Diagnostic Performance of a prototype Dual-Energy Chest Imaging System

by

Hany Mehdizadeh Kashani

A thesis submitted in conformity with the requirements for the degree of Master of Science
Institute of Medical Science
University of Toronto

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Abstract

**Purpose:** To assess the performance of a Dual-Energy chest radiography system.

**Methods:** A cohort of 129 patients was recruited from population referred for CT guided biopsy of a lung lesion. Digital radiography (DR) and Dual Energy (DE) images were acquired. Receiver operating characteristic (ROC) tests were performed to evaluate performance of DE images compared to DR. Five chest radiologists scored images. Performance was analyzed for all cases pooled and sub groups based on gender, nodule size, density, location, and chest diameter.

**Results:** There was no significant difference between DE and DR for all cases (p = 0.61). There was a significant advantage for DE imaging of small nodules, and nodules located in right-upper lobe. (p = 0.02 and 0.01)
Conclusions: DE imaging demonstrated significant improvement in diagnosis of sub-centimeter lung nodules and lesions in the upper lung zones which are common characteristic of early stage lung malignancy.
TO ALL THAT I DEARLY LOVE,

.........
Acknowledgments

I once read a statement by Elwyn Brooks White to which I felt a deep connection:

“I arise in the morning torn between a desire to improve the World and a desire to enjoy the world. This makes it hard to plan the day.”

And I wondered……., is there a way to enjoy and improve the World at the same time?

I have found my passion, in research, in the field of Medical Imaging. I believe that research plays an important role in improving and evolving technology. There is a keener awareness of potential health risks, even from small doses of radiation, than ever before, however, increased utilization of imaging modalities has ignited a grand revolution in detection and management of a wider spectrum of pathology. One of the most important aims for researchers in this field is to create techniques with less radiation exposure and better image quality, and that is what I have and will be involved with, something that makes me feel quite useful. Of course, I have two strong-minded men to thank for it, Dr. Narinder Paul and Dr. Jeff Siewerdsen and it has been an honor to be their pupil.

Jeff, I would like to thank you for trusting me with the DE project even though my engineering and physics experience were close to zero. It was very exciting to be in charge of the Dual Energy X-ray system, to optimize the images, perform clinical trials and to finally see the results of our experiment. I became familiar with Dual Energy innovative technology, troubleshooting, programming, Physics, Math and much more. A whole new World, so different from what I knew before and I absolutely fell in love with it. I also like to thank you for your support and guidance throughout this project that has given me the opportunity to win scientific prizes, publish and present our work in peer reviewed scientific journals and conferences.

Narinder, you have been my guiding light and my mentor since that fateful day in October of 2004 when I stepped through the doors of Toronto General Hospital. I was very
confused but your nice attitude took away all my anxiety and gave me peace of mind and confidence. I was also quite amazed by all the advanced imaging modalities in your Department as well as your research projects involving low dose and very low dose CT. Suddenly it occurred to me! I would like to be a part of all this innovation, creativity and improvement. You are the reason that I could find my true path in life and I thank you for that. Thank you for teaching me, trusting me and motivating me. Your enthusiasm, discipline and eagerness to improve the world of Imaging have been quite encouraging for me. Your help and guidance made it possible for me to achieve a lot and I owe you the success in all my endeavours.

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Chapter 1
Introduction

1 Anatomy

1.1 The lungs

There are two lungs located on each side of the thoracic cavity and covered by the rib cage and separated by mediastinal structures which include the heart. The superior boundary of the lungs is located just above the first rib, inferiorly the lungs rest on the diaphragm which separates the chest from the abdominal cavity. (Fig 1.1) The lungs are covered by two thin membranes: the parietal and visceral pleura. The visceral pleura covers the lungs directly and the parietal pleura covers the inside of the chest wall. The hilum of each lung is the point of attachment for the root of the lung. It contains the bronchi, pulmonary, bronchial and lymphatic vessels, and nerves.
The right lung is divided by the oblique and horizontal fissure into three lobes: upper, middle and lower. The left lung contains only the oblique fissure that divides the lung into upper and lower lobes. The lingua of the upper lobe corresponds to the middle lobe of the right lung. Each lobe consists of multiple segments and each segment has multiple lobules.

1.2 The Lung Interstitium

The lung interstitium is a network of connective tissue fibers with several components that support the lung. These components include: peribronchovascular interstitium, centrilobular interstitium, interlobular interstitium, interlobular septa, and sub-pleural interstitium.

The peribronchovascular interstitium is a system of fibers that surrounds bronchi and pulmonary arteries to form a strong supportive sheath of connective tissue. The subpleural interstitium invests the lungs in a fibrous sac and is located deep to the visceral pleura. From this fibrous sac, connective tissue septa penetrate into the lung parenchyma.

The interlobular septum is part of these septa. The intralobular interstitium consists of thin fibers that form a mesh in the alveolar walls, connecting the centrilobular interstitium located within the centre of lobules to the interlobular septa and subpleural interstitium in the lobular periphery. These fibers integrate to form a protective fibrous skeleton for the lungs.

1.3 Arterial and Venous Supply

The right and left pulmonary arteries arise from the pulmonary trunk. The pulmonary arteries deliver deoxygenated blood to the lungs from the right side of the heart (Fig1.3. a). The bronchial arteries arise from the descending thoracic aorta to supply the bronchi and the non-
respiratory portion of the lung (Fig1.3. b). There are four pulmonary veins: superior right and left and inferior right and left. The pulmonary veins carry oxygenated blood to the left atrium of the heart (Fig1.3. a). The bronchial veins drain to the azygos system. They share drainage from the bronchi with the pulmonary veins. (Fig1.3. b)

(a) Pulmonary circulation.

Modified from http://webschoolsolutions.com/patts/systems/lungs.htm
(b) Bronchial circulation.

Modified from http://webschoolsolutions.com/patts/systems/lungs.htm

Figure 1.2 (a, b) Pulmonary and bronchial circulation of the respiratory system. Pulmonary arteries carry the deoxygenated blood from the right side of the heart and to the lung. Pulmonary veins carry oxygenated blood to the left side of the heart. (a) The bronchial arteries supply bronchi and the non-respiratory part of the lungs. They drain to the azygos. (b)
2 Histology

The respiratory system is divided into an air conducting portion, including the nasal cavity, pharynx, larynx, trachea, bronchi and bronchioles, that carries the gases during inspiration and expiration, and a respiratory portion, including alveoli, that provides for gas exchange between air and blood. The trachea leads to the main terminal and respiratory bronchioles. In the transition from the trachea to the respiratory bronchioles, the epithelium changes from pseudostratified ciliated columnar cells to simple columnar cells. (4)

Terminal bronchioles lead to respiratory bronchioles that contain alveoli and branches to form two to three alveolar ducts, which are long sinuous tubes that often terminate in alveolar sacs. Alveolar sacs are spaces formed by two or more conjoined alveoli. They are lined by simple squamous alveolar epithelia. Alveoli are the terminal thin-walled sacs of the respiratory tree that are responsible for gas exchange. There are approximately 300 million alveoli per lung, each of which is 20 to 30 mm in diameter. (5)

The alveoli epithelium contains two cell types. Type I cells cover almost the entire alveolar luminal surface and provide a thin surface for gas exchange. Type II cells are cuboidal-like cells that sit on the basal lamina of the epithelium and contain membrane-bound granules of phospholipids and protein. These contents are secreted onto the alveolar surface to provide a coating of surfactant that reduces alveolar surface tension. (6) (Fig 1.4)
Figure 1.3 Structure of alveolus. The alveoli epithelium contains two different cell types, type I and type II. Type I cells are responsible for gas exchange and type II secrete the surfactant coating to reduce surface tension.

Modified from herkules.oulu.fi/isbn9514270584/html/c273.html

3 Epidemiology

3.1 Lung Cancer

Cancer is the leading cause of death worldwide, accounting for 7.4 million deaths or around 13% of all deaths in 2004. (7) Among the different types of cancer, lung cancer remains the leading cause of cancer-related deaths, killing 1.3 million people each year. (7) Lung cancer is the second-most frequently diagnosed cancer in both men and women; after prostate and breast cancers which are the most frequent in men and women, respectively. (8) In Canada, lung cancer continues to be the leading cause of cancer mortality with an estimated 9,400 deaths occurring among women and an estimated 11,200 deaths among men in 2009. (9) In 2010, an estimated 24,200 Canadians will be diagnosed with lung cancer and 20,600 will die of it. Lung cancer remains the leading cause of cancer death for both men and women. (10)
Lung cancer usually presents in patients aged between 40 and 70 years, with the peak incidence in the sixth decade. (11) There is evidence that cigarette smoking is a major causal factor. Clinical studies have observed the changes in the epithelium lining of the respiratory tract in smokers and non-smokers. (12) Loss of ciliated cells, basal cell hyperplasia, squamous metaplasia and atypia of cells are the changes that have been noticed among smokers. (13) Also, the risk of developing lung cancer is increased by industrial exposure to carcinogens such as radiation, uranium, asbestos, nickel, chromates, coal, mustard gas, arsenic, beryllium, and iron. (14) Genetic predisposition is also known to be associated with bronchogenic carcinoma. Approximately 10% of these cancers occur in non-smokers; although it should be mentioned that bronchogenic carcinoma in non-smokers is usually of a different subgroup than in smokers’ type. Occasionally, lung cancers arise in scar tissue and are termed scar cancers. Histologically, these are usually of an adenocarcinoma subtype. These scars are usually due to old infarcts, wounds, foreign bodies or granulomatous infections such as tuberculosis.

3.2 Solitary Pulmonary Nodules

A solitary pulmonary nodule is defined as a discrete lesion < 3 cm in diameter that is surrounded by lung parenchyma (i.e., does not touch the hilum, mediastinum, or pleura). Solitary pulmonary nodules are most often detected incidentally when a chest x-ray is taken for other reasons. Every pulmonary nodule could be an early stage of a lung cancer; therefore it is vital to distinguish a benign nodule from a cancerous one. Solitary pulmonary nodules have many causes, including inflammatory lesions, granulomas and infections. (15) Patient age and risk factors play an important role in assessing the likelihood of malignancy. The primary goal in evaluation of any nodule should be to differentiate benignity from malignancy and active infection. Although a thorough history and physical exam is the first step in approaching a patient with a pulmonary
nodule, the first imaging test that is usually performed is a chest x-ray followed by a thoracic CT scan if necessary. There are some important radiographic characteristics that help to distinguish a benign from a malignant nodule:

1) Growth rate, which is determined by comparing the current images with previous ones. No enlargement in more than 2 years suggests a benign etiology. Tumors with volume doubling times from 21 to 400 days have a higher probability of being malignant.

2) Presence of calcification if it is central, concentric or in a popcorn configuration suggests benign disease.

3) Margins that are spiculated or irregular are more indicative of malignancy.

4) Diameter less than 1.5 cm strongly suggests a benign etiology and more than 5.3 cm strongly suggests cancer although there are some exceptions, which include lung abscess, Wegener granulomatosis, and hydatid cysts.

Since the clinical stage at the time of diagnosis is the major factor in survival rate after therapy, there has been a lot of interest in early lung cancer screening lately using the low dose CT which is a highly sensitive modality for detection of lung nodules.

3.3 Lung Cancer Screening

As advances in computed tomography allow detection of very small pulmonary nodule at lower radiation doses low dose computed tomography seems to play a very important role in early detection of lung cancer. The International Early Lung Cancer Action Project (I-ELCAP) (16) with over 35,000 subjects enrolled is currently one of the most prominent studies of base line screening for lung cancer. It involves a baseline and an annual repeat CT following the I-ELCAP
regimen of screening. New nodules that are identified are investigated within a specified protocol incorporating frequent CT scans, positron emission tomography scans, antibiotics or lung biopsy. Patients found to have lung cancer undergo surgical resection.

According to IELCAP data, the 10 year survival rate for these patients is 88 -92%. Although these results seem very favorable and encouraging, there are concerns: according to the Mayo Clinic experience (17) only 1.3 % of the detected nodules are malignant therefore ~ 99% of the detected nodules include other abnormalities which had unnecessary further work up. Although not all of these patients had lung biopsies performed, all of them were subjected to high levels of anxiety while waiting for the result of their tests. (18) A second methodological issue relates to the inherent biases on the observed results: selection bias, lead-time bias, length bias, and over diagnosis bias. Selection bias is the error in choosing the subjects to take part in a study. Ideally, they should be very similar to one another and to the larger population from which they are drawn (for example, all individuals with the same disease or condition). If there are important differences, the results of the study may not be valid. It can affect the mortality as well as other measures of outcome, but in a randomized trial, it is eliminated. All the observed results from the lung cancer screening studies, except mortality were subject to lead-time bias, length bias, and over diagnosis bias. (19) None of these studies demonstrated any significant improvements in the mortality rate for the screened population.

