Imaging extracellular potassium dynamics in brain tissue using a potassium-sensitive nanosensor

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1 Introduction

During the repolarization phase of action potentials, neurons release potassium (K$^+$), causing an increase of the extracellular K$^+$ concentration ([K$^+$]$_o$). Since K$^+$ is one of the key membrane potential determinants and, thus, fundamentally affects electrical signaling, the central nervous system has several homeostatic mechanisms to control [K$^+$]$_o$. In addition to passive diffusion through the extracellular space (ECS), K$^+$ is taken up and released by both glia and neurons, which is accomplished by various K$^+$ selective channels and transporters. In addition to its fundamental effects on neuronal excitability, changes in [K$^+$]$_o$ have been shown to influence long-term potentiation, vascular tone and glial morphology. Impaired [K$^+$]$_o$ homeostasis is associated with a wide range of pathological states, such as epilepsy, cortical spreading depression, and ischemic stroke/anoxia. It is yet unclear whether pathological changes in [K$^+$]$_o$ are a result or a cause of some of these diseases. To address the role of [K$^+$]$_o$ in physiological and in pathological conditions, monitoring the spatial and temporal dynamics of [K$^+$]$_o$ changes is of crucial importance.

To date, [K$^+$]$_o$ has been measured predominantly using K$^+$-sensitive microelectrodes. Although still the methodological standard for measuring [K$^+$]$_o$, they are nonetheless limited by only providing a single point measurement. They also introduce a substantial dead space in the tissue, which may alter local structure and thus function. This makes their application limited in scope, especially from the perspective of monitoring the spatiotemporal flux of K$^+$, which is needed to address the standing questions regarding K$^+$ kinetics and dynamics.

Alternatively, one may attempt to image extracellular ion dynamics with ion sensitive dyes. This, however, is a far less widespread practice due to methodological problems, such as immediate diffusion of the dye through the ECS and a lack of dyes with a suitable affinity for their respective ion concentrations found in the ECS. Due to these methodological difficulties, published attempts to image K$^+$ in the ECS are limited to only a few studies. For instance, in one study, the authors utilized a long wavelength fluorescent sensor based on the tetra-zachlorine moiety, which was applied to the exposed cortex of anesthetized mouse to visualize K$^+$ waves in the cortex. Others injected the K$^+$ sensitive dye APG2 into the cisterna magna, and, after the diffusion of the dye throughout the brain, performed in vivo fluorescence imaging of the cortex. Both studies utilize in vivo fluorescence imaging strategies that, while highly valuable, only permit the imaging of superficial cortical regions.

To image various brain regions that are not accessible with current in vivo optical imaging methods, acute brain slice preparations can be used. This approach is also benefited by the ability to place microelectrodes more accurately and in the desired brain region for electrophysiology, as well as the ability to control the environment of the tissue to a greater degree. Imaging extracellular ion dynamics in acute slices is more difficult, however, due to the lack of an enclosed compartment that the skull and dura naturally provide, as well as the constant superfusion of artificial cerebrospinal fluid (ACSF) over the slice. Together, these cause a rapid washout of small molecule dyes from the ECS, with little to no time to gather useful data.

**Abstract.** Neuronal activity results in the release of K$^+$ into the extracellular space (ECS). Classically, measurements of extracellular K$^+$ ([K$^+$]$_o$) are carried out using K$^+$-sensitive microelectrodes, which provide a single point measurement with undefined spatial resolution. An imaging approach would enable the spatiotemporal mapping of [K$^+$]$_o$. Here, we report on the design and characterization of a fluorescence imaging-based K$^+$-sensitive nanosensor for the ECS based on dendrimer nanotechnology. Spectral characterization, sensitivity, and selectivity of the nanosensor were assessed by spectrofluorimetry, as well as in both wide-field and two-photon microscopy settings, demonstrating the nanosensor efficacy over the physiologically relevant ion concentration range. Spatial and temporal kinetics of the nanosensor responses were assessed using a localized iontophoretic K$^+$ application on a two-photon imaging setup. Using acute mouse brain slices, we demonstrate that the nanosensor is retained in the ECS for extended periods of time. In addition, we present a ratiometric version of the nanosensor, validate its sensitivity in brain tissue in response to elicited neuronal activity and correlate the responses to the extracellular field potential. Together, this study demonstrates the efficacy of the K$^+$-sensitive nanosensor approach and validates the possibility of creating multimodal nanosensors.

