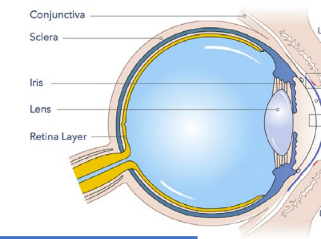


Novel *in vitro*, *ex vivo* and *in vivo* assessment of ophthalmic semi-solid drug products

Xiuling Lu, Ph.D.
Professor
University of Connecticut

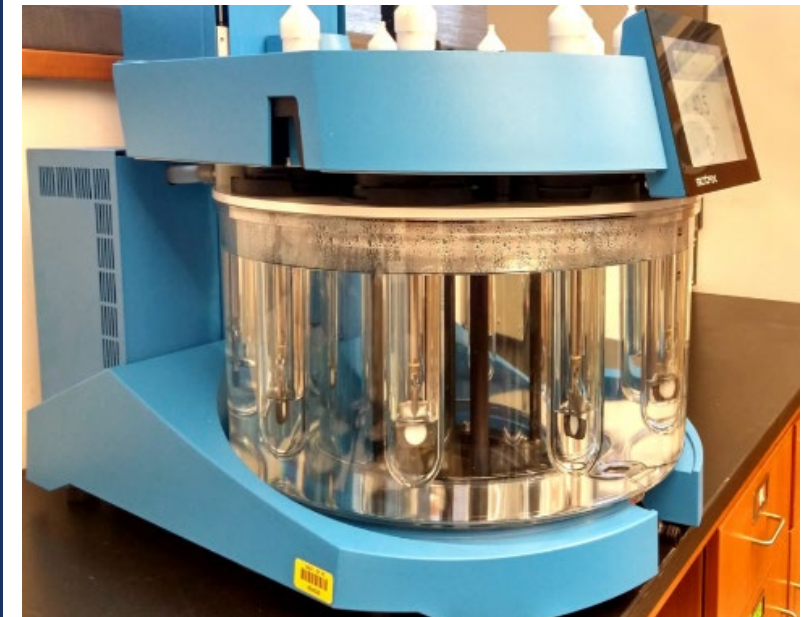
Challenges in Assessing Ophthalmic Ointments



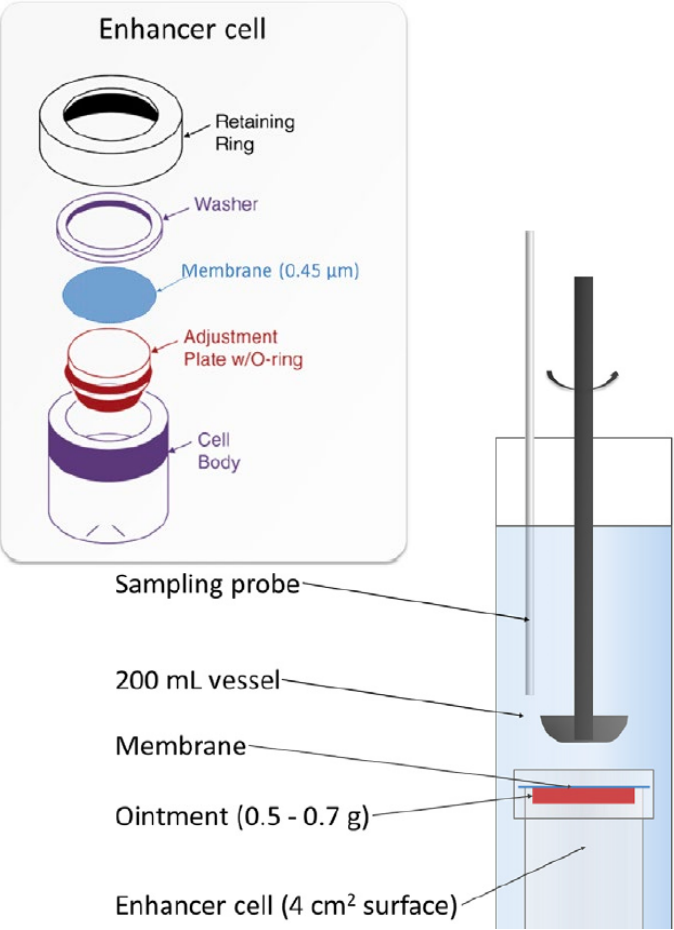
Challenges	Considerations
Low drug content requiring sensitive analytical approaches	LC/MS
Limited drug release, mainly from surface layer	Large surface area for in vitro release testing, membrane binding
Lack of compendial in vitro drug release testing methods	Apparatus setup, sample adaptors
Incomplete understanding on the impact of formulation properties on drug release profiles	Drug hydrophilicity, crystallinity, polymorphism, source, particle size
Lack of correlation between <i>in vitro</i> release, <i>ex vivo</i> test, and <i>in vivo</i> performance	Appropriate <i>in vitro</i> release, <i>ex vivo</i> studies, understand the correlation with <i>in vivo</i>



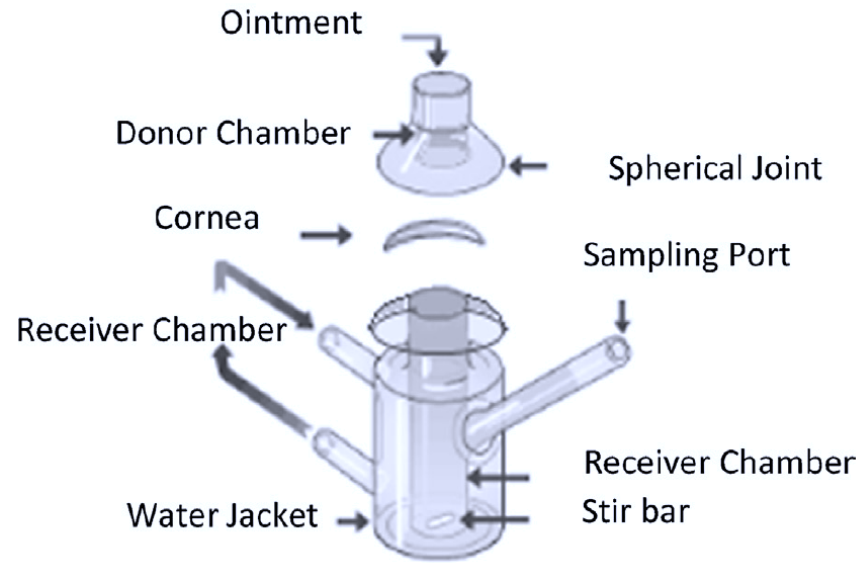
EVALUATION OF INVITRO RELEASE APPARATUS SETUPS



In vitro release testing (IVRT) for ophthalmic ointments

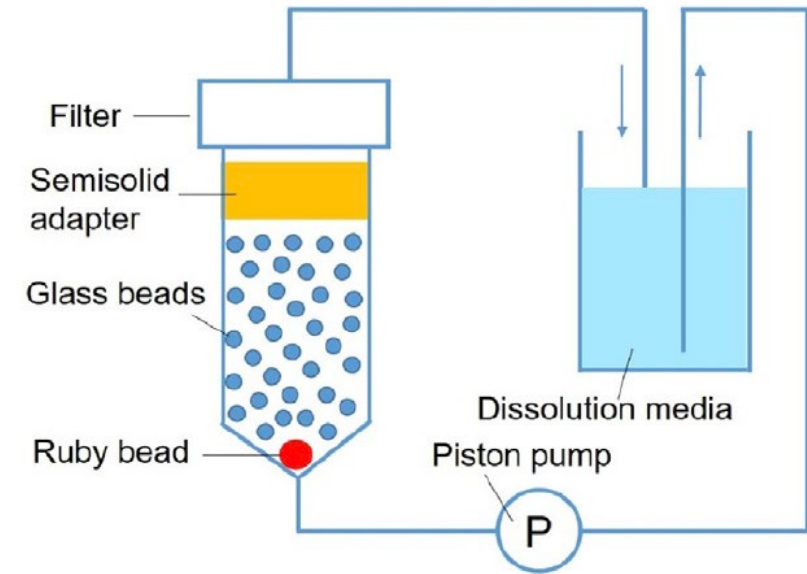


A: USP apparatus 2 with enhancer cells



B: Franz diffusion cells

Al-Ghabeish et al., 2015



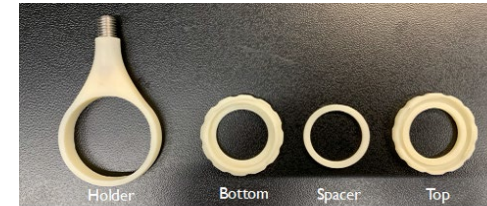
C: USP apparatus 4 with semisolid adapters



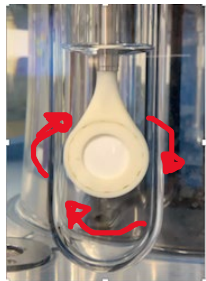
Dissolution Apparatus Optimization

5

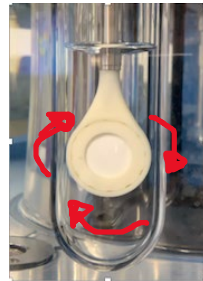
Objective: To define optimal dissolution apparatus to study drug release from ophthalmic ointments and discriminate formulations.



