

FDA Type C Meetings on ISS Safety Analysis Strategy and Related Data Requirements

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Learning Objectives

- Understand the types of formal meetings with FDA
- Describe how Type C ISS meeting can be used to support drug development programs
- Describe how to request a Type C ISS Meeting

Formal Meetings with FDA



What: 6 types of formal meetings under PDUFA



Who: Sponsor/Investigators and FDA



Why: Provide advice on drug/biologics development



When: Critical points in product development

Types of Formal Meetings with FDA

Type A

Type B

Type B (end
of phase)

Type C

Type D

INTERACT

INTERACT: Initial Targeted Engagement for Regulatory Advice on CDER and CBER Products

Types of Formal Meetings with FDA

Type A

- Stalled development e.g., clinical holds, dispute resolutions

Type B & Type B (EOP)

- preIND, preNDA/BLA
- EOP2

Type C

- Any meeting that does not fit in the other categories

Type D

- Focused topic meetings

INTERACT

- Unique challenges in early development

Type C Meetings on ISS Safety Analysis Strategy and Related Data Requirements (Type C ISS Meetings)



Precede the Pre-NDA meeting

- \approx 1 year before pre-NDA meeting
- After drafting analysis plan for ISS
- Before programming work for pooled or other safety analysis



Discuss safety analysis strategy for the ISS and related data requirements

Type C ISS Meetings

- Opportunity to discuss with FDA approach to aggregate safety analysis (e.g., before the pre-NDA meeting)
- Topics:
 - Data pooling for safety analysis
 - Analytic methods
 - Approach to AESI
 - Data standards
 - Submission requirements for datasets

Example Sponsor Questions



- *Does the Agency agree with the proposed pooling strategy?*
- *Does the Agency agree with the proposed Integrated Safety Analysis Plan?*
- *Does the Agency agree with the proposed grouping of MedDRA preferred terms for evaluation of ...*
- *Does the Agency agree with the proposed Study Data Standardization Plan?*
- *Are legacy data acceptable if SDTM and ADaM datasets are not available?*
- *Does the Agency agree with the proposal for narratives and case reports forms (CRFs)?*

Meeting Submission Content



- Tabular listing of clinical trials
- Draft ISS statistical analysis plan (SAP)
 - Pooling strategy
 - Rationale for inclusion or exclusion of trials
 - Planned analytic strategies to manage differences in trial designs (e.g., length, randomization ratio imbalances, study populations)
- Method of assignment of study events to a specific study period
- Previously observed and anticipated safety issues

How to Request a Type C ISS Meeting?

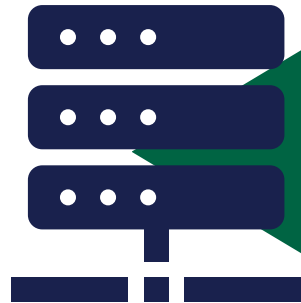


Cover Letter: “Discuss Safety Analysis Strategy for the ISS”

Type C Meeting Formats

- Sponsor may request:
 - In-person
 - Virtual
 - Teleconference
 - Written response only (WRO)
- Agency may determine that WRO is appropriate

How to Submit a Sample Standardized Dataset?



Cover Letter: “Clinical/Safety Sample Datasets”

Sample Standardized Datasets



- Reviewed for conformance to:
 - Standards
 - Structure
 - Format
- Are **NOT** reviewed as part of marketing application

Can safety analysis strategy and data requirements be discussed during the pre-NDA meeting?



Yes, BUT...

Advantages of Type C ISS Meeting



REGULATORY
GUIDANCE



MORE TIME TO PLAN
AND MAKE
REVISIONS



DEDICATED
MEETING ON
IMPORTANT TOPIC

Summary

- Request Type C ISS meetings
- Early discussion with Agency on approach to aggregate safety analysis
- Timing: Before preNDA meeting
- Timing: After drafting ISS SAP

Challenge Question

Which of the following statements regarding Type C ISS Meeting is NOT true?

- A. They are optional
- B. Can be conducted via WRO, in person, or teleconference
- C. Best conducted before the preNDA meeting
- D. Best conducted before drafting the analysis plan for ISS

Resources



- Guidance for Industry: Formal Meetings Between the FDA and Sponsors or Applicants of PDUFA Products (Draft Guidance, September 2023). Available at: <https://www.fda.gov/media/172311/download>
- Study Data Standards Resources. Available at: <https://www.fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources>
- Study Data Technical Conformance Guide - Technical Specifications Document. Available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/study-data-technical-conformance-guide-technical-specifications-document>

