Optical Drug Monitoring: Photoacoustic Imaging of Nanosensors to Monitor Therapeutic Lithium in Vivo

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ABSTRACT  Personalized medicine could revolutionize how primary care physicians treat chronic disease and how researchers study fundamental biological questions. To realize this goal, we need to develop more robust, modular tools and imaging approaches for in vivo monitoring of analytes. In this report, we demonstrate that synthetic nanosensors can measure physiologic parameters with photoacoustic contrast, and we apply that platform to continuously track lithium levels in vivo. Photoacoustic imaging achieves imaging depths that are unattainable with fluorescence or multiphoton microscopy. We validated the photoacoustic results that illustrate the superior imaging depth and quality of photoacoustic imaging with optical measurements. This powerful combination of techniques will unlock the ability to measure analyte changes in deep tissue and will open up photoacoustic imaging as a diagnostic tool for continuous physiological tracking of a wide range of analytes.

KEYWORDS: diagnostic · continuous monitoring · bipolar · nanoparticle · nanomedicine

Medical imaging and diagnostics are two key pillars of personalized medicine and the study of disease. However, in order to advance these fields, we need to develop novel approaches and technologies for in vivo imaging and monitoring of physiologically important analytes. Conventional optical imaging techniques suffer from the limited penetration depth of light and the loss of spatial resolution in deep tissue due to scattering. Photoacoustic imaging1–3 mitigates many of these shortcomings and achieves imaging depths of up to several centimeters with acoustically defined spatial resolution. Many photoacoustic techniques focus on imaging either endogenous signals such as hemoglobin4 or exogenous contrast agents such as carbon nanotubes,5 gold nanoparticles,6 and iron oxide nanoparticles.7 These measurements are particularly valuable for examining local oxygenation or tissue structure, but there is immense value in the untapped use of photoacoustics as a measurement tool to monitor changes in analyte concentrations deep within intact tissue. Some development work has focused on the combination of photoacoustics with molecular imaging probes for distribution,8 pH,9,10 oxygen/reactive oxygen species,11,12 or glucose13 but these probes are available for limited targets and do not incorporate the ability to tune sensor response to desired physiological ranges.

Fluorescent nanosensors are well-established, and multiple distinct designs have proven capable of monitoring a wide range of analytes in vitro14,15 and in vivo16–19. These nanosensors offer several key advantages that make them suitable for continuously tracking analyte levels. A great deal of development work has established methods for both producing nanosensors and tuning a nanosensor’s analytical performance to physiological concentration ranges. These nanosensors are particularly appealing because they have the added benefit of a modular design. With the modular optode design, the nanosensor’s target analyte, spectral properties, and response range arise from straightforward design choices such as component ratios. However,
balances the charge inside the sensor. An additive (R) extracted from bulk solution into the nanosensor, altering the protonation state of a chromoionophore (CH$^+$), changing the optical properties of the nanosensor. An additive (R$^-$) balances the charge inside the sensor.

In this paper, we formulate, characterize, and validate lithium-sensitive nanosensors for photoacoustic measurements and compare to the results obtained with fluorescence measurements. Lithium is a valuable model system for photoacoustic imaging because continuous measurement technologies such as photoacoustics can provide a method to track systemic concentrations of therapeutic drugs and improve disease management. Lithium is a common treatment for bipolar disorder, but it has both a narrow therapeutic window (0.6–1.2 mM) and a low toxic dose (~2 mM), demonstrating a need for continuous monitoring.

RESULTS

The core mechanism for lithium recognition and signaling in these nanosensors is based on the well-established optode approach, demonstrated previously for several ionic and small-molecule analytes. In this instance, lithium is recognized by a lithium selective crown ether ionophore, lithium ionophore VI, and extracted from bulk solution into the hydrophobic polymer core of the nanosensor (Figure 1). This extraction of a cation into the core of the polymer causes a shift in the internal pH of the nanosensor, altering the protonation state of a chromoionophore, which alters the optical properties of the nanosensor. Both the absorption spectrum and fluorescence intensity of the nanoparticles change as a function of lithium concentration. These optical changes occur at visible wavelengths, which present difficulties in obtaining good fluorescent signals for in vivo imaging, but represent an opportunity for photoacoustic imaging.