Compartments of two-side semisolid adapter
surface area = 3.15 cm^2 /each side



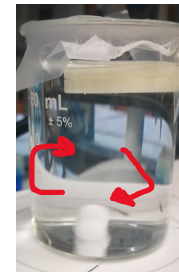
VS



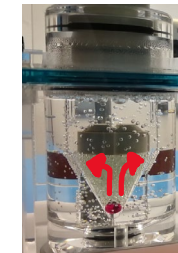
VS



VS



VS



Factors:

- 1) Surface area
- 2) Flow direction, agitation

USP 1
2- side adapter
Surface area = 6.30 cm^2

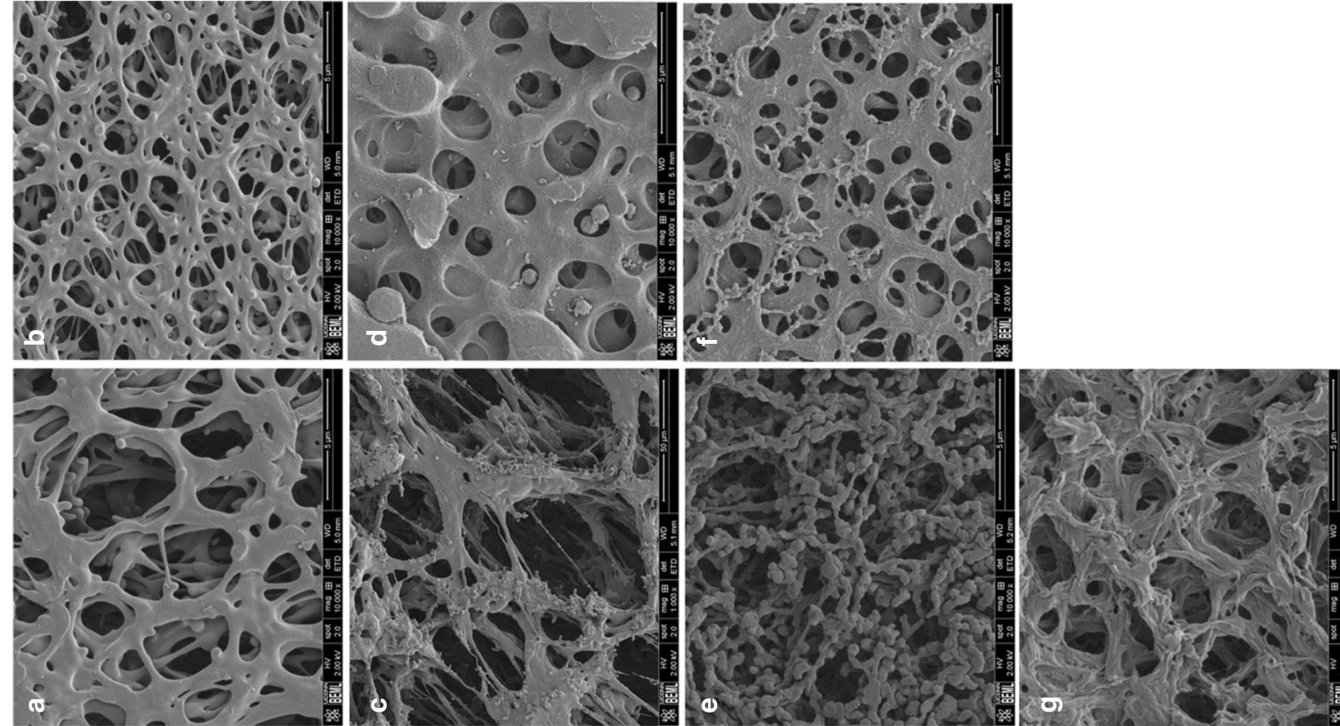
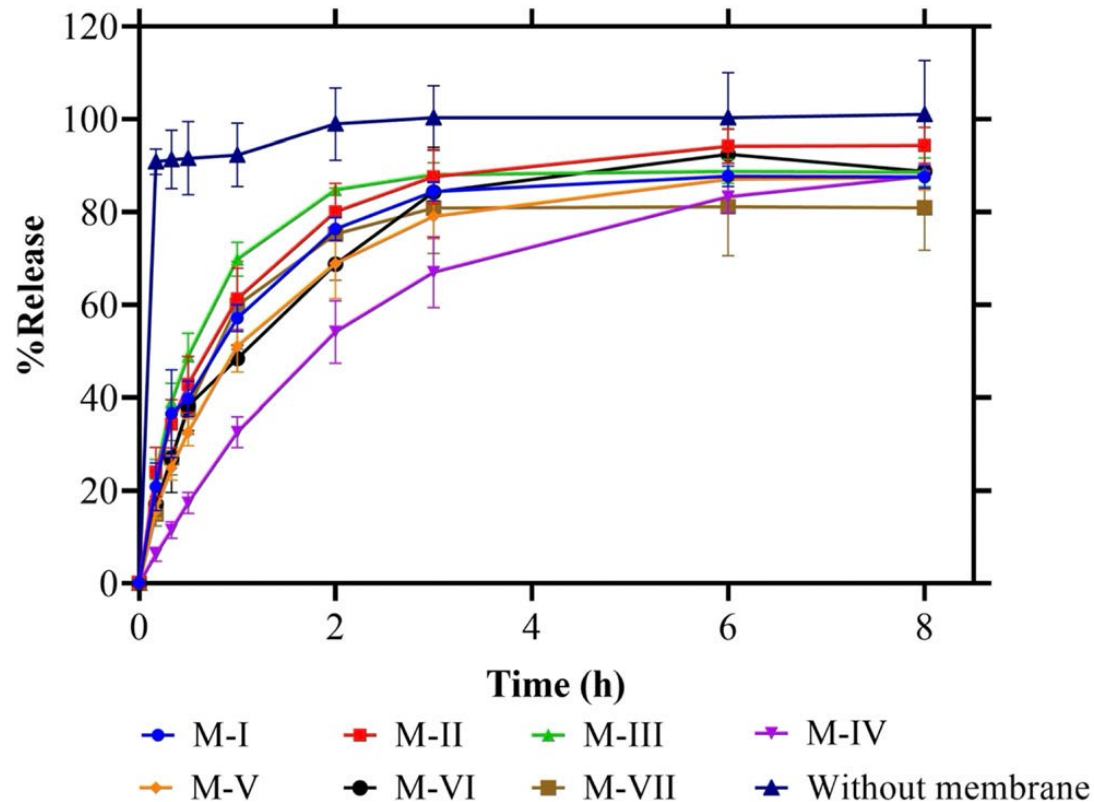
USP 1
2- side adapter using
plastic block one side
Surface area = 3.15 cm^2

USP II
1- side adapter
Surface area = 1.77 cm^2

2- side adapter: MB
facing to media using
magnetic stirrer
Surface area = 3.15 cm^2

USP IV
1- side adapter
Surface area = 1.77 cm^2

Membrane Impact on Drug release

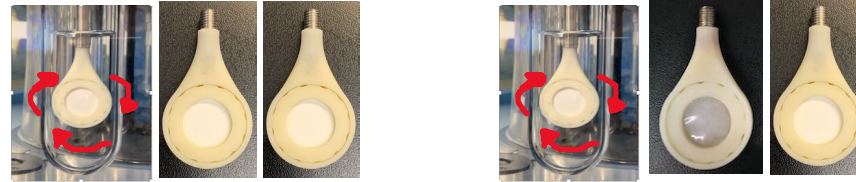


1.2 μm PES member was selected due to low drug binding and high differentiating ability