Lead time bias is one screening feature that could explain increased incidence, better survival and no decrease in mortality rate. If a screen detected cancer results in death exactly at the same time as it would have without screening, the only influence of screening was an earlier detection of the same fatal cancer. It can also explain “increased incidence” by leading to excess of cases in the screening population due to early detection. Over diagnosis bias happens when screening
detects lesions that are not clinically important and would not make any difference to the patient’s life span. Length bias can be explained by the fact that most indolent tumors have a long preclinical phase and therefore are more likely to be diagnosed during screening; whereas more aggressive tumors progress and present relatively quickly and therefore are less likely to be detected by screening. This could explain the observed increased survival in screening populations compared to the general population as most of the malignancies detected during screening are indolent, slow growing tumors.

The National Lung Screening Trial (NLST) is a multicenter study, and the largest randomized trial of lung cancer screening to date and it compares low dose helical computed tomography with chest radiography in an at-risk population. This study has enrolled 53,456 participants over a period of almost two years from September 2002 to April 2004. Participants had one baseline and two annual screenings, using either chest radiography or low dose thoracic CT. The end point of this trial was to assess the utility of low dose CT on lung cancer mortality. The five years survival rate approached 70% in patients with surgical resection of stage IA disease but in patients with metastatic disease the survival rate was less than 5%. (20). Recently, the National Cancer Institute (NCI) issued a press release of trial results that demonstrate a significant mortality benefit from screening with low-dose CT. There are twenty percent fewer lung cancer deaths among those who were screened with low dose helical CT as compared with chest X-ray. (21)

4 Lung Cancer Pathology

4.1 Lung Cancer Types

A wide spectrum of benign and malignant tumours can occur in the lungs, and bronchogenic carcinoma is the most common with an incidence rate of 90–95% of all lung tumours. (22) It is
followed in prevalence by bronchial carcinoid (5%), mesenchymal tumours and other miscellaneous neoplasms (2–5%). (23) This means that any lung primary malignancy can be assumed to be bronchogenic carcinoma until proven otherwise. Bronchogenic refers to the origin in the bronchial and sometimes bronchiolar epithelium. It usually arises in and around the hilus of the lung. About 75% of the lesions take origin from the lower trachea, first, second and third order bronchi. A small percentage of carcinomas are located in the lung periphery. Histologically, bronchogenic carcinoma is divided into four different subtypes:


Although each lesion usually consists of one specific cell type, there might be a mixture of histological patterns in a single lesion. It is important to know that squamous cell carcinoma has the strongest relationship to smoking among the different histological types. There is also another classification based on the lesion response to therapy, which includes:

A) Small cell carcinomas with high initial response to chemotherapy

B) Non-small cell carcinoma which is less responsive to chemotherapy.

In its developmental pathway, bronchogenic carcinoma begins as an area of in situ cytologic atypia. Then, over an unknown interval of time, it yields a small area of thickening of the bronchial mucosa. This small area usually grows to about 1 cm. It usually appears as an irregular warty area that erodes the lining epithelium. The lesion may then follow one of the following paths:

A) It may fungate into the bronchial lumen to produce an intraluminal mass.
B) It may rapidly penetrate the wall of the bronchus to infiltrate along the peribronchial tissue into the adjacent region of the carina or mediastinum. In his fashion, the tumour may even extend to about the pericardium or to the pleural surface and then within the pleural cavity.

C) It may also grow along a broad front to produce a cauliflower intraparenchymal mass that appears to push lung substance ahead of it.

The lesion is usually gray-white and firm to hard and may contain focal areas of hemorrhage and necrosis with cavitations, especially when the tumours are bulky. Almost 50% of cases spread to the tracheal, bronchial and mediastinal nodes. Distant metastasis usually spread widely through hematogenous or lymphatic pathways. Occasionally, the first manifestation of the underlying carcinoma is due to metastatatic disease. The most commonly involved organs include the adrenals (50%), liver (30%), brain (20%), bone (20%), and the kidneys (15%).

4.1.1 Squamous Cell Carcinoma

Squamous cell carcinoma has the strongest relationship to cigarette smoking and is more common in men than women. This tumour arises in the larger, more central bronchi, tends to spread locally and metastasizes later than the other cell types. A squamous cell lung tumour doubles its size every four months and usually requires approximately nine years to achieve a mass of 2 cm in diameter.

4.1.2 Adenocarcinoma

Adenocarcinoma usually presents in two forms: 1) bronchial-derived adenocarcinoma, and 2) bronchoalveolar carcinoma, arising from terminal bronchioles or alveolar walls. Although there is overlap between these two forms, bronchoalveolar carcinoma has distinctive gross, microscopic and epidemiologic features. Adenocarcinoma is the most common type of
bronchogenic carcinoma in women and non-smokers. Lesions are more peripherally located and tend to be smaller. They tend to grow more slowly than squamous cell carcinoma. Adenocarcinoma, including bronchoalveolar carcinoma, is less associated with a history of smoking than are squamous cell or small cell carcinoma.

4.1.3 Small Cell Carcinoma

Small cell carcinomas are highly malignant tumours. The epithelial cells are usually small and have little cytoplasm. Cells are round or oval, which is characteristic of oat cell carcinoma. Other small cells are spindle-shaped or polygonal-shaped and may be classified as spindle or polygonal small cell carcinoma. The cells grow in clusters that exhibit neither glandular nor squamous organization. Electron microscope studies show dense core neurosecretory granules in some of these tumours cells. These suggest derivation of these tumours from neuroendocrine cells of the bronchial epithelium lining. These tumours have a strong relationship to cigarette smoking. Only about 1% occurs in non-smokers. These tumours are usually located in hilar or central regions and are the most aggressive type of lung tumours. They metastasize widely and generally are incurable by surgical means.

4.1.4 Small Cell Carcinoma

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about 1% occurs in non-smokers. These tumours are usually located in hilar or central regions and are the most aggressive type of lung tumours. They metastasize widely and generally are incurable by surgical means.

4.1.5 Large Cell Carcinoma

Large cell carcinoma has larger, more polygonal cells and vesicular nuclei. They probably represent undifferentiated squamous and adenocarcinoma. Some tumours contain mucin, some have a larger number of multinucleated cells (giant cell carcinoma), some have clear cells and are termed clear cell carcinoma, and others have a distinctly spindly histological appearance (spindle cell carcinoma).

4.2 Secondary Pathology

Bronchogenic carcinomas cause related anatomical changes in the lung distal to the point of bronchial involvement. Partial obstruction may cause marked focal emphysema and total obstruction may lead to atelectasis. The impaired drainage of the airways is a common cause for severe suppurative or ulcerative bronchitis or bronchiectasis. Pulmonary abscesses sometimes draw attention to an occult carcinoma that has become infected. Compression or invasion of the superior vena cava can cause venous congestion, dusky head and arm edema, and ultimately circulatory compromise, known as the superior vena cava syndrome. Extension to the pericardium or pleural sac may cause pericarditis or pleuritis with development of significant effusions.

4.3 Lung cancer Staging

A uniform TNM system for staging cancer according to its anatomical extent at the time of diagnosis is important. The “T” descriptor indicates the extent of the primary tumour, “N” the
extent of lymph node involvement, and “M” the presence of metastases. (23, 25) Table 1.1 below shows the staging system in its current form:

<table>
<thead>
<tr>
<th>Diameter</th>
<th>Bronchoscope</th>
<th>Atelectasis</th>
<th>Invasion</th>
<th>Nodules</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T1</strong></td>
<td>T1a: &lt; 2cm</td>
<td>No invasion,</td>
<td>Lobar or Bronchus</td>
<td></td>
</tr>
<tr>
<td></td>
<td>T1b: 2-3 cm</td>
<td>Lobar atelectasis or obstructive pneumonia to hilus</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>T2</strong></td>
<td>T2a: 3-5 cm</td>
<td>&gt; 2 cm to carina</td>
<td>Chest wall</td>
<td>Nodules in same lobe</td>
</tr>
<tr>
<td></td>
<td>T2b: 5-7 cm</td>
<td></td>
<td>Diaphragm</td>
<td></td>
</tr>
<tr>
<td><strong>T3</strong></td>
<td>&gt; 7 cm</td>
<td>&lt; 2 cm to carina</td>
<td>Mediastinum</td>
<td>Nodules in other ipsilateral lobes</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Pleura</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Pericardium</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Heart</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Great vessels</td>
<td></td>
</tr>
<tr>
<td><strong>T4</strong></td>
<td>Tumour in carina</td>
<td>Whole lung</td>
<td>Trachea</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Esophagus</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>spine</td>
<td></td>
</tr>
</tbody>
</table>

**N0**
No lymph nodes involved

**N1**
Involvement of ipsilateral bronchopulmonary or hilar nodes

**N2**
Involvement of ipsilateral mediastinal, ipsilateral supraclavicular or subcarinal nodes

**N3**
Involvement of contralateral mediastinal, hilar, or supraclavicular lymph nodes

**M0**
No distant metastasis

**M1**
Distant metastasis

**M1a**: Separate tumour in a contralateral lobe or tumour with plural nodules or malignant plural or pericardial effusion

**M1b**: Distant metastasis
Table 1.1 TMN staging. The TNM staging system takes into account the degree of spread of the primary tumour, represented by T; the extent of regional lymph node involvement, represented by N; and the presence or absence of distant metastases, represented by M.

The TNM staging is commonly grouped into subsets with similar prognosis and treatment options. These are grouped into four stages, as described in table 1.2

<table>
<thead>
<tr>
<th>Stage</th>
<th>Tumour</th>
<th>Lymph Node</th>
<th>Metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage IA</td>
<td>T1a,T1b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IB</td>
<td>T2a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIA</td>
<td>T1a,T1b</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T2a</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T2b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIB</td>
<td>T2b</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIA</td>
<td>T1a,1b,T2a,T2b,T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3, T4</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIB</td>
<td>Any T</td>
<td>N3</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IV</td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

Table 1.2 Subtypes of lung cancer based on therapeutic and prognostic implication.

4.4 Clinical Course

Lung cancer is one of the most insidious neoplasms in the world of oncology. It may present with six to seven months of symptom duration. The major presenting complaints are a cough (75%), weight loss (40%), chest pain (40%) and dyspnea (20%). Increased sputum production is common and often contains diagnostic tumour cells when examined as cytologic specimens.
Similarly cytologic examination from a fine needle aspiration (FNA) of a mass can often provide the diagnosis.

The outlook is generally poor for most patients with bronchogenic carcinoma. Despite all the efforts at early diagnosis by frequent radiological exams of the chest, cytological examination of the sputum, bronchial washing and brushing and many other improvements in thoracic surgery, radiotherapy, and chemotherapy, the overall five-year survival rate in united state is about 15% and in Europe and developing country is only 8% (25). In general, adenocarcinoma and squamous cell carcinoma tend to remain localized longer and have slightly better prognosis than undifferentiated tumours. Surgery is generally not curative for small cell carcinoma. If untreated, survival length is about 6–17 weeks, but this cancer is particularly sensitive to chemotherapy therefore providing temporary symptomatic relief in suitable patients. Unfortunately, most of these patients have distant metastasis at the time of diagnosis. However, earlier diagnosis of symptomatic patients with an earlier disease stage and use of multimodality therapy has shown promising outcomes in selective sub groups of patients.

Bronchogenic carcinoma has been associated with a number of paraneoplastic syndromes. The hormones or hormone-like factors elaborated include 1) antidiuretic hormones (ADH) inducing hypothermia, due to inappropriate ADH secretion, 2) adrenocorticotropic hormone (ACTH), producing Cushing syndrome, 3) parathormone, parathyroid hormone-related peptide, prostaglandin E, and some cytokines, all of which are implicated in hypercalcemia and sometimes seen in lung cancer, 4) calcitonin causing hypocalcemia, 5) gonadotropin causing gynecomastia, and 6) serotonin, associated with carcinoid syndrome. Any one of the histological types may occasionally produce any ectopic hormone, but ACTH and ADH are predominantly produced by small cell carcinoma, whereas hypercalcemia is usually related to squamous cell
tumours. Carcinoid syndrome is rarely associated with small cell carcinoma but is more common with bronchial carcinoid.

Other systemic manifestations of lung tumours include Lambert-Eaton myasthenic syndrome, where muscle weakness is caused by autoimmune antibodies directed to t-neural calcium channels, peripheral neuropathy, usually purely sensory, dermatologic abnormalities, including acanthosis nigricans, hematologic abnormalities such as a leukemoid reaction, and finally a peculiar abnormality of connective tissue called hypertrophic pulmonary osteoarthropathy, associated with clubbing of the fingers.

Apical lung cancer in the superior pulmonary sulcus tends to invade the neural structures around the trachea, including the cervical sympathetic plexus, and produces a group of clinical findings that includes severe pain in the distribution of the ulnar nerve and Horner’s syndrome (exophthalmos, ptosis, miosis, and anhidrosis) on the same side as the lesion. These tumours are also referred to as Pancoast tumours.

5 Chest Imaging Technologies

5.1 Chest Radiography (CXR)

Chest radiography (CXR) is usually the first imaging step in evaluating patients with signs and symptoms of lung cancer. It presents a low-cost, minimal radiation dose modality that remains the mainstay of non-invasive examination and diagnosis of all chest pathologies in large health centers and rural hospitals alike.
Figure 1.4 The anteroposterior (a) and lateral (b) views of male chest x-ray. Main anatomical regions are labeled.

