

Change From Baseline in Patient Reported Outcomes Measurements Information Systems (PROMIS) Pain Intensity 3a Score at Week 52 in Arm A: Prophylaxis

Statistical analysis title	Arm A: Prophylaxis
Statistical analysis description	Testing according to hierarchical testing procedure (only performed if previous endpoint [Change From Baseline in Haem-A-QOL Physical Health Score at Week 52: Prophylaxis Arm] was statistically significant for considered dosing regimen). Least square (LS) mean difference, standard error and 95% confidence interval were estimated by mixed-effect model with repeated measures (MMRM) using visit as fixed effect and Baseline PROMIS Pain Intensity 3a as covariate.
Comparison groups	Arm A: Prophylaxis - Baseline data versus Week 52 data
Number of subjects included in analysis	119
Analysis specification	Pre-specified
Analysis type	Superiority
Analysis type comment	
P-value	0.0042
Method	Unstructured covariance matrix
P-value comment	An unstructured covariance matrix within a subject was used.
Parameter type	LS mean difference
Point estimate	-1.94
Confidence interval	
level	95%
sides	2-sided
Lower limit	-3.26
Upper limit	-0.63
Variability estimate	Standard error of the mean
Dispersion value	0.67