

Single Day Event

From Protocol to SDTM: Automating Trial Design
with Large Language Models

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Disclaimer

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Agenda

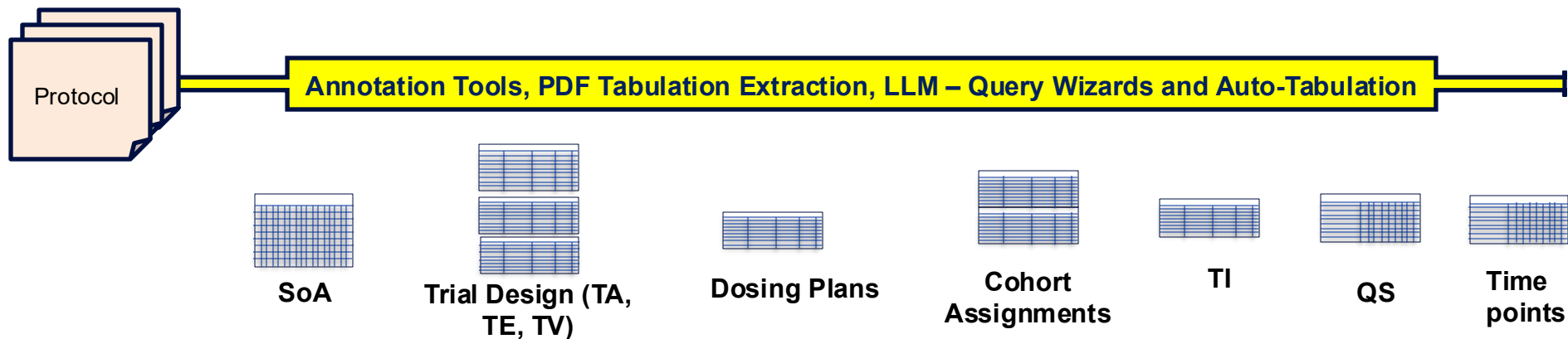
- ✓ Protocol Digitization
- ✓ Why & How Large Language Models are used?
- ✓ Experience & Learnings
- ✓ Our Approaches & Methodologies
- ✓ Few Examples of Targeted Prompt Execution
- ✓ Human in Loop for Review & Approvals
- ✓ Automatic generation of SDTM Trial Design Domains
- ✓ Conclusion



Protocol Digitization

The protocol is the central source of study truth, guiding data collection and analysis. Manual extraction into systems is slow and error-prone, and frequent amendments often lead to inconsistencies across platforms.

Digitization converts the protocol from a narrative document into a structured, machine-readable model. This allows automated configuration of downstream systems (e.g., EDC, CDISC outputs), improving both accuracy and consistency.



Key study elements (TS, TE, TA, TV, TI, Dosing, DM, Cohorts, SoA) become machine-readable objects.



Why & How Large Language Models are Used?

Understanding Domain-Specific Language

Interprets clinical terminology and varied protocol phrasing to consistently extract accurate study concepts.

Handling Variability

Clinical documents vary vastly in styles and LLMs adapt to these variations.

Few-shot Learning

LLMs can be guided with just a few examples to extract specific information, making them much faster to deploy.

Chunking & Context Management

Breaks long protocols into logical sections while preserving relationships across design, visits, and endpoints.

Extracting Unstructured/Tabular Data

Converts narrative text and complex tables into clean, structured data formats like JSON or CSV.

Summarizing Complex Study Design Logic

Simplifies dose escalation, randomization, and visit rules into clear summaries or rule-based representations.

Building Traceability

Links extracted data back to exact protocol locations for transparency, validation, and audit readiness.