Chest radiography has significant limitations due to the superimposition of anatomical structures. This is one of the main reasons why CXR lacks sensitivity in detection of small lung nodules, and has not shown itself to be suitable for screening purposes. (26) Technical advances in digital radiography (DR) have yielded remarkable improvements in diagnostic quality through the introduction of high-performance flat-panel detectors (FPDs). (27-29)

Today, conventional analogue chest imaging has been replaced by digital imaging at most medical centers. Digital image acquisition allows immediate electronic distribution of images through the use of a picture archiving and communication system (PACS). Digital imaging modalities include computed radiography (CR) and direct capture radiography (DR) imaging of lung disease. Still, the fundamental limitation to diagnostic performance remains which is the compression of three-dimensional anatomy into a two-dimensional projection image, (27) with the visual clutter associated with superimposed structures believed to be a major limiting factor.
These limitations make planar radiography an unreliable technique for pulmonary nodule detection. Austin et al. (30) reviewed 27 cases of resectable lung cancer that were missed on previous chest radiographs. The majority of missed nodules were located in upper lobes and in females. Muhm et al (31) showed that 90% of peripheral nodules detected during a lung cancer screening program in retrospect were visible on earlier radiographs.

5.2 Low Dose CT

Computed tomography (CT), especially with helical and multi-detector technology, is superior to chest x-radiography for detecting pulmonary nodules. However, there is concern for the higher radiation exposure to the patient, compared with chest radiography, and this creates anxiety regarding the potential induction of malignancy (32).

The reported effective radiation dose in one routine computed tomography chest exam is several hundred times more than that of a single frontal chest radiograph (8 mSv versus 0.02 mSv) (33). However, with modern CT units and more awareness of radiation dose issues, the radiation dose associated with thoracic CT has significantly decreased but still remains significantly higher than chest radiography (1.0 mSv for low dose CT). Low dose computed tomography has been shown to be a useful tool in the early detection of lung cancer (34). However, a low radiation dose causes increased image noise and image artifacts, which degrades image quality, especially in larger patients. Development of reconstruction software like special frequency filters, artifact correction and noise reduction algorithms have reduced such effects to some degree (35).

Although recent technological developments like volumetric scanning in a single gantry rotation offer faster scan times, reduced motion artifact and radiation dose, there are still some obstacles in using CT, which include high number of false positives and very high financial cost.
Figure 1.5 Comparison of the anatomical detail displayed on a frontal chest radiograph and an axial projection from a thoracic CT study taken at the level of the main pulmonary trunk.

5.3 Positron Emission Tomography

18F-fluorodeoxyglucose (FDG) positron emission tomography (PET) is a nuclear medicine imaging technique with evolving potential. 18F-fluorodeoxyglucose is a radiolabeled glucose analogue that is injected intravenously as the tracer. It is taken up by metabolically active cells, including cancer cells, and is phosphorylated in the intracellular space to FDG-6-Phosphate. This compound cannot get further catabolized and therefore remains trapped inside the cell. The trapped FDG decays by positron emission, the positron collides with an electron and produces energy in the form of a gamma ray, which can be detected by a tomography camera. The resultant images can be displayed in sagittal, axial or coronal planes. FDG-PET measures the metabolic activity in a lung nodule/mass and can be very useful in determining the presence of cancer cells in indeterminate lung nodules and the presence of occult metastases. Therefore,
FDG-PET is useful in detecting and staging lung cancer in earlier stages, however, it is associated with a high radiation dose especially when it is combined with CT as usually is the case in modern practice (37) Major reductions in radiation doses from PET/CT scans can be achieved by modifying the acquisition parameters for CT. Also, PET has the potential to assess tumour responsiveness to chemotherapy, which could serve as a prognostic factor and could influence further management of the disease. (38)

Figure 1.6 Projection images from a FDG PET scan (lower row) demonstrate intense uptake within a right lower lobe tumour. The top row shows the corresponding CT images (lung reconstruction (left) and mediastinal reconstruction, (right)). The tumour is seen at the same location. Modified from www.chestjournal.org/.../124/3/893/F1.expansion
5.4 Dual Energy Chest Imaging

Dual Energy (DE) imaging acquires two projections of the patient at different x-ray energies and selectively decomposes the image into soft tissue and bone components. The former presents soft tissue structures in a context that is largely free of the main source of anatomical clutter (the ribs and clavicles), thus improving the conspicuity of subtle nodules. Furthermore, because the presence of calcification is an important indicator of benignancy, DE imaging could help characterize benign lesions with a higher level of specificity. Dual energy imaging may mitigate the limitations of anatomical clutter found in regular chest x-rays and also increase specificity by obtaining tissue-specific images. (39)

The general form of weighted log-subtraction decomposition of a DE image is as follow:

\[
\text{Log}(I_{DE}) = \log(I_H) - w \log(I_L)
\]

Where \(I_H\) is the high energy image and \(I_L\) is the low energy image. \(w\), the weighting factor is the ratio of the attenuation coefficient of the cancelled material at low and high energy. This is described in details in the next chapter.

Commercially available DE imaging systems include storage phosphor and (Flat panel detector) FPD designs. Single-shot DE imaging (e.g., FCR XU-D1; Fujifilm, Tokyo, Japan) involves two CR storage phosphors as front and back detectors separated by an intermediate filter, with the image produced by the front plate representing a low-energy image and that by the back plate a high-energy image. As detailed below, soft tissue and bone images are decomposed from these low- and high-energy images using various subtraction algorithms. Double-shot DE images are generated using two x-ray exposures at two kVp settings (e.g., QX/I Revolution, GE Healthcare, Chalfont St. Giles, UK). The advantage of this system is the improved energy separation (and, therefore, contrast-to-noise ratio) compared to single shot DE designs but the potential
disadvantage of two acquisitions lies in the patient related motion artifacts (breathing, moving and cardiac) which happen due to the time interval between the two exposures, even for FPDs with rapid readout. Although a compliant patient may be able to hold their breath and not move between the two exposures, there is usually a change in heart rate, even with a short breath hold, therefore accurate and robust cardiac gating would be a useful feature for two shot DE acquisitions.

5.5 Radiation Hazard

Potential long term health effects, such as cancer induction and hereditary genetic damage, caused by exposure to ionizing radiation are not well understood but have been identified. It is assumed that the relationship between radiation exposure and adverse effect is linear, meaning that a doubling of radiation dose results in twice the risk of damage. In addition, many health scientists assume that there is no radiation dose threshold for these ill effects, meaning that there is no level of energy at which the health risk is assumed to be zero. (40) The advent of computed tomography has revolutionized diagnostic radiology but CT involves larger radiation doses than conventional chest radiography. Multiple factors are involved in determining the radiation dose to a particular organ during CT imaging, including the tube potential in kilovolts (kVp), the tube current in milliamperes (mA), the scan time in second (s) and the size of the patient. The scan parameters should be tailored to the study protocol in order to minimize patient radiation exposure.

The principle reason that ionizing radiation is a biological hazard relates to the unique energy of the X-ray photons which is sufficient to overcome the binding energy of the electrons orbiting the atomic nucleus and thereby displace them from their orbits. The residual atomic nucleus becomes a positively charged ion and this entity can interact with DNA. The most commonly
created ions are hydroxyl radicals from interaction of X-ray photons with water molecules. These radicals can cause breakage of the DNA double helix strands or can ionize DNA proteins directly. The break in the double stranded DNA is not easily repaired and occasionally the mis-repair can induce point mutations and chromosomal translocations which potentiate cancer cells. (41)

Most of the available data for estimating the risk of radiation induced cancer comes from extensive studies performed on the Japanese survivors of atomic bombs explosions. The risk from exposure to ionizing radiation was quantitatively consistent in all of these studies. Children are at greater risk than adults from a given dose of radiation since they are more radiosensitive and they also have more remaining years of life, in which to develop induced cancer, compared to adults. (42) Overall it is possible to estimate the cancer risks associated with radiation exposure by estimating the organ doses involved and applying organ specific cancer incidence or mortality data.

The estimated number of CT scans that lead to cancer, varies depend on the type of the scan and the patient sex and age. For example according to one study published by Smith et al (43) the estimated risk of cancer induction in women undergoing CT coronary angiography at the age of 40 years, is 1 in 270. The analysis resulted in the rate of 1 in 600 for men. For head CT the same analysis resulted in an estimated cancer rate of 1 in 8100 for women and 1 in 11080 for men. For 20 year old patients the overall risk was almost doubled and for 60 year old patients, it was 50 % lower.

Bearing this in mind, an important pre-requisite for this research was to ensure that the radiation dose associated with the dual energy exposure was equivalent to that from a conventional frontal chest radiography (0.02 mSv).
Chapter 2  
Dual Energy Prototype and Patient Cohort

1 DE Imaging System

The experimental prototype DE imaging system was based on a Kodak RVG 5100 digital radiography chest stand (Carestream Health Inc., Rochester, NY). An acquisition workstation controls generator technique setting, filter selection, detector acquisition parameters, and data transfer. The system was modified for the purposes of DE imaging to include: 1) a flat-panel detector (Trixell Pixium-4600, Moirans, France) with a ~250 mg/cm$^2$ CsI:Tl scintillator and 143 μm pixel pitch, 2) a fingertip pulse oximeter to trigger x-ray exposure during diastole for reduction of cardiac motion artifacts, (44) and 3) a multi-position filter wheel within the collimator to change the added filtration between the low-kVp (2.5 mm Al) and high-kVp (2 mm Al + 0.6 mm Ag) exposures, as guided by previous studies of DE image quality performed in our laboratory. (44-46)

The X-ray tube produces a polychromatic beam with spectrum of beam energies distributed around the preset target tube potential. Therefore, a high-energy filter was selected to “harden” the high energy beam, to reduce spectral overlap between the low- and high-energy projections. This was the subject of considerable investigation in our laboratory (45, 46, 47). Differential filtration between low- and high-kVp projections was selected and an extensive analysis of contrast, Noise Equivalent Quanta (NEQ) and nodule detectability for low- and high-kVp added filtration was performed. A greater than 50% improvement in contrast and a boost in NEQ were achieved using an additional 0.6 mm Ag in the high-kVp projection. (45) The added filtration for the DR image was typical of that used in conventional digital chest radiography. Three different
filters for the low-energy, high-energy, and DR exposures were implemented using a computer-controlled, multi-position filter wheel within the collimator.

Optimal acquisition techniques including kVp pair and dose allocation ($A\epsilon$) were identified in previous work by experimental measurements in phantoms (46) and theoretical analysis. (48) A chest phantom with simulated lung nodules was used. For thick (30 cm), average (24 cm) and thin (18 cm) patients, various thickness of acrylic was used. Multiple acquisition were taken using a range of low-kVp (60-90 kVp), high-kVp (120-150 kVp) and dose allocation ($A\epsilon = \sim 0-1$). Patient dose, characterized as total imparted energy ($\epsilon_{\text{Total}} = \epsilon_L + \epsilon^H$) was kept equal to that of a single DR radiograph (within $\pm 5\%$) for the same chest thickness. DE images were acquired at various low-kVp, high-kVp, and allocations. For each patient thickness, and kVp pair, combinations of $\epsilon^L$ (mAs) and $\epsilon^H$ (mAs) that yielded a given total dose ($\epsilon_{\text{Total}}$) equal to one DR (within $\pm 5\%$) were identified. The total of ten combinations were identified for each patient thickness and kVp pair, since for each mAs combination, a specific allocation resulted in total dose equal to one DR. To identify the optimal kVp pair and allocation for each patient thickness, Signal difference to noise ratio (SDNR) was measured in DE soft tissue images of the phantom for a total of 16 kVp pairs and three phantom thicknesses. (46) For each combination, the peak SDNR was found at an allocation of $A\epsilon = 0.33$, suggesting optimal image quality when one third of the total dose is imparted by the low-kVp beam. Also a significant improvement in SNRD was observed with increasing the spectral separation, by reducing the low-kVp. The SDNR was found to be highest at 60/120 kVp. DR techniques for thin, average and thick patient sizes were obtained from the clinical technique chart in our institution. (46)
The technique charts for DE and DR (Table 2.1, 2.2) were interpolated across patient chest thicknesses ranging 18–28 cm in increments of 2 cm, with the total dose dependent on patient thickness (e.g., 0.11 mGy for 24 cm chest thickness).

