EXMC GROUND-BASED SPACE RADIATION ANALOG PILOT DRUG STABILITY STUDY:

PRELIMINARY DATA REVIEW

Vernie Daniels, MS, RPh; Tina Bayuse, PharmD, RPh; Rebecca Blue, MD, MPH; Erik Antonsen, MD, PhD; Kris Lehnhardt, MD

Presentation Outline

- >Introduction:
 - Historical Significance
 - Purpose
 - Research Objectives
- >Materials / Methods

- >Results
 - Time-point I Review
 - Time-point II Update
- > Preliminary Conclusions

Historical Significance

➤ Historical NASA drug stability studies suggested that spaceflight conditions compromise medication safety and efficacy (Putcha et al, 2001 – 2011).

➤ Historical NASA ground analog experiments designed to simulate the effects of high-energy radioactive particles on medications during spaceflight, suggested that radiation exposure during spaceflight could threaten drug quality and potency on long-duration exploration missions (Putcha et al, 2006).

➤ Follow-on NASA flight studies revealed reduced active pharmaceutical ingredient (API) concentrations, and altered drug release; when compared to matching ground controls (Putcha et al, 2006 – 2011).

Purpose

- Uncertainty remains regarding space radiation impacts on drug stability and shelf life
- ➤ Space environmental analog and ground-based targeted radiation research could reveal valuable insight into drug safety and effectiveness
 - In 2017, the Exploration Medical Capability (ExMC) Element designed a three-year pilot analog experiment to expose medications to a series of simulated Galactic Cosmic Radiation (GCR) mixed-species beam exposures at the NASA Space Radiation Laboratory (NSRL) at Brookhaven National Laboratory (BNL)
 - First time-point analysis completed 2018; presented IWS 2019
 - Second time-point analysis completed 2019; presented IWS 2020

Research Objectives

- Evaluate if the effects of ground-based rapid-switching radiation beam exposures can effectively reproduce previously observed effects of spaceflight radiation on drug stability and shelf life.
- Further evaluate the utility of simulated GCR beam exposures as an effective ground-based analog for predicting the impacts of GCR exposure on drug stability and shelf life during spaceflight.

Study Drugs:

- > Four medications were prioritized and selected based on:
 - Pharmaceutical stability profiles confirmed by previous research / literature
 - Clinical relevance for exploration spaceflight

Table A.	Experimental	Drug	List
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Test Product	Drug	Expiration Date		
Α	Acetaminophen 500 mg Tablets	01/31/21		
В	Amoxicillin 500 mg Capsules	12/31/19		
С	Ibuprofen 400mg Tablets	11/30/19		
E	Promethazine 25mg Tablets	02/29/20		

- Sets (identical brands / lots) of each drug product procured for each experimental arm
 - Sufficient quantities to provide a statistically significant number of replicates
 - ❖ 50-100 dosage units / package
 - ❖ 4 different drugs x 2 packages each x 4 different study conditions = 32 packages of drugs
- Packaged (as closely as possible) to resemble flight medical systems operational packaging (e.g. drug flight bottles / plastic bags / unit-dose strips, etc.).

Study Design: Four Experimental Arms

- Non-irradiated JSC Control Group
- 2. Non-irradiated Traveling Control Group
- 3. Irradiation Group I (Mixed-beam 0.5Gy Total Dose)
- 4. Irradiation Group II (Mixed-beam 1.0Gy Total Dose)

Environmental Monitoring

> Temperature / RH:

- Shipment / Storage: USP <659> "Packaging and Storage Requirements" defined conditions for "controlled room temperature" (15 - 30° C, 30 - 65% RH)
 - Environmental condition tracking
 - o Environmentally controlled storage chambers

> Radiation:

- Detection and Monitoring: Thermoluminescence Dosimeters (TLD-100 LiF:Mg,Ti)
 - o TLDs enclosed in clear gelatin capsules, attached to front and / or back, of each drug product package

Irradiation:

- First experiment at NSRL to utilize the mixedspecies simulator:
- > Exposure dose: Two mixed-beam radiation doses
 - 0.5 Gy
 - 1.0 Gy
- ➤GCR-like beam profile:
 - ¹H, ⁴He, ¹²C, ¹⁶O, ²⁸Si, ⁴⁸Ti, and ⁵⁶Fe
- ➤ Dose detection and monitoring: Thermoluminescence Dosimeters (TLD-100 LiF:Mg,Ti)
 - TLDs enclosed in clear gelatin capsules, attached to front and / or back, of each drug product package

						ion	(MeV/n)	Range (cm)	LE1 (κeV/μm)	(mGy)
			,			¹ H	20.0	0.43	2.59	30.4
Ion	Energy	Range		Dose		1H	23.3	0.56	2.29	6.7
ion	(MeV/n)	(cm)	(keV/µm)	(mGy)		1H	27.2	0.75	2.02	7.4
¹H	100	Poly	ethylene degrad	ler to		1H	31.7	0.98	1.79	8.0
1H	150	15.9	0.54	35.0		1H	37.0	1.30	1.58	8.7
1H	250	38.1	0.39	68.9		1H	43.2	1.72	1.39	9.3
1H	1000	325.6	0.22	123.6		1H	50.3	2.26	1.23	10.0
	100		ethylene degrad		V	*H	58.7	2.99	1.09	10.6
⁴ He						1H	68.5	3.95	0.97	11.1
⁴ He	150	16.0	2.17	7.5		1H	79.9	5.20	0.86	11.2
⁴ He	250	38.3	1.56	16.4		IH.	100.0	7.76	0.73	27.2
⁴ He	1000	327.8	0.88	24.9		Same 3	Energy		Commence and the same and	Dose
12C	1000	110.1	7.95	11.7		lon	(MeV/n)	Range (cm)	LET (keV/μm)	(mGy)
160	350	17.0	20.8	15.4		⁴ He	20.0	0.43	10.34	11.0
28Si	600	22.7	50.2	8.1		⁴ He	23.3	0.57	9.14	2.1
48Ti	1000	32.5	109.5	4.5		⁴ He	27.2	0.75	8.06	2.2
56Fe	600	13.1	175.1	4.1		⁴ He	31.7	0.99	7.12	2.3
Total				500.0		⁴ He	37.0	1.31	6.29	2.5
				500.0		4He	43.2	1.73	5.56	2.6
						⁴ He	50.3	2.28	4.92	2.7
						⁵ He	58:7	3.01	4.36	2.7
							SPACE T	- PARE	9539	617
					1	4He	68.5	3.97	3.86	2.7

Figure 2.0: NSRL GCR Simulation Beam Composition



Figure 3.0: Irradiation Dose Measurement_TLD Placement

Drug Stability Analyses: USP monograph Test methods developed for all analyses

- ➤ API chemical content (Liquid Chromatography: UPLC H-Class System with PDA Detector)
 - Trial runs to validate USP method suitability
 - Assay methods validated using commercial chemical reference standards

