INDICATIONS/FORMULATIONS DOSING AND ADMINISTRATION CLINICAL PATIENT SUPPORT SERVICES

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# BIMZELX® (BIMEKIZUMAB-BKZX) FREQUENTLY ASKED QUESTIONS (FAQs) FOR SPECIALTY PHARMACIES













# **INDICATIONS AND FORMULATIONS**



#### What is BIMZELX indicated for?

BIMZELX® (bimekizumab-bkzx) is indicated for the

- Treatment of adult patients with **moderate-to-severe plaque psoriasis (PSO)** who are candidates for systemic therapy or phototherapy
- Treatment of adult patients with active psoriatic arthritis (PsA)
- Treatment of adult patients with active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation
- Treatment of adult patients with active ankylosing spondylitis (AS)
- Treatment of adult patients with moderate-to-severe hidradenitis suppurativa (HS)



#### What are the available dosing forms and strengths for BIMZELX?

BIMZELX is available in the following strengths and forms:

#### **BIMZELX Autoinjector BIMZELX Prefilled Syringe** Carton of two (2) Carton of two (2) 160 mg/mL single-dose prefilled 160 mg/mL single-dose autoinjectors syringes Dermatology NDC: 50474-0780-79 **NDC:** 50474-0781-85 Carton of one (1) Carton of one (1) 320 mg/2 mL (160 mg/mL) 320 mg/2 mL (160 mg/mL) single-dose prefilled syringe single-dose autoinjector NDC: 50474-0782-84 NDC: 50474-0783-78 Rheumatology Carton of one (1) Carton of one (1) 160 mg/mL single-dose 160 mg/mL single-dose prefilled autoinjector syringe NDC: 50474-0781-84 NDC: 50474-0780-78

# **DOSING AND ADMINISTRATION**

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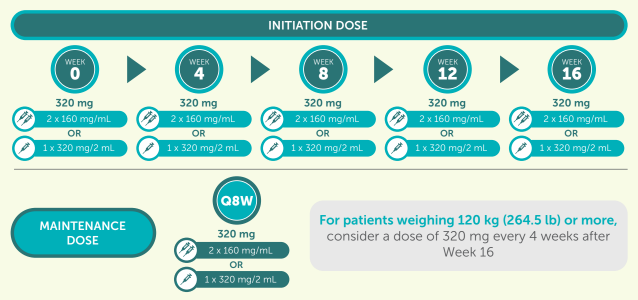


Dermatology

#### What is the dosing regimen for BIMZELX?

#### **Dosing for Patients With Moderate-to-Severe Psoriasis (PSO)**

The recommended dosage of BIMZELX is 320 mg (can be given as either 2 subcutaneous injections of 160 mg each or 1 subcutaneous injection of 320 mg) at Weeks 0, 4, 8, 12, and 16, then every 8 weeks thereafter.



## Dosing for Patients With Moderate-to-Severe Hidradenitis Suppurativa (HS)

The recommended dosage of BIMZELX is 320 mg (can be given as either 2 subcutaneous injections of 160 mg each or 1 subcutaneous injection of 320 mg) at Weeks 0, 2, 4, 6, 8, 10, 12, 14, and 16, then every 4 weeks thereafter.





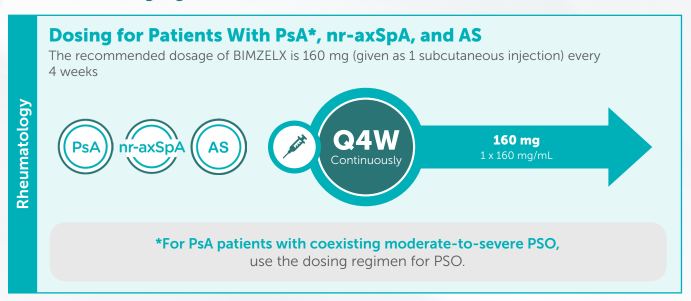
# **DOSING AND ADMINISTRATION**

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What is the dosing regimen for BIMZELX (continued)?