<table>
<thead>
<tr>
<th>Chest thickness (cm)</th>
<th>Range (cm)</th>
<th>mA</th>
<th>ms</th>
<th>mAs</th>
<th>mA</th>
<th>ms</th>
<th>mAs</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>&lt;15</td>
<td>400</td>
<td>6.25</td>
<td>2.5</td>
<td>400</td>
<td>25</td>
<td>10.0</td>
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<tr>
<td>16</td>
<td>15 - 16.9</td>
<td>400</td>
<td>6.25</td>
<td>2.5</td>
<td>400</td>
<td>32</td>
<td>12.8</td>
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<tr>
<td>18</td>
<td>17 - 18.9</td>
<td>400</td>
<td>8</td>
<td>3.2</td>
<td>400</td>
<td>40</td>
<td>16.0</td>
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<tr>
<td>20</td>
<td>19 - 20.9</td>
<td>400</td>
<td>10</td>
<td>4.0</td>
<td>500</td>
<td>40</td>
<td>20.0</td>
</tr>
<tr>
<td>22</td>
<td>21 - 22.9</td>
<td>500</td>
<td>10</td>
<td>5.0</td>
<td>630</td>
<td>40</td>
<td>25.2</td>
</tr>
<tr>
<td>24</td>
<td>23 - 24.9</td>
<td>400</td>
<td>12.5</td>
<td>5.0</td>
<td>630</td>
<td>40</td>
<td>25.2</td>
</tr>
<tr>
<td>26</td>
<td>25 - 26.9</td>
<td>400</td>
<td>16</td>
<td>6.4</td>
<td>630</td>
<td>50</td>
<td>31.5</td>
</tr>
<tr>
<td>28</td>
<td>27 - 28.9</td>
<td>630</td>
<td>12.5</td>
<td>7.9</td>
<td>630</td>
<td>63</td>
<td>39.7</td>
</tr>
<tr>
<td>30</td>
<td>29 - 30.9</td>
<td>630</td>
<td>16</td>
<td>10.1</td>
<td>630</td>
<td>63</td>
<td>39.7</td>
</tr>
</tbody>
</table>

Table 2.1 Optimal technique chart for Dual energy projections based on patient thickness.

<table>
<thead>
<tr>
<th>Chest Thickness (cm)</th>
<th>Range (cm)</th>
<th>kVp</th>
<th>mA</th>
<th>ms</th>
<th>mAs</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>&lt;15</td>
<td>120</td>
<td>32</td>
<td>50</td>
<td>1.6</td>
</tr>
<tr>
<td>16</td>
<td>15 - 16.9</td>
<td>120</td>
<td>32</td>
<td>50</td>
<td>1.6</td>
</tr>
<tr>
<td>18</td>
<td>17 - 18.9</td>
<td>120</td>
<td>40</td>
<td>50</td>
<td>2</td>
</tr>
<tr>
<td>20</td>
<td>19 - 20.9</td>
<td>120</td>
<td>40</td>
<td>50</td>
<td>2</td>
</tr>
<tr>
<td>22</td>
<td>21 - 22.9</td>
<td>120</td>
<td>40</td>
<td>63</td>
<td>2.5</td>
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<tr>
<td>24</td>
<td>23 - 24.9</td>
<td>120</td>
<td>40</td>
<td>80</td>
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<tr>
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<td>25 - 26.9</td>
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<td>50</td>
<td>80</td>
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<tr>
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<td>27 - 28.9</td>
<td>120</td>
<td>63</td>
<td>80</td>
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</tr>
<tr>
<td>30</td>
<td>29 - 30.9</td>
<td>120</td>
<td>80</td>
<td>80</td>
<td>6.4</td>
</tr>
</tbody>
</table>

Table 2.2 Optimal technique chart for digital radiograph projection based on patient thickness.
DE images were processed and decomposed as illustrated in Fig. 2.1. Offset and gain corrections were based upon 25 averaged dark-fields and 25 averaged flood-fields (acquired at ~50% sensor saturation), respectively. Prior to DE decomposition, the high-energy image was automatically registered to the low-energy image by means of deformable registration based on mutual information maximization and a morphological pyramid. (49)

**Figure 2.1** Illustration of dual-energy (DE) image processing, registration (Reg), decomposition (Decomp) and display. Processing steps include offset, gain, and defect correction; deformable registration; weighted log-subtraction decomposition; and the transformation of pixel values to log-exposure space for display. The photograph on the right shows the experimental setup for observer studies, with images displayed on the left and the rating scale on the right. Bone: bone decomposition; Comp: composite equivalent radiograph; HE: high-energy projection; LE: low-energy projection; Soft: soft tissue decomposition.

The low-energy image and the (registered) high-energy image were decomposed into soft tissue and bone images by log-weighted subtraction of the basic form:

\[ \log(I_{\text{soft}}) = \log(I_H) - w_s \log(I_L) \]  
\[ \log(I_{\text{bone}}) = -\log(I_H) + w_b \log(I_L) \]  
\[ \log(I_{\text{comp}}) = \log(I_H) + w_c \log(I_L) \]
Where $I_H$ is the high energy image and $I_L$ is the low energy image. $w$ is the tissue cancellation parameter and ideally is the ratio of the attenuation coefficient ($\mu$) of the cancelled material at low and high energy. For example:

$$w_s = \frac{\mu_{\text{bone}}}{\mu_{\text{soft}}},$$

$$w_b = \frac{\mu_{\text{bone}}}{\mu_{\text{soft}}}.$$

This gives three components to each DE image: the soft-tissue image (DE soft), the bone image (DE bone), and the composite “equivalent radiograph” image (DE comp). The basic log subtraction technique of Eq. (1) was modified to include noise reduction techniques optimal to each image. (50) Specifically, the soft tissue image was decomposed using an anti-correlated noise reduction (ACNR) algorithm, (48, 50, 51) and the bone image was decomposed using a simple smoothing of the high-energy image (SSH) algorithm. (48, 52) The ACNR algorithm is based on the fact that quantum noise in the soft tissue image and bone only image is anti-correlated. In decomposing a soft tissue image ACNR applies a high pass filter to the complementary image (the bone only image) which removes the gross anatomical structures from the complementary image and leave it only with some residual edge artifact and quantum noise which is anti-correlated to the quantum noise in the original DE image. The original DE image and the filtered complementary image are then added and weighted by a parameter $w_n$. Therefore the soft tissue image decomposed by ACNR is:

$$I_{\text{ACNR}} = I_{\text{Soft}} + w_n I_{\text{Bone}} * h_{\text{HPF}}$$

Where $I_{\text{Bone}} = -I_H + w_b I_L$, is the complementary image and $w_b$ is the tissue cancellation parameter. $h_{\text{HPF}}$ is the high pass filter used for complementary image. $w_n$ can be determined qualitatively or quantitatively through the minimization of quantum noise.
SSH is based on the proven fact that the greater contributor of the quantum noise in the resulting DE images is the high energy projection. This effect can be alleviated by applying a low pass filter to the high energy projection before log weighted subtraction. The unfiltered low energy projection preserves the high special frequency information. Therefore the resulting image is expressed as:

\[ I_{SSH} = -I_H * h_{LPF} + w I_L \]

Where \( h_{LPF} \) is the low pass filter, \( I_H \) is the high energy image, \( I_L \) is the low energy image and \( w \) is the cancellation parameter.

All decomposition parameters (ws, wb, wc, etc.) were manually selected for each case by a reader (who was not one of the observers in the observer tests). The reader was trained by a radiologist prior to modifying the images to learn the overall radiologists’ preferences for the resulting soft and bone images. The main image quality criterion considered by the expert reader involved the overall completeness of tissue cancellation (e.g., extinction of bone in the soft tissue image and vice versa). The resulting soft tissue images were judged to maximize conspicuity of pulmonary structures (and minimize visibility of bone), and conversely for the bone image. The composite image was selected in a manner to give image quality, tissue contrast, etc. as close as possible to that exhibited in the DR image. The adjustments to each image were qualitative in nature, this was necessary in order to allow for variables including patient body shape and positioning. Example images are shown in Fig. 2.2

The DR image was acquired a few seconds after the DE image pair on the same imaging prototype. The technique for the DR image used a tube potential of 120 kVp (1 mm Al + 0.2 mm
Cu filtration), and the tube current (mAs) was set according to the PA chest thickness and determined from a published technique chart. (46)

**Figure 2.2** Example dual energy (DE) and digital radiography (DR) images. DE components include: (A) the soft-tissue image, (B) the bone image, and (C) the composite image. The corresponding DR image is shown in (D). The location of a 2.4 cm nodule in the left upper lobe is marked by an arrow.

## 2 Patient cohort

Patients were accrued under informed consent in a prospective, non-randomized trial with approval from the institutional research ethics board and with Health Canada Investigational Testing Authorization. The total study cohort of 220 patients was drawn from the patient population referred for a percutaneous computed tomography-guided biopsy of suspicious lung nodules between September 2006 and April 2008. Based on the study design, all patients had to have a previously detected indeterminate lung nodule, a standard dose thoracic CT and histology. Therefore, all of the patients had at least one suspicious lung nodule detected either on an initial computed tomography study or on chest radiography.
Patients were prospectively accrued into five arms differing in DE imaging technique to evaluate different DE imaging parameters. These included differential x-ray spectra, cardiac gating, dose allocation, and total dose. (53) The main study cohort of 129 patients (80 M, 49 F, 62% male) with a mean age of 64.8 year was allocated to the optimal imaging technique. This gender distribution is reflective of the patient population referred to our Institution for lung biopsy during the period of the study (Sep 2006-Apr 2008, 56 % males). The remaining patients were equally allocated to the other four groups. Figure 2.3 below shows the different group allocations.

**Figure 2.3** Patient accrual into five different research arms to evaluate variable DE parameters.
The following data was gathered for each patient in the trial: standard-of-care image data (standard dose CT, ultra low-dose CT, acquired just prior to lung biopsy, and a post-biopsy CR image), experimental protocol image data (a frontal chest DR image and a DE image, each described below), and biopsy data. Percutaneous CT guided transthoracic biopsies were performed immediately following the DE/DR imaging exam to provide a definitive diagnosis of the lung lesion. Biopsies were performed either by fine-needle aspiration and cytologic examination or by core biopsy and histologic examination. (54)

Two different observer studies were performed. The first study incorporated a diagnostic satisfaction rating test using 55 cases (accrued at the time of the observer study) and the second study utilized Receiver Operating Characteristic (ROC) analysis for pulmonary nodule detection, using 129 cases. (53, 55) All images used in the two studies reported herein were taken from Group 1 (the “optimal” DE imaging technique described above).

3 Meta Data

Different parameters were measured for each patient. These parameters were grouped in five different categories:

A. Patient Characteristics

B. Disease Characteristics

C. Exam Characteristics

D. DE Image Techniques

E. Multi-Scale, Multi-Resolution Registration (MSMR) Characteristics
3.1 Patient Characteristics

3.1.1 Gender

Patient gender was acquired from electronic patient information accessible to hospital employees. A total of 91 females and 129 males were accrued for the study.

3.1.2 Age

Patient age at the time of admission was acquired from the patient information system. The mean age was 64.9 years, ranging from 25 to 90 years; the frequency of cases in each age range is demonstrated in Figure 2.4. Most patients were in the 55–75 years age range and this accurately reflects the presenting age of patients with lung cancer.

![Age Distribution](image)

**Figure 2.4** A histogram describing the number of cases for each age group. The majority of patient with suspicious lung nodules were between 55 and 65 years of age (70 Cases) followed by those between 65 and 75 years of age (69 cases).
3.1.3 Region

Primary lung cancer most commonly presents in the upper lobes (56). To determine the most common location for the study population, the location of the lesion undergoing biopsy was recorded using the diagnostic CT axial projections that were taken prior to the lung biopsy.

Regions were divided into 7 different areas based on the anatomy of the lung: right upper lobe, right middle lobe, right lower lobe, left upper lobe, lingula, left lower lobe and mediastinum. Both lungs have an oblique fissure that separates the upper lobe, from the lower lobe. In addition, the right lung has a horizontal fissure that creates a middle lobe from the upper lobe. The middle lobe has an anatomical equivalent in the left lung, the lingual which has a dedicated bronchial anatomy but does not have a separate fissure. The mediastinum lies between the right and left pleura and is surrounded by loose connective tissue. It contains the heart, the great vessels of the heart, esophagus, trachea, phrenic nerve, cardiac nerve, thoracic duct, thymus, and lymph nodes of the central chest.

The number of cases in each category was plotted against the region. As demonstrated in Figure 2.5 below, the frequency of cases was highest in the right upper lobe and lowest in the left middle lobe.
Figure 2.5 A histogram showing the anatomical location of the lung nodule undergoing biopsy. As shown in the histogram, the most frequent location was the right upper lobe. Anatomical lobes of lungs: 1. Right Upper Lobe=RUL, 2. Right Middle Lobe=RML, 3. Right Lower Lobe=RLL, 4. Left Upper Lobe=LUL, 5. Left Middle Lobe=Lingula, 6. Left lower Lobe=LLL, 0=Meditastinum.
3.1.4 Chest Thickness

Chest thickness was measured for each patient with two different methods. Firstly, a chest caliper was used to manually measure the anteroposterior (AP) diameter of the patients’ thorax before taking the DE projections. Secondly, the skin-skin maximum thoracic AP diameter was measured from the lateral CT scout projection using electronic calipers available on the Picture Archiving and Computer System (PACS). The measurement was made at the level of the T9 vertebra. These measurements were plotted and the resulting histograms are shown in Figure 2.6, with the majority of patients having a thoracic AP diameter 24–26 cm.
Figure 2.6 Histograms of chest measurements versus frequency in total patient and in each gender individually. The majority of cases have the anteroposterior diameter of the average size patient, being 24-26 cm.

Figure 2.7 below demonstrates the association between the two different methods of AP measurements using regression analysis and measuring the R-squared value. With an R-square
value of 0.73, the association is quite strong. This analysis was performed to correlate the accuracy of manual and electronic chest AP diameter measurement.

