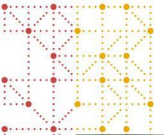
The TransCelerate / CDISC Digital Data Flow Project: Practical Electronic Study Designs

PHUSE EU Connect 2023 (DS02)

Dave Iberson-Hurst, CDISC Product Owner 7th November 2023, Version 3

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Meet the Speaker

Dave Iberson-Hurst

Title: Partner

Organization: d4k, Copenhagen

Dave has over 40 years' experience across several industries with the last 20 years spent in the pharmaceutical industry combining his technology and software development experience with clinical data standards.

During this time, he has served as the CDISC CTO, worked on, and led, several CDISC teams, presented in many forums in Europe, the US, and elsewhere across the globe. He has worked closely with the FDA, EMA, HL7, ISO, and other standards organizations and was was a member of CDISC's Blue Ribbon commission. He is currently the CDISC Product Owner for the Digital Data Flow project.

He is a partner at data4knoweldge in Copenhagen and is focused on getting greater value and utility from clinical trial data.



Disclaimer and Disclosures

- The views and opinions expressed in this presentation are those of the author(s) and do not necessarily reflect the official policy or position of CDISC.
- On contract to CDISC for the DDF work





Abstract

Over the last two years CDISC, in collaboration with Transcelerate, have been working on the Digital Data Flow (DDF) initiative. This initiative aims to *"modernize clinical trials by enabling a digital workflow to allow for the automated creation of study assets and configuration of study systems to support clinical trial execution."*. The work is focused on the protocol and associated study designs and manifests itself in a new CDISC standard, the Unified Study Definitions Model (USDM), and an open-source implementation of the USDM known as the Study Definitions Repository (SDR).

Now coming to the end of the second phase, with the third phase about to commence, the DDF project delivers a new standard that allows for the digitization of study designs and the foundation of the digital protocol.

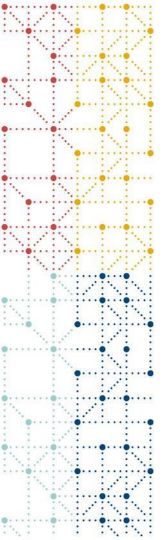
This presentation will detail:

- The work performed in phases one and two.
- The work planned as part of phase three.
- The use cases supported by the model.
- How the model/standard can enable protocol creation, automated data flow and interoperability between systems.
- How the model/standard can be deployed and implemented today.



Agenda

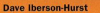
- 1. Introduction
- 2. Digital Data Flow The Project
- 3. Use Cases
- 4. Phase Three: USDM Meets M11
- 5. Summary

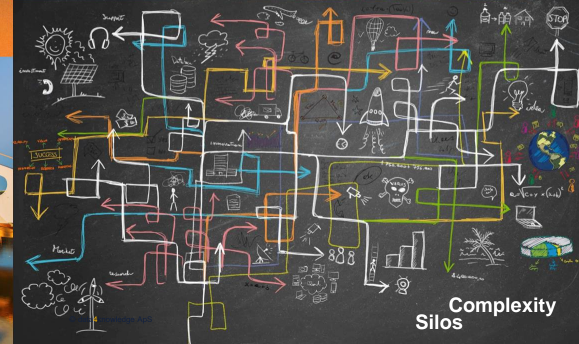


Introduction

ACT 101

The CDISC Operational Data Model: Ready to Roll?