Experience & Learnings

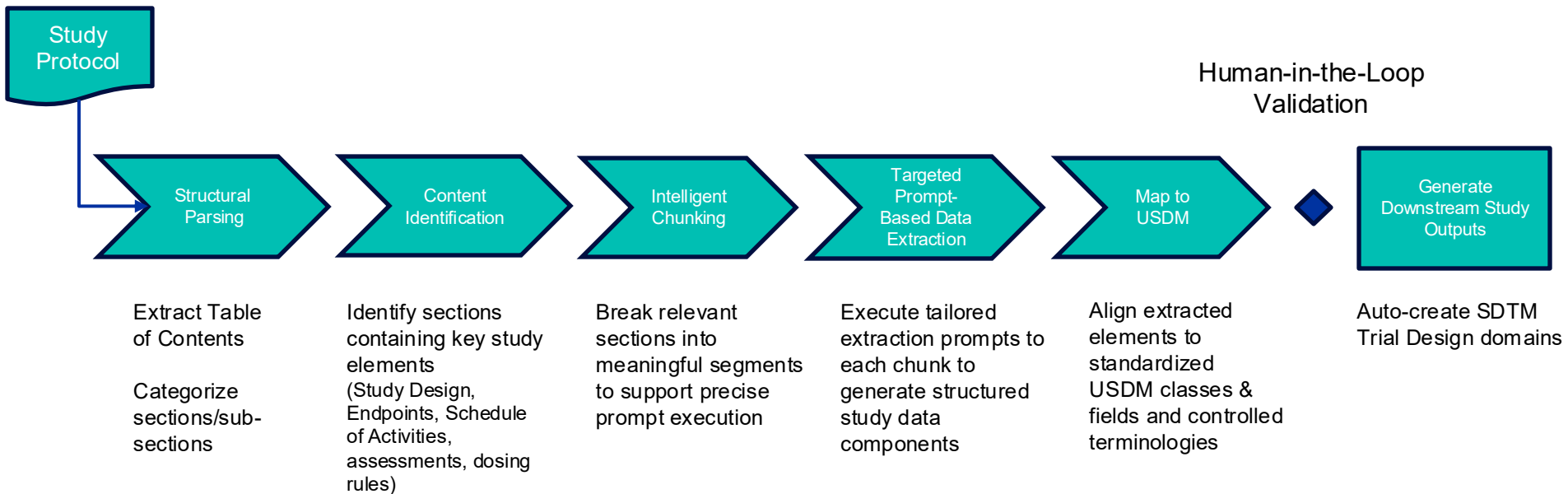
Using public or proprietary AI LLM providers with clinical study information raises significant regulatory risks. Concerns center on PHI/PII security, Intellectual Property exposure, and the inherent lack of GxP/regulatory compliance within these AI models.

To address this, we shifted to experimenting with secure private models:

- Nov 2024: Initial LLaMA training on study protocols produced unreliable, hallucinated outputs, not viable for production.
- Jan 2025: Switched to DeepSeek inference, resulting in a major improvement in quality and reliability.
- Mar 2025: Transitioned to Gemma, achieving stable, accurate, and consistently better performance.
- Expanded to fully private deployments on AWS SageMaker and Bedrock using large models like Anthropic Sonnet to ensure compliance, scalability, and secure enterprise-grade AI operations.



