Interdisciplinary Technical Journal Club:
Special series on Laboratory Animal Science recognized by the Veterinary Office of the Canton of Zurich

*In vivo imaging*

- In accordance with 3R policies

02.08.2016

Regina R. Reimann

Institute of Neuropathology
The Three R’s

Small-animal models a bridge between basic science and clinic application

Human animal research (Russell, 1957):

Methods which avoid or replace the use of animals  
**Replacement**

Methods which minimise the number of animals used per experiment  
**Reduction**

Methods which minimise suffering and improve animal welfare  
**Refinement**

https://www.nc3rs.org.uk
Reduction: Longitudinal *in vivo* studies

- Reduction of variance: each mouse is its own control
- Monitoring of disease progression in the individual mouse
- Enabling a systematic collection of tissue (in further studies): Time course and the anatomic hot-spots of the disease (model) are known
Refinement by the use of in vivo imaging

‘It constitutes a way of assessing biological structures and function in vivo by non invasive means, allowing the collection of quantitative information, both in health and disease states’ (Zanzonico, 2011)

• Acquisition of an impressive amount of unique often multi-modular information (without interfering with the biological process under study)
• Real time studying of disease in a quantitative way
• Macroscopic level and molecular level
• Repeatedly and non-invasively monitor disease progression or response to treatment
• Translational aspect : nearly identical settings than used in clinic
In vivo imaging: main modalities

Table 1 Summary of general properties of diagnostic imaging modalities

<table>
<thead>
<tr>
<th>Imaging modality</th>
<th>Physical basis</th>
<th>Information supplied</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positron emission tomography (PET)</td>
<td>Gamma-radiation (derived from positron emission)</td>
<td>Tracer uptake</td>
</tr>
<tr>
<td>Single photon emission computed tomography (SPECT)</td>
<td>Gamma-radiation</td>
<td>Tracer uptake</td>
</tr>
<tr>
<td>Optical imaging (OI)</td>
<td>Light emissions (ex: fluorescence)</td>
<td>Probe uptake</td>
</tr>
<tr>
<td>Computed tomography (CT)</td>
<td>X-rays</td>
<td>Tissue density</td>
</tr>
<tr>
<td>Magnetic resonance imaging (MRI)</td>
<td>Magnetic properties</td>
<td>Tissue composition</td>
</tr>
<tr>
<td>Ultrasound (US)</td>
<td>Sound reflection of high-frequency sound waves</td>
<td>Internal movements and flows, differences of tissues</td>
</tr>
</tbody>
</table>

Electromagnetic radiation:

<table>
<thead>
<tr>
<th>Radiation / Wavelength</th>
<th>Modality</th>
<th>Medical information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radio Wave</td>
<td>MRI image</td>
<td>Chemical composition</td>
</tr>
<tr>
<td>10^{-3}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microwave</td>
<td>Ultrasound</td>
<td>Tissue structure characteristics, flow</td>
</tr>
<tr>
<td>10^{-2}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infrared</td>
<td>Infrared</td>
<td>Surface temperature</td>
</tr>
<tr>
<td>10^{-5}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visible Light</td>
<td>Arthroscopy</td>
<td>Anatomy Intraarticular structure, inflammation</td>
</tr>
<tr>
<td>10^{-6}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ultraviolet</td>
<td>UV-radiation</td>
<td>Skin, chronic</td>
</tr>
<tr>
<td>10^{-8}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-Ray</td>
<td>X-Ray</td>
<td>Anatomy Bone injuries</td>
</tr>
<tr>
<td>10^{-10}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gamma ray</td>
<td>Scintigraphy</td>
<td>Inflammation, metabolism, metabolism of the bone</td>
</tr>
<tr>
<td>10^{-12}</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Partial radiation:

Hildebrandt, 2010
X-ray computed tomography

Application:
- Assessment of skeletal and lung abnormalities
- Heart function
- Tumor growth and angiogenesis

Advantages:
- High spatial resolution with a relative low time required for scanning

