Regional distribution of high-attenuation areas on chest computed tomography in the Multi-Ethnic Study of Atherosclerosis

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ABSTRACT High-attenuation areas (HAA) are a computed tomography-based quantitative measure of subclinical interstitial lung disease (ILD). We aimed to validate HAA in lung regions that are less subject to artefacts, such as extravascular lung water or dependent atelectasis. We examined the associations of HAA within six lung regions (basilar, non-basilar, peel, core, basilar peel, basilar core) with serum biomarkers of lung remodelling, forced vital capacity (FVC), visually-assessed interstitial lung abnormalities (ILA), and all-cause and ILD-specific mortality.

We performed cross-sectional and longitudinal analyses of participants in the Multi-Ethnic Study of Atherosclerosis, a prospective cohort of 6814 adults aged 45–84 years without known cardiovascular disease who underwent cardiac computed tomography.

Median regional HAA ranged from 3.8% in the peel to 4.8% in the basilar core. Doubling of regional HAA was associated with greater serum matrix metalloproteinase-7 (range 3.8% to 10.3%; p<0.01), higher odds of ILA (OR 1.42 to 2.20; p<0.03), and a higher risk of all-cause mortality (hazard ratio 1.20 to 1.47; p<0.001). Doubling of regional HAA was associated with greater serum interleukin-6 (4.9% to 10.3%; p<0.005) and higher risk of ILD-specific mortality (hazard ratio 3.30 to 3.98; p<0.001), except in the basilar core. Doubling of regional HAA was associated with lower FVC in the non-basilar, core and basilar core (113 mL to 186 mL; p<0.001).

Associations of HAA with lung remodelling biomarkers, ILA risk and all-cause mortality were consistent across all regions of the lung, including dependent areas where atelectasis may be present. These findings support the validity of HAA as a measure of pathologic subclinical ILD.

Cite this article as: Choi B, Kawut SM, Raghu G, et al. Regional distribution of high-attenuation areas on chest computed tomography in the Multi-Ethnic Study of Atherosclerosis. ERJ Open Res 2020; 6: 00115-2019 [https://doi.org/10.1183/23120541.00115-2019].

This article has supplementary material available from openres.ersjournals.com

Received: 9 May 2019 | Accepted after revision: 4 Dec 2019

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Introduction

The interstitial lung diseases (ILD) are a heterogeneous group of disorders characterised by alveolar injury, inflammation and extracellular matrix remodelling [1, 2]. Adults with subclinical ILD have radiological, histopathological and functional changes that precede the development of clinically evident ILD [3]. Studies of subclinical ILD may lead to improved detective, preventative and treatment strategies for ILD.

Subclinical ILD can be identified using computer-based, automated approaches to detect early interstitial changes in adults without symptoms or a prior diagnosis of ILD [4–7]. High-attenuation areas (HAA) are one example of a validated computer tomography (CT)-derived, quantitative phenotype of subclinical ILD [8–10]. We have previously shown that HAA is independently associated with inflammatory biomarkers, reduced lung function, visually identified interstitial lung abnormalities (ILA) and higher risk of all-cause and ILD-specific mortality [8, 11]. We have used HAA and other measures to identify novel risk factors for ILD, including obstructive sleep apnoea and high-density lipoprotein cholesterol [12, 13].

Given the lower specificity of HAA as a measure of early lung fibrosis compared to other automated approaches, it has been suggested that HAA may be an artefact caused by pulmonary oedema or dependent atelectasis [14–16]. Therefore, in this study, we aimed to validate HAA in lung regions that are less subject to these artefacts. We examined associations of HAA within six lung regions (basilar, non-basilar, peel, core, basilar peel and basilar core) with matrix metalloproteinase (MMP)-7, interleukin (IL)-6, surfactant protein (SP)-A, forced vital capacity (FVC), ILA and all-cause and ILD-specific mortality. MMP-7 and SP-A were chosen as biomarkers of lung remodelling and injury and IL-6 was chosen as a biomarker of systemic inflammation [8, 17–23]. We hypothesised that HAA in the non-basilar regions would be more strongly associated with biomarkers of injury and remodelling, lower FVC, higher odds of ILA, and mortality. Secondarily, we aimed to examine whether HAA in the outer regions of the lung are more strongly associated with ILA.