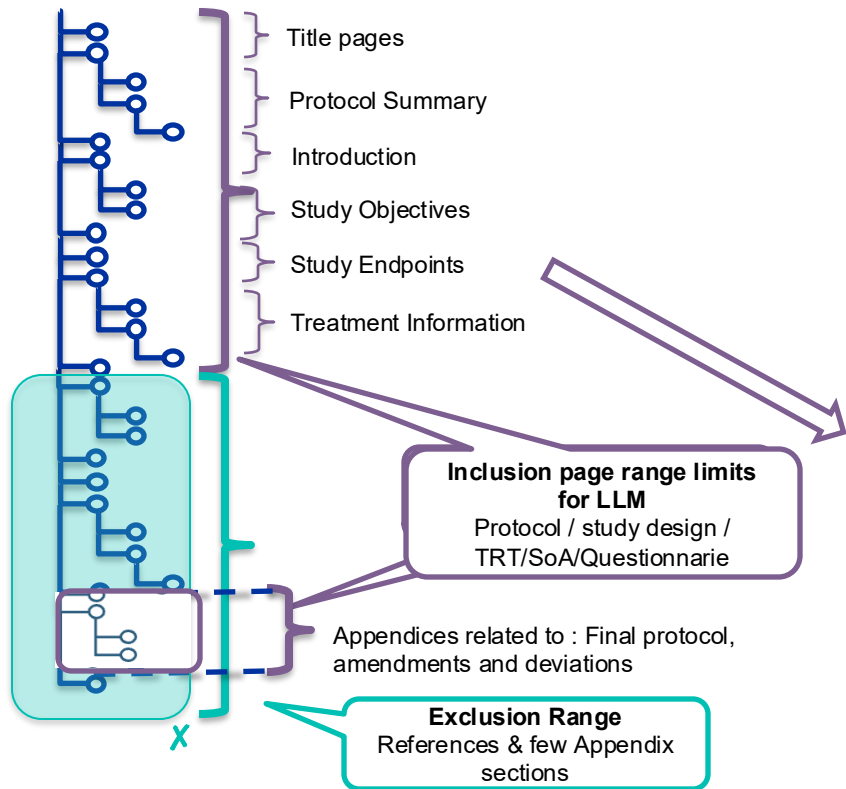
Our Approach





Intelligent Parsing of Table of Contents

Study Protocol ToC



Matrix of **Parameters** vs **Curated section** names where parameter information likely to be found

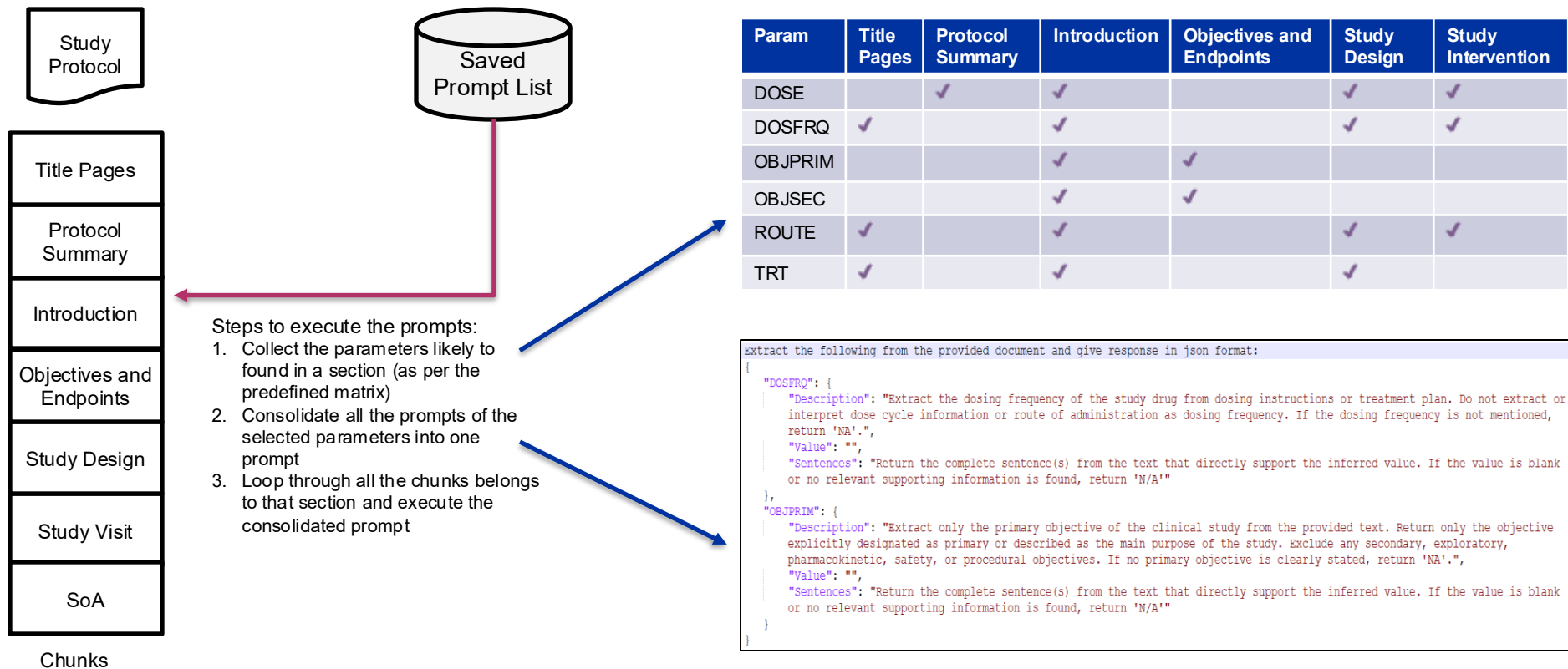
Param	Title Pages	Protocol Summary	Introduction	Objectives and Endpoints	Study Design	Study Intervention
DOSE		✓	✓		✓	✓
DOSFRQ	✓		✓		✓	✓
OBJPRIM			✓	✓		
OBJSEC			✓	✓		
ROUTE	✓		✓		✓	✓
TRT	✓		✓		✓	

ToC extracted page ranges

		Start	End
1	Title Pages	1	3
2	Protocol Summary	6	8
3	Introduction	16	20
4	Study Objectives	22	22
5	Study Endpoints	23	24
6	Study Design	25	26
7	...		

