



Change in Neighborhood Characteristics and Change in Coronary Artery Calcium

A Longitudinal Investigation in the MESA (Multi-Ethnic Study of Atherosclerosis) Cohort

Editorial, see p 514

BACKGROUND: Although some evidence shows that neighborhood deprivation is associated with greater subclinical atherosclerosis, prior studies have not identified what aspects of deprived neighborhoods were driving the association.

METHODS: We investigated whether social and physical neighborhood characteristics are related to the progression of subclinical atherosclerosis in 5950 adult participants of the MESA (Multi-Ethnic Study of Atherosclerosis) during a 12-year follow-up period. We assessed subclinical disease using coronary artery calcium (CAC). Neighborhood features examined included density of recreational facilities, density of healthy food stores, and survey-based measures of availability of healthy foods, walking environment, and social environment. We used econometric fixed-effects models to investigate how change in a given neighborhood exposure is related to simultaneous change in subclinical atherosclerosis.

RESULTS: Increases in density of neighborhood healthy food stores were associated with decreases in CAC (mean changes in CAC Agatston units per 1-SD increase in neighborhood exposures, -19.99 ; 95% confidence interval, -35.21 to -4.78) after adjustment for time-varying demographic confounders and computed tomography scanner type. This association remained similar in magnitude after additional adjustment for time-varying behavioral risk factors and depression. The addition of time-varying biomedical factors attenuated associations with CAC slightly (mean changes in CAC per 1-SD increase in neighborhood exposures, -17.60 ; 95% confidence interval, -32.71 to -2.49). Changes across time in other neighborhood measures were not significantly associated with within-person change in CAC.

CONCLUSIONS: Results from this longitudinal study provide suggestive evidence that greater access to neighborhood healthy food resources may slow the development of coronary atherosclerosis in middle-aged and older adults.

Jeffrey J. Wing, PhD,
MPH*

Ella August, PhD*
Sara D. Adar, ScD, MHS
Andrew L. Dannenberg,
MD, MPH

Anjum Hajat, PhD, MPH
Brisa N. Sánchez, PhD
James H. Stein, MD
Matthew C. Tattersall, DO
Ana V. Diez Roux, MD,
PhD, MPH

*Drs Wing and August contributed equally to this article.

Correspondence to: Jeffrey J. Wing, PhD, MPH, Department of Public Health, Grand Valley State University, 545 Michigan St NE, Ste 300 Grand Rapids, Michigan 49503. E-mail wingje@gvsu.edu

Sources of Funding, see page 511

Key Words: atherosclerosis
■ coronary disease

© 2016 American Heart Association, Inc.

Clinical Perspective

What Is New?

- Neighborhood deprivation is thought to affect subclinical atherosclerosis, but prior studies have not identified which aspects of deprived neighborhoods were driving the association.
- This is the first prospective study to investigate associations of changes in neighborhood factors with the progression of subclinical atherosclerosis.
- Using 5950 adult participants from the MESA (Multi-Ethnic Study of Atherosclerosis) during a 12-year follow-up period, this study investigated whether social and physical neighborhood characteristics are related to the progression of subclinical atherosclerosis measured by coronary artery calcium.

What Are the Clinical Implications?

- Our analysis identified that greater access to neighborhood healthy food resources may slow the development of coronary atherosclerosis in middle-aged and older adults.
- Our findings have important implications for the need for policy changes to support the increase in neighborhood healthy food stores.

Neighborhood deprivation, including high neighborhood poverty level, low education level, high unemployment, poor housing conditions, and other indicators, has been linked to greater prevalence and incidence of coronary heart disease.^{1,2} There is some evidence that specific neighborhood features such as fewer neighborhood physical activity resources^{3,4} and less safety^{5,6} and social connectedness⁷ are associated with greater coronary heart disease prevalence, incidence, and mortality, but questions remain as to whether these associations reflect causal processes.

A limitation of focusing on clinical cardiovascular outcomes (especially in the cross-sectional context) is that it is not possible to determine the direction of causality (ie, whether neighborhood characteristics have caused poorer cardiovascular health or poorer cardiovascular health has caused participants to live in certain kinds of neighborhoods). In addition, studies that focus on clinical outcomes are unable to determine whether neighborhood exposures are related to the triggering of cardiovascular events in persons with underlying atherosclerotic disease or to the development of atherosclerosis itself.

A few studies have reported that neighborhood deprivation is associated with greater subclinical atherosclerosis^{8–12}; however, prior studies were not able to identify what aspects of deprived neighborhoods were driving the association. Investigation of specific neighborhood

exposures beyond neighborhood deprivation is of key public health relevance because this approach provides insight into specific pathways for intervention. Furthermore, the investigation of whether changes in neighborhood characteristics are related to changes in subclinical disease strengthens causal inferences.

We investigated whether social and physical neighborhood characteristics are related to the progression of subclinical atherosclerosis in healthy older male and female participants of the MESA (Multi-Ethnic Study of Atherosclerosis) during a 12-year follow-up period. Because a range of neighborhood factors could be important, we examined several characteristics previously hypothesized to be linked to the development of coronary heart disease.¹³ The study design allowed us to investigate how change in a given neighborhood exposure related to simultaneous change in subclinical atherosclerosis.

METHODS

Study Sample

The study sample consisted of 6814 participants from MESA, a longitudinal study of subclinical cardiovascular disease. Participants who were 45 to 84 years of age and free of clinical cardiovascular disease were enrolled in the study from 6 field sites (Baltimore, MD; Chicago, IL; Forsyth County, North Carolina; Los Angeles, CA; New York, NY; and St. Paul, MN) from 2000 to 2002.¹⁴ The study was approved by the institutional review boards at each site, and all participants gave written informed consent. Our analysis used data from 5 MESA examinations spanning 2000 to 2011 and included only data from those who participated in the neighborhood ancillary study (n=6191). We further restricted our sample to include only those that had multiple coronary artery calcium (CAC) measurements, yielding an analytic sample of 5950 participants.

CAC Outcome

We used CAC as a measure of subclinical disease; it has been shown to independently predict coronary events in diverse ethnic groups. CAC was assessed by chest computed tomography (CT) and quantified with the continuous Agatston score, measured in Agatston units.¹⁵ Cardiac CT was performed on all participants at baseline; a second CAC measurement was performed on half of the cohort at examination 2; and the other half of the participants were scanned at examination 3. Additional CAC measurements were taken on a subset of the MESA cohort at examinations 4 and 5. Four percent of the sample had only 1 CAC measurement (and were excluded); 38% had 2, 45% had 3, and 13% had 4 CAC measurements. The median number of years since baseline for the CAC measurement at each examination is as follows: for examination 2, 1.6 years (interquartile range, 1.4–1.8 years); for examination 3, 3.1 years (interquartile range, 3.0–3.4 years); for examination 4, 4.8 years (interquartile range, 4.6–5.0 years); and for examination 5, 9.4 years (interquartile range, 9.1–9.7 years).

Neighborhood Exposure Measurements

The neighborhood environment was characterized with the use of both commercially available food store and recreational facility data, as well as surveys on healthy food, walking, safety, and social cohesion.

Geographic Information Systems-Based Measures: Recreational and Healthy Food Resources

The availability of healthy food stores and recreational facilities within 1 mile of participants' homes was characterized with ArcGIS 9.3.¹⁶ Normal kernel estimation¹⁷ was used to calculate the densities such that facilities closer to participants' homes were given more weight than those farther away. Densities are expressed in units per square mile (henceforth, density indicates units per square mile). Relevant classifications and locations of stores and facilities were identified from Dun and Bradstreet data as compiled by Walls and Associates in the National Establishment Time Series database.¹⁸ Recreational facilities included 114 standard industrial classification codes that classified an establishment as having potential for physical activity or recreation such as indoor conditioning, dance, bowling, golf, team and racquet sports, and water activities; this classification was derived from previous studies.^{19,20} Healthy food stores included fruit and vegetable markets and supermarkets (defined as food stores with at least \$2 million in annual sales or at least 25 employees). Additional data were included from the Nielsen TDLinX Service Supermarket Retail Category Database to further capture area supermarkets as described by Auchincloss and coauthors.²¹ Densities were created for each calendar year between 2000 and 2010.