Methods

Study design and participants

We performed cross-sectional and longitudinal analyses of participants enrolled in the Multi-Ethnic Study of Atherosclerosis (MESA). MESA is a prospective cohort of 6814 adults aged 45–84 year without known cardiovascular disease at baseline, drawn from six communities in the USA. Enrollment criteria have been described previously; there were no selection criteria based on lung disease, respiratory symptoms or smoking history [24]. At the baseline visit (years 2000–2002), participants underwent cardiac CT scans and received a blood draw for measurements of the biomarkers MMP-7, IL-6 and SP-A. Participants were followed longitudinally with contact every 9–12 months and with serial follow-up visits. At the examination 3 and 4 follow-up visits (years 2004–2007), a random sample of participants underwent spirometry. At the examination 5 follow-up visit (years 2010–2012), a subset of participants underwent full-lung CT scans that were then assessed for ILA. The National Death Index (NDI) was reviewed to assess mortality. Death due to ILD was adjudicated as previously described [11].

Regional HAA

HAA was assessed on full inspiratory cardiac CT scans obtained at the baseline visit. The scans were performed twice and the scan with the lowest density was used to assure the best inspiratory effort. HAA was defined as the percentage of voxels with attenuation between −600 and −250 Hounsfield units, as previously described [8]. Regional HAA was calculated for each of the lung regions described below. All participants received complete cardiac CT scans but regional HAA was available for 6671 to 6810 participants (>97%); the missing data were removed for poor quality, incomplete image or excessive artefact.

To define the lung regions, CT scans were divided into three lung zones. These three zones on cardiac CT scans do not correspond with upper, middle (or lingula) and lower lobes of the lungs. Rather, they are the upper, middle and lower thirds of the craniocaudal extent of the cardiac CT scan. Previous studies have shown good correlation between the caudal eighth of cardiac CTs and the caudal third of full-lung CTs for percentage emphysema [25]. The basilar (dependent) region included the caudal third of the imaged lung. The non-basilar region included the remaining two-thirds. The peel region included the outer 20 mm of the lung, and the core region included the remaining inner axial portions. With these definitions, we analysed six distinct axial and craniocaudal lung regions: basilar, non-basilar, peel, core, basilar peel and basilar core (figure 1).

Biomarkers, spirometry and ILA

The biomarkers MMP-7, IL-6 and SP-A were measured from participants’ serum obtained at examination 1 using quantitative sandwich ELISA assays, as previously described [8].
Spirometry was performed at examination 3 according to the American Thoracic Society/European Respiratory Society guidelines, as previously described [26].

ILA was assessed on full-lung CT scans performed at examination 5 (approximately 10 years after the baseline cardiac CTs), as previously described [27, 28]. The CTs were read by one of five trained readers, who scored the presence of ILA if the images contained ground-glass opacity, reticular pattern, nonemphysematous cysts, honeycombing, traction bronchiectasis or centrilobular nodules in at least 5% of nondependent portions of the lung [4, 6, 7]. Indeterminate scans were excluded from analyses.

Statistical analysis

We used generalised linear models to examine the associations of regional HAA with MMP-7, IL-6, SP-A, FVC and ILA. Regional HAA, MMP-7, IL-6 and SP-A were binary log-transformed, and we report effect estimates per doubling of regional HAA. We used Cox hazards models to examine the associations of regional HAA with time to death and ILD death. All models were adjusted for age, sex, race, BMI, height, waist circumference, mAs (milliampere-seconds), regional volume of imaged lung, smoking status, pack-years, regional percentage emphysema, educational attainment, glomerular filtration rate and study site. Due to low event rates, models for ILD death were adjusted for a generalised propensity score (GPS), as previously described [11]. This included the above covariates, as well as alcohol use, total intentional exercise (metabolic equivalent min·wk$^{-1}$), coronary artery calcium, diabetes medication use, insulin use, fasting glucose, hypertension, antihypertensive medication use, systolic and diastolic blood pressures, cholesterol medication use, total and high-density lipoprotein cholesterol levels, C-reactive protein, D-dimer and cancer history. We also performed analyses stratified by BMI category and smoking status. We used likelihood ratio tests to evaluate for effect modification by BMI and smoking status. All analyses were performed using Stata version 14.2 (StataCorp, College Station, TX, USA).