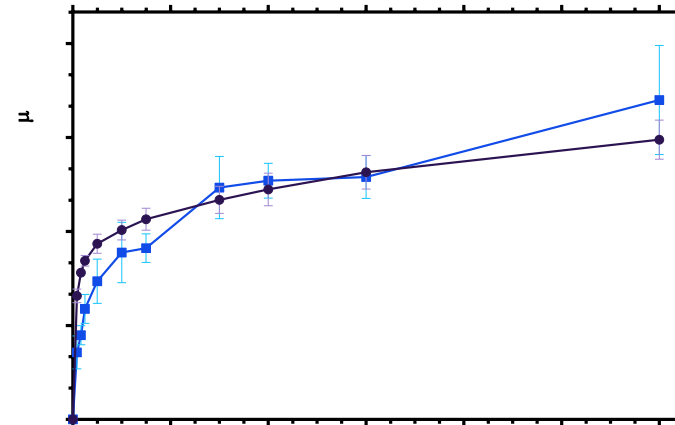
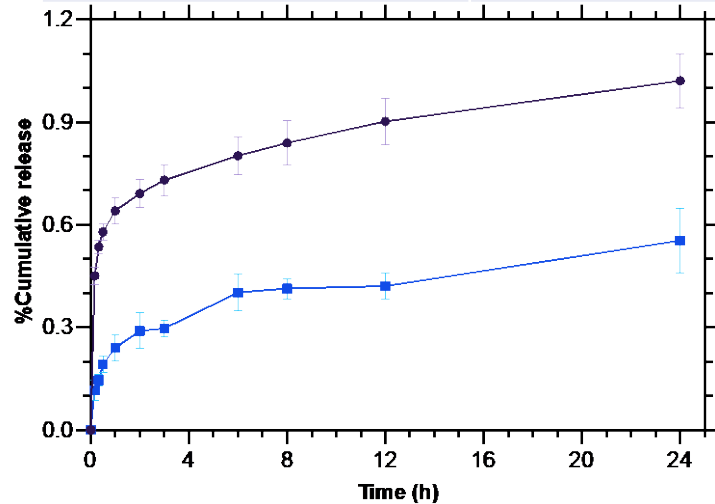
Mekjaruskula C, Berings AO, Luo W, Xu Q, Halquist M, Qin B, Wang Y, Lu X. Impact of Membranes on In Vitro Release Assessment: a Case Study Using Dexamethasone. AAPS PharmSci Tech. 2021 Jan 10;22(1):42.

FACTORS: 1) SURFACE AREA

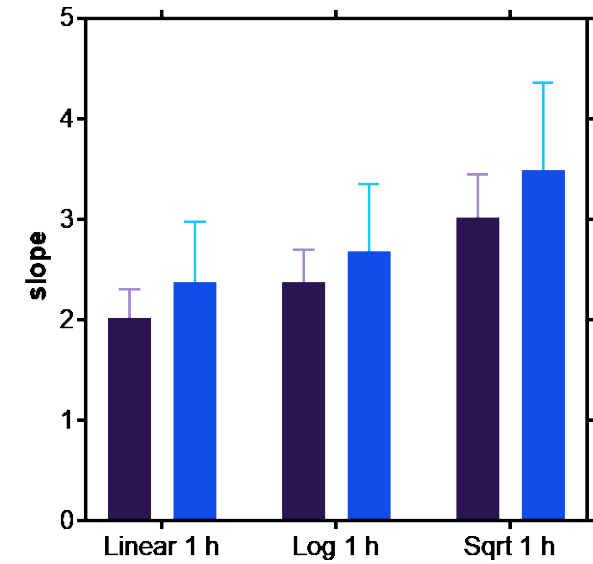
7



Parameter	USP Apparatus I	USP Apparatus I
Membrane	1.2 μm PES + 1.2 μm PES	1.2 μm PES + Plastic
Surface area	6.30 cm^2	3.15 cm^2
Release medium	STS with 0.1% Tween [®] 80, 37°C	
Ointments	1% DEX in IGI [®] 386	



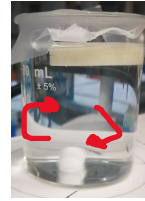
—●— 2-side (PES&PES) using USP1 —■— 2-side (PES&Plastic) using USP1



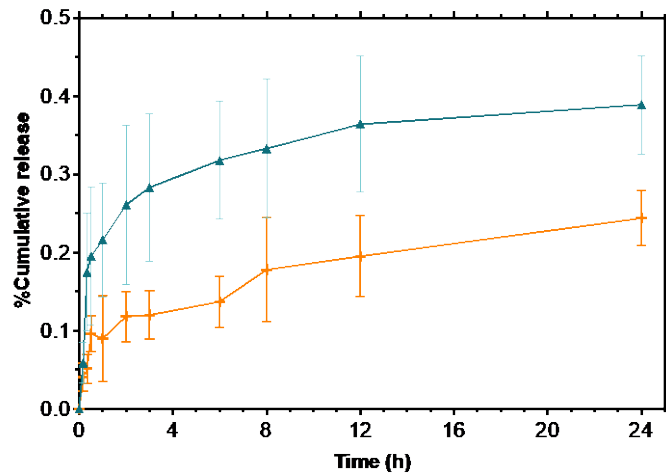
There was no significant difference between 2 models.

FACTORS: 1) SURFACE AREA

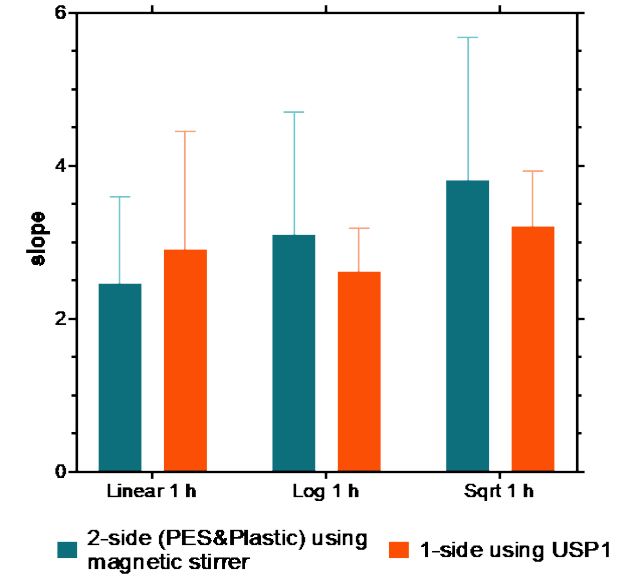
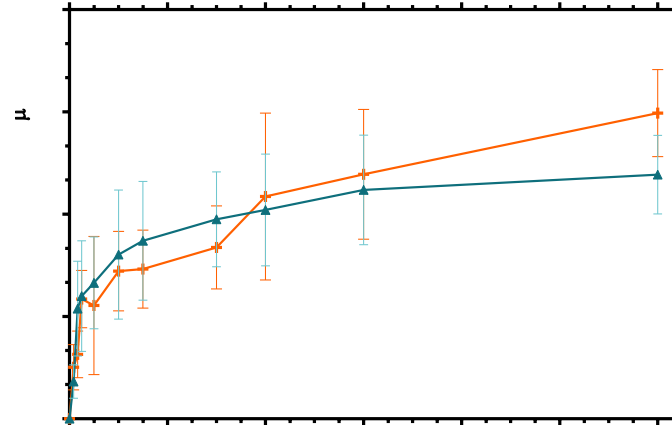
8



Parameter	USP Apparatus II	Magnetic stirrer
Adapter	1-side	2-side
Membrane	1.2 μm PES	1.2 μm PES + Plastic
Surface area	1.77 cm^2	3.15 cm^2
Release medium	STS with 0.1% Tween [®] 80, 37°C	
Ointments	1% DEX in IGI [®] 386	

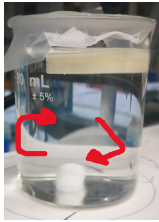
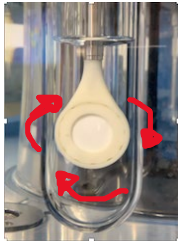


