



An Overview of Biologic and Biosimilar Products

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- ✓ I have no conflicts of interest or financial relationships to disclose
- ✓ I will not discuss off label use and/or investigational use in this presentation

Learning Objectives

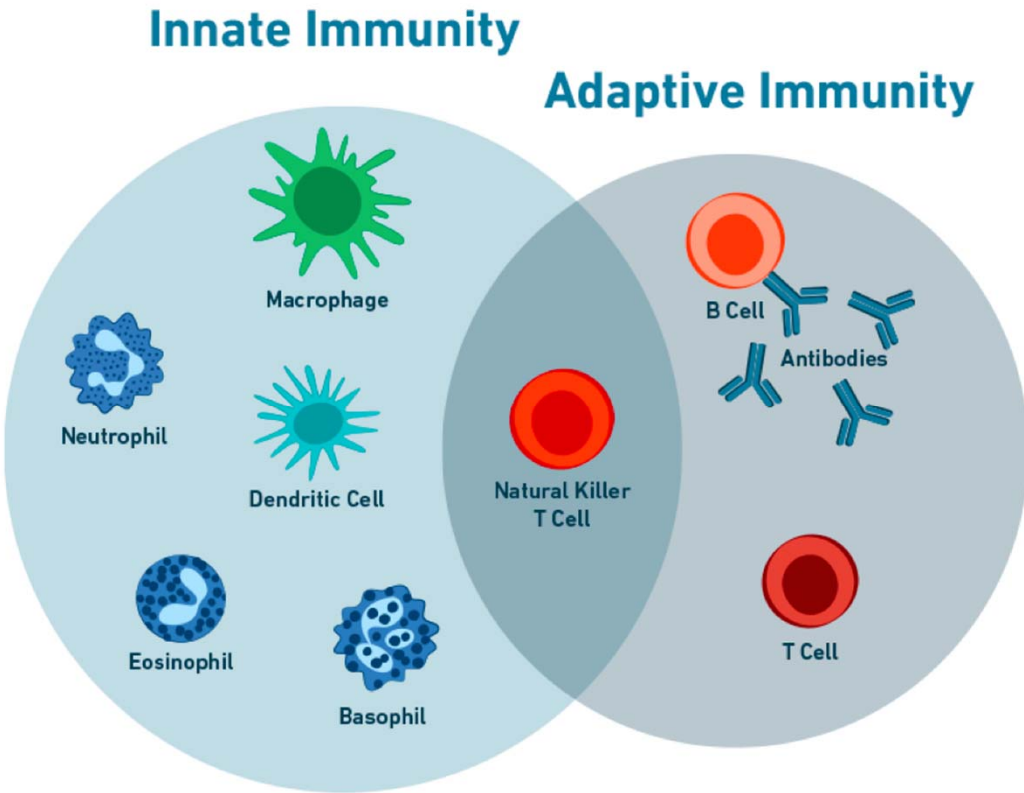
Pharmacist Learning Objectives

1. Identify the most common diseases in which immunomodulating biologics are used and their mechanism of action in those disease states.
2. List the common and severe adverse effects, precautions, contraindications, black box warnings, testing, and screening considerations associated with immunomodulating biologics.
3. Recognize the key differences between reference biologic products and biosimilars and the FDA guidance and Illinois laws on their interchangeability.
4. Explain the 2021 nomenclature update from the World Health Organization (WHO) and International Nonproprietary Names (INN)