## How should patients be instructed to inject BIMZELX?

BIMZELX is administered under the skin (subcutaneously) by autoinjector or prefilled syringe. BIMZELX is intended for use under the guidance and supervision of a healthcare professional. Patients may self-inject after training in subcutaneous injection technique.

BIMZELX can be self-administered in the thighs or abdomen, or administered by an HCP or caregiver in the back of the upper arm. BIMZELX should not be injected into areas where the skin is tender, bruised, red, hard, thick, scaly, or affected by psoriasis. Choose a different site on the body for each injection. Do not use the same injection site two times in a row.



Instruct patients and/or caregivers to review injection training resources at https://www.bimzelx.com/about-bimzelx/dosing

For full injection instructions, refer patients to the Instructions for Use.



It's important to instruct patients switching from the 1 mL autoinjector to the 2 mL autoinjector that there are differences in the administration instructions and to carefully read the Instructions for Use for the 2 mL autoinjector.



# **DOSING AND ADMINISTRATION (CONTINUED)**





#### How should BIMZELX be stored?



- Cartons should be refrigerated between 2°C and 8°C (36°F and 46°F). When necessary, BIMZELX prefilled syringes or autoinjectors may be stored at room temperature up to 25°C (77°F) in the original carton for a single period of up to 30 days. Once BIMZELX prefilled syringes or autoinjectors have been stored at room temperature, do not place back in refrigerator.
- Keep in original carton to protect from light. Do not freeze. Do not use beyond expiration date.



#### What is the shelf life for BIMZELX?



• The shelf life of BIMZELX is 36 months.



#### Is BIMZELX available for me to order?

BIMZELX is operating with an open distribution model. Any Specialty Pharmacy can order and dispense BIMZELX.



Click or scan to download the **BIMZELX Wholesale Order Entry Form** 



# **CLINICAL**



# When were the BIMZELX indications approved?

UCB announced that BIMZELX was approved for the treatment of moderate-to-severe HS on November 20, 2024. The BIMZELX rheumatology (PsA, nr-axSpA, AS) indications were approved on September 21, 2024. The BIMZELX moderate-to-severe PSO indication was approved in October 2023.



# Are there drug interactions with BIMZELX?

Avoid use of live vaccines in patients treated with BIMZELX. Patients treated with BIMZELX may receive inactivated or non-live vaccinations.

- Healthy individuals who received a single dose of 320 mg of BIMZELX 2 weeks before receiving vaccination with an inactivated seasonal influenza vaccine had similar antibody responses as compared to individuals who did not receive BIMZELX before vaccination.
- The effectiveness of inactivated seasonal influenza vaccines and other inactivated and non-live vaccines has not been evaluated in patients treated with BIMZELX.



# What is the mechanism of action of BIMZELX?

IL-17A and IL-17F are naturally occurring cytokines that can **drive chronic inflammation and damage across multiple tissues**.

- BIMZELX selectively inhibits both IL-17F and IL-17A and provides more inhibition of inflammation.
- Please see Section 12.1 (Mechanism of Action) in full Prescribing Information at BIMZELX.com for more information.



# What are the most common side effects of BIMZELX?

Most common (≥ 1%) adverse reactions in plaque psoriasis and hidradenitis suppurativa include upper respiratory infections, oral candidiasis, headache, injection site reactions, tinea infections, gastroenteritis, herpes simplex infections, acne, folliculitis, other candida infections, and fatique.

Most common (≥ 2%) adverse reactions in psoriatic arthritis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, and urinary tract infections.

Most common (≥ 2%) adverse reactions in non-radiographic axial spondyloarthritis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, cough, fatigue, musculoskeletal pain, myalgia, tonsillitis, transaminase increase, and urinary tract infections.

Most common (≥ 2%) adverse reactions in ankylosing spondylitis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, injection site pain, rash, and vulvovaginal mycotic infection.



