

POLICY AND PROCEDURES (ポリシーおよび手順)

OFFICE OF GENERIC DRUGS

Conversion of ANDA Approval to Tentative Approval Because of Court Order

裁判所命令による ANDA 承認の暫定承認への変更

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PURPOSE (目的)

This Manual of Policies and Procedures describes the policies and procedures of the Office of Generic Drugs for converting the approval status of an abbreviated new drug application (ANDA) from final approval to tentative approval (TA) following a court order issued under 35 U.S.C. 271(e)(4)(A) for patent infringement.

本ポリシーおよび手順のマニュアルでは、271(e)(4)(A)の下で発出される特許侵害の裁判所命令を受け、ジェネリック医薬品局における簡略新薬承認申請 (ANDA) の承認ステータスを最終承認から暫定承認 (TA) に変更するためのポリシーおよび手順について記述する。

BACKGROUND (背景)

The timing of ANDA approval depends on, among other things, the patent and exclusivity protections for the reference listed drug (RLD) on which the applicant relies in seeking approval. An applicant must provide, in its ANDA, information related to any patents listed for the RLD in the Food and Drug Administration's *Approved Drug Products With Therapeutic Equivalence Evaluations* (the Orange Book).¹ In particular, an ANDA applicant generally must submit to the Food and Drug Administration (FDA) one of four specified certifications regarding the patents for the RLD under section 505(j)(2)(A)(vii) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 355(j)(2)(A)(vii)).

With respect to each patent listed in the Orange Book for the RLD, the ANDA applicant's patent certification must state one of the following:

ANDA 承認のタイミングは、とりわけ、申請者が承認を求める際に使用する参照リスト掲載医薬品 (RLD) の特許および独占権の保護に依存する。

申請者は、ANDA において、FDA の治療的同等性評価のある承認医薬品 (オレンジブック1) において RLD にリスト掲載されている特許に関連する情報を提供する必要がある。特に、ANDA 申請者は通常、FD&C 法のセクション 505(j)(2)(A)(vii) に基づく RLD の特許に関する 4 つの指定された証明のうちの 1 つを FDA に提出する必要がある。

RLD のオレンジブックに記載されている各特許について、ANDA 申請者の特許認定には、次のいずれかが記載されている必要がある :

- That such patent has expired (a paragraph II certification),
- The date on which such patent will expire (a paragraph III certification), or
- That such patent is invalid, unenforceable, or will not be infringed by the manufacture, use, or sale of the new drug for which the application is submitted (a paragraph IV certification)²

- そのような特許が失効していること (パラグラフ II 証明)、
- そのような特許が失効する日付 (パラグラフ III 証明)、または
- そのような特許が無効であり、行使不能であるか、または申請された新薬の製造、使用、または販売によって侵害されないこと (パラグラフ IV 証明)²

¹ オレンジブックの入手 : <https://www.accessdata.fda.gov/scripts/cder/ob/>.

² Section 505(j)(2)(A)(vii) of the FD&C Act; see also 21 CFR 314.94(a)(12)(i)(A).

Once FDA has received an ANDA for review,³ an applicant that submitted a paragraph IV certification to a listed patent must provide the NDA holder and each patent owner notice of its paragraph IV certification, including a description of the legal and factual basis for the ANDA applicant's assertion that the patent is invalid, unenforceable, or will not be infringed.⁴ If a patent is listed at the time an original ANDA is submitted and, in response to a notice of a paragraph IV certification, the NDA holder or patent owner initiates a patent infringement action against the ANDA applicant within 45 days of receiving the required notice, approval of the ANDA generally will be stayed for 30 months from the later of the date of receipt of the notice by any owner of the patent or the NDA holder or such shorter or longer time as the court might order.⁵

FDA が審査のために ANDA を受け取ると³、リストに掲載された特許に関するパラグラフIV証明を FDA に提出した ANDA 申請者は、NDA 所有者と各特許所有者に、当該特許が無効であるか、行使不可能である、または侵害されないという ANDA 申請者の主張の法的小よび事実上の根拠の説明を含めてパラグラフIV証明について通知する必要がある⁴。

オリジナルの ANDA が提出されたときに特許がリストされており、要求される通知の受領後 45 日以内に、パラグラフIV証明の通知に対して、NDA 保有者または特許所有者が ANDA 申請者に対して特許侵害訴訟を提起した場合、ANDA の承認は通常、特許の所有者または NDA 保有者が通知を受領した日から 30 か月間または裁判所の命令が発せられるまでは行われない⁵。

FDA may issue final approval to an ANDA at the conclusion of a 30-month stay if a patent infringement lawsuit about the drug product at issue in that ANDA is pending, the ANDA does not contain any paragraph III certifications, the ANDA is not blocked by any unexpired exclusivities, and all other requirements for approval have been met. However, after the ANDA is approved, the NDA holder or patent owner may be successful in its patent infringement lawsuit against the ANDA holder. In such a case, the district court may order that the patent is infringed and that the approval of the ANDA is not effective before expiration of the infringed patent pursuant to 35 U.S.C. 271(e)(4)(A).

