

Imaging by Listening to Molecules: From translational research to drug development

Presented by Rui Wang, PhD Cold Spring Biotechnology 24.09.2015



Agenda



Company overview

- MSOT technology
- Selection of MSOT applications
- Handheld system and its clinical outlook

Co-founder: Vasilis Ntziachristos

Major professional appointments

- 2007- Professor & Chair for Biological Imaging
 Technische Universität München, Germany
 School of Medicine and School of Electrical Engineering
 Director, Institute for Biological and Medical Imaging (IBMI)
 Helmholtz Zentrum München, Munich, Germany
- 2002-2007 Assistant Professor Director, Laboratory for Bio-optics and Molecular Imaging (LBMI) Harvard University, School of Medicine & Massachusetts General Hospital, Boston MA

Selected professional activities

- 2013 Established Photoacoustic editorial

 2010 Advisory Board, Journal of Contrast Media & Molecular Imaging

 2008
 Council Member Society for Molecular Imaging
- 2008- Council Member, Society for Molecular Imaging
- 2006- Topical Editor for **Optics Letters**, Optical Society of America
- 2005- Associate Editor, IEEE Transactions on Medical Imaging
- 2005- Associate Editor, International Journal of Biomedical Imaging





<u>NEW JOURNAL – to promote a Photoacoustics community</u>

http://ees.elsevier.com/pacs/





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Medical

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iThera Medical – overview

- Founded in 2010 as a spin-off from Helmholtz Centre Munich
- Developing and marketing novel molecular imaging technology
- Launched first optoacoustic preclinical scanner in 2010
- Supported by BMBF, awarded multiple times as innovative young company







MSOT-based publication each year





24/09/2015

MSOT: Next-gen biomedical imaging







Alexander Graham Bell first published on the photoacoustic effect in 1881

(A. G. Bell, "The Production of Sound by Radiant Energy," Science, vol. 2, pp. 242-53, May 28 1881)



- Incident sunlight was reflected to and rapidly interrupted by a rotating slotted disc
- A thin absorbing solid in the path of the filtered light was connected to a hearing tube
- Bell demonstrated that the strength of the acoustic signal depended on the intensity of the incident light

Availability of adequate lasers, ultrasound detectors, acquisition electronics, algorithms and computing performance now enable practical use

Agenda



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- Handheld system and its clinical outlook

Technology and benefits of "MSOT"





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Advancing biology AND medicine!



Clinical

Preclinical



- Small animal scanner "inVision"
- Proven technology
- Wide range of applications

Handheld system "EIP"

- CE and FDA mark in 2015
- Clinical studies ongoing

Nov 2014

Preclinical imaging with "MSOT inVision" iTheraMedical

Animal preparation



- Defined / repeatable positioning
- Nose cone with anesthesia supply
- Foil membrane for signal coupling

Image acquisition



- Chamber filled with water
- Stages for x-y-z positioning
- Ultra-fast image acquisition

Data processing



- Quantitative image reconstruction
- Multispectral / kinetic data analysis
- Export of images, videos, graphs

Clinical imaging with "MSOT EIP"



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Two modes of EIP operation Handheld Stage-controlled



1140 O R G



Information acquired with MSOT



Anatomical information	Optical absorption of tissue	
Functional information	Perfusion, oxygenation	
Molecular information	(Targeted) probes / FPs	
Kinetic information	PK/biodistribution data	





Implications of wavelength range for NIR optoacoustic imaging

- 1. Organic dyes: Cy5.5, IRDye, ICG, AF750 and similar dyes absorb in the NIR
- 2. Bioinorganic nanoparticles can be synthesized to absorb in the NIR (e.g. AuNR)
- 3. Fluorescent proteins: iRFP* can be detected using a NIR laser; blue-shifted fluorescent proteins (e.g. GFP, YFP, CFP, mCherry) require illumination at wavelengths in the visible range

