DENOsumab (Prolia®), every 6 months







Printed: 13/May/2020

Name: Nationali Gender/A	-		File #: Civil ID: DOB:	Ht (cm): Wt (Kg): 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				
Parameters: Firstly, Dental clearance. CrCl > 45 ml/min				
Standard Protocol:				
DRUG		DOSE	ADMINISTRATION	DAYS
DENOsumab (Prolia@		®) 60 mg	Subcutaneous injection	n 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#				
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				