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Hyperpolarized 129Xe Magnetic Resonance Imaging of Radiation-Induced Lung Injury

Ozkan Doganay  
*The University of Western Ontario*

Supervisor  
Giles Santyr  
*The University of Western Ontario*

Graduate Program in Medical Biophysics

A thesis submitted in partial fulfillment of the requirements for the degree in Doctor of Philosophy

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Hyperpolarized $^{129}\text{Xe}$ Magnetic Resonance Imaging of Radiation-Induced Lung Injury

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by

Ozkan Doganay

Graduate Program in Medical Biophysics

A thesis submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy

The School of Graduate and Postdoctoral Studies
The University of Western Ontario
London, Ontario, Canada

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Abstract

Lung cancer is the largest contributor to cancer-related mortality worldwide. Only 20% of stage III non-small cell lung cancer patients survive after 5-years post radiation therapy (RT). Although RT is an important treatment modality for lung cancer, it is limited by Radiation-Induced Lung Injury (RILI). RILI develops in two phases: (i) the early phase (days-weeks) referred to radiation pneumonitis (RP), and (ii) the late phase (months). There is a strong interest in early detection of RP using imaging to improve outcomes of RT for lung cancer. This thesis describes a promising approach based on $^{129}$Xe gas as a contrast agent for Magnetic Resonance Imaging (MRI) of the lung airspace due to the large increase in signal possible by spin exchange optical pumping, or hyperpolarization (Hp). Additionally, $^{129}$Xe provides unique functional information due to its relatively high solubility and significant chemical shift in pulmonary tissue (PT) and red blood cell (RBC) compartments. In this thesis, a specialized Hp $^{129}$Xe MRI method was developed for detection of gas exchange abnormalities in the lungs associated with thoracic RT. In particular, the feasibility of quantifying the early phase of RILI is demonstrated in a rat model of RILI two weeks post-irradiation with a single fraction dose of 18 Gy. The challenge of low signal-to-noise ratio (SNR) in the dissolved phases was addressed in this work by development and construction of a Transmit-Only/Receive-Only radiofrequency coil. Another challenge addressed in the thesis was the lack of imaging techniques that provide sufficient spatial and temporal information for gas exchange. Therefore, a novel Hp $^{129}$Xe MRI technique was developed based on the multi-point IDEAL pulse sequence. The combination of these two developments enabled investigation of regional gas exchange changes associated with RP in the rat lung two weeks post-irradiation to assess the feasibility of early detection of RILI. Theoretical analysis of the gas exchange curves enabled measurements of average PT thickness ($L_{PT}$) increases consistent with histology and relative blood volume ($V_{RBC}$) reductions in the irradiated animal cohort compared to a non-irradiated cohort, and between irradiated right lungs compared to unirradiated left lungs in the irradiated cohort.

Keywords

Radiation-induced lung injury, pneumonitis, Hyperpolarized $^{129}$Xe, MRI, lung imaging, IDEAL, dissolved xenon, pulmonary tissue, red blood cell, gas exchange, spiral, point spread function, rat, radiofrequency coil.
Co-Authorship Statement

This thesis is based on work published in the following three manuscripts:


For this manuscript, Ozkan Doganay was responsible for optimizing the sensitivity using the numerical COMSOL simulations as well as building both transmit and receive coils and corresponding electronic circuits, bench top measurements, experimental data acquisition and analysis of data including MATLAB coding. Dr. Kundan Thind provided meaningful discussion. Dr. Trevor Wade and Dr. Alexei Ouriadov helped with MRI calibration experiments and *in vivo* imaging. Dr. Giles Santyr provided guidance throughout the project and assistance with the manuscript preparation. Additionally, Elaine Hegarty helped with animal preparation and Krzysztof Wawrzyn helped with Hp $^{129}$Xe gas production.


For this manuscript, Ozkan Doganay was responsible for the theoretical analysis and derivation of the corresponding equations as well as design and optimization of spiral trajectories and corresponding gradient waveforms, experiment design, data acquisition
and analysis of data, including MATLAB coding. Dr. Trevor Wade, Dr. Charles McKenzie and Dr. Rolf F. Schulte provided meaningful discussion. Additionally, Elaine Hegarty assisted with animal preparation, Andrew Wheatley and Krzysztof Wawrzyn helped with Hp $^{129}$Xe gas production. Dr. Giles Santyr provided guidance throughout the project and assistance with the manuscript preparation.

(3) Ozkan Doganay, Elaine Stirrat, Charles McKenzie, Rolf F. Schulte, Giles E. Santyr, Quantification of Regional Gas Exchange Changes using Hyperpolarized $^{129}$Xe MRI in a Rat Model of Early Stage Radiation-induced Lung Injury, Medical Physics 2015 (in Revision).

For this manuscript, Ozkan Doganay was responsible for experiment design including installation and optimization of the insert gradient and RF coils, experimental data acquisition, and analysis of data including post-processing with MATLAB and segmentation. Elaine Stirrat assisted with animal preparation and performing histological preparation and interpretation. Dr. Charles McKenzie and Dr. Rolf F. Schulte provided meaningful discussion. Dr. Giles Santyr provided guidance throughout the project and assistance with the manuscript preparation.
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<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>$^{129}$Xe</td>
<td>Xenon-129</td>
</tr>
<tr>
<td>$^{13}$C</td>
<td>Carbon-13</td>
</tr>
<tr>
<td>ADC</td>
<td>Apparent Diffusion Coefficient</td>
</tr>
<tr>
<td>$B_0$</td>
<td>Main static magnetic field</td>
</tr>
<tr>
<td>$B_1$</td>
<td>Intensity of magnetic field of applied RF pulse</td>
</tr>
<tr>
<td>$B_{1r}$</td>
<td>Receive RF field obtained by the receive coil</td>
</tr>
<tr>
<td>$B_{1+}$</td>
<td>Transmit RF field created by the transmit coil</td>
</tr>
<tr>
<td>CSSR</td>
<td>Chemical Shift Saturation Recovery</td>
</tr>
<tr>
<td>CT</td>
<td>Computed Tomography</td>
</tr>
<tr>
<td>FGRE</td>
<td>Fast Gradient Recalled Echo</td>
</tr>
<tr>
<td>FID</td>
<td>Free Induction Decay</td>
</tr>
<tr>
<td>FOV</td>
<td>Field Of View</td>
</tr>
<tr>
<td>Gy</td>
<td>Gray</td>
</tr>
<tr>
<td>Hp</td>
<td>Hyperpolarized</td>
</tr>
<tr>
<td>IDEAL</td>
<td>Iterative Decomposition of water and fat with Echo Asymmetry and Least squares estimation</td>
</tr>
<tr>
<td>LC</td>
<td>Capillary average thickness</td>
</tr>
<tr>
<td>$L_{PT}$</td>
<td>Pulmonary tissue average thickness</td>
</tr>
<tr>
<td>$M_0$</td>
<td>Equilibrium magnetization</td>
</tr>
<tr>
<td>MR</td>
<td>Magnetic Resonance</td>
</tr>
<tr>
<td>MRI</td>
<td>Magnetic Resonance Imaging</td>
</tr>
<tr>
<td>$M_{xy}$</td>
<td>Transverse magnetization</td>
</tr>
<tr>
<td>$M_z$</td>
<td>Longitudinal magnetization</td>
</tr>
<tr>
<td>NMR</td>
<td>Nuclear Magnetic Resonance</td>
</tr>
<tr>
<td>NSCLC</td>
<td>Non-Small Cell Lung Carcinoma</td>
</tr>
<tr>
<td>PET</td>
<td>Positron Emission Tomography</td>
</tr>
<tr>
<td>PFTs</td>
<td>Pulmonary Function Tests</td>
</tr>
<tr>
<td>PSF</td>
<td>Point Spread Function</td>
</tr>
<tr>
<td>PT</td>
<td>Pulmonary Tissue</td>
</tr>
<tr>
<td>PTA</td>
<td>Percent Tissue Area</td>
</tr>
<tr>
<td>RBC</td>
<td>Red Blood Cell</td>
</tr>
<tr>
<td>RF</td>
<td>Radiofrequency</td>
</tr>
<tr>
<td>RILI</td>
<td>Radiation-Induced Lung Injury</td>
</tr>
<tr>
<td>ROI</td>
<td>Region of interest</td>
</tr>
<tr>
<td>RP</td>
<td>Radiation Pneumonitis</td>
</tr>
<tr>
<td>RT</td>
<td>Radiation Therapy</td>
</tr>
<tr>
<td>SBRT</td>
<td>Stereotactic Body Radiotherapy</td>
</tr>
<tr>
<td>SCLC</td>
<td>Small Cell Lung Carcinoma</td>
</tr>
<tr>
<td>SPECT</td>
<td>Single-Photon Emission Computed Tomography</td>
</tr>
<tr>
<td>SRN</td>
<td>Signal-to-Noise Ratio</td>
</tr>
<tr>
<td>T</td>
<td>Tesla</td>
</tr>
<tr>
<td>$T_1$</td>
<td>Longitudinal relaxation time</td>
</tr>
<tr>
<td>$T_2$</td>
<td>Transverse relaxation time</td>
</tr>
<tr>
<td>$T_2^*$</td>
<td>Apparent transverse relaxation time</td>
</tr>
<tr>
<td>Symbol</td>
<td>Definition</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
</tr>
<tr>
<td>TE</td>
<td>Echo Time</td>
</tr>
<tr>
<td>TO/RO</td>
<td>Transmit-Only / Receive-Only</td>
</tr>
<tr>
<td>TR</td>
<td>Repetition Time</td>
</tr>
<tr>
<td>T\text{read}</td>
<td>Read-out time</td>
</tr>
<tr>
<td>\alpha</td>
<td>Flip angle</td>
</tr>
<tr>
<td>\alpha_G</td>
<td>Flip angle (delivered to gas phase of $^{129}$Xe)</td>
</tr>
<tr>
<td>\alpha_{PT}</td>
<td>Flip angle (delivered to dissolved phase of $^{129}$Xe in pulmonary tissue)</td>
</tr>
<tr>
<td>\gamma</td>
<td>Gyromagnetic ratio</td>
</tr>
<tr>
<td>V_{RBC}</td>
<td>Relative RBC Volume</td>
</tr>
<tr>
<td>\omega</td>
<td>Carrier frequency of the RF pulse</td>
</tr>
<tr>
<td>\omega_0</td>
<td>Larmor frequency</td>
</tr>
</tbody>
</table>
Chapter 1: Introduction

This chapter is intended to provide the introduction and motivation for the work presented in chapters 2, 3 and 4. First, an overview of lung physiology, lung cancer, and radiation induced lung injury (RILI), including the importance of early detection, will be explained. Furthermore, the advantages and limitations of clinical imaging modalities, are discussed with respect to lung imaging and RILI. A brief description of the principles of hyperpolarized (Hp) Magnetic Resonance Imaging (MRI), radiofrequency (RF) coils, and pulse sequence developments will be given. Finally, the hypothesis and specific objectives of the thesis project will be presented.

1.1 Overview and Motivation

Lung cancer is the largest contributor to cancer related deaths worldwide due to its low survival rate. Over 26,000 new lung cancer patients were reported in 2014 within Canada alone (1). Most cases are diagnosed as non-small cell lung cancer (~85%), for which radiotherapy (RT) is the most prominent treatment method (2). Although RT is successful in treating the cancer, there are undesirable consequences to the quality of life of the patient due to inadvertent injury of the normal lung tissue, known as Radiation-Induced Lung Injury (RILI). Current methods for assessing the severity of RILI include pulmonary function tests (PFTs), computed tomography (CT), single-photon emission computed tomography (SPECT), and positron emission tomography (PET). Although great advances
in these fields have been made, there is still a need for a non-ionizing imaging technique to assess regional functional (e.g. gas exchange) changes associated with RILI.

Novel Hp MRI techniques have been shown to be more sensitive to the early phase of RILI than other clinical imaging modalities and PFTs (3). Particularly, there is a strong interest in using Hp $^{129}$Xe gas as a contrast agent for MRI of lung due to the large increase in signal possible by spin exchange optical pumping (factor up to 100,000), known as hyperpolarization. Hp $^{129}$Xe has many applications in imaging of the lung airspace and in diagnosis of lung diseases, including chronic obstructive pulmonary disease, and asthma. In addition to imaging the lung airspace, $^{129}$Xe promises to provide unique functional information due to its relatively high solubility and significant chemical shift in pulmonary tissue (PT) and red blood cell (RBC) compartments. Additionally, dynamic Hp $^{129}$Xe MR spectroscopy has been shown to be sensitive in detection of RILI in a rat model as early as two weeks post-irradiation (4). The research described in this thesis is directed mainly to the development of hyperpolarized MR diagnostic tools and investigation of a pre-clinical models of RILI.

In this thesis, a novel Hp $^{129}$Xe MRI technique of the dissolved phases (i.e. PT and RBC) is described and the feasibility of quantifying the early phase of RILI in a rat model two weeks post-irradiation is shown. Low Signal-to-Noise Ratio (SNR) was the largest limitation for imaging dissolved phases of Hp $^{129}$Xe in the PT and RBC, therefore, a Transmit-Only/Receive-Only (TO/RO) RF coil is developed and constructed to increase the sensitivity. Another challenge of imaging the dissolved phases of $^{129}$Xe, was the lack of fast imaging techniques that provide simultaneous spatial and spectral information to follow the gas exchange between alveolar air space and PT and RBC. Therefore, a novel
The lungs are the respiratory organ responsible for gas exchange, consisting primarily of (i) airways (trachea, bronchi and terminal airways) and (ii) gas-exchange units known as alveoli (5) as shown in Figure 1.1. The lung facilitates the gas exchange of oxygen and carbon dioxide between the bloodstream and air. The lung start with the trachea, which is composed of cartilage rings and divides into two main bronchi. Each bronchus sequentially branches up to 23 bronchioles until the alveoli are reached. Inhaled air travels through the trachea into the bronchi and finally into the alveoli, respectively. Human lungs have three right lobes and two left lobes, while rat lungs consist of four right lobes and one left lobe (6). Human lungs contain a collection of alveoli in the range of 400 to 500 million (7,8). The alveolus is the smallest subunit of the lungs, which facilitates the site of the gas exchange, encompassing a total gas exchange surface are of approximately 130 m² (9). Oxygen in the alveolus air sac diffuses into PT and finally into RBCs in the surrounding capillaries. Then, oxygenated blood is carried to the heart via the pulmonary veins. The total volume of the trachea, bronchi and alveolar air sacs encompasses approximately 80% of the lungs (i.e. gas space). The remaining 20% (i.e. non-gas space) includes PT and surrounding capillaries.
Figure 1.1: (a) Sketch of human lungs showing the trachea and major bronchi of the lungs (Lung rendering adapted from: http://www.londonlungcancer.co.uk/lungcancer/). (b) Sketch of alveoli showing the alveolus and capillaries (Alveoli rendering adapted from: Figure 44-19, Biological Science 2/e, Pulmonary system ©2005 Pearson Prentice Hall, Inc http://35.9.122.184/lecture-slides/LB145-lecture21-S2013.pdf).

1.3 Lung Cancer

Lung cancer remains the major leading cause of cancer related death in Canada and worldwide (1,10-12). The Canadian Cancer Society estimates 26100 new lung cancer cases in 2015, and reports by sex, one in 12 males and one in 14 females are expected to develop lung cancer in their life-times (13). According to the Ontario Lung Association, approximately 310,000 people in Canada will develop lung cancer over the next 30 years, resulting in a total health care cost of approximately $33.6 billion (14).

Based on the microscopic appearance of the tumor cell, lung cancer is classified into three major categories. Non-small cell lung carcinoma, NSCLC, is the most common type compromising 85-90% of incident rates (15). Small cell lung carcinoma, SCLC, and lung carcinoid tumor type account for the remaining 10-15% of all lung cancer (16). NSCLC is
graded based on three criteria, known as TNM: (i) the size and location of tumor (T primary tumor), (ii) whether the tumor has spread to lymph nodes (N-regional lymph nodes) and (iii) tissues inside or outside the chest (M-distant metastasis) (17,18). After the TNM classification stage, a grouping is assigned using the numerical subset ranging from 0 to 4 to indicate the progressive extent of the malignant cancer (18). Together, the TNM classification and staging is used to select the most effective course of therapy and estimate the prognosis (19).

1.4 Advances in Treatment of Lung Cancer

The treatment options for lung cancer include chemotherapy, surgical resection, and RT (19). Chemotherapy has been the standard care of treatment in the clinic for patients with NSCLC and stage I-III, yielding a median overall survival of nine to twelve months (20-23). Platinum-based chemotherapy drugs (Cisplatin, Carboplatin, Oxaliplatin) are generally used for treatment of NSCLC patients. While the platinum-based chemotherapy drugs demonstrate substantial benefits for advanced NSCLC, the ideal number of therapy cycles still remains unclear, resulting in a dramatic change in the quality of life due to side effects (24).

Recent human trials using epidermal growth factor receptor tyrosine kinase (EGFR-TK) inhibitors have shown that the median overall survival for lung cancer has increased significantly, extending up to 30 months (25,26). The EGFR-TK molecular-based targeting agents (i.e., gefitinib, erlotinib) have been investigated as alternatives to chemotherapy owing to improved health outcomes while reducing previously observed toxic effects (27,28). The clinical use of molecular-based therapeutic strategies is limited by the increase
in the cost of gene mutation screen tests (Bradbury et al., 2010). EGFR-TK inhibitors have been primarily restricted to specific patient groups who are female and/or who have no smoking history (26,28).

Surgical resection is also regarded as a treatment option for 20-30% of NSCLC and SCLC, particularly for stages I and II (29-31). However, surgical resection often results in pain, discomfort and decreased quality of life, including significant decline in physical, social and mental states (30). Surgery with adjuvant chemotherapy approaches have also been offered to patients with stage I NSCLC to provide additional therapeutic benefits, with a reported 4% improvement in 5-year survival to 29% (32). Recently, post-operative radiotherapy for patients with stage I NSCLC, after complete surgical resection with adjuvant chemotherapy, seems to confer an additional improvement in overall survival compared to adjuvant chemotherapy alone (33).

1.5 Radiation Therapy

Radiation therapy (RT) plays an important role for lung cancer treatment (19). RT has been recommended in approximately 75% of patients with NSCLC and 79% of patients with SCLC by Dalaney et al. in Australia (34) and 61% of all patients with lung cancer by Tyldesley et al. in Canada (35). However, only 16-43% of patients with SCLC and 40-51% of patients with NSCLC receive RT in the world (34). This discrepancy may be because the clinical use of radiation therapy is limited by many confounding factors, such as different health care policies, availability of RT units, low referral rates and limited information about toxic effects. However, the growing clinical interest in RT has led to more prominent guideline recommendations for RT as part of the treatment plan (36).
The primary goal of RT is to deliver a high radiation dose to the tumor while minimizing the damage to surrounding normal tissues. There are two types of radiation dosing schedules for lung cancer; (i) conventional fractionated radiation and (ii) stereotactic body radiotherapy (SBRT). Conventional radiation therapy involves the delivery of total dose of radiation (40-50 Gy) over many (25-30) small fractions within three to five weeks (37). Usually, these small fractions are administered every day with radiation doses of less than two Gy per fraction. Neoadjuvant RT followed by surgery has also improved the three-year survival rate to 45% for Stage III NSCLC (38). SBRT uses a high dose of radiotherapy to treat small gross tumor volumes (~60 Gy), based on using several radiation beams that intersect at the site of the tumor. Particularly, SRBT with 3D image guidance allows high precision with a relatively high fraction (20 Gy per fraction). A growing number of SBRT studies have demonstrated an improved 2-year survival for medically inoperable NSCLC patients (39,40). Although RT is mostly used for stage III NSCLC patients (40-66%), recent studies have also reported improved survival rates of stage IV NSCLC with SBRT and adjuvant chemotherapy treatment approaches (41). Early-phase toxic effects (i.e. RILI) are significantly worse for SBRT compared to conventional radiation therapy (19,42), though overall outcome is improved, for reasons that are not clear.

1.6 Radiation-Induced Lung Injury (RILI)

Irradiation of normal lung tissues during the course of RT results in RILI which includes a range of complications such as acute toxicities, chronic symptoms, and fibrosis that can all lead to organ failure. RILI is categorized into two phases: (i) early-phase and (ii) late-phase depending on the development of clinical symptoms, though one can occur without the
other. The early-phase effect, also known as radiation pneumonitis (RP), is an acute inflammatory response that typically occurs within weeks following RT and is characterized with symptoms of dry cough and dyspnea (43). According to histopathology, RP includes extensive vasculature damage as well as tissue thickening (i.e. inflammation), alveolar edema and macrophage accumulation (44). After conventional RT, the overall incidence rates of moderate RP ranges from 15-50%, depending on the dose of the irradiated volume (45,46). While the incidence of severe RP is low, 5-35%, the mortality rates are high after SBRT (47,48). Vascular injury in the early-phase progresses with an extensive increase in collagen, which then transforms to fibrosis if the RP is not treated (49). Late-phase RILI represents collagen deposition and fibrosis after three months irradiation (43). Although radiation dosing strategies and adjuvant therapies have been developed to reduce short and long-term complications, RILI remains a significant problem for RT (50).

1.7 Diagnosis of Radiation-Induced Lung Injury

Monitoring of RP, during the course of RT, would allow for patient specific alterations to the therapy plan based on the severity of RILI. These alterations may include modification of radiation dose, or the use of adjuvant therapy, including chemotherapy and radio-protective and anti-inflammatory drugs (50,51). Detection of RP in its early phase would also help monitor the effectiveness of radio-protective changes. Minimizing normal tissue toxic effects after RT can increase the overall patient survival rate and improve the quality of life (50).
1.7.1 Pulmonary Function Tests

Pulmonary function tests (PFTs) are used to assess global lung function using measurements of forced expiratory volume (FEV), vital capacity, and carbon monoxide diffusing capacity (DLCO) (52). Changes induced in the lungs by thoracic irradiation are commonly evaluated by spirometry (53). Thoracic irradiation typically results in a reduction in the DLCO. However, FEV₁ (FEV measured at 1 second) may show inconsistently an increase or no changes in patients with centrally located lung lesions (54-56). Additionally, PFT measurements depends on the patients’ breathing efforts and need to be corrected for anemia (49). Furthermore, changes in whole lung function do not reveal regional information, limiting the ability of PFTs to resolve regional disease heterogeneities (57). Consequently, PFT can only detect RILI when a relatively large dose of 25 Gy is given and only three to six months post irradiation (54).

1.7.2 Chest Radiography / Computed Tomography

Computed tomography (CT) is an imaging modality that detects X-rays taken at different angles around the body to form cross-sectional images. Clinically, plane film chest X-ray and CT imaging are commonly utilized to assess RILI based on corresponding changes in lung density. Irreversible density gain and structural changes detected by X-rays in the lungs due to RILI occurs mainly in late phase due to the tissue remodeling. CT is capable of detecting late-phase fibrosis, two-three months following radiation doses of 40 Gy and above (53,58,59). CT perfusion is a relatively new technique which measure the perfusion by comparing the density changes in the region of interest during the intravenous injection of iodinated contrast material (60,61). Recent imaging developments in CT perfusion have
shown that perfusion-based imaging techniques are more sensitive to early-phase RILI than conventional CT (62). Hu et al. reported that CT perfusion was a promising technique for detection of early-stage RILI after four weeks following a radiation dose of 60-62 Gy. However, both conventional RT and SBRT use treatment plans that are administered in two to three weeks, therefore detection of RILI four weeks post-radiation could be too late to adjust the therapy plan (e.g. radiation fractionation) or use pharmaceuticals (radio-protectors and anti-inflammatory drugs) (50,63,64).

1.7.3 SPECT

Single photon emission computed tomography (SPECT) is a nuclear imaging modality that images the distribution of gamma-emitting radionuclides in the body. SPECT is widely used to map both perfusion and ventilation changes due to RILI (53,65-67). SPECT ventilation is performed by inhaling radioactive tracers such as radioactive gas (e.g. $^{81m}$Kr and $^{133}$Xe), particulate aerosols labeled with radioactive compounds (i.e. $^{99m}$Tc-DTPA) and ultrafine graphite particles labeled with $^{99m}$Tc (i.e. Technegas) (68). One of the main concerns with detecting ventilation defects by means of ventilation SPECT is the inconsistency in intensity distribution of the radioactive gas when different radioactive gases are used within the same isogravitational planes (69). Particulate aerosols can diffuse to the alveolar capillary network and lead to ventilation image distortions. Furthermore, the clinical use of Technegas has not yet been approved by the US Food and Drug and Administration (69).

