

# Bladder Cancer (Urothelial) Pathways

Patient Name: \_\_\_\_\_

Date of Birth: \_\_\_\_\_

Member Number: \_\_\_\_\_

Treatment Start Date: \_\_\_\_\_

**Pathology:** \_\_\_\_\_

**Stage:** \_\_\_\_\_

**Line of Therapy:** \_\_Neoadjuvant/Pre-Op \_\_Adjuvant/Post-Op  
\_\_1<sup>st</sup> Line \_\_2<sup>nd</sup> Line \_\_3<sup>rd</sup> Line \_\_3<sup>rd</sup> Line+ \_\_Maint

**ECOG Performance Status:** \_\_\_\_\_ **ICD-10 Code:** \_\_\_\_\_

**Goal of Treatment:** \_\_Curative \_\_Non-Curative

**Biomarkers/Characteristics:** (select all that apply)

Microsatellite instability: \_\_dMMR/MSI-H \_\_MSI-L \_\_Not reported

Platinum Resistant/Refractory? \_\_ Yes \_\_ No \_\_ Not reported

NTRK Fusion: \_\_Positive \_\_Negative \_\_Not reported

## Neoadjuvant Therapy | Clinical Stage II, III, or IV Without Evidence of Metastases (cT2, cT3, cT4a, cT4b, M0)

**CMV:** cisplatin, methotrexate, and vinblastine 3 cycles

Gemcitabine (Gemzar) and cisplatin 4 cycles

## Adjuvant Therapy | Stage 0 (Ta, Tis) or Stage I | After TURBT\* or Following Resection of Recurrent or Persistent Disease

**BCG:** bacillus calmette-guerin, intravesical

Gemcitabine (Gemzar), intravesical (**low-grade histology only**)

## Metastatic Disease | First Line of Therapy (1st Line)

Gemcitabine (Gemzar) and cisplatin†

## Metastatic Disease | Second Line of Therapy (2nd Line)

Gemcitabine (Gemzar)

Paclitaxel

Pembrolizumab (Keytruda)‡

\* TURBT: Transurethral resection of bladder tumor

† In the setting of recurrent/metastatic disease, a substitution of carboplatin for cisplatin will be considered a pathway option

‡ Administered at a dose of 200 mg every 3 weeks per the FDA label OR 2 mg/kg (up to a maximum of 200 mg) every 3 weeks, as clinically appropriate

**Note: Pathways are independent of specific health plan medical policy coverage criteria. Health plan medical policy/clinical guidelines should be consulted to determine whether proposed services will be covered. Biosimilars of reference products listed are considered "on pathway." However, reimbursement for biosimilar products may be impacted by health plan specific formularies, medical policy and preferred product rules.**



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