

## Product Characterization and In Vitro Testing for Establishing Equivalence of Complex Products

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> SESSION 1: Equivalence of Complex Products FY 2017 GDUFA Regulatory Science Initiatives Public Workshop

## **Complex Products**



- Complex active ingredients
  - Complex mixtures of APIs, polymeric compounds, peptides
- Complex formulations
  - Liposomes, suspensions, emulsions, gels
- Complex routes of delivery
  - Locally acting such as dermatological and inhalational drugs
- Complex dosage forms
  - Long acting injectables and implantables, transdermals, MDIs
- Complex drug-device combinations

## Scope of this Session



- Complex active ingredients
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# **Complex Active Ingredients**



- Research activities
  - External: grants/contracts on pentosan polysulfate sodium and crofelemer
  - Internal: peptide related impurity analysis and immunogenicity evaluations, sucralfate, high dimensional/multivariate data comparison

### Regulatory outcomes

- Product Specific Guidance: colesevelam, omega-3 carboxylic acids, glatiramer acetate, ethiodized oil
- Guidance agenda 2017: Submission of ANDAs for Certain Highly Purified Synthetic Peptide Drug Products That Reference Peptide Drug Products of rDNA Origin

### LC-MS and MS/MS of Salmon Calcitonin



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# LC-HRMS vs USP LC-UV



- For the calcitonin RLD LC-HRMS identified 12 impurities for a total of 2.6% (Area%)
- The same sample analyzed by the USP HPLC-UV method observe 6 impurities with a 2.0% total
- Detection limits for the 2 identified peptide impurities were below 0.1% (Area %) by LC-HRMS

## Cell Based Assays to Detect IIRMIs in Drug Products





IIRMIs: innate immune response modulating impurities Haile LA, Puig M, Kelley-Baker L, Verthelyi D (2015) PLoS ONE 10(4)

## **Complex Formulations**











Characterizations of Complex Formulations

- Development of advanced analytical techniques
  - Characterize critical attributes for product equivalence, functional excipients, and bioanalytical methods for different forms of drugs *in vivo*

170 160 150 140 130 120 110 100 90



#### From product-specific guidance of risperidone injection

The proposed parenteral drug product should be qualitatively (Q1) and quantitatively (Q2) the same as the reference product for all strengths (12.5 mg/vial, 25 mg/vial, 37.5 mg/vial, and 50 mg/vial). Please provide characterization data on poly(lactide-co-glycolide) (PLGA) for both the test and reference product including polymer composition (ratio between glycolic acid and lactic acid), molecular weight and weight distribution, and PLGA architecture (e.g., linear or star-branched PLGA). Additional data on PLGA characterization may be requested during the review of the ANDA.

#### Physiochemical Equivalence Assessment of Reference and Generic Sodium Ferric Gluconate Complex

#### Dynamic Light Scattering (DLS):

Drug product (Lot #)	Z-average diameter (nm)	Intensity-weighted diameter (nm)	Volume-weighted diameter (nm)	PDI Value
Ferrlecit <sup>®</sup> (D2C283A)	11.5	13.9	9.0	0.163
Ferrlecit <sup>®</sup> (D2C593A)	12.1	14.5	8.8	0.158
Generic SFG (132296.1)	10.5	12.1	8.1	0.123

#### Cryogenic Transmission Electron Microscopy (Cryo-TEM):



Atomic Force Microscopy (AFM):





FDA internal study

#### Physiochemical Equivalence Assessment of Reference and Generic Sodium Ferric Gluconate Complex

Gel Permeation Chromatography (GPC):

Drug product (Lot #)	M <sub>w</sub> (kDa)	
Ferrlecit (D2C283A)	384.7 ± 5.1	
Ferrlecit (D2C593A)	393.4 ± 1.9	
Ferrlecit (A5075)	467.7 ± 3.0	
Generic SFG (132996.1)	387.4 ± 2.1	
Generic SFG (142241.1)	365.9 ± 5.4	
Generic SFG (142290.1)	363.7 ± 1.9	