SPECT perfusion is a highly specialized lung perfusion imaging approach largely used in research settings and capable of assessing the reduction in lung perfusion associated with
RILI (70). SPECT perfusion lung imaging is accomplished by intravenous administration of albumin macroaggregates (MMA) labeled with $^{99m}$Tc. Using SPECT perfusion imaging, Marks L.B. et al. demonstrated a 20-30% drop in blood perfusion in RT patients who received a radiation dose of 15-30 Gy (49). Despite SPECT perfusion imaging being more sensitive to RILI than CT, correction for attenuation, scatter, is required for quantitative imaging (71). Downscatter effects, which are caused from simultaneous ventilation and perfusion imaging using different radiotracers is also required to correct for interactions between ventilation and perfusion. Quantitative assessment of total blood volume is difficult since the sensitivity of SPECT scans significantly depends on patient positioning (69). SPECT is hampered by the poor spatial resolution (10-20 mm for 140 keV gamma rays from $^{99m}$Tc) and long scan times (10-30 minutes).

1.7.4 PET

Positron emission tomography (PET) is a radionuclide imaging modality that images the distribution of intravenously injected positron emitting tracer in the region of interest. PET utilizes collinear 511 keV gamma rays offering a better spatial resolution (5-10 mm) than SPECT. PET with [$^{18}$F] fluorodeoxyglucose (FDG) provides functional information about metabolic activity due to the accumulation of FDG in inflamed tissue (72). Increased $^{18}$F-FDG activity has been reported and correlated with clinical symptoms of RP in patients undergoing thoracic radiotherapy (3-12 weeks) receiving radiation doses of as low as 20 Gy (73). Hart et al. showed corelation between RP clinical symtoms and measured $^{18}$F-FDG activity with an accuracy of approximately 50% (73). McCurdy et al. noted a linear relationship between post-RT $^{18}$F-FDG uptake and radiation dose response 1-3 months
after RT (74). Although high $^{18}$F–FDG uptake corresponded to RP in these studies, the tumor growth was also associated with enhanced $^{18}$F-FDG uptake restricting the sensitivity to detection of early-phase normal tissue changes (75,76). PET scans are limited by the lack anatomical references, therefore PET imaging frequently requires another imaging modality to provide anatomical images for the image registration such as CT (77). Another limitation with chest X-ray, CT, SPECT and PET are the radiation exposure of 2-3mSv for a typical ventilation and perfusion study.

1.7.5 Magnetic Resonance Imaging Methods

Conventional MRI provides images depicting the spatial distribution of hydrogen nuclei (i.e. proton) within water molecules in the body. MRI achieves high sensitivity due to the high molar concentration of water, which provides excellent soft tissue contrast with high temporal and spatial resolution without the need for ionizing radiation. However, lung imaging with MRI presents a challenge. The lungs have a low proton density and therefore low SNR since the pulmonary tissue (PT) volume comprises only ~ 20% of the total lung volume. Contrast-enhanced Gd-DTPA MRI techniques can provide perfusion characteristics of RILI (78). Ogasawara et al. investigated the time course of the contrast enhancement in the lungs using Gd-DTPA kinetics between irradiated- and non-irradiated lungs. This study showed altered Gd-DTPA kinetics one-month post irradiation with a single dose of 40 Gy in eight dogs compared to normal lungs due to a perfusion reduction. In this study, dynamic Gd-DTPA MRI was significantly affected by long scan times (~3 minutes) that affect the measurement due to blurring from respiratory motion. Gd-DTPA MRI lung perfusion and volume measurements are difficult to correct for quantitative
measurement of perfusion (79). Conventional $T_2^*$ MRI techniques are sensitive to the mobility of water molecules and therefore can assess RILI by monitoring changes in $T_2^*$ decay (80). A prolongation of the $T_2^*$ two-weeks post-irradiation (20 Gy) was observed in *ex vivo* rat lungs (80). Zhang *et al.* have demonstrated in a rabbit model that the apparent diffusion coefficient (ADC) in tumor lesions decreases 1-5 weeks post-irradiation of the lungs (81). However, in that study it was difficult to obtain ADC values from the normal PT due to low SNR and short $T_2^*$ because of air/tissue magnetic susceptibility differences.

1.7.6 Hyperpolarized MRI

Hyperpolarized (Hp) MRI has the ability to assess the anatomical, functional and metabolic changes associated with RILI using $^3$He, $^{129}$Xe, and $^{13}$C respectively (3). MR spectroscopic imaging can detect changes due to RILI with intravenous injection of Hp contrast agent labeled with [1-$^{13}$C] pyruvate (82). Thind *et al.* reported the lactate-to-pyruvate signal ratio (Lac/Pyr) in the irradiated rat lung with a radiation dose of 18.5 Gy was significantly different from the non-irradiated rat lung two-weeks post irradiation (83). The increase in the Lac/Pyr ratio was correlated with changes in macrophages density due to RP measured histologically. The Lac/Pyr measurements were not able to directly measure the anatomical and functional changes due to RP. Additionally, MR spectroscopic imaging using Hp [1-$^{13}$C] pyruvate was challenging due to low tissue concentration, requiring relatively large voxels.

Hp $^3$He gas MRI has also been explored to quantify RILI. Hp $^3$He gas MRI can provide Apparent Diffusion Coefficient (ADC) maps of lungs with high SNR, temporal and spatial resolution. The first study of RILI using a rat model has shown a significant increase in
ADC values of Hp $^3$He associated with three months-post irradiation dose of 40 Gy (84). Increases in the regional lung $^3$He ADC values have been associated with fibrosis in patient groups 8-months post-RT (85). Ireland et al. investigated the feasibility of detecting the ventilation changes in the lungs between pre- and post-RT using Hp $^3$He gas (86). In this work, Ireland et al. correlated the degree of emphysema in the lungs using CT and Hp $^3$He MRI in regions of RP three-months post-RT. Although Hp $^3$He gas, particularly ADC, shows feasibility for detection of fibrosis. Hp $^3$He gas MRI is restricted to ventilated regions of the lung since $^3$He has negligible solubility. Furthermore, $^3$He is an exceedingly rare gas with limited abundance and unlikely to develop into a clinical tool. $^{129}$Xe on the other hand is plentiful and has relatively high tissue solubility and chemical shift. Hp $^{129}$Xe diffusion-weighted MR imaging has been used to measure changes in lung morphometric parameters associated with RILI, specifically the mean airspace chord length ($L_m$) in a RILI rat model (87). In this work, Ouriadov et al., measured a decrease in $L_m$ two weeks post-irradiation with a radiation dose of 18 Gy and successfully correlated the changes to histology measurement of mean linear intercept.

### 1.7.7 Hp $^{129}$Xe MR Spectroscopy

In addition to lung ventilation imaging, Hp $^{129}$Xe solubility properties allow spectroscopy techniques to be used for the measurement of exchange from the alveolar air space to the PT and RBC compartments (88,89). There are two dissolved phase peaks relative to gas signal in MR spectra as shown in Figure 1.2. The largest peak occurs at -6950 Hz corresponds to Hp $^{129}$Xe in the gas phase. The peak at the center of the MR spectrum (0 Hz) (in Figure 1.2) is due to PT which also includes a small and negligible fraction of blood
plasma at approximately the same frequency (90). The peak at -520 Hz (19 ppm) is the xenon bound to hemoglobin inside the RBC (91,92). Dissolved phase signal is approximately 2% of the gas signal and saturates in the lungs very rapidly within a gas transfer time range of 50-150 ms. Therefore, the use of small RF excitation (i.e. flip angles) of the gas phase magnetization in the alveolar air pool, allows the dynamics of the dissolved phase signal (i.e. gas exchange curves) to be assessed using subsequent MR spectroscopy acquisitions. This can be done by changing the time delay between data acquisitions. This MR spectroscopy technique is called chemical shift saturation recovery (CSSR) (93,94). Gas transfer dynamics measured from the whole lung using CSSR technique can be translated into assessment of structural and functional information of lung by fitting with numerical gas exchange models (90,95,96).

Figure 1.2: MR spectrum of rat lungs after inhaling Hp$^{129}$Xe using a spectrally selective RF pulse that excites the gas phase with a low flip angle and the dissolved phases with a high flip angle.

Fox et al. have investigated early-phase RILI using the CSSR technique (4). In this study, average gas transfer time for PT was observed to be elevated significantly with a whole
lung irradiated rat model two weeks following 16 Gy irradiation compared to a control group. In this study, correlation between early-phase RILI changes in whole-lung and RBC gas transfer curve was not significant. This was an unexpected finding as the vascular network is expected to be affected by RILI, and may have been due to the fact whole lung CSSR measurement cannot identify regional changes. Saturation of $^{129}$Xe in the RBC compartment is expected to vary regionally and temporally due to perfusion and gas exchange of $^{29}$Xe in RBC within the capillary network. Although the Hp $^{129}$Xe gas MR spectroscopy CSSR technique provides functional information about the global lung physiology, regional information of the Hp $^{129}$Xe gas exchange process is expected to increase the sensitivity for detection of RILI effects when using partial (or conformal) radiotherapy.

1.7.8 Hp $^{129}$Xe Dissolved Phase imaging

Imaging the dissolved phases of $^{129}$Xe (i.e. PT, RBC) has been explored using three-point Dixon approaches based on Cartesian k-space sampling (97), following a one-point Dixon approach based on radial k-space sampling fashion to address short $T_2^*$ (98). Cleveland et al. extended the one-point Dixon approach to include 3D imaging (99). On the other hand, Qing et al. also developed a 3D imaging technique using hierarchical IDEAL (Iterative Decomposition of water and fat with Echo Asymmetry and Least-square estimation) reducing the breath-hold time down to approximately 11 s (99,100). However, the dissolved phase imaging techniques have yet to be explored for RILI.
1.8 Principles of MRI

The physical phenomenon called nuclear magnetic resonance (NMR) is concerned with the intrinsic angular momentum (spin) of nuclei that contain unpaired protons or neutron. The nuclei of $^1$H in water (and fats) serve as the primary source at the signal in MRI since water is highly abundant in the body. The source of the NMR signal originates from the precession of the bulk spin system in the presence of a strong static magnetic field and subsequently excitation of spins by application of an externally applied radiofrequency (RF) field. The NMR signal is spatially localized by MR imaging using spatially-varying magnetic field gradients (i.e. frequency and phase encoding gradients). The type of RF excitation (i.e. the selection of flip angle); timing in regards to acquisition of data, and data acquisition (i.e. k-space) trajectory describe the MR pulse sequence strategy. In this section, basic concepts of MRI are introduced including conventional and Hp MRI relevant to the thesis. Interested readers are referred to Haacke et al. and Bernstein et al. for more details (101,102).

1.8.1 Magnetic Moment in the Presence of a Magnetic Field

On closer inspection, a nuclear spin, which has a magnetic moment, $\mu$, experiences a torque when an external magnetic field, $B_o$, is applied, resulting in the precession of spins about the direction of the magnetic field. The precession angular frequency, $\omega_0$, is called the Larmor frequency and given by the following equation:

$$\omega = \gamma B_o$$

[1.1]
where $\gamma$ is the gyromagnetic ratio which is usually expressed as $\gamma/2\pi$ in SI units of MHz/T. The gyromagnetic ratio for $^{129}$Xe and $^1$H are -11.78 MHz/T and 42.58 MHz/T respectively. Thus, the Larmor frequencies for $^{129}$Xe and $^1$H are -35.34 MHz and 127.7 MHz respectively at a field strength of 3T.

1.8.2 Conventional MR Magnetization

Application of an external magnetic field, $B_0$, aligns the spins either parallel or anti-parallel to give a net magnetization of $M$ as shown in Figure 1.3 (a). Conceptually, the conventional MR signal arises from $M$, which aligns along the external magnetic fields as shown in Figure 1.3 (b). Application of an RF pulse, $B_1$, at the Larmor frequency along the $x$-axis will provide a torque that tips $M$ from the $z$-axis to the transverse plane ($xy$-plane) (Figure 1.3 (c)). Therefore, following RF pulse application, $M$ can be decomposed into a longitudinal component $M_z$ and a transverse component $M_{xy}$. $M_{xy}$ precesses at the Larmor frequency while inducing an exponentially decreasing time-varying voltage signal in the receiver RF coil (Figure 1.3 (d)), or Free Induction Decay (FID). The signal received by the RF coil resembles a damped oscillation, governed by an apparent transverse relaxation time ($T_2^*$). On the other hand, while the transverse magnetization decreases via transverse relaxation, the $M_z$ re-grows via longitudinal relaxation, re-establishing the net magnetization to thermal equilibrium along the $z$-axis.
Figure 1.3: (a) Bulk magnetization vector, $\mathbf{M}$, of a spin population in the external magnetic field, $\mathbf{B}_0$; (b) Tipping the magnetization from $z'$ direction to $xy$ plane using an RF pulse, $\mathbf{B}_1$, applied in the $x'$ direction; (c) Precession of transverse magnetization, $\mathbf{M}_{xy}$, in the laboratory frame (d) Time-varying voltage signal induced in the receive coil.

For a spin 1/2 system, magnetization in a spin population arises from two quantum states. In the presence of $\mathbf{B}_0$, the nuclear spins split into low energy, $S_{-1/2}$, and high energy, $S_{+1/2}$, states called Zeeman levels. The energy difference between these state is given by:

$$E = -\mu \cdot \mathbf{B}_0$$ \hspace{1cm} [1.2]

where, $\mu$ is the magnetic moment of the spin, $1/2\gamma \hbar$ for the $S_{+1/2}$ and $-1/2\gamma \hbar$ for the $S_{-1/2}$.

The number of spins in the $S_{+1/2}$ state which are aligned with $\mathbf{B}_0$, contributes to the net magnetization $\mathbf{M}$, and thereby the thermally-polarized signal for conventional MR. Therefore, the longitudinal magnetization $M_z$ is given by:

$$M_z = P \cdot N \cdot \frac{\gamma \hbar}{2}$$ \hspace{1cm} [1.3]

where $\hbar$ is the Planck’s constant ($\hbar = 1.054571800(13) \times 10^{-34} \text{ (J s)}$), $N$ is the total number of atoms, and $P$ is the polarization. The magnetization, and thus the NMR signal depends on the polarization levels. The polarization, $P$, is given by:
\[ P = \frac{N^+ - N^-}{N} \]  \[ 1.4 \]

where, \( N^+ \) and \( N^- \) are the number of spins in the high and low Zeeman levels respectively.

Ideally, the condition of \( N^+ \gg N^- \) corresponds to the highest possible polarization, \( P \approx 1 \), and therefore provides the highest signal. However, for thermal equilibrium, polarization is very low due to the small difference between the \( N^+ \) and \( N^- \) states (Figure 1.4 (a)).

Therefore, \( P \) can be re-written using the Boltzmann distribution as follows:

\[ P_{\text{thermal}} \approx \frac{1}{2} \left[ \frac{N^+}{N^-} - 1 \right] \approx \frac{\gamma \hbar B_0}{2kT} \]  \[ 1.5 \]

where, \( k \) is the Boltzmann constant \( (k = 8.6173324 \times 10^{-5} \text{ (eV K}^{-1})) \), and \( T \) is temperature \( (T=300 \text{ K at laboratory temperature}) \). For a magnetic field strength of \( B_0 = 3 \text{T} \), the achievable thermal polarization fraction for \(^{129}\text{Xe} \) is approximately \( 2.8 \times 10^{-6} \). At room temperatures, the percentage polarization for \(^{129}\text{Xe} \) is therefore very small, however this can still detected if there are a large number of \(^{129}\text{Xe} \) nuclei within the volume. Therefore, thermally-polarized \(^{129}\text{Xe} \) phantoms containing pressurized \(^{129}\text{Xe} \) gas are generally used to achieve a measurable thermal MR signal.
Figure 1.4: Zeeman energy levels at thermal equilibrium (a) and in the hyperpolarized state (b). $S_{-1/2}$ and $S_{+1/2}$ are low and high energy levels.

Substituting Eqn. [1.5] in Eqn. [1.3], the longitudinal magnetization for conventional MR can be re-written as follows:

$$M_z = \frac{N\gamma^2\hbar^2 B_o}{4kT}.$$  \[1.6\]

Equation [1.6] provides the basis of the MR signal, once factors including relaxation time dependencies, and the RF coil sensitivity are considered. Conceptually, Eqn. [1.6] can also be related to the achievable conventional MR signal ($^1$H) from lung. Since $N$ is low in the lungs due to high fraction of air space, the thermally-polarized conventional MR signal is very low. This signal can be artificially improved using hyperpolarized gas techniques as explained in the next section.
1.8.3 Hyperpolarized MR Magnetization

Hyperpolarization is a pre-magnetized form of MR in which the percentage polarization achievable is in a range of 10-40%, typically using a spin-exchange optical pumping (SEOP) polarization approach (103). The population of $N^+$ nuclei is artificially increased up to a factor of $10^5$ compared to thermally polarized $^{129}$Xe gas as shown in Figure 1.4 (b). While the initial magnetization is significantly higher, the Hp $^{129}$Xe gas differs from conventional MR since the longitudinal magnetization is not renewable and the magnetization depends on the polarization from the SEOP approach. The following sections briefly explain the process of SEOP polarization.

1.8.4 Hyperpolarized $^{129}$Xe Gas Production

Hyperpolarization is achieved by the combination of two different stages: (i) transfer of the angular momentum from the circularly polarized laser photons to the valence electrons of Rubidium-87 atoms ($^{87}$Rb) via optical pumping, and (ii) transfer of the electron polarization of $^{87}$Rb atoms to $^{129}$Xe atoms via spin exchange (Fermi-contact hyperfine) interactions (104). This is usually achieved in a continuous flow of $^{87}$Rb vapour of a mixture of $^{129}$Xe (1-2%) and other buffer gases (usually 97% N$_2$ and 1-2% ³He) through a polarizer cell containing $^{87}$Rb vapor as shown in Figure 1.5 (a) (105,106). The cell is placed in a weak magnetic field (60-70 Gauss) in order to split the valence electrons of $^{87}$Rb into different Zeeman energy levels. A circularly polarized laser ($\sigma^+, \Delta m_j = +1$) with a wavelength of 795nm polarizes the $^{87}$Rb valence electrons. The electrons in the $5S_{1/2}$ ($m_j = -1/2$) ground level is excited to the higher energy level $5P_{1/2}$ ($m_j = +1/2$) upon absorption of photons as shown in Figure 1.5 (b). Since the laser light is circularly polarized, the transition only
allows jumps from $m_j = -1/2$ to $m_j = +1/2$ levels, which represents azimuthal spin quantum number. The $^{87}\text{Rb}$ electrons de-excite back to the ground state due to either the quenching by $\text{N}_2$ gas or radiatively (life time less than 30 ns). Finally, spin exchange takes place causing a transfer of the spin angular momentum of the $^{87}\text{Rb}$ valence electrons to the $^{129}\text{Xe}$ nucleus as shown Figure 1.5 (c). The achievable $^{129}\text{Xe}$ polarization is given by:

$$P_{\text{Xe}} = P_{\text{Rb}} \cdot \frac{\gamma_{SE}}{\gamma_{SE} + \Gamma} \left[1 - \exp(-\tau \cdot (\gamma_{SE} + \Gamma))\right]$$  \hspace{1cm} [1.7]$$

where $P_{\text{Rb}}$ is the polarization of $^{87}\text{Rb}$, $\gamma_{SE}$ is the spin exchange rate, $\Gamma$ is the depolarization rate of the $^{129}\text{Xe}$ gas, and $\tau$ represents the time required for $^{129}\text{Xe}$ to reach maximum polarization due to the spin exchange interaction between the optically pumped $^{87}\text{Rb}$ and $^{129}\text{Xe}$ within the SEOP cell. The use of high power narrowed laser sources (60-150 W) provides significantly high polarization of $^{87}\text{Rb}$ along the SEOP cell. Maintaining the appropriate cell temperatures in the SEOP cell is one of the main criteria to reduce $\Gamma$. However, after extended use, temperature variations and cold spots on the SEOP cell due to the inappropriate thermal management of polarizer cell causes the $^{87}\text{Rb}$ vapour to condense on the cell surface. The $^{87}\text{Rb}$ buildup layer due to the cold spots on the cell diminishes the polarization of the $^{129}\text{Xe}$ gas. Another important limiting factor is the separation of the Hp $^{129}\text{Xe}$ gas from other gas mixtures. After polarization in the SEOP cell, the gas mixture flows into a trap submerged in liquid nitrogen at temperature of ~77 K. Hp $^{129}\text{Xe}$ is frozen and accumulated in the trap for approximately one hour (107). The frozen mixture is sublimated to the gas state for use during experiments. This transition from the frozen state to gas state is the major source polarization loss since the decay of polarization is rapid during the phase transition (on the order of 30-50%). Consequently,
10-40% is the typical level of polarization achievable; however this depends on a number of factors, including laser power, the geometry of the SEOP cell, the thermal management of the SEOP cell, the gas flow and fraction of the gas mixture, and the design of the xenon freeze out trap (108).

Figure 1.5: (a) SEOP cell is shown including the Helmholtz coils, cell, and circularly polarized \((\sigma^+)\) laser beam. (b) Optical pumping for polarizing the valence electrons of \(^{87}\text{Rb}\). (c) The spin exchange interactions between the \(^{87}\text{Rb}\) and \(^{129}\text{Xe}\). \(\tau\) is the time required for \(^{129}\text{Xe}\) to reach saturation due to the spin exchange interaction.

Substituting Eqn [1.7] in Eqn [1.3], the longitudinal magnetization is given by:

\[
M_z = \frac{1}{2} N \cdot \gamma \cdot h \cdot P_Rb \cdot \frac{\gamma_{SE}}{\gamma_{SE} + \Gamma} [1 - \exp(-\tau \cdot (\gamma_{SE} + \Gamma))] \tag{1.8}
\]

Another distinguishing feature of Hp MR can be seen by comparing Eqn. [1.8] to Eqn. [1.6]. The hyperpolarized magnetization does not depend on the main magnetic field and is approximately linearly proportional to gyromagnetic ratio of the nuclei. This feature has implications for low field MRI that are not described in this thesis.
1.8.5 Chemical Shift

Although the initial Hp signal strength is independent of the main magnetic field \( B_o \), \( B_o \) still plays an important role in Hp \(^{129}\)Xe MRI. The precession frequency of spins depends on \( B_o \) as already shown in Eqn [1.1] assuming the \( B_o \) is the same for all spins. In reality, the spins in different chemical environments experience different amounts of de-shielding resulting in variations of local \( B_o \). The local magnetic field, \( B_{\text{local}} \), experienced by the nuclear spin is given by:

\[
B_{\text{local}} = (1 - \sigma) B_o
\]

[1.9]

where \( \sigma \) is the de-shielding constant that depend on the intrinsic chemical properties of the electron cloud on the nuclei and its environment. Therefore, Eqn. [1.1] can be rewritten as follows:

\[
\omega = \gamma B_{\text{local}}
\]

[1.10]

The variation in \( B_{\text{local}} \) is referred to as the chemical shift of the Larmor frequency. Sufficiently high magnetic field strength permits separation of distinct chemical shifts of \(^{129}\)Xe. The phenomenon of chemical shift is shown in Figure 1.2 from rat lungs at 3T. \(^{129}\)Xe in the PT and RBC compartments experiences a different chemical environment and \( B_{\text{local}} \). The frequency differences due to the chemical shift between the \(^{129}\)Xe gas and the dissolved \(^{129}\)Xe phases in the PT and RBC compartments are approximately 6950 Hz and 7475 Hz at 3 T respectively.
1.8.6 RF Pulse

The concept of the radiofrequency (RF) magnetic field ($B_1$) was mentioned in section 1.8.2 and Figure 1.3 (c). The application of $B_1$ tips the magnetization vector into the transverse plane by an angle called the flip angle. The flip angle, $\alpha$, is given by:

$$\alpha = \gamma \int_0^T B_1(t) \, dt$$

[1.11]

where $T$ is the pulse duration. A flip angle of 90° will produce the maximum MR signal by tipping all the available magnetization into the transverse plane. Since the Hp magnetization is non-renewable, a second RF pulse would result in no signal. Therefore, to optimize the available magnetization the selection of both the amplitude and duration of RF pulse are equally important. Another important concept is the spatial homogeneity of the applied $B_1$ field. A homogenous $B_1$ can be achieved by using a birdcage RF coil as described in Chapter 2.