![Correlation of two different methods for measuring chest AP diameter](image)

Coeficient of determination $R^2 = 0.7930$  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Coefficient</th>
<th>Std. Error</th>
<th>95% CI</th>
<th>t</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
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<td>1.2236</td>
<td>-2.9531 to 1.8956</td>
<td>-0.4321</td>
<td>0.6665</td>
</tr>
<tr>
<td>Slope</td>
<td>1.0461</td>
<td>0.05051</td>
<td>0.9460 to 1.1462</td>
<td>20.7120</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

**Figure 2.7** Regression analysis for measuring the anteroposterior diameter on the CT axial projection and by using a manual caliper. R-squared is quite high, showing the strong relationship between the two variables.

As seen in the scatter plot above, measured chest diameters using the two different methods are highly correlated. ($R=0.79$)
3.2 Disease Characteristics

3.2.1 Density

The density (Hounsfield unit, HU) of each lung nodule was measured using an electronic measuring tool on the PACS workstation (Fusion E-film 2.1; Merge Healthcare, Milwaukee, WI) in By convention, all lesions measuring $\geq$20 HU are considered solid, and any lesion $<20$ HU is considered non-solid. The histogram in Figure 2.8 demonstrates the frequency of nodule density measured from the diagnostic thoracic CT scan, performed using 120kVp, 50-200mAs and reconstructed with 5mm slice thickness using 50% overlap. The mean density was 12 HU with a range of -592 to 180 HU. All the measurements were performed using the standard dose thoracic CT, performed on a 64MDCT slice scanner, with 120 kVp, modulated tube current, with 5mm reconstructed slice thickness.

![Density](image)

**Figure 2.8** A histogram showing the number of cases in each density group.
3.2.2 Size

The diameter of each nodule was measured using electronic calipers on the axial CT projection. The largest diameter of the lesion in the cross-sectional image, displayed on a wide window and level setting (W1600 L -600) was measured and recorded. Figure 2.9 demonstrates a histogram of the frequency of lung nodules in each size range. The average size was 2.7 cm and sizes ranged from 0.7 to 13.9 cm.

![Histogram of Lung Nodule Sizes](image)

*Figure 2.9* A histogram showing the number of cases in each nodule size group.

3.2.3 Pathology

Percutaneous CT-guided transthoracic biopsies were performed immediately following the DE/DR imaging exam to provide a definitive diagnosis of the lung lesion. Biopsies were performed either by fine-needle aspiration and cytologic examination or by core biopsy and
histologic examination. The principal lung nodule pathology was non small cell lung cancer (Figure 2.10).

![Pathology Diagram](image)

**Figure 2.10** Illustration of lung nodule pathology. The most common pathology was non-small cell carcinoma and the most common type of non-small cell carcinoma was adenocarcinoma.

The most common type of pathology was non-small cell carcinoma (NSCC), with adenocarcinoma representing the most frequent subtypes of NSCC.

### 3.3 Exam Characteristics

#### 3.3.1 Repeat Breath Hold

Patients were instructed not to breathe during the exams between the low- and high-energy projections, but in 14/220 patients (6%) technical issues created longer than anticipated delay
times (6–8s) between the two projections, and therefore there was a second breath hold. Occasionally breathing artifacts were seen in the resultant images.

### 3.3.2 Time between the Exposures

As mentioned above, a 6–8-second delay usually occurred between the two projections due to detector acquisition delay. The delay time was different from cases to case and in some cases was longer due to technical difficulties. To determine how many cases had more than the usual delay, the frequency of cases in each delay time interval is plotted in Figure 2.11.

![Time Between Exposures](image)

**Figure 2.11** Projections acquisitions frequency for different time delay intervals. As observed in the histogram above, most of the studies were performed with an average delay of 6–9

### 3.3.3 Heart Rate

Patients’ heart rates (HR) varied during low- and high-energy projections. Figure 2.12 below shows patients’ HR variations in high- and low-energy projections.
Figure 2.12 HR variation during high- and low-energy projections. As shown here, most of the patients had a HR of about 60–80 (beat/sec) during projections. But there is a shift towards a lower rate during the high-energy projection, the second acquisition. This could be explained by...
the effect of resulting hypoxemia on the chemo receptors of the carotid body and the following vagal stimulation.

Figure 2.16 below shows the association between HR during low and high projection acquisitions.

![Correlation of heart rates during high and low energy projections](image)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Coefficient</th>
<th>Std. Error</th>
<th>95% CI</th>
<th>t</th>
<th>P</th>
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<td>Intercept</td>
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<td>-5.0894 to 5.0812</td>
<td>-0.001593</td>
<td>0.9987</td>
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<tr>
<td>Slope</td>
<td>0.9684</td>
<td>0.03269</td>
<td>0.9035 to 1.0332</td>
<td>29.6233</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Figure 2.13 Regression analysis. The coefficient of determination for heart rate during low- and high-energy projections. The HRs are highly correlated (R2 = 0.69).
3.4 DE Image Characteristics

3.4.1 Tissue Cancellation Parameters: Ws, Wb, Wc, Wn

Tissue cancellation parameters were selected manually during image decomposition in consensus with an observer who was trained by an experienced thoracic radiologist regarding the optimal presentation of the important anatomical landmarks. Future work will include the use of more detailed, patient-specific, automated and spatially varying parameter selection and will consider incorporation of $w_s$ and $w_b$ as parameters that may be freely varied by the radiologist in a manner analogous to varying the display window/level.

![Bar chart showing distribution of Ws values across different patients.](image)
Figure 2.14 (a, b, c, d) Frequency of cases in different Weighting factor (W) ranges for different decompositions.

3.5 Multi Scale Multi Resolution (MSMR) Registration Algorithm

As mentioned previously, occasional delay between two dual energy projections acquisition and the nature of dual energy techniques (log-weighted subtraction) causes image quality degradation in the resulting dual energy images. This is mainly due to different type of motion artifact including breathing, cardiac and patient motion. Different techniques were used before acquiring the projections to reduce these artifacts as much as possible. These included, instructing patients to hold their breath and hold still, during the high- and low-energy projection acquisitions, and also using cardiac gating techniques (pulse oximeter). For residual problems, registration techniques were used to maximize the coordination of the two images. The second projection (high-energy image) was aligned (registered) onto the low-energy image by means of a deformable registration technique. The registration algorithm operated in multiple passes at
progressively smaller scales and increasing resolution (57). Different variables of the registration technique are discussed below.

### 3.5.1 Mean Vector displacement (MVD) and Standard Deviation Displacement (SDD)

As mentioned above the registration algorithm operates on multiple scales and at multiple resolutions to transform the HE image. A total of four iterative passes were used in the reported registration technique herein where in each pass a series of translation vectors was calculated. A spatial transformation inferred from these vectors, was interpolated and applied to the high energy image in a pixel–wise manner. The transformed HE image and the original LE image then made the input for the next pass where the process was repeated. The mean vector displacement (MVD, mm) for each case was calculated. To do so, for each single ROI, overall X and Y coordinates displacement at the end of final pass were measured the total vector displacement was calculated by means of Pythagorean Theorem: $V_{D} = \sqrt{X^2 + Y^2}$ where $V_{D}$ is the vector displacement and $X$ and $Y$ the overall coordinates displacement.

To calculate the mean vector displacement (MVD) for each single case, all vector displacements for all ROIs (mm) were averaged, over the number of ROIs. The MVD and standard deviation (SD) for each case were plotted against the number of cases in each range as shown below.
Figure 2.15 (a, b) Histograms showing frequency of cases in different ranges of MVD and SD of MVD.
Chapter 3
Satisfaction Test

4 Observer Study

For the observer study described below, the DE images were presented as a “two-slice volume” that could be viewed by mouse wheel scrolling between the soft-tissue and bone images. Five expert observers (3 radiology fellows, 2 radiology staff, each a specialist in chest imaging) were independently presented with the images, one at a time in a randomized order, and asked to rate each image according to the 9-point diagnostic satisfaction scale shown in Table. 3.1

<table>
<thead>
<tr>
<th>Score</th>
<th>Rating</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>Very satisfied</td>
<td>The abnormality is perfectly obvious and easily characterized.</td>
</tr>
<tr>
<td>8</td>
<td>Satisfied</td>
<td>The abnormality is visible and can be well characterized.</td>
</tr>
<tr>
<td>7</td>
<td>Neither satisfied nor</td>
<td>The abnormality is reasonably well seen and characterized.</td>
</tr>
<tr>
<td>6</td>
<td>dissatisfied</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Dissatisfied</td>
<td>The abnormality is visible, but detection and characterization of subtle</td>
</tr>
<tr>
<td></td>
<td></td>
<td>features are a bit challenging.</td>
</tr>
<tr>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Very dissatisfied</td>
<td>The abnormality could be overlooked or mischaracterized.</td>
</tr>
</tbody>
</table>

Table 3.1 Satisfaction rating scale. This table was displayed to observers on a second monitor during all tests as shown in table 3.1.
Scales with fewer (5) and more (up to 100) points were considered in preliminary studies, with 9 found to be tolerated well by observers and consistent within reasonably fine levels of image quality discrimination, as described by Van Metter and Foos. (58) The scale was visible to observers at all times on a second monitor (illustrated in Fig. 2.1).

The satisfaction or preference test performed here is essentially a subjective rank-order evaluation. This method is usually used for answering questions that pertain to different image processing (rendering) variations where the choices tend to be heavily weighted in terms of observer preference, and where detection performance differences is not necessarily expected. (58) An advantage of the approach is that the assessment of overall impression allows for a greater variety of image types to be included more easily into the study, and consequently the results from the study tend to be robust in that can be applied to an extended population of images. This is because the reader response provides an overall impression of the interpretability of the image, and is not just focused for any single specific task. A disadvantage of this approach is that it does not provide objective performance results as there is no "truth" associated with the image that the response is compared against, so the results in a sense are softer. The proponents of this methodology advocate this approach as it has generated consistent results (responses) among readers in their studies. (59) However, a wider validation of this approach has not been performed.

Preceding each observer test, a training session involving 8 DE and 7 DR images was conducted to familiarize the observers with the software and standardize their understanding of the rating scale. The training images were drawn from the pool of cases available at the time of the study and did not overlap with those used as test images. To examine intra-reader consistency during the actual test, the first 7 images displayed in each test were displayed again at the end of the test.
(without informing the observers that the images were repeated), with differences examined in terms of the Wilcoxon signed rank p-value, as described below. The first 7 images were rejected from the study, except for purposes of intra-reader variability; therefore, the analysis pertains to each DE and DR image shown only once (no repeats). Repeat readings were found to be highly reproducible for expert observers in preliminary studies ($p < 0.001$, Wilcoxon signed rank test) and would not add to the statistical power of the study. The study was conducted in a clinical radiology reporting room with subdued lighting on diagnostic-quality; monochrome LCD monitors (AM-QX21-A9300, National Display, San Jose, CA)

5 Statistical Analysis

5.1 Fraction of Responses at or above a Given Rating

The rating scale responses constitute ordinal data. The fraction of observer responses ($F$) at or above a given rating ($R$) was plotted versus the rating scale, giving curves that range 0 to 1 on the vertical axis plotted versus the ordinal rating scale (1 to 9) on the horizontal axis – essentially a cumulative histogram of responses. The fraction at a rating of 1 is 1 by definition. A higher curve for a given modality indicates a greater fraction of higher ratings (a greater degree of diagnostic satisfaction) for that modality.

Error bars on such plots reflect a two-sided 95% confidence interval computed according to a binomial distribution as described below. The corresponding error bars are asymmetric and appropriately bounded between 0 and 1. Each image was scored as either (i) at or above a certain rating, or (ii) below that rating, giving two mutually exclusive outcomes such that $F$ (the ‘fraction at or above a given rating’) follows a binomial distribution. The upper and lower bounds of the confidence intervals were calculated as:
Upper bound: \( \sum_{\kappa=0}^{N} \binom{N}{\kappa} p_U^\kappa (1 - p_U)^{N-\kappa} = \frac{\alpha}{2} \) \hspace{1cm} (2a)

Lower bound: \( \sum_{\kappa=0}^{N-1} \binom{N}{\kappa} p_L^\kappa (1 - p_L)^{N-\kappa} = \frac{1-\alpha}{2} \) \hspace{1cm} (2b)

Where \( N \) is the total sample size, \( N_d \) is the number of samples with the outcome of interest, \( \alpha \) is the level of significance (5%), \( p \) is the proportion of samples with the outcome of interest, \( p_U \) is the upper bound of \( p \), and \( p_L \) is the lower bound of \( p \). The upper and lower bounds were calculated using Newton-Raphson method in Matlab (The Mathworks, Natick MA) to a precision of \( 1 \times 10^{-6} \).

5.2 Statistical Significance: P-value from Wilcoxon Signed Rank Test

The statistical significance in differences observed between DE and DR scores was evaluated in terms of the p-value at a 95% level of significance from a Wilcoxon signed rank test – a non-parametric test suitable to paired ordinal data, assuming all observations within a given modality are independent. (60) The p-value was calculated using the Matlab function ‘sign rank’ accounting for both the sign and magnitude of the difference in ratings.

