Ionizing Radiation in Abdominal CT: Unindicated Multiphase Scans Are an Important Source of Medically Unnecessary Exposure

Kristie M. Guite, MDa, J. Louis Hinshaw, MDa, Frank N. Ranallo, PhDab, Mary J. Lindstrom, PhDc, Fred T. Lee Jr, MDa

Purpose: CT radiation exposure has come under increasing scrutiny because of dramatically increased utilization. Multiphase CT studies (repeated scanning before and after contrast injection) are a potentially important, overlooked source of medically unnecessary radiation because of the dose-multiplier effect of extra phases. The purpose of this study was to determine the frequency of unindicated multiphase scanning and resultant excess radiation exposure in a sample referral population.

Methods: Abdominal and pelvic CT examinations (n = 500) performed at outside institutions submitted for tertiary interpretation were retrospectively reviewed for (1) the appropriateness of each phase on the basis of clinical indication and ACR Appropriateness Criteria® and (2) per phase and total radiation effective dose.

Results: A total of 978 phases were performed in 500 patients; 52.8% (264 of 500) received phases that were not supported by ACR criteria. Overall, 35.8% of phases (350 of 978) were unindicated, most commonly being delayed imaging (272 of 350). The mean overall total radiation effective dose per patient was 25.8 mSv (95% confidence interval, 24.2-27.5 mSv). The mean effective dose for unindicated phases was 13.1 mSv (95% confidence interval, 12.3-14.0 mSv), resulting in a mean excess effective dose of 16.8 mSv (95% confidence interval, 15.5-18.3 mSv) per patient. Unindicated radiation constituted 33.3% of the total radiation effective dose in this population. Radiation effective doses exceeding 50 mSv were found in 21.2% of patients (106 of 500).

Conclusions: The results of this study suggest that a large proportion of patients undergoing abdominal and pelvic CT scanning receive unindicated additional phases that add substantial excess radiation dose with no associated clinical benefit.

Key Words: Radiation, CT, abdomen, pelvis

J Am Coll Radiol 2011;8:756-761. Copyright © 2011 American College of Radiology

INTRODUCTION

CT scanning has become ubiquitous in medicine. Recent technical advances, including faster scan times, improved spatial resolution, and advanced multiplanar reconstruction techniques, have increased the usefulness of CT for virtually every anatomic abnormality. Concomitantly, a rise in defensive medicine and ownership interest in CT centers by referring physicians have resulted in a dramatic increase in utilization [1-3]. Approximately 3 million scans were performed annually in the United States in 1980, and by 2008, that number had grown to 67 million [4]. Along with this increased number of scans, an increasing awareness of medical radiation has permeated the popular and scientific press. More than two-thirds of all medical radiation can now be attributed to CT, with the majority resulting from examinations of the chest, abdomen, and pelvis [5-7].

Although there is no doubt that radiation exposure from CT has been increasing rapidly, the significance of this exposure remains unclear. High levels of ionizing radiation exposure are known to increase cancer risk [8-10], but the data for lower doses of radiation are less clear.
and remain controversial [11-13]. Therefore, in the absence of clarity on this topic, the ACR, Health Physics Society, and other interested organizations have adopted the principle of ALARA, whereby physicians should minimize the amount of radiation exposure to only what is medically necessary [7,14,15].

Most strategies to reduce radiation associated with CT have focused on vetting CT as the appropriate diagnostic test, limiting the examination to the anatomic area in question, and optimizing scanning parameters (particularly in pediatric patients) [2,16-18]. Applying optimized technical parameters alone can decrease radiation exposure by up to 65% [15,16]. However, an important but potentially overlooked source of medically unnecessary radiation is the use of multiphase examinations when a single or lesser number of phases would suffice [16]. The different phases that are possible with state-of-the-art CT scanners are myriad and include scanning before and after contrast administration, delayed imaging, venous and arterial phases, and others. Considering the dose-multiplication effect of extra phases, it is possible that inappropriate multiphase CT could be an important source of excess radiation exposure. Recognizing the need for guidelines addressing multiphase examinations, the College has developed evidence-based ACR Appropriateness Criteria® describing scanning protocols with specific phase selections for various clinical conditions [19].