Survey-Based Measures

Survey-based neighborhood scales included the availability of healthy food (2 items), walking environment (4 items), safety (2 items), and social cohesion (4 items). The participants were asked to rate their neighborhoods within a mile of their homes (the items are listed in Table I in the online-only Data Supplement). These aspects of the neighborhood environment were selected on the basis of conceptual links to cardiovascular disease²² and developed from prior work.^{23,24} Survey data collected from MESA participants were combined with data from participants in an external survey (the Community Survey) administered to other residents of neighborhoods in which MESA participants lived. Scales were based on a 1-mile buffer around the participant's residence and were created by taking the crude mean of the responses for all respondents living within a 1-mile buffer, excluding themselves. Only respondents who answered all questions within the scale were included. Each score has a total possible range from 1 to 5, with a higher score representing a more favorable environment.

Covariates

Time-invariant demographic covariates included highest education level completed, race/ethnicity, and sex, all assessed via questionnaire at the baseline examination. All other covariates were time varying, and for examinations in which a given measure was not assessed, the value from the closest examination

was used. Demographic covariates included age, marital status, family income level, and current employment status. Moderate and vigorous physical activity levels were based on summed minutes of reported relevant physical activities multiplied by the metabolic equivalent of the activity, and cigarette smoking status was classified as current, former, or never. Depressive symptoms were assessed with the 20-item Center for Epidemiological Studies Depression Scale.²⁵ Biomedical factors, measured by physical examination and laboratory tests, included diabetes mellitus, defined as fasting glucose ≥ 126 mg/dL or the use of hypoglycemic medications; hypertension, defined as a systolic blood pressure ≥ 140 mmHg, diastolic blood pressure ≥ 90 mmHg, or taking antihypertensive medication; body mass index (BMI), calculated as weight in kilograms divided by height in meters squared; and ratio of total to high-density lipoprotein cholesterol (both measured from blood samples obtained after a 12-hour fast). Self-reported medications for elevated lipids were also included. CT scanner model was also included as a time-varying covariate because scanner type varied across site and time.

Statistical Methods

We examined the distribution of demographic characteristics and risk factors across levels of baseline CAC (participants with baseline CAC equal to 0 [ie, no discernible calcium on the coronary artery], those with baseline CAC between 0 and 86.5 Agatston units [the median of those with CAC >0], and those with CAC ≥ 86.5 Agatston units). ANOVA was used to compare continuous variables, and χ^2 tests were used for categorical variables. We also examined associations of categories of neighborhood characteristics (based on quartiles) with percent with baseline CAC >0, mean baseline CAC, and mean annual change in CAC after adjustment for age, sex, and race/ethnicity (χ^2 tests, linear tests for trend, and time-interaction terms were used to test these comparisons). On the basis of prior MESA analyses²⁶ and analytic recommendations and given the known robustness of results to violations of the normality assumption,²⁷ CAC was treated as a continuous variable on the native scale to facilitate model fitting and interpretation of associations.

The association of CAC with each of the 6 neighborhood measures (density of recreational facilities, density of healthy food stores, and survey-based measures of availability of healthy foods, walking environment, and social cohesion and safety) was initially examined separately. The survey-based social environment measures (ie, safety and social cohesion) were highly correlated and similarly related to CAC and therefore were combined into a summary social environment index (Cronbach $\alpha=0.77$) by standardizing and then summing their standardized scores. To facilitate comparison across estimates, all neighborhood characteristics were standardized and included in regression models in standard deviation units. Each neighborhood variable was tested in a separate model.

Econometric fixed-effects models²⁸ were used to estimate associations of within-person changes in neighborhood characteristics with within-person changes in CAC. This approach allowed us to examine whether changes in exposures were related to simultaneous changes in the outcome. Because they condition on the study participant, these models have the important advantage of tightly adjusting for measured

and unmeasured time-invariant characteristics of individuals²⁸; thus, they were adjusted only for time-varying covariates. We used a sequence of 4 models that are based on the hypothesized causal pathways linking neighborhood exposures to the outcome (the models progressively adjust for variables more proximal to the outcome and hence more likely to be in the causal pathway). Model 1 adjusted for demographic covariates (age, marital status, income, and working status) and CT scanner type; model 2 additionally adjusted for health behaviors (moderate/vigorous physical activity, cigarette smoking status); model 3 additionally adjusted for depressive symptoms (Center for Epidemiological Studies Depression Scale); and model 4 additionally adjusted for biomedical factors (ratio of total to high-density lipoprotein cholesterol, BMI, hypertension, diabetes mellitus, and lipid-lowering medication). Each of the 5 neighborhood characteristics was investigated separately. Because it has been suggested that neighborhood socioeconomic status may confound associations of neighborhood physical and social environments with health outcomes, additional adjustment for neighborhood socioeconomic status using a summary measure²⁹ was investigated in sensitivity analyses. We also explored effect modification of the neighborhood characteristics by sex and race/ethnicity in an exploratory analysis.

Main analyses included both movers and nonmovers (hence, changes over time in neighborhood exposures may result from moving to a different neighborhood or from changes over time in a neighborhood). In sensitivity analyses, we adjusted for moving by including a time-varying indicator of whether the participant had moved between the prior visit and the current visit and an interaction term between the moving indicator and the neighborhood variable.

Results are presented as the mean difference in CAC for a 1-SD increase in neighborhood score and 95% confidence interval. All statistical tests were 2 sided. Values of $P < 0.05$ were considered statistically significant and were not adjusted for multiple testing. All analyses were conducted in SAS version 9.3 (SAS Institute Inc, Cary, NC).

RESULTS

Of the 5950 participants in our sample, 86% contributed to at least 3 time points, with a mean of 2.7 time points ($SD=0.7$) per participant and a mean follow-up time between CAC measurements of 3.5 years ($SD=3.1$ years). Average CAC in the total sample at baseline was 135 ($SD=380$). Just over half of participants (51%) had a CAC=0 at baseline (Table 1). A nonzero CAC at baseline was associated with being older, male, and white; having a lower income and less education; and being a former smoker. Thirty-two percent of participants moved at some point during the follow-up period.

The percent of participants with CAC>0 at baseline was lower (45%) in the highest quartile of healthy food density than in other quartiles (49%–50%) after adjustment for age, sex, and race/ethnicity (Table 2). Residents of neighborhoods with higher levels of social cohesion and safety were slightly more likely to have CAC>0 at baseline than those of neighborhoods with lower social

cohesion and safety (51% versus 44% for highest versus lowest quartiles) after adjustment.

Mean baseline CAC was lower in neighborhoods with a higher density of healthy food stores and with higher reported availability of healthy foods after adjustment for age, sex, and race/ethnicity, although trends across quartiles were not statistically significant (Table 2). No clear pattern in mean CAC was apparent across quartiles of recreational facility density, the walking environment, or the social environment. Mean annual CAC progressed fastest in neighborhoods with a lower density of recreational facilities, a lower density of healthy food stores (although no dose response was observed), and lower reported availability of healthy foods after adjustment for age, sex, and race/ethnicity. The patterns of mean annual change were inconsistent across quartiles of other survey-based neighborhood characteristics. Of the total variance in CAC, 17% was within-person variance. Of the total variance in neighborhood characteristics, 10%, 5%, 31%, 21%, and 16% were within-person variances for recreational facility density, healthy food store density, availability of health food, walking environment, and social environment, respectively.