Photoacoustic imaging is based on the absorption of photons by a molecule and the conversion to ultrasonic waves via the photoacoustic effect. When the absorption spectrum of the chromoionophore changes, the photoacoustic spectrum changes accordingly (Figures 1a, 2a). As the lithium concentration increases, the absorbance and photoacoustic amplitude at 515 nm increase, and the absorbance and photoacoustic amplitude at 660 nm decrease. Importantly, these changes in photoacoustic amplitude result from the nanosensor and chromoionophore, because the analyte itself does not possess any intrinsic photoacoustic properties. From the measurements at these two peak wavelengths (Figure 2b), we can form a ratiometric index that increases linearly with lithium concentration but is not susceptible to common artifacts such as variation in nanosensor concentration or number. Photoacoustics can also be used to image these nanosensors under ~1.5 mm of chicken muscle tissue (Figure 2c,d). The signal attenuation at 515 nm is larger than at 660 nm, which decreases the contrast at that wavelength and reduces the ratiometric values for the data while remaining linear over the physiologically relevant range of lithium.

Lithium selective nanosensors injected into the skin of mice can be imaged with photoacoustic...
tomography (Figure 3a), but a key advantage of photoacoustics is the ability to collect data from the entire tissue volume (Figure 3b, Supporting Video 1). Importantly, the data collected in the surface projection is a result of the entire sensor injection, rather than the nanosensors closest to the skin surface, as is the case with fluorescence imaging. Upon intraperitoneal (ip) administration of lithium to the animals at a physiological concentration of 38 mg/kg (~2.5 mM blood concentration),\textsuperscript{26} the nanosensor photoacoustic ratio increased by 25%. This clear response to the increasing lithium has a time to peak concentration of 14 min (lithium $n = 3$, vehicle $n = 1$).

These multimodal nanosensors generate both fluorescent and photoacoustic signals, which allow us to compare the two imaging modes for the sensors. Traditionally, these sensors have relied on the fluorescence of the chromoionophore directly, which is in the visible range. In this work, we shifted the fluorescence to near-IR (NIR) wavelengths to minimize optical effects from imaging in tissue (Figure 1b). To accomplish this shift, we incorporated an NIR fluorophore (DiR) into the nanosensor formulation used for photoacoustics. This dye serves as an internal reference signal and a FRET acceptor from the chromoionophore. When directly excited, the NIR intensity from DiR is not responsive to changes in lithium concentration, providing a reference signal that controls for a variety of factors, such as nanosensor number, injection depth, and sensor migration. When the chromoionophore is excited, energy transfer to DiR generates an NIR intensity that changes with lithium concentrations, because the fluorescence of the donor (chromoionophore) is responsive to lithium. When the ratio of these two signals is calculated, the resulting index is correlated with the lithium concentration and controls for other variables, similar to the photoacoustic index.

In vivo monitoring of lithium levels is drastically improved relative to traditional fluorescence imaging through the use of NIR fluorescence. Figure 4a shows example fluorescent images of mice with lithium selective nanosensors injected into their skin. Following administration of lithium via ip injection under the same conditions as the photoacoustic experiments above, the nanosensor fluorescence ratio increases in a dose-dependent manner with increasing systemic lithium concentrations (Figure 4b, Supporting Figures 15 and 16). The time to peak concentration, 18 min, is similar to that of photoacoustics, although the magnitude of the signal change is 8% as opposed to 25% for photoacoustics at the same lithium dose, again highlighting the advantages from photoacoustic imaging and improved imaging depth even when compared with NIR fluorescent imaging.