**Keywords:** extracellular space; potassium; potassium buffering; fluorescence microscopy; two-photon imaging; fluorescent indicator; nanotechnology.

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To overcome these challenges, we designed a $K^+$ potassium nanosensor based on dendrimer nanotechnology. For this, we used a larger molecule [generation 5, polyethylene glycol (PEG)-functionalized, PAMAM dendrimer] as a scaffold for the small molecule $K^+$ sensitive dye Asante Potassium Green 4 (APG4) through a noncovalent interaction. This strategy of dye encapsulation has been previously applied in our laboratory for intracellular imaging with the sodium (Na$^+$) sensitive dye, corona green. Here, we show that this new APG4-based nanosensor is well retained in the ECS, is selective and sensitive enough to measure $[K^+]_o$, at ionic concentration ranges relevant for ECS, and can be used to measure rapid local changes in $[K^+]_o$ in response to iontophoretic pulses of $K^+$. Furthermore, we show changes in $[K^+]_o$ after the induction of neuronal activity with glutamate and correlate this to local field potential. Overall, we demonstrate that this APG4-based $K^+$ sensitive nanosensor is suitable for imaging $[K^+]_o$ in acute brain slices and can be used to monitor the spatiotemporal dynamics of $[K^+]_o$ in the brain.

2 Materials and Methods

2.1 Nanosensor Synthesis

Asante Potassium Green 4 (APG4), (Teflabs, Texas, USA) and Generation 5 PAMAM dendrimers with a 1,4-diaminobutane core and a poly (ethyleneglycol) methyl ether surface (Andrews ChemServices, Michigan) were used to prepare the nanosensor. The preparation of PAMAM dendrimer and APG4 was performed at a 1:1.2 molar ratio to ensure at least one dye molecule per dendrimer. For the ratiometric nanosensor, a second dye, AlexaFluor 568 (AF568) was incorporated, with a final 1.2:1.2:1 APG4 to AF568 to dendrimer ratio. To remove any unbound dye molecules after their integration into the dendrimer complex, the solution was passed through an ultrafiltration column with a 3kDa cutoff (Millipore, Massachusetts). Filtration was repeated several times to ensure a thorough washout of unbound dye. After being prepared, the nanosensor was stored at $-20^\circ$C in aqueous HEPES buffer at pH 7.5. Under these conditions, the nanosensor remains stable for >1 year.

2.2 Spectral Characterizations

Fluorimetric analysis was performed in quartz cuvettes using a using a spectrofluorimeter (LS55 Luminescence, PerkinElmer, Massachusetts). First, the nanosensor was diluted to 100 nM in an extracellular-like solution containing (mM): NaCl 160, KCl/K$^+$/glucurate 0-50, CaCl$_2$ 1.3, MgSO$_4$ 0.81, NaH$_2$PO$_4$ 0.78, HEPES 20 adjusted to pH 7.4 with NaOH. For $K^+$ calibration curves, fluorescence intensities of the emission maxima were plotted against known $K^+$ concentrations. For this, K-glucurate (1M) was titrated in the cuvette over the range of 0 to 50 mM, and the results were corrected for dilution. For Na$^+$ experiments, extracellular-like solution, as described above, was used but with 5 mM KCl and 135 mM NaCl. NaCl (2M) was then titrated to gather data in the range of 135 to 165 mM Na$^+$ and were also corrected for dilution. For Mg$^{2+}$ and Ca$^{2+}$ experiments, MgSO$_4$, CaCl$_2$, and Ba(OH)$_2$ (adjusted to pH 7.4) solutions were titrated in extracellular-like solution (5 mM $K^+$), as described above. All spectral characterizations are an average of three separate scans.

