What You Want to Know About Generic Drugs

Generic Drugs: Safe. Effective. FDA-Approved.
What do you want to know about ALL the medicines you use.... brand-name and generic?

• safe and effective
• affordable and available
• use with confidence
Myths about Generic Drugs

- Generics...are not as safe
- Generics...are not as potent
- Generics...take longer to act in the body
- Generics...are made in sub-standard facilities
What You Want to Know About Generic Drugs

I. How FDA assures that generics:
   • are Safe. Effective. FDA-approved.
   • are an affordable alternative
   • can be used with confidence
What You Want to Know About Generic Drugs

II. How FDA and the administration are working to make more generic medicines available to the American public
# What You Want to Know About Generic Drugs

<table>
<thead>
<tr>
<th>Brand-name drug</th>
<th>Generic drug</th>
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<tbody>
<tr>
<td>• supplied by one drug company</td>
<td>• may be supplied by more than one company</td>
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<tr>
<td>• sold under drug company’s trademarked name</td>
<td>• may be sold under active ingredient(s) name(s)</td>
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What is a generic drug?

A copy of a brand-name drug, which must have the:

- same quality
- same safety
- same strength
What You Want to Know About Generic Drugs

Both brand name and generics drugs:

• are approved by the FDA
• must meet the same FDA standards for quality
Generic Competition

It is essential to have brand-name and generic drugs available.
Generic Competition

- helps keep drug costs down
- encourages research
- helps keep insurance premiums down
- saves consumers $8 to $10 billion yearly
Patent Protection

A patent:

- protects the investment of the drug company that developed the drug (the manufacturer)
- gives the drug company the sole right to sell the drug while the patent is in effect
When the patents on a brand-name drug near expiration, drug companies that want to manufacture a generic can apply to the FDA to sell a generic version of the drug.
Generic Drug Review

- Much the same as new, brand name drug review
- 8 major parts
Generic Drug Review

1. FDA-approved generic drugs must have:
   • same active ingredient(s)
   • same labeled strength
   • same dosage form
   • same administration
Generic Drug Review

2. The drug company must show the generic drug is "bioequivalent" to the brand-name drug.
   • active ingredient works in the same way
   • active ingredient works in the same amount of time
3. The generic drug’s labeling must be basically the same as that of the approved brand-name drug.
Generic Drug Review

4. The drug company must:
   - fully document the generic drug’s chemistry, manufacturing steps, and quality control measures
   - detail each step of the process
5. The raw materials and the finished product must meet USP specifications, if these have been set.

USP-United States Pharmacopeia
Generic Drug Review

6. The drug company must:

• show that its generic drug maintains stability as labeled before it can be sold
• continue to monitor drug’s stability
Generic Drug Review

7. **The drug company must:**
   - comply with federal regulations for current good manufacturing practices
   - give a full description of the facilities it uses to manufacture, process, test, package, label, and control the drug
8. Inspection at the proposed manufacturing site ensures that the firm:

- is capable of meeting commitments of the application
- can manufacture the product consistently
FDA Requirements for Brand-Name and Generic Drugs

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Brand Name Drug</th>
<th>Generic Drug</th>
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<tr>
<td>For reformulations of a brand-name drug or generic versions of a drug, FDA reviews data showing the drug is bioequivalent to the one used in the original safety and efficacy testing.</td>
<td>✔️</td>
<td>✔️</td>
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<tr>
<td>FDA evaluates the manufacturer's adherence to good manufacturing practices before the drug is marketed.</td>
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<td>✔️</td>
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<tr>
<td>FDA reviews the active and inactive ingredients used in the formulation before the drug is marketed.</td>
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<td>✔️</td>
</tr>
<tr>
<td>FDA reviews the actual drug product.</td>
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<td>✔️</td>
</tr>
<tr>
<td>FDA reviews the drug's labeling.</td>
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</tr>
<tr>
<td>Manufacturer must seek FDA approval before making major manufacturing changes or reformulating the drug.</td>
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<tr>
<td>Manufacturer must report adverse reactions and serious adverse health effects to the FDA.</td>
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<tr>
<td>FDA periodically inspects manufacturing plants.</td>
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<tr>
<td>FDA monitors drug quality after approval.</td>
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What You Want to Know About Generic Drugs

II. What the FDA and the administration are doing to make more generic medicines available to the American public
Making More Generics Available

A. Hatch-Waxman Act, or the Patent Term Restoration Act of 1984

B. President’s 2003 budget increased FDA’s funding to speed up generic drug reviews (more reviewers, etc.)
Making More Generics Available

C. “Improving Access to Generic Drugs” initiative

- new regulatory processes to reduce time and cost of generic drug approvals
Making More Generics Available

C. “Improving Access to Generic Drugs” initiative (continued)

- enhanced public and professional education
C. “Improving Access to Generic Drugs” initiative (continued)

• enhanced scientific study of generic drugs
Making More Generics Available

C. “Improving Access to Generic Drugs” initiative (continued)

- enhanced monitoring of the safety of generic drugs
Print Public Service Ads for Consumers

You know that question
that goes through your mind
when you take your
generic drug?
Here's the answer.

FDA ensures that your generic drug is safe and effective. All generic drugs are put through a rigorous, multi-step approval process. From quality and performance to manufacturing and labeling, everything must meet FDA's high standards. We make it tough to become a generic drug in America so it's easy for you to rest assured.

Visit www.fda.gov/cder/ or call 1-888-INFO-FDA to learn more.

Generic Drugs: Safe. Effective. FDA Approved.