TEMPER BAR

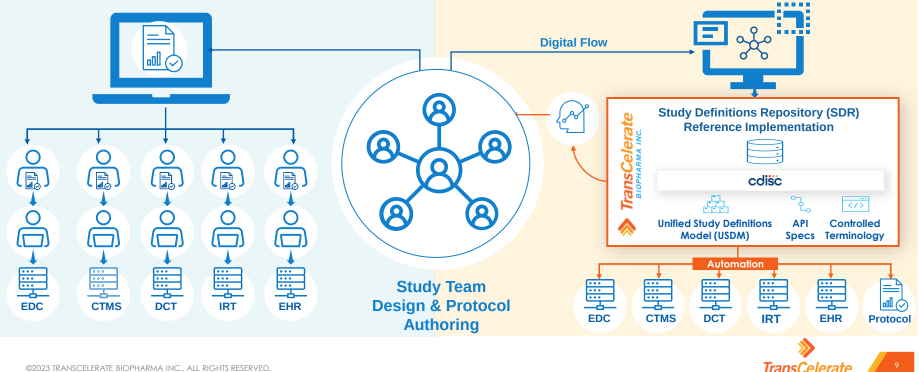
Digital Data Flow - The Project

https://www.transceleratebiopharmainc.com/assets/digital-data-flow-solutions/

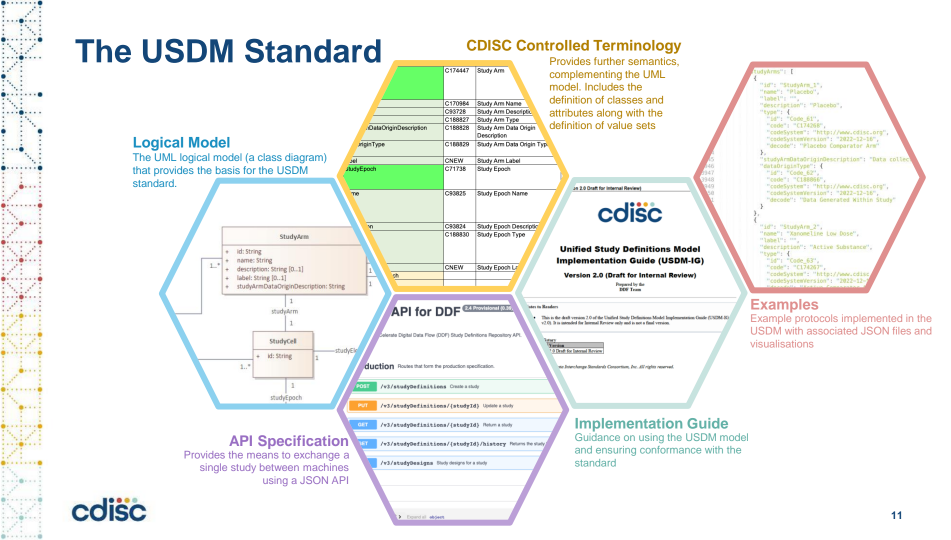
TransCelerate Digital Data Flow (DDF) Ambition Write Once, Read Many

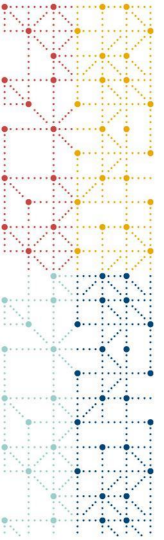
TODAY: Document-based paradigm for protocol creation, interpretation, and transcription into consuming systems

TOMORROW: Digital paradigm for protocol creation, with fully automated data flow and interoperability between systems







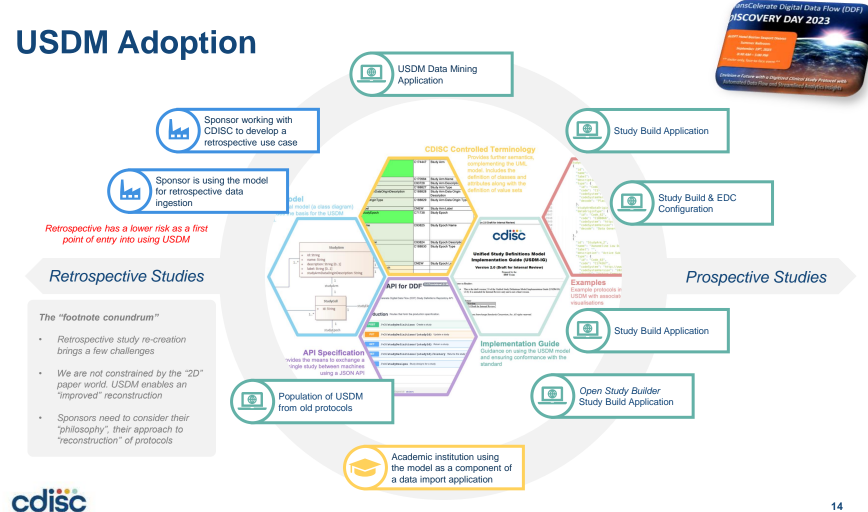


Use Cases

Use Cases: USDM with BCs allows for ...

