

Scan for Author Video Interview

Association Between Marijuana Exposure and Pulmonary Function Over 20 Years

Mark J. Pletcher, MD, MPH
Eric Vittinghoff, PhD
Ravi Kalhan, MD, MS
Joshua Richman, MD, PhD
Monika Safford, MD
Stephen Sidney, MD, MPH
Feng Lin, MS
Stefan Kertesz, MD

xposure to tobacco smoke causes lung damage with clinical consequences that include respiratory symptoms, chronic obstructive pulmonary disease, and

obstructive pulmonary disease, and lung cancer. ^{1,2} Chronic obstructive pulmonary disease and lung cancer are leading causes of death, ^{2,3} and smoking tobacco cigarettes is the most important preventable cause of death in the United States. ^{4,5}

Marijuana smoke contains many of the same constituents as tobacco smoke,6 but it is unclear whether smoking marijuana causes pulmonary damage similar to that caused by tobacco. Prior studies of marijuana smokers have demonstrated consistent evidence of airway mucosal injury and inflammation7-9 as well as increased respiratory symptoms such as cough, phlegm production, and wheeze, similar to that seen in tobacco smokers. 10-12 However, analyses of pulmonary function and lung disease have failed to detect clear adverse effects of marijuana use on pulmonary function. 10-13 It is possible that cumulative damage to the lungs from years of marijuana use could be masked by short-term

Author Video Interview available at www.jama.com.

Context Marijuana smoke contains many of the same constituents as tobacco smoke, but whether it has similar adverse effects on pulmonary function is unclear.

Objective To analyze associations between marijuana (both current and lifetime exposure) and pulmonary function.

Design, Setting, and Participants The Coronary Artery Risk Development in Young Adults (CARDIA) study, a longitudinal study collecting repeated measurements of pulmonary function and smoking over 20 years (March 26, 1985-August 19, 2006) in a cohort of 5115 men and women in 4 US cities. Mixed linear modeling was used to account for individual age-based trajectories of pulmonary function and other covariates including tobacco use, which was analyzed in parallel as a positive control. Lifetime exposure to marijuana joints was expressed in joint-years, with 1 joint-year of exposure equivalent to smoking 365 joints or filled pipe bowls.

Main Outcome Measures Forced expiratory volume in the first second of expiration (FEV_1) and forced vital capacity (FVC).

Results Marijuana exposure was nearly as common as tobacco exposure but was mostly light (median, 2-3 episodes per month). Tobacco exposure, both current and lifetime, was linearly associated with lower FEV₁ and FVC. In contrast, the association between marijuana exposure and pulmonary function was nonlinear (P<.001): at low levels of exposure, FEV₁ increased by 13 mL/joint-year (95% CI, 6.4 to 20; P<.001) and FVC by 20 mL/joint-year (95% CI, 12 to 27; P<.001), but at higher levels of exposure, these associations leveled or even reversed. The slope for FEV₁ was -2.2 mL/joint-year (95% CI, -4.6 to 0.3; P=.08) at more than 10 joint-years and -3.2 mL per marijuana smoking episode/mo (95% CI, -5.8 to -0.6; P=.02) at more than 20 episodes/mo. With very heavy marijuana use, the net association with FEV₁ was not significantly different from baseline, and the net association with FVC remained significantly greater than baseline (eg, at 20 joint-years, 76 mL [95% CI, 34 to 117]; P<.001).

Conclusion Occasional and low cumulative marijuana use was not associated with adverse effects on pulmonary function.

JAMA. 2012;307(2):173-181

www.jama.com

effects; prior analyses have not attempted to disentangle these factors. Smoking marijuana is increasingly common in the United States, ¹⁴ and understanding whether it causes lasting damage to lung function has important implications for public health messaging and medical use of marijuana. ^{15,16}

The Coronary Artery Risk Development in Young Adults (CARDIA) study collected repeated measures of tobacco and marijuana smoking as well as pulmonary function over the course

Author Affiliations: Department of Epidemiology and Biostatistics (Drs Pletcher and Vittinghoff and Mr Lin) and Division of General Internal Medicine, Department of Medicine (Dr Pletcher), University of California, San Francisco; Asthma-COPD Program, Division of Pulmonary and Critical Care Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois (Dr Kalhan); Department of Surgery (Dr Richman) and Division of Preventive Medicine (Drs Safford and Kertesz), University of Alabama at Birmingham; Center for Surgical, Medical and Acute Care Research and Transitions, Veterans Affairs Medical Center, Birmingham (Drs Richman and Kertesz); and Division of Research, Kaiser Permanente of Northern California, Oakland (Dr Sidney).

Corresponding Author: Mark J. Pletcher, MD, MPH, Department of Epidemiology and Biostatistics, University of California, San Francisco, 185 Berry St, Ste 5700, San Francisco, CA 94107 (mpletcher@epi.ucsf.edu).

of 20 years (March 26, 1985-August 19, 2006) in more than 5000 study participants. We estimated both current intensity and lifetime cumulative exposure to tobacco and marijuana smoking and analyzed their associations with spirometric measures of pulmonary function over the 20 years of follow-up.

METHODS

Study Design and Sample

CARDIA is a longitudinal study designed to measure risk factors for coronary artery disease in a cohort of black and white women and men (n=5115) aged 18 through 30 years and healthy at enrollment in 1985. ^{17,18} Participants were sampled from 4 US communities without selection for smoking behaviors and comprise a broad cross-section of typical tobacco and marijuana use patterns.

With the written informed consent of participants and the approval of institutional review boards at each study center (Oakland, Chicago, Minneapolis, and Birmingham), participants underwent a baseline examination and 6 follow-up examinations, with 69% retention at year 20. Pulmonary function testing was performed at years 0, 2, 5, 10, and 20. For this investigation, we included all visits for which pulmonary function, smoking behavior, secondhand smoke exposure, height, and waist circumference were available.