2.3 Sol-Gel Synthesis

To immobilize the nanosensor for in vitro spatiotemporal imaging, silica sol-gels were synthesized in two steps: hydrolysis and condensation. The hydrolysis consists of the following reaction: Si(OMe)$_3$ + MeOH + H$_2$O $\rightarrow$ MeOH + Si-OH, while the condensation follows the reaction: Si-OH + Si-OH $\rightarrow$ Si-O-Si + H$_2$O. Hydrolysis was performed under a fume hood at room temperature by mixing MeOH, H$_2$O, and Si(OMe)$_3$ in molar ratios of 7.5:10:1. After hydrolysis, a small volume of fluorescent nanosensor preparation was diluted in <100 µl of the sol-gel mix and 5 µl drops of the fluorescent sol-gel mix were pipetted onto glass microscope coverslips. Drops were allowed to condense and dry and were used for experiments after 24 h.

2.4 K$^+$ Iontophoresis

Iontophoresis was carried out using a Neurophore BH-2 (Digitimer, Hertfordshire, United Kingdom). Borosilicate glass pipettes (~5 or 25 MΩ) containing 1M KCl were used. A DC holding current of 50 nA was applied before and after iontophoretic pulses to ensure $K^+$ retention. 500 nA pulses were used for rapid $K^+$ ejection.

2.5 Imaging

Single-photon fluorescence imaging of sol-gels was performed on an inverted epifluorescence microscope (Axiovert 100M, Carl Zeiss, Jena, Germany) using a 40 × 1.3 N.A. oil-immersion objective lens. Selection of the fluorescence excitation wavelength was performed using a monochromator (Till Photonics, Gräfelfing, Germany), and fluorescence was detected using an EM CCD camera (Andor, Belfast, United Kingdom).
acquisition was computer-controlled using the Metafluor imaging software (Molecular Devices, California).

Two-photon imaging of sol-gels and acute brain slices was carried out using a custom-built two-photon microscope with a 40×0.8 N.A. water-dipping objective (Olympus). Fluorescence excitation was performed using a Chameleon Vision S femtosecond infrared laser including group velocity dispersion compensation (Coherent, California). Captured emission wavelengths were 525 ± 25 nm and 620 ± 30 nm for the green and red channels, respectively. Image acquisition and linescan was performed using custom-written software in the Labview environment.

**Fig. 2** Spectral properties and sensitivity of potassium nanosensors. (a) Single-photon excitation and emission spectra of the APG4 PAMAM-PEG nanosensor measured in a spectofluorimeter. (b) In vitro emission spectra (λex 515 nm) of APG4 PAMAM-PEG with a K⁺ titration in the range 0 to 50 mM, corrected for dilution. (c) Comparison of sensitivity of three different K⁺-sensitive fluorescent indicators encapsulated in PAMAM dendrimers. The respective responsivity of APG4, APG2, and PBFI is shown for the same K⁺ concentration range. (d) Differential sensitivities of the K⁺-indicator PBFI to K⁺ in solutions mimicking the extracellular or the intracellular milieu (λex 330 nm) in response to a K⁺ titration ranging from 0 to 150 mM. Intracellular-like solutions contained 15 mM Na⁺, whereas extracellular-like solution contained 150 mM Na⁺. (e) APG4 PAMAM-PEG fluorescence in response to an increasing Na⁺ concentration spanning the extracellular physiological range, and (f) to the divalent cations Ca²⁺, Mg²⁺, and Ba²⁺ measured at a fixed 5 mM K⁺ concentration emission (λex 515 nm, λem 537 nm).
2.6 Brain Slice Preparation
After decapitation of 2.5- to 3-week old mice, brains were extracted and 300-μm thick hippocampal slices were prepared with a vibratome (VT 1000S, Leica, Wetzlar, Germany), as previously described. Slices were imaged in an open chamber superfused with ACSF, bubbled with 95% O₂ 5% CO₂. ACSF solution contains (mM): NaCl 125, KCl 3, NaHCO₃ 25, CaCl₂ 1.2, MgSO₄ 1, NaH₂PO₄ 1.25, glucose 10. Local delivery of the APG₄ PAMAM-PEG nanosensor was achieved by applying pressure pulses of 5 to 60 s at 1 to 2 psi to the back of a glass pipette of tip diameter 1 to 2 μm using a Pressure System II (The Toohey Company, Fairfield, New Jersey). Pipettes were retracted out of the tissue after satisfactory dye application.