Analytical Ultracentrifugation (AUC):



Asymmetric filed flow fractionation – multi-angle laser scattering (AFFF-MALS):

Drug product (Lot #)	Run	M <sub>n</sub> [kDa]	M <sub>w</sub> [kDa]	M <sub>w</sub> /M <sub>n</sub>
Ferrlecit <sup>®</sup> (D2C283A)	1	83.5 ± 2.3	316.7 ± 0.9	3.8
Ferrlecit <sup>®</sup> (D2C283A)	2	88.8 ± 2.6	317.8 ± 1.3	3.6
Ferrlecit <sup>®</sup> (D2C283A)	3	87.4 ± 2.1	319.1 ± 1.3	3.6
Ferrlecit <sup>®</sup> (D2C593A)	1	98.9 ± 1.5	329.1 ± 0.7	3.3
Ferrlecit <sup>®</sup> (D2C593A)	2	92.7 ± 2.4	329.9 ± 1.6	3.6
Ferrlecit <sup>®</sup> (D2C593A)	3	92.7 ± 2.5	330.7 ± 1.3	3.6
Generic SFG (132296.1)	1	218.4 ± 0.7	415.6 ± 1.2	1.9
Generic SFG (132296.1)	2	219.6 ± 0.7	418.3 ± 1.3	1.9
Generic SFG (132296.1)	3	222.2 ± 0.7	417.7 ± 1.3	1.9



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Characterizations of Complex Formulations

- Study impact of manufacturing and formulation processes on the end product's critical quality attributes
  - Liposomes
  - Microspheres
  - Implants/inserts



Sample	Solvent	Preparation method	Porosity (%)	105
Risperdal Consta			43.97 ± 4.60	90 - (%) 95 -
F1	DCM	Homogenization & dry sieving	43.19 ± 4.60	one Relea
F2	DCM	Homogenization & wet sieving	46.04 ± 42.90	sperido
F3	EA	Vortex & wet sieving	54.98 ± 1.25	10 - 00 -
F4	EA	Homogenization & wet sieving	61.75 ± 1.08	15 -

Shen J, et al. J Control Release. 2015 Nov 28;218:2-12 Shen J, et al. Int J Pharm. 2016 Feb 10;498(1-2):274-82



10 mM PBS (pH 7.4), 37°C

## In Vitro Release Testing



- Development of new methods for *in vitro* release testing
  - Quality control
  - In vitro in vivo correlation
  - Various products: ophthalmic suspensions/ointments, periodontal inserts, parenteral suspensions, microspheres and implants, intrauterine systems...
  - Different methodologies: pulsatile microdialysis (PMD), modified USP II, USP IV, macro-fabricated flow cells

# Critical Attributes and In Vitro Tests for Ophthalmic Drug Products



Urtti A, et al. AAPS 2016; Grant 1U01FD005180-01 Sailor MJ, et al. CRS 2016; Grant 1U01FD005173-01



In vivo animal tests to measure how formulation properties affect local pharmacokinetics



14

# Cage model to assess in vivo release of microspheres



15

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## **IVIVC of Risperidone Microspheres**



Shen J, et al. J Control Release. 2015 Nov 28;218:2-12

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## Summary



- Access to complex generics is accelerated by analytical advances that:
  - Ensure equivalence of critical attributes
  - Enable alternatives to in vivo BE studies
- Two categories of advances
  - Characterization
    - New technology and new characteristics
    - New analysis methods for complex data
  - In vitro performance testing
    - Biological tests to ensure equivalence of proposed generic products
    - Release tests under similar physiological conditions

## **Priorities for the Panel**



- New advanced analytics for characterization of chemical compositions, molecular structures and distributions in complex active ingredients
- Predictive in silico, in vitro and animal studies to evaluate immunogenicity risk of formulation or impurity differences in generic products
- Particle size, shape and surface characterization based bioequivalence for suspended and colloidal drug products
- Predictive in vitro BE methods for long-acting injectables