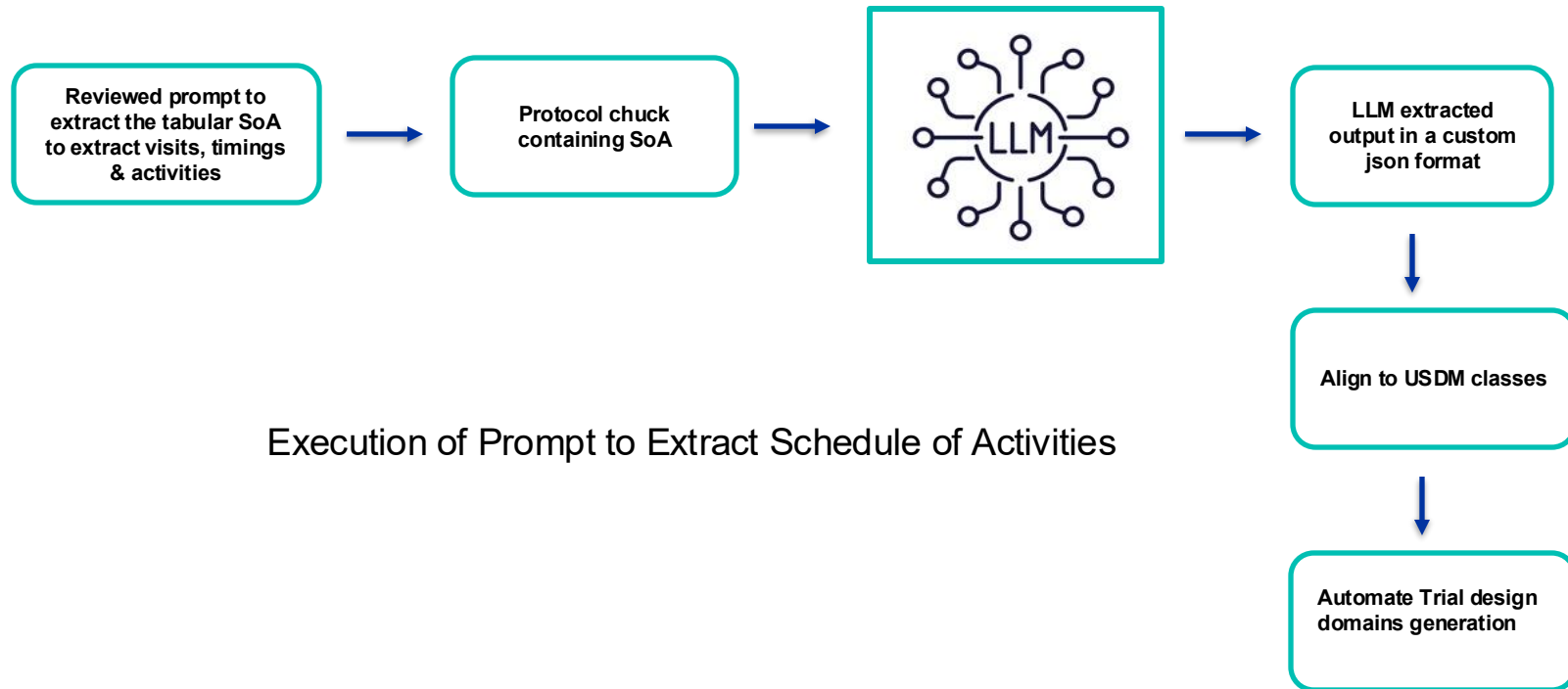


Trial Summary Prompts Execution





Schedule of Activities Prompt Execution



phuse Schedule of Activities Prompt Execution

Precise prompt to extract the tabular SoA to extract visits, timings & activities