> Presence of impurities or degradation products

- Assessment of chromatographic peak percentages
- Drug formulation component chromatogram overlays
- > Dissolution testing to determine API release characteristics
 - Hanson Vision Elite 8 dissolution apparatus
 - Ultraviolet—visible (UV / Vis) Spectrophotometer to assist with dissolution assessments

>Irradiation Dose Measurements

- Entrance dose for irradiated drugs at the 500 mGy dose: 422.7 ± 5.7 -465.3 ± 6.3 mGy
 - a measured dose of 7-15% lower than the expected nominal dose (500 mGy)
- Entrance dose for irradiated drugs at the 1000 mGy dose: 856.8 ± 11.6 -932.4 ± 12.7 mGy
 - a measured dose of 7-14% lower than the expected nominal dose (1000 mGy)
- A dose-decreasing trend between the front and back TLDs of 7 – 16% was observed for each drug group.

Table C: Summary of TLD-100 Dose Measurement Results

Drug Type	Exposure	TLD-100	TLD-100	TLD-100	Nominal
		Measured	Mean Dose	Ratio	NSRL
		Dose	(mGy)	Back/Front	Dose
		(mGy)			(mGy)
Acetaminophen	A3a_Front	465.3 ± 6.3	448.1 ± 6.1	0.93 ± 0.02	500
500mg	A3a_Back	431.0 ± 5.9	440.1 ± 0.1		500
	A3b_Back	412.7 ± 5.6	412.7 ± 8.8	N/A	500
Acetaminophen	A4a_Front	932.4 ± 12.7	899.2 ± 9.8	0.93 ± 0.02	1000
500mg	A4a_Back	866.0 ± 11.8	699.2 ± 9.6		1000
	A4b_Back	843.9 ± 11.5	843.9 ± 11.5	N/A	1000
Amoxicillin	B3a_Front	436.2 ± 5.9	400 7 1 5 5	0.84 ± 0.02	500
500mg	B3a_Back	365.2 ± 5.0	400.7 ± 5.5		500
	B3b_Back	371.9 ± 5.1	371.9 ± 5.1	N/A	500
Amoxicillin	B4a_Front	864.4 ± 11.7	804.4 ± 9.0	0.86 ± 0.02	1000
500mg	B4a_Back	744.4 ± 10.1	804.4 ± 9.0		1000
	B4b_Back	747.0 ± 10.2	747.0 ± 10.2	N/A	1000
Ibuprofen	C3a_Front	422.7 ± 5.7	405.7 ± 5.5	0.92 ± 0.02	500
400mg	C3a_Back	388.8 ± 5.3	405.7 ± 5.5		500
	C3b_Back	394.4 ± 5.4	394.4 ± 5.4	N/A	500
Ibuprofen	C4a_Front	871.5 ± 11.8	822.6 ± 9.2	0.89 ± 0.02	1000
400mg	C4a_Back	773.7 ± 10.5	022.0 ± 3.2		1000
	C4b_Back	733.3 ± 10.0	733.3 ± 10.0	N/A	1000
Levofloxacin	D3a_Front	432.0 ± 5.9	412.6 ± 5.6	0.91 ± 0.02	500
500mg	D3a_Back	393.2 ± 5.3	412.0 ± 3.0		500
	D3b_Back	384.0 ± 5.2	384.0 ± 5.2	N/A	500
Levofloxacin	D4a_Front	856.8 ± 11.6	855.5 ± 9.0	1.00 ± 0.02	1000
500mg	D4a_Back	854.2 ± 11.6	633.3 ± 9.0		1000
	D4b_Back	711.0 ± 9.7	711.0 ± 9.7	N/A	1000
Promethazine	E3a_Front	448.4 ± 6.1	413.8 ± 5.6	0.85 ± 0.02	500
25mg	E3a_Back	379.2 ± 5.2	415.6 ± 5.0		500
	E3b_Back	400.4 ± 5.4	400.4 ± 5.4	N/A	500
Promethazine	E4a_Front	923.6 ± 12.6	847.5 ± 9.7	0.84 ± 0.02	1000
25mg	E4a_Back	771.5 ± 10.5	847.5 ± 9.7		1000
	E4b_Back	769.4 ± 10.5	769.4 ± 10.5	N/A	1000

Note: The TLD measured dose values include the control dose subtraction, no additional corrections needed.

API Content Analysis: API content for all irradiated and control study medications tested at time-points (t₁- t₂) met the USP acceptance criteria for potency, or percentage of label claimed API content:

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SAMPLE	PRODUCT NAME	STUDY ARM	% LABEL CLAIM API 2018	% LABEL CLAIM API 2019	% CHANGE IN POTENCY (t2-t1 / t1)	% API USP	RESULT OUTCOME
A1A	Acetaminophen 500 mg Tablet	Non-Irradiated JSC Control	95.3	103.22	↑8.31	90-110	Pass
A1B	Acetaminophen 500 mg Tablet		100.4	101.85	↑1.44	90-110	Pass
A2A	Acetaminophen 500 mg Tablet	Non-Irradiated Travel Control	97.08	102.18	↑5.25	90-110	Pass
A2B	Acetaminophen 500 mg Tablet	Non-Irradiated Travel Control	97.73	102.81	↑5.2	90-110	Pass
АЗА	Acetaminophen 500 mg Tablet	Irradiation Group I (Mixed-beam 0.5 GY	100.18	102.45	↑2.27	90-110	Pass
A3B	Acetaminophen 500 mg Tablet	Irradiation Group I (Mixed-beam 0.5 GY	96.51	99.86	↑3.47	90-110	Pass
A4A	Acetaminophen 500 mg Tablet	Irradiation Group II (Mixed-beam 1.0 GY	95.76	102.4	↑6.93	90-110	Pass
A4B	Acetaminophen 500 mg Tablet	Irradiation Group II (Mixed-beam 1.0 GY	103.67	99.32	↓4.2	90-110	Pass

