Notification of Medical Marijuana Product Recall

The Marijuana Regulatory Agency (MRA) is issuing this health and safety advisory bulletin due to the presence of Vitamin E Acetate in several vape cartridges which failed safety compliance testing in January 2020. These cartridges were manufactured prior to the Emergency Rules for marihuana products intended for inhalation filed on November 22, 2019.

All affected vape cartridges will have a label that indicates the license number of the marijuana facility and the METRC number assigned to the product in the statewide monitoring system.

Recalled products are listed below with the dates of sale in parenthesis.

**Plan B Wellness**

This recall affects the following marijuana products sold from Plan B Wellness – License PC-000137 – located at 20101 8 Mile RD, Detroit, MI 48219:

**Individual Package # 1A405010000426A000000015** - SAVAGE STICK 1G Concentrate (October 3, 2019 and November 22, 2019)

**Individual Package # 1A405010000426A000000743** - 1g-SAVAGE-BLACKBERRY KUSH CARTRIDGE (January 16, 2020)

**Individual Package # 1A405010000426A000000744** - 1g-SAVAGE-GG#4 CARTRIDGE (January 16, 2020)

**Individual Package # 1A405010000426A000000746** - 1g- SAVAGE-RUNTZ CARTRIDGE (January 16, 2020)

Patients or caregivers who have these affected vape cartridges in their possession should return them to Plan B Wellness for proper disposal. Plan B Wellness must notify patients or caregivers that purchased these vape cartridges of the recall.

Patients who have experienced symptoms after using these products should report their symptoms and product use to their physician.

Patients and caregivers are requested to report any adverse product reactions to the MRA via email: MRA-Enforcement@michigan.gov or via phone: 517-284-8599.