— 2-side (PES&Plastic) using magnetic stirrer — 1-side using USP1

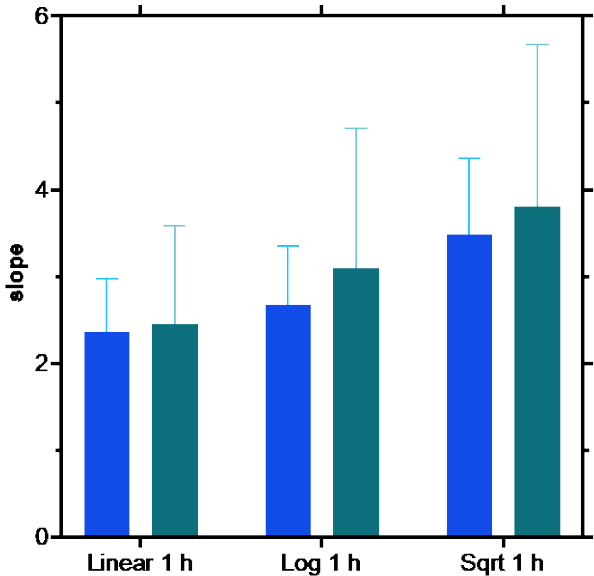
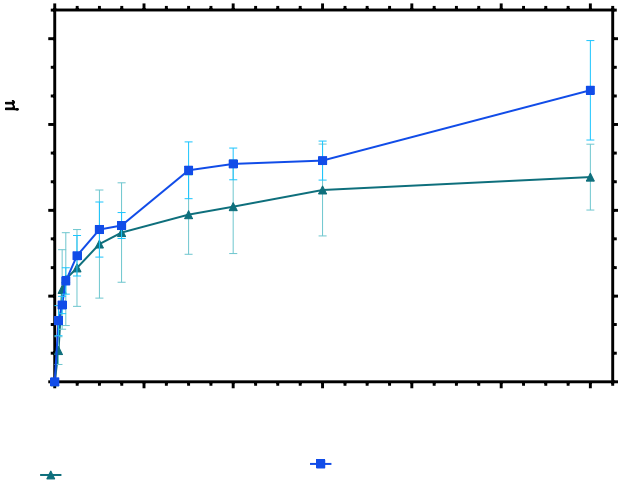
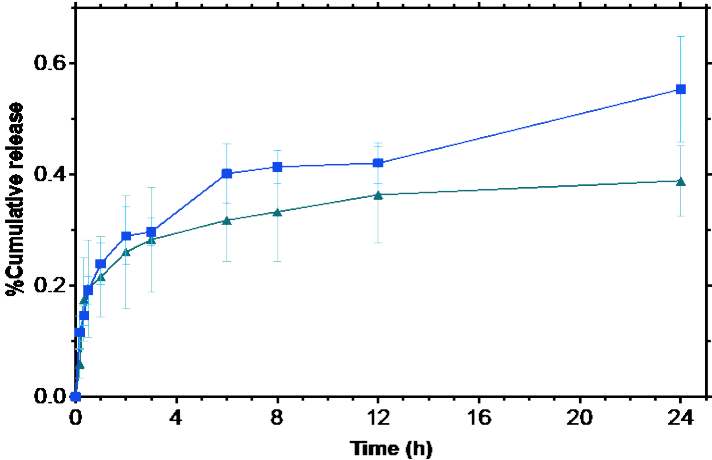


There was **no significant difference** between 2 models.

FACTORS: 2) FLOW DIRECTION

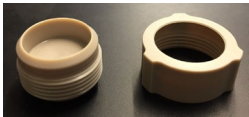


Parameter	USP Apparatus I	Magnetic stirrer
Membrane	1.2 μm PES + Plastic	1.2 μm PES + Plastic
Surface area	3.15 cm^2	3.15 cm^2
Release medium	STS with 0.1% Tween [®] 80, 37°C	
Ointments	1% DEX in IGI [®] 386	

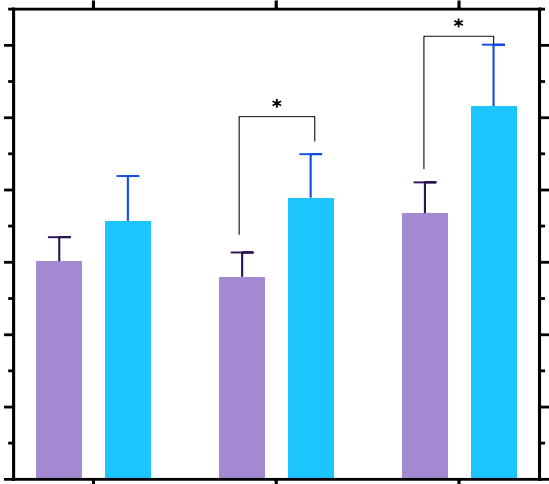
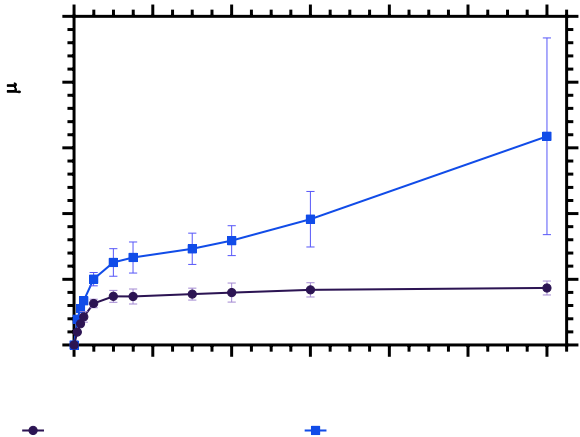
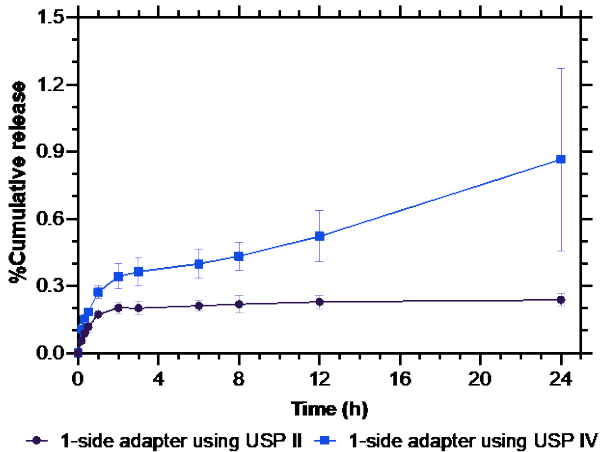


There was no significant difference between 2 models.

FACTORS: 2) FLOW DIRECTION

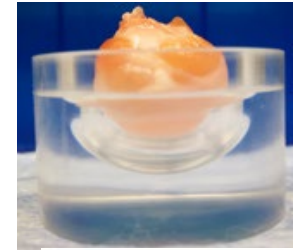
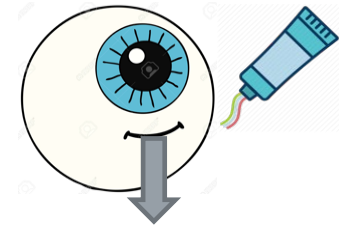


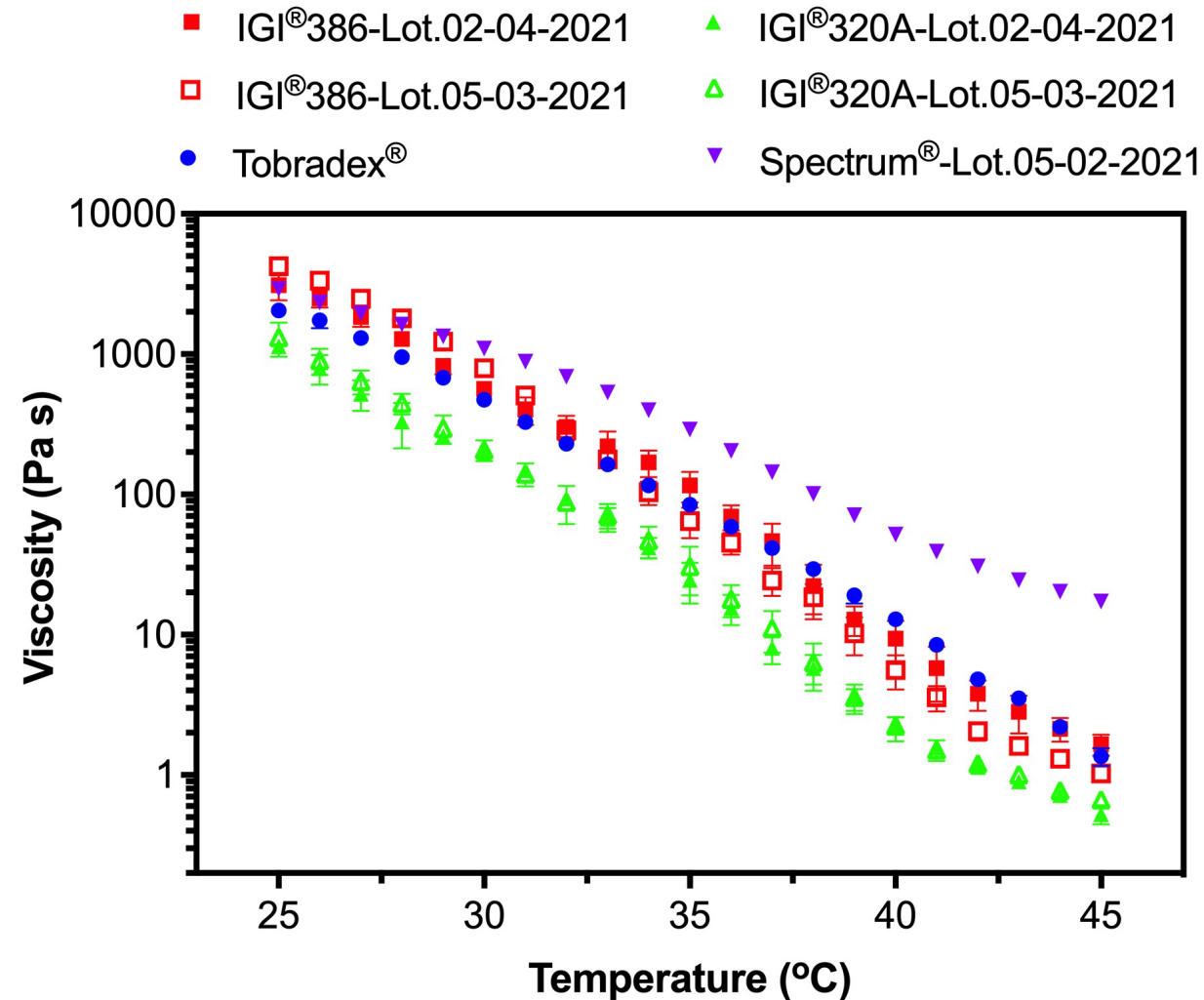
Parameter	USP Apparatus II	USP Apparatus IV
Adapter	1-side	1-side
Membrane	1.2 µm PES	1.2 µm PES
Surface area	1.77 cm ²	1.77 cm ²
Release medium	STS with 0.1% Tween® 80, 37°C	
Ointments	10% DEX in IGI® 386	



- Larger surface area, higher percent cumulative drug release
- Release amount and rate per surface area were consistent
- Agitated flow of USP IV **enhanced** hydrophobic drug (dexamethasone) release from the ointments and increased its release rates when compared to the immersion cells.