SAMPLE	PRODUCT NAME	STUDY ARM	% LABEL CLAIM API 2018	% LABEL CLAIM API 2019	% CHANGE IN POTENCY (t2-t1) / t1	% API USP REQUIREMENT	RESULT OUTCOME
B1a	Amoxicillin 500 mg Capsules	Non-Irradiated JSC Control	100.16	102.08	↑1.92	90-120	Pass
B1b	Amoxicillin 500 mg Capsules	Non-Irradiated JSC Control	97.44	98.58	↑1.17	90-120	Pass
B2a	Amoxicillin 500 mg Capsules	Non-Irradiated Travel Control	100.96	101.51	↑0.54	90-120	Pass
B2b	Amoxicillin 500 mg Capsules	Non-Irradiated Travel Control	100.04	100.02	↑0.02	90-120	Pass
ВЗа	Amoxicillin 500 mg Capsules	Irradiation Group I (Mixed-beam 0.5 GY	101.57	99.68	↓1.86	90-120	Pass
B3b	Amoxicillin 500 mg Capsules	Irradiation Group I (Mixed-beam 0.5 GY	99.31	97.11	↓2.25	90-120	Pass
B4a	Amoxicillin 500 mg Capsules	Irradiation Group II (Mixed-beam 1.0 GY	98.74	98.97	↑0.23	90-120	Pass
B4b	Amoxicillin 500 mg Capsules	Irradiation Group II (Mixed-beam 1.0 GY	102.42	93.72	↓8.49	90-120	Pass

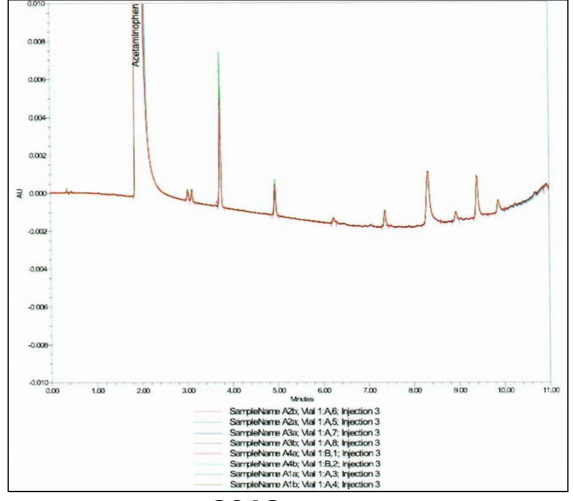
	, cooking release	(IVIIAOU DOUITI 1.0 OT					
SAMPLE	PRODUCT NAME	STUDY ARM	% LABEL CLAIM API 2018	% LABEL CLAIM API 2019	% CHANGE IN POTENCY (t1-t2 / t1)	% API USP REQUIREMENT	RESULT OUTCOME
C1a	lbuprofen 400 mg Tablets	Non-Irradiated JSC Control	103.85	98.24	↓5.40	90-110	Pass
C1b	lbuprofen 400 mg Tablets	Non-Irradiated JSC Control	106.6	102.94	↓3.43	90-110	Pass
C2a	lbuprofen 400 mg Tablets	Non-Irradiated Travel Control	109.32	97.21	↓11.08	90-110	Pass
C2b	lbuprofen 400 mg Tablets	Non-Irradiated Travel Control	103.84	101.37	↓2.38	90-110	Pass
СЗа	lbuprofen 400 mg Tablets	Irradiation Group I (Mixed-beam 0.5 GY	106.6	96.98	↓9.02	90-110	Pass
C3b	lbuprofen 400 mg Tablets	Irradiation Group I (Mixed-beam 0.5 GY	109.31	96.96	↓11.3	90-110	Pass
C4a	lbuprofen 400 mg Tablets	Irradiation Group II (Mixed-beam 1.0 GY	104.38	95.15	↓8.84	90-110	Pass
C4b	lbuprofen 400 mg Tablets	Irradiation Group II (Mixed-beam 1.0 GY	106.43	95.37	↓10.39	90-110	Pass

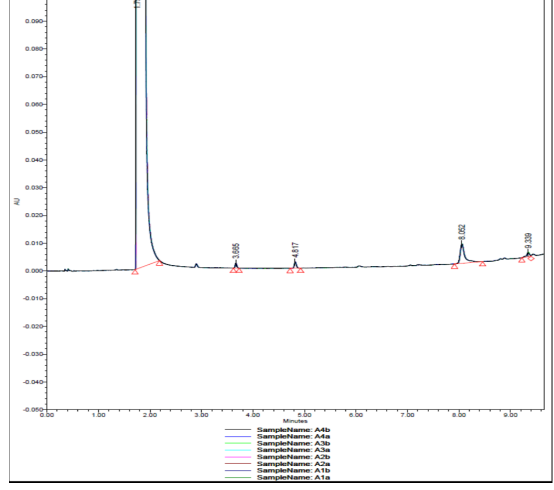
SAMPLE	PRODUCT NAME	STUDY ARM	% LABEL CLAIM API 2018	% LABEL CLAIM API 2019	% CHANGE IN POTENCY (t2-t1 / t1)	% API USP REQUIREMENT	OUTCOME
E1a	Promethazine 25 mg Tablets	Non-Irradiated JSC Control	99.17	100.2	↑1.04	95-110	Pass
E1b	Promethazine 25 mg Tablets	Non-Irradiated JSC Control	104.66	101.39	↓3.12	95-110	Pass
E2a	Promethazine 25 mg Tablets	Non-Irradiated Travel Control	107.32	100.09	↓6.73	95-110	Pass
E2b	Promethazine 25 mg Tablets	Non-Irradiated Travel Control	104.33	100.68	↓3.49	95-110	Pass
E3a	Promethazine 25 mg Tablets	Irradiation Group I (Mixed-beam 0.5 GY	103	104.02	↑0.99	95-110	Pass
E3b	Promethazine 25 mg Tablets	Irradiation Group I (Mixed-beam 0.5 GY	109.53	101	↓7.79	95-110	Pass
E4a	Promethazine 25 mg Tablets	Irradiation Group II (Mixed-beam 1.0 GY	108.33	102.3	↓5.57	95-110	Pass
E4b	Promethazine 25 mg Tablets	Irradiation Group II (Mixed-beam 1.0 GY	107.3	100.53	↓6.31	95-110	Pass

The specification limit for change in potency usually ≤ 10%. (Waterman KC, Swanson JT, Lippold BL. A scientific and statistical analysis of accelerated aging for pharmaceuticals. Part 1: accuracy of fitting methods. J Pharm Sci 2014 Oct;103(10):3000-6).