All experimental procedures were carried out in strict accordance with the recommendations of the Swiss Ordinance on Animal Experimentation and were specifically approved for this study by the Veterinary Affairs Office of the Canton of Vaud, Switzerland.

2.7 Extracellular Field Recordings
Borosilicate glass pipettes (~4.5 MΩ) were filled with an ACSF-like solution. Recordings were obtained in the zero current clamp configuration using a Multiclamp 700B amplifier (Molecular Devices) in gap-free mode. Data were acquired at 10 kHz by a Digidata 1440 analog-to-digital converter, controlled with the pCLAMP 10 software.

3 Results
3.1 Nanosensor Design, Preparation, and Characterization
The K⁺ sensitive nanosensor is composed of a generation 5 poly (amidoamine) (PAMAM) type dendrimer with a poly(ethylene-glycol) functionalized surface. With our strategy, this G5 PAMAM-PEG dendrimer, which has been measured as 6.57nm in diameter, becomes a host for small molecules, such as the K⁺ sensitive dye APG₄, via electrostatic interaction (Fig. 1). Tertiary amines that compose the dendrimer core are thought to play a key role in this incorporation process, since electrostatic interaction of any anionic dye with core amines will determine its integration pattern or lack thereof, as reported for other guest–host dendrimer interactions.

A dye-to-dendrimer ratio of 1.2:1 was selected to ensure at least one dye per dendrimer molecule. After incorporation of the dye, the complex was repeatedly passed through a centrifugal filter with a 3-kDa cutoff to eliminate nonencapsulated dye. While each dendrimer most likely encapsulates a single APG₄ molecule, the possibility that a single given dendrimer may contain fewer or more APG₄ molecules has not been ruled out, as one has to consider the statistical nature of the nanoparticles produced.

Emission and excitation spectrum of the APG₄ PAMAM-PEG nanosensor in extracellular-like solution are presented in Fig. 2(a), along with the excitation spectrum of the free APG₄ indicator. The spectra exhibit an excitation maximum around 515 nm and show a slightly red-shifted maximum excitation wavelength compared with free APG₄. The emission maximum was found to be ~540 nm. The K⁺ dependent fluorescence was analyzed over a K⁺ concentration range of 0 to 50 mM [Fig. 2(b)] and showed that the K⁺-nanosensor fluorescence increases stepwise with increasing [K⁺], and did not present spectral shifts in response to the increasing concentration. The peak emission intensity was plotted against [K⁺] and compared to two other nanosensors, APG₂ PAMAM-PEG and PBFI PAMAM-PEG, which only differ in regards to their encapsulated fluorescent indicator [Fig. 2(c)]. The APG₄-based nanosensor exhibits a fivefold higher fluorescent yield when compared to the APG₂ and PBFI nanosensors in an extracellular-like milieu. It is noteworthy that the classical K⁺-indicator PBFI behaves drastically differently in solutions mimicking the intracellular and extracellular milieu (Fig. 2(d)), which may explain the poorer performance of PBFI PAMAM-PEG compared with the other two nanosensors.