Within-person increases in neighborhood healthy food stores density were associated with within-person decreases in subclinical atherosclerosis (Table 3). In model 1, which adjusted for time-varying demographic confounders and CT scanner type, increases in density of neighborhood healthy food stores were associated with decreases in CAC (mean changes in CAC per 1-SD increase in neighborhood exposures, -19.99 ; 95% confidence interval, -35.21 to -4.78). This association remained similar in magnitude after additional adjustment for time-varying behavioral risk factors and depression (see models 2 and 3 in Table 3). The addition of time-varying biomedical factors (model 4) attenuated associations with CAC slightly (-17.60 ; 95% confidence interval, -32.71 to -2.49). Within-person increases in recreational facilities density were associated with decreases in calcium, but associations were not statistically significant. In exploratory analyses, the association of healthy food density with calcium was stronger in women than in men (interaction $P=0.02$), but no effect modification by race/ethnicity was observed.

When survey-based measures were used, changes in availability of healthy foods, in walking environments, and in the social environment were not significantly associated with changes in CAC over time. In sensitivity analyses, associations of healthy food densities with changes in calcium were robust to additional adjustment for neighborhood socioeconomic status. A sensitivity analysis that included adjustment for moving status did not find heterogeneity in estimates by moving status (all interactions $P > 0.47$).

Table 1. Characteristics of Participants by Levels of CAC at Baseline (n=5950), MESA, 2000 to 2002

Characteristic	CAC*=0, %	CAC >0, %	
		CAC<86.5 (Median Value)	CAC≥86.5 (Median Value)
n	3065	1451	1512
Age, mean (SD), y	57.9 (9.1)	63.6 (9.6)	68.6 (8.6)
46–56	43.4	21.6	8.0
57–66	30.5	30.3	21.1
67–76	21.1	33.4	43.2
77–86	5.0	14.8	27.7
Sex			
Female	63.0	47.7	34.5
Male	37.0	52.3	65.5
Race/ethnicity			
White	34.3	39.8	50.8
Chinese	11.7	13.4	10.5
Black	31.2	25.2	20.1
Hispanic	22.8	21.6	18.6
Annual family income, \$			
<12 000	9.3	12.6	12.6
12 000–24 999	18.2	20.2	20.6
25 000–39 999	19.4	18.9	18.5
40 000–74 999	28.6	26.2	25.8
75 000+	24.6	22.2	22.5
Education			
Completed high school/GED or less	32.6	36.4	36.5
Some college, technical or associate degree	29.8	27.4	27.5
Bachelor's degree or higher	37.6	36.3	35.9
Smoking			
Never	56.0	48.2	40.8
Former	31.1	38.9	47.5
Current	12.9	12.9	11.7

CAC indicates coronary artery calcium; GED, General Educational Development; and MESA, Multi-Ethnic Study of Atherosclerosis. *CAC was measured by the Agatston score. The median value for CAC is 86.5 Agatston units.

DISCUSSION

Our study found that increases in the density of healthy food stores around the home were related to simultaneous decreases in subclinical atherosclerosis as characterized by coronary calcium after adjustment for age, marital status, income, working status, and CT scanner type. These associations were reduced slightly but persisted after additional adjustment for changes in moderate/vigorous physical activity, cigarette smoking status, Center for Epidemiological Studies Depression Scale score, ratio of total to high-density lipoprotein

cholesterol, BMI, hypertension, diabetes mellitus, and any lipid-lowering medication. The effect size associated with a 1-SD increase in density was similar to the effect on CAC of a 1-year increase in age (17- to 20-point increase in CAC). Although a number of studies have linked neighborhood characteristics to coronary heart disease incidence and mortality,^{6,7,30,31} we are aware of no studies that have investigated associations of changes in neighborhood factors with the progression of subclinical atherosclerosis. Our results suggest that effects of neighborhood food context on the development of subclinical disease may be one of the mecha-

Table 2. Percent of Participants With CAC>0 at Baseline, Mean (SE) at Baseline, and Annual Change in CAC by Neighborhood Characteristic at Baseline (n=5950)

Characteristic	Percent of All Participants With CAC>0* at Baseline†	Mean (SE)‡ CAC at Baseline Among All Participants§	Mean (SE)‡ Annual Change in CAC Among All Participants
Density of recreational facilities¶			
Quartile 1 (lowest)	48.2	133.3 (7.5)	3.11 (2.20)
Quartile 2	49.3	169.5 (7.3)	0.89 (2.10)
Quartile 3	49.5	163.7 (7.4)	0.22 (2.15)
Quartile 4 (highest)	47.2	140.4 (7.4)	0.35 (2.17)
P value	0.52	0.67	0.32
Density of healthy food stores¶			
Quartile 1 (lowest)	50.1	160.8 (7.5)	3.31 (2.21)
Quartile 2	49.2	154.5 (7.3)	0.19 (2.11)
Quartile 3	50.0	160.5 (7.7)	0.50 (2.24)
Quartile 4 (highest)	44.7	132.4 (7.4)	1.10 (2.15)
P value	0.008	0.02	0.51
Availability of healthy foods#			
Quartile 1 (lowest)	49.9	158.8 (7.7)	4.09 (2.22)
Quartile 2	45.6	164.9 (7.7)	1.33 (2.12)
Quartile 3	49.8	156.6 (7.4)	0.36 (2.23)
Quartile 4 (highest)	48.8	130.0 (7.6)	-0.97 (2.21)
P value	0.07	0.001	0.09
Walking environment#			
Quartile 1 (lowest)	46.7	141.4 (7.7)	-0.30 (2.24)
Quartile 2	49.7	165.9 (7.5)	1.47 (2.19)
Quartile 3	48.3	160.3 (7.4)	2.28 (2.12)
Quartile 4 (highest)	49.4	143.4 (7.6)	1.54 (2.20)
P value	0.35	0.99	0.54
Social environment#			
Quartile 1 (lowest)	44.2	140.4 (7.9)	1.58 (2.27)
Quartile 2	49.2	161.9 (7.5)	-0.14 (2.15)
Quartile 3	50.0	158.7 (7.6)	0.41 (2.23)
Quartile 4 (highest)	50.7	158.3 (7.6)	3.03 (2.19)
P value	0.002	0.25	0.58

CAC indicates coronary artery calcium.

*CAC was measured by the Agatston score.

†P value from χ^2 test.

‡Adjusted for age at baseline, sex, and race/ethnicity.

§P value from linear test for trend.

||P value from interaction term between time by neighborhood variable (as quartiles) in a model with CAC as the outcome variable.

¶Measured with Geographic Information Systems.

Measured by questionnaire.

nisms through which these factors affect the incidence of coronary heart disease.

The use of fixed-effects models allowed us to investigate whether a within-person change in the neighborhood attribute was related to a within-person change in CAC while

tightly adjusting for the time-invariant person characteristics that could confound associations of neighborhood characteristics with CAC. This is a major advance over prior work, which can be subject to between-person confounding. Although residual confounding by time-varying

Table 3. Mean Within-Person Differences in CAC Associated With a 1-SD Within-Person Increase in Selected Neighborhood Characteristics*

Parameter	Mean Difference (95% CI)			
	Model 1†	Model 2‡	Model 3§	Model 4
Geographic information systems measures				
Density of recreational facilities¶	-8.42 (-17.36 to 0.52)	-8.39 (-17.32 to 0.54)	-8.43 (-17.34 to 0.49)	-7.68 (-16.57 to 1.21)
Density of healthy food stores¶	-19.99 (-35.21 to -4.78)	-18.99 (-34.19 to -3.79)	-19.41 (-34.59 to -4.23)	-17.60 (-32.71 to -2.49)
Survey-based measures				
Availability of healthy foods#	6.84 (-0.46 to 14.15)	6.79 (-0.51 to 14.09)	6.84 (-0.45 to 14.13)	6.44 (-0.77 to 13.65)
Walking environment#	5.99 (-2.01 to 14.00)	5.92 (-2.07 to 13.92)	5.82 (-2.16 to 13.80)	4.95 (-2.96 to 12.86)
Social environment#	7.86 (-0.74 to 16.45)	8.00 (-0.59 to 16.59)	7.94 (-0.63 to 16.51)	6.88 (-1.62 to 15.37)

CAC indicates coronary artery calcium; and CI, confidence interval.