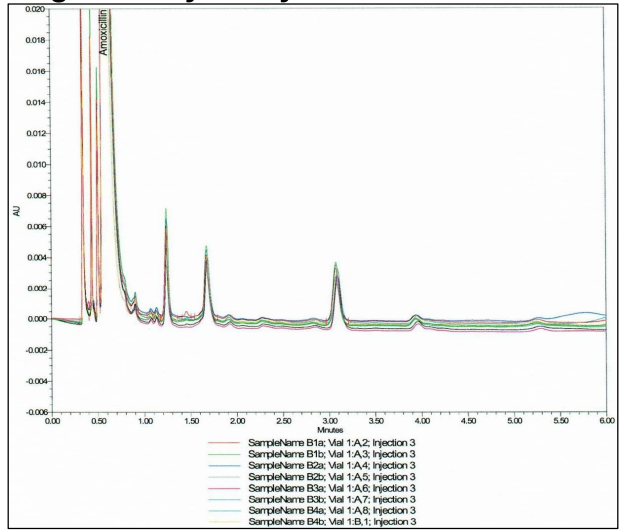
Drug Stability Analyses: Assessment of drug component chromatograms at $t_1 - t_2$ revealed no new or foreign peaks in any of the irradiated drug product samples

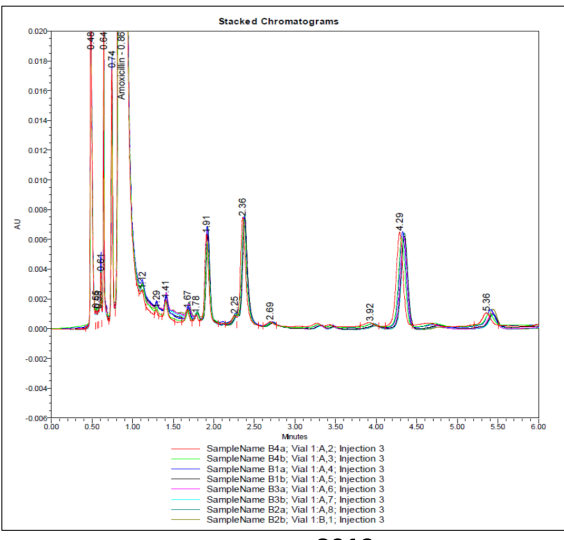
Acetaminophen



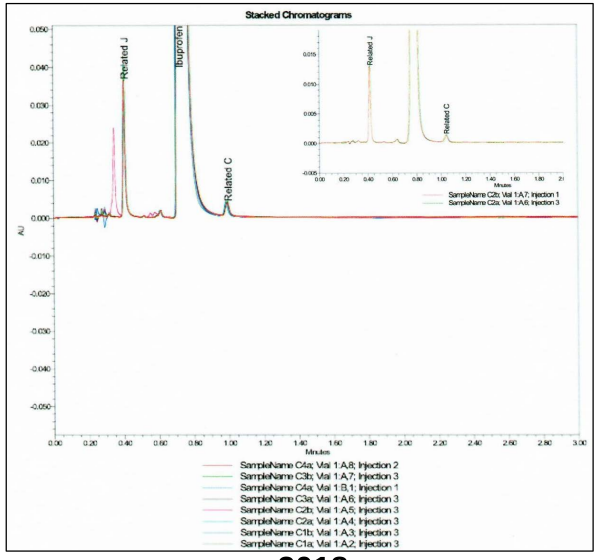


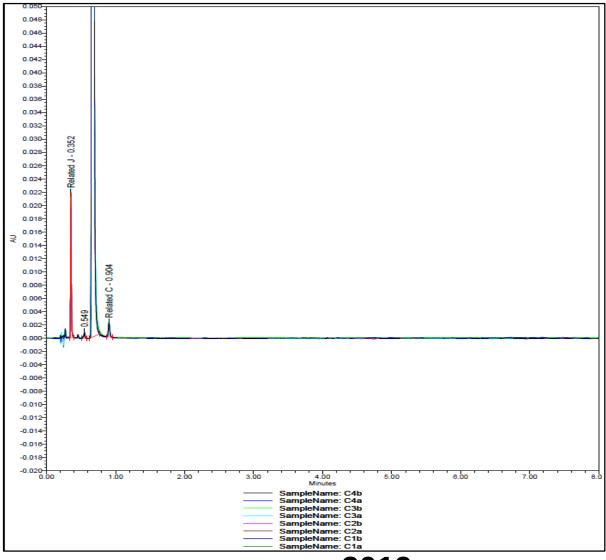
Drug Stability Analyses Continued: Amoxicillin





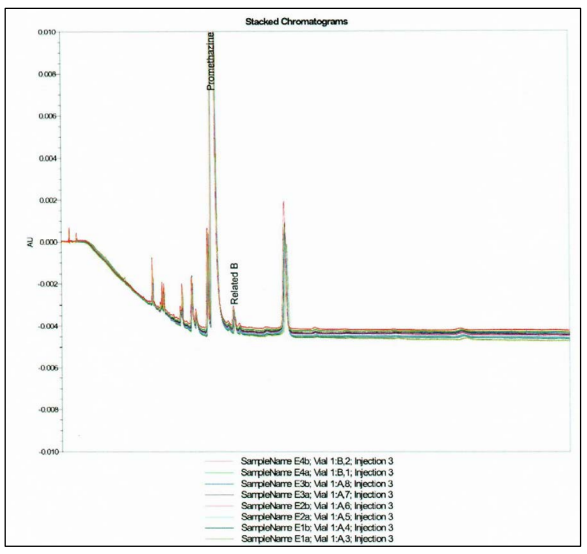
Drug Stability Analyses Continued: Ibuprofen

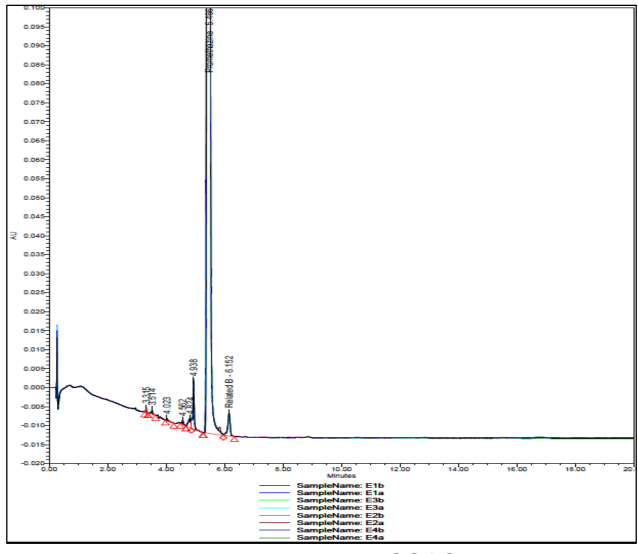




2018

Drug Stability Analyses Continued: Promethazine





Dissolution: All samples met the USP requirement for Dissolution.