VITRO, EX-VIVO AND IN VIVO ASSESSMENT COMPARISON





IN VITRO RELEASE TESTING

Dissolution Parameters	USP Apparatus I
Samples	1) Tobradex [®] ointments 0.3% TOB/0.1% DEX 2) 0.3% TOB/0.1% DEX in IGI [®] 386 3) 0.3% TOB/0.1% DEX in IGI [®] 320A 4) 0.3% TOB/0.1% DEX in Spectrum [®]
Weight of sample	0.58 g/adaptor
Release medium	80 mL of STS <u>without</u> 0.1% Tween 80 80 mL of STS <u>plus</u> 0.1% Tween 80
Temperature	40°C
pH	7.4
Stirring Speed	200 rpm
Membrane	1.2 µm PES from Sterlitech [®]
Dissolution apparatus	3D-Printed Two-side Adapter in USP apparatus I
Aliquot removed	1 mL (<u>reconstitute replaced</u> with 1 mL <u>fresh medium</u> after withdrawn)
Sampling times	5, 10, 15, 30, 45 min, 1, 2, 4, 7 h



3D-Printed Two-side Adapter in USP apparatus I

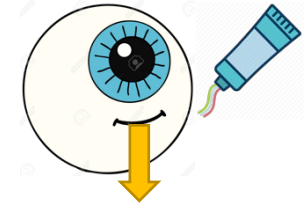
0.3% TOB/0.1% DEX ointments

Rx

TOB	0.3%
DEX	0.1%
Mineral oil	5%
Chlorobutanol	0.5%
Petrolatum qs to	100%

Ex vivo release study

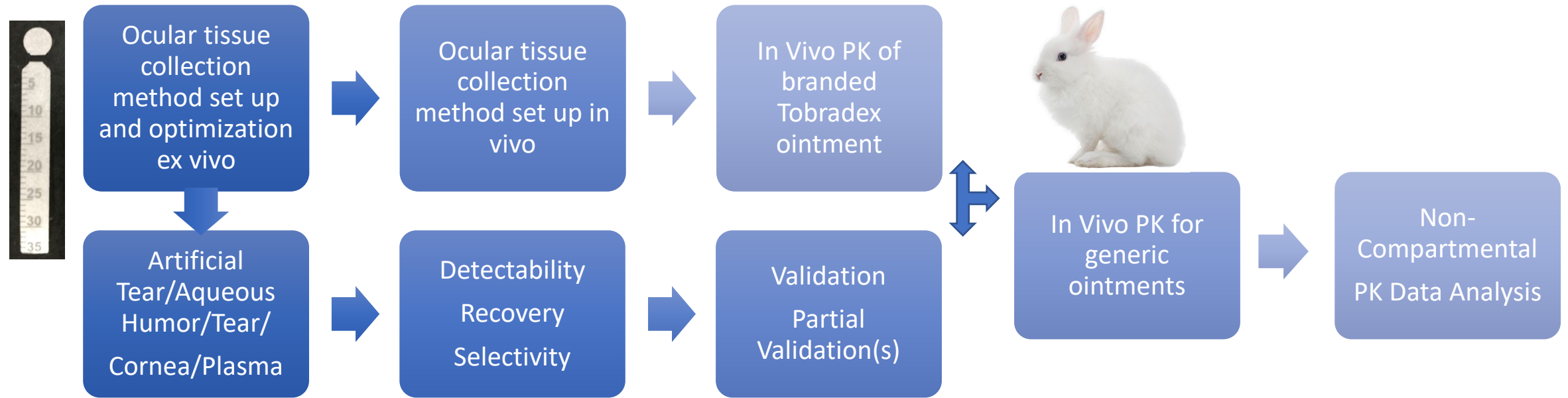
Eyes	Frozen rabbit eyes, thaw & incubate in STS 1 h (n=3)
Ointments	1) Tobradex® ointments 0.3% TOB/0.1% DEX 2) 0.3% TOB/0.1% DEX in IGI® 386 3) 0.3% TOB/0.1% DEX in IGI® 320A 4) 0.3% TOB/0.1% DEX in Spectrum® Apply 30 mg on the eye
Media	STS without Tween® 80 (0.1%) 650 µL
Temperature	34°C
Timepoints	5, 10, 15, 30, 45 min, 1, 2, 4 h



- Release media
- Aqueous humor
- Cornea

[Tissue collection and extraction were performed following VCU method.]

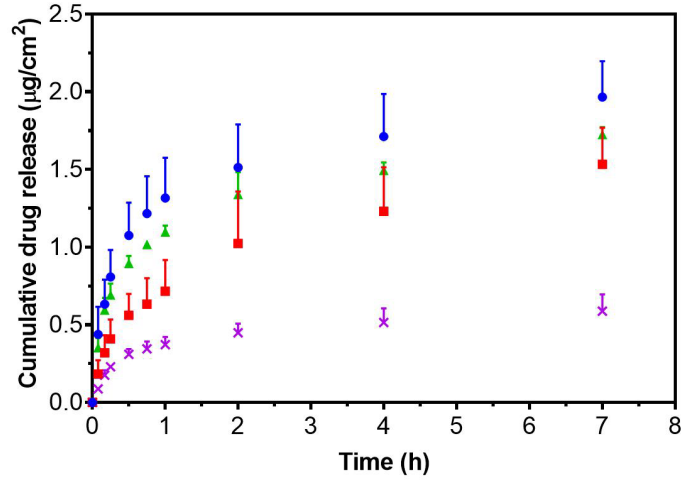
Pharmacokinetics studies in rabbits



Meng T, Kosmider L, Chai G, Moothedathu Raynold AA, Pearcy AC, Qin B, Wang Y, Lu X, Halquist MS, Xu Q. LC-MS/MS method for simultaneous quantification of dexamethasone and tobramycin in rabbit ocular biofluids. J Chromatogr B. Analyt Technol Biomed Life Sci. 2021 Apr 30;1170:122610. Epub 2021 Mar 1.

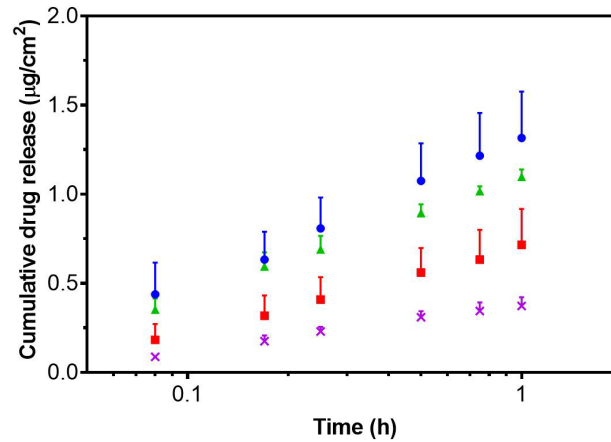
STS without Tween80

In vitro release testing_DEX
STS without Tween 80



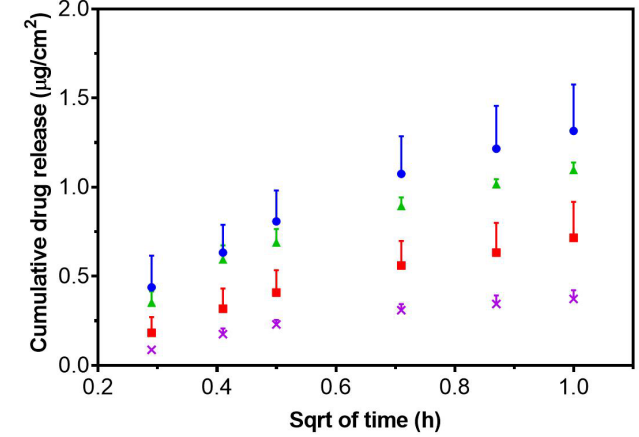
● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

