Heart dose and coronary artery calcification in patients receiving thoracic irradiation for lung cancer

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Background: Thoracic irradiation (TIR) is associated with an increased risk of coronary artery disease (CAD) and coronary-related death. Lung cancer patients receive considerable doses of TIR, making them a high-risk population that may benefit from post-therapy surveillance. Coronary artery calcium (CAC) is a known biomarker of CAD development and may serve as a useful indicator of disease progression in this population. We hypothesized greater CAC progression in lung cancer patients subjected to higher whole heart radiation doses.

Methods: CAC progression (pre- and >2 years post-TIR) from chest CT scans of lung cancer patients were evaluated. A 2:1 matched control population was established controlling for age, gender, race, and CT scan interval. Vessel-specific CAC presence, progression, and extension in pre- and post-interval CT studies was evaluated by two blinded reviewers using the ordinal method. Dosimetric treatment files were restored and contours of the whole heart and proximal left anterior descending artery (LAD) were created within existing plans to compute radiation doses (Pinnacle Treatment Planning Software). Binary logistic regression analysis identified factors predictive for CAC development. Multiple logistic regression analysis with hierarchal method was used to assess covariates.

Results: Thirty-five patients and 65 controls (50% female) were evaluated; mean age 57 years, mean follow-up post-radiation 4.9±2.2 years. Average mean and maximum left anterior descending coronary artery (LAD) radiation doses were 19.9 Gy (95% CI, 14.1–25.7) and 30.7 Gy (95% CI, 23.8–37.5), respectively; 91.6% inter-observer variability. There was greater incidence of coronary calcification in irradiated patients (48.6% vs. 24.6%; P=0.01). In interval CT scans, a greater proportion of radiated patients demonstrated new coronary calcification (P=0.007) and extension within the LAD (P=0.003). Radiation exposure was the only independent predictor of new calcification (OR 3.1; 95% CI: 1.09–9.2).

Conclusions: We identified both an increase in the development and progression of CAC in lung cancer patients receiving TIR. Future studies utilizing alternative cancer populations and larger sample sizes are necessary to further correlate radiographic and dosimetric observations to cardiovascular events.

Keywords: Coronary artery calcification (CAC); thoracic irradiation (TIR); lung cancer

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Introduction

Heart disease remains the primary non-malignant cause of death in cancer survivors, with recent evidence suggesting that up to 50% of cancer patients will develop some degree of cardiovascular complication (1). Thoracic irradiation (TIR) is known to cause microvascular damage and is associated with an increased risk of coronary artery disease (CAD) and coronary related death (2-4). As long-term survival rates and patient lifespan improve, the long-term effects of cancer treatment, specifically cardiotoxicity, are becoming more evident.

Coronary artery calcium (CAC) is associated with coronary plaque accumulation and is a strong predictor of CAD, cardiovascular events, and overall mortality (1-5). CAC scoring has been shown to exceed other generally accepted screening methodologies for assessing cardiovascular disease risk (6). Alluri et al. demonstrated improved health outcomes in patients that received CAC imaging as a CAD screening modality during times of lung cancer surveillance, typically in patients with concerns for recurrence (4).

Despite improvements in modern radiotherapy techniques, cancer patients receiving TIR still sustain considerable doses to surrounding organs, including the heart (7). A study of 2,168 female patients that received radiotherapy for breast cancer demonstrated that the rate of major coronary events was directly related to mean radiation dose to the heart. Each additional gray (Gy) to the heart was associated with a 7.4% increase in coronary events (95% CI, 2.9–14.5, P<0.001) (7). Additionally, the anterior location of the left anterior descending (LAD) coronary artery suggests that radiation to the LAD may also serve as an independent predictor of cardiovascular events (8).

The goal of this study is to characterize the presence, severity and progression of CAC in cancer patients post TIR. We hypothesized that CAC presence, severity and progression is increased in lung cancer patients who received high levels of TIR in comparison to those that had received less radiation or no radiation at all.

Methods

Study design

We conducted a retrospective cohort study with an approximate 2:1 matched control population. All protocols were pre-approved by the institutional review board at Rush University Medical Center (RUMC). Patient group was identified from the RUMC cancer registry and included patients diagnosed with lung cancer between 2007 and 2012, age 18–80 years old; and whom received TIR. We excluded patients that did not receive follow-up care at RUMC, those without serial chest computed tomography (CT) surveillance at RUMC following their lung cancer treatment, patients who died within 2-years of cancer diagnosis.