RF pulses can be used to excite chemical shifts with different flip angles. The previously described CSSR technique (4) uses a frequency selective rectangular (or ‘hard’) RF pulse; with a low flip angle applied to the $^{129}$Xe gas phase, and a large flip angle applied to the dissolved phases of $^{129}$Xe. In RF pulse terminology, a hard pulse has the RF profile shape of a rect function in the time domain. The Fourier transform of a hard pulse leads to an RF profile with a sinc function in the frequency domain. Selective excitation of the gas and dissolved phases of $^{129}$Xe can be achieved in the frequency domain with a sinc function. The central lobe of the sinc function provides the necessary high flip angle to the dissolved
phase of $^{129}$Xe and the side lobes provide the low flip angle excitation to the gas phase of $^{129}$Xe.

The RF pulse and magnetic moment are analyzed in a rotating reference frame to simplify the mathematical analysis of the temporal progression of the magnetization. An effective magnetic field is derived to explain the interaction between the $B_1$ and spin system in the rotating frame as follows:

$$B_{\text{eff}} = (B_o - \frac{\omega}{\gamma})\hat{z} + B\hat{x}$$  \[1.12\]

where, $\omega$ is the carrier frequency of the RF pulse. Thus, the effective magnetic field has two components in the rotating frame. On-resonance ($\omega = \omega_o$), $B_{\text{eff}}$ is in the same direction as $B_1$.

1.8.7 MR Signal

The phenomenological Bloch equations describe the behavior of the magnetization after the application of an RF pulse, including the relaxation terms as follows:

$$\frac{dM}{dt} = \gamma M \times B_{\text{eff}} + \frac{1}{T_1} (M_o - M_z) - \frac{1}{T_2^*} M_{xy}$$  \[1.13\]

The first term in the Bloch equation represents the relationship between the application of the RF pulse and spin system including the off-resonance effects. The second term describes the relaxation of the longitudinal magnetization, characterized by the spin-lattice relaxation time ($T_1$). The third term describes the relaxation of the transverse magnetization, characterized by the apparent spin-spin relaxation time ($T_2^*$). The solution
to the Bloch equation for thermally polarized signal has been discussed previously (101,102). In particular, the solution to the Bloch equation for Hp $^{129}$Xe imaging after the application of an RF pulse of flip angle of $\alpha_1$ is given by(109):

\[ M_z = M_i \cos(\alpha_1) \left(1 - \exp(-t/T_1)\right) \]  \hspace{1cm} [1.14] 

\[ M_{xy} = M_i \sin(\alpha_1) \left(1 - \exp(-t/T_1)\right) \exp(-t/T_2^*) \]  \hspace{1cm} [1.15] 

where $M_i$ is the initial magnetization that depends on the polarization of the $^{129}$Xe gas. $T_1$ describes the exponential decay of the longitudinal magnetization toward thermal equilibrium. $T_1$ mechanisms include the interaction with paramagnetic molecules such as diatomic oxygen. However, the effects of $T_1$ are generally negligible since $T_1$ is on the order of ~30s for $^{129}$Xe gas in the lungs and the MR acquisition is on the orders of a few seconds.

The spin-spin relaxation time ($T_2^*$) describes the exponential decay of the transverse magnetization. Among other factors, $T_2^*$ depends on the uniformity of $B_0$, as such the $T_2^*$ for Hp $^{129}$Xe in the gas phase and dissolved phases are approximately 6 ms and 2 ms at 3T respectively. Apart from the non-renewable nature of Hp magnetization, $M_{xy}$ and thereby the SNR of dissolved phases of $^{129}$Xe strongly depends on $T_2^*$. To deal with the signal loss due to short $T_2^*$, the signal is optimally sampled in as short a read-out time as possible using a high flip angle. The selection of the flip angle and optimization of read-out time will be discussed in Chapter 3 in more detail.
1.8.8 RF Coils

As discussed in the previous section 1.8.6, $B_1$ tips the longitudinal magnetization into the transverse plane initiating the MR signal. While the transverse magnetization is precessing in the $xy$ plane, it induces an exponentially decaying voltage signal (i.e. FID) that is used to reconstruct the MR image. An RF coil is used to transmit the excitation RF field, $B_{1+}$, and receive the RF field, $B_{1-}$, radiated by the spin system. To achieve a homogenous flip angle distribution ($B_{1+}$) over the imaging region, RF coils are often designed in shape of cylinder (e.g. birdcage coils). The $B_{1+}$ homogeneity generated by the transmit coil can be derived using the Biot-Savart law for a unit current flowing in the coil elements. A receive coil that is placed close to the object is known as a surface coil which improves the sensitivity. The sensitivity of the receive coils can be determined using the principle of reciprocity, where $B_{1-}$ for a unit current flow, $I$, in the receive coil element is considered to arise from a sample associated with the density of the spin population. The MR signal for a given volume of sample is expressed using the following equation:

$$ S(t) = A \omega_0 \int \frac{(B_{1-} \cdot (\vec{r})/I)}{M_{xy}(\vec{r}, t)} dV $$

[1.16]

where $A$ is a constant that depends on coil filling factor, gain, and frequency deviations due to $B_0$ and $B_{1+}$ inhomogeneities. A detailed discussion about optimization of the $B_+$ and $B_-$ field and other technical details including the geometry and electrical components of transmit and receive RF coils can be found in the Chapter 2.
1.8.9 Spatial Localization of the MRI Signal

Imaging requires spatial localization of the precession frequencies as a function of position over the imaging region. A precise variation in precession frequencies is achieved by applying linearly changing magnetic field gradients using gradient coils. If the gradient coils are switched on, the net magnetic field is given by:

\[ B(x,y,z) = (B_o + G_x x + G_y y + G_z z) \hat{z} \]  \[1.17\]

where, \( G_x \), \( G_y \), and \( G_z \) are the gradient amplitudes produced by the \( x \), \( y \), and \( z \) gradient coils respectively. Typically, the maximum gradient amplitude for a clinical scanner is on the order of 50 mT/m. Additionally, the rate by which the achievable gradient amplitude can be obtained, is called the slew rate. Typical clinical values for slew rate are on the order of 200mT/m/ms for a clinical scanner. Both the slew rate and gradient amplitudes are significantly important for establishing the temporal and spatial resolution of MR imaging and will be discussed in detail in Chapter 3.

In this thesis, MR images were acquired in 2D projection fashion. 2D imaging is performed without slice-selection, using a phase encoding gradient applied in \( y \) direction and frequency encoding gradient applied in \( x \) direction. A hard pulse with a small duration (or large bandwidth) was used to excite the whole lung as a single slab in the \( z \) direction. Neglecting relaxation and off-resonance effects, the acquired 2D MR data is given by:

\[ s(k_x, k_y) = \int \int m(x, y) \exp(-i2\pi(xk_x + yk_y)) dx dy \]  \[1.18\]
where \( m(x, y) \) is the magnetization in the voxel in the imaging plane corresponding to the spin density, \( k_x \) and \( k_y \) are the location in the k-space along the frequency and phase encoding directions respectively. A 2D image is computed from the inverse Fourier transform \( \mathcal{F}^{-1} \) of the data in k-space as:

\[
m(x, y) = \mathcal{F}^{-1}\{s(k_x, k_y)\} = \int \int s(k_x, k_y) \exp(i2\pi(xk_x + yk_y)) dk_x dk_y
\]

The sampled k-space points define the scan trajectory. The geometry of the scan trajectory has a first-order impact on the quality of the reconstructed image. The selection of optimum k-space trajectories must consider the trade-off between the signal dynamics and SNR considering the gradient performance of the imaging system. Procedures for sampling of k-space data and resulting SNR and resolution are investigated in Chapter 3 using a point spread function (PSF) analysis as shown in Figure 1.6. Briefly, calculation of the PSF allows for quantitative evaluation of image blurring, through the full-width at half maximum (FWHM), and SNR for a corresponding read-out time \( T_{\text{read}} \), field-of-view (FOV) and imaging matrix.
Gradient Echo is a common frequency encoding method and considered a 1D MR signal, which encodes spatial information in the FID signal by frequency encoding. A 1D MR gradient echo signal is accomplished by applying a magnetic field gradient during the FID signal. This gradient spatially encodes the FID signal by first de-phasing (with a negative gradient lobe) and re-phasing the transverse magnetization signal (with a positive polarity gradient) as shown in Figure 1.7. As already mentioned in the previous section, the transverse magnetization is produced after an RF pulse application. Following the RF pulse, the application of the negative lobe of the $G_x$ gradient causes a very fast de-phasing of the FID signal. Then, the positive lobe of the $G_x$ gradient reverses the FID signal known as re-focusing, thereby creating an ‘echo’ at the echo time (TE) Consequently, the associated FID (or 1D k-space) data consists of a range of frequencies along the frequency-encoding gradient direction. This can be translated to 2D k-space data using an additional phase encoding gradient as explained in the next section.
Figure 1.7: 1D gradient echo timing diagram. Transverse magnetization signal re-phasing and de-phasing is shown for the 1D frequency encoding process. TE is the echo time.

1.8.11 2D Cartesian k-space Sampling

Cartesian sampling (i.e. uniform grid) is the conventional way MR data is sampled in k-space. Fast gradient recalled echo (FGRE) imaging is the most commonly used pulse sequence based on Cartesian k-space sampling as shown in Figure 1.8. A single line of k-space along the frequency encoding ($G_x$) direction is acquired after each RF excitation. The phase encoding gradient ($G_y$) and another RF pulses are then repeated to cover the next line of k-space in the y direction. The pulse repetition time (TR) is the time duration between the RF pulse applications between k-space lines. For 2D proton FGRE imaging, TR is often set to a value greater than $4 \times T_1$ for maximum signal
intensity. Thus, the total scan time for a $32 \times 32$ matrix size would be $32 \times 4 \times T_1$, resulting in a long scan time ($128 \times T_1 = 128$ s for $T_1 = 1$ s). Faster imaging is possible with Hp $^{129}$Xe gas imaging since the time for recovery of the magnetization is not needed. Higher temporal resolution is achieved for Hp imaging: approximately $32 \times TR$ with TR on the order of 10-20 ms. Thus the total scan time for Hp imaging is approximately 300 ms.

A representative coronal rat lung FGRE image obtained using 64 RF pulses is shown in Figure 1.9. The high number of RF pulse is a limitation for Hp imaging due to the non-renewable nature of hyperpolarized imaging. To deal with the problem of using a high number of RF pulses, each acquisition along the frequency encoding direction uses a small flip angle ($\alpha \approx 10^5$). Alternatively, more efficient k-space trajectories (e.g. spiral) can substantially reduce the number of RF pulses, allowing for higher flip angles and hence

Figure 1.8: 2D Gradient echo pulse sequence timing diagram (a) and Cartesian k-space sampling trajectory (b). TR represents the pulse repetition time. TE is the echo time. $G_x$ and $G_y$ represent the frequency and phase encoding gradients respectively.
increasing SNR. Additionally, faster k-space acquisition also allow for capturing of the gas exchange of $^{129}$Xe.

![Figure 1.9: Coronal 2D MR FGRE gas phase image of in vivo rat lung. Acquisition parameters include: 64 $\times$ 64 matrix size, and 6 cm FOV.](image)

1.8.12 Spiral k-space Sampling

An efficient approach for mapping the entire k-space following an RF excitation utilizes a single-shot spiral acquisition. Single-shot spiral is based on acquiring the k-space data in a spiral pattern using gradients ($G_x$ and $G_y$) applied as an increasing sinusoidal function of time as shown in

**Figure 1.10.** The duration of the data acquisition window is called the read-out time, $T_{\text{read}}$. The selection of $T_{\text{read}}$ on the order of $T_2^*$ (~2 ms) is desirable for increasing SNR and reducing blurring artifacts. Short $T_{\text{read}}$ is usually performed using high gradient amplitudes and slew rates. However, high gradient power can cause image artifacts including Eddy
current induced k-space distortions and time delay errors arising from gradient hardware for long $T_{read}$. Fortunately, these image artifacts can be corrected using k-space trajectory measurement methods (110). The single-shot k-space sampling provides a better tool for imaging pulmonary functions including perfusion and gas diffusion imaging (111,112). Additionally, $TR$ values can be used to create functional images for detection of gas exchange dynamics. The build-up of dissolved $^{129}$Xe signal between sequential images can be controlled by changing the TR values taking advantage of rapid single-shot imaging with short TE. Furthermore, obtaining TR-weighted rapid single-shot spiral images of PT, RBC and gas compartments separately can be achieved using advanced pulse sequence approaches such as Iterative Decomposition of water and fat with Echo Asymmetric and Least-squares estimation (IDEAL) described next.

Figure 1.10: Single-shot spiral pulse sequence timing diagram, including the RF pulse, gradient and read-out window (a) and the spiral k-space sampling trajectory (b).
1.8.13 IDEAL Approach for Imaging Chemical Species

IDEAL is a Dixon-based imaging method commonly used to exploit the difference in chemical shifts between fat and water to separately image fat and water in conventional proton MRI (113). The three-point Dixon method is based on acquisition of three consecutive images with a phase difference of 0, π, 2π. The phase differences are achieved by selecting echo times based on the chemical shift between fat and water. Three echo images are acquired to estimate the $B_0$ inhomogeneity, water and fat compartments (114,115). On closer inspection, the complex fat/water MR signal strength after Fourier transformation for a voxel at a chosen $TE$ value can be expressed as follows:

$$S(TE_n) = [W + F \exp(i\Delta w_{FW}TE_n)]\exp(i\gamma B_i TE_n)$$

[1.20]

where, $W$ and $F$ are water and fat fraction respectively, and $\Delta w_{FW}$ is the chemical shift difference between the fat and water. For each voxel, the signal is a mixture of signals from the fat and water as a function of echo time ($TE$). Because the chemical shift is known and $TE$ is controlled by the timing of the data acquisition, the corresponding signal strength equations for each image can be simplified as follows:

$$S(TE_n) = [W + F \cdot C(TE_n)]H(TE_n)$$

[1.21]

where $C(TE_n)$ is a constant which depends on TE and $H(TE_n)$ is the $B_0$ inhomogeneity factor. This equation contains three unknown contributions from $F$, $W$ and $H(TE_n)$. Therefore, acquisition of three echo images with different echo shifts allows separation of these three unknowns to purposely-separate fat and water. Since the chemical shift depends
on the $B_o$ inhomogeneity, three-point Dixon method correct for the effect of $B_o$ inhomogeneity. However, selection of echo shift with a phase difference of $\pi$ causes image artifact, reducing the accuracy of three-point Dixon for separating the water and fat (113). Imperfect selection of echo shifts can result in inaccurate calculation of the fraction of water to fat within each voxel. IDEAL combines asymmetrically acquired echoes with an iterative least-squares decomposition algorithm to maximize noise performance and address the dependence of water/fat separation on echo shift. Asymmetrically acquired echoes helps to reduce the scan times for substances with large chemical shift. For example, to obtain a phase difference of $\pi$ between compartments with small chemical shift (~10 ppm) requires a long echo time spacing (~340 ms) but large chemical shifts (~200 ppm) require a much shorter echo time spacing (50 ms). The iterative least-squares decomposition part of IDEAL stands for estimation of the fraction of each chemical shift compartments iteratively in multi steps (116). Additional improvement for estimation of fat/water decomposition has been achieved using phase difference of $2\pi /3$ by analyzing the noise performance (113). The noise performance of the a water/fat separation has been shown to be proportional to the SNR performance of the pulse sequence which is estimated based on the effective number of signal averages (NSA) (117). NSA for three point Dixon is given by:

$$NSA = \frac{2(1 - \cos \theta)^2}{1 + 2\cos^2 \theta}$$

[1.22]

where $\theta$ is the phase encoding increment. Thus the maximum SNR can be achieved for $NSA = 3$ at $\theta$ of $2\pi /3$ for three-point IDEAL. Therefore, the selection of echo times can be optimized using the NSA approach.
Optimization of NSA and other imaging parameters for IDEAL imaging will be discussed in detail in Chapter 3. To achieve the measurement of the gas exchange curves for PT and RBC compartments separately in this thesis, IDEAL is used in spiral k-space sampling fashion following a single RF pulse application. Four-point IDEAL is used to produce gas, PT and RBC images by varying TR which controls the build-up of the dissolved phase $^{129}$Xe signal. Thereby, gas exchange curves have been calculated from the variation in the image intensity as a function of TR.

1.8.14 Gas Exchange Models of Hyperpolarized $^{129}$Xe

Gas exchange curves represent the diffusion of $^{129}$Xe from gas phase to the PT compartment and from the PT to the RBC compartment as a function of time respectively. A representative gas exchange curve is shown in Figure 1.11(a). The concentration of $^{129}$Xe in each of the compartments is proportional to the build-up $^{129}$Xe signal which is usually obtained from whole lungs using the CSSR technique (89). Gas exchange curves are usually analyzed by fitting CSSR experimental data points by the Mansson exponential function (95). In this study, the IDEAL imaging technique is used to obtain gas exchange curves, which have been fitted by the Mansson’s gas exchange model to explore lung function regionally. The gas exchange curves provide information about the concentration of $^{129}$Xe gas that diffuses from alveolar air space into the PT and RBC compartments. As a function of time the concentration of $^{129}$Xe build-up in PT compartment increases exponentially giving rise to MR image intensity within approximately 100 ms (Figure 1.11 (a)). The linear slope after 100 ms corresponds to the linear slope ($S_1$) due to the saturation of $^{129}$Xe concentration in the PT compartment. The dynamics of $^{129}$Xe transfer from
alveolar air space to the PT compartment is modelled using one-dimensional diffusion equations. The diffusion from the PT to RBC is also included in the blood perfusion with a constant flow as shown by the red color in Figure 1.11 (b).

The exponential fitting function from the Mansson model is given as follows:

\[ S(TR) = S_o (1 - \exp(TR/\tau)) + S_1 TR \]  

where, \( S_o \) is the y-intercept of the linear slope, \( \tau \) is the exponential gas transfer time constant corresponding to the fast exponential increase in the dissolved phase concentration due to the diffusion and \( S_1 \) is the linear slope corresponding to the accumulation of \(^{129}\text{Xe}\) concentration after fast diffusion. \( S_o, S_1, \) and \( \tau \) are subsequently used to calculate the average pulmonary tissue thickness, \( L_{PT} \), and capillary thickness, \( L_C \). Detailed information about the mathematical derivation can be found in Mansson et al. (95). Additionally, the relative blood volume, \( V_{RBC} \), can be calculated by \( L_C / (L_C + L_{PT}) \).

Figure 1.11: (a) A representative gas exchange curve is obtained by fitting experimental data points by the Mansson exponential function Eqn. [1.23]. \( S_o \) is the y-intercept of the linear slope, \( S_1 \) is the linear slope, \( TR \) represents the pulse repetition time. (b) Geometry of the Mansson’s gas exchange model is shown. \( L_{PT} \) and \( L_C \) are the pulmonary tissue thickness and capillary thickness, respectively.
1.9 Thesis Hypothesis and Objectives

The hypothesis of this research is that hyperpolarized $^{129}$Xe MRI will provide measurement of gas exchange changes, specifically $L_{PT}$ and $V_{RBC}$, in an early-phase rat model of RILI which will correlate with histology. To address this hypothesis, new Hp $^{129}$Xe approaches are required to increase signal-to-noise ratio (SNR), especially from the dissolved phases of $^{129}$Xe in the PT and RBC compartments. The first specific goal of thesis was to increase the SNR of dissolved phase $^{129}$Xe images by developing a Transmit-Only / Receive-Only (TO/RO) RF coil. The second specific goal was to develop an MRI technique with sufficient temporal and spatial resolution for quantification of the regional changes in the dynamics of the $^{129}$Xe gas exchange in the RBC and PT compartments. Therefore, a 2D IDEAL MRI pulse sequence using single-shot spiral k-space sampling was optimized and implemented for simultaneous Hp $^{129}$Xe imaging of gas, PT, and RBC compartments. Finally, the regional gas exchange abnormalities in a rat model of RILI associated with partial thorax irradiation was investigated and correlated with histology measurements.

1.10 Thesis Outline

Chapter 2 describes the design and construction of a novel TO/RO RF coil for rat imaging. Chapter 2 is a reproduction of the accepted manuscript by Concepts in Magnetic Resonance Part B: Magnetic Resonance Engineering 2014 (DOI: 10.1002/cmrb.201288). The coil sensitivity was optimized through numerical simulations. Performance of the new TO/RO coil was investigated using a thermally-polarized phantom and in vivo experiments.
Chapter 3 describes the development and optimization of a novel imaging technique (IDEAL with spiral k-space sampling) that provides regional quantification of the dynamics of the gas exchange. Chapter 3 is a reproduction of the accepted manuscript by *Magnetic Resonance in Medicine* 2015, Manuscript ID: MRM-15-16025. A theoretical analysis was performed to optimize the spiral k-space read-out time, echo time, flip angles, spatial resolution and required gradient slew rate and amplitude respectively. The feasibility of imaging the $^{129}$Xe gas, PT and RBC, and measuring the dynamics of the gas exchange was investigated *in vivo* in healthy rat lungs.

Chapter 4 is a reproduction of a submitted manuscript to *Medical Physics* 2015. Chapter 4 investigates the feasibility of detecting early RP in a rat model of RILI involving right lung irradiation 2-weeks post-irradiation. Taking advantages of developments made in Chapter 2 and 3, the gas exchange curves were measured separately for PT and RBC compartments for the left and right lungs. Average PT thickness, ($L_{PT}$) and RBC relative volumes ($V_{RBC}$) were measured from the left and right lungs separately. The average PT and RBC volumes were statistically compared between the irradiated and non-irradiated cohorts.

Chapter 5 includes a summary of discussion of the thesis, including the contribution of this work to the detection of RILI in its early stage. Additionally, future work is discussed including further improvements in polarization, modeling of gas exchange, and MR pulse sequence design for transition to clinical imaging.
1.11 References


Chapter 2: RF Coil Configuration for Hyperpolarized $^{129}$Xe

This chapter is adapted from the published paper: Doganay O, Thind K, Wade T, Ouriadov A, Santyr GE. Transmit-Only/Receive-Only Radiofrequency Coil Configuration for Hyperpolarized $^{129}$Xe MRI of Rat Lungs. Concepts Magn Reson. doi:10.1002/cmr.b.21288

2.1 Introduction

Hyperpolarized $^{129}$Xe is a useful inhaled contrast agent for Magnetic Resonance Imaging (MRI) of the lung air space and promises to provide unique functional information due to its solubility and significant chemical shift (200 ppm) when dissolved in lung tissues (1). In particular, MRI of the gas and dissolved phases of $^{129}$Xe can be used to quantify the rate of transfer of xenon between the air space and the lung tissue and the red blood cells (RBC), providing regional information about pulmonary gas exchange (2). Hyperpolarized $^{129}$Xe MRI has been useful for development of biomarkers in rodent (i.e. rat and mouse) models of lung injury and disease, including emphysema (3,4). Rodent models provide a powerful approach for modeling of lung diseases and developing new therapeutic targets (5), but the tools are only beginning to be established.