The purpose of this study was to determine the frequency with which the ACR Appropriateness Criteria for abdominal and pelvic CT are being followed, the frequency with which the ACR Appropriateness Criteria for multiphase examinations when a single or lesser number of phases would suffice [16]. The different phases that are possible with state-of-the-art CT scanners are myriad and include scanning before and after contrast administration, delayed imaging, venous and arterial phases, and others. Considering the dose-multiplication effect of extra phases, it is possible that inappropriate multiphase CT could be an important source of excess radiation exposure. Recognizing the need for guidelines addressing multiphase examinations, the College has developed evidence-based ACR Appropriateness Criteria® describing scanning protocols with specific phase selections for various clinical conditions [19].

The purpose of this study was to determine the frequency with which the ACR Appropriateness Criteria for abdominal and pelvic CT are being followed, the frequency of unindicated phases, and the magnitude of excess radiation exposure for patients when unindicated phases are performed.

**METHODS**

**Selection and Description of Participants**

This study was approved by the human subjects committee of our institutional review board, with a waiver of the requirement for informed consent. The patient group consisted of 708 consecutive abdominal and pelvic CT scans performed at outside institutions during a 4-month period (February 26, 2008, to June 6, 2008) and submitted to our institution for an official “overread.” Excluded were nondigitized images; pelvis-only examinations; specialty examinations, including CT colonography, CT-guided biopsies, and vascular studies; and studies for which the clinical indication was unknown. The final cohort was composed of 500 patients with a median age of 60 years (range, 9 months to 90 years). There were 263 female (53%) and 237 male (47%) patients, with 18 patients aged ≤18 years. The studies were primarily from referring institutions in Wisconsin and Illinois, with a smaller number coming from Michigan, Minnesota, Iowa, Ohio, Florida, Missouri, and Alaska.

**Appropriateness Criteria**

CT examinations were reviewed by one of two experienced abdominal radiologists (F.T.L. or J.L.H.) to determine which phases were indicated for the given clinical indication. ACR Appropriateness Criteria [19] were used as the gold standard. A CT phase was considered to be appropriate (indicated) if the ACR Appropriateness Criteria score was ≥4 (on a scale ranging from 1 to 9, scores of 4 to 6 indicate that studies “may be appropriate”, and scores of 7 to 9 indicate that studies are “usually appropriate”) and unindicated if the score was <4. Each examination that had an unindicated phase or phases was reviewed to determine if there was an incidental finding on the scan that could justify additional scanning for further characterization (eg, incidental liver mass necessitating delayed imaging). If so, these phases were categorized as “unindicated but justified.”

**Technical Information**

**Radiation Effective Dose Calculations.** The clinical history, indication, phases performed, scanning parameters (including CT scanner make and model, tube current, kilovoltage, slice thickness, collimation, rotation time, and pitch), and body part were all recorded. CT scanner models from GE (Milwaukee, Wis), Siemens (Erlangen, Germany), Toshiba (Tokyo, Japan), and Phillips (Andover, Massachusetts) were represented. The collected parameters were used to calculate effective dose for each phase using the ImPACT CT Patient Dosimetry Calculator (version 0.99x 20/01/06), and the effective dose in millisieverts was recorded [20]. For patients with more than one phase, doses were added together to obtain a total dose per patient. Patients with unindicated but justified phases were analyzed with the unindicated group. These patients were initially identified to determine which patients had incidental findings that if noted on the CT scanner could warrant additional phases. However, because it was impossible to determine if these findings were identified before or after the patient left the CT scanner (with the latter thought to be more likely), we analyzed these patients as part of the unindicated group.

Rotation time and pitch were unavailable for 56 of 500 subjects. For these patients, the rotation time and pitch were estimated using the mean values obtained from all other scans that used the same CT scanner model.

**Statistical Analysis**

The distribution of total effective dose, indicated effective dose, and excess effective dose were all skewed, so that a log transformation was necessary to obtain approximate Gaussian distributions. Differences be-
tween effective doses in various groups were assessed using two-sample t tests after transformation to the log scale. All reported means and 95% confidence intervals (CIs) were calculated in the log scale and transformed back to the original units. Comparisons of proportions were done using χ² tests. P values < .05 were considered statistically significant. Mean radiation effective doses (95% CIs) are reported in millisieverts.

RESULTS

Scan Phases
The majority of patients (307 of 500 [61.4%]) had multiphase CT examinations, with 264 of 307 of these patients (86.0%) having at least 1 phase that was not indicated (Table 1). Overall, 264 of 500 of the total patient population (52.8%) had at least 1 unindicated phase, and 350 of 978 of all phases (35.8%) were unindicated. The majority of the unindicated phases were delayed phase imaging (272 of 350 [78%]), with the remainder being a combination of arterial phase (37 of 350 [11%]) and noncontrast imaging (41 of 350 [12%]).

