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**Reframing pharmacy's leadership,  
relationship, and scholarship**

**Drug therapy during cardiac arrest**

**Methods used by pharmacy departments  
to identify drug interactions**

**Annual index**

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**2458 News**

After developing bladder tumor, pharmacist calls for national cancer registry  
States surpass OBRA '90 patient-counseling requirements  
International guidelines for pharmacy practice approved at 53rd FIP congress in Tokyo  
Patient care gaining importance in Japanese pharmacy practice (*Audrey St. Jean Wittenburg*)  
Alcohol rivals myocardial infarction as cause of hospitalization in elderly people  
Jump in cigarette taxes would lower demand, NCI panel concludes  
More than 1 million report using steroids  
Medicare project increases influenza vaccination rate, predicts lower hospital costs  
Pharmacy school applications, enrollments grew in 1992; entry-level degrees dropped slightly  
MedWatch update: Pharmacists submit over half of product problem reports  
HMO spending for outpatient pharmaceuticals held steady in 1992  
Contract pharmacy services grew in 1992

**2484 ASHP Affiliates**

Vermont Society helps promote adverse drug reaction reporting  
Hands-on patient care explored at Nevada Society meeting  
Montana Society annual meeting features psychopharmacy, management, clinical skills  
Raehl, Bond discuss pharmaceutical care at Idaho Society meeting  
New Mexico Society Balloon Fiesta meeting draws attendees from 18 states

**2497 ASHP Foundation Reports**

Atlanta Midyear activities kick off Foundation's 25th anniversary  
Literature awards to be presented in Atlanta  
Foundation awards Demonstration Project Grants, New Pharmacy Practice Researcher Grant  
Pharmacists complete dialysis traineeship  
Anticoagulation traineeship program expands in 1993  
Funding available for state conferences on pharmaceutical care

**2506 Questions and Answers**

Outpatient drug discounts for hospitals with a disproportionate share of indigent patients

**2511 Student & Resident Forum**

ASHP summer internship

**2515 Frontline Pharmacist**

Treatment for ailing staff-management communication

**2518 Letters**

Precipitation of paclitaxel during infusion by pump (*Robert W. Pfeifer, Karen N. Hale; Suzanne Emerson Cronquist, Michelle Daniels*)  
Incompatibility of labetalol hydrochloride and furosemide (*John Zeisler, Cheri Alagna*)  
Sports pharmacy clerkship (*Sally J. Sato, Donna L. Weingart, Kenneth L. Brier*)  
Topical polyhexamethylene biguanide (pool cleaner) for treatment of acanthamoeba keratitis (*Edmond Yee, Richard Fiscella, Tien Kiat Winarko*)  
Advice on submitting papers (*Gregory Gousse; Thomas G. Burnakis*)  
Difficulties in assessing appropriate ondansetron use (*Lewis R. Schwarz; Gerald M. Higa; Mark D. Peters II, Jeffrey A. Reitz, Lois M. Jessen, Kimberly S. Long*)  
Perceived usefulness of supplement on rising pharmaceutical costs (*Michael J. Melby; Fred D. Chatelain; John P. Santell*)

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Joseph A. Oddis  
Executive Vice President

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- Editorial**
- 2529** Compassionate dispensing
- Special Feature**
- 2533** Reframing pharmacy's leadership, relationship, and scholarship  
*Calvin H. Knowlton*
- Reports**
- 2538** Drug therapy during cardiac arrest in two hospitals  
*Barry E. Bleske, Wendy K. Wheatley, Kevin A. Townsend, Marc L. Guzzardo, John E. Billi, and Michael J. Shea*
- 2546** Methods used by pharmacy departments to identify drug interactions  
*Sona Elanjian, Mary Lea Gora, and Laura Roeder Symes*
- 2550** Interaction of sucralfate with antibiotics used for selective decontamination of the gastrointestinal tract  
*Barbara Feron, Colin G. Adair, Sean P. Gorman, and Barry McClurg*
- Notes**
- 2554** Procedure for evaluating nonformulary drug orders  
*Gene T. Jay, Robert A. Quercia, Gregory Gousse, Moses S. S. Chow, and Richard Quintiliani*
- 2556** Comparison of concurrent and retrospective methods of detecting adverse drug reactions  
*Janet J. Madsen*
- 2558** Medication administration records for drugs given during surgery  
*Patti R. Hawkins*
- 2559** Stability of cimetidine hydrochloride and of clindamycin phosphate in water for injection stored in glass vials at two temperatures  
*Milap C. Nahata, Richard S. Morosco, and Thomas F. Hipple*
- 2561** Stability of nitroglycerin as nitroglycerin concentrate for injection stored in plastic syringes  
*Paul S. Driver, Eric J. Jarvi, and Peggy L. Gratzler*
- ASHP Report**
- 2564** Acknowledgment to reviewers—November 1, 1992, to October 31, 1993
- AJHP Continuing Education**
- 2566** ASHP journal continuing-education instructions and enrollment form
- 2569** Career Opportunities
- 2584** CP Digest
- 2604** Current Literature  
Journal References  
Book Reviews: *Bad Prescription for the First Amendment: FDA Censorship of Drug Advertising and Promotion* (Kaplar), William L. Fong  
*Geriatric Dosage Handbook, 1993* (Semia, Beizer, Higbee), Nathan Rawls  
*Manual for Pharmacy Technicians* (American Society of Hospital Pharmacists), Darla Gallo  
Books Received
- 2615** Advertising Index
- At Large**
- 2616** Context of a transition
- 2617** Annual Index