In vitro design, fabrication, and characterization of both photoacoustic and fluorescent nanosensors are
essential for effective in vivo application. These steps are similar to previous reports\textsuperscript{18} and are also detailed thoroughly in the Supporting Information (Supporting Figures 2–14). Two key characteristics were tuned during the formulation process: sensitivity in the physiological range for lithium (0.5–4 mM)\textsuperscript{29} and selectivity for the nanosensors over sodium. Sensitivity is primarily controlled through selection of the chro-
omionophore and the ratio of the sensor components. Selectivity is controlled through the selection of the lithium ionophore and the addition of TOPO (triocetylphosphine oxide), which has been shown to increase lithium selectivity in macroscale optodes.\textsuperscript{20} We found a similar effect of TOPO on the selectivity of nanosensors (Supporting Figure 3), increasing selectivity over sodium to a level suitable for monitoring changes in lithium concentration in the background of physiological sodium. As a result, changes of sodium in the physiological range do not affect the nanosensor signal. Details on tuning sensor response, final formulations, further spectral data, and additional characterization are provided in the Supporting Information. The calibration of the response to lithium for the final formulation of nanosensors in the background of physiological sodium is shown in Supporting Figure 6. The EC\textsubscript{50} for the fluorescent sensors is 3.7 mM with a sensitivity of 45%/log at 2 mM lithium compared with 10 mM and 29%/log for photoacoustic imaging. These results are similar, indicating that the mechanism underlying sensor response to lithium is decoupled from the readout mechanism as expected. Due to the small size of the nanosensors (27 nm by Dynamic Light Scattering (DLS), Supporting Figure 9), the nanosensors are able to respond faster than 15 s (Supporting Figure 11), significantly faster than necessary for the dynamics of lithium administration. The nanosensors are also reversible (Supporting Figure 12), which is an essential property for in vivo monitoring so the sensors can continuously monitor increases and decreases in lithium concentration.

**DISCUSSION AND CONCLUSION**

This is the first report of photoacoustic imaging to measure therapeutic drug concentrations in vivo. The results show a promising improvement over fluorescence imaging of the same nanosensors and can image a depth profile with \sim 200 \mu m resolution. Imaging nanosensor injections in three dimensions solves several problems with current fluorescence-based approaches because the depth profile separately measures sensor responses in regions closest to the capillary bed. For larger molecules such as glucose, the differences between values near the capillary bed and skin surface may differ significantly,\textsuperscript{30} and photoacoustic imaging may provide superior diagnostic power. Fluorescence and photoacoustic measurements yielded the same basic response in vitro and lithium kinetics in vivo, demonstrating that the multimodal imaging does not alter the mechanism of the sensors and highlighting the robustness of the platform and earlier nanosensor development work. This validation is a key step toward the application of these nanosensors for imaging in deeper organs such as the brain.

A common limiting factor for time resolution in basic and clinical pharmacokinetic research is the need for blood sampling. This is the first in vivo real-time tracking of lithium levels without blood sampling. Lithium is a common and potent treatment for bipolar disorder, but its narrow therapeutic window and low toxic dose make it more difficult to dose than other pharmaceutical therapies. A minimally invasive continuous monitor for blood lithium concentrations would be highly useful for dose administration and management.\textsuperscript{21} Nanosensors produce a ratiometric photoacoustic index and a ratiometric fluorescent index that both respond to lithium concentrations in vivo. These multi-wavelength indexes are less affected by sensor concentration and sensor injection depth than single-wavelength intensity measurements, although still suffer from biases resulting from differential tissue attenuation at the two wavelengths. Commonly, it is believed that shifting fluorescence wavelengths to the NIR will alleviate the effect of tissue absorption and scatter, but the improved index change from photoacoustic imaging indicates that this approach is only partially effective. In the larger context of continuous physiologic monitoring, this modular nanosensor platform translates directly to other electrolytes\textsuperscript{16,31} and small molecules such as histamine,\textsuperscript{17} with future work in our groups directed toward photoacoustic monitoring of these analytes with nanosensors. Additionally, this approach should work with other optical and photoacoustic imaging configurations (nonlinear optical, tomography, etc.) to obtain necessary resolution or imaging depth for particular applications outside of drug monitoring. Future work in our groups will determine the maximum penetration depth where we can image nanosensor function as well as image with higher spatial resolution. The incorporation of other imaging approaches such as SERS (surface-enhanced Raman spectroscopy),\textsuperscript{32} SPECT (single-photon emission computed tomography),\textsuperscript{33} CT (computed tomography),\textsuperscript{34} MRI (magnetic resonance imaging),\textsuperscript{35} and fluorescence deconvolution\textsuperscript{36} will also yield value to these sensors.