Agenda



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- MSOT technology

Selection of MSOT applications

Handheld system and its clinical outlook

Imaging anatomy with MSOT



Experiment	Top row shows single-	Brain	Liver	Kidney/Spleen
	images at different characteristic cross- sections. Bottom row shows histology for reference.	A B C 3mm	A B E D C	A C B D Smm
Application	MSOT can visualize optical contrast at high resolution throughout the entire animal cross-	A: superior sagittal sinus B: posterior cerebral art. C: temporal artery	A: spinal cord B: vena cava C: liver D: stomach E: aorta	A: spinal cord B: right kidney C: vena cava D: intestines E: spleen
	The main absorber in tissue is blood. This yields an excellent contrast especially for highly perfused regions such as kidney, liver, and spleen.	A B C	C D	E C C

Anatomy

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Cardio Anatomy Cancer

INJECTION OF MICROSPHERES INTO IIVER			
Experiment	100µm microspheres were injected into the ileo colic vein to introduce small spheres with high absorbance into the liver. The mouse was then scanned <i>in vivo</i> via MSOT, visualizing the distribution of these spheres.	Single slice	Fly-through video
Application	The visualization of microspheres of 100µm diameter demonstrates the capacity of MSOT to detect structures the size of micro-metastases throughout the mouse, especially in hemoglobin- rich organs such as the liver which provide high background absorption.	In collaboration with Lacey R. McNally PhD, University c	of Louisville, USA

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Nov 2014



Assessing probe delivery to tumor area



Courtesy of Helmholtz Zentrum Muenchen (Hertzog et.al. 2011, submitted for publication)



Quantifying perfusion heterogeneity





Experiment	Mice with orthotopic 4T1 tumors were injected with 40 nmoles of liposomal ICG and images were acquired for 30 mins. ROI analysis was performed on a per-pixel basis and MSOT-signal vs. time data (red circles) was fitted (blue curve) to a PK model (cartoon). Derived PK parameters (e.g. Cmax and Tmax) can now be visualized in parametric images and areas of reduced perfusion can be compared to areas of relative hypoxia (blue areas, right panel).
Application	Fast dynamic processes, such as perfusion heterogeneity throughout tumors, can be visualized by MSOT using dynamic contrast enhancement (DCE). This is important for the analysis of tumor perfusion, to compare the extent of EPR effect during tumor growth, or to predict compound delivery to tumor tissue.
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nv Cancer	Cardio Brain Kinetics Inflam.

Analysis of $\alpha\nu\beta$ 3-integrin targeting





Study performed in collaboration with Florian Rechenmacher, Stefanie Neubauer, Prof. Dr. Horst Keller (Technische Universität München)

Experiment	Female BALB/c nude mice were injected with 0.5×10^6 4T1 mouse breast tumor cells in the abdominal mammary fat pad. Ten days post implantation, 5 nMoles of Cy5.5-labeled $a_v\beta_3$ -ligand was administered by tail vein injection and allowed to circulate for 1 hour. Tumor accumulation of probe was visualized (A) and quantified (C) by applying linear regression component analysis using ViewMSOT TM software. Localization of MSOT signal was validated by post mortem whole animal cryoslicing (B). The fluorescent signals (B) were in good accordance with MSOT images (A).
Application	Using MSOT technology receptor expression can be assessed <i>in vivo</i> using short imaging regimens. The targeting to expressed receptors can accurately be visualized and quantified.

Cancer

Cardio >

2

Cell tracking: 巨噬细胞靶向





Application

Cancer

Using MSOT technology cells can be tracked *in vivo* over prolonged periods of time. The biodistribution of labeled cells can be visualized and quantified using spectral unmixing.

Imaging apoptosis markers

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Cancer

Cardio



Analysis of apoptotic process heterogeneity in vivo



- Caspase-targeting apoptosis detection reagent injected i.v.
- Hypoxic regions within the tumor identified by spectral Hb/HbO₂ unmixing
- Strong apoptosis signals detected in more hypoxic regions in the tumor

Imaging of apoptosis in tumor research iThera Medical

Cardio

Cancer

Simultaneous injection of caspase-targeting apoptosis probe and control probe

- Spectral unmixing of biodistribution and tumor targeting of both probes
- Accumulation of targeted probe vs. wash-out of control probe

EGFR-targeting in pancreatic tumor

Hudson SV et al., **Targeted non-invasive imaging of EGFR-expressing orthotopic pancreatic cancer using Multispectral Optoacoustic Tomography (MSOT)**, Cancer Res November 1, 2014 74:6271-6279. DOI: 10.1158/0008-5472.CAN-14-1656.