Under these circumstances, FDA must determine whether it is appropriate to convert the approval status of the ANDA to TA⁶ and, if that conversion is appropriate, the timing of such conversion. In order for FDA to timely convert the approval status of an ANDA to TA, ANDA applicants are required to submit any and all documents pursuant to 21 CFR 314.107(e) within 14 days of the date of entry by the court or the date of appeal or expiration of the time for appeal.⁷

ANDA で問題となっている医薬品に関する特許侵害訴訟が係属中であり、ANDA にパラグラフ III 証明が含まれておらず、ANDA が期限が切れていない独占権によって阻害されず、および承認のための他のすべての要件が満たされている場合、FDA は 30 か月のペンディング期間の終了時に ANDA に最終承認を発行することができる。ただし、ANDA が承認された後、NDA 保有者または特許所有者が、ANDA 保有者に対する特許侵害訴訟で勝訴する可能性がある。このような場合、地方裁判所は、特許が侵害されていること、および ANDA の承認が 35 U.S.C. 271(e)(4)(A)に従って侵

³ 21 CFR 314.101(b).

⁴ Section 505(j)(2)(B) of the FD&C Act.

⁵ Section 505(j)(5)(B)(iii) of the FD&C Act and 21 CFR 314.107(b)(3)(i).

害された特許の有効期限が切れる前に有効にならないように命じることができる。

このような状況においては、FDAはANDAの承認ステータスをTA⁶に変更することが適切かどうか、およびその変更が適切な場合は当該変更のタイミングを決定する必要がある。

FDAがANDAの承認ステータスをタイムリーにTAに変更するために、ANDAの申請者は、21 CFR 314.107 (e) に従い、裁判所による申立の日または控訴日または控訴の期限切れの時から14日以内にすべての書類を提出する必要がある⁷。

POLICY (ポリシー)

FDA considers certain factors when determining whether it is appropriate to convert the approval status of an approved ANDA to TA. Upon receipt of a federal district court judgment that the patent is infringed and the approval of the ANDA is not effective before expiration of the infringed patent, as described in 35 U.S.C. 271(e)(4)(A), FDA will consider the judgment and will also consider any documents showing (1) that the district court judgment has been stayed or (2) that there is a pending motion for stay of the district court judgment.

FDAは、承認されたANDAの承認ステータスをTAに変更することが適切かどうかを判断するときに、特定の要因を考慮する。35 U.S.C. 271(e)(4)(A)に記載されているように、特許が侵害されているという連邦地方裁判所の判決がなされた場合、ANDAの承認は侵害された特許の有効期限が切れるまで有効とはならない。FDAは判決及び(1) 地方裁判所の判決が留まったこと、または(2) 地方裁判所の留保の係属中の申立てがあることを示す文書を検討する。

⁶ 21 CFR 314.107(g) (“If FDA issues an approval letter in error or a court enters an order requiring, in the case of an already approved 505(b)(2) application or ANDA, that the date of approval be delayed, FDA will convert the approval to a tentative approval if appropriate.”).

⁷ 21 CFR 314.107(e).

RESPONSIBILITIES (責任)

● Office of Generic Drug Policy Patent and Exclusivity Team

- Receives information from the Office of Regulatory Operations (ORO) Division of Project Management (DPM) concerning patent infringement lawsuits, which may be submitted by electronic submission by the ANDA holder or by paper submission by or on behalf of the NDA holder;
- Verifies that any patents subject to the patent infringement lawsuits are listed in the Orange Book
- Assesses court orders and motions for stay related to patent infringement lawsuits to determine whether it is appropriate to convert the approval status of an ANDA from final approval to TA
- Notifies the Deputy Director of the Office of Generic Drug Policy (OGDP) Division of Legal and Regulatory Support (DLRS) when information concerning patent infringement lawsuits is received, provides the Deputy Director the information received on the status of the patent infringement lawsuits (e.g., copies of the order issued by the district court that finds patent infringement pursuant to 35 U.S.C. 271(e)(4)(A)), and provides a recommendation of approval status to the Deputy Director
- Drafts “Conversion to ANDA Tentative Approval” letters