Drawbacks:
- Radiation burden
- Volume of contrast agent (animal research)

Spiral CT scanner:

Micro CT:

http://www.le.ac.uk/richardiii/science/microct.html

**Magnetic Resonance Imaging**

**Micro-MRI:**
- Require stronger magnets (at least 4.7 T), specific receiver, stronger gradient sets
- 23% of all small-animal imaging

**Application:**
Multiple due to the big variety of MRI techniques

**Advantages:**
- Non-ionizing, use of tissue properties
- Excellent contrast and spatial resolution
- Variety of MRI techniques (signal weighting, contrast agents, DWI, fMRI)

**Drawbacks:**
- Very expensive
- Longer acquisition time
Positron emission tomography
Single-photon emission computed tomography

Application:
- SPECT: cardiology, neurology (brain perfusion, neurotransmission)
- PET: metabolism, angiogenesis, hypoxia, amyloid imaging (11C-PiB PET)
Optical imaging

Modalities:
- Bioluminescence (luciferin substrate)
- Fluorescence and near-infrared fluorescence (NIR)
- Diffuse optical tomography (DOT)

Application:
- Molecular imaging of reporters
- Enzymatic imaging, tumor angiogenesis

Advantages:
- High sensitivity
- Low costs
- Relatively high throughput
- Short acquisition time

Drawbacks:
- Low resolution
- Limited depth
Photoacoustic imaging: Listen to absorption

Is the combination of optic imaging and US.

a, Pulsed light of time-shared multiple wavelengths illuminates the tissue of interest

b, In the tissue there are absorbing elements. In response to the fast absorption of light by this elements acoustic responses are generated. They can then be detected with acoustic detectors.

Advantage: Combination of high optical contrast and submillimeter ultrasound resolution

V. Ntziachristos, Nature America 2010
Contrast agent versus molecular imaging probe

**Anatomical imaging:** = Contrast agent

CT: Iodine-based, barium
MRI: Gd, Mn, SPIO

**Molecular imaging:** = imaging probes

PET: $^{18}$F, $^{11}$C, $^{13}$N, $^{68}$Ga
SPECT: $^{99m}$Tc, $^{123}$I, $^{111}$In
OI: Organic fluophores, inorganic semiconductor nanoparticles
### Table 6: Summary of the main characteristics of small-animal imaging modalities