Radiation Effective Dose
The mean effective dose per patient for the entire patient population was 25.8 mSv (95% CI, 24.2-27.5 mSv; range, 3.5-144 mSv), with a mean effective dose per CT phase of 14.1 mSv (95% CI, 13.6-14.7 mSv; range, 2.1-71.0 mSv). In patients who received the correct number of phases (n = 236), the mean effective dose per patient was 17.5 mSv (95% CI, 16.1-19.0 mSv). When unindicated phases were performed (n = 264), the total mean effective dose per phase was 13.1 mSv (95% CI, 12.3-14.0 mSv), and the mean effective dose per patient was substantially higher at 36.5 mSv (95% CI, 34.0-39.3 mSv), with 18.4 mSv (95% CI, 17.0-19.9 mSv) being indicated and 16.8 mSv (95% CI, 15.5-18.3 mSv) being unindicated (Figure 1). The total excess effective dose over all patients was 5,484.2 mSv, and the total effective dose over all patients was 16,449.1 mSv. Thus, 33.3% of the total radiation effective dose to the patient population was due to unindicated phases.

Patients who had unindicated phases had a mean effective dose that was significantly higher than for patients who had appropriate imaging protocols (36.5 mSv [95% CI, 34.0-39.2 mSv] vs 17.5 mSv [95% CI, 16.1-19.0 mSv], P < .001). For patients who had imaging protocols not supported by ACR criteria, a mean of 46.1% (95% CI, 44.4%-47.9%) of their radiation effective dose was attributable to unindicated phases. Notably, 106 of 500 patients (21.2%) received radiation effective doses >50 mSv, and 7 of 500 patients (1.4%) received radiation effective doses >100 mSv for a single examination, levels that are not appropriate in any setting. Patients were more likely to have greater overall radiation effective doses if unindicated phases were performed (add doses, P < .001). No significant difference in the total indicated radiation effective dose was seen between the patients who had unindicated phases and those who had appropriate imaging protocols (P = .37).

### Table 1. Number of phases per patient

<table>
<thead>
<tr>
<th>Number of Phases per Study</th>
<th>Number of Patients (n = 500)</th>
<th>Total Number of Phases</th>
<th>Number of Unindicated Phases</th>
<th>% of Total Phases That Were Unindicated</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>192</td>
<td>192</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>176</td>
<td>350</td>
<td>152</td>
<td>43.4</td>
</tr>
<tr>
<td>3</td>
<td>101</td>
<td>303</td>
<td>146</td>
<td>48.2</td>
</tr>
<tr>
<td>4</td>
<td>23</td>
<td>92</td>
<td>36</td>
<td>39.1</td>
</tr>
<tr>
<td>5</td>
<td>8</td>
<td>40</td>
<td>16</td>
<td>40</td>
</tr>
</tbody>
</table>

Fig 1. The mean total radiation dose for the entire population was 25.8 mSv. The mean dose for patients who received unindicated phases was substantially higher than for patients who received only indicated phases (P < .001). The indicated dose for both groups was not significantly different (P = .37).
who had unindicated phases and those who received only indicated phases (add doses, \( P = .37 \)).

**Radiation Effective Dose by Age**

Patients aged < 50 years were less likely to receive excess radiation (57 of 128 [44.5%]) compared with those aged ≥ 50 years (207 of 372 [55.6%], \( P = .044 \); Table 2). However, excess radiation effective dose was seen in patients in all age groups, and patients aged 30 to 39 years had the highest mean excess radiation. Patients aged 30 to 39 who had unindicated phases had a total mean radiation effective dose of 41.7 mSv (95% CI, 32.5-53.4 mSv), of which 23.2 mSv (95% CI, 27.6-30.5 mSv) was not indicated (Table 2).

**Radiation Effective Dose for Benign and Malignant Indications**

Patients being evaluated for underlying malignancies (ie, known malignancies, palpable abdominal masses, suspicious lesions or carcinomatosis identified on another imaging modality, or painless jaundice) constituted 238 of 500 of the patient population (47.6%). The remaining 262 patients were being evaluated for likely benign indications. Overall, individuals being evaluated for malignancies were significantly more likely to receive excess effective doses than those being evaluated for benign indications (148 of 262 [56.5%] vs 88 of 238 [37.0%], \( P < .001 \)).