#### What are the warnings related to BIMZELX?

BIMZELX® (bimekizumab-bkzx) may increase the risk of suicidal ideation and behavior (SI/B). A causal association between treatment with BIMZELX and increased risk of SI/B has not been definitively established. Prescribers should weigh the potential risks and benefits before using BIMZELX in patients with a history of severe depression or SI/B. Advise monitoring for the emergence or worsening of depression, suicidal ideation, or other mood changes. If such changes occur, instruct to promptly seek medical attention, refer to a mental health professional as appropriate, and re-evaluate the risks and benefits of continuing treatment.

BIMZELX may increase the risk of infections, including serious infections. Do not initiate treatment with BIMZELX in patients with any clinically important active infection until the infection resolves or is adequately treated. In patients with a chronic infection or a history of recurrent infection, consider the risks and benefits prior to prescribing BIMZELX. Instruct patients to seek medical advice if signs or symptoms suggestive of clinically important infection occur. If a patient develops such an infection or is not responding to standard therapy, monitor the patient closely and do not administer BIMZELX until the infection resolves.

Evaluate patients for tuberculosis (TB) infection prior to initiating treatment with BIMZELX. Avoid the use of BIMZELX in patients with active TB infection. Consider anti-TB therapy prior to initiation of BIMZELX in patients with a past history of latent or active TB in whom an adequate course of treatment cannot be confirmed. Closely monitor patients receiving BIMZELX for signs and symptoms of active TB during and after treatment.

Elevated serum transaminases were reported in clinical trials. Test liver enzymes, alkaline phosphatase, and bilirubin at baseline, periodically during treatment, and according to routine patient management. If treatment-related increases in liver enzymes occur and drug-induced liver injury is suspected, interrupt BIMZELX until a diagnosis of liver injury is excluded. Permanently discontinue use of BIMZELX in patients with causally-associated combined elevations of transaminases and bilirubin. Avoid the use of BIMZELX in patients with acute liver disease or cirrhosis.

Cases of inflammatory bowel disease (IBD) have been reported in patients treated with IL-17 inhibitors, including BIMZELX. Avoid use of BIMZELX in patients with active IBD. During BIMZELX treatment, monitor patients for signs and symptoms of IBD and discontinue treatment if new onset or worsening of signs and symptoms occurs.



# **PATIENT SUPPORT SERVICES**





#### What is BIMZELX Navigate<sup>®</sup>?

BIMZELX Navigate provides access, affordability, and treatment support for eligible patients.





## What type of patient access and treatment support services are available?\*

BIMZELX Navigate offers a wide range of resources—from quick product access for eligible patients to personalized onboarding support—to get patients on treatment as smoothly as possible.



#### BIMZELX Navigate Bridge<sup>1,‡</sup>

Eligible, commercially insured patients whose prescription is initially denied or delayed by insurance may be eligible to receive BIMZELX at \$15 per dose for up to 2 years or until insurance coverage is approved, whichever comes first.



#### **BIMZELX Navigate Savings<sup>®</sup>**

Once insurance approval is obtained, commercially insured patients may be eligible to receive BIMZELX for as little as \$5 per dose.



<sup>\*</sup>Please see full eligibility and terms at www.bimzelx.com/patient-support/navigate-benefits.

To facilitate enrollment into BIMZELX Bridge, two (2) prescriptions are requested when prescribing to non-Bridge dispensing Specialty Pharmacy; alternatively submit BIMZELX Patient Enrollment Form.

For eligible, commercially insured patients only. Eligible, commercially insured patients whose prescription is initially denied or delayed by insurance may be eligible to receive BIMZELX at \$15 per dose for up to 2 years or until insurance coverage is approved, whichever comes first. Please see full eligibility and terms at <a href="https://www.bimzelx.com/patient-support/navigate-benefits">www.bimzelx.com/patient-support/navigate-benefits</a>.

Eligible, commercially insured patients may pay as little as \$5 per dose. Please see full eligibility requirements and terms at www.bimzelx.com/patient-support/navigate-benefits.