Photoacoustic imaging is much less established than fluorescence imaging, and several technical challenges obstruct the widespread adoption of the technique. First, photoacoustic imaging instrumentation, not available in most research groups, is more complex than fluorescence imaging instrumentation. However, improved light sources and more portable imaging setups are under development, and commercial
photoacoustic imaging systems are starting to appear on the market for preclinical applications. Current research efforts in the photoacoustics community have begun to focus on increasing the photoacoustic contrast from exogenous probes. Future work in our groups will focus on developing nanosensors with active wavelengths in the NIR region. This will benefit both fluorescence and photoacoustic imaging by eliminating much of the background signal and enhancing sensitivity of the approach. Despite these current shortcomings, the results herein show that photoacoustics is a measurement mode for nanosensors that can provide previously unattainable measurement depths, which can have profound effects on chronic disease management.

**METHODS**

Poly(vinyl chloride), high molecular weight (PVC), bis[2-ethylhexyl] sebacate (DOS); sodium tetrasodium-3,5-bistrifluoromethyl-phosphonic acid (NaTPFB), 9-dimethylamino-5-[4-[(15-buty1)-1,3-dioxo-2,14-dioxanonoacyl]phenylimino][benzol][p]oxazoline (chromoionophore VII; CHVII), 6,6-dibenzy1-1,4,8,11-tetraacycloclotetradecane (lithium ionophore VI; Ll VI), tricyclophosphine oxide (TOP0), tetrahydrofuran (THF), 4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES), dichloromethane (DCM), 9-dimethylamino-5-[4-[(15-buty1)-1,3-dioxo-2,14-dioxanonoacyl]phenylimino][benzol][p]oxazoline (chromoionophore II; CHII), N,N-dicyclohexyl-N'N'-diobutyl-cis-cyclohexane-1,2-dicarboxamide (lithium ionophore III; Ll III), 2-nitrophenyl octyl ether (NP0E), 2-nitrophenyl phenyl ether (NPPE), and lithium chloride were purchased from Sigma-Aldrich (St. Louis, MO, USA), 1,2-Dioleoyl-sn-glycero-3-phosphoethanolamine-N-[methy1(polyethylene glycol)-550] amminum salt in chloroform (DSPE-PEG) was purchased from Avanti Polar Lipids (Alabaster, AL, USA), 1,1-Dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine iodide (DiR) was purchased from Life Technologies (Grand Island, NY, USA). 2-Amino-2-hydroxypropylpropane-1,3-diol, 2 M solution (TRIS, 2M) was purchased from Fisher Scientific (Waltham, MA, USA). Phosphate-buffered saline (PBS with Ca²⁺ and Mg²⁺; pH = 7.4) was purchased from Boston Bioproducts (Ashland, MA, USA).

**Nanosensor Fabrication.** Protocols used in this report are based on those previously described. In brief, the process of fabricating optodes and nanosensors starts with formulation of an optode cocktail comprising 500 μL of THF containing PVC, DOS, and the sensing components. The ratio of these components is tuned to control the response of the nanosensors. The formulation of the nanosensors used here is 60 mg of PVC, 120 μL of DOS, 20 mg of TOP0, 20 mg of Li VI, 22 mg of NaTPFB, and 2 mg of CH VII dissolved in 1 mL of THF. The formulation for fluorescence nanosensors also includes 0.5 μg of DiR in each batch.

