

FDA's Study Data Policy Framework and Recent Activities

Helena Sviglin, OSP Data Standards Staff
Fall 2022



Current Contributions:

- Internal policy development and study data governance
- Chair, FDA Study Data Technical Conformance Guide (sdTCG)
- Chair, FDA Business Rules
- eData responses
- FDA Data Standards Catalog, Chair
- I AM NOT A LAWYER



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Office of Strategic
Programs



FDA Study Data Policy Framework Overview

Providing Regulatory Submissions in
Electronic Format — Submissions Under
Section 745A(a) of the Federal Food,
Drug, and Cosmetic Act

Guidance for Industry

U.S. Department of Health and Human Services
Food and Drug Administration
Center for Drug Evaluation and Research (CDER)
Center for Biologics Evaluation and Research (CBER)

December 2014
Electronic Submissions

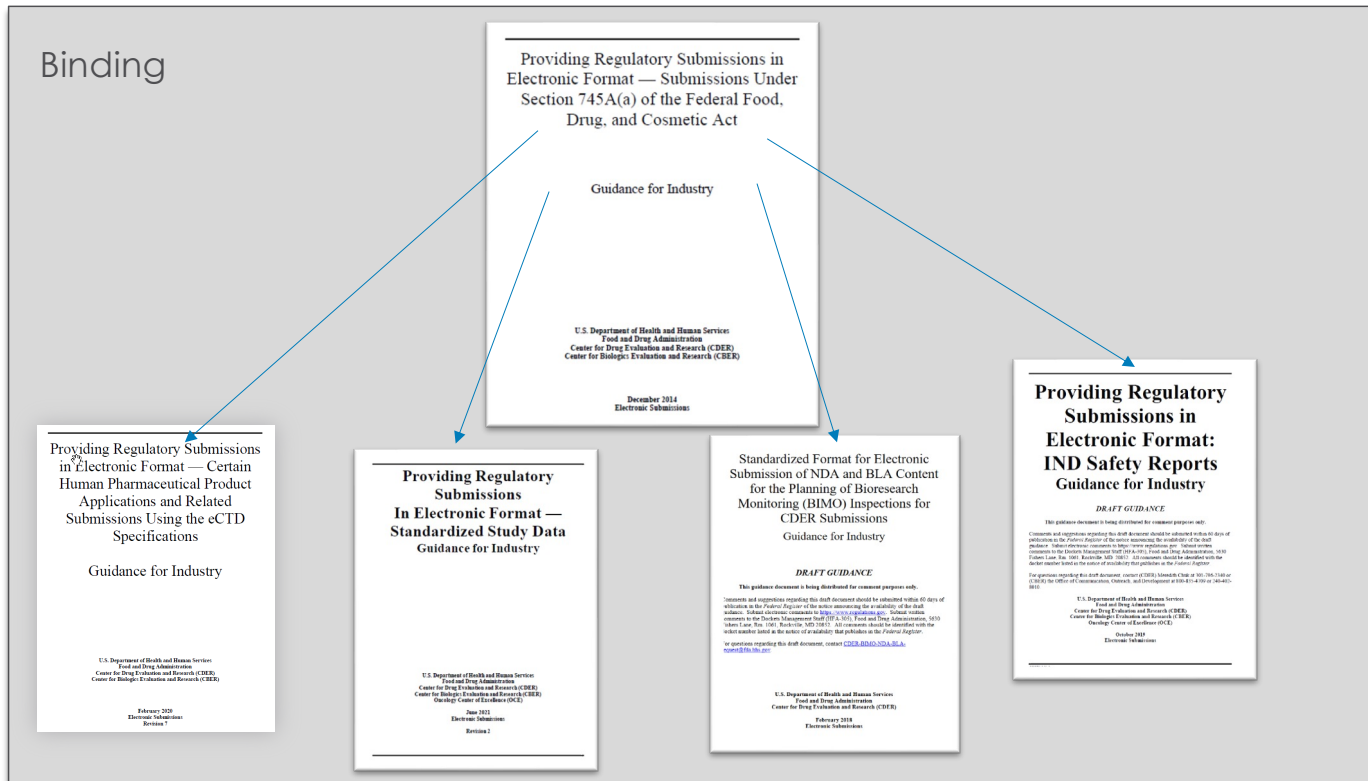
Primary Statute



[Search for FDA Guidance Documents | FDA](https://www.fda.gov/regulatory-information/search-fda-guidance-documents)
([fda.gov/regulatory-information/search-fda-guidance-documents](https://www.fda.gov/regulatory-information/search-fda-guidance-documents))