Results

The baseline characteristics of all participants are shown in table 1. Participants had a mean age of 62, 53% were female, 13% were current smokers and 37% were former smokers. Participants had a mean BMI of 28 kg·m$^{-2}$; 39% were overweight (BMI 25–30 kg·m$^{-2}$) and 32% were obese (BMI >30 kg·m$^{-2}$). The median (IQR) regional HAA ranged from 3.8% (3.1–5.0%) in the peel to 4.8% (3.8–6.6%) in the basilar core region (figure 2 and Table S1). Correlation coefficients amongst different areas ranged from 0.68 to 0.90 (Table S2).

Biomarkers

Greater regional HAA was associated with higher serum MMP-7 and IL-6 levels within each lung region and the magnitude of these associations was smallest in the basilar core region (table 2). For each doubling of regional HAA, there was an associated increment in MMP-7 ranging from 3.8% (95% CI 0.8–7.0%) in the basilar core to 10.3% (95% CI 4.7–16.3%) in the core regions, and an associated increment in IL-6 ranging from 1.5% (95% CI −0.9–4.0%) in the basilar core to 10.3% (95% CI 6.5–14.2%) in the non-basilar regions. There were no significant associations of regional HAA with SP-A, except for the core region, where each doubling of HAA was associated with an 8.7% increment (95% CI 0.2–17.8%) in SP-A.
There was evidence of effect modification by smoking status on the association between regional HAA and MMP-7 (p for interaction <0.005 for all lung regions), with stronger associations within each lung region among ever-smokers compared to never smokers (Table S3). There was evidence of effect modification (p for interaction <0.05) by BMI on the association between regional HAA and MMP-7 in the

![Baseline regional high-attenuation areas (HAA).](https://doi.org/10.1183/23120541.00115-2019)

**FIGURE 2** Baseline regional high-attenuation areas (HAA).

| Table 1: Baseline characteristics of Multi-Ethnic Study of Atherosclerosis (MESA) participants |
|-----------------------------------------------|
| **Age years** | 62±10 |
| **Female** | 3600 (53) |
| **Race/ethnicity** |  |
| White | 2622 (39) |
| African American | 1892 (28) |
| Hispanic | 1494 (22) |
| Chinese American | 804 (12) |
| **BMI kg·m^−2** |  |  |
| <25 | 28±5 |
| 25–30 | 32% |
| >30 | 32% |
| **Height cm** | 166±10 |
| **Waist circumference cm** | 98±4 |
| **Glomerular filtration rate** | 81±18 |
| **Smoking** |  |  |
| Never smoker | 50% |
| Former smoker | 37% |
| Current smoker | 13% |
| Data are presented as mean±sd or n (%), unless otherwise stated. All parameters collected at MESA baseline visit in years 2000–2002. BMI: body mass index. |
| Basilar | Non-basilar | Peel | Core | Basilar peel | Basilar core |
|---------|-------------|------|------|--------------|-------------|
| **MMP-7** | | | | | |
| 6.6 (2.6–10.8) | 0.001 | 6.4 (2.5–10.4) | 0.001 | 5.9 (2.2–9.8) | 0.002 |
| 4.9 (1.5–8.4) | 0.005 | 10.3 (6.5–14.2) | <0.001 | 7.3 (4.0–10.6) | <0.001 |
| 4.9 (1.1–11.3) | 0.113 | 2.7 (3.0–8.8) | 0.362 | 5.2 (0.5–11.1) | 0.073 |
| **IL-6** | | | | | |
| 4.9 (1.5–8.4) | 0.005 | 10.3 (6.5–14.2) | <0.001 | 7.3 (4.0–10.6) | <0.001 |
| 4.9 (1.1–11.3) | 0.113 | 2.7 (3.0–8.8) | 0.362 | 5.2 (0.5–11.1) | 0.073 |
| **SP-A** | | | | | |
| 4.9 (1.1–11.3) | 0.113 | 2.7 (3.0–8.8) | 0.362 | 5.2 (0.5–11.1) | 0.073 |
| **FVC** | | | | | |
| −28 (−57–0) | 0.051 | −13 (−151–75) | <0.001 | −26 (−58–5) | 0.099 |
| **ILA** | | | | | |
| 2.20 (1.66–2.90) | <0.001 | 1.54 (1.17–2.02) | 0.002 | 1.79 (1.20–2.30) | <0.001 |
| **Hazard ratio per doubling HAA** | | | | | |
| 1.25 (1.10–1.42) | <0.001 | 1.47 (1.29–1.66) | <0.001 | 1.41 (1.26–1.58) | <0.001 |
| **ILD mortality** | | | | | |
| 3.98 (2.10–7.55) | <0.001 | 3.44 (2.13–5.56) | <0.001 | 3.30 (1.83–5.93) | <0.001 |

