3rd JSPFI & 5th IWPFI

The Interaction between Medical Engineering and Clinical Medicine

January 28(Fri)-30(Sun), 2011
Awaji Island, Japan

Venue
Awaji Yumebutai International Conference Center

President
Yoshiharu Ohno, M.D., Ph.D.
Joint Meeting combining
The 3rd meeting of the Japanese Society of Pulmonary Functional Imaging and 5th International Workshop of Pulmonary Functional Imaging

“The Interaction between Medical Engineering and Clinical Medicine”

Date: January 28 [Fri] - 30 [Sun], 2011
Venue: Awaji Yumebutai International Conference Center, Hyogo, Japan
1 Yumebutai, Awaji City, Hyogo 656-2306 Japan
Phone: +81-799-74-1020 Fax: +81-799-74-1021

President: Yoshiharu Ohno, M.D., Ph.D
(Department of Radiology, Kobe University Graduate School of Medicine)

Secretariat of 3rd JSPFI & 5th IWPFI
C/o Department of Radiology, Kobe University Graduate School of Medicine
7-5-2 Kusunoki-cho, Chuo-ku, Kobe, 650-0017, Japan
Phone: +81-78-382-6104 Fax: +81-78-382-6129
E-mail:jspfi3@med.kobe-u.ac.jp
Secretariat Chair: Daisuke Takenaka M.D.
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Dear Colleagues and Friends,

It is our great pleasure to invite you to a joint meeting, combining the 3rd meeting of the Japanese Society of Pulmonary Functional Imaging and the 5th International Workshop for Pulmonary Functional Imaging, scheduled for January 28-30, 2011. This joint meeting aims to serve emerging clinical needs for improved imaging diagnostics for various lung diseases by bringing together a multidisciplinary group of researchers and clinicians working at the forefront of image-based approaches to lung function analysis. The scientific program offers an attractive combination of invited keynote lectures, selected scientific papers and poster presentations on several specific topics.

The inaugural meeting of the Japanese Society of Pulmonary Functional Imaging in Kyoto in 2009 was followed by a second meeting in Okinawa in 2010, while the inaugural meeting of the International Workshop for Pulmonary Functional Imaging, was started in Philadelphia in 2002 and followed up with meetings in Awaji in 2004, in Heidelberg in 2006 and in Boston in 2009. This time, we plan to hold the 1st joint meeting of the two organizations, one academic and the other concerned with clinical activities in this field, at the Awaji Yumebutai International Conference Center and Westin Hotel, where the 2nd International Workshop for Pulmonary Functional Imaging was held in 2004. As you may know, Awaji Island is located just off the coast of Honshu about 20 kilometers west of Kobe, 40 kilometers west of Osaka and 60 kilometers west of Kyoto.

We cordially invite you to join us on this beautiful island where you can experience warm hospitality as well as both modern and traditional Japanese culture while sharing and adding to your knowledge of Pulmonary Functional Imaging. We are expecting a large number of researchers, physicians and students at this exciting meeting. We look forward to seeing you on Awaji Island in January, 2011.

Yoshiharu Ohno, M.D., Ph.D.
President of the Organizing Committee
Joint Meeting combining
The 3rd meeting of the Japanese Society of Pulmonary Functional Imaging and 5th International Workshop for Pulmonary Functional Imaging
By Car

You can access Awaji Yumebutai in:
- 30 min. from Kobe (Sannomiya) - 30 km
- 60 min. from Osaka (Umeda) - 60 km
- 90 min. from Kansai International Airport (via the Hanshin Highway Wangan Line and Akashi Kaikyo Bridge) - 100 km
- 50 min. from Osaka International Airport (via the Chugoku Expressway, Hanshin Highway Kitakobe Line, and Akashi Kaikyo Bridge) - 75 km
- 45 min. from Kobe Airport - 45 km
- 70 min. from Tokushima Airport - 85 km
- 5 min. along Route 28 from Awaji IC exit or Higashiura IC exit on Kobe Awaji Naruto Expressway.
Floor Guide

- Awaji Yumebutai -

Route from Conference Center

- The Westin Awaji Island -

Connect to International Conference Center

Joint Meeting Dinner on January 29

Board Meeting of IWPFI on January 29
- Awaji Yumebutai International Conference Center -
Information for Participants

1. Registration

On-site Registration
Registration Desk
Registration Desk will be opened the following hours at the meeting venue.

---Open Hours---
17:00-18:00 Thursday, January 27, 2011
8:00-18:30 Friday, January 28, 2011
8:00-18:30 Saturday, January 29, 2011
8:00-14:30 Sunday, January 30, 2011

1) Registration fee (Only Japanese Yen)

<table>
<thead>
<tr>
<th></th>
<th>On-site Registration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>JSPFI Member</td>
</tr>
<tr>
<td>Medical Doctor</td>
<td>¥28,000</td>
</tr>
<tr>
<td>Non-Medical Doctor</td>
<td>¥25,000</td>
</tr>
<tr>
<td>Student* (Includes Graduate Student)</td>
<td>¥15,000</td>
</tr>
</tbody>
</table>

JSPFI: Japanese Society for Pulmonary Functional Imaging

※JSPFI member except student must pay the annual membership fee 2,000yen. The annual membership fee is able to pay at the Registration Desk.

*Student(Includes Graduate student) : Student participants are required to present a valid student ID upon the Registration Desk.

The all type of registration fee includes: 1. Admission to scientific programs
2. Admission to the exhibition

2) Payment Method (Only Japanese Yen)
Payment must be made in Japanese Yen, by cash or with a credit card at Registration Desk. Please note that neither personal checks nor any other currencies will be accepted.

Credit card: American Express, Visa, MasterCard, Diners Club, and JCB are acceptable.
2. Name Badges
Please wear your name badges at all times during the meeting for identification and security purposes. Only registered participants wearing a name badge will be allowed access to the session rooms, exhibition and social programs.

3. Commercial Exhibition
Commercial Exhibition will be held at the “Reception Hall B” during the meeting. Exhibitors will be eager to demonstrate and explain their latest products, and answer your question.

4. Internet Access Service & Drink Service
Internet access service and drink service are available at the “Reception Hall B Foyer”.

5. Travel Desk
Travel Desk will be opened at the venue operated by JTB.

6. Mobile Phone
During the session, you are prohibited from using a mobile phone. Please turn off or switch to the silent mode.

7. Message Board
To prevent from interrupting the sessions, we do not call up anyone in the meeting venue. Please use message board. Leave your message for your friends and colleagues and look for messages left for you.

8. Cloakroom
The Cloakroom is on 2F near by Front Desk of the Westin:

9. Be warned against pickpockets or stolen
Please keep valuables in your possession at all times. The meeting cannot responsible for lost or stolen items.

10. Social Events
The organizing committee takes pleasure in presenting the following social events that offer all participants and accompanying persons.

Welcome Reception

Joint Meeting Dinner at The Westin
19:30-21:30, Saturday, January 29, 2011 at The Westin Awaji Island, “STELLA”
Instruction for Speakers and Moderators / Discussers

Instruction for Speakers

Presentation Time (except Invited Speakers)

<table>
<thead>
<tr>
<th>Session</th>
<th>Presentation(min.)</th>
<th>Discussion(min.)</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Core Session</td>
<td>7</td>
<td>3</td>
<td>PC</td>
</tr>
<tr>
<td>Scientific Presentation</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Poster Session</td>
<td>5</td>
<td>4</td>
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</tr>
</tbody>
</table>

For Invited Speakers: Please follow the instruction for presentation and discussion time from secretariat and moderators.

Oral Presentation Information
(Special Lecture, Luncheon Lecture, Evening Lecture, Core Session)

Equipment: Computers

<table>
<thead>
<tr>
<th>OS</th>
<th>Software for Presentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Windows 7</td>
<td>PowerPoint ver. 2000 to 2010</td>
</tr>
<tr>
<td>Mac OSX10.5</td>
<td>PowerPoint ver. 2004 &amp; 2008, Keynote ver. 5</td>
</tr>
</tbody>
</table>

All meeting rooms are equipped with the following audio-visual equipment:

Preview Center (2F, “Foyer”)

---Open Hours---
17:00-18:00  Thursday, January 27, 2011
8:00-18:30   Friday, January 28, 2011
8:00-18:30   Saturday, January 29, 2011
8:00-14:30   Sunday, January 30, 2011

• All Speakers (except Poster Session) are requested to come to the Preview Center at least 30 minutes in advance of their presentations to verify that the data functions properly on the equipment provided.

• If you bring your own laptop PC, after checking-in at the Preview Center, please come to the PC operation desk in front of the podium located at the left-front in the session room by 15 minutes prior to your presentation and hand your PC to the staff. (You can come to the PC operation desk while the previous session in proceeding.)

PowerPoint Presenters

• Bring your presentation in a USB flash Drive or CD-ROM. In case you use animation, you should to bring your own laptop PC.
• Make sure that you close or “finalize” your presentation file when create a CD. If you omit this step, you cannot access the CD from any other computers.
• Only the standard font (e.g., Times New Roman, Arial, Century) will be available. Unusual fonts may not be displayed properly on the computers in session rooms.
• Please name the file as: “Date of presentation_presenter’s name, ppt(x)” e.g., 0128_TaroAWAJI.pptx
• In order to avoid virus infection, please scan your data with updated anti-virus software beforehand.
• Please note that you cannot make any modification at the Preview Center and session room.
• After the meeting, all presentation data will be erased by secretariat.

Laptops

• Speakers using their own laptops must have a Mini D-sub 15pin female output. Special video output cable is required for some laptops to use Mini D-sub 15pin to connect to external monitors and data projectors. Please note that we are not equipped with that special cable and you must bring it in case it is necessary.

---Mini D-sub 15pin---

• Please be sure to bring the power adapter.
Poster Presentation Information

Guidelines for Poster Presenters:

Presenters are requested to follow the schedule below in mounting poster on their assigned board. The poster number for your presentation can be found in the program book. Please follow the instructions provided.

- Schedule

<table>
<thead>
<tr>
<th>Date</th>
<th>Mounting</th>
<th>Poster Viewing</th>
<th>Presentation</th>
<th>Removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thursday January 27, 2011</td>
<td>17:00-18:00</td>
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</tr>
<tr>
<td>Friday January 28, 2011</td>
<td>8:00-11:00</td>
<td>11:00-12:00</td>
<td>17:40-18:52 Last</td>
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</tr>
<tr>
<td>Saturday January 29, 2011</td>
<td>/</td>
<td>11:00-12:00</td>
<td>17:40-18:52 Last</td>
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<tr>
<td>Sunday January 30, 2011</td>
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<td>/</td>
<td>8:00-15:00</td>
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</tbody>
</table>

Poster Board Information

- Your poster number will already be on your assigned board.
- Please prepare a label showing the title, institution (includes country) and the author’s name. You cannot use pins or adhesive tape.
- Hook - and - loop fastener for mounting will be available from the Poster Reception Desk at the Poster Session Area.
- Poster should be brought by yourself to the meeting and not mailed, as the organizing committee cannot be responsible for loss or mishandling.
- Presenters are responsible for posting and removing their own materials.
- Audio-visual equipment may not be used. Please refer to the poster image for your poster.

The Best Poster Award

This award will be established committee of the meeting to encourage the high performance of the presentation in the poster session. The winner will be selected during the poster presentation time by the moderators.

For the winner

Accommodation ceremony by the president of the meeting during the Joint Meeting Dinner at The Westin, 19:30-, January 28.

Instructions for Moderators and Discussers

All moderators and discussers are asked to ensure that each sessions start on time and finish punctually as scheduled.

Oral Sessions

- Please come to the “Next moderator”’s seat of the session room (the front row on your right side) no later than 15 minutes prior to the beginning of the session and identify yourself to the staff.

Poster Sessions

- All moderators for poster session are requested to come to the “Poster Reception Desk” near by Poster Session Area no later than 15 minutes prior to the beginning of the session.
- Please select the best poster presentation of the session (one person per session), and tell the staff of “Poster Reception Desk”.

# Time Table

## January 27, 2011 [Thu]

<table>
<thead>
<tr>
<th>Time</th>
<th>Room A</th>
<th>Room B</th>
<th>Poster Exhibition</th>
<th>Commercial Exhibition</th>
<th>Receptions</th>
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</thead>
<tbody>
<tr>
<td>16:00</td>
<td>Main Hall (2F)</td>
<td>Event Hall (B1)</td>
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<td>17:00</td>
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<tr>
<th>Time</th>
<th>Room A</th>
<th>Room B</th>
<th>Poster Exhibition</th>
<th>Commercial Exhibition</th>
<th>Receptions</th>
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</thead>
<tbody>
<tr>
<td>8:00</td>
<td>Main Hall (2F)</td>
<td>Event Hall (B1)</td>
<td>Lobby (B1)</td>
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<td>9:00</td>
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## January 28, 2011 [Fri]

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<tr>
<th>Time</th>
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<th>Commercial Exhibition</th>
<th>Receptions</th>
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<tbody>
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<td>8:00</td>
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<td>Event Hall (B1)</td>
<td>Lobby (B1)</td>
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<tr>
<td>9:00</td>
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<td>9:00-18:00 Poster Mounting</td>
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<td>10:00</td>
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<th>Time</th>
<th>Room A</th>
<th>Room B</th>
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<th>Commercial Exhibition</th>
<th>Receptions</th>
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<tbody>
<tr>
<td>8:45-9:00</td>
<td>Opening Remark</td>
<td></td>
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<tr>
<td>9:00-10:50</td>
<td>Core Session 1</td>
<td></td>
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</tr>
<tr>
<td>11:00-11:45</td>
<td>Organizer Meeting of JSPFI</td>
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<tr>
<td>12:00-13:00</td>
<td>Luncheon Lecture 1</td>
<td>Functional Imaging for Pediatrics</td>
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<tr>
<td>13:30-14:20</td>
<td>Special Lecture 1</td>
<td>From Morphology to Function and Metabolism</td>
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<tr>
<td>14:30-16:20</td>
<td>Core Session 2</td>
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<tr>
<td>16:30-17:30</td>
<td>Evening Lecture 1</td>
<td>Functional Imaging at High-Field MR System</td>
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<td>17:40-18:52</td>
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</table>

19:30-21:30 Welcome Reception at Botanical Gardens “The Miracle Planet Museum of Plants”
<table>
<thead>
<tr>
<th>Time</th>
<th>Room A</th>
<th>Room B</th>
<th>Poster Exhibition</th>
<th>Commercial Exhibition</th>
<th>Receptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>8:00</td>
<td>Main Hall (2F)</td>
<td>Event Hall (B1)</td>
<td>Lobby (B1)</td>
<td>Reception Hall B (2F)</td>
<td></td>
</tr>
<tr>
<td>8:00-8:40</td>
<td>Board Meeting of IWPFI</td>
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<td></td>
<td>8:00-15:00</td>
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<tr>
<td>9:00-10:50</td>
<td>Core Session 3</td>
<td></td>
<td></td>
<td></td>
<td>9:00-18:00</td>
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<tr>
<td>11:00-11:45</td>
<td>General Meeting of JSPFI</td>
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<tr>
<td>12:00-13:00</td>
<td>Poster Viewing</td>
<td></td>
<td></td>
<td></td>
<td>Commercial Exhibition</td>
</tr>
<tr>
<td>13:20-14:20</td>
<td>Special Lecture 2</td>
<td></td>
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<tr>
<td>14:30-16:15</td>
<td>Core Session 4</td>
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<tr>
<td>17:40-18:52</td>
<td>Last Evening Lecture 2</td>
<td></td>
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<tr>
<td>19:30-21:30</td>
<td>Joint Meeting Dinner</td>
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</table>

**January 30, 2011 [Sun]**

<table>
<thead>
<tr>
<th>Time</th>
<th>Room A</th>
<th>Room B</th>
<th>Poster Exhibition</th>
<th>Commercial Exhibition</th>
<th>Receptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>8:00</td>
<td>Main Hall (2F)</td>
<td>Event Hall (B1)</td>
<td>Lobby (B1)</td>
<td>Reception Hall B (2F)</td>
<td></td>
</tr>
<tr>
<td>8:30-10:20</td>
<td>Core Session 5</td>
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<tr>
<td>10:30-12:15</td>
<td>Core Session 6</td>
<td></td>
<td></td>
<td>Poster Removal</td>
<td>Commercial Exhibition</td>
</tr>
<tr>
<td>12:30-13:30</td>
<td>Luncheon Lecture 3</td>
<td></td>
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<tr>
<td>13:50-15:05</td>
<td>Special Lecture 3</td>
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<tr>
<td>15:05-15:15</td>
<td>Closing Remark</td>
<td></td>
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</tr>
</tbody>
</table>
Special Lecture


1. From Morphology to Function and Metabolism

Moderator: Tadashi Abe  Tokai University
           Kiyoshi Murata  Shiga University of Medical Science
           Edwin J.R. van Beek  University of Edinburgh

13:20-13:50  Interaction between Functional and Morphological Imaging:
             3D Airway Analysis in Chronic Obstructive Pulmonary Disease
           Invited Speaker 1: Masaharu Nishimura  Hokkaido University

13:50-14:20  Interaction between Metabolic and Morphological Imaging
           Invited Speaker 2: Kyung Soo Lee  Samsung Medical Center,
                                Sungkyunkwan University School of Medicine

January 29 [Sat]  13:20-14:20  Sponsored by PHILIPS

2. Basics and Future Direction for Quantitative Functional CT Assessment

Moderator: Michiaki Mishima  Kyoto University
           Kenya Murase  Osaka University
           James P. Butler  Harvard Medical School

             ~Basics of Interpretation of HRCT of the Lung~
           Invited Speaker 1: Harumi Itoh  University of Fukui

13:50-14:20  Current Status and Future Direction of Quantitative Analysis for Functional CT
           Invited Speaker 2: Eric A. Hoffman  University of Iowa Carver College of Medicine

January 30 [Sun]  13:50-15:05  Sponsored by TOSHIBA MEDICAL SYSTEMS CORPORATION

3. Future Direction for Pulmonary Functional and Molecular Imaging

Moderator: Yasutaka Nakano  Shiga University of Medical Science
           Yoshiharu Ohno  Kobe University
           Hiroto Hatabu  Brigham and Women’s Hospital, Harvard Medical School

13:50-14:15  From Basic Science Side
           Invited Speaker 1: Hisataka Kobayashi  National Institute of Health

14:15-14:40  From Radiology Side
           Invited Speaker 2: Hans-Ulrich Kauczor  Heidelberg University Medical Center

14:40-15:05  From Pulmonology Side -Terminal Air Space of X-ray CT Images in COPD-
           Invited Speaker 3: Michiaki Mishima  Kyoto University
Luncheon Lecture

January 28 [Fri]  12:00-13:00  Sponsored by TERUMO

1. Functional Imaging for Pediatrics
   Moderator: Atsushi Nagai  Tokyo Women’s Medical University

   12:00-12:30  Functional Thoracic CT for Pediatric Patients
      Invited Speaker 1: Hyun Woo Goo  Asan Medical Center, University of Ulsan College of Medicine

   12:30-13:00  Pediatric Applications of Lung MRI
      Invited Speaker 2: Talissa A. Altes (Presented by Lucia Flors)
      University of Virginia Medical Center

January 29 [Sat]  12:00-13:00  Sponsored by Astellas Pharma Inc. / AstraZeneca K.K.

2. Imaging for Asthmatics and COPD
   Moderator: Shoji Kudoh  Double-Barred Cross Hospital

   12:00-12:30  CT Assessment of Asthma
      Invited Speaker 1: Gregory G. King  The Woolcock Institute of Medical Research

   12:30-13:00  Hyperpolarized Noble Gas MRI for Asthma and COPD
      Invited Speaker 2: Mitchell S. Albert  University of Massachusetts Medical School

January 30 [Sun]  12:30-13:30  Sponsored by DAIICHI SANKYO COMPANY, LIMITED

3. Hyperpolarized Noble Gas MRI
   Moderator: Toshihiro Nukiwa  Tohoku University

   12:30-13:00  Development of Polarizer for Future Clinical Works
      Invited Speaker 1: F. William Hersman  University of New Hampshire, Xemed LLC

   13:00-13:30  Hyperpolarized Noble Gas MRI: Clinical Applications
      Invited Speaker 2: Hiroto Hatabu  Brigham and Women’s Hospital, Harvard Medical School

Evening Lecture

January 28 [Fri]  16:30-17:30  Sponsored by Bayer Yakuhin, Ltd.

1. Functional Imaging at High-Field MR System
   Moderator: Yukihiko Sugiyama  Jichi Medical University

   16:30-17:00  Animal Studies for Functional Imaging at High-Field System: Ultra-short TE (UTE) MRI of the Lung
      Invited Speaker 1: Masaya Takahashi  University of Texas Southwestern Medical Center at Dallas

   17:00-17:30  Clinical Functional Imaging at High-Field MRI
      Invited Speaker 2: Christian Fink  University Medical Center Mannheim

January 29 [Sat]  16:30-17:30  Sponsored by Eisai Co., Ltd.

2. New Techniques for Pulmonary Vascular Disease
   Moderator: Kazuro Sugimura  Kobe University

   16:30-17:00  Dual-Energy CT Application for Pulmonary Vascular Disease
      Invited Speaker 1: Joon Beom Seo  University of Ulsan College of Medicine, Asan Medical Center

   17:00-17:30  Quantitative MR Assessment for Pulmonary Vascular Disease
      Invited Speaker 2: David L. Levin  Mayo Clinic
Core Session

January 28 [Fri] 9:00-10:50

1. Pulmonary Hypertension

Moderator: Hiroshi Kimura  Nara Medical University
Noriyuki Tomiyama  Osaka University
Discuss: Mark L. Schiebler  University of Wisconsin

9:00-9:20  Etiology and Pathophysiology of Pulmonary Hypertension
Invited Speaker 1: Koichiro Tatsumi  Chiba University

9:20-9:35  Nuclear Medicine Assessment for Pulmonary Hypertension
Invited Speaker 2: Takayuki Shinkai  Nara Medical University

9:35-9:50  CT Assessment of Pulmonary Hypertension
Invited Speaker 3: Joon Beom Seo  University of Ulsan College of Medicine, Asan Medical Center

9:50-10:05  MR Assessment for Pulmonary Hypertension
Invited Speaker 4: David L. Levin  Mayo Clinic

10:05-10:20  Update of Treatment for Pulmonary Hypertension
Invited Speaker 5: David G. Kiely  Royal Hallamshire Hospital

10:20-10:50  Scientific Presentation 1-3

January 28 [Fri] 14:30-16:20

2. Thoracic Malignancy: Lung Cancer Treatment

Moderator: Michio Kono  Kobe University
Yoichi Nakanishi  Kyushu University
Discuss: Sadayuki Murayama  University of the Ryukyus

14:30-14:50  Newest Evidences of Drug Therapy for Lung Cancer
Invited Speaker 1: Yoichi Nakanishi  Kyushu University

14:50-15:05  Update for New Molecular Targeted Therapy for Lung Cancer
Invited Speaker 2: Kazuhisa Takahashi  Juntendo University School of Medicine

15:05-15:20  Molecular Imaging for Lung Cancer Treatment
Invited Speaker 3: Koji Murakami  Keio University

15:20-15:35  MR Imaging for Lung Cancer Treatment
Invited Speaker 4: Juergen Biederer  University Hospital Schleswig-Holstein, Campus Kiel

15:35-15:50  CAD for Lung Cancer Treatment –Evaluation of Tumor Response to Treatment
Invited Speaker 5: Jin Mo Goo  Seoul National University Hospital

15:50-16:20  Scientific Presentation 1-3
### January 29 [Sat] 9:00-10:50

#### 3. Chronic Obstructive Pulmonary Disease

Moderator : Toyohiro Hirai  
Kyoto University  
Kiminori Fujimoto  
Kurume University School of Medicine  
Discusser : Hans-Ulrich Kauczor  
Heidelberg University

9:00-9:20 **Etiology and Pathophysiology of COPD**  
Invited Speaker 1 : Kazutetsu Aoshiba  
Tokyo Women’s Medical University

9:20-9:35 **PET Assessment for COPD: Changes in Pulmonary Perfusion Distribution**  
Invited Speaker 2 : José G. Venegas  
Harvard Medical School, Massachusetts General Hospital

9:35-9:50 **CT Assessment for COPD-Phenotype on CT in COPD**  
Invited Speaker 3 : Keisaku Fujimoto  
Shinshu University School of Health Sciences

9:50-10:05 **Proton MR Assessment of COPD**  
Invited Speaker 4 : Yoshiharu Ohno  
Kobe University

10:05-10:20 **Hyperpolarized Noble Gas MRI for COPD**  
Invited Speaker 5 : Edwin J.R. van Beek  
University of Edinburgh

10:20-10:50 **Scientific Presentation 1-3**

### January 29 [Sat] 14:30-16:15

#### 4. Interstitial Lung Disease

Moderator : Yukihiko Sugiyama  
Jichi Medical University  
Masashi Takahashi  
Shiga University of Medical Science  
Discusser : Kyung Soo Lee  
Samsung Medical Center,  
Sungkyunkwan University School of Medicine

14:30-14:55 **Update for Diagnosis of IIP**  
Invited Speaker 1 : Takeshi Johkoh  
Kinki Central Hospital of Mutual Aid Association of  
Public School Teachers

14:55-15:15 **Update for Treatment of Idiopathic Interstitial Pneumonias (IIPs)**  
Invited Speaker 2 : Kingo Chida  
Hamamatsu University School of Medicine

15:15-15:30 **Smoking-related Interstitial Pneumonia Associated with Emphysema**  
Invited Speaker 3 : Minoru Kanazawa  
Saitama Medical University

15:30-15:45 **Hyperpolarized Noble Gas MRI for ILD**  
Invited Speaker 4 : Hiroto Hatabu  
Brigham and Women’s Hospital, Harvard Medical School

15:45-16:15 **Scientific Presentation 1-3**
### January 30 [Sun] 8:30-10:20
#### 5. Asthma and Airway Disease

**Moderator:** Yuji Tohda  Kinki University  
Shigeru Kosuda  National Defense Medical College  

**Discuss:** Shu Hashimoto  Nihon University  

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Speaker(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>8:30-8:50</td>
<td>Pathophysiology of Asthma</td>
<td>Invited Speaker 1: Hiromasa Inoue  Kagoshima University</td>
</tr>
<tr>
<td>8:50-9:05</td>
<td>Evaluation of Airway Obstruction and Altered Pulmonary Permeability with Lung Scintigraphy in Young Patients</td>
<td>Invited Speaker 2: Mayuki Uchiyama  The Jikei University School of Medicine</td>
</tr>
<tr>
<td>9:05-9:20</td>
<td>CT Assessment for Asthma</td>
<td>Invited Speaker 3: Hiroshi Tanaka  Sapporo Medical University School of Medicine</td>
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<tr>
<td>9:20-9:35</td>
<td>MR Assessment of Asthma; Hyperpolarized $^3$He Magnetic Resonance Imaging to Probe Temporal and Spatial Airway Functional Changes</td>
<td>Invited Speaker 4: Grace Parraga  The University of Western Ontario</td>
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<tr>
<td>9:35-9:50</td>
<td>Update for Treatment of Asthma</td>
<td>Invited Speaker 5: Kazuto Hirata  Osaka City University</td>
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<tr>
<td>9:50-10:20</td>
<td>Scientific Presentation 1-3</td>
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### January 30 [Sun] 10:30-12:15
#### 6. Respiratory Motion Analysis

**Moderator:** Ichiro Kuwahira  Tokai University Tokyo Hospital  
Shuji Sakai  Tokyo Women’s Medical University  

**Discuss:** James Gee  University of Pennsylvania  

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Speaker(s)</th>
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</thead>
<tbody>
<tr>
<td>10:30-10:45</td>
<td>Aims of Respiratory Motion Assessment for Anesthesiology</td>
<td>Invited Speaker 1: Klaus Markstaller  Medical University of Vienna</td>
</tr>
<tr>
<td>10:45-11:00</td>
<td>MR Assessment of Respiratory Motion</td>
<td>Invited Speaker 2: Tae Iwasawa  Kanagawa Cardiovascular and Respiratory Center</td>
</tr>
<tr>
<td>11:00-11:15</td>
<td>Nuclear Medicine Assessment for Respiratory Lung Motion</td>
<td>Invited Speaker 3: Kazuyoshi Suga  Yamaguchi University School of Medicine</td>
</tr>
<tr>
<td>11:15-11:30</td>
<td>Dynamic Volume Scan Using 320-row ADCT for the Assessment of Respiratory Motion</td>
<td>Invited Speaker 4: Hiroshi Moriya  Sendai Kousei Hospital</td>
</tr>
<tr>
<td>11:30-11:45</td>
<td>Respiratory Motion Assessment to Increase the Accuracy of Non-contact Screening System to Prevent the Spread of Infectious Diseases</td>
<td>Invited Speaker 5: Takemi Matsui  Tokyo Metropolitan University</td>
</tr>
<tr>
<td>11:45-12:15</td>
<td>Scientific Presentation 1-3</td>
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Program

January 28, 2011 [Fri]
Room A
Main Hall (2F)

8:45-9:00  Opening Remark

9:00-10:50  Core session 1
Pulmonary Hypertension

Moderator  Hiroshi Kimura  Nara Medical University
           Noriyuki Tomiyama  Osaka University

Discussser  Mark L. Schiebler  University of Wisconsin

Invited Speaker

9:00-9:20  1. Etiology and Pathophysiology of Pulmonary Hypertension
           Koichiro Tatsumi  Chiba University

9:20-9:35  2. Nuclear Medicine Assessment for Pulmonary Hypertension
           Takayuki Shinkai  Nara Medical University

9:35-9:50  3. CT Assessment of Pulmonary Hypertension
           Joon Beom Seo  University of Ulsan College of Medicine, Asan Medical Center

9:50-10:05 4. MR Assessment for Pulmonary Hypertension
           David L. Levin  Mayo Clinic

10:05-10:20 5. Update of Treatment for Pulmonary Hypertension
           David G. Kiely  Royal Hallamshire Hospital

Scientific Presentation

10:20-10:30 1. Optimized Flowchart of Multi-technical MR Protocol for Pulmonary Circulation and Right Heart Function on the Busy Clinical Schedule
           Shu-Hui Peng  Chung Shan Medical University Hospital

10:30-10:40 2. V-Q Imbalance in Primary/Passive Pulmonary Hypertension on V/Q SPECT
           Kazuyoshi Suga  St. Hill Hospital

10:40-10:50 3. Estimation of Pulmonary Vascular Resistance by the Calculation of Contrast Transit Time from Gadolinium Contrast Enhanced Pulmonary MRI
           Adam G Telfer  University of Sheffield
13:20-14:20  Special Lecture 1  Sponsored by Siemens Japan K.K.

From Morphology to Function and Metabolism

Moderator  Tadashi Abe  Tokai University
Kiyoshi Murata  Shiga University of Medical Science
Edwin J.R. van Beek  University of Edinburgh

13:20-13:50
1. Interaction between Functional and Morphological Imaging:
   3D Airway Analysis in Chronic Obstructive Pulmonary Disease

   Masaharu Nishimura
   Hokkaido University

13:50-14:20
2. Interaction between Metabolic and Morphological Imaging

   Kyung Soo Lee
   Samsung Medical Center, Sungkyunkwan University School of Medicine
14:30-16:20  **Core session 2**

**Thoracic Malignancy: Lung Cancer Treatment**

**Moderator**  Michio Kono  Kobe University
Yoichi Nakanishi  Kyushu University

**Discuss** Sadayuki Murayama  University of the Ryukyus

---

**Invited Speaker**

14:30-14:50

1. Newest Evidences of Drug Therapy for Lung Cancer  
   Yoichi Nakanishi  Kyushu University

14:50-15:05

2. Update for New Molecular Targeted Therapy for Lung Cancer  
   Kazuhisa Takahashi  Juntendo University School of Medicine

15:05-15:20

3. Molecular Imaging for Lung Cancer Treatment  
   Koji Murakami  Keio University

15:20-15:35

4. MR Imaging for Lung Cancer Treatment  
   Juergen Biederer  University Hospital Schleswig-Holstein, Campus Kiel

15:35-15:50

5. CAD for Lung Cancer Treatment - Evaluation of Tumor Response to Treatment  
   Jin Mo Goo  Seoul National University Hospital

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**Scientific Presentation**

15:50-16:00

1. Comparative Reading System for Lung Cancer CT Screening  
   Hidenobu Suzuki  The University of Tokushima

16:00-16:10

2. The Evaluation of Lung Perfusion in a Radiation Field Through the Radiation Therapy for Lung Cancer - Using Lung Perfused Blood Volume Imaging with Dual Energy CT-  
   Sachiko Miura  Nara Medical University

16:10-16:20

3. Therapy Response Assessment in Neoadjuvant Chemotherapy for Esophageal Cancer: Comparison of PET Response Criteria in Solid Tumors (PERCIST) and Response Evaluation Criteria in Solid Tumors (RECIST)  
   Masahiro Yanagawa  Osaka University
12:00-13:00  Luncheon Lecture 1
Functional Imaging for Pediatrics
Moderator  Atsushi Nagai  Tokyo Women's Medical University

12:00-12:30
1. Functional Thoracic CT for Pediatric Patients
Hyun Woo Goo
Asan Medical Center, University of Ulsan College of Medicine

12:30-13:00
2. Pediatric Applications of Lung MRI
Talissa A. Altes
(Presented by Lucia Flors)
University of Virginia Medical Center

16:30-17:30  Evening Lecture 1
Functional Imaging at High-Field MR System
Moderator  Yukihiko Sugiyama  Jichi Medical University

16:30-17:00
Masaya Takahashi
University of Texas Southwestern Medical Center at Dallas

17:00-17:30
2. Clinical Functional Imaging at High-Field MRI
Christian Fink
University Medical Center Mannheim
Poster Session
Event Hall Lobby (B1)

17:40-18:25  Poster Session 1
Pulmonary Circulation 1
Moderator  Toshiko Hoshi  Saitama Cardiovascular and Respiratory Center
Nobuhiro Tanabe  Chiba University

P1-1. Follow-up Cases with Perfusion Scintigraphy in Patients with Pulmonary Thromboembolism
Shigeru Tominaga
Juntendo University Urayasu Hospital

P1-2. Most Optimum Display Condition(Height,Center) of Dual Energy Perfusion CT for Diagnosing Pulmonary Embolism
Tomohiro Suzuki
Nagoya City University

P1-3. Pulmonary CT Perfusion Imaging in Assessment of Pulmonary Vascular Disease -First Experience Using 320-MDCT.
Edwin J.R. van Beek
University of Edinburgh

P1-4. Reevaluation of the Tc-99m MAA Whole Body Imaging for Detecting Right-to-left Shunt
Takayuki Shinkai
Nara Medical University School of Medicine

P1-5. Nonembolic Causes of Perfusion Defect on Contrast-enhanced Dual-Energy CT: Review of 85 Consecutive Cases
Chae-Hun Lim
University of Ulsan College of Medicine, Asan Medical Center

17:40-18:52  Poster Session 2
Chronic Obstructive Pulmonary Disease
Moderator  Shin Matsuoka  St. Marianna University School of Medicine
Hiroshi Kimura  Nara Medical University

P2-1. Correlation between Airway Dimensions and Emphysema Scores in Patients with Chronic Obstructive Pulmonary Disease: Patient-based versus Lobe-based Analysis
Ji-Young Ko
Seoul National University Hospital

P2-2. Comparison of Airway Remodelling Assessed by Three-dimensional Computed Tomography between Bronchial Asthma and Chronic Obstructive Pulmonary Disease
Kaoruko Shimizu
Hokkaido University School of Medicine

P2-3. Airway Wall Remodeling in Different Anatomic Locations Assessed by Computed Tomography in Patients with Stable Asthma and COPD
Tsuyoshi Oguma
Kyoto University
P2-4. Quantitative Assessment of Chronic Obstructive Pulmonary Disease (COPD) with Multi-detector row CT (MDCT): Comparison with Lung Structure and Pulmonary Function Test

Jun Sato
Iwata City Hospital

P2-5. Quantitative Assessment of Low Attenuation Volume Percentage(LAV%) and Mean Lung Density(MLD) in Chronic Obstructive Pulmonary Disease Using Inspiratory and Expiratory MDCT

Naoko Kawata
Chiba University

P2-6. Relationship between Low Attenuation Areas Evaluated by Thurlbeck Panel Method and Body Weight in Patients with Stable COPD

Takanobu Shioya
Akita University

P2-7. Effects of Chronic Obstructive Pulmonary Disease on Thoracic Vascular Calcifications

Bertram J Jobst
University of Heidelberg

P2-8. Analysis of Ventilation Abnormalities in Patients with Obstructive Lung Diseases on Xenon-enhanced CT According to Lung Lesion Patterns: Multicenter Study

Eun Jin Chae
Ulsan College of Medicine, Asan Medical Center

17:40-18:43 Poster Session 3

Diffuse Lung Disease
Moderator
Nobuyuki Tanaka
Yamaguchi University School of Medicine
Takashi Hajiro
Tenri Hospital

P3-1. Evaluation of the Extent of Ground-glass Opacity on High-Resolution CT in Patients with Interstitial Pneumonia: Comparison between Quantitative and Qualitative Analysis

Hidetake Yabuuchi
Kyushu University

P3-2. Correlation between Pirfenidone Effect and the Initial CT Findings in the Patients with Idiopathic Pulmonary Fibrosis

Tae Iwasawa
Kanagawa Cardiovascular and Respiratory Center

P3-3. Relation between Radiological Abnormalities and Functional Impairment Assessed from Distributional Analysis on Ventilation and Diffusing Capacity to Perfusion in Idiopathic Interstitial Pneumonias

Kazuhiro Yamaguchi
Tokyo Women's Medical University Medical Center East

P3-4. Clinical Considerations of Postoperative Acute Exacerbation in Patients with Interstitial Pneumonia Combined with Lung Cancer

Takahiro Omori
Kanagawa Cardiovascular and Respiratory Center
P3-5. Thoracic and Abdomino-pelvic Radiologic Findings of Japanese Patients with Lymphangioleiomyomatosis

Kuniaki Seyama
Juntendo University School of Medicine

P3-6. Clinical and Radiographic Features of Pulmonary Aspergillosis in Patients with Chronic Fibrosing Interstitial Pneumonia

Akiko Kaga
Saitama Medical University

P3-7. Clinical and Radiographic Features of Pneumothorax and Pneumomediastinum in Patients with Chronic Fibrosing Interstitial Pneumonia

Ayako Shiono
Saitama Medical University

17:40-18:34 Poster Session 4

Respiratory Motion 1

Moderator
Masaki Hara Nagoya City University
Keisaku Fujimoto Shinshu University

P4-1. Effect of Pulmonary Rehabilitation on Parenchymal Lung Motion Evaluated with Dynamic MRI in COPD Patients

Takeshi Shinohara
Kanagawa Cardiovascular and Respiratory Center

P4-2. Diaphragmatic Motion before and after Plication of Central Tendon in Patients with Diaphragmatic Eventration

Koji Chihara
Shizuoka City Shizuoka Hospital

P4-3. Dynamic MRI Measured Respiratory Motion of the Internal Structures of the Lung. Differences between Healthy Subjects and COPD Patients.