**Unindicated But Justified Phases**

Additional scanning for further characterization could be justified on the basis of scan findings in 20 of 307 patients (6.5%). It is not known whether these phases were obtained in response to the findings or if they were performed as standard practice at the referring institutions. The radiation effective dose from these phases represented 3.9% of the total effective dose seen. The incidental findings identified included renal lesions (\( n = 12 \)), unsuspected liver lesions (\( n = 3 \)), and hydronephrosis (\( n = 5 \)).

**DISCUSSION**

The motivation for this study was the anecdotal observation that a large proportion of CT studies submitted to our tertiary center for reinterpretation were performed with multiphase scan protocols that were not appropriate for the clinical indication. Because extra phases effectively multiply radiation dose, we hypothesized that unindicated multiphase examinations were an important source of excess radiation, particularly if extrapolated across larger populations. The results of this study confirm the hypothesis: >50% of patients were exposed to at least 1 unindicated phase, resulting in a mean excess dose of 16.8 mSv of medically unnecessary radiation. The overall mean radiation dose in our population was 25.8 mSv, which far exceeds [21] the national benchmark of 10 to 15 mSv per CT examination [20,22,23], and the majority of this difference was due to unindicated phases. If patients had received only phases indicated by the ACR Appropriateness Criteria, the mean radiation dose would have been 17.9 mSv. This suggests that the high radiation doses seen in this population (and potentially throughout the country) are correctable with simple changes in practice. Conforming to ACR guidelines in this patient population would have reduced the radiation exposure by 46.1% for patients who received inappropriate multiphase examinations or 33.3% for the entire population. Thus, it seems that inappropriate multiphase scanning could be an even more important source of medically unnecessary radiation than nonoptimized technical scanner settings. Prior studies of scanner settings have suggested that a potential decrease in radiation dose of 20% to 65% [15,16] is possible with parameter optimization [2,16-18], but given recent attention in the medical and lay press, much of this improvement has likely already been realized.

Data from experimental models suggest that a carcinogenic risk secondary to high-dose ionizing radiation is real [24,25]. However, the risk at low doses of radiation typical in medical imaging and the threshold required for

---

**Table 2. Age distribution of indicated and unindicated phases and dose**

<table>
<thead>
<tr>
<th>Age (y)</th>
<th>Indicated</th>
<th>Unindicated</th>
<th>% Indicated Phases</th>
<th>Mean Dose per Patient (mSv)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Indicated</td>
<td>Excess*</td>
<td></td>
</tr>
<tr>
<td>0-9</td>
<td>15</td>
<td>5</td>
<td>68.4</td>
<td>8.5</td>
</tr>
<tr>
<td>10-19</td>
<td>11</td>
<td>2</td>
<td>63.2</td>
<td>11.8</td>
</tr>
<tr>
<td>20-29</td>
<td>14</td>
<td>9</td>
<td>60.9</td>
<td>18.3</td>
</tr>
<tr>
<td>30-39</td>
<td>34</td>
<td>24</td>
<td>67.8</td>
<td>25.5</td>
</tr>
<tr>
<td>40-49</td>
<td>81</td>
<td>39</td>
<td>71.8</td>
<td>27.3</td>
</tr>
<tr>
<td>50-59</td>
<td>149</td>
<td>82</td>
<td>64.9</td>
<td>27.4</td>
</tr>
<tr>
<td>60-69</td>
<td>184</td>
<td>108</td>
<td>62.4</td>
<td>27.3</td>
</tr>
<tr>
<td>70-79</td>
<td>89</td>
<td>50</td>
<td>60.1</td>
<td>26.3</td>
</tr>
<tr>
<td>80-89</td>
<td>50</td>
<td>31</td>
<td>62.4</td>
<td>27.5</td>
</tr>
<tr>
<td>≥90</td>
<td>1</td>
<td>0</td>
<td>100.0</td>
<td>26.0</td>
</tr>
<tr>
<td>All patients</td>
<td>628</td>
<td>350</td>
<td>64.2</td>
<td>25.8</td>
</tr>
</tbody>
</table>