Data are presented as mean (95% CI). All analyses adjusted for age, sex, race/ethnicity, body mass index (BMI), waist circumference, smoking status, pack-years, educational attainment, renal function, total volume of imaged lung (not in FVC analysis), percentage emphysema, mA dose and study site. HAA: high attenuation areas; MMP-7: matrix metalloproteinase-7; IL-6: interleukin-6; SP-A: surfactant protein-A; FVC: forced vital capacity; ILA: interstitial lung abnormalities. Adjusted for generalised propensity score, which includes age, sex, race/ethnicity, BMI, waist circumference, smoking status, pack-years, educational attainment, renal function, total volume of imaged lung, percentage emphysema, mA dose, study site, alcohol use, total intentional exercise (metabolic equivalent min·wk<sup>−1</sup>), coronary artery calcium, diabetes medication use, insulin use, fasting glucose, hypertension, antihypertensive medication use, systolic and diastolic blood pressures, cholesterol medication use, total and high-density lipoprotein cholesterol levels, C-reactive protein, a-dimer and cancer history.

https://doi.org/10.1183/23120541.00115-2019
non-basilar, core and basilar core regions. Associations were strongest in the overweight group (BMI 25–30 kg·m$^{-2}$) (Table S4). There was no evidence of effect modification by smoking or BMI on the association between regional HAA and IL-6 or SP-A in any of the lung regions (p for interaction >0.06, data not shown).

**Forced vital capacity**
Higher regional HAA was associated with lower FVC in all lung regions, but this association was not statistically significant in the basilar, peel and basilar peel regions (table 2). The association between HAA and lower FVC was of the greatest magnitude in the core region, where each doubling of HAA was associated with a 186 mL (95% CI 145–226 mL) decrement in FVC.

There was evidence of effect modification by smoking status on the association between regional HAA and FVC, with the association strongest among ever-smokers (Table S3). The effect of smoking status was greatest in the core and least in the basilar regions. There was no evidence of effect modification by BMI on the association between regional HAA and FVC in any of the lung regions (p for interaction >0.10; Table S4).

**Interstitial lung abnormalities**
Greater regional HAA in all lung regions was associated with higher odds of overall ILA (figure 3). Doubling of regional HAA in each region was associated with higher odds of ILA, with OR ranging from 1.42 (CI 1.14–1.76) in the basilar core to 2.20 (CI 1.66–2.90) in the basilar regions (table 2). There was no evidence of effect modification by smoking or BMI on the association between regional HAA and ILA in any of the lung regions (p for interaction >0.10; data not shown).

**Mortality**
Greater regional HAA in all lung regions was associated with increased mortality (figure 4). Doubling of regional HAA in each region was associated with higher hazard ratio of all-cause mortality ranging from 1.20 (CI 1.09–1.32) in the basilar core to 1.47 (CI 1.29–1.66) in the non-basilar regions (table 2). There was no evidence of effect modification by smoking or BMI on the association between regional HAA and mortality in any of the lung regions (p for interaction >0.682, data not shown).

There were 20 ILD deaths in the cohort. Greater regional HAA was associated with increased ILD-specific mortality. Doubling of regional HAA in each region except the basilar core was associated with a higher hazard ratio of ILD death ranging from 3.30 (CI 1.83–5.93) in the core to 3.98 (CI 2.10–7.55) in the basilar regions (table 2).