Fig. 3 Single-photon fluorescence imaging of APG₄ PAMAM-PEG immobilized in a silica sol-gel. (a) Image of a condensed sol-gel preparation containing the APG₄ PAMAM-PEG nanosensor. During the process of condensation, the thin layer of silica sol-gel attached to the coverslip cracks into small fragments. (b) Fluorescence recording of the nanosensor embedded in sol-gels during superfusion of extracellular-like solutions containing 0, 5, 10, 15, or 20 mM K⁺, and showing the responsiveness and reversibility of sensor response.
experimentally, caution must be employed when using the nanosensor.

After demonstrating the sensitivity of the nanoprobe in a cuvette, we then investigated its spatiotemporal characteristics using single and two-photon microscopy in vitro. For this purpose, we incorporated the nanosensor into a porous silica glass using a sol-gel process, providing us with an in vitro platform in which we could image its performance, all the while keeping it immobilized, as well as closely controlling the local environment via iontophoresis or superfusion of solutions.

After preparation of the sol-gel silica structures on cover slips [Fig. 3(a)], we performed single-photon imaging in a closed perfusion chamber, in which we could measure the stepwise increase in fluorescence following increasing concentrations of $K^+$. Superfusion and imaging of the sol-gel fragments demonstrated that its porosity is sufficient for allowing fast diffusion of solutes through their matrix, while being able to immobilize the nanosensor. The stepwise increase in fluorescence and the reversibility of the signal upon washout of the high $K^+$ solution demonstrate that the dye is stable and responsive over time [Fig. 3(b)].

Because two-photon microscopy is significantly more advantageous for imaging in thick tissue and it is primarily the setting in which this nanosensor is ultimately intended to be employed, we analyzed the APG4 PAMAM-PEG containing silica glass in a two-photon microscope. This allowed us to verify its efficacy in a two-photon excitation setting, as well as to gather additional data regarding spatial and temporal kinetics of the sensor. To demonstrate the excitation maxima in a two-photon setting, wavelength tuning from 770 to 840 nm was performed. The emission intensity was recorded for each wavelength, correcting for laser output power as measured in the beam path. Figure 4(a) shows the measured two-photon excitation spectrum, showing a maximal peak two-photon excitation wavelength for the nanosensor at ~815 nm.

To analyze the spatial and temporal kinetics of the sensor, iontophoretic $K^+$ ejection was performed. A 1M KCl filled borosilicate glass pipette was lowered to a distance of 5 $\mu$m from the surface of the sol-gel, and iontophoretic pulses were used to expel $K^+$, thereby increasing the local $K^+$ concentration in a radial manner from the pipette tip [Fig. 4(b)]. A local fluorescence increase that was maximal at the tip of the pipette opening and radiating outward could be observed. Figure 4(c) shows

![Fig. 4](https://www.spiedigitallibrary.org/journals/Neurophotonics on 1/22/2018 Terms of Use: https://www.spiedigitallibrary.org/terms-of-use)
the waveform of fluorescence at regions of varying distance from the epicenter, demonstrating spatial variation of the $K^+$ concentration increase and the resulting fluorescence. The characteristically slow increase to steady-state and fast decay of signal also matches the kinetics of iontophoretic release, as demonstrated previously by others. Taken together, these results indicate that the temporal and spatial characteristics of fluorescence of the nanosensor are favorable and that the signal is an accurate representation of the $K^+$ fluxes induced. To further explore the temporal kinetics of the nanosensor, line-scanning experiments were performed during iontophoresis. The pipette was placed on the surface of the sol-gel, and simultaneous iontophoresis of sulforhodamine 101 (SR101) and $K^+$ was performed, causing a measurable increase in fluorescence of both the red channel (SR101 expelled from the pipette) and the green channel [APG4 PAMAM-PEG embedded in the sol-gel: Fig. 4(d)]. The presence of the two fluorescent dyes in the pipette allowed us to take into account the nonlinear kinetics of iontophoresis. The inflection points of the fluorescence signals were compared as an indication of rapid kinetics of the sensor. The slope of the fluorescence of the red channel reaches its inflection point 33.6 ms before the slope of the green channel. This delay is a good estimate of the responsiveness of the nanosensor in the presence of a sudden $K^+$ rise. By comparison, a fast version of the genetically encoded calcium indicator, GCaMP6F, has an activation time of 74 ms. Synthetic organic ion indicators are generally faster. A range of hypothetical calcium dyes with different $K_a$ values binding Ca$^{2+}$ over a range of concentrations will have a response time between 2 to 40 ms, while the widely used dye fura-2 demonstrates a response time estimated at 11 ms in-vitro and 40 ms in-vivo.