<table>
<thead>
<tr>
<th>Imaging modality</th>
<th>Spatial resolution (mm)</th>
<th>Sensitivity</th>
<th>Depth</th>
<th>Imaging period (min)</th>
<th>Radiation exposition (Gy)</th>
<th>Maximum number of animals/same study</th>
<th>Equip. cost (M€)</th>
<th>Procedure relative cost</th>
<th>Type of probe</th>
<th>Major advantages</th>
<th>Major limitations</th>
<th>Major applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPECT</td>
<td>0.5–2</td>
<td>pM</td>
<td>No limit</td>
<td>30–90</td>
<td>0.1–1.0</td>
<td>4</td>
<td>0.6–0.8</td>
<td>$$</td>
<td>Variety of molecules labeled with low-energy gamma emitters</td>
<td>Longer physical half-lives; multiple radionuclides can be detected simultaneously; physiological and molecular information</td>
<td>Less sensitive than PET</td>
<td>Oncology; cardiology; neurology (brain perfusion and neurotransmission)</td>
</tr>
<tr>
<td>PET</td>
<td>1–2</td>
<td>pM</td>
<td>No limit</td>
<td>20–60</td>
<td>0.1–1.0</td>
<td>4</td>
<td>0.6–0.8</td>
<td>$$$</td>
<td>Variety of molecules labeled with positron emitters</td>
<td>High sensitivity; accurate quantification; diversity of biological probes available; physiological and molecular information</td>
<td>Short-lived radionuclides; impracticability of distribution of some of them; expensive equipment and overall procedure</td>
<td>Oncology (tumor metabolism/proliferation, angiogenesis, hypoxia); neurology</td>
</tr>
<tr>
<td>CT</td>
<td>0.05</td>
<td>mM</td>
<td>No limit</td>
<td>10–15</td>
<td>0.1–0.6</td>
<td>1</td>
<td>0.2–0.4</td>
<td>$$</td>
<td>Radiopaque contrast agents</td>
<td>Spatial resolution, particularly for lung and bone imaging; morphological and physiological information</td>
<td>Poor soft tissue contrast; radiation exposure</td>
<td>Bone, lungs, vascular imaging</td>
</tr>
<tr>
<td>MRI</td>
<td>0.1</td>
<td>pM–mM</td>
<td>No limit</td>
<td>60</td>
<td>None</td>
<td>10</td>
<td>1.0</td>
<td>$$$</td>
<td>Paramagnetic metal chelates/superparamagnetic iron oxide</td>
<td>Spatial resolution; high soft tissue contrast; morphological, physiological and molecular information</td>
<td>Low sensitivity and long acquisition times</td>
<td>Oncology (tumor metabolism and oxygenation); cardiology (heart perfusion)</td>
</tr>
<tr>
<td>OI</td>
<td>1–2</td>
<td>pM–nM</td>
<td>mm</td>
<td>1–10</td>
<td>None</td>
<td>5</td>
<td>0.1–0.4</td>
<td>$</td>
<td>Fluorophores or β-emitters</td>
<td>High sensitivity, high throughput; low cost; physiological and molecular information</td>
<td>Limited depth of penetration</td>
<td>Oncology (tumor angiogenesis, and enzymatic activity)</td>
</tr>
<tr>
<td>US</td>
<td>&lt;0.1</td>
<td>–</td>
<td>cm</td>
<td>60</td>
<td>None</td>
<td>1</td>
<td>0.2–0.3</td>
<td>$</td>
<td>Microbubbles</td>
<td>Vascular and soft tissue imaging; morphological and physiological information</td>
<td>Difficult to image hollow organs and bone</td>
<td>Vascular imaging</td>
</tr>
</tbody>
</table>

*CT* computed tomography, Equip equipment, Gy Gray, *μm* micromolar, *mm* millimeter, MRI magnetic resonance imaging, M€ million euros, min minutes, nM nanomolar, OI optical imaging, PET positron emission tomography, pM picomolar, SPECT single photon emission computed tomography, US ultrasound.
How are these images made? 
Nice pictures .... Are they biomedical useful? 
And from the view of a small animal?

Ultrafast ultrasound localization microscopy for deep super-resolution vascular imaging
Claudia Errico, Juliette Pierre, Sophie Pezet, Yann Desailly, Zsolt Lenkei, Olivier Couture*, Mickael Tanter* 
Nature, November 2015

- Unprecedented spatiotemporal resolution
- Deep penetration & super-resolution
- Images are acquired in 150s !
- In-vivo, „Minimal-invasive“: Catheterized jugular vein for the administration of the contrast agent (conventional microbubbles also used in clinical applications), thinned-skull imaging window
Basic principle of Ultrasound

Sonar imaging, Langevin (20th century)

Acoustic waves: wavelength ($\lambda$), frequency (f), amplitude (A)

- Array of multiple piezoelectric elements
- Line-per-line
- Transmission of a slightly defocused US beam and parallel processing of 4 US beam in the receive mode (25 fps)

Tanter and Fink, 2015
Ultrafast Ultrasound Imaging

- Transmission of plane (or unfocused) waves
- Tilted plane waves with different angles
- Full region of interest, using all array elements
- Higher amplitudes
- Short acquisition (hundreds of microseconds) : functional imaging (tissue motion, brain activity via blood flow)

Tanter and Fink, 2015

Leaving sonar principles: Plane wave insonifying

Bercoff, 2011
Ultrasound Contrast agent

Primarily designed to detect small blood vessels
Biocolloid: Colloid particles made from biocompatible materials
Smaller than the wavelength of diagnostic ultrasound (100 – 1000 μm)