Data Capture	Automate the setup of data capture systems, incl. RWE, and capture the data.	CTMS, TMF	The provision of protocol information to downstream systems needing "study" information.
SoA	Use the study design to build the FHIR SoA message.	Query	Having multiple studies that have a common structure allows for data export and query across the set of studies.
Data Import	Import data from a variety of sources. Can be re-exported thus allowing for conversion across versions.	SDTM	Automate the generation of SDTM datasets using the study design and BCs, including the "T" Domains.
Common Protocol Template (CPT)	Generation of the CPT from a study design.	DRAFT DDF Con	sceptual Architecture - Enhanced Study Builder and Repository, Additional Integrations (R2) -Hestoral and Actual Data
Data Decay	Re-import data using the USDM as a framework to rebuild a study design & data using the SDTM Trial Design Domains.	Sponsor Operational Data	
Scoring	The "scoring" of a study for such purposes as site impact, subject impact, environmental impact etc.	External (Public) Study Dua	Story Rudor Composition Composite Composition Composition Composition Composition
Feasibility	The use of the design to determine study feasibility including subject recruitment. A study data template.	Real World Data Addisonal Sources	Wash Learning
CT Registry	The provision of study information to a CT registry.	Study Definition Data Sources Build Buttons - Click b More Foundation Rt Insegued R	Outward Workflow, Security Oversitran Oversitran Study Definition Solution Components Oversitran Oversitran Oversitran determine Quarkstran Outcomponents Oversitran Oversitran determine Data Flow (brieflections) Outcomponents Oversitran Image: Components 02 determine Data Flow (brieflections) Image: Components Image: Components Image: Components
FAIR Data	The use of the design to aid Findability, Accessibility, Interoperability, and Reusability.	Repository now supports importing d sources, advanced Builder enaberom systems supported via additional ma	relgins from external data etta, more downitraam oprimge/integrations services





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Phase Three: USDM Meets M11

Next Steps – Phase Three





- Baseline model for specifying a study in digital format
- Model supports use of a CRF link to specify which forms to use in EDC
- Handles simple study designs

- Consume digitized study specification from an upstream source e.g., study builder)
- Store, view and search study concepts
- Downstream EDC systems may pull study specification to aid in set-up

 Improved support for complex study designs with a fully specified digitized Schedule of Activities (SoA)

2

- Model supports the identification of the appropriate CRFs for data collection to enable automated, faster configuration via use of Biomedical Concepts
- Improved CPT alignment
- Initial 'T' Domain support
- Downstream vendors can readily consume the SoA from the SDR
- Sponsor system admins can perform a visual check that SoA data received from an upstream system displays an accurate, human-readable SoA table
- Opportunity to aggregate robust historical protocol information to support analytics to drive smart design and assess risk

Focus for Phase 3 is currently being determined. Current expectations are:

Slide from May 2023

- Expand ability to handle increasingly complex studies
- ICH M11 & CPT alignment



M11 Is ...

ICH CLINICAL ELECTRONIC STRUCTURED HARMONISED PROTOCOL (CeSHarP)

https://www.ich.org/page/multidisciplinary-guidelines



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M11 Simple Example

\mathbf{N}	mnia Evampi	\frown			rechnical Specification
	mple Exampl	E	Term (Variable)	Trial Phase	
	· · · · · ·		Data Type	Pick list	
Protocol Full Title:	Template Specification	1	Topic, Value or Header	D	
Trotocorr un ruc.	The protocol should have a descriptive title that identifies the		Definition		
	scientific aspects of the trial sufficiently to ensure it is		User Guidance	For trials combining investigational drugs of	r vaccines with devices,
	immediately evident what the trial is investigating and on whom, and to allow retrieval from literature or internet			classify according to the phase of drug deve	elopment.
	searches.		Conformance	Required	•
Sponsor Confidentiality	[Sponsor Confidentiality Statement]		Cardinality		
Statement:	Insert the Sponsor's confidentiality statement, if applicable, otherwise delete.		Relationship content from ToC	Title Page	
Protocol Number:	[Protocol Number]		representing the		
	A unique alphanumeric identifier for the trial, designated by the Sponsor, is a standard part of trial data, and should be included		protocol hierarchy		
	for most trials.		Relationship		
Version:	[Version]		(reference to high level conceptual		
	An optional field for use by the Sponsor at their discretion.		model)		
Amendment Number:	[Amendment Number]		Value	Early Phase 1	
i	Enter the amendment number. If this is the original instance of			Phase 1	
rial Phase:	I Phase: [Trial Phase] [Description of Trial Phase Other]			Phase 1/Phase 2	
	Acceptable entries are: "Early	Phase 1", "Phase 1	", "Phase	Phase 2	
	1/Phase 2", "Phase 2", "Phase 2	2/Phase 3". "Phase	e 3". "Phase 4".	Phase 2/Phase 3	
				Phase 3	
Compound Number(s):	[Compound Number]			Phase 4	
Compound Number(s).	Enter the Sponsor's unique identifier for investigational			Other	
	compound(s) in the trial. Add or delete additional fields as		Business rules	Value Allowed: yes	
	needed.			Relationship: n/a	
Compound Name(s):	[Nonproprietary Name], [Proprietary Name], [Additional Proprietary Name]			Concept: Protocol short title	
	Delete this line from the table if a nonproprietary name has not		Duplicate field in		
	yet been assigned. Omit proprietary name fields if not yet		other sections		
Trial Phase:	[Trial Phase] [Description of Trial Phase Other]				
	Acceptable entries are: "Early Phase 1", "Phase 1", "Phase				
	1/Phase 2", "Phase 2", "Phase 2/Phase 3", "Phase 3", "Phase 4",				