Experiment	Mice were orthotopically implanted with S2VP10 (derived from human pancreas carcinoma) luciferase cells. A peptide-based probe targeting the EGF receptor was injected i.v., and the mice were analyzed by optical imaging and optoacoustic tomography.
Application	Multiple modalities can track optically labeled tumors. However, MSOT offers high resolution spatial imaging at depth in orthotopic models of pancreas cancer. Tumor volumes can be calculated by 3D rendering of sequential axial data sets.

Cancer

Cardio

27

Imaging probe targeting to tumor

pH-sensitive insertion peptide for tumor targeting

Kimbrough CK, Khanal A, Zeiderman M, Khanal BR, Burton NC, McMasters KM, Grizzle WE and McNally LR. Targeting Acidity in Pancreatic Adenocarcinoma Multispectral Optoacoustic Tomography Detects pH-low Insertion Peptide Probes in vivo. Submitted and under review process

- Mice were implanted with human pancreatic cancer cells and a pH-low insertion peptide V7-750 or control K7-750 were injected i.v.
- Multiple optical imaging methods were used to track V7 and K7 bio-distribution.
- MSOT accurately shows the location of the tumor in deep tissue

Inflam.

Cancer

心脏区域血红蛋白分析

Experiment	Left, single-wavelength MSOT image of the heart and multispectrally processed image showing HbO ₂ and Hb. <u>Right</u> , reference anatomy	Single-WL MSOT image	Reference cryoslice
Application	Spectral decomposition of HbO ₂ and Hb shows regions of highly oxygenated Hb in the heart, allowing a functional characterization of cardiac activity <i>in vivo</i> . Myocardial infarction can thereby be visualized in real time.	Multispectral image HbO ₂ Hb	lung atrium rib ventricle septum sternebra

10 vs. 50 Hz analysis of the heart beat iThera Medical

Experiment	The heart of a 12 day old mouse was scanned by MSOT at 10 and 50Hz. Panel A shows the heart imaged at 50Hz, with a blue ROI drawn within the left ventricle. Panel B shows ROI analysis from A, with the blue ROI data emphasizing signal change from the heart beat, and the red ROI emphasizing breathing and heart beat. Panel C shows multispectral analysis of Hb and HbO2 in the heart at 50Hz (top), or by averaging multiple sequential slices (bottom), which blurs the image. Panel D shows a single-wavelength video of the heart beat at 10 and 50Hz	A () () () () () () () () () ()	t=0.06 sec 3 mm
Application	50Hz acquisition allows the observation of the full cardiac cycle via MSOT, enabling the calculation of the heart and breathing rate. Faster acquisition also obviates the need for averaging, allowing maximal spatial resolution of multispectral absorbers such as Hb and HbO ₂ .	A sveraged	50 Hz acquisition
Nov 2014			30

Cancer

iTheraMedical Imaging vascular anatomy in the brain Listening to Molecules

Experiment	Non-invasive coronal brain showing superficial vasculature and blood vessels within the brain, at multiple levels of the mouse brain Note: all images acquired through intact skin and skull Solid arrow: Superior sagittal sinus; dotted arrow: Temporal artery; long dash: inferior cerebral vessel; short dash: 3 rd ventricle	MSOT signal (a.u) 5 mm
Application	MSOT can be used to accurately reveal the vascular anatomy of the brain through intact skin and skull. MSOT enables the visualization of anatomical and structural changes associated with the presence of lesions, tumors and hydrocephalus.	Burton NC et al., Multispectral Optoacoustic Tomography (MSOT) Brain Imaging and Characterization of Glioblastoma, Neuroimage, 2012 Sep 28; pii: S1053-8119(12)00963-
Nov 2014		