### 3.2 Acute Brain Slice Measurements

The second issue related to imaging $K^+$ in the ECS is the retention of the dye in the tissue. Injecting a small water-soluble molecule such as APG4 into acute brain slices results in a fast disappearance of the dye that makes it impossible to use effectively. Therefore, the dendrimer must serve the function of stabilizing the dye in the ECS for extended periods of time. Figure 5(a) displays the nanosensor’s distribution in the ECS after several microinjections demonstrating concurrent fluorescence of the nanosensor (green), staining of astrocytes with SR101 (red), and unlabeled cell bodies likely belonging to neurons. A comparison between the injections of free APG4 and APG4 PAMAM-PEG demonstrates that the dendrimer greatly improves the nanosensor retention in tissue and that the dendrimer is successfully being retained in the ECS (Fig. 5(b)).

The marked difference in dye signal over time demonstrates the efficacy of the nanosensor design. Occasionally, observed decay of signal after injection in the tissue was found to be mainly due to bleaching, with minimal contribution of washout of the dye, demonstrating that the dye is very well stabilized by the dendrimer in the ECS (data not shown).

To demonstrate sensitivity of the nanosensor in tissue, iontophoresis of $K^+$ in acute slices was carried out with or without the presence of 4-aminopyridine (4-AP, 50 μM) in the perfusion solution (Fig. 5). The amplitude of signal increase was higher for slices perfused with 4-AP. This observation is consistent with the fact that 4-AP is known to block a number of $K^+$ channels and therefore affects $K^+$ buffering out of the ECS. This likely slows the process of $K^+$ uptake by cells in response to the extracellular concentration increase via iontophoresis.

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**Fig. 5** Retention of the potassium nanosensor in the ECS. (a) Fluorescence of the nanosensor (green) injected in the ECS as measured by two-photon microscopy in a mouse acute brain slice ($\lambda_{ex}$ 800 nm) with concurrent SR101 staining of glial cells (red). White arrows indicate microinjection sites. (b) Fluorescence retention of APG4 of free APG4 locally applied compared with the retention of the APG4 PAMAM-PEG nanosensor in the same conditions.

**Fig. 6** $K^+$ iontophoresis in acute brain slices. Measurement of fluorescence of the APG4 PAMAM-PEG nanosensor after iontophoresis of $K^+$ in acute brain slices. Measurements were performed in the absence and in the presence of the $K^+$ channel blocker 4-AP (50 μM, $n = 3$).
One challenge related to imaging brain tissue in a two-photon setting is that it is very sensitive to small movements or drift, which greatly affects the thin focal plane. This is especially true in regards to imaging the ECS. While individual cells show relatively stable signal even with small focal plane shifts, the fine and heterogeneous structures of the ECS are, in our experience, extremely prone to such shifts. Such movement artifacts cannot be avoided entirely, especially with long duration recordings. To counteract this problem, we designed another version of the APG4-based nanosensor. This improved version includes a second, potassium insensitive dye, AlexaFluor 568 (AF568), and is thus dubbed the APG4/AF568 PAMAM-PEG nanosensor [Fig. 7(b)]. Its emission spectral scans [Fig. 7(b)] demonstrate that the sensitivity of the APG4 is not diminished by the addition of another molecule in the dendrimer. The AF568 does not respond to potassium and emits in the red spectrum. This allows for more precise ratiometric imaging of $K^+$ dynamics and demonstrates the general feasibility of the ratiometric, multidaye, and multimodal nanosensor design.