➤ Some amoxicillin samples revealed "significant changes" in release between t₁ and t₂

Acetaminophen:

Amoxicillin:

Sample	Product Name	Sample Name	% Dissolved	2018 Standard	% Dissolved	2019 Standard	% Change in	USP Standard	Sample	Product Name	Sample Name	% Dissolved	2018 Standard	% Dissolved	2019 Standard	% Change in	USP Standard
			2018	Deviation (n=6)	2019	Deviation (n=6)	Dissolution	(≥ 80%)				2018	Deviation (n=6)	2019	Deviation (n=6)	Dissolution	(≥ 80%)
A1a	Acetaminophen	Non-irradiated JSC	99.51	1.10%	102.54	1.07%	3.04	Pass	B1a	Amoxicillin	Non-irradiated JSC	100.16	5.78%	93.43	2.12%	↓6.72	Pass
711%	500 mg Tablets	Control Group								500 mg Capsules	Control Group						
A1b	Acetaminophen	Non-irradiated JSC	100.71	3.56%	100.4	1.24%	0.31	Pass	B1b	Amoxicillin	Non-irradiated JSC	97.44	5.06%	92.18	4.53%	↓5.4	Pass
7112	500 mg Tablets	Control Group							510	500 mg Capsules	Control Group						
A2a	Acetaminophen	Non-irradiated	100.12	2.95%	101.09	1.49%	0.97	Pass	B2a	Amoxicillin	Non-irradiated	100.96	4.63%	89.69	3.16%	↓11.16	Pass
/ La	500 mg Tablets	Traveling Control Group							DZa	500 mg Capsules	Traveling Control Group						
A2b	Acetaminophen	Non-irradiated Traveling	100.77	4.48%	99.47	2.08%	1.29	Pass	B2b	Amoxicillin	Non-irradiated	100.04	4.70%	92.80	1.65%	↓7.24	Pass
AZU	500 mg Tablets	Control Group							DZN	500 mg Capsules	Traveling Control Group						
A3a	Acetaminophen	Irradiation Group I	102.75	4.01%	100.49	1.67%	2.2	Pass	B3a	Amoxicillin	Irradiation Group I	101.57	6.17%	91.25	3.89%	↓10.16	Pass
Asa	500 mg Tablets	(Mixed-beam 0.5Gy							DJa	500 mg Capsules	(Mixed-beam 0.5Gy Total						
٨٥١	Acetaminophen	Irradiation Group I	100.85	2.19%	101.19	0.86%	0.34	Pass	B3b	Amoxicillin	Irradiation Group I	99.31	5.46%	91.05	5.43%	↓8.32	Pass
A3b	500 mg Tablets	(Mixed-beam 0.5Gy							D30	500 mg Capsules	(Mixed-beam 0.5Gy Total						
140	Acetaminophen	Irradiation Group II	99.51	2.81%	100.43	1.56%	0.92	Pass	B4a	Amoxicillin	Irradiation Group II	98.74	4.53%	86.13	2.77%	↓12.78	Pass
A4a	500 mg Tablets	(Mixed-beam 1.0Gy							D4a	500 mg Capsules	(Mixed-beam 1.0 Gy Total	90.74	4.33%				
1 1 h	Acetaminophen	Irradiation Group II	95.45	4.47%	100.74	2.08%	5.54	Pass	B4b	Amoxicillin	Irradiation Group II	400.40	0.400/	88.59	5.18%	↓13.5	Pass
A4b	500 mg Tablets	(Mixed-beam 1.0Gy							D40	500 mg Capsules	(Mixed-beam 1.0 Gy Total	102.42	2.49%				
	<u> </u>	,									· · · · · · · · · · · · · · · · · · ·						

Drug Stability Analyses Continued:

Ibuprofen:

Promethazine:

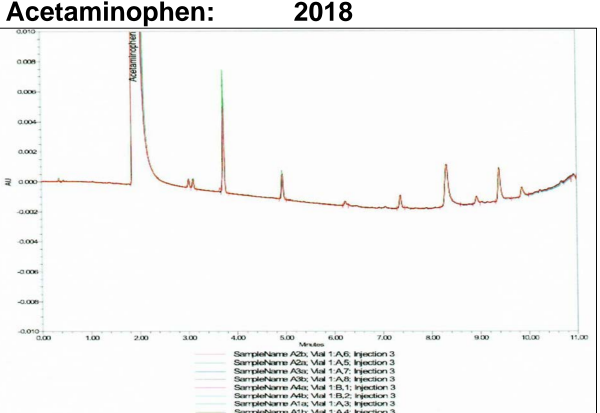
Sample	Product Name	Sample Name	2018%	2018 Standard	2019%	2019Standard	% Change in	USP Standard	Sample	Product Name	Sample Name	2018 %	2018 Standa
			Dissolved	Deviation (n=6)	Dissolved	Deviation (n=6)	Dissolution	(≥ 80%)				Dissolved	Deviation (N=
04-	lbuprofen	Non-irradiated	100.64	1.32%	98.23	0.20%	↓2.39	Pass	E1a	Promethazine	Non-irradiated	98.48	0.92%
C1a	400 mg Tablets	JSC Control Group								25 mg Tablets	JSC Control Group		
041	lbuprofen	Non-irradiated	100.97	0.95%	98.17	0.16%	↓2.77	Pass	E1b	Promethazine	Non-irradiated	98.38	0.58%
C1b	400 mg Tablets	JSC Control Group								25 mg Tablets	JSC Control Group		
00.	lbuprofen	Non-irradiated	100.38	1.52%	98.11	0.00%	↓2.26	Pass	E2a	Promethazine	Non-irradiated	98.21	2.13%
C2a	400 mg Tablets	Traveling Control Group					,			25 mg Tablets	Traveling Control Group		
001	lbuprofen	Non-irradiated	100.58	2.39%	98.55	0.38%	↓2.02	Pass	E2b	Promethazine	Non-irradiated	98.69	1.35%
C2b	400 mg Tablets	Traveling Control Group								25 mg Tablets	Traveling Control Group		
00	lbuprofen	Irradiation Group I	100.49	1.92%	98.74	0.40%	↓1.74	Pass	E3a	Promethazine	Irradiation Group I	98.12	1.69%
C3a	400 mg Tablets	(Mixed-beam 0.5Gy					,			25 mg Tablets	(Mixed-beam 0.5Gy Total		
001	lbuprofen	Irradiation Group I	100.59	3.26%	98.86	0.42%	↓1.72	Pass	E3b	Promethazine	Irradiation Group I	98.58	0.80%
C3b	400 mg Tablets	(Mixed-beam 0.5Gy								25 mg Tablets	(Mixed-beam 0.5Gy Total		
- 04-	lbuprofen	Irradiation Group II	100.53	1.36%	98.99	0.71%	↓1.53	Pass	E4a	Promethazine	Irradiation Group II	98.41	1.47%
C4a	400 mg Tablets	(Mixed-beam 1.0 Gy								25 mg Tablets	(Mixed-beam 1.0 Gy Total		
C4h	lbuprofen	Irradiation Group II	100	2.66%	99.05	0.86%	↓0.95	Pass	E4b	Promethazine	Irradiation Group II	98.48	0.62%
C4b	400 mg Tablets	(Mixed-beam 1.0 Gy								25 mg Tablets	(Mixed-beam 1.0 Gy Total		