**Discussion**
High attenuation areas (HAA), the number of lung voxels between $-600$ and $-250$ Hounsfield units, were previously validated as a quantitative measure of subclinical ILD on chest CT. In the current study, we found that regional HAA calculated from portions of the cardiac CTs, divided into regions, was consistently associated with biomarkers of lung injury and inflammation, reduced lung function, visually-assessed interstitial lung abnormalities (ILA), and higher risk of all-cause and ILD mortality. We had hypothesised that HAA in the non-basilar region would be less subject to artefacts such as pulmonary oedema or dependent atelectasis and, therefore, would be more strongly associated with our outcomes. Interestingly, and somewhat contrary to our hypothesis, we found similar associations when comparing HAA in the basilar (dependent) regions to the non-basilar regions and peel compared with core

**FIGURE 3** Odds ratios (OR; with 95% confidence intervals) of interstitial lung abnormalities (ILA) per doubling of high-attenuation areas (HAA).

| Lung region  | n     | p-value |
|--------------|-------|---------|
| Basilar      | 2413  | <0.001  |
| Non-basilar  | 2413  | 0.002   |
| Peel         | 2413  | <0.001  |
| Core         | 2413  | 0.03    |
| Basilar peel | 2412  | <0.001  |
| Basilar core | 2368  | 0.001   |

https://doi.org/10.1183/23120541.00115-2019
This may not be surprising when considering that many of the idiopathic interstitial pneumonias have a basilar predominance [1]. Our findings provide additional support for HAA as a measure of subclinical lung injury, remodelling and inflammation. Given the consistency of our findings across the different lung regions and across multiple measures of subclinical ILD, HAA is less likely to represent significant pulmonary oedema or atelectasis, even when found within the dependent portions of the lung. Furthermore, these findings suggest that even small amounts of HAA detected in just a portion of the lung on cardiac CT have significant implications for the development of ILA and for survival.

The associations of regional HAA with MMP-7, IL-6, ILA and all-cause and ILD-specific mortality were weakest in the basilar core region. This finding may be explained by the fact that this region does not capture typical patterns of ILD found on high-resolution chest CT (HRCT). As seen in figure 1, the basilar core on the cardiac CT correlates with neither the basal nor subpleural portions of the lung where usual interstitial pneumonia (UIP) and nonspecific interstitial pneumonia (NSIP) are typically seen, nor the upper and central portions where hypersensitivity pneumonitis or sarcoidosis may be found [29–31]. The basilar core is also the region most subject to pulmonary oedema.

The associations of regional HAA with MMP-7 and FVC were strongest in the core region of the lung. This was also the only region where we found an association between HAA and SP-A. The core region captures ILDs that have a more central or airway-centred pattern on chest CT, such as hypersensitivity pneumonitis, sarcoidosis and NSIP. Parenchyma that is closer to the bronchioles is also more susceptible to inhalational exposures and aspiration injury, which may be particularly relevant in early disease in many subtypes of pulmonary fibrosis, including idiopathic pulmonary fibrosis (IPF) [14]. Our findings suggest that the core region may be involved in the earliest steps in ILD pathogenesis. The recognition of the \textit{MUC5B} variant as the strongest known risk factor for IPF has led to a new focus on the airways, where \textit{MUC5B} is expressed, in the pathogenesis of pulmonary fibrosis [32]. \textit{MUC5B} overexpression has been shown to lead to impaired mucociliary clearance [33]. Others have also shown that airway abnormalities are associated with ILA and may play a role in early pulmonary fibrosis, but further mechanistic studies are still lacking [34].

On the other hand, regional HAA was most strongly associated with ILA in the basilar, peel and basilar peel portions of the lung. ILA was assessed on full lung CTs performed 10 years after the HAA measurement. ILA is a well-validated measure of subclinical ILD and represents the same visually assessed patterns seen in more advanced ILD [3–7]. Many cases of ILA represent peripheral- and basilar-predominant reticularity and ground glass opacities; more than half have a subpleural predominance [8]. It is reassuring that basilar, peel and basilar peel HAA strongly correlate with ILA. This suggests that, in many cases, the quantitative (HAA) and qualitative (ILA) assessments are measuring the same phenotype.