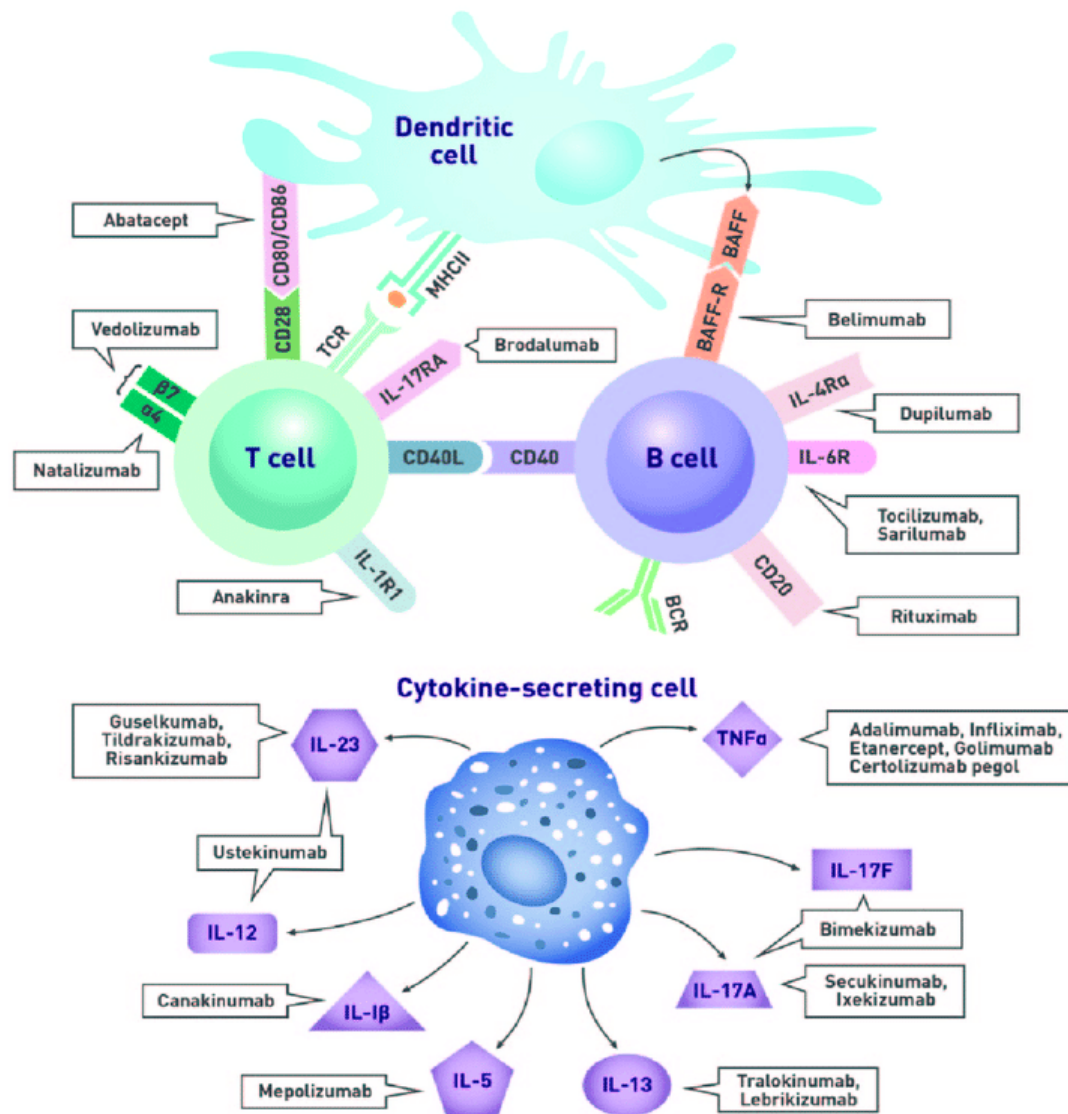
Pharmacy Technician Learning Objectives

1. Recall the history, types of molecules, process of manufacturing, regulation, naming and unique properties of biologics versus biosimilars
2. Identify common disease states in which biological products are used.
3. Describe the FDA guidance and state of Illinois regulation regarding interchangeability of reference biologic products for their biosimilars

Immunology Review



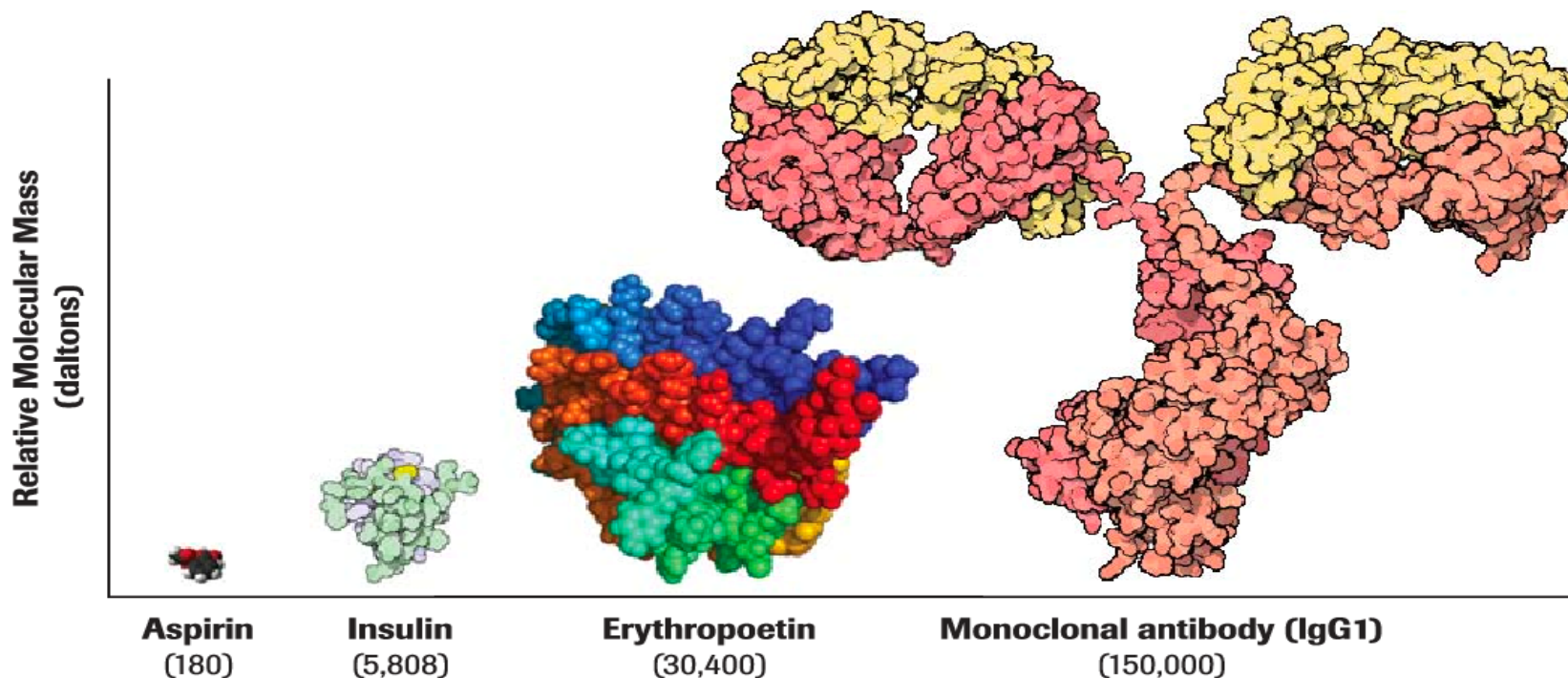
Immunological Targets of Biologics



What are Biologic Products?

- Protein-based therapies derived from living organisms or their cells including plants, animals, and microorganisms such as yeast and bacteria
- Genetically engineered for a specific medical purpose
- Typically larger, more complex molecules when compared to traditional drugs
- Examples: vaccines, monoclonal antibodies, therapeutic proteins
- The Public Health Service (PHS) Act definition:
 - “virus, therapeutic serum, toxin, antitoxin, vaccine, blood, blood component or derivative, allergenic product, or analogous product applicable to the prevention, treatment, or cure of a disease or condition of human beings.”