Sample	Product Name	Sample Name	2018 %	2018 Standard	2019 %	2019 Standard	% Change in	USP Standard
			Dissolved	Deviation (N=6)	Dissolved	Deviation (N=6)	Dissolution	(≥ 80%)
E1a	Promethazine	Non-irradiated	98.48	0.92%	103.46	0.53%	↑5.05	Pass
	25 mg Tablets	JSC Control Group						
E1b	Promethazine	Non-irradiated	98.38	0.58%	103.95	0.68%	↑5.66	Pass
	25 mg Tablets	JSC Control Group						
E2a	Promethazine	Non-irradiated	98.21	2.13%	102.94	0.46%	↑4.82	Pass
	25 mg Tablets	Traveling Control Group						
E2b	Promethazine	Non-irradiated	98.69	1.35%	103.93	0.36%	↑5.31	Pass
	25 mg Tablets	Traveling Control Group						
E3a	Promethazine	Irradiation Group I	98.12	1.69%	103.90	0.32%	↑5.89	Pass
	25 mg Tablets	(Mixed-beam 0.5Gy Total						
E3b	Promethazine	Irradiation Group I	98.58	0.80%	104.03	0.59%	↑5.53	Pass
	25 mg Tablets	(Mixed-beam 0.5Gy Total						
E4a	Promethazine	Irradiation Group II	98.41	1.47%	103.50	0.51%	↑5.17	Pass
	25 mg Tablets	(Mixed-beam 1.0 Gy Total					,	
E4b	Promethazine	Irradiation Group II	98.48	0.62%	103.46	0.53%	↑5.05	Pass
	25 mg Tablets	(Mixed-beam 1.0 Gy Total						

Drug Degradation Products / Impurities:

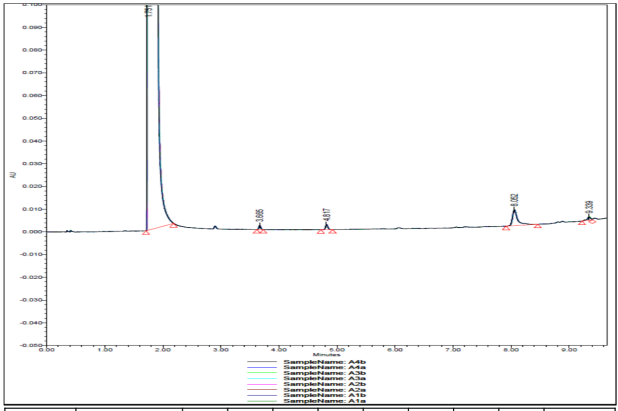
Impurities peak percent calculations, and overlay chromatograms revealed no foreign or new peaks in any of the irradiated samples during the first two time-point analyses.

Acetaminophen:



	Samperane Alb, viai 1,44, iljection 3												
Degradation	Retention time	A1a	A1b	A2a	A2b	A3a	A3b	A4a	A4b	Standard			
Peak#										Deviation			
1	P-Aminophenol	ND	-										
2	3.738	0.11	0.12	0.15	0.11	0.11	0.12	0.1	0.130	0.016			
3	4.946	0.03	0.03	0.04	0.03	0.03	0.03	0.03	0.04	0.005			
4	7.369	0.02	0.02	0.02	0.02	0.02	0.03	0.02	0.03	0.005			
5	8.32	0.15	0.15	0.15	0.15	0.14	0.15	0.15	0.13	0.007			
6	8.935	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.000			
7	9.4	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.000			
8	9.872	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.000			

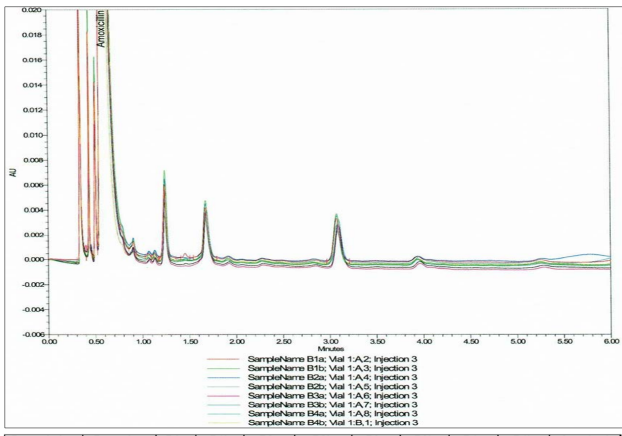
ND -	"Not	Detected"

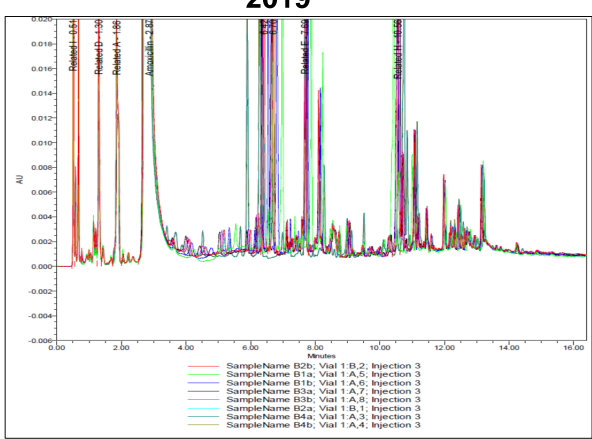


Degradati	Retention time	A1a	A1b	A2a	A2b	A3a	A3b	A4a	A4b	Standard
on Peak#										Deviation
1	P-Aminophenol	ND	ND	ND	ND	ND	ND	ND	ND	
2	3.665	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.00
3	4.817	0.02	0.03	0.03	0.025	0.02	0.02	0.02	0.02	0.0045
4	6.241	ND	ND	ND	ND	ND	0.01	ND	0.01	0.000
5	8.052	0.163	0.16	0.166	0.163	0.166	0.163	0.16	0.163	0.0052
6	9.339	0.02	0.02	0.02	0.02	0.023	0.023	0.02	0.02	0.0015

Drug Degradation Products / Impurities:

Amoxicillin: 2018 2019



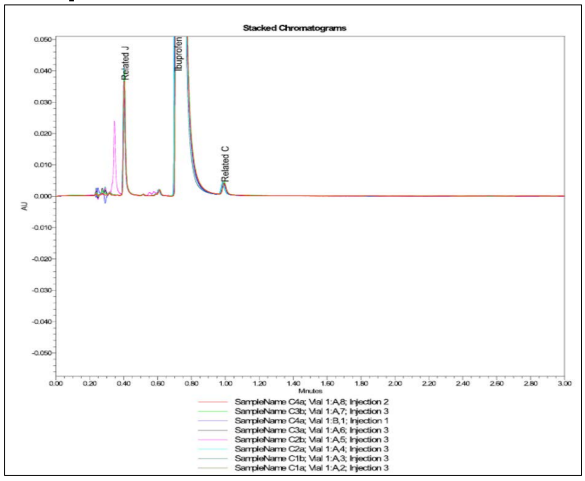


Degradation	Retention	B1a	B1b	B2a	B2b	B3a	B3b	B4a	B4b	Standard
Peak#	Time (Avg.)			l		l	l			Deviation
Related I	0.52	0.5	0.48	0.49	0.49	0.46	0.47	0.42	0.45	0.32
Related D	1.322	0.22	0.23	0.22	0.23	0.26	0.24	0.34	0.26	0.040
Related A	1.945	0.44	0.45	0.44	0.45	0.49	0.47	0.61	0.48	0.056
Related B	2.26	ND								
Related E	7.592	0.48	0.49	0.48	0.48	0.51	0.49	0.6	0.49	0.041
Related G	8.139	0.11	0.12	0.12	0.11	0.13	0.13	0.17	0.13	0.019
Related C	9.167	0.04	0.04	0.04	0.04	0.05	0.05	0.08	0.05	0.014
Related H	10.708	0.69	0.71	0.7	0.69	0.76	0.72	0.98	0.72	0.097
Dimer	11.899	0.07	0.08	0.06	0.08	0.09	0.08	0.12	0.09	0.018

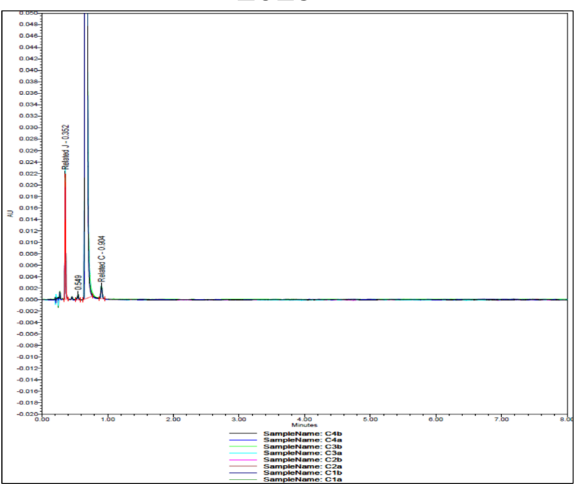
Degradation Peak#	Retention Time (Average)	B1a	B1b	B2a	B2b	ВЗа	ВЗЬ	B4a	B4b	Standard Deviation
Related I	0.504	0.46	0.47	0.46	0.46	0.46	0.47	0.51	0.51	0.022
Related D	1.322	0.32	0.33	0.29	0.29	0.28	0.28	ND	ND	0.020
Related A	1.945	0.42	0.43	0.42	0.42	0.41	0.43	0.38	0.43	0.016
Related B	2.26	ND								
Related E	7.782	0.59	0.61	0.60	0.59	0.59	0.59	0.72	0.72	0.084
Related G	8.139	Z	Z	ND	ND	Z	ND	ND	ND	-
Related C	9.3	ND	-							
Related H	10.567	0.59	0.62	0.61	0.61	0.63	0.62	0.69	0.68	0.036
Dimer	11.899	ND	ND	ND	ND	ND	ND	2	ND	-

Drug Stability Analyses Continued: Drug Degradation Products / Impurities

Ibuprofen: 2018 2019



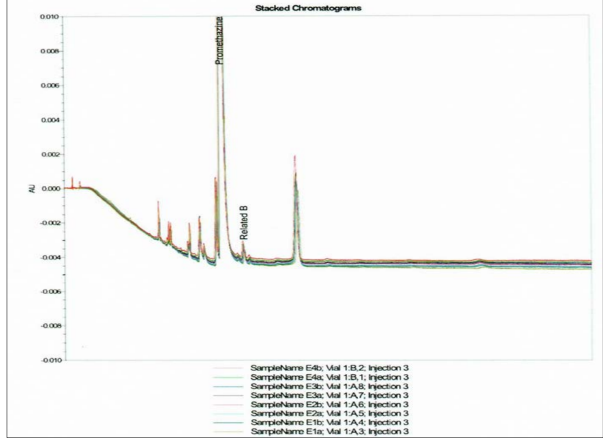
Degradation Peak#	Retention Time (min)	C1a	C1b	C2a	C2b	СЗа	C3b	C4a	C4b	Standard Deviation
1	ND	ND	ND	ND	ND	0.86	ND	ND	ND	ND
Related J	0.41	1.55	1.54	1.57	1.51	1.57	1.57	1.55	1.57	0.034
3	0.622	0.17	0.13	0.16	0.12	0.15	0.23	0.14	0.17	0.279
Related C	1.011	0.22	0.27	0.25	0.27	0.26	0.27	0.26	0.25	0.021



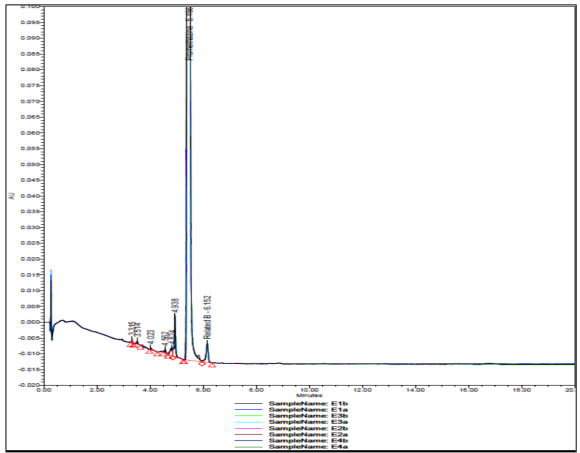
Degradation Peak#	Retention Time (min)	C1a	C1b	C2a	C2b	СЗа	C3b	C4a	C4b	Standard Deviation
Related J	0.351	1.79	1.77	1.82	1.78	1.71	1.77	1.81	1.76	0.034775
3	0.548	0.12	0.115	0.125	0.12	0.11	0.11	0.12	0.12	0.0047
Related C	0.902	0.29	0.34	0.35	0.35	0.33	0.33	0.34	0.33	0.0191

