

Council for Federal Cannabis Regulation February Webinar to Focus on Implications for Research and Regulatory Science

*Next CFCR Event Scheduled for Thursday, February 24, 1:00 - 2:00
p.m. EST.*

WASHINGTON D.C., UNITED STATES, February 17, 2022
/EINPresswire.com/ -- The [Council for Federal Cannabis
Regulation](#) (CFCR) announced today the fifth installment of
their monthly web series. February's virtual conversation,
entitled The Endocannabinoid System: Implications for
Research and Regulatory Science, is designed to educate the
public, policy makers, and industry about the research and
regulatory science surrounding the endocannabinoid system –
including the cannabis-derived products that impact the
human body and their current and potential future use when
infused into everything from pharmaceuticals to cosmetics.

Moderated by CFCR's Executive Director Sarah A. Chase along
with Council Advisory Board Member Jack Jacobson, a
cannabis, energy and education practices advisor for
Thompson Coburn LLC, the panel will consist of FDA Scientists
Cassandra Taylor, Ph.D. and Dan Mellon, Ph.D., Dr. Vicki
Seyfert-Margolis, Chairwoman of CFCR's Science and
Regulatory Affairs Committee, molecular geneticist Dr. Reggie
Gaudino, also a member of that same committee and Mara
Gordon, co-founder of Aunt Zelda's, a pioneering, data-driven developer of cannabis-based plant
medicines.

Offering in-depth subject knowledge of the still under-studied and often misunderstood
endocannabinoid system, each of the three panelists will bring a uniquely varied area of
expertise from molecular genetics, immunology, and patient advocacy. This webinar will help to
unpack some of the complexity surrounding the long-term considerations for researchers and
regulators.

The event will take place over WebEx Video Conference, Thursday, February 24, from 1:00 until



Cassandra Taylor, Ph.D.,
joined FDA in December 2014
as a primary BRT reviewer and
has evaluated over 100
botanical drug submissions
from all CDER's clinical
divisions, with a focus on
reviewing cannabis
submissions.

2:00 p.m. EST.

With roughly 125 monthly listeners already since debuting last September, CFGR has explored various facets of the cannabis industry in their monthly webinar series – including the regulatory impacts of a groundbreaking CBD study, as well as Federal and state cannabis policies.

Ongoing webinars will take place on the second or third Thursday of every month (depending on holidays), each to spotlight a wide range of industry professionals and focus on preparing to regulate and regulating today's legal cannabis molecules and products. These virtual sessions can be viewed on WebEx. Each webinar will consist of a 40-minute interactive conversation followed by approximately 20-minutes of moderated audience Q&A for a total run-time of approximately one hour.

Speaker Profiles for the February 24 Webinar:

Cassandra Taylor, Ph.D., Chemist, US Food and Drug Administration

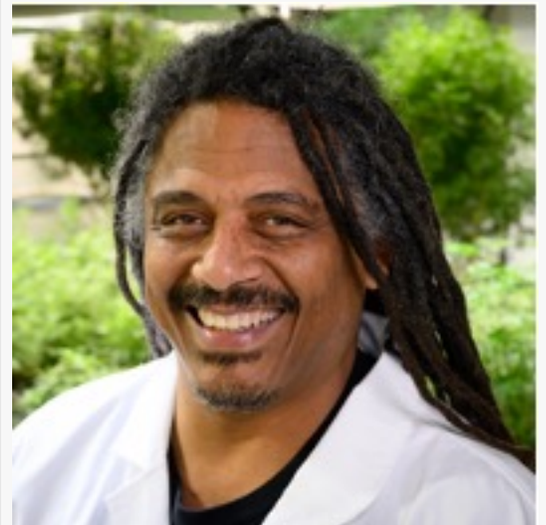
Cassandra Taylor, Ph.D. is a Chemist at U.S. Food and Drug Administration within the Center for Drug Evaluation and Research (CDER) and is a member of the Botanical Review Team (BRT) which resides within the Office of Pharmaceutical Quality (OPQ). BRT collectively serves as an expert resource for CDER on all botanical issues. Dr. Taylor received her B.S. in Chemistry with a minor in Forensics from St. Francis University in Loretto, PA (2005), and her Ph.D. in Analytical Chemistry from the University of Maryland in College Park, MD (2014).