To fabricate nanosensors from this optode cocktail, 2 mg of DSPE-PEG (80 μL of a 25 mg/mL solution in chloroform) was dried in a 4 dram scintillation vial and then resuspended in 5 mL of PBS with a probe tip sonicator for 30 s at 20% intensity (Branson, Danbury, CT, USA). A 50 μL amount of the optode cocktail was combined with 50 μL of dichloromethane and added to the PBS/PEG–lipid solution under probe tip sonication (3 min, 20% intensity). The nanosensor solution was added to the PBS/PEG–lipid solution under probe tip sonication (3 min, 20% intensity). The nanosensor solution was added to the PBS/PEG–lipid solution under probe tip sonication (3 min, 20% intensity). The nanosensor solution was added to the PBS/PEG–lipid solution under probe tip sonication (3 min, 20% intensity).

The bulk of nanosensor formulation and screening experiments examine only the chromoionophore fluorescence rather than a ratio of two fluorescent signals. Additionally, screening experiments used 10 mM HEPES and 6 mM TRIS as the buffer solution instead of PBS. This is to enable quantification of lithium selectivity over sodium. Nanosensors were added to a 96-well plate, lithium (or sodium) solutions were then added to the wells to final concentrations of 0 mM through 1 mM, and the plate was scanned with a Spectr��nd M3 plate reader. End point fluorescence values were obtained at an excitation wavelength of 665 nm, emission at 705 nm, and a cutoff filter at 665 nm. In experiments where DiR was included in the formulation, fluorescence was measured with excitation/emission:cut-off at 640/780:695 for FRET and 740/780:NA for DiR fluorescence. Fluorescence spectra were acquired using similar settings. Absorbance spectra were taken between 400 and 850 nm.

To characterize nanoparticle diameter and morphology, scanning electron microscopy (SEM) was performed on fluorescent nanosensors. Nanosensors were dried under vacuum on conductive adhesive tape attached to 25 mm aluminum SEM mounts from Electron Microscopy Sciences (Hatfield, PA, USA). SEM images were taken using a Hitachi S4800 HRSEM operating at an accelerating voltage of 5.0 kV. Nanoparticle diameter was measured using ImageJ.

**Photacoustic System.** Two photoacoustic tomography systems were used for this research. For in vivo phantom experiments, photoacoustic computed tomography (PACT) was used as previously detailed. Briefly, a tunable optical parametric oscillator laser (basiScan 120; Spectra-Physics) pumped by a Nd:YAG laser (Brilliant b; Quantel) with 10 Hz pulse repetition rate was used to excite photoacoustic waves. Light exiting the laser system was homogenized by an optical diffuser and then illuminated the sample. The photoacoustic waves were detected by a 512-element circular transducer array (iLasonic) with 5 MHz central frequency. The imaging system has 0.10 mm radial resolution and 0.25 mm tangential resolution. The image was reconstructed based on back projection algorithm.

**Fluorescent Nanosensor Characterization.** Nanosensors were calibrated in vitro utilizing a Lumina II in vivo imaging system (Caliper Life Sciences, Hopkinton, MA, USA). This plate was imaged with two channels: DiR: high lamp power, excitation filter centered at 745 nm (30 nm bandpass), emission filter from 810 to 875 nm, and a 1 s exposure. CHVII/FRET: high lamp power, excitation filter centered at 640 nm (30 nm bandpass), emission filter from 810 to 875 nm, and a 1 s exposure. For data analysis, regions of interest were drawn over each well using Living Image 4 software (Caliper Life Sciences), and total fluorescent intensity values were obtained for each well. The ratio of the intensities of the two channels (DiR/FRET) was calculated and then converted to alpha by normalizing to the ratio obtained in the presence of 0.25 mM HCl and then from that the maximum protonated and deprotonated states for the sensors. The data for the calibration curve was fit to a four-parameter logistic curve using Prism 6 to determine EC₅₀ and sensitivity.
(5072PR, Panametrics-NDT) and digitized by an oscilloscope (ZT4421, ZTEC Instruments). The imaging system was mounted on a linear translation stage (XY-6060, Danaher Motion) to acquire three-dimensional images. The spatial resolutions of the system are approximately 0.17 mm and 0.29 mm in the axial and lateral directions, respectively.42

