### **FOOD AND DRUG ADMINISTRATION (FDA) Center for Drug Evaluation and Research (CDER)**

Bone, Reproductive, and Urologic Drugs Advisory Committee (BRUDAC) Meeting Tommy Douglas Conference Center, 10000 New Hampshire Ave, Silver Spring, MD December 6, 2016

#### **AGENDA**

The committee will discuss appropriate clinical trial design features, including acceptable endpoints for demonstrating clinical benefit, for drugs intended to treat secondary hypogonadism while preserving or improving testicular function, including spermatogenesis.

8:00 a.m.	Call to Order and Introduction of Committee	Vivian Lewis, MD (Chairperson), BRUDAC
8:10 a.m.	Conflict of Interest Statement	CDR LaToya Bonner, PharmD, NCPS Acting Designated Federal Officer, BRUDAC
8:15 a.m.	FDA Opening Remarks	Hylton V. Joffe, MD, MMSc Director, Division of Bone, Reproductive and Urologic Products (DBRUP) Office of Drug Evaluation III (ODE III) Office of New Drugs (OND), CDER, FDA
8:30 a.m.	GUEST SPEAKER PRESENTATION	
	Treatment of Secondary Hypogonadism	Sergio Oehninger, MD, PhD Director, Division of Reproductive Endocrinology The Jones Institute for Reproductive Medicine Eastern Virginia Medical School
9:10 a.m.	Clarifying Questions to the Guest Speaker	
9:25 a.m.	INDUSTRY PRESENTATIONS	
	Introduction	Michael Wyllie, PhD Managing Director, Global Pharma Consulting, Ltd. Introduction
	Treatment Considerations for Secondary Hypogonadism	Mohit Khera, MD Associate Professor of Urology Baylor College of Medicine
	Sperm Concentration is an Acceptable Endpoint for Demonstrating Clinical Benefit in Men who Have Hypogonadotropic Hypogonadism and Oligozoospermia (Impaired Spermatogenesis) as a Cause of Male Infertility	Edward Kim, MD Professor of Surgery University of Tennessee Graduate School of Medicine

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#### AGENDA (cont.)

	INDUSTRY PRESENTATIONS (CONT.)		
	Human Chorionic Gonadotropin	Mohit Khera, MD	
	Diagnostic Categories of Hypogonadism and Secondary Hypogonadal Population	Frederick Wu, MD Professor of Medicine and Endocrinology University of Manchester	
	Weight Associated, Secondary Hypogonadism: An acquired Estrogen- Dependent Disorder	Andrew McCullough, MD Director of Male Sexual Health, Urology Department Lahey Health and Medical Center	
	Summary and Conclusions	Michael Wyllie, PhD	
10:25 a.m.	Clarifying Questions to Industry		
10:45 a.m.	Break		
11:00 a.m.	FDA PRESENTATIONS		
	FDA Clinical Perspective on Development of Non-Testosterone Products to Treat Male Secondary Hypogonadism	Olivia Easley, MD Medical Officer DBRUP, ODE III, OND, CDER, FDA	
	Regulatory Approach to Clinical Outcome Assessment Review for Drug Development	Selena Daniels, PharmD, MS Team Leader, Clinical Outcome Assessments Staff OND, CDER, FDA	
11:40 a.m.	Clarifying Questions to the FDA	OND, CDER, PDA	
12:00 p.m.	LUNCH		
1:00 p.m.	OPEN PUBLIC HEARING		
2:00 p.m.	Clarifying Questions to the Guest Speaker, Industry or FDA		
2:30 p.m	Break		
2:45 p.m.	Questions to the Committee/Committee Discussion and Voting		
5:00 p.m.	ADJOURN		