Imaging of $^{129}$Xe dissolved in lung tissues is challenging due to modest solubility and relatively low tissue density in the lung, resulting in a relatively low signal, which is approximately two percent of the hyperpolarized $^{129}$Xe gas signal (6-8). Furthermore, lung volumes in rats are typically small (~ 5 to 10 mL) (9), requiring sensitive radiofrequency (RF) receiver coils for detection. Another important consideration is that hyperpolarized
MRI benefits by the use of accurate flip angles (i.e. transmit $B_{1+}$ field) delivered uniformly over the imaging volume, particularly to exploit variable flip angle approaches which make most efficient use of the non-renewable magnetization (10,11). Hyperpolarized $^{129}$Xe MRI of the rat lung therefore would benefit from an RF coil configuration which combines high sensitivity in receive mode with a uniform transmit field ($B_{1+}$) over the entire lung volume. RF sensitivity can be improved by using a surface coil to maximize sensitivity ($B_1$) over a smaller region of interest (12-14). However, when using this same coil to transmit, surface coil sensitivity decreases with increasing distance from the coil elements leading to an inhomogeneous $B_{1+}$ field (15,16). A potential solution to this problem, is to use a surface coil for receive and a birdcage transmit coil for transmission of the RF pulse, the latter providing a homogeneous transmit field across the lungs. Such transmit-only/receive-only (TO/RO) coil configurations have been demonstrated for clinical hyperpolarized $^{129}$Xe and $^3$He gas MRI of human lungs (17-19). Additionally, the use of TO/RO coil configurations has been motivated by the desire to improve SNR or reduce image acquisition time as previously shown for other nuclei including $^{13}$C (20) and $^{23}$Na (21) for small animal imaging.

Despite the significant interest in hyperpolarized $^{129}$Xe MR imaging of rat lungs, the literature mostly describes the use of birdcage transmit/receive (T/R) volume coils (9,22), with little attention to TO/RO approaches. A custom-made TO/RO coil configuration was developed for hyperpolarized $^{129}$Xe imaging of rat lungs at low field (0.07T) and optimized by reducing the noise using Litz wire and geometric de-coupling between saddle-shaped transmit coils and multi-turn saddle-shaped receive coils (23,24). Another TO/RO design has been proposed at high field based on the use of two volumetric birdcage coils for
imaging of dissolved phase hyperpolarized $^{129}$Xe in rat brain (25). In this study, we present a novel RF transmit-only/receive-only (TO/RO) coil configuration providing excellent transmit uniformity as well as high SNR for hyperpolarized $^{129}$Xe MR lung imaging of rat lungs at 3T (35.34 MHz). Transmission is accomplished by a birdcage coil with uniform $\mathbf{B}_{1+}$ field uniformity and reception is performed by a closely-coupled saddle-shaped surface coil conforming to the rat lung and accommodating ventilation, surgical access and drug delivery. SNR and $\mathbf{B}_{1}$-uniformity improvement of the TO/RO coil configuration compared to a commercially-made transmit/receive (T/R) birdcage volume coil are demonstrated theoretically and verified experimentally in $^{129}$Xe gas phantoms.

2.2 Methods

A TO/RO coil configuration was designed and constructed consisting of three components: (i) a high-pass birdcage transmit coil that produced a homogeneous $\mathbf{B}_{1+}$ magnetic field which could also be used in T/R mode, (ii) a saddle-shaped receive-only surface coil that couples closely to the rat lung, (iii) a fast switching circuit and RF shielding as shown in Figure 2.1 (a). Switching between transmit and receive modes was achieved as previously described (20). Briefly, a fast MOSFET switch circuit was gated using the DC (5 V) component of the transmit signal from the MRI system. Since the gate threshold voltage of the MOSFET circuit (FDP8030L, Fairchild corp., California, USA) was 2V, the 5 V bias was more than adequate to switch between the transmit and receive modes. On transmit, the MOSFET switch circuit was DC biased, and a fast switch power supply (U8002A, Agilent Technologies, Malaysia) provided a 5V DC bias to activate PIN diodes in order to switch on the transmit coil and de-tune the receive coil using a de-tuning circuit. On receive
(no MOSFET bias), the receive coil was returned to resonance (35.34 MHz) and the transmit coil was deactivated through the use of PIN diodes (um9415, Microsemi corp., Watertown, MA), located in each rung element. DC bias from the constant current power supply to the PIN diodes passed through two chokes (Axial RF choke 3.9 µH, Bourns Inc., Riverside, CA) to isolate the high frequency noise of the power supply as shown in a simplified circuit diagram in Figure 2.1 (b).

Figure 2.1: Schematic view of the TO/RO configuration (a) consisting of: (i) high-pass birdcage transmit coil, comprised of two end-rings of 12 cm diameter and eight rungs with a length of 16.5 cm each, (ii) saddle-shaped surface coil, and (iii) RF shielding. Tuning capacitors are placed in the end-rings. Simplified circuit diagram of transmit coil (b). CB denotes blocking capacitor to block DC component of the transmit signal, $C_M$ and $C_T$ are tuning and matching capacitors, respectively. Simplified circuit diagram of saddle-shaped, receive surface coil (c). $U$ denotes the aperture angle. The distance from points A to B is 55 mm, from C to D is 60 mm and from E to F is 23 mm.
For comparison purposes, the transmit birdcage coil dimensions were chosen to be similar to a commercially-made rat T/R birdcage coil (Morris Instruments, Ottawa, Canada) with a similar RF shield for both coil configurations. The constructed birdcage transmit coil consisted of two end-rings of 120 mm diameter and eight rungs with a length of 165 mm. The tuning capacitor, $C_T \approx 400 \text{ pF}$ (100B11160FT500X, American Technical Ceramics corp, Jacksonville, FL) values were roughly estimated using a birdcage builder program (Birdcage Builder Version 1.0 Software, Penn State Hershey Medical Center, Hershey, PA). A network analyzer (Agilent E5061B, Agilent, Santa Clara, CA) and a pickup coil were used to measure the resonance properties of each rung separately, as well as the entire transmit coil. Fine tuning was done by adding smaller (a few pF) capacitors until each rung resonated exactly at 35.34 MHz. The Smith chart on the network analyzer was used to perform impedance matching to a 50 ± 2 ohm transmission line, and S-parameters after the coil was loaded with 300 mL 50 mM NaCl solution (26). The Q factor values were measured from the $S_{12}$ response curve at the -3dB level from the TO/RO coil in both transmit and receive modes and compared to the commercial T/R coil (26-28). Two pickup coils, each with diameter of 3cm were placed 4cm away from either side of the RF coils. Loading factors factor ($L_f$) were calculated within measurement uncertainty of ±10% as previously described (23,27).

A saddle-shaped, receive-only surface coil was designed and constructed around a half cylinder acrylic tube with a diameter of 55 mm as shown in Figure 2.1 (c). Based on measurements of the rat lung and thorax anatomy (diameter and length), fixed lengths of 60 mm (points C to D in Figure 2.1 (c)) and 55 mm (points A to B in Figure 2.1 (c)) were chosen to conform to the rat lungs. The aperture angle, $\Phi$, of the saddle coil was optimized
through numerical simulations to maximize SNR and $B_1$-uniformity across the lungs (from point A to B in Figure 2.1 (a)) as described in the next section. The coil was built from 5 mm-wide copper foil and attached to the acrylic tube with adhesive. Tuning was accomplished using capacitors connected in series as shown in the simplified circuit diagram in Figure 2.1 (c). The receive coil was actively de-tuned during transmission using the same DC bias used to de-couple the transmit coil on receive. A capacitance circuit was used for fine-tuning and matching when the coil was loaded with a rat.

The mechanical ventilation of the rat lungs has been described previously (29,30). A large surgical incision area was required across the neck for intubating the trachea and to secure the intubation catheter. The choice of a saddle-shaped surface coil was determined based on accommodating ventilation, surgical access and drug delivery. Due to the complexity of acquiring forced-ventilation hyperpolarized $^{129}$Xe MRI (i.e. dislocation of the catheter, localization of the rat lungs, bleeding), imaging of the rat in the supine position using a saddle-shaped RF coil provides the necessary access to the surgical area during extensive imaging time.

2.2.1 Numerical Modeling

SNR improvement was achieved by maximizing the $B_1$-magnetic field for a unit current flow through the receiving coil and computing sensitivity maps, according to the reciprocity theorem. In order to estimate the sensitivity maps, the $B_1$-magnetic field was computed by solving the Biot-Savart equation in 3D using finite element modeling (Comsol Multiphysics, COMSOL 4.2a) for both the T/R (shown in Figure 2.3 (a)) and TO/RO geometries shown in Figure 2.1 (a). The model geometries including birdcage coil,
receive coil and shielding were meshed using a free tetrahedral mesh with a minimum element size of 0.01 mm. The shielding was modelled as a magnetic shielding boundary condition with the relative permeability of 7000 and thickness of 1 mm (based on shielding material). Rungs and rings were modeled corresponding to copper foil’s electrical properties. The model was solved in the magnetic field interface under the AC/DC branch (31). Next, sensitivity maps were calculated by normalizing $B_1$ by the surface area of each receiver coil in order to: (i) compare coil performances and (ii) calculate the image uniformity. A percentage uniformity, $U$, was calculated from simulations and axial images for both coil configurations for four identical circular regions-of-interest (ROIs) to approximate the coverage of the lungs as shown in Figure 2.4(b) (32). The percentage uniformity was calculated for each ROI as:

$$U = 100 \times \left(1 - \frac{(S_{\text{max}} - S_{\text{min}})}{(S_{\text{max}} + S_{\text{min}})}\right)$$

where $S_{\text{max}}$ and $S_{\text{min}}$ are the maximum and minimum signal intensities within the ROI.

Considering that the rat lung occupies a volume at the center of the transmit coil with dimensions of 4cm x 4cm x 4cm, the geometry of the saddle-shaped coil was optimized for a body weight up to 400 g (33). $B_1$ field profiles from point A to B, C to D and E to F were investigated as a function of aperture angle ($\Phi$). The total sensitivity was quantified as a function of aperture angle using of the area under simulated $B_1$ field profiles from point A to B as shown in Figure 2.2 (b). The SNR was expected to be proportional to the area under the sensitivity profiles as shown in Figure 2.2 (c) for the same RF shielding and the phantom. The noise variations including conductivity, radiation, and sample inductive loses, for each aperture angle was considered by scaling the area under the sensitivity
profiles to the coil surface area (34). Finally, the uniformity was maximized to determine the best aperture angle.

![Graph](image)

Figure 2.2: Plot of theoretical $B_1$ (a) from point E to G (in Figure 2.1), and (b) from point A to B (in Figure 2.1) for the receive coil at aperture angles of 100°, 120°, and 140°, respectively. Plot of the area under the simulated $B_1$ field profiles from point A to B as a function of aperture angle (b). The SNR profile from A to B, from C to D, and from E to G as a function of aperture angle in log scale (c).

### 2.2.2 Experimental measurements

A thermally-polarized $^{129}$Xe gas phantom was used to compare the signal-to-noise ratios of the TO/RO coil configuration and the T/R coil since the signal from thermal phantom did not depend on differences in polarization and relaxation time effects between measurements. The thermally-polarized phantom consisted of a glass cylinder of length 52 mm and diameter 35 mm, with a volume of approximately 50 ml. The glass phantom was filled with 200 ml of $^{129}$Xe gas (80% enriched) and 200 ml O$_2$ gas (to reduce $T_1$ relaxation time down to ~1 s) at room temperature in order to provide a measurable $^{129}$Xe gas phase axial image. 2D projection images of the thermally-polarized phantom in the axial view were obtained for both coil configurations at 3T (MR750, GEHC) using a 2D fast gradient-recalled echo (FGRE) pulse sequence (TR = 4.0 s, TE = 6 ms, FOV = 10 cm × 10 cm, matrix of 64 × 64, BW = 2.0 kHz, number of averages = 16, flip angle = 90 ± 1 degrees).
A free induction decay (FID) was used to find the 90° flip angle by adjusting the transmit gain to achieve a maximum signal. Image SNR was calculated using the dual acquisition technique (35). The mean value of a circular region of interest (15 pixel diameter) covering most of the axial image was divided by the standard deviation from the same ROI within a subtraction image. A B\textsubscript{1+} map was obtained from the TO/RO coil using the thermally-polarized phantom and a non-inverted double angle Look-Locker (niDALL) technique with nominal flip angles of α\textsubscript{1} = 60°±1° and α\textsubscript{2} = 2α\textsubscript{1} sequence (TR = 5.0 s, FOV = 8 cm × 8 cm, matrix of 32 × 32, BW = 2.0 kHz, number of averages = 8) (36-38).

For uniformity measurements, a 2D fast gradient-recalled echo sequence (FGRE) pulse sequence implementing a variable flip angle (39) used to acquire projection images of a hyperpolarized \textsuperscript{129}Xe gas phantom in the axial view for both coil configurations (TR = 100 ms, TE = 4 ms, FOV = 10 cm × 10 cm, matrix of 64 × 64, BW = 5.0 kHz). The hyperpolarized \textsuperscript{129}Xe gas phantom consisted of a 125 mL plastic bottle (2401-0250 Wash bottles, Thermo Scientific Nalgene, Rochester, NY) of diameter 55 mm filled with hyperpolarized, naturally-abundant \textsuperscript{129}Xe gas (polarized to approximately 5%) using a home-built polarizer described previously (11).

Imaging of a healthy Sprague Dawley rat (380 g, Charles River Laboratories, Saint-Constant, Canada) was performed using a MRI-compatible mechanical ventilator (GE Healthcare, Malmo, Sweden) for delivery of Hp \textsuperscript{129}Xe to the lung following an animal care protocol approved by the Animal Care and Veterinary Service of Western University (30). 2D projection MR images were obtained from the same rat in the supine position following five breaths of Hp \textsuperscript{129}Xe. The FGRE pulse sequence parameters were as follows: TR = 20 ms, TE = 4 ms, FOV = 7 cm × 7 cm, matrix of 48 × 48, BW = 4.0 kHz. Coronal images
were repeated six times in intervals of two minutes. A fresh bag of Hp $^{129}$Xe was used for each coil within 15 minutes following the preparation of polarized gas to ensure that the polarization (approximately 5%) was the same for each coil.

2.3 Results

The electrical properties of the TO/RO in both transmit and receive modes and the commercial T/R birdcage coils using the network analyzer with a span of 5 MHz are shown in Table 2.1. The reflected power, $S_{11}$, measurements showed that the variations in resonance frequencies for both transmit and receive modes (loaded) were less than 20 kHz at the MR scanner (3 T) for the TO/RO coil. The loading factors were 0.27 and 0.24 for the TO/RO in receive mode and transmit mode respectively. For comparison, the loading factor for the commercial T/R birdcage coil was 0.23. Isolation between quadrature channels, $S_{12}$, was measured to be greater than 20 dB for the phantom and rat loading. This indicates that there is adequate decoupling between quadrature channels in transmit. To minimize the coupling between birdcage coil and saddle-shaped receive coil for the TO/RO configuration, the resonance frequency for the saddle-shaped surface coil was detuned to 5 MHz away from the resonance frequency of transmit coil during the transmit mode.

Table 2.1: Network analyzer measurement of the electrical properties of the TO/RO coil in receive mode and transmit mode respectively and the commercial T/R birdcage coil configurations.

<table>
<thead>
<tr>
<th></th>
<th>Q-factor Unloaded</th>
<th>Q-factor Loaded</th>
<th>$S_{11}$ (dB)</th>
<th>Isolation (dB)</th>
<th>Resonant Frequency (MHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TO/RO receive mode</td>
<td>220</td>
<td>160</td>
<td>41.2</td>
<td>−30.1</td>
<td>35.34</td>
</tr>
<tr>
<td>TO/RO transmit mode</td>
<td>143</td>
<td>108</td>
<td>29.7</td>
<td>−24.2</td>
<td>35.34</td>
</tr>
<tr>
<td>Commercial T/R birdcage coil</td>
<td>170</td>
<td>130</td>
<td>34.1</td>
<td>−24.8</td>
<td>35.34</td>
</tr>
</tbody>
</table>
The effect of varying the aperture angle, $\Phi$, on the simulated uniformity of the saddle coil is shown in Figure 2.2 (a) and (b) where $B_1$-field profiles from point A to B and E to G are compared for different aperture angle values of the TO/RO receive coil as well as the T/R coil configuration. The SNR profiles as a function of aperture angle in Figure 2.2 (c) from C to D and E to F are negligible compared to profile A to B since the SNR is dominated by the rungs of the saddle shaped coil at the ROI. Optimization the aperture angle to achieve the highest SNR is dominated by the choice of AB profile. The optimum aperture angle was determined to be approximately $120^\circ$ from Figure 2.2 (c) for the highest $B_1$-sensitivity and uniformity in agreement with a previous study (40).

Table 2.2: Comparison of SNR measured in the thermally-polarized gas phantom using the TO/RO and commercial T/R birdcage coil configurations respectively.

<table>
<thead>
<tr>
<th></th>
<th>Signal</th>
<th>Noise</th>
<th>SNR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Commercial T/R</td>
<td>$1.4 \times 10^4$</td>
<td>$4.4 \times 10^3$</td>
<td>4.5</td>
</tr>
<tr>
<td>birdcage coil</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TO/RO coil</td>
<td>$4.8 \times 10^4$</td>
<td>$5.1 \times 10^3$</td>
<td>13.3</td>
</tr>
</tbody>
</table>
Figure 2.3 shows a schematic of commercial T/R birdcage and TO/RO coil set-up and corresponding FGRE axial images from the thermal gas phantom. The corresponding SNR values are shown in Table 2.2. The TO/RO coil design provided an SNR improvement of a factor of three compared to the commercial T/R birdcage coil configuration. Table 2.3 shows the simulated and experimental uniformity values for four ROIs, U, obtained from the axial images of hyperpolarized gas phantom shown in Figure 2.4 (a) and (d), and the simulated B1-field in Figure 2.4 (b) and (e) for both the TO/RO coil and commercial T/R birdcage coil. Figure 2.4 (c) and (f) show coronal MR images of rat lungs obtained from the same rat using TO/RO and commercial T/R birdcage coil configurations. Although the image SNR is not optimal due to low polarization, the improvement in the SNR with the TO/RO coil is apparent. In particular, the SNR improvement provided by the TO/RO coil allows visualization of the right lobe of lungs otherwise not apparent using the commercial T/R birdcage coil. SNR values were 19.7/0.63 and 7.5/0.62 for the TO/RO and T/R coil configurations respectively.

Table 2.3: Experimental and Simulated Axial Percentage B1-ROI Uniformity (U) for the Hyperpolarized Gas Phantom for both TO/RO and Commercial T/R Birdcage Coil Configurations [as shown in Fig. 2.4(b)].

<table>
<thead>
<tr>
<th></th>
<th>First ROI</th>
<th>Second ROI</th>
<th>Third ROI</th>
<th>Fourth ROI</th>
</tr>
</thead>
<tbody>
<tr>
<td>(U_{TO/RO}(Simulation))</td>
<td>70.8</td>
<td>82.0</td>
<td>78.7</td>
<td>70.1</td>
</tr>
<tr>
<td>(U_{TO/RO}(Experiment))</td>
<td>68.9</td>
<td>79.6</td>
<td>76.8</td>
<td>69.4</td>
</tr>
<tr>
<td>(U_{T/R}(Simulation))</td>
<td>98.0</td>
<td>97.2</td>
<td>97.7</td>
<td>97.4</td>
</tr>
<tr>
<td>(U_{T/R}(Experiment))</td>
<td>95.3</td>
<td>95.0</td>
<td>95.1</td>
<td>93.0</td>
</tr>
</tbody>
</table>
Figure 2.3: Schematic of thermal gas phantom and commercial T/R birdcage coil configuration (a) consisting of two end-rings of 12 cm diameter and eight rungs of length 18.0 cm each. Also shown is the corresponding axial image from the phantom (b). The saddle-shaped surface coil is shown schematically in (c) (transmit coil has been omitted) along with the phantom. Also shown is the corresponding axial image from the phantom (d).
Figure 2.4: Axial images of hyperpolarized gas phantom using the commercial T/R birdcage coil (a) and the corresponding simulated B1- sensitivity map (b). Coronal image of rat lungs using the commercial T/R birdcage coil (c). Axial image of the hyperpolarized gas phantom using the TO/RO coil configuration (d) and the corresponding simulated B1- sensitivity map (e). Coronal image of rat lungs using the TO/RO coil (f).

The calculated $B_{1+}$ map and flip angle profile for the TO/RO coil is shown in Figure 2.5 (a) and (b) respectively. The flip angle distribution was relatively homogenous with a mean flip angle of 60.2±1.6°. As expected, the mean flip angle was approximately the same as the prescribed nominal flip angle $\alpha_1 = 60\pm1^\circ$. 
Figure 2.5: (a) B$_1$ map and (b) the flip angle profile (from A to B as shown in Figure 2.1) obtained from the TO/RO coil using the thermally polarized phantom.

2.4 Discussion

The TO/RO coil configuration provides a homogeneous excitation field and a high sensitivity for $^{129}$Xe MRI of the rat lung. MR images from a thermally-polarized phantom confirm that a novel TO/RO configuration provides an improvement in SNR of up to a factor of approximately three compared to a commercial T/R birdcage coil of comparable geometry. The Q factor measurements allow a preliminary evaluation of the coil sensitivity depending on the various conditions including the coil dimensions (i.e. the B$_1$-sensitivity and inductance), shielding, resonance frequency and loading conditions. As expected, the Q factor of the TO/RO coil in transmit mode was 20% that of the commercial T/R birdcage coil due to the additional required pin diodes and chokes (as shown in Figure 2.1 (b)) for the switch circuits. We also noticed that the SNR performance of the TO/RO coil in T/R mode reduced after adding the necessary components and receive coil. Considering the same birdcage coil dimension used in our TO/RO and the commercial T/R coil configurations, the SNR of TO/RO coil in T/R mode was expected to be lower than the commercial T/R coil proportional to the square root of the Q factor (23). Although the
quality factors between the commercial T/R birdcage coil and our saddle-shaped receive TO/RO coil cannot be directly compared due to the difference in dimensions, the loading factors can be compared to estimate the ohmic and electromagnetic losses. Based on the benchtop experiments, $L_f$ was approximately the same for both coil configurations confirming that the coil is operating in the coil noise dominated regime and loading the coil with a rat-sized conductor (300 gr, 0.48S/m) does not significantly affect the SNR.

The transmit coil of the TO/RO configuration used here was designed to take advantage of the highly homogeneous transmit field of the birdcage geometry for imaging of hyperpolarized $^{129}$Xe gas and dissolved phases, known to be sensitive to flip angle variations. Therefore, investigation of the TO/RO configuration in a “T/R” mode was not considered in the imaging and SNR comparison due to the expected lower efficiency compared to the purpose-built commercial birdcage T/R coil. The commercial T/R birdcage coil used here did not require the additional switch circuits but had the same birdcage coil dimensions as the TO/RO configuration. Therefore, it was considered a reasonable gold standard for the purposes of comparison between the TO/RO mode and a best-case T/R mode. Nonetheless, one of the limitations of the TO/RO design used here was the lower efficiency in T/R mode which could presumably be improved in future if required.

When the aperture angles of receive coils are compared, both 120° and 140° showed a high sensitivity (Figure 2.2 (c)). However, the $B_1$-sensitivity profile at 120° is much more uniform compared to 140° with a strong increasing signal toward the edges and decreasing at the center. This confirms that the choice of receive aperture angle of 120° was a good trade-off between overall $B_1$-sensitivity and uniformity. Further SNR improvements with
the TO/RO approach should be possible using a multichannel receiver coil design (17,19,41).

The numerical simulations of $\mathbf{B}_{1+}$ uniformity for the TO/RO configuration shows reasonable agreement with the experimental values (Table 3). Simulated $\mathbf{B}_{1+}$ uniformity was approximately 70% at ROI one and four, and approximately 80% at ROI three and four for the axial images (Figure 2.4 (e) and Table 2.3). As expected, the T/R coil had better uniformity (~95%) for the same ROIs (Fig 4 (b)). The difference between experiments and simulated uniformity (~3%) are likely due to the small variations in excitation flip angle. The percentage variation in the simulated $\mathbf{B}_{1+}$ uniformity is less than 3% at each of the ROIs, which was in agreement with the early experimental measurement of $\mathbf{B}_{1+}$ for the commercial T/R coil (39). Although the $\mathbf{B}_{1+}$ maps of the TO/RO in T/R mode were not measured in this work, they are expected to be better than $\mathbf{B}_{1+}$ maps of the TO/RO since it includes the $\mathbf{B}_{1+}$ inhomogeneity due to the receive coil. $\mathbf{B}_{1+}$ profile obtained using TO/RO coil exhibited a uniform flip angle distribution with flip angle variations less than 3% of the nominal flip angle which was similar to the previously published study (20).