# **PATIENT SUPPORT SERVICES (CONTINUED)**

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# How can eligible, commercially insured patients enroll in BIMZELX Navigate Bridge?

If the healthcare provider sends the prescription directly to BIMZELX Navigate (Hub), they will automatically enroll eligible patients into BIMZELX Navigate Bridge. Select Specialty Pharmacies within the enhanced network are contracted to dispense BIMZELX on behalf of the BIMZELX Navigate Bridge program directly to patients.

Non-Bridge Dispensing Specialty Pharmacies can facilitate enrollment into BIMZELX Navigate Bridge using one of the following options:



Submit to BIMZELX Navigate:

Transfer prescription to the BIMZELX Hub Partner (Phone: 412-250-4407; Fax: 412-774-9652)



Specialty Pharmacy Rx Transfer:
 Transfer prescription to the Specialty Pharmacy
 Bridge-Dispensing Network\*

#### **Bimzelx Navigate Bridge-Dispensing Specialty Pharmacies\***

| Specialty Pharmacy <sup>†</sup> | Phone Number   | Fax Number     |
|---------------------------------|----------------|----------------|
| altScripts Specialty Pharmacy   | 1-414-385-9500 | 1-414-385-7200 |
| Amber Specialty Pharmacy        | 1-888-370-1724 | 1-402-896-3774 |
| Ardon Health Specialty Pharmacy | 1-855-425-4085 | 1-855-425-4096 |
| BioPlus Specialty Pharmacy      | 1-888-292-0744 | 1-800-269-5493 |
| Encore Pharmacy                 | 1-817-335-5712 | 1-866-326-9731 |
| Meijer Specialty Pharmacy       | 1-855-263-4537 | 1-734-391-2365 |
| Polaris Specialty Rx            | 1-855-797-0857 | 1-800-781-3364 |
| Publix Specialty Pharmacy       | 1-855-797-8254 | 1-863-413-5723 |
| Senderra Specialty Pharmacy     | 1-855-460-7928 | 1-888-777-5645 |
| Sterling Specialty Pharmacy     | 1-888-618-4126 | 1-866-588-0371 |

<sup>\*</sup>Any specialty pharmacy can order/dispense BIMZELX.



<sup>&</sup>lt;sup>†</sup>These pharmacies are subject to change.

# **PATIENT SUPPORT SERVICES (CONTINUED)**

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## How can eligible, commercially insured patients enroll in BIMZELX **Navigate Savings?**

There are a few ways for eligible, commercially insured patients to receive BIMZELX Navigate Savings:



If the healthcare provider (HCP) sends the prescription directly to BIMZELX Navigate (Hub), they will automatically enroll eligible patients into the BIMZELX Navigate Savings Program



All patients may self-enroll at **BIMZELX.com** 



Specialty Pharmacies can enroll eligible patients into **BIMZELX Navigate Savings by** visiting **BIMZELXhcp.com** or **UCBSavings.com** 

Certain restrictions apply. Please see full Eligibility Criteria on page 9.



### What are the eligibility qualifications for BIMZELX Navigate Bridge?

• For initial enrollment into the program, eligible patients must be experiencing a delay in, or have been denied, coverage for BIMZELX by their commercial insurance plan. To maintain eligibility in the program, the following are required: (1) a prior authorization request has been submitted and/or coverage remains unavailable for the patient; and (2) if the prior authorization is denied by the payer, the prescriber must submit an appeal within the first sixty (60) days of the prior authorization denial and a prior authorization request (or documentation as may otherwise be required by the payer) must be submitted every six (6) months thereafter.

Certain restrictions apply. Please see full Eligibility Criteria on page 9. This offer is not available for patients eligible for Medicare, Medicaid, or any other form of government insurance coverage. Uninsured patients may be eligible for the UCB Patient Assistance Program.



