



Guidelines for Authors

SUPPLEMENTS

Information for Authors.

Pharmaceutical Technology publishes topical supplements on various topics throughout the year. Topics scheduled for 2009 include:

- Excipient performance for solid dosage forms (April)
- Technology and regulatory update for sterile manufacturing (May)
- Bioprocessing (July)
- Outsourcing resources (August)
- Contract manufacturing: API synthesis and formulation (September)
- Quality by Design (October)
- Injectable drug delivery (November)

Manuscripts for consideration of publication should be submitted to Editor-in-Chief Michelle Hoffman or Managing Editor Angie Drakulich at the address on the next page, or via email at mhoffman@advanstar.com or adrakulich@advanstar.com, respectively.

General guidelines. Before submitting a paper, authors are urged to review manuscripts for clarity of expression, grammar, and typographical accuracy. Acronyms and abbreviations used in manuscripts should be defined on first reference and within tables and figures. The author is responsible for all statements in his or her work. All accepted manuscripts are subject to editing. Manuscripts are reviewed with the understanding that they have not been published previously, are not ghostwritten, and are not under consideration for publication elsewhere, including on the Internet. The author and any coauthors will be required to sign a license agreement before a manuscript is accepted for publication.

Preparing the electronic file. The text should be formatted in Microsoft Word and labeled with the corresponding author's last name. Do not use any special text formatting codes or control characters. Strip out any special-character codes and font-change codes. Please include tables and figure captions in the electronic file at the end of the article (see below for submission of images). Beginning with the first page of the text, each page of the hard copy should be numbered consecutively. Be sure to include a title page (with authors' name, titles, company affiliation, address, phone, fax, and e-mail) and a synopsis or abstract. Paper titles should be short and specific and should accurately reflect the content of the article.

Illustrations. Photographs, line drawings, and other illustrations may be submitted in color or black and white and must be referred to in the text in consecutive order: Figure 1, Figure 2, and so forth. Be sure to include descriptive captions for all figures and images. All illustrations that do not increase the reader's understanding of the text should be omitted. In addition to the substantive figures and images provided, a four-color photograph may be used on the opening page of the article. Color photos that an author wishes to submit for consideration should be submitted with the article and should be in the form of a TIFF (Mac), TIF (PC), JPEG, or EPS/EPSF (Mac or PC) file. Images must have a minimum resolution of 300 dpi.

Figures and tables. Line drawings, graphs, and tables or charts must be identified with the figure number and principal author's name. Figure legends should be collected on a separate page, each identified by its proper number. All symbols, acronyms, and abbreviations used in figures and tables should be explained. In the text, tables should be referred to by Roman numerals in consecutive order: Table I, Table II, and so forth.

Credits. All images, figures, graphs, and tables must be accompanied by the source (creator) name so that the editors can provide appropriate credit. If the author(s) or the author's company created and/or own the rights to the images, figures, tables or graphics, please note that this is the case. If a third party created and/or owns the rights to such illustrations, the author is responsible for acquiring rights permissions and providing related documentation. Please note that *Pharmaceutical Technology* will only publish original tables, graphics, and figures.

References. Literature citations in the text should be numbered consecutively and indicated by Arabic numerals in parentheses after appropriate sentences and/or paragraphs. Please do not use endnotes or footnotes. References should be grouped at the end of the manuscript and arranged in order of their appearance in the text — not alphabetically. **Please adhere to our reference style below:**

Chapter in a book:

-E.F. Fiese and T.A. Hagen, "Preformulation," in *Theory and Practice of Industrial Pharmacy*, L. Lachman, H.A. Lieberman, and J.L. Kanig, Eds. (Lea & Febiger, Philadelphia, PA, 3rd ed., 1986), pp. 171–194.

Article in a journal:

-G.M. Golden, J.E. McKie, and R.O. Potts, "Role of Stratum Corneum Lipid Fluidity in Transdermal Drug Flux," *J. Pharm. Sci.* 76 (1), 25–28 (1987).

Published conference proceedings:

-J.B. Dressman, "Predicting Release Kinetics from Enteric-Coated Dosage Forms," in *Proceedings of Pharm Tech Conference* (Aster Publishing Corporation, Eugene, OR, 1988), pp. 239–250.

Oral presentations:

-B.L. Hawkins, A. Baxter, and G.E. Masters, "NMR Preview: A Look to the 1990s," presented at the 25th Experimental NMR Conference, Wilmington, DE, Feb. 1984.

-J. Woodcock, MD, Statement, House Committee on Energy and Commerce Hearing, Apr. 29, 2008, available at http://energycommerce.house.gov/cmte_mtgs/110-oi-hrg.042908, accessed May 8, 2008.

Government publications:

-*Code of Federal Regulations*, Title 21, Food and Drugs (Government Printing Office, Washington, DC), Part 121, pp. 270–290.