Tobacco and Marijuana Exposure

Current intensity of tobacco use (cigarettes smoked per day) was assessed at each examination. These data, along with baseline examination data on past years of smoking, were used to estimate cumulative lifetime exposure to cigarettes in terms of pack-years, with 1 pack-year of exposure equivalent to 7300 cigarettes (1 year × 365 days/y × 1 pack/d × 20 cigarettes/pack). Misclassification of smoking exposure by self-report, measured by comparisons with serum cotinine levels, is uncommon.¹⁹

Current intensity of marijuana use (episodes in the last 30 days) was also

assessed at each examination. Using baseline examination data on past lifetime exposure to marijuana, current intensity of marijuana use, and another question designed to assess number of joints or filled pipe bowls smoked per episode (eMethods, available at http://www.jama.com), we calculated total lifetime exposure to marijuana joints in joint-years, with 1 joint-year of exposure equivalent to 365 joints or filled pipe bowls smoked (1 year × 365 days/y × 1 joint/d), as described previously.²⁰

Outcome Measures

Study outcomes were forced expiratory volume in the first second of expiration (FEV₁) and forced vital capacity (FVC) measured by forced spirometry. These were collected using a Collins Survey 8-L water-sealed spirometer and an Eagle II microprocessor (years 0, 2, 5, and 10) and then an OMI rolling seal spirometer (year 20). A comparability study performed among 25 participants demonstrated an average difference of less than 1% for both measurements. Standard quality control and testing procedures were maintained according to established guidelines.21,22

Other Covariates

CARDIA was designed to recruit approximately equal numbers of selfidentified "black, not Hispanic" and "white, not Hispanic" men and women to ensure an adequate sample of the largest minority group in the United States at that time. Height and waist circumference were measured at each examination. As a proxy for socioeconomic status, we used the maximum educational grade attained for each participant. Secondhand smoke exposure in hours per week (sum of exposure in the home, small enclosed spaces, and large spaces) was assessed at each examination, with linear interpolation for missing data. Asthma was self-reported at each examination; we used the baseline assessment. We obtained average annual city-specific levels of airborne

particulate matter less than 10 microns and less than 2.5 microns in size²³ around the 4 CARDIA study centers from the Environmental Protection Agency²⁴ (eMethods).

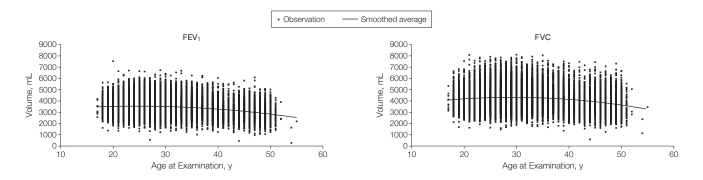
Statistical Analysis

Participants were categorized by whether they ever reported current use of tobacco, marijuana, or both at a CARDIA examination and compared across these categories using descriptive statistics. We then categorized participants according to degree of current and lifetime tobacco and marijuana exposure at each examination and described pulmonary function (FEV₁ and FVC) across categories before and after adjustment. Tests of trend and interaction were performed in fully adjusted models.

The categorized exposure models described above represent a standard approach to multivariable-adjusted association testing. Categorization models, however, use necessarily arbitrary category thresholds and do not take full advantage of the continuous exposure measurements for estimation or adjustment purposes. To fully explore and test potential nonlinear associations, we modeled tobacco and marijuana exposure variables as flexible cubic splines (eMethods) in adjusted models to allow associations with pulmonary function to take different shapes at lower vs higher levels of exposure.25

For each adjusted analysis described above, we used mixed models accounting for repeated measures of pulmonary function within participants, with a random intercept and a random 3-knot age spline within each individual and an unstructured variance-covariance structure. Fully adjusted models included fixed effects for year, center, and center-year (their interaction), race-sex category, education, and asthma; cubic splines for age, height, waist circumference, secondhand smoke exposure, and exposure to airborne particulate matter less than 10 microns and less than 2.5 microns in size; and interactions between the agespline variables and race-sex, asthma,

Figure 1. Pulmonary Function Measurements by Age



Participants (n=5017) contributed an average of 3.9 measurements per person (n=19705 total) over the course of 20 years. A lowess smoother was used to calculate the smoothed average. FEV₁ indicates forced expiratory volume in first second of expiration; FVC, forced vital capacity.

waist-spline variables, and heightspline variables to allow for differing flexible age-based trajectories of pulmonary function for participants with differing characteristics. Models were queried to produce adjusted estimates of slope (reflecting the incremental difference in pulmonary function observed with additional tobacco or marijuana smoking) and net association (reflecting the net observed difference between persons with a particular level of consumption and persons with none) at various points along the association curve. All analyses were performed using Stata version 11 and used 2-sided tests for significance at the .05 level, with 95% CIs.

RESULTS

The 5115 CARDIA participants recruited in 1985-1986 contributed 20777 total visits that included pulmonary function testing. Of these, 959 visits were excluded for lack of complete information on smoking behavior, 114 for lack of height or waist measurements, and 1 for an unknown visit date, leaving 19703 visits (95%) with complete data from 5016 participants (98%). Participants contributed 3.9 visits/participant on average; attrition was more common in tobacco smokers but not associated with marijuana use. FEV₁ and FVC varied across participants, increased slightly with age through the

late 20s, and declined slowly thereafter (FIGURE 1).

More than half of participants (54%; mean age at baseline, 25 years) reported current marijuana smoking, tobacco smoking, or both at 1 or more examinations (TABLE 1). Smoking patterns differed by race and sex, with black women most likely to smoke tobacco only, white men most likely to smoke marijuana only, and black men most likely to smoke both. Tobacco smokers tended to have lower education and income and to be slightly shorter and less active, whereas marijuana smokers tended to be taller and more active. The median intensity of tobacco use in tobacco smokers was substantially higher (8-9 cigarettes/d) than the median intensity of marijuana use in marijuana smokers (2-3 episodes in the last 30 days). Although marijuana and tobacco exposures were strongly correlated, our sample included 91 participants with no tobacco exposure and more than 10 joint-years of marijuana exposure (contributing 153 observations of pulmonary function), 40 (56 observations) of whom had more than 20 joint-years of exposure.