# What are the enrollment qualifications for patient assistance programs (PAPs)?

A PAP may be available to patients with limited or no health insurance coverage who meet specific eligibility requirements. To be eligible for a PAP, patients must:

- Have a valid prescription from a US HCP for on-label indication
- Reside within the United States, including the District of Columbia or US Territory
- Have a total household income that does not exceed 500% of the federal poverty limit (see https://www.healthcare.gov/glossary/federal-poverty-level-FPL)

Contact UCBCares® by visiting askucbcares.com or calling 1-844-599-CARE (2273) for more information.



# Is patient service mandated through BIMZELX Navigate?

No, HCPs may also submit the prescription to their SP of choice.



For additional information regarding patient support services, visit BIMZELXhcp.com or call BIMZELX Navigate at 1-866-4-BIMZELX (1-866-424-6935).





Full Terms and Conditions for BIMZELX Navigate® Bridge
BIMZELX Navigate Bridge (the "Program") provides BIMZELX® (bimekizumab-bkzx) to eligible patients for \$15 per dose for up to two (2) years or until the patient's commercial insurance plan approves coverage for the drug, whichever comes first. Eligible patients must be 18 years of age or older with commercial insurance and a valid prescription consistent with FDA-approved product labeling. For initial enrollment into the Program, the patient must be experiencing a delay in, or have been denied, coverage for BIMZELX by their commercial insurance plan. To maintain eligibility in the Program, the following are required: (1) a prior authorization request has been submitted and/or coverage remains unavailable for the patient; and (2) if the prior authorization is denied by the payer, the prescriber must submit an appeal within the first sixty (60) days of the prior authorization denial and a prior authorization request (or documentation as may otherwise be required by the payer) must be submitted every six (6) months thereafter. Program is not available (1) to patients whose prescriptions are reimbursed, in whole or in part, under Medicare (including Medicare Part D), Medicaid, or any other federal- or state-funded healthcare programs (including any state prescription drug assistance programs and the Government Health Insurance Plan available in Puerto Rico), (2) where a patient's insurance covers the drug, (3) to uninsured or cash-paying patients, or (4) where otherwise prohibited by law. Product shall be dispensed pursuant to Program rules and federal and state laws. Patients may be asked to re-verify insurance coverage status during participation in the Program. No purchase necessary. Program is not health insurance, nor is participation a guarantee of insurance coverage. Limitations may apply. This Program cannot be combined with any other savings, free trial, or similar offer for the specified prescription. The patient, or healthcare provider on the patient's behalf, must not submit any claim for reimbursement for product provided under this Program to any third-party payer. UCB, Inc. reserves the right to end or amend this Program without notice

Full Terms and Conditions for BIMZELX Navigate® Savings

BIMZELX Navigate Savings (the "Program") provides BIMZELX® (bimekizumab-bkzx) to eligible patients with commercial insurance coverage for as little as \$5 per dose. Eligible patients must be 18 years of age or older with commercial insurance coverage with a valid prescription consistent with FDA-approved product labeling. The Program is not available for (1) for prescriptions that are reimbursed, in whole or in part, under Medicare (including Medicare Part D), Medicaid, or any other federal- or state-funded healthcare programs (including any state prescription drug assistance programs and the Government Health Insurance Plan available in Puerto Rico), (2) where a patient's commercial insurance plan reimburses for the entire cost of the drug, (3) for uninsured or cash-paying patients, or (4) where otherwise prohibited by law. Product shall be dispensed pursuant to Program rules and federal and state laws. The value of the Program is exclusively for the benefit of patients and is intended to be credited in full towards patient out-of-pocket obligations and maximums, including applicable copayments, coinsurance, and deductibles. Patient may not seek reimbursement for the value of this Program from other parties, including third-party payers (i.e., any health insurance program or plan, or public payers (i.e., and bol). Patient is responsible for complying with any applicable limitations and requirements of their health plan related to the use of the Program. This Program cannot be combined with any other savings, free trial, or similar offer for the specified prescription. UCB, Inc. reserves the right to amend or end this Program at any time without notice. Subject to the prior sentence, this Program expires at 11:59 p.m. on December 31. Patients that meet the above requirements may re-enroll in the Program each year.