**Photoacoustic in Vitro Characterization.** Mixtures of nanosensors and different concentrations of lithium were injected into silicone tubing. Then, these pieces of tubing were sealed by rubber at both sides. The sealed tubing was embedded in agar gel and covered by chicken breast tissue for imaging. On the basis of these initial results, we chose imaging parameters for later in vivo imaging to minimize signal from vasculature and tissue to focus on the signal from nanosensors alone.

**In Vivo Studies.** All in vivo studies were approved by the Institutional Animal Care and Usage Committee (IACUC) of Northeastern University. In vivo photoacoustic studies were also approved by the Institutional Animal Care and Usage Committee of Washington University in St. Louis.

In Vivo Photoacoustic Studies. The mice used in this research were Hsd:Athymic Nude-Foxn1tm1 Harlan Laboratories. Photoacoustic imaging experiments were conducted using the system described above. Animals were anesthetized with 1.5% isoflurane in oxygen, and 40 μL of nanosensors were injected intradermally (id) into the mouse on the back. The animals were imaged with two channels at excitation wavelengths 515 and 660 nm (the peak wavelengths for the PA signal). Baseline raster scan images were acquired at both wavelengths, and continuous monitoring was done with both wavelengths at the center of the nanosensor injection. Data points were acquired approximately every minute after an ip injection of 38 mg/kg lithium (administered as lithium chloride in PBS, experimental) or a matching volume of PBS (control). Images were acquired every minute for approximately 1 h. Animals were imaged one at a time. All animals were sacrificed after experiments were completed. For data analysis of each experiment, an index of the two photoacoustic intensities was generated by dividing the 515 nm signal by the 660 nm signal. These ratios were normalized to that at the first time point after injection of lithium. These data were linearly interpolated to align time and amplitude points before averaging. Error bars for the lithium data set represent the standard deviation of three animals.

**Fluorescent in Vivo Studies.** The mice used in this research were male SKH: I-E nude mice from Charles River (Wilmington, MA, USA). Fluorescent imaging experiments were conducted using a Lumina II in vivo imaging system. Animals were anesthetized with 2.25% isoflurane in oxygen and placed in the animal imager. A 30 μL amount of nanosensors was injected id into the mouse on the back. The animals were imaged with two channels. DiR high lamp power, excitation filter centered at 745 nm (30 nm bandpass), emission filter from 810 to 875 nm, and a 1 s exposure. CHVI-FRET to DiR: high lamp power, excitation filter centered at 640 nm (30 nm bandpass), emission filter from 810 to 875 nm, and a 1 s exposure. Baseline images were acquired for approximately 30 min, followed by an ip injection of 12 or 38 mg/kg lithium (administered as lithium chloride in PBS, experimental) or a matching volume of PBS (control). Images were acquired every minute for approximately 1 h. The imaging equipment could not image six animals simultaneously, so animals were imaged in pairs. All animals were sacrificed after experiments were completed. For data analysis of each experiment, a region of interest encompassing the injection area was selected and total fluorescent intensity for each channel was recorded. At each time point, an index was generated by dividing the intensity from the DiR channel by the CHVI-FRET channel. These ratios were normalized to the first time point before injection of lithium. These data were then averaged together across three experimental animals for each lithium concentration and six control animals using linear interpolation to align time and intensity points before averaging. Error bars represent the standard deviation of these animals.

**Conflict of Interest:** The authors declare the following competing financial interest(s): L.V.W. has a financial interest in Microphotoacoustics, Inc., and Endra, Inc., which, however, did not support this work.

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**Supporting Information Available:** Additional information on optimization and in vitro characterization of nanosensors as well as full data sets for in vivo experiments are presented. This material is available free of charge via the Internet at http://pubs.acs.org.

**REFERENCES AND NOTES**


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