Hirofumi Shibata
Yokohama City University School of Medicine

P4-4. Respiratory Motion Analysis with Correlation Function and Frequency Decomposition from MR Image Sequences of COPD Patients and Normal Subjects

Yew Kurabayashi
Takasaki University of Commerce

P4-5. Evaluation of Airway Dimension During Respiration

Hiroshi Wada
Shiga University of Medical Science


Norihisa Motohashi
Fukujuji Hospital, JATA
17:40-18:34  Poster Session 5
New MR Technique

Moderators:
Mitchell S. Albert  University of Massachusetts Medical School
Atsoumi Kimura  Osaka University

P5-1.  Hyperpolarized 3He Magnetic Resonance Imaging Biomarkers of Bronchoscopic Airway Bypass in COPD
Lindsay Mathew  Robarts Research Institute

P5-2.  Line-Narrowed Diode Lasers for High-Efficiency Continuous Production of Hyperpolarized Xe-129 using Optimized Cell for High Concentration of Optically Pumped Rubidum
Mineyuki Hattori  Photonics Research Institute, National Institute of Advanced Industrial Science and Technology

P5-3.  Development of a Novel Method for Evaluating Pulmonary Disease Using Hyperpolarized Xe-129 Ultrashort Echo-time MRI
Hirohiko Imai  Osaka University

P5-4.  Non-contrast Enhanced MR Angiography Using Volumetric T2-weighted Imaging Technique on 3T
Mitsutomi Ishiyama  St.Luke’s International hospital

Norio Yoshizaki  Kousei Hospital

P5-6.  Quantitative Analysis of the Lung using Hyperpolarized Helium-3 Imaging
James C. Gee  University of Pennsylvania

17:40-18:43  Poster Session 6
Basic Study 1

Moderators:
Sumiaki Matsumoto  Kobe University
Toyohiro Hirai  Kyoto University

P6-1.  The Phantom Study for Analyzing the Small Airway Disease
Noriyuki Yanagawa  Chiba University

P6-2.  Integral-based Half-Band Method in Measuring Airway Wall Thickness on CT: Comparison with Conventional Full-Width Half Maximum Method in Actual and Simulated Phantoms
Kim, Namkug  Ulsan University College of Medicine, Asan Medical Center
P6-3. Enhancement of Computerized Differentiation Accuracy of Region Pattern of Diffuse Infiltrative Lung Disease Using Local Context Sensitive SVM Classifier
Kim, Namkug
Ulsan University College of Medicine, Asan Medical Center

P6-4. Newly Developed Denoising Technique: Utility for Quantitative Assessment of Pulmonary Emphysema on Low-Dose CT Examination
Mizuho Nishio
Kobe University

P6-5. Comparative Study of Respiratory Function for Pulmonary Lobectomy Based on Computer Simulation
Yasushi Hirano
Yamaguchi University

P6-6. Lung CT Image Registration Using Diffeomorphic Transformation Models
Gang Song
University of Pennsylvania School of Medicine

P6-7. Airway Tree Segmentation by Removing Paths of Leakage
Gang Song
University of Pennsylvania School of Medicine

17:40-18:43 Poster Session 7
Clinical Study
Moderator
Koji Takahashi Asahikawa Medical University
Keisuke Tomii Kobe City Medical Center General Hospital

P7-1. A Low Dose CT Lung Screening Method for Smoking-related Thoracic Abnormalities Using 3D Automatic Exposure Control: Diagnostic Capability and Effectiveness in Radiation Dose Reduction (iLEAD Study)
Takeshi Kubo
Kyoto University

Wolfram Stiller
University Hospital Heidelberg

P7-3. Proposal of New Method: Single Breath Xenon Ventilation CT
Norinari Honda
Saitama Medical Center, Saitama Medical University

P7-4. Relationship between Lung Morphologic and Functional Abnormality Assessed by Breath-Hold Perfusion SPECT-CT Fusion images
Kazuyoshi Suga
St. Hill Hospital

P7-5. Diagnostic Value of Initial Computed Tomography in Critically Ill Patients with Pulmonary Edema
Kosaku Komiya
Oita University Faculty of Medicine
P7-6. Pulmonary Lymphatic Drainage to the Mediastinum Based on CT Observations of Primary Complex of Pulmonary Histoplasmosis

Koji Takahashi
Asahikawa Medical University

P7-7. Evaluation of Cross-bridging Structures in Pneumothorax Patients and Correlation with Aging and Pulmonary Inflammation

Tomoaki Sasaki
Asahikawa Medical University

17:40-18:25  Poster Session 8

Case Report 1  Moderator  Shuichi Ono  Hirosaki University
Yoshihiro Nishimura  Kobe University

P8-1. A Case of Thromboembolism in Peripheral Pulmonary Artery, Diagnosed by Dual-energy Multi-detector CT

Takahiko Kanokogi
Nara Medical University

P8-2. CT-guided Pericardiocentesis for Recurrent Cardiac Tamponade due to Carcinomatous Pericarditis in a Patient with Adenocarcinoma of the Lung

Yoshihiro Kanemitsu
Takatsuki Red Cross Hospital

P8-3. A Case of Pleomorphic Carcinoma of the Lung Producing Multiple Cytokines and Forming Rapidly Progressive Mass-like Opacity

Masataka Matsumoto
Kitano Hospital

P8-4. A Case of Malignant Mesothelioma with Diffuse Swelling Mediastinum Lacking of Apparent Pleural and Pericardial Lesion

Hiroki Kimura
The University of Tokyo

P8-5. A Case of Pulmonary Tumor Thrombotic Microangiopathy Presenting Progressive Pulmonary Hypertension

Hiromi Tomioka
Kobe City Medical Center West Hospital
Program
January 29, 2011 [Sat]
9:00-10:50  Core session3

Chronic Obstructive Pulmonary Disease

Moderator  Toyohiro Hirai  Kyoto University
            Kiminori Fujimoto  Kurume University School of Medicine
Discusser  Hans-Ulrich Kauczor  Heidelberg University

Invited Speaker

9:00-9:20
1. Etiology and Pathophysiology of COPD
Kazutetsu Aoshiba  
Tokyo Women’s Medical University

9:20-9:35
2. PET Assessment for COPD: Changes in Pulmonary Perfusion Distribution
José G. Venegas  
Harvard Medical School, Massachusetts General Hospital

9:35-9:50
3. CT Assessment for COPD-Phenotype on CT in COPD
Keisaku Fujimoto  
Shinshu University School of Health Sciences

9:50-10:05
4. Proton MR Assessment of COPD
Yoshiharu Ohno  
Kobe University

10:05-10:20
5. Hyperpolarized Noble Gas MRI for COPD
Edwin J.R. van Beek  
University of Edinburgh

Scientific Presentation

10:20-10:30
1. Hyperpolarized Noble Gases and COPD: The Frontier
James P. Butler  
Brigham and Women’s Hospital, Harvard Medical School

10:30-10:40
2. Regional Quantification of Hyperpolarized Helium-3 Magnetic Resonance Imaging Apparent Diffusion Coefficients: What Dose Bronchodilator Administration in COPD Accomplish?
Miranda Kirby  
The University of Western Ontario

10:40-10:50
3. Airway Disease, Emphysema, and Airflow Limitation in Patients with Chronic Obstructive Lung Disease
Hironi Makita  
Hokkaido University Hospital
13:20-14:20 Special Lecture 2
Basics and Future Direction for Quantitative Functional CT Assessment
Moderator Michiaki Mishima Kyoto University
Kenya Murase Osaka University
James P. Butler Harvard Medical School

13:20-13:50
1. Architecture of Lung Parenchyma –Basics of Interpretation of HRCT of the Lung–
Harumi Itoh University of Fukui

13:50-14:20
2. Current Status and Future Direction of Quantitative Analysis for Functional CT
Eric A. Hoffman University of Iowa Carver College of Medicine

14:30-16:15 Core session 4
Interstitial Lung Disease
Moderator Yukihiko Sugiyama Jichi Medical University
Masashi Takahashi Shiga University of Medical Science
Discussser Kyung Soo Lee Samsung Medical Center, Sungkyunkwan University school of Medicine

Invited Speaker
14:30-14:55
1. Update for Diagnosis of IIP
Takeshi Johkoh Kinki Central Hospital of Mutual Aid Association of Public School Teachers

14:55-15:15
2. Update for Treatment of Idiopathic Interstitial Pneumonias (IIPs)
Kingo Chida Hamamatsu University School of Medicine

15:15-15:30
3. Smoking-related Interstitial Pneumonia Associated with Emphysema
Minoru Kanazawa Saitama Medical University

15:30-15:45
4. Hyperpolarlized Noble Gas MRI for ILD
Hiroto Hatabu Brigham and Women’s Hospital, Harvard Medical School

Scientific Presentation
15:45-15:55
1. Prognostic Significance of High-Resolution Computed Tomography in Nonspecific Interstitial Pneumonia
Hironao Hozumi Hamamatsu University School of Medicine

15:55-16:05
2. Automatic Classification of Regional Disease Pattern of Diffuse Lung Disease at HRCT: Cross-Vendor Study
Kim, Namkug Ulsan University College of Medicine, Asan Medical Center

16:05-16:15
3. FDG Positron Emission Tomography Imaging of Drug-induced Pneumonitis
Miwa Morikawa University of Fukui
Room B  
Event Hall (B1)

12:00-13:00  **Luncheon Lecture 2**  
*Imaging for Asthmatics and COPD*  
Moderator: **Shoji Kudoh**  
Double-Barred Cross Hospital  
Sponsored by **Astellas Pharma Inc. / AstraZeneca K.K.**

12:00-12:30
1. **CT Assessment of Asthma**  
**Gregory G. King**  
The Woolcock Institute of Medical Research

12:30-13:00
2. **Hyperpolarized Noble Gas MRI for Asthma and COPD**  
**Mitchell S. Albert**  
University of Massachusetts Medical School

16:30-17:30  **Evening Lecture 2**  
*New Techniques for Pulmonary Vascular Disease*  
Moderator: **Kazuro Sugimura**  
Kobe University  
Sponsored by **Eisai Co., Ltd.**

16:30-17:00
1. **Dual-Energy CT Application for Pulmonary Vascular Disease**  
**Joon Beom Seo**  
University of Ulsan College of Medicine, Asan Medical Center

17:00-17:30
2. **Quantitative MR Assessment for Pulmonary Vascular Disease**  
**David L. Levin**  
Mayo Clinic
Poster Session
Event Hall Lobby (B1)

17:40-18:43 Poster Session 9
Pulmonary Circulation 2

Moderator Christian Fink University Medical Center Mannheim
Minoru Kanazawa Saitama Medical University

P9-1. Changes in Regional Pulmonary Perfusion and Morphology as a Result of Lung Volume Assessed by Technetium-99m Macroaggregated Albumin SPECT and MiniP with MDCT

Hiroko Tomita National Defense Medical College

P9-2. Estimation of Pulmonary Arterial Pressure in Pulmonary Hypertension Based on Interventricular Septal Configuration Obtained by Electrocardiogram-gated 320 Slice CT

Toshihiko Sugiura Chiba University

P9-3. Right Ventricle to Left Ventricle Volume Ratio by Electrocardiogram-gated 320 Slice CT is a Predictor of Right Ventricle Pressure Load in Pulmonary Hypertension

Toshihiko Sugiura Chiba University

P9-4. Correlation of Right Ventricular Ejection Fraction with Tricuspid Annular Plane Systolic Excursion by Electrocardiogram-gated 320 Slice CT in Pulmonary Hypertension

Toshihiko Sugiura Chiba University

P9-5. Evaluation of Hemodynamic Changes in Pulmonary and Systemic Arterial Circulation in Patients with Pulmonary Fibrosis in Relation to Breathing Methods Using Phase Contrast MRI

Nanae Tsuchiya University of the Ryukyus Hospital

P9-6. Alteration of Pulmonary Perfusion after CT Density-Based Non-uniform Attenuation Correction on DIBrH SPECT-CT Fusion Images

Kazuyoshi Suga St. Hill Hospital

P9-7. Comparative Study of Tc-99m MAA Lung Perfusion SPECT and Multidetector CT Pulmonary Angiography in Patients with Chronic Pulmonary Embolism-Multicenter Trial

Shigeru Kosuda National Defense Medical College
Poster Session 10

Thoracic Malignancy

Moderator: Hidetake Yabuuchi, Kyushu University
Yoshinobu Iwasaki, Kyoto Prefectural University of Medicine

P10-1. CT Diagnosis of Recurrence of Lung Cancer
Fumiyasu Tsushima, Hirosaki University

P10-2. Comparison of Quantitative Diagnostic Capability of Lymph Node Metastasis in Non-small Cell Lung Cancer Patients between STIR Turbo SE Imaging, Diffusion-Weighted Imaging and FDG-PET/CT
Daisuke Takenaka, Kobe University

P10-3. Capability for Quantitative Differentiation of Small Cell Lung Cancer from Non-Small Cell Lung Cancer by Diffusion-Weighted Imaging and STIR Turbo SE Imaging
Hisanobu Koyama, Kobe University

P10-4. Comparison of Capability for Pulmonary Nodule Detection; 1.5T vs 3.0T Systems
Keiko Matsumoto, Yamanashi Hospital of Social Insurance

P10-5. Utility of Nodule Type Assessment for Improving Diagnostic Capability in Patients with Solitary Pulmonary Lung Nodules on Low-Dose CT at FDG-PET/CT
Yumiko Onishi, Kobe University

P10-6. Estimation of Increase of Pulmonary Ground-Glass Opacity: Integrated evaluation of detected GGOs using early indicator of growth
Shuji Yamamoto, LISIT, Co., Ltd.

Ayano Kamiya, University of the Ryukyus

P10-8. Prediction of Postoperative Lung Function in Patients with Lung Cancer: Comparison of Dual-energy Perfusion CT and Perfusion Scintigraphy
Eun Jin Chae, Ulsan College of Medicine, Asan Medical Center
17:40-18:43  Poster Session 11
Chronic Obstructive Pulmonary Disease and Others
Moderator  Osamu Honda  Osaka University
            Masaharu Nishimura  Hokkaido University

P11-1. Quantitative Analysis of Pulmonary Cysts and Emphysema on Computed Tomography: Lymphangioleiomyomatosis, Birt-Hogg-Dubé Syndrome and Chronic Obstructive Lung Disease
Kazunori Tobino  Juntendo University

P11-2. In Vivo Monitoring of Cystic Fibrosis-like Lung Disease in Mice by Volumetric Computed Tomography
Mark. O. Wielpuetz  University Hospital Heidelberg

P11-3. Using Hyperpolarized 3He MRI for Treatment Evaluation in Cystic Fibrosis
Yanping Sun  University of Massachusetts Medical School

P11-4. Assessment of Paraseptal Emphysema in the Inner Lung Field by Using Inflated-Fixed Lung Specimens
Katashi Satoh  Kagawa Prefectural College of Health Sciences

P11-5. I-123-MIBG Lung Kinetic Abnormality in Pulmonary Emphysema
Kazuyoshi Suga  St. Hill Hospital

P11-6. The Epithelial-Mesenchymal Transition of Progenitor Cells Lead to Low Attenuation Areas of the Pulmonary Parenchyma: An Autopoietic Model Analysis of Alveolar Epithelial Maintenance
Kyongyob Min  Itami City Hospital

P11-7. Oscillatory Mechanics within Respiratory Cycles Related to COPD Severity
Junichi Ohishi  Tohoku University School of Medicine

17:40-18:34  Poster Session 12
Asthma, Airway Disease and Others
Moderator  Jin Mo Goo  Seoul National University Hospital, Seoul National University College of Medicine
            Shigeo Muro  Kyoto University

P12-1. Hyperpolarized 3He MRI Apparent Diffusion Coefficients as a Probe of Airway Function in Asthma After Methacholine Challenge and Recovery
Stephen E. Costella  Robarts Research Institute

P12-2. 4-Dimentional Imaging of the Central Airway Using a Low Dose Technique: Initial Experience with 320-row Multidetector CT
Ryo Sakamoto  Kyoto University
P12-3. Dynamic Property of Central Airway Walls as Assessed by Computed Tomography: Correlation with Asthma Pathophysiology
Masafumi Yamaguchi
Shiga University of Medical Science

P12-4. Comparison of 3-D-Visualized Respiratory Impedance Patterns and Structural Change Detected by CT in Smokers
Haruko Shinke
Kobe University

P12-5. Usefulness of SD-101, a Non-restrictive Device Developed for Screening Sleep Apnea-hypopnea Syndrome
Keisaku Fujimoto
Shinshu University School of Health Sciences

P12-6. Vibration Response Imaging (VRI) in the Detection of Asynchrony between Left and Right Lung for Main Bronchial Stenosis
Hirotaka Kida
St. Marianna University School of Medicine

17:40-18:34 Poster Session 13
Respiratory Motion 2

Moderator
Hideaki Haneishi Chiba University
Koji Chihara Shizuoka City Shizuoka Hospital

P13-1. Analysis of Lung Motion Due to Respiration and Its Application to PET Images
Hideaki Haneishi
Research Center for Frontier Medical Engineering

P13-2. A 4D Model Generator of the Human Lung: Lung CataChiCaraCure-er, Alias Lung4Cer
Hiroko Kitaoka
JSOL Corporation

P13-3. Alveolar Recruitment Analysis in Acute Respiratory Distress Syndrome (ARDS) Using End-inspiratory and End-expiratory Breath-hold 3D-CT Images
Hiroko Kitaoka
JSOL Corporation

P13-4. Velocity Vector Imaging in Assessing the Regional Diaphragmatic Movement of Patients with Chronic Diaphragmatic Hernia- Preliminary Results
Teng-Fu Tsao
Chung Shan Medical University Hospital

P13-5. Usefulness of Ultrasonography in Preoperative Diagnosis of Solitary Fibrous Tumor of the Pleura
Mitsuaki Sekiya
Juntendo University School of Medicine

P13-6. A Method for Reduction of Subtraction Artifacts on Temporal Subtraction Images by Use of Generalized Gradient Vector Flow Technique
Hyoungseop Kim
Kyushu Institute of Technology
17:40-18:34  Poster Session 14
Basic Study 2

Moderator  Takatoshi Aoki  University of Occupational and Environmental Health School
           Gregory G. King  The Woolcock Institute of Medical Research

P14-1. Towards a Research Software Platform for Diagnosis and Therapy Monitoring for COPD and Asthma with Pulmonary Magnetic Resonance Images
Peter Kohlmann  Fraunhofer MEVIS

P14-2. Clinical Chest CAD System for Lung Cancer, COPD, and Osteoporosis Based on MDCT Images
Shinsuke Saita  The University of Tokushima

P14-3. Developments of Thrombosis Detection Algorithm using the Contrast Enhanced CT Images
Jun Oya  The University of Tokushima

P14-4. Segmentation of Thoracic Organs from Multi-slice CT Images
Mikio Matsuhiro  The University of Tokushima

P14-5. Iterative Algorithm for Airway Segmentation in CT Images Using a Region-based Active Contour Model
Sumiaki Matsumoto  Kobe University

P14-6. A Comparative Study of HRCT Image Metrics and PFT Values for Characterization of ILD and COPD
Gang Song  University of Pennsylvania School of Medicine

17:40-18:16  Poster Session 15
Case Report 2

Moderator  Fumito Okada  Oita University
           Kazuya Fukuoka  Hyogo College of Medicine

P15-1. Unexplained Desaturation in the Sitting Position with Stroke
Shigeki Nanjo  Kobe City Medical Center General Hospital

P15-2. Acute-onset Sarcoidosis with Polyarthralgia and Hilar Lymphadenopathy
Hiroshi Santo  Kinki University

P15-3. Pulmonary Function and Computer Tomography of a Relapsing Polychondritis with Bronchial Lesion
Takahiro Tsuji  Tenri Hospital

P15-4. A Case of Toxocariasis Who Presented Abnormal Tubular Structures in Lung Fields
Yuko Nakase  Teikyo University School of Medicine
Program

January 30, 2011 [Sun]
Room A  
Main Hall (2F)

8:30-10:20  
**Core session 5**

**Asthma and Airway Disease**

**Moderator**  
Yuji Tohda  Kinki University

Shigeru Kosuda  National Defense Medical College

**Discusser**  
Shu Hashimoto  Nihon University

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**Invited Speaker**

8:30-8:50  
1. Pathophysiology of Asthma  
Hiromasa Inoue  Kagoshima University

8:50-9:05  
2. Evaluation of Airway Obstruction and Altered Pulmonary Permeability with Lung Scintigraphy in Young Patients  
Mayuki Uchiyama  The Jikei University School of Medicine

9:05-9:20  
3. CT Assessment for Asthma  
Hiroshi Tanaka  Sapporo Medical University School of Medicine

9:20-9:35  
4. MR Assessment of Asthma; Hyperpolarized $^3$He Magnetic Resonance Imaging to Probe Temporal and Spatial Airway Functional Changes  
Grace Parraga  The University of Western Ontario

9:35-9:50  
5. Update for Treatment of Asthma  
Kazuto Hirata  Osaka City University

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**Scientific Presentation**

9:50-10:00  
1. The Effect of Methacholine Challenge on Ventilation Heterogeneity in Asthma Measured by SPECT  
Gregory G. King  The Woolcock Institute of Medical Research

10:00-10:10  
2. Surface Irregularity of Airway Walls as Assessed by MDCT in Asthmatic Patients  
Tsuyoshi Oguma  Kyoto University

10:10-10:20  
3. Intrathoracic Tracheal Volume and Collapsibility on Inspiratory and End-expiratory CT Scans: Correlations with Lung Volume and Pulmonary Function in 85 Smokers  
Tsuneo Yamashiro  University of the Ryukyus
10:30-12:15  Core session 6
Respiratory Motion Analysis

Moderator  Ichiro Kuwahira  Tokai University Tokyo Hospital
           Shuji Sakai  Tokyo Women’s Medical University
Discusser  James Gee  University of Pennsylvania

Invited Speaker

10:30-10:45  1. Aims of Respiratory Motion Assessment for Anesthesiology
            Klaus Markstaller  Medical University of Vienna

10:45-11:00  2. MR Assessment of Respiratory Motion
            Tae Iwasawa  Kanagawa Cardiovascular and Respiratory Center

11:00-11:15  3. Nuclear Medicine Assessment for Respiratory Lung Motion
            Kazuyoshi Suga  Yamaguchi University School of Medicine

11:15-11:30  4. Dynamic Volume Scan Using 320-row ADCT for the Assessment of Respiratory Motion
            Hiroshi Moriya  Sendai Kosei Hospital

11:30-11:45  5. Respiratory Motion Assessment to Increase the Accuracy of Non-contact Screening System
to Prevent the Spread of Infectious Diseases
            Takemi Matsui  Tokyo Metropolitan University

Scientific Presentation

11:45-11:55  1. Temporal Stability of the 4D-CT Pulmonary Ventilation Imaging Method
            Tokihiro Yamamoto  Stanford University

11:55-12:05  2. Analysis of Respiratory Movement Using 4-Dimensional Chest CT image
            Seiji Tani  The University of Tokushima

12:05-12:15  3. Asymmetry of the Mechanical Property in the Human Respiratory System Causes the
             Asymmetry of the Lingual Center in the Brain: Diaphragmatic Motion Analysis with Dynamic
             MR Images
            Hiroko Kitaoka  JSOL Corporation
13:50-15:05  **Special Lecture 3**  
**Future Direction for Pulmonary Functional and Molecular Imaging**  
Sponsored by TOSHIBA MEDICAL SYSTEMS CORPORATION

**Moderator**
- **Yasutaka Nakano**  
  Shiga University of Medical Science  
- **Yoshiharu Ohno**  
  Kobe University  
- **Hiroto Hatabu**  
  Brigham and Women's Hospital, Harvard Medical School

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<th>Time</th>
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<th>Speaker</th>
<th>Institution</th>
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| 13:50-14:15| 1. From Basic Science Side                      | **Hisataka Kobayashi**  
  National Institute of Health                    |
| 14:15-14:40| 2. From Radiology Side                          | **Hans-Ulrich Kauczor**  
  Heidelberg University Medical Center            |
| 14:40-15:05| 3. From Pulmonology Side - Terminal Air Space of X-ray CT Images in COPD- | **Michiaki Mishima**  
  Kyoto University                                  |
| 15:05-15:15| Closing Remark                                    |                                              |
Room B
Event Hall (B1)

12:30-13:30  Luncheon Lecture 3  Sponsored by DAIICHI SANKYO COMPANY, LIMITED
Hyperpolarized Noble Gas MRI
Moderator  Toshihiro Nukiwa  Tohoku University

12:30-13:00
1. Development of Polarizer for Future Clinical Works  
F. William Hersman
University of New Hampshire, Xemed LLC

13:00-13:30
2. Hyperpolarized Noble Gas MRI: Clinical Applications  
Hiroto Hatabu
Brigham and Women’s Hospital, Harvard Medical School
Abstract

Special Lecture
Interaction between Functional and Morphological Imaging: 3D Airway Analysis in Chronic Obstructive Pulmonary Disease

Masaharu Nishimura, M.D., Ph.D.
Division of Pulmonary Medicine, Department of Internal Medicine, Hokkaido University Graduate School of Medicine, Japan

Chronic obstructive pulmonary disease (COPD) is a disease characterized by airflow limitations that are not fully reversible and consists of small airways disease (obstructive bronchiolitis) and parenchymal destruction (emphysema), the relative contributions of which vary among patients. Thin-section computed tomography (CT) has been used to quantify emphysema by detecting low-attenuation areas (LAAs), and the role of CT in diagnosing emphysema is well established. However, airflow limitations as evaluated by forced expiratory volume in 1 s (FEV₁) do not necessarily correlate well with severity of emphysema as evaluated by CT, since small airways disease appears to contribute more significantly to airflow limitations in COPD.

Recent progress in CT technology has enabled the detection and quantification of airway dimensions, including airway wall thickness and airway luminal area. The use of CT to evaluate airway dimensions in a variety of diseases has thus been suggested. However, only a few studies had attempted to measure airway dimensions in patients with COPD until 2006, when our study was first published in Amer J Respir Crit Care Med in 2006. Most previous reports had, unfortunately, suffered from inherent technological problems with respect to the measurement of airway dimensions. First, accurate cross-sectional images could not always be obtained using conventional two-dimensional images, since airways run in various directions in the lung. Second, since investigators did not obtain longitudinal airway images, which generation of the bronchus was actually being measured, could not be recognized. The idea of three-dimensional airway analysis was not entirely new, but had not previously been applied to measuring airway dimensions in patients with COPD.

In this presentation, I would like to show you how we validated our original software using phantoms, and how we applied this software to a series of clinical studies in COPD. Firstly, we have demonstrated, in a relatively large patients with COPD, that the percentage of predicted forced expiratory volume in 1 s (% FEV₁), an airflow limitation index, correlated highly with airway luminal area (Ai) and, to a lesser extent, with percentage wall thickening (WA%) from the 3rd to the 6th generation of both the apical upper bronchus (B1) and the anterior lower bronchus (B8). More interestingly and importantly, we showed that correlation coefficients improved with decreasing airway size in both airways. We then examined the relationship between airway dimensions and emphysema severity at the upper lung and the lower lung separately and how these pathological processes contributed to overall pulmonary function. Secondly, we attempted to determine the relationship between improved pulmonary function and changes in airway caliber at various sites among the airways in response to inhaled anticholinergics in patients with COPD. We performed CT at deep inspiration and detailed pulmonary function tests before and 1 week after daily inhalations of tiotropium bromide in 15 patients with clinically stable COPD. We analyzed the airway luminal area at the 3rd (segmental) to the 6th generations of 8 bronchi in the right lung. Bronchodilation was demonstrated as an overall average of a 39% increase in the inner luminal area, and the mean forced expiratory volume in 1 sec (FEV₁) increased from 1.23 ± 0.11 to 1.47 ± 0.13 (SE). The magnitude of bronchodilation closely correlated with improved pulmonary function, particularly with that of FEV₁, \( r = 0.843, p < 0.001 \). Such correlations were significant at the 4th to the 6th, but not at the 3rd generation of bronchi, and the slope of regression lines became steeper from the 3rd to the 6th generation. In conclusion, inhaled anticholinergics induce overall bronchodilation in proportion to improvements in FEV₁ in patients with COPD, and bronchodilation at distal, rather than proximal airways is the determinant of functional improvement.

These studies clearly demonstrated functional importance of the smaller airways among the 3rd to the 6th generation of airways in COPD. Although we are not in a stage where we can visualize and measure “small airways” which are physiologically defined as those whose inner diameter is less than 2 mm, we have come close to those areas. In this presentation, I will also present potential differences in airway remodeling which might exist between COPD and bronchial asthma. In short, airway remodelling was more prominent in asthma compared with age- and sex-matched COPD subjects at the 3rd- to the 6th-generation airways when airflow limitations as assessed by FEV₁/FVC and % FEV₁ were mild to moderate to similar extent under stable clinical conditions.

In the near future, with further advances in both CT technology and software, this kind of approach will become even more attractive. However, at the same time, we must be cautious about potential problems in the application of three-dimensional airway analysis to longitudinal follow-up studies and/or large-scale multi-center studies, which will be also discussed in this presentation.
Interaction between Metabolic and Morphological Imaging

Kyung Soo Lee, M.D.
Department of Radiology
Samsung Medical Center, Sungkyunkwan University School of Medicine, Korea

In the era of fusion (e.g., PET-CT or MR-PET) imaging, the representative morphologic imaging tools are CT and MRI, whereas PET and SPECT may be the representative of metabolic imaging. By integrating morphologic and metabolic imaging, synergistic effect of both modalities can be acquired.

The substantial and expected effect of combining metabolic and morphologic imaging devices includes 1) precise localization of lesions detected on metabolic imaging scans within the anatomic frame provided by morphologic imaging tool (seeking a lesion of morphologic changes that matches with a metabolic-imaging lesion), thus increasing specificity of the examination, 2) a supply of morphologic information of target lesions (e.g., attenuation value including calcification at CT and signal intensity of target lesions at MRI) by morphologic imaging and thus preventing observers from making false positive interpretations, 3) an insight on the necessity of additional morphologic information acquisition in organs where normal metabolism of radioisotopes do occur (like brain, myocardium, liver and the kidneys) at FDG PET, and 4) information of a need for development of various radiopharmaceuticals other than FDG and for anatomically-guided partial volume correction in metabolic imaging.

In this talk, the speaker will elaborate on above-mentioned problems by showing illustrative cases.
1. Introduction

This special lecture will describe three basic structures of the lung. The first is three dimensional features of alveoli and alveolar ducts. The second is a respiratory bronchiole and its relation to adjacent parenchyma. The third is small pulmonary vasculatures and lymphatic vessels. These topics are based on the results of radiologic-anatomical-pathologic correlations on lung specimens obtained from surgery and autopsy. Contact radiography and stereomicroscopic observations were especially useful to study detailed anatomy of peripheral portions of the lung.

2. Normal morphologic features of lung periphery

(1) Spatial configuration of alveoli and alveolar ducts

Lung parenchyma is known to occupy 90% of total lung volume, however it looks empty or zero in normal radiological images (chest radiograph, HRCT) because of remarkable radiolucent nature of the normal lung parenchyma. Macroscopically normal lung parenchyma seen in lung specimens looks compact like a liver or brain. These facts implies that radiologic zero-looking lung parenchyma is packed with air and specially designed real tissue which forms lung parenchyma. Contact radiograph and HRCT of normal lung specimen shows the lung parenchyma has weak radio-opacity which is definitely greater than air.

Stereomicroscope is useful to observe lung parenchyma on the cut surface of inflated and fixed lung specimens. With weak magnification the lung parenchyma resembles a sponge. The numerous small holes correspond to alveolar ducts, not the alveoli. With high power magnification, the alveolus appears polyhedral in shape, and two types of alveolar wall are identified, that is, lateral wall and dome of alveolus. Each alveolar entrance is polygonal in shape and is packed like a honeycomb which surrounds the lumen of alveolar ducts.

One set of alveoli facing the adjacent two alveolar ducts shows a back to back arrangement. This back to back arrangement of the alveoli is recognized both in profile and from front under the stereomicroscope. The zigzag pathway of alveolar dome seen in histological section corresponds to this arrangement seen in profile. The superimposed back to back alveoli show fine reticular pattern in a contact radiograph of lung specimens.

Stereomicroscope is also useful to observe the bronchioles in post mortem bronchograph. It shows the diameter of terminal and respiratory bronchioles is smaller than that of more peripheral air passages. Since cross section of such peripheral air passages consists of an alveolar duct and marginal alveoli, its whole diameter becomes bigger than that of the central bronchioles. This change of diameter gives a tree-in-bud pattern made by a contrast medium or by pathologic lesions filling those portions of the lung.

(2) Double and single faced alveolar walls

Each alveolar wall included in the back to back alveoli is called double faced alveolar wall, because gas exchange is performed in both sides of the alveolar wall. In contrast the alveolar walls attaching to the airways, pulmonary vessels, interlobular septum and pleura are single faced alveolar walls, since gas exchange is limited to one direction. Ninety percent of total alveolar walls are the double faced alveolar wall.

Respiratory bronchiole is characterized by scattered alveoli opening from the bronchiolar wall. The important point is how the alveoli of the respiratory bronchiole are related to those of surrounding pulmonary parenchyma. Histological study shows the alveolar domes of both sides of alveoli are held in common, therefore, respiratory bronchiole and adjacent parenchyma are connected by the back to back to alveoli, instead the alveolar dome surrounding the bronchiolar wall is the single faced alveolar wall and separated from the bronchial lumen.

The back to back to alveoli extending from the wall of the respiratory bronchiole may be responsible for producing centriacinar or centrilobular inflammatory nodules seen in pulmonary infectious diseases.

(4) Arterioles and venules

Pulmonary vessels are one of the non parenchymatous structures of the lung, and occupy less than 10% of total lung volume. However, we tend to overestimate the volume of the pulmonary vessels in HRCT, angiogram, and even in chest radiograph, because the lung parenchyma superimposed to those vessels may not be recognized correctly.

Small pulmonary vessels whose diameter is below 70 microns are called arterioles and venules. Within
the secondary lobule of the lung, the arterioles divide from muscular arteries which run with bronchioles, and distribute along the alveolar domes.

An interesting feature of the arterioles is a peculiar branching pattern which is demonstrated by post mortem angiograph. A number of arterioles divide directly from the axial muscular artery like a twig. Such a small lateral branch provides blood flow to the lung parenchyma adjacent to the respiratory bronchioles.

Arterioles and venules are equipped with lymphatic vessels in the perivascular interstitial space. These lymphatics are related to the pulmonary vessels and do not exist in the alveolar wall.

(5) **Alveolar capillaries**

The double faced alveolar wall is equipped with denser network of capillaries than that of single faced alveolar wall. A post mortem micro-angiograph shows alveolar capillaries appearing like a polygonal network. However, the microphotograph of the same specimen shows alveolar capillaries are not distributed in a smooth plane but undulated within the alveolar wall, which may contribute to increase alveolar surface area for gas exchange.

(6) **Lymphatic vessels**

Lymphatic vessels are distributed along non-parenchymatous structures such as airways, pulmonary vessels, interlobular septa and pleura. The lymphatic vessels are usually observed by histological sections, however, those of visceral pleura are able to observe with a naked eye. Three dimensional images of the pleural lymphatic vessels are obtained from HRCT of lung specimens inflated with air. The pleural lymphatics form variously sized polygonal network. The large network borders the secondary pulmonary lobe with interlobular septum and pulmonary vein.

3. **Disease patterns of lung parenchyma**

Typical disease patterns are mentioned here, which is suitable for establishing morphological concepts specified to lung parenchyma and related structures.

(1) **Disease of alveolar wall**

Radiologic-pathologic correlations done in case of adenocarcinoma prove thickening of alveolar walls is responsible for ground glass attenuation in HRCT. Histological studies showed all alveolar walls within the tumor are thickened by cancer invasion and interstitial reaction, and the tumor is surrounded by normal back to back alveoli. These observations suggest that adenocarcinoma may extend continuously along the back to back alveoli.

Usual interstitial pneumonia involves primarily the alveolar walls of back to back alveoli. Histological studies show that in addition to honeycomb lung isolated fibrotic lesions are surrounded by normal back to back alveoli.

(2) **Disease of alveoli and alveolar duct**

A tree-in-bud pattern seen in HRCT of pulmonary tuberculosis is formed by compact caseous materials filling sequentially the respiratory bronchiole, alveolar ducts and alveoli, while accompanying arteries and arterioles are kept in normal. In HRCT these lesions observed in pulmonary tuberculosis may mimic to those of pulmonary sarcoidosis. In sarcoidosis the disease tends to be localized around the pulmonary vessels.

(3) **Centriacinar (Centrilobular) disease**

Centriacinar disease involves respiratory bronchioles, peribronchiolar interstitial tissue and adjacent pulmonary parenchyma simultaneously. The back to back alveoli anchored to the wall of respiratory bronchiole may provide bypass route of disease extending to adjacent lung parenchyma. Centriacinar inflammatory nodules are bigger than the diameter of respiratory bronchiole and such lesions are seen typically in diffuse panbronchiolitis, pneumoconiosis and pulmonary infection.

(4) **Perivascular disease**

Perivascular disease pattern is derived from pathologic lesions distributing along lymphatic vessels related to pulmonary vasculatures. Since pathologic lesions surround pre-existing pulmonary vessels, the vascular images in HRCT become thickened and irregular in contour. Typical images are shown in case of pulmonary sarcoidosis, lymphangiosis carcinomatosa and lymphoma.

4. **Summaries**

The key structure of the lung parenchyma is the alveolar wall. Most of the alveolar walls are intervened by air both in the alveoli and alveolar ducts except the alveolar walls attaching to the non parenchymatous pulmonary structures. The lateral walls and domes of alveoli are connected to form back to back alveoli and distribute among adjacent alveolar ducts. The superimposed alveolar walls show fine reticular patterns in contact radiograph of sliced lung specimens. The arterioles and venules do not accompany the bronchial trees and are laid within interstitial space of alveolar domes. The respiratory bronchioles are equipped with a bypass route for some diseases extending from the bronchial wall to adjacent lung parenchyma through the back to back alveoli.
Imaging-based metrics are playing a central role in the quest to identify COPD phenotypes, serving to establish homogeneous sub-populations to: 1) aid in the search for genotypes; 2) provide targets for the development of new therapeutic interventions; and 3) serve as outcomes measures for the evaluation of therapeutic interventions. CT has emerged as a tool of choice for assessing structural-based changes in the lung associated with COPD including measures of lung density distributions at full inspiration as an index of emphysema, lung hyperlucency at expiration as an index of air trapping together with measures of airway wall remodeling. While measures of lung destruction and airway remodeling are providing metrics which are more sensitive than standard pulmonary function tests, *CT phenotypes of lung disease must move towards providing insights into the etiology of the disease, not just road maps of destruction.*

Recent findings have lead us to believe that CT derived perfusion (PBF) and mean transit time (MTT) measures within inflamed lung parenchyma provide for a functional phenotype of emphysema which may be directly tied to the etiology of the pathologic process leading to emphysema in a subset of the smoking population susceptible to centrilobular emphysema. **Our central hypothesis is that smokers prone to emphysema have abnormal vasoregulation in that regional hypoxic pulmonary vasoconstriction (HPV) continues despite inflammation.** This failure to block vasoconstriction alters the repair response and leads to tissue injury and eventually smoking-associated centrilobular (and paraseptal) emphysema in those smokers with abnormal vasoregulation. The normal response to regional hypoxia in the lung is to shunt blood towards better-ventilated regions. However, if small, regional infiltrates associated with a heterogeneous smoking-associated process induces local hypoxia, HPV in regions of smoking-induced infiltrates will interfere with defense mechanisms serving to clear the irritant and will interfere with mechanisms of repair. We (1) have recently demonstrated that smokers with normal pulmonary function tests (PFTs) but subtle visual CT evidence of smoking-associated centrilobular emphysema (SCE) have increased heterogeneity (coefficient of variation: CV) of CT-derived measures of PBF and MTT as compared with smokers with normal PFTs and normal CT images (SNI). This is consistent with reduced perfusion in hypoxic (alveolar flooded) regions. A corollary to the above hypothesis is that the enhanced CV of perfusion in SCE subjects will appear throughout the lung (smoking is associated with increased parenchymal density on CT) while the destructive effects resultant from HPV will be magnified at the lung apices where gravitational forces resisting flow are the greatest, compounding effects of increased vascular resistance. We thus provide what has been an elusive explanation for the apical distribution of centrilobular emphysema.