SoA

LLM extracted output to flat structure

Align to USDM classes to automate Trial design domains generation

Study Period	Screen	Treatment Period															Post-treatment		
		Cycle 1					Cycle 2					C3+					EOT	SFU	LTFU ^a
Study Day	-28 to -1	1 ^b	2	3	5	8	15	1	8	15	1	2	3	5	8	15	≤7d post last dose	>28d post last dose	
Window		+1d	+1d	+1d	+1d	+1d	n/a	+1d	+1d		n/a				+1d	+1d			
Informed Consent ^a	X																		
Inclusion/Exclusion	X																		
Enrollment/Dose Assignment	X																		
Demography/Medical History	X																		
ECOG	X							X			X								X
Weight, Height ^d	X							X			X						X ^e	X ^e	X
Physical Examination ^a	X	X				X	X	X	X	X	X				X ^e	X ^e	X	X	X
Vital Signs ^f	X	X	X	X		X	X	X	X	X	X				X ^e	X ^e	X	X	X
Pulse Oximetry ^g	X	X				X	X	X	X	X	X				X ^e	X ^e	X	X	X
12-Lead ECG ^h	X							X			X						X		X
Hematology ^b	X	X	X	X		X	X	X	X	X	X						X		X
Chemistry ⁱ	X	X	X	X		X	X	X	X	X	X						X		X
Coagulation ^j	X	X						X			X						X		X
Urinalysis ^k	X							X			X						X		X
Pregnancy Test ^l	X	X						X			X						X		X
Serum Cytokines ^m	X	X	X	X		X	X	X			C3, C6 ⁿ								
Immunophenotyping (blood) ^o	X	X	X	X		X	X	X			C3, C6 ^m	C6	C6						
Serum BCMAs ^p	X							X	X	X	X ^o	X ^o							X
PK Samples ^q	X	X	X	X	X	X	X	X	X	X	X ^o	C6	C6	C6					X
Anti-Drug Antibodies ^r	X							X	X	X	X ^o								X
Bone Marrow Biomarkers ^s	X							X			C3, C6 ⁺¹								X
Disease Assessment ^t	X	X						X			X						X	X ^o	
Hospital Admission ^u	X	X																	

Extract the Schedule of Activities data from the following text and return it in tabulated format. Capture all relevant information from tables, footnotes, and inline citations—even if multiple tables are present. Different treatment arms, dose escalation phases, or varied schedules. Ensure all activity-note pairs are extracted accurately and completely. Do not extract data under the following columns, based on the availability.

Table - The title of the table.

Activity - This refers to the specific tasks, assessments, or procedures that are conducted during a study visit.

Cycle - A repeating period of treatment and evaluation in a clinical trial.

Cycle Window - The total length of a single treatment cycle. It defines the start and end days for that specific block of the study.

Visit - The specific label assigned to each clinical trial visit.

Visit Day - This specifies the exact day or range of days when a visit is scheduled to happen, relative to the start of the study treatment. This provides clarity on how long after enrollment a specific visit is planned.

Visit Window - This column defines the permissible timeframe around a target visit day, allowing for some flexibility in scheduling. If not explicitly provided, skip this field.

Epoch - A period of time that serves a purpose in the trial as a whole. This refers to a named study segment that reflects a distinct period in the study timeline. E.g., Screening, Baseline, etc.

Notes - Extract all footnotes linked to the Schedule of Activities table by identifying superscript/subscript markers (e.g., ^a, ^b, ^c, ^d, ^e, ^f, etc) that are directly tagged to the activity names, visit timepoints (column headers), or specific cell values. For each activity that has a superscript/subscript, consolidate all related footnotes—including inline notes, cell-level citations, column-level notes, and table-level footnotes—into a single entry formatted as: "Activity name" = " [Note Marker 1, Note 1] | [Note Marker 2, Note 2] ...". If no note is present or no superscript/subscript is tagged to the activity, return: "Activity name" = "None". Ensure each superscript/subscript is accurately mapped to its corresponding footnote, and avoid duplicating identical notes across activities unless they are explicitly referenced.

Do not miss any notes attached as they are critical information to understand the activities specifically performed during the visits. Do not infer missing values, only return explicitly stated values. Ensure accurate mapping of the appropriate column and maintain consistent formatting across all fields. Follow the instructions provided above strictly and return only the tabulated response.