-"Potable Water, Control of Communicable Diseases," in *Code of Federal Regulations*, Title 21, Food and Drugs (Government Printing Office, Washington, DC), Part 1240.3 (m), pp. X–X. (Note: same style as chapter in a book, but without an author).

-FDA, "Human and Veterinary Drugs, Good Manufacturing Practices and Proposed Exemptions for Certain OTC Products," *Fed. Regist.* 43 (190), 45013–45089 (Sept. 29, 1978).

-"Minimizing the Risk of Transmitting Animal Spongiform Encephalopathy Agents Via Medicinal Products," in *European Pharmacopoeia* (European Directorate for the Quality of Medicines, Strasbourg, France, YEAR), General Text 5.2.8, pp–pp.

-ICH and FDA, Q1B *Photostability Testing of New Drug Substances and Products* (Geneva, Switzerland, and Rockville, MD, Nov. 1996), fda.gov/cder/guidance/1318.htm, accessed Sept. 15, 2005.

-*Photostability Testing of New Drug Substances and Products* (Geneva, Switzerland, and Rockville, MD, Nov. 1996), available at fda.gov/cder/

guidance/1318.htm, accessed Sept. 15, 2005. (Note: same as book style, add web url only if entire document is online, update access date)

-FDA, *Guidance Title* (Rockville, MD, Nov. 2004).

[Note: Add a URL if entire document is available online.]

USP publications

-*USP 27–NF 22* (US Pharmacopeial Convention, Rockville, MD, 2003), pp. X–X.

[Note: If the above is a reference for an entire General Chapter or other titled section in *USP*, then it follows the style of a book chapter]

- USP Proposed General Test <429>, “Light Diffraction Measurement of Particle Size,” *Pharmacopeial Forum* 28 (4), 1293–1298.

Patents

-Author or Company name, “Transmitter Switch for Wireless ID,” US patent 125356, Dec. 2003.

Keywords, weblinks, and online tools. Please provide a list of keywords relevant to the article, as well as websites or additional online references and materials that might be of interest to readers. *Pharmaceutical Technology* is specifically interested in publishing video demonstrations, audio tutorials, and downloadable tools that are related to the article. Such interactive items would published on its website, PharmTech.com, with the full article at the same time it appears in the print magazine. Please include mention of such items, if applicable and available, when you submit your paper.

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Frequently Asked Questions.

What kinds of articles do you look for?

Manuscripts should be sufficiently novel to be of interest to an experienced audience. Articles should be data driven and provide sufficient detail. Topics should be timely and useful.

What are your deadlines for submission?

Manuscripts should be submitted 2–4 months before the supplement’s scheduled publication month.

What does it take to get accepted?

Manuscripts must be interesting, timely, and original. Manuscripts should be 2000–4000 words.

How many readers do you have?

Pharmaceutical Technology readers (38,600 BPA-qualified) are chemists, engineers, and scientists involved in pharmaceutical manufacturing, including formulation, development, analytical testing, quality control/quality assurance, and synthesis of active ingredients. Readership also includes individuals responsible for purchasing or making purchasing decisions for machinery and equipment, as well as outsourced services. Please refer to our media kit (PharmTech.com) for a breakdown of readers’ job functions.

What else does *Pharmaceutical Technology* publish?

Pharmaceutical Technology covers all aspects of applied research and development, scale-up, and manufacturing technologies for the pharmaceutical industry. Topics include analytical testing, ingredients, formulation, drug delivery systems, information technology, manufacturing, formulation and API synthesis, packaging, outsourcing, and regulatory affairs. See our separate “Guidelines for Authors” and “Guidelines for Authors: Short Papers (Technical Notes & Case Studies)” to contribute.

• Buyers Guide and Resource Directory. The comprehensive annual guide to suppliers of products, services, and resources of all kinds, from the most complete and accurate sourcing database in the industry.

• Corporate Capabilities. The annual indispensable compendium of detailed information on suppliers, products, and services, mailed to *Pharmaceutical Technology*’s entire circulation.

What about *Pharmaceutical Technology*’s online offerings?

- *ePT*—the Electronic Newsletter of *Pharmaceutical Technology*. The early warning system for drug manufacturing, an easy-to-use weekly summary of drug-making news.
- *Sourcing and Management*. A monthly electronic newsletter for pharmaceutical technology management, with original articles on outsourcing, and case histories.
- *Equipment & Processing Report*. This monthly electronic newsletter covers manufacturing trends, new products, and equipment updates.
- PharmTech.com. The magazine’s home on the Internet, with searchable news, features, products, and columns.
- Blog.PharmTech.com. Our daily blog covers news, live events, and industry trends.
- Online Buyers Guide. The Buyers Guide in real time—searchable, comprehensive, and constantly updated.

Where can I find your editorial calendar?

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