

State of California—Health and Human Services Agency California Department of Public Health



Food and Drug Branch – Device Recalls

CALIFORNIA DEVICE RECALL INFORMATION SHEET

Universal Meditech Inc. Recalls Prestibio Breast Milk Alcohol Test Strip For Inability To Continue Fulfilling Any Post-Market Reponsibilities

Recall Date	Product Description	Recalling Firm	Recall Reason
11/1/2023	Prestibio Breast Milk Alcohol Test Strip REF 910-10 25 TESTS	UNIVERSAL MEDITECH INC. Reedley, California	Universal Meditech Inc. was violatively distributing PrestiBio" Breastmilk Alcohol Test Strip, DiagnosUSÂ; SARS-CoV-2 Antibody (IgG/IgM) Test, HealthyWiser KetoFast" Ketone Test Strips and PrestiBio" Ketone Test Strips without marketing authorization. UMI is also recall their tests because they are going out of business and would not be able to continue fulfilling any post- market responsibilities of these distributed products. UMI claims to hold 510(k)s for the other devices, purchased the intellectual property in 2015 from the previous 510(k) holder, but FDA has been unable to verify this information. The devices may have unknown performance

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Recall Class	Product Identification	Distribution	Affected Dates
11	PrestiBio BREAST MILK ALCOHOL TEST STRIP Model Number: 910-10 UDI-DI Code: None Lot Numbers: All products manufactured after March 2021	25 Units Nationwide including California	22 May 2023 and prior

FOR ADDITIONAL INFORMATION, PLEASE VISIT THE FDA WEBSITE