Drug Stability Analyses Continued: Drug Degradation Products / Impurities

Promethazine: 2018 2019



Degradation Peak#	Peak Retention Time (min)	E1a	E1b	E2a	E2b	E3a	E3b	E4a	E4b	Standard Deviation
1	1.71	0.08	0.03	ND	ND	ND	ND	ND	ND	0.035
2	3.577	0.21	0.19	0.2	0.18	0.15	0.15	0.19	0.22	0.026
3	3.651	0.05	0.05	0.06	0.05	0.03	0.06	0.07	0.05	0.012
4	3.956	0.12	0.03	0.14	0.03	0.12	0.16	0.11	0.19	0.057
5	4.026	0.02	0.12	0.03	0.13	ND	0.03	0.06	0.04	0.045
6	4.734	0.04	0.04	0.04	0.04	0.06	0.05	0.05	0.06	0.009
7	5.115	0.11	0.09	0.09	0.08	0.07	0.08	0.2	0.09	0.042
8	5.274	0.1	0.1	0.1	0.1	0.1	0.09	0.11	0.03	0.025
Related B	6.77	0.21	0.21	0.2	0.21	0.2	0.21	0.21	0.22	0.006
11	8.791	0.18	0.2	0.2	0.19	0.18	0.17	0.21	0.23	0.019
12	10.003	0.02	0.02	ND	0.03	0.03	ND	ND	ND	0.006



Degradation Peak#	Peak Retention Time (min)	E1a	E1b	E2a	E2b	E3a	E3b	E4a	E4b	Standard Deviation
1	3.318	0.02	0.02	0.02	0.2	0.02	0.02	0.04	0.02	0.063
2	3.516	0.03	0.03	0.03	0.3	0.03	0.03	0.1	0.03	0.095
3	4.029	0.02	0.02	0.02	0.01	0.02	0.02	0.03	0.02	0.005
4	4.571	0.03	0.03	0.02	0.025	0.02	0.02	0.16	0.02	0.048
5	4.818	0.12	0.12	0.11	0.11	0.11	0.11	0.236	0.12	0.043
6	4.945	0.33	0.32	0.33	0.33	0.33	0.33	0.566	0.32	0.085
Related B	6.142	0.45	0.45	0.45	0.45	0.45	0.45	0.48	0.45	0.01

Preliminary Conclusions

- Results revealed that the simulated GCR exposure did not facilitate noncharacteristic degradation two years post radiation exposure.
 - Two study drugs (Amoxicillin, Ibuprofen) approached labeled expiration dates, <u>none</u> had expired prior to t₂ stability testing (09/16/19).
 - "Lag-time" degradation is characteristic of some solid dosage forms.
- ➤ Uncertainties regarding the extent and rate of drug degradation for the tested medications may be further clarified by t₃ testing.
- ➤ The observed results from t₁ and t₂ drug stability analyses, concur with those from previous JSC stability studies:
 - Differences in API potency between spaceflight and ground-controlled drug samples(Du et al, 2011)
 - Differences in API potency between irradiated and non-irradiated control drug samples simulated single-beam radiation ground-analog studies (Putcha et al, 2006).

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- ➤ NASA Shipping and Receiving
- ExMC Clinical & Science Team Lead
- > KBR Human Health and Performance Contracts
 - Logistics Team
 - Task Order Management Teams

Backup Slides

Introduction

- > Pharmaceutical "Drug" Stability: The chemical and physical integrity of a drug dosage unit, or finished pharmaceutical product (FPP).
 - Drug stability testing evaluates how drug quality varies as a function of time and storage conditions (e.g., temperature, humidity, radiation)
 - FDA Monographs for all approved drugs are in the United States Pharmacopeia (USP); which includes acceptance criteria for the API (ICH Q1A R2, "Stability Testing of New Drug Substances")
 - Failure to meet the acceptance criteria for potency, presence of degradation products,
 and dissolution is considered a "significant change" for an FPP
 - Chromatographic methods provide quantitative and qualitative analysis of drug substances
 - The most common chromatographic method for stability studies uses HPLC with UV detection
 - A dissolution or API release test measures the extent and rate of solution formation from a solid (e.g. tablet, capsule) or semi-solid (e.g. cream, ointment) FPP
 - Changes in API release from a FPP can influence bioavailability and therapeutic effectiveness

Introduction

- > Photostability refers to how a drug compound responds to radiation exposure.....(Glass et al., 2004)
 - Exposure to high-intensity electromagnetic radiation may cause significant loss of the API, and initiate formation of degradation products (M Jamrógiewicz, 2016)
 - Drug photodecomposition can lead to:
 - Loss of API potency which could lead to a reduction in therapeutic activity
 - Degradation product contamination leading to adverse drug experiences (van Henegouwen, 1997; Moan, 1996; Kullavanijaya and Lim, 2005)

Study Design: Four Experimental Arms

- Non-irradiated JSC Control Group
- 2. Non-irradiated Traveling Control Group
- 3. Irradiation Group I (Mixed-beam 0.5Gy Total Dose)
- 4. Irradiation Group II (Mixed-beam 1.0Gy Total Dose)

Environmental Monitoring

> Temperature / RH:

- Shipment / Storage: USP <659> "Packaging and Storage Requirements" defined conditions for "controlled room temperature" (15 - 30° C, 30 - 65% RH)
 - o Courier tracking: Sensitech Temp Tale®4 temperature tracker
 - o Project tracking: HOBO U12-012 environmental tracking device
 - o JSC storage: Environmental chambers (Darwin, Model KB0303-AA-DA, Sanyo, model MLR-350H
 - o Analytical vendor storage: Caron Environmental Chamber, Model 7000-10

> Radiation:

- Detection and Monitoring: Thermoluminescence Dosimeters (TLD-100 LiF:Mg,Ti)
 - TLDs enclosed in clear gelatin capsules (Lilly, No. 0, NDC 00002240702); and attached to front and / or back, of each drug product package

- > Dissolution testing to determine API release characteristics
 - Hanson Vision Elite 8 dissolution apparatus
 - Ultraviolet—visible (UV / Vis) Spectrophotometer to assist with dissolution assessments
 - UV/Vis refers to absorption spectroscopy or reflectance spectroscopy in part of the ultraviolet and the full, adjacent visible spectral regions. Direct UV/VIS spectrophotometric determination of absorbance has been the traditional analytical method for dissolution testing

Environmental Control:

- ➤ Transport / Storage temperatures / RH on average remained within USP limits for "controlled room temperature" throughout the experimental timeline.
 - Temperatures: 18.9°C 28.8°C
 - RH: 4% 79% (transport from JSC to analytical vendor only)
 - Only brief excursions (< 24 hours) above RH upper and lower limits

➤ Irradiation Dose Measurements

- Entrance dose for irradiated drugs at the 0.5 Gy dose: 422.7 ± 5.7 465.3 ± 6.3 mGy
 - o a measured dose of 7-15% lower than the expected nominal dose (500 mGy)
- Entrance dose for irradiated drugs at the 1.0 Gy dose: 856.8 ± 11.6 932.4 ± 12.7 mGy,
 - o a measured dose of 7-14% lower than the expected nominal dose (1000 mGy)
- A dose-decreasing trend between the front and back TLDs of 7 16% was observed for each drug group.