

## Teleconference Date:

- September 19, 2008
- 12:00PM EST

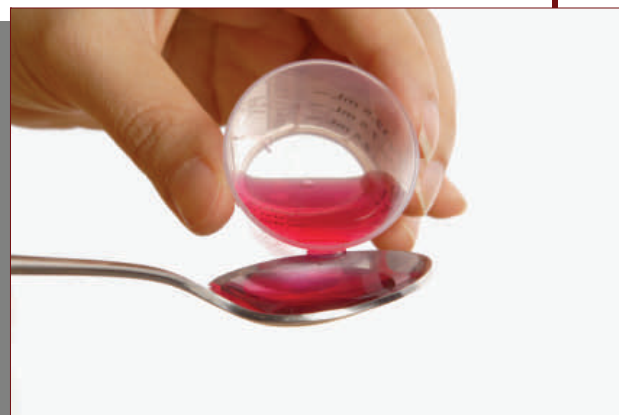
Audio recording  
available

# Over-the-Counter Drugs

An Emord & Associates, P.C. Teleconference

FDA's Center for Drug Evaluation and Research (CDER) regulates over 80 classes of Over-the-Counter (OTC) drug products. In this informative teleconference, the attorneys at Emord & Associates explain:

1. The significance of prior-marketed drugs and OTC drug monographs
2. Specific labeling requirements applicable to OTC drug products
3. Permitted claims concerning an OTC product's efficacy
4. FDA's New Drug Approval process as it pertains to OTC drugs
5. The new Adverse Event Reporting requirements for OTC drug products



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*To participate, call Emord & Associates, P.C. (202) 466-6937 or email us at [jemord@emord.com](mailto:jemord@emord.com). The fee for participation (payable by credit card) is \$225. A pass code and phone number for the September 19th program will be supplied to you upon paying the fee.*

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