If you're experiencing anxiety
about taking your
generic drug,
read this ad and repeat as needed.

FDA ensures that your generic drug is safe and effective. All generic drugs are put through a rigorous, multi-step approval process. From quality and performance to manufacturing and labeling, everything must meet FDA's high standards. We make it tough to become a generic drug in America so it's easy for you to rest assured.

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Generic Drugs: Safe. Effective. FDA Approved.
Public Transportation Ads in U.S. Cities
What is a generic drug?

- Consistent labeling
- Quality control
- Same drug
- Performance evaluation
- Rigorous manufacturing standards

When a brand-name drug's patent protection expires, generic versions of the drug can be created and sold. The generic version works like the brand-name drug in dosage, strength, performance and use, and must meet the same quality and safety standards. All generic drugs must be approved by FDA.

How does FDA ensure that my generic drug is as safe and effective as the brand-name drug?

All generic drugs are put through a rigorous, multi-step approval process that includes a review of scientific data on the generic drug's ingredients, performance and effectiveness. FDA also conducts continuous inspections of the manufacturing plant, and monitors drug quality—even after the generic has been approved.

If generic drugs and brand-name drugs have the same active ingredients, why do they look different?

The drugs look different because certain inactive ingredients—like colors and flavorings—may be different. These ingredients do not affect the performance of the generic drug in any way, but trademark laws in the U.S. do not allow a generic drug to look exactly like drugs already on the market.

Is my generic drug made by the same company that makes the brand-name drug?

Quite possibly, but not always. Brand-name companies are responsible for manufacturing approximately 20 percent of generic drugs. They frequently make versions of their own or other brand-name drugs. There are also other approved companies that produce generic drugs.

Are generic drugs always made in the same kind of facilities as brand-name drugs?

Yes. Both brand-name and generic drug facilities must meet the same standards of good manufacturing practices. FDA will not permit drugs to be made in substandard facilities. FDA conducts about 5,000 inspections a year to ensure standards are met.

¿Qué son los medicamentos genéricos y por qué son importantes para usted? Todo lo que necesita saber sobre los medicamentos genéricos.

FDA makes it tough to become a generic drug in America so you can feel confident about taking generic drugs. If you still want to learn more, talk with your doctor, pharmacist, medical provider or insurance company. Or call 1-888-INFO-FDA today.

Generic Drugs: Safe, Effective, FDA Approved.
You know that question that goes through your mind when you take your **generic drug?**

**Here's the answer.**

FDA ensures that all generic drugs are put through a rigorous, multi-step review process. From manufacturing to labeling, everything must meet FDA's high standards. We make it tough to become a generic drug in America so you can feel confident. **Generic Drugs: Safe. Effective. FDA Approved.**

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¿Recuerda esa preguntita que siempre se hace cuando toma **medicamentos genéricos?**

Aquí tiene la respuesta.

La Agencia FDA del gobierno federal se asegura que todos los medicamentos genéricos sean sometidos a un riguroso proceso de revisión. Desde su fabricación hasta la colocación de la etiqueta, todo debe cumplir con las más altas normas de calidad de la FDA. Y hacemos el proceso exigente para que en los Estados Unidos, todos podamos confiar en los medicamentos genéricos. **Medicamentos genéricos: Seguros. Efectivos. Aprobados por la FDA.**
Blue Cross Blue Shield of Michigan Billboards

Generic drugs:
Safe. Effective. FDA approved.

Food and Drug Administration
Posters Appearing in 4,000 Walgreens Pharmacies Nationwide
Making More Generics Available

D. FDA’s generic drug Final Rule, 8-18-03

• seeks to close legal loopholes in the Hatch Waxman Act that delay generic drug approval
Making More Generics Available

D. FDA’s generic drug Final Rule, 8-18-03 (continued)

• implements an FTC recommendation to tighten the patent submission and listing process
D. FDA’s generic drug Final Rule, 8-18-03 (continued)

• clarifies the types of patents that must be submitted to the FDA
E. House and Senate passed a bill, “Greater Access to Affordable Pharmaceuticals Act”

• complements FDA’s final rule.
• FDA is working with Congress
Myths and Facts about Generic Drugs
Myths and Facts about Generic Drugs

MYTH #1

Generics are not as safe as brand-name drugs.
Myths and Facts about Generic Drugs

FACT #1

Generics use the same ingredients, and
• work the same in the body
• have the same risk-benefit profile
Myths and Facts about Generic Drugs

MYTH #2

Generics are not as potent as brand-name drugs.
Myths and Facts about Generic Drugs

FACT #2

Generic drugs have the same quality, strength, purity and stability.
Myths and Facts about Generic Drugs

**MYTH #3**

Generics take longer to act in the body.
Myths and Facts about Generic Drugs

FACT #3

The generic drug delivers the same amount of active ingredient in the same time as the original drug.
Myths and Facts about Generic Drugs

MYTH #4

Brand-name drugs are made in modern manufacturing facilities, and generics are often made in sub-standard facilities.
Myths and Facts about Generic Drugs

FACT #4

Sub-standard facilities are not permitted by the FDA.
What you want to know about Generic Drugs

For more information on generic drugs, visit the FDA website at:

http://www.fda.gov/cder/ogd/index.htm
What you want to know about Generic Drugs

Contact your physician, pharmacist, or insurance company for more information about your generic drugs.
Generic Drugs:
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(Name of presenter)

>Title of presenter

November 19, 2003
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FDA Acts to Lower Drug Prices By Implementing New Regulation to Improve Generic Drug Competition, June 12, 2003 - TODAY'S ACTION

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