In fully adjusted models that considered 4-level categorizations of current and lifetime exposure to tobacco and marijuana, tobacco smoking (both

current and lifetime) was associated with a lower FEV₁ and current smoking with a lower FVC (TABLE 2). For example, compared with zero exposure, FEV₁ was 63 mL lower (95% CI, -89 to -36; P<.001 for trend) and FVC was 69 mL lower (95% CI, -97 to -41; P<.001 for trend) with current tobacco exposure of more than 20 cigarettes per day and 101 mL lower (95% CI, -136 to -65; P<.001 for trend) with lifetime tobacco exposure of more than 20 pack-years.

In contrast, exposure to marijuana (both current and lifetime) was associated with higher FVC and lifetime exposure with higher FEV₁. For example, compared with zero exposure, FVC increased with greater lifetime exposure in joint-years (P=.01)for trend) and FEV1 increased with greater lifetime exposure of up to 10 joint-years and then declined to 36 mL (95% CI, -6.5 to 79) greater than the zero exposure level (P=.049 for trend). FVC increased with smoking intensity up to 20 marijuana smoking episodes in the past 30 days and then declined to 20 mL greater than the zero exposure level (P = .03 for trend). We found no statistically significant interactions between tobacco and marijuana exposure for either FEV₁ or FVC.

When we modeled current and lifetime tobacco and marijuana exposure as continuous exposures and permitted flexible nonlinear associations (via splines), we again found strong, dose-related associations (P < .001) between increasing exposure to tobacco and lower FEV₁ and FVC (FIGURE 2), with no evidence of nonlinearity (TABLE 3). Declining slopes ranged as steep as -2.8 mL (95% CI, -4.8 to -0.7; P=.007) per additional cigarette smoked per day and -7.0 mL (95% CI, -10 to -3.7; P < .001) per additional pack-year for FEV₁ and were of similar magnitude for FVC (Table 3). At 50 pack-years of exposure, FEV₁ was on average 332 mL lower (95% CI, -401 to -263; P < .001)

and FVC was 229 mL lower (95% CI, -310 to -147; P < .001), compared with no exposure.

For marijuana, we found strong statistical evidence that associations between marijuana use and pulmonary function were nonlinear (Figure 2, Table 3). At low lifetime exposure levels, increasing marijuana use was associated with a steep increase in both FEV₁ (13 mL/joint-year higher [95% CI, 6.4 to 20], P<.001) and FVC (20 mL/joint-year higher [95% CI, 12 to 27], P<.001), but at higher levels of exposure (>7 joint-years), the slope leveled or even turned downward. At more

than 10 joint-years of lifetime exposure, we found a nonsignificant decline in FEV₁ (-2.2 mL/joint-year [95% CI, -4.8 to 0.3], P = .08) but a significant decline in FEV1 at more than 20 episodes of marijuana use per month (-3.2 mL/episode [95% CI, -5.8 to -0.6], P=.02). Although net associations with FEV₁ became negative at very high exposure levels (>40 joint-years or >25 episodes/mo), these negative deflections were not statistically significant (Table 3). FVC remained significantly elevated in even heavy users (eg, 76 mL [95% CI, 34 to 117; P < .001] at 20 joint-years).

Table 1.	 Characteristics of 	CARDIA Participants	vvitn Pulmonary	Function	rest Results	s, by Smoking E	enavior
					Marij	uana/Tobacco l	Jse ^a

	Marijuana/Tobacco Use ^a					
Baseline Characteristics ^b	Neither (n = 2305)	Tobacco Only (n = 851)	Marijuana Only (n = 795)	Both (n = 1065)	<i>P</i> Value ^c	
Age, mean (SD), y	25 (4)	25 (4)	25 (4)	25 (4)	<.001	
Race-sex, No. (%) ^d White men	525 (23)	133 (16)	251 (32)	249 (23)		
White women	672 (29)	266 (31)	186 (23)	172 (16)	<.001	
Black men	399 (17)	167 (20)	185 (23)	367 (34)	<.001	
Women	709 (31)	285 (33)	173 (22)	277 (26)		
College educated at any examination, No. (%) ^b	1291 (56)	245 (29)	381 (48)	240 (22)	<.001	
Income >\$50 000/y at any examination, No. (%)	1414 (68)	324 (46)	429 (60)	344 (35)	<.001	
Body mass index, mean (SD) ^e	25 (5)	25 (5)	24 (4)	25 (5)	.22	
Height, mean (SD), cm	170 (10)	169 (9)	172 (9)	171 (9)	<.001	
Waist circumference, mean (SD), cm	77.4 (11.9)	77.6 (11.5)	78.0 (10.6)	78.8 (11.2)	.009	
History of asthma at the baseline visit, No. (%)	89 (4)	45 (5)	39 (5)	43 (4)	.001	
Secondhand smoke exposure, median (IQR), h/wk	7 (3-25)	28 (10-56)	12 (4-38)	33 (12-62)	<.001	
Airborne particulate matter exposure, mean (SD), μg/m³f PM10	86 (19)	85 (20)	87 (21)	84 (19)	.006	
PM2.5	33 (8)	35 (8)	33 (8)	33 (8)	.002	
Average intensity of tobacco use, median (IQR), cigarettes/d ^b		8 (3-15)		9 (4-15)	.37	
Average intensity of marijuana use, median (IQR), episodes in last 30 d ^b			2 (1-6)	3 (1-9)	<.001	
Lifetime tobacco use, median (IQR), pack-years ^b		7 (3-15)		9 (3-16)	.07	
Lifetime marijuana use, median (IQR), joint-years ^b			0.9 (0.2-2.8)	1.5 (0.6-4.3)	<.001	
CARDIA examinations with PFT results recorded, No. (SD)	4.0 (1.1)	3.6 (1.2)	4.0 (1.2)	3.9 (1.1)	<.001	
Attended year 20 examination, No. (%)	1442 (63)	357 (42)	492 (62)	516 (48)	<.001	

Abbreviations: CARDIA, Coronary Artery Risk Development in Young Adults study; COPD, chronic obstructive pulmonary disease; IQR, interquartile range; PFT, pulmonary function test; PM10, airborne particulate matter less than 10 microns in size; PM2.5, airborne particulate matter less than 2.5 microns in size.