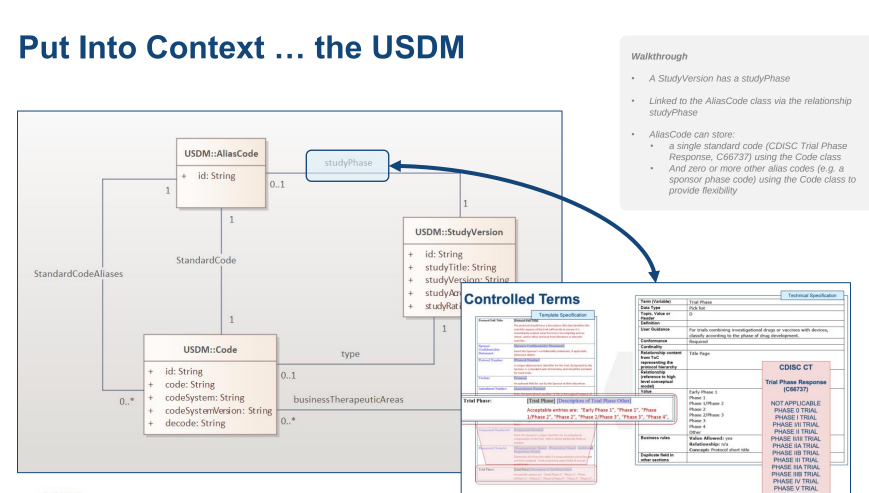
Technical Specification

Controlled Terms

				r connical opconicatio
υπηο	lied lerms	Term (Variable)	Trial Phase	
		Data Type	Pick list	
	Template Specification	Topic, Value or Header	D	
Protocol Full Title:	[Protocol Full rate] The protocol should have a descriptive title that identifies the	Definition		
	scientific aspects of the trial sufficiently to ensure it is	User Guidance	For trials combining investigationa	al drugs or vaccines with devices
	immediately evident what the trial is investigating and on whom, and to allow retrieval from literature or internet		classify according to the phase of	5
	searches.	Conformance	Required	
Sponsor	[Sponsor Confidentiality Statement]	Cardinality	Required	
Confidentiality Statement:	Insert the Sponsor's confidentiality statement, if applicable,	Relationship content	Title Page	
	otherwise delete.	from ToC	Inde Fage	
Protocol Number:	Protocol Number A unique alphanumeric identifier for the trial, designated by the	representing the		
	Sponsor, is a standard part of trial data, and should be included	protocol hierarchy		CDISC CT
	for most trials.	Relationship (reference to high		
Version:	[Version]	level conceptual		Trial Dhase Desnense
	An optional field for use by the Sponsor at their discretion.	model)		Trial Phase Response
Amendment Number:	[Amendment Number]	Value	Early Phase 1	(C66737)
			Phase 1	
l Phase:	[Trial Phase] [Description of Tr	I Phase Other]	Phase 1/Phase 2	NOT APPLICABLE
	Acceptable entries are: "Early I	ase 1". "Phase 1". "Phase	Phase 2	PHASE 0 TRIAL
			Phase 2/Phase 3	
	1 / Phase 2" "Phase 2" "Phase "			
	1/Phase 2", "Phase 2", "Phase 2		Phase 3	PHASE I TRIAL
	field.		Phase 3 Phase 4	PHASE I/II TRIAL
Compound Number(s):	field. [Compound Number]			
Compound Number(s)	field.	Business rules	Phase 4 Other	PHASE I/II TRIAL PHASE II TRIAL
Compound Number(s)	field.		Phase 4 Other Value Allowed: yes	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL
Compound Number(s):	field.		Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL
	field.		Phase 4 Other Value Allowed: yes	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL
	field.	Business rules	Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL
Compound Name(s):	field. [Compound Number] Enter the Sponsor's unique identifier for investigational compound(s) in the trial. Add or delete additional fields as needed. [Nonproprietary Name]. [Proprietary Name]. [Additional Proprietary Name] Delete this line from the table if a nonproprietary name has not yet been assigned. Omit proprietary name fields if not yet established. [Additional proprietary name fields if not yet established.	Business rules Duplicate field in	Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL PHASE IIB TRIAL PHASE III TRIAL
	field. Field. Enter the Sponsor's unique identifier for investigational compound(s) in the trial. Add or delete additional fields as needed. [Nonproprietary Name]. [Proprietary Name]. [Additional Proprietary Name] Delete this line from the table if a nonproprietary name has not yet been assigned. Omit proprietary name fields if not yet established. [Trial Phase] [Description of Trial Phase Other]	Business rules Duplicate field in	Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL PHASE IIB TRIAL PHASE III TRIAL PHASE IIIA TRIAL
Compound Name(s):	field. [Compound Number] Enter the Sponsor's unique identifier for investigational compound(s) in the trial. Add or delete additional fields as needed. [Nonproprietary Name]. [Proprietary Name]. [Additional Proprietary Name] Delete this line from the table if a nonproprietary name has not yet been assigned. Omit proprietary name fields if not yet established. [Additional proprietary name fields if not yet established.	Business rules Duplicate field in	Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL PHASE IIB TRIAL PHASE III TRIAL PHASE IIIA TRIAL PHASE IIIB TRIAL
Compound Name(s):	field.	Business rules Duplicate field in	Phase 4 Other Value Allowed: yes Relationship: n/a	PHASE I/II TRIAL PHASE II TRIAL PHASE II/III TRIAL PHASE IIA TRIAL PHASE IIB TRIAL PHASE III TRIAL PHASE IIIA TRIAL



