



GDUFA III Redesigned Pre-Submission (PSUB) Meetings

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A Deep Dive: GDUFA III Scientific Meetings – May 15, 2023

Learning Objectives

- Clarify the purpose of the redesigned pre-submission (PSUB) meetings under GDUFA III
- Describe new GDUFA III timeline and changes to overall meeting goal and process
- Identify what, when, where, and how to submit your PSUB meeting request

Goals of Pre-ANDA Program

- Clarify regulatory expectations for prospective applicants early in product development
- Assist applicants in developing more complete submissions
- Promote a more efficient and effective ANDA assessment process
- Reduce the number of assessment cycles needed to obtain ANDA approval

Pre-ANDA Meetings

Pre-ANDA meetings were introduced in GDUFA II to facilitate pre-submission communications between the FDA and a prospective applicant related to a ***complex*** product and/or complicated drug development questions

- Product Development (PDEV) – No change from GDUFA II
- Pre-submission (PSUB) – *Redesigned for GDUFA III*

Purpose of Redesigned PSUB Meetings

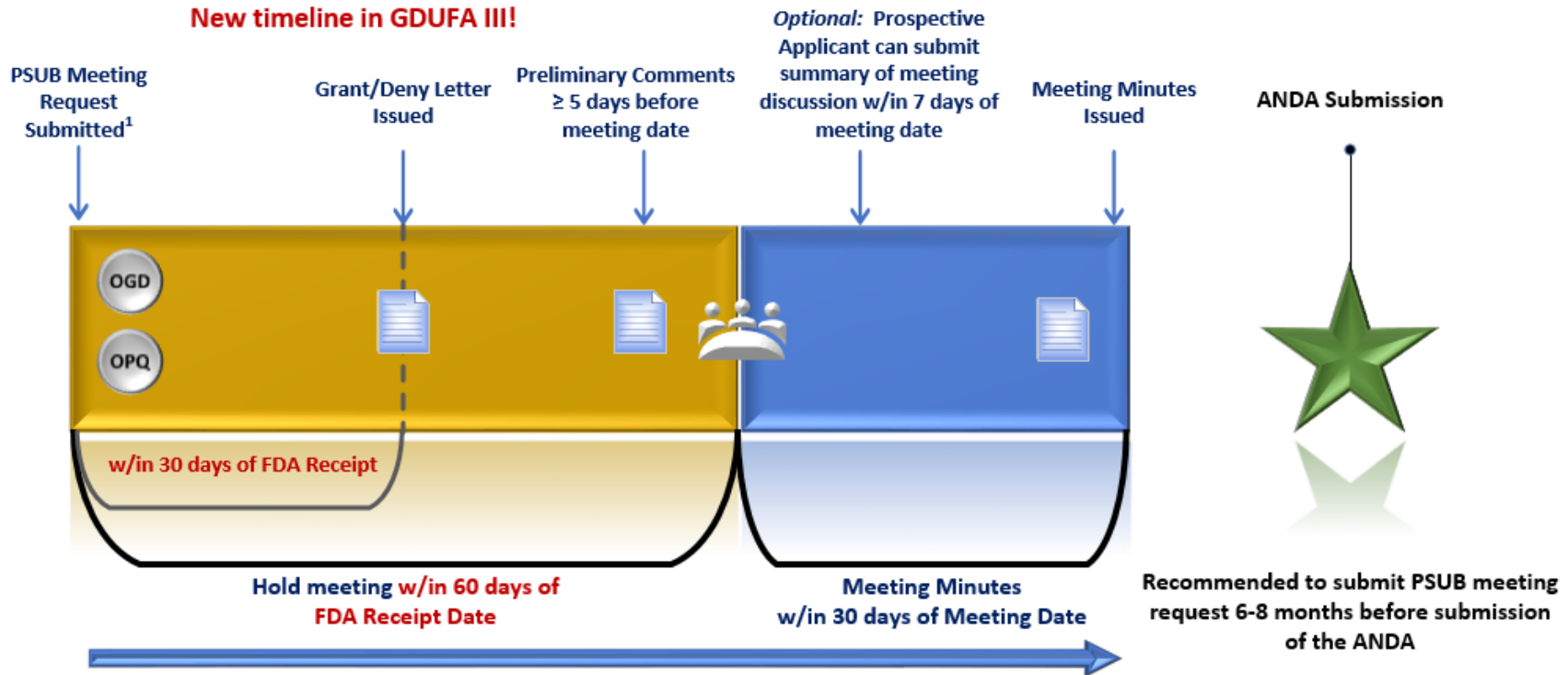
- Provides a prospective applicant the opportunity to present *unique* or *novel* data or information that will be included in the ANDA submission, such as
 - Formulation
 - Key studies
 - Justifications
 - Methods used in product development
 - Interrelationship of the data and information in the ANDA
- ***DOES NOT*** include substantive assessment of summary data or full study reports
- ***IS NOT*** an opportunity to determine whether the ANDA is acceptable for receipt