While increased heterogeneity of perfusion in SCE subjects is consistent with our above hypothesis, further supportive evidence is required. Additionally, to translate our laboratory research into multi-center studies, we have demonstrated the equivalence of perfused blood volume (PBV) vs. PBF/MTT CV. Using dual energy CT, a PBV protocol avoids placement of a central contrast line and significantly reduces radiation dose. In addition to the use of PBV as an index of regional perfusion, we are seeking to evaluate the equivalency of a single breath / dual energy Xenon CT scan as an alternative to a washin protocol for the assessment of regional ventilation to derive matched PBV / Ventilation images.
The development of imaging technologies that have sufficient specificity and sensitivity to enable early and accurate detection of cancer as well as offer the ability to monitor early tumor response to therapy has long been a goal of oncologic research. Numerous radiological techniques have been used for the diagnosis of diseases including cancer as well as for the surveillance of recurrence. These include: x-ray (plain film and computed tomography [CT]), magnetic resonance imaging (MRI), radionuclide imaging (single photon emission computed tomography (SPECT)/ positron emission tomography (PET)), and ultrasonography (US). It is evident that imaging has revolutionized various fields including in the lung diseases. However, despite the widespread use of these technologies, the ability of currently available modalities to facilitate the early detection, precise characterization, and accurate localization of diseases remains suboptimal. For example, the current detection threshold for imaging solid tumors of 1cm³ or approximately 10⁹ cancer cells exemplifies this, as achieving early and accurate diagnosis clearly requires detecting cancer far earlier in the course of disease. Multiplexed imaging offers great potential for improving cancer detection and treatment. In a broad sense, multiplexed imaging refers to techniques that yield multiple distinct data sets from a single imaging session, however, few primitive methods have recently been used in the clinic. Four general strategies for multiplexed imaging have been proposed.

1) **Multimodality** in which two (or more) distinct imaging modalities are simultaneously or consecutively employed; PET/CT is an example that is currently in clinical use.

2) **Multicolor contrast agents** in which multiple different agents, with distinct energies (colors) are simultaneously imaged using a single modality; SPECT has been used in this manner and can simultaneously detect two imaging agents, i.e., Tc-99m and Tl-201, by utilizing the differences in energy emission of distinct radioisotopes.

3) **Multiple signal collection** in which a single modality can detect and interpret multiple signals obtained with distinct signaling technologies; MRI techniques such as T1/T2 and diffusion-weighted imaging as well as dual-energy X-ray CT are examples of multiple signal imaging currently used in clinic.

4) **Activatable “smart” reagents**, which generate data regarding both tumor location and molecular profile using targeted probes that emit signal in target tissue only after cellular internalization or other triggering events. These imaging methods based on the multiplexed strategies would overcome the limitation of each one of conventional modalities, which can be broadly categorized into primarily anatomical and primarily molecular imaging techniques, generally with an inverse relationship between spatial resolution and their sensitivity to detect small changes of disease. Primarily anatomic imaging technologies include CT, MRI, and ultrasound. These modalities are limited by their ability to detect disease only after tumor size or associated structural changes are significant enough to be detected grossly. At the other end of the spectrum are primarily molecular imaging modalities, which offer the ability to detect molecular and cellular changes of disease before disease can be detected grossly. Although extremely promising, currently available molecular modalities are often limited by depth of signal penetration as well as poor spatial resolution. Within current clinic, the most widely used functional modalities include PET and SPECT, which can provide information regarding tissue activity at the expense of the anatomic detail due to poor spatial resolution.
Of the desired improvements in clinical imaging, earlier detection of disease, more accurate molecular characterization of tumors and the tumor microenvironment, earlier detection of response to therapy, are the most highly sought after advances. In order to achieve these important imaging objectives, the limitations of individual technologies must be minimized and new innovations developed with these endpoints in mind. Towards this end, the simultaneous use of two or more modalities, contrast reagents, signaling methods, or the coupling of agent and tissue properties, to achieve so called “multiplexed imaging,” is a promising approach. For example, combining PET with CT simultaneously yields data regarding tissue metabolism and structural anatomy. A parallel example in molecular imaging is the smart probe, which utilizes both molecular targeting and intracellular probe processing to provide information about the surface molecular profile of a tumor, thus improving sensitivity and specificity compared to conventional “always on” imaging agents. In this lecture, I will provide a broad overview of multiplexed imaging technologies ranging from those currently available to those still under development especially for oncologic imaging.

References:
Lung imaging has shown a series of groundbreaking technological developments in recent years. They involve the major radiological imaging techniques such as computed tomography (CT), magnetic resonance imaging (MRI), positron emission tomography (PET) and to some degree single photon emission computed tomography (SPECT) and ultrasonography. They are mainly related to high imaging speed; gating techniques, either respiratory, cardiac or intrinsic; signal enhancement technologies including new contrast agents and mechanisms, as well as new probes and targets. These modalities are available for clinical studies, but also for translational and preclinical research. The complexity increases even more with so-called hybrid imaging systems, such as PET/CT, SPECT/CT, MRI/PET but also combinations with optical imaging systems. For small animal imaging additional approaches such as high-speed video microscopy, multiphoton microscopy, darkfield and (auto-)fluorescence microscopy, as well as molecular confocal live-cell imaging and other techniques of optical imaging are also available or under investigation.

Some years ago, MRI paved the way for comprehensive imaging of structure and function in lung disease. Such protocols include perfusion, ventilation, V/Q ratio and match; gas exchange, vascular permeability, blood flow, shunt; hemodynamics, heart function, lung volumes, respiratory motion, and mechanics. Recent developments, such as "interpolated volume imaging” with high spatial resolution; contrast enhancement; single and multiphasic MR angiography; diffusion-weighted and diffusion tensor MRI sequences; multidirectional blood flow measurements; angiogenesis imaging and high-resolution ventilation imaging with hyperpolarized $^{129}$Xenon gas are the main drivers for broader translational and clinical implementation.

This comprehensive imaging assessment is an asset when compared to CT, which is further complemented by the fact that MRI does not carry any exposure to ionizing radiation which makes it especially advantageous in children, young adults and for follow-up examinations either in disease surveillance or therapy monitoring.

In the meantime, CT has demonstrated a surprising development towards a complex quantitative imaging technology with a substantial increase of measurable variables. Progress in iterative image reconstruction, post-processing, registration algorithms as well as multi-energy, spectral CT and potentially photon counting CT are the current examples. Especially spectral and photo-counting CT will also induce a renaissance of research for CT contrast agents. The most obvious for pulmonary CT being inhaled Xenon gas for imaging of ventilation distribution and trapped air.

Clinical PET/CT of the lung is still mainly based on FDG as an indicator of increased glucose metabolism, especially in malignant disease. Current developments, however, focus more on specific probes targeting dedicated functions or receptors, e.g. in pulmonary inflammation.

Beyond the modality-driven radiological view onto the future direction of pulmonary functional and molecular imaging, the objectives and endpoints of imaging have to be revisited. It is no longer mainly on morphology, e.g. size, volume, shape and a single descriptor of tissue composition, but on function, metabolism and molecular interactions as well as on multilevel analysis of tissue composition.

Thus, future endpoints of pulmonary imaging might be:

1) Inflammation

Infectious inflammatory pulmonary diseases pose a major challenge to public health and healthcare systems worldwide. Although imaging is currently not in a pivotal role for the detection, characterization and treatment decision making, there is a clear unmet clinical need to improve these aspects, especially within the scope of new micro-organisms, pandemics, resistant strains, such as in mycobacteriosis, and opportunistic infections. Besides, the incidence of non-infectious inflammatory diseases is increasing and detection and characterization as well as quantitative disease surveillance and treatment monitoring are becoming increasingly important. Potential future directions will be:

- Differentiation between infectious and non-infectious inflammation
- Estimation of inflammatory activity in pneumonitis, such as pulmonary fibrosis, granulomatosis and pulmonary manifestations of collagen vascular disease
- Airway inflammation in chronic bronchitis, chronic obstructive pulmonary disease (COPD) and asthma
- Imaging cell trafficking
- Imaging pulmonary immune responses and inflammatory signaling events
2) Homeostasis and energy balance
Imaging of water and salt represents a potential novel field for imaging. It will focus on ions such as sodium, potassium, chlorine by dedicated MRI or even MRS techniques at ultrahigh fields. Another important aspect will be the definition of a methodology for measurements of transepithelial transport mechanisms, e.g. gas exchange, but also transport of liquid and mucus along the airway surfaces.
Energy balance imaging also comprises the visualization of the respiratory muscles during breathing, e.g. respiratory mechanics, and their metabolism.

3) Real-time visualization of lung function: “Watch the lungs work”
The concept from structural to functional imaging opens the new field of real-time imaging of lung function, such gas exchange, cell trafficking and airway clearance. These developments are on the way for preclinical applications especially using live cell imaging, optical imaging and video-microscopy, but also in the clinical arena as 4D CT and MRI of pulmonary motion become widely available. Among other techniques, these will include spectral CT and MRI approaches to measure ventilation distribution and ventilation/perfusion ratios using inhaled contrast agents, such as xenon gas; or multidirectional measurements of air and blood flow. Such investigations might be extended to imaging of the lung during pharmacological stress or exercise (treadmill).

4) Imaging tumor biology
Different features of tumor biology may be visualized by imaging. Fluorescent imaging of the airways and early detection of dysplastic and neoplastic cells are important tools for early detection as are different morphologic features of early cancers at CT: solid, semi-solid, non-solid. Characterization of tumor biology will also comprise imaging of tumor angiogenesis as currently available and apoptosis in the future.

5) Phenotyping and genotype correlation
In the future, radiology will be increasingly used for phenotyping disease subtypes, e.g. in airway disease such as COPD. Phenotyping will be an essential counterpart of genotyping and other read-outs from molecular medicine. These activities will pave the way for individualized therapies and concepts for active surveillance in the future.
1. Introduction
We have been engaged in the digital analysis of X-ray CT image. Nakano et al. reported that LAA% (percent ratio of low attenuation area to whole airway area) and WA% (percent ratio of airway wall area to whole airway area of rt. B1) could complementally explain the obstructive disorders of ventilation in COPD [1].

We now focus on the importance of the analysis of the terminal airspace detected by CT image. The role of “Distribution of terminal airspace” and “Comorbidity of emphysema” will be presented.

2. Distribution of terminal airspace.
Mishima et al. examined the statistical properties of LAA clusters in COPD patients and in healthy subjects [2]. In COPD patients, the percentage of the lung field occupied by LAAs (LAA%) ranged from 2.6 to 67.6. In contrast, LAA% was always <30% in healthy subjects. The cumulative size distribution of the LAA clusters followed a power law characterized by an exponent D (fractal dimension). They show that D is a measure of the complexity of the terminal airspace geometry. The COPD patients with normal LAA% had significantly smaller D values than the healthy subjects, and the D values did not correlate with pulmonary function tests except for the diffusing capacity of the lung. We interpret these results by using a large elastic spring network model and find that the neighboring smaller LAA clusters tend to coalesce and form larger clusters as the weak elastic fibers separating them break under tension. This process leaves LAA% unchanged whereas it decreases the number of small clusters and increases the number of large clusters, which results in a reduction in D similar to that observed in early emphysema patients. These findings suggest that D is a sensitive and powerful parameter for the detection of the terminal airspace enlargement that occurs in early emphysema.

Seyama et al. found that the LAA distribution of LAM had poor fractal properties, and the mechanism of low attenuation formation in LAM may be quite different from that in emphysema.

Sato et al. applied the concept of fractal geometry to investigate the structural mechanism of the development of pulmonary emphysema in the klotho mouse, which, after milk feeding, exhibits characteristics resembling aging and develops emphysema [3]. They calculated the relationships between perimeter and size characterizing shape and between cumulative frequency and size of the terminal air spaces identified from histologic slides and found that both relations followed a power law with fractal properties. However, the fractal dimensions related to the shape and size (Dsn) in the klotho mice were significantly lower than in controls. Additionally, in the klotho mice, Dsn decreased with age without significant change in mean linear intercept. These abnormal morphological changes were restored when the klotho mice were fed with a vitamin D-deficient diet. Previously undescribed morphological model simulations showed that a random destruction, in which the destruction process occurs homogeneously in the lungs, was more consistent with the data than a correlated destruction that is usually seen in smoking-related human emphysema. These results suggest that the pathological changes in the lungs of the klotho mice are derived not from localized causes, but from systemic causes that are related to abnormal activation of vitamin D. The morphogenesis of emphysema in the klotho mice and morphological analyses using fractal geometry may contribute to the understanding of the progressive nature and cause of parenchymal destruction in human emphysema.

3. Comorbidity of emphysema:
COPD has recently been announced to be “a systemic disease”. It is clinically important that COPD and comorbid diseases influence together on the condition and prognosis of the patients. Emaciation has been known to be associated with COPD. Ogawa et al. evaluated emphysema using percentage of LAA% and airway by percentage wall area (WA%) [4]. Body mass index (BMI) was significantly lower in the higher LAA% phenotype. BMI correlated with LAA% but not with WA%. BMI was significantly lower in the emphysema dominant phenotype than in the airway dominant phenotype, while there was no difference in forced expiratory volume in 1 s %predicted between the two. These results support the concept of different COPD phenotypes and suggest that there may be different systemic manifestations of these phenotypes.

Osteoporosis is one of the important systemic features of COPD. However, the relationship between the extent of emphysema and reduced bone density has been unclear. Using CT images, the CT scan density of the thoracic and lumbar vertebrae (T4, T7, T10, and L1) and the LAA% were measured quantitatively, and their correlations were analyzed. Linear regression analyses revealed that LAA% had a significant negative correlation with bone mineral density (BMD). In addition, multiple regression analysis showed
that only LAA% and body mass index (BMI) were predictive of BMD among age, BMI, smoking index, FEV(1), arterial blood gas, and LAA%. These results suggested that COPD itself could be a risk factor for osteoporosis [5].

Haruna et al. analyzed the relationship between CT scan assessment and COPD mortality in patients with various stages of COPD [6]. Univariate Cox analysis revealed that emphysematous change as assessed by CT scan, lung function, age, or BMI were significantly correlated with mortality. Multivariate analysis revealed that emphysematous change as assessed by CT scan had the best association with mortality. They concluded that the emphysematous change as assessed by CT scan predicts respiratory mortality in outpatients with various stages of COPD.

4. Conclusion
Analysis of the distribution of terminal airspace using fractal analysis offers the information about the cause and process of the diseases. The emphysema has a tight relationship with the comorbidity. It is a key pathological change which prescribes the prognosis of COPD. "respiratory CT imaging" is always "respiratory functional imaging", and it plays an important role in the development of respiratory medicine.

References:
Abstract

Luncheon Lecture
Recent technical developments of multi-slice CT characterized by faster speed and longer coverage facilitate functional thoracic CT evaluation in children by reducing motion artifacts. Changes in lung density and airway caliber throughout the respiratory cycle have been used for the detection of air trapping and tracheobronchomalacia, respectively. Paired end-inspiratory and end-expiratory CT is used for cooperative patients, whereas free-breathing cine-CT that needs only sedation without any respiratory maneuvers or general anesthesia is a useful alternative for uncooperative patients. Dynamic expiratory CT increases sensitivity in detecting tracheobronchomalacia in cooperative patients. Because of longer coverage, 4D analysis of airways puts to our clinical practice. New respiratory-triggered CT acquisition of the whole lung with non-rigid 3D registration enables us to produce density difference map and perform quantitative analysis. Dual-energy CT opens up a new horizon for the qualitative and quantitative evaluation of lung perfusion and lung ventilation. Of interest, collateral ventilation can be visualized and quantified with dynamic xenon-enhanced dual-energy lung ventilation CT. This experience paradoxically underscores the importance of lung anatomy by recognizing that the incomplete and accessory fissures substantially influence regional lung ventilation. In addition to the separate use of lung perfusion or lung ventilation CT, these two emerging methods can be used together like V/Q scan. This combined evaluation is particularly useful for the evaluation of pulmonary embolism. We, pediatric imagers, should keep in mind that clinical benefits should outweigh radiation risks of functional thoracic CT. Therefore, we should try every effort to minimize CT radiation dose.
Pediatric Applications of Lung MRI

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Lung disease, including asthma, cystic fibrosis and bronchopulmonary dysplasia, is the most common chronic disorder of childhood. However, the assessment of the severity of lung disease can be difficult in the pediatric population. Young children are typically unable to perform the breathing maneuvers required to perform spirometry, the most commonly used method to assess lung function in adults. CT, which provides detailed images of lung structure, exposes the patient to relatively high doses of ionizing radiation. Children are at greater risk of developing a radiation induced malignancy than adults because children are inherently more radiation sensitive (more rapidly dividing cells) and typically have a long life expectancy following the exposure in which to develop a radiation induced malignancy. Thus the role of CT in pediatric lung disease is limited. MRI, which is thought to be very low risk, has the potential to have an important role in the assessment of pediatric lung disease.

Conventional proton MRI in the absence of inhaled or intravenous contrast currently has a limited role in the assessment of lung disease due to the relatively low proton density of the lung. There have been a few studies assessing the utility of proton MRI in cystic fibrosis (CF), essentially as a substitute for CT, to detect lung structural changes such as bronchiectasis, mucus plugging, and consolidation. Not surprisingly, CT has better image quality but important structural alterations can be detected with MRI. Further, there are techniques under development, such as ultra-short TE and Fourier decomposition, which have the potential to greatly increase the type of information about the lung that can be obtained with proton MRI.

Hyperpolarized noble gases (helium-3 and xenon-129) are used as inhaled gaseous contrast agents for MRI, and provide high temporal and spatial resolution images of the lungs. Most of the human imaging with hyperpolarized gas has employed helium-3 as the contrast agent. In the past few years, the available supply of helium-3 has decreased dramatically due to a spike in demand for helium-3 for use in neutron detectors. This coupled with recent advances in xenon polarization technology makes xenon-129 an attractive alternative to helium-3. Since little work has been done in pediatric lungs diseases with xenon-129, the following discussion will be focused on results obtained with helium-3. However, there is every reason to believe similar results could be obtained with xenon-129.

Images of lung ventilation can be obtained by performing spin density imaging during a breath hold following the inhalation of hyperpolarized gas. Areas of the lung that are well ventilated appear bright on the MR images and those that are poorly ventilated appear dark. Asthmatics have more ventilation defects, that is more poorly ventilated regions with their lungs, than healthy subjects. Further there is a moderate correlation between the number of ventilation defects and asthma severity as measured by spirometry or the frequency of symptoms. However the correlation is imperfect with some asthmatics having many ventilation defects but few symptoms and vice versa. Hyperpolarized helium-3 MRI gas has yielded a new insight into the pathophysiology of asthma, that the distribution of ventilation abnormalities in the lung is not random as would be expected if the whole lung was equally affected. Instead, some regions of the lung are more likely to appear abnormal on ventilation imaging than others, suggesting the severity of asthma within the lung varies regionally. This may have important implications for regional treatments such as bronchial thermoplasty in which the airway wall smooth muscle is RF ablated during bronchoscopy. Since asthma is not randomly distributed throughout the lung, it is possible that better results with this treatment will be obtained if imaging is used to guide the procedure.

Hyperpolarized helium-3 MR ventilation defects are also found in patients with CF, even those with normal spirometry. Similar to the results in asthma, there is some correlation between the number of ventilation defects and CF disease severity on spirometry. The number of ventilation defects has been shown to change with treatment including mechanical airway clearance, bronchodilator, and intensive antibiotic treatment. However, both mechanical airway clearance and intensive antibiotic treatment have been shown to increase rather than decrease the number of ventilation defects in some CF patients. The reason for this somewhat paradoxical response is not yet known but is certainly an interesting avenue for further research.

Diffusion imaging with hyperpolarized gas MRI provides information about the size and connectedness of the alveoli. In adults with COPD and in animal models of emphysema, the hyperpolarized gas apparent
diffusion coefficient (ADC) has been shown to correlate with disease severity in humans and alveolar size in animals. Bronchopulmonary dysplasia is a chronic lung disease that occurs in children with a history of premature birth and is also known as chronic lung disease of prematurity. Initially, this lung disease was thought to be the result of lung damage from the oxygen concentrations and high pressure ventilation used in the perinatal period to achieve adequate oxygenation in infants with very immature lungs. Improvements in the treatment of premature infants have decreased the mortality from lung disease but have not decreased the incidence of BPD. Most of alveolar development occurs during 25 to 40 weeks post conception and during the first few years of life. It is now thought that in addition to direct lung injury, in some premature infants something occurs that alters the development of the alveoli resulting in lung with too few alveoli that are too large in size. This has been confirmed in a study with hyperpolarized helium-3 diffusion imaging in which children with a history of BPD had an elevated ADC but similar to smaller lung volumes as compared with age matched healthy children.

In conclusion, MRI of the lung is already yielding new insights into pediatric lung diseases and has the potential to be a clinically useful tool in the assessment and treatment of children with lung disease.
Asthma is a common disease that comprises a large burden of disease in many countries. In Australia, it occurs in 10% of adults and 20% of children, a prevalence that has been stable for approximately 15 years. Asthma is a highly heterogeneous disease and may in fact be a cluster of distinct pathophysiological entities, which may be the basis of the so-called asthma ‘phenotypes’. The difficulty in better characterising asthma is that there are few tools that can distinguish clinically important differences in pathophysiology. However, Computed Tomography (CT) is a tool that has been able provide useful information with which to characterise the pathophysiology of asthma and as such, has provided a large amount of important information with which asthma can be characterised.

Computed Tomography provides structural information of the airways and lung parenchyma. The majority of CT studies have involved measurement of airway dimensions and of airway narrowing. Airway wall dimensions are difficult technically because of the thinness of the airway walls in relation to CT resolution and the variability of measurements is a limitation. Therefore, high-resolution technique is required to enable detection of the wall boundaries. To this end, both axial scanning and thin-slice helical scanning have been used. Despite the limitations of the measurements, results of studies have clearly and consistently shown asthmatic airway walls to be thicker compared with the wall thickness in COPD and with controls. This is in keeping with the findings from morphometric studies of post mortem lungs which show that asthma is characterised by marked airway wall thickening of small and large airways. Furthermore, wall thickening has been related directly to asthma severity, airway hyperresponsiveness and response to inhaled steroid treatment.

Acute airway narrowing induced by bronchial challenge has also been measured using thin-slice CT. Since subjects are required to move between the baseline scans and post-challenge scans to allow the challenge test to be done, the added technical complexity is an added source of measurement variation. Nevertheless, important insights can be gained from knowledge of the distribution of airway narrowing which has implications for the understanding of the pathophysiologic basis of airway hyperresponsiveness.

The combination of CT with functional measures has been used and will likely be the way forward in the study of asthma. Functional imaging provides data on regional ventilation in asthma. The pattern of ventilation disruption is relevant because it is apparent that both airway closure and heterogeneity of ventilation are relevant to asthma, potentially from both mechanistic and clinical expression points of view. CT does have the ability to provide some functional information, i.e. on ventilation and gas trapping. How functional information from different modalities compare with CT and to what extent they are complementary are still to be undetermined.

References:


Introduction

Lung diseases, including asthma and COPD, cause high morbidity and mortality, and immense economic expenditures. A component of effective diagnosis and management of chronic lung diseases is testing pulmonary structure and function. However, existing clinical tests—including radiography and high resolution CT, which both utilize ionizing radiation; conventional MRI, which sometimes employs chemical contrast agents; and global spirometry measurements taken at the mouth, which provide no regional information on lung function—all have serious limitations. An innovative MRI technology for lung imaging, hyperpolarized (HP) gas MRI, overcomes many of the limitations of these other modalities, and it offers an extremely promising complimentary tool for diagnosis, disease staging, and treatment evaluation for asthma and COPD.

HP gas MRI, which was pioneered in the 1990s (Albert et al. 1994), can elucidate structure and function previously undetectable with radiography or conventional MRI, and it uses no ionizing radiation or chemical contrasts agents, making it non-invasive and safe. This safety is particularly important for imaging asthmatics, because asthma disproportionately affects children, in whom exposure to radiation should be minimized. HP gas MRI also allows physicians to safely monitor how defects in ventilation progress over time, and how effectively specific treatments work over time.

Conventional MRI detects protons in water in tissues; HP gas MRI, on the other hand, detects atoms of \(^{3}\)He or \(^{129}\)Xe gas that are inhaled by the subject. Protons are rare in the airways and alveoli of the lungs, and thus conventional MRI of pulmonary ventilation is not viable. Although \(^{3}\)He and \(^{129}\)Xe are normally below the detection threshold for MRI, when these atoms are hyperpolarized through spin exchange with rubidium that has been optically excited by a diode laser, their detection increases about 100,000 times! With inhalation of HP \(^{3}\)He or \(^{129}\)Xe, high-resolution HP MR imaging of the lung airways and periphery is possible, providing a complete picture of lung ventilation with spectacular detail.

HP \(^{3}\)He MRI provides images of the lungs far superior to conventional MRI. Figure 1 compares conventional MRI with HP \(^{3}\)He MRI. Note the high level of detail in the \(^{3}\)He MR images. Also note that the \(^{3}\)He MR images clearly show differences in ventilation between the healthy (top right) and asthmatic (bottom right) subjects. Areas where gas is not reaching (known as ventilation defects) can be seen as dark areas in the image of the asthmatic subject on bottom right. This technique precisely pinpoints where gas is reaching and where it is not reaching, something other measurement modalities cannot do.

HP Gas MR Imaging of Asthma

The most commonly employed tools to assess lung function in patients with asthma are spirometry measures of global airflow and resistance. Because of its sensitivity, HP gas MRI can detect asthma at earlier stages, allowing treatment to be provided earlier, thus reducing the loss of lung function from which many asthmatics suffer (Altes et al. 2001). HP gas MRI is also totally changing the way we view asthma. The existing paradigm has been that some asthmatics have reversible obstruction, while others go on to develop fixed obstruction. However, HP \(^{3}\)He MRI studies suggest that asthmatics may have fixed obstruction in some lung regions much earlier in the course of their disease than earlier believed, and that disruption of airflow in asthmatics is regional, not diffuse.
HP gas MRI investigations have established that ventilation defects in asthmatics can persist for long periods in the same location. Figure 2 illustrates data from our group showing defects that persisted for over two months at baseline in a mild-moderate asthmatic. Note that pronounced defects (dark areas) persist between the scans. de Lange et al. (2007) gave 10 asthmatics a methacholine challenge to create bronchoconstriction, and then gave a second methacholine challenge a mean of 185 days later. Sixty nine percent of ventilation defects were in the same locations during both challenges. de Lange et al. (2009) found, in a study of 43 asthmatics, that 67% of defects persisted from the first baseline measurement to the second over a median interval of 41 days, and 38% of defects persisted from the first baseline measurement to the third over a median interval of 85 days.

Similarly, we looked at the number of defects that remained in the same location, and how they changed in size, over about 45 days at baseline and following administration of methacholine (Sun et al. 2010). We found that the total number of ventilation defects increased post-methacholine, as predicted, and that defects were found to be much more consistent in location during methacholine challenges than at baseline. (Post-methacholine, 96% remained in the same location between Day 1 and Day 2, whereas at baseline, only 75% of ventilation defects remained in the same location.) Data from all of these studies support the hypothesis that respiratory dysfunction in asthma has an important localized component.

Several studies have also shown that HP gas MRI measures positively correlate with asthma severity. Altes et al. (2001) found that defect number generally correlated with severity of asthma symptoms. We observed that HP gas MRI ventilation heterogeneity and ventilation defect volume correlated with asthma severity (Sun et al. 2009). Similarly, de Lange et al. (2006) observed that the number of defects was greater in patients with moderate and severe asthma than in patients with mild asthma. de Lange et al. (2006) also reported that many asthmatics with ventilation defects had normal spirometry readings (except for forced expiratory flow between 25% and 75% from the beginning of forced vital capacity); this provides dramatic evidence that HP gas MRI can detect ventilation impairment undetectable with traditional measurements, an invaluable clinical capability.

**Figure 2.** HP $^3$He MRI images of a mild-moderate asthmatic subject. Ventilation defects can be seen as dark spots.

**Figure 3.** HP $^3$He MR images of a mild-moderate asthmatic subject at baseline and following methacholine (Mch) challenges.

**Figure 4.** Ventilation maps ($b_1$–$b_3$) and heterogeneity maps ($c_1$–$c_3$) in a healthy subject. $b_1$ & $c_1 =$ pre-methacholine challenge; $b_2$ & $c_2 =$ post-methacholine challenge; and $b_3$ & $c_3 =$ post deep inhalations.

**Figure 5.** Ventilation maps ($b_1$–$b_3$) and heterogeneity maps ($c_1$–$c_3$) in an asthmatic subject. $b_1$ & $c_1 =$ pre-methacholine challenge; $b_2$ & $c_2 =$ post-methacholine challenge; and $b_3$ & $c_3 =$ post deep inhalations.
An important area of ongoing improvement in HP gas MR imaging is increasing quantification of signal characteristics. We have developed advanced techniques to quantify ventilation distribution and heterogeneity throughout $^3$He MR images (Tzeng et al. 2009). Figures 4 and 5 show ventilation maps and heterogeneity maps pre-methacholine, post-methacholine, and post-deep inhalations. In ventilation maps, colors represent ventilation levels relative to the maximum $^3$He per pixel (1.0) as seen in the trachea; in heterogeneity maps, the color represents an index associated with the coefficient of variation in ventilation levels for a region of interest around a given pixel. We found that in healthy subjects (Figure 4), ventilation heterogeneity appeared after the methacholine challenge, but was resolved by deep inhalations. In asthmatic subjects (Figure 5), ventilation heterogeneity appeared after the methacholine challenge, but was not resolved by deep inhalations.

Another important quantitative advance we reported was providing information about the percent ventilation in specific regions (Tzeng et al. 2009). Previous studies used only visual inspection to classify regions as defects, assuming that ventilation in a given region was either 100% ventilated or 100% unventilated. We now know that ventilation is not an either-or phenomenon—it varies along a continuum, and thus it is essential to provide spatial quantification of ventilation. Our group developed a technique to estimate the percentage volume occupied by fresh gas in each voxel of the lungs, providing greatly improved accuracy in estimating ventilation.

We also used dynamic $^3$He MRI to differentiate small ventilation defects (closure of only one terminal airway) from clustered defects (closure of two or more terminal airways). We measured 751 ventilation defects and found that 390 were small and 361 were clustered (Mullally et al. 2009). In addition, whereas previous studies used only visual inspection to classify defect location, we developed a technique to quantify ventilation defect location, a significant advance for the field of HP gas MRI.

A significant new direction in pulmonary imaging is using HP gas MRI to assess the physiological effects of pulmonary therapeutics. We have investigated the efficacy of several medicines for treating patients with lung disease. Figure 6 illustrates the power of HP gas MRI in detecting improvement in ventilation following administration of albuterol, a short-acting, beta-adrenergic agonist commonly used for treatment of asthma and COPD. This figure shows $^3$He MRI color-coded ventilation scans of a severe asthmatic at baseline (A), following bronchoconstriction induced by methacholine (B), and following administration of albuterol (C). Ventilation defects were increased by methacholine, and then reversed by albuterol.

**Figure 6.** $^3$He MRI color-coded ventilation maps from a subject with severe asthma. A = pre-methacholine challenge; B = post-methacholine challenge; C = post-albuterol administration.

**Figure 7.** $^3$He apparent diffusion coefficient (ADC) images (from Wang et al. 2008). Note the dramatic differences between the ADC images of healthy and COPD subjects at both short and long time scales.

**Figure 8.** HP $^3$He MR images of a subject before (left) and after (right) receiving arformoterol.
As with asthma, one of HP gas MRI’s most valuable capabilities is the information it provides on lung function before and after therapy. We performed HP ³He MR imaging on patients with COPD to assess the effects of arformoterol tartrate on ventilation function. Arformoterol is a long-acting, beta2 adrenergic agonist bronchodilator. We observed qualitative increases in ventilation in the lower half of the lungs in some subjects receiving arformoterol (see Figure 8). This illustrates how useful HP gas MRI can be in assessing the effect of drug administration on regional lung ventilation function.

**Future Directions in HP gas MR Imaging**

The future for HP gas MRI is exciting, with the prospect of wide adoption and increased translation to clinical applications. One new direction in HP gas MRI is further development of low-field MR scanners, which could have wide applications in instances where portability and/or low cost are essential. Another noteworthy new direction is a transition from ³He to ¹²⁹Xe. Xenon offers many advantages over helium: (1) it is cheaper and more widely available; (2) it can measure lung ventilation like helium, but because it readily dissolves in the blood, it can also be used as a perfusion tracer; (3) it is lipid soluble, making it ideal for imaging lipid-rich organs like the brain. Xenon is also heavier than helium, so it has a lower diffusion value. We look forward to seeing HP gas MRI help improve the lives of countless people with chronic pulmonary diseases.

**References:**

Over a period of fourteen years, the University of New Hampshire hyperpolarized gas academic group has been dedicated to improving technologies and methodologies for producing hyperpolarized 129-Xenon (HXe) [1-8] and acquiring high-quality images of pulmonary function [9-20]. Several technical obstacles to producing large quantities of highly polarized xenon were overcome with innovative developments. Our initial contribution was to recognize that significant improvement can be realized by pre-saturating the xenon and buffer gases with alkali vapor, then flowing the mixed gases through a long polarization cell opposite to the direction of laser propagation, and finally extracting the alkali vapor on a cold surface in the same cell in the presence of the laser. This three step process allowed beneficial utilization of the full polarizing power of the laser while simultaneously avoiding destructive non-linear thermal runaways. Our second contribution was devising a rising-dewar method for distributing the cryogenic extraction of polarized xenon from the gas mixture and subsequent thawing without losing polarization. Our third contribution was a stepped-mirror optical system to create multi-kilowatt, highly directional, spectrally narrowed laser beams for optical pumping. Our fourth contribution was demonstrating a finned-channel copper enclosure to efficiently polarize xenon in the presence of such intense laser beams while efficiently extracting the deposited heat. These four innovations combined to allow assembly and operation of xenon polarizers with productivity that exceeded earlier technologies by a factor of one hundred.

The first of our xenon polarizer prototypes, completed in 2004 at UNH, incorporated the first two of these innovations [2]. This prototype yielded a liter of 50% polarized xenon, sufficient to initiate human studies. At that time in 2004 we also founded a spin-off company Xemed to undertake development of additional polarizer prototypes. A second polarizer prototype built by UNH and Xemed together integrated the capability we had already demonstrated into a compact, partially automated package [1].

The third generation, “Einstein”, was implemented with two 12 bar stacks of 100W diodes and a copper polarizing column [3]. Although capable of free-running power of 2400W, we designed the laser for spectrally narrowed operation at 1400W. An afocal telescope with two converging lenses and magnification of four produces an image of the emitter bars on a grating in Littrow configuration [4]. Path lengths between off-axis bars and the angled grating are corrected using the step-mirror, allowing a perfect focus for all emitters [7]. The zeroth order beam is transported to the polarizing cell via another afocal pair, with cylindrical lenses providing different horizontal and vertical magnifications. The collimated beam is reconfigured using periscopes to match the square format of the polarizing cell. Flow of gases undergoing kilowatt heating will not be stable without effective heat removal. Cylindrical glass cells are insufficient for the task. We implemented a copper polarization cell with an internal finned structure to provide a factor four increase in heat removal surface area [8]. Because the directionality of the lasers is very high, the polarization can be performed in sixteen narrow long channels. The polarization cell is configured with two thermal zones along its length. As the gas flows through the warmer section, it interacts with the laser and increases its polarization. After reaching a polarization maximum, the gas moves into the cooler region where it loses its rubidium. With a high power laser and finned copper polarization cell, Einstein is capable of producing several liters of hyperpolarized xenon in twenty minutes.

During four months in the fall of 2009, we tested the clinical utility of this third generation xenon polarizer prototype at the University of Virginia. For these measurements we assembled a 32-element parallel receive coil and mated it to a highly-uniform birdcage transmit coil with an asymmetric shape to fit into the 3T Siemens scanner [9]. Six hundred measurements were performed on 64 subjects, including healthy volunteers and patients with COPD, asthma, and cystic fibrosis. Roughly a dozen imaging protocols were demonstrated, some for the first time. We acquired ventilation images with resolution in the range 2mm to 3mm within a ten second breath hold using GRAPPA acceleration [10, 11]. We studied diffusion weighted imaging at short and long distance scales in lungs of COPD patents [12]. Image sets with a breath hold were used to determine xenon T1 longitudinal relaxation in order to map regional variations in alveolar oxygen partial pressure [13]. In addition to these gas-phase protocols,
versatile suite of imaging protocols was developed to interrogate the xenon dissolved phase [14-18]. The Xenon magnetization Transfer Contrast (XTC) protocol senses xenon exchange between the gas and dissolved phase by repeatedly quenching the dissolved phase and imaging the changes in the gas phase polarization [21]. We studied the XTC protocol for various choices of frequency range of the dissolved xenon, allowing us to isolate xenon exchanging between alveolar gas and red blood cells [14, 15]. We also acquired sets of images using the XTC protocol for a range of choices of the exchange time, extracting the rate of saturation of the xenon compartments [16, 17]. Finally, we also performed imaging of the gas and dissolved phase directly, together in a single pulse sequence [18]. A region of hyperintensity on the dissolved-phase image of the lung of a healthy volunteer, which corresponded to hypointensity on the xenon ventilation image, was later attributed to an inflammatory lesion, confirmed by proton MRI.

Over the past year (2010) we assembled our fourth generation prototype, "Hawking," improving on operational methodology and incorporating new technology. The sixteen channel finned copper column remains unchanged, however heating and cooling systems are implemented with improvements in stability and robustness. The 1400W spectrally narrowed laser is redesigned, with a new optical layout and a mechanically stabilized step mirror. We also incorporated deflection and focusing corrective optics for each of the 1440 emitters. The cryogenic separation system incorporates larger volume helixes, a diaphragm pump to scavenger polarized xenon from the freeze-out volume and deliver it into the bag, and a four-port bagging system.