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# LETTERS

## **Precipitation of paclitaxel during infusion by pump**

2518 Robert W. Pfeifer,  
Karen N. Hale;  
Suzanne Emerson Cronquist,  
Michelle Daniels

## **Incompatibility of labetalol hydrochloride and furosemide**

2521 John Zeisler,  
Cheri Alagna

## **Sports pharmacy clerkship**

2522 Sally J. Sato,  
Donna L. Weingart,  
Kenneth L. Brier

## **Topical polyhexamethylene biguanide (pool cleaner) for treatment of acanthamoeba keratitis**

2522 Edmond Yee,  
Richard Fiscella,  
Tien Kiat Winarko

## **Advice on submitting papers**

2523 Gregory Gousse;  
Thomas G. Burnakis

## **Difficulties in assessing appropriate ondansetron use**

2523 Lewis R. Schwarz;  
Gerald M. Higa;  
Mark D. Peters II,  
Jeffrey A. Reitz,  
Lois M. Jessen,  
Kimberly S. Long

## **Perceived usefulness of supplement on rising pharmaceutical costs**

2526 Michael J. Melby;  
Fred D. Chatelain;  
John P. Santell

## Precipitation of paclitaxel during infusion by pump

**P**aclitaxel (Taxol, Bristol-Myers Squibb Oncology) is administered as a 24-hour continuous i.v. infusion. The use of an infusion-control device has been recommended. When diluted to a final concentration of 0.3 to 1.2 mg/mL, paclitaxel has been reported to be visually and chemically stable for up to 27 hours at room temperature and under ambient light.<sup>1</sup>

Investigational use of paclitaxel at our institution began in September 1991. Doses of paclitaxel were prepared in 1000 mL of 5% dextrose injection in glass bottles and infused through vented, polyethylene-lined solution administration sets (2C7552s, Baxter Healthcare Corporation) and 0.22- $\mu$ m filters (Millex-GV, Millipore). Volumetric infusion pumps (Flo-Gard models 6200 [single chamber] and 6300 [dual chamber], Baxter) were used; these pumps infuse fluid by linear peristalsis—a series of rollers move against a small

segment of polyvinyl chloride (PVC) tubing in a peristaltic motion that resembles pushing the fluid along the tubing path.

In August 1992, we received sporadic reports of fluffy, white precipitate in paclitaxel-exposed solution administration sets. The precipitate appeared throughout the section of tubing distal to the pump chamber but not in the glass bottle or the tubing leading to the chamber. The precipitation apparently began in the section of PVC tubing that came into direct contact with the peristaltic rollers of the pump. When precipitation was observed, the solution administration set and filter were changed and the balance of the infusion was administered. Partial doses were not remade, because precipitation was not found in the glass bottles. In most cases, precipitation was observed near the end of the 24-hour infusion period. The concentration of these solutions ranged from 0.2

*Continued on page 2521*

The Letters column is a forum for rapid exchange of ideas among readers of AJHP. Liberal criteria are applied in the review of submissions to encourage contributions to this column.

The Letters column includes the following types of contributions: (1) comments, addenda, and minor updates on previously published work, (2) alerts on potential problems in practice, (3) observations or comments on trends in drug use, (4) opinions on public health issues of interest to pharmacists in institutional settings, (5) comments on ASHP activities, and (6) human interest items about life as a pharmacist.

Short papers on practice innovations and other original work are included in the Notes section rather than in Letters.

Letters need not be submitted with AJHP's manuscript checklist. The following conditions, however, must be adhered to: (1) the body of the letter must be no longer than two typewritten pages, (2) the use of references and tables should be minimized, (3) the authors' names, affiliations, and mailing addresses must be typed at the end of the letter in the format used by AJHP, and (4) the entire letter (including references, tables, and authors' names) must be typed double-spaced on plain white paper (not letterhead). Following acceptance of a letter, the authors are required to sign an exclusive publication statement and a copyright transferal form. All letters are subject to revision by the editors. Authors do not receive proofs of edited letters.

Continued from page 2518

to 0.3 mg/mL.