Name	Description	Label	From	Type	To	ToFrom	Value	Value Label	Window Label	Window Lower	Window Upper
TM1	Screening period - Informed consent	Screening	SCREEN	BEFORE	C1D1	S2S	P28D	Day -28 to -1	0-28 to -1	P1D	P27D
TM2	Cycle 1 Day 1 - First fixed dose admin	C1D1 - First Dose	C1D1	FIXED	C1D1	S2S	P1D	Day 0			
TM3	Cycle 1 Day 2 - First post-dose	C1D2	C1D1	AFTER	C1D2	S2S	P1D	Day 2			
TM4	Cycle 1 Day 3 - Continued safety and	C1D3	C1D2	AFTER	C1D3	S2S	P1D	Day 3			
TM5	Cycle 1 Day 8 - Safety and lab evaluat	C1D5	C1D3	AFTER	C1D5	S2S	P2D	Day 5	1 day after planned visit	P1D	P1D
TM6	Cycle 1 Day 8 - Second weekly fixed d	C1D6	C1D5	AFTER	C1D6	S2S	P3D	Day 8	1 day before to 1 day after pl	P1D	P1D
TM7	Cycle 1 Day 15 - End-of-cycle fixed do	C1D15	C1D6	AFTER	C1D15	S2S	P7D	Day 15	1 day before to 1 day after pl	P1D	P1D
TM8	Cycle 1 Day 15 - Start of next cycle dose	C2D1	C1D15	AFTER	C2D1	S2S	P7D	Day 22			
TM9	Cycle 2 Day 8 - Mid-cycle fixed dose	C2D8	C2D1	AFTER	C2D8	S2S	P7D	Day 29	1 day before to 1 day after pl	P1D	P1D
TM10	Cycle 2 Day 15 - End-of-cycle dose	C2D15	C2D8	AFTER	C2D15	S2S	P7D	Day 36	1 day before to 1 day after pl	P1D	P1D
TM11	Cycle 3 Day 1 - Maintenance weekly	C3D1	C2D15	AFTER	C3D1	S2S	P7D	Day 43			
TM12	Cycle 3 Day 2 - Early post-dose	C3D2	C3D1	AFTER	C3D2	S2S	P7D	Day 49			
TM13	Cycle 3 Day 3 - Continued safety and P	C3D3	C3D2	AFTER	C3D3	S2S	P7D	Day 55			
TM14	Cycle 3 Day 5 - Safety and lab evaluat	C3D5	C3D3	AFTER	C3D5	S2S	P7D	Day 61			
TM15	Cycle 3 Day 8 - Third weekly fixed dose	C3D8	C3D5	AFTER	C3D8	S2S	P7D	Day 67	1 day before to 1 day after pl	P1D	P1D

Table	Activity	Cycle	Cycle Win	Visit	Win	Epoch	Notes
Schedule of Assessments - Study Visits: Single-F	Informed -	Screen	-28 to -1	-	Screen	Informed Consent	• [Superscript 1] Long term follow-up telephone calls or visits to occur monthly (x7 days) for 6 months, the
Schedule of Assessments - Study Visits: Single-F	Inclusion -	Screen	-28 to -1	-	Screen	Inclusion/Exclusion	• [Superscript 2] Brackets designate visits/assessments that be completed within x1d screening assess
Schedule of Assessments - Study Visits: Single-F	Enrollmen -	Screen	-28 to -1	-	Screen	Enrollment (Dose Assignment)	• None
Schedule of Assessments - Study Visits: Single-F	Demograp -	Screen	-28 to -1	-	Screen	Demography/Medical History	• None
Schedule of Assessments - Study Visits: Single-F	ECOG	Screen	-28 to -1	-	Screen	ECOG	• None
Schedule of Assessments - Study Visits: Single-F	ECOG	Cycle 1	1	1	Treatment	ECOG - None	
Schedule of Assessments - Study Visits: Single-F	ECOG	Cycle 2	1	n/a	Treatment	ECOG - None	
Schedule of Assessments - Study Visits: Single-F	ECOG	-	EOT	s7d post -	Post-treat	ECOG - None	
Schedule of Assessments - Study Visits: Single-F	Weight, H -	Screen	-28 to -1	-	Screen	Weight, Height	• [Superscript 3] Height is measured at Screening only.
Schedule of Assessments - Study Visits: Single-F	Weight, H -	Cycle 1	1	1	Treatment	Weight, Height	• [Superscript 3] Height is measured at Screening only.
Schedule of Assessments - Study Visits: Single-F	Weight, H -	Cycle 2	1	n/a	Treatment	Weight, Height	• [Superscript 3] Height is measured at Screening only.
Schedule of Assessments - Study Visits: Single-F	Weight, H -	EOT	s7d post -	-	Post-treat	Weight, Height	• [Superscript 3] Height is measured at Screening only.
Schedule of Assessments - Study Visits: Single-F	Physical I -	Screen	-28 to -1	-	Screen	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 1	1	1	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 2	8	8 +1d	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 3	15	15 +1d	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 4	1	n/a	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 5	5	5 +1d	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	Cycle 6	8	8 +1d	Treatment	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Physical I -	EOT	s7d post -	-	Post-treat	Physical Examination	• [Superscript 4, 5] Full physical examination & Screening: abbreviated physical exams at indicated vis
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Screen	-28 to -1	-	Screen	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 1	1	1	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 2	2	2	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 3	3	3	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 4	8	8 +1d	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 5	15	15 +1d	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d
Schedule of Assessments - Study Visits: Single-F	Vital Sign -	Cycle 6	1	n/a	Treatment	Vital Signs	• [Superscript 6] Vital signs: blood pressure, heart rate, respiratory rate, and body temperature call or test as at d