In addition to these technical improvements, the fourth generation polarizer offers improvements in operation, safety, stability, and reproducibility. All processes are fully automated including the production sequence, polarization measurement, warm-up, cool-down, and self-check. Quality control is incorporated throughout. Each bolus bag is bar-coded and labeled with its Investigational Drug status, its quantity of hyperpolarized xenon, the polarization, and a time-stamp. On-board diagnostics are incorporated into every subsystem and data are logged and stored archivally. The polarizer can be controlled locally or remotely. The laser path is fully enclosed and interlocked, permitting Class 1 designation. The frame is designed to be aesthetic, robust, easily transportable on its wheels, or able to be packed for shipping. Operator training will no longer require advanced degree in physics. The technician simply pushes a soft button on the polarizer’s touch screen or schedules a bag of gas with the polarizer online.

Xemed is expanding partnerships with research sites for validating clinical indications towards regulatory approval of hyperpolarized xenon and seeking funding mechanisms to support those activities. In the US, the National Institutes of Health is continuing its support to Xemed for refinement of polarization and imaging infrastructure, development of new pulse sequences, and investigations of clinical indications. A recently approved grant will add a clinical site to the network of Xemed’s academic partners. Xemed is pursuing joint development agreements with pharmaceutical and medical device companies. One such project recently funded will develop quantitative hyperpolarized xenon imaging biomarkers for lung disease and compare with other metrics. With the recent completion of our latest polarizer, we now have a new resource for pursuing future clinical work and advancing the regulatory status of hyperpolarized xenon worldwide.

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Hyperpolarized noble gas magnetic resonance imaging has been explored extensively as a promising tool for the quantitative evaluation of regional pulmonary pathophysiology. This noninvasive technique is capable of providing both structural information down to the level of the alveolar microstructure and functional information, such as dynamic ventilation, intrapulmonary partial pressure of oxygen, and alveolar surface area. In this lecture, the role of hyperpolarized 3-helium and 129-xenon magnetic resonance imaging for clinical applications, current and in the future, will be reviewed.

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Abstract

Evening Lecture
Understanding how the mammalian respiratory system works and how it changes in disease states and under the influence of drugs is frequently pursued in model systems such as small rodents. Indeed, the field of lesion modeling is progressing at an accelerated pace and creating mouse models that accurately reproduce the genetic and histopathological alterations present in human diseases. Animal models of human pulmonary disorders have been very valuable tools for proof of concept studies, for investigating respiratory biology and for increasing our knowledge about disease mechanisms. Given the heterogeneity of disease and the slow progression of chronic obstructive pulmonary disease and interstitial lung disease, a pilot clinical study would be difficult to design and costly to carry out. More importantly, histopathological correlation is requisite for development of new technologies to test the sensitivity and specificity of the methods. For those reasons, animal study is important particularly in this field of research. Hence, one needs the ability to assess morphology and function in vivo systems.

High-resolution computed tomography (HR-CT) is the standard for assessing pulmonary disorders, which yields in detailed anatomical information but incurs radiation exposure. It has been identified that an important challenge in pulmonary disorders is the development of more powerful, multi-variate methods for predicting regional outcome and individual responsiveness to particular therapies on the basis of clinical and laboratory characteristics.

One of our major research focuses is to exploit the recent advances in magnetic resonance imaging (MRI) to image the lung in vivo. MR imaging of the lung is extremely challenging due to multiple inherent difficulties associated with the properties of the lung. These challenges, including low proton density, severe magnetic field susceptibility, and respiratory and cardiac motion artifact, have resulted in the delay in refining lung imaging as compared to MRI progress with other major organs. However, the recent development of fast MR imaging techniques along with more powerful hardware has made possible detailed, non-invasive imaging of pulmonary parenchyma, and pulmonary MRI is currently an active area of research. We have during the past years been dedicated to the development of new acquisition and processing methods by means of MRI in animals and humans, permitting quantitative characterization of the pathophysiological change in pulmonary disorders, including lung cancer.

The utility of ultra-short TE (UTE) MRI in conjunction with projection acquisition of the free induction decay (FID) allows us to reduce TE to less than 100 μsec to minimize signal decay caused by short transverse relaxation time (T2*). It brings higher MR signals from tissues which have short T2* and thus are unobservable in conventional methods with short TE (c.a. 2 msec). On the subtraction image between the images with an UTE (c.a. ~500 μsec) and a conventional range of TEs, the biological materials/tissues, such as tendon, bone or periostea are enhanced. In early 90s, it was demonstrated that UTE imaging with projection reconstruction provided much higher signal intensity (SI) in the lung parenchyma and finer pulmonary structures in human. Using cardiac and respiratory gating, it was showed excellent UTE images of the lung in normal rats using a small animal imaging system. It was also reported the ultra-high quality UTE images of the normal rat lung in vivo that resolve the thin membranes that separate lung lobes. Another study using a single point imaging method that could achieve an ultra-short TE (200 μs) demonstrated lower MR signal and shorter T2* in a mouse model of emphysema than those in the control. However, utilization of UTE MRI for lung imaging is still sparse in these days.

In this lecture, we will first discuss the principal of UTE MRI. Subsequently, our systemic approach to use UTE MRI for pulmonary imaging will be introduced. Using an UTE sequence, we have demonstrated in normal and a transgenic murine model that the parenchyma with airspace enlargement exhibits reduced SI and T2* that closely relate to lung tissue density. Further, the capability to acquire endogenous MR signal of the lung parenchyma also enables us to assess change in SI due to inhalation of molecular oxygen (O2) or intravenous injection of Gd, which demonstrates regional ventilation and perfusion, respectively. The method could detect the regions with ventilation/perfusion mismatch in a model of pulmonary embolism. The method can be implemented in a standard clinical system and does not require any special equipment and is directly translatable to clinical use. The requisite image processing is simpler compared with currently existed methods. The UTE MRI has the new potential to assist in the diagnosis of early lung disease and in monitoring of response to treatment without incurring the risks of radiation exposure.
Magnetic resonance imaging (MRI) of the lung has been considered experimental for many years. With ongoing technical improvements such as multichannel MRI, systems with powerful gradients as well as the development of innovative pulse sequence techniques e.g. implementing parallel imaging, MRI of the lungs was finally clinically implemented. Nowadays MRI is routinely used for lung imaging in clinical practice as a radiation-free alternative to computed tomography (CT), e.g. in children with lung disease. In addition to a morphologic assessment, MRI offers the great potential of functional lung imaging. In clinical medicine this may help in the differential diagnosis of pulmonary disorders, improve disease stratification – and thus improving therapy selection, and finally giving insights into pathophysiology. Several functional MRI techniques have been developed, among which contrast-enhanced perfusion imaging, diffusion weighted imaging (DWI), and dynamic MRI of respiratory mechanics can be clinically implemented on almost every modern MR system. Less commonly used or more complex MR techniques are oxygen-enhanced MRI and hyperpolarized noble gas imaging.

For a long time imaging of lung perfusion was the domain of nuclear medicine. With the development of time-resolved MR angiography (MRA) techniques perfusion imaging of the lung became clinically feasible. Perfusion imaging has been used for the visualization of regional pulmonary perfusion in patients with different lung diseases such as lung cancer, airway disease or pulmonary vascular disease. In combination with morphologic MRI perfusion imaging allows a comprehensive assessment of those lung diseases thereby improving differential diagnosis (e.g. the differentiation between chronic obstructive pulmonary hypertension and other forms of pulmonary hypertension). Using perfusion as a surrogate parameter of lung function perfusion MRI can be used to objectively quantify the functional impairment in pediatric lung disease.

After two decades of diffusion-weighted imaging for the evaluation of intracranial diseases, DWI of the thorax has become feasible by implementation of MRI techniques with significantly reduced acquisition times, thus minimizing artifacts from respiratory and cardiac motion. In chest imaging, DWI has been mainly suggested for the characterization of lung cancer, lymph nodes and pulmonary nodules. DWI visualizes the microscopic movement of water molecules within tissues which is highly influenced by the cellular environment of water, intracellular organelles and macromolecules. As water molecules move within tissues, they encounter various restrictions and hindrances. Thus, DWI allows for a functional assessment of microstructure in relation to gross anatomy. The movement of the spinning water molecules causes phase dispersion, which results in a signal intensity loss. This signal intensity loss can be quantified by calculating the apparent diffusion coefficient (ADC). Malignant tumors are characterized by increased cellularity, larger nuclei with more abundant macromolecular proteins, a larger nuclear/cytoplasmic ratio and less extracellular space relative to normal tissue. Because of these characteristics, the diffusion of water molecules in malignant tumors is restricted, resulting in decreased ADC. DWI has been used to differentiate lung cancer from benign lesions as well as for the subcharacterization of different histologic subtypes of lung cancer. Moreover, DWI has been used to differentiate central tumor from the surrounding lung consolidation, and as alternative imaging test for the detection of nodal and distant metastasis.

Dynamic CINE MRI techniques have traditionally been used for the assessment of cardiac function. By applying these techniques to the entire chest respiratory mechanics can be visualized in vivo. Several applications of dynamic MRI in pulmonary imaging have been described such as radiotherapy planning, cancer staging or assessment of airway and lung motion in airway disease.

Compared to standard field strength (i.e. 1.5 Tesla) high field MRI (e.g. 3 Tesla) has the advantage of an improved SNR, since the signal scales linearly with field strength. Moreover, for contrast-enhanced applications, the sensitivity to injected Gd agents is heightened since the longitudinal relaxation time (T1) of unenhanced blood increases with field strength. At MR angiography shortening of T2* causes a reduction of background noise, thus resulting in a better discrimination of enhancing blood vessels. In the lungs, however, the higher magnetic susceptibility gradients associated with high magnetic field can be limiting, resulting in signal loss and potentially lower image quality when compared to standard field strength MRI. The increasing number of installation of high field MR systems necessitates functional MRI of the lungs to be implemented at high field. This lecture will give an overview of the current status of functional MRI of the lungs at high field and present some clinical examples.
Dual-energy CT (DECT) is a new, emerging, imaging technique which can differentiate such materials as iodine, xenon, bone, and air, based on their specific attenuation differences at high and low x-ray energies. Dual-source CT (DSCT) allows acquisition of both high-energy CT data and low-energy CT data during a single scanning by applying two different tube voltage settings, typically 140 kVp and 80 kVp. Clinical DECT application using first generation of DSCT was somewhat limited due to several technical factors including limited field-of-view and energy differentiation. These limitations have been significantly improved by the introduction of second generation dual source CT with larger field-of-view and better dual-energy contrast. Recently, alternative approach of DECT has been introduced by different manufactures, which is based on rapid kV switching method using single x-ray tube. Although this new technique is somewhat different from dual-source approach in image processing and evaluation algorithm, this approach will also allow the similar clinical applications.

For the past few years, there have been several studies on the thoracic applications of this interesting technique. One of typical applications is perfusion or blood volume imaging, which is based on the ability of extraction of iodine content from contrast enhanced CT images. With application of the material decomposition theory, dual-energy CT can be used to differentiate the iodine component from lung parenchyma enhanced with iodine contrast material. Materials can be differentiated by analysis of changes in attenuation coefficients in materials at two energies. Owing to the photoelectric effect, the material decomposition theory is especially applicable to materials with large atomic numbers, such as iodine, and iodine is generally known to have stronger enhancement at low tube voltage settings. At dual-energy CT, pulmonary CTA and the iodine enhancement representing lung perfusion can be performed simultaneously in a single acquisition after contrast injection.

Studies have shown that perfusion images obtained at dual-energy CT may be useful adjunct in diagnosing acute pulmonary embolism. The perfusion defect was found in the corresponding area of the occluded vessel, and visualization of the perfusion defect with dual-energy CT had good agreement the findings at perfusion scintigraphy. In addition, Chae et al, reported that the degree and extent of perfusion defects can be sued in severity assessment of acute pulmonary embolism (AJR 2010). In segment-based analysis in their study, they found interesting mismatch between the vascular occlusion and perfusion defect. Perfusion was preserved in the segments in which there was partial occlusion above the segmental artery. Therefore, these segments may represent partial occlusion with normal perfusion. Conversely, there were segments having perfusion defects more severe than the degree of vascular obstruction. One assumption that may explain the mismatched areas is that they represented areas of microcirculation disturbance.

Another way to use DECT in assessment of acute pulmonary embolism is so called ‘lung vessel’ application, allowing differentiation of small thrombosed small vessels from contrast-filled non-thrombosed vessels even with the presence of partial volume effects. Despite the presence of partial volume effects, the ratio of the respective low and high kV CT-values relative to air (negative 1000 HU) is only a function of the iodine content but not of the vessel diameter. Using this concept, the ‘Lung Vessels’ application software can divide the pixels of lung vessels into those with iodine and those without iodine by means of a straight line in the CT-value diagram. Lee, et al has recently reported that the diagnostic performance of pulmonary CTA in diagnosing acute PE can be improved by applying this interesting application (Eur Radiol 2010). This effect was more pronounced in detecting small, peripheral emboli.

DECT pulmonary perfusion map can also allow for the depiction of perfusion defects in patients with chronic thrombembolic pulmonary hypertension. Perfusion imaging is also useful in the assessment of other vascular diseases including congenital vascular anomalies, vasculitis and so on. This technique also may replace perfusion scintigraphy for preoperative assessment of lung cancer patient, because perfusion CT was comparable to perfusion scintigraphy in the prediction of postoperative lung function.

In addition, perfusion imaging can be also applied in assessing the perfusion status of various diffuse lung diseases including COPD, smoker’s lung, interstitial lung diseases, and so on. More importantly, xenon ventilation DECT imaging for the simultaneous assessment of anatomy and ventilation has been introduced (Chae et al, Radiology 2008 / Invest. Radiol 2010). Possible clinical indications of this new
approach include the assessment of COPD, asthma, and other diffuse airway and vascular disease. Simultaneous assessment of anatomy, perfusion and ventilation using DECT technique may open new fields of functional CT imaging of various pulmonary diseases.

In this talk, firstly, the basic principles and theory of dual energy CT imaging will be introduced. Early experiences in using dual energy CT in various pulmonary vascular diseases will be discussed.
Quantitative assessment of pulmonary blood flow may be able to provide tremendous insight into the mechanisms of pulmonary vascular disease. In this presentation, we will discuss aspects of normal pulmonary physiology that affect the quantitative assessment of pulmonary blood flow, including respiration and gravitational affects on lung structure and blood flow. We will then examine both contrast and non-contrast methods that can be used to quantify the various parameters that define pulmonary perfusion. Lastly, we will examine the application of these methods to the study of pulmonary vascular disease using the study of high-altitude pulmonary edema (HAPE) as a model that has been investigated using both contrast and non-contrast techniques.
Abstract
Core Session
Pulmonary hypertension (PH) has been defined as a resting mean pulmonary arterial pressure > 25 mm Hg. The subgroup of PH known as pulmonary arterial hypertension (PAH) adds the criterion that the pulmonary arterial wedge pressure (PAWP) is ≤ 15 mm Hg. The prevalence of PH varies substantially depending on the type, etiology and underlying condition, such as idiopathic PAH (iPAH), scleroderma, portal hypertension, pulmonary embolism and so on. Right heart catheterization with the measurement of pulmonary arterial pressure (PAP), cardiac output (CO), right arterial pressure (RAP) and PAWP remains the gold standard for the diagnosis of PAH.

Although the tools used to diagnose and assess patients with PAH have improved, the number and complexity of therapeutic interventions have increased, making it even more challenging to accurately predict prognosis. With the introduction of Doppler echocardiography, approximate evaluation of PAP became feasible. Although this method is helpful in detecting or excluding significant PH, its intrinsic and operator-dependent limitations make early PH diagnosis and screening challenging. In addition, the sensitivity and specificity of echocardiography for the detection of PH remain undefined.

The usual cause of death in PAH is right ventricular (RV) failure. Both diastolic and systolic dysfunction are likely contributors to RV failure. Diastolic RV dysfunction is thought to be related to RV hypertrophy and chronic pressure overload, with prolonged isovolumetric relaxation time. With systolic RV dysfunction, isovolumetric contraction time is prolonged and ejection time is shortened. Magnetic resonance imaging is useful in the noninvasive assessment of the right heart in PH, although it cannot be used as a screening test.

CTEPH should be considered for all patients with unexplained PH because there are many treatment strategies for it. Pulmonary perfusion and ventilation scintigraphy has been used for the differential diagnosis of chronic pulmonary thromboembolism from other variable diseases which cause PH. Ventilation-Perfusion (V/Q) lung scan of patients with CTEPH generally show multiple segmental-sized or larger mismatched perfusion defects. SPECT acquisition is recommended to detect a small perfusion defect. Perfusion scan for CTEPH have a tendency to underestimate the severity, although negative scan results are highly specific for absence of thromboembolism. A normal V/Q scan means that CTEPH is unlikely cause of PH.

In Japan, latest and advanced CT scanners have been widely used across the country, and most of them are available 24 hours. This accessibility made SPECT unnecessary in the APTE diagnosis. Ventilation scan are recently often left out in the diagnostic procedure. However, the diagnosis on perfusion SPECT without ventilation SPECT may cause false positive results of other all pathological conditions which cause regional alveolar hypoxemia –COPD, for example. In such cases, comparison with CT for each perfusion defect is requisite to avoid such misdiagnosis. Simultaneous evaluation using SPECT/CT is considered desirable.

V/Q SPECT has advantages over CTPA in terms of free complications induced by contrast medium or an excessive radiation exposure to the breast. Diagnosis in accordance with proper plans should be made: V/Q scanning must be performed in all patients with unexplained PH to rule out CTEPH. If V/Q scan reveals suggestion of CTEPH, pulmonary angiography or CTPA is required for an accurate diagnosis.
Pulmonary hypertension may be idiopathic or arise in association with chronic pulmonary embolism, parenchymal lung disease, liver disease, vasculitis, congenital left-to-right cardiac shunt, pulmonary veno-occlusive disease, left-sided cardiac disease and others. With the advance of multi-detector technology, CT has become one of essential test to evaluate pulmonary hypertension. One of the key roles of CT pulmonary angiography is to differentiate the causes of pulmonary hypertension because each disease has somewhat different imaging features. CT also can be used to assess the severity of the diseases. In addition, CT can be used in to predict the effect of vasodilators in treating idiopathic pulmonary hypertension. CT angiography also can be used in selecting candidates for thromboendarterectomy in treating chronic thromboembolic pulmonary hypertension. In this talk, I will be focusing on the technical aspects of CT imaging for assessing pulmonary hypertension and the typical and atypical CT features of various causes of pulmonary hypertension. In addition, recommendations for clinical use will be given. Lastly, recently introduced pulmonary perfusion CT imaging with dual energy CT technique will be introduced and early experience in using dual energy CT in pulmonary hypertension will be discussed.

In this presentation, we will discuss various methods that have been used in the evaluation of pulmonary arterial hypertension (PAH). These methods can be divided into those that evaluate the proximal pulmonary arteries and those that examine the peripheral circulation. Within the proximal pulmonary arteries, correlations have been made between patterns of arterial blood flow and pulmonary artery pressure. In addition, the mechanical properties of the pulmonary artery have also been shown to correlate with pulmonary arterial pressures. Quantitative evaluation of the peripheral pulmonary circulation may identify differences in pulmonary blood flow, blood volume, or mean transit time between normal subjects and patients with PAH. Qualitative evaluation of the peripheral pulmonary circulation has been shown to be useful in distinguishing chronic thromboembolic PAH from non-thrombotic causes.
Pulmonary hypertension (PH), is defined at cardiac catheterisation as a mean pulmonary artery pressure of ≥ 25 mmHg. It has a variety of mechanisms and causes which are classified into 5 major clinical groups (Dana Point). In patients with pulmonary arterial hypertension (PAH) and chronic thromboembolic pulmonary hypertension (CTEPH) life expectancy is significantly reduced. Since the first randomised controlled trial in 1996 in idiopathic PAH approximately 30 RCT’s have been published. Currently licensed therapies for PAH include drugs that target prostanoid, nitric oxide and endothelin pathways. For a selected group of patients with CTEPH, pulmonary endarterectomy can offer the prospect of cure. Drug therapies for PAH are expensive and although they have been shown to improve exercise capacity, quality of life and slow progression of disease there remains no cure. Long term data is emerging from various registries and suggest that these therapies also improve survival. A number of therapies targeting other important pathways are currently under development / trial. In the 3 other forms of PH such as that owing to, lung disease / hypoxia (PH-Lung) to left heart disease (PH-LHD) and that with unclear / multifactorial mechanisms (PH-Multifactorial) although PH is associated with a worse outcome and functional status there is lack of efficacy and safety data. Therapies suitable for PAH are not recommended in these patients. Radiological assessment is crucial in the systematic evaluation of patients with unexplained PH, in the accurate classification of PH and therefore in defining patients most likely to benefit from specific interventions.
Core Session 1 - Scientific Presentation 1 -

**Optimized Flowchart of Multi-technical MR Protocol for Pulmonary Circulation and Right Heart Function on the Busy Clinical Schedule**

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**Purpose:**
There are various techniques of current MR imaging to noninvasively evaluate the pulmonary circulation and right heart function. The evaluation for this specific purpose based on electively one or two techniques may be difficult in clinical practice because of the lack of full information available. The combination of multiple scanning techniques without an adequately designed flowchart, however, may expand the scan time for the overloaded MR schedule. The purpose of our study was to evaluate the feasibility of a multi-technical full set MR protocol for clinical usage.

**Material and Methods:**
To resolve various clinical problems about pulmonary circulation and right heart function, we designed a MR protocol with multiple techniques using the 1.5 Tesla MR scanner. This set included (1) tissue characteristics study using steady-state precession (true FISP) and black-blood fast low angle shot gradient echo (FLASH) (2) dynamic gadolinium-enhanced MR pulmonary perfusion evaluation using FLASH (3) right heart function evaluation using serial short-axis cine true FISP (4) gadolinium-enhanced 3D MR angiography, and (5) phase contrast MR pulmonary flow measurement. The expected scanning time to cooperative patients was established by four experienced MR technicians after inspecting the protocol.

**Results:**
The expected scanning time from four technicians was 30 minutes, 30 minutes, 35 minutes, and 40 minutes respectively. All of them were within the default setting (40 minutes per MR case) in our hospital.

**Conclusion:**
Our optimized multi-technical full set MR protocol for the evaluation of pulmonary circulation and right heart function is feasible with acceptable scanning time.

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Core Session 1 - Scientific Presentation 2 -

**V-Q Imbalance in Primary/Passive Pulmonary Hypertension on V/Q SPECT**

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Cross-sectional lung ventilation (V)-perfusion (Q) im balance in primary pulmonary arterial hypertension (PAH) and passive pulmonary hypertension (PH) was characterized by automated V/Q SPECT.

Technegas/MAA SPECT-derived V/Q SPECT and V/Q profile were automatically built to characterize cross-sectional lung V-Q imbalance in 12 patients with primary (idiopathic or familial) PAH and 15 patients with passive PH associated with left ventricular dysfunction or failure. The abnormality of V/Q distribution in these patients was correlated with PaO2 and pulmonary arterial pressure and with lung morphologic changes on CT.
Markedly low V/Q ratios (reverse V-Q mismatch) in the background lungs with heterogeneous V/Q distribution was seen in 12/12 (100%) patients with primary PAH and in 10/15 (66%) patients with passive PH, which were predominantly seen in upper lung zone. Including these regions with reverse V-Q mismatch, V/Q profile frequently showed flattened peaks with asymmetric and broadened V/Q distribution in all patients, with significant correlation between the standard deviation (SD) of V/Q ratios in the entire lungs and PaO2 and mean pulmonary arterial pressure (both; P < 0.01). At the lung regions with reverse V-Q mismatch, bronchial lumens compressed by dilated pulmonary arteries and heterogeneous lung attenuations were frequently seen on CT.

Patients with primary PAH and passive PH appear to characteristically have a high prevalence of reverse V-Q mismatch indicative of an inadequate hypoxic vasoconstriction reflex on V/Q SPECT, frequently accompanied with heterogeneous lung attenuations and compressed airways on CT.

Cardiac Output (CO), Pulmonary Artery Pressures (PAP) and Pulmonary Vascular Resistance (PVR) are key diagnostic variables in Pulmonary Hypertension (PH). The current gold standard technique for the measurement of these variables is Right Heart Catheterisation (RHC). Well validated non-invasive alternative techniques using Echocardiography or MRI exist for estimating both CO and PAP, however at the present time no validated method exists for the non-invasive estimation of PVR.

In this study 21 patients with CTEPH, IPAH or Systemic Sclerosis with or without PH were selected who had undergone RHC and Gadolinium Contrast Enhanced Pulmonary MR imaging within a 72 hour period. Using each patients MRI data functional maps were created showing the passage of contrast through each pixel and Regions of Interest drawn outlining the complete lung field on three sample coronal slices. The Contrast Transit Time (CTT) from the contrast bolus being detected in the Superior Vena Cava (SVC), Right Atrium (RA), Right Ventricle (RV) and Pulmonary Artery (PA) to the time of peak MR signal in each sample slice was recorded and compared with the PVR using a Pearson’s Rank Correlation.

Correlation Coefficients were highest in the most posterior sample slice (level of the Descending Aorta) with a correlation coefficient between PVR and CTT from the SVC of r=0.922, p<0.001, no correlation coefficients of less than 0.801 were recorded in any sample.

CTT appears to correlate strongly with PVR and may serve as a non-invasive measurement of this clinically important physiological variable in patients with PH.
Lung cancer is the leading cause of cancer death worldwide. Despite progress of platinum-based chemotherapy, the median survival time of patients with inoperable non-small cell lung cancer (NSCLC) still remains poor. The median survival time is approximately 14 months, and has improved only marginally over the last few decades. A recent advance in the treatment of inoperable lung cancer is the development of molecular-targeting drugs including gefitinib and erlotinib targeting the epidermal growth factor receptor (EGFR). Anti-angiogenic drug, such as bevacizumab, in a combination with chemotherapy has become clinical use in the treatment for NSCLC. Moreover, discovery of EML4-ALK made the great progress in cancer research in NSCLC. In this lecture, the development of molecular-targeting drugs, such as EGFR-TKI, anti-angiogenic drug, and EMLA-ALK inhibitors will be reviewed.

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Lung cancer is the number one cause of cancer death. Most cases are found after distant metastasis, and outcome of drug therapy for these patients used to be disappointing. However, we have faced a new paradigm shift, i.e., the molecular targeted therapy and the individualized therapy. Many promising data have reported from not only western countries but also Asian countries such as the efficacy of EGFR TKI to tumors with mutated EGFR, that of ALK inhibitor to tumor with EML4-ALK fusion protein, and that of pemetrexed to non-small non-squamous cell lung cancers. New questions have emerged from these new evidences derived from some important clinical trials. Among them, questions regarding with ethnic difference would be one of the most important issues. Are survival data same between Asian and Caucasian? (Data from Japan Lung Cancer Registry Study as well as some global trials have shown survival of Asian patients with lung cancer appears to be obviously better than that of Caucasian.) Why EGFR mutation is frequent in Asian patients? Is only EGFR gene status related with prognosis of lung cancer? Is the criteria of pathological diagnosis for lung cancer same between Asian countries and western countries?

Here, newest evidences for treatment of lung cancer will be presented, and importance of ethnic difference and Asian trials will be discussed.

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Lung cancer is the leading cause of cancer death worldwide. Despite progress of platinum-based chemotherapy, the median survival time of patients with inoperable non-small cell lung cancer (NSCLC) still remains poor. The median survival time is approximately 14 months, and has improved only marginally over the last few decades. A recent advance in the treatment of inoperable lung cancer is the development of molecular-targeting drugs including gefitinib and erlotinib targeting the epidermal growth factor receptor (EGFR). Anti-angiogenic drug, such as bevacizumab, in a combination with chemotherapy has become clinical use in the treatment for NSCLC. Moreover, discovery of EML4-ALK made the great progress in cancer research in NSCLC. In this lecture, the development of molecular-targeting drugs, such as EGFR-TKI, anti-angiogenic drug, and EMLA-ALK inhibitors will be reviewed.
In current practice, staging and follow-up of lung cancer is based on MDCT (morphology), PET (metabolism), scintigraphy (ventilation/perfusion/bone metastases) and MRI (brain). This arsenal of modalities is challenged by MRI. It offers superior soft tissue contrast for differentiation of tumour/atelectasis, mediastinal involvement or chest wall invasion. MRI is superior to X-ray and matches CT in detection of nodular and infiltrative lung disease. In whole-body staging, the diagnostic accuracy of MRI matches even PET-CT - plus comprehensive functional imaging capacities without radiation exposure to patients or hospital personnel.

Functional MRI of the lung presently comprises tumour- and lung contrast-enhanced perfusion studies, diffusion-weighted imaging and analysis of respiration dynamics. Non-contrast enhanced, motion-corrected V/Q-MRI and arterial spin labelling are under development. Their value for monitoring tumour viability and metabolism in response to treatment is subject to current investigation. In radiotherapy, refinements of target definition and dose delivery based on regional lung tissue characterization with ventilation/perfusion MRI are being discussed. Furthermore, concepts for radiotherapy of organs with respiratory motion have raised the need for dynamic volume lung imaging. The CT-approach, so-called 4D-CT in helical or cine mode, either with prospective acquisition or retrospective reconstruction, leads to long exposure times and high radiation dose. It is therefore limited to very few indications. Instead, 4D-MRI with parallel imaging and echo sharing allows for repeated and prolonged measurements of lung motion and tumour displacement without radiation exposure, which makes it the future tool for patient selection in motion adapted radiotherapy and for scientific work in the field.
Oncologic imaging has made tremendous progress over the past decades and targeted treatment strategies have been suggested to improve patients’ outcome. Objective criteria defining treatment responses are essential to evaluate the treatment effect of anticancer drugs. Currently imaging biomarkers approved for drug development are based on measurements of size in one dimension. Volume-based size change using the volumetric imaging with MDCT and the application of computerized segmentation of tumors might provide more reproducible measurements and a more sensitive scale for change assessment compared with unidimensional measurements.

Tracer kinetic modeling of dynamic imaging data enables estimates of blood volume, blood flow, and capillary endothelial permeability. With the development of new targeted therapy, selection of eligible patients for these targeted drugs and patient monitoring during the treatment require the availability of noninvasive tools that are capable of assessing tumoral neovascularization. Studies have shown significant changes in perfusion parameters after treatment and have suggested that early prediction for the treatment response may be possible. The availability of rapid imaging with MDCT and commercial analysis software has made perfusion imaging with CT feasible. Perfusion imaging can be performed as an adjunct to a conventional CT examination and is therefore particularly appropriate when a conventional CT is part of routine clinical protocols.

These new techniques need to be validated for their reproducibility and should be standardized for multicenter trials.
Comparative Reading System for Lung Cancer CT Screening

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Lung cancer kills more people than any other cancer worldwide. Lung cancer screening using low-dose CT have been performed in many countries. Comparative reading of current and past CT images is important for evaluation of pulmonary nodules in lung cancer CT screening. However, primary problem in comparative reading is the mismatch of slice and nodule positions caused by lung variation. It is hard for physicians to manually match slice positions, nodule positions, and evaluate the nodule's degree of change. A system to assist smooth comparative reading is necessary. We proposed a comparative reading system for lung cancer CT screening. It's distinctive feature is a highly accurate matching method of region of interest based on thoracic organs registration. Pulmonary blood vessels registration using analysis of the tree structure is performed. The system is evaluated using 1mm and 2mm slice thickness CT images obtained from lung cancer CT screening. We show its usefulness for lung cancer CT screening.

The Evaluation of Lung Perfusion in a Radiation Field Through the Radiation Therapy for Lung cancer - Using Lung Perfused Blood Volume Imaging with Dual Energy CT -

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Purpose
To evaluate the changes of lung perfusion in a radiation therapy (RT) field using lung perfused blood volume (PBV) imaging with dual-energy CT (DECT).

Materials and methods
19 patients receiving RT for primary lung cancers underwent dual-source DECT (SOMATOM Definition Flash). Lung PBV images were generated at 4 points, before, during, 1 week after and 1 month after RT. Follow-up periods was 3-11 months (median 7 months). Radiation dose was 60-70Gy/30-35 fraction. Normalized lung PBV (nPBV) images were created by considering heterogeneous structure of lung parenchyma. The changes of nPBV were analyzed: 1) inside and outside of RT-field, 2) in patients with radiation pneumonitis within 1 month after RT and in patients without pneumonitis.

Results
Radiation pneumonitis was observed in 12/19 patients (63%) within 1 month after RT and in 3/19 (16%) later, and no radiation pneumonitis in 4/19 (21%) in all periods. 1) nPBV decreased 1 month after RT compared with before RT (p=0.0002) inside of RT-field, but didn't change outside in all periods. 2) For patients with early radiation pneumonitis, nPBV increased 1 week after RT compared with before RT (p=0.034) and decreased 1 month after RT compared with during RT (p=0.02) or 1 week after RT (p=0.002). For patients without radiation pneumonitis, no relation was found.

Conclusion
The changes of nPBV decreased 1 month after RT for all patients, and increased 1 week after RT for early radiation pneumonitis patients. nPBV has the potential to detect radiation pneumonitis earlier than other methods described in previous reports.
Core Session 2 - Scientific Presentation 3 -

Therapy Response Assessment in Neoadjuvant Chemotherapy for Esophageal Cancer: Comparison of PET Response Criteria in Solid Tumors (PERCIST) and Response Evaluation Criteria in Solid Tumors (RECIST)

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Purpose:
To evaluate therapeutic response to neoadjuvant chemotherapy for esophageal cancer, compared PERCIST to RECIST.

Materials and Method:
Thirty-one patients with esophageal cancers were scanned by 18F-FDG PET/CT before and after chemotherapy. Tumor size was measured in the longest diameter on CT. Standardized uptake value on PET was normalized to lean body mass (SUL). SULpeak was measured using maximal 1.2-cm diameter volume region of interest placed on the hottest lesion of tumor on PET. Objective therapeutic responses were decided according to the RECIST [complete response (CR), partial response (PR), stable disease (SD), and progressive disease (PD)] and the PERCIST [complete metabolic response (CMR), partial metabolic response (PMR), stable metabolic disease (SMD), and progressive metabolic disease (PMD)].

Results:
The reduction ratio of tumor sizes was 30.3 ± 23.9%: tumor size before chemotherapy, 7.0 ± 2.4cm; after, 5.1 ± 2.9cm. Objective therapeutic response on the RECIST was as follows: CR(n=0), PR(n=13), SD(n=18), and PD(n=0). The reduction ratio of SULpeak was 53.9 ± 24.3%: SULpeak before chemotherapy, 8.9 ± 2.9; after, 4.1 ± 2.6. Objective therapeutic response on the PERCIST was as follows: CMR(n=8), PMR(n=17), SMD(n=6), and PMD(n=0). The results between the RECIST and PERCIST criteria were significantly different in all 31 patients (Mann-Whitney’s U-test, p=0.0002). Twenty-one of 31 patients were finally judged as clinical complete responders because of no recurrence in the progress after an operation. In these 21 patients, the results of PERCIST [CMR(n=6), PMR(n=11), SMD(n=4), and PMD(n=0)] were more accurate than those of RECIST [CR(n=0), PR(n=9), SD(n=12), and PD(n=0)] (p=0.002).

Conclusion:
PERCIST criteria were more suitable for the evaluation of chemotherapy response to esophageal cancer than RECIST criteria.
The progression of COPD involves changes in vasculature and parenchyma. Changes in parenchyma tissue are well characterized by HRCT and MR images of apparent diffusion of hyperpolarized gases are sensitive to early alterations in distal lung structures. However, not much is known about the changes in regional perfusion (Qr) in COPD, particularly in early disease stages. Studies using HRCT and PET suggest that the pattern of distribution of Qr may be altered early in the disease. Here I will discuss those studies and present the PET imaging methodology used to compare patients with mild-to-moderate COPD against healthy controls. The $^{18}$F-NH$_3$-labeled saline injection method was used to generate images of Qr and ventilation (Vr), while tissue density was estimated from the PET transmission scans. The mean-normalized variance was used to characterize the heterogeneity of Qr, and to quantify the contributions to heterogeneity caused by regions ranging from less than 12 mm to more than 108 mm. Compared with healthy subjects, COPD subjects had larger heterogeneity in Qr, and in Qr normalized by tissue density, with higher contribution from large length scales, and lower contribution from small length scales. Healthy subjects showed higher Vr in dependent regions, whereas no gradients in Vr were shown by COPD. CONCLUSION: Qr heterogeneity was greater in COPD than in healthy subjects and this was caused by changes in relatively large lung regions not explained by changes in tissue density or Vr. The redistribution pattern of Qr in COPD may be an early biomarker of the disease.
We have compared the clinical characteristics of COPD patients visually classified according to the dominancy of emphysema or the presence of bronchial wall thickening (BWT), and also examined combined pulmonary fibrosis and emphysema (CPFE). Little-emphysema phenotype showed a higher prevalence of patients who had never smoked, maintained body mass index, milder decreases in DLco and SpO₂ during exercise as compared with the emphysema-dominant phenotype. COPD with BWT showed a higher prevalence of patients complaining of a large amount of sputum, productive cough, and wheezing, and higher rate of exacerbation or hospitalization, and higher rate of patients having sputum eosinophilia who showed greater reversibility of airflow limitation responsive to the treatment with inhaled corticosteroids as compared with the COPD without BWT. The prevalence of CPFE was low at 0.96% in health screening study. We have compared clinical characteristics with CPFE and emphysema-dominant COPD without fibrosis. Forty-six of the 47 CPFE were male. Paraseptal emphysema was particularly common in the CPFE. Honeycombing, ground-glass opacities and reticular opacities were present in 75.6% and 84.4% of CPFE, respectively. Twenty-two patients (46.8%) were complicated with lung cancer. Pulmonary function tests showed that the CPFE had milder airflow limitation and lower diffusing capacity. Desaturation during exercise was more severe than those in emphysema-dominant COPD without fibrosis. It has been demonstrated the patients with CPFE having pulmonary hypertension is poor prognosis. These findings suggest that the morphological phenotypes of COPD show several clinical characteristics. It should be noteworthy that the prevalence of lung cancer might be high in CPFE phenotype.

According to the American Thoracic Society/ European Respiratory Society consensus and the Global Initiative for Chronic Obstructive Lung Disease guidelines, chronic obstructive pulmonary disease (COPD) is described as a slowly progressive disease characterized by airflow limitation, cough, sputum production, and, at later stages, dyspnea. Currently, computed tomography (CT) is most widely used for radiological disease severity assessment in COPD in clinical and academic practice. Although CT provides morphological information with high spatial resolution, increased iodinated radiation exposure and no direct assessment of pulmonary function are major its’ drawback in routine clinical practice. To overcome these drawbacks, several investigators have been suggested the potential advantages of magnetic resonance imaging (MRI) from the results of basic and clinical researches in 1990’s.