A 2:1 matched control population was created consisting of lung cancer patients that did not receive therapeutic TIR. Matched parameters included age, gender, race, and comparable CT study time intervals. Hospital records of all the patients were retrospectively reviewed for clinical characteristics and comorbidities including obesity (BMI >30 kg/m²), hypertension, diabetes mellitus, dyslipidemia, and smoking history (9). Additionally, a medication review was conducted, specifically examining for the use of angiotensin converting enzyme inhibitors (ACE-I), beta-blockers, nitrates, aspirin and statins at the time of their first CT date.

CAC quantification

Chest CT scans were obtained using the GE BrightSpeed CT system. Axial acquisitions were obtained through the chest, and multiple reformatted images were generated; including coronal and sagittal multi-planar reformation (MPR) images; coronal and axial maximum intensity projection (MIP) images. The CT scans were performed with attention to patient radiation dose reduction, as low as reasonably achievable (ALARA), while maintaining diagnostic image quality. At least one of the following dose reduction techniques was used: automated exposure control, adjustment of the mA and/or kV according to patient size, use of iterative reconstruction technique. Radiation Shields were used.

High-resolution CT images obtained prior to cancer diagnosis were utilized for baseline CAC measurements. Each patient received 2 CT scans. Post-treatment chest CT studies following TIR were utilized for computation of CAC progression metrics throughout the imaging interval. These post-treatment CT scans were as far apart as possible from at least 2 years, to a maximum time of 7 years apart. CT images for control patients were selected within one year of their study patient counterpart’s pre- and post-interval dates.

Chest CT images were assessed for presence, severity and extension of CAC by 2 independent physician
The Student’s t-test was used to compare quantitative variables between patient group and controls, and continuous data were presented as mean ± SD. The Chi square test or Fisher’s exact test was used to compare qualitative variables, and categorical variables were summarized as percentages. Binary logistic regression analysis was used to identify factors associated with and predictive for development of CAC. Multiple logistic regression analysis with hierarchal method was used to assess the predictive effect of variables like smoking, radiation exposure, dyslipidemia, and duration between CT on CAC development after correction for baseline confounding factors. Interobserver variability coefficient was computed for both independent CAC reviewers. SPSS IBM 23 was used for all statistical analysis and value of <0.05 was considered statistically significant.

**Results**

We identified a total of 35 patients based on set inclusion and exclusion criteria, matched with 65 controls in an approximately 2:1 ratio. The number of participants in the control group were limited in the study and 5 patients were unable to be matched in a 2:1 fashion due to a lack of similar CT time interval. The average mean and maximum radiation doses to the whole heart were 11.61 Gy (95% CI: 8.0–15.2) and 52.1 Gy (95% CI: 41.8–55.9); while the average mean and maximum doses to the proximal LAD were 19.9 Gy (95% CI: 14.1–25.7) and 30.7 Gy (95% CI: 23.8–37.5) respectively. Baseline characteristics including use of known cardio-protective medications for patients and controls are shown in Table 1. The mean age for the study population was 57 years. There were significantly more smokers (P<0.001) and lower BMI in the treatment group vs. controls (26±6.9 vs. 29±8.2; P=0.03). The control group was comprised of more patients utilizing aspirin (P=0.01) and ACE-I (P<0.001).

The presence, extent, and severity of CAC progression in the patient group (post TIR), and controls, are included in Tables 2, 3 and 4, respectively. A higher number of the patient group developed CAC in one or more major coronary arteries compared with controls (48.6% vs. 24.6%; P=0.01) (Table 2). In particular, CAC development was significantly higher in the LAD (P=0.007) and Cx (P=0.01). Additionally, there was significant CAC extension in the patient group for LAD (P=0.003), Cx (P=0.03), aortic arch (P=0.003) and descending aorta (P=0.009), compared with controls (Table 3). Similarly, progression in severity of CAC was significantly increased in the LAD (P=0.005) and Cx (P=0.01) of the patient group, compared with the control group.

**Radiation dosimetry**

Treatment planning records from the patient group were retrieved from MOSAIQ the electronic radiation oncology records database. Dosimetric treatment records were de-archived and restored in the treatment planning system (TPS), Pinnacle (Phillips; Amsterdam, Netherlands). Whole heart contours were created within existing plans and the TIR doses were computed. Heart contours began one slice above the aortic valve, at the approximate level of the ostia, and continued inferiorly to the cardiac apex.