Cancer

Brain

31

3D vasculature of head / brain

Cardio

Brain

Detecting probes deep in the brain

Cancer

Cardio

Brain

Experiment	Left: Grayscale background is an optoacoustic image taken at 900nm. Green overlay represents multispectrally resolved probe; <u>Right:</u> Cryosection with green overlay showing fluorescence from NIR dye797 SSS: superior sagittal sinus D3V: dorsal 3rd ventricle LV: lateral ventricle 3V: 3rd ventricle	Multispectral MSOT image	Reference cryoslice
Application	MSOT can be used to accurately determine the spatial biodistribution of probes in the brain through an intact skin and skull. In combination with specific probes, this provides the capacity to study molecular features of neurological disease <i>in vivo</i> .	2mm Lozano N et al., Liposome-gold Nanorod Hybrid Tissues , J Am Chem Soc, 2012 Aug 15;134(32):13	s for High-resolution Visualization Deep in 3256-8.
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iTheraMedical Monitoring perfusion in the brain Listening to Molecules Experiment ICG was injected via the **MSOT** perfusion video **Reference anatomy** tail vein and the accumulation in the blood Superior Temporal vessels of the head and sagittal arterv sinus brain was monitored in real time. **Application** The accumulation and clearance of probes in the brain can be monitored in ICG level: 03.8% := 15.0s real time, allowing the **ICG spectrum ICG signal intensity** direct visualization and calculation of the extinction coefficient (cm^-1/M) pharmacokinetics of 1.5x10⁵ MSOT value (a.u) 0.75 molecular probes in the 1.0x10⁴ brain. 0.50 5.0x10⁴ 0.25 MSOT can be used to 0 00determine the half-life of 5.0 7.5 10.0 12.5 15.0 0.0 2.5 700 750 800 850 900 Time (minutes) Wavelength (nm) drugs and the localization of disease markers in the Burton NC et al., Multispectral Optoacoustic Tomography (MSOT) Brain Imaging and brain. Characterization of Glioblastoma, Neuroimage, 2012 Sep 28; pii: S1053-8119(12)00963-9 34

Brain

2D monitoring oxygenation in the brain is the brain is the brain in the brain is th

Imaging iRFP-transfected brain tumor

Adapted from: Deliolanis et al. 2014, Deep-Tissue Reporter-Gene Imaging with Fluorescence and Optoacoustic Tomography: A Performance Overview, Mol Imaging Biol. 2014 Mar 8.

Experiment	Mice were implanted with U87 glioblastoma cells transfected with a near-infrared-absorbing protein and imaged <i>in vivo</i> by multiple modalities, including epi-illumination / trans-illumination fluorescence and MSOT, and compared to <i>ex vivo</i> cryo-sectioning with fluorescence detection.	
Application	Compared to established optical imaging modalities, MSOT more accurately determines the distribution of fluorescent protein in the brain <i>in vivo</i> and with high resolution, thus providing 3D information about the tumor volume, without losing resolution with increasing imaging depth.	
Nov 2014	36	
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MSOT anatomy of metabolic organs

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Cancer





Preclinical imaging in drug discovery

MSOT imaging in the drug discovery stage



Molecular imaging:

- Target validation (localization and expression levels)
- Imaging drug localization and efficacy (characterization of pathology and response to treatment)

Pharmacokinetics and biodistribution imaging:

- Identification of PK issues in Hit-to-lead stage and optimization of animal PK profiles
- PK/PD relationships in preclinical studies
- Absorption, Distribution, Metabolism, Excretion (ADME) analysis
- Toxicity screening and TK

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Listening to Molecule

Measuring probe PK properties with MSOT



Experiment	After i.v. injection of probe (Cy-X; 25nMoles) the jugular veins of mice were monitored over 30 minutes by MSOT. After ROI analysis concentration-time curves were determined and non- compartmental analysis was performed. This allows for the calculation of important pharmacokinetic parameters such as half-life, clearance and AUC; all without having to bleed or sacrifice the animals.	After injection	5 Minutes	5 Minutes 10 Minutes 5 Minutes 6 Minutes		30 Mi	nutes ROI Unit Unit unin µg/m
Applicatio n Dec 2012	By MSOT, PK parameters of NIR-absorbing agents can be determined non-invasively, thus reducing the number of animals and saving time. PK analysis can aid in the opti- mization of dosing schedules and imaging time points	Study performe	10 ne after injec ed in collaborati	20 30 tion (min)	CI MRT	0.06845 30.55 STAR, SBI	mL/min min C, Singapore

Determining probe pharmacokinetics





Application Using MSOT technology pharmacokinetic behavior can be studied non-invasively. PEGylated liposomal ICG is an excellent intravascular contrast agent.

