

PHARMOL 10

Complies with the following specifications:

- US FDA 21 CFR 172.878 & 21 CFR 178.3620(a), White Mineral Oil
- USP 40/ NF35 (US Pharmacopoeia/ National Formulary), Light Mineral Oil

S. No.	Characteristic	Unit	Test Method	Typical Data	
				Min.	Max.
1	Appearance	-	Visual	A Colourless, transparent, oily liquid free from fluorescence in day light. Practically insoluble in water, slightly soluble in ethanol (96%), miscible with hydrocarbons.	
2	Colour, Saybolt	-	ASTM D 156	+30	
3	Odour	-	Olfactory	Almost odourless	
4	Kinematic Viscosity at 40 °C	cSt	ASTM D 445/D 7042	9.0	13.0
5	Relative Density at 20°C	-	BP/EP	0.810	0.875
6	Specific Gravity at 25°C	-	NF/ USP	0.815	0.880
7	Flash Point	°C	ASTM D 92	150	
8	Pour Point	°C	ASTM D 97		-15
9	Acidity	-	USP	Not more than 1 ml of 0.01N NaOH	
10	Limit of Polycyclic Aromatic Hydrocarbons	-	NF/USP/BP/ EuP/IP	Pass	
11	Readily Carbonisable Substances	-	NF/USP/BP/ EuP/IP	Pass	
12	Solid Paraffin	-	NF/USP/BP/ EuP/IP	Pass	
13	Sulphur Compounds	-	NF/USP/IP	Pass	

Pharmol 10 – Light Mineral Oil – NF/USP is a highly pure grade of White Mineral Oil specially formulated from chosen severely hydrotreated and highly refined Paraffinic Oils, thus qualifying for the severe requirements stipulated under NF/ United States Pharmacopoeia. The absence of Sulphur compounds, metal impurities and aromatic compounds render these oils very suitable for use in formulation of cosmetics and personal care products.

Disclaimer: Manufacturer makes no warranties, representation or conditions of any kind, expressed or implied, for use with respect to these products. Final determination of suitability of these products for the application contemplated by the user is solely the user's responsibility.