The thermally-polarized phantom experiments were more consistent for comparing the SNR between TO/RO and T/R coil configurations since the $in vivo$ signal strongly depends on animal ventilation conditions, including polarized bag pressure, ventilation or surgical related defects of the lungs while transferring from one coil to another, and the amount of polarization. Although the signal is sensitive to the experimental conditions, similar noise values were observed when the coils were loaded with rat and thermally-polarized phantoms. The results show the SNR improvement corresponds to an increased sensitivity for the surface coil for both the thermal phantom and $in vivo$ (loaded) configurations.
2.5 Conclusion

The use of a TO/RO radiofrequency coil configuration is advantageous for MRI of hyperpolarized $^{129}$Xe providing an approximate three-fold SNR improvement, with a $\mathbf{B}_{1+}$ transmit field variation of less than three percent. Although the $\mathbf{B}_{1}$ reception sensitivity is less uniform (approximately 30 %) compared to a T/R (i.e. birdcage) coil, this can be corrected by post-processing based on the simulated or measured reception sensitivity profile of the receive coil.
2.6 References


11. Ouriadov AV, Lam WW, Santyr GE. Rapid 3-D mapping of hyperpolarized He-3 spin-lattice relaxation times using variable flip angle gradient echo imaging with


31. Multiphysics C. MRI Birdcage Coil: Comsol Model Library Version. 3.5


Chapter 3: Hyperpolarized Dissolved $^{129}\text{Xe}$ MRI of the Rat Lung


3.1 Introduction

Hyperpolarized (Hp) $^{129}\text{Xe}$ gas can provide an increase in NMR signal of up to a factor of 100,000 compared to thermally polarized $^{129}\text{Xe}$ gas. MRI of Hp $^{129}\text{Xe}$ gas following inhalation enables unique functional and microanatomical lung imaging measurements, including ventilation defects (1) and apparent diffusion coefficient (ADC) (2). Although most of the inhaled Hp $^{129}\text{Xe}$ lung signal originates from the gas phase, approximately 2% of the signal originates from $^{129}\text{Xe}$ dissolved in the pulmonary tissue (PT, or barrier tissues) and red blood cell (RBC) compartments (3). These dissolved phase $^{129}\text{Xe}$ signals can be separated using MR spectroscopy approaches, taking advantage of the significant chemical shifts, relative to the gas phase, of $^{129}\text{Xe}$ dissolved in PT (197 ppm) and RBC (217 ppm) compartments, with respect to the gas phase (4). Furthermore, by measuring the time course of the lung RBC and PT $^{129}\text{Xe}$ signals following spectrally-selective RF saturation (e.g. chemical shift saturation recovery, CSSR), measures of perfusion (5) and pulmonary gas exchange (6) respectively can be derived. Pulmonary gas exchange measured with Hp $^{129}\text{Xe}$ CSSR spectroscopy in whole lungs has been shown to be sensitive to alveolar tissue
thickening, which is associated with early pneumonitis (i.e. inflammation) in radiation-induced lung injury (RILI) (7) and in idiopathic pulmonary fibrosis (8).

MRI pulse sequences for imaging dissolved phases of Hp $^{129}$Xe in the lung need to address the following specific challenges: (i) low lung $^{129}$Xe $T_2^*$ (~2 ms at 2 T (9)) leading to rapid signal loss and image blurring, (ii) poor SNR due to low density of $^{129}$Xe in the RBC and PT compartments and (iii) short breath-hold time (~ 10 s). An image acquisition scheme that acquires k-space with the fewest number of RF pulses, and a read-out time on the order of $T_2^*$ would enable high flip angles to be used, thus significantly increasing the SNR, spatial and temporal resolution. Improved temporal resolution would also allow improved SNR via increased replenishment by the gas phase within a short breath-hold interval, as well as quantification of the exchange between gas, PT and RBC compartments.

A 3-point Dixon approach has been reported for separation of gas and dissolved phases within a single breath-hold at both 0.2 T (10) and 3 T (11); however, dissolved phase SNR was limited in that study by the use of relatively long echo times and low flip angles. PT and RBC $^{129}$Xe signals have been spectrally resolved using a one-point variant of the Dixon technique for 2D imaging of both rats (9) and humans (12). This approach has also been extended to 3D rat imaging within a breath-hold time of approximately 16 seconds by using the replenishment of xenon from the gas phase between k-space acquisitions (13). A recent study by Qing et. al. took advantage of the use of hierarchical IDEAL (Iterative Decomposition of water and fat with Echo Asymmetry and Least-square estimation) to image the gas, PT and RBC phases simultaneously within a scan time of a single breath-hold (approximately 11 s) based on the independent excitation of the gas and dissolved phase compartments using a 3D radial pulse sequence (14). In this work, however, the
dissolved phase magnetization was excited with a relatively low flip angle and without substantial replenishment of the dissolved phase compartments by the gas phase.

A robust and efficient IDEAL imaging technique based on a single-shot spiral acquisition and echo-time shifting in between excitations has been reported for Hp $^{13}$C-pyruvate metabolic imaging (15). Additionally, Hp [1-$^{13}$C] pyruvate metabolic imaging using a single-shot spiral acquisition within the $T_2^*$ relaxation time (~40 ms) and a 90° selective RF pulse to maximize SNR has been demonstrated in phantoms and *in vivo* (16).

In this study, a single-shot spiral IDEAL approach was designed and implemented for rapid imaging of Hp $^{129}$Xe gas, PT and RBC compartments in the rat lung. The spiral read-out time was optimized to maximize both the spatial resolution and SNR of the dissolved phase Hp $^{129}$Xe images. The echo time spacing was also optimized using the effective number of averages (NSA) approach. The optimal flip angles were calculated through a theoretical Bloch equation analysis including the effect of gas exchange. The feasibility of the method was demonstrated for three-point IDEAL imaging of gas/PT compartments and four-point IDEAL imaging of gas/PT/RBC compartments in rat lungs. Two-dimensional IDEAL images using separate breath-holds at different TR values were acquired to demonstrate the feasibility of regional measurement of gas exchange.
3.2 Theory

3.2.1 Point Spread Function (PSF) Analysis

The PSF was calculated to investigate the effects of read-out time, $T_{read}$, (as shown in Figure 3.1) on image SNR and spatial resolution. For spiral k-space sampling, the PSF was calculated using the inverse Fourier transform of the modulation transfer function in Matlab (R2013a, The Mathworks Inc., Massachusetts), which incorporates the effects of sampling, truncation, and $T_2^*$ (17) as described in detail in Appendix A (see Calculation of Spiral PSF) (18) and (19). For comparison, the PSF for conventional Cartesian k-space sampling was calculated analytically along the frequency encoding direction as described in Appendix B (Calculation of Cartesian PSF). Spatial resolution was approximated using the FWHM of the calculated PSF function as a function of $T_{read}/T_2^*$ to demonstrate the effects of $T_2^*$ blurring on resolution. SNR was approximated from the corresponding PSF amplitude, $A(T_{read}/T_2^*)$ using the following expression:

$$SNR_N \propto A(T_{read}/T_2^*) \times \sqrt{N/BW} \times \sin(\alpha)$$ \hspace{1cm} [3.1]

where, $\alpha$ is the flip angle, $N$ is number of points in k-space and $BW$ is the read-out bandwidth. The $\sin(\alpha)$ term accounts for the difference in the flip angle used for Cartesian and spiral, depending on the number of RF pulses and/or views. The $N/BW$ term accounts for the effect of bandwidth on image noise. The $BW$ was varied to change the read-out time, $T_{read} = N/BW$ for a fixed $N$. The effect of varying the $BW$ on the noise was proportional to the square root of the $BW$. 
Figure 3.1: Pulse sequence timing diagram for the spiral acquisition of a single echo. A hard pulse is followed by read-out gradients ($G_x$ and $G_y$) during which data acquisition occurs. The echo time $TE = \Delta TE \times m$ is defined as the echo time spacing multiplied by the number of echoes. Following the data acquisition, transverse magnetization is spoiled with gradients prior to the next acquisition at an echo time shifted according to the IDEAL approach.

Eqn. [3.1] was used for optimization of SNR with respect to $T_{read}$ for the spiral k-space sampling. It was also used to predict the SNR of the single-shot spiral k-space sampling in comparison to Cartesian. For fair comparison, the $A(T_{read}/T_2^*)$ were calculated using the same nominal resolution for both the Cartesian and spiral k-space sampling. Thus, for the Cartesian sampling, a matrix size of $32 \times 32$, $FOV$ of 10 cm and $BW = 16$ kHz was chosen similar to that used previously for in vivo Hp $^{129}$Xe rat lung imaging (11). A constant flip angle of 15° was used for the corresponding matrix size of the Cartesian sampling scheme to achieve the maximum SNR (20,21). A flip angle of 90° was used for the spiral SNR
calculation for thermally-polarized gas phantom experiments (as described in the Phantom MRI Experiments section below).

3.2.2 Spiral Gradient Waveform Design

Spiral k-space trajectory and gradient waveforms \((G_x(t), G_y(t))\) required to achieve a given \(T_{\text{read}}\) were calculated using an algorithm similar to that described by Lee et al. (22). A surface plot of \(T_{\text{read}}\) as a function of both maximum gradient amplitude \((G_{\text{max}})\) and gradient slew rate \((SR)\) was generated for a \(FOV = 70\) mm and in plane voxel size of 3 mm corresponding to typical rat imaging parameters. For the calculation, \(G_{\text{max}}\) was varied between 0 and 500 mT/m in steps of 20 mT/m and \(SR\) was varied between 0 to 3000 T/m/s in steps of 100 T/m/s, corresponding to the capabilities of the high performance gradient coil used in the study (23). This was also compared to a trajectory optimized for the performance of the conventional clinical gradient system \((G_{\text{max}} = 50\) mT/m, \(SR = 200\) mT/m) to investigate the feasibility of human imaging.

3.2.3 Optimization of Echo-Time spacing

The optimal echo times, \(TE\), for IDEAL were determined theoretically using the effective number of signal averages (NSA) approach by including the short \(T_2^*\) effect as previously described for C-13 metabolites (16,24). This approach accounts for the noise performance as a function of echo time spacing, \(\Delta TE\), and chemical shift (24). From these analyses, the optimal \(TE\) for each echo image, \(m\), were calculated with \(TE = \Delta TE \times m\) for the corresponding \(\Delta TE\) from Figure 3.4. \(\Delta TE\) was 50 us for 3-point IDEAL \((m = 1, 2, 3)\) gas-PT imaging and 340 us for 4-point IDEAL \((m = 1, 2, 3, 4)\) gas-PT-RBC imaging based on
the known chemical shift differences \((\Delta f_{G-PT} = 6950 \text{ Hz}, \Delta f_{PT-RBC} = 525 \text{ Hz})\) at a field strength of 3 T). This choice of \(TE\) was based on measured \(T_2^*\) values of gas and dissolved phases as described in the results section.

### 3.2.4 Calibration of Flip Angles

In delivering the RF excitation to each phase (e.g., gas versus dissolved phase), it was desirable to apply a sufficiently large flip angle (e.g. 10-20 degrees) to the gas phase to achieve reasonable gas phase image SNR without exhausting the gas phase pool, while at the same time applying a large flip angle (i.e. 90°) to the dissolved phases. This provided the additional benefit of improved dissolved phase SNR via replenishment from the gas phase between each image acquisition. The dependence of \(^3\text{He}\) gas image SNR on flip angle for different varying RF pulse number (i.e., phase-encoding step) has been discussed by Miller et al. (20). Using a similar approach, gas signal strength, \(S_G\), and PT signal strength, \(S_{PT}\), equations were derived from the analytic solutions to the Bloch equations including the effect of exchange of magnetization between the PT and gas compartments (Appendix). The gas and PT signal dependence following the \(n^{th}\) RF pulse are respectively given by the following two equations:

\[
S_G = M_G \left[ \cos(a_G) \right]^{n-1} \sin(a_G) \tag{3.2}
\]

\[
S_{PT} = M_{PT} \left[ M(TR) \left[ \cos(a_G) \right]^{n-1} \sin(a_{PT}) + \left[ \cos(a_{PT}) \right]^{n-1} \sin(a_{PT}) \right] \tag{3.3}
\]

where \(a_G\) and \(a_{PT}\) are the flip angles applied to the gas phase and PT phase respectively. 
\(M_G\) is a scaling factor based on the gas phase image, which accounts for differences in
polarization and ventilation conditions between scans, $M(TR)$ represents the percentage replenishment of dissolved phase magnetization by the gas phase as a function of the time between RF pulses (TR). $M(TR)$ was calculated by fitting the Mansson gas exchange model to CSSR experimental data previously published by Fox et al. (7). $M(TR)$ was approximately 0.5 for the TR used in this study (30 ms) (see Appendix [C.6] for more details). $M_{PT}$ is a scaling factor based on the dissolved phase image, which accounts for solubility, tissue density and $M_G$. $T_1$ relaxation of $^{129}$Xe in both the gas and dissolved phases (25) is ignored in Eqn. [3.2] as it was assumed to be much longer than the total imaging time. Eqn. [3.3] shows that $S_{PT}$ is reduced with the use of larger $\alpha_G$ and increased with the use of larger $\alpha_{PT}$ and larger $TR$, the latter due to replenishment. The SNR of the gas image, $SNR_G$, and SNR of the PT image, $SNR_{PT}$, were optimized as a function of $\alpha_G$ and $\alpha_{PT}$, including the SNR improvement due to the number of averages.

### 3.3 Methods

All images were acquired using a 3 Tesla MRI system (MR750, GEHC, Wisconsin, USA), a custom-built transmit-only/receive-only birdcage coil (26) and high performance insertable gradient system (23). Images were reconstructed in Matlab as described in Schulte et al. (15,16).

#### 3.3.1 Phantom MRI Experiments

A thermally-polarized phantom, containing 200 ml of $^{129}$Xe gas (80% enriched) and 200 ml $O_2$ gas (to reduce $T_1$ relaxation to ~1 s) at a pressure of 8 atmosphere at room temperature, was used to experimentally confirm the theoretical dependence of spatial
resolution on $T_{\text{read}}/T_2^*$. $T_2^*$ for the thermal phantom was measured to be 28±1 ms. 2D projection images of the thermally-polarized phantom in the axial view were obtained with the following eight values of $T_{\text{read}}/T_2^* = 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0$ using a 3-point IDEAL spiral approach with the following parameters: $TR = 1.0$ s, $\Delta TE = 50$ us, $FOV = 10$ cm $\times$ 10 cm, with a real pixel size of 32, $BW = 62.5$ kHz, number of averages = 8, flip angle = 90°. $T_{\text{read}}$ was adjusted experimentally by changing the SR of the spiral gradient waveforms ($G_x(t)$, $G_y(t)$). Experimentally measured resolution including $T_2^*$ blurring were calculated from thermal phantom images by the FWHM of the PSF as described in Appendix (19).

3.3.2 *In vivo* MRI experiments

All procedures followed animal use protocols approved by Western University’s Animal Use Subcommittee and were consistent with the guidelines written by the Canadian Council on Animal Care (CCAC) (27). Five Sprague Dawley rats (390±7 g, Charles River Laboratories, Saint-Constant, Canada) were imaged in the supine position following three breaths of Hp $^{129}$Xe using a MRI-compatible mechanical ventilator (GE Healthcare, Malmo, Sweden) for delivery of Hp $^{129}$Xe to the lung. Hp $^{129}$Xe gas (approximately 80% enriched with 10% polarization) was obtained from a commercial xenon polarizer system (Polarean 9800, Durham, NH) yielding polarizations of ~10%. *In vivo* images were acquired with a three-point IDEAL approach for imaging of gas-PT and four-point IDEAL approach for imaging of gas-PT-RBC using the spiral pulse sequence as shown in Figure 3.1. A set of IDEAL coronal projection gas-PT images were obtained using a single-shot spiral three-point IDEAL technique and three RF pulses ($n = 3$) and echo images with echo-
times of 50 us × \( m \), \( m = 1, 2, 3 \), resulting in a total scan time of 90 ms for \( TR = 30 \) ms). IDEAL images of gas, PT and RBC were obtained with a single-shot four-point IDEAL technique using four RF pulses \( n = 4 \) and four echo images with echo-times of 340 us × \( m \), \( m = 1, 2, 3, 4 \), resulting in a total scan time of 120 ms for \( TR = 30 \) ms). Other pulse sequence parameters included: \( FOV = 70 \) mm, \( BW = 125 \) kHz, a nominal matrix resolution of 22, number of points in the k-space equal to 660 and spiral trajectories consisting of 11 turns.

To validate Eqns. [3.2] and [3.3], multiple IDEAL gas and PT images (21 pairs) were acquired using 63 RF pulses \( n = 63 \) with TR fixed at 30 ms within a total scan time of 1.92 s during a single breath-hold interval. A mask that corresponded to the left lung of each rat was created from the first gas phase image and applied to the all proceeding tissue images. \( S_G(j) \) and \( S_{PT} (j) \) were determined by the mean values of regions of interest placed on the gas and PT images as a function of image number \( j = 1, \ldots, n/3 = 21 \). For comparison between rats, the multiple \( H_p^{129}\)Xe IDEAL images were normalizing by \( S_{PT}(j = 1:21) \) with \( S_G(j = 1) \), since the absolute values of \( S_{PT}(j) \) depend on experiment conditions (e.g., polarization, ventilation) through the scaling factor \( M_{PT} \) (as explained in Eqn. [3.3]). The mean of \( S_G(j) \) and normalized \( S_{PT}(j) \) and their standard deviations were also calculated to investigate the reproducibility of the IDEAL images.

Ten sets of coronal IDEAL gas, PT, RBC images \( j = 1:10, n = 40 \) were obtained using a four-point IDEAL approach within a total scan time of approximately 4 s in a single-breath hold (with a \( TR \) of 100 ms) in the rat lung. IDEAL gas, PT, RBC images were reconstructed in Matlab. The ten reconstructed gas, PT, and RBC images were then averaged in Matlab to generate a final IDEAL gas, PT, and RBC images. Similarly, representative average
coronal *in vivo* IDEAL gas and PT images were obtained using three-point spiral IDEAL 
\((j = 1:10, n = 30)\) within a total scan time of approximately 3 s.

### 3.3.3 RF pulse calibration

To achieve the optimum \(\alpha_{PT}\) and \(\alpha_G\), a hard pulse was centered at the PT resonance in order to maximize the signal from the tissue phase. The gas signal was excited using the side band of a rectangular (i.e., hard) RF pulse (28). When \(\alpha_{PT} = 90^\circ\), the side band excitation, \(\alpha_G\) was much higher than the optimal. Therefore, the transmit power was decreased so that \(\alpha_{PT} = 75^\circ\) resulting in \(\alpha_G = 18^\circ\). These flip angles were achieved using a pulse width of 0.18 ms to minimize the side band excitation of the gas phase and \(T_2^*\) amplitude loss. The actual \(\alpha_{PT}\) and \(\alpha_G\) values were calibrated by observing the free induction decay (FID) signal following the RF pulse. For this purpose, the hard pulse was centered on the gas phase and required power was adjusted to generate a 90° excitation. Based on the power used to generate a 90° excitation, the actual \(\alpha_{PT}\) and \(\alpha_G\) were calibrated to be approximately 75° and 18° respectively. For comparison, \(\alpha_{PT}\) and \(\alpha_G\) were calculated from multiple three-point IDEAL images *in vivo* using Eqns. [3.2] and [3.3].

As a proof-of-concept demonstration of the ability of the spiral IDEAL approach to measure gas exchange, four-point IDEAL images of gas, PT, RBC were obtained at five different TR values \((30, 50, 100, 200, 300 \text{ ms})\) using five separate breath-holds respectively. Signal from PT images, \(S_{PT}(j = 2:5, TR)\) and RBC images \(S_{RBC}(j = 2:5, TR)\) were normalized with \(S_G(j = 2:5, TR)\). Dissolved phase signal from the first image \((j = 1)\) was ignored since it was not weighted by the TR. \(S_{PT}(TR)\) and \(S_{RBC}(TR)\) were calculated
for the left and right lungs of five healthy rats and averaged to provide mean signals as a function of increasing $TR$.

3.4 Results

Theoretical and experimental dissolved phase spatial resolution from the PSF analysis and the thermal phantom for the eight $T_{\text{read}}/T_2^*$ values respectively are shown in Figure 3.2 (a). As expected, measured blurring was diminished as read-out time was reduced and plateaued for $T_{\text{read}} \sim 1.5 \times T_2^*$ in good agreement with theory. Theoretical SNR from Eqn. [3.1] is shown in Figure 3.2 (b) over a similar range of $T_{\text{read}}/T_2^*$ as in Figure 3.2 (a). The maximum SNR was obtained for $T_{\text{read}}$ values between approximately $T_2^*$ and $2 \times T_2^*$. The optimum value of $T_{\text{read}}/T_2^*$ was chosen to be 1.8 to achieve the maximum SNR for the dissolved phase image while maintaining acceptable spatial resolution. $T_{\text{read}}/T_2^* = 1.8$ corresponded to a 33% decrease in resolution from the nominal pixel size. In particular, for a nominal pixel size of 3.1 mm, the spatial resolution was expected to be 4.1 mm due to $T_2^*$ blurring. The theoretical maximum Cartesian SNR$_N$ was found to occur at $T_{\text{read}}/T_2^* = 0.9$ similar to that previously reported (18). Compared to the Cartesian data acquisition with the use of flip angle of 15°, the single-shot spiral showed a theoretical improvement in the SNR of 3.3 fold. It should be noted that this SNR improvement was expected to be less than 3.3-fold due to the dissolved signal replenishment between phase encoding steps (i.e. TR) of the Cartesian k-space sampling. The Cartesian acquisition with a TR time of 4 ms, would allow replenishment of dissolved phase signal of approximately 10 % (see Appendix Eqn. [C.5]). Therefore, considering the replenishment of the dissolved phase signal, the SNR improvement would be expected to be approximately 3-fold.
Figure 3.2: Theoretical spatial resolution (a) and normalized \( SNR_N \) (b) as a function of \( \frac{T_{\text{read}}}{T_2^*} \) spiral k-space trajectories. Experimental results are also shown for the measurement of FWHM in (a).

Figure 3.3 (a) shows the gradient amplitude and slew rate required to achieve a given \( T_{\text{read}} \) as well as the resulting spatial resolution for a given \( T_{\text{read}} \) at the optimal gradient amplitude and slew rate for in vivo imaging. For \( T_{\text{read}} \) of 2.8 ms, corresponding to \( \frac{T_{\text{read}}}{T_2^*} = 1.8 \) and the measured \( T_2^* \) values of 1.6±0.1 ms and 1.6±0.2 ms for PT and RBC respectively, the required \( G_{\text{max}} \) and \( SR \) were estimated from Figure 3.3 (a) to be 300 mT/m and 3000 T/m/s for fixed pixel size of 3 mm. It was noted that for \( T_{\text{read}} \) values less than 3.4 ms (\( G_{\text{max}} = 200 \) mT/m and \( SR = 3000 \) T/m/s), the gradient requirements did not change significantly.

Repeating this process with the high-performance gradient parameters fixed at the optimum values \( G_{\text{max}} = 200 \) mT/m, \( SR = 3000 \) T/m/s, and \( FOV = 70 \) mm, the relationship between spatial resolution and \( T_{\text{read}} \) is shown in Figure 3.3 (b) (circle symbols). A read-out time of
2.8 ms was obtained for the nominal pixel size of 3.3 mm. The corresponding relationship for the conventional clinical gradient coil configuration with parameters fixed at the maximum values ($G_{\text{max}} = 50 \text{ mT/m}, SR = 200 \text{ T/m/s}$ and $FOV = 40 \text{ cm}$) for a six interleaved spiral acquisition is also shown in Figure 3.3 (b) (square symbols).