Technical Specification



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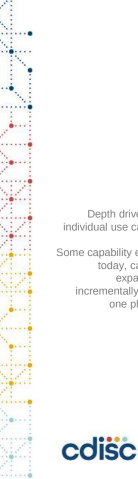
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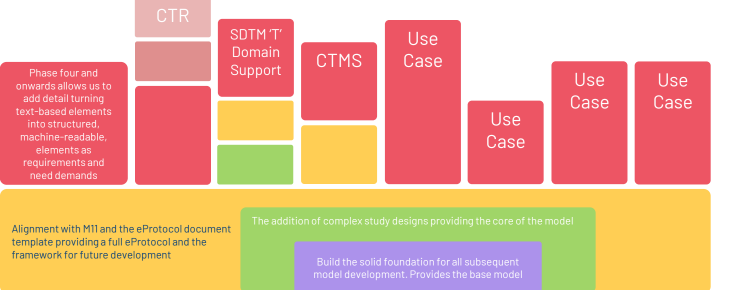


Breadth versus Depth



Depth driven by individual use cases.

Some capability exists today, can be expanded incrementally or in one phase.

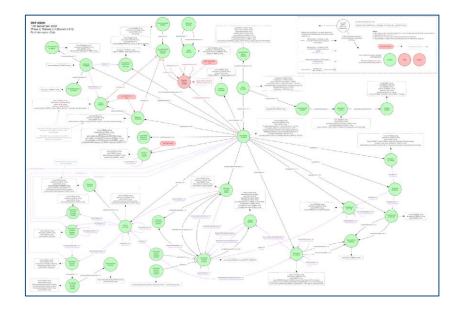


Breadth driven by the bounds of the M11 technical Specification

Shift of Focus

Phases One & Two

- Focused on the structured elements of the protocol, e.g. the Schedule of Activities (SoA)
- The protocol document was an external entity into which the structured content could be exported
- Phase Three
 - Now contains structured and unstructured elements
 - The entire protocol document is held within the USDM
 - Allows for the protocol document to be generated from the model



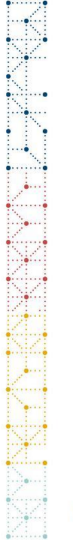


M11 Template Example Document

- First attempt to create a protocol document from the USDM, both structured [non-narrative] and unstructured [narrative text] content
- Functionality has been added to the Excel test data tool
- More work is needed, this is very much a first draft