*All estimates are derived from fixed-effects models. Each neighborhood characteristic is investigated in a separate model.

†Model 1: adjusted for age, marital status, income, working status, and CT scanner type.

‡Model 2: adjusted as for model 1 plus moderate/vigorous physical activity and cigarette smoking status.

§Model 3: adjusted as for model 2 plus Center for Epidemiological Studies Depression Scale score.

||Model 4: adjusted as for model 3 plus ratio of total to high-density lipoprotein cholesterol, body mass index, hypertension, diabetes mellitus, and any lipid-lowering medication.

¶|Measured with geographic information systems. Higher scores represent more density: 1 SD of density of recreational facilities=8.4 U/sq mile, and 1 SD of density of healthy food stores=4.3 U/sq mile.

#Measured via questionnaire; higher scores represent a more favorable environment: 1 SD of the availability of healthy foods score=0.54, 1 SD of the walking environment score=0.33, and 1 SD of the social environment score=1.65.

characteristics (including socioeconomic factors) cannot be completely ruled out in our study, we adjusted for several time-varying confounders in the models. If moving status is related to changes in calcium and to changes in neighborhood exposures, it could confound the associations of interest. However, adjusting for moving status and allowing associations to differ by moving status did not materially affect our conclusions. A limitation of fixed-effects models is that they rely exclusively on within-person variability and can be inefficient when within-person variability in exposures or outcomes is very low. As expected, within-person variability in neighborhood characteristics was substantially smaller than between-person variability, possibly limiting our ability to detect statistically significant associations with some neighborhood exposures.

Neighborhood food environments may affect cardiovascular risk by shaping dietary intake. Studies have linked food environments with limited choices or a high prevalence of fast food restaurants to poorer diet quality and greater fast food consumption.^{32,33} Greater neighborhood availability of healthier foods has also been linked to higher fruit and vegetable consumption.³⁴ Diets higher in fruits, vegetables, antioxidants, whole grains, and fish have been shown to reduce the progression of subclinical heart disease in prospective observational studies³⁵ and randomized trials,^{36,37} and some evidence from randomized trials shows a reversal of carotid atherosclerosis through adherence to a heart

healthy diet.³⁸ In our study, greater access to healthier foods may have promoted healthier diets and less coronary plaque formation in neighborhood residents. The limited dietary data available in MESA at the time of these analyses limited our ability to examine the mediating effects of diet.

Greater neighborhood availability of recreational resources has been associated with a greater physical activity level in residents,^{39,40} and longitudinal studies have shown that greater physical activity reduces the progression of subclinical coronary disease.^{41,42} Other studies have shown that favorable neighborhood physical activity environments are associated with better cardiometabolic risk factors,⁴³ lower BMI and smaller waist size,⁴⁴ lower risk of type 2 diabetes mellitus,⁴⁵ and decreased risk of coronary events.^{30,31} Although the associations of changes in recreational facilities with changes in CAC were in the hypothesized direction, they were not statistically significant. It could be that recreational facility densities are not important drivers of overall physical activity (which may be more relevant to CAC change than only leisure activity).

Survey-based measures of the neighborhood food, physical activity, and social environments were not significantly related to CAC progression. It has been argued that survey-based measures of neighborhoods may capture information that is not reflected in objective locational data.⁴⁶ The social environment in particular is more readily measured with survey measures. In earlier work,

we documented associations of baseline survey-based measures of the physical environment (especially the food environment) with the incidence of diabetes mellitus and obesity.⁴⁷⁻⁴⁹ An important limitation of survey-based measures in this study, however, is that their time-varying nature was much more limited than it was for the density measures (which were updated every year). In contrast, the survey-based measures relied on substantial interpolation to create time-varying measures. This added measurement error and reduced within-person variability may have seriously limited our ability to detect associations of change with change in the fixed-effects models.

We hypothesized that a favorable social environment would improve stress and depressive symptoms, favorably affecting CAC. Although only 1 identified cross-sectional study has evaluated neighborhood social characteristics in relation to CAC, reporting that better social environment was related to lower CAC,¹¹ other studies have found that a better social neighborhood environment is associated with reduced myocardial infarction incidence.^{6,7,31} We also hypothesized that an improvement in the walking environment would be related to reduced progression of subclinical disease. It has been shown that changes in walking environments are related to change in physical activity and changes in BMI over time.^{50,51} However, these studies relied on objective measures of the built environment features rather than survey measures, as used in the analyses reported here. Studies with improved measurement of the social and walking environments are needed to draw firmer conclusions on the possible effects of these domains on changes in subclinical disease.

Our study has several strengths. It is the first study to examine whether changes in neighborhood characteristics influence the progression of CAC. It included a diverse sample from 6 different sites across the United States. It included detailed time-varying measures of neighborhood environments and state-of-the-art assessments of subclinical atherosclerosis. The study also has several limitations. We used a 1-mile buffer size for our neighborhood measures; however, relevant buffer sizes could be different for different exposures, for example, healthy food versus physical activity versus social engagement, and different distances may be relevant for different individuals.⁵² Additionally, the study does not have information on whether participants actually use the nearby healthy food or physical activity resources, and we did not include other neighborhood environment variables such as green space, esthetic quality, or objectively measured crime rates. Workplace environments may have been more relevant for those participants who worked, and we did not characterize the neighborhood environment around participant workplaces. We explored a range of neighborhood-level variables; however, correlations between variables and power limitations inherent in the fixed-effects approach (which relies only on

within-subject variability) precluded meaningful analyses of their independent effects. We also did not account for multiple testing in our models and comparisons.

The fixed-effects models that we used adjust for measured and unmeasured person-specific covariates related to residential location, but residual confounding resulting from unmeasured time-varying covariates cannot be ruled out. Changes in densities may also be proxying other environmental changes related to subclinical disease. Prior work has documented changes in CAC associated with aging comparable to the strength of the associations that we observed between a 1-SD increase in healthy food availability and changes in CAC.⁵³ Thus, our estimates of effects seem plausible. However, caution should be used in the interpretation of our results as causal, and further replication is needed before major public health implications can be drawn from this study.

The results from this longitudinal study provide new evidence that greater access to neighborhood healthy food resources may slow the development of coronary atherosclerosis in middle-aged and older adults. Our findings support the need to consider neighborhood and environmental interventions in the prevention of cardiovascular disease. Future research should examine the impact on cardiovascular risk of specific interventions such as promoting the location of healthy food stores and how neighborhood characteristics may interact with individual-level factors, including genetic predispositions.

ACKNOWLEDGMENTS

This publication was developed under STAR research assistance agreement No. RD831697 (MESA Air) awarded by the US Environmental Protection Agency. It has not been reviewed formally by the US Environmental Protection Agency. The views expressed in this document are solely those of the authors, and the US Environmental Protection Agency does not endorse any products or commercial services mentioned in this publication.

We thank the other investigators, the staff, and the participants of the MESA study for their valuable contributions. A full list of participating MESA investigators and institutions can be found at <http://www.mesa-nhlbi.org>.

SOURCES OF FUNDING

This work is supported by R01 HL071759 (Dr Diez Roux). This research was supported by contracts N01-HC-95159, N01-HC-95160, N01-HC-95161, N01-HC-95162, N01-HC-95163, N01-HC-95164, N01-HC-95165, N01-HC-95166, N01-HC-95167, N01-HC-95168, and N01-HC-95169 from the National Heart, Lung, and Blood Institute and by grants UL1-TR-000040 and UL1-TR-001079 from the National Center for Research Resources. Dr August (formerly Dr Tomey) was supported by a National Institutes of Health grant K01 AG039554-04.

DISCLOSURES

None.