Current use reported at 1 or more CARDIA examinations at which pulmonary function was measured.

DUnless otherwise noted, values at the first available examination at which pulmonary function was measured are presented. For average smoking intensity, an average across all examinations was calculated, and the median (IQR) of these averages is presented. For lifetime smoking exposure, the maximum (last) value was used, and the median (IQR) of these maximums is presented.

^cP values are from a 1-way analysis of variance test for age, body mass index, height, waist circumference, PM10 and PM2.5 exposure, and number of CARDIA examinations; from a x² test for race-sex, education, income, and asthma; and from a Kruskal-Wallis nonparametric test for smoking variables, limiting comparisons to smokers of the relevant substance for each test.

d By design, the CARDIA study sampled white men, white women, black men, and black women in roughly equal numbers for participation in the study (see "Methods").

Calculated as weight in kilograms divided by height in meters squared

^fMeasured at the level of the city or metropolitan area.

COMMENT

In this 20-year study of marijuana and pulmonary function, we confirmed the expected reductions in FEV₁ and FVC from tobacco use. In contrast, marijuana use was associated with higher FEV₁ and FVC at the low levels of exposure typical for most marijuana users. With up to 7 joint-years of lifetime exposure (eg, 1 joint/d for 7 years or 1 joint/wk for 49 years), we found no evidence that increasing exposure to marijuana adversely affects pulmonary function. This association, however, was nonlinear: at higher exposure levels, we found a leveling off or even a reversal in this association, especially for FEV₁. Although our sample contained insufficient numbers of heavy users to confirm a detrimental effect of very heavy marijuana use on pulmonary function, our findings suggest this possibility.

The associations we found between tobacco and pulmonary function are consistent with a large body of prior research on the adverse pulmonary consequences of tobacco smoking. The high prevalence of tobacco smoking, the wide range of exposure intensity among smokers, and the legality of tobacco have made tobacco smoking an easy target for observational epidemiology. Exposure predicts reduced expiratory flow and air trapping, gas-exchange abnormalities, and emphysema,1 and smoking cessation interventions reduce the rate of FEV₁ decline in smokers²⁶ (ie, these associations are likely causal). Our findings of a linear dose-response relationship showing lower FEV1 and FVC with increasing tobacco expo-

		FEV ₁			FVC			
Smoking Exposure Category	No. ^a	Mean (SD), L	Adjusted Difference (95% CI), mL ^c	<i>P</i> Value ^b	Mean (SD), L	Adjusted Difference (95% CI), mL ^c	<i>P</i> Value ^t	
Overall	19704	3420 (810)			4.23 (1.0)			
Current tobacco/marijuana smoking status								
Neither	12 288	3.41 (0.80)	1 [Reference]		4.19 (1.03)	1 [Reference]		
Tobacco only	3483	3.27 (0.77)	-24 (-38 to -11)	.003	4.07 (.97)	-19 (-33 to -4.6)	.004	
Marijuana only	2021	3.73 (0.81)	0.7 (-12 to 13)	.003	4.60 (1.04)	8.2 (-5 to 22)	.004	
Both	1912	3.52 (0.79)	−13 (−29 to 3)		4.39 (1.02)	2.7 (-14 to 20)		
Current tobacco smoking intensity, cigarettes/d								
0	14313	3.45 (0.81)	1 [Reference]	<.001	4.24 (1.04)	1 [Reference]	<.001	
1-10	2972	3.28 (0.76)	-13 (-27 to 1.0)		4.05 (.95)	-15 (-30 to -0.4)		
11-20	1852	3.41 (0.79)	-36 (-53 to -19)		4.27 (1.00)	-30 (-49 to -12)		
>20	567	3.63 (0.82)	-63 (−89 to −36) J		4.60 (1.05)	-69 (−97 to −41)		
Current marijuana smoking intensity, episodes in the last 30 d								
0	15771	3.38 (0.80)	1 [Reference]		4.16 (1.02)	1 [Reference]		
1-10	2784	3.59 (0.81)	0.8 (-10 to 11)	.32	4.44 (1.03)	5.8 (-5.4 to 17)	.03	
11-20	665	3.68 (0.80)	16 (-3.5 to 35)	.32	4.57 (1.03)	35 (15 to 55)	.00	
>20	484	3.75 (0.77)	−18 (−42 to 6.1)		4.75 (1.01)	20 (-5.2 to 49)		
Lifetime exposure to tobacco, pack-years d								
0	11 183	3.44 (0.82)	1 [Reference]		4.22 (1.05)	1 [Reference]	.047	
1-10	6458	3.44 (0.77)	3.2 (-18 to 25)	<.001	4.24 (.99)	37 (12 to 61)		
11-20	1447	3.35 (0.83)	-41 (-38 to -14)		4.24 (1.07)	11 (-20 to 41)		
>20	616	3.29 (0.85)	-101 (−136 to −65) <u></u>		4.27 (1.09)	−35 (−76 to 5.0)		
Lifetime exposure to marijuana, joint-years ^d								
0	5619	3.28 (0.79)	1 [Reference]		4.00 (1.00)	1 [Reference]		
1-5	13 493	3.49 (0.80)	38 (15 to 62)	.049	4.31 (1.03)	41 (14 to 67)	.01	
6-10	371	3.57 (0.78)	66 (32 to 100)	.010	4.50 (1.02)	54 (16 to 91)		
>10	221	3.45 (0.86)	36 (-6.5 to 79)		4.44 (1.08)	59 (12 to 107)		

Abbreviations: FEV₁, forced expiratory volume in first second of expiration; FVC, forced vital capacity.