- ANDA 保有者による電子提出、または NDA 保有者によるまたは NDA 保有者に代わる紙による提出により提出される可能性のある特許侵害訴訟に関して、規制管理局 (ORO) プロジェクト管理部門 (DPM) から情報を受け取る。
- 特許侵害訴訟の対象となっている特許がオレンジブックに掲載されていることを確認する。
- ANDA の承認ステータスを最終承認から TA に変更することが適切かどうかを判断するために、特許侵害訴訟に関連する滞在の裁判所命令と動議を評価する。
- 特許侵害訴訟に関する情報が受け取られたときにジェネリック医薬品政策局 (OGDP) 法規制サポート部門 (DLRS) の副局長に通知し、特許侵害訴訟の状況について受け取った情報を副局長に提供する (例: 35 U.S.C. 271(e)(4)(A)) に従って特許侵害を認定し、副局長に承認ステータスの推奨を提供する地方裁判所が発行した命令のコピー)
- 「ANDA 暫定承認への変更」レターをドラフトする

● OGDG DLRS Deputy Director (or designee)

- Reviews assessment conducted by the Patent and Exclusivity Team (PET) for concurrence
- Reviews “Conversion to ANDA Tentative Approval” letter
- 同意を得るために特許独占権チーム (PET) が実施した評価のレビューを行う
- 「ANDA 暫定承認への変更」レターのレビューを行う

● DLRS Director (or designee)

- Performs a secondary review of the “Conversion to ANDA Tentative Approval” letter
- 「ANDA 暫定承認への変更」レターの 2 次レビューを行う

● **ORO DPM**

- Sends information received electronically about a patent infringement lawsuit to PET
 - Issues the “Conversion to ANDA Tentative Approval” letter
 - Updates the status of the ANDA that has undergone conversion from final approval to TA status in the relevant FDA databases and informs the Orange Book staff of the conversion
- 特許侵害訴訟について電子的に受け取った情報を PET に送付する
 - 「ANDA 暫定承認への変更」レターを発行する
 - 関連する FDA データベースで、最終承認から TA ステータスに変更された ANDA のステータスを更新し、オレンジブックスタッフに変更について通知する。

● **ORO Immediate Office**

- Provides the final signature on the “Conversion to ANDA Tentative Approval” letter
- 「ANDA 暫定承認への変更」レターの最終署名を提供する

PROCEDURES

手順

● **Receipt of court decisions or other documents related to patent infringement lawsuits (裁判所の決定または特許侵害訴訟に関連するその他の文書の受領)**

Submissions related to patent infringement lawsuits may be submitted by or on behalf of either (1) the patent owner or NDA holder or (2) the ANDA holder. Depending on the submitter, the document may be received in electronic or paper format to the ANDA or to an office or person at FDA.

特許侵害訴訟に関連する申請は、(1) 特許所有者または NDA 保有者、または (2) ANDA 保有者のいずれかによって、またはその代理として申請される。申請者に応じて、文書は電子形式または紙形式で ANDA または FDA の事務所や担当者へ送付される。

● **Triage of submissions related to patent infringement lawsuits (特許侵害訴訟に関連する提出のトリアージ)**

All submissions related to patent infringement lawsuits are reviewed by the PET. If the submission is made by the ANDA holder and submitted electronically to its application, the regulatory project manager in the ORO DPM notifies the PET and provides either an electronic link to or an electronic copy of the submission. If the submission is made by the NDA holder or patent owner to the Office of Generic Drugs on paper, the submission is forwarded to the PET by the document room. The PET will verify that any patents identified in the lawsuit are listed in the Orange Book.

特許侵害訴訟に関連するすべての提出物は、PET によって審査される。申請が ANDA 保有者によって行われ、当該申請が電子的になされた場合、ORO DPM の規制プロジェクトマネージャーは PET に通知し、申請への電子リンクまたは電子コピーのいずれかを提供する。申請が NDA 保有者または特許所有者によって書面でジェネリック医薬品局に行われた場合、申請はドキュメントルームによって PET に転送される。PET は、訴訟で特定された特許がオレンジブックに掲載されていることを確認する。

- **Decision to convert the approval status of an ANDA from final approval to TA (ANDA の承認ステータスを最終承認から TA に変更する決定)**

Once the verification process is complete, the PET provides the submitted documentation and verification results to the DLRS Deputy Director and makes a recommendation regarding approval status. The DLRS Deputy Director determines whether it is appropriate to convert by considering the factors noted in the Policy section above. If the approval status of the ANDA is converted from final approval to TA, any unapproved supplements and/or annual report changes submitted to the ANDA will be considered withdrawn.