In the last decade, hyperpolarized noble gas MRI, oxygen-enhanced MRI (O₂-enhanced MRI) and contrast-enhanced or non-contrast-enhanced perfusion MRI (CE or non-CE perfusion MRI) have also been suggested as useful for assessment of regional morphological and functional changes in COPD patients. Although hyperpolarized noble gas MRI is warranted several special equipments such as laser polarizer, specialized transmitter and receiver coils etc, CE and non-CE perfusion MRIs and O₂-enhanced MRI are able to perform by using conventional proton MR scanners.

In this lecture, I will talk about 1) the basics of CE and non-CE perfusion MRIs and O₂-enhanced MRI; 2) summaries of clinical studies by using these MR techniques in COPD subjects; and 3) advantages and disadvantages of these MR technique for clinical setting. We hope this lecture will encourage to clinical setting of proton MR techniques for COPD in clinical and academic practices.
Chronic obstructive pulmonary diseases are a group of lung diseases that are intimately linked to smoking and other types of exposures. The prevalence is increasing, and as a result there is a need to increase the understanding of the disease process, its natural progression and the way in which we can quantify it appropriately. Regional information is heavily reliant on imaging methods, including CT and MRI. This information is hopefully going to be transformed into novel treatment modalities, such as improved delivery methods, interventional methods to decrease hyperinflation and methods which will result in anti-inflammatory effects - all these treatments are aimed at preservation of lung function.

Hyperpolarized noble gas MRI has been shown to hold great promise to probe both the extent of functional impairment in COPD, its underlying physiological and morphological properties and the possible quantification of these measures. Thus far, mainly due to restrictions in availability of methodology and also due to a lack of a reliable 3-Helium gas supply, the work has not propagated into mainstream clinical practice. However, with the resurgence of 129-Xenon gas imaging and the improvements in polarizer equipment, there is a new opportunity to apply the technology into a wider community. Whether the technology will become mainstream practice remains to be seen (MDCT is also increasingly being developed and has wider availability), but it is quite likely that hyperpolarized noble gas MRI will result in improved understanding of the disease and may be helpful in the development of new treatment opportunities.
Structure and function are two sides of the same pulmonary coin of physiology and pathophysiology. Structure exists to service the lung’s function, and the lung’s function is dictated by its structure. Hyperpolarized noble gases have the unique capability of probing both, in a non-invasive manner, with imaging to obtain regional information, and without ionizing radiation.

Specific to structure, the use of 3He or 129Xe, through the apparent diffusion coefficient (ADC), probes lung structure at various length scales, depending on the times between dephasing and rephasing MR pulses. This provides a direct measure of the mean squared diffusive displacement of gas molecules. Restrictions due to parenchymal septal tissue, specifically alveolar and ductal architecture, lead to systematic changes in ADC, from which length scales can be assessed. Loss of septal surface for gas exchange, a hallmark of emphysema in COPD can be measured through this approach.

Specific to function, the solubility of 129Xe, together with its NMR chemical shift in tissue, makes 129Xe an ideal surrogate for oxygen transport. Saturation pulses kill the tissue polarization, and recovery measured. Uptake dynamics can be directly related to structural parameters with immediate functional consequences: (1) alveolar surface area loss, (2) septal thickening and remodelling, and (3) capillary transit time and links to vascular co-morbidities; all three biomarkers of COPD.

This, then, is the frontier - hyperpolarized noble gases have the potential for broadening our understanding of basic physiology and its derangements in COPD, and represent novel tools for diagnosis, patient management, and follow-up care.

Hyperfused helium-3 (3He) magnetic resonance imaging (MRI) apparent diffusion coefficients (ADC) enable direct quantification of lung microstructure based on the 3He signal in gas ventilated regions. In serial or acute interventional studies there exists the potential for functional lung changes that might allow for different microstructural environments to be probed, leading to altered ADC values. Here we evaluated ten GOLD stage II - IV COPD subjects before and after administration of a bronchodilator (salbutamol) and interrogated the ADC signal in regions of interest that were newly ventilated. 3He ADC was calculated from MRI diffusion-weighted imaging, and ADC maps were generated for the five center slices, using in-house software, for calculation of ADC in the whole lung (WL), the regions of pre/post overlap (RO), and the regions of pre/post non-overlap (RNO) using two-dimensional landmark-based registration. The subsets of pre/post overlap were identified and mean ADC were calculated in the RO and then used as a mask on pre/post images to calculate mean ADC in RNO. 3He ADC was not significantly different pre - and post-salbutamol for the WL (pre=0.51 cm²/s, post=0.51 cm²/s, p=.35) or the RO (pre=0.51 cm²/s, post=0.51 cm²/s, p=.35).
Rationale: Understanding of the relative contributions of airway disease and emphysema to airflow limitation in chronic obstructive lung disease (COPD) has been a subject of interest for many years.

Objectives: To examine the relationships between airway dimensions and emphysema severity in the upper and lower parts of the right lung separately using 3D computed tomography, and then to explore the relationships of both pathologic processes to airflow limitation indices on a regional basis.

Methods: Seventy male patients with clinically stable COPD and 10 male smokers at risk served as subjects who underwent pulmonary function tests and computed tomography at full inspiration. Wall area % (WA%) was measured at the 3rd (segmental bronchus) to 6th generation of the two bronchi as a marker of airway disease, and low attenuation volume (LAV%) was measured as a marker of emphysema severity.

Measurements and main results: The relationship between airway dimensions and LAV% was not significant in the upper lung and was weakly significant in the lower lung. The significant correlations observed between WA% and FEV1 (% predicted) were better at the more distal generations of both airways. LAV% correlated significantly with FEV1 (% predicted) only for the lower lung (r=0.621, p<0.001) but not for the upper lung (r=-0.178, p=0.127).

Conclusions: Relationship of airway disease and emphysema may be different in the upper lung and the lower lung and thus relative contribution of both pathologic processes to airflow limitation may be different according to region of the lung in male patients with smoking-related COPD.
Core Session 4 - Invited Speaker 1 -
Update for Diagnosis of IIP

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Idiopathic interstitial pneumonias (IIPs) are a heterogeneous group of diseases characterized histologically by inflammation and fibrosis of the parenchymal interstitial tissue. Their causes are unknown and some degree of airspace disease commonly accompanies the interstitial abnormality in all these conditions. IIPs are a heterogeneous group of diseases characterized histologically by inflammation and fibrosis of the parenchymal interstitial tissue. IIPs are classified acute interstitial pneumonia (AIP), cryptogenic organizing pneumonia (COP), nonspecific interstitial pneumonia (NSIP), usual interstitial pneumonia (UIP), desquamative interstitial pneumonia (DIP), respiratory bronchiolitis-associated interstitial pneumonia (RB-ILD) and lymphoid interstitial pneumonia (LIP). Each IIP has characteristic CT findings including extensive areas with ground-glass attenuation, patchy areas of airspace consolidation and architectural distortion in AIP, bilateral peripheral areas of airspace consolidation in COP, areas with ground-glass attenuation with lower and peribronchiolar predominance in NSIP, intralobular reticular opacities and honeycombing distributing mainly in the subpleural regions and lung bases in UIP etc. However, there are several overlaps of their characteristic CT findings. Characteristic CT findings of LIP diagnosed by recent criteria have been unclear. Some cases show lower lobe predominance and areas with ground-glass attenuation. In this lecture, imaging finding of each IIP will be presented paying special attention to its characteristic CT findings and pathologic backgrounds. Moreover, the recent diagnostic criteria of IPF/UIP by ATS-ERS-JRS will be mentioned paying special attention to the drawbacks.

Core Session 4 - Invited Speaker 2 -
Update for Treatment of Idiopathic Interstitial Pneumonias (IIPs)

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The ATS/ERS proposal approaches the classification of IIPs by identifying the pathologic patterns seen on lung biopsy in addition to clinico-radiologic findings, thereby dividing into 7 categories. According to corticosteroid (CS) resposivity, the IIPs are broadly divided into two groups: cases such as NSIP, COP and DIP that respond to CS, and cases with IPF and AIP that poorly respond to CS. Among IIPs, IPF is the most common type of IIPs, and the underlying concept that IPF is primarily an epithelial-fibroblastic disease explains the failure of anti-inflammatory therapy, and consequently, introduction of antifibrotic agents has been expected to improve survival rates of IPF. In this session new treatments for IPF are summarized, including interferon-γ, pirfenidone, antioxidants, anti-TNF-α antibody, endothelin receptor antagonist and BIBF 1120 (triple kinase inhibitor).

Abbreviations:
NSIP: nonspecific interstitial pneumonia
COP: cryptogenic organizing pneumonia
DIP: desquamative interstitial pneumonia
IPF: idiopathic pulmonary fibrosis
AIP: acute interstitial pneumonia
Smoking-related Interstitial Pneumonia Associated with Emphysema

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Cigarette smoking affects many diseases such as chronic obstructive pulmonary disease, lung cancer and cardiovascular diseases. Cigarette smoking may also prime the lung toward a fibrotic response and participates in the etiology of a spectrum of fibrosing lung diseases, including idiopathic interstitial pneumonias (IIPs). The purpose of the presentation is to clarify clinical and radiographic features of chronic fibrosing IIPs, especially in view of the influence of smoking history. One hundred and eight consecutive patients with chronic IIPs were studied. They were divided into three groups according to smoking history and presence of centrilobular emphysema (CLE) on HRCT. The three groups were: fibrosing IIP with CLE and smoking history (IPE, n=54), fibrosing IIP without CLE but with smoking history (IPS, n=33) and fibrosing IIP without smoking (IPNS, n=21). Forty two out of 54 patients with IPE showed thick-walled large cysts (TWLC) more than 2cm in diameters. Ten out of 33 showed TWLC in IPS group and none demonstrated TWLC in the IPNS group. The mean %VC was 90% in IPE, 89% in the IPS, 59% in the IPNS, and was decreased in IPNS. The mean FEV1% was 77%, 83%, 88%, and was decreased in the IPE. The mean %DLCO was 56%, 73%, 67%, respectively, and was decreased in the IPE. TWLC is characteristic of chronic fibrosing IIPs with smoking history. UIP pattern may be highly associated with the progression as well as the development of TWLC in subjects with smoking history.

Hyperpolarized Noble Gas MRI for ILD

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Hyperpolarized ^129Xe NMR was utilized to measure pulmonary alveolar surface area per unit gas volume $S_a/V_{gas}$, alveolar septal thickness $h$, and capillary transit time. The global $S_a/V_{gas}$, $h$, and capillary transit time were measured in normal subjects and subjects with mild interstitial lung disease. These are the first noninvasive, non-radiation based, quantitative measurements of alveolar septal thickness and capillary transit time in patients with pulmonary disease. This new approach may open a new window to monitor disease progression and treatment response in patients with interstitial lung disease.
**Core Session 4 - Scientific Presentation 1 -**

**Prognostic Significance of High-Resolution Computed Tomography in Nonspecific Interstitial Pneumonia**

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**Purpose:** To assess the prognostic implications of high-resolution computed tomography (HRCT) findings for patients with biopsy-proven nonspecific interstitial pneumonia (NSIP).

**Materials and Methods:** 59 patients with biopsy-proven NSIP (25 idiopathic NSIP, 34 collagen-vascular disease-associated NSIP) were included in this retrospective study. Data and HRCT images were obtained before surgical lung biopsy and treatment. Two chest radiologists independently evaluated the extent, presence and distribution of various HRCT findings of NSIP patients. Cox proportional hazards regression analysis was used to assess the relationship between HRCT findings and survival outcome.

**Results:** The median observation period was 1477 days, ranging from 289 to 4526. Among all patients, the 5-year survival rate was 83% and the 10-year survival rate was 66%. Univariate analysis revealed that the extent of GGO without bronchi-bronchiolectasis (hazard ratio = 0.78, P < 0.01) and total consolidation (hazard ratio = 0.83, P = 0.04) were associated with favorable outcome whereas the extent of intralobular reticular opacity (hazard ratio = 1.23, P < 0.01) was associated with worse prognosis. Multivariate analysis revealed that the extent of total consolidation was an independent prognostic factor of favorable outcome (hazard ratio = 0.59, P = 0.03).

**Conclusion:** In patients with biopsy-proven NSIP, detailed analysis of HRCT is useful for predicting outcome.

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**Core Session 4 - Scientific Presentation 2 -**

**Automatic Classification of Regional Disease Pattern of Diffuse Lung Disease at HRCT: Cross-Vendor Study**

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To investigate the effect of HRCT images from different vendors on classification accuracy for regional disease pattern analysis of diffuse lung disease (DLD).

Using HRCT images from vendor A and B, two sets of 600 rectangular regions of interest (ROIs) with 20x20 pixels, comprising of each 100 ROIs representing six regional disease patterns (normal, GGO, reticular opacity, honeycombing, emphysema, and consolidation) were marked by two experienced radiologists with consensus at HRCT images of various DLDs. Each ROI image, then, was represented by 28 features (histogram, gradient, run-length, GLCM, LAA cluster, and top-hat transform). For automatic classification, support vector machine was employed to generate a classification model. Firstly, classification accuracies using each vendor’s data were estimated by a 10-folding cross validation with 20 repetitions (internal-vendor study). Secondly, A and B data were used for training and testing, respectively, and vice versa (cross-vendor study). Finally, all ROI data were integrated, trained and tested by 2-folding cross validation with 20 repetitions (integrated data study).

On internal-vendor study, the accuracies of vendor A and B were 89.4 ± 0.98% and 89.8 ± 0.79%,
respectively. The classification accuracies of cross-vendor study were 45.8 ± 0.85% when training vendor A data and testing vendor B data, and 42.7 ± 0.81% vice versa (t-test, p=0.000). On integrated data study, accuracy was 88.3 ± 1.48%. Selected features were different among the studies.

It is recommended to train automatic classifiers for each vendor separately or to build an automatic classifier with integrated data set if needed.

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**FDG Positron Emission Tomography Imaging of Drug-induced Pneumonitis**

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Fluorine-18-labeled fluoro-2-deoxy-D-glucose positron emission tomography (FDG-PET) is increasingly used for diagnosing lung cancer, but FDG accumulation is not specific to malignancy. There are numerous causes of FDG uptake in benign processes, such as inflammatory lung disease. However, ⁶⁷Ga scintigraphy has been widely used for the evaluation of interstitial pneumonitis. Several studies have reported the findings of FDG-PET in benign lung disease with diffuse pulmonary injury, but the characteristics and effectiveness of FDG-PET imaging for interstitial pneumonitis have not been substantiated. We report two cases of drug-induced pneumonitis in two patients treated for breast cancer who were diagnosed by FDG-PET examination.

Both of these cases showed diffuse interstitial infiltration in the bilateral lungs on CT, but the degree of FDG accumulation was different. It is probable that the degree of FDG accumulation reflected the activity of the drug-induced pneumonitis. The present cases show very interesting FDG-PET imaging findings of diffuse lung disease.
**Core Session 5 - Invited Speaker 1 -**

**Pathophysiology of Asthma**

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Bronchial asthma is characterized by chronic inflammation, eosinophilic infiltration, reversible airway narrowing, airway hyperresponsiveness (AHR) to non-specific stimuli, hyperplasia/metaplasia of goblet cells, hypertrophy/hyperplasia of airway smooth muscle cells and subepithelial fibrosis. It is now considered that airway inflammation and structural changes of the airways are important features. T helper 2 (Th2) cells are the predominant lymphocyte population that infiltrates the airways of asthmatics, and Th2 cytokines, such as IL-4, IL-5, and IL-13, play essential roles in airway eosinophilia, AHR, and serum IgE in animal models. These cytokines bind to receptors at the cell surface to activate complex signal transduction pathways, including JAK-STAT and Ras-ERK signaling pathways. This presentation will cover the function of cytokine signaling pathway in pathophysiology of asthma and discuss new therapeutic targets. The suppressor of cytokine signaling (SOCS) family proteins has been shown to regulate JAK-STAT, and the Sprouty-related EVH1-domain-containing protein (SPRED) family proteins regulate the Ras-ERK pathway. In the inflammatory/immune cells, SOCSs contribute to helper T cell differentiation during immune responses, and SPRED-1 negatively controls the eosinophil numbers and functions by modulating IL-5 signaling. In the airway structural cells, SOCS1 induction by IL-13 is critical to negatively control asthma. Airway cells from asthmatics had impaired upregulation of SOCS1 after IL-13 stimulation. Therefore, modulators of Th2 responses including the negative regulators of cytokine signals may improve our understanding of these processes to asthma and hold therapeutic potential in asthma.

**Core Session 5 - Invited Speaker 2 -**

**Evaluation of Airway Obstruction and Altered Pulmonary Permeability with Lung Scintigraphy in Young Patients**

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The $^{133}$Xe ventilation scan shows three distinct phases. The initial phase is the wash-in or ventilation phase. The second phase is the equilibrium phase representing the gas volume. Finally, the wash-out phase, when the $^{133}$Xe has been disconnected, gives information about the possibility of any gas trapping, which may occur in obstructive airways disease. In addition to imaging with $^{99m}$Tc-DTPA aerosol of the disappearance rate of tracer from regions of the lung can provide further information about lung permeability and may degrade the ventilation images very rapidly.

In contrast to the adult form of rheumatoid disease, follicular bronchiolitis appears to be the most common manifestation of pulmonary involvement in the juvenile form. Bronchiolitis obliterans appears in patients with chronic graft-versus-host disease. The $^{133}$Xe ventilation scan and the $^{99m}$Tc-DTPA aerosol inhalation scan can be performed in young patients with bronchiolitis obliterans safety and evaluated in quantitative analysis. These lung scans are considered useful as a method for diagnosing in early stage and a follow-up to bronchiolitis and interstitial pneumonitis, provided that pulmonary function tests and CT show normal patterns and images.

Metered dose inhalation therapy of beta agonist or steroid is gaining popularity in the treatment of bronchial asthma, and the spacer is applied to distribute the drug to the lung stably and diffusely. The inhalation scan is useful to assess the effect of spacer on deposition rate and distribution of the medicine in the lung, besides response to the therapy.
Asthma is diagnosed in an estimated 300 million people worldwide and it claims the lives of 250,000 people annually. Clinical diagnosis is typically achieved using methods that report global lung function such as spirometry measurements of the forced expiratory volume in 1 s (FEV1), and low attenuation area without destruction of interlobular septa is correlated with RV/TLC. Central airway diseases are almost even, however peripheral airway diseases are uneven and ventilation heterogeneity is apparent during asthma exacerbation. Previous reports have provided evidence that area of peripheral airway diseases move in each exacerbation using radioisotope or CT scans. In general, heterogeneity of peripheral airway disease is measured by closing volume, slope of phase III of single N2 washout test. Pathological findings of open lung biopsy specimens during asthma exacerbation demonstrate that 1) peripheral airways are filled with sputum, 2) peripheral airways are dilated by accumulated sputum, and 3) the degrees of these diseases are uneven in the lung. Pathological-radiological correlation shows that ground glass opacities are peripheral airway disease under the diameter of 500 micrometer, centrilobular nodules are dilated peripheral airways filling with sputum. And all findings disappear after asthma treatment. However, severe persistent asthma remains centrilobular nodules for a long time, which may contribute to persistent severe obstructive change. CT scans of patients with aspirin induced asthma during asthma exacerbation reveal that thickening of interlobular septa and pleural effusion in addition to ordinary asthma exacerbation, suggesting edema of pulmonary parenchyma. In this session, I focus on heterogeneity of peripheral airway disease, using pulmonary function test, CT finding and pathological features.
Medication plans should be selected on the basis of the severity of asthma, availability of antiasthma medications, conditions of health care system, and individual patient circumstances to achieve best control. Therapeutic strategy for asthma, the basically standard medication is inhaled corticosteroids (ICS). ICS reduces the risk of hospitalization or mortality in elderly asthmatic patients as same as in younger adult asthmatic patients. The addition of a long-acting beta-agonist (LABA) including combination LABA and ICS or the addition of a leukotriene modifier improves symptoms in patients with inadequately controlled asthma by ICS. Recently, It is reported that the addition a long-acting anticholinergic agent, tiotropium improved symptoms and lung function in these patients. But a careful treatment is required in some asthmatic patients who cannot inhale drugs or cannot adhere to their treatment regimens well.

Smoking, obesity, elderly asthma and comorbidity with chronic sinusitis are well known risk factors of uncontrolled asthma by ICS. Especially smoking is a risk factor for accelerated lung function deterioration, impaired corticosteroid response, and increased mortality, so smoking cessation is strongly recommended. Diet modification and weight loss intervention are also recommended for obesity. For severe asthma with fixed airway obstruction, an anti-IgE therapy is recommended in the patients with persistent eosinophilic airway inflammation, and macrolids antibiotics and PDE4 inhibitor may improve non eosinophilic inflammation, particularly neutrophilic inflammation.
Introduction: In asthma ventilation is heterogeneous and airway closure is increased, both being associated with greater asthma severity and airway hyperresponsiveness. Heterogeneous ventilation has been demonstrated by various imaging techniques but the changes in spatial distribution during bronchoconstriction, and the predictors of that change are unknown.

Hypothesis: that the spatial heterogeneity of ventilation in Technegas SPECT correlates with global measures of ventilation distribution and that it increases with methacholine challenge.

Methods: 14 Asthmatic subjects underwent Technegas SPECT/CT before and after bronchial challenge. 1L breaths of Technegas were inhaled from FRC, in upright posture before SPECT and CT acquisition in supine posture. Technegas distribution was measured by plotting the ventilated volume determined from 3 different thresholds vs those thresholds, in log-log space (TechnegasDIST). Ventilation distribution was also measured by multiple breath nitrogen washout (MBNW) from which Sacin and Scond were calculated, as indices of heterogeneity in diffusion dependent airways and in small, conducting airways, respectively.

Results: FEV1 was 87.5 $\pm$ 15.8 % predicted and 7 subjects had AHR. Methacholine challenge decreased FEV1 by 23 $\pm$ 5% of baseline. TechnegasDIST increased from -0.35 $\pm$ 0.03 to -0.47 $\pm$ 0.15 (p=0.008). TechnegasDIST at baseline was unrelated to spirometry or MBNW indices. However, the post-methacholine TechnegasDIST and the change from baseline, correlated with baseline Sacin (r=-0.62, p=0.17 and r=-0.64, p=0.013, respectively).

Conclusions: The spatial distribution of ventilation measured by Technegas SPECT increases with methacholine challenge which in turn, is predicted by baseline heterogeneity in diffusion dependent airways. The anatomical basis of ventilation distribution requires further study.

Core Session 5 - Scientific Presentation 1 -

The Effect of Methacholine Challenge on Ventilation Heterogeneity in Asthma Measured by SPECT

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Core Session 5 - Scientific Presentation 2 -

Surface Irregularity of Airway Walls as Assessed by MDCT in Asthmatic Patients

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[Backgrounds] In asthmatic patients, airway wall remodeling is observed both at large and small airways. Injury to the epithelium is a common finding in asthma, and loss of alveolar attachments at small airways was reported in severe fatal asthma. But few reports have described structural changes in outer layer of large airways.

[Objectives] To analyze inner and outer surface irregularity of central airways using multi-detector row CT (MDCT) and examine its correlations with clinical variables.

[Methods] CT scans were performed in treatment-naive asthmatic patients. Then, inner radius and wall thickness of all segmental bronchi were calculated by using three-dimensional reconstruction, and coefficient of variation (CV) of distance from center line to inner and outer walls at each bronchus were calculated. Finally, correlations between clinical variables and averaged CV of all right segmental bronchi were calculated.

[Results] Averaged CV of both inner and outer wall showed negative correlation with FEV1, FEF25-75 of spirometry and airway hyperresponsiveness estimated by Dmin of acetylcholine using Astograph, and positive correlation with R5 R20 R5-R20 and AX of impulse oscillometry. They showed no significant correlation with exhaled NO or sputum eosinophils.

[Conclusions] Surface irregularity of airway walls may have some relations with disease severity of asthma, although its pathological mechanism is not unclear.
Objectives: To evaluate the correlations of tracheal volume and collapsibility on inspiratory and end-expiratory computed tomography (CT) with lung volume and with lung function in smokers.

Materials and Methods: An institutional review board approved this study at each institution. 85 smokers (age range 45-87 years), enrolled in the Lung Tissue Research Consortium, underwent pulmonary function tests and chest CT at full inspiration and end-expiration. On both inspiratory and end-expiratory scans, intrathoracic tracheal volume (from the apex to the carina) and lung volume were measured by open-source software. Collapsibility of the trachea and the lung was expressed as expiratory/inspiratory (E/I) ratios of these volumes. Correlations of the tracheal measurements with the lung measurements and with lung function were evaluated by the linear regression analysis.

Results: Tracheal volume correlated with lung volume on both inspiratory (r=0.661, p<0.0001) and end-expiratory (r=0.749, p<0.0001) scans. The E/I ratio of tracheal volume also correlated with the E/I ratio of lung volume (r=0.711, p<0.0001). The E/I ratio of tracheal volume negatively correlated with the ratio of forced expiratory volume in the first second to forced vital capacity (FEV1/FVC) (r=−0.436, p<0.0001). A weak, positive correlation was observed between the E/I ratio of tracheal volume and the ratio of residual volume to total lung capacity (RV/TLC) (r=0.253, p=0.02).

Conclusions: Tracheal volume and collapsibility, measured by inspiratory and end-expiratory CT scans, is related to lung volume and collapsibility. The highly collapsed trachea on end-expiratory CT does not indicate more severe airflow limitation or air-trapping in smokers.
From time immemorial, lung physiology has been explained in concepts assuming steady state conditions. Within the last decade clear evidence has been found, that lung ventilation and perfusion are highly dynamic processes, with regional inhomogeneities in their kinetics. This holds specifically true in the injured lung.

Cyclic recruitment of atelectasis (“atelectotrauma”), and regional lung overdistention due to high airway pressures (“barotrauma”) or high tidal volumes (“volutrauma”) have been shown to lead to an inflammatory response of lung tissue (“biotrauma”). These iatrogenic mechanisms during mechanical ventilation are summarized in the definition of “ventilator associated lung injury”.

Thus, novel therapeutic strategies require dynamic and regional information of lung ventilation and perfusion. Dynamic computed tomography and magnetic resonance imaging with inhalative contrast agents have brought new insights and became an emerging research field. However, up to this point, these imaging modalities are not routinely applied in the treatment of patients with acute lung injury. Non-invasive and bedside imaging technologies like electrical impedance tomography or vibration response imaging could serve as a bridge to bring the imaging information from the bench to the bedside.

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MR is a suitable tool for observing respiratory motion noninvasively. From 1980s dynamic images applies to assess the chest wall motion. Recent advances in MR allow to visualize internal motion of the lung. Voorhees A, et al applied 3D tagging method and showed the lung internal motion in tidal breathing [1]. Plathow C et al. measured the motion of the lung tumor in 3D-dynamic MRI [2]. We developed a system to trace the pulmonary vessels on 2D dynamic MRI [3]. In this presentation I will introduce recent results using this system. Shibata H et al. showed the motion of the pulmonary vessels was restricted in the patients with chronic obstructive lung disease (COPD), and movement of the vessels on MR significantly correlated with airflow limitation evaluated by spirometry [4]. Shinohara T et al. evaluated the COPD patients using 2D dynamic MRI before and after rehabilitation [5]. After rehabilitation, the movement pattern of the pulmonary vessels changed to similar pattern observed in normal volunteers. These results suggest that motion of the pulmonary vessels reflects the expansion and deflation of the lung parenchyma. This method will be a tool to investigate the respiratory mechanics.

3. Gotoh T, et al. 18th IFAC world Congress.
4. Shibata H, et al. 5th IWPFI
5. Shinohara T, et al. 5th IWPFI
The development of 320-row area detector CT (ADCT) has made it possible to perform dynamic volume scanning. A range of 160 mm can be scanned in a single rotation in a short time of 0.35s. ADCT provides superior performance in terms of both temporal resolution and spatial resolution. The acquired data can be obtained in the same time phase regardless of the position in the axial direction. Using dynamic volume scanning, temporal uniformity and four-dimensional data can be acquired.

In the chest region, dynamic volume scanning is a simple method to evaluate the various respiratory mechanics. For example, diaphragm motion, chest wall motion, intrathoracic tumor mobility, deformity of trachea, bronchus and lung tissue during the respiration cycle.

Controlling the exposure dose is one of the main issues that must be addressed. Thus, mainly 3 types of scanning protocols (intermittent scan, respiration-gated scan, and continuous scan) with low-dose exposure have been used. By adjusting the number of scan, tube voltage, tube current, and noise-reduction reconstruction method, we have to try to obtain the four-dimensional data without compromising image quality.
Pandemic influenza A(H1N1) virus emerged worldwide in 2009. WHO provided guidance on public health measures, including entry screening at airports to reduce the risk of transmission. A number of countries have already applied thermography at international airports in order to detect infected passengers. However, the effectiveness of the conventional screening system using thermography is limited in detecting asymptomatic or sub-clinical infections. In order to conduct more accurate and fast screening, we have developed a non-contact screening system that can perform human medical inspections within ten seconds. The system monitors not only temperature but also heart and respiratory rates, which leads to an increase in accuracy in detecting infected individuals. The system is operated by a screening program via a linear discriminant analysis using non-contact derived variables, i.e., palmar pulse derived from a laser Doppler blood-flow meter, respiration rate determined by a 10-GHz microwave radar, and average facial temperature measured by thermography. In our case control study, the discriminant function of this system is found to be effective in distinguishing influenza patients regardless of their temperatures. This can be attributed to the fact that even in the cases of influenza patients with normal temperatures, their heart and respiratory rates are relatively high compared with normal subjects. The system enables PPV rate of 93%, which is much higher than the conventional method using only thermography. It can be improved for future applications to reduce the risk of transmission of infectious diseases.

Core Session 6 - Scientific Presentation 1 -

Temporal Stability of the 4D-CT Pulmonary Ventilation Imaging Method

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Purpose: A novel pulmonary ventilation imaging method based on four-dimensional (4D) computed tomography (CT) has advantages over existing methods and could be used for functional avoidance in lung cancer radiotherapy as ‘free’ information. The purpose of the present study was to quantify the temporal stability of the 4D-CT ventilation imaging method from repeat scans.

Methods: Two 4D-CT ventilation images from repeat scans (interval range, 9-13 days) in the absence of therapeutic intervention were created for three radiotherapy patients with lung cancer. Both the Hounsfield-unit (HU) change-based metric (V_HU) and Jacobian-based metric (V_Jac) were used to create the ventilation images from the displacement vector fields. The second 4D-CT ventilation image was deformed into the domain of the first ventilation image for the analysis. The voxel-based Spearman’s rank correlation coefficients between two ventilation images were quantified.

Results: From visual inspection, repeat 4D-CT ventilation images showed regions of agreement and disagreement for both V_HU and V_Jac. The correlation coefficients ranged from 0.30 to 0.47 for V_HU, which were consistently lower than those for V_Jac ranging from 0.38 to 0.66. It was found that the large differences between two ventilation images mainly manifested around the interfaces between two adjacent 4D-CT data segments, and also around the 4D-CT artifacts (especially diaphragm).

Conclusions: The temporal stability of 4D-CT pulmonary ventilation imaging was weak to moderate and varied between patients and ventilation metrics. Future work will investigate the spatial characteristics of the stability of 4D-CT ventilation images for more patients.

Core Session 6 - Scientific Presentation 2 -

Analysis of Respiratory Movement Using 4-Dimensional Chest CT image

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Recent improvements in radiotherapy technology have enabled the respiratory gating system. The system tracks patient’s respiratory cycle using camera and reflective marker. In this study, we quantitatively analyze respiratory movement of pulmonary organs and nodules using 4-dimensional chest Computed Tomography images, aim for the therapeutic application.

Firstly, we apply automatic segmentation of pulmonary organs, lung, pulmonary blood vessel, trachea, bronchus, interlobar fissure, lung lobe, rib, and spine. Secondly, lung nodules are segmented. Thirdly, volumetric variation and locational motion of the organs and nodules with respiration are visualized and quantified.
Asymmetry of the Mechanical Property in the Human Respiratory System Causes the Asymmetry of the Lingual Center in the Brain: Diaphragmatic Motion Analysis with Dynamic MR Images

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The reason why the lingual center is located in the left hemisphere has been unknown. Here, we propose a hypothesis that asymmetric diaphragmatic motion during vocalization leads the motor lingual center to the left side. Recent comparative neurophysiologic research strongly suggests that animal vocalization is the origin of the language. Meaningful vocalization is performed not only by acoustic source organs such as vocal cords, but also by the respiratory system which generates and controls expiratory airflow. During vocalization, abdominal muscles work so as to control diaphragmatic motion and to control expiratory airflow rate. There is an extreme difference in organs located beneath the diaphragm between the right and the left sides. The solid liver is expected to convert horizontal force generated by the abdominal muscles into vertical force much more precisely than the air-containing stomach. Consequently, the information from the right pleura is expected to be used for control of vocalization. In order to validate the hypothesis, we acquired dynamic 2D MR images for a normal volunteer on the frontal section at the center of the thorax at supine posture, and measured diaphragmatic motions both at rest and during singing. The result indicated that there was significant difference between right and left diaphragmatic motions only during singing. The right diaphragmatic motions were completely coherent to vocalization but the left diaphragmatic motions were often paradoxical. The importance of abdominal breathing should be investigated more intensively in relation to the lingual function as well as the respiratory physiology.
Abstract

Poster Session
**Poster Session 1 - 1**

Follow-up Cases with Perfusion Scintigraphy in Patients with Pulmonary Thromboembolism

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We investigated the recovery course of pulmonary perfusion on Tc-99m-MAA pulmonary perfusion scintigraphy in nineteen patients with pulmonary thromboembolism. We classified 19 patients into 2 groups in whom the onset day was evident (Group A) and those in whom the onset day was not evident (Group B). Group A consisted of 10 patients in which 7 patients had underlying disease, such as malignant and autoimmune diseases, and thrombolytic and anticoagulant therapies were performed in 3 and all patients, respectively. Perfusion defects on scintigraphy were completely and partially recovered within 2 weeks of therapy in 3 and 5 patients, respectively. Since responses to therapy were good, re-examination for course observation should be performed within 12 weeks in cases meeting the group A conditions. Group B consisted of 9 patients in which 6 patients had underlying diseases, and thrombolytic and anticoagulant therapies were performed in 1 and all patients, respectively. Only 1 and 2 cases were completely and partially recovered within 2 weeks after the initiation of therapy, respectively, showing low response. This group included cases with delayed discovery and diagnosis due to the slow progression of symptoms. No improvement of pulmonary perfusion was noted in 3 cases of chronic pulmonary thromboembolism. One case completely recovered after 4 years, showing that long term follow-up by perfusion scintigraphy is necessary to investigate whether partial recovery case meeting the condition of 2 groups completely recover or enter chronic state.

**Poster Session 1 - 2**

Most Optimum Display Condition(Width,Center) of Dual Energy Perfusion CT for Diagnosing Pulmonary Embolism

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**Purpose:** Recently computed tomography pulmonary angiography (CTPA) becomes an alternative to conventional perfusion scintigraphy for diagnosing pulmonary embolism (PE). Using dual energy (DE) technique of dual source CT (DSCT) it becomes possible to visualize the information of pulmonary perfusion in addition to thrombus in pulmonary arteries. Although dual energy iodine value (DEIV) calculated on DECT is an absolute value, the perfusion images using color scale might be affected by window settings. In this study we investigated the optimum display settings of DECT using DSCT for diagnosing PE.

**Material and Method:** Thirty-one lesions diagnosed with pulmonary artery embolism by perfusion scintigraphy and 66 regions evaluated as normal in 13 patients were enrolled. Scanners were SOMATOM Definition (SIEMENS) using DE mode (80 and 140 kVp), pitch: 0.5 and rotation time: 0.33 seconds and E. CAM (SIEMENS) using 185 MBq of 99mTc-MAA.

**Result:** DEIVs (average ± SD) of lesions and normal areas were 12.3 ± 10.6 and 34.9 ± 6.9, respectively. The optimum display settings to separate the both groups would be expected as window center of 20-25 and window width of 15-20. The results of CT and SPECT were correlated well in all cases except for one location that showed normal on CT with a defect on scintigraphy.

**Conclusion:** The appropriate display settings to differentiate the areas with perfusion defect from those with normal blood flow were determined on DE mode of DSCT. DECT might replace perfusion scintigraphy.
**Poster Session 1-3**

**Pulmonary CT Perfusion Imaging in Assessment of Pulmonary Vascular Disease-First Experience Using 320-MDCT.**

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**Introduction:** There is increasing evidence that pulmonary embolism may lead to significant vascular bed alterations, including the development of pulmonary hypertension in 5-7% of patients. MDCT using 320-slice system can provide volumetric perfusion data in a single rotation at acceptably low-dose radiation to make this a possible clinically useful application.

**Materials and Methods:** We studied 5 subjects with suspected pulmonary hypertension (one with liver cirrhosis and four following large pulmonary emboli). Patients underwent expiratory breath-hold CT following 70 ml Iomeron 400 mg I/ml (Bracco) and 30 ml saline flush at 9 ml/s i.v.

**Scan parameters:** 100 kv, 100 mA, a dynamic field-of-view of 400L and a rotation time of 0.5s. Scans were obtained from 5 s to 20 s after start of injection at 2 s intervals with 160 mm z-axis coverage.

**Initial Results:** We demonstrated expected gravity dependent perfusion as expected in a normal distribution pattern in 2 subjects, minor peripheral defects in 1 subject and significant peripheral defects in a further 2 subjects.

**Conclusion:** Large z-axis, low-dose CT perfusion may be a useful method to study physiological effects of lung and pulmonary vascular diseases. This work is to be expanded in the near future.

**References:**

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**Poster Session 1-4**

**Reevaluation of the Tc-99m MAA Whole Body Imaging for Detecting Right-to-left Shunt**

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We retrospectively evaluated examinations for R/L shunt index measurement by Tc-99m MAA whole body imaging.

**Subjects & Methods:** 26 cases with suspicious of R/L shunt performed a whole body perfusion scan. 4 of 26 perfusion scans were performed with Xe ventilation scan (group-Xe), 8 with Tc-99m Technegas inhalation scan (group-TcG). Remaining 14 cases were performed perfusion scan only; whole body scan was performed directly after injection of Tc-99m MAA (group-Dir). The site of extrapulmonary accumulation and the shunt index were analyzed and compared among the 3 groups.

**Results:** 13 scans showed the brain accumulation. 11of 13 cases were diagnosed as having R/L shunt. All of 4 cases in group-Xe showed brain accumulation. Group-TcG showed higher rate of the shunt index than other groups. 7cases of group-Dir were diagnosed as having R/L shunt, and uptake for muscle or intestine were considered characteristic. Remaining 7cases of group-Dir diagnosed as having no shunt, but these cases showed extrapulmonary accumulations for thyroid, stomach or urinary bladder.

**Conclusion:** R/L shunt index measurement by Tc-99m MAA whole body imaging has many pitfalls. For precise evaluation of the shunt, whole body perfusion scan is recommended to perform separately from ventilation scan. The evaluation of extrapulmonary accumulations is important for the diagnosis of R/L shunt.
**Nonembolic Causes of Perfusion Defect on Contrast-enhanced Dual-Energy CT: Review of 85 Consecutive Cases**

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**PURPOSE**

To identify nonembolic causes of perfusion defects on contrast enhanced dual-energy pulmonary CT angiography (DECTPA) and to analyze the defect pattern for each cause.