PSUB Meeting Eligibility

✓ FDA *will* grant a PSUB:

- If prospective applicant was granted a prior product development (PDEV) meeting for the same complex generic product
- If FDA believes in its sole discretion that a PSUB meeting would improve assessment efficiency
- Prospective ANDA applicants may request a PSUB meeting whether they had a PDEV meeting or not

PSUB Meeting Timeline



¹Day 0 = FDA Receipt Date
Days = calendar days

PSUB Meeting Package



- Refer to the “Formal Meeting Guidance” before submitting
- Submit concurrently with the PSUB meeting request
- In general, high-level information is sufficient
- In general, ***should not*** include questions

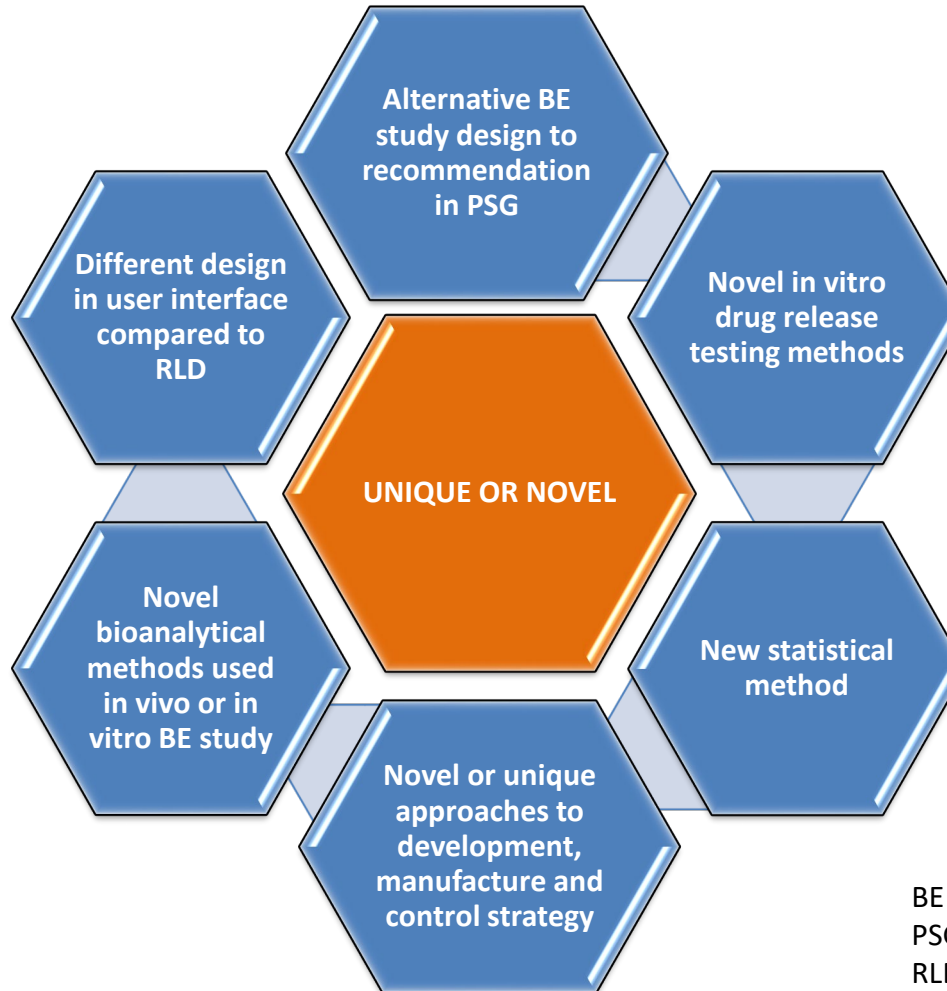
“Formal Meeting Guidance”: Guidance for industry [Formal Meetings Between FDA and ANDA Applicants of Complex Products Under GDUFA Guidance for Industry](#)

PSUB Meeting Package



- Meeting package should include the following, among other items:
 - If there were any prior PDEV meetings for the same proposed complex generic product
 - Event IDs for the prior PDEV meeting(s)
 - Summary of the advice provided from PDEV meeting(s)
 - *If no prior PDEV meeting(s) were held*, an explanation for why a PSUB meeting should be granted
 - Estimated timeline for submission of the ANDA
 - Unique or novel data or information to be included in the ANDA submission

PSUB Meeting Package



BE = Bioequivalence

PSG = Product-specific guidance

RLD = Reference listed drug

PSUB Meeting Package



- PSUB meeting package can be submitted in the format of a draft meeting presentation
 - For a suggested presentation outline template with recommendations on information that should be included, see **Appendix B** of the “Formal Meetings Guidance”