In vitro release testing_DEX_Logarithm
STS without Tween 80



● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

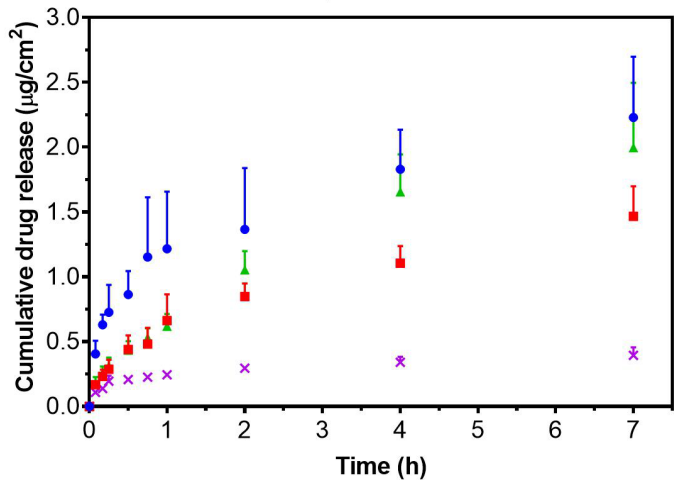
In vitro release testing_DEX_Sqrt
STS without Tween 80



● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

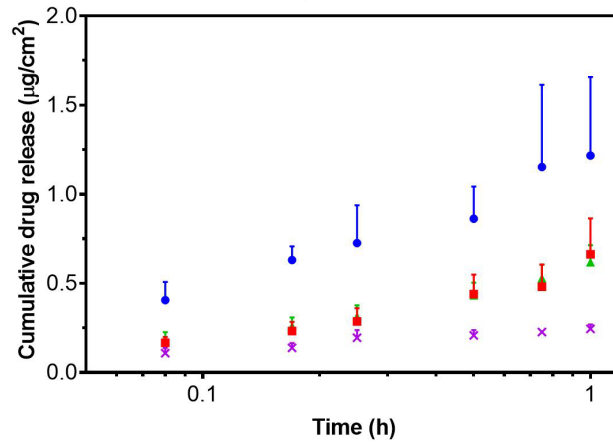
STS plus Tween80

In vitro release testing_DEX
STS plus Tween 80



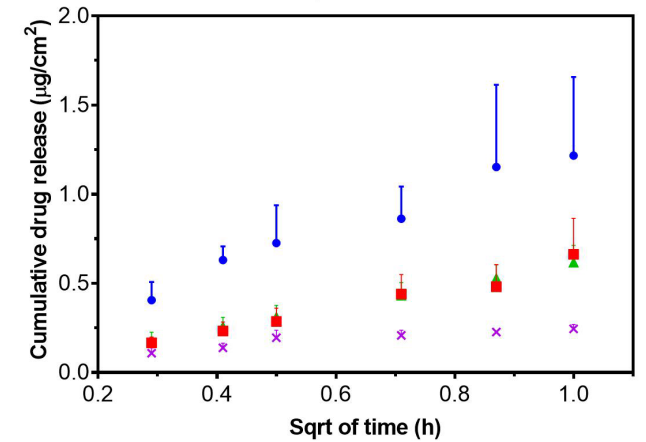
● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

In vitro release testing_DEX_Logarithm
STS plus Tween 80



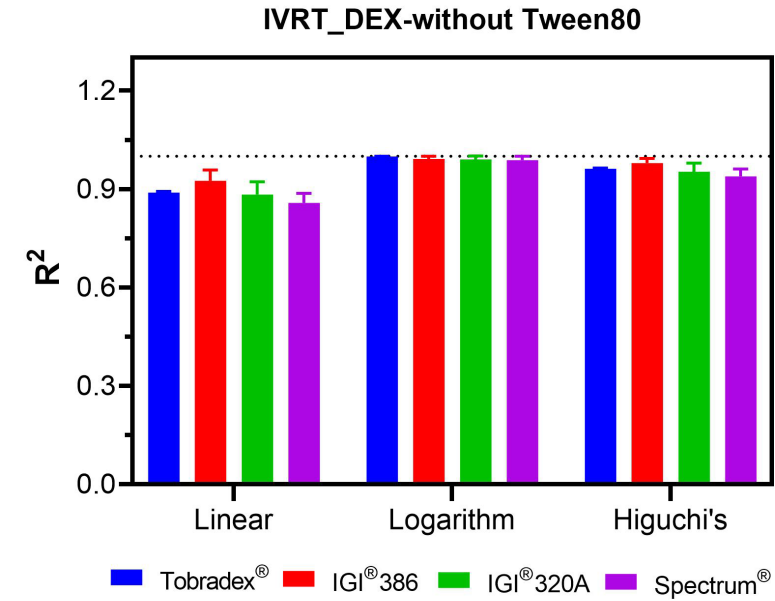
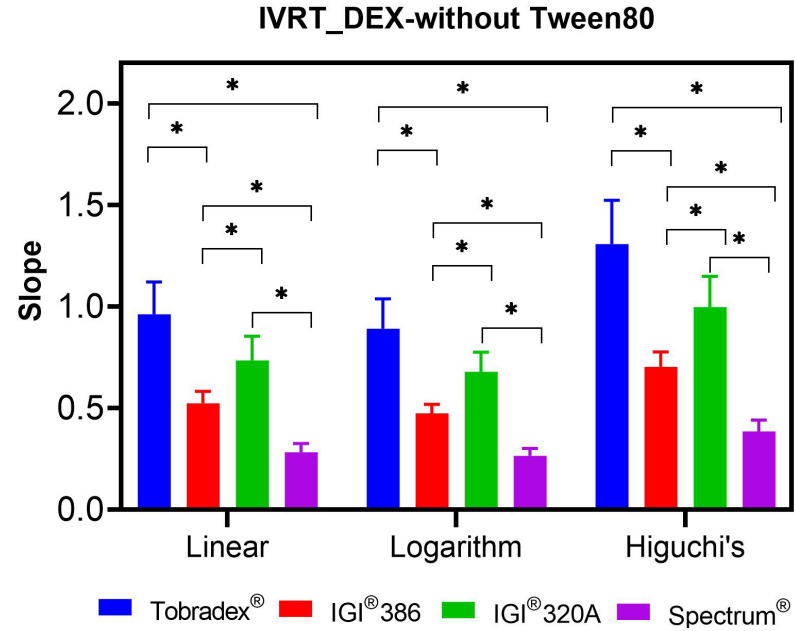
● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

In vitro release testing_DEX_Sqrt
STS plus Tween 80

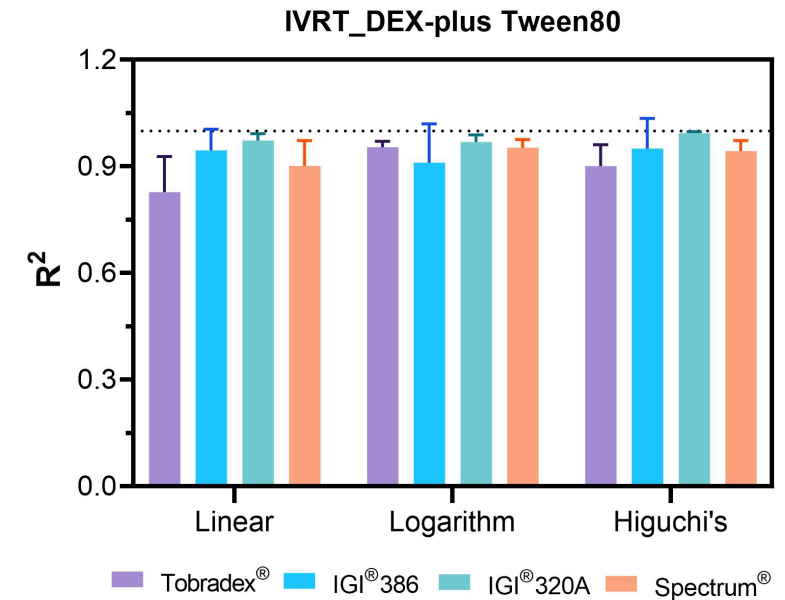
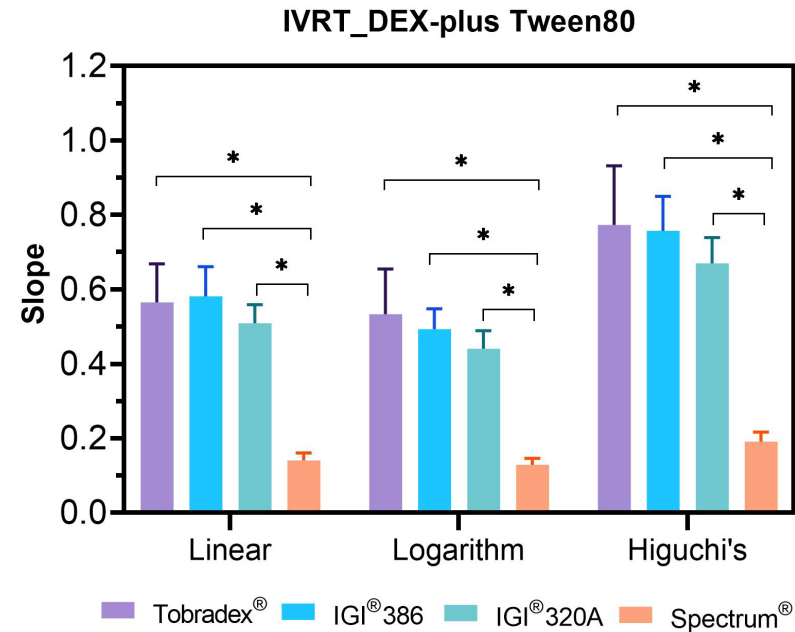