**METHOD AND MATERIALS**

152 consecutive patients underwent DECTPA for clinical suspicion of embolism (PE). Consequently, 85 patients with 510 lobes were included in the study (M:F=42:43; mean age, 60.64 yr). DECTPA scanning (Somatom Definition, Siemens) was performed at 140 kV and 80 kV. Color-coded perfusion images were obtained with the workstation software (Syngo Dual Energy). The presence, and pattern of perfusion defects (PD) in each lobe were evaluated and their causes were assessed by two radiologists.

**RESULTS**

Out of 510 lobes, 37 lobes demonstrated artifacts accounted for more than 30% of the each lobe area. With the right middle lobes being the most common sites (27%), these lobes were waived from further evaluation. PDs were seen in 114 out of 510 lobes in total (22%). Nonembolic vascular causes of PDs were as follows; extrinsic occlusion/stenosis of PA at hilum by fibrosing inflammation, congenital PA hypoplasia, venous occlusion, and pulmonary angiosarcoma. PD areas were matched with vascular territory in most cases. Nonvascular causes include: mosaic lung attenuation, bronchitis, emphysema, bronchiectasis, cellular bronchiolitis, bronchial obstruction by mucus and bronchopneumonia. PDs was geographic and mismatched with anatomic extent in cases with bronchitis, bronchiectasis and bronchopneumonia, whereas PDs by cellular bronchiolitis, mosaic attenuation and emphysema were well matched with anatomic extent.

**CONCLUSION**

PDs without is not uncommon. Various nonembolic vascular and nonvascular diseases can develop PDs in different pattern.
Poster Session 2-1

Correlation between Airway Dimensions and Emphysema Scores in Patients with Chronic Obstructive Pulmonary Disease: Patient-based versus Lobe-based Analysis

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PURPOSE
The purpose of this study was to correlate airway dimension and emphysema score (ES) using patient-based and lobe-based analysis.

METHODS AND MATERIALS
Forty-one patients with COPD (39 men, 2 women; mean age, 66.75 years; range, 51-78 years) were included in this study. ES (relative area under the threshold of -950 HU), airway dimensions (wall area, WA; total airway area, TAA) at the level of segmental bronchi, and lobe segmentation were obtained with a commercial software package (Thoracic VCAR, General Electric Medical System). For each patient and lobe, summation of WA to summation of TAA (WA/TAA) was calculated. Correlation analysis was performed between software measurements and pulmonary function test results, and between airway dimensions and ES. The correlation coefficients within subjects were also calculated using ANCOVA.

RESULTS
ES showed significant correlations with FEV1 (%predicted) (r=-0.523, p<0.01) and GOLD stage (r=0.497, p<0.01), whereas airway dimensions did not. On patient-based analysis, WA/TAA did not show a significant correlation with ES. On lobe-based analysis, however, a significant correlation between WA/TAA and ES was observed (r=-0.251, p<0.01). On ANCOVA, there was a significant effect of WA/TAA on ES (p=0.048) after controlling for the effect of subjects and a significant correlation between WA/TAA and ES (r=-0.154, p<0.01)

CONCLUSION
Our results suggest that regional changes in airways at the level of segmental bronchi have a significant correlation with parenchymal destruction in patients with COPD.

Poster Session 2-2

Comparison of Airway Remodelling Assessed by Three-dimensional Computed Tomography between Bronchial Asthma and Chronic Obstructive Pulmonary Disease

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Objective: There are few studies of direct comparison between asthma and COPD on airway remodelling assessed by computed tomography (CT). We hypothesized that differences might exist in airway dimensions between them showing similar levels of airflow limitation, under clinically stable condition.

Methods: We recruited 19 older asthmatic patients showing FEV1/FVC<70% with no or 5 pack-year of smoking history, randomly selected 28 age-matched COPD patients demonstrating similar airflow limitation from participants to Hokkaido COPD cohort study. 13 age and sex-matched healthy never-smokers were enrolled. Using our software, we selected 8 airways in the right lung, and measured wall area percent (WA%) and airway luminal area (Ai) at the mid-portion of the 3rd to the 6th generation of each airway. For comparison, we took the average of 8 measurements per generation.

Results: Both FEV1%predicted and FEV1/FVC were similar between asthma and COPD (82.3 ± 3.3%SE vs. 77.6 ± 1.8% and 57.7 ± 1.6% vs. 57.9 ± 1.4%). At any generation, WA% was larger and Ai/BSA was smaller in asthma, followed by COPD, then by controls. Significant differences were observed only between asthma and controls in WA% of the 3rd to the 5th generation and Ai/BSA of any generation, but not in either variable between COPD and controls. There were significant differences in Ai/BSA of any generation between two diseases.

Conclusions: Airway remodelling assessed by CT may be more prominent in older asthmatic patients compared with age-matched COPD patients at the airways of the 3rd to the 6th generation showing similar levels of airflow limitation, in clinically stable condition.
Purpose: To test severity indices derived from three-dimensional reconstruction images of lungs using multi-detector row CT (MDCT) are correlated with the data of pulmonary function tests in chronic obstructive pulmonary disease (COPD).

Material and Methods: This study included 23 patients (23 men; mean age, 71.8 years; age range, 51 to 85 years) who were diagnosed as COPD, and examined by MDCT (Brilliance iCT, Philips) in our hospital. Their values for predicted forced expiratory volume in 1 second (FEV1.0) ranged from 24.9 to 103.6% (mean, 64.1%). The lungs were reconstructed as 3D volume model, and were assessed with the application (Lung Emphysema, Philips). Relative volumes of lungs with attenuation coefficients lower than -950 HU as the threshold were calculated, and the value was defined as low attenuation volume (LAV)%.

Results: The mean LAV% in these patients was 11.8% (range, 0.35 to 38.89%), and the mean CT attenuation value was -871.23 HU (range, -919.39 to -814.85 HU). Significant correlations were observed between LAV% and FEV1.0 (%predicted) (r=-0.622, p<0.01), as well as between the mean CT attenuation value and FEV1.0 (%predicted) (r=-0.539, p<0.01).

Conclusions: Relative lung volume with attenuation coefficients lower than -950 HU and the mean CT attenuation value of the whole lungs measured with MDCT may be useful quantitative indicators reflecting the severity of the pulmonary emphysema.

Poster Session 2-4

Quantitative Assessment of Chronic Obstructive Pulmonary Disease (COPD) with Multi-detector row CT (MDCT): Comparison with Lung Structure and Pulmonary Function Test

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Purpose: To test severity indices derived from three-dimensional reconstruction images of lungs using multi-detector row CT (MDCT) are correlated with the data of pulmonary function tests in chronic obstructive pulmonary disease (COPD).

Material and Methods: This study included 23 patients (23 men; mean age, 71.8 years; age range, 51 to 85 years) who were diagnosed as COPD, and examined by MDCT (Brilliance iCT, Philips) in our hospital. Their values for predicted forced expiratory volume in 1 second (FEV1.0) ranged from 24.9 to 103.6% (mean, 64.1%). The lungs were reconstructed as 3D volume model, and were assessed with the application (Lung Emphysema, Philips). Relative volumes of lungs with attenuation coefficients lower than -950 HU as the threshold were calculated, and the value was defined as low attenuation volume (LAV%). Also, the mean CT attenuation value of the whole lungs was measured as an index of the severity of emphysema.

Results: The mean LAV% in these patients was 11.8% (range, 0.35 to 38.89%), and the mean CT attenuation value was -871.23 HU (range, -919.39 to -814.85 HU). Significant correlations were observed between LAV% and FEV1.0 (%predicted) (r=-0.622, p<0.01), as well as between the mean CT attenuation value and FEV1.0 (%predicted) (r=-0.539, p<0.01).

Conclusions: Relative lung volume with attenuation coefficients lower than -950 HU and the mean CT attenuation value of the whole lungs measured with MDCT may be useful quantitative indicators reflecting the severity of the pulmonary emphysema.
Poster Session 2-6

Quantitative Assessment of Low Attenuation Volume Percentage (LAV%) and Mean Lung Density (MLD) in Chronic Obstructive Pulmonary Disease Using Inspiratory and Expiratory MDCT

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Purpose: To quantify low attenuation volume percentage (LAV%) and mean lung density (MLD) in chronic obstructive pulmonary disease (COPD) using inspiratory and expiratory multi-detector row CT (MDCT) scans and analyze the relationship between airflow limitation and these parameters.

Materials and Methods: Forty-two consecutive patients with COPD were enrolled and underwent 64-MDCT (Aquilion ONE, Toshiba) and pulmonary function tests (PFTs). Images were obtained using 120kV and 200mA (CT-AEC) and 0.35sec/rotation. CT images were reconstructed with 0.5mm slice thickness using standard and bone algorithm. Using commercial software (Virtual Place, Aze), the whole lung volume was segmented with attenuation between 1024 and 740 HU. Low attenuation volume percentage (LAV%) was segmented with attenuation less than 960HU, and we also quantified mean lung density (MLD). Then, we assessed the relationship between these parameters and PFTs.

Results: LAV% in inspiratory and expiratory CT scan had an inverse correlation with FEV1/FVC and FEV1 (%predicted). The correlation coefficients of LAV% in inspiratory CT with FEV1/FVC and FEV1 (%predicted) were -0.66 (p < 0.001) and -0.60 (p < 0.001), respectively. The correlation coefficients of MLD ratio with FEV1/FVC and FEV1 (%predicted) were -0.58 (p < 0.001) and -0.68 (p < 0.001), respectively.

Conclusion: This study suggests that LAV% and MLD ratio can be useful for the evaluation of the severity of COPD.

Poster Session 2-6

Relationship between Low Attenuation Areas Evaluated by Thurlbeck Panel Method and Body Weight in Patients with Stable COPD

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Purpose: The purpose of this study was to verify the effectiveness of Thurlbeck panel method in the evaluation of emphysema in COPD patients. The second purpose of this study was to clarify the relationship between low attenuation areas (LAA) and body weight, exercise capacity and HRQOL in these patients.

Methods: Emphysema was evaluated in 32 COPD patients (30 males, 2 females, mean age 76.6yrs, BMI: 21.4kg/m², FEV1/FVC: 46.0%, FEV1pred: 55.5%). These patients were divided into three groups; under body weight group (%IBW < 90%), normal body weight group (90% < %IBW < 110%) and over weight body weight group (%IBW > 110%). High resolution CT was used to quantify LAA with -960 HU at the cut-off level. CT sections were analyzed at bilateral upper, middle and lower lung fields. LAA was scored using Goddard method and Thurlbeck panel method by two pulmonologists and one radiologist independently. Spearman's rank correlation was used to evaluate the relationship among LAA, IBW and exercise capacity. Results: CT scores evaluated by Goddard method and Thurlbeck panel method were significantly correlated (r=0.83, p<0.0001). There was a significant negative correlation between CT scores and %IBW (r=-0.505, p<0.01). CT scores in low body weight group were smallest in three groups.

Conclusion: A low IBW is associated with the presence of emphysema suggesting the concept of different COPD phenotype. Our data also suggest effectiveness of Thurlbeck panel method in evaluation of LAA in COPD.
**Poster Session 2-8**

**Effects of Chronic Obstructive Pulmonary Disease on Thoracic Vascular Calcifications**

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**Background:** Although some of the associations between COPD and atherosclerosis may be the result of common risk factors such as smoking, recent epidemiological evidence suggest that impaired lung function and emphysema are risk factors for cardiovascular disease due to systemic inflammation. We tested the hypothesis that COPD is a risk factor for cardiovascular calcifications, independent of tobacco use.

**Subjects:** 72 patients referred for MDCT lung cancer screening were included and grouped according to the severity of pulmonary disease. 25 subjects without pulmonary disease (mean age 58 ± 5 years), 25 subjects with mild COPD (GOLD I/II) (mean age 59 ± 4 years), and 22 subjects with severe COPD (GOLD III/IV) (mean age 62 ± 6 years). Overall volume of thoracic vascular calcifications (thoracic aorta, coronary arteries, supraaortic branches, and aortic valve) was determined by applying a volume score on low dose un gated MDCT scans. Effects of FEV1%, smoking status, gender, age, BMI, emphysema index and 15th percentile point on vascular calcifications were determined by stepwise multiple linear regression analysis.

**Results:** There was no difference in age between the different severity groups. Median values of overall calcium in severe COPD were 1875 ± 2577 mm3, in mild COPD 721 ± 1082 mm3 and in patients without COPD 880 ± 1793 mm3. Multiple linear regression analysis revealed that only age was linked to the overall volume of calcifications (R2=0.162; p ≤ 0.05).

**Conclusion:** In our cohort, pulmonary disease with concomitant systemic inflammation was not independently associated with thoracic vascular calcifications. Only age could be considered as a significant contributor to vascular disease.

**Funding Acknowledgement:** The work was supported by the Competence Network Asthma/COPD funded by the German Federal Ministry of Education and Research (FKZ 01GI0884).

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**Poster Session 2-8**

**Analysis of Ventilation Abnormalities in Patients with Obstructive Lung Diseases on Xenon-enhanced CT According to Lung Lesion Patterns: Multicenter Study**

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**Purpose:** To analyze ventilation abnormalities on xenon-enhanced CT in patients with obstructive lung diseases according to lung lesion patterns.

**Method and Materials**

Forty-seven subjects (M:F=35:12, mean age, 61.8 yrs) with obstructive PFT abnormality were enrolled from two centers. Patients’ diagnoses were COPD (32), asthma (4), bronchiolitis obliterans syndrome (3), diffuse panbronchiolitis (2), bronchitis (2), bronchiectasis (2), bronchopneumonia (1), and anthracofibrosis (1). All subjects underwent single phase, thoracic dual-energy CT using dual-source CT (Definition and FLASH, Siemens) when they inhaled 30% xenon for 90 seconds. Lung lesion pattern was evaluated on weighted average image and ventilation defect (VD) was evaluated on color-coded xenon map. In each lobe, lung lesion pattern in point of both airway and parenchyma, extent of ventilation defect per lesion (VD/lesion, %), and extent of ventilation defect per lobe (VD/lobe, %) were evaluated.

**Results:** There was no noticeable change in monitored parameters in all subjects. Among 235 lobes of 47 patients, 231 lobes were evaluated and 4 lobes were excluded due to total collapse. Mean VD/lobe was 37.2%. In point of parenchyma, lesion patterns included centrilobular emphysema (CLE, n=82), panlobular emphysema (PLE, 8), hypoattenuation (78), bullous emphysema (1), ground glass opacity (1), and normal (61). In point of airway, lesion patterns included diffuse bronchial wall thickening (n=135), cellular bronchiolitis (56), large airway obstruction (6), bronchiectasis (4) and normal (30). The results of VD/lesion in point of parenchyma were: mild CLE, 34.6%; moderate CLE, 71.2%; severe CLE, 80.9%; moderate PLE, 100%; hypoattenuation, 96.4%. The results of VD/lesion in point of airway were: bronchial wall thickening, 35.0%; cellular bronchiolitis, 80.4%; large airway obstruction, lobar, 85%; segmental/subsegmental, 100%; bronchiectasis, 25.0%.

**Conclusion:** Areas of hypoattenuation, cellular bronchiolitis, segmental/subsegmental airway obstruction, and PLE showed relatively matched areas of ventilation defect, whereas areas of CLE, bronchial wall thickening, and bronchiectasis showed smaller extent of ventilation defect than anatomical extent.
**Poster Session 3-1**

Evaluation of the Extent of Ground-glass Opacity on High-resolution CT in Patients with Interstitial Pneumonia: Comparison between Quantitative and Qualitative Analysis

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**Purpose:** To compare quantitative and qualitative analysis in the evaluation of the extent of ground-glass opacity on high-resolution CT (HRCT) in patients with interstitial pneumonia (IP) with systemic sclerosis (SSc).

**Method and Materials:** Fifteen patients with refractory IP associated with SSc received peripheral blood stem cell transplantation, and HRCT was performed before and 12 months after treatment. We evaluated the activity of the IP by monitoring serum KL-6 and %VC. Five readers analyzed the extent of GGO (%GGO) on HRCT obtained before and 12 months after treatment using qualitative analysis (estimated to the nearest 10% of parenchymal involvement). Quantitative analysis of the %GGO was also performed using a commercially available workstation. We also assessed the therapeutic effect by calculating KL-6 and %VC during one year after treatment. Pearson’s correlation coefficients were used to examine the relationship between pretreatment %GGO, %GGO, and KL-6 and %VC. Thereafter, we compared the correlation coefficient value (r) between quantitative and qualitative analysis.

**Results:** The %GGO was correlated negatively with KL-6 and correlated positively with %VC on both quantitative and qualitative analysis. However, evaluation with quantitative analysis showed stronger correlation than the qualitative analysis.

**Conclusion:** Quantitative analysis of the extent of ground-glass opacity (GGO) on high-resolution CT (HRCT) using a workstation could aid in making a correct decision in predicting or monitoring the therapeutic response of interstitial pneumonia (IP) compared with qualitative analysis.

**Poster Session 3-2**

Correlation between Pirfenidone Effect and the Initial CT Findings in the Patients with Idiopathic Pulmonary Fibrosis

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**Objectives:** Pirfenidone is a new anti-fibrotic drug for idiopathic pulmonary fibrosis (IPF). We evaluated the prognostic value of CT image for the response to pirfenidone.

**Materials and Methods:** In this retrospective cohort study, written informed consent was waived. The subjects were 51 consecutive patients (37 males, 14 females; mean age, 69.9 ± 8.3, ± SD) with IPF. The pattern of usual interstitial pneumonia was confirmed histopathologically using surgical biopsy in 33 patients. About a half of patients were receiving the combination therapy with corticosteroid (n=20) and immunosuppressants (n=11). We checked side effects and pulmonary function tests (PFTs). In PFTs after 12 months, 10% or greater decline of VC in predicted value was defined deterioration. We also analyzed initial CT quantitatively using originally-developed computed-assisted system. This system can divide the lungs into normal (N-pattern) and each lesion pattern.

**Results:** Twelve patients discontinued pirfenidone because of adverse effects; severe appetite loss (n=7), rash (photosensitivity) (n=3), liver dysfunction (n=1) and others (n=1). In 39 patients who were continuing pirfenidone, 7 patients died and 8 patients deteriorated (non-responder). Other 24 patients were stable in PFTs (responder). Initial vital capacity (VC) %predicted was significantly smaller in non-responder (63.6 ± 20.4% vs 75.5 ± 14.0%, p=0.037). In quantitative CT results, initial N-pattern volume was significantly smaller in non-responder (25.2 ± 13.9 vs 37.6 ± 12.9%predTLC, p=0.006), and Consolidation-pattern volume was larger in non-responder (1.1 ± 0.8 vs 0.6 ± 0.5%predTLC, p=0.021).

**Discussion and Conclusion:** Pirfenidone clinically stabilized disease in 62% of patients for one-year follow-up. The initial CT findings were correlated with response to pirfenidone.
Poster Session 3-3

Relation between Radiological Abnormalities and Functional Impairment Assessed from Distributional Analysis on Ventilation and Diffusing Capacity to Perfusion in Idiopathic Interstitial Pneumonias

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We attempted to clarify the simultaneous distributions of VA/Q and diffusing capacity to perfusion (D/Q) in patients with idiopathic interstitial pneumonias (IIP) with varied pathological abnormalities. We categorized the patients with IIP into three groups based on HRCT findings; group I (n=6): ground-glass with no obvious reticular pattern, group II (n=9): reticular pattern associated with honeycombing cysts but without distinct ground-glass opacities, and group III (n=12): a mixture of ground-glass and reticular patterns. To assess functional abnormalities in their lungs, we determined VA/Q and D/Q distributions by using O₂, CO₂ and CO together with six foreign inert gases as indicator gases. We transformed the measured data on indicator gases in arterial blood into a continuous distribution of Q in the VA/Q-D/Q field. Group I subjects exhibited bimodal Q distributions along the D/Q axis but fairly homogeneous Q distributions along the VA/Q axis. Group II subjects showed markedly abnormal VA/Q distributions but unimodal Q distributions along the D/Q axis. Group III subjects showed broad but unimodal Q distributions along the D/Q axis. Q distributions along the VA/Q axis in group III were qualitatively similar to those observed in group II. In IIP, active alveolitis or acinitis played a major role in producing low D/Q regions impeding gas exchange by a diffusion limitation, while extensive fibrosis with minimal inflammation accounted for low D/Q as well as low VA/Q regions. These findings suggest that hypoxemia in IIP patients may be caused by inhomogeneous distributions of D/Q in combination with those of VA/Q.

Poster Session 3-4

Clinical Considerations of Postoperative Acute Exacerbation in Patients with Interstitial Pneumonia Combined with Lung Cancer

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Objective
The purpose of this study was to examine the predictive factors of postoperative acute exacerbation of interstitial pneumoniap Patient with lung cancer.

Methods
Subjects were 530 patients who received operation in our hospital. We reviewed CT images obtained before surgery to evaluate presence of IP. We classified IP into three patterns; UIP pattern, NSIP pattern and combined pulmonary fibrosis and emphysema(CPFE) pattern. We also analyzed other clinical data such as background, hematological examinations, respiratory function tests and operation-related factors.

Results
We found 87 patients with IP in total 530 patients. The 15 patients had UIP pattern, 13 patients had NSIP pattern and 59 patients had CPFE pattern. The postoperative acute exacerbation of IP was observed 5 of 87 patients(5.7%); 4 out of 15 patients with UIP pattern(26.7%), 1 out of 13 patient with NSIP pattern(7.7%). Non of patients with CPFE pattern developed acute exacerbation. In background and hematological examination, Hugh-Jones score, preoperative oxygen use, preoperative steroid use and WBC were significantly higher in the patients developed acute exacerbation(AE group) than in the patients did not developed acute exacerbation(NAE group). In respiratory function test and operation-related factors, %FVC and tidal volume during one lung ventilation were significantly lower in the AE group than in the NAE group.

Conclusion
The UIP pattern on preoperative CT was a risk factor of postoperative acute exacerbation.
Pulmonary aspergillosis has been regarded as an important complication of patients with chronic fibrosing interstitial pneumonia (CFIP) with or without immunosuppressive treatment, however, their clinicoradiographical features have not been fully described.

Objective
To clarify the clinical and radiographic features of pulmonary aspergillosis in patient with CFIP of unknown etiologies, we retrospectively analyzed medical records of 196 patients with CFIP who visited our department between September 2007 and May 2010.

Results
Out of 196 patients with CFIP, 6 cases of pulmonary aspergilloma and 6 cases of chronic necrotizing pulmonary aspergillosis were found. Each lesion of aspergillosis developed in the pulmonary lesions of airspace enlargement, such as honeycombing, emphysema, or thick-walled large cyst. In patients with pulmonary aspergillosis, the mean smoking index was high, pulmonary emphysematous changes were highly prevalent, and the mean diffusion capacity for gas exchange was markedly decreased. Ever-smokers with combined pulmonary fibrosis and emphysema were more likely to be complicated with pulmonary aspergillosis, compared to ever-smokers without emphysema as well as never-smokers.

Conclusion
Patients with smoking-related interstitial pneumonia with emphysema was likely to be suffered from pulmonary aspergillosis. Airspace enlargement of the lung was the site of infection, possibly due to insufficiency in local innate immunity.

Aim:
To analyze thoracic and abdomino-pelvic radiologic findings of Japanese patients with lymphangioleiomyomatosis.

Subjects and methods:
The chest CT images of 138 patients with LAM (125 sporadic LAM and 13 TSC-associated LAM) and also abdomino-pelvic CT images of 76 patients (70 sporadic and 6 TSC) among them were studied by 4 radiologists. All were female. Although this is a cross-sectional study, the frequency of “lymphatic lesion” in each part of the body may answer to the question.

Results:
Pulmonary cysts were demonstrated in all patients and its mean extent in the lung field was 39%. Small nodules in the lung field (4%), dilatation of thoracic duct (4%), and pleural effusion (11%) were identified with the similar frequency with those reported by NIH group. In abdomino-pelvic scan, renal angiomyolipomas (AMLs) (26%), AMLs in liver (24%), ascites (12%), lymphangioleiomyomas (46%), and lymph node enlargement (20%) were identified. Lymphatic involvement was more frequently demonstrated pelvic cavity than thoracic cavity.

Conclusion:
Radiologic findings and their frequencies in the pulmonary manifestation was similar to that of NIH group, but this may not be the case in the extrapulmonary manifestations: lower incidence of renal AML, increased detection of liver AML, and frequent lymphatic involvement in the pelvic cavity in Japanese LAM patients.

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Conclusion:
Radiologic findings and their frequencies in the pulmonary manifestation was similar to that of NIH group, but this may not be the case in the extrapulmonary manifestations: lower incidence of renal AML, increased detection of liver AML, and frequent lymphatic involvement in the pelvic cavity in Japanese LAM patients.
(Background)
Pneumomediastinum as well as pneumothorax has been regarded as a distressing problem in patients with chronic fibrosing interstitial pneumonia (CFIP). Some are intractable and they are sometimes life-threatening. However, their clinicoradiographical features have not been fully described.

(Objective)
To clarify the clinical and radiographic features of pneumothorax and pneumomediastinum developed in patients with CFIP, we retrospectively analyzed medical records of patients with CFIP who visited our department between January 2007 and May 2010.

(Results)
Out of 498 subjects with CFIP, 37 were found to have pneumothorax, pneumomediastinum or both. Each number of cases was 12, 8, and 17, respectively. The overall mean smoking index was 24 pack years. By high-resolution computed tomography, 21 subjects demonstrated the usual interstitial pneumonia pattern of interstitial pneumonia, 21 demonstrated emphysema or thick-walled large cyst, and 9 subjects demonstrated bilateral pneumothorax. 14 out of 16 subjects, who could be estimated the severity of CFIP by SpO2, %FVC, and %DLco, showed moderate to severe CFIP. The severity of pneumothorax was usually up to the second degree. The development of pneumomediastinum was associated with glucocorticosteroid administration.

(Conclusion)
Pneumothorax and pneumomediastinum were clinically significant complications during the course of CFIP.
Poster Session 4-1

Effect of Pulmonary Rehabilitation on Parenchymal Lung Motion Evaluated with Dynamic MRI in COPD Patients

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Background:
Pulmonary rehabilitation does not improve lung function, but it increases exercise tolerance and improves quality of life; the underlying mechanism is uncertain. Recently, parenchymal lung motion in COPD patients is recognized to be different from it in healthy subjects evaluated with dynamic MRI.

Purpose:
To analyze parenchymal lung motion evaluated with dynamic MRI before and after pulmonary rehabilitation in COPD patients.

Methods:
A total of 12 subjects were examined, including 7 COPD patients and 5 control individuals. Rehabilitation programs feature 2 sessions per week, each lasting 1 hour for 12 weeks. Dynamic MRI and clinical assessments including lung function test, 6-minute walk test and QOL test were performed before and after rehabilitation. MRI was performed using 10 sample points for motion analysis during deep slow breathing. The directional ratio of the sample points (the ratio of the cranio-caudal change to the dorsal-ventral change) was calculated as parenchymal lung motion.

Results:
In patients with COPD, the directional ratio increased compared with that in control (1.98 ± 0.58 vs 1.33 ± 0.35, respectively). After pulmonary rehabilitation, the directional ratio decreased (1.98 ± 1.33 vs 1.31 ± 0.40, respectively).

Conclusions:
The parenchymal lung motion was altered from cranio-caudal to dorsal-ventral direction after pulmonary rehabilitation in patients with COPD and was suggested to become similar to that in control individuals.

Poster Session 4-2

Diaphragmatic Motion before and after Plication of Central Tendon in Patients with Diaphragmatic Eventration

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As diaphragmatic eventration is a chronic disorder, it is rather difficult for both a patient and a surgeon to select operative treatment with few evidences. There are few reports on the diaphragmatic motion before and after surgery. We performed plication of a thin and extended central tendon of the left diaphragm in two elder patients with symptoms such as palpitation, or exertional dyspnea during daily activities. Dynamic MRI during deep breathing of the chest and the upper abdomen in the mid-coronal and sagittal slices were obtained with a 1.5T MR instrument using fast spoiled GRASS method. In coronal slice, the central part of the left diaphragm was pulled down to the right side during inspiration while the lateral part did not move like a piston but did like a hinge. In sagittal slice, the posterior part of the left diaphragm moved a little resulting smaller excursion than the anterior part. These impairment were basically same but lesser after operation. Although the dome shape of the left floppy diaphragm could be fairly corrected, tension yielded by the healthy right diaphragm during contraction might be transferred through the central tendon, and affect strongly the left diaphragmatic motion due to strength difference between the two hemidiaphragms.
Dynamic MRI Measured Respiratory Motion of the Internal Structures of the Lung. Differences between Healthy Subjects and COPD Patients.

Objective
The aim of this study was to evaluate respiratory motion of the internal structure of the lung using dynamic MRI in healthy subjects and patients with chronic obstructive pulmonary disease (COPD).

Methods
The subjects were 5 normal volunteers and 21 patients diagnosed cigarette smoking-related COPD. Imaging sequence of the dynamic MRI was a balanced fast field echo (BFFE). We sequentially captured 80 frames of the mid sagittal portion of the right lung while repeating forced deep breathing. We set 15 tracking points for pulmonary vessels, 5 tracking points in the upper lobe and 10 tracking points in the middle and lower lobes and measured maximal motion distance of each tracking point for ventrodorsal direction (X-axis) and craniocaudal direction (Y-axis). We compared these parameters with spirometric data and evaluated the correlation between airflow limitation and the respiratory motion of the internal structure of the lung.

Results
Respiratory motion distance (X and Y-axis of middle and lower lobe) and spirometric data such as forced expiratory volume (FEV) /forced vital capacity (FVC), FEV1 and %FEV1 showed a highly significant positive correlation. %residual volume (RV) showed a highly significant negative correlation.

Conclusion
Dynamic MRI using BFFE sequences is a highly precise technique to directly and noninvasively visualize dynamic respiratory motions of the internal pulmonary structures. Respiratory motions of the internal structure of the lung correlate spirometric data which express airflow limitation in normal volunteers and patients with COPD.

Respiratory Motion Analysis with Correlation Function and Frequency Decomposition from MR Image Sequences of COPD Patients and Normal Subjects

We examine motions of some points on a diaphragm with their correlation and frequency analysis. We expect that the information of correlation and frequency can be used to measure quality of respiratory motion of a lung and for image diagnosis on COPD and so on.

Each part of a lung of normal subject, especially his or her diaphragm, extends toward outside and shrinks toward inside together, while that of a patient sometimes moves unevenly. Recording positions of some points on a diaphragm, we can get sequences of each of them. Frequency analysis involved with correlation is performed to these sequences and the spectrum is used to obtain features of a respiratory motion of a subject. These features are used as measures of quality of respiratory motion.

Bispectrum is a Fourier transformation of correlation function of three sequences. We got three sequences of different positions of points on diaphragms and calculated bispectrum of them in our previous study. There was a difference between the phases of bispectrum of patients and ones of normal subjects. It suggests that the phase of bispectrum can be used as a measure of respiratory motion.

We verify that the difference shows up again by using points of positions different from previous study on each diaphragm in this time. In addition, we present results from correlation analysis with wavelet transform for sequences of points on diaphragms.
Poster Session 4-5

Evaluation of Airway Dimension During Respiration

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Introduction
It is widely known that the lung volume changes during respiration. However, it is unknown how the airway dimensions change during normal respiration.

Objectives
We hypothesized that, in normal subjects, airway length and luminal area extend from expiration to inspiration. To investigate this hypothesis, we used inspiratory and expiratory CT of normal subjects.

Methods
We recruited 13 normal and healthy subjects and they underwent both inspiratory and expiratory CT. We analyzed volumetric CT data and measured the airway length, luminal area, wall thickness, wall area, and WA% (percentage of wall area to total airway area) of right apical bronchus (the fourth generation) at inspiration and expiration.

Results
The airway length was longer at inspiration than expiratory (mean (SD) 12 (1.83) vs 10 (1.35) mm, respectively; p<0.001). The airway luminal area at inspiration was greater than expiration (mean (SD) 21.2 (5.53) vs 16.1 (4.77) mm², respectively; p<0.001). There was no difference in the wall thickness between inspiration and expiration. The airway wall area was greater at inspiration than expiration (mean (SD) 34.1 (5.20) vs 32.1 (5.38) mm², respectively; p=0.03). WA% was smaller at inspiration than expiration (mean (SD) 62.6 (3.46) vs 67.1 (3.18) %, respectively; p<0.001).

Conclusion
The airway length of right apical bronchus in normal healthy subjects extended from expiration to inspiration. Luminal area and airway wall area increased but WA% decreased from expiration to inspiration.

Poster Session 4-6

New Method for Ventilation Function Using Dynamic Chest X-ray Examination: Evaluation of Regional Changes in Inspiratory / Expiratory Flow Rate Ratio

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Purpose: Evaluation of a regional ventilation function with use of dynamic chest X-ray. The differential value of density between expiratory and inspiratory phases has possibilities to reflect the regional ventilation function on dynamic chest X-ray.

Subjects: Dynamic chest X-ray from 11 ventilation impairment patients with COPD and 5 normal volunteers were obtained in the upright position in about 10 seconds of tidal breathing at rest. Image data captured at 7.5 frames per second was synchronized with the pulsed X-ray.

Methods: We developed an automatic analyzing software for a regional ventilation function, which includes:
1) Lung fields extraction from dynamic chest X-ray.
2) Identification of small local areas.
3) Calculation of the maximal differential values in each ventilation phase at the corresponding small local area in the series of dynamic chest X-ray.
4) Calculation of the regional relative flow rate ratio as the ratio of peak inspiratory phase to peak expiratory phase.
5) Display a color map for the ratio on a representative chest image from the dynamic chest X-ray.

Results: The regional changes in inspiratory / expiratory flow rate ratio had larger values and broader distribution in patients with COPD compared to those of normal volunteers. The average of the ratio in patients and in normal volunteers was 1.26 ± 0.25 and 1.03 ± 0.07 (mean ± SD, P = 0.013), respectively.

Conclusion: The newly proposed method using dynamic chest X-ray examination enabled us to noninvasively analyze the regional flow rate.
**Poster Session 5-1**

**Hyperpolarized 3He Magnetic Resonance Imaging Biomarkers of Bronchoscopic Airway Bypass in COPD**

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We explored and examined hyperpolarized 3He magnetic resonance imaging (MRI) acquired longitudinally, both pre- and post-airway bypass and compared these to measurements of pulmonary function, the six-minute walk distance (6MWD), St. George’s respiratory questionnaire (SGRQ) and modified medical research council (mMRC) score. A single subject underwent pulmonary function tests, 3He MRI diffusion-weighted, static ventilation, and proton MRI at 3.0T. 3He MRI apparent diffusion coefficients were calculated and as well 3He MRI ventilation and ventilation defect measurements using two-dimensional landmark-based registration of 3He and 1H pulmonary images, and subsequent application of an automated K-means cluster algorithm to generate 3He ventilation defect volume (VDV), and 3He ventilation volume (VV).

Four days prior to bronchoscopic introduction of the airway bypass (Exhale Drug Eluting Stent, Broncus Technologies, Mountainview CA) and six months post-stent, the subject reported mMRC scores of 2 and 1, 6MWD of 288m and 366m and SGRQ scores of 65 and 27 respectively. At eight months pre-stent, whole lung 3He MRI VV was 1.7L and at six months post-stent VV was 4.0L- an increase of 2.3L, whereas the RV/TLC ratio was 57% and 56%. FEV1 was 1.09L and 1.11L, and FVC was 3.15L and 3.51L, pre- and six months post-stent respectively. In conclusion, for a single COPD subject, the high sensitivity of 3He MRI provided a way to detect regional functional ventilation changes that were suggestive of ventilation improvements and that were in accordance with improvements in mMRC score and 6MWD and were not detected by established pulmonary function tests.

**Poster Session 5-2**

**Line-Narrowed Diode Lasers for High-Efficiency Continuous Production of Hyperpolarized Xe-129 using Optimized Cell for High Concentration of Optically Pumped Rubidium**

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A flow-through-type apparatus for the polarization of Xe-129 using four line-narrowed high-power InGaAsP diode laser arrays, with fast and slow axis collimation lenses (Ingeneric) and a volume holographic grating (Ondax PowerLocker™, PLR794.5-22 5-13-1.75-1.5) (794.73 ± 0.15 nm, 40 W 4 = 160 W) was constructed. The obtained maximum polarization of Xe-129 for the natural abundance of Xe gas was 20 ± 5 % when the temperature of the cell was ~ 220 oC and the flow rate of the mixture gas was 40-100 sccm. We have developed a flow-through-type apparatus with a cell structure that is optimized for the hyperpolarization of Xe-129. In this equipment, to increase the laser adsorption coefficient, heating temperatures in the polarization cell are set at 180 to 300 oC. Heating causes the vapor pressure of Rb to increase, and the irradiation cell gap is set at 1 mm, resulting in a polarization rate of ~ 20% and a high rate of polarization of Xe-129 per unit of time at a flow rate of 40-100 sccm. The newly developed equipment makes it possible to realize the continuous production of hyperpolarized Xe-129 with high productivity.
Poster Session 5-3

Development of a Novel Method for Evaluating Pulmonary Disease Using Hyperpolarized Xe-129 Ultrashort Echo-time MRI

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Hyperpolarized (HP) Xe-129 is capable of wide application in lung MRI by the use of both gas- (air space) and dissolved-phase (tissue and blood) signals, offering a powerful tool for the physiological evaluation. However, it is still difficult to obtain the dissolved-phase image due to the extremely short T2* and the low spin density. In the present study, we intended to apply ultrashort echo-time (UTE) sequence in acquiring the dissolved-phase image and to develop a novel method to visualize and evaluate the pulmonary function.

HP Xe-129 gas was produced by a home-built polarizer and continuously supplied to the mouse, which breathed the gas spontaneously. Measurements were performed with a 9.4-T MRI system (Varian). Gas- and dissolved-phase images were acquired using the UTE sequence with frequency selective excitation. We proposed the novel marker, xenon distribution ratio, which is the ratio of magnetization of the dissolved-phase Xe-129 relative to that of the gas-phase Xe-129. The distribution ratio maps were compared between healthy mice and elastase-induced mouse models of emphysema.

The application of the UTE sequence enabled to obtain the dissolved-phase Xe-129 lung image as well as the gas-phase one with relatively high spatial resolution. As a result, we could evaluate the local xenon distribution ratio in the mouse lung by combining these images. It was found that the distribution ratio in elastase-treated mice was reduced and widely distributed compared to that of healthy mice, reflecting impaired diffusing capacity caused by emphysema.

This work was supported in part by JST (a210017 and AS2124004F).

Poster Session 5-4

Non-contrast Enhanced MR Angiography Using Volumetric T2-weighted Imaging Technique on 3T

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Purpose.
To investigate the clinical potential of volumetric T2-weighted MR imaging technique in depicting pulmonary arteries on 3T.

