



مركز الكويت لمكافحة السرطان
Kuwait Cancer Control Center

DENOSumab (Prolia®), every 6 months



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Ministry of Health

Name:

File #:

Ht (cm):

Nationality:

Civil ID:

Wt (Kg):

Gender/Age:

DOB:

BSA (m²):

Indication(s): Androgen deprivation-induced bone loss in men with prostate cancer.
 Armotase inhibitor-induced bone loss In women With breast cancer.
 Postmenopausal osteoporosis.

Central line: Available NA

Allergies: NKA Yes, specify; _____

Parameters: Firstly, Dental clearance. CrCl > 45 ml/min

Standard Protocol:

DRUG	DOSE	ADMINISTRATION	DAYS
DENOSumab (Prolia®)	60 mg	Subcutaneous injection	D1
To be repeated every 6 months until intolerable toxicity.			

Special instructions: Prior To administration, keep the ampule in room temperature in its original container for ~15 to 30 minutes; Do not warm by any other method.

Treatment Description:

Cycle	Date	Prolia®	Physician	Consultant
C# __				
C# __				
C# __				
C# __				
C# __				
C# __				

Important Notes:

Reported grade 3/4 toxicities: None Hematological Non-Hematological

If yes; Did it indicate hospitalization? Yes No

Did it indicate chemo-delay for ≥ 7 days? Yes No

Did it indicate dose reduction? Yes No

Did it indicate G-CSF support? Yes No