● Tobradex® ■ IGI®386 ▲ IGI®320A × Spectrum®

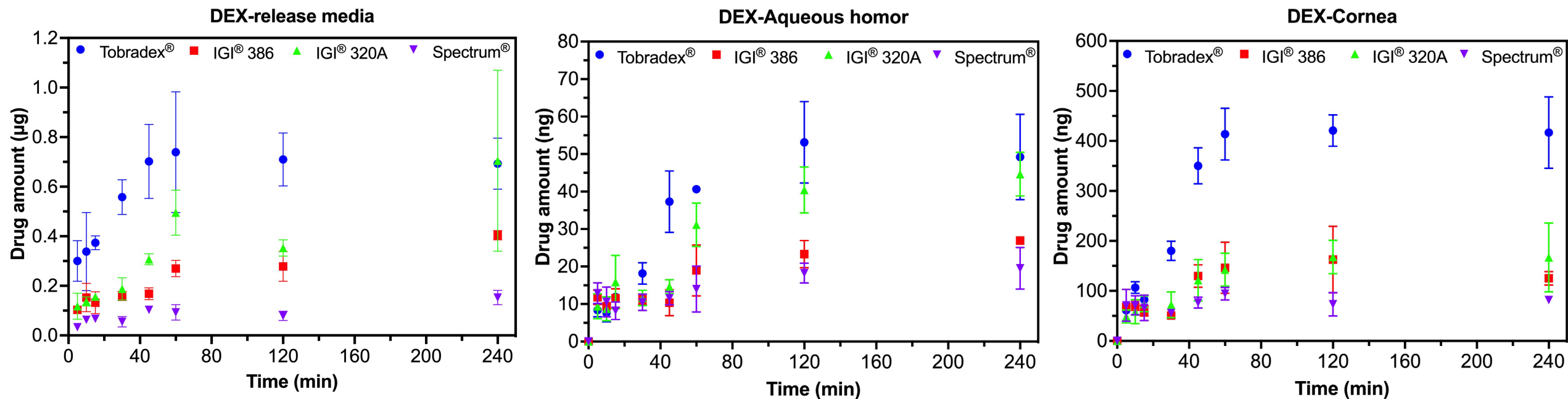
STS without Tween80



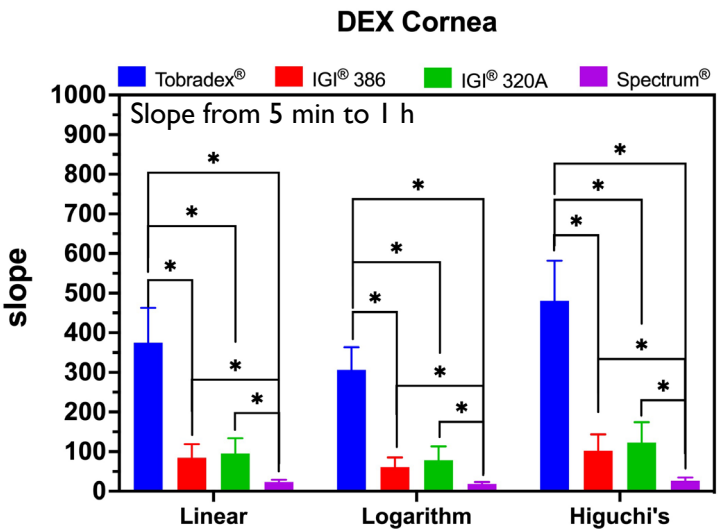
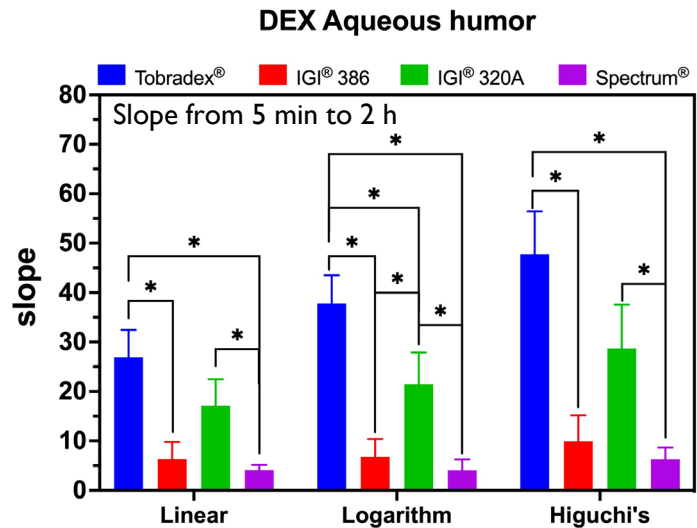
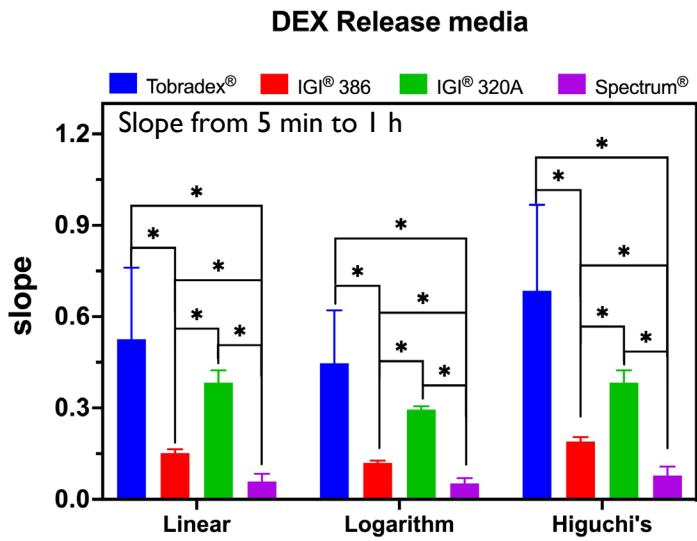
STS plus Tween80



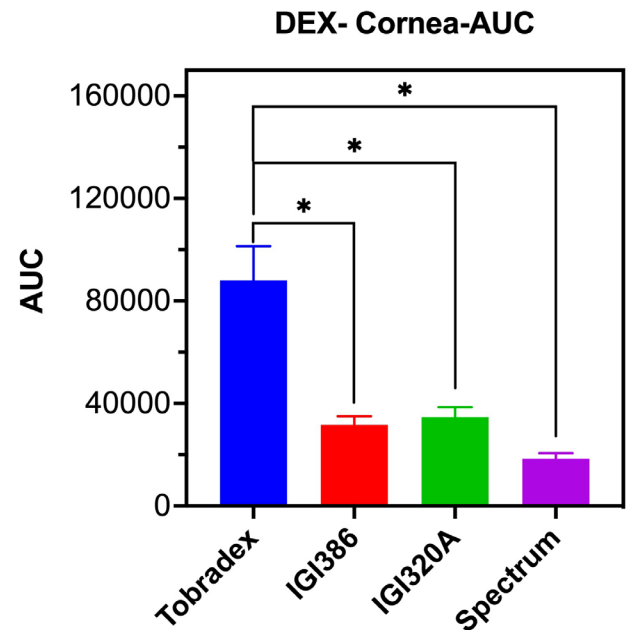
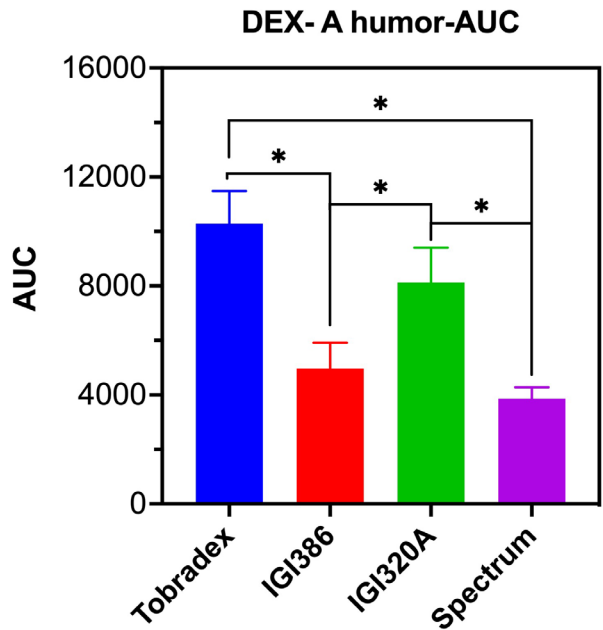
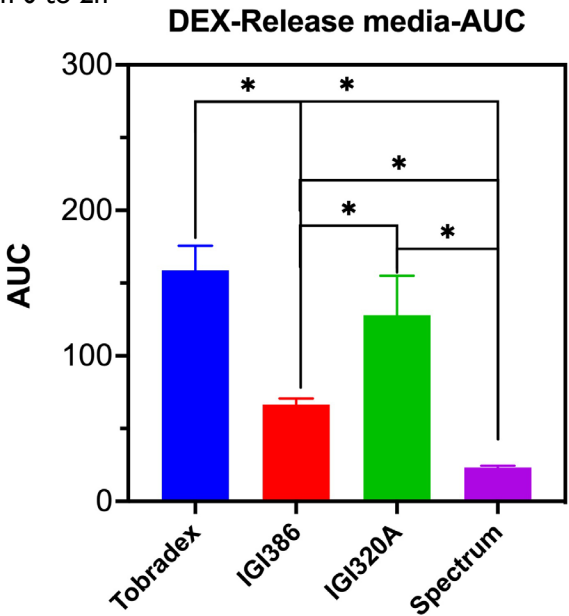
EX VIVO RELEASE : DEXAMETHASONE



