February 28, 2019

## VIA UPS

Mr. Abed Asker
Access Vapor, LLC
6550 International Drive, Suite 103
Orlando, FL 32819
Submission Tracking Number: TC0004806
Dear Mr. Asker:
It has come to our attention that Access Vapor, LLC may be importing new finished tobacco products including, but not limited to, Cali Pods and Delicious Pods without premarket authorization, as required by $\S 910$ of the Federal Food, Drug, and Cosmetic Act (FD\&C Act). ${ }^{1}$ Cali Pods and Delicious Pods appear to meet the definition of a tobacco product as set forth in $\S 201(\mathrm{rr})$ of the FD\&C Act. Although FDA has extended the compliance deadlines for the premarketing requirements for deemed products, FDA's compliance policy applies only to those deemed products that were on the market as of August 8, 2016. FDA has received complaints that Access Vapor, LLC may have first commercially marketed Cali Pods and Delicious Pods in the United States after August 8, 2016.

Please provide FDA with the following information:

1. For the brand names referenced above, please list all sub-brands under the brand name, including all applicable sizes, flavors, nicotine strengths, and other variations. For each sub-brand, confirm whether you are currently marketing the product in the United States, and the date that the product was first commercially marketed in the United States. FDA suggests that you use a chart similar to the following example to help us understand your response:

| Product Name <br> (Brand and Sub- <br> brand) | Currently Marketed (Y/N) | Date Product First <br> Commercially Marketed <br> in U.S. |
| :---: | :---: | :---: |
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[^0]2. For each of the products listed above, please provide the following information, as applicable:
a. Evidence that your product was commercially marketed (other than for test marketing) as of February 15, 2007; ${ }^{2}$
b. Evidence that your product is a deemed product that was on the market on August 8, 2016, and has not been modified since that date; ${ }^{3}$
c. Evidence that your product was first introduced or delivered for introduction into interstate commerce for commercial distribution after February 15, 2007, and prior to March 22, 2011, and for which a $905(\mathrm{j})$ (or substantial equivalence) report was submitted no later than March 22, 2011; ${ }^{4}$
d. Evidence that FDA has issued an order permitting the marketing of this product; and/or
e. Evidence, which may include a statement from the firm, that the firm is currently not marketing the above-listed product(s).

We request that a written response be submitted within 30 days of receipt of this letter. The response and any further correspondence regarding this matter should reference the Submission Tracking Number listed above. We encourage you to send your response electronically via the CTP Portal ${ }^{5}$ using eSubmitter. ${ }^{6}$ However, you may also send your response by mail to our Document Control Center:

Food and Drug Administration
Center for Tobacco Products
Document Control Center
10903 New Hampshire Avenue
Building 71, Room G335
Silver Spring, MD 20993-0002

[^1]If you have any questions concerning this matter, please contact the Center for Tobacco Product's Office of Compliance and Enforcement via email at CTP-OCEPostmarket@fda.hhs.gov.

Sincerely yours,

Ann Simoneau, J.D.<br>Director<br>Office of Compliance and Enforcement<br>Center for Tobacco Products


[^0]:    ${ }^{1}$ https://www.fda.gov/downloads/TobaccoProducts/Labeling/RulesRegulationsGuidance/UCM557716.pdf

[^1]:    ${ }^{2}$ https://www.fda.gov/TobaccoProducts/Labeling/RulesRegulationsGuidance/ucm416495.htm
    ${ }^{3}$ See "Effective and Compliance Dates Applicable to Retailers, Manufacturers, Importers and Distributors of Newly Deemed Products, located here for more information. https://www.fda.gov/downloads/tobaccoproducts/guidancecomplianceregulatoryinformation/ucm501016.pdf
    ${ }^{4}$ https://www.fda.gov/downloads/TobaccoProducts/Labeling/RulesRegulationsGuidance/UCM436468.pdf
    ${ }^{5}$ http://www.fda.gov/TobaccoProducts/GuidanceComplianceRegulatoryInformation/Manufacturing/ucm515047.htm FDA's Electronic Submission Gateway (ESG) is still available as an alternative to the CTP Portal. ${ }^{6}$ http://www.fda.gov/ForIndustry/FDAeSubmitter