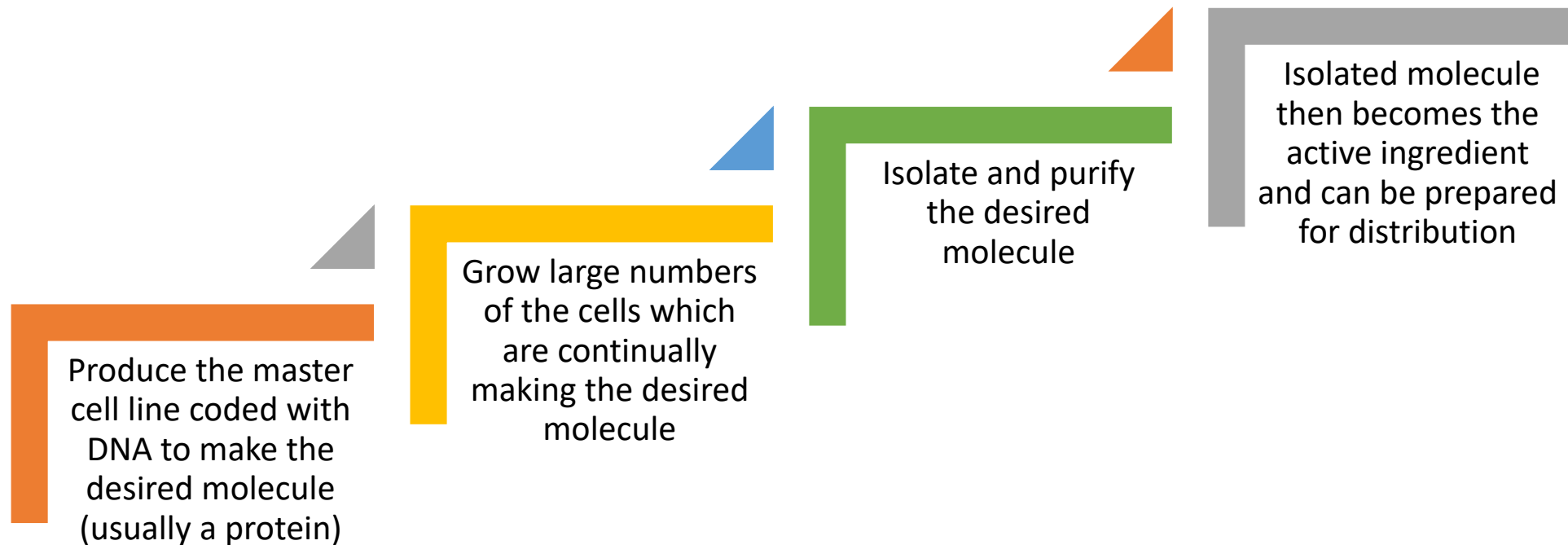
Traditional small-molecule drugs vs. Biologics



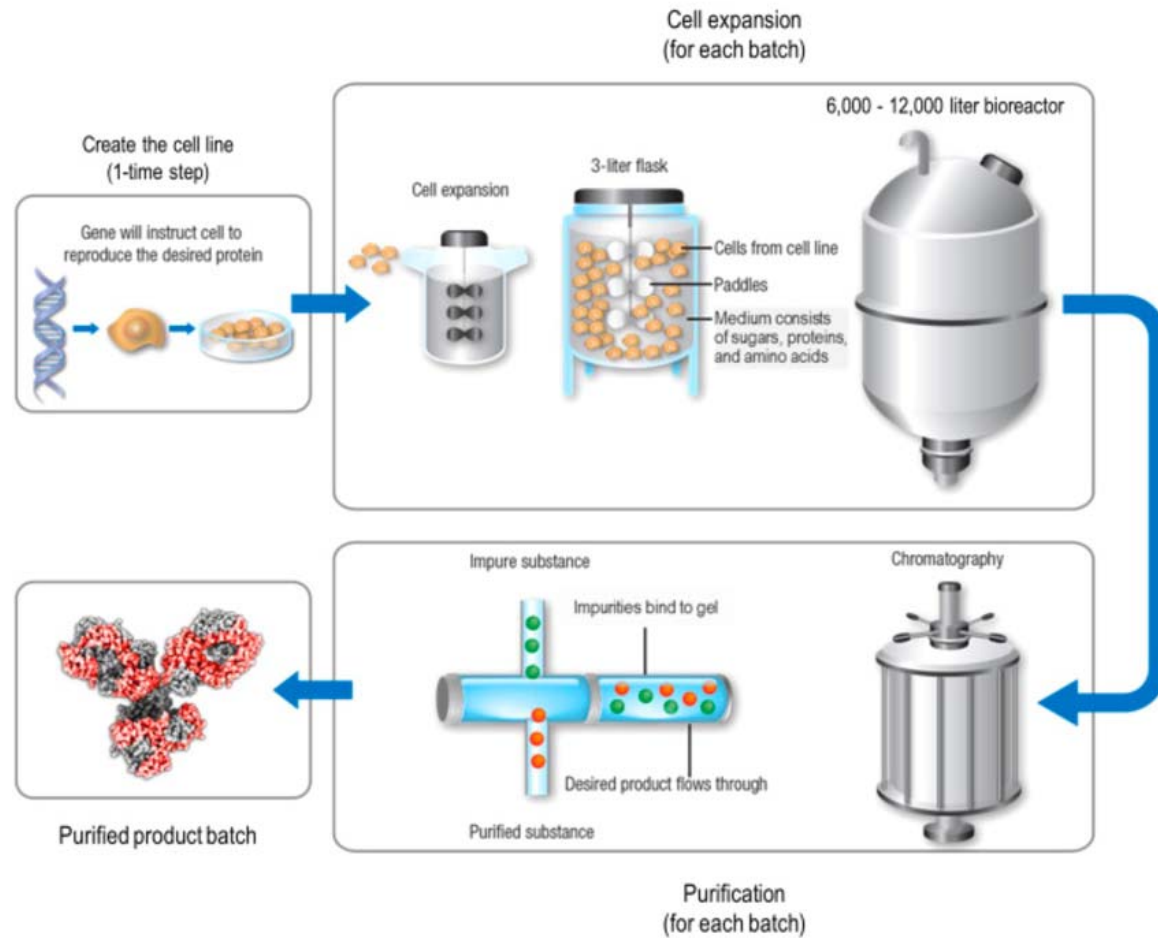
Molecular size comparison of a small molecule drug (aspirin) to 3 different classes of biologics

(Relative molecular masses shown in parenthesis)

How Are Biologics Manufactured?