Dec 2012

Anatomy

Cardio

Quantitative analysis of kidney function Istening to Molecules



Cancer

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Whole body biodistribution imaging



ICG-labeled fatty acid distribution throughout the mouse



Mouse scanned from head to base of tail (10 minute scans)

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Assessment of renal damage



Experiment	Renal clearance of	Control	Adriamycin	ROIs
	by MSOT in saline and adriamcyin-treated (6.5mg/kg) mice.			Boom (a.u.)
Application	ROI analysis of dye clearance can differentiate	4 min	Зтт	0 0 5 min 3 mm
	kinetics in the renal cortex and pelvis, allowing the observation of a Tmax delay that provides a surrogate marker for glomerular function. This correlates with histological markers	12 Tito Telais 6 Col 0.5 Col 0.5 Time (minutes)	x — Pelvis ntrol, min 0 11 12 0 11 12 0 11 12	Cortex Pelvis Adr, 2.5 min 9 10 11 12 inutes
	(Picrosirius Red) of renal damage.	Control Ac	ar 3.5 Con	trol Adr
Scarfe L et al., Nove measurements of	el techniques for non-invasive glomerular kidney damage.	Contraction of		

Submitted for publication.

Cancer

Cardio

Nov 2014

Kinetics

Analysis of urinary excretion



Experiment	A catheter was inserted into the tail vein of a nude mouse, and the mouse was imaged by MSOT. During image acquisition, an organic NIR- absorbing dye (AlexaFluor 750) conjugate was injected i.v., and the accumulation of this dye in the bladder was monitored over a two hour period. Shown are individual time points, with a single wavelength MSOT image (800nm) in greyscale and the spectrally unmixed AlexaFluor750 overlaid	T = 0:00:00 min T = 0:12:28 min Image: Constraint of the second secon
	in jet. Also shown is a time-lapse video showing the accumulation over a two hour period.	
	The graph shows experimental values (black dots) as well as the modeled data (red line) in the ROI indicated in the first panel.	180 160 - 140 - 140 - 140 -
Application	The high temporal resolution of MSOT combined with high spatial resolution imaging through the entire cross-section of the mouse allows an investigator access to organs involved in excretion as well as the ability to quantify these physiological processes.	20 0 0 0 0 0 0 0 0 0 0 0 0 0

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Cancer

Cardio

Kinetics

In vivo assessment of gastric emptying iThera Medical

Experiment	 10 nMoles of ICG was administered by oral gavage to BALB/c nude mice. MSOT images were acquired every 8-10 minutes post administration for 120 minutes. ICG signals are superimposed onto single-wavelength optoacoustic images (850nm; grayscale) and quantified (panel a). Clearance kinetics are determined by fitting the MSOT data to a 2-compartmental pharmacokinetic model that reveals a clearance half-life from the stomach of ~22 minutes (panel b). Postmortem validation by fluorescence imaging is in good correspondence with MSOT (panel c). Panel (d) shows real-time distribution and clearance of ICG 	a iiver stomach b t = 15 min d head> iiver stomach	t = 5 min	t = 40 min C 1 (rv) reußg	t = 75min • MSOT • Validation • Validation • Validation	iodel
Application	signals in the stomach and liver. MSOT technology offers the potential to monitor physiological regulation and pharmacological modulation of gastric emptying.	intestines <tai⊳ t = 5 min</tai⊳ 	Mo mu tor PK Pho	rscher S et al S Iltispectral opt mography (MSC imaging of gas ptoacoustics. 202	Semi-quantitativ oacoustic DT) for volumetr stric emptying. 14 2(3):103-110.	e ic
Nov 2014			~			