Refer to guidance for industry [*Formal Meetings Between FDA and ANDA Applicants of Complex Products Under GDUFA Guidance for Industry*](#)

Contains Nonbinding Recommendations

B. Pre-Submission Meeting Presentation Outline Template for Prospective ANDA Applicants

The pre-submission meeting presentation outline template provided below is intended to assist prospective ANDA applicants in preparing pre-submission meeting presentations, and it includes suggested items from the Agency for prospective ANDA applicants to present at the pre-submission meetings to help orient the discussion. Suggested items for the pre-submission meeting presentation include, but are not limited to: (1) formulation; (2) new analytical methods; (3) new statistical methods; (4) novel in vitro drug release testing methods; (5) alternative bioequivalence study design to the recommendations in the product-specific guidance with justification for the alternative study design; (6) regulatory history; and (7) summary of generic development.

Prospective ANDA applicants should address the suggested items, as applicable, and provide responses/information as appropriate in a concise and clear manner.

Note that the information included below is not an exhaustive list of the information that prospective ANDA applicants should consider including in their pre-submission meeting presentation. There may be additional items that should be included in the pre-submission meeting presentation.

Presentation Outline Template:

1. Pre-Submission Meeting Request Summary
 - a. Applicant name
 - b. Anticipated ANDA submission date
 - c. Reference Listed Drug (RLD)
 - i. Information on drug substance, dosage form, route of administration
 - ii. RLD information (RLD number, approval date, application holder)
 - iii. Indication(s)
 - iv. Dose and route of administration
 - d. Reference Standard
 - i. Indicate if the Reference Standard is the same as the RLD
 - ii. When the Reference Standard is different from the RLD, include the Reference Standard information (application number, approval date, application holder)
 - e. Complex drug as defined by the GDUFA III commitment letter (indicate all that apply)
 - i. Complex active ingredient
 - ii. Complex formulation
 - iii. Complex route of delivery
 - iv. Complex dosage form

Submitting Your Meeting Request



- Obtain a pre-assigned ANDA number if no prior PDEV meeting
<https://www.fda.gov/Drugs/DevelopmentApprovalProcess/FormsSubmissionRequirements/ElectronicSubmissions/ucm114027.htm>
- Submit via the CDER Direct NextGen Collaboration Portal

<https://edm.fda.gov>



Submitting Your Meeting Request



ANDA Information

* What is the type for this Pre-ANDA Meeting Request?
ANDA Pre-Submission Meeting

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ANDA Pre-Submission Meeting

Pre-ANDA Product Development – Discuss new or alternative approaches to demonstrating equivalence early in product development

ANDA Presubmission Meeting – Discuss the content and format of unique, novel or complex components of an upcoming ANDA submission

* Has the ANDA for which you are submitting a Meeting Request been granted a Competitive Generic Therapy Designation?
 Yes
 No

* Has a Product Development Meeting been held between you and FDA?
 Yes
 No

* What is the Event ID of the previously held product development meeting?
[Text input field]

* What is the Event ID of the previously held product development meeting?
[Text input field]

* Has a Product Development Meeting been held between you and FDA?
 Yes
 No

* What is the Event ID of the previously held product development meeting?
[Text input field]

Within 30 Days of Receipt



- Parallel grant/deny assessments performed by OGD and OPQ
 - Will reach one unified decision
- If granting, FDA will:
 - Identify ANDA assessment team members who will attend the PSUB meeting
 - In the grant letter, identify additional topics that prospective applicant should address during the meeting, when applicable
- If denying, FDA will provide the reason for denial
- By Day 30, issue grant/deny letter

Your Meeting Was Granted



- PSUB offered as in-person face-to-face (FTF) or videoconference meetings
 - As of March 27, 2023, prospective applicants can request in-person FTF meetings for PDEV or PSUB meetings
- If your request meets the criteria in the GDUFA III commitment letter, FDA will generally grant the requested meeting format
- Format of the meeting and scheduling information will be included in your grant letter
- A project manager (PM) from the Office of Research and Standards (ORS) will be your primary point of contact

Within 60 Days of Receipt

- FDA will generally not issue information requests for PSUB meetings
- Draft presentations may be updated ***up to 21 days*** prior to the meeting so that FDA may provide preliminary comments at least 5 days before the meeting date
 - Final presentations should be submitted ***at least 48 hours prior*** to the scheduled meeting date
- FDA will issue preliminary written comments no later than 5 days prior to the meeting date
 - FDA may indicate there are no comments
- Preliminary comments should not result in cancellation of a PSUB meeting
- By Day 60, PSUB meeting is held

