

Howard C. Ansel
Nicholas G. Popovich
Loyd V. Allen, Jr.

ションドコニコドラス





Executive Editor: Donna M. Balado Developmental Editor: Frances M. Klass Production Coordinator: Peter J. Carley Project Editor: Jessica Howie Martin

Copyright © 1995 Williams & Wilkins 200 Chester Field Parkway Malvern, PA 19355 USA



All rights reserved. This book is protected by copyright. No part of this book may be reproduced in any form or by any means, including photocopying, or utilized by any information storage and retrieval system without written permission from the copyright owner.

Accurate indications, adverse reactions, and dosage schedules for drugs are provided in this book, but it is possible they may change. The reader is urged to review the package information data of the manufacturers of the medications mentioned.

Printed in the United States of America

Library of Congress Cataloging in Publication Data

94 95 96 97 98 1 2 3 4 5 6 7 8 9 10

Ansel, Howard C., 1933-

Pharmaceutical dosage forms and drug delivery systems / Howard C. Ansel, Nicholas G. Popovich, Lloyd V. Allen, Jr.—6th ed.

p. cm.

Includes bibliographical references and index.

ISBN 0-683-00193-0

1. Drugs—Dosage forms. 2. Drug delivery systems.

I. Popovich, Nicholas G. II. Allen, Loyd V. III. Title.

[DNLM: 1. Dosage Forms. 2. Drug Delivery Systems. QV 785 A618i

1995]

RS200.A57 1995

615'.1---dc20

DNLM/DLC

for Library of Congress

94-22471

CIP

The use of portions of the text of USP23/NF18, copyright 1994, is by permission of the USP Convention, Inc. The Convention is not responsible for any inaccuracy of quotation or for any false or misleading implication that may arise from separation of excerpts from the original context or by obsolescence resulting from publication of a supplement.

PRINTED IN THE UNITED STATES OF AMERICA

Print No. 4 3 2 1



#### **Drug Stability**

One of the most important activities of preformulation work is the evaluation of the physical and chemical stability of the pure drug substance. It is essential that these initial studies be conducted using drug samples of known purity. The presence of impurities can lead to erroneous conclusions in such evaluations. Stability studies conducted in the preformulation phase include solid state stability of the drug alone, solution phase stability, and stability in the presence of expected excipients.

Initial investigation begins through knowledge of the drug's chemical structure which allows the preformulation scientist to anticipate the possible degradation reactions.

Chemical instability of medicinal agents may take many forms, because the drugs in use today are of such diverse chemical constitution. Chemically, drug substances are alcohols, phenols, aldehydes, ketones, esters, ethers, acids, salts, alkaloids, glycosides, and others, each with reactive chemical groups having different susceptibilities toward chemical instability. Chemically, the most frequently encountered destructive processes are hydrolysis and oxidation.

Hydrolysis is a solvolysis process in which (drug) molecules interact with water molecules to yield breakdown products of different chemical constitution. For example, aspirin or acetylsalicylic acid combines with a water molecule and hydrolyzes into one molecule of salicylic acid and one molecule of acetic acid:

The process of hydrolysis is probably the most important single cause of drug decomposition mainly because a great number of medicinal agents are esters or contain such other groupings as substituted amides, lactones, and lactams, which are susceptible to the hydrolytic process.

Another destructive process is oxidation. The oxidative process is destructive to many drug types, including aldehydes, alcohols, phenols, sugars, alkaloids, and unsaturated fats and oils.

Chemically, oxidation involves the loss of electrons from an atom or a molecule. Each electron lost is accepted by some other atom or molecule, thereby accomplishing the reduction of the recipient. In inorganic chemistry, oxidation is accompanied by an increase in the positive valence of an element—for example, ferrous (+2) oxidizing to ferric (+3). In organic chemistry, oxidation is frequently considered synonymous with the loss of hydrogen (dehydrogenation) from a molecule. The oxidative process frequently involves free chemical radicals, which are molecules or atoms containing one or more unpaired electrons, as molecular (atmospheric) oxygen (•O—O•) and free hydroxyl (•OH). These radicals tend to take electrons from other chemicals, thereby oxidizing the donor. Many of the oxidative changes in pharmaceutical preparations have the character of autoxidations. Autoxidations occur spontaneously under the initial influence of atmospheric oxygen and proceed slowly at first and then more rapidly as the process continues. The process has been described as a type of chain reaction commencing by the union of oxygen with the drug molecule and continuing with a free radical of this oxidized molecule participating in the destruction of other drug molecules and so forth.

In drug product formulation work, steps are taken to reduce or prevent the occurrence of drug substance deterioration due to hydrolysis, oxidation, and other processes. These techniques are discussed in a later section.

### **Pharmaceutic Ingredients**

In order to prepare a drug substance into a final dosage form, pharmaceutic ingredients are required. For example, in the preparation of pharmaceutic solutions, one or more solvents are utilized to dissolve the drug substance, preservatives may be added to prevent microbial growth, stabilizers may be used to prevent drug decomposition, and colorants and flavorants added to enhance product appeal. In the preparation of tablets, diluents or fillers are commonly added to increase the bulk of the formulation, binders to cause the adhesion of the powdered drug and pharmaceutic substances, antiadherents or lubricants to assist the smooth tableting process, disintegrating agents to promote tablet break-up after administration, and coatings to improve stability, control disintegration, or to enhance appearance. Ointments, creams, and suppositories achieve their characteristic features due to the pharmaceutic bases which are utilized. Thus, for each dosage form, the pharmaceutic ingredients



establish the primary features of the product, and contribute to the physical form, texture, stability, taste and overall appearance.

