

VHA Pharmacy Benefits Management Strategic Healthcare Group
and The VA Medical Advisory Panel
with the acknowledgements from the
Associate Chief Consultant Pharmacy Benefits Management
Adverse Drug Event Reporting Programs
and VHA Patient Care Services

presents:

VIEWING AND ACCREDITATION INFORMATION

View live on your desktop: Content Distribution Network (CDN)
<http://vawww.vakncdn.lrn.va.gov>

Broadcast Information and Materials:
Log on to: <http://vawww.sites.lrn.va.gov/vacatalog>
Keyword: Adverse Drug Event

**Contact the Education Coordinator at your facility
for required Scantron Forms**

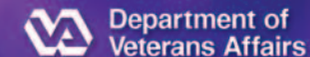
THE ABCs OF ADEs:

Adverse Drug Events . . . *Ask, Document, and Enter*

**Wednesday, September 6
1:00-2:00 pm ET**



**PHARMACY BENEFITS MANAGEMENT
MEDICAL ADVISORY PANEL**



FACULTY

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VAKN SATELLITE BROADCAST INFORMATION:

Wednesday, September 6, 1:00-2:00 pm ET

Satellite Broadcast (on Channel 1)
Veterans Affairs Knowledge Network (VAKN)

The September 6 broadcast will be followed at 2:00 pm ET with a
VANTS conference call which will provide an opportunity to discuss
issues covered in the broadcast with the faculty.

Dial-in number 1-800-767-1750, Access Code 14945#

Program Re-Broadcast Dates:

September 8	6:00 pm (ET)	CH 1
September 11	9:00 pm (ET)	CH 1
September 12	4:00 pm (ET)	CH 1
September 14	8:00 am (ET)	CH 1
September 20	3:00 am (ET)	CH 1
September 26	3:00 pm (ET)	CH 1
September 28	10:00 am (ET)	CH 1

View live on your desktop: Content Distribution Network (CDN)
<http://vawww.vakncdn.lrn.va.gov>

This program is supported by an unrestricted educational grant from Abbott Laboratories.



PURPOSE:

To give VHA healthcare providers and allied health personnel up-to-date and a broader understanding of the importance of asking, documenting and entering adverse drug events into the VHA adverse drug reporting systems.

TARGET AUDIENCE:

Primary Care Physicians, Pharmacists, Nurses, Dietitians, Interventional Radiologists, and other clinical staff

LEARNING OBJECTIVES:

1. Define the terms ADE and ADR
2. Be able to document a meaningful ADR in CPRS package
3. Identify at least two reporting systems for the reporting of ADEs
4. Discuss the importance of documenting ADEs associated with off-label medication use
5. Describe utilization by VHA of ADE data reporting at the local, VISN and national level

BENEFITS:

- Nationally recognized faculty presenters
- Interactive Q&A session to enrich educational experience
- Current information relevant to VHA patient populations
- Live training with minimal time investment
- Quality educational experience offering continuing education credits without travel or extended time away from regular duties

BROADCAST SLIDE PRESENTATIONS

The broadcast slide presentations can be downloaded and printed from <http://vawww.pbm.va.gov/pbm/dislearning.htm/>
Or
<http://www.pbm.va.gov/pbm/dislearning.htm/>

PROGRAM INFORMATION AND ACCREDITATION REQUIREMENTS

Complete information on this satellite broadcast, including handouts, is available at <http://vawww.sites.lrn.va.gov/vacatalog>
Key word: Adverse Drug Event

Completed paperwork must be received at the EES, Birmingham Employee Education Resource Center no later than October 30, 2006.

Contact the Education Coordinator at your facility for the required Scantron Evaluation Forms which includes:

- Satellite Participant Form
- Faculty Evaluation Form

The completed forms must be mailed to:
Birmingham Employee Education Resource Center
ATTN: SDU
950 North 22nd Street, Suite 500
Birmingham, AL 35203

Further information may be obtained from:

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DISCLOSURE(S)

The VA Employee Education System (EES) must insure balance, independence, objectivity, and scientific rigor in all its individually sponsored or jointly EES sponsored educational activities. All prospective faculty & planning committee members participating in an EES activity must disclose any relevant financial interest or other relationship with: (a) the manufacturer(s) of any commercial product(s) and/or provider(s) of commercial services discussed in an educational presentation, and (b) any commercial supporters of the activity. Relevant financial interest or other relationship includes but is not limited to such things as personal receipt of grants or research support, employee or consultant status, stock holder, member of speaker's bureau, within the prior 12 months. EES is responsible for collecting such information from prospective planners and faculty, evaluating the disclosed information to determine if a conflict of interest is present and, if a conflict of interest is present, to resolve such conflict. Information regarding such disclosures and the resolution of the conflicts for planners and faculty shall be provided to activity participants. *When an unlabeled use of a commercial product or an investigational use not yet approved by the FDA for any purpose is discussed during an educational activity, EES shall require the speaker to disclose that the product is not labeled for the use under discussion or that the product is still investigational.*

Each faculty and planning committee member (author, facilitator, moderator) reported having no relevant* financial relationships with any commercial interest. This activity includes no discussion of uses of FDA regulated drugs or medical devices which are experimental or off-label.

**The ACCME defines "relevant financial relationships" as financial relationships in any amount occurring within the past 12 months that creates a conflict of interest.*

THE REHABILITATION ACT OF 1973, AS AMENDED:

The Employee Education System wishes to ensure no individual with a disability is excluded, denied services, segregated or otherwise treated differently from other individuals participating in its educational activities, because of the absence of auxiliary aids and services. If you require any special arrangements to participate in this educational activity, please contact:

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