Figure 3.3: (a) A surface plot of $T_{\text{read}}$ as a function of both maximum gradient amplitude ($G_{\text{max}}$) and gradient slew rate (SR) generated for a $FOV = 70 \text{ mm}$ and resolution of 3 mm. In (b) the read-out time as a function of resolution is shown for two gradient performance scenarios: (i) the high performance gradient for rat imaging ($G_{\text{max}} = 300 \text{ mT/m}, SR = 3000 \text{ T/m/s}, FOV = 70 \text{ mm}$) and (ii) the clinical gradient performance for human imaging ($G_{\text{max}} = 50 \text{ mT/m}, SR = 200 \text{ T/m/s}, FOV = 400 \text{ mm}$ and six interleaves).

Figure 3.4 (a) shows NSA values as a function of $\Delta TE$ for 3-point and 4-point IDEAL approaches. Ignoring the effects of $T_2^*$ amplitude loss, the maximum theoretical NSA was three and four for three-point and four-point IDEAL respectively. The optimum $\Delta TE$ values were found to be 50 us for the three-point IDEAL gas-PT imaging in Figure 3.4 (a). The corresponding NSA was found to be 2.8 due to $T_2^*$ amplitude loss. Thus, it is reasonable to
expect that the $T_2^*$ amplitude loss was negligible for the three-point IDEAL approach. For four-point IDEAL, the optimum $\Delta T E$ that maximized NSA of all components and allowed separation of PT and RBC signals was found to be relatively long (340 us) in Figure 3.4 (b). These relatively long echo times are required due to the relatively smaller chemical shift difference between PT and RBC ($\Delta f_{PT-RBC} = 525$ Hz) leading to drop in the SNR of a factor of approximately two compared to the scheme without the $T_2^*$ amplitude loss. Conceptually, the SNR performance of PT images from three-point IDEAL (Figure 3.4 (a)) is better than four-point IDEAL by a factor of 1.4 owing to relatively short echo times.

![Figure 3.4](image)

**Figure 3.4:** The NSA as a function of $\Delta T E$ for (a) 3-point IDEAL and (b) 4-point IDEAL.

Figure 3.5 (a) shows the measured mean $S_G(j)$ including standard deviation from the five healthy rats as well as the best fit to the data based on Eqn. [3.2] yielding estimated flip angles of $18^\circ \pm 0.4^\circ$ and $R^2$ value of 0.998. Figure 3.5 (b) shows the PT signal (i.e. normalized $S_{PT}(j)$) as a function of pulse number and the fit of function given by Eqn. [3.3]
including gas exchange, yielding an $R^2$ value of 0.997. In the first PT image ($j=1$), the percentage gas exchange percentage rate ($M(TR)$) was determined to be 0.1 (7) since the first PT image ($TR = 30$ ms) is not weighted by gas exchange. $M(TR) = 0.5$ was used for fitting the remaining 20 images. The flip angle delivered to the PT phase was determined to be $75^\circ \pm 2^\circ$ using Eqn. [3.3], in good agreement with the calibrated value. Normalized $S_G$ and $S_{PT}$ as a function of pulse number were highly reproducible for the five healthy rats with the maximum standard deviation value of 2.6%.

![Graph](image)

Figure 3.5: Experimentally measured gas (a) and pulmonary tissue (b) signals as a function of image number. The solid line shows the theoretical fit to the data from Eqns. [3.2] and [3.3].
Figure 3.6 shows the theoretical SNR performance of the gas and PT images as a function of both $\alpha_{PT}$ and $\alpha_G$, after averaging 10 sets of IDEAL images. Figure 3.6 (a) shows that the use of $\alpha_G$ between 15° and 20° provides sufficiently high SNR for the gas phase image, but results in a reduction in the SNR of the tissue phase image in Figure 3.6 (b). This is because the Hp$^{129}$Xe gas in the alveolar air space acts as a reservoir to replenish the dissolved phase image signal in the time between averages. Notably, $\alpha_G = 18°$ was used to obtain a sufficiently high SNR for the gas phase of approximately 23 as shown in Figure 3.6 (a) and $\text{SNR}_{PT}$ was approximately 5.8 as shown in Figure 3.6 (b). The theoretical dependences of $\text{SNR}_{PT}$ on $\alpha_{PT}$ for $TR = 30\text{ms}$ (circles) and $TR = 100\text{ms}$ (square) are shown in Figure 3.6 (c), for $\alpha_G$ fixed at 18°. $\text{SNR}_{PT}$ increases with $\alpha_{PT}$ but displays no significant increase beyond $\alpha_{PT}=75°$ which corresponds to the use of 99% of available PT signal with a $TR = 30\text{ms}$. Furthermore, Figure 3.6 (b) shows an SNR improvement of a factor of approximately 2 for $TR = 100\text{ms}$ due to the increased time for gas exchange.
Figure 3.6: Theoretical SNR\textsubscript{G} (a) and SNR\textsubscript{PT} (b) as a function of $\alpha\textsubscript{G}$ for $\alpha\textsubscript{PT}$ fixed at 90\textdegree. SNR\textsubscript{PT} as a function of $\alpha\textsubscript{PT}$ (c) for the $\alpha\textsubscript{G}$ fixed at 18\textdegree is also shown for TR = 30 ms (circles) and TR = 100 ms (squares).

Figure 3.7 shows representative coronal \textit{in vivo} IDEAL gas and PT images obtained using three-point spiral IDEAL with a $TR$ of 100 ms in the rat lung within a total scan time of 3 s. The SNR of the gas and PT images were 22.8 and 10.2 respectively and satisfactory given the polarization of ~10\%. Figure 3.8 shows representative coronal \textit{in vivo} IDEAL gas, PT, and RBC images obtained using four-point spiral IDEAL with a $TR$ of 100 ms in the rat lung within a total scan time of 4 s. The SNR of the gas image (SNR\textsubscript{Gas} = 23.0) was similar to the gas image from three-point IDEAL. The SNR of the PT image (SNR\textsubscript{PT} = 8.6) was higher than the RBC image (SNR\textsubscript{RBC} = 4.7) due to the tissue-blood partition based on the different of xenon solubility and density in the dissolved compartments (5).
Figure 3.7: Coronal gas (a) and PT (b) images of a representative rat lung obtained using the three-point spiral IDEAL approach.

Figure 3.8: Coronal gas (a), PT (b), and RBC (c) images obtained using the four-point spiral IDEAL approach.

Figure 3.9 shows mean PT (a) and RBC signals (b) from the left and right lungs as a function of TR. A strong correlation was observed in xenon gas exchange between the left and right lungs (Pearson’s correlation statistics: $r = 0.98$ and $P = 0.0023$ from PT image; $r = 0.997$ and $P = 0.0002$ from RBC image) showing that repeated acquisitions in five healthy rats were regionally consistent between the left and right lungs.
Figure 3.9: Mean signals measured from all rats for PT (a) and RBC images (b) as a function of TR values: 30, 50, 100, 200, and 300 ms.

3.5 Discussion

In this work, a single-shot spiral k-space encoding strategy was applied to IDEAL imaging for separation of hyperpolarized $^{129}$Xe signals originating from gas, PT and RBC compartments in the rat lung within a breath-hold interval (4 s). To achieve the necessary spatial resolution (3.3 mm) with minimal $T_2^*$ blurring, a relatively short spiral read-out time, $T_{\text{read}}$, (2.8 ms) on the order of $T_2^*$ was found to be important and was facilitated in this study using a high performance insertable gradient system to achieve the required $G_{\text{max}}$ and $SR$. Although the theoretical PSF is an approximation, good agreement was shown with experimental spatial resolution data obtained with thermally-polarized $^{129}$Xe phantom experiments. The single-shot spiral read-out also permitted the use of fewer, larger flip angle pulses to improve SNR compared to Cartesian approaches. Based on the theoretical
PSF analysis, spiral IDEAL approaches that utilize the entire dissolved phase magnetization should give an SNR improvement of a factor of approximately 3.3 for $\alpha_G = 90^\circ$ compared to Cartesian approaches with similar spatial resolution ($\alpha_G = 15^\circ$).

For *in vivo* imaging, SNR$_{PT}$ depends significantly on the choice of both $\alpha_G$ with $\alpha_{PT}$ and polarization. The optimum value for $\alpha_G$ was found to be between $15^\circ$ to $20^\circ$ for the use of 10% polarization. Although the optimum value for $\alpha_{PT}$ was found to be $90^\circ$, a $75^\circ$ pulse was used to reduce the side band excitation of the gas phase. In future, the use of spectrally-selective RF pulses (28) that provide $\alpha_G = 3^\circ$ and $\alpha_{PT} = 90^\circ$ with higher polarization on the order of 30% would significantly improve the SNR of the dissolved phases. Less consumption of Hp $^{129}$Xe gas magnetization with $\alpha_G = 3^\circ$ would be sufficient to generate an SNR$_G$ value of approximately 23 due to the three-fold improvement of polarization Figure 3.6 (a)). The use of $\alpha_G = 3^\circ$ and a 3-fold improvement in polarization would also result in an improvement of SNR$_{PT}$ from $5.8$ (for $\alpha_G = 18^\circ$) to $10.3 \times 3 = 30.9$ (for $\alpha_G = 3^\circ$) based on Figure 3.6 (b).

Obtaining gas images in concert with dissolved phase images is expected to be helpful for identification of corresponding ventilation defects. Gas images from both 3-point and 4-point IDEAL match well from the same rat. The reproducibility was significantly high between five healthy rats. Gas phase images are also useful for optimization of flip angle (Eqn.[3.2]) from multiple IDEAL images and for normalization purposes (e.g. to account for polarization differences) for quantitative dynamic Hp $^{129}$Xe gas exchange imaging.

In comparison to the three-point IDEAL PT image, four-point IDEAL PT has lower SNR (by a factor of 1.2) and temporal resolution performance (by a factor of 1.3) owing to the
use of longer echo time spacing and more RF pulses and fewer images in the reconstruction.

In principle, four-point IDEAL has the advantage of separating the PT compartment from the RBC compartment. RBC signal intensity was strong in the region of left atrium and ventricle due to the relatively high density of blood compared to other parts of the thorax.

The effect of short $T_2^*$ and differences in $T_2^*$ between the dissolved phases of Hp $^{129}$Xe (~1.6 ms) and gas phase of Hp $^{129}$Xe (~6.0 ms) requires further discussion. On the IDEAL acquisition side, the appropriate $T_2^*$ values for the different species is explicitly accounted for in the NSA plot (Figure 3.4) when choosing the echo times which gives the optimum separation of the IDEAL signal between the gas and dissolved phases. The relatively large chemical shift between the gas phase and the dissolved phases (~ 200 ppm) means the echo times can be quite short (~ 50 us), thereby reducing $T_2^*$ amplitude losses, since a separate excitation is used for each IDEAL echo signal. However, the longer echo time spacing (~ 340 us) required for the 4-point IDEAL approach results in a more significant drop in SNR due to $T_2^*$ amplitude losses compared to the 3-point IDEAL approach. To address the smaller chemical shift between the dissolved phases and short echo time, an unequally spaced short echo approach could be used in future (29). On the reconstruction side, the respective $T_2^*$ values are not explicitly considered in this work which could affect the separation of the PT and RBC images from the gas phase images (30).

The use of the high gradient amplitude and slew rate, timing delays in the gradient waveforms, and eddy currents associated with the high performance gradient insert used in this study may lead to characteristic spiral image artifacts, including streaking and ghosting (31,32). The accuracy of the gradient encoding fields ($G_X$ and $G_Y$ in Figure 3.1), and corresponding geometric distortions and image quality have not been explicitly evaluated.
in this study. However, hyperpolarized phantom experiments showed good agreement with the geometry of the phantom and images suggesting that the k-space distortions were negligible in our previous studies (33). Image distortion and ghosting were likely minimized in the present study owing to the use of a relatively larger field-of-view. While there was some streaking artifact visible, this was only apparent at the edge of the gas phase images since the gas signal was much higher than the signal from dissolved phases. Some spiral artifacts were visible in the RBC images, likely due to the low SNR owing to the low polarization.

In this work, we have demonstrated the feasibility of measuring regional gas exchange curves for the left and right rat lungs separately. The ability to image $^{129}$Xe gas exchange at various TR values (e.g. from 30 ms to 300 ms) in separate breath-holds may permit quantitative measurement of gas exchange similar to that obtained using the conventional CSSR technique but with the advantage of regional information. In particular, the slopes and intercepts of gas exchange curves (8,34-36) such as shown in Figure 3.9, allow mapping of alterations in lung structure and function associated with pulmonary disease. In particular, four-point IDEAL may provide quantitative measures of both the vascular damage and tissue thickening (i.e. pneumonitis) corresponding to the early stages of RILI following thoracic irradiation (7). By choosing smaller regions-of-interest instead of the whole left and right lungs, the gas exchange curves could be mapped using higher polarization (e.g. 50%) in future.

For \textit{in vivo} imaging of rat lungs with the single-shot spiral IDEAL approach, high \textit{SR} (3000 T/m/s) and $G_{\text{max}}$ (300 mT/m) were used to increase the spatial resolution within an optimum read-out time. IDEAL spiral imaging could be potentially translatable to human imaging
with the use of an interleaved spiral approach. Conceptually, the imaging of Hp\textsuperscript{129}Xe gas exchange at TR values (e.g. 15, 30, 50, 100, 250 ms) in a single breath-hold of 10.7 s should be possible using six interleaved spirals. Figure 3.3 (b) shows the relationship between the spatial resolution and read-out time with the use of clinical gradient for six RF pulses and interleaved spiral with FOV of 40 cm. Furthermore, at lower field strength (e.g. 1.5 T) the spatial resolution could be improved, even at longer read-out times due to the increase in $T_2^*$ (12). In future, 2D imaging can also be extended to multi-slice imaging using parallel imaging approaches (37) and a short duration slice-selective pulse (28). Although, five TR values would be achievable for 2D human imaging, the total scan time required for obtaining the gas exchange curves would be challenging for the case of multi-slice imaging. However, the total scan time for multi-slice human imaging might be reduced by using lower TR values to obtain the gas exchange curves. Furthermore, 2D imaging can be extended to an isotropic 3D MR imaging using an additional phase encoding gradient, undersampling approach (38) and cylindrical (39) or radial (40) k-space sampling techniques.

3.6 Conclusion

Single-shot spiral IDEAL imaging for separation of Hp\textsuperscript{129}Xe gas and dissolved phase images is feasible in the rat lung. The spiral approach offers the benefits of fewer, higher flip angle RF pulses to increase the signal from the dissolved phases of Hp\textsuperscript{129}Xe. An SNR improvement of approximately 3 times is expected from spiral IDEAL compared to the conventional Cartesian k-space sampling IDEAL approach. The read-out time of approximately 1.8 time $T_2^*$ was found to be optimal based on a point spread function
analysis to minimize $T_2^*$ blurring and maximize the SNR. The rapid acquisition times of spiral IDEAL permits multiple gas/PT and gas/PT/RBC images to be acquired within a breath-hold time of a few seconds. This allows flip angle calibration and optimization and permits imaging of exchange between the gas, PT and RBC compartments for regional characterization of lung microstructure and pulmonary and vascular function. Although, the imaging approach use here was optimized for rat imaging, it could be translated for human imaging using an interleaved spiral approach in future.
3.7 Reference


36. Doganay O, Fox M, Santyr GE. Measurement of Pulmonary Perfusion and Gas Exchange using Hyperpolarized 129Xe in a Rodent Model of Radiation-Induced


Chapter 4: Early Detection of RILI using $^{129}\text{Xe}$ Gas Exchange

This chapter is adapted from the submitted paper: Doganay O, Stirrat E T, McKenzie C, Schulte RF, Santyr GE. Quantification of Regional Gas Exchange Changes using Hyperpolarized $^{129}\text{Xe}$ MRI in a Rat Model of Early Stage Radiation-induced Lung Injury. Medical Physics, 2015 (in revision).

4.1 Introduction

Radiation therapy is a primary treatment method for thoracic malignancies (2). However, the lung is a very radiosensitive organ (3-7), and Radiation-induced Lung Injury (RILI) is an important factor limiting thoracic radiotherapy. The incidence for moderate and severe RILI in patients after radiation therapy is 50% and 5-35% respectively (8,9). RILI generally progresses in two phases: (i) early-phase, radiation pneumonitis (RP) and (ii) late-phase, pulmonary fibrosis. RP is acute and occurs within 12 weeks after radiation therapy, mainly comprising of inflammation with symptoms of dry cough and dyspnea. Pulmonary fibrosis is irreversible and can develop as early as three months after radiation therapy. Severe pulmonary fibrosis may lead to organ failure and even death (10,11). Detection of early-stage RILI (i.e. RP) can improve outcomes by reducing normal tissue injury and late-stage effects (8,12) through adjustments in therapy plan (e.g. radiation fractionation) or use of pharmaceuticals, including radioprotectors and anti-inflammatory drugs (13-15). Pulmonary function tests (PFTs), such as spirometry, are often used for monitoring RILI but provide measures of whole-lung function only, without regional information (16). Furthermore, PFT measurements depend strongly on patient positioning and effort,
obstructive disease and are sensitive to hyperfractionated therapy (total dose > 25 Gy) (17-19).

To overcome some of the limitations of PFTs and to provide regional detection of RILI, imaging techniques have been investigated. The most commonly used imaging approach for detection of RILI involves computed tomography (CT). CT is sensitive to increases in lung density (ie. microanatomy) due to RILI (16,20-22), but these increases are typically detected during the late irreversible phase of fibrosis. Furthermore, CT lung density measurements are variable and depend on the degree of inspiration and patient posture (23). CT perfusion imaging with intravenous injection of contrast agents has been shown to be sensitive to vessel damage associated with RILI as soon as 4 weeks following irradiation (6,24), but does not provide ventilation information. Single-photon emission computed tomography (SPECT) imaging is sensitive to radiation-induced reductions in regional perfusion and ventilation (25-27). Ventilation SPECT of the lungs can be based on administration of radioactive gas (e.g. $^{81m}$Kr and $^{133}$Xe), particulate aerosols labeled with radioactive compounds (i.e. $^{99m}$Tc-DTPA) and ultrafine graphite particles labeled with $^{99m}$Tc. Perfusion SPECT of lungs typically uses intravenous administration of macroaggregates of albumin labeled with $^{99m}$Tc (28). Lung perfusion SPECT has been shown to be more sensitive than CT to early-phase RILI (6 weeks post-irradiation), capable of detecting approximately 20% perfusion reduction in 3D dose distributions over the range 15 to 30 Gy (29). However, quantitative SPECT images have inherently poor spatial resolution and require corrections for image-degrading factors (i.e., photon scatter and attenuation), leading to underestimation of ventilation-to-perfusion ratios (28,30,31). Positron Emission Tomography (PET) using $^{18}$F-FDG offers higher spatial resolution than
SPECT and has been shown to detect metabolic changes associated with RILI within 1-3 months after radiotherapy (32). Hybrid nuclear imaging approaches such as SPECT/CT and PET/CT further improve diagnostic accuracy and sensitivity to RILI by reducing registration problems and improving corrections for image degradation (33); however, CT, SPECT and PET are all limited by radiation exposure (~3 mSV) and SPECT and PET are additionally limited by long scan times (10-30 minutes).

Although conventional proton Magnetic Resonance Imaging (MRI) of the lungs is limited by low signal, the addition of hyperpolarized (Hp) nuclei including: $^{129}$Xe, $^3$He and $^{13}$C can significantly improve image quality and provide functional information (34). Hyperpolarization permits increases in signal of up to 105 from these nuclei, allowing rapid and high spatial resolution MRI of the effects of radiation on the lung. Advances in Dynamic Nuclear Polarization of $^{13}$C-pyruvate (35) have enabled identification of early (days – weeks) metabolic changes in lung lactate-to-pyruvate ratios in a rat model of whole thorax irradiation (36) and right thorax irradiation which correlates well with increases in inflammatory cells (ie. macrophages) as confirmed by histology (37). The first case study on rat model has shown that there is a significant increase in ADC maps of Hp $^3$He associated with three months-post irradiation dose of 40 Gy (38). Hp $^3$He gas MRI has been used to investigate human subjects undergoing radiotherapy suggesting that assessment of ADC and ventilated defect percent can reveal late-phase (i.e. 6-8 months) RILI (39). Another Hp $^3$He study showed the feasibility of detecting ventilation changes in image intensity due to late RP (40). Recently, Hp $^{129}$Xe gas MRI morphometry has been shown to be sensitive to microstructural changes in lung parenchyma in rat model of RILI after two weeks irradiation (41).
Due to the relatively high solubility of xenon in biological tissues, Hp $^{129}$Xe has been shown to be a promising approach for quantification of gas exchange between the alveolar airspace and the dissolved phases, principally the pulmonary tissue (PT) and red blood (RBC) compartments. Dissolved phase $^{129}$Xe exhibits a large chemical shift in PT (197 ppm) and RBCs (218 ppm) relative to the gas phase signal (42). Furthermore, the Chemical Shift Saturation Recovery (CSSR) technique has been used for assessment of gas exchange curves in the whole lung (43). Fitting the CSSR data to gas exchange models permits quantification of lung function (i.e. gas exchange time constants and perfusion) as well as microstructural (i.e. alveolar wall thickness and relative blood volume) changes (1,44-46). Recently, whole lung CSSR has been shown to be sensitive to RILI at two weeks following whole thorax $^{60}$Co irradiation in a rat model (47).

The above spectroscopic CSSR approaches, while quantitative, do not provide regional information, which is expected to reflect regional heterogeneity in RILI, particularly in the case of conformal irradiation approaches. MRI of Hp $^{129}$Xe in the dissolved phase can be accomplished using multiple echo time MRI approaches, which exploits the phase evolution of the MRI echo signal as a function of echo time to reconstruct images corresponding to each dissolved compartment. In particular, IDEAL (Iterative Decomposition of water and fat with Echo Asymmetry and Least-square estimation) MRI has been shown to be efficient for obtaining gas and dissolved phase $^{129}$Xe images in a breath-hold duration (48-51).

Recently, we have shown that an IDEAL approach utilizing single-shot spiral read-outs can provide approximately a factor of ten improvement in speed, allowing measurement of regional gas exchange curves in healthy rats (52). The purpose of this work was to
investigate early-stage regional RILI changes using Hp $^{129}$Xe and the rapid single shot-spiral IDEAL approach. A rat model of RILI involving unilateral exposure of the right lung to $^{60}$Co radiation followed by two weeks incubation period was employed. Using this technique, separate gas, PT and RBC images were acquired at various gas transfer times in order to obtain gas exchange curves for the left and right lungs. The ability of this approach to distinguish regional changes between an irradiated rat cohort from a non-irradiated rat cohort is demonstrated. The differences are correlated with histology and reconciled in terms of pulmonary tissue thickening (i.e. RP) and vascular damage due to RILI using a theoretical gas exchange model.

4.2 Methods

4.2.1 Animal Irradiation

All procedures followed animal use protocols approved by Western University’s Animal Use Subcommittee and were consistent with the guidelines written by the Canadian Council on Animal Care (CCAC). Animal preparation details for irradiation were described previously (41). Two cohorts of five age-matched Sprague Dawley rats (374±26 g, Charles River Laboratories, Saint-Constant, Canada) were prepared for experiments. The animals in the irradiated group were exposed to a total dose of 18 Gy delivered uniformly to the right lung using a $^{60}$Co source at the London Regional Cancer Program centre. A semi-circular collimator with a diameter of 4 cm was used to irradiate the right thorax only. To achieve a uniform dose distribution, 9 Gy was delivered to the rat in the supine position (at a rate of 95.1 cGy/min) and 9 Gy was delivered with the rat in the prone position (at a rate of 80.3 cGy/min), with the collimator appropriately re-positioned to
irradiate only the right lung and spare the left lung. The second group of rats were not irradiated and served as age-matched controls. The irradiated and non-irradiated cohorts were housed in HEPA-filtered cages for two weeks prior to MRI. The image analyst (O. D.) was blinded to the identity of the cohorts until after the analysis of the MRI results.