- Gas spheres (perfluocarbon)
- Gas core with a low density basic for resonation
- A clinical US system is in principle capable of detecting the signature from a single microbubble (resolution ...)
- Shell: in clinical use phospholipid (older albumin, protein shelled)
- A clean microbubble is inherently unstable

Solid and liquid nanoparticles:
- Less echogenic than gas bubbles (incompressible)
- Stable; Advantageous pharmacokinetic properties
- Enhanced permeability and retention principle
The PALM approach applied to ultrasound

The problem to solve:

Conventional ultrasound contrast imaging is limited by the classical wave diffraction theory and corresponds roughly to the ultrasonic wavelength (200 μm – 1 mm).

Solution:

Pinpointing the location of the few, well-separated microbubbles that degraded between each image (“blinking of bubbles”)

Viessmann et al. 2013: Achievement of the necessary separation by sufficient dilute of microbubbles. However, hour-long imaging acquisition.
The principle of ultrafast ultrasound localization microscopy

a, Average stack of 250 beamformed images
b, Frames separated by 44 ms and filters (to remove the tissue signal)
c, three independent microbubbles blinking over several milliseconds
Red cross: Exact position of centroid (deconvolution with point-spread function)

http://www.nature.com/nature/journal/v527/n7579/fig_tab/nature16066_SV1.html
Spatial resolution and bubble velocity maps

a, Microbubble density maps with a spatial resolution of $\lambda/10$
Resolution in depth and lateral direction: 8 $\mu$m x 10 $\mu$m
b, Same area in a conventional power Doppler image
c, Interpolated profiles the lines marked in a
d, dynamic tracing of bubbles separates vessels in two populations
e, f, Velocity lines associated with d (n = number of bubbles)
Thinned skull window versus intact skull

a, uULM preformed through shinned skull (8 μm x 10 μm)

b, corresponding velocity map

http://www.nature.com/nature/journal/v527/n7579/fig_t ab/nature16066_SV2.html

c, uULM performed through the intact skull: attenuation of the ultrasound wave in the presence of bone (12.5 μm x 10 μm)
d, corresponding velocity map
Nice pictures .... Are they biomedical useful?

**Possible fields:**
- Normal and diseased blood-vessel function,
- identification of microvessel-related disorders,
- angiogenesis in neoplasms,
- vascular dementia etc
- functional imaging in neuroscience (combining this technique with functional ultrasound)

**Conventional clinical ultrasound:**
Resolution inversely correlates to penetration (f)

**In ultrafast ultrasound localization microscopy:**
Resolution is related to:
- SNR, - BW of backscattered echoes, -N of array elements
= high resolution deep into organs could be reached
= clinical application’s (liver, kidney, breast)

**Human brain?** By the use of longer wavelength maybe the challenge of the thick human skull can be circumvented
And from the view of a small animal?

**Replacement**: Realistic non-invasive human application

**Reduction**: Longitudinal (functional) studies

**Refinement**: Fast acquisition, minimal invasive
Are these bicolloids of further use?

Molecular Imaging,
Weisleder et al, 2010
Theranostics: Agents for diagnosis and Therapy

Theranostic: The combination of diagnostic and therapeutic entities into one drug delivery vehicle for simultaneous diagnosis and treatment of disease

A large variety of inorganic and organic based nanoparticles

Nanoporphyrin:
- All-in-one porphyrin-based organic nanoconstruct (nanoporphyrin NP)
- NP platform which integrates a variety of imaging and therapeutic functions

Yuanpei Li et al., 2014
From micro to nano

In situ conversion of porphyrin microbubbles to nanoparticles for multimodality imaging

- Drug delivery by using ultrasound to implode microbubbles into nanoparticles
- Bacteriochlorophyll-lipid (BChl-lipid) shell; porphyrins confer **photoacoustic** and fluorescent properties
- In-vivo anatomical guidance by the microbubble US contrast
- Nanodroplets are better able to penetrate fenestrated or compromised vessel
- Interaction of ultrasound with drug-loaded microbubbles causing non-lethal transient pores in blood vessels
The echogenity of microbubbles

