Nivolumab + ipilimumab

CheckMate142



Nivolumab + ipilimumab CheckMate142 Nivolumab + ipilimumab CheckMate142 PRFLIMINARY SCORE SCORE CURATIVE CURATIVE Overall Survival / Disease-Free Survival / Pathological Complete Response NON-CURATIVE NON-CURATIVE ORR **Overall Survival ADJUSTMENTS** Quality of life Progression-Free Survival Not qualified for an ESMO-MCBS credit Non-inferiority (Improved Quality of Life or Reduced Adverse Events) / Response Rate Serious and disabling adverse effects 3 Overall Response Rate / Duration of Response Overall Survival / Disease-Free Survival / Pathological Complete Response Other adjustments INFORMATION Tumour type: Gastrointestinal Cancers Therapeutic Indication: FDA: Treatment of adult patients with dMMR or MSI-H mCRC after prior fluoropyrimidine-based combination ChT. EMA: Nivolumab in combination with ipilimumab for the treatment adult patients with dMMR or MSI-H colorectal cancer after prior fluoropyrimidine based combination chemotherapy Experimental Arm: Nivolumab + ipilimumab Control Arm: Single arm



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