検証プロセスが完了すると、PETは提出された文書と検証結果をDLRSの副局長に提供し、承認ステータスに関する推奨を行う。DLRSの副局長は、上記の「ポリシー」セクションに記載されている要素を考慮して、変更することが適切かどうかを判断する。ANDAの承認ステータスが最終承認からTAに変更された場合、ANDAに送信された未承認の補足や年次レポートの変更は撤回されたものと見なされる。

- **Creation and issuance of “Conversion to ANDA Tentative Approval” letter (「ANDA 暫定承認への変更」レターの作成と発行)**

If the DLRS Deputy Director determines that conversion is appropriate, the PET drafts the “Conversion to ANDA Tentative Approval” letter to the ANDA holder and sends the letter to the DLRS Deputy Director for review. Upon completion of the DLRS Deputy Director review, the DLRS Director performs a secondary review of the “Conversion to ANDA Tentative Approval” letter. Upon approval of the conversion letter by the DLRS Director, the PET provides the final letter to the appropriate regulatory project manager in the ORO DPM. The letter is generally transmitted for review and clearance by email. Once the letter is signed by the ORO Immediate Office, the “Conversion to ANDA Tentative Approval” letter is issued to the ANDA holder by the ORO DPM. The ORO DPM will update the status of an ANDA that has undergone conversion from final approval to TA status in the relevant FDA databases and inform the Orange Book staff of the conversion so that the application can be removed from the Orange Book.

DLRSの副局長が変更が適切であると判断した場合、PETはANDA保有者に対する「ANDA 暫定承認への変更」レターをドラフトし、DLRSの副局長にレビューのために送付する。DLRSディレクターのレビューが完了すると、DLRSディレクターは「ANDA 暫定承認への変更」レターの2次レビューを行う。DLRSディレクターによる変更レターの承認後、PETはORO DPMの適切な規制プロジェクトマネージャーに最終レターを提供する。レターは通常、レビューとクリアランスのために電子メールで送信される。レターがORO Immediate オフィスによって署名されると、「ANDA 暫定承認への変更」レターがORO DPMによってANDA保有者に対して発行される。ORO DPMは、関連するFDAデータベースで最終承認からTAステータスに変更されたANDAのステータスを更新し、オレンジブックスタッフに変更を通知して、アプリケーションをオレンジブックから削除できるようにする。

DEFINITIONS (定義)

- **Tentative approval:** notification that an NDA or ANDA otherwise meets the requirements for approval under the FD&C Act, but cannot be approved because there is a 7-year period of orphan exclusivity for a listed drug under section 527 of the FD&C Act and § 316.31 of this chapter, or that a 505(b)(2) application or ANDA otherwise meets the requirements for approval under the FD&C Act, but cannot be approved until the conditions in § 314.107(b)(1)(iii), (b)(3), or (c) are met; because there is a period of exclusivity for the listed drug under § 314.108; because there is a period of pediatric exclusivity for the listed drug under section 505A of the FD&C Act; because there is a period of exclusivity for the listed drug under section 505E of the FD&C Act; or because a court order pursuant to 35 U.S.C. 271(e)(4)(A) orders that the NDA or ANDA may be approved no earlier than the date specified. A drug product that is granted tentative approval is not an approved drug and will not be approved until FDA issues an approval letter after any necessary additional review of the NDA or ANDA.⁸

- **暫定承認:** NDA または ANDA が FD&C 法に基づく承認の要件を満たしているが、FD&C 法のセクション 527 および本章の § 316.31 に記載されている医薬品には 7 年間のオーファン独占権があるため、承認できない、または、505(b)(2)申請または ANDA が FD&C 法に基づく承認の要件を満たしているが、314.107(b)(1)(iii)、(b)(3) または(c)の条件が満たされるまで承認できない。 § 314.108 に基づいてリストされた医薬品には独占権の期間があるため。FD&C 法の § 505A の下にリストされている薬物には小児独占期間があるため; FD&C 法のセクション 505E に基づいてリストされた医薬品には独占権の期間があるため。または、U.S.C. 35 U.S.C. 271(e)(4)(A)に基づく裁判所命令は、NDA または ANDA が指定された日付より前に承認されないように命じる。暫定的な承認が与えられた医薬品は承認された医薬品ではなく、FDA が NDA または ANDA の必要な追加審査を経て承認書を発行するまで承認されない⁸。

EFFECTIVE DATE (発効日)

This MAPP is effective upon date of publication.
 本 MAPP は、発行日から有効。

CHANGE CONTROL TABLE (改訂管理表)

Effective Date	Revision Number	Revisions
6/11/2020	Initial	N/A

⁸ 21 CFR 314.3(b).