For additional information, contact UCBCares® at 1-844-599-CARE (2273).

#### IMPORTANT SAFETY INFORMATION

#### Suicidal Ideation and Behavior

BIMZELX® (bimekizumab-bkzx) may increase the risk of suicidal ideation and behavior (SI/B). A causal association between treatment with BIMZELX and increased risk of SI/B has not been definitively established. Prescribers should weigh the potential risks and benefits before using BIMZELX in patients with a history of severe depression or SI/B. Advise monitoring for the emergence or worsening of depression, suicidal ideation, or other mood changes. If such changes occur, instruct to promptly seek medical attention, refer to a mental health professional as appropriate, and re-evaluate the risks and benefits of continuing treatment.

#### **Infections**

BIMZELX may increase the risk of infections, including serious infections. Do not initiate treatment with BIMZELX in patients with any clinically important active infection until the infection resolves or is adequately treated. In patients with a chronic infection or a history of recurrent infection, consider the risks and benefits prior to prescribing BIMZELX. Instruct patients to seek medical advice if signs or symptoms suggestive of clinically important infection occur. If a patient develops such an infection or is not responding to standard therapy, monitor the patient closely and do not administer BIMZELX until the infection resolves.

#### **Tuberculosis**

Evaluate patients for tuberculosis (TB) infection prior to initiating treatment with BIMZELX. Avoid the use of BIMZELX in patients with active TB infection. Initiate treatment of latent TB prior to administering BIMZELX. Consider anti-TB therapy prior to initiation of BIMZELX in patients with a past history of latent or active TB in whom an adequate course of treatment cannot be confirmed. Closely monitor patients for signs and symptoms of active TB during and after treatment.

#### **Liver Biochemical Abnormalities**

Elevated serum transaminases were reported in clinical trials with BIMZELX. Test liver enzymes, alkaline phosphatase, and bilirubin at baseline, periodically during treatment with BIMZELX, and according to routine patient management. If treatment-related increases in liver enzymes occur and drug-induced liver injury is suspected, interrupt BIMZELX until a diagnosis of liver injury is excluded. Permanently discontinue use of BIMZELX in patients with causally associated combined elevations of transaminases and bilirubin. Avoid use of BIMZELX in patients with acute liver disease or cirrhosis.

#### **Inflammatory Bowel Disease**

Cases of inflammatory bowel disease (IBD) have been reported in patients treated with IL-17 inhibitors, including BIMZELX. Avoid use of BIMZELX in patients with active IBD. During BIMZELX treatment, monitor patients for signs and symptoms of IBD and discontinue treatment if new onset or worsening of signs and symptoms occurs.

#### **Immunizations**

Prior to initiating therapy with BIMZELX, complete all age-appropriate vaccinations according to current immunization guidelines. Avoid the use of live vaccines in patients treated with BIMZELX.

#### MOST COMMON ADVERSE REACTIONS

Most common (≥ 1%) adverse reactions in plaque psoriasis and hidradenitis suppurativa include upper respiratory tract infections, oral candidiasis, headache, injection site reactions, tinea infections, gastroenteritis, herpes simplex infections, acne, folliculitis, other candida infections, and fatigue.

Most common (≥ 2%) adverse reactions in psoriatic arthritis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, and urinary tract infections.

Most common (≥ 2%) adverse reactions in non-radiographic axial spondyloarthritis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, cough, fatigue, musculoskeletal pain, myalgia, tonsillitis, transaminase increase, and urinary tract infections.

Most common (> 2%) adverse reactions in ankylosing spondylitis include upper respiratory tract infections, oral candidiasis, headache, diarrhea, injection site pain, rash, and vulvovaginal mycotic infection.

Please refer to the full Prescribing Information provided by the UCB representative, and visit BIMZELXhcp.com. For more information on BIMZELX, contact UCBCares® at 1-844-599-CARE (2273).



Bimzelx® (bimekizumab-bkzx)