR) — In general, a WR is a document from FDA, signed by the applicable office director(s), requesting submission of a certain study or studies to determine whether the use of a drug could provide a meaningful health benefit in the pediatric population that is issued under section 505A of the FD&C Act (21 U.S.C. 355a) or section 351(m) of the PHS Act (42 U.S.C. 262(m)).


² See section 505B(a)(5) of the FD&C Act (21 U.S.C. 355c(a)(5)).

³ We update guidances periodically. To make sure you have the most recent version of a guidance, check the FDA guidance web page at https://www.fda.gov/regulatory-information/search-fda-guidance-documents.