How Are Biologics Manufactured?



How Are Biologics Administered?

Small Molecule Drugs

Can be taken orally
(i.e. tablets or capsules)

Disintegrates in the GI
tract, then absorbed into
the bloodstream

Typically self
administered

Biologics

Cannot be given orally

Most are given parenterally
via SQ, IM, or IV routes

Administered at prescriber's
office, infusion center, or
self-administered



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Checkpoint 1: Which of the following is TRUE regarding biologics when compared with traditional small-molecule drugs?



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Biologics are given orally due to high oral bioavailability

0%



Biologics are less sensitive to light and temperature variations

0%



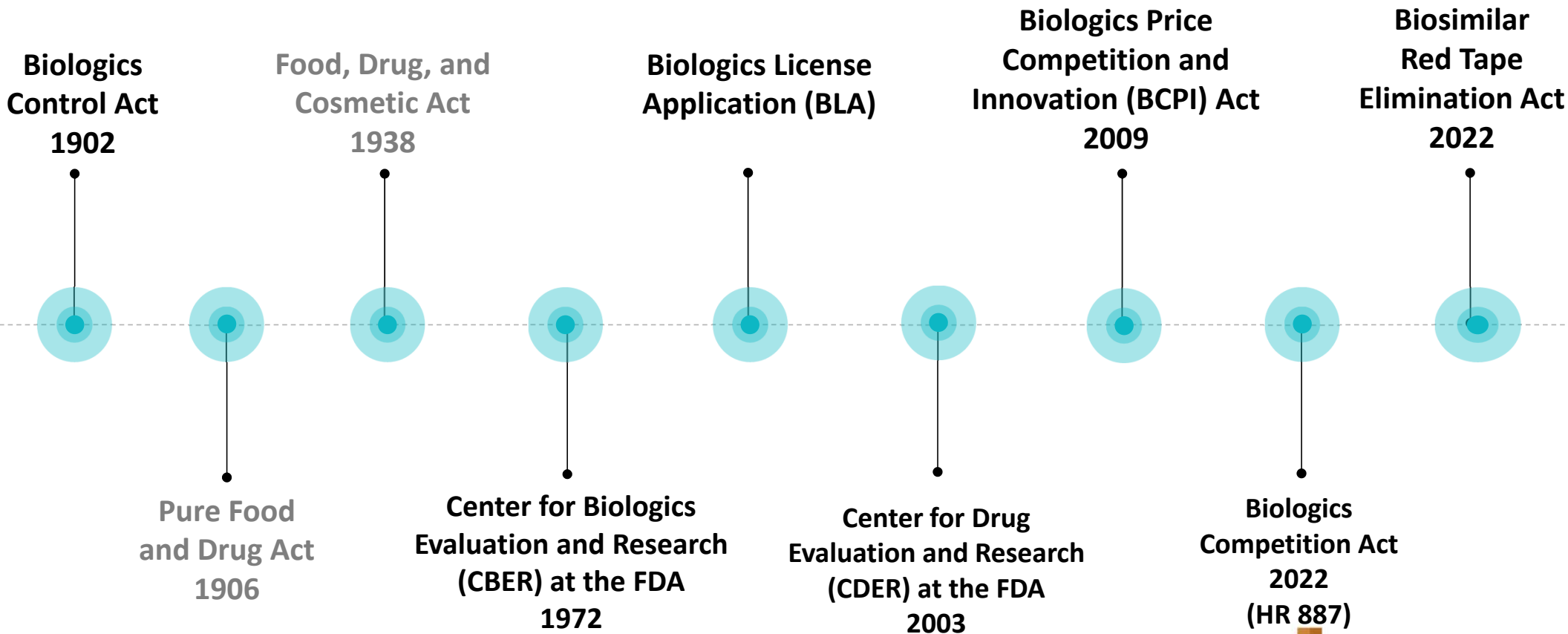
Biologics are larger in molecular size and have more complex and fragile structures

0%



Biologics are less costly to manufacture

Biologics Regulation Timeline



Kennedy, D. Public Health Rep. 1978.
US Pharmacopeia. Biologics Control Act. 2010.
Junod SW. US Food and Drug Administration. 2023
GaBI Online - Generics and Biosimilars Initiative. 2023

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Test Your Knowledge #1: Pharmacists & Techs



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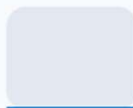
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1. (P&T) Which of the follow department(s) regulate biological products for human use under applicable federal laws?



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Center for Drug Evaluation and Research (CDER)

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World Health Organization (WHO)

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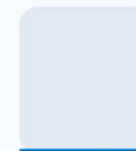
Center for Biologics Evaluation and Research (CBER)

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United States Pharmacopeia (USP)