AFFILIATIONS

From Department of Public Health, Grand Valley State University, Grand Rapids, MI (J.J.W.); Departments of Epidemiology (E.A., S.D.A.) and Biostatistics (B.S.), University of Michigan School of Public Health, Ann Arbor; Departments of Environmental and Occupational Health Sciences (A.L.D.) and Epidemiology (A.H.), University of Washington School of Public Health, Seattle; Department of Medicine, University of Wisconsin School of Medicine and Public Health, Madison (J.H.S., M.C.T.); and Department of Epidemiology and Biostatistics, Dornsife School of Public Health, Drexel University School of Public Health, Philadelphia, PA (A.V.D.R.).

FOOTNOTES

Received November 22, 2015; accepted July 11, 2016.

The online-only Data Supplement is available with this article at <http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIRCULATIONAHA.115.020534/-/DC1>.

Continuing medical education (CME) credit is available for this article. Go to <http://cme.ahajournals.org> to take the quiz.

Circulation is available at <http://circ.ahajournals.org>.

REFERENCES

- Diez Roux AV, Merkin SS, Arnett D, Chambless L, Massing M, Nieto FJ, Sorlie P, Szklo M, Tyroler HA, Watson RL. Neighborhood of residence and incidence of coronary heart disease. *N Engl J Med*. 2001;345:99–106. doi: 10.1056/NEJM200107123450205.
- Winkleby M, Sundquist K, Cubbin C. Inequities in CHD incidence and case fatality by neighborhood deprivation. *Am J Prev Med*. 2007;32:97–106. doi: 10.1016/j.amepre.2006.10.002.
- Mobley LR, Root ED, Finkelstein EA, Khavjou O, Farris RP, Will JC. Environment, obesity, and cardiovascular disease risk in low-income women. *Am J Prev Med*. 2006;30:327–332.
- Pereira G, Foster S, Martin K, Christian H, Boruff BJ, Knuiaman M, Giles-Corti B. The association between neighborhood greenness and cardiovascular disease: an observational study. *BMC Public Health*. 2012;12:466. doi: 10.1186/1471-2458-12-466.
- Augustin T, Glass TA, James BD, Schwartz BS. Neighborhood psychosocial hazards and cardiovascular disease: the Baltimore Memory Study. *Am J Public Health*. 2008;98:1664–1670. doi: 10.2105/AJPH.2007.125138.
- Sundquist K, Theobald H, Yang M, Li X, Johansson SE, Sundquist J. Neighborhood violent crime and unemployment increase the risk of coronary heart disease: a multilevel study in an urban setting. *Soc Sci Med*. 2006;62:2061–2071. doi: 10.1016/j.socscimed.2005.08.051.
- Chaix B, Lindström M, Rosvall M, Merlo J. Neighbourhood social interactions and risk of acute myocardial infarction. *J Epidemiol Community Health*. 2008;62:62–68. doi: 10.1136/jech.2006.056960.
- Dragano N, Hoffmann B, Stang A, Moebus S, Verde PE, Weyers S, Möhlenkamp S, Schmermund A, Mann K, Jöckel KH, Erbel R, Siegrist J; Heinz Nixdorf Recall Study Investigative Group. Subclinical coronary atherosclerosis and neighbourhood deprivation in an urban region. *Eur J Epidemiol*. 2009;24:25–35. doi: 10.1007/s10654-008-9292-9.
- Murray ET, Diez Roux AV, Carnethon M, Lutsey PL, Ni H, O'Meara ES. Trajectories of neighborhood poverty and associations with subclinical atherosclerosis and associated risk factors: the Multi-Ethnic Study of Atherosclerosis. *Am J Epidemiol*. 2010;171:1099–1108. doi: 10.1093/aje/kwq044.
- Petersen KL, Bleil ME, McCaffery J, Mackey RH, Sutton-Tyrrell K, Muldoon MF, Manuck SB. Community socioeconomic status is associated with carotid artery atherosclerosis in untreated, hypertensive men. *Am J Hypertens*. 2006;19:560–566. doi: 10.1016/j.amjhyper.2005.12.008.
- Kim D, Diez Roux AV, Kiefe CI, Kawachi I, Liu K. Do neighborhood socioeconomic deprivation and low social cohesion predict coronary calcification? The CARDIA study. *Am J Epidemiol*. 2010;172:288–298. doi: 10.1093/aje/kwq098.
- Lemelin ET, Diez Roux AV, Franklin TG, Carnethon M, Lutsey PL, Ni H, O'Meara E, Shrager S. Life-course socioeconomic positions and subclinical atherosclerosis in the multi-ethnic study of atherosclerosis. *Soc Sci Med*. 2009;68:444–451. doi: 10.1016/j.socscimed.2008.10.038.
- Diez Roux AV. Residential environments and cardiovascular risk. *J Urban Health*. 2003;80:569–589. doi: 10.1093/jurban/jtg065.
- Bild DE, Bluemke DA, Burke GL, Detrano R, Diez Roux AV, Folsom AR, Greenland P, Jacob DR Jr, Kronmal R, Liu K, Nelson JC, O'Leary D, Saad MF, Shea S, Szklo M, Tracy RP. Multi-Ethnic Study of Atherosclerosis: objectives and design. *Am J Epidemiol*. 2002;156:871–881.
- Agatston AS, Janowitz WR, Hildner FJ, Zusmer NR, Viamonte M Jr, Detrano R. Quantification of coronary artery calcium using ultrafast computed tomography. *J Am Coll Cardiol*. 1990;15:827–832.
- ArcGIS [computer program]. Version 9.3. Redlands, CA: Environmental Systems Research Institute; 2011.
- Silverman B. *Density Estimation for Statistics and Data Analysis*. New York, NY: Chapman and Hall; 1986.
- Walls & Associates. National Establishment Time-Series (NETS) Database: Database Description. 2012. <http://exceptionalgrowth.org/downloads/NETSDatabaseDescription2013.pdf>. Accessed July 23, 2013.
- Powell LM, Chaloupka FJ, Slater SJ, Johnston LD, O'Malley PM. The availability of local-area commercial physical activity-related facilities and physical activity among adolescents. *Am J Prev Med*. 2007;33(suppl):S292–S300. doi: 10.1016/j.amepre.2007.07.002.
- Gordon-Larsen P, Nelson MC, Page P, Popkin BM. Inequality in the built environment underlies key health disparities in physical activity and obesity. *Pediatrics*. 2006;117:417–424. doi: 10.1542/peds.2005-0058.
- Auchincloss AH, Moore KA, Moore LV, Diez Roux AV. Improving retrospective characterization of the food environment for a large region in the United States during a historic time period. *Health Place*. 2012;18:1341–1347. doi: 10.1016/j.healthplace.2012.06.016.
- Diez Roux AV. Residential environments and cardiovascular risk. *J Urban Health*. 2003;80:569–589. doi: 10.1093/jurban/jtg065.
- Echeverria SE, Diez-Roux AV, Link BG. Reliability of self-reported neighborhood characteristics. *J Urban Health*. 2004;81:682–701. doi: 10.1093/jurban/jth151.
- Mujahid MS, Diez Roux AV, Morenoff JD, Raghunathan T. Assessing the measurement properties of neighborhood scales: from psychometrics to ecometrics. *Am J Epidemiol*. 2007;165:858–867. doi: 10.1093/aje/kwm040.
- Radloff L. The CES-D scale: a self-report depression scale for research in the general population. *Appl Psychol Meas*. 1977;1:385–401.
- Kwon Y, Duprez DA, Jacobs DR, Nagayoshi M, McClelland RL, Shahar E, Budoff M, Redline S, Shea S, Carr JJ, Lutsey PL. Obstructive sleep apnea and progression of coronary artery calcium:

- the Multi-Ethnic Study of Atherosclerosis study. *J Am Heart Assoc*. 2014;3:e001241. doi: 10.1161/JAHA.114.001241.
27. Lumley T, Diehr P, Emerson S, Chen L. The importance of the normality assumption in large public health data sets. *Annu Rev Public Health*. 2002;23:151–169. doi: 10.1146/annurev.publhealth.23.100901.140546.
 28. Allison PD. *Fixed Effects Regression Methods for Longitudinal Data Using SAS*. Cary, NC: SAS Press; 2005.
 29. Christine PJ, Auchincloss AH, Bertoni AG, Carnethon MR, Sánchez BN, Moore K, Adar SD, Horwich TB, Watson KE, Diez Roux AV. Longitudinal associations between neighborhood physical and social environments and incident type 2 diabetes mellitus: the Multi-Ethnic Study of Atherosclerosis (MESA). *JAMA Intern Med*. 2015;175:1311–1320. doi: 10.1001/jamainternmed.2015.2691.
 30. Griffin BA, Eibner C, Bird CE, Jewell A, Margolis K, Shih R, Ellen Slaughter M, Whitsel EA, Allison M, Escarce JJ. The relationship between urban sprawl and coronary heart disease in women. *Health Place*. 2013;20:51–61. doi: 10.1016/j.healthplace.2012.11.003.
 31. Kershaw KN, Diez Roux AV, Bertoni A, Carnethon MR, Everson-Rose SA, Liu K. Associations of chronic individual-level and neighborhood-level stressors with incident coronary heart disease: the Multi-Ethnic Study of Atherosclerosis. *J Epidemiol Community Health*. 2015;69:136–141. doi: 10.1136/jech-2014-204217.
 32. Moore LV, Diez Roux AV, Nettleton JA, Jacobs DR, Franco M. Fast-food consumption, diet quality, and neighborhood exposure to fast food: the multi-ethnic study of atherosclerosis. *Am J Epidemiol*. 2009;170:29–36. doi: 10.1093/aje/kwp090.
 33. Boone-Heinonen J, Gordon-Larsen P, Kiefe CI, Shikany JM, Lewis CE, Popkin BM. Fast food restaurants and food stores: longitudinal associations with diet in young to middle-aged adults: the CARDIA study. *Arch Intern Med*. 2011;171:1162–1170. doi: 10.1001/archinternmed.2011.283.
 34. Zenk SN, Lachance LL, Schulz AJ, Mentz G, Kannan S, Ridella W. Neighborhood retail food environment and fruit and vegetable intake in a multiethnic urban population. *Am J Health Promot*. 2009;23:255–264. doi: 10.4278/ajhp.071204127.
 35. Millen BE, Quatromoni PA, Nam BH, O'Horo CE, Polak JF, D'Agostino RB. Dietary patterns and the odds of carotid atherosclerosis in women: the Framingham Nutrition Studies. *Prev Med*. 2002;35:540–547.
 36. Ellingsen I, Seljeflot I, Arnesen H, Tonstad S. Vitamin C consumption is associated with less progression in carotid intima media thickness in elderly men: a 3-year intervention study. *Nutr Metab Cardiovasc Dis*. 2009;19:8–14. doi: 10.1016/j.numecd.2008.01.006.
 37. Niebauer J, Hambrecht R, Schlierf G, Marburger C, Kälberer B, Kübler W, Schuler G. Five years of physical exercise and low fat diet: effects on progression of coronary artery disease. *J Cardiopulm Rehabil*. 1995;15:47–64.
 38. Shai I, Spence JD, Schwarzfuchs D, Henkin Y, Parraga G, Rudich A, Fenster A, Mallett C, Liel-Cohen N, Tirosh A, Bolotin A, Thiery J, Fiedler GM, Blüher M, Stumvoll M, Stampfer MJ; DIRECT Group. Dietary intervention to reverse carotid atherosclerosis. *Circulation*. 2010;121:1200–1208. doi: 10.1161/CIRCULATIONAHA.109.879254.
 39. Ranchod YK, Diez Roux AV, Evenson KR, Sánchez BN, Moore K. Longitudinal associations between neighborhood recreational facilities and change in recreational physical activity in the Multi-Ethnic Study of Atherosclerosis, 2000–2007. *Am J Epidemiol*. 2014;179:335–343. doi: 10.1093/aje/kwt263.
 40. Adams MA, Ding D, Sallis JF, Bowles HR, Ainsworth BE, Bergman P, Bull FC, Carr H, Craig CL, De Bourdeaudhuij I, Gomez LF, Hagströmer M, Klasson-Heggebø L, Inoue S, Lefevre J, Macfarlane DJ, Matsudo S, Matsudo V, McLean G, Murase N, Sjöström M, Tomten H, Volbekiene V, Bauman A. Patterns of neighborhood environment attributes related to physical activity across 11 countries: a latent class analysis. *Int J Behav Nutr Phys Act*. 2013;10:34. doi: 10.1186/1479-5868-10-34.
 41. Pahkala K, Heinonen OJ, Simell O, Viikari JS, Rönkä T, Niinikoski H, Raitakari OT. Association of physical activity with vascular endothelial function and intima-media thickness. *Circulation*. 2011;124:1956–1963. doi: 10.1161/CIRCULATIONAHA.111.043851.
 42. Kozakova M, Palombo C, Morizzo C, Nolan JJ, Konrad T, Balkau B. Effect of sedentary behaviour and vigorous physical activity on segment-specific carotid wall thickness and its progression in a healthy population. *Eur Heart J*. 2010;31:1511–1519.
 43. Müller-Riemenschneider F, Pereira G, Villanueva K, Christian H, Knuiman M, Giles-Corti B, Bull FC. Neighborhood walkability and cardiometabolic risk factors in Australian adults: an observational study. *BMC Public Health*. 2013;13:755. doi: 10.1186/1471-2458-13-755.
 44. Van Dyck D, Cerin E, Cardon G, Deforche B, Sallis JF, Owen N, de Bourdeaudhuij I. Physical activity as a mediator of the associations between neighborhood walkability and adiposity in Belgian adults. *Health Place*. 2010;16:952–960. doi: 10.1016/j.healthplace.2010.05.011.
 45. Booth GL, Creatore MI, Moineddin R, Gozdyra P, Weyman JT, Matheson FI, Glazier RH. Unwalkable neighborhoods, poverty, and the risk of diabetes among recent immigrants to Canada compared with long-term residents. *Diabetes Care*. 2013;36:302–308. doi: 10.2337/dc12-0777.
 46. Diez Roux AV. Neighborhoods and health: where are we and where do we go from here? *Rev Epidemiol Sante Publique*. 2007;55:13–21. doi: 10.1016/j.respe.2006.12.003.
 47. Auchincloss AH, Diez Roux AV, Mujahid MS, Shen M, Bertoni AG, Carnethon MR. Neighborhood resources for physical activity and healthy foods and incidence of type 2 diabetes mellitus: the Multi-Ethnic study of Atherosclerosis. *Arch Intern Med*. 2009;169:1698–1704. doi: 10.1001/archinternmed.2009.302.
 48. Auchincloss AH, Mujahid MS, Shen M, Michos ED, Whitt-Glover MC, Diez Roux AV. Neighborhood health-promoting resources and obesity risk (the Multi-Ethnic Study of Atherosclerosis). *Obesity (Silver Spring)*. 2013;21:621–628.
 49. Christine PJ, Auchincloss AH, Bertoni AG, Carnethon MR, Sánchez BN, Moore K, Adar SD, Horwich TB, Watson KE, Diez Roux AV. Longitudinal associations between neighborhood physical and social environments and incident type 2 diabetes mellitus: the Multi-Ethnic Study of Atherosclerosis. *JAMA Intern Med*. 2015;175:1311–1320. doi: 10.1001/jamainternmed.2015.2691.
 50. Hirsch JA, Moore KA, Barrientos-Gutierrez T, Brines SJ, Zagorski MA, Rodriguez DA, Diez Roux AV. Built environment change and change in BMI and waist circumference: Multi-ethnic Study of Atherosclerosis. *Obesity (Silver Spring)*. 2014;22:2450–2457. doi: 10.1002/oby.20873.
 51. Hirsch JA, Moore KA, Evenson KR, Rodriguez DA, Diez Roux AV. Walk Score® and Transit Score® and walking in the Multi-Ethnic Study of Atherosclerosis. *Am J Prev Med*. 2013;45:158–166. doi: 10.1016/j.amepre.2013.03.018.
 52. Diez Roux AV, Mair C. Neighborhoods and health. *Ann NY Acad Sci*. 2010;1186:125–145. doi: 10.1111/j.1749-6632.2009.05333.x.
 53. Budoff MJ, Young R, Lopez VA, Kronmal RA, Nasir K, Blumenthal RS, Detrano RC, Bild DE, Guerci AD, Liu K, Shea S, Szklo M, Post W, Lima J, Bertoni A, Wong ND. Progression of coronary calcium and incident coronary heart disease events: MESA (Multi-Ethnic Study of Atherosclerosis). *J Am Coll Cardiol*. 2013;61:1231–1239. doi: 10.1016/j.jacc.2012.12.035.