Similarly, proximal LAD contours were generated to compute the respective radiation doses. Specifically, contours were determined by most recent, pre-treatment chest CT scan. Pinnacle CT-to-CT fusion software was used to fuse these images with the simulation CT using a mutual information-based algorithm. The heart/LAD volumes, maximum doses, mean doses and $V_{25%}-V_{70}$ were determined based on the original plan.

**Statistical analysis**

Descriptive statistics were calculated for the study variables. The Student’s t-test was used to compare quantitative variables.
The calculated inter-observer reliability between both CT readers was 91.6%.

Unadjusted logistic regression was used to analyze the characteristics of our study population with newly developed CAC in at least one coronary artery during follow up (Table 5); and showed that radiation exposure (P=0.01) and longer time duration between CT scans (P=0.02) are two independent variables that significantly correlated with development of new CAC. With an adjusted model (Table 6), radiation exposure was the only independent variable that significantly predicted new CAC development (OR 3.1; 95% CI: 1.09–9.2).

Discussion

While CAC based models have been shown to predict future cardiovascular events and mortality beyond the traditional Framingham risk criteria (6), the use of CAC in assessing cardiac risk in cancer patients having undergone cancer therapy remains largely unknown. In our study, we identified both an increase in the presence and progression of both severity and extent of CAC in lung cancer patients receiving TIR. Furthermore, our study showed that after multivariable adjustments, radiation exposure was the only independent variable that significantly predicted new CAC development. A strength of our study is that we evaluated patients with lung cancer (whom usually receive higher doses of TIR), and identified increased development, progression, and severity of CAC in these patients, compared with non-irradiated controls. Our study, unlike others, did not compare baseline CAC data to published values for CAC, but instead utilized a rigorously matched control population without any reported TIR. This not only increased the power of the study, but also allowed us to include a common population with similar exposures. To the best of our knowledge, this is the first study to examine the development, extension and progression of CAC in lung cancer patients exposed to TIR. This data suggests
that despite improvements in TIR techniques, lung cancer patients still receive a radiation dose that may lead to the development of CAC in as little as 2 years after treatment. We particularly utilized lung cancer patients for our study because of the higher radiation dose to the chest and heart, which would allow us observe results in a small number of patients within a shorter period of time.

Micro- and macro-vascular injury in tissues receiving radiotherapy is considered the main cause of radiation-induced heart disease (7). Radiotherapy causes acute endothelial inflammation leading to the mobilization of granulocytes, monocytes, and macrophages that may lead to fibrosis and marked thickening of the adventitia and media of the arteries (8). Eventually this leads to stenosis and calcifications and accelerates progression to unstable atherosclerosis.

We aimed to measure this atherosclerosis by utilizing a visually assessed CAC score. As previously mentioned, calculated CAC scores have been shown to exceed other generally accepted screening methodologies for assessing cardiovascular disease risk (10,11). In comparison to the general population, cancer survivors experience an increase in the risk of coronary arterial atherosclerotic-related events including CAD, angina, and myocardial infarction (12). This begged the question of whether CAC would also develop in cancer patients as a result of TIR;

### Table 2 Presence of CAC after TIR

<table>
<thead>
<tr>
<th>Variables of CAC development in cardiovascular structures</th>
<th>Patient group (N=35)</th>
<th>Control group (N=65)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presence of CAC in one or more coronary arteries</td>
<td>17 (48.6)</td>
<td>16 (24.6)</td>
<td>0.01</td>
</tr>
<tr>
<td>Presence of CAC LM</td>
<td>6 (17.1)</td>
<td>5 (7.7)</td>
<td>0.1</td>
</tr>
<tr>
<td>Presence of CAC LAD</td>
<td>13 (37.1)</td>
<td>9 (13.8)</td>
<td>0.007</td>
</tr>
<tr>
<td>Presence of CAC Cx</td>
<td>8 (22.9)</td>
<td>4 (6.2)</td>
<td>0.01</td>
</tr>
<tr>
<td>Presence of CAC RCA</td>
<td>5 (14.3)</td>
<td>7 (10.8)</td>
<td>0.6</td>
</tr>
<tr>
<td>Presence of CAC in aortic arch</td>
<td>7 (20.0)</td>
<td>9 (13.8)</td>
<td>0.4</td>
</tr>
<tr>
<td>Presence of CAC in descending aorta</td>
<td>10 (28.6)</td>
<td>9 (13.8)</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Table demonstrates CAC presence after TIR in patient group and baseline presence of CAC developed in matched CT interval times in control group that did not receive TIR. Data showed in number (percentage) for patients with new developed CAC (P values for chi-square test). CAC, coronary artery calcification; TIR, thoracic irradiation; Cx, left circumflex coronary artery; LAD, left anterior descending coronary artery; LM, left main coronary artery; RCA, right coronary artery.