To examine intra-reader consistency, the Wilcoxon signed rank test was also used in relation to the first and last 7 images in the reading study, repeated at the beginning and end of the test as described above. In this case, the alternative hypothesis was that ‘the two sets of scores are not equivalent;’ therefore, a two-sided p-value was calculated directly from ‘sign rank’. Across 5 observers, all p-values assessing intra-reader consistency were greater than 0.05 (specifically, p-value = 0.125, 1, 0.0625, 0.5313, and 1 for observers A-E, respectively), suggesting that there was no significant difference in observer readings at the beginning and end of the test.
5.3 Stratification of the Results

Performance was analyzed for all cases pooled, as well as by post-hoc stratification of the data according to lesion size, lesion density, chest thickness, gender, and location of the lesion. Lesion size was characterized as the greatest linear dimension as measured on CT, and the results were stratified as “nodule” (≤3 cm) or “mass” (>3 cm), consistent with typical clinical terminology. Lesion density was measured using the attenuation measurement tool on a PACS workstation (Fusion E-film 2.1, Merge Healthcare, Milwaukee, WI), and the results were stratified as “Solid” (≥20 HU) or “Non-solid” (<20 HU). Chest thickness was characterized as the anterior-posterior distance measured from the xiphoid process to the T9 thoracic vertebra taken from the axial CT image at this level, and the results were stratified as “Average” (≤26 cm) or “Thick” (>26 cm). Lesion location was determined according to the anatomical position with respect to lung zones (or mediastinum), and the results were stratified as “Right-Upper,” “Left-Upper,” “Right-Middle,” “Left-Middle,” “Right-Lower,” “Left-Lower,” and “Mediastinum.” In terms of stratification analysis, since multiple comparisons (five) were performed on the same set of data, the threshold level for a significant p value was considered to be less than 0.01 (0.05/5). (61) The number of cases overall and within each stratum is summarized in Table 2.1
Table 3.2 Summary of Cases and Stratifications of Data. The P values relate to the statistical significance in the difference between satisfaction with dual-energy and digital radiographic images, regarded as significant for P < .01.

6 Result
6.1 All Cases Pooled
The results for the 275 total ratings (5 radiologists * 55 cases) for each of the two modalities (DE and DR) are summarized in Fig. 3.1. Individual case-by-case comparison of DE and DR for each patient is evaluated in Fig. 3(a). In 41.5% (114/275) of cases, the DE image was rated superior to
DR by at least a difference of R=1. In 38.9% (107/275) of cases, the DE and DR images were rated equal. In 19.6% (54/275) of cases, the DR image was rated superior. Further to this case-by-case examination, the proportion of cases for which one modality was superior (or equal) to the other, as judged by 3 or more of the 5 observers, is plotted in Fig. 3.1(b). Of these: 36.4% (20/55) scored DE superior to DR; 36.4% (20/55) rated DE and DR equivalent; and 5.5% (3/55) rated DR superior to DE. In the remaining 12 cases (21.8%), a majority could not be reached regarding the superiority / equality / inferiority between the two modalities. The fractions of images rated at or above a given rating score [Fig. 3.1(c)] shows that DE rated consistently higher than DR (p-value < 0.001) in the detection and characterization of lung nodules.
Figure 3.1 Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings (all cases pooled): (a) percentage of image pairs for which the DE image was rated superior, equal, or inferior to the DR image; (b) percentage of image pairs for which the DE image was rated superior, equal, or inferior to the DR image as agreed by three or more observers; (c) fraction of observer responses at or above a given rating score. The error bars represent 95% confidence intervals.
6.2 Stratification by Lesion Size

The data were subsequently analyzed in terms of cases for which the lesion size was ≤ 3 cm and > 3 cm (36 and 19 cases, respectively, as shown in Table II). Results are shown in Fig. 3.2. A statistically significant improvement in diagnostic satisfaction was observed for DE imaging for nodules (lesion size ≤ 3cm, p-value < 0.001). The advantage of DE is more pronounced for nodules, as seen from the distinctly separated curves in Fig. 3.2(a) and the correspondingly smaller p-value.

**Figure 3.2** Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings stratified by lesion size. The curves show the fraction of responses at or above a given rating score for (a) lesion size ≤ 3 cm and (b) lesion size > 3 cm. The error bars reflect 95% confidence intervals.
6.3 Stratification by Lesion Density

Cases were stratified according to lesion density as solid (≥20 HU) and non-solid (<20 HU), with 46 and 9 cases, respectively (Table 3.2). A statistically significant improvement in diagnostic performance was observed for DE imaging of solid lesions (p-value < 0.001). For non-solid lesions, DE and DR scores were not significantly different overall (p-value = 0.0968) however, this finding may have been influenced by the low sample size of non-solid lesions in the pilot study. This finding will be more thoroughly investigated in future work using the entire patient cohort.

![Graphs showing diagnostic satisfaction for non-solid and solid lesions](image)

**Figure 3.3** Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings stratified by lesion density. The curves show the fraction of responses at or above a given rating score for (a) nonsolid lesions (<20 Hounsfield units [HU]) and (b) solid lesions (≥20 HU). The error bars reflect 95% confidence intervals.
6.4 Stratification by Chest Thickness

Results grouped according to PA chest thickness are shown in Fig. 3.4. A statistically significant boost was evident for DE imaging in the detection and characterization of lesions in both categories (viz., p-value < 0.0001 for “average” and p-value <0.002 for “thick”). The curves indicate a fairly uniform improvement in diagnostic satisfaction for cases of “average” thickness (i.e., a uniform boost across all ratings), whereas for the “thick” cases, the curves appear to suggest a boost at the higher ratings (i.e., more conspicuous lesions). This observation suggests that the benefit of DE images may be somewhat less in “thick” than in “average” cases, presumably because images for the former are limited, at least in part, by quantum noise, contrast, x-ray scatter, etc., rather than anatomical clutter.

![Figure 3.4](image)

**Figure 3.4** Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings stratified by chest thickness. The curves show the fraction of responses at or above a given rating score for (a) thickness< 26 cm and (b) thickness > 26 cm. The error bars reflect 95% confidence intervals.
6.5 Stratification by Gender

Cases were further grouped based on gender – a total of 31 male and 24 female patients. Results in Fig. 3.5 suggest a significant improvement in diagnostic performance for DE in each case (p-value < 0.001 for male and p-value = 0.013 for females). Such is likely consistent with the trend for improved performance overall for DE (Fig. 3.1). The smaller level of improvement suggested for the female sub-cohort (while still a statistically significant improvement) is possibly related to that observed for larger chest thickness (in this case, breast tissue), which correlated with a smaller improvement in diagnostic performance.

Figure 3.5 Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings stratified by patient gender. The curves show the fraction of responses at or above a given rating score for (a) men and (b) women. The error bars reflect 95% confidence intervals.
6.6 Stratification by Region

Results grouped according to 7 regions of the chest (left and right apex, left and right middle, left and right lower, and mediastinal regions) are shown in Fig. 3.6. Although the number of nodules is relatively small in some of the sub-groups, therefore the findings should be viewed with some degree of caution, there is a trend for improved diagnostic performance for DE imaging in the left apex, right apex, left-middle, and left-lower regions. The results for the apical regions are consistent with the hypothesis that DE imaging improves diagnostic quality by removing anatomical noise – in this case, the clavicles and 1st and 2nd ribs, which exhibit complex anatomical clutter on frontal chest radiographs and can significantly diminish conspicuity. The significant improvement observed for DE imaging in the left-lower region is somewhat surprising, given that this region is challenged by a preponderance of soft-tissue structures (the heart) and is most susceptible to cardiac motion artifacts. Figures 3.7, 3.8 and 3.9 illustrate significant improvement of DE compared to DR in detection of a small solid nodule and nodules in right and left upper lobes. Images a, b and c are illustrating examples of soft only, bone only and DR images.
Figure 8. Diagnostic satisfaction in DE (Dual energy) and DR (Digital radiography) image readings (by lesion location). The curves show the fraction of responses at or above a given rating score for lesions located in (a) Right-Upper, (b) Left-Middle, (d) Left-Middle, (e) Right-Lower, and (g) Mediastinal regions. The error bars reflect a 95% confidence interval.

- Right-Upper (n=100) p-Value = 0.0012
- Left-Upper (n=25) p-Value = 0.0067
- Right-Middle (n=20) p-Value = 0.18
- Left-Middle (n=10) p-Value = 0.0313
- Right-Lower (n=40) p-Value = 0.3625
- Left-Lower (n=45) p-Value = 0.001
- Mediastinal (n=35) p-value = 0.2455
Figure 3.6 Diagnostic satisfaction in dual-energy (DE) and digital radiographic (DR) image readings stratified by lesion location. The curves show the fraction of responses at or above a given rating score for lesion located in (a) right upper, (b) left upper, (c) right middle, (d) left middle, (e) right lower, (f) left lower, and (g) mediastinal regions. The error bars reflect 95% confidence intervals. For lower satisfaction scores (≤5) reflecting lower satisfaction, there is no clear pattern however for higher satisfaction scores (>5), DE images demonstrate an advantage over DR.

Figure 3.7 Illustration of (a) dual-energy soft tissue, (b) dual-energy bone, and (c) digital radiographic images for a case exhibiting a small, solid nodule (0.8 cm, 58.4 Hounsfield units, benign lymphoid tissue), marked by the arrows. The bone image exhibits some residual soft tissue attributable to incomplete cancellation as well as motion artifact, which is more obvious in this case because of the cardiac motion artifact.
Figure 3.8 Illustration of (a) dual-energy soft tissue, (b) dual-energy bone, and (c) digital radiographic images for a case exhibiting a nodule (2 cm, 67.5 Hounsfield units, adenocarcinoma) in the right upper lobe, marked by the arrows.

Figure 3.9 Illustration of (a) dual-energy soft tissue, (b) dual-energy bone, and (c) digital radiographic images for a case exhibiting a nodule (2.8 cm, 35.1 Hounsfield units, adenocarcinoma) in the left apex, marked by the arrows.
Chapter 4
ROC Test

1 Study Population

As all 129 subjects were recruited from a population referred for biopsy of suspicious lung lesions (confirmed by CT prior to biopsy), all patients in the study cohort had at least one lung nodule. A pre-requisite of ROC analysis is to present a random series of images that contain the target and some that do not. Therefore, given the study cohort, the chest radiographs were divided to provide half-chest images, where each image was cropped near the vertical midline of the spine to present a complete image of just one lung. Although the presentation of half-chest images somewhat perturbs the usual presentation of chest images and therefore the usual radiologist reading pattern, observers reported no difficulty in assessing the cropped images.

Of the 129 cases, 29 exhibited bilateral disease; therefore, the 129 cases provided 258 half-chest images of which 158 contained nodules (“disease”), and 100 did not (“normal”). Of the 158 disease cases, 55 had multiple lesions, and 103 had only one lesion. The average size of the lesions (maximum diameter measured on CT) was 2.7 cm, ranging from 0.7 to 14 cm. The average density of lesions (measured in CT) was 11.5 HU, ranging from -769 HU to +180 HU.

Visualization software (termed “DEvice”) was developed by the research team, to allow viewing of all three DE image components in a single, spatially co-registered view – i.e., soft-tissue, bone, and composite images viewed as a three-slice “stack”. The software provided the ability to “scroll” between DE image components (using the mouse wheel) and to “see through” any given component to another (using a digital “looking glass” loupe invoked on the display) in addition to common image manipulation tools (e.g., window/level, magnification, pan, etc.). These software tools allowed fast, convenient visualization and comparison of anatomy presented
among various image components. The composite image was the initial view (presenting a familiar radiographic scene with anatomical landmarks, etc.); via the mouse wheel (or the “looking glass” loupe), the reader could scroll to the soft-tissue and bone component images to more closely evaluate subtle image characteristics without the confounding influence of overlying anatomical noise from other tissue components. In this way, the display software better exposed the intrinsic potential advantage associated with multiple tissue components in DE imaging (without forcing the reader to pan between side-by-side images).

Five expert observers (3 staff radiologists and 2 radiology fellows, each a specialist in thoracic imaging) were independently presented with the images. The reading order was randomized, with one image presented at a time [either a DR image or a DE image (with all 3 components co-registered in DEvice as described above)]. A diagnostic-quality monochrome LCD monitor (AM-QX21-A9300, National Display, San Jose, CA) calibrated to meet the DICOM grayscale standard was used. Studies were conducted in a dark-controlled radiology reading room (~0.15 Cd/m² ambient light). Each observer was asked to rate each image in terms of the detection of abnormal lung lesions according to the 5-point scale in Table. 4.1. The scale was visible to observers on a second monitor. As the test was not a localization-specific ROC (LROC), observers were asked to score the lung image as a whole and did not record the exact location of the abnormalities. This imparts a potential limitation of the current study with respect to multiple nodules within one lung and the possibility of true-positive responses based on mistaken identification of a normal structure within an abnormal lung and not necessarily the abnormality itself. The possible error associated with this limitation is an overestimation of sensitivity, believed to be small (in part due to the conspicuity of actually-positive cases, as noted below). A training session involving 84 images (42 DE and 42 DR) was conducted prior to each observer study to familiarize the observers with the software. The training images were drawn from Group
3 (non-cardiac-gated sub-group), which were the same as Group 1 in image acquisition and decomposition technique but did not overlap with cases used in the actual observer study. A viewing distance of ~50 cm was encouraged but not enforced, and observers were free to adjust window/level, zoom, etc. as well as to scroll between DE image components or use the DE “looking glass.” Tests required an average of 150 min to complete (18 sec per image), with one break permitted mid-way through the test.