^aRefers to the number of observations; the 5016 participants contributed an average of 3.9 observations per participant.

^bFor trend, except for "current tobacco/marijuana smoking status," for which a nonordered test is used.

CAdjusted differences represent comparisons of average pulmonary function (FEV1 and FVC), in mL, between persons in the given smoking exposure category and the reference category. Mixed models with a random intercept and a random 3-knot age spline were used to adjust for repeated measures, and fixed effects were included for year, center and center-year (their interaction), race-sex category, education, and asthma; cubic splines for age, height, waist circumference, secondhand smoke exposure, and exposure to airborne particulate matter less than 10 microns and less than 2.5 microns in size; and interactions between the age spline variables and race-sex, asthma, waist spline variables, and height spline variables. Except for in the first subsection (current tobacco/marijuana smoking status), all 4 smoking variables (4 categories each) were included in the same model, including current

and lifetime smoking intensity for both tobacco and marijuana. d One pack-year of exposure to tobacco smoke equals 7300 cigarettes (1 pack/d×20 cigarettes/pack×365 d/y); 1 joint-year of exposure to marijuana smoke equals 365 joints of marijuana (1 joint/d × 365 d/y).

sure, consistent with prior findings, represent a positive control for our study of the association between marijuana smoking and pulmonary function.

Prior studies of marijuana smoking and pulmonary function have yielded apparently conflicting results. ¹⁰⁻¹³ Many studies have focused on FEV₁:FVC ratio, lower values of which suggest the presence of airway obstruction, and have found either no association ^{10,20,27} or lower FEV₁:FVC ratios with marijuana use. ²⁸⁻³² Lower FEV₁:FVC ratios in marijuana smokers, however, can be explained at least partly by a tendency toward higher FVC or total lung capacity. ^{28,29,32} A recent longitudinal study,

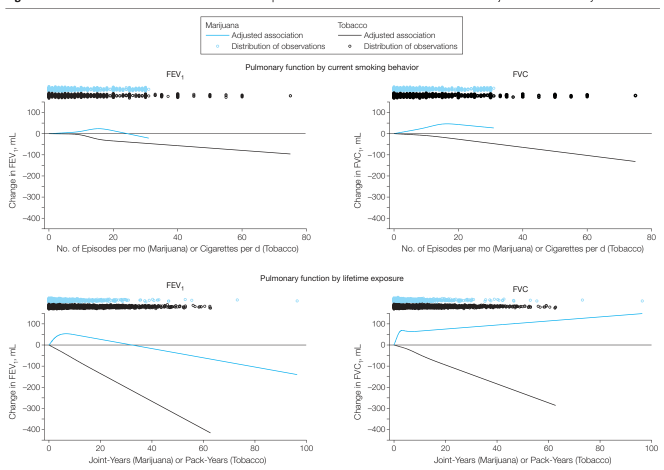
which demonstrated significantly higher FVC and total lung capacity with marijuana exposure, strongly supports this notion, ^{13,20} as does our study.

The potential association of marijuana smoking with FEV₁ has been even less clear. Tobacco smoking reduces FEV₁, but despite the similarities in the constituents of marijuana smoke and tobacco smoke and our a priori expectations that marijuana smoking might have similar effects, prior research has not demonstrated this. In studies that report FEV₁ in association with marijuana use, findings have mostly been null,^{20,28,32-35} although one study reported the apparently paradoxical find-

ing of a lower FEV₁ with past marijuana use but a nonsignificantly higher FEV₁ with current use.²⁹

Our study suggests a way to reconcile these findings. Because of the many thousands of measurements obtained over 20 years among more than 5000 participants with a wide range of smoking habits, we could simultaneously account for levels of current and past lifetime use of both marijuana and tobacco and test for nonlinearity in their associations with pulmonary function to disentangle short-term and long-term effects. We found highly significant nonlinearity, with a positive association for both FEV₁ and FVC at low

Figure 2. Associations Between Continuous Smoothed Exposure to Current and Lifetime Tobacco and Marijuana and Pulmonary Function



Associations between continuous current and lifetime exposure measurements and pulmonary function were modeled via cubic splines (see "Methods"). All 4 exposure measurements were included in each model (one model each for forced expiratory volume in the first second of expiration [FEV₁] and forced vital capacity [FVC]). Mixed models with a random intercept and a random 3-knot age spline were used to adjust for repeated measures, and fixed effects were included for year, center and center-year (their interaction), race-sex category, education, and asthma; cubic splines for age, height, waist circumference, secondhand smoke exposure, and exposure to airborne particulate matter less than 10 microns and less than 2.5 microns in size; and interactions between the age spline variables and race-sex, asthma, waist spline variables, and height spline variables. Point estimates and confidence intervals for slopes and net associations at different exposure levels are provided in Table 3.

Table 3. Estimated Slopes and Net Associations Between Continuous Smoothed Exposure to Current and Lifetime Tobacco and Marijuana and **Pulmonary Function**