Change in Neighborhood Characteristics and Change in Coronary Artery Calcium: A Longitudinal Investigation in the MESA (Multi-Ethnic Study of Atherosclerosis) Cohort

Jeffrey J. Wing, Ella August, Sara D. Adar, Andrew L. Dannenberg, Anjum Hajat, Brisa N. Sánchez, James H. Stein, Matthew C. Tattersall and Ana V. Diez Roux

Circulation. 2016;134:504-513

doi: 10.1161/CIRCULATIONAHA.115.020534

Circulation is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231

Copyright © 2016 American Heart Association, Inc. All rights reserved.

Print ISSN: 0009-7322. Online ISSN: 1524-4539

The online version of this article, along with updated information and services, is located on the World Wide Web at:

<http://circ.ahajournals.org/content/134/7/504>

Data Supplement (unedited) at:

<http://circ.ahajournals.org/content/suppl/2016/08/09/CIRCULATIONAHA.115.020534.DC1>

Permissions: Requests for permissions to reproduce figures, tables, or portions of articles originally published in *Circulation* can be obtained via RightsLink, a service of the Copyright Clearance Center, not the Editorial Office. Once the online version of the published article for which permission is being requested is located, click Request Permissions in the middle column of the Web page under Services. Further information about this process is available in the [Permissions and Rights Question and Answer](#) document.

Reprints: Information about reprints can be found online at:
<http://www.lww.com/reprints>

Subscriptions: Information about subscribing to *Circulation* is online at:
<http://circ.ahajournals.org/subscriptions/>

SUPPLEMENTAL MATERIAL

Table I. Neighborhood dimensions of the Multi-Ethnic Study of Atherosclerosis Neighborhood Study

Neighborhood Dimension	Question Item
Availability of Healthy Foods	“A large selection of fresh fruit and vegetables is available in my neighborhood” “A large selection of low fat foods is available in my neighborhood”
Safety	“I feel safe walking in my neighborhood day or night” “Violence is a problem in my neighborhood”
Social Cohesion	“People around here are willing to help their neighbors”, “People in my neighborhood generally get along with each other” “People in my neighborhood can be trusted” “People in my neighborhood share the same values”
Walking Environment	“It is pleasant to walk in my neighborhood” “In my neighborhood it is easy to walk to places” “I often see other people walking in my neighborhood” “I often see other people exercise in my neighborhood”

Carolyn:

Welcome to Circulation on the Run. Your weekly podcast summary and backstage pass to the journal and its editors. I'm Dr. Carolyn Lam, associate editor from the National Heart Center and Duke National University of Singapore. I am so pleased to be joined this week by Dr. Judd Hollander and Dr. Deborah Diercks to discuss a problem that all of us, as cardiologists and emergency department physicians will recognize. This is a feature paper on the state of the art approach to the patient presenting to the emergency department with symptoms and signs suggestive of an acute coronary syndrome, but first here are the highlights of this week's issue.

The first study is from first author's Dr. Wing and Dr. August from Grand Valley State University in Michigan who investigated whether social and physical neighborhood characteristics are related to progression of subclinical atherosclerosis measured by coronary artery calcium. They studied this in almost six thousand adult participants of Mesa, a multi-ethnic study of atherosclerosis, followed over twelve years. The main result was that increases in density of neighborhood healthy food stores were associated with decreases in coronary artery calcium. This was significant even after adjusting for time-varying demographic confounders, time-varying behavioral risk factors and depression.

The next study from Dr. Hess and colleagues from the University of Colorado School of Medicine characterized rates of implantable cardioverter defibrillator or ICD counseling and ICD use among more than twenty-one thousand potentially ICD eligible hospitalized heart failure patients in the Get With the Guidelines heart failure program. This study had several notable findings. First, only twenty-two point six percent of patients received ICD counseling. This means that up to four out of five hospitalized heart failure patients, eligible for ICD counseling, did not receive it. Women were counseled less often than men and racial or ethnic minorities were counseled less frequently than white patients.

Among counseled patients, a total of sixty-two point six percent of patients received an ICD or had a documented plan for ICD placement. Women were just as likely as men to receive an ICD, however, ICD use differences by race and ethnicity persisted. The clinical implications of this study are that future quality improvement initiatives should incorporate culturally competent ICD counseling and elevating ICD counseling to a full performance measure and publicly reporting it by sex or race or ethnicity may need to be considered.

The next paper is from first author Dr. Rescovey and corresponding author, Dr. Catalucci and colleagues from the Institute of Genetic and BioMedical Research in Milan, Italy. These authors looked at the voltage dependent [inaudible 00:03:31] calcium channel which is a key mediator of interest [inaudible 00:03:34] calcium entry associated with various cardiovascular conditions such as hypertrophy, atrial fibrillation, hypertension and diabetic cardio myopathy. The author's aim to address the problem that [inaudible 00:03:47] approaches aimed at enhancing calcium current and inotropism in heart failure have also frequently been found to favor arrhythmogenesis and diastolic dysfunction. Thus, limiting their clinical use.

The novel hypothesis addressed in this study is that a peptidome emetic therapeutic approach may overcome the arrhythmogenic limitations of current channel activator inotropes. To test this hypothesis, the author's used a whole host of methods to dissect new regulatory pathways modulating the [inaudible 00:04:24] tight calcium channel life cycle. This included yeast, two hybrid screenings, biochemical and molecular evaluations, protein interaction essays, fluorescence, microscopy, and structural molecular modeling and functional studies. Having uncovered a novel mechanism involving the [inaudible 00:04:44] tight calcium channel, calcium beta two chaperon, the author's then generated a mimetic peptide that specifically targets this calcium beta two chaperon. Thereby controlling the channel assembly and density of the plasma membrane while preserving its physiological channel function.

Finally, they showed that delivery of this mimetic peptide into a mouse model of diabetic cardiomyopathy restored calcium balance and recovered cardiac function. This study is so significant because it provides the proof of concept for the exploitation of novel therapy based on mimetic peptide technology. Really opens the field to mimetic peptides being used as innovative therapeutic tools for the treatment of cardiac disease.

The last study is from Dr. Cammel from the Feil Family Brain and Mind Research Institute in New York and colleagues who studied the association between pregnancy and aortic complications such as dissection or rupture. They used data on all emergency department visits and acute care hospitalizations at nonfederal health care facilities in California and New York between the period of 2005 to 2013. This was a cohort crossover study where they authors defined the period of risk as six months before delivery until three months after delivery. Compared each patient's likelihood of aortic complications during this high risk period to an equivalent control period of two hundred and seventy days exactly one year later.

Among more than six and a half million pregnancies in almost five million women, they identify thirty-six cases of aortic dissectional rupture during the high risk pregnancy period and nine cases during the control period. This gives the rate of aortic complications a five point five per million patients during pregnancy compared to one point four per million during the equivalent period one year later. Thus, pregnancy was associated with a significantly increased risk of aortic dissectional rupture with an incidence rate ratio of four compared to the control period one year later.