Microbubble passage of an acoustic wave
A) Bubble with an initial radius of 1.5 μm
B-E) Expanding and Contraction
E-G) Collaps

Basis for a strong and unique echo, microbubble resonate at frequencies typical used in US imaging

Effect of pulse driving pressure on bubble radial oscillation
A and C) Driving (acoustic) pressure; A = 50 kPa and C = 500 kPa
B and D) Expansion ratio in response to either A or C

B represents a linear dynamic
D represents a nonlinear dynamic

Molecular Imaging, Weisleder et al, 2010
Pulse-inversion imaging: Contrast between bubbles and tissue

A and B) Transmitted pulses (A: 50 kPa, B: 100 kPa); B is inverted (180°)

C and D) Corresponding echoes from 1-micron bubble and tissue
- Bubble echoes: With the higher pressure non-linear dynamic

E) Summation of 2 times echo in C plus echo in D
- The linear echoes from tissue are cancelled while non-linear echoes from the bubble are acquired
Basic properties of BChl-lipid microbubbles

a) Acoustic attenuation measurement of pMBs, Resonance attenuation peak at 4.5 MHz
b) Linear and nonlinear ultrasound properties of pMBs
Tissue-mimicking flow phantom (agar and graphite)

Microbubbles generate both linear and non-linear ultrasound signals
“Conversion ultrasound”

“Conversion ultrasound”: 1 MHz, high-duty-cycle (50 %), ultrasound (2 W cm\(^{-2}\))

f, light microscopy image of microbubbles before conversion ultrasound

g, Electron microscopy of porphyrin nanoparticles after ten ultrasound pulses

c, Size distribution of microbubbles before and after conversion ultrasound

d, Concentration of microbubbles and nanoparticles before and after conversion ultrasound

e, Size distribution of nanoparticles before and after conversion ultrasound

Elizabeth Huynh, 2015
Multimodal imaging of Microbubbles (pMBs) and resulting nanoparticles (pNPs) following ultrasound-induced conversion

a, Acrylamide gel phantom imaged with ultrasound, photoacoustic and fluorescence

1) PBS 2) without conversion US 3) one pulse of conversion US 4) three pulses 5) ten pulses

b-d, Quantified signals (b: ultrasound, c: photoacoustic, d: fluorescence

Elizabeth Huynh, 2015
Conversion of pMBs to pNPs in vivo in mice

Mouse model: Subcutaneous inoculation of 2x10^6 KB cells (HeLa derivative) into the right flank of athymic nude mice. Experiments were performed when tumors reached a surface of 5-7 diameter.

a, Cross section of the tumor (linear and non-linear imaging)
Non con. US: After injection, the pMBs circulate into the tumor, reaching a peak in circulation at 20 s and could be observed continuously in the circulation beyond 40 s
With con. US: A decrease of non-linear signal after the 20 s time point

b, ROI analysis without conversion ultrasound (top) and with conversion ultrasound (bottom)
Successful delivery of pNPs into the tumor xenograft

c, Retention of pNPs in the tumor xenograft enabled by conversion from pMBs to pNPs
d, Normalized photoacoustic signal over time in the tumor, indicative for a successful delivery

Elizabeth Huynh, 2015
Nice pictures .... Are they biomedical useful?

Not addressed:
- Penetration of the conversion ultrasound?  
  (1 MHz = high penetration)
- Limitations for the drug delivery? 
  (size, chemical properties etc)

Possible fields:
- Translational medicine (drug delivery)
- The use of nanoporphyrin offer an expansion to other imaging modalities
And from the view of a small animal?

**Replacement**: (further investigation needed, before application in humans are possible)

**Reduction**: Longitudinal studies are possible (Reaches the drug the target tissue?)

**Refinement**: Multimodality imaging (multiple information)

**Drawbacks**: Setup “experimental”, photoacoustic longitudinal studies performed over 2 h
Thank you for your attention!

- Questions?
- Get your forms signed!