	FEV ₁	FVC			
Smoking Exposure Estimate Type ^a	Adjusted Estimate (95% CI) ^b	P Value	Adjusted Estimate (95% CI) ^b	P Value	
Current marijuana exposure Test of overall association		.06		<.001	
Test of nonlinearity		.02		.04	
Slope, mL per episode per mo At 5 episodes/mo	0.8 (-1.4 to 3.1)	.47	2.8 (0.4 to 5.1)	.02	
At 10 episodes/mo	2.6 (-0.3 to 5.4)	.07	3.7 (0.7 to 6.6)	.02	
At 20 episodes/mo	-3.2 (-5.8 to -0.6)	.02	-1.5 (-4.2 to 1.3)	.30	
At 40 episodes/mo	NA ^c	NAc	NAc	NAc	
Net association, mL At 5 episodes/mo	4.1 (-7.1 to 15)	.47	14 (1.9 to 26)	.02	
At 10 episodes/mo	11 (-6.2 to 29)	.21	29 (11 to 48)	.002	
At 20 episodes/mo	14 (-4.7 to 32)	.14	43 (23 to 63)	<.001	
At 40 episodes/mo	NAc	NA ^c	NA°	NAc	
Lifetime marijuana exposure Test of overall association		<.001		<.001	
Test of nonlinearity		<.001		<.001	
Slope, mL per joint-year At 2 joint-years	13 (6.4 to 20)	<.001	20 (12 to 27)	<.001	
At 7 joint-years	-0.4 (-2.6 to 1.8)	.74	0.0 (-2.4 to 2.5)	.97	
At 20 joint-years	-2.2 (-4.6 to 0.3)	.08	1.0 (-1.8 to 3.7)	.49	
At 50 joint-years	-2.2 (-4.6 to 0.3)	.08	1.0 (–1.8 to 3.7)	.49	
Net association, mL At 2 joint-years	30 (8.4 to 53)	.007	59 (35 to 83)	<.001	
At 7 joint-years	53 (28 to 79)	<.001	64 (36 to 92)	<.001	
At 20 joint-years	27 (-10 to 64)	.16	76 (34 to 117)	<.001	
At 50 joint-years	-39 (-141 to 64)	.46	104 (-12 to 220)	.08	
Current tobacco exposure Test of overall association		<.001		.003	
Test of nonlinearity		.29		.73	
Slope, mL per cigarettes/d At 5 cigarettes/d	-0.2 (-2.3 to 1.9)	.85	-0.8 (-3.1 to 1.4)	.46	
At 10 cigarettes/d	-2.8 (-4.8 to -0.7)	.007	-1.3 (-3.4 to 0.9)	.25	
At 20 cigarettes/d	-1.1 (-2.7 to 0.5)	.16	-1.9 (-3.6 to -0.2)	.02	
At 40 cigarettes/d	-1.1 (-2.7 to 0.5)	.16	-1.9 (-3.6 to -0.2)	.02	
Net association, mL At 5 cigarettes/d	-1.0 (-11 to 9.4)	.85	-4.2 (-15 to 6.9)	.46	
At 10 cigarettes/d	-6.3 (-23 to 11)	.47	-9.1 (-27 to 8.9)	.32	
At 20 cigarettes/d	-34 (-53 to -16)	<.001	-26 (-46 to -7.0)	.008	
At 40 cigarettes/d	-57 (-92 to -22)	.001	-65 (-102 to -28)	.001	
Lifetime tobacco exposure Test of overall association		<.001		<.001	
Test of nonlinearity		.98		.85	
Slope, mL per pack-year At 2 pack-years	-6.5 (-12 to -1.2)	.02	-3.5 (-9.3 to 2.4)	.25	
At 7 pack-years	-7.0 (-10 to -3.7)	<.001	-5.5 (-9.3 to -1.8)	.004	
At 20 pack-years	-6.6 (-8.4 to -4.7)	<.001	-4.5 (-6.6 to -2.3)	<.001	
At 50 pack-years	-6.6 (-8.4 to -4.7)	<.001	-4.5 (-6.6 to -2.3)	<.001	
Net association, mL At 2 pack-years	-13 (-23 to -2.4)	.02	-6.9 (-19 to 4.8)	.25	
At 7 pack-years	-46 (-72 to -21)	<.001	-28 (-57 to 0.1)	.05	
At 20 pack-years	-135 (-166 to -104)	<.001	-95 (-130 to -59)	<.001	
At 50 pack-years	-332 (-401 to -263)	<.001	-229 (-310 to -147)	<.001	

Abbreviations: FEV₁, forced expiratory volume in first second of expiration; FVC, forced vital capacity; NA, not available.

^aAssociations between continuous current and lifetime exposure measurements and pulmonary function were modeled via cubic splines (see "Methods"), and the estimates presented here describe the same analyses illustrated in Figure 2. The estimates presented are for slope (reflecting the incremental difference in pulmonary function observed with

sented niet describe the same analyses indistrated in Figure 2. The estimates presented are to slope (reflecting the internet and inter

^CData not available at this exposure level.

levels of exposure that reversed in direction toward a possibly negative association for FEV1 at higher levels of exposure (Figure 2 and slopes in Table 3). These findings could explain the paradox previously noted regarding past and current use29 and are also consistent with the average null association reported in studies^{20,28,32-35} that either dichotomized marijuana exposure (user/nonuser)^{28-31,33,36} or constrained the association to be linear across all levels of exposure. 10,20,32,35 When we looked at "marijuana only" smokers (Table 2), we also found a null association with FEV₁ and FVC. Only after parsing the association at different levels of exposure, with careful control for confounding, did the suggestion emerge of a negative association for FEV1 at high levels of exposure.

These findings suggest that marijuana smoking could influence pulmonary function via multiple mechanisms. To explain the higher FVC previously observed in marijuana smokers, 20,32 some investigators have proposed that the deep inspiratory maneuvers practiced by marijuana smokers could stretch the lungs, ^{13,20} resulting in larger lung volumes. ^{20,32} Another speculative possibility is strengthening of chest wall musculature or another "training" effect that allows marijuana users to inspire more fully (closer to total lung capacity) on spirometry testing. A nondestructive stretch or training effect is consistent with previously reported findings in marijuana smokers of lower lung density32 and a lack of emphysematous change³² or diminished diffusion capacity. 20,27,32,36 This mechanism would explain our FVC results and could explain the positive deflection of FEV1. The functional effects of this association on lung health or respiratory function in daily life are unclear.13 An alternate explanation is the acute bronchodilatory effect of marijuana use that has been directly observed in some studies.11 This effect. however, is transient (lasting approximately 60 minutes11) and seems unlikely to explain higher lung volumes measured during the CARDIA examination unless many marijuana users smoked immediately before the examination.