4.2.2 MR Image Acquisition and Analysis:

Imaging was performed using a 3T MRI system (MR750, GEHC, Waukesha, WI), a custom-built transmit-only/receive-only birdcage coil (53) and a high performance insertable gradient system as described previously (54). Hp\textsuperscript{129}Xe gas (approximately 80% enriched with 10% polarization) was obtained from a commercial xenon polarizer system (Polarean 9800, Durham, NH). All animals were imaged in the supine position following three breaths of Hp\textsuperscript{129}Xe using a MRI-compatible mechanical ventilator (GE Healthcare, Malmo, Sweden) for delivery of Hp\textsuperscript{129}Xe to the lung as described previously (47).

**Four-point IDEAL:** Images corresponding to gas, PT and RBC compartments were acquired using a single-shot spiral, four-point IDEAL pulse sequence described previously (1). Briefly, four-point IDEAL images were obtained using four echoes with a low flip angle on the gas phase, high flip angle on the dissolved phases per echo and echo-times of 340 us × m, (m=1, 2, 3, 4). The sequence was averaged 10 times during a breath-hold period of 3 s. The gas, PT and RBC images were reconstructed off-line using a Matlab routine (MathWorks, Natick, MA) as described in Schulte et al. (2,3).

To obtain \textsuperscript{129}Xe gas exchange curves, the acquisition above was repeated for TR = 30, 50, 100, 200, 300 ms using separate breath-holds. Other pulse sequence parameters included: \textit{FOV} = 70 mm, \textit{BW} = 125 kHz, a nominal matrix resolution of 22. The four-point gas
images were used to define ROIs corresponding to the left and right lungs of each rat (as described in Figure 4.1). ROIs were used as masks to identify the lungs on all subsequent four-point IDEAL images. The first acquired PT and RBC images were discarded since they were not weighted by TR. The PT and RBC images were normalized by the Hp $^{129}$Xe gas image to account for any systematic differences between breath-holds, including polarization and ventilation. $^{129}$Xe gas exchange signal curves ($S_{PT}$ and $S_{RBC}$ vs. TR) were then calculated corresponding to each lung (i.e. right vs. left) for the PT and RBC compartments respectively for further analysis as described below.

**Three-point IDEAL**: Compared to four-point, higher SNR, temporal and spatial resolution can be achieved using three-point IDEAL (1). However, three-point IDEAL is a less practical imaging method for processing large cohorts since it results in long scan times and gas consumption per animal. Therefore, for demonstration and conceptual purposes, representative gas-PT and gas-RBC images were acquired for separate breath-holds for only two rats in the cohort. Figure 4.1 shows a representative set of three-point IDEAL images explaining the segmentation method (used also for the four-point IDEAL gas exchange analysis above). Three-point IDEAL images of gas-RBC and gas-PT were acquired for a $FOV = 60\, mm$ and $TE = 50\, us \times m$, ($m = 1, 2, 3$) from an irradiated animal in the cohort study using two separate breath-holds. To achieve adequate SNR and spatial resolution, the representative three-point IDEAL images were averaged 15 times during each breath-hold taking advantage of higher temporal resolution.
4.2.3 Histology

Following MRI, rats were euthanized and lungs were extracted for histological analysis. The rat lungs were fixed by intratracheal infusion of 9 mL of 10% neutral-buffered formalin, and embedded in paraffin wax (57,58). The tissue was then cut into 5 µm thick tissue sections and stained with haematoxylin-eosin (H&E). The sections were chosen to avoid major airways and vessels and be evenly distributed between the apex and base of each lung. Five images were acquired from both the left and right lungs at 40× magnification corresponding to a pixel resolution of 0.23 µm using a Zeiss Axio Imager microscope with a Retiga EXi Digical CCD camera (Q Imaging, Vancouver, BC, Canada). Each image was then binarized using Matlab image processing functions (graythresh.m, Matlab). The average percent tissue area (PTA), was calculated from each image by counting the number of pixels with signal corresponding to all non-airspaces (e.g. parenchyma, vessels etc.) and dividing by the total number of pixels in the field using an automated binary thresh-holding approach (59). The PTA was calculated separately for both the left and right lungs for each rat of both cohorts.
4.2.4 Gas Exchange Curve Analysis

The theoretical gas exchange model of Mansson et al. (1) was fitted to the $S_{PT}$ and $S_{RBC}$ data as a function of $TR$ for the left and right lungs. This analysis was similar to the fitting of CSSR data from whole lungs described previously (46,47) and used the following exponential function:

$$S(TR) = S_o \cdot (1 - \exp( TR / \tau )) + S_i \cdot TR$$  \[4.1\]

where $S_o$ is the y-intercept of the linear slope, $\tau$ is the exponential gas transfer time constant and $S_i$ is the linear slope. From this fitting, the parameters $S_o$, $S_i$, and $\tau$ were obtained separately for $S_{PT}$ and $S_{RBC}$ for each of the right and left lungs of all rats. The average alveolar tissue thickness ($L_{PT}$) and the relative RBC volume ($V_{RBC}$) were then calculated for both left and right lungs of all rats as described in (1). Additionally, a total PT and RBC signal, $S_{total} = S_{PT} + S_{RBC}$, for each of the right and left lungs of each rat was calculated based on the respective normalized image signal intensities at $TR = 300$ ms, for comparison with $PTA$.

4.2.5 Statistical Analysis

Statistical analysis was performed using Tukey’s multiple comparison as part of a two-way ANOVA analysis (Graphpad Prism software, version 6, La Jolla, CA, USA) within a confidence interval of 95% to assess whether the differences in $L_{PT}$ and $V_{RBC}$ between left and right lungs were significant for the irradiated and non-irradiated groups. Additionally, Fisher’s Least Significant Difference (LSD) multiple comparison test was used to compare the mean $PTA$ of the irradiated and non-irradiated groups within a 95% confidence interval.
Unlike the Tukey’s multiple comparison test, Fisher’s LSD test does not correct for multi-
comparison of the left and right lungs, since the PTA values were not found to differentiate
between the left and right lungs of either the irradiated or unirradiated cohorts, as discussed
later. Finally, a correlation analyses between $S_{total}$ and $PTA$ was performed using two-tailed
nonparametric Spearman correlation within a confidence interval of 95%.

4.3 Results

Figure 4.1 shows coronal three-point IDEAL images of (a) gas, (b) PT and (c) RBC
compartments from a representative irradiated animal. The gas compartment image
indicated in grey scale corresponds to the ventilation of $^{129}$Xe gas in the airways and in the
alveoli. The false color scale of the PT and RBC images allows the dissolved $^{129}$Xe signal
to be compared qualitatively and quantitatively between the left and right lungs taking
advantage of higher SNR of three-point IDEAL. In particular, the PT image shows
excellent correspondence with the gas image. The RBC image displayed signal both within
and outside the lung and is notably higher at the center of the image, corresponding to the
location of the heart and main pulmonary vessels. Figure 4.1 (d) and (e) show the RBC and
PT images respectively in false color overlaid on the corresponding gas image and cropped
to the contour lines obtained from the gas image. Regional intensity variations could be
seen between the right (irradiated) and left (unirradiated) lungs. An increase in the $^{129}$Xe
tissue image signal from the irradiated right lung compared to the unirradiated left lung is
apparent in Figure 4.1 (b) and (d). By contrast, the RBC image signal of the irradiated right
lung in Figure 4.1 (c) and (e) is reduced compared to the un-irradiated left lung.
Figure 4.1: Representative coronal gas (a), PT (b), and RBC (c) images from an irradiated animal obtained using three-point IDEAL for gas-PT and gas-RBC separately. The white contour lines represent the masks for the left and right lungs used for region-of-interest (ROI) analysis. The left and right lungs of PT (d) and RBC (e) images (in color) and underlying gas images (in grey scale) with contour lines are shown for visual confirmation of the PT and RBC. The strong signal in the center of RBC image (c) as shown by the arrow appears to correspond to the left atrium, ventricle and arch of the aorta.

Experimentally measured gas exchange curves ($S_{PT}$ and $S_{RBC}$ as a function of $TR$) for both the left and right lungs of the non-irradiated and irradiated cohorts are shown respectively in Figure 4.2 (a), (b), (c), and (d). In Figure 4.2, the points represent the mean values based on all animals in the two cohorts and the error bars represent the standard deviation based on all animals in the respective cohorts. The smooth curves are the best fits to the data based on Eqn. [4.1], yielding R-square values $> 0.9979\pm0.0018$, for all animals.
Figure 4.2: Mean gas transfer curves and corresponding fits to Eqn. [4.1] for the left and right lungs of the unirradiated group (N=5) for PT (a) and RBC (c). Mean gas transfer data and corresponding fits to Eqn. [4.1] for the left and right lungs of the irradiated group for PT (b) and RBC (d). The smooth curves represent the best fits to the data and the points and error bars represent the mean values calculated over all the animals in each respective cohort for each lung.

The individual $L_{PT}$ and $V_{RBC}$ values are shown in a scatter plot in Figure 4.3 and summarized in Table 4.1 for both the left and right lungs of both cohorts. The mean $L_{PT}$ values of the left and right lungs (Figure 4.3 (a)) were determined to be 6.46±0.2 µm and 6.40±0.2 µm respectively for the non-irradiated group (Table 4.1). The irradiated cohort demonstrated a significant increase in $L_{PT}$ compared to the non-irradiated cohort of approximately 8.2% ($P = 0.0002$) and 5.0% ($P = 0.0497$) for the right lungs and the left lungs respectively. As expected, $L_{PT}$ was not statistically significantly different ($P = 0.9682$) between the left and right lungs of the non-irradiated cohort. $L_{PT}$ was observed to be statistically significantly
different between the irradiated right lungs and unirradiated left lungs of the irradiated cohort (P = 0.0210). The irradiated cohort exhibited a reduction in $V_{RBC}$ compared to the non-irradiated cohort as shown in Figure 4.3 (b). As expected, the mean $V_{RBC}$ values were not statistically different for the left and right lungs (P = 0.9353) of the non-irradiated cohort. Compared to the non-irradiated cohort, $V_{RBC}$ values of the irradiated cohort decreased by 36.1% (P = 0.0002) and 11.7% (P < 0.0354) for the right and left lungs respectively. $V_{RBC}$ values were statistically significantly different (P = 0.0223) between the irradiated right lungs and unirradiated left lungs of the irradiated group and showed variations similar to $L_{PT}$ in Figure 4.3 (a).

Figure 4.3: Scatter plots of calculated $L_{PT}$ (a) and $V_{RBC}$ (b) values for all animals for both left and right lungs from fit to Eqn. [4.1] in Figure 4.2. Corresponding p values are also shown.
Table 4.1: Summary of mean PT thickness ($L_{PT}$) and relative blood volume ($V_{RBC}$) calculated for all rats for both left and right lungs.

<table>
<thead>
<tr>
<th>Rat number</th>
<th>Right Lung $L_{PT}$a [µm]</th>
<th>Left Lung $L_{PT}$ [µm]</th>
<th>Right Lung $V_{RBC}$b</th>
<th>Left Lung $V_{RBC}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>NON-IRRADIATED</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>6.54±0.5</td>
<td>6.43±0.5</td>
<td>0.36±0.07</td>
<td>0.39±0.08</td>
</tr>
<tr>
<td>2</td>
<td>6.34±0.3</td>
<td>6.25±0.6</td>
<td>0.41±0.08</td>
<td>0.44±0.10</td>
</tr>
<tr>
<td>3</td>
<td>6.19±0.4</td>
<td>6.51±0.4</td>
<td>0.45±0.10</td>
<td>0.37±0.07</td>
</tr>
<tr>
<td>4</td>
<td>6.47±0.4</td>
<td>6.26±0.5</td>
<td>0.38±0.11</td>
<td>0.44±0.08</td>
</tr>
<tr>
<td>5</td>
<td>6.78±0.6</td>
<td>6.54±0.5</td>
<td>0.30±0.12</td>
<td>0.36±0.09</td>
</tr>
<tr>
<td>M±SD</td>
<td>6.46±0.2</td>
<td>6.40±0.2</td>
<td>0.38±0.04</td>
<td>0.40±0.04</td>
</tr>
<tr>
<td>IRRADIATED</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>7.65±0.4</td>
<td>6.74±0.5</td>
<td>0.14±0.07</td>
<td>0.31±0.07</td>
</tr>
<tr>
<td>2</td>
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<td>0.16±0.08</td>
<td>0.17±0.05</td>
</tr>
<tr>
<td>3</td>
<td>7.18±0.6</td>
<td>6.50±0.6</td>
<td>0.20±0.07</td>
<td>0.37±0.08</td>
</tr>
<tr>
<td>4</td>
<td>7.08±0.5</td>
<td>6.64±0.6</td>
<td>0.23±0.08</td>
<td>0.34±0.10</td>
</tr>
<tr>
<td>5</td>
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<td>6.89±0.7</td>
<td>0.19±0.05</td>
<td>0.29±0.08</td>
</tr>
<tr>
<td>M±SD</td>
<td>7.30±0.2</td>
<td>6.82±0.3</td>
<td>0.18±0.03</td>
<td>0.30±0.03</td>
</tr>
</tbody>
</table>

*a$L_{PT}$ and b$V_{RBC}$ are the pulmonary tissue thickness and the relative blood volume respectively obtained from the Mansson model (4).

Figure 4.4 shows binarized histology images of (a) a non-irradiated and (b) an irradiated rat lung chosen to visually accentuate the differences in pulmonary tissue thickening. Figure 4.4 (c) shows a scatter plot of $PTA$ values between the left and right lungs in the non-irradiated and irradiated groups. The $PTA$ values from the left (23.2 ± 4.8 %) and right (22.2 ± 3.1 %) lungs in the non-irradiated group were not statistically significantly different
(P = 0.855) and close to expected values for normal animals (59) confirming that the rats in this cohort were healthy. The PTA values of both the irradiated right lungs and unirradiated left lungs of the irradiated cohort were observed to be significantly elevated (~26.8 %) compared to the non-irradiated cohort for both the left (P = 0.048) and right lungs (P = 0.0089), confirming the presence of radiation pneumonitis. However, there was no statistically significant difference in PTA measurements between the irradiated right lungs and unirradiated left lungs of the irradiated cohort (P = 0.524) showing that histology was not able to distinguish differences in RP between the irradiated right lung and the unirradiated left lung of the irradiated cohort. Figure 4.4 (d) shows a scatter plot of the measured PTA values for all rats versus $S_{total}$. The PTA measurements showed a significant correlation with $S_{total}$ for both the non-irradiated ($r = 0.795$, P = 0.0081) and irradiated groups ($r = 0.911$, P = 0.0005) and the two cohorts could be distinguished on the basis of $S_{total}$. 
Figure 4.4: Binarized histology images of a non-irradiated (a) and an irradiated (b) rat lung obtained at 10× magnification chosen to accentuate changes in pulmonary tissue thickening (i.e. PTA). Scatter plot (c) showing PTA values for all rats and p values from Fisher’s LSD multi-comparison test. Scatter plot (d) showing PTA and \( S_{total} \) for all rats. The corresponding Spearman correlation coefficients were (\( r=0.911, p=0.0005 \)) and (\( r=0.795, p=0.0081 \)) for the irradiated and non-irradiated groups respectively.
4.4 Discussion

The results of this work confirm the feasibility of measuring changes in gas exchange curves associated with RILI regionally using Hp $^{129}$Xe MRI and a single-shot spiral four-point IDEAL approach. Regional mapping of gas exchange curves based on imaging of PT and RBC compartments separately enables regional differences in RILI to be resolved, allowing contributions from RBC signal (i.e. heart and main pulmonary vessels), to be excluded from the PT signal, and thereby provide insight into radiation pneumonitis (RP) spatially, based on the choice of region of interest. Specifically, this work demonstrates that $S_{PT}$ and $S_{RBC}$ gas exchange curves are significantly different for irradiated rats compared to non-irradiated rats at two weeks following irradiation. Based on these differences in gas exchange curves, the calculated $L_{PT}$ and $V_{RBC}$ were found to be statistically larger and smaller respectively for the irradiated cohort compared to the non-irradiated cohort. The $L_{PT}$ thickening was shown to correlate well with RP based on $PTA$ determined using histology. Furthermore, differences were more significant for the irradiated right lung compared to the unirradiated left lung of the irradiated cohort, as expected.

The sufficiently high SNR of PT images enables precise measurement of $S_{PT}$ gas exchange curves from the left and right lungs separately with a standard error of less than 5% of the mean $S_{PT}$ ($N = 5$). As expected, the $S_{PT}$ curves and calculated PT thickness in the non-irradiated group were approximately the same for the left and right lungs. The signal from blood plasma, which has the same chemical shift (197 ppm) as the PT signal, was expected to be highest from the heart and corresponding arteries on PT image. However, it did not
appear to be as strong as the RBC signal from the heart. Therefore, we assume that the contribution of blood plasma to the PT signal is not significant since the Ostwald solubility of xenon in the blood plasma is approximately three times less than the RBC (60). Furthermore, since the ROIs were also chosen such that the heart was excluded, the contribution of plasma from the heart and great vessels to the tissue signal from the left and right lungs can be safely ignored. However, it should be noted that this assumption may not hold for the whole lung CSSR spectroscopy technique as discussed further below.

Both $S_{PT}$ and $S_{RBC}$ curves had a standard error less than reported for a whole lung CSSR spectroscopy technique (~15%) (47) confirming the ability of the IDEAL imaging approach to more accurately localize signal contributions. Furthermore, including the signal from $^{129}$Xe-rich blood pool (i.e. heart and pulmonary arteries) leads to overestimation of the linear slope ($S_1$) and thereby the gas transfer time constant ($\tau_{RBC}$) with CSSR techniques for long gas transfer times (or $TR$) on the order of seconds. The direct comparison of fitting parameters obtained in this study to previously-published whole lung CSSR spectroscopy results is not straight-forward due to the confounding effects of multiple regional PT and RBC signals affecting CSSR. As discussed previously by Chang et. al. (44) implementing overestimated fitting parameters would also cause inaccurate calculation of physiological parameters.

Nevertheless, the $S_{PT}$ from the irradiated group showed a higher linear slope compared to the non-irradiated group, which was consistent with findings previously reported by Fox et. al. (47) for a whole lung irradiated rat model within the transfer time interval of 300 ms (61). The signal amplitude was correlated to increased tissue volume (48). A higher signal amplitude on the injured right lung compared to the non-irradiated left lung was also noted
in the representative PT image (Figure 4.1). Although the pulmonary tissue thickening developed in both lungs, the impact of the injury was more severe on the irradiated right lung leading to image intensity variations when compared to the unirradiated left lung. Presumably this is due to the development of RP in the early stages of RILI (after two weeks of irradiation) resulting in tissue thickening from inflammation and/or edema. Correspondingly, the time course of exchange of Hp $^{129}$Xe from the gas phase to PT will be elevated due the increased PT thickness leading to a steeper gas exchange curve (or increased $S_i$) in the early TR data points. The PT gas exchange curves from the irradiated right lungs were also statistically different from those of the unirradiated left lungs of the irradiated cohort confirming the presence of more significant RP in the right lung. This agrees with the findings previously reported by Thind et al. for a partially irradiated rat lung model (37).

Despite the approximations inherent to the theoretical gas exchange model used here and designed for global lung function with the CSSR technique, the average $L_{PT}$ values agree well with previously published healthy rat pulmonary tissue thicknesses (5-6 µm) obtained using optical microscopy techniques (57). Although the mean PTA measurements made in this study using histology (Figure 4.4 (a) and (b)) were not capable of separating the contributions of PT and vessels (including RBC) from each other and any other tissue present, the mean PTA of the irradiated cohort was still found to be statistically significantly different from the non-irradiated cohort confirming that the irradiated animals had developed RP. This is perhaps not surprising since the tissue thickening due to inflammation is likely more significant than any losses in blood vessels (and RBC) due to irradiation, thus giving rise to a net increase in PTA (62). Nevertheless, in contrast to the
MRI approach, the $PTA$ was not found to be different between the left and right lungs of the irradiated cohort, suggesting that regional differences in RP and vessel damage cannot be discriminated using histology. The densities of vessels in the histology images and choice of the slice location could help explain discrepancies in measured $PTA$. Further studies using vessel and RBC specific techniques (e.g. autofluorescence) are needed to histologically measure the pulmonary tissue and RBC separately for comparison with the $L_{PT}$ and $V_{RBC}$ values obtained from MRI measurements.

The selection of ROIs based on the three-point IDEAL gas images permits the assessment of RBC gas exchange curves, $S_{RBC}$, primarily from the pulmonary capillaries, excluding the confounding signal from the heart and other great vessels. Interestingly, there was approximately a 5% difference in $S_{RBC}$ observed between the left and right lungs in the non-irradiated group which may due to the inherent difference between the normal total blood volumes in the healthy right and left lungs of the rat (3). Another possible explanation for the difference between the left and right lung in the non-irradiated group might be to low signal, variation in the regional pulsation (60) and the asymmetry due to the strong signal from the heart and corresponding arteries in the region of interest. Nevertheless, a relatively large standard error in $S_{RBC}$ (~10%) compared to the $S_{PT}$ (~5%) was observed and the $S_{RBC}$ differ, on average, approximately two-fold in the irradiated cohort compared to the non-irradiated group.

The reduction of arterial perfusion and obstruction of capillaries in the irradiated lungs due to RILI has been observed histologically in other studies (3–5). However, reduced perfusion has only previously been reported after 30 days post-irradiation using SPECT. The early and significant reductions in $S_{RBC}$ with irradiation observed in this study could potentially
provide another method by which to quantify and discern the mechanisms of RILI, for example RP versus vascular changes known to depend on the method of radiation fractionation. Stereotactic body radiation therapy (SBRT) which is based on giving a large total dose (60 Gy) using fewer fractions (20 Gy/fraction) has been shown to provide better local tumor control and higher rates of local progression-free survival (63,64). Despite the improved survival rates with high dose approaches, lung toxicity is still a limiting factor (65) and detection of RILI after two weeks irradiation may help to reduce incidence of radiation pneumonitis. In particular, SBRT could benefit from early identification of RP and vascular changes using the approach described in this study. The two-week post-irradiation time interval chosen for this study should allow detection of the early stage pneumonitis and re-evaluation of the dose fractionation scheme or indicate the need for pharmaceuticals (i.e. radioprotective drugs). Further studies are needed to identify how could this new imaging technique would help to improve RT planning for patients.

In this study, the theoretical model of Mansson et al. (1) was used to extract measures of average pulmonary tissue thickness and relative blood volume regionally. This model is based on one-dimensional radial diffusive exchange in a 2D compartmental geometry. It is likely that a more sophisticated diffusive gas exchange which accounts for the 3D structure of alveoli including blood flow within surrounding capillaries would provide more physiologically insight and help to reveal RILI mechanisms in more detail, including vascular damage (61). In particular, measurement of changes in surface area between the capillaries and alveolar walls independent of changes in perfusion would help to assess early RILI. This approach could also be applied to perfusion of other organs specifically the heart using a dedicated TO/RO RF coils optimized for heart imaging (53).
The four-point IDEAL method used here should be translatable to clinical early-phase RILI detection within a single breath-hold time of 10.7 seconds using an interleaved spiral approach with $TR$ values of 15, 30, 50, 100, 250 ms and would benefit by acceleration provided by parallel MRI. Higher polarizations (30-40%) should also allow measurement of regional gas exchange curves and mapping on a voxel-by-voxel basis and, potentially, isotropic 3D imaging allowing resolution of early-phase RILI regional effects associated with complex dose distributions in the lung.