0%



Both A & C



Immunomodulating Biologic Therapies

Diseases In Which Biologics Are Used

Dermatology

- Eczema, plaque psoriasis

Ophthalmology

- Uveitis, ocular surface disease, thyroid eye disease

Rheumatology

- Rheumatoid arthritis, psoriatic arthritis, ankylosing spondylitis

Gastrointestinal diseases

- Crohn's disease, ulcerative colitis

Oncology

- Many types of cancers

Biologic Therapeutic Targets

Cytokine modulators

- TNF- α inhibitors
- IL inhibitors

Lymphocyte modulators

- B lymphocyte depletion therapy

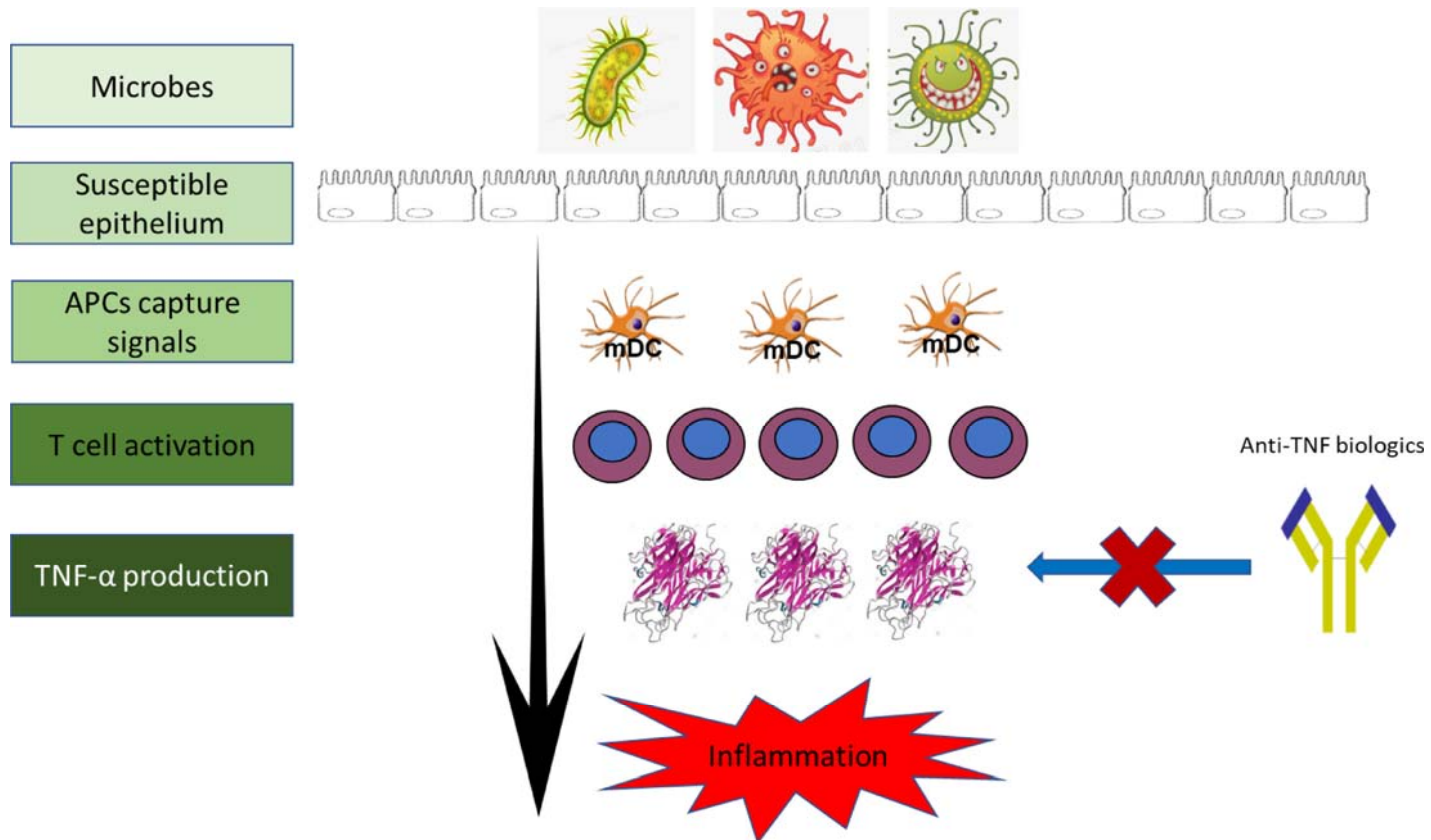


Cytokine Modulators

Cytokine modulators: TNF- α inhibitors

- ▮ TNF- α is a proinflammatory cytokine
- ▮ Inappropriate or excessive activation of TNF- α signaling leads to chronic inflammation and disease
- ▮ TNF- α inhibitors bind to the TNF receptor and blocks receptor and thus reduces the inflammatory process
- ▮ Commonly used in multiple autoimmune diseases
- ▮ Currently five agents on the market
 - ▮ Adalimumab (Humira[®]), infliximab (Remicade[®]), certolizumab (Cimzia[®]), golimumab (Simponi[®]), etanercept (Enbrel[®])

Mechanism of Action, Example Inflammatory Bowel Disease (IBD)



TNF- α Inhibitors: Clinical Pearls

Common Adverse Effects

- Injection site or infusion reactions
- Upper respiratory infections
- Abdominal pain, nausea, diarrhea
- Rash
- Anemia
- Headaches
- Elevated LFTs

Serious Adverse Effects

- Increased risk of malignancies
- Serious infections: bacterial, viral, & fungal infections
- CHF exacerbation
- Demyelinating disorders (e.g., multiple sclerosis)

Contraindications

- CHF class III or IV
- Hypersensitivity to any of part of the biologic product
- Live vaccines

Black Box Warning

- Increased risk of serious infections that can lead to death
- Increased risk of malignancies, especially lymphomas