The suggestion of a negative association with FEV1 at higher exposure levels could reflect mixing of this putative stretch/training effect with a second mechanism operating on a different time-exposure scale. A negative association with heavy exposure to marijuana smoke aligns with our a priori hypothesis that marijuana smoking should produce damage to the airways and accelerated loss of lung function similar to that caused by tobacco smoking. Hypothetically speaking, a positive effect from marijuana in the short term (the stretch/training effect) and a negative effect in the long term (damage from smoke exposure) should result in a nonlinear association such as the one we observed. According to this explanation, the predominant effect for FEV₁ at very high exposure (more than 40 joint-years) reflects cumulative damage; the predominant effect for FVC at all levels of exposure is from the stretch/training mechanism.

Our study has limitations. Although CARDIA offers longitudinal spirometry measurements, it lacked body plethysmographic measurements of static lung volumes (total lung capacity and residual volume) and measures of diffusing capacity and radiographic emphysema. A minority of our participants reported very high levels of marijuana exposure (and a smaller minority of these were nonsmokers of tobacco), so our estimates at high marijuana exposure levels are imprecise. The self-reported measures of marijuana and tobacco smoking are certain to include recall error, both random and systematic, and do not include any indication of smoking method (joint, pipe, "bong", etc). It is unlikely, however, that such error would differentially occur in association with pulmonary function, and nondifferential error would most likely bias results toward the null. Our mixed modeling approach is ideal for filtering out random error and taking advantage of individual-level correlations in the data.

As with any observational analysis, unmeasured or inadequately modeled confounding effects could be mixed with our estimates, but the extensive covariate measurements and large sample in our study permitted more extensive efforts to control confounding than were possible in previous studies. This study addressed respiratory exposure to marijuana and not exposure by ingestion. Recent increases in the potency of marijuana are unlikely to have influenced our estimates, because we did not detect an interaction of marijuana and pulmonary function by calendar time.

Marijuana may have beneficial effects on pain control, appetite, mood, and management of other chronic symptoms. 15,16 Our findings suggest that occasional use of marijuana for these or other purposes may not be associated with adverse consequences on pulmonary function. It is more difficult to estimate the potential effects of regular heavy use, because this pattern of use is relatively rare in our study sample; however, our findings do suggest an accelerated decline in pulmonary function with heavy use and a resulting need for caution and moderation when marijuana use is considered.

Author Contributions: Dr Pletcher had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Pletcher, Richman, Safford. Acquisition of data: Sidney.

Analysis and interpretation of data: Pletcher, Vittinghoff, Kalhan, Richman, Safford, Lin, Kertesz. Drafting of the manuscript: Pletcher, Safford.

Critical revision of the manuscript for important intellectual content: Pletcher, Vittinghoff, Kalhan, Richman, Safford, Sidney, Lin, Kertesz.

Statistical analysis: Pletcher, Vittinghoff, Richman, Lin.
Obtained funding: Pletcher, Sidney, Kertesz.
Administrative technical or material synnort: Kertesz.

Administrative, technical, or material support: Kertesz. Study supervision: Kertesz

Conflict of Interest Disclosures: The authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Dr Kalhan reported serving as a consultant for Boehringer-Ingelheim, Forest Laboratories, and AstraZeneca; receiving honoraria for lectures from GlaxoSmithKline and AstraZeneca; receiving honoraria for development of educational materials from Quantia Communications and Medscape Education; and receiving industry-sponsored grants from GlaxoSmithKline and Boehringer-Ingelheim. Dr Kertesz reported chairing a committee that advised the Drug Treatment Task Force for the Chief Justice of the State of Alabama and that he is an employee of the Department of Veterans Affairs. No other authors reported disclo-

Funding/Support: This study was supported by the National Institute on Drug Abuse (R01-DA-025067) and the National Heart, Lung, and Blood Institute (NO1-HC-95095 and N01-HC-48047).

Role of the Sponsors: The National Institute on Drug Abuse funded this analysis but did not participate in CARDIA or review the manuscript. The National Heart, Lung, and Blood Institute helped design CARDIA and funds data collection, supports a Publications and Presentations Committee that reviews and approves all publications, and provides a representative who sits on that committee, but does not otherwise control publication

Disclaimer: The views expressed in this article do not reflect positions of the Department of Veterans Affairs or of any other entity of the federal govern-

Online-Only Material: The eMethods, eTable, eFigures 1 and 2, and Author Video Interview are available at http://www.jama.com.

REFERENCES

- 1. Kamholz SL. Pulmonary and cardiovascular consequences of smoking. Med Clin North Am. 2004; 88(6):1415-1430.
- 2. Pauwels RA, Rabe KF. Burden and clinical features of chronic obstructive pulmonary disease (COPD). Lancet. 2004;364(9434):613-620.
- 3. Number of deaths from each cause, by 10-year age groups, race, and sex: United States, 2005. Centers for Disease Control and Prevention National Vital Statistics Web site. http://205.207.175.93/VitalStats /TableViewer/tableView.aspx?ReportId=26044. Accessed November 30, 2011.
- 4. Danaei G, Ding EL, Mozaffarian D, et al. The preventable causes of death in the United States: comparative risk assessment of dietary, lifestyle, and metabolic risk factors. PLoS Med. 2009;6(4): e1000058.
- 5. Mokdad AH, Marks JS, Stroup DF, Gerberding JL. Actual causes of death in the United States, 2000. JAMA, 2004:291(10):1238-1245.
- 6. Novotny M, Merli F, Weisler D, Fencl M, Saeed T. Fractionation and capillary gas chromatographicmass spectrometric characterization of the neutral components in marijuana and tobacco smoke condensates. J Chromatogr A. 1982;238(1):141-150.
- 7. Fligiel SE, Roth MD, Kleerup EC, Barsky SH, Simmons MS, Tashkin DP. Tracheobronchial histopathology in habitual smokers of cocaine, marijuana, and/or tobacco. Chest. 1997;112(2):319-
- 8. Barsky SH, Roth MD, Kleerup EC, Simmons M,