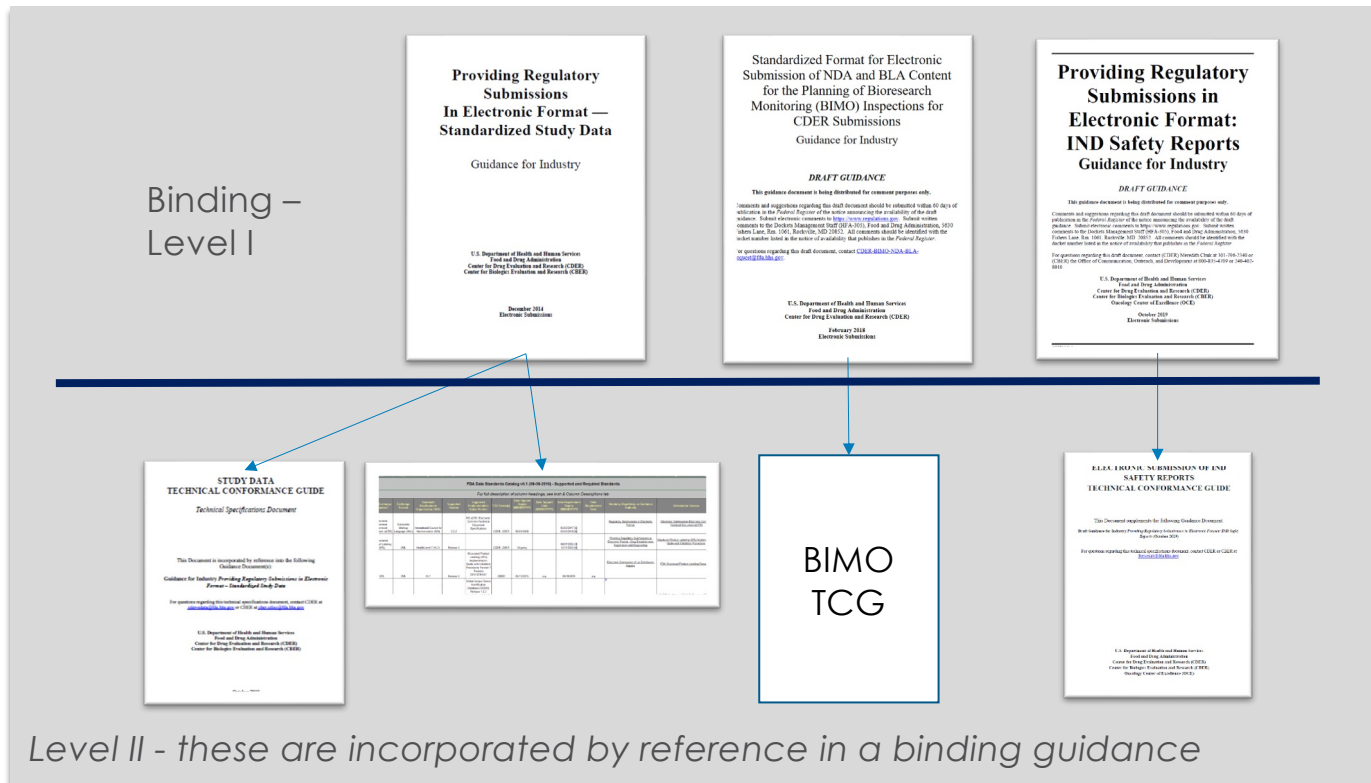
[Study Data Standards Resources | FDA](https://www.fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources)
([fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources](https://www.fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources))