Kinetics

Cardio

Cancer

Nanoparticle distribution



Accumulation of dendrimer particles in tumor-tissue



- ¹⁸F PET imaging offers sensitive quantification, but poor resolution and limited ability for longitudinal studies
- Optical imaging offers the ability for longitudinal studies, but with poor resolution

Nanoparticle distribution



Accumulation of dendrimer particles in tumor-tissue



MSOT offers high-resolution longitudinal imaging in 3D

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Nanoparticle distribution



Heterogenous accumulation of dendrimer particles in tumor-tissue





- By MSOT distribution imaging it is possible to monitor nanoparticle accumulation in tumors longitudinally
- In contrast to ¹⁸F PET imaging, the accumulation can be tracked for multiple days, in this case the more crucial time points, as the dendrimer continues to accumulate post 6hrs
- In contrast to optical imaging, MSOT imaging offers high resolution imaging in 3D
- Uniquely, MSOT offers the capability to combine the imaging of injected probes with functional imaging (Hb and HbO₂)

Photosensitizer-based MSOT contrast



in vitro analysis allows for quick prioritization



 ϕ F=fluorescence QY; ϕ \Delta=singlet oxygen QY; ϕ P=MSOT QY

Ho & Balasundaram et al. Scientific Reports (2014)

Five PDT agents of different classes were evaluated for MSOT signal generation:

 Zinc phthalocyanine (ZnPc), protoporphyrin IX (PpIX), 2,4-bis[4-(N,N-dibenzylamino)-2,6dihydroxyphenyl] squaraine (Sq), chlorin e6 (Ce6) and methylene blue (MB)

In vivo biodistribution analysis





- MSOT imaging allows visualization and quantification of distribution of injected agents
 - ZnPC reaches a maximum Tumor-Background ratio at T=60mins
- Quick optimization of theranostic approaches

Upconversion Nanoparticles as a Contrast Agent for Photoacoustic Imaging in Live Mice





PAI of UCNP in live mouse







9/24/2015

Simultaneously resolving two agents





Imaging of inflammation in the knee

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Cancer

Cardio



Experim	ent 12 mice were injected with Freund's Complete Adjuvant (FCA) into the left knee. Three weeks later, mice were imaged by MRI (Gd-enhanced, T1-weighted). Mice showing induction of FCA-induced inflammation were then further injected with dPGS-NIR, which targets P and L selectins indicative of inflammation, and imaged by MSOT.	$Control \begin{array}{c c} Single \lambda & dPGS-NIR & HbO_2 & MRI \\ \hline A & An & B & C & C & G & G & G & G & G & G & G & G$
Applicati	on MSOT offers structural and functional imaging of joints involved in arthritis. Further, molecular imaging via MSOT enables visualization and quantification of P and L selectins, allowing MSOT- guided staging of inflammation in a mouse model of arthritis.	FCA FCA by by b

Inflam.

Agenda



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- Selection of MSOT applications

Handheld system and its clinical outlook

MSOT Experimental Imaging Platform (EIP) istening to Molecules



Two modes of EIP operation

Handheld

Stage-controlled





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Current EIP detector options



# Elements	Geometry	Central frequency	Resolution	Field of view (xyz)
128	2D	8 MHz	115 µm	5 x 10 mm
256	2D	4 MHz	200 µm	20 x 30 mm
384	3D	2.5 MHz	300 µm	15 x 15 x 25 mm
384	3D	4 MHz	200 µm	10 x 10 x 15 mm
512	3D	10 MHz	80 µm	5 x 5 x 7 mm

3D MSOT





2D MSOT



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Clinical applications for MSOT EIP					
Shortlist of currently as	sessed app	olications	Clinical study ongoing Clinical study upcoming		
Clinical application	Incidence	MSOT value			
Malignant melanoma	230K	Non-invasive, improved detection			
Sentinel node detection	>2.000K	Non-radioactive procedure			
Peripheral vascular disease	5.000K	Assess tissue hypoxia as primary burden			
Rheumatoid arthritis	1.000K	Point-of-care inflammation monitoring			
Breast cancer	1.700K	Non-invasive lesion assessment			
Head and neck cancer	640K	Plan radio-/chemotherapy, monitor ablation			
Diabetic wound healing	15.000K	Assess progress of chronic ulcer healing			
Alopecia	>100.000K	Assess follicle composition for hair growth			
Colon anastomosis	>1.000K Predict anastomotic leakage		akage		
Neonatal brain injury	>1.000K	Accurate hemorrhage / ischemia detection			