In-Person Meetings



- ***At least 2 weeks prior to the scheduled meeting date*** provide the assigned PM:
 - Name, title, and company affiliation of all meeting participants
 - Relevant information (e.g., country of citizenship, passport information, etc.) for any Foreign National (FN) visitors participating in person
 - Foreign Visitor Data Request Form will be included with the Meeting Information Letter
- ALL visitors must present a valid government-issued photo identification on the day of the meeting
 - FN visitors **must** present the passport that matches information provided to the PM or will be denied entry
 - Lawful Permanent Residents (LPR) of the U.S. **must** present a valid LPR card. Other forms of identification will not be accepted

Meeting Day



- Meetings are typically 60 minutes
- Prospective applicant's presentation will help orient the discussion
- FDA attendees will include:
 - Members of the ANDA assessment team
 - Participants in prior PDEV meetings, if applicable
- At the meeting, FDA will identify items or information that should be clarified before submission of the ANDA

Meeting Minutes

- If a prospective applicant would like FDA to consider their meeting summary:
 - Submit within 7 calendar days of the meeting via the portal
- FDA will issue meeting minutes within 30 calendar days after the meeting date
- FDA-issued minutes are considered the official record of the meeting

Dispute of Meeting Minutes



- A prospective applicant requesting additional clarification of the meeting minutes issued by FDA should contact the assigned FDA point of contact (POC)
- FDA recommends any concerns about the meeting minutes be submitted in writing to FDA within 10 calendar days of receipt

Challenge Question #1

FDA recommends that a pre-submission meeting request be submitted how far in advance of the ANDA submission?

- A. 12 months
- B. 6-8 months
- C. 10-12 months
- D. 2-3 months

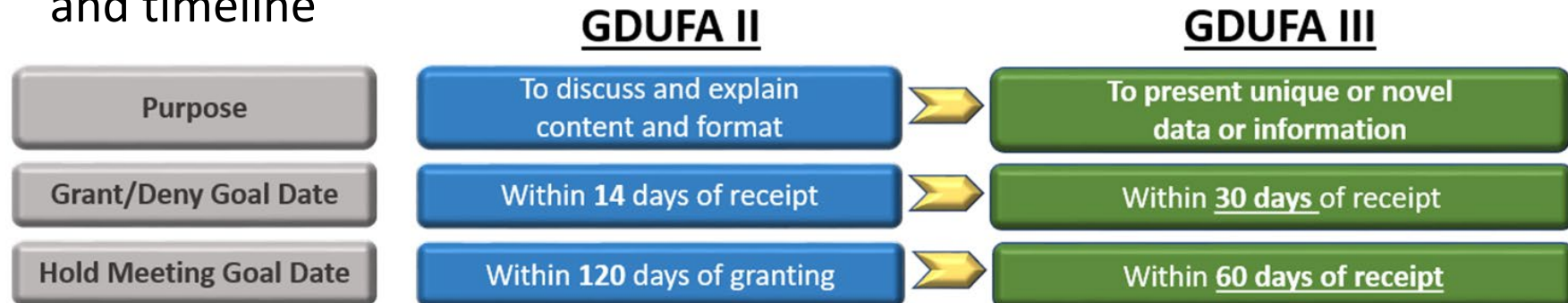
Challenge Question #2

Which of the following statements is NOT true?

- A. In GDUFA III, PSUB meetings provide prospective applicants the opportunity to present unique or novel data or information that will be included in the ANDA submission
- B. For a PSUB meeting, FDA will not provide a substantive assessment of summary data or full study reports
- C. A PSUB meeting will be granted as a teleconference or written response only if requested by the prospective applicants
- D. The meeting package for a PSUB meeting can be submitted in the format of a draft presentation

Summary

- PSUB meeting has been redesigned for GDUFA III with a new scope and timeline



- A request for a PSUB meeting should be made approximately 6-8 months before the ANDA submission
- Seek FDA's input via a PDEV meeting so that FDA has knowledge of your development program at the time of the PSUB meeting

Resources



- [GDUFA Reauthorization Performance Goals And Program: Enhancements Fiscal Years 2023-2027 \(GDUFA III Commitment Letter\)](#)
- Guidance for industry [Formal Meetings Between FDA and ANDA Applicants of Complex Products Under GDUFA](#) (October 2022)
- MAPP 5220.8 (Rev 1): [Evaluating Requests for and Conducting Product Development and Pre-Submission Pre-ANDA Meetings](#) (October 2022)
- Infographic: [GDUFA III – Summary of Teleconferences and Meetings](#)
- MAPP 5240.10: [Classifying Approved New Drug Products as Complex Products for Generic Drug Development Purposes](#) (April 2022)
- Draft guidance for industry [Controlled Correspondence Related to Generic Drug Development](#) (December 2022)
- [GDUFA III Enhancement to the Pre-ANDA Program](#)

