

GDUFA II: Pre-ANDA Meetings for Complex Generic Products

Kris Andre

Associate Director for Regulatory Affairs Office of Research and Standards Office of Generic Drugs (OGD) Center for Drug Evaluation and Research U.S. Food and Drug Administration



Pre-ANDA Program Goals

- Clarify regulatory expectations for prospective applicants early in product development
- Assist applicants to develop more complete submissions
- Promote a more efficient and effective ANDA assessment process
- Reduce the number of review cycles required to obtain ANDA approval, particularly for *complex* products

Complex Products

Complex active pharmaceutic al ingredient (API)	 Any drug product containing a complex API, regardless of administration routes and dosage forms. e.g., Conjugated Estrogen Tablet, Glatiramer Acetate Injection 	
Complex routes of delivery	 Any non-solution drug product with a non-systemic site of action (e.g., topical, ophthalmic, local gastrointestinal (GI) action) e.g., Cyclosporine Emulsion, Acyclovir Cream 	
Complex dosage forms/formu- lations	• Any non-oral complex formulation/dosage form product where there are often two or more discrete states of matter within the formulation e.g., Doxorubicin HCl Liposomes, Leuprolide Acetate for Depot Suspension	
Complex drug- device combinations	 Where the drug constituent part is pre-loaded in a product-specific device constituent part or is specifically cross-labeled for use with a specific device, in which the device design affects drug delivery to the site of action and/or absorption e.g., Epinephrine Injection (autoinjector) 	
Other products	 Any solid oral opioid drug products with FDA approved labeling for that show properties (and thus gaining their labeling) to meaningfully deter drug abuse e.g., Hydrocodone Bitartrate ER Tablet 	
www.fda.gov	Lionberger R. Innovation for Generic Drugs: Science and Research Under the Generic Drug User Fee Amendments of 2012, Clinical Pharmacology & Therapeutics (CPT), 2019, Vol.105(4), p.878-885	

GDUFA II Meetings: Before ANDA Submission

Product Development (PDEV)

- <u>Scientific exchange</u> to discuss specific issues or questions (e.g., a proposed study design, alternative approach, or additional study expectations)
- <u>Targeted advice</u> regarding ongoing ANDA development program

Pre-submission (PSUB)

- Discuss and explain <u>content and</u> <u>format of the ANDA to be submitted</u>
- Advice to <u>enable efficient review</u> and improve chances of first cycle approval
- Does *not* include substantive review of summary data or study reports
- ANDA is anticipated to be submitted ~6 months of meeting date



GDUFA II Meetings: After ANDA Submission

Mid-Review-Cycle Meeting (MRCM)

- For applicants with prior PDEV and/or PSUB meetings
- Generally within 30 days after the mid-point
- Update on status of review and next steps

FDA <u>will</u> grant a PDEV or PSUB meeting for a complex product, if:

- No PSG available
- Proposing an alternative BE approach to the PSG
 - Change in study type (e.g., in vitro instead of in vivo approach)
- Meeting package is complete
- Questions could not be adequately addressed through a controlled correspondence (CC)
- A meeting would significantly improve ANDA review efficiency



Depending on available resources, FDA <u>may</u> grant if, in FDA's judgment:

- Concerns complex product development issues
- Meeting package is complete
- Questions could not be adequately addressed through a CC, and
- A meeting would significantly improve ANDA review efficiency

Submitting Your Meeting Request

FDA

• Obtain a pre-assigned ANDA number

https://www.fda.gov/Drugs/DevelopmentApprovalProcess/FormsSubmissionRequirements/ElectronicSubmissions/ucm114027.htm

• Submit via the CDER Direct NextGen Collaboration Portal

Create Pre-ANDA Meeting Request			
Pre-ANDA Meeting Request Information			
* What is the Pre-assignment Number for this Pre- ANDA Meeting Request? ANDA Select One Select One Abbreviated New Drug Application (ANDA)			
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 Note: Applicants that have requested and received a competitive generic therapy designation under section 506H of the Federal Food, Drug, and Cosmetic Act may select either of these meeting types.			
* What is the type for this Pre-ANDA Meeting Request?			
* Has the ANDA for which you are submitting a Pre- ANDA Meeting Request been granted a Competitive Generic Therapy Designation?	Select One Pre-ANDA Product Development ANDA Presubmission		

Submitting Your Meeting Request

- Meeting package for PDEV
 - Provide specific proposals and questions supported by appropriate data and scientific justification
- Meeting package for PSUB
 - Outline the unique, novel, or complex aspects of your upcoming submission
 - If you have specific questions, provide appropriate background material and data related to those questions

FDA

Meeting Package Format and Content FDA

- Refer to the draft Guidance for Industry (October 2017)
 - Formal Meetings Between FDA and ANDA Applicants of Complex Products Under GDUFA
- Each question is followed by a corresponding justification, rationale or data to support discussion as applicable
- List of questions grouped by discipline (e.g., BE, CMC, etc.)
- Each question clearly numbered (e.g., 1,2,3 without subquestions)

Meeting Request Evaluation

FDA

- Parallel assessments of the meeting request by Office of Generic Drugs (OGD) and Office of Pharmaceutical Quality (OPQ)
 - Assessment team reviews the product details, contents and submitted questions
 - OGD and OPQ coordinate to provide a unified response

My Meeting Was Granted

- FDA
- Typically granted as face-to face meeting, though the applicant can request a written response or teleconference
- Written responses and teleconferences still qualify you for a mid-review-cycle meeting
- A project manager from the Office of Research and Standards (ORS) is assigned as the point of contact

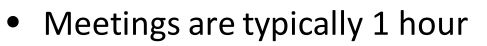
Pre-ANDA Meeting Package Assessment

- FDA staff will review the meeting package, request consults and send information requests (if needed)
- Information Requests (IR)
 - Sent to prospective applicant through the portal
 - FDA strives to send early in the process, but can be sent at any point
 - Applicant responds to the IR through the portal
- Preliminary responses are based upon the Agency's current thinking and knowledge
 - May change with available data or research, etc.

Preliminary Responses

- Preliminary written responses from the FDA will be sent via the portal approximately 5 days before your scheduled meeting
- Your opportunity to focus your meeting
 - Submit presentation materials (not required)
 - Submit a revised agenda
 - Submit these items through the portal <u>at least 48 hours prior to scheduled</u> meeting
- Should <u>NOT</u> generate the submission of new questions
- You can cancel your meeting if you feel the preliminary responses adequately address your questions
 - Still be eligible for a MRCM

Meeting Day



- Discussion should be focused on clarification of the Agency's preliminary written responses
- Meeting participants discuss the data, questions, and the responses provided to assist the prospective ANDA applicant's complex product development program
- FDA <u>will not</u> address or discuss new data or questions not presented in the original meeting package

Post-Meeting



- If prospective ANDA applicants would like the FDA to consider their meeting summary:
 - Submit within 7 calendar days of the meeting via the portal
- FDA will issue official minutes within 30 calendar days of the meeting

Competitive Generic Therapy



- New pathway for drugs with "inadequate generic competition"
- Eligible for PDEV and PSUB meetings
 - Includes both complex and non-complex products
 - Provide documentation of Competitive Generic Therapy (CGT) designation with meeting request
 - Does <u>not</u> provide for an expedited meeting timeline
- FDA will consider the following, among other factors, to determine whether to grant or deny a meeting request with CGT:
 - Complexity of developing an ANDA for a specific drug
 - Potential public health impact (e.g., severity of the condition treated, size of impacted patient population)
 - Impact on FDA resources and other workload commitments

