ICD-10 diagnostic codes consistent with the ORBACTIV® (oritavancin) Indication

ICD-10-CM Diagnosis Codes

Diagnosis



Staphylococcus	
A49.01	Methicillin susceptible staphylococcus aureus, unspecified site
A49.02	Methicillin resistant staphylococcus aureus infection, unspecified site
B95.62	Staphylococcus aureus as the cause of diseases classified elsewhere (methicillin resistant)
B95.61	Staphylococcus aureus as the cause of diseases classified elsewhere (methicillin susceptible)
B95.8	Unspecified staphylococcus as the cause of diseases classified elsewhere

Streptococcus	
A49.1	Streptococcus infection, unspecified site
B95.0-B95.2, B95.4	Streptococcus, as the cause of disease classified elsewhere

Other infections	
A46	Erysipelas
L08.0-L08.1, L08.81-L08.89, L08.9	Other local infections of skin and subcutaneous tissue

Cellulitis	
L03.211	Cellulitis of face
K12.2	Cellulitis and abscess of mouth
H05.011-H05.019	Cellulitis of orbit, abscess of orbit
H60.10-H60.13	Cellulitis of external ear
J34.0	Cellulitis and abscess of external nose
L03.221	Cellulitis of neck
L03.113-L03.114	Cellulitis of upper limb
L03.111-L03.114	Cellulitis of axilla and upper limb
L03.011-L03.019	Cellulitis of finger
N61	Inflammatory disorders of breast (includes cellulitis/abscess breast)
L03.311-L03.316, L03.319	Cellulitis of trunk
L03.317	Cellulitis of buttock
L03.119	Cellulitis of unspecified part of limb
L03.115-L03.116	Cellulitis of lower limb
N48.22	Cellulitis of corpus cavernosum and penis
L03.031-L03.039	Cellulitis of toe
L03.811-L03.818	Cellulitis of other sites
L03.90	Cellulitis, unspecified

INDICATION AND USAGE

ORBACTIV® (oritavancin) for injection is indicated for the treatment of adult patients with acute bacterial skin and skin structure infections (ABSSSI) caused or suspected to be caused by susceptible isolates of the following gram-positive microorganisms: Staphylococcus aureus (including methicillin-susceptible [MSSA] and -resistant [MRSA] isolates), Streptococcus pyogenes, Streptococcus agalactiae, Streptococcus dysgalactiae, Streptococcus anginosus group (includes S. anginosus, S. intermedius, and S. constellatus), and Enterococcus faecalis (vancomycin-susceptible isolates only).

To reduce the development of drug-resistant bacteria and maintain the effectiveness of ORBACTIV® and other antibacterial drugs, ORBACTIV® should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptible bacteria.

Please see reverse for Important Safety Information.



ICD-10 diagnostic codes (continued)

Abscess

L02.01 Cutaneous abscess of face H00.031–H00.039 Abscess and furuncle of eyelid

H60.00–H60.03 Abscess of external earK12.2 Submandibular abscessL02.11 Cutaneous abscess of neck

L02.411–L02.414 Cutaneous abscess of axilla and upper limb

L02.511–L02.519 Cutaneous abscess of hand
L02.211–L02.219 Cutaneous abscess of trunk
L02.31 Cutaneous abscess of buttock

L02.419 Cutaneous abscess of limb, unspecified

L02.415–L02.416 Cutaneous abscess of lower limb
L02.611–L02.619 Cutaneous abscess of foot

N48.21 Abscess of corpus cavernosum and penis

N76.4 Abscess of vulva

K61.0–K61.4 Abscess of anal and rectal regions
L02.811–L02.818 Cutaneous abscess of other sites
L02.91 Cutaneous abscess, unspecified

Furuncle

L02.02 Furuncle of face L02.12 Furuncle of neck

L02.421–L02.424 Furuncle of axilla, upper limb

L02.521–L02.529 Furuncle of hand
L02.221–L02.229 Furuncle of trunk
L02.32 Furuncle of buttock

L02.429 Furuncle of limb, unspecified

L02.425–L02.426 Furuncle of lower limb
L02.621–L02.629 Furuncle of other sites

L02.92 Furuncle, unspecified

Carbuncle

L02.03 Carbuncle of face

J34.0 Carbuncle and furuncle of external nose

L02.13 Carbuncle of neck

L02.431–L02.434 Carbuncle of axilla, upper limb

L02.531–L02.539 Carbuncle of hand L02.231–L02.239 Carbuncle of trunk L02.33 Carbuncle of buttock

L02.439 Carbuncle of limb, unspecified

L02.435–L02.436 Carbuncle of lower limb
L02.631–L02.639 Carbuncle of foot
L02.831–L02.838 Carbuncle of other sites
L02.93 Carbuncle, unspecified

This resource identifies diagnosis codes that are likely to be most relevant to healthcare provider claims for the administration of ORBACTIV®. Billing and coding information is illustrative and is not intended to assist providers in obtaining reimbursement for any specific claim.

Healthcare providers are responsible for selecting appropriate codes for in the submission of claims consistently with health plan requirements and applicable standards of care. Actual clinical diagnosis and coding should be done to the highest level of specificity based on the patient's condition and the items and services that are actually furnished.

IMPORTANT SAFETY INFORMATION

Contraindications

Use of intravenous unfractionated heparin sodium is contraindicated for 120 hours (5 days) after ORBACTIV® administration because the activated partial thromboplastin time (aPTT) test results may remain falsely elevated for approximately 120 hours (5 days) after ORBACTIV® administration.

ORBACTIV® is contraindicated in patients with known hypersensitivity to oritavancin products.

Warnings and Precautions

Coagulation test interference: Oritavancin has been shown to artificially prolong aPTT for up to 120 hours and may prolong PT and INR for up to 12 hours, and ACT for up to 24 hours. Oritavancin has also been shown to elevate D-dimer concentrations up to 72 hours. For patients who require aPTT monitoring within 120 hours of oritavancin dosing, consider a non-phospholipid dependent coagulation test such as a Factor Xa (chromogenic) assay or an alternative anticoagulant not requiring aPTT.

Serious hypersensitivity reactions, including anaphylaxis, have been reported with the use of oritavancin products, including ORBACTIV®. Discontinue infusion if signs of acute hypersensitivity occur. Closely monitor patients with known hypersensitivity to glycopeptides.

Infusion Related Reactions: Administer ORBACTIV® over 3 hours to minimize infusion-related reactions. Infusion reactions characterized by chest pain, back pain, chills and tremor have been observed with the use of oritavancin products (e.g. ORBACTIV®), including after the administration of more than one dose of oritavancin during a single course of therapy. Stopping or slowing the infusion may result in cessation of these reactions.

Clostridioides difficile-associated diarrhea: Evaluate patients if diarrhea occurs.

Concomitant warfarin use: Oritavancin has been shown to artificially prolong PT/INR for up to 12 hours. Patients should be monitored for bleeding if concomitantly receiving ORBACTIV® and warfarin.

Osteomyelitis: Institute appropriate alternate antibacterial therapy in patients with confirmed or suspected osteomyelitis.

Prescribing ORBACTIV® in the absence of a proven or strongly suspected bacterial infection or a prophylactic indication is unlikely to provide benefit to the patient and increases the risk of development of drug-resistant bacteria.

Adverse Reactions

The most common adverse reactions (≥3%) in patients treated with ORBACTIV® were headache, nausea, vomiting, limb and subcutaneous abscesses, and diarrhea.

Please see accompanying Full Prescribing Information.