Pretreatment Screening

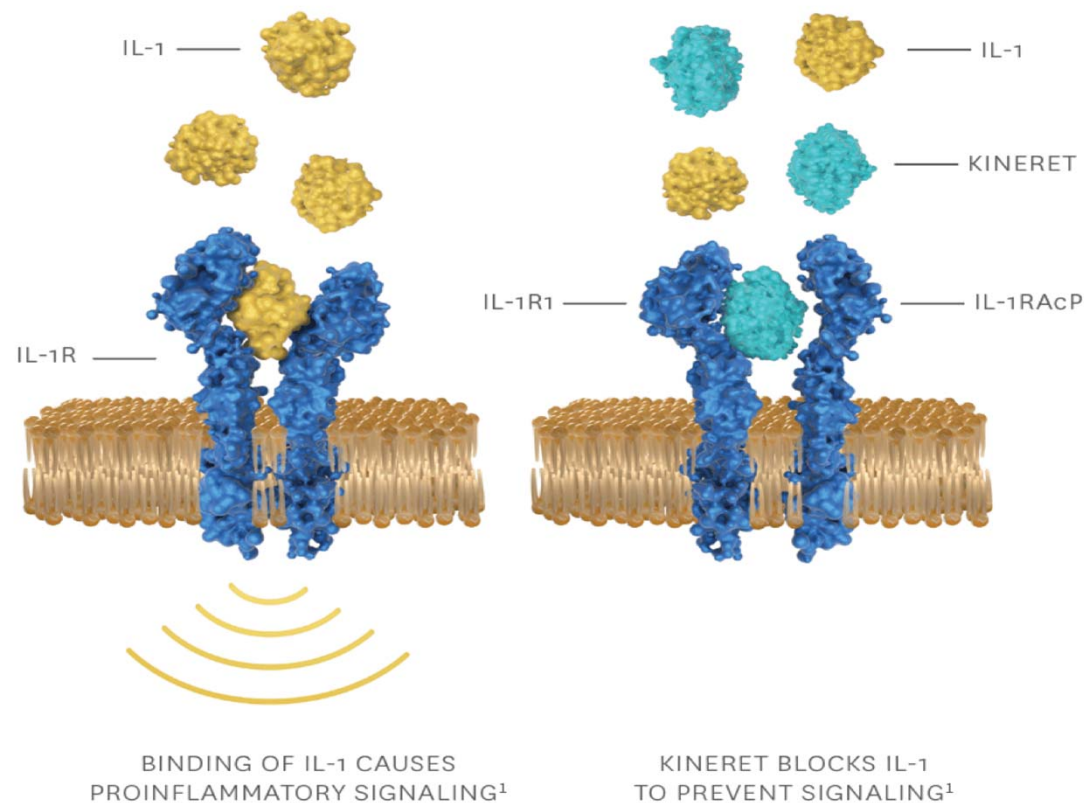
- Hepatitis B & C, HIV, tuberculosis
- CBC, BMP, LFTs
- Vaccination status

Cytokine modulators: IL-inhibitors

- ▣ Naturally occurring large group of cytokines
- ▣ Many interleukin types with varying effects
 - ▣ Pro-inflammatory and anti-inflammatory
- ▣ Targeting specific proinflammatory IL receptors to treat diseases
 - ▣ Travels to target cell and binds it via the receptor on the cell's surface
- ▣ Example: IL-6 receptor antagonist, tocilizumab (Actemra[®]) used in the treatment of rheumatoid arthritis

Mechanism of Action, Example

IL-1 receptor antagonist, Anakinra (Kineret[®])



IL-inhibitors: Clinical Pearls

Common Adverse Effects

- Neutropenia
- Injection site or infusion reactions
- Abdominal pain
- URIs
- Headaches
- Elevated LFTs

Serious Adverse Effects

- Anaphylaxis
- GI adverse effects (e.g. diverticulitis, Crohn's disease exacerbation)

Contraindications

- Hypersensitivity to any of part of the biologic
- Live vaccines

Black Box Warning

- Increased risk of serious infections that can lead to death, with some IL-inhibitors

Pretreatment Screening

- TB screening
- LFTs
- Baseline CBC



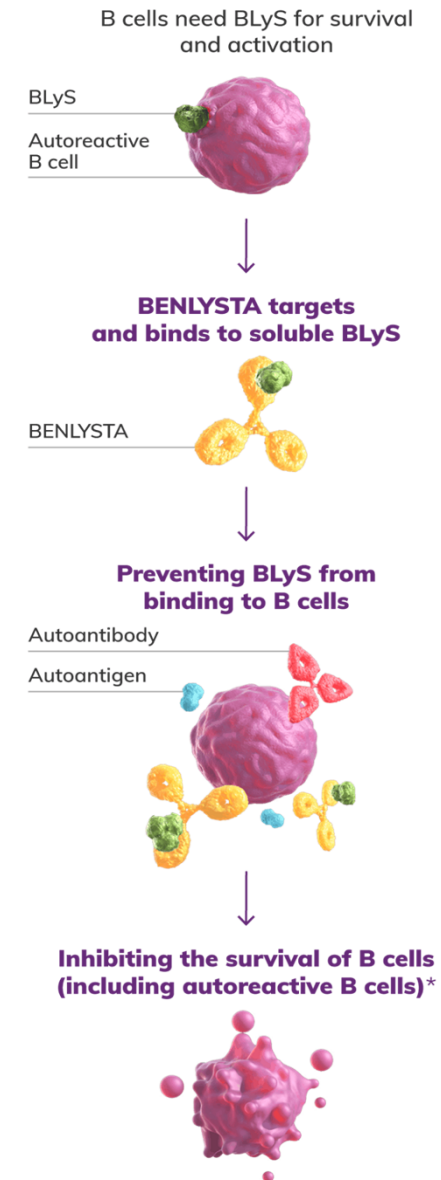
Lymphocyte Modulators

B Lymphocyte Depletion Therapy

- ▣ B cells play a key role in the pathogenesis of autoimmune diseases
 - ▣ Production of autoantibodies
 - ▣ Presentation of autoantigens to autoreactive T cells
 - ▣ Secretion of pro-inflammatory cytokines
- ▣ The main goal of B cell depletion therapy is to destroy harmful B cells while retaining protective B cell immunity
- ▣ Depletion is achieved through two main mechanisms:
 - ▣ Inhibition of the B-lymphocyte stimulator protein (BLyS)
 - ▣ Direct target by monoclonal antibodies against B-cell surface molecules, e.g. CD20 (most common)


Mechanism of Action, Example

B Lymphocyte Depletion Therapy Belimumab (Benlysta®)



B Lymphocyte Depletion Therapy: Clinical Pearls

Examples	Belimumab (Benlysta®) Anti-BLyS	Rituximab (Rituxan®) Anti-CD20
Dosing	10 mg/kg IV q 2wks x 3, then q 4 wks OR 400 mg inj SQ wkly x 4, then 200 mg qwk	375 mg/m ² BSA IV weekly x 4 OR 500-1,000 mg IV on days 1 and 15
Common adverse effects	Nausea, diarrhea, fever, bronchitis, nasopharyngitis, insomnia, depression, migraine, pharyngitis, infusion reaction	Lymphopenia/neutropenia, infection, asthenia, fever
Precautions	Serious infections, anaphylaxis, infusion reactions, psychiatric illnesses (e.g., depression, suicidal ideation), cancers	Tumor lysis syndrome, infections, cardiac adverse events, renal toxicity, bowel obstruction/perforation
	Live vaccines, concurrent use with other biologics	
Black Box Warnings	None	Fatal infusion-related reactions, severe mucocutaneous reactions, reactivation of hepatitis B virus, progressive multifocal leukoencephalopathy (PML) and death
Pretreatment screening	Depression and suicide risk	Hepatitis B, CBC



