

DEVICE DETAILS				
NAME OF DEVICE	RESTYLANE SKINBOOSTERS VITAL LIGHT LIDOCAINE			
ESTABLISHMENT NAME	GALDERMA MALAYSIA SDN. BHD.			
ROLE OF ESTABLISHMENT	AUTHORIZED REPRESENTATIVE			
REGISTRATION NO	GD38162959818			
BRAND NAME	RESTYLANE			
DEVICE GROUPING TYPE	FAMILY			
DEVICE DESCRIPTION	RESTYLANE SKINBOOSTERS VITAL LIGHT LIDOCAINE IS A STERILE, TRANSPARENT GEL OF STABILIZED HYALURONIC ACID OF NON-ANIMAL ORIGIN WITH THE ADDITION OF 0.3% LIDOCAINE HYDROCHLORIDE. IT IS SUPPLIED IN A GLASS SYRINGE. THE PRODUCT HAS A BUILT IN DOSE-GUIDE, SMART CLICK SYSTEM, WHICH WHEN ACTIVATED CREATES A CLICKING SOUND TO INDICATE EACH INJECTED DOSE. THE 1 ML SYRINGE GIVES PPROXIMATELY 100 DOSES. THE CONTENTS OF THE SYRINGE ARE STERILIZED USING MOIST HEAT. THE PRODUCT IS FOR SINGLE USE ONLY. DISPOSABLE 29G TW (THINWALLED) NEEDLES, STERILIZED USING ETHYLENE OXIDE, ARE PROVIDED. TO ENSURE TRACEABILITY THE PATIENT RECORD LABEL (PART OF SYRINGE LABEL) SHOULD BE ATTACHED TO PATIENT RECORDS.			
DEVICE INTENDED PURPOSE	THIS PRODUCT IS INTENDED TO RESTORE SKIN HYDROBALANCE, IMPROVE SKIN STRUCTURE AND THE ELASTICITY OF THE SKIN. IT SHOULD BE INJECTED IN THE DERMAL LAYER OF THE SKIN. THE ADDITION OF LIDOCAINE PROVIDES INCREASED OVERALL TREATMENT COMFORT. BEFORE THE FIRST TREATMENT SESSION IT IS RECOMMENDED TO CONTACT YOUR LOCAL GALDERMA REPRESENTATIVE OR RESTYLANE DISTRIBUTOR FOR MORE INFORMATION ABOUT INJECTION TECHNIQUES AND TRAINING OPPORTUNITIES. THIS PRODUCT IS ONLY INTENDED TO BE ADMINISTERED BY AUTHORIZED PERSONNEL IN ACCORDANCE WITH LOCAL LEGISLATION.			
VALIDITY DATE OF REGISTRATION	31/05/2023 - 30/05/2028			

LIST OF DEVICE			
NO	NAME OF DEVICE	IDENTIFIER	
1	Restylane Skinboosters Vital Light Lidocaine	011635	
2	Terumo Short K-pack II	N/A	
3	BD Hypoint	N/A	

NO	NAME OF DEVICE	IDENTIFIER
4	Syringe barrel	N/A
5	Tip cap	N/A
6	Luer-lock adapter	N/A
7	Shell	N/A
8	Plunger	N/A
9	Finger grip	N/A
10	Plunger rod	N/A