ATTACHMENT 1

ANDA #####

**CONVERSION TO ANDA
TENTATIVE APPROVAL**

APPLICANT NAME
ADDRESS

Attention: CONTACT NAME
TITLE

Dear CONTACT:

This is in reference to your abbreviated new drug application (ANDA) for ESTABLISHED NAME DOSAGE FORM AND STRENGTH(S), approved on APPROVAL DATE pursuant to section 505(j) of the Federal Food, Drug, and Cosmetic Act (FD&C Act). We are writing to inform you that, consistent with the COURT DECISION issued on DATE OF COURT ORDER, the Agency hereby converts the final approval of ANDA ##### to a tentative approval, and considers that this conversion occurred on the date of that decision. Thus, FDA regards ANDA #####, as having been tentatively approved as of DATE OF COURT ORDER. This action conforms the ANDA's status to the court's final judgement, as described in detail below.

The reference listed drug (RLD) upon which your ANDA is based, PROPRIETARY DRUG NAME DOSAGE FORM AND STRENGTH(S), of RLD HOLDER, is subject to periods of patent protection. The following patents and expiration dates are listed in the Agency's publication titled *Approved Drug Products with Therapeutic Equivalence Evaluations* (the "Orange Book"):

U.S. Patent No.

Expiration Date

Your ANDA contained a paragraph IV certification(s) under section 505(j)(2)(A)(vii)(IV) of the FD&C Act stating that the PATENT #(s) patent(s) is/are invalid, unenforceable, or will not be infringed by your manufacture, use, or sale of ESTABLISHED NAME DOSAGE FORM AND STRENGTH(S), under this ANDA.ⁱ As noted in your APPROVAL DATE approval letter, you notified the Agency that ANDA HOLDER complied with the requirements of section 505(j)(2)(B) of the FD&C Act, and that litigation was initiated against ANDA HOLDER for infringement of the PATENT #(s) patent(s) within the statutory 45-day period in the COURT [INCLUDE COURT CASE NAME Plaintiff v. Defendant, Civil Action No. CASE #].

INCLUDE HISTORY OF COURT CASEⁱⁱ

Section 505(j) of the FD&C Act does not expressly provide for a change in approval status of an approved ANDA when the patent litigation results in a finding that one or more listed patents is infringed; however, when a court orders that the approval of an ANDA is not effective before a certain date pursuant to 35 U.S.C. 271(e)(4)(A), FDA may convert an approved ANDA to tentative approval status to reflect the court's order.ⁱⁱⁱ Further, pursuant to 21 CFR 314.107(g), if a court enters an order requiring, in the case of an already approved ANDA, that the date of approval be delayed, FDA will convert the approval to a tentative approval, if appropriate.

Therefore, after consideration of the **COURT'S ORDER DATE ORDER** that the date of approval for ANDA **#####**, is not effective before **DATE OF PATENT EXPIRATION**, FDA is converting the **APPROVAL DATE**, final approval of **ANDA HOLDER'S ANDA #####** for **ESTABLISHED NAME DOSAGE FORM AND STRENGTHS(S)**, to a tentative approval.

Please be aware that any approved supplemental ANDAs filed to this ANDA since **APPROVAL DATE** are considered tentatively approved. Any unapproved supplemental ANDAs and any Annual Report changes filed to the ANDA are considered WITHDRAWN and should be resubmitted in full as either a "MINOR/MAJOR Amendment to the Original," or as a new supplemental ANDA once final approval has been obtained again. We note that this is an administrative conversion only and does not reflect any review of the ANDA subsequent to its original approval or approval of any supplements.