4.5 Conclusion

Regional gas exchange changes associated with early (two week) radiation pneumonitis in a rat model of Radiation-Induced Lung Injury (RILI) can be measured using $Hp^{129}$Xe MRI and a single-shot spiral IDEAL approach. The pulmonary tissue (PT) and red blood cell (RBC) compartments are significantly different in irradiated rats compared to non-irradiated rats. The novel approach was also capable of distinguishing irradiated lung from the non-irradiating lung suggesting that the early-phase changes can be measured regionally. The measured pulmonary tissue thickness ($L_{PT}$) of non-irradiated rats is in agreement with previously reported histology measurements, and the $L_{PT}$ from irradiated rat lungs is significantly elevated compared to unirradiated lung and correlates with histology. In addition, the RBC volume ($V_{RBC}$) is reduced in the irradiated lung. These preliminary in vivo results confirm that this new approach can be used to quantify regional gas exchange changes associated with a rat model of early stage RILI two weeks post irradiation and potentially could be translated to humans for improving thoracic radiotherapy outcomes.
4.6 References


Chapter 5 : Thesis Discussion and Future Work

5.1 Thesis Discussion

Various imaging modalities such as SPECT, PET, CT, Hp MRI, and non-imaging techniques (eg. PFTs) are capable of providing functional information about RILI. Among them, Hp $^{129}$Xe MR imaging technique may exceed the capacities of established modalities for detection of early phase RILI. The reason for this is $^{129}$Xe diffuses in the lung air space, PT and RBC in a way similar to $O_2$ providing relevant functional information about the gas exchange (ie. gas exchange curves). Additionally, $^{129}$Xe is a chemically inert and metabolically inactive gas in the PT and RBC. $^{129}$Xe gas exchange curves can be quantitatively used to explore regional lung function when fitted to appropriate theoretical models.

In this work, a new approach using Hp $^{129}$Xe MRI was developed to quantitatively measure the early phase of RILI in a rat model two weeks post-irradiation. The design and development of this approach required the establishment of multi-disciplinary interactions, including technical developments for improved Hp MRI tools. Firstly, SNR was increased by designing, constructing and optimizing a custom transmit-only / receive-only (TO/RO) coil consisting of two components: (i) a high-pass birdcage transmit-only coil that produced a homogenous $B_1$ magnetic field, and (ii) a saddle-shaped receive-only surface coil that allowed high sensitivity for the lung. Next, a single-shot spiral IDEAL imaging technique was implemented and optimized for Hp $^{129}$Xe dissolved phase imaging. In vivo results using a rat model showed regional detection of xenon exchange curves. Finally,
with the use of the TO/RO coil in combination with the single-shot spiral IDEAL pulse sequence, the feasibility of early detection of RILI in a rat model two weeks-post irradiation (18 Gy) was investigated. Lung tissue for histological examinations were obtained from both an irradiated (N=5) and a non-irradiated (N=5) cohort on order to confirm whether the rats developed RILI at two weeks following irradiation. The following sections provide an overall discussion about each chapter and addresses the potential of translating this Hp $^{129}$Xe MRI technique to clinical imaging.

5.2 SNR Improvement

Chapter 2 described the study performed to improve the SNR by developing a TO/RO coil configuration for rat imaging. In general, Hp $^{129}$Xe gas phase imaging is conventionally achieved using birdcage volume coils, which provide acceptable transmit RF field and image homogeneity for gas imaging. However, improved SNR is necessary for imaging the dissolved phases of $^{129}$Xe as they represent only 2% of the total signal available. To demonstrate the efficiency of our novel TO/RO RF coil configuration, a thermally-polarized phantom was imaged in Chapter 2. Since there was not an available dissolved phase MR sequence and the polarization of the $^{129}$Xe gas was limited (~5%), the feasibility of the TO/RO RF coil was initially tested for imaging only the gas phase in vivo in rat lungs. FGRE coronal images showed an SNR improvement factor of approximately three with the new TO/RO coil configuration compared to the conventional T/R coil configuration. Although this SNR improvement was encouraging, further development was required for imaging the dissolved phases of $^{129}$Xe.
In Chapter 3, an additional improvement of approximately three-fold in SNR was accomplished using a single shot-spiral IDEAL pulse sequence. The total improvement in SNR, using both the new TO/RO RF coil configuration and the single-shot IDEAL spiral imaging sequence, enabled imaging of gas, PT and RBC compartments separately. Even though polarization levels of 10% with enriched $^{129}$Xe gas was used during the study, the SNR was not sufficient to achieve the necessary image quality to capture the gas exchange curves within a single breath-hold (6-10s). To address this problem, 10 sets of IDEAL images were acquired for a specific TR value within a single breath-hold time. Apart from the non-renewable nature of the noble gas polarization, the dissolved phase signal replenishes during the TR value due to transfer from the gas phase reservoir. The IDEAL sequence used this property to improve SNR by additional factor of three. Altogether, a total improvement in SNR was approximately 27-fold for each breath-hold image. Consequently, the measurement of gas exchange curves was achievable by analyzing the image intensity variations as a function of TR.

5.3 Early Detection of RILI in a Rat Model

Chapter 4 discussed the use of the new TO/RO RF coil and single-shot spiral IDEAL pulse sequence for detection of RILI in a partially irradiated rat thorax. The dynamics of gas exchange significantly changed within the two weeks post-irradiated group compared to the control group. This was consistent with previously published $^{129}$Xe CSSR MR results in a whole-lung irradiated rat model (1). In particularly, we investigated whether the new imaging approach would allow detect of regional changes due to the partial-irradiation of
the right thorax. This new technique was able to distinguish the irradiated lung from the unirradiated lung in the irradiated cohort group.

Compared to previous published studies using SPECT, PET, and CT, Hp $^{129}$Xe MRI was also capable of differentiating the effects of RILI on the PT and RBC compartments. An increase in $L_T$ was observed in both the left and right lungs in the irradiated cohort compared to the non-irradiated cohort (Table 4.1). Although, the increase in $L_T$ was less in the unirradiated (left) lungs compared to the irradiated right lungs of the irradiated cohort, this result was not surprising, since it was also observed in the histology and has been reported previously in other imaging studies such as Hp $^{13}$C (2). One hypothesis to explain this effect is that the whole organ responded to the irradiation delivered to one side. A significant drop in $V_{RBC}$ was observed in the irradiated lungs, which was similar to observations in studies using SPECT lung perfusion (3,4). However, SPECT was only able to detect RILI with doses greater than 20 Gy and four weeks post-irradiation. Therefore, Hp $^{129}$Xe MRI was highly sensitive to the regional changes of RILI as early as to two weeks post-irradiation with 18 Gy in the rat model.

The PTA values derived from the histology of the total pulmonary tissue and surrounding capillaries were unable to distinguish between the irradiated and unirradiated lungs in the irradiated cohort as discussed in Chapter 4. However, PTA measurements showed a statistically significant difference between the irradiated and non-irradiated rat cohorts. In the future, the histological measurements could be improved using better techniques to validate $L_T$ and $V_{RBC}$ measurements separately. For example, autofluorescent microscopy techniques could be used in future to verify changes in $V_{RBC}$ (5).
5.4 Analysis of Gas Exchange Curves

The quantification of lung function using theoretical 1D gas exchange models as used in this thesis are often based on assumptions that over-simplify the alveolar geometry as shown in Figure 1.11 (b) (6). Therefore, the 1D-gas exchange model used in this thesis can only be used to calculate the variations in the 1D (i.e. radial) diffusion path length (i.e. $L_T$ and $L_c$) separately for the PT and RBC compartments. Changes in the diffusion path length are then interpreted in terms of other estimations of global function of the lungs, including the relative blood volume by integration of the 1D-geometry over the entire alveolar cell volume. The 1D circularly symmetric gas exchange model excludes information about the gas exchange volume and diffusion surface area between (i) the alveolar air space and the PT compartment, and (ii) the PT and RBC compartments. This information is more appropriately extracted using a 3D-gas exchange model of the alveolus.

Another 1D-gas exchange model, the MOXE model, also interprets CSSR gas exchange data through diffusion path lengths by solving diffusion equations (7). In the MOXE model, the missing link between the diffusion path length and the CSSR measurement is provided using two normalization factors including the surface-to-volume ratio and barrier-to-septum ratio. The normalization factors are empirically obtained from the fitting parameters of the gas exchange curves. The first normalization factor is used to scale the solution of the diffusion equations between the gas and PT (8). The second normalization factor is used to scale the solution of the diffusion equations between the PT and RBC (7).

Overall, all the existing analytical gas exchange models treat the lung as a simplified uniform 1D model to fit to the CSSR data from the whole lung. In this study, the Mansson
model was used to analyze the gas exchange curves because the Mansson model was easy to implement and did not require additional normalization factors. However, as already discussed in Chapter 4 (Figure 4.1), the RBC image intensity distribution was significantly different regionally from both the gas and PT compartment images. Therefore, an improved theoretical model with a more sophisticated 3D geometry may be important to consider for explaining the spatial variation in RBC signal intensity. Additionally, the dynamics of the gas diffusion from the PT to RBC has to consider the diffusion area, possible using a 3D numerical model (9). In future, using a 3D model, the RBC compartments could be simulated without using a scaling factor from the PT signal. In this way, a study using a 3D numerical model may reveal mechanisms of RILI changes in the RBC compartment independently from the PT compartment.

5.5 Future Work

Although SNR improvement accomplished in this thesis with the TO/RO RF coil and single-shot spiral IDEAL pulse sequence design, was able to capture the dynamics of gas exchange within a single breath-hold, this was still challenging with the relatively modest 10% polarization and required multiple breath-holds to perform averaging, especially for the low TR values. The SNR can be significantly elevated using new commercially available polarizers (i.e. Polarean 9820 $^{129}$Xe hyperpolarizer, Polarean Inc., Durham) that can produce polarization levels of approximately 45%. Further improvement in polarization can be achieved by addressing the unwanted thermal gradients along the SEOP cell, improving the spin exchange interaction time between $^{129}$Xe and polarized $^{87}$Rb, and improving the filtration of $^{129}$Xe process traps. The use of higher polarizations (~30% or
higher) in future would significantly improve the $^{129}$Xe dissolved phase signal. In this case, smaller flip angle values, on the order of 1-2 degrees, would be sufficient to produce enough SNR for the gas phase images. Since the low flip angles do not consume the gas reservoir magnetization, a further increase in dissolved phase SNR would be achieved due to the replenishment of dissolved phase magnetization. This could be accomplished by using a selective excitation RF pulse based on a composite pulse approach \((10)\).

As discussed previously in Chapter 3, the read-out time \(T_{\text{read}}\) of the spiral k-space sampling scheme should be as low as possible to achieve the desired resolution. In this thesis, \(T_{\text{read}}\) values of 2-3 ms were obtained using a high performance insertable gradient system. Unfortunately, typical clinical gradient coil strengths are not strong enough to sample the whole k-space data within such a short \(T_{\text{read}}\). Therefore, interleaved spiral approaches could be used to address this limitation in future. With interleaved spirals, the whole k-space data could be divided into six interleaves and sampled after the application of independent RF pulses. Using an interleaved spiral approach, the dependence on high slew rates and gradient amplitudes could be eliminated for clinical MR imaging. This could also be tested using a healthy rat.

Imaging at low magnetic field strengths (0.5 T – 1.5 T) could significantly improve the spatial resolution. The \(T_2^*\) is inversely proportional to the magnetic field strength allowing for the use of longer \(T_{\text{read}}\). The longer \(T_{\text{read}}\) would be beneficial for improving spatial resolution or reducing the number of interleaved acquisitions required for clinical gradient configurations. Therefore, translation of the technique presented in this thesis towards human imaging would benefit from the use of an interleaved spiral approach, higher polarization, and a selective excitation RF pulse.
The MRI technique developed in this thesis can potentially be used to diagnose gas exchange abnormalities resulting in other lung diseases such as chronic obstructive pulmonary disease (COPD), asthma, and cystic fibrosis. Corresponding changes in the gas exchange curves can be helpful to detect regional physiological abnormalities. Additionally, the strong dissolved phase signal from the RBC compartment of the heart can be translated to a perfusion image by taking advantage of high polarization and optimized RF coil configurations. Similarly, white matter, grey matter and blood flow of the brain can be imaged by taking advantage of the different chemical shifts using a spiral IDEAL imaging approach in the future.

5.6 Thesis Summary

A novel Hp$^{129}$Xe MRI technique was developed and used to successfully detect Radiation-Induced Lung Injury (RILI) in a rodent model as early as two-weeks post-irradiation. To test the hypothesis, firstly, a TO/RO radiofrequency coil configuration was optimized and built for hyperpolarized$^{129}$Xe MRI providing a three-fold SNR improvement compared to the commercially available RF coils. Next, a single-shot spiral IDEAL MRI pulse sequence, capable of separating the Hp$^{129}$Xe gas and dissolved phase images, was implemented to image rat lungs. This new imaging approach offered benefits of fewer and higher flip angle RF pulses to increase the SNR of the PT and RBC images. Additionally, the short read-out times of single-shot spiral IDEAL permitted multiple gas, PT, and RBC images to be acquired within a single breath-hold time of a few seconds. Substantially improved SNR of the PT and RBC images and short scan time permitted imaging of the exchange between the gas, PT and RBC compartments for regional characterization of lung
microstructure and pulmonary and vascular function. Finally, regional gas exchange changes associated with radiation pneumonitis in a rat model of RILI was measured using a single-shot spiral IDEAL approach. An in vivo proof-of-concept approach was able to distinguish the irradiated lung from the non-irradiating lung. The measured pulmonary tissue thickness ($L_{PT}$) of non-irradiated rats were found to increase in the irradiated cohort and was in agreement with previously reported studies. Additionally, in the irradiated cohort, the $L_{PT}$ from the irradiated rat lungs was significantly elevated compared to unirradiated lung. Furthermore, again in the irradiated cohort, the RBC volume ($V_{RBC}$) was significantly reduced in the irradiated lung compared to unirradiated lungs. A histological measurement of the percent tissue area (PTA) was correlated to the pathology of the irradiated and non-irradiated cohorts. The novel Hp $^{129}$Xe MRI method presented in this thesis showed regional change in gas exchange curves associated with a rat model of early-phase RILI two-weeks post-irradiation. The method is a promising approach for detection of early-phase RILI and has considerable potential for translation clinic, including radiation therapy, for management of RILI.
5.7 References


Appendix A: Calculation of Spiral PSF

$H(k_r,k_\theta)$ is the modulation transfer function in polar coordinates for the spiral k-space trajectory given by:

$$H(k_r,k_\theta) = H_w(k_r,k_\theta) \times \exp\left(-\frac{\Delta r T_{\text{read}}}{2\pi T^*} (k(r,\theta))\right)$$  \[A.1\]

where, $H_w(k_r,k_\theta)$ is a rect function used for truncation of k-space invoking circular symmetry and can be reduced to $H_w(k_r)$ for 1D in the radial direction, $\Delta r$ is the nominal resolution in k-space, $T_{\text{read}}$ is k-space readout time. The point spread function, $h(r,\theta)$, is calculated by the inverse Fourier transform of the modulation transfer function, $H(k_r,k_\theta)$:

$$h(r,\theta) = \mathcal{F}^{-1}\{H(k_r,k_\theta)\} = \int_0^{2\pi} \int_0^\infty H(k_r,k_\theta) \exp(-i2\pi k_r r \cos \theta) k_r dk_r dk_\theta.$$  \[A.2\]

Substituting Eqn [A.1] in Eqn. [A.2], using the convolution identity and invoking circular symmetry, Eqn. [A.2] can be re-written as follows:

$$h(r) = H_o(2\pi f_1(k_r)f_2(k_r)) = F_1(r) * F_2(r)$$  \[A.3\]

where $F_1(r)$ is the Hankel transform of the rect function, $f_1(k_r)$, and integral transformations whose kernels are Bessel functions. $F_1(r)$ was calculated using besselj.m function for solution of zero-order Bessel functions of the first kind in Matlab (Natick MA, USA) as follows:
In Eqn. [A.4], $F_2(r)$ is the Hankel transform of $f_2(k_r)$ which is dependent on $T_{\text{read}}/T_2$ and can be solved by differentiation in the Laplace transform domain as follows:

$$F_2(r) = H_o\{2\pi f_2(k_r)\} = 2\pi \int_0^\infty H_w(k_r)k_r J_o(2\pi k_r, r) dk_r = 2\pi \int_0^{k_{r\text{max}}} k_r J_o(2\pi k_r, r) dk_r$$

$$= 2\pi \int_0^{k_{r\text{max}}} \frac{d}{2\pi r} [k_r J_o(2\pi k_r, r)] dk_r = 2\pi \int_0^{k_{r\text{max}}} \frac{1}{2\pi r} d[k_r J_o(2\pi k_r, r)] dk_r \tag{A.4}$$

$$F_1(r) = \frac{k_{r\text{max}}}{r} J_1(2\pi rk_{r\text{max}})$$

$$F_2(r) = H_o\{f_2(r)\} = \int_0^\infty \exp(-Bk_r)k_r dk_r J_o(rk_r) = \frac{B}{[r^2 + B^2]^{3/2}} \tag{A.5}$$

where, $f_2(k_r) = \exp(-\frac{\Delta r k_r T_{\text{Read}}}{2\pi T_2^*}), \quad B = \frac{\Delta r T_{\text{Read}}}{2\pi T_2^*}$. $F_2(r)$ was normalized by the global minimum of $F_2(r)$ (for $T_{\text{read}}/T_2^* = 0.01$) before the convolution.
Appendix B: Calculation of Cartesian PSF

$H(k)$ is the modulation transfer function in Cartesian coordinates where $k$-space trajectories are sampled along the frequency encoding direction, $k_x$ from $-k_{x,\text{max}}$ to $k_{x,\text{max}}$ and is given by (1,2):

$$H(k) = \text{rect}(k/W) \times \exp(-t(k)/T_2^*)$$  \[B.1\]

;where, $t(k) = \frac{\Delta x T_{\text{Read}} (k_x)}{2\pi}$ and $\Delta x$ is the nominal resolution. Since the signal decay is dominated by $T_2^*$ effect along the frequency encoding direction, $h(x)$ is calculated analytically along $x$, as follows:

$$h(x) = \exp(-A k_{\text{max}}) \frac{1}{(ix + A)} (\exp(k_{\text{max}} (ix + A)) - \exp(-k_{\text{max}} (ix + A)))$$  \[B.2\]

;where $A = \frac{\Delta x T_{\text{Read}}}{2\pi T_2^*}$.


Appendix C: Dissolved Phase Signal Following Multiple Pulses

A Bloch equation analysis was performed including the replenishment of dissolved tissue magnetization between RF pulses. Assuming the signal from Hp$^{129}$Xe is much larger than the thermal equilibrium magnetization and ignoring $T_1$ relaxation, the transverse ($M_{xy}$) and longitudinal ($M_z$) magnetizations following the first RF pulses ($n = 1$) are given respectively by:

$$M_z(n=1) = M_i \cos(\alpha_G) \quad [C.1]$$

$$M_{xy}(n=1) = M_i \lambda \sin(\alpha_{PT}) \quad [C.2]$$

where, $\lambda$ is the solubility of xenon, $\alpha_G$ and $\alpha_{PT}$ are the flip angles delivered to the gas and pulmonary tissue phases respectively and $M_i$ is the initial gas phase magnetization dependent on the polarization.

The corresponding magnetizations following the second RF pulse ($n = 2$) are given by:

$$M_z(n=2) = M_z(n=1) \cos(\alpha_G) \quad [C.3]$$

$$M_{xy}(n=2) = M(TR) \ M_z(n=1) \lambda \sin(\alpha_{PT}) + M_i \lambda \cos(\alpha_{PT}) \sin(\alpha_{PT}) \quad [C.4]$$

where, $M(TR)$ is the percentage replenishment of $^{129}$Xe in pulmonary tissue depending on TR. $M(TR)$ was calculated by dividing the gas exchange curves by its maximum value thereby making it independent of the MRI parameters. Mansson’s gas exchange model was fitted to previously published CSSR data set of the gas transfer curve for healthy rats. Then, $M(TR)$ was calculated as follows:

$$M(TR) = (S_o \left[1 - \exp(-\Delta / \tau_1)\right] + S_i \Delta) / M(TR_{max}) \quad [C.5]$$
where, the pulse repetition time, \( TR \), corresponds to \( \Delta \) in Mansson’s model which is the time duration for the replenishment of \( \text{Hp}^{129}\text{Xe} \). \( \tau_1, S_0, S_1 \) were the fitting parameters for the healthy animal respectively. Thus, \( M(TR=30 \text{ ms}) \) and \( M(TR=4 \text{ ms}) \) were found to be 0.5 and 0.1 respectively within a measurement uncertainty of \( \pm 5\% \). In other word, \( M(TR) \) represents the percentage concentration of \( \text{Hp}^{129}\text{Xe} \) molecules in the pulmonary tissue pool at the end of each TR period and independent from the use of excitation flip angle.

Extending to the \( n^{\text{th}} \) excitation, the dissolved signal is given by the following:

\[
S_{PT} = M_{PT} \left[ M(TR) \left[ \cos(\alpha_G) \right]^{n-1} \sin(\alpha_{PT}) + \left[ \cos(\alpha_{PT}) \right]^{n-1} \sin(\alpha_{PT}) \right]
\]

where \( M_{PT} \) term represents the total contribution from polarization \( M_i \), solubility of xenon \( \lambda \), and other imaging parameters (e.g., resolution, coil sensitivity etc.). The first term is the signal contribution from the kinetics of gas exchange including the effect of the flip angle on the gas phase pool and the pulmonary tissue phase pool. The second term is magnetization in the pulmonary tissue phase, which was left over after the \( n^{\text{th}}-1 \) excitation. Additionally, RF pulse width is short (\( <<1 \text{ ms} \)) with respect to the relevant exchange times (tens to hundreds of ms). Therefore, exchange during the application of the RF pulse is ignored.
Appendix D: Animal Use Protocol Approval

2007-083-09

AUP Number: 2007-083-09
AUP Title: Assessment of Lung Function and Structure in Anaesthetized and Ventilated Rats Using Hyperpolarized Noble Gas Magnetic Resonance Imaging and Xenon-enhanced Computed Tomography

Yearly Renewal Date: 11/01/2013

The YEARLY RENEWAL to Animal Use Protocol (AUP) 2007-083-09 has been approved, and will be approved for one year following the above review date.

1. This AUP number must be indicated when ordering animals for this project.
2. Animals for other projects may not be ordered under this AUP number.
3. Purchases of animals other than through this system must be cleared through the ACVS office.

Health certificates will be required.

REQUIREMENTS/COMMENTS

Please ensure that individual(s) performing procedures on live animals, as described in this protocol, are familiar with the contents of this document.

The holder of this Animal Use Protocol is responsible to ensure that all associated safety components (biosafety, radiation safety, general laboratory safety) comply with institutional safety standards and have received all necessary approvals. Please consult directly with your institutional safety officers.

Submitted by: Kinchlea, Will D
on behalf of the Animal Use Subcommittee
University Council on Animal Care
Curriculum Vitae

DEGREES

Ph.D. Candidate
September 2011 to present
Medical Biophysics
University of Western Ontario, London, ON, CANADA

M.Sc.
August 2010
Biomedical Physics
Ryerson University, Toronto, ON, CANADA
Thesis: Monitoring electric field-induced changes in biological tissues by using ultrasound.

B.Sc.
June 2007
Physics
Ege University, Izmir, TURKEY
Thesis: Sub-Nuclear Particles, Weak Interactions and Unitary Symmetry U(1), SU(2), SU(3).

PROFESSIONAL EXPERIENCE

Teaching Assistant

Western University, Department of Mathematics, London, ON., from Sep 2014 to Apr 2015.
Duties: Tutorial, invigilation, and marking for following undergrad courses:
   Calculus I (1000a)
   Applied Math (1413)

Ryerson University, Department of Physics, Toronto, ON., from Sep 2008 to May 2011.
Duties: Tutorial, laboratory instruction, invigilation, and marking for following undergrad and graduate courses:
   Medical Imaging (PCS405)
   Solid State Physics (PCS224)
   Physics I and II (PCS130)
Research Assistant

Western University, Imaging Department, London, ON., from Jun 2011 to Sep 2014.
Duties: MR technical support, RF coils design, MR sequence design, animal preparation and surgery, etc.

Ryerson University, Ultrasound Imaging, Toronto, ON., from Sep 2010 to Jun 2011.
Duties: Design magneto acoustic electrical signal experiments and analyze data, design single element ultrasound experiments, collect and analyze data, etc.

PUBLICATIONS

Refereed Journal Articles

Submitted and/or Accepted (pending revisions)

Published or In Press

Published Proceedings of Conferences
Refereed Proceedings of Conferences:


AWARDS

1. CIHR in Cancer Research and Technology Transfer (CaRTT), London, ON. Amount: $19100/year, Sep 2013 – Present.