We found very weak (and statistically non-significant) association between FVC and HAA in the basilar, peel and basilar peel portions of the lung. This may in part reflect variation in lung volume between these regions since the basilar region has a lower lung volume than the non-basilar regions (Table S1), and there are correlations between FVC, lung volume and HAA. However, this does not explain the stronger association in the core region, which has a lower lung volume than the peel region. It is possible that, in the core region, airway collapse and resistance play bigger role in decreasing FVC and contributing to HAA. It is also possible that some of the HAA in the peripheral and basilar regions reflects dependent atelectasis, which does not contribute to reduced FVC. Interestingly, we did not find that BMI modified the association between HAA and FVC, as would be expected if artefact from obesity and compressive atelectasis was a significant contributor to HAA.
We found that smoking was an effect modifier of the associations between HAA, MMP-7 and FVC across all lung regions [8]. Consistent with our prior work, it is likely that HAA, at least in part, reflects smoking-induced lung injury and remodelling, which can affect any lung region. Given that MMP-7 is a biomarker of lung injury, inflammation and fibrosis, it is likely elevated both in participants with subclinical ILD and in smokers [17–19]. Furthermore, as HAA is an early measure of lung injury and remodelling and can be considered a risk factor for ILD, it is likely that smoking has an additive effect on disease pathogenesis.

We found that BMI was an effect modifier of the association between HAA and MMP-7 in most lung regions, with strongest associations among overweight participants and weaker in the obese participants. Interestingly, several studies have also shown an inverse relationship between obesity and MMP-7, with downregulation of MMP-7 expression seen in obese mice [35–37]. The effect modification suggests that increased body weight may contribute to subclinical ILD and lung inflammation, but that obesity has a complex and perhaps even protective effect on subclinical ILD.

There are some limitations to our study. First, we examined both cross-sectional and longitudinal associations of exposures and outcomes collected at various time points. Thus, it is difficult to fully conclude a causative relationship amongst these measurements. Secondly, our study used cardiac CTs, which did not image the whole lung. Furthermore, these were divided into arbitrary regions that do not have anatomical correlations and may not necessarily be matched with the distribution of ILD seen in the clinical setting. Thirdly, because the MESA cohort excluded patients with underlying cardiovascular disease at baseline, the study participants were likely healthier and more ambulatory than the average population. Last, while we accounted for a number of potential confounders in our analyses, there remains the possibility of residual confounding.

**Conclusions**

Associations of HAA with biomarkers of lung remodelling, ILA risk, and all-cause mortality were consistent across all regions of the lung. These findings support the validity of HAA in all lung regions as a measure of subclinical ILD.

Author contributions: B. Choi drafted the initial manuscript. All authors approved the final version of the manuscript. All authors agree to be accountable for all aspects of the work.

Conflict of interest: B. Choi has nothing to disclose. S.M. Kawut reports grants from the NIH during the conduct of the study; travel support from the ATS and the Pulmonary Hypertension Association, grants for CME courses from Actelion, United Therapeutics, Gilead, Lung Biotech, Bayer and Mallinkrodt, and grants and nonfinancial support from the Cardiovascular Medical Research and Education Fund, outside the submitted work. In addition, S.M. Kawut has served in an advisory capacity (for grant review and other purposes) for United Therapeutics, Akros Pharmaceuticals, GlaxoSmithKline and Complexa Inc. without financial support or in-kind benefits. G. Raghur has nothing to disclose for this work E. Hoffman is a founder and shareholder in VIDA Diagnostics, a company commercialising lung imaging analysis software developed, in part, at the University of Iowa. R. Tracy has nothing to disclose. P. Madahar has nothing to disclose. E.J. Bernstein reports acting as a site PI of a clinical trial for Corbus Pharmaceuticals and grants from Pfizer, outside the submitted work. R.G. Barr reports grants from the NIH during the conduct of the study and royalties from UpToDate outside the submitted work. D.J. Lederer reports personal fees for consulting on IPF from Genentech, Pharmakea and Veracyte, personal fees for consulting on IFP and institutional funding for research from Boehringer Ingelheim and Gilead, personal fees for consulting on PF and institutional support for consultation from the Pulmonary Fibrosis Foundation, personal fees for consulting on PDF education and speaking fees from the France Foundation, and personal fees for adjudication panel membership from Degge Group, outside the submitted work. A. Podolanczuk has nothing to disclose.

Support statement: The work was funded by the National Institutes of Health contracts HHSN268201500003I and N01-HC-95159–N01-HC-95169, and grants UL1-TR-000040, UL1-TR-001079, R01-HL077612, RC1-HL100543, R01-HL103676, K24-HL131937, K24-HL103844, K23AR075112 and K23-HL140199. Funding information for this article has been deposited with the Crossref Funder Registry.

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