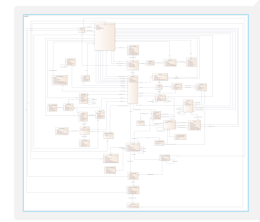
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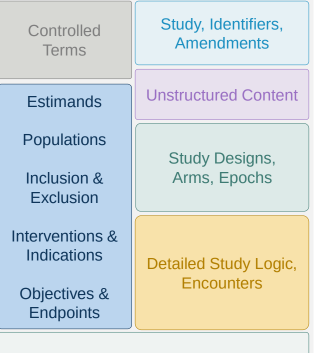


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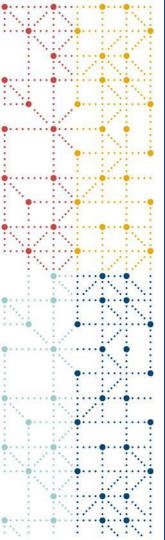
USDM Summary

• Structured content along with the ability to hold unstructured content



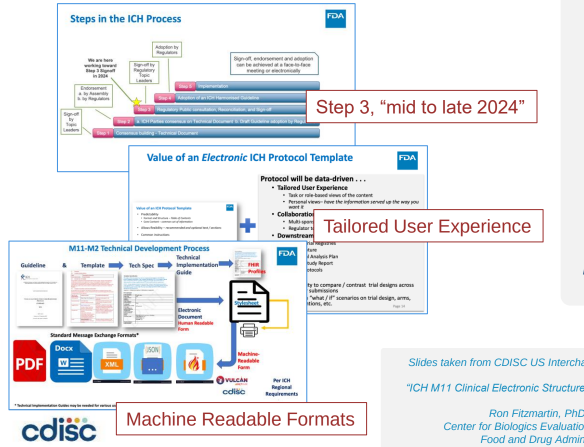


Procedures, Biomedical Concepts



Summary

ICH & The CDISC Project



Phase Three Timeline

January 2024 Phase 3 development sprints complete

> February 2024 Phase 3 public review

April 2024 Version 3 USDM published

Dates may be adjusted to align with ICH M11 publication dates.

Slides taken from CDISC US Interchange 2023 Presentation

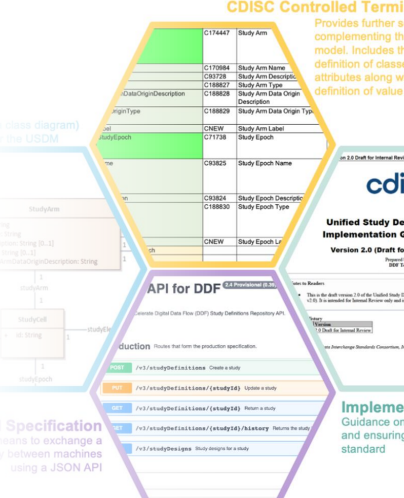
"ICH M11 Clinical Electronic Structured Harmonized Protocol"

Ron Fitzmartin, PhD, MBA Center for Biologics Evaluation and Research Food and Drug Administration



Summary

- We are "understanding" the complexity
- We can start to remove the silos and join the dots
- Digital Data Flow / Unified Study Definitions Model (DDF/USDM) fills an important gap
- USDM is but one building block, but an important one
- USDM alignment with ICH M11 will be an important step forward
- Can support various use cases, the prospective versus the retrospective
- We are only limited by our imagination



> Expand all object





Thank You

Contacts:

Dave IH: <u>diberson-hurst.external@cdisc.org</u> John Owen: <u>jowen@cdisc.org</u>

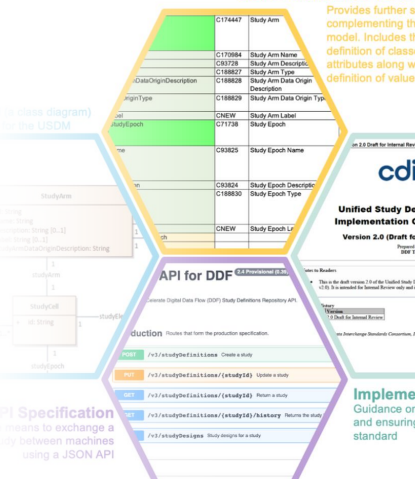
Link:

Github: https://github.com/cdisc-org/DDF-RA

CDSIC Team:

- Gerry Campion
- Drew Mills
- Erin Muhlbradt
- John Owen
- Jared Schreibman
- Berber Snoeijer
- Craig Zwickl

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> Expand all object

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CDISC Controlled Termi