DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Potential Tobacco Product Violations Report

Form Approved: OMB No.: 0910-0716 Expiration Date: 07/31/2020 (See page 3 for PRA Statement)

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Use this form to report potential tobacco-related violations of the Federal Food, Drug, and Cosmetic Act and associated regulations. These submissions are reviewed by FDA's Center for Tobacco Products.

WHO can report? - Any member of the public.

Tell us:

WHEN did you see the potential violation?

WHERE did the potential violation occur?

WHAT is the potential violation?

WHY report? - Information we receive from the public is often very helpful in identifying problems with marketed products and possible violations of the laws that we enforce.

To submit your report, complete the form below:

Date and State Where Violation Occurred						
Date potential violation occurred (mm/	(dd/yyyy)	I do not recall the date this potential violation occurred		State in which potential violation occurred		
	Desc	ription of Product				
Туре		Tobacco Brand				
Potential violation type	Sales to minor	S		Free samples		
(choose all that apply)	Flavored cigare			Self-service display/direct access to cigarette or smokeless tobacco		
		Advertising/promotion/marketing Vending machine/direct access to cigarette		Sale of cigarettes in packs of less than 20		
	or smokeless to products	obacco or covered tobacco		Unsure		
Type of potentially	Newspaper			Price signage		
violative promotional materials <i>(choose all</i>	Magazine			Posters		
that apply)	Periodicals			Coupons		
	Billboard			Internet		
	Direct mail			Unsure		
	In-store advert	isements				
Who potentially violated?	Retailer			Distributor		
(choose all that apply)	Manufacturer			Unsure		
	☐ Importer					

Potential Tobacco Product Violations Report
Description of potential violation
Name and physical address of the potential violator, if known
Retailer, manufacturer, importer, or distributor name
Street Address
Street Address Line 2
Street Address Line 2
City State/Province/Region Postal/Zip Code
If report is about a website, insert website address:
All reports will remain private to the extent allowed by law. For more information about FDA's
internet policies, please visit: http://www.fda.gov/AboutFDA/AboutThisWebsite/WebsitePolicies/default.htm
May we contact you if we No, I want my report to be anonymous. (Please note that if you submit this form by email, need additional information? No, I want my report to be anonymous. (Please note that if you submit this form by email, address. However, if you choose "no" FDA will not contact you.)
need additional information? FDA will receive your email address. However, if you choose "no" FDA will not contact you.) Yes, FDA may contact me. (Please fill in contact information below.)
Name
Affiliation (such as appropriate and arrange)
Affiliation (such as company, school, or group)
Street Address
Street Address Line 2
(continued on next page)

Potential Tobacco Product Violations Report				
City		State/Province/Region		
Postal/Zip Code		Phone Number		
Email				
Please email me to notify me that FDA got my complaint	☐ No ☐ Yes	In order to receive a response, please configure your email spam/junk filter to allow messages from ctpcompliance@fda.hhs.gov. In most cases, this is solved by adding our email address to your address book.		

If you would rather submit your report to us in writing, along with any attachments, please do so at the following address:

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

To reach us by telephone, please call 1-877-CTP-1373, and select option 3. You may also email us at ctpcompliance@fda.hhs.gov.

An email message automatically will be produced when you click the SUBMIT BY EMAIL button. In the resulting email message, please don't forget to click the "Send" button or its equivalent when you are ready to send the email.

OMB Paperwork Reduction Act Statement

This section applies only to requirements of the Paperwork Reduction Act of 1995.

DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.

The burden for this collection of information is estimated to average 0.25 hour per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to the following address:

Department of Health and Human Services Food and Drug Administration Office of Operations Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."