2 ROC Analysis

2.1 ROC Curves and Area under the Curve (AUC)

Receiver Operating Characteristic (ROC) analysis which is frequently used to compare the diagnostic qualities of imaging systems (62) was performed using MedCalc (version 11.3, MedCalc Software bvba, Belgium). The resulting ROC curves plot the true-positive fraction (TPF, or sensitivity) versus the false-positive fraction (FPF, or 1-specificity), with each point along the curve representing TPF and FPF at a different decision threshold. For a five-point ROC rating scale as in Table 4.1, the corresponding four decision thresholds yield four measured control points on the ROC plot.(62)

<table>
<thead>
<tr>
<th>Receiver-Operating Characteristic Rating Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Definitely normal (or almost definitely normal)</td>
</tr>
<tr>
<td>2. Probably normal</td>
</tr>
<tr>
<td>3. Possibly abnormal</td>
</tr>
<tr>
<td>4. Probably abnormal</td>
</tr>
<tr>
<td>5. Definitely abnormal (or almost definitely abnormal)</td>
</tr>
</tbody>
</table>

Table 4.1 Five-point Rating Scale for Receiver-Operating Characteristic Studies
The majority of ROC analysis used in radiological studies uses a discrete rating scale with five or six categories. The discrete five point rating scale has been compared to a continuous confidence scale by Rockette et al (63) and the measured area under the curve (AUC) was not significantly different between the two different methods, indicating that either could be used in image evaluation studies. Hadgiiski et al (65) examined the effects of the number of categories in the rating scale on the result of ROC. They concluded that the fitted curve and the performance indices do not vary significantly with different rating scales, only if the estimated operating points, obtained from the data, are relatively evenly distributed over the entire range of TPF and FPF. According to their results, ROC analysis of discrete confidence rating scales with few and unevenly distributed data points may cause unreliable estimation. The area under the curve is basically created by varying the cut point (decision making threshold) that is used to determine the values of the observed variables. If a test has a perfect discrimination capability, it would have a threshold value that would separate the entire abnormal population from the entire normal population (or vice versa). A graphical display would plot this kind of curve through (0, 1) (62). This would be the ideal point and as the observed curve conforms closer to this ideal form, it demonstrates improved discrimination between the normal and abnormal variables. If a test cannot discriminate at all it would follow a diagonal course within the grid. The area under the curve represents the probability that, if a randomly selected individual from the abnormal population and a randomly selected individual from the normal population are selected the observed values would be in correct order. Parametric assumptions are usually applied and the maximum likelihood method are used to estimate the area under the curve (65, 66) Figure 4.1 below shows the distribution of the observed values based on the 5 points rating scale, by all 5 observers and for two modalities, DE and DR. Since the observed values from our study did not follow a normal distribution, a non-parametric approach was chosen to estimate the AUC and to
compare the areas under the correlated ROC curves with a non-parametric approach; trapezoidal rules apply to measure the area under the points comprising an empirical ROC curve. (67) The Mann-Whitney statistical test was used to compare the area under the two correlated curves.

Figure 4.1 The distribution of rating scores among all observers for Dual energy (DE) and DR. As observed on the graphs, the distribution is non-parametric and therefore non-parametric ROC was applied.
To measure the inter rater agreement, weighted kappa was used for rating scores of 5 observers and for each modality separately. The average weighted kappa for DE was calculated as $k = 0.51$ and for DR as $k = 0.52$. These numbers considered to fall under moderate agreement category.

Performance was analyzed for all 258 cases pooled, and by post-hoc stratification of the data according to lesion size, lesion density, chest thickness, lesion location, and patient gender. Table 4.2 shows a summary of the strata. Results were analyzed for all strata by individual observer and for all observer responses pooled.

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Description</th>
<th>DE</th>
<th>DR</th>
<th>How Determined</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pooled</td>
<td>All cases</td>
<td>258</td>
<td>258</td>
<td>N/A</td>
</tr>
<tr>
<td>Lesion size</td>
<td>Mass (&gt;3 cm)</td>
<td>44</td>
<td>44</td>
<td>Maximum linear dimension measured in CT</td>
</tr>
<tr>
<td></td>
<td>Nodule (≤3 cm)</td>
<td>114</td>
<td>114</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nodule (≤1 cm)</td>
<td>18</td>
<td>18</td>
<td></td>
</tr>
<tr>
<td>Lesion density</td>
<td>Solid (≥20 HU)</td>
<td>128</td>
<td>128</td>
<td>Mean HU measured in CT</td>
</tr>
<tr>
<td></td>
<td>Non-solid(&lt;20 HU)</td>
<td>30</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Chest thickness</td>
<td>Large(&gt;26 cm)</td>
<td>68</td>
<td>68</td>
<td>AP distance (T9-xiphoid) measured in axial CT</td>
</tr>
<tr>
<td></td>
<td>Average(≤26 cm)</td>
<td>190</td>
<td>190</td>
<td></td>
</tr>
<tr>
<td>Lesion location</td>
<td>Right-upper lobe</td>
<td>48</td>
<td>48</td>
<td>Belonging to a given anatomical region as visualized in CT</td>
</tr>
<tr>
<td></td>
<td>Left-upper lobe</td>
<td>29</td>
<td>29</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Right-middle lobe</td>
<td>8</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lingula</td>
<td>4</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Right-lower lobe</td>
<td>26</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Left-lower lobe</td>
<td>30</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mediastinum</td>
<td>13</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>Gender</td>
<td>Male</td>
<td>160</td>
<td>160</td>
<td>Patient chart</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>98</td>
<td>98</td>
<td></td>
</tr>
</tbody>
</table>
Table 4.2 Summary of cases overall and within each post-hoc stratification.

3 Results

3.1 Individual Observers

The results of ROC tests for 5 individual observers (258 DE, 258 DR each) are summarized in Table 4.3, and the ROC curves for each observer are illustrated in Fig. 4.2, showing the fitted ROC as well as the coordinates (empiric ROC).

<table>
<thead>
<tr>
<th>Observer</th>
<th>AUC&lt;sub&gt;DE&lt;/sub&gt;</th>
<th>AUC&lt;sub&gt;DR&lt;/sub&gt;</th>
<th>P Value In AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observer 1</td>
<td>0.779</td>
<td>0.777</td>
<td>0.9345</td>
</tr>
<tr>
<td>Observer 2</td>
<td>0.778</td>
<td>0.756</td>
<td>0.4279</td>
</tr>
<tr>
<td>Observer 3</td>
<td>0.769</td>
<td>0.782</td>
<td>0.6519</td>
</tr>
<tr>
<td>Observer 4</td>
<td>0.758</td>
<td>0.763</td>
<td>0.7633</td>
</tr>
<tr>
<td>Observer 5</td>
<td>0.854</td>
<td>0.833</td>
<td>0.1842</td>
</tr>
</tbody>
</table>

Table 4.3 The area under the curve (AUC) for each observer is shown, found to be superior for dual energy (DE) imaging in three of five observers but not to a significant extent (two-tailed P value based on AUC).

The AUC for DE imaging was superior to DR for 3 out of 5 observers (two staff and one fellow) although the difference was not statistically significant. As described below, a similar result was found by pooling over all observers.
Dual energy (DE) outperformed digital radiography (DR) for three of five readers (a, b, and e), although the difference in performance was not statistically significant ($P > .05$).

**Figure 4.2** Receiver-operating characteristic (ROC) curves for five observers. Dual energy (DE) outperformed digital radiography (DR) for three of five readers (a, b, and e), although the difference in performance was not statistically significant ($P > .05$).
3.2 All Observers Pooled

Results for the 1290 total ratings (5 radiologists x 258 cases) for each of the two modalities (DE and DR) are summarized in Table 4.4. The statistical significance in observed differences was evaluated in terms of the entire AUC (total area under the curve). Overall, the AUC for DE imaging was higher than DR in the following cases: all cases pooled; small chest thickness; female; nodule diameter ≤ 3 cm; solid nodules; and nodules located in the right-upper lobe (RUL) and right-lower lobe (RLL). However, the difference was statistically significant only for cases of nodule diameter ≤ 1 cm (p-value=0.02), and RUL (p-value =0.002). A statistically significant difference was also seen in the LLL but here DR outperformed DE (p-value=0.05).

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Description</th>
<th>AUC&lt;sub&gt;DE&lt;/sub&gt;</th>
<th>AUC&lt;sub&gt;DR&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pooled</td>
<td>All cases</td>
<td>0.781</td>
<td>0.776</td>
</tr>
<tr>
<td>Lesion size</td>
<td>Mass (&gt;3 cm)</td>
<td>0.797</td>
<td>0.822</td>
</tr>
<tr>
<td></td>
<td>Nodule (≤3 cm)</td>
<td>0.775</td>
<td>0.758</td>
</tr>
<tr>
<td></td>
<td>Nodule (&lt;1 cm)</td>
<td>0.758</td>
<td>0.689</td>
</tr>
<tr>
<td>Lesion density</td>
<td>Solid (≥20 HU)</td>
<td>0.805</td>
<td>0.798</td>
</tr>
<tr>
<td></td>
<td>Non-solid (&lt;20 HU)</td>
<td>0.678</td>
<td>0.682</td>
</tr>
<tr>
<td>Chest thickness</td>
<td>Large (&gt;26 cm)</td>
<td>0.741</td>
<td>0.758</td>
</tr>
<tr>
<td></td>
<td>Average (≤26 cm)</td>
<td>0.798</td>
<td>0.784</td>
</tr>
<tr>
<td>Lesion location</td>
<td>Right upper lobe</td>
<td>0.804</td>
<td>0.759</td>
</tr>
<tr>
<td></td>
<td>Left upper lobe</td>
<td>0.801</td>
<td>0.808</td>
</tr>
<tr>
<td></td>
<td>Right middle lobe</td>
<td>0.808</td>
<td>0.839</td>
</tr>
<tr>
<td></td>
<td>Lingula</td>
<td>0.898</td>
<td>0.933</td>
</tr>
<tr>
<td></td>
<td>Right lower lobe</td>
<td>0.770</td>
<td>0.738</td>
</tr>
<tr>
<td></td>
<td>Left lower lobe</td>
<td>0.766</td>
<td>0.806</td>
</tr>
<tr>
<td></td>
<td>Mediastinum</td>
<td>0.578</td>
<td>0.609</td>
</tr>
<tr>
<td>Gender</td>
<td>Male</td>
<td>0.773</td>
<td>0.775</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>0.798</td>
<td>0.782</td>
</tr>
</tbody>
</table>
The total number of readings for each modality was 1290 (equal to 258 cases - 5 radiologists), with 790 actually positive and 500 actually negative cases. Results were analyzed in terms of the AUC for dual energy (DE) imaging and digital radiography (DR) (area under the curve AUC DE and AUC DR, respectively). Statistical significance in the observed differences were analyzed in terms of the P value associated with the entire AUC. Statistically significant results are highlighted in bold text.

The ROC curves for the two strata with statistically significant difference between the two modalities are illustrated in Fig. 4.3. For small nodules (diameter ≤ 1 cm, Fig. 4.3a), DE imaging exhibited improved AUC (p-value=0.03). This observation suggests a clinical advantage in discriminating small, benign lesions with DE. The improvement in AUC for DE demonstration of lesions in the RUL (Fig. 4.3b) is consistent with the notion that DE imaging improves delectability in regions dominated by anatomical clutter; in this case the ribs and clavicles. The lack of a similar finding in the LUL may be due to the smaller number of cases in that stratum. Figures 4.4 and 4.5 show example images for which DE imaging clearly outperformed DR, cases of a small nodule (diameter ≤ 1 cm) and a nodule in the RUL, respectively.