Table 4–2 presents the principal categories of pharmaceutic ingredients, with examples of some of the official and commercial agents currently used. Additional discussion of many of the pharmaceutic ingredients may be found in the chapters where they are most relevant; for example, pharmaceutic materials used in tablet and capsule formulation are discussed in Chapter 5, Peroral Solids, Capsules, Tablets, and Controlled-Release Dosage Forms.

The reader should also be aware of the Hand-

Table 4-2. Examples of Pharmaceutic Ingredients

Ingredient Type	Definition	Examples
Acidifying Agent	Used in liquid preparations to provide acidic medium for product stability.	acetic acid citric acid fumaric acid hydrochloric acid nitric acid
Alkalinizing Agent	Used in liquid preparations to provide alkaline medium for product stability.	ammonia solution ammonium carbonate diethanolamine monoethanolamine potassium hydroxide sodium borate sodium carbonate sodium hydroxide triethanolamine trolamine
Adsorbent	An agent capable of holding other molecules onto its surface by physical or chemical (chemisorption) means.	powdered cellulose activated charcoal
Aerosol Propellant	An agent responsible for developing the pressure within an aerosol container and expelling the product when the valve is opened.	carbon dioxide dichlorodifluoromethane dichlorotetrafluoroethane trichloromonofluoromethane
Air Displacement	An agent which is employed to displace air in a hermetically sealed container to enhance product stability.	nitrogen
Antifungal Preservative	Used in liquid and semi-solid preparations to prevent the growth of fungi. The effectiveness of the parabens is usually enhanced when they are used in combination.	benzoic acid butylparaben ethylparaben methylparaben propylparaben sodium benzoate sodium propionate
Antimicrobial Preservative	Used in liquid and semi-solid preparations to prevent the growth of microorganisms.	benzalkonium chloride benzethonium chloride benzyl alcohol cetylpyridinium chloride chlorobutanol phenol phenylethyl alcohol phenylmercuric nitrate thimerosal



Table 4–2. Continued

Ingredient Type	Definition	Examples
Antioxidant -	An agent which inhibits oxidation and thus is used to prevent the deterioration of preparations by the oxidative process.	ascorbic acid ascorbyl palmitate butylated hydroxyanisole butylated hydroxytoluene hypophophorous acid monothioglycerol propyl gallate sodium ascorbate sodium bisulfite sodium formaldehyde sulfoxylate sodium metabisulfite
Buffering Agent	Used to resist change in pH upon dilution or addition of acid or alkali.	potassium metaphosphate potassium phosphate, monobasic sodium acetate sodium citrate anhydrous and dihydrate
Chelating Agent	A substance that forms stable, water soluble complexes (chelates) with metals. Chelating agents are used in some liquid pharmaceuticals as stabilizers to complex heavy metals which might promote instability. In such use they are also called sequestering agents.	edetate disodium edetic acid
Colorant	Used to impart color to liquid and solid (e.g., tablets and capsules) pharmaceutical preparations.	FD&C Red No. 3 FD&C Red No. 20 FD&C Yellow No. 6 FD&C Blue No. 2 D&C Green No. 5 D&C Orange No. 5 D&C Red No. 8 caramel ferric oxide, red
Clarifying Agent	Used as a filtering aid because of adsorbent qualities.	bentonite
Emulsifying Agent	Used to promote and maintain the dispersion of finely subdivided particles of a liquid in a vehicle in which it is immiscible. The end product may be a liquid emulsion or semisolid emulsion (e.g., a cream).	acacia cetomacrogol cetyl alcohol glyceryl monostearate sorbitan monooleate polyoxyethylene 50 stearate
Encapsulating Agent	Used to form thin shells for the purpose of enclosing a drug substance or drug formulation for ease of administration.	gelatin cellulose acetate phthalate
Flavorant	Used to impart a pleasant flavor and often odor to a pharmaceutical preparation. In addition to the natural flavorants listed, many synthetic flavorants are also used.	anise oil cinnamon oil cocoa menthol orange oil peppermint oil vanillin



# DOCKET

# Explore Litigation Insights



Docket Alarm provides insights to develop a more informed litigation strategy and the peace of mind of knowing you're on top of things.

# **Real-Time Litigation Alerts**



Keep your litigation team up-to-date with **real-time** alerts and advanced team management tools built for the enterprise, all while greatly reducing PACER spend.

Our comprehensive service means we can handle Federal, State, and Administrative courts across the country.

## **Advanced Docket Research**



With over 230 million records, Docket Alarm's cloud-native docket research platform finds what other services can't. Coverage includes Federal, State, plus PTAB, TTAB, ITC and NLRB decisions, all in one place.

Identify arguments that have been successful in the past with full text, pinpoint searching. Link to case law cited within any court document via Fastcase.

## **Analytics At Your Fingertips**



Learn what happened the last time a particular judge, opposing counsel or company faced cases similar to yours.

Advanced out-of-the-box PTAB and TTAB analytics are always at your fingertips.

### API

Docket Alarm offers a powerful API (application programming interface) to developers that want to integrate case filings into their apps.

#### **LAW FIRMS**

Build custom dashboards for your attorneys and clients with live data direct from the court.

Automate many repetitive legal tasks like conflict checks, document management, and marketing.

#### **FINANCIAL INSTITUTIONS**

Litigation and bankruptcy checks for companies and debtors.

## **E-DISCOVERY AND LEGAL VENDORS**

Sync your system to PACER to automate legal marketing.