Dynamic 5D mouse brain imaging





Science&Applications 2014

- Dynamic contrast enhancement via ICG injection enables quantification of fast kinetics in the brain
- 5D imaging allows spatial and kinetic imaging with specificity



4T1 tumors imaged with the 512-element cup (10MHz)



- Mice bearing orthotopic 4T1 tumors were imaged using the EIP 100 system with a cup-shaped transducer with 512 US elements (10MHz central frequency)
- The laser was operated at a frequency of 25Hz, allowing for fast multispectral data collection (5Hz, for 5 wavelengths)

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Live spectral analysis in 2D



Spectral unmixing of blood oxygenation and melanin



- Besides real-time reconstruction, tissue chromophore distribution can be spectrally unmixed in real time
- Oxy- and deoxy-Hb as well as melanin are among the most interesting intrinsic markers for MSOT imaging
- Chromophore concentration is visualized as overlay on background tissue absorption

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3D high-resolution vascular imaging





Deán-Ben et al., **Portable, Spherical array probe for volumetric real-time optoacoustic imaging at centimeter-scale depths**, Opt Express. 2013 Nov 18

- Structures can currently be visualized up to approx. 2 cm depth
- 10 MHz detector 80µm resolution and visualization of microvasculature

Functional angiography with MSOT





Deán-Ben et al., **Functional optoacoustic human angiography with handheld video rate three dimensional scanner**, Photoacoustics 2013: 1(3-4):68-73.

- Oxygen saturation is quantified via spectral unmixing of Hb and HbO₂
- Changes in perfusion and oxygenation can be visualized in real time



- A melanoma patient was scanned with MSOT before undergoing surgical sentinel lymph node (SLN) biopsy
- MSOT enables the detection of SLNs via subcutaneous ICG injection and the assessment of melanin presence, potentially indicative of LN metastasis

Multispectral imaging of hair follicles



 The morphology of hair follicles as well as their micro-environment are determinants for continued hair growth

Listening to Molecules

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- Endogenous chromophoric substances including hemoglobin, melanin and lipids are critical for hair growth
- MSOT can assess morphology and endogenous chromophore distribution related to hair growth and alopecia

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MSOT inVision 512-echo

Optoacoustic UltraSound (OPUS)



Hybrid Optoacoustic/Ultrasound (OPUS) iTheraMedical

MSOT inVision 512-echo

OPUS imaging system



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Imaging chamber



Technical parameter	Specifications		
Center frequency	5 MHz		
Number of elements	512		
Angular coverage	270°		
Radius / focus	40 / 37 mm		
Height / pitch	15 / 0.37 mm		
Resolution OA / US	150 / 300 µm		

Interleaved image acquisition

Single-wavelength OA and US images



Imaging mouse anatomy with OPUS





24/09/2015

Tumor imaging with OPUS





- 4T1 mammary tumors were implanted in the mammary fat pad
- Ultrasound imaging shows clear tumor boundaries, allowing segmentation
- MSOT imaging allows functional oxygenation analysis within the tumor

Imaging probe accumulation in bladder



- The bladder was imaged during i.v. injection of IRDye-800CW
- Ultrasound imaging allows identification of the bladder, invisible in MSOT in naïve mice, prior to injection
- Bladder uptake of dye reflects glomerular filtration and excretion, enabling functional analysis of the kidney