Materials and Methods.
Six healthy volunteers who underwent lung MRI were enrolled in this retrospective study. MR protocol included volumetric T2-weighted imaging (TR/TE 4300-6500/49) with ECG and respiratory gating. To depict pulmonary arteries as bright, MR data acquisition was performed during diastolic phase which was determined by cardiac cine study using a steady state free precession sequence. Two radiologists performed a consensus evaluation of reformatted axial and coronal images on a per-vessel basis (main pulmonary artery (PA), lobar PA, segmental PA, and subsegmental PA). The image quality was graded into three categories (poor, fair, and good).

Results.
In six subjects, the image quality on a per-vessel basis was good in 11 of 12 main arteries (91.7%), 30 of 36 lobar arteries (83.3%), 96 of 108 segmental arteries (88.9%), and 216 of 252 subsegmental arteries (85.7%), respectively.

Conclusion.
MR angiography using volumetric T2-weighted imaging technique appears to be feasible in depicting pulmonary arteries. It may be applicable to patients with renal impairment or allergy to contrast media.
Non-Contrast-Enhanced Pulmonary MR Angiography Using Electrocardiography-triggered Three-Dimensional Half-Fourier FSE with a Swap Phase Encode Extended Data (SPEED) Acquisition and Image Processing

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Three variations of a non-contrast-enhanced MR angiography technique using electrocardiography (ECG)-triggered three-dimensional (3D) half-Fourier FSE fresh blood imaging sequences were acquired to visualize the pulmonary arterial vasculature. Systolic and diastolic images with the phase encode directions swapped between the first and second acquisitions were obtained in ten consecutive healthy volunteers. Prior to the 3D acquisition, an ECG preparation scan was acquired to determine the appropriate diastolic and systolic ECG delay times. For the pulmonary arterial imaging, automatic subtraction of the systolic images from the diastolic images is performed for all of the paired acquisitions to produce arterial images. The two sets of arterial images are then added together to produce the swap phase encode extended-data (SPEED) acquisition data. The subtracted arterial images and the SPEED calculated images then undergo maximum intensity projection (MIP) processing.

The image quality of the pulmonary arterial segments in the three MR angiography techniques were assessed quantitatively by measurements of signal-to-noise ratio (SNR) and contrast-to-noise ratio (CNR). For all pulmonary arterial segments, the MR angiography using SPEED acquisition and image processing technique produced significantly higher SNR and CNR values compared with the MR angiography techniques acquired in the right-left or superior-inferior RO direction alone. Our proposed non-contrast-enhanced MR angiographic technique allows depiction of the pulmonary arteries without dephasing signals due to the relationship of the RO direction and the vessel orientation.

Quantitative Analysis of the Lung using Hyperpolarized Helium-3 Imaging

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A computational framework was developed for quantitative analysis of hyperpolarized helium-3 MR lung ventilation image data. Application to a cohort of 55 subjects (47 asthmatic and eight normal) included the generation of multiple image features comprised of first and second-order statistical measures calculated both regionally and globally. A mutual information feature subset selection algorithm was used to determine which imaging features best characterized clinical diagnosis. The performance of these imaging features were compared with spirometry features. It was found that the top image features perform much better compared with the current clinical gold-standard spirometric values when considered individually. Interestingly, it was also found that spirometric values are relatively orthogonal to these image feature values in terms of informational content.

Related was the development of a segmentation algorithm designed to quantify ventilation defects specifically in these lung image data. Comparative assessment was performed using 43 subjects (8 normal and 35 asthmatic) in which the poorly ventilated regions identified by the computational algorithm were compared to two independent human readers (R1 and R2) who were blinded to all clinical information. Inter-rater agreement between our algorithm and R1/R2 was characterized using Bland-Altman test plotting and calculation of the intra-class correlation coefficient (ICC) between pairings (ICC(R1,R2) = 0.97, ICC(R1,AP) = 0.86, and ICC(R2,AP) = 0.85). The findings indicate the proposed approach may be a potentially useful method for further lung research.
Poster Session 6-2

Integral-based Half-Band Method in Measuring Airway Wall Thickness on CT: Comparison with Conventional Full-Width Half Maximum Method in Actual and Simulated Phantoms

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To compare accuracy and robustness of a new integral-based half-band method with them of a conventional full-width half maximum method (cFWHM) in measuring airway wall thickness on CT using actual and simulated phantoms.

Images were magnified ten-times and boundary of segmented lumen was used as an initial band. The band enlarged outward to the peak of density profile. The integrals of average density was accumulated and doubled for final measurement. The phantom was scanned in 0.75mm collimation with a 16-multi detector row CT scanner (Siemens Sensation 16). CT images were reconstructed by various reconstruction kernels. To mimic complex anatomy of bronchi, simulated phantoms of attached vessel and incomplete airway wall were generated. Measured wall thickness was compared with real thickness and that of cFWHM method using Bland-Altman and paired t-test.

Measured wall thickness on phantom CT images with soft and standard kernels using a new method showed better agreement with actual thickness than cFWHM method (Bland-Altman 95% CI: -0.17 ~-0.01, -0.12 ~0.14, -0.32 ~0.18mm, vs. -1.03 ~0.32, -0.67 ~0.37, -0.28 ~0.44mm, respectively). Using the new method, the accuracy was higher in measuring wall thinner than 1.2mm (all p<0.05). However, on images reconstructed with sharp kernels (B70f, B80f), the measurement errors were larger (all p<0.05). In measuring simulated airway phantoms of attached vessel and incomplete wall, the new method was more accurate (all p<0.001).
Poster Session 6-3

Enhancement of Computerized Differentiation Accuracy of Region Pattern of Diffuse Infiltrative Lung Disease Using Local Context Sensitive SVM Classifier

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PURPOSE
To propose a novel local context sensitive SVM classifier (CSSVM) to improve differentiation of regional disease patterns of diffuse infiltrative lung disease (DILD) at HRCT: Comparison with that of radiologist’s decision.

METHOD AND MATERIALS
900 rectangular regions of interest (ROIs) with 20x20 pixels, consisting of each 150 ROIs representing six regional disease patterns (normal, NL; ground-glass opacity, GGO; reticular opacity, RO; honeycombing, HC; emphysema, EMPH; and consolidation, CONS) were marked by two experienced radiologists with consensus at HRCT images of various DILD. 21 textural and shape features (textural features: histogram, gradient, run-length, co-occurrence matrix and shape features: cluster, top-hat) were calculated and employed to characterize the ROIs. A proposed CSSVM classified the center ROI by using probability (SVM’s decision value) of each class and feature distance and class difference from eight neighborhood ROIs to the current ROI, simultaneously. Sequential forward selection was used for feature selection method. In addition, five folding cross validation with twenty repetitions were performed for cross validation. To validate the results, 900 ROIs and only consensus region of 67 slices drawn by two radiologists were compared, independently.

RESULTS
The accuracy of the proposed method of ROIs and whole lung region (89.88 ± 0.02, and 62.18 ± 0.15, respectively) was higher than that of the conventional SVM classifier (87.39 ± 0.02, and 59.72 ± 0.12, respectively) (paired t test, p = 0.000, and p=0.000, respectively).

CONCLUSION
The proposed local context sensitive SVM classifier showed more accurate differentiation of regional patterns of DILD.

Poster Session 6-4

Newly Developed Denoising Technique: Utility for Quantitative Assessment of Pulmonary Emphysema on Low-Dose CT Examination

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PURPOSE: To evaluate the utility of a new denoising technique for quantitative assessment of emphysema on low-dose CT (LDCT) as compared with the assessment on standard-dose CT (SDCT).

MATERIALS AND METHODS: Twenty-six patients (20 men, 6 women; age, 57-84; smoking history, 0-180 pack-years) underwent standard-dose (150 mAs) and low-dose (25 mAs) CT examinations using a 16-detector-row CT scanner. We utilized a new denoising technique, called adaptive iterative dose reduction (AIDR), for LDCT and reconstructed thin-section images with and without AIDR. As quantitative indices of emphysema, we calculated emphysema index and numbers of four emphysema clusters (cluster 1, 2-8 mm³; cluster 2, 8-65 mm³; cluster 3, 65-120 mm³; cluster 4, >120 mm³) on LDCT without AIDR, LDCT with AIDR, and SDCT. To evaluate the effect of AIDR, we compared Bland-Altman 95% limits of agreement (LOA) between SDCT and LDCT without AIDR as well as SDCT and LDCT with AIDR.

RESULTS: The LOA of emphysema index was -0.084 to 0.055 between SDCT and LDCT without AIDR and -0.046 to 0.076 between SDCT and LDCT with AIDR. The bias of emphysema index on LDCT was slightly improved by using AIDR. The LOA of four emphysema clusters was also improved by using AIDR, especially in small clusters. The LOA of clusters 1-2 were: -36200.6 to 5395.9 and -8974.5 to 1754.0 between SDCT and LDCT without AIDR; -7242.1 to 4224.5 and -1787.4 to 1471.1 between SDCT and LDCT with AIDR.

CONCLUSION: The new denoising technique can be useful for quantitative assessment of emphysema on LDCT.
Lung CT Image Registration Using Diffeomorphic Transformation Models

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Given the importance of quantifying kinematics in the lung and segmenting its complex structure, various algorithms have been proposed in the literature for inter-subject and intra-subject lung registration. In this work, we describe the use of our open source Advanced Normalization Tools (ANTS) software package in the EMPIRE10 challenge. Specifically, we describe the results of applying a subset of possible algorithmic configurations focusing on two ANTS diffeomorphic transformation models: greedy Symmetric Normalization (piscI-gsyn) and exponential mappings (piscI-exp), which placed 1st and 10th respectively in the EMPIRE10 challenge evaluating 24 total algorithms. The registration pipeline begins with an affine alignment using lung masks, which precedes each of the two diffeomorphic models for deformable registration using normalized cross correlation integrated over local windows. The results show that our general image registration algorithm is able to achieve the state-of-the-art performance and is reproducible by using the same set of parameters in a fully automatic and open source pipeline.
This paper describes a new method for airway segmentation from CT images. We propose a form of adjusted image gradients for thin airways segmentation and apply it in multi-stencil fast marching. A graph of airway path segments is constructed from the arrival time function of fast marching. Instead of detecting leakage during segmentation, our method verifies each path segment in a separate step, using a novel leakage cost function defined on the whole path. Our scheme of path removal can be viewed as complementary post-processing for existing region growing methods. Experiments show that the proposed method can remove leakage regions while keeping most thin airways.
Poster Session 7-1

A Low Dose CT Lung Screening Method for Smoking-related Thoracic Abnormalities Using 3D Automatic Exposure Control: Diagnostic Capability and Effectiveness in Radiation Dose Reduction (iLEAD Study)

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PURPOSE: To determine the utility of 3D automatic exposure control system of 64-detector row CT system for low-dose chest CT screening examination of smoking-related thoracic diseases.

METHOD AND MATERIALS: 75 consecutive enrollees (40 male, 35 female) underwent a chest non-contrast MDCT examination consisting of two scans; (1) 3D AEC (automatic exposure control) scan and (2) non-AEC scan (fixed tube current).

RESULTS: The mean kappa values were 0.70, 0.81, 0.72, 0.43, 0.63 and 0.67 for aortic aneurysm, coronary calcification, emphysema, ground-glass opacity, honeycombing and nodules, respectively.

CONCLUSION: 3D AEC-assisted low dose CT protocol can produce the screening results consistent with standard dose CT protocol.

Poster Session 7-2


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Objectives: To quantitatively assess image quality relative to radiation dose for standard and low-kVp CT of the lungs with different state-of-the-art MDCT systems, and relate these findings to qualitative evaluation of lung structures and tissues.

Materials and Methods: For three MDCT systems, a 32- and a 64-row detector dual-source CT (Somatom© Definition DS/Flash, Siemens Medical Solutions, Forchheim, Germany), and a 128-row detector CT (Brilliance iCT, Philips Medizin Systeme GmbH, Hamburg, Germany) image data of a cylindrical water (Ø =40cm), a low-contrast, and a spatial resolution phantom (Siemens Cone-Beam Phantom, QRM GmbH, Moehrendorf, Germany) as well as a formalin-fixed porcine lung explant have been acquired using standard (120kVp/150mAs) and low-kVp (80kVp/300mAs) protocols. Primary collimation varied from 256, 128, 64, 32, and 8x0.5/0.6mm, reconstructed slice thicknesses were 2.0, 1.0, and 0.5mm.

Other scanning parameters (fixed): 300/600mA tube current, 0.5s rotation time, helical pitch 1, 10cm scan length, 500mm FOV. Apart from qualitative evaluation of image quality in the lung explant, image noise, homogeneity, low-contrast and spatial resolution have been determined from geometric phantom image data. For each dataset, CTDIvol and DLP have been determined and compared to console prediction and scan summary values.

Results and Conclusions: For lung CT imaging, state-of-the-art MDCT systems differ in the efficiency of their radiation dose usage with regard to achievable image quality. Independent of the systems’ dose efficiency, low-kVp protocols at elevated tube current-time products show potential to allow for future radiation dose savings, although spatial and low-contrast resolution are affected by increasing image noise.

Acknowledgements: This research project is funded by the German Federal Ministry of Education and Research (BMBF) within the collaboration project: “Innovative Methods for the Optimization of Radiological Applications in Biomedical Imaging”, grant numbers 02NUK008G/B.
Poster Session 7-3

Proposal of New Method: Single Breath Xenon Ventilation CT

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[Purpose] CT ventilation images in clinical studies have been acquired by re-breathing of stable xenon (Xe) exclusively. Though the method is suitable for quantitative assessment of ventilation in normal tidal breathing, it requires relatively large amount of Xe leading to side effect of anesthesia. The method is not suited for some diseased lung where the analytic model does not fit. We propose a new method to obtain Xe concentration (Xe-C) images by a single breath of Xe.

[Materials and Methods] A phantom of an acryl cylinder of 4 L capacity inserted with six small cylinders of variable apertures of 1 - 12 mm diameter was created (Anzai Medical). The phantom was pictured by a dual energy, dual source CT (SOMATOM Definition flash, Siemens) after aspirating 3 L of 35%Xe-65%O₂ mixture prepared by Xe control system (AZ-726, Anzai Medical) into the phantom. Four volunteers underwent dual energy CT by the dual source CT during a breath hold after a single vital capacity breath of 35%Xe-65%O₂ mixture. Three-material decomposition was used to obtain Xe-C images.

[Results] The Xe-C images showed expected Xe distribution in the phantom and in the volunteers: Xe concentrations were parallel with the aperture sizes of the inserted cylinders. In healthy volunteers, Xe concentration was higher in the anterior parts of the lung. In one smoking volunteer with emphysema, the emphysematous portion has lower Xe concentration.

[Conclusion] Single vital capacity breath of Xe followed by dual energy CT and three-material decomposition can create ventilation image.

Poster Session 7-4

Relationship between Lung Morphologic and Functional Abnormality Assessed by Breath-Hold Perfusion SPECT-CT Fusion images

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Automated deep-inspiratory breath-hold (DIBrH) perfusion and ventilation SPECT-CT fusion images were used to objectively assess relationship between lung CT morphologic and functional abnormality in various lung diseases.

The fusion images showed a good correlation between lung CT attenuation changes and perfusion heterogeneity at CT mosaic attenuation areas, as perfusion was persistently decreased at low CT attenuation areas and preserved at intervening high CT attenuation areas in vascular or airway obstructive diseases and chronic obstructive lung diseases, and as perfusion was persistently decreased at high CT attenuation areas in interstitial lung diseases and bronchial pneumonia. These images frequently showed perfusion defects even in normal lung areas on CT in patients with vascular obstructive diseases, hepatopulmonary syndrome and pulmonary arteriovenous fistulas.

Assessment of relationship between lung CT morphologic and functional abnormality on the fusion images contributes to clarification of mechanisms and pathophysiology of abnormal lung CT appearance in various lung diseases.
Poster Session 7-6

Pulmonary Lymphatic Drainage to the Mediastinum Based on CT Observations of Primary Complex of Pulmonary Histoplasmosis

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PURPOSE: To assess mediastinal lymphatic drainage patterns from each pulmonary segment/lobe using computed tomographic (CT) observations of calcified primary complex pulmonary histoplasmosis.

MATERIALS AND METHODS: We assessed 585 CT studies of patients with primary complex histoplasmosis consisting of a pulmonary lesion in single segment and hilar and/or mediastinal nodal disease. We assessed the distribution of hilar and mediastinal nodal involvement depending on pulmonary segment and lobe with focus on the incidence of N1 and skip N2 involvement, the mean number of involved mediastinal nodal stations in non-skip N2 and skip N2 group, and preferentially involved mediastinal node stations. Institutional review board approval was obtained without requirement for informed consent for this retrospective study.

RESULTS: Skip involvement was common in apical segment (9/45, 20.0%), posterior segment (7/31, 22.6%), and medial basal segment (13/20, 65.0%) in the right lung and in apical posterior segment (16/47, 34.0%) in the left lung. The incidence of skip involvement in each segment of both lungs shows a significant inverse correlation with that of N1 involvement (r=-0.51, P<0.05). The mean number of involved mediastinal nodal stations in non-skip N2 and skip N2 group in all segments of both lungs were 1.4 (434/301) and 1.2 (93/77) and the former was significantly larger than the latter (p<0.01). Preferentially involved mediastinal node was shown in relation to each pulmonary segment and lobe.

CONCLUSION: Our data showed a predictable pattern of segmental and lobar lymphatic drainage to the mediastinum, including skip invasion.
Objective: In T4 stage of non-small cell lung cancers, specific mediastinal invasion has better prognosis, which presumes direct pathways could be present other than the pulmonary hilum. We evaluated whether cross-bridging structures between each lung lobe and the mediastinum in the pneumothorax patients other than the pulmonary hilum exist and what factor affects these structures.

Material and Methods: The studies were approved by the Ethics Committee. We analyzed multidetector low computed tomographic images of 81 patients with pneumothorax for the incidence and distribution pattern of cross-bridging structures in the pleural cavities.

Result: Cross-bridging structures were observed in 34 of 54 patients (63%) in the right pneumothoraces, in 19 of 32 patients (59%) in the left pneumothoraces. The positive correlation was noted between the number of cross-bridging structures and aging or with pulmonary diseases. The cross-bridging structures were also present in specific distribution patterns, a part of them were common in their locations regardless of pulmonary disease or age.

Conclusion: Cross-bridging structures in pneumothoraces existed more than 50% of patients and were more frequently in older patients and with pulmonary diseases. These structures indicated specific distribution pattern and could play an important role as a direct pathway from each lung to the mediastinum.
Poster Session 8-2

CT-guided Pericardiocentesis for Recurrent Cardiac Tamponade due to Carcinomatous Pericarditis in a Patient with Adenocarcinoma of the Lung

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A 60 year-old man with dyspnea on effort was admitted to our hospital because of an abnormal lesion on Chest X-ray.

The next day of admission, he rapidly developed dyspnea on rest.

Chest-X-ray showed cardiomegaly and transthoracic echocardiogram (TTE) indicated massive pericardial effusion (PE).

Ultrasound-guided pericardiocentesis through the subxyphoid approach was performed. Cytology of pericardial effusion and histology of lung specimens from bronchoscopy revealed adenocarcinoma of the lung.

Despite of Erlotinib treatment, the patient developed cardiac tamponade again.

PE at apex was minimum with pericardial adhesion, which was resulted from the previous puncture.

Chest CT demonstrated massive PE located at left and right paraventricular spaces.

We chose CT-guided pericardiocentesis from the right parasternun and successfully placed a dorainage catheter.

CT enables to visualize intrathoracic anatomy including intercostal and internal mammary arteries, and thereby allows safe access to the pericardial space in case that subxyphoid or apical approach by TTE is difficult.
A Case of Malignant Mesothelioma with Diffuse Swelling Mediastinum Lacking of Apparent Pleural and Pericardial Lesion

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We report a case of malignant mesothelioma with very unusual radiological feature. A 67-year-old man visited our hospital with complaint of exertional dyspnea. This patient was a retired craftsman of fabric items and gave a negative history for both environmental and occupational asbestos exposure. Chest X-ray showed bilateral pleural effusion and swelling of mediastinum. CT scan revealed remarkable swelling of mediastinal soft tissue density, but apparently no pleural or pericardial lesion was detected. The pleural effusion was exudates, and hyaluronic acid of pleural effusion was markedly increased to 302 mcg/ml. The diagnosis of malignant mesothelioma was established by histopathological findings from cell block preparation of pleural effusion, including the immunohistochemical profile. So far, a small number of cases of malignant mesothelioma with mediastinal lesion have been reported. However, to our knowledge, this is the first case of malignant mesothelioma with diffuse swelling of mediastinum lacking of apparent pleural and pericardial lesion.
Pulmonary tumor thrombotic microangiopathy (PTTM) is characterized by widespread fibrocellular intimal proliferation of the small pulmonary arteries and arterioles in patients with metastatic carcinoma. Microscopic pulmonary tumor emboli have frequently occurred in patients with malignant tumors, however, few cases of PTTM have been reported. We report a case of PTTM presenting progressive pulmonary hypertension and lethal respiratory failure.

A nonsmoking 64-year-old woman visited our hospital because of cough, chest pain and dyspnea on effort. High-resolution CT showed subpleural patchy small consolidation with ground-glass opacity. Right ventricular systolic pressure (RVSP) by echocardiography predicted mild pulmonary hypertension (48mmHg). Tc-99m MAA lung perfusion images showed multiple small defects in both lungs. Sputum cytology revealed adenocarcinoma cells, and FDG-positron emission tomography image showed increased uptake at right lower lobe and vertebral body of the Th7 level. A video-assisted lung biopsy was performed, and histology revealed well differentiated adenocarcinoma combined with PTTM, lymphangitis carcinomatosis and necrosis due to the pulmonary infarction. The patient was diagnosed as having PTTM associated with clinical stage IV primary lung adenocarcinoma. In spite of chemotherapy with cisplatin and vinorelbine, respiratory failure combined with pulmonary hypertension was progressed (RVSP=150 mmHg) and she died of respiratory failure. Generally it is difficult to diagnose PTTM in patients during life. This was an important case of PTTM associated with lung cancer and diagnosed antemortem by video-assisted lung biopsy.
Purpose: To estimate pulmonary artery (PA) pressure in subjects with pulmonary hypertension (PH), we retrospectively evaluated the interventricular septal configuration obtained by ECG-gated conventional scanning with 320-slice computed tomography (CT). The configuration of the interventricular septum was compared with systolic PA pressure (sPAP) and mean PA pressure (mPAP) obtained by right heart catheterization (RHC).

Materials and Methods: 38 subjects (10 male, 56±12 yrs) with proven PH (23 chronic thromboembolic pulmonary hypertension and 15 pulmonary arterial hypertension) underwent enhanced 320-slice CT (Aquilion one) and RHC one month apart with no change in clinical status between the examinations. CT images were reconstructed every 5% from 0-95% of the R-R interval and a series of short-axis images was acquired covering the entire left and right ventricles (Zio Station). We compared the leftward shift of the interventricular septum (LSIVS) and septal bowing expressed as curvature (reciprocal of radius in cm; a leftward curvature was denoted as a negative value) at end systole (ES) with both sPAP and mPAP.

Results: sPAP and mPAP from RHC were 70.8±20.0 mmHg and 42.0±11.6 mmHg, respectively. LSIVS at ES was detected in 19 subjects. sPAP and mPAP were significantly higher in subjects with LSIVS (84.3±14.0 vs 57.3±15.3 mmHg [sPAP] (P<0.001), and 49.4±8.0 vs 34.6±9.7 mmHg [mPAP] (P<0.001)). The correlation coefficients of septal curvature with sPAP and mPAP were -0.73 (P<0.001) and -0.74 (P<0.001), respectively.

Conclusions: Septal curvature based on ECG-gated 320-slice CT can be used to obtain an accurate estimate of sPAP and mPAP in subjects with PH.
**Poster Session 9-3**

**Right Ventricle to Left Ventricular Volume Ratio by Electrocardiogram-gated 320 Slice CT is a Predictor of Right Ventricular Pressure Load in Pulmonary Hypertension**

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**Purpose:** To estimate predictors of right ventricular (RV) pressure in subjects with pulmonary hypertension (PH), we measured right (RV) and left ventricular (LV) volumes by ECG-gated 320-slice CT and compared the RV to LV volume ratio with pulmonary arterial pressure (PAP) and pulmonary vascular resistance (PVR) obtained by right heart catheterization (RHC).

**Materials and Methods:** 38 subjects (10 male, 56 ± 12 yrs) with PH (23 chronic thromboembolic pulmonary hypertension and 15 pulmonary arterial hypertension) underwent enhanced retrospective ECG-gated conventional 320-slice CT (Aquilion ONE) and RHC 1 month apart between the two examinations. CT images were reconstructed every 5% from 0-95% of the R-R interval. We quantified RV ES (RVESV) and ED volumes (RVEDV), LV ES (LVESV) and ED volumes (LVEDV), the ratios of RVESV/LVESV and RVEDV/LVEDV. These parameters were compared with the results of RHC.

**Results:** RVESV, RVEDV, LVESV, and LVEDV were 96 ± 47, 164 ± 48, 31 ± 10, and 96 ± 26ml, respectively. In RHC, mean PAP (mPAP), systolic PAP(sPAP) and PVR were 42 ± 12mmHg, 71 ± 20mmHg and 780 ± 342dyne. sec.cm-5, respectively. The correlation coefficients of RVESV/LVESV with mPAP, sPAP and PVR were 0.60 (P<0.001), 0.55 (P<0.001) and 0.69 (P<0.001), respectively. The correlation coefficients of RVESV/LVESV with mPAP, sPAP and PVR were 0.60(P<0.001), 0.55(P<0.001) and 0.69(P<0.001), respectively. The correlation coefficients of RVEDV/LVEDV with mPAP, sPAP and PVR were 0.51(P=0.001), 0.44(P=0.005) and 0.70(P<0.001), respectively.

**Conclusions:** The RV to LV volume ratio by ECG-gated 320-slice CT significantly correlated with PVR acquired by RHC in subjects with PH, and it may estimate mPAP, sPAP and PVR acquired by RHC in these subjects.

**Poster Session 9-4**

**Correlation of Right Ventricular Ejection Fraction with Tricuspid Annular Plane Systolic Excursion by Electrocardiogram-gated 320 Slice CT in Pulmonary Hypertension**

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**Purpose:** There is a strong correlation between right ventricular ejection fraction (RVEF) and tricuspid annular plane systolic fraction (TAPSE) determined by echocardiography in subjects with pulmonary hypertension (PH). However, it is unknown whether there is a correlation between RVEF and TAPSE determined by 320-slice CT. We tested whether TAPSE measured by ECG-gated 320-slice CT correlates with RVEF, and correlates with pulmonary arterial pressure (PAP) and pulmonary vascular resistance (PVR) obtained by right heart catheterization (RHC) in PH subjects.

**Materials and Methods:** 38 subjects (10 male, 56 ± 12 yrs) with PH (23 chronic thromboembolic pulmonary hypertension and 15 pulmonary arterial hypertension) underwent enhanced retrospective ECG-gated conventional 320-slice CT (Aquilion ONE) and RHC, with 1 month between the two examinations. CT images were reconstructed every 5% from 0-95% of the R-R interval and a series of apical 4-chamber images. TAPSE was measured from systolic displacement of the RV freewall and tricuspid annular plane junction (Zio Station). RV end-systolic and end-diastolic true volumes were measured from 3-dimensional reconstruction (Virtual Place Fujin, AZE) and used to calculate RVEF.

**Results:** TAPSE and RVEF were 14.9 ± 3.6 mm and 46.1 ± 12.6%, respectively. In RHC, mean PAP (mPAP) and PVR were 42 ± 12 mmHg and 680 ± 342 dyne sec cm-5, respectively. The correlation coefficient of TAPSE with RVEF was 0.81 (P<0.001). The correlation coefficients of TAPSE with mPAP and PVR were 0.67 (P<0.001) and 0.62 (P<0.001), respectively.

**Conclusions:** TAPSE by ECG-gated 320-slice CT correlated strongly with RVEF and significantly with mPAP and PVR acquired by RHC in subjects with PH.
**Poster Session 9-5**

**Evaluation of Hemodynamic Changes in Pulmonary and Systemic Arterial Circulation in Patients with Pulmonary Fibrosis in Relation to Breathing Methods Using Phase Contrast MRI**

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**PURPOSE:** Blood flow volume of the pulmonary and systemic arterial circulation in patients with pulmonary fibrosis is not well understood. The purpose of this study was to evaluate the hemodynamic changes in relation to breathing methods in patients with pulmonary fibrosis using phase-contrast MRI.

**METHODS:** Eleven patients with pulmonary fibrosis and 15 healthy volunteers were examined by a 1.5T MR apparatus. Phase-contrast flow measurements with free breathing and inspiratory breath-holding were performed in the ascending aorta and pulmonary trunk. Blood flow volume per minute measured from the time-velocity curve was compared. For statistical analysis of each blood flow volume between the patients and the healthy subjects, the Mann-Whitney U test was used. Wilcoxon’s signed-rank test was used to assess significant differences in both the pulmonary and systemic flow volume between free breathing and breath-holding in patients and volunteers.

**RESULTS:** The pulmonary and systemic blood flow in patients with pulmonary fibrosis was significantly reduced compared with those in healthy subjects during both free breathing and inspiratory breath-holding. However, shunt volume between pulmonary and systemic blood flow in patients with pulmonary fibrosis was maintained during both free breathing and inspiratory breath-holding.

**CONCLUSION:** Physiological shunt volume was confirmed to be maintained without influence of breathing methods even in patients with low blood flow circulation affected by pulmonary fibrosis.

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**Poster Session 9-6**

**Alteration of Pulmonary Perfusion after CT Density-Based Non-uniform Attenuation Correction on DIBrH SPECT-CT Fusion Images**

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Alteration of pulmonary perfusion after CT density-based, nonuniform photon attenuation correction (AC) on deep-inspiratory breath-hold (DIBrH) SPECT-CT fusion image was evaluated.

Forty patients with pulmonary emphysema, 32 with pulmonary thromboembolism, 23 with lung cancer and 6 normal controls underwent DIBrH Tc-99m-MAA SPECT, using a dual-head SPECT system and a respiratory tracking device. Scatter-corrected DIBrH SPECT with the triple energy window method was automatically coregistered with DIBrH CT. AC of DIBrH SPECT was performed using an attenuation coefficient map of a variable-effective linear coefficient calculated from CT pixel density of the co-registered DIBrH CT. Alteration of pulmonary perfusion after AC was evaluated by comparison with uncorrected DIBrH SPECT.

Compared with non-AC DIBrH SPECT, the maximum pixel counts in normal lungs were predominantly increased at deep lung portions near the mediastinum and vertebrae and at the upper and middle lungs, regardless of systematic increases (145% ± 28), with significant enhancement of physiological gravitational ventral-dorsal gradient of pulmonary perfusion (P < 0.01). In the affected lungs of all the lung diseases, perfusion defect clarity and/or heterogeneity were significantly enhanced (P < 0.001), without noticeable artifacts. The correlation between perfusion heterogeneity and the lung diffusing capacity for carbon monoxide was significantly improved in patients with emphysema (P < 0.005), and postoperative respiratory function (FEV1) could be better predicted in patients with lung cancer (P < 0.005).

Alteration of pulmonary perfusion after AC on DIBrH SPECT-CT fusion images appears to better reflect physiologic perfusion in normal lungs and perfusion impairment in the lung diseases.
Chronic pulmonary embolism (CPE) leads to pulmonary hypertension that is a progressive disease with a poor prognosis.

**Objectives:** To retrospectively compare Tc-99m MAA lung perfusion SPECT study with multidetector CT pulmonary angiography (CTPA) regarding the number of affected segmental pulmonary arteries in patients with CPE.

**Methods:** We retrospectively reviewed the results of Tc-99m MAA SPECT and CTPA in CPE patients at our hospitals between 2001 and 2008. A total of 50 patients (27 males, 23 females; age range, 32-79 y; mean age 65.0 y) were enrolled. The SPECT and CTPA images were interpreted by three NM physicians. The CTPA criteria of PTE were as follows; pouch defect, abrupt narrowing, obstruction, intimal irregularity, webs and bands. Semi-quantification was done by using a scoring system with a maximal score of 17 segments.

**Results:** Seven (23%) patients had a final diagnosis of chronic PTE hypertension. Only 4 (13%) patients had a complete improvement of acute PTE, though 18 (60%) patients exhibited a partial improvement of PTE. SPECT showed improvement and aggravation of the disease more accurately than CTPA. However, CTPA provided us with morphological information such as PA dilation, RV enlargement.

**Conclusions:** Our results demonstrated that SPECT and CTPA were complementary to each other and Tc-99m MAA SPECT was more accurate than CTPA in monitoring PTE patients. SPECT should be performed as a baseline study at the initial diagnosis of acute PTE.
**Poster Session 10-1**  
**CT Diagnosis of Recurrence of Lung Cancer**

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The purpose of this study was to determine recurrence of lung cancer predictors based on postoperative CT findings. Among 203 patients with primary lung cancer who underwent thoracotomy since January 2003 until December 2004, 61 patients had recurrence, 46 of them were found to have recurrent tumor, LN swelling or pleural dissemination. In 43 of 46 patients, recurrent tumor, LN swelling, pleural nodules or thickening were detected on postoperative CT.

From these results, we conclude that CT is essential for detecting recurrence of postoperative lung cancer.

**Poster Session 10-2**  
**Comparison of Quantitative Diagnostic Capability of Lymph Node Metastasis in Non-small Cell Lung Cancer Patients between STIR Turbo SE Imaging, Diffusion-Weighted Imaging and FDG-PET/CT**

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**Purpose:** To prospectively compare quantitative diagnostic capabilities of lymph node metastasis among short inversion time inversion recovery (STIR) turbo spin-echo imaging, diffusion-weighted imaging (DWI), and FDG-PET/CT in non-small cell lung cancer (NSCLC) patients.

**Material and Methods:** 62 consecutive NSCLC patients underwent STIR-TSE imaging, DWI, integrated FDG-PET/CT and surgical resections. For quantitative differentiation of metastatic from non-metastatic lymph nodes, ratio of signal intensity between lymph node and muscle (LMR) on STIR, apparent diffusion coefficient (ADC) on DWI and SUV on PET/CT at each lymph node were measured by ROI measurement. Then, each index was statistically compared between metastatic and non-metastatic lymph nodes by Student’s t-test. To determine the feasible threshold value of each index, ROC-based-positive test was performed on a per node basis. Finally, sensitivity, specificity and accuracy of each index were compared each other on a per node basis and on a per patient basis by McNemar’s test.

**Results:** LMR and SUV had significant difference between metastatic and non-metastatic lymph nodes. When each feasible threshold value revealed with ROC-positive test was adapted on a per node basis analysis, specificities and accuracies of STIR was significantly higher than those of DWI and PET/CT on a per node basis assessment. When corresponding values were adapted on a per patient analysis, specificity and accuracy of STIR were significantly higher than those of PET/CT.

**Conclusions:** STIR turbo spin-echo imaging is more accurate method for quantitative diagnosis of metastatic lymph node than PET/CT, and would be better to utilized rather than DWI in NSCLC patients.
Objective: To determine the capability of quantitative differentiation between small cell lung cancer (SCLC) and non-SCLC (NSCLC) of diffusion-weighted image (DWI) and short TI inversion recovery turbo spin-echo imaging (STIR).

Materials and Methods: In this study, the pathological diagnoses of all subjects were proved from surgical specimens for purposes of accuracy. Four patients with SCLC (mean age; 66.3 years) and 20 NSCLC patients selected from our database who were matched to SCLC patients in age and in the diameter of cancer were enrolled. All MR examinations were performed on a 1.5 T scanner, and DWI and STIR were acquired. For quantitative assessments of histological classification, apparent diffusion coefficient (ADC) values on DWI and contrast ratios between cancer and muscle (CRs) on STIR were measured. ADC values and CRs were then compared, and diagnostic capabilities were obtained statistically.

Results: There were significant differences between the mean ADC values of SCLC and those of NSCLC (p<0.05); however, there were no significant differences between the mean CRs of SCLC and those of NSCLC (p>0.05). With adopted threshold values, the specificity and accuracy of DWI were 90.0% (18/20) and 87.5% (21/24), respectively. Those diagnostic capabilities were higher than those of STIR (p<0.05). In addition, the combination of both sequences could improve diagnostic accuracy (specificity; 100% [20/20], and accuracy; 95.8% [23/24]).

Conclusion: DWI is a sensitive sequence for the differentiation of SCLC from NSCLC, and the combination of DWI and STIR had a reliable diagnostic indicator of differentiation.

Poster Session 10-4
Comparison of Capability for Pulmonary Nodule Detection ; 1.5T vs 3.0T Systems

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[PURPOSE] To prospectively and directly compare the detection and differentiation capabilities of non-contrast-enhanced (non-CE) pulmonary MR imaging between 1.5 T and 3T systems.

[MATERIALS AND METHOD] A total of 40 patients (20 male, 20 female; mean age 66.4 years) with 70 pulmonary nodules detected on thin-section MDCT underwent non-CE MR imaging on 1.5T and 3T systems. Black-blood T1WI, T2WI and STIR turbo spin-echo imaging were obtained on both systems. The gold standard was determined from the results of thin-section MDCT. The probabilities of nodule presence were assessed by using 5-point scoring system. To determine for nodule detection on appropriate sequences, ROC analyses were performed to determine the feasible threshold values. Overall nodule detection rates of all sequences were compared by using McNemar’s test. For determination of the practical threshold values to differentiate malignant from benign nodules, ROC-based positive tests were performed. Finally, sensitivity, specificity and accuracy were directly compared by McNemar’s test.

[RESULTS] Although ROCs of all sequences had no significant difference for qualitative nodule detection (p>0.05), overall nodule detection rates of T1WI, T2WI and STIR at 1.5 T MR scanner and STIR at 3.0 T MR scanner were significantly lower than the standard reference (p<0.05).

[CONCLUSION] Non-CE MR imaging on 3T system was more useful for detection and diagnosis of pulmonary nodules than that on 1.5T system.
A GGO is a finding on thin-section CT images of the lung that has been described as a hazy area of increased attenuation with preservation of the bronchial and vascular margins. The major teaching points in our presentations are: 1. To demonstrate an optimal segmentation method of localized Ground Glass Opacity (GGO) both pure GGO and mixed GGO. 2. To present the effective quantitative analysis to evaluate the characteristics of GGOs on original developed 2D and 3D approached application software. 3. To understand the limitation of low dose HRCT with simulated control of signal to noise ratio for GGO physically. In this presentation, we suggest the integrated three approaches (clinical, experimental and computational approach) for lung cancer screening.
### Poster Session 10-7

**How Much Can the Radiation Dose be Reduced in Automated Volumetry of Pulmonary Nodules? A Comparison of 16 cm Volume Scans and 64-row Helical Scans with 320-row MDCT.**

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**Purpose:** To determine how much the radiation dose can be reduced in automated volumetry of pulmonary nodules with 16 cm volume scans and 64-row helical scans using a chest phantom with artificial pulmonary nodules.