How FDA communicates technical requirements for submitting study data



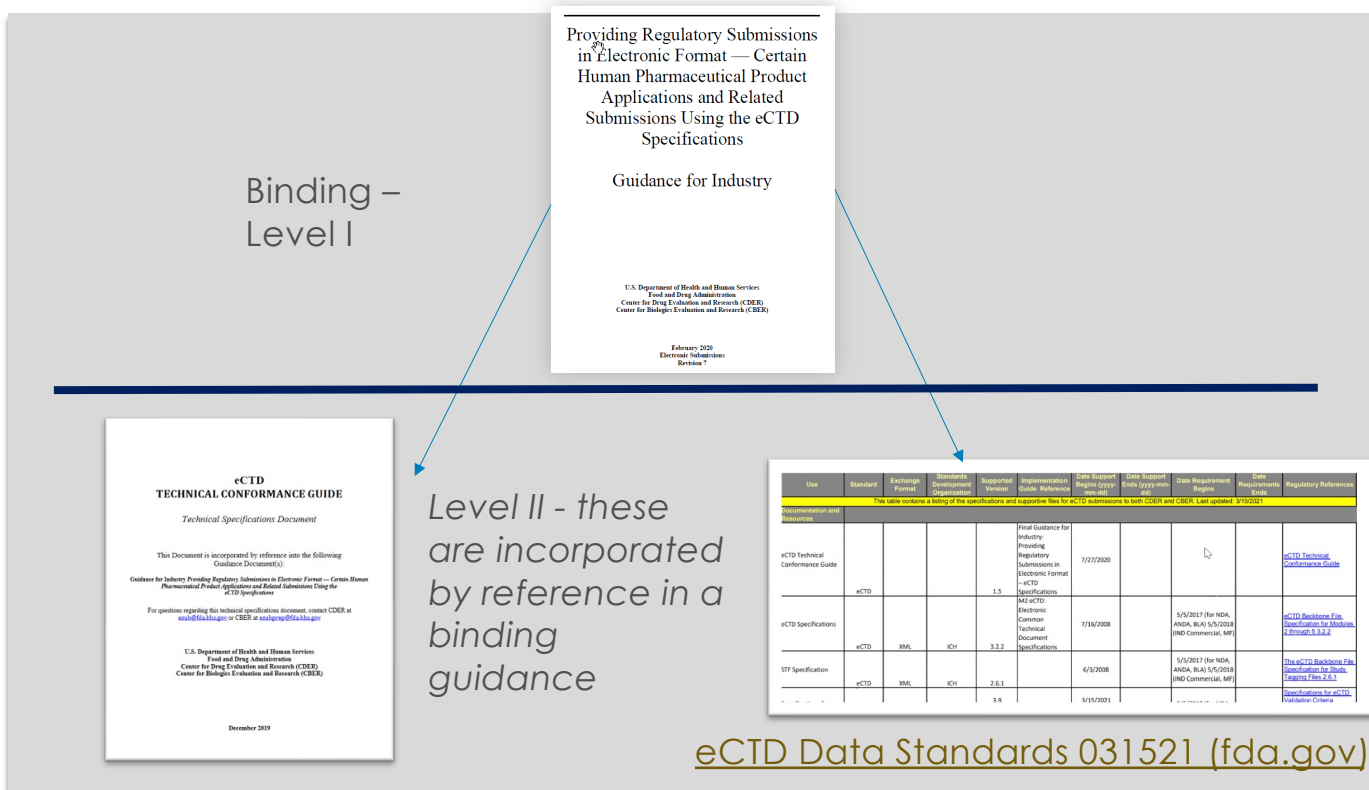
Fall 2022 Svigin

How FDA communicates technical requirements for submitting study data





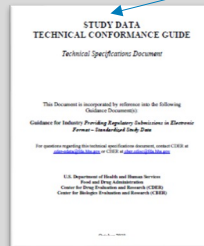
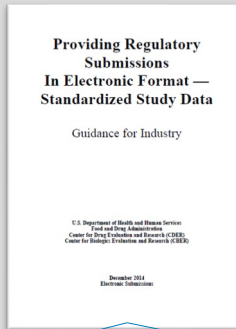
Related guidance - eCTD



Focus on study data



Binding –
Level I



FDCA Data Standards Catalog v4.1 (09-09-2016) - Supported and Required Standards
For full descriptions of column headings, see Part 2: Column Descriptions tab

Standard Number	Package Format	Standard Description (CDER/CBER)	Standard Version	Supports CDER/CBER Submission	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)	CDER/CBER Standard (CDER/CBER)
000001	Electronic	Electronic Submission (XML)	1.0	Yes	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER
000002	Electronic	Electronic Submission (XML)	1.0	Yes	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER
000003	Electronic	Electronic Submission (XML)	1.0	Yes	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER
000004	Electronic	Electronic Submission (XML)	1.0	Yes	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER
000005	Electronic	Electronic Submission (XML)	1.0	Yes	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER	CDER/CBER

Level II - these are incorporated by reference in a binding guidance

Level I vs. Level II Guidance



Level 1 guidance set forth the agency's initial interpretations of new significant regulatory requirements; describe substantial changes in FDA's earlier interpretation or policy; and deal with complex scientific or highly controversial issues.

Level 2 guidance usually address existing practices or minor changes in FDA's interpretation or policy.

