



Bioavailability and Bioequivalence of Topical Formulations

Guest Editor:

Prof. Dr. Isadore Kanfer

Leslie Dan College of Pharmacy,
University of Toronto, Toronto,
ON M5S 3M2, Canada

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Message from the Guest Editor

Dear Colleagues,

The market approval of a generic product requires the demonstration of bioequivalence (BE) against an innovator reference product in order to ensure safety and efficacy. Unlike extravascular dosage forms where well-established methods and regulatory guidelines are available for BE assessment, procedures or guidelines for the BE of topical dosage forms intended for local action, apart from the US FDA's vasoconstrictor assay (VCA), are conspicuously absent. Most regulatory authorities usually require lengthy and expensive comparative clinical endpoint studies in patients. More recently, considerable efforts have been directed towards the development and validation of surrogate models to demonstrate the BE of topical products for local action and facilitate the faster entry of generic products into the market.

This Special Issue on the bioavailability (BA) and bioequivalence (BE) of topical formulations intended for local action will focus on research and review papers discussing applications of validated methods used to assess topical dosage forms not intended for absorption, and will include both in vivo and in vitro methods.





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Editor-in-Chief

Prof. Dr. Patrick J. Sinko

Department of Pharmaceutics,
Ernest Mario School of
Pharmacy, Rutgers University,
Piscataway, NJ 08854, USA

Message from the Editor-in-Chief

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Pharmaceutics Editorial Office
MDPI, St. Alban-Anlage 66
4052 Basel, Switzerland

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