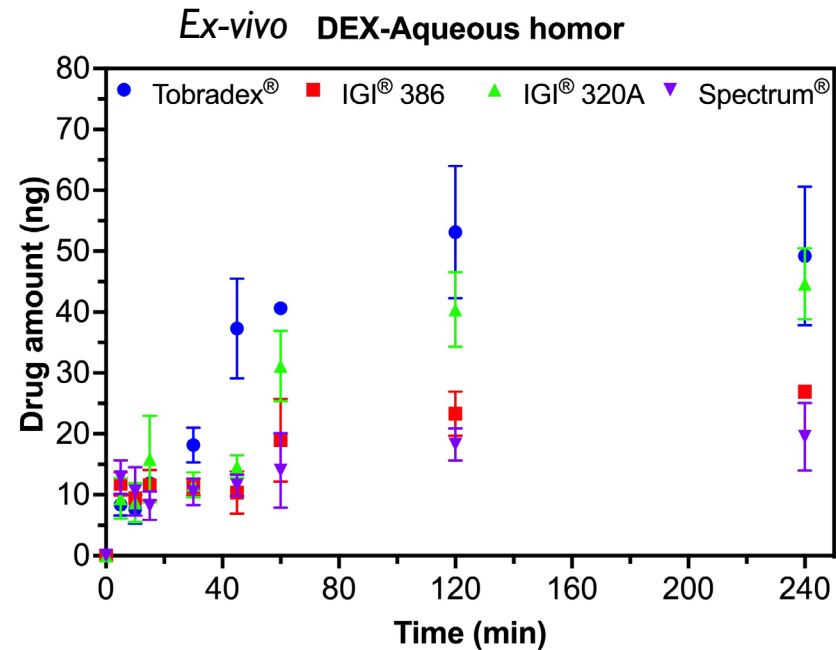
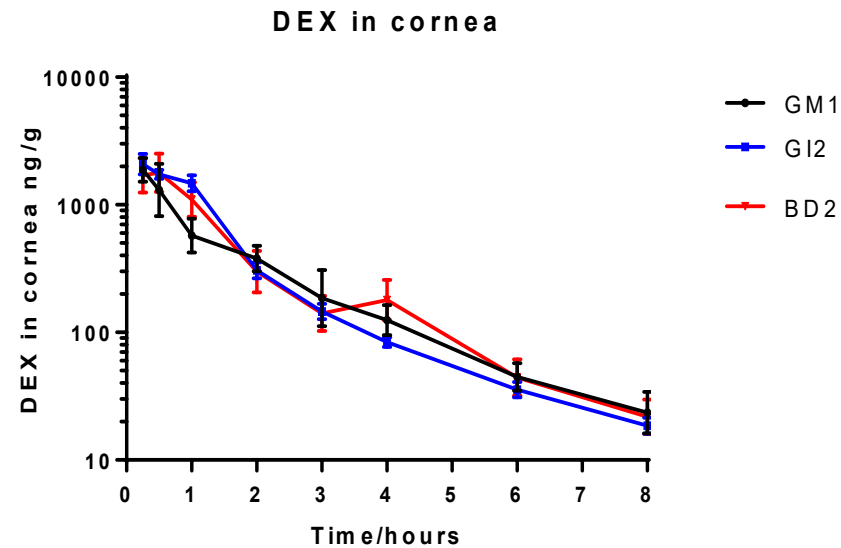
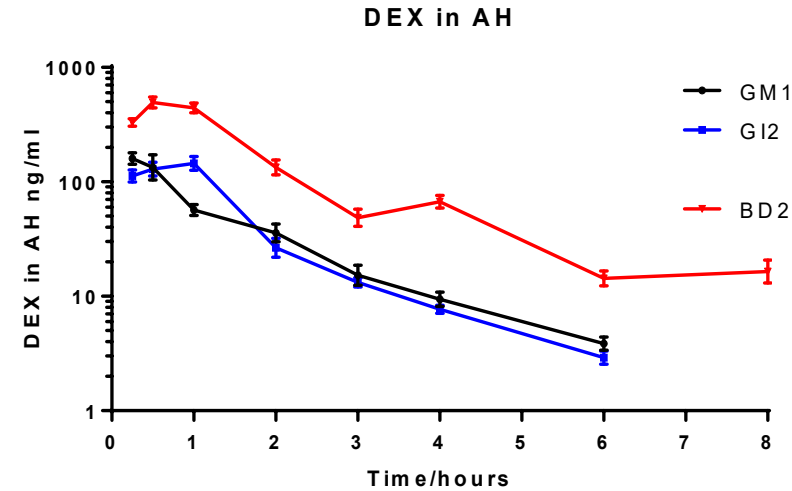
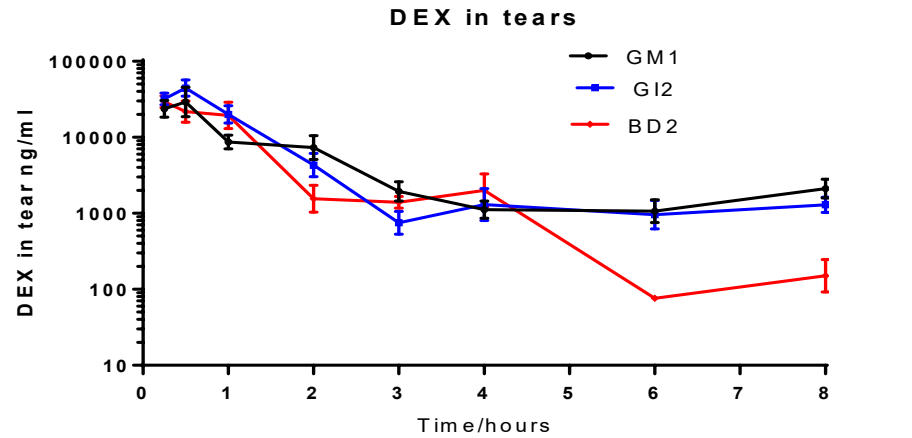
- Release media and aqueous humor : Tobradex > IGI 320A > IGI 386 > Spectrum
- Cornea: Tobradex > IGI 320A = IGI 386 > Spectrum



*AUC from 0 to 2h



In vivo (rabbits) results



GM1: 1st generic formulation, Spectrum

GI2: 2nd generic formulation, IGI386

BD: brand name

SUMMARY ON THE ASSESSMENT METHODS

Method	Dexamethasone	Sensitivity for Differentiation
IVRT using STS without Tween80	Tobradex [®] > IGI [®] 386, Tobradex [®] > Spectrum [®] at p<0.05	Very sensitive
IVRT using STS with Tween80	Tobradex [®] > Spectrum [®] at p<0.05	Sensitive
<i>Ex vivo</i> study	- Release media and aqueous humor :Tobradex > IGI 320A > IGI 386 > Spectrum - Cornea:Tobradex > IGI 320A = IGI 386 > Spectrum	Sensitive and closer to <i>in vivo</i> results and also can cover the variations <i>in vivo</i>
<i>In vivo</i> study in rabbits	- No difference between generics - AH:Tobradex 2> generics - Cornea:Tobradex 1>generics	Can be variable



THANK YOU!

Yan Wang, Ph.D. (FDA)
Bin Qin, Ph.D. (FDA)
Catheleeya Mekjaruskul, Ph.D