[Fact Sheet: FDA Good Guidance Practices | FDA](#)

([fda.gov/about-fda/transparency-initiative/fact-sheet-fda-good-guidance-practices#:~:text=In%20general%3A,scientific%20or%20highly%20controversial%20issues.](https://www.fda.gov/about-fda/transparency-initiative/fact-sheet-fda-good-guidance-practices#:~:text=In%20general%3A,scientific%20or%20highly%20controversial%20issues.))

Level I vs. Level II Updates to Guidance



Level 1 updates to guidances describe substantial changes in FDA's earlier interpretation or policy; and deal with complex scientific or highly controversial issues.

Level 2 guidance updates usually address changes to existing practices or minor changes in FDA's interpretation or policy.

[Fact Sheet: FDA Good Guidance Practices | FDA](#)

([fda.gov/about-fda/transparency-initiative/fact-sheet-fda-good-guidance-practices#:~:text=In%20general%3A,scientific%20or%20highly%20controversial%20issues.](https://www.fda.gov/about-fda/transparency-initiative/fact-sheet-fda-good-guidance-practices#:~:text=In%20general%3A,scientific%20or%20highly%20controversial%20issues.))



FDA's Study Data Policy Framework

Level I Guidance (sitting under 745A(a))

- eStudy Data Guidance (eStudy)
- Real World Data (RWD) (currently in Draft)

Level II Guidance

- Technical Conformance Guides (TCGs)
- FDA Data Standards Catalog (Catalog)
- Certain Technical Specifications (Tech Specs)
- FDA Business Rules

[Search for FDA Guidance Documents | FDA \(fda.gov/regulatory-information/search-fda-guidance-documents\)](https://www.fda.gov/regulatory-information/search-fda-guidance-documents)

[Study Data Standards Resources | FDA \(fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources\)](https://www.fda.gov/industry/fda-data-standards-advisory-board/study-data-standards-resources)

eStudy Data Guidance Updates



Two Level II Updates were made since 2020

1. Technical Rejection Criteria (TRC) considerations, 2020
2. Scope of SEND, 2021

Providing Regulatory Submissions in Electronic Format
-- Standardized Study Data | FDA



RWD Guidance Update

- Establishes that RWD is considered study data at the point of submission and falls under 745A(a)
- Draft Published late 2021
- Review of public comments is still underway
- This review will inform the final guidance

[Providing Regulatory Submissions in Electronic Format -- Standardized Study Data | FDA](https://www.fda.gov/regulatory-information/search-fda-guidance-documents)
([fda.gov/regulatory-information/search-fda-guidance-documents](https://www.fda.gov/regulatory-information/search-fda-guidance-documents))

Technical Conformance Guide (TCG) Recent Updates



- Study Data TCG (sdTCG) updated October 2022 (pending)
- Biomedical Monitoring (BIMO) TCG updated April 2022
- IND Safety Reports TCG updated April 2022

[Study Data Standards Resources | FDA](#)



FDA Data Standards Catalog Recent Updates

Catalog continues to be updated to address the current state of requirements for study data at other Centers:

- Annotated ECG (aECG) standard
- Existing open docket added
- Added CVM as an involved center for UNII Add USCDI+ to the catalog
- S-CAP for CDRH
- ISCR R2 for CDRH
- CDISC Standards ADaM IG v1.2 and 1.3

[FDA Data Stds Catalog 8.01.2022 \(v8.2\).xlsx \(live.com\)](#)

Let's take a look at the Catalog



FDA Data Stds Catalog_8.01.2022 (v8.2).xlsx
(Navigate to the site:

[Study Data Standards Resources | FDA](#))

FDA Study Data Technical Specifications (Tech Spec) Recent Updates



1. Submitting Nonclinical Datasets for Evaluation of Rodent Carcinogenicity Studies of Pharmaceuticals, Guidance for Industry, Technical Specifications Document v. 1.0 (May 2021)
2. Submitting Next Generation Sequencing Data to the Division of Antiviral Products v. 1.0 (July 2019)
3. QT Studies Technical Specification Document v. 1.0
4. Bioanalytical Methods Validation (BMV) Tech. Spec. v1.0
5. HIV Technical Specifications Guidance v. 1.0 (March 2018) – [being considered for an update](#)
6. Vaccines Technical Specification Guidance v2.1
7. Clinical Endpoint BE Studies v1.0
8. Technical Specifications for Submitting Clinical Trial Data Sets for Treatment of Noncirrhotic Nonalcoholic Steatohepatitis (NASH) (Jan 2022)

Questions?

If you think if something later, please feel free to email:
helena.sviglin@fda.hhs.gov or
edata@fda.hhs.gov (cder)