**Material and Methods:** A lung phantom containing four nodules of different sizes (5 and 10 mm diameters) and densities (solid and ground glass opacity; GGO) with three different nodule categories (intraparenchymal, attached to blood vessels or the pleura) was scanned using both 16 cm volume scans and 64-row helical scans with a 320-row MDCT scanner. Nodules were imaged using varying tube currents and collimations. Data sets were analyzed with a dedicated workstation using volumetry software. Absolute percentage error of estimated volume to known real volume < 20% was defined as eligible.

**Results:** Nodules with 10 mm diameter on thinner section CT were evaluated in detail, as measurement eligibility was high. All solid nodules were measured accurately by both scans, even with tube currents as low as 3.5 and 5 mAs. GGO nodules could be accurately measured using 17.5 mAs with a volume scan. Intraparenchymal nodules and those attached to the pleura were not measured accurately with tube current lower than 35 mAs in a helical scan.

**Conclusions:** Volumetry of 10 mm solid nodule was eligible with a very low tube current when a thin section was applied. For GGO nodules, volumetry with a tube current of 17.5 ms was eligible in a volume scan, which was lower than in a helical scan.

### Poster Session 10-8

**Prediction of Postoperative Lung Function in Patients with Lung Cancer: Comparison of Dual-energy Perfusion CT and Perfusion Scintigraphy**

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**Purpose:** To assess the potential of a dual-energy perfusion CT as a substitute for a pulmonary perfusion scintigraphy in predicting postoperative lung function in patients undergoing lung resection.

**Materials and Methods:** Forty-five patients (M:F = 35:10; mean age, 63.2 years) were prospectively enrolled and they underwent dual-energy perfusion CT, perfusion scintigraphy, and pulmonary function test (PFT) before surgery and 6 months after surgery. Lobectomy in 42, bilobectomy in two, and pneumonectomy in one patient. CT was scanned at 40 second after contrast injection using dual-source CT (Definition, Siemens) with dual-energy technique. On dual-energy CT data, each lobe was segmented using in-house software. Regional perfusion of each lobe was measured by multiplying lobe volume and iodine value per voxel. The proportion of perfusion of each lobe per whole lung was obtained from both perfusion CT and scintigraphy. Predicted postoperative FEV1 (po-FEV1) was calculated by multiplying preoperative FEV1 and (1 - proportion of perfusion of lobe(s) to be resected). Agreements between predicted po-FEV1 and real po-FEV1 were assessed with Bland-Altman plot for both modalities. Precision of prediction of two modalities was assessed by comparing percentage errors of predicted po-FEV1 to real po-FEV1.

**Results:** On Bland-Altman method, the limits of agreement (± 2SD) between real po-FEV1 and predicted po-FEV1 in liter, were -0.88 and 0.85 in scintigraphy and -0.88 and 0.56 in CT. The limits of agreement (± 2SD) between real po-FEV1% predicted and predicted po-FEV1% predicted, were -30.1 and 27.7 in scintigraphy and -30.0 and 18.0 in CT. Percentage errors of FEV1 in liter and FEV1% predicted by perfusion CT were comparable to those by scintigraphy (15.8% and 16.0% by CT; 18.5% and 18.7% by scintigraphy).

**Conclusion:** Dual-energy perfusion CT was comparable to perfusion scintigraphy in the prediction of postoperative lung function.
### Poster Session 11-1

**Quantitative Analysis of Pulmonary Cysts and Emphysema on Computed Tomography: Lymphangioleiomyomatosis, Birt-Hogg-Dubé Syndrome and Chronic Obstructive Lung Disease**

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**Objectives:** To characterize the pulmonary cysts in patients with cystic lung diseases (i.e. Lymphangioleiomyomatosis [LAM] and Birt-Hogg-Dubé syndrome [BHDS]) and emphysema in patients with chronic obstructive lung disease (COPD) by quantification of low attenuation areas (LAAs) on CT and to assess the usefulness of the methods of characterizing fractals of LAAs to analyze the pathophysiology in cystic lung diseases.

**Subjects and Methods:** This study included 52 LAM patients, 18 BHDS patients and 40 COPD patients who had undergone CT scans at Juntendo University Hospital between January 2002 and August 2009. The number and size of discrete continuous low attenuation areas (CLA) on CT were quantitatively assessed by computer software, and then the fractal properties of LAAs were evaluated in these diseases.

**Results:** The characteristics of LAAs differed between these diseases, and the fractal properties of CLA varied with a value of LAA% in LAM and BHDS, but in COPD. By comparing these results, the way of the development of LAAs in these diseases were suggested.

**Conclusion:** The computer-assisted, quantitative analysis of LAAs on CT images is useful to elucidate the pathogenesis of cystic lung diseases including LAM, BHDS, and COPD.

### Poster Session 11-2

**In Vivo Monitoring of Cystic Fibrosis-like Lung Disease in Mice by Volumetric Computed Tomography**

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**Purpose**

The onset and spontaneous progression of cystic fibrosis (CF) lung disease is poorly understood. Aim of our study was to longitudinally characterize βENaC-transgenic (βENaC-TG) mice, an established in vivo model of CF lung disease, by flat-panel volumetric computed tomography (VCT) as a new method to identify initial lesions and monitor spontaneous disease progression as compared to wild-type (WT) littermates.

**Methods and Materials**

Using a VCT scanner prototype (80kV, 50mAs, scan time 19s, spatial resolution 200 µm) 20 βENaC-TG and 23WT mice were examined longitudinally at 10 different time points from the age of 3 to 42 days. Studies were performed in inhalative sedation and free breathing. Blinded for the genotype, VCT images were evaluated with respect to qualitative (airway-obstruction [trachea, bronchi], diffuse infiltrates, atelectasis, air-trapping, parenchyma homogeneity), and quantitative parameters (lung density in Hounsfield units [HU]).

**Results**

The presence of airway-obstruction, diffuse infiltrates, atelectasis, and air-trapping were significant signs for βENaC-TG mice (p-values from <0.05 to <0.001). Obstruction of the trachea occurred exclusively in βENaC-TG mice and proved to be a predictive factor for early mortality. Lung parenchyma showed inhomogeneous texture as well as significantly lower HU in βENaC-TG compared to WT mice at all time-points (p<0.001).

**Conclusion**

We conclude that VCT is a sensitive technique for non-invasive detection of early lesions, and longitudinal intra- and inter-individual monitoring of spontaneous progression of CF-like lung disease in mice. Our data suggest that VCT may provide a useful tool for pre-clinical evaluation of novel treatment strategies for CF and other chronic lung diseases.
Objectives: There are several morphological subtypes of pulmonary emphysema based on their location in the secondary pulmonary lobule. Although morphological features of paraseptal emphysema usually involve abnormalities in the periphery of the pulmonary lobule, typical findings on CT images show them only adjacent to the pleura including along the interlobar fissures. The objective of the present study was to examine paraseptal emphysema in the inner lung field.

Materials and Methods: Emphysematous lesions and anthracosis adjacent to the interlobular septa or large vessels and bronchi and their predominant areas within lung parenchyma were examined in the inflated-fixed lung specimens.

Results: Lesions adjacent to interlobular septa were seen in places ranging from continuous to pleural surfaces to the inner zone of the lung and mild to severe in degree. They were also seen in the areas adjacent to large vessels and bronchi. The lesions existed not only in one pulmonary lobule but also in both lobules adjoining each other. In cases of co-existence of centrilobular and paraseptal types, the former type was more predominant than the latter. However, only few cases were observed where both types were present equally. Similar to centrilobular type, paraseptal emphysema lesions were seen in the upper lobe as well.

Conclusions: The present study suggests that in routine clinical examination, HRCT could depict paraseptal emphysema adjacent to interlobular septa in the inner zone of the lung in some cases. For understanding of pulmonary function, it is important to distinguish emphysema types of different origins.
**Poster Session 11-6**

**The Epithelial-Mesenchymal Transition of Progenitor Cells Lead to Low Attenuation Areas of the Pulmonary Parenchyma: An Autopoietic Model Analysis of Alveolar Epithelial Maintenance**

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**BACKGROUND AND PURPOSE:** A group of progenitor cells maintains alveolar epithelial integrity in the pulmonary parenchyma. The existence of progenitors which have the ability to differentiate into the alveolar epithelium and the fibroblast (EMT) may be important in the genesis of pulmonary fibrosis. However, no reports refer to the role of EMT in case of pulmonary emphysema. In this paper, we obtained a set of equations to describe the role of EMT in the genesis of pulmonary emphysema.

**METHOD:** An autopoietic system is a network of processes that produces the components that produce those processes. We defined a network of bronchiole-alveolar surfaces in the pulmonary lobule as an autopoietic system by using Bourgine and Stewart’s mathematical theory (Artificial Life 10:327,2004).

**RESULTS:** We obtained two equations for emphysema: $\frac{dCM}{dt} = kC CM + k_F(C1 -CM)Pr(CM)$, and $LAAR\% = \frac{\{a_1[a] (S/V)d/dt[a\ b]\}/kb[a]}{kb[b]}$. CM was the number of alveolar type I cells (C). $a$ and $b$ were densities of stem cells (A) and of progenitor cells (B), respectively. $S/V$ was the density of the alveolar capillary surfaces. $ka$, $kb$ and $kc$ were kinetic parameters of the transition from A to B, from B to C, and from C to apoptosis, respectively. $Pr$ was the engrafting ratio of A. Fibroblasts (F) catalyzed A to B. The process of B F was EMT, which could produce a stable attractor of CM=0 in the phase space.

**CONCLUSION:** Loss of the progenitor cells by EMT would induce emphysematous loss of the bronchiole-alveolar surfaces.
Oscillatory Mechanics within Respiratory Cycles Related to COPD Severity

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Oscillatory properties of respiratory system vary considerably within respiratory cycle (RC) in patients with COPD. As seen in the representative features of airway narrowing at tidal expiration, the oscillatory flow resistance tends to be higher at expiration than inspiration. The relationship between the RC dependency and COPD severity is not fully investigated. We examined the relationship by means of our original 3-D color images.

Oscillatory properties were assessed in patients with COPD (n=39, mean 70-yrs) using Impulse Oscillometry (MS-IOS; Cardinal Health, USA). Impulse Oscillometry was performed in 40 seconds under tidal breathing at sitting position. The oscillatory pressure and flow relationships, resistance (Rrs) and reactance (Xrs), were calculated every 0.2 second between 5 and 35 Hz. All data were evenly divided into 4 sections at inspiration and expiration respectively, and were averaged on 8 respiratory phases. Rrs and Xrs were lined up and assigned color gradients, which resulted in colored 3-D graphs. COPD severity was estimated by %FEV1.

In patients with severe COPD, the main characteristics at lower frequency were the increases in Rrs and the shifts of Xrs curves toward negative on whole respiratory phases, especially during expiration. Those RC dependencies were more remarkably with decrease in %FEV1. The variations of Rrs and Xrs within respiratory phases were not necessarily similar. These physiological features were visualized on 3-D graphs.

The oscillatory flow resistance can be high on whole respiratory phases and have RC dependency with COPD severity. Our 3-D color images can be an effective tool to understand those mechanics.
**Poster Session 12-1**

**Hyperpolarized 3He MRI Apparent Diffusion Coefficients as a Probe of Airway Function in Asthma After Methacholine Challenge and Recovery**

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We measured hyperpolarized 3He Apparent Diffusion Coefficients (ADC) to quantitatively evaluate the asthmatic lung before, immediately after methacholine challenge and after recovery from methacholine challenge, 30 minutes post bronchodilatation with salbutamol. We hypothesized that functional imaging measurements such as 3He MRI ADC that are reflective of airway and airspace microenvironments would be altered after induction of bronchoconstriction in a methacholine challenge. Hyperpolarized 3He magnetic resonance imaging (MRI) was performed at 3.0 T to acquire diffusion weighted images (b=1.6 s/cm², 14-16 slices at 30mm slice thickness) in the coronal plane using a fast gradient-recalled echo (FGRE) method with centric k-space sampling. Plethysmography and spirometry were also performed with MRI on subjects with a clinical diagnosis of asthma (n=13, age = 32.3 ± 9.2y) three times within an hour (just prior to methacholine challenge, 5 minutes after achieving PC20 and finally 30 minutes after bronchodilator administration following PC20 dose and MRI). A two-way ANOVA showed significant interactions between mean whole lung ADC and patient/timepoint with significant differences between all patients post-methacholine and after salbutamol recovery. A region of interest (ROI) analysis identified that these significant differences were specific to the posterior slices and posterior ROI (p < 0.0001). The increased ADC post-methacholine with significant reversal after salbutamol administration suggests that 3He MRI ADC sensitively reflects the presence of gas-trapping in asthma that is relieved after administration of salbutamol.

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**Poster Session 12-2**

**4-Dimensional Imaging of the Central Airway Using a Low Dose Technique: Initial Experience with 320-row Multidetector CT**

Ryo Sakamoto, Satoshi Noma², Takeshi Kubo¹, Hiroyuki Sekiguchi¹, Shigeaki Umeoka¹, Takanori Higashino², Yuko Nishimoto², Yutaka Emoto³, Kaori Togashi¹

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BACKGROUND: 320-row multidetector CT (MDCT) provides not only volumetric image acquisition capability but also wide cranio-caudal coverage of 160 mm without table movement. Therefore, it has become possible to obtain continuous volumetric images of central airway from trachea to proximal bronchi during a respiratory maneuver, at a cost of higher radiation exposure.

MATERIALS AND METHODS: Five adult patients with suspected central airway disorders were scanned during an exhalation. Scanning area was set to cover intrathoracic trachea to proximal lower lobe bronchi. Low dose technique was used while performing a continuous image acquisition of the central airway until full expiration. Consecutive volumetric images were reconstructed at 0.5 second intervals, thus producing 4-dimensional image data of the central airway. In each frame, images were reconstructed in the same plane containing a particular bronchus of interest. The reconstructed images were presented in a cine mode.

RESULTS: With a low dose technique, radiation exposure during 4-D acquisition was reduced nearly to the level of routine whole lung helical CT scan. Production of cine images was possible with acceptable image noise and motion artifacts which allow us to evaluate real-time airway morphology. Cine imaging demonstrated the dynamic deformation of airways, representing pathological changes of each clinical condition.

CONCLUSION: Cine CT using 320-row MDCT with low dose technique provides continuous evaluation of airway motion during respiratory maneuvers. Application of this new diagnostic method to various respiratory diseases is expected to disclose the dynamic change in central airway, thus clarifying pathophysiological mechanism of these diseases.
Poster Session 12-3
Dynamic Property of Central Airway Walls as Assessed by Computed Tomography: Correlation with Asthma Pathophysiology

Masafumi Yamaguchi, Akio Niimi, Hisako Matsumoto, Isao Ito, Toyohiro Hirai, Yasutaka Nakano, Michiaki Mishima

Background: A subset of asthmatic patients suffers from frequent exacerbations despite continuous medication, possibly due to various features of remodeling and associated elastic properties of airway walls. Objective: To examine the fragility of central airways and sputum biomarkers of airway remodeling in different classes of asthmatics, classified according to disease stability.

Methods: We studied moderate-to-severe asthmatics with ≤1 asthma worsening during the previous year requiring beta 2-agonist therapy (stable, n=18), those with ≥2 such episodes (unstable, n=11), and 11 healthy controls. Wall thickness and luminal area of a segmental bronchus were measured by computed tomography at full-inspiration. Percentage decrease in luminal area from full-inspiration to full-expiration was calculated as a measure of airway fragility. Sputum biomarkers associated with fibrosis [TGF-beta 1 and matrix metalloproteinase (MMP)-9/tissue inhibitors of metalloproteinase (TIMP)-1 molar ratio] and those associated with angiogenesis/edema [vascular endothelial growth factor (VEGF) and vascular permeability index (sputum/serum ratio of albumin levels)] were examined.

Results: The airway wall was similarly thickened in both asthmatic subgroups as compared with controls. Airway fragility was greater in unstable asthmatics than in stable asthmatics and controls but was similar in the latter two groups. Sputum TGF-beta 1 levels were higher and MMP-9/TIMP-1 molar ratios were lower in stable asthmatics than in controls. Sputum VEGF levels and vascular permeability index were higher in unstable asthmatics than in controls.

Conclusion: Fragility of thickened airway walls, which in turn may depend on whether fibrosis or angiogenesis/edema predominates in the airways, may be associated with stability of asthma.

Poster Session 12-4
Comparison of 3-D-Visualized Respiratory Impedance Patterns and Structural Change Detected by CT in Smokers

Haruko Shinke, Masatsugu Yamamoto, Nobuko Hazeki, Kazuyuki Kobayashi, Yasuhiro Funada, Yoshihiko Kotani, Yoshiharu Ohno, Yoshihiro Nishimura

Background: We previously reported the influence of smoking on respiratory impedance using MostGraph-01 (CHEST, Tokyo), which enables us to evaluate the variance of respiratory resistance and reactance as 3-D graph along time axis. Further studies are needed to clarify the relations hip between structural change and impedance change in respiratory system. To examine that, we analyzed the structural changes using CT scan to compare estimated parameters from CT images and 3D image pattern of respiratory impedance.

Methods: We examined total of 15 smokers who visited the out-patient clinic of respiratory medicine in Kobe University hospital. The subjects were measured pulmonary function test (PFT). Low attenuation area of the lung (LAA) and airway parameters were measured using CT scan. The respiratory impedance patterns were measured under spontaneous tidal breathing by impulse signal using MostGraph-01. The patterns on 3D graph were categorized to compare with other clinical parameters.

Results: Resistance at 5Hz (R5) was 0.27 ± 0.11kPa s/L, Reactance at 5Hz (X5) was -0.726 ± 0.64 kPa s/L, Mean LAA% was 8.056 ± 8.938%. Some smokers with normal PFT showing frequency dependency of resistance or negative shift of reactance showed increase of %LAA as COPD patients. The impedance variances were seen dominantly in expiratory phase.

Conclusion: The variance of respiratory impedance reflects early change of pulmonary airway disorder induced by smoking. Using this method may provide easy screening strategy to detect occult structural changes and flow limitation in smokers. Now we are accumulating clinical data.
Poster Session 12-5

Usefulness of SD-101, a Non-restrictive Device Developed for Screening Sleep Apnea-hypopnea Syndrome

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Background and objective: Kentzmedico in Japan has developed the SD-101 as a nonrestrictive, sheet-like medical device with an array of pressure sensors, to automatically detect sleep-disordered breathing (SDB) by sensing gravitational alterations in the body corresponding to respiratory movements. The purpose of this study was to examine the clinical usefulness of the SD-101 for screening sleep apnea-hypopnea syndrome (SAHS).

Methods: Nocturnal polysomnography (PSG) and SD-101 recordings were run simultaneously and compared for 201 subjects attending our hospital with suspected SAHS (suspected SAHS group) and in all 165 male employees of a transport company (screening group).

Results: PSG revealed apnea-hypopnea index (AHI) < 5, 5 < AHI < 15, 15 < AHI < 30, 30 < AHI < 60, and AHI ≥ 60 events/h in 39, 35, 38, 68 and 21 subjects in the suspected SAHS group and 103, 34, 12, 12 and 4 subjects in the screening group, respectively. Central SAHS and obstructive SAHS were subsequently diagnosed in 11 (5.5%) and 135 (67.2%) subjects in the suspected SAHS group and 5 (3.0%) and 39 (23.6%) subjects in the screening group, respectively. Significant correlations were apparent between AHI and respiratory disturbance index (RDI) measured with the SD-101 in both the suspected SAHS group (r = 0.88) and screening group (r = 0.92). Receiver operating characteristic analysis revealed 89.5% sensitivity and 85.8% specificity in identifying SAHS, using an RDI of 14.0 events/h.

Conclusion: These findings suggest that the SD-101 is a useful device for screening SAHS.

Poster Session 12-6

Vibration Response Imaging (VRI) in the Detection of Asynchrony between Left and Right Lung for Main Bronchial Stenosis

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INTRODUCTION:

The VRI is an imaging modality that records energy generated by the vibrations of the lungs during respiration cycle and creates an image that can be viewed dynamically or sequentially. The timing of inspiratory and expiratory vibration energy peaks (VEPs) of right versus left lungs can be analyzed. We defined the average of absolute value of the gaps between VEPs of both lungs during 12 sec as VEPgap, and observed left and right lung asynchrony in patients with main-stem bronchial stenosis.

PURPOSE:

In this study we planed to elucidate the threshold level of stenosis that cause left and right lung synchrony using a lung model, then validate this threshold level in patients with main-stem bronchial stenosis.

METHODS:

We created a lung model using a pig lung in a sealed box connected to a ventilator. We measured airway pressure of right and left bronchus at the same time using two catheters to manually narrow the airways. We validated the relation between rate of stenosis and time delay. The stenosis rate was measured by computed tomography using ZAIO soft and VEPgap in 10 patients with unilateral main-stem bronchial stenosis.

RESULTS:

Time delay increased sharply from 90% in the pig model (P<0.05). As in the Pig model, VEPgap increased sharply from 90% in the patients with unilateral main-stem bronchial stenosis.

CONCLUSIONS:

In this study, the VRI was able to identify severe stenoses above 90% in patients. We found VRI was useful in assessing treatment outcomes and decisions on interventional indication.
**Poster Session 13-1**

**Analysis of Lung Motion Due to Respiration and Its Application to PET Images**

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2. St. Hill Hospital, Japan

Lung motion due to respiration causes image degradation in medical imaging, especially in nuclear medicine which requires long acquisition time. We have developed a method for image correction between the respiration-gated (RG) PET images in different respiration phases. The method offers the motion field vector of lung from the PET image set and the motion field vector plays a role for image summation of the RG PET images. This technique is applied to a set of breath-hold (BH) PET images in this paper. In the BH method, a patient is asked to hold his/her breath for 10 to 30 s as the image acquisition is performed. This is repeated several times and the obtained images are summed. In practice, however, a patient cannot necessarily hold his/her breath at the same timing of breathing. In such a case, the summed images still has a blur. We propose to apply our image registration method developed before for RG images to BH images. The BH images are nonlinearly motion-corrected so as to match a reference BH image and summed. Through the experiment with nine patient BH image data, we confirmed that the proposed method is effective in obtaining a non-blurred image.

**Poster Session 13-2**

**A 4D Model Generator of the Human Lung: Lung CataChiCaraCure-er, Alias Lung4Cer**

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Division of Engineering Technology, JSOL Corporation, Japan

We have developed a free software which generates various kinds of 4D lung models for the purpose of studying structure and function of the human broncho-alveolar system. The name of "CataChiCaraCuri-er" is arisen from Japanese words, CataChi (= form, structure) and CaraCuri (= machine, mechanism). The last "er" indicates a doer in English and its pronunciation also means a doer in Japanese (Ya). The alias of "4Cer" means a 4D model generator like a farmer who grows vegetables rapidly. Lung4Cer makes dynamic air pathway models from the trachea to alveoli based on four algorithms published previously. Breathing motion is given by parameters which assign respiratory modes and body postures. Lung4Cer contains a newly developed algorithm which converts geometric models into 4D finite element models which allow airflow simulation by means of computational fluid mechanics. Visualization of generated models is expected to use a free popular software "ParaView", which can be easily obtained by internet. Potentially, Lung4Cer can make a whole lung model with 300 million alveoli. However, it costs incredibly huge computer resources. Instead, suitable model types can be selected according to user’s interest and available computer resource, such as (1) airway tree only, (2) airway tree with air-supplied parenchymal regions, (3) air pathway from the trachea to a subacinus with alveolar structure, and (4) alveolar system only. Lung4Cer also contains Origami models for alveolar ducts nearly equivalent to the computer models so that users may handle them in reality. Lung4Cer will be a powerful basic tool for studying pulmonary functional imaging.
Poster Session 13-3

Alveolar Recruitment Analysis in Acute Respiratory Distress Syndrome (ARDS) Using End-inspiratory and End-expiratory Breath-hold 3D-CT Images

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3. Department of Radiology, Osaka University Graduate School of Medicine, Japan

Purpose: Open lung strategy for ARDS is an artificial ventilation which intends to increase persistent alveolar recruitment (= recruited through respiratory cycle) and decrease cyclic recruitment (= recruited at end-inspiration but collapsed at end-expiration) without hyperinflation of intact alveoli. We have developed a quantitative image analysis technique for detecting recruited alveolar regions in 3D CT images by the use of nonrigid image registration between expiratory and inspiratory CT images for the purpose of estimating the optimal artificial ventilation mode.

Methods: Five patients of ARDS under artificial ventilation were subjected (approved by the in-hospital ethics committee). A set of iBH-CT and eBH-CT images was obtained under the same settings of mechanical ventilation in ICU. Another set was obtained ten minutes after increasing PEEP by 5cmH2O. Two sets of eBH-CT and iBH-CT at respective PEEP levels and a set of eBH-CTs at different PEEP levels were subjected to nonrigid image registration by which CT value difference between corresponding points in iBH-CT and eBH-CT were computed. The last set was to assess the direct effect of PEEP level. A voxel cluster whose CT values changed more than 400 HU was defined as a recruited region.

Results: Recruitments were detected at dependent zones in all cases. FiO2 deference between different PEEP levels was in strong correlations to volumes of persistent recruitment and volume reduction of cyclic recruitment.

Conclusions: Our method can evaluate alveolar recruitments both visually and quantitatively. It is expected to assist open lung strategy for ARDS.

Poster Session 13-4

Velocity Vector Imaging in Assessing the Regional Diaphragmatic Movement of Patients with Chronic Diaphragmatic Hernia: Preliminary Results

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Purpose:
To evaluate the regional diaphragmatic movement in patients with chronic diaphragmatic hernia using the velocity vector imaging (VVI) technique, a two-dimensional angle-independent ultrasound speckle tracking method.

Material and Methods:
Two-dimensional cine-loop ultrasound images of right lateral intercostal view were obtained in 5 patients with chronic right diaphragmatic hernia and 3 healthy volunteers under quiet respiration. The regional movements of right hemidiaphragm were analyzed with VVI offline software. The diaphragmatic contours were manually identified in a single frame of a cine-loop and the contours in other frames were automatically generated with the ability for the operator to alter any of those contours. After performing calculation process, the velocity vectors superimposed on two-dimensional cine-loop ultrasound images to describe the pattern of diaphragmatic movement.

Results:
Of the 5 cases with chronic right diaphragmatic hernia, 3 (60%) cases were successfully analyzed with VVI software, while another 2 cases were excluded due to poor acoustic window. Among the 3 successfully analyzed cases, 2 cases showed dysynchronous movement with opposite direction in both inspiratory and expiratory phases at the free edge of diaphragm close to the defect of the hernia. The other case with hernia showed synchronous movement during whole respiration phases that were typically noted in the healthy volunteers.

Conclusion:
VVI could recognize and quantify the regional diaphragmatic movement in cases with adequate acoustic windows. We demonstrate the regional dysynchronous movement of diaphragm in two cases of chronic diaphragmatic hernia. This is an interesting phenomenon and has not well documented in the previous lecture.
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<td><strong>Usefulness of Ultrasonography in Preoperative Diagnosis of Solitary Fibrous Tumor of the Pleura</strong></td>
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Mitsuki Sekiya, Kaku Yoshimi¹, Keiko Muraki¹, Noriyuki Homma¹, Teruhiko Sato¹, Takashi Dambara², Kenji Suzuki², Toshimasa Uekusa³, Kazuhsa Takahashi⁴  
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⁴. Department of Pathology, Kanto Rosai Hospital, Japan

**Background:** Solitary fibrous tumor of the pleura (SFTP) is a rare pleural tumor of mesenchymal origin, accounting for 5% to 10% of primary pleural tumors. Although usefulness of thoracic CT and MRI in the evaluation of SFTP has been well documented, ultrasonographic evaluation has not been reported. We reviewed the ultrasonographic features of three pathologically proven cases of SFTP.

**Methods and results:** Patients consisted of two men and one female (age 55±19 years old). Thoracic CT demonstrated a well-defined, pleural-based lesion that has obtuse angles and tapering margin with adjacent pleura. The patients underwent thoracic ultrasonography (US) using the commercially available ultrasonic unit (SSA-550A; Toshiba, Tokyo, Japan) with the convex type linear array probe (PLM-805AT 8MHz, Toshiba, Tokyo, Japan). Thoracic US demonstrated solitary, homogeneous, hypo-echoic, hemicycle lesions. All lesions abut on the hemicycler line originating from visceral pleura of the adjacent lung. However, they showed respiratory motion together with adjacent lung. On the basis of the above findings, pedunculated tumors arising from the visceral pleura, especially SFTP, were suspected. We performed surgical resection and confirmed pedunculated tumors of visceral pleura. Then, pathological diagnosis of SFTP was established in all cases.

**Conclusion:** Preoperative thoracic US could be useful in the evaluation of the patients with SFTP.

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<td><strong>A Method for Reduction of Subtraction Artifacts on Temporal Subtraction Images by Use of Generalized Gradient Vector Flow Technique</strong></td>
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Hyoungseop Kim¹, Noriaki Miyake¹, Joo Kooi Tan¹, Seiji Ishikawa¹, Seiichi Murakami¹², Takatoshi Aoki²  
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**Introduction:** A temporal subtraction (TS) image, which is obtained by subtraction of a previous image from a current one, can be used for enhancing interval changes on images by removing most of the normal structures. In this paper, we propose a registration method for TS using GGVF.

**Methods:** In the first stage, the vector fields are described using GGVF from the previous and current images, respectively. Next, some VOIs are set up on the CT image pairs. Furthermore, the shift vectors are calculated using nearest neighbor matching of the vector fields in these VOIs. The previous CT voxel which resemble standard the current voxel is detected by voxel value and vector of the GGVF in the kernel. Finally, a TS image is made by subtraction of a warping image from a current one.

**Results:** We employed 6 databases obtained from different time-series on same patients and evaluate the effect of reducing the subtraction artifacts based on histogram and satisfactory results are obtained. In the experimental results, there was a trend toward a decrease in contrast value on the TS image.

**Conclusions:** In this paper, we have developed a new image registration method to reduce subtraction artifacts in TS images. Subtraction artifacts on temporal subtraction image were substantially removed. We believe that the TS images can be introduced as a useful tool to analyze an interval changes.

**Acknowledgement:** This work was supported in part by a research grants from Kitakyushu Foundation for the Advancement of Industry Science and Technology.
Cancer is the leading cause of death with lung cancer as the number one cause. Since the survival rates of those with lung cancer are low, earlier detection and early treatment are important. In addition, chronic obstructive pulmonary disease (COPD) appears to be in the top ten cause of death in 2000, and more than 5.3 million Japanese older than 40 years old have this disease. Specifically, treatment for pulmonary emphysema is difficult, therefore early detection of the disease at an early stage is important. The patients of osteoporosis comprised of about 10 million people in Japan and it is one of the problems the aging society has. We developed clinical chest CAD system for lung cancer, COPD, and osteoporosis based on MDCT images.

Clinical chest CAD system could quantitatively detect lung cancers of many size, type, and texture variations. In addition, our CAD system could detect lung cancers, COPD, and osteoporosis at early stage.

Acknowledgments: The work was supported by the Competence Network Asthma/COPD funded by the German Federal Ministry of Education and Research (FKZ 01GI0881-0888).
Poster Session 14-3

Developments of Thrombosis Detection Algorithm using the Contrast Enhanced CT Images

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In the diagnosis of thrombosis with no specific clinic symptoms, diagnostic imaging plays a greater role. Particularly, contrast Enhanced CT is low invasive diagnostics, and the thrombus in the pulmonary artery can be detected as a low density without the contrast effect. Moreover, because describing the change of concentration in lung field and the decline in lung blood vessel shadow is also possible, it is indispensable to diagnose of thrombosis. As the image diagnosis support, it is necessary to classify the pulmonary artery and vein that relate to the thrombosis, and to analyze the lung blood vessel quantitatively. The technique for detecting the thrombosis by detecting the position of the thrombus has been proposed so far. In this study, it aims to focusing on the dilation of the main pulmonary artery and to detect the thrombosis. The effectiveness of the method is shown by measuring the pulmonary trunk diameter by using the extracted pulmonary artery from contrast Enhanced CT through semi-automated method, and comparing it with a normal case.

Poster Session 14-4

Segmentation of Thoracic Organs from Multi-slice CT Images

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With the development of multi-slice CT technology, obtaining an accurate 3D image of lung field in a short time becomes possible. To support that, a lot of image processing methods need to be developed. To diagnose lung cancer in the clinical field, it is important to study and analyse lung structure. Therefore, classification of lung lobe and classification of lung segment provide useful information for lung cancer analysis. In this report, we describe algorithms which segment thoracic organ, classify lungs into lung lobes and lung segments from multi-slice CT images. The classification algorithm of lung lobes is efficiently carried out using information of lung blood vessel, bronchus, and interlobar fissure. On the other hand, the classification algorithm of lung segments is carried out using information of bronchus and pulmonary artery.
**Poster Session 14-5**

**Iterative Algorithm for Airway Segmentation in CT Images Using a Region-based Active Contour Model**

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**Purpose:** To develop and evaluate a new algorithm for airway segmentation in CT images using a region-based active contour model.

**Materials and methods:** Being based on level sets, the region-based active contour model requires no intrinsic parameter concerning gray levels. The new algorithm gradually executes airway segmentation through the iterative use of the active contour model. Initially, a central trunk of airway leading from the right (or left) main bronchus to the ipsilateral lower lobe bronchus is extracted. In this trunk, regions near the origins of its branches are identified. On the other hand, a FIFO queue of seed regions is set up and initialized with these regions. Subsequently, a seed region retrieved from the queue is elongated distally along airways according to the evolution of the active contour model; this elongation is stopped when new candidate seed regions are identified. Each candidate seed region, if not rejected as a leak outside airways by a classifier, is then put into the queue. The above process is repeated until the queue becomes empty. Using 15 thin-section standard-close CT datasets, the results of airway segmentation obtained with this algorithm were compared with those obtained with a benchmark (recently-proposed centricity-based region growing algorithm).

**Results:** The number of extracted airway branches was $103 \pm 19$ and $65 \pm 13$ when using the new algorithm and the centricity-based algorithm, respectively. The difference between these numbers was significant ($P<0.01$, paired t-test).

**Conclusion:** The new algorithm for airway segmentation in CT images was superior to the benchmark algorithm.

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**Poster Session 14-6**

**A Comparative Study of HRCT Image Metrics and PFT Values for Characterization of ILD and COPD**

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Several image metrics have been proposed in the literature for pulmonary assessment via thoracic high-resolution computed tomography for various diseases. This article describes a systematic analysis of the utility of such metrics for characterizing interstitial lung disease and chronic obstructive pulmonary disease in comparison to data from pulmonary function testing. 14 patients with ILD and 11 with COPD were retrospectively identified who had undergone both HRCT and PFT within 3 days of each other. Using a custom-made computational pipeline, 93 different image-based features were computed for each patient.

The relationships between these image metrics and the 21 clinically-used PFT parameters were analyzed using a minimal-redundancy-maximal-relevance statistical framework. The results showed that not only are some image metrics good discriminators for characterization of ILD and COPD compared with PFT, but also that image metrics are not redundant for differentiation between ILD and COPD groups when PFT values are provided. Image metrics of the attenuation histogram statistics and of more sophisticated texture descriptions are both selected, which suggests that both types of image metrics may be valuable for further investigation in computer assisted diagnosis.
Poster Session 15-1

Unexplained Desaturation in the Sitting Position with Stroke

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A 87 year old woman admitted to our hospital with speech disturbance was found to have unexplained desaturation in the sitting position. Her brain MRI showed multiple infarctions in bilateral cerebellum, left occipital lobe, and right thalamus. Oxygen saturation at O2 4L/min fell precipitously from 93% in the supine position to 70% in the sitting position. Chest enhanced CT revealed no findings suggestive of thromboembolism. Micro bubble test in the sitting position was positive, and pulmonary perfusion scintigraphy suggested 23% right-to-left shunt. A transesophageal echocardiogram was taken in the supine and sitting positions. The images taken in the supine position showed patent foramen ovale with atrial septal aneurysm, and mainly left-to-right shunt. However, the images taken in the sitting position showed further opening of foramen ovale with massive right-to-left shunt across the defect. We diagnosed her with platypnea-orthodeoxia syndrome, which was characterized by dyspnea and hypoxemia induced by orthostatism and relieved by supine position. Multiple infarctions were thought to be caused by paradoxical embolism due to this shunting. Anatomical and functional components are necessary to cause platypnea-orthodeoxia. In this case, foramen ovale was open anatomically, and a dilated aortic root functionally changed the blood flow to right-to-left shunt through the foramen in the sitting position.

Poster Session 15-2

Acute-onset Sarcoidosis with Polyarthralgia and Hilar Lymphadenopathy

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A 33 years old woman was admitted because of high grade of fever, polyarthralgia and coughing. Chest Computed tomography showed multiple lymphadenopathy of the mediastinum, hila and left collarbone fossa, and also showed nodele with the cavity in the left S6 domain. Bronchial alveolar lavage showed a slightly high number of lymphocyte and a high CD4/CD8 cell ratio. Although transbronchial lung biopsy of the lung nodule did not show noncaseating granuloma. Gallium scintigraphy showed Accumulation to hilar lymph nodes, eye sockets, and paroilds. Video-assisted thoracic surgery and lymph node biomicroscopy were performed for diagnosis. In the pathology organization, both lymph node and lungs organization accepted a noncaseating granuloma. Fever up and the arthralgia were improved from preoperation, but those symptoms comes out again from postoperative the sixth day. Gallium scintigraphy showed accumulation to the same part as the last time and an and both knees, an ankle. We diagnosed as an Löfgren syndrome without the erythema nodosum. Oral administration of prednisolone(20mg/ day) was initiated. A symptom was improved afterward and became the discharge. The Löfgren syndrome is one model of the acute sarcoidosis to assume 3 the main sign, erythema nodosum, polyarthralgia, and bilateral hilar lymphadenopathy. It is reported that a ratio with the erythema nodosum is high with a woman in comparison with the man. We add it and report consideration from literatures about relations with this case.
Helminthic infections are relatively rare in respiratory medicine, but we must keep in mind the possibility of parasitic disorders when there are pulmonary infiltrations or distinctive abnormal structures whose diagnosis is not obvious. Here, we report a 65-year-old male with toxocariasis who presented with unusual tubular structures in both lung fields in CT images. His chief complaint was transient hemosputa, but no fever or eruptions were noted. Chest plain radiograph showed ground glass opacity in his right lower lung. Axial and multiplanar reconstructed CT images showed irregularly curving tubular structures, approximately ~10 cm long, that passed along the subpleural area of bilateral lower lobes beyond the segments. The tubular structures connected to lower lobe bronchioles and consolidations on the diaphragm. Based on these radiographic findings, we speculated that the distinctive images might be due to migration scars of some parasite. Although there was no blood eosinophilia or elevation of serum IgE levels, IgG ELISA demonstrated the presence of serum antibodies against Toxocara canis ES antigen. Raw foods such as beef liver were suspected sources. Three months after 4-week administration of albendazole, the specific IgG level declined, indicating that the toxocariasis became inactive. Although the radiographic findings do not necessarily equate with toxocariasis and may be evidence of some other previously migrated parasites, the observed patterns are strongly suggestive of scars of helminth migration. We believe that curved multiplanar reconstruction technique of CT scan is a powerful tool for clear depiction of the full-length scars left by migrated parasites.