Dan Mellon, Ph.D., Director of Pharmacology/Toxicology, U.S. Food and Drug Administration

Dr. Mellon holds the lengthy title of Deputy Director for the Division of Pharmacology/Toxicology for Neuroscience and Pharmacology Toxicology Supervisor supporting the Division of Anesthesiology, Addiction



Dan Mellon, Ph.D., began his career at FDA as a Pharmacology Toxicology Reviewer in the Division of Anesthetics, Critical Care, and Addiction Drug Products in 2001. He is currently the Deputy Director for the Division of Pharmacology Toxicology for Neuroscience.



Dr. Gaudino spent 22 years working as a patent scientist or Patent Agent with a number of national law firms, and was the Lead Scientist on the Litigation support team in an early, highly publicized, plant biotechnology patent litigation.

Medicine, and Pain Medicine Office of New Drug Center for Drug Evaluation and Research.

Vicki Seyfert-Margolis, Chairwoman, CFCR Science & Regulatory Affairs Committee

Vicki Seyfert-Margolis, Ph.D. founded My Own Med in January 2013, based on over two years of work on a database, web and mobile application platform technology for family-based co-management of health. Previously, Vicki was the Senior Advisor for Science Innovation and Policy in the Office of the Commissioner of the US Food and Drug Administration. While at the FDA, she oversaw the development and execution of an agency wide strategic plan for regulatory science.

Dr. Reggie Gaudino, Science & Regulatory Affairs Committee

Dr. Reggie Gaudino is a molecular geneticist focused on biochemical networks in plant phytochemistry. After finishing his PhD in molecular genetics at the University of Buffalo in New York, investigating the biochemistry of transcription initiation and termination in E.coli, he started his postdoc at Washington University in St. Louis investigating control of gene expression, including showing specific Cytokinin responsive RNA Polymerase I transcription initiation in isolated A.thaliana nuclei.

During a particularly austere funding period in the mid-nineties Dr. Gaudino, pursued a legal path to incorporate biotechnology with intellectual property - having been involved in many incarnations of recombinant DNA technologies, DNA sequencing, RNA analysis, in vitro promoter analysis and codon optimization for reporter gene systems in organisms ranging from bacteria to plants. Dr. Gaudino spent the next 22 years working as a patent scientist or Patent Agent with a number of national law firms, and was the Lead Scientist on the Litigation support team in an early, highly publicized, plant biotechnology patent litigation.

Mara Gordon, Co-Founder, Aunt Zelda's

Mara Gordon is a cannabis advocate, entrepreneur, and researcher. She has harnessed her background as a process engineer to create therapeutic dosing regimens for thousands of patients around the world, drastically improving their health, quality of life and longevity. Mara openly shares her knowledge about the therapeutic benefits of the cannabis plant – whether consulting with medical teams, through TEDx Talks or calling out hyperbole in the industry about which she cares so deeply.

To register for this CFCR's webinar series, please visit: <https://www.uscfc.org/events>. For additional information, please visit [uscfc.org](https://www.uscfc.org).

ABOUT CFCR

The Council for Federal Cannabis Regulation (CFCR) is 501(c)(3) non-profit based in Washington, DC. The mission of CFCR is to assist the government, and specifically federal regulatory agencies, to rethink, develop, and implement evidence-based cannabis regulations. Our overarching goal is the de-stigmatization, normalization, and legitimization of cannabis on behalf of consumers, the professions, organizations, and businesses who support and serve them. We do this by serving as a conduit for informed scientific research, inclusive education, and by mainstreaming the best practices that enable the industry to maximize its potential. For additional information please visit www.uscfcf.org and follow us on social media @USCFCR.

MEDIA CONTACTS

Steve Winter, Brotman | Winter | Fried, swinter@aboutbwf.com, 202-468-8100

Kim Prince, Proven Media, kim@provenmediaservices.com, 480-221-7995

Heather Mahoney, Brotman | Winter | Fried, hmahoney@aboutbwf.com, 240-271-5762

Sarah Chase, Executive Director

Council for Federal Cannabis Regulation

+1 202-518-5359

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