- Tashkin DP. Histopathologic and molecular alterations in bronchial epithelium in habitual smokers of marijuana, cocaine, and/or tobacco. J Natl Cancer Inst. 1998;90(16):1198-1205.
- 9. Roth MD, Arora A, Barsky SH, Kleerup EC, Simmons M, Tashkin DP. Airway inflammation in young marijuana and tobacco smokers. Am J Respir Crit Care Med. 1998;157(3, pt 1):928-937.
- 10. Tashkin DP, Baldwin GC, Sarafian T, Dubinett S, Roth MD. Respiratory and immunologic consequences of marijuana smoking. J Clin Pharmacol. 2002; 42(11)(suppl):71S-81S.
- 11. Tetrault JM, Crothers K, Moore BA, Mehra R, Concato J, Fiellin DA. Effects of marijuana smoking on pulmonary function and respiratory complications: a systematic review. Arch Intern Med. 2007; 167(3):221-228.
- 12. Hall W, Degenhardt L. Adverse health effects of non-medical cannabis use. Lancet. 2009;374(9698): 1383-1391
- 13. Tashkin DP. Does cannabis use predispose to chronic airflow obstruction? Eur Respir J. 2010; 35(1):3-5
- 14. Results from the 2008 National Survey on Drug Use and Health: National Findings. HHS Publication SMA 09-4434. Substance Abuse and Mental Health Services Administration Web site. http://www.oas .samhsa.gov/NSDUH/2k8NSDUH/2k8results.cfm. 2009. Accessed November 30, 2011.
- 15. Joy JE, Watson SJ, Benson JA, eds. Marijuana and Medicine: Assessing the Science Base. Washington, DC: National Academies Press; 1999.
- 16. Cannabis and cannabinoids. National Cancer Institute Web site. http://www.cancer.gov/cancertopics /pdg/cam/cannabis/healthprofessional. 2011. Accessed November 30, 2011.
- 17. Hughes GH, Cutter GR, Donahue R, et al. Recruitment in the Coronary Artery Disease Risk Development in Young Adults (CARDIA) study. Control Clin Trials. 1987;8(4)(suppl):68S-73S.
- 18. Friedman GD, Cutter GR, Donahue RP, et al. CARDIA: study design, recruitment, and some characteristics of the examined subjects. J Clin Epidemiol. 1988:41(11):1105-1116.
- 19. Wagenknecht LE, Burke GL, Perkins LL, Haley NJ. Friedman GD. Misclassification of smoking status in the CARDIA study: a comparison of self-report with serum cotinine levels. Am J Public Health. 1992; 82(1):33-36
- 20. Hancox RJ, Poulton R, Ely M, et al. Effects of cannabis on lung function: a population-based cohort study. Eur Respir J. 2010;35(1):42-47.
- 21. American Thoracic Society. Standardization of spirometry, 1994 update. Am J Respir Crit Care Med. 1995:152(3):1107-1136.
- 22. Miller MR, Hankinson J, Brusasco V, et al; ATS/

- ERS Task Force. Standardisation of spirometry. Eur Respir J. 2005:26(2):319-338.
- 23. Kelly FJ, Fussell JC. Air pollution and airway disease. Clin Exp Allergy. 2011;41(8):1059-1071.
- 24. Air quality monitoring information: air quality statistics by city, 2009. US Environmental Protection Agency Web site. http://www.epa.gov/airtrends /factbook.html. Accessed November 30, 2011.
- 25. Marrie RA, Dawson NV, Garland A. Quantile regression and restricted cubic splines are useful for exploring relationships between continuous variables. . J Clin Epidemiol. 2009;62(5):511-517, e1.
- 26. Anthonisen NR, Connett JE, Kiley JP, et al. Effects of smoking intervention and the use of an inhaled anticholinergic bronchodilator on the rate of decline of FEV1: the Lung Health Study. JAMA. 1994; 272(19):1497-1505.
- 27. Sherman MP, Roth MD, Gong H Jr, Tashkin DP. Marijuana smoking, pulmonary function, and lung macrophage oxidant release. Pharmacol Biochem Behav. 1991;40(3):663-669.
- 28. Bloom JW, Kaltenborn WT, Paoletti P, Camilli A, Lebowitz MD. Respiratory effects of non-tobacco cigarettes. Br Med J (Clin Res Ed). 1987;295(6612): 1516-1518.
- 29. Sherrill DL, Krzyzanowski M, Bloom JW, Lebowitz MD. Respiratory effects of non-tobacco cigarettes: a longitudinal study in general population. Int J Epidemiol. 1991;20(1):132-137.
- 30. Taylor DR, Poulton R, Moffitt TE, Ramankutty P, Sears MR. The respiratory effects of cannabis dependence in young adults. Addiction. 2000;95(11): 1669-1677
- 31. Moore BA, Augustson EM, Moser RP, Budney AJ. Respiratory effects of marijuana and tobacco use in a U.S. sample. J Gen Intern Med. 2005;20(1): 33-37.
- 32. Aldington S, Williams M, Nowitz M, et al. Effects of cannabis on pulmonary structure, function and symptoms. Thorax. 2007;62(12):1058-1063.
- 33. Tashkin DP, Calvarese BM, Simmons MS, Shapiro BJ. Respiratory status of seventy-four habitual marijuana smokers. Chest. 1980;78(5):699-706.
- **34.** Tashkin DP, Simmons MS, Chang P, Liu H, Coulson AH. Effects of smoked substance abuse on nonspecific airway hyperresponsiveness. Am Rev Respir Dis. 1993:147(1):97-103.
- 35. Tashkin DP, Simmons MS, Sherrill DL, Coulson AH. Heavy habitual marijuana smoking does not cause an accelerated decline in FEV1 with age. Am J Respir Crit Care Med. 1997:155(1):141-148.
- **36.** Tashkin DP, Coulson AH, Clark VA, et al. Respiratory symptoms and lung function in habitual heavy smokers of marijuana alone, smokers of marijuana and tobacco, smokers of tobacco alone, and nonsmokers. Am Rev Respir Dis. 1987;135(1):209-216.