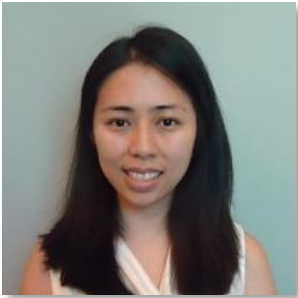


Update on the Development and Revision of USP Compounding Chapters

The presentation will provide an overview of USP's standard setting process specifically as it relates to the development and revision of compounding general chapters. The presentation will provide an overview of the proposed revisions to General Chapter <795> Pharmaceutical Compounding – Nonsterile Preparations and <797> Pharmaceutical Compounding – Sterile Preparations and how these chapters relate to <800> Hazardous Drugs – Handling in Healthcare Settings. USP continues to request stakeholders to provide input on revising these standards.



Jeanne Sun, PharmD

United States Pharmacopeia

Jeanne Sun, Pharm.D. is a Scientific Liaison and Manager at the United States Pharmacopeial Convention in Rockville, MD. In her position, she is responsible for coordinating the volunteer efforts of the Compounding Expert Committee in the development and revision of standards for nonsterile, sterile, and veterinary

compounding as well as for handling hazardous drugs. She also works with analytical laboratories in the development and validation of stability-indicating methods to establish beyond-use dates for extemporaneous preparations published in the USP-NF. She assists in the outreach efforts to inform stakeholders of new standards development related to compounding.

Prior to joining USP, she held various positions including a research associate for Advancis Pharmaceuticals, a biological aid for the United States Department of Agriculture, and a summer intern in Quality of Compounded Medicines at USP. She has over 10 years of experience in a community pharmacy and has received certification for immunization delivery. She earned a Bachelor of Science in Biochemistry from the University of Maryland, College Park and a Pharm.D. from the University of Maryland Baltimore School of Pharmacy. She is currently pursuing a J.D. at Georgetown University Law Center.