The use of paclitaxel vials and solution administration sets from other lots did not prevent precipitation. We notified the National Cancer Institute (NCI) and sent samples of tubing with precipitate for analysis. At that time, NCI had received no similar report of precipitation. In November 1992, following an amendment to NCI treatment referral center protocol 9103, we decreased the volume of a 24-hour dose of paclitaxel to 500 mL; we also began dividing the total daily dose into two 12-hour infusions (250 mL each) to decrease the length of time during which the drug could precipitate. Over the next six weeks, 18 incidents of precipitation (in approximately 150 doses) occurred in solution administration sets used to infuse paclitaxel 0.38–0.58 mg/mL.

Similar problems have been reported with teniposide (Vumon, Bristol-Myers Squibb Oncology),<sup>2</sup> which also contains polyoxyethylated castor oil and alcohol. According to the manufacturer, supersaturated solutions of teniposide rapidly precipitate after formation of seed crystals, a process that may be accelerated by agitation.<sup>3</sup> Agitation sufficient to induce precipitation could be produced by a mechanical pumping action. We suggest that a similar event occurs with our method of paclitaxel administration.

Further studies have been performed. Various types of agitation and manipulation appeared to induce precipitation earlier than was seen in the static samples. Also, higher paclitaxel concentrations began to precipitate sooner than lower concentrations. However,

precipitation was not seen when another group repeated the procedure with the solutions and equipment that we routinely use to infuse paclitaxel (Trissel LA, personal communication, 1993 Feb 12).

We have not completely resolved our problem with precipitation, despite interventions to minimize agitation of paclitaxel solutions. Another, more costly option we are considering is to use cassette-type, positive-displacement, volumetric pump systems (e.g., Flo-Gard model 8000, Baxter) for administering paclitaxel. We would appreciate receiving suggestions or hearing about others' experiences with paclitaxel infusion solutions.

1. Bristol-Myers Squibb Company. Taxol package insert. Princeton, NJ: 1992 Dec.
2. Strong DK, Morris LA. Precipitation of teniposide during infusion. *Am J Hosp Pharm.* 1990; 47:512,518. Letter.
3. Bogardus JB, Kaplan MA, Carpenter JP. Precipitation of teniposide during infusion [response]. *Am J Hosp Pharm.* 1990; 47:518-9. Letter.

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**B**efore we received a copy of the above letter, Bristol-Myers Squibb was not aware of clinical or laboratory conditions in which paclitaxel precipitated during infusion by peristaltic pump or with a specific type of infusion device.

The paclitaxel package insert cautions against the use of plasticized polyvinyl

chloride (PVC) equipment or devices. The infusion set described by Mr. Pfeifer and Ms. Hale contains a small amount of PVC tubing in a segment where the PVC tubing comes in direct contact with the peristaltic rollers of the pump; however, a laboratory simulation at another institution did not produce paclitaxel precipitation.

Since paclitaxel became commercially available in December 1992, the company has received fewer than a dozen reports of precipitation. Information on the drug concentration, infusion set, and infusion pump used when the drug precipitated has not always been provided. In most cases when this information was provided, precipitation occurred in a PVC-containing segment of tubing.

For the mechanism of

the reported precipitation to be identified, precipitation would have to be reproducible under controlled conditions. Perhaps the sample size used by Mr. Trissel during simulation was too small to represent the small number of occurrences in clinical practice.

Finally, we would like to emphasize the selection of infusion sets appropriate for the infusion pump chosen and the avoidance of PVC equipment, as directed in the paclitaxel package insert.

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## Incompatibility of labetalol hydrochloride and furosemide

**W**e report the apparent incompatibility of labetalol hydrochloride and furosemide injections.

A 60-year-old woman was receiving labetalol hydrochloride (Normodyne, Schering Laboratories, Kenilworth, NJ, lot 2 DDF 102) 1.6 mg/mL in 5% dextrose injection by continuous i.v. infusion at 30 mg/hr for treatment of systolic hypertension. Furosemide 40 mg (10 mg/mL, Elkins-Sinn, Cherry Hill, NJ, lot 033021) was to be given by i.v. push. Less than 0.5 mL of furosemide had been injected into the port of the labetalol i.v. line when white precipitation was noted. No further furosemide was injected, and the i.v. line was flushed with 0.9% sodium chloride injection.

The patient suffered no detectable adverse drug reaction from this event. The incompatibility was confirmed in the pharmacy department's laminar-airflow hood by using injectable solutions from the same lots and in the same concentrations as the patient received.

Schering Laboratories could provide no previous reports of this incompatibility (Newandee K, Drug Information Manager, personal communication, 1993 Jul 16). This incompatibility is not described in Trissel's *Handbook on Injectable Drugs*.<sup>1</sup> We suspect the incompatibility is due to the difference in solution pH between labetalol hydrochloride (pH 3–4) and furosemide (pH 8–9.3).<sup>2</sup>