Test Your Knowledge #2: Pharmacists



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2. (P) Which of the following cytokine or lymphocyte modulators is contraindicated in patients with CHF Class III or IV

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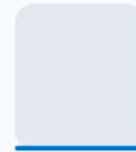
Interleukin (IL) inhibitors

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TNF-alpha inhibitors

0%



B cell depletion therapy

0%



T cell activator



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Checkpoint 2: Prior to initiating most biologics, which of the following screenings should be conducted? (select all that apply)

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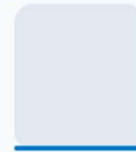
Hepatitis B and C

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Tuberculosis

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


Liver Function Tests (LFTs)

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Complete Blood Count (CBC)



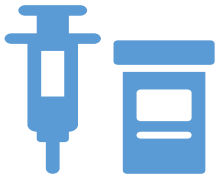
Biologics → Biosimilars



Biologics Price Competition and Innovation (BCPI) Act

- Created an abbreviated licensure pathway for biological products shown to be *biosimilar* to or *interchangeable* with an FDA-licensed reference product
- The FDA provides guidance to industry regarding scientific considerations in demonstrating biosimilarity
 - MUST show comparable clinical effectiveness, safety, and purity to be approved as a biosimilar
- Helps to address the high cost of biologics and provides increased access to safe and effective, lower cost biosimilar alternatives

BCPI Act: Key Definitions



Reference Product

Reference Product

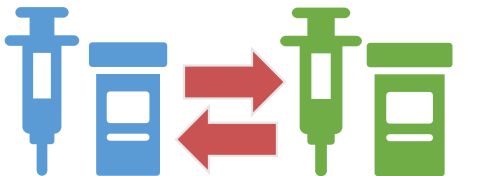
A reference product is the single biological product, already approved by the FDA, against with a proposed biosimilar product is being compared



Biosimilar Product

Biosimilar Product

A biosimilar is a biological product that is highly similar to and has no clinically meaningful differences from an existing FDA-approved reference product



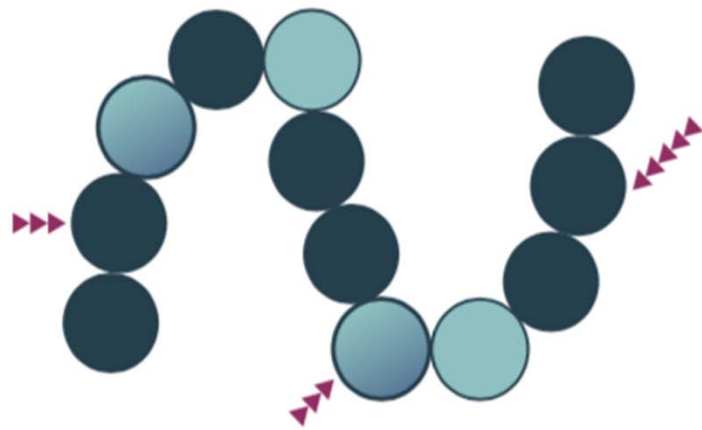
Reference Product

Interchangeable Product

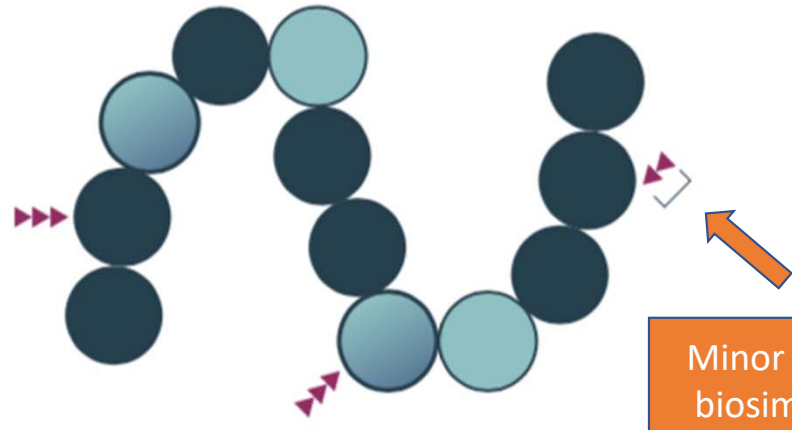
Interchangeable Product

- A biosimilar expected to produce the same clinical result as the reference product
- There is no safety risk of reduced efficacy associated with switching between the reference and interchangeable product
- May be substituted for reference product depending on state pharmacy laws

Reference Biologic vs. Biosimilar



Reference Biologic Product



Biosimilar Product

Minor difference in biosimilar product compared to reference product



Test Your Knowledge #3: Pharmacists & Techs



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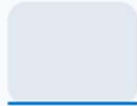
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3. (P&T) The difference between a biologic and a biosimilar is that a biosimilar is **MUST:**

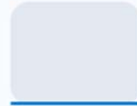


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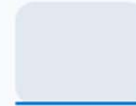
Show sight differences in clinical effectiveness

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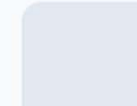
Have similar safety

0%



Meet purity standards

0%



B and C

0%



All of the above



Biologics and Biosimilars Nomenclature

Biologics Nomenclature

- The World Health Organization (WHO) is responsible for and manages the International Nonproprietary Names (INN) drug list
 - Provides a standardized, globally recognized structure for the naming of biologics
 - Reflect unique active ingredient of the drug
 - Typically includes a prefix, sub-stem and an official suffix, designated for that class of drugs