*The mean excess dose was calculated using patients with excess dose only.
carcinogenesis remain unclear [10,11]. The linear-no-threshold model championed by some authors suggests proportionality between dose and cancer risk (even at very low doses) and represents the worst possible scenario for low-dose exposure [26,27]. There is serious doubt about the validity of the linear-no-threshold theory [10,28], but it is often quoted as a means to estimate cancer risk for patients exposed to low-dose radiation [29]. There are several studies that argue against the linear-no-threshold model, and no clear evidence for increased cancer risk with low-dose radiation exposure has been identified to date [30-32]. However, in light of the current uncertainty and the unknown “true” risk for low-dose radiation, the concept of ALARA should be followed. Notably, 21.2% of the patients in this study received ≥50 mSv, and 1.4% received ≥100 mSv for a single examination, levels that meet or exceed thresholds for increased cancer risk proposed by various studies and are certainly higher than is acceptable by any current published standard for diagnostic CT [8-11]. Interestingly, the patients with doses >100 mSv all had at least 1 unindicated phase.

Although younger patients being evaluated for benign indications were less likely to receive excess phases, these patients still received large amounts of medically unnecessary radiation. Overall, the greatest excess radiation dose was in patients aged 30 to 39 years. This is important because excess radiation in younger patients has a higher potential for adverse outcomes than in older adults [33] because of more radiosensitive tissues [8] and a longer life span with more opportunities for any radiation-induced genetic mutations to be expressed as neoplasia. Although the retrospective nature of this study did not allow us to investigate the reasons behind the performance of inappropriate multiphase examinations, we suspect that a lack of focus on performing protocols tailored for the individual patient and clinical indication is the predominant factor, with most patients being prospectively scanned with extra phases to reduce callbacks. On the basis of the results of this study, the risk for callbacks seems to be negligible, with a paucity of cases in which inappropriate extra phases were justified. In addition, it is likely that if given the choice, most patients would accept the inconvenience of a callback for additional scanning rather than routine exposure to unindicated series. As professional societies, accrediting bodies, insurers, and health care institutions increasingly use radiation exposure as a measure of health care quality, the routine use of multiphase examinations will become increasingly difficult to defend. Already, radiologists are being asked to dictate radiation exposure into reports as a pay-for-performance measure [34]. However, currently we are unaware of payments for inappropriate multiphase examinations being systematically denied because of excess radiation exposure (excluding contrast and non-contrast phases, for which an inappropriate increase in charges could be an issue). The results of this study make it clear that there is substantial room for improvement, and specialty societies can aid in this effort by providing parameters and protocols that set diagnostic dose references, radiation safety education beginning in medical school, and other tools to support optimization efforts [26].

This study had several limitations. The small sample size and limited geographical area make it difficult to be certain that our results can be extrapolated across the entire US population. Because of the small number of pediatric patients in the study, strong conclusions about excess radiation exposure in children caused by multiphase CT are not possible. Perhaps this issue could be better addressed with a similar study based at a large children’s hospital.

An additional limitation of our study is a possible selection bias toward highly ill patients on the basis of the referral to a tertiary care center. On the basis of a nonquantitative evaluation of the patient population, this does not seem to be the case. The most frequent indications for scanning in our study were relatively routine: abdominal pain (50 of 500), flank pain (41 of 500), follow-up of prostate cancer (24 of 500), and follow-up of colon cancer (19 of 500).

An additional potential limitation is that this study used the ACR Appropriateness Criteria as the sole adjudication method to determine the appropriateness of each CT phase. Although it is our belief that these criteria are the most widely accepted guidelines available for this purpose, they do not cover every clinical situation, and they are not being continuously updated. In terms of radiation dose calculators, an exhaustive description of the various methods to calculate radiation dose is beyond the scope of this study. However, the methods used in this study have been previously validated, have been used in many peer-reviewed publications, and are widely accepted [35]. A different calculator may have resulted in different overall radiation exposures, but the relative impact of medically inappropriate scanning would likely not be changed.

CONCLUSIONS

In summary, our study suggests that a large proportion of patients who undergo abdominal and pelvic CT scanning receive medically unnecessary multiphase examinations, resulting in substantial excessive radiation exposure. This source of excess radiation seems almost entirely correctable by widespread adoption of individual scan protocling tailored specifically for the patient’s clinical condition and guided by the ACR Appropriateness Criteria or other evidence-based criteria. Last, the routine use of “one-size-fits-all” multiphase protocols for abdominal and pelvic CT should cease immediately.
REFERENCES

17. Paterson A, Frush D, Donnelly LF. Helical CT of the body: are settings adjusted for pediatric patients? Am J Radiol 2001;176:297-301.