Human in Loop - Review & Approvals

- Compare LLM-extracted data side-by-side with source protocol text and line references
- Study team reviews and approves extracted elements before processing
- Approved data automatically supports SDTM Trial Design generation, and study build outputs

The screenshot displays the Phuse Human in Loop interface, which is used for reviewing and approving trial design parameters. The interface is divided into two main sections: a left sidebar for trial design parameters and a right pane for the clinical protocol document.

Left Sidebar (Trial Design Parameters):

- Trial Summary:** Shows the trial title, study phase, and other key information.
- TD Generation Form:** Displays the trial design parameters, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Elements:** Lists the trial elements, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Arms:** Displays the trial arms, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Visits:** Displays the trial visits, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Inclusion/Exclusion:** Displays the trial inclusion/exclusion criteria, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Disease Assessments:** Displays the trial disease assessments, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Trial Disease Milestones:** Displays the trial disease milestones, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Demographics:** Displays the demographics, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Subject Elements:** Displays the subject elements, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Subject Disease Milestones:** Displays the subject disease milestones, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Subject Visits:** Displays the subject visits, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.
- Time Points:** Displays the time points, including trial arms, trial visits, trial inclusion/exclusion, trial disease assessments, trial disease milestones, demographics, subject elements, subject disease milestones, subject visits, and time points.

Right Pane (Clinical Protocol Document):

The right pane displays the clinical protocol document, which is titled "CLINICAL PROTOCOL: PC202301". The document is a Phase 1/2 Open-label, Multicenter, Dose Escalation and Dose Expansion Study of the Safety, Tolerability, and Pharmacokinetics of PC001 in Patients with Advanced Cancers Associated with Expression of Delta-Like Canonical Notch Ligand 3 (DLL3) Who Have Failed Standard Available Therapy. The document is dated 20 August 2020, Version 2.0.

The document is divided into several sections, including:

- Study Title:** A Phase 1/2 Open-label, Multicenter, Dose Escalation and Dose Expansion Study of the Safety, Tolerability, and Pharmacokinetics of PC001 in Patients with Advanced Cancers Associated with Expression of Delta-Like Canonical Notch Ligand 3 (DLL3) Who Have Failed Standard Available Therapy
- Study Number:** PC202301
- Study Phase:** 1/2
- Product Name:** PC001
- Indication:** Advanced Solid Tumors Associated with Expression of Delta-Like Canonical Notch Ligand 3
- Investigator:** Multicenter
- Sponsor:** Papatrans Life Sciences, USA
- Medical Monitor:** Neeraj J. Executive Medical Director, Papatrans Life Sciences

The document also includes a table of document versions:

Document	Version	Date
Original Protocol:	Version 1.0	07 July 2020
Amendment 1	Version 2.0	20 August 2020



Conclusion

Targeted prompt-driven NLP and semantic modeling enable precise extraction of key study elements directly from the protocol.

Human-in-the-loop review safeguards scientific intent while reducing manual effort and interpretation variability.

This digital solution accelerates workflows, enhances data quality, and enables full automation via USDM structure for study build and SDTM trial design with full traceability.

Thank you

XbiomTM makes data useful



**The Global Healthcare
Data Science Community**

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