<table>
<thead>
<tr>
<th></th>
<th>AUC</th>
<th>SE</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DE</strong></td>
<td>0.758</td>
<td>0.0293</td>
<td>0.700 to 0.815</td>
</tr>
<tr>
<td><strong>DR</strong></td>
<td>0.689</td>
<td>0.0310</td>
<td>0.628 to 0.750</td>
</tr>
<tr>
<td><strong>Difference between areas</strong></td>
<td>0.0684</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Standard Error</strong></td>
<td>0.0295</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>95% Confidence Interval</strong></td>
<td>0.0105 to 0.126</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>z statistic</strong></td>
<td>2.317</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Significance level</strong></td>
<td>P = 0.0205</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
b) **Figure 4.3 a, b.** Receiver-operating characteristic (ROC) curves for five nodules smaller than 1 mm (a) and nodules located in the Right Upper Lobe. (b) DE outperformed DR in these two categories (P < .05)

![Figure 4.3](image)

<table>
<thead>
<tr>
<th></th>
<th>AUC</th>
<th>SE</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>DE</td>
<td>0.804</td>
<td>0.0206</td>
<td>0.763 to 0.844</td>
</tr>
<tr>
<td>DR</td>
<td>0.759</td>
<td>0.0220</td>
<td>0.716 to 0.802</td>
</tr>
</tbody>
</table>

**Difference between areas**

<table>
<thead>
<tr>
<th>Standard Error</th>
<th>0.0176</th>
</tr>
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**95% Confidence Interval**

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**z statistic**

2.538

**Significance level**

P = 0.0112

**Figure 4.4** Example of dual energy (DE) and digital radiography (DR) images for a small nodule. The location of a 0.9 cm nodule in the right upper lobe with density of 28.5 HU is marked by an arrow.
3.3 Diagnostic Accuracy and Clinical Significance

Table 4.5 shows the overall accuracy of DE and DR imaging in discriminating abnormal and normal cases. For the purposes of this study, accuracy was defined as the capability of the reader to correctly identify a case as true-positive or true-negative (i.e., a rating of 1 or 2 in normal cases or 3-5 in abnormal cases). The term “accuracy” as defined here is equivalent to the fraction of correctly identified cases as commonly defined \([\frac{(TP+TN)}{(TP+TN+FP+FN)}]\), a reasonable figure of merit for tests in which the number of normal and abnormal (disease) cases are approximately the same. For disease cases, \(~0.31\) (49/158 readings) were missed in DR but were properly identified in DE. Conversely, 0.26 (41/158) of disease cases were missed in DE but were properly identified in DR [DE and DR agreed (i.e., both \(\geq 3\) or both \(\leq 2\)) in the remainder of cases.] These data suggest slightly improved sensitivity for DE imaging to an extent that could avoid false negative findings in \(~5\%\) (8/158) of cases. For normal studies, 0.24 (24/100) were false positive cases in DR but were properly identified in DE. Conversely, 0.33 (33/100) were false positive cases in DE but were properly identified in DR. These data suggest a reduction in
specificity associated with the increased sensitivity of DE imaging to an extent that would cause false-positive readings in 9% (9/100) cases. While such tentative results would benefit from a dedicated multi-center trial (in which the importance of the bone image in detecting subtle calcification might become more evident), they suggest the extent to which DE imaging could indicate a clinically distinct diagnostic path for patients, suggestive of a clinically significant difference in up to ~10% of patients, with ~5% of patients benefiting from early nodule detection.

Table 4.5 also compares the overall definitiveness of DE and DR imaging in identifying disease and normal cases. The definitiveness is defined as the highest level of reader confidence in identifying or rejecting an abnormality (i.e., a rating of 1 or 5). The number of cases “definitely” identified in one modality was analyzed (i.e., a rating of 1 or 5 for TP or TN, respectively) relative to a less definitive score for the other modality (i.e., a rating ≥2 or ≤4 for disease or normal cases, respectively). For disease cases, ~0.29 (46/158 readings) were more definitively identified in DE, compared to 0.18 (29/158) for DR. These data suggest an improvement in the definitiveness of positive diagnosis for DE compared to DR, where DE imaging gave more definitive scores in ~11% of cases (17/158) that were either missed or were less certainly scored using DR. Normal cases, 0.46 (46/100) were more definitively rejected (ROC score = 1) by DE imaging than in DR, compared to 0.44 (44/100) more definitively identified in DR. These data suggest a slight improvement in definitive specificity for DE to an extent that avoided false positive readings in 2% of cases (2/100). Improved definitiveness of diagnosis (in both normal and disease cases) may be particularly important in clinics without access to CT, and it may present a cost-effectiveness advantage by reducing the transfer of suspicious cases to better equipped centers.
Table 4.5 Accuracy, Definitiveness, and Clinical Significance, Dual energy (DE) imaging was more accurate than digital radiography (DR) in identifying actually positive cases but less accurate in ruling out actually negative cases. DE imaging was more definitive in properly identifying both actually positive and actually negative cases.
Chapter 5
Conclusion and Future Work

DR imaging of the chest represents a cost-effective, widely available, low-dose modality used for a broad spectrum of applications, ranging from bedside exams to the initial examination and diagnosis of lung disease. Still, it is known to suffer in sensitivity for the detection of subtle lesions, limited primarily by a lack of conspicuity caused by superimposition of anatomic structures. DE imaging, which reduces anatomic clutter by selectively removing material specific components from the image, showed a significant improvement in satisfaction ratings associated with the detection and characterization of pulmonary nodules. DE imaging by selective decomposition and visualization of soft tissue and bone images reduces anatomical noise and demonstrates an increased sensitivity for the detection of lung nodules especially in the areas that are challenging for DR, such as the lung apex (68). This study evaluated the performance of DE imaging using a preference test and an ROC study involving 5 observers. The preference test was initially evaluated in 55 patients drawn from an ongoing trial; it provided initial investigation of diagnostic performance and supported the hypothesis that DE imaging boosts lesion conspicuity.

The differences in satisfaction between DE and DR imaging are shown in Figures 3.1 to 3.6, with qualitative differences illustrated in the patient images in Figures 3.7 to 3.9. For example, the difference observed for small, solid nodules (<3 cm diameter, 20 HU density) in Figures 3.2 and 3.3 is illustrated qualitatively in Figure 3.7. The improved visualization of small, solid nodules by virtue of rib cancellation is clear. Figures 3.8 and 3.9 similarly illustrate improved nodule visibility in the lung apex, in which the clavicles, first and second ribs present confounding clutter that is significantly reduced in the DE soft tissue image. That the difference in performance was greatest for small, apical solid nodules is particularly valuable, because these characteristics represent precisely the most challenging (i.e., small), important (i.e., solid, more
likely malignant), and frequent (i.e., location in lung apex) cases and the areas most in need of improvement in chest imaging.

The initial results were encouraging, although the study was not without its limitations. First, the number of cases was low, particularly for certain stratifications and subgroup analysis (e.g., anatomic region), because these early data constitute an initial study from an ongoing trial. Furthermore, the stratifications within the data were post hoc, and although the overall study was prospectively designed to investigate the difference in performance between DE and DR imaging, the data pertaining to individual strata (i.e., nodule size, nodule density, chest thickness, gender, and nodule location) should be considered “hypothesis generating” in the sense of retrospective analysis. Furthermore, the strata were distinct and independent (e.g., grouping small or solid or apical nodules, but not small, solid, and apical nodules), and joint grouping was not examined. Second, the satisfaction test on the basis of a nine-point scale has not been validated in a wider range. Although the intra observer agreement was testes for all observers at the beginning of the study and the repeat readings were found to be highly reproducible, the inter observer variability and validity of the test was not quantified. Two previous studies have found the same approach to generate consistent results between the readers but a thorough evaluation of the overall strength of the psychometric analysis of this method has never been performed. The Statistical significance was evaluated in terms of P values obtained from Wilcoxon’s signed-rank tests. A limitation associated with clustering effects within ratings for the same patient across observers is possible. A third limitation of the study involves the DE image processing and decomposition. Although the DR images were post processed according to techniques established for optimal clinical DR imaging, the DE image post processing was fairly simple. The DE images used in the observer study used the simple image processing and decomposition techniques specifically, single-point offset and gain corrections, non-optimized registration of
low- and high-energy projections, and simple log weighted subtraction according to a scalar tissue cancellation parameter.

Overall, DE imaging had a superior diagnostic performance in the satisfaction test than DR imaging at equivalent radiation dose. This was mainly attributed to the soft tissue images that present subtle lesions more conspicuously by virtue of reduced anatomic clutter. Although the observer response (rating) pertained to the combined DE image set (i.e., the soft tissue and bone images considered together, rather than each rated individually), it was clear that the soft tissue image was the more important in nodule detection, while the bone image presented complementary information regarding characterization (e.g., calcification). For this patient cohort in particular (drawn from a clinical patient population referred for a lung nodule biopsy), there were few cases exhibiting calcified nodules. Therefore, the bone images were likely used less than would be the case in a general screening population (in which the frequency of calcified nodules would presumably be greater). Furthermore, the bone image could provide diagnostic value regarding bony pathology, for example in differentiation of rib metastasis from rib fracture and improved visualization of fine bony detail to exclude cortical invasion.

The ROC study evaluated the performance of DE imaging involving 5 observers and 129 cases with half-chest images, resulting in 258 images. DR images were compared to DE images consisting of three components: a soft-tissue image, a bone image, and a composite image comparable to DR. The techniques were compared by measuring the total area under the curve (AUC) in an ROC analysis. Overall, DE outperformed DR in detection of nodules smaller than 1 cm, and nodules in the RUL. Superior diagnostic performance for DE imaging in the RUL was likely due to the removal of anatomical noise related to the clavicle, first, and second ribs in this region which cause significant limitations in detection of parenchymal abnormality by DR. This
problem is reduced through tissue decomposition techniques (i.e., bone cancellation) in DE imaging. It is therefore somewhat surprising that similar results were not realized for lesions in the LUL, which could reflect the smaller number of cases with LUL lesions in this study cohort.

The results were promising, but the study was not without limitations. First, the study subjects were recruited from a population suspicious for lung cancer and referred for CT-guided percutaneous lung biopsy. Most of these cases had lung nodules that were initially detected on CR or DR chest radiographs and therefore had fairly conspicuous nodules; this observation might have limited the potential advantage of DE imaging. However, DE imaging demonstrated a relatively high sensitivity and specificity for small (<1 cm) nodules. This technique may be useful in evaluating high-risk populations for lung malignancy – for example, cigarette smokers, who exhibit a high incidence of small nodules on low dose CT, which is known to be sensitive but non-specific. The vast majority of nodules detected on low-dose CT are benign, but detection of these nodules generates significant expense, the studies are time-intensive to read, and are performed at a relatively high radiation dose. Dual-energy imaging has the potential to overcome many of these deficiencies and also demonstrates superior performance to DR imaging for RUL nodules, an area where early lung cancers are commonly detected late due to obscuration by the overlying rib or clavicle.

Second, the observers as chest radiologists were all aware of the modality they were evaluating during the study (due to specific features of each modality) and the use of half-chest images, although necessary for the purposes of this study, is a distortion of the usual clinical reading pattern used by thoracic radiologists. These aspects of the study could have biased the results, presumably to an equivalent degree for both DE and DR.
The third limitation relates to the relatively small numbers of cases available for retrospective subset analysis, and this might have limited the ability of this study to distinguish real differences between DE and DR imaging. In addition, although the study was designed as a prospective trial, the stratification was performed post hoc and therefore was retrospective.

Overall (all cases and readers pooled), DE imaging performance was equal to DR at equivalent radiation dose; Although DE imaging requires the assessment of a greater number of images compared to DR imaging, this will be compensated for by an increased level of diagnostic confidence in lesion detection, which in turn should translate into the earlier detection of disease. In addition, the characteristics of DE imaging offer promise in other areas of thoracic disease, such as the earlier detection of airspace disease (pneumonia in patients with fever of unknown origin), the improved demonstration of airway disease (bronchiectasis in patients with chronic productive cough), and the improved visualization of catheters, tubes, and pneumothoraces in patients in intensive care units. (69)

The clinical role of DE is yet to be fully understood. The lateral view, for example, is an important aspect of diagnostic chest imaging (e.g., to visualize retrohepatic and retrocardiac lung), but the performance of DE imaging in the lateral view remains to be evaluated. Similarly, the equivalence of the DE “composite” image and a DR image obtained at equivalent total dose is yet to be established. Clinically, high-performance DE imaging at dose equivalent to DR may yield a new normal means of chest radiography, but it does not completely resolve the lack of sensitivity exhibited by DR as a screening modality; rather, the value of DE imaging is likely to be a better first read. Finally, the performance and clinical role of DE imaging with respect to low-dose CT (LDCT) is yet to be fully assessed. While not intended in this work as a
replacement to LDCT, the use of DE imaging as an adjuvant examination that could help resolve the lack of LDCT specificity is a potentially promising avenue to be investigated.

As an extension of this work, the application of DE imaging in portable radiography is currently under investigation. While maintaining the positive characteristics of DR – i.e., low cost and accessibility – DE imaging could potentially improve the image quality in portable exams. The removal of anatomical noise has even greater importance in this clinical situation where there are often limitations in lesion detectability due to extremely poor image quality in existing portable imaging technologies. Such situations include patients that are unable to be transferred to the radiology department, are obtunded, often examined in a sub-optimal position (supine or rotated), and exhibit limited inspiratory effort, causing under inflation of the lungs. In this study, thoracic DE imaging, performed at equivalent radiation dose to DR, demonstrated superior performance for the detection of small lung nodules that are usually challenging to detect, and the right upper lobe is a common area for missed lung cancers. (68) DE imaging also demonstrated a clinically significant advantage (i.e., a correct change in the diagnostic trajectory of positive cases) as well as more definitive diagnosis of both positive and negative cases.
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42. Berenner DJ, Hall EJ. “Computed tomography- an increasing source of radiation exposure” N Engl J Med. 2007; 357(22):2277-84


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