Furthermore, absolute risks were particularly elevated in those with a documented diagnosis of hypertension or a connective tissue disease. These findings have clinical implications for the counseling of patients at high base line risk of aortic complications and they also further suggest that clinicians may need to have a lower threshold for initiating diagnostic testing for symptoms of a possible aortic dissection or rupture in pregnant or postpartum patients and especially in those with connective tissue disorders or hypertension.

Our feature paper this week discusses a problem that impacts twenty million patients in North America and Europe every year. What am I talking about? These are patients presenting to the emergency department with symptoms and signs suggestive of an acute coronary syndrome. Who am I talking with? Well, today we have first author Dr. Judd Hollander from Thomas Jefferson University and Dr. Deborah Diercks associate editor from UT Southwestern. Welcome Judd and Deborah.

Dr. Deborah: Thank you.

Dr. Judd: Thank you.

Carolyn: Let's start with a behind the scenes look at this paper. It's an in depth review that was invited by the editorial team. Deborah, can you tell us how this idea came about?

Dr. Deborah: I think one of the goals of the editorial board of circulation is really to provide great clinical reviews that really could benefit the members. I have a unique aspect in that I'm an emergency physician. This idea was really brought about by discussion of really what can we merge cardiology and emergency medicine with. What would be the most clinically issue we're challenged with right now? You can't get two emergency physicians in a cardiologist's room together without some discussion and challenge around the [inaudible 00:09:11].

There's been so many changes in the last decade and so much more information about how we can use these in a clinically relevant way. It really fit nicely into a really great review article and I am really happy that we are able to invite Judd who's well known to the US and one of the leaders in the United States in this area and also an international group inviting a cardiologist from Europe and also an emergency physician from New Zealand to participate in it.

Carolyn: Judd, what is the take home message of this in depth review from your point of view?

Dr. Judd: I think the biggest take home message is we have known for decades and decades that if we rely on our clinical judgement we miss too many patients. We send home people that will be having acute coronary syndromes and acute myocardial infraction and the challenge over the last decades of trying to find ways where we're not going to spend a ton of money over admitting people to the hospital because of a fear of missing an event that may happen five percent of the time.

The beauty of the advances in troponins is we now have troponins that now have increasing sensitivity whether they be the non high sensitivity troponins used in the US or the high sensitivity troponins that are actually used in Europe and the rest of the world. We can use those better [inaudible 00:10:29] and combine them with clinical decision rules to create accelerated diagnostic pathways which is a big term. For now, if we put a blood test together with a structured clinical decision rule, we

can, with more than ninety-nine percent negative predictor value, find patients who are safe to send home.

Carolyn: Judd, I really have to congratulate you on such a beautiful paper. You really did cover all of that but what I love most is the way that you've managed to summarize very clearly a whole wealth of information because when you talk about biomarkers, there's so many out there and there's zero hour, one hour, two hours, this score and that score. I'm actually looking at table one now where you show a summary of the biomarkers strategies and then, in table two, you show a summary of the risk scores and then the performance measures of each of these scores. That must have taken quite a lot to put together.

Dr. Judd: I think that's why Deb was very smart and invited authors from around the world. We have Christian Muller from Switzerland and Martin Tann from New Zealand which, literally, means we're all on different time zones and we were able to work around the clock to do that. There as always somebody awake. Getting more series, the nice thing is that my colleagues on this paper are some of the leaders in doing this kind of research. In fact, they are the leaders in doing this kind of research.

What I think is very challenging for the average cardiologist or the average emergency physician is there have been so many different approaches and many of them actually work. The challenge for us was to try and make it relatively simple so you can choose the approach at your institution and put it into a structured pathway and pick the one that works best for you. You can get a ninety-nine percent negative predicted value using the right essays with samples that the time of presentation and one hour later, you can get a ninety-nine percent negative predictor value at zero and two hours. You can combine it with an accelerated diagnostic pathway and do that at zero and two hours and zero and three hours.

I think the important thing is you need to figure out what will your clinicians use? Certain clinicians may be very comfortable with one risk score and not another and then they need to combine the timing of testing with the risk score their comfortable with in order for us to achieve the great possibilities we have with these new tasks. I think when you try and do a one size fits all, there are going to be people who push back because they don't like one component of the risk score. Really what we're trying to do and we didn't say everybody should do A, B or C but we present the data on five or six different options and let people choose what is most feasible for them.

Carolyn: How wonderful. Deborah, what were you thinking when you were reviewing this paper and trying to structure it for the clinician out there who wants to use this information?

Dr. Deborah: I think that, overall, we were really impressed by the clarity and the ease that a reader can take this information home. There is so much information out there and there are so many different ways to apply it that we're really impressed how the authors put it in a really pretty clear manner so you can actually see the risk

stratification tools that are out there, what they're used with and what type of troponins. Think about your own clinical practice and what you can adapt really based on the evidence that is out there.

Carolyn: I couldn't agree more. Judd, how about this issue of the coronary CT angiogram and where that falls?

Dr. Judd: That's really an interesting question because there's been a lot of publicity and a lot of editorializing in recent years that maybe you can make a decision with your two troponins and your biomarkers and decrease the number of people that need downstream testing. One of the dilemma with this, like I said before, is we know we're not really good at predicting who has acute coronary syndrome based on clinical things and for that reason the European Society guidelines as well as the American AHAACC guidelines have always said you need to do two things. You need to rule out acute myocardial infraction and you need to risk stratify patients for underlying coronary disease. When a patient comes into the emergency department, if I'm going to be guideline compliant with the recommendations in the world, I need to do both things.

The paper, we summarize really clearly ways you can get out of the woods with biomarker testing and clinical pathways but then you still want to risk strategy for coronary disease. There are sometimes where you might not need that downstream testing but what coronary CTA really lets us do is it makes us more efficient than a stress test. A stress test I like to say is a next day test; although there is data that you can do it when the patient's in the emergency department rapidly. It certainly is not the standard practice.

There are people afraid of putting people on the treadmill too soon in case they have unstable angina but a coronary CTA lets me look at the coronary arteries, immediately, when they're in the emergency department. There's very few areas in emergency medicine where there are three large randomized control trials that all give the same results. It doesn't say coronary CTA is better than a next day stress test but it does say you can avoid admission and, hence, save some dollars. It says you can send patients home sooner and, hence, save some angst that the patients may feel while they're in that diagnostic indecision area.

Carolyn: That's such a practical summary and, in fact, it really reflects the entire paper which is really so clearly presenting the information. Judd, one last thing, could I check is this correct, in my understanding, that the main difference between this and say the guidelines that you just measured is that what you do here is really give the readers all the information? As you say, allow the readers to choose what suits them best. This is not making recommendations, it's summarizing all the information. Is that right?

Dr. Judd: Yeah, that's exactly right. If you look, I think it's table number four, where we go through each one of the decision aids and how many or what percent of patients actually fit into that decision aid and what the negative predictive value is for that

decision aid combined with troponin. Then what type of troponin was used to achieve those results, you'll see that about half the studies are done with, what we call, the contemporary troponin or just the regular sensitivity troponin that we use in the United States. The other half of the data we show is with high sensitivity troponins. It would not be a good idea for somebody creating their quality program in their emergency department to take something that was tested with a high sensitivity troponin and validate it there and then apply it in an emergency department in the United States where we don't have those [inaudible 00:17:18].

We thought it was critically important to lay out the data and as the high sensitivity troponins come on the market, hopefully in the next year in the US, people can begin with something now and switch to something else later if they want. If we made a recommendation that was firm, the world changes too fast. I don't think we would be doing the best for our patients.

Carolyn: That is such a great statement to end this on. Thank you so much Judd and Deborah. This was an excellent discussion.

Dr. Judd: Thank you.

Dr. Deborah: Thank you.

Carolyn: You've been listening to Circulation on the Run. Thank you for joining us this week and don't forget to tune in next week for more exciting cardiology needs from all over the world.