Please note that if FDA requires a Risk Evaluation and Mitigation Strategy (REMS) for a listed drug, an ANDA citing that listed drug also will be required to have a REMS. See section 505-1(i) of the FD&C Act. **[DELETE REMS SENTENCE IF ANDA HAS REMS OR IF OTC ANDA]**

RISK EVALUATION AND MITIGATION STRATEGY (REMS) REQUIREMENTS (if applicable)**Send to REMS Coordinator Team for language****RESUBMISSION**

To request final approval, please submit an amendment titled "FINAL APPROVAL REQUESTED" with enough time to permit FDA review prior to the date you believe that your ANDA will be eligible for final approval. A request for final approval that contains no new data, information, or other changes to the ANDA generally requires a period of 90 days for Agency review. Accordingly,

such a request for final approval should be submitted no later than 90 days prior to the date on which you seek approval. A request for final approval that contains substantive changes to the ANDA or changes in the status of the manufacturing and testing facilities' compliance with CGMPs will be classified and reviewed according to OGD policy in effect at the time of receipt. Applicants should review available Agency guidance for industry related to amendments under the generic drug user fee program to determine the duration of Agency review needed to review the changes submitted. The submission of multiple amendments prior to final approval may also result in a delay in the issuance of the final approval letter.

The amendment requesting final approval should provide the legal/regulatory basis for your request for final approval and should include a copy of the court decision, settlement or licensing agreement, or other information described in 21 CFR 314.107, as appropriate. It should also identify changes, if any, in the conditions under which the ANDA was tentatively approved, e.g., updated information such as final-printed labeling, chemistry, manufacturing, and controls data, as appropriate. This amendment should be submitted even if none of these changes were made, and it should be designated clearly in your cover letter as a "FINAL APPROVAL REQUESTED."

In addition to the amendment requested above, the Agency may request, at any time prior to the date of final approval, that you submit an additional amendment containing information as specified by the Agency. Failure to submit either or, if requested, both types of amendments described above may result in a delay in the issuance of the final approval letter.

This drug product may not be marketed without final Agency approval under section 505(j) of the FD&C Act. The introduction or delivery for introduction into interstate commerce of this drug product before the final approval date is prohibited under section 301 of the FD&C Act. Also, until the Agency issues the final approval letter, this drug product will not be deemed approved for marketing under section 505(j) of the FD&C Act, and will not be listed in the Orange Book. Should you believe that there are grounds for issuing the final approval letter prior to **DATE OF PATENT EXPIRATION**, you should amend your ANDA accordingly.

Annual Facility Fees

The Generic Drug User Fee Amendments of 2012 (GDUFA) (Public Law 112-144, Title III) established certain provisions^{iv} with respect to self-identification of facilities and payment of annual facility fees. Your ANDA identifies at least one facility that is subject to the self-identification requirement and payment of an annual facility fee. Self-identification must occur by June 1st of each year for the next fiscal year. Facility fees must be paid each year by the date specified in the *Federal Register* notice announcing facility fee amounts.

All finished dose forms (FDFs) or active pharmaceutical ingredients (APIs) manufactured in a facility that has not met its obligations to self-identify or to pay fees when they are due will be deemed misbranded. This means that it will be a violation of federal law to ship these products in interstate commerce or to import them into the United States. Such violations can result in prosecution of those responsible, injunctions, or seizures of misbranded products. Products misbranded because of failure to self-identify or pay facility fees are subject to being denied entry into the United States.

In addition, we note that GDUFA requires that certain non-manufacturing sites and organizations listed in generic drug submissions comply with the self-identification requirement. The failure of any facility, site, or organization to comply with its obligation to self-identify and/or to pay fees when due may raise significant concerns about that site or organization and is a factor that may increase the likelihood of a site inspection prior to approval. FDA does not expect to give priority to completion of inspections that are required simply because facilities, sites, or organizations fail to comply with the law requiring self-identification or fee payment.

For further information on the status of this ANDA, or prior to submitting additional amendments, please contact **NAME**, Regulatory Project Manager, at **(XXX) XXX-XXXX**.

Sincerely,

{See appended electronic signature page}

NAME
Director/Deputy Director
Office of Regulatory
Operations
Office of Generic Drugs
Center for Drug Evaluation
and Research

ⁱ The Agency notes the X patent(s) were submitted to the Agency after the submission of your ANDA. Litigation, if any, with respect to these patents would not create a statutory stay of approval.

ⁱⁱ *List relevant court decision(s), as appropriate*

ⁱⁱⁱ *Mylan Laboratories, Inc. v. Thompson, 389 F.3d 1272, 1281-82 (D.C. Cir. 2004).*

^{iv} Some of these provisions were amended by the Generic Drug User Fee Amendments of 2017 (GDUFA II) (Public Law 115-52, Title III).