2021 Update: New Naming of Monoclonal Antibodies

- WHO's INN program and the United States Adopted Names (USAN) have revised the nomenclature scheme for monoclonal antibodies
- **The suffix “-mab” has been replaced with four new suffixes (-tug, -bart, -mig,-ment), which further divide the classification of mAbs by their immunoglobulin (Ig) structure**
 - Now names using distinct random prefix, two substems, and a suffix
 - *Prefix* carries no specific meaning, but should be unique
 - *Substem A* relates to the target of the mAb
 - *Substem B* relates to the species used to manufacture the mAb
 - *Suffix* reflects the immunoglobulin structure of the molecule
- Applies to newly named monoclonal antibodies starting in December 2021 (drugs named prior will keep their original name)

New Naming of Monoclonal Antibodies

Prefix	Substem A		Substem B		Suffix	Stem Structure
	Sub-stem	Target class	Sub-stem	Source species		
Random	-b(a)-	bacterial	-a-	Rat	-tug	<u>U</u> nmodified Ig
	-c(i)-	cardiovascular	-e-	Hamster		
	-f(u)-	fungal	-i-	Primate	-bart	Antib <u>o</u> dy <u>a</u> rtificial
	-gr(o)-	skeletal muscle mass related growth factors and receptors	-o-	Mouse		
	-k(i)-	interleukin	-u-	Human		
	-l(i)-	immunomodulating	-xi-	Chimeric	-mig	<u>M</u> ulti <u>I</u> g
	-n(e)-	neural	-xizu-	Chimeric- humanized		
	-tox(a)-	toxin	-zu-	Humanized	-ment	<u>F</u> ragment
	-t(u)-	tumor				
-v(i)-	viral					

**Example:
Nepuvibart**

Naming of Biosimilars

- The FDA provides specific guidance on the naming of biosimilars
- Biologics and biosimilars share a core name with the reference product hyphenated with four unique and meaningless lowercase letters as a distinguishing suffix
- Sharing a core name does NOT automatically infer interchangeability

Examples of Select Biosimilars

Reference Biologic	Select Biosimilar (FDA-approved products)
Adalimumab (Humira®)	Adalimumab-fkjp (Hulio®) Adalimumab-adbm (Cyltezo®)*
Infliximab (Remicade®)	Infliximab-abda (Renflexis®) Infliximab-dyyb (Inflectra®)
Etanercept (Enbrel®)	Etanercept-ykro (Eticovo™) Etanercept-szsz (Erelzi®)
Ranibizumab (Lucentis®)	Ranibizumab-nuna (Byooviz™)
Rituximab (Rituxan®)	Rituximab-arrx (Riabni™)
Filgrastim (Neupogen®)	Filgrastim-ayow (Releuko™) Filgrastim-sndz (Zarxio®)‡

‡ Zarxio® is the first FDA approved biosimilar product

*Cyltezo® is the first FDA approved monoclonal antibody biosimilar to be FDA approved as interchangeable

Purple Book Database of Licensed Biological Products

- Contains information on all FDA-licensed biological products regulated by CDER, including licensed biosimilar and interchangeable products, and their reference products
- Also includes information about FDA-licensed allergenic, cellular, gene therapy, hematologic, and vaccine products regulated by CBER
- The database can be accessed via PurpleBookSearch.fda.gov

Purple Book
Database of Licensed Biological Products

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[FAQs](#)
[Patent List](#)
[Download Purple Book Data](#)

The Purple Book database contains information on all FDA-licensed (approved) biological products regulated by the Center for Drug Evaluation and Research (CDER), including licensed biosimilar and interchangeable products, and their reference products.

The Purple Book also contains information about all FDA-licensed allergenic, cellular and gene therapy, hematologic, and vaccine products regulated by the Center for Biologics Evaluation and Research (CBER).

Enter a product's proprietary (brand) name or the nonproprietary (proper) name to find biological products. As you type, a list of potential results will begin to appear below the search box based on what you are typing. Click on a product from the auto-populated results list below to view the results page. The results page for your selected product will include all biological products that share a core name (i.e., biosimilar, interchangeable, reference, and related biological products).

[Advanced Search](#) Database last updated: April 12, 2023



Test Your Knowledge #4: Pharmacists



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4. (P) Newer monoclonal antibodies manufactured after 2021 will now use the new suffixes. Which of the following is NOT one of the new suffixes?



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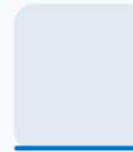
“-tug”

0%



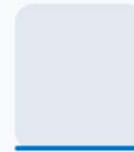
“-bart”

0%



“-cel”

0%



“-mig”

Biosimilar Interchangeability Laws – Illinois



Senate Bill 455 (2015) amended section 5 of the Pharmacy Practice Act

A pharmacist may substitute an interchangeable biosimilar for the biologic product ONLY if:

- The biosimilar has been FDA-approved as interchangeable with the biologic product
- The prescribing physician does not designate orally, in writing, or electronically that substitution is prohibited
- The pharmacy informs the patient of the substitution



Test Your Knowledge #5: Pharmacists & Techs



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5. (P&T) Which of the following should be referenced to determine the interchangeability of a biosimilar with a reference biologic product?



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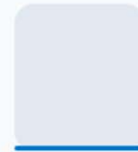
Drug package insert

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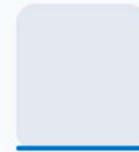
FDA Biologic Database

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Orange Book

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Purple Book

Summary

Biologic therapeutics is a constantly evolving field, providing life-saving therapies to severe disease conditions

Since biologics are made from living organisms, their manufacturing process is complex and costly

Nomenclature for monoclonal antibodies have recently changed from "-mab" to "-tug, -bart, -mig, and -ment"

Biosimilars are an affordable alternative to biologics, but interchangeability is based on the guidance from the FDA

Increased awareness of the interchangeability is important to improve access, reduce cost, and allow clinicians to be better advocates for their patients



Moderate

Visual settings

Edit



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Questions?

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Nobody has responded yet.

Hang tight! Responses are coming in.

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Questions?

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