

FDA-Industry GDUFA Reauthorization Meeting
July 7, 2016, 10:00 am – 2:45 pm
FDA White Oak Campus, Silver Spring, MD
Building 51, Room 1219

Purpose

To continue discussions from the July 6 negotiation meeting on issues pertaining to Abbreviated New Drug Applications (ANDAs) and Drug Master Files (DMFs).

Participants

FDA

Donald Beers
Robert Berlin
Ashley Boam
Mary Beth Clarke
Keith Flanagan
Michael Jones
Robert Lionberger
Edward Sherwood
Martin Shimer

OC/OCC
OC/OPPLA
CDER
CDER
CDER
CDER
CDER
CDER
CDER

Industry

John DiLoreto
David Gaugh
Marcie McClintic Coates
Alan Nicholls
Molly Rapp
Gil Roth
Cornell Stamoran
Rich Stec
Terri Stewart
Scott Tomsy
Keith Webber

BPTF
GPhA
GPhA (Mylan)
BPTF
GPhA (Fresenius-Kabi)
PBOA
PBOA (Catalent)
GPhA (Perrigo)
GPhA (Teva)
GPhA (Teva)
GPhA (Perrigo)

FDA Supporting Staff

Carter Beach, Heather Brown, Derek Griffing, Martha Nguyen, Trang Tran, Lucie Yang

Industry Supporting Staff

Mark Hendrickson (GPhA)

Discussion

FDA and Industry continued discussions from the July 6 negotiation meeting on issues pertaining to ANDAs and DMFs. Topics included review goals, ANDA review program enhancements, facility evaluations, resource management enhancements, and a pre-ANDA process (pre-ANDA meetings, product-specific guidance, and controlled correspondence).

Next Meeting

The next negotiation meeting is planned for Tuesday, July 12, 2016.