Selection of MSOT publications



- Shanice V Hudson et al., Targeted Non-invasive Imaging of EGFR-expressing Orthotopic Pancreatic Cancer using MSOT, Cancer Res. 2014 Sep 12. DOI: 10.1158/0008-5472.CAN-14-1656.
- N. Beziere et al., Optoacoustic Imaging and Staging of Inflammation in a Murine Model of Arthritis, Arthritis Rheumatol. 2014 Aug;66(8):2071-8. DOI: 10.1002/art.38642
- X. Luís Deán-Ben et al., Adding fifth dimension to optoacoustic imaging: volumetric time-resolved spectrally enriched tomography, Light: Science & Applications (2014) 3, e137; doi:10.1038/lsa.2014.18.
- Stritzker J et al., Vaccinia Virus-mediated Melanin Production Allows MR and Optoacoustic Deep Tissue Imaging and Laser-induced Thermotherapy of Cancer, PNAS February 26, 2013 vol. 110 no. 9 3316-3320.
- Buehler A et al., **Real-time handheld multispectral optoacoustic imaging,** Opt Lett. 2013 May 1;38(9):1404-6. doi: 10.1364/OL.38.001404.
- Burton NC et al., Multispectral Opto-acoustic Tomography (MSOT) Brain Imaging and Characterization of Glioblastoma, Neuroimage, 2012 Sep 28; pii: S1053-8119(12)00963-9.
- Herzog E at al., Optical Imaging of Cancer Heterogeneity with MSOT, Radiology. 2012 May;263(2):461-8.
- Taruttis A et al., Fast Multispectral Optoacoustic Tomography (MSOT) for Dynamic Imaging of Pharmacokinetics and Biodistribution in Multiple Organs, PLoS ONE 2012, 7(1):e30491.
- Razansky D et al., Volumetric Real-time Multispectral Optoacoustic Tomography (MSOT) of Biomarkers, Nature Protocols 6, 1121-1129 (2011).

Current Users

- 1. Institute for Biological and Medical Imaging (IBMI),Helmholtz Zentrum München, Munich, Germany
- 2. Institute for Radiology,Klinikum rechts der Isar, Technische Universität München, Munich, Germany
- 3. ZMB/Faculty of Medicine, University Hospital Esssen, Germany
- 4. Department of Surgery, University Medical Center Groningen, The Netherlands
- 5. Bioorganic Chemistry and Molecular Imaging (LCBIM),École Polytechnique Fédérale de Lausanne, Switzerland
- 6. Centre for Drug Delivery Research,UCL School of Pharmacy, University College London
- 7. Department of Imaging, Merck Research Laboratories, Merck Inc., Philadelphia, USA
- 8. Roche Diagnostics, Penzberg,

24.09.2015



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- 2. Chang Chun Institute of Applied Chemistry Chinese Academy of Sciences
- 3. Institute of Automation, CAS
- 4. Beijing University of Chemistry technology
- 5. Institute of Materia Medica
- 6. Henan University of Traditional Chinese Medicine
- 7. Soochow Univeristy
- 8. Beijing University of Technology
- 9. IMPLAD
- **10. SCUT**
- 11. The Hong Kong Polytechnic University
- 12. West China Hospital
Conclusions, outlook



- Multispectral Optoacoustic Tomography(MSOT) provides:
 - ...anatomical, functional and molecular information
 - ...in real time
 - ...at high spatial resolution, in deep tissue
 - The range of preclinical/research applications in pharmacokinetic research spans wide...
 - Plasma-concentration time curves of absorbers
 - Whole body biodistribution at high spatial and temporal resolution
 - Longitudinal studies of heterogeneous accumulation allow for the visualization and quantification of ADME processes
 - Toxicity assessment
 - Dynamic Contrast Enhanced(DCE) MSOT
- Clinical translation is within reach!
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- OPUS is the world's first hybrid tomographic optoacoustic / ultrasound imaging technology, featuring unparalleled highresolution image quality, user-independent and whole-body
- Incorporation of ultrasound into MSOT was achieved through innovation of the detector and by developing novel acquisition and image reconstruction algorithms for reflection-mode ultrasound computed tomography (R-UCT)
- The addition of anatomical ultrasound contrast expands the utility of MSOT in a wide range of applications, including cancer and pharmacology

PARTICIPATE IN AN IMAGING REVOLUTION. Introducing MSOT, the next generation in molecular imaging.