### Table 3 Extension of CAC after TIR

<table>
<thead>
<tr>
<th>Cardiovascular structures</th>
<th>Patient group (N=35)</th>
<th>Control group (N=65)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>None</td>
<td>Mild</td>
<td>Mod</td>
</tr>
<tr>
<td>LM</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>29 (82.9)</td>
<td>5 (14.3)</td>
<td>1 (2.9)</td>
<td>0</td>
</tr>
<tr>
<td>LAD</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 (45.7)</td>
<td>9 (25.7)</td>
<td>7 (20.0)</td>
<td>3 (8.6)</td>
</tr>
<tr>
<td>Cx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24 (68.6)</td>
<td>6 (17.1)</td>
<td>4 (11.4)</td>
<td>1 (2.9)</td>
</tr>
<tr>
<td>RCA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>29 (82.9)</td>
<td>5 (14.3)</td>
<td>1 (2.9)</td>
<td>0</td>
</tr>
<tr>
<td>Aortic arch</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21 (60.0)</td>
<td>6 (17.1)</td>
<td>6 (17.1)</td>
<td>2 (5.7)</td>
</tr>
<tr>
<td>Descending aorta</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>17 (48.6)</td>
<td>7 (20.0)</td>
<td>4 (11.4)</td>
<td>7 (20.0)</td>
</tr>
</tbody>
</table>

Table demonstrates CAC extension after TIR in patient group and baseline extension of CAC developed in matched CT interval times in control group that did not receive TIR. Data showed in number (percentage) for patients with progression of CAC extension (P values for chi-square test). Scale for extension of CAC: None, No changes; Mild, 1 degree changes (none to mild; mild to moderate, moderate to severe); Moderate: 2 degree changes (none to moderate, mild to severe); Severe: 3 degree changes (none to severe). CAC, coronary artery calcification; TIR, thoracic irradiation; Cx, left circumflex coronary artery; LAD, left anterior descending coronary artery; LM, left main coronary artery; RCA, right coronary artery.
thereby forming a useful screening tool for this population. Whitlock et al. demonstrated in the large multi-ethnic atherosclerosis (MESA) study, that the development of CAC occurs in many types of cancer patients even after accounting for other atherosclerotic risk factors (13). However, they were unable to show a significant progression of CAC in individuals they found with pre-existing CAC. This led them to question the potential mechanisms for the development of CAC in cancer survivors. In particular, their analysis indicated that the progression of atherosclerosis that can be detected on imaging. It is noteworthy that their study evaluated all comers with cancer who received various treatments. Conversely, our study focused on patients particularly post (high dose) TIR.

Several other studies have investigated the potential impact of cancer and its therapies on CAC. In a cohort of breast cancer patients prior to chemotherapy or radiation, Mast et al. demonstrated increased CAC in middle aged women as compared to age-matched controls (14). Another case series of 47 Hodgkin’s lymphoma patients treated with

Table 4 Progression of CAC severity after TIR

<table>
<thead>
<tr>
<th>Cardiovascular structures</th>
<th>Patient group (N=35)</th>
<th>Control group (N=65)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>None</td>
<td>Mild</td>
<td>Mod</td>
</tr>
<tr>
<td>LM</td>
<td>28 (80.0)</td>
<td>5 (14.3)</td>
<td>0</td>
</tr>
<tr>
<td>LAD</td>
<td>16 (45.7)</td>
<td>11 (31.4)</td>
<td>6 (17.1)</td>
</tr>
<tr>
<td>Cx</td>
<td>23 (65.7)</td>
<td>11 (31.4)</td>
<td>0</td>
</tr>
<tr>
<td>RCA</td>
<td>28 (80.0)</td>
<td>4 (11.4)</td>
<td>3 (8.6)</td>
</tr>
<tr>
<td>Aortic arch</td>
<td>20 (57.1)</td>
<td>10 (28.6)</td>
<td>5 (14.3)</td>
</tr>
<tr>
<td>Descending aorta</td>
<td>15 (42.9)</td>
<td>12 (34.3)</td>
<td>5 (14.3)</td>
</tr>
</tbody>
</table>

Table demonstrates CAC progression after TIR in patient group and baseline progression of CAC developed in matched CT interval times in control group that did not receive TIR. Data showed in number (percentage) for patients with progression of CAC severity (P values for chi-square test). Scale for extension of CAC: None: No changes; Mild: One-degree changes (none to mild; mild to moderate, moderate to severe); Moderate: Two-degree changes (none to moderate, mild to severe); Severe: Three-degree changes (none to severe). CAC, coronary artery calcification; TIR, thoracic irradiation; Cx, left circumflex coronary artery; LAD, left anterior descending coronary artery; LM, left main coronary artery; RCA, right coronary artery.

Table 5 Binary logistic regression analysis to predict new CAC development

<table>
<thead>
<tr>
<th>Variables</th>
<th>B</th>
<th>P value</th>
<th>OR (EXP B)</th>
<th>95% confidence interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time between CTs (years)</td>
<td>0.23</td>
<td>0.02</td>
<td>1.2</td>
<td>1.03–1.5</td>
</tr>
<tr>
<td>Radiation exposure (Gy)</td>
<td>1.06</td>
<td>0.01</td>
<td>2.8</td>
<td>1.2–6.9</td>
</tr>
<tr>
<td>Smoking</td>
<td>1.1</td>
<td>0.059</td>
<td>3.08</td>
<td>0.95–9.9</td>
</tr>
<tr>
<td>Age</td>
<td>−0.02</td>
<td>0.3</td>
<td>0.9</td>
<td>0.91–1.03</td>
</tr>
<tr>
<td>BMI</td>
<td>0.009</td>
<td>0.7</td>
<td>1</td>
<td>0.95–1.06</td>
</tr>
<tr>
<td>Dyslipidemia</td>
<td>0.8</td>
<td>0.057</td>
<td>2.3</td>
<td>0.97–5.5</td>
</tr>
<tr>
<td>DM</td>
<td>0.49</td>
<td>0.3</td>
<td>1.6</td>
<td>0.6–4.2</td>
</tr>
<tr>
<td>HTN</td>
<td>0.09</td>
<td>0.8</td>
<td>0.9</td>
<td>0.3–2.3</td>
</tr>
<tr>
<td>Statin use</td>
<td>0.8</td>
<td>0.07</td>
<td>2.2</td>
<td>0.9–5.4</td>
</tr>
<tr>
<td>ACEI use</td>
<td>0.86</td>
<td>0.05</td>
<td>2.3</td>
<td>1–5.6</td>
</tr>
</tbody>
</table>

CAC, coronary artery calcification; CT, computed tomography; BMI, body mass index; Gy, Gray; DM, diabetes mellitus; HTN, hypertension; ACEI, angiotensin converting enzyme inhibitor.
radiation found abnormally high CAC scores as compared to the published values for CAC (15). A similar earlier series of Hodgkin's lymphoma patients at a median age of 45 years treated with mediastinal radiotherapy approximately 26 years earlier were also reported to have CAC above the 90th percentile for age and gender reference values (16). Despite this evidence supporting the development of CAC at various stages of diagnosis or remission, other studies suggest no changes in CAC. For example, a recent cohort of 236 breast cancer survivors who had undergone radiation and/or chemotherapy compared CAC with published general population CAC scores, and showed a similar age-matched distribution (10). Our study is unique as described below. Unlike these published literatures that have matched patients with historical controls, our study compared a patient group with a carefully selected population of controls, and likely explains our significant results. In our study, both men and women with lung cancer diagnosed between 2007 to 2013 were included if they had survived both diagnoses and treatment for at least 24 months. The reason for this timeline is due to the perceived effects of TIR on the heart. Most evidence estimated aggregate incidence of radiotherapy-induced ischemic heart disease to be 10–30% by 5–10 years post treatment (2). However, a population-based case control study of breast cancer patients by Darby et al. outlined that increased exposure of the heart to ionizing radiation during TIR increased the rate of subsequent ischemic heart disease proportional to the mean dose received by the heart, and can begin within 2 years after exposure and continue for at least 20 years (7). Furthermore, the risk of major coronary events increased linearly with mean dose of radiation to the heart, with the magnitude of the risk increasing by 7.4% per gray (7,17). It should be noted that TIR regimens have changed since the women in these trials were irradiated and the doses of radiation to which the heart is exposed are now generally lower. During Darby's study, the mean dose to the whole heart was 4.9 Gy (range, 0.03–27.72 Gy), compared with 11.61 Gy in our study. These results support Darby's study by showing that higher radiation doses continue to contribute to an increased risk of CAD. Our study utilized the development and progression of CAC as a surrogate for future CAD.

On average, lung cancer patients receiving radiotherapy treatment since 2007 unintentionally receive a 4.1 Gy dose to the heart (18). Considering the high doses of TIR administered to lung cancer patients, this population lends itself to the study of CAC, a biomarker of CAD and predictor of major coronary events (19). Additionally, patients with lung cancer require lifelong surveillance for recurrence at intervals ranging from 3–12 months. Although the images acquired are considered low dose/quality, Dirriches et al. in 2015 was able to adequately evaluate CAC (20). The value of CAC seen in routine screening/surveillance CT scan to predict clinical outcomes was also seen in 3,559 male current or former smokers who underwent a CT chest scan as part a lung cancer screening program (9,21). CAC was associated with increased risk of both fatal and non-fatal cardiovascular events over a median follow up of 2.9 years. The detection of extensive CAC (Agatston >400) in this population was associated with an increased hazard ratio of 12.58 (95% CI: 5.42–29.16) in comparison to patients with no coronary calcification (17,20).

The proximal LAD was of particular interest due to the anatomical localization and reference in previous investigations (21,22). According to Darby et al., the mean heart dose was a better predictor of the rate of major coronary events than the mean dose to the LAD (7). However, despite the historical changes in radiotherapy techniques and heart doses, the LAD lying on the anterior myocardial wall still received an average dose of 20 Gy during breast cancer treatment in the 1980s and has been linked to a risk of ischemic heart disease for years after radiotherapy (8). In our study, the average mean and maximum doses to the proximal LAD were approximately 19.86 and 30.68 Gy,
respectively. These results may indicate relevant late cardiac toxicity and may support routine delineation of the vessel during TIR planning to further avoid non-targeted radiotherapy; particularly because lung cancer patients are living longer with improved targeted cancer therapies.

Lung cancer patients are ideal for detecting CAC due to recommended surveillance for recurrence. Moreover, CAC detection is relatively safe and inexpensive when compared to other common CAD detecting techniques such as a stress echocardiography, coronary CT angiography or stress myocardial perfusion studies (17,19). Specifically, these alternative detection modalities pose a greater risk to patients; often involving higher radiation dose (which may increase the risk of subsequent malignancy), and intravenous contrast (which can induce nephropathy and renal damage) (23). In this study, we showed that we could possibly substitute other more-involved, time-consuming, more expensive detection modalities for simple CAC screening on the already necessary surveillance chest CT scans in lung (and likely other) cancer survivors (24). Of note, 74.3% (26 patients) of the patients undergoing TIR quit smoking by the conclusion of their radiation treatment course.

Our study was limited by the small size of the cohort (n=35), made stronger by a meticulously matched selection of 2:1 controls to increase the power of the study. Additionally, this is a single-center study limiting the variability of the population type and potential external validity. In order to elucidate the effect of TIR, strong exclusion criteria were included such as those who did not have post CT images at least 2 years after initial treatment. This may have introduced selection bias; but if anything, should have diluted our study findings. Furthermore, the current study only included lung cancer patients rather than other cancer patients receiving TIR. Despite these limitations, the adjusted logistic regression analysis did not suggest that there were significant differences between patients and controls in our study. Although the control group did have more dyslipidemia (59.6% vs. 40%) and were receiving more cardiovascular medications including aspirin, ACE-I and beta blocker, these variables were adjusted for in our regression analysis, and TIR still independently predicted new development of CAC. The retrospective nature of our study meant that we were unable to determine a true baseline CAC particularly in the control population; however, our patient and control groups were matched based on time intervals between CT scans. Further study limitations include the use of ordinal visualized CAC scoring instead of gold standard Agatston score. This was necessary to prove our case that routine screening/surveillance chest CT scans could be utilized for CAC assessment in patients post TIR. In addition, prior studies have demonstrated that a simple, visually assessed CAC score was strongly correlated with the Agatston score and had similar predictive values for CAD death (24).

Conclusions

We found increased and newly developed CAC in lung cancer patients exposed to TIR compared with matched controls who were not irradiated. In multiple logistic regression analysis, the strongest independent predictor of new development of CAC was TIR. This study identifies CT-based CAC screening as a potential modality for CAD screening in cancer patients following TIR. Larger trials linking these study findings to major cardiovascular events are required.

Acknowledgments

None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All protocols were pre-approved by the institutional review board at Rush University Medical Center (No. 15041514).

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