While the basal level of $[K^+]_o$ in the mammal brain rests around 2.5 to 3 mM, periods of intense neuronal activity can raise that value to a physiological ceiling level of around 10 to 12 mM. It was also shown recently with $K^+$-sensitive microelectrodes that bath application of 500-$\mu$M glutamate induces a rise of $[K^+]_o$ in the range of several millimolar in the “stratum radiatum” of the CA1. Using the APG4/AF568 PAMAM-PEG nanosensor, we performed ratiometric imaging of $K^+$ in the stratum radiatum of the CA1 in response to intermittent bath application of 500-$\mu$M glutamate (Fig. 8). In this experiment, spatial resolution was privileged by selecting one small, 5-$\mu$m diameter region of interest. The signal-to-noise ratio of the trace would increase for larger regions. An application time of 30 s was chosen since $[K^+]_o$ elevations typically follow sustained and persistent neuronal activity.

With each sequential application, the ratio of fluorescence of the two channels rises and then drops accordingly. The corresponding field potential was simultaneously recorded, demonstrating a similar waveform. Individual peaks exhibiting common characteristics in both the field potential and fluorescence can be observed, further supporting the efficacy of the nanosensor design.

**4 Discussion**

During neuronal activity, ion fluxes across cell membranes are associated both with the generation of action potentials and with the postsynaptic events. $[K^+]_o$ levels have been shown to undergo substantial fluctuations in brain tissue during physiological and pathological conditions. In particular, neuronal excitation leads to elevations of $[K^+]_o$ that have to be rapidly regulated to enable subsequent activity. Once locally released in the ECS by neurons, $[K^+]_o$ elevations are rapidly dissipated by passive diffusion across the ECS, reuptake into neurons and astrocytes, and redistribution across the astroglial syncytium.

The measurement of $[K^+]_o$ has been so far mainly performed using ion-selective electrodes. These approaches are relatively invasive and do not provide direct spatial information about $K^+$ changes. Fluorescence-based detection of ECS $K^+$ offers the benefits of being both noninvasive and capable of assessing the spatiotemporal pattern of $K^+$ flux within the tissue. The advantages of spatiotemporal data acquisition in regards to ion dynamics are evident when one considers the wealth
of knowledge uncovered, for instance, in the field of \( \text{Ca}^{2+} \) signaling.

The ECS in the brain occupies 15% to 30% of the volume, and its geometry is made of an interconnected network of <100 \( \text{nm} \) tube- or sheet-like conduits. Due to the limited volume and diffusion constraints of the ECS, the local neuronal \( \text{K}^{+} \) efflux (e.g., occurring during the repolarization phase of action potentials) is sufficient to significantly increase \( [\text{K}^{+}]_{o} \). While the basal \( [\text{K}^{+}]_{o} \) concentration is \( \sim 3 \text{ mM} \) somewhat lower than the plasma concentration, \( [\text{K}^{+}]_{o} \) may rise to 10 to 12 \( \text{mM} \) during sustained neuronal activity. While the ECS presents some diffusion barriers due to its constrained geometry, diffusion of small molecules in the ECS is relatively fast. In particular, organic fluorescent molecules routinely used for ion imaging have molecular weights typically <1500 Da. Local application of such dyes in the ECS leads to disappearance of fluorescent signal within seconds, as seen in Fig. 5. The strategy that we followed was to incorporate a \( \text{K}^{+} \)-selective indicator inside a larger scaffold consisting of a generation 5 PAMAM dendrimer. The size of this construct, estimated to be around 6.6 nm in diameter with a molecular mass above 100 kDa, considerably slows down its diffusion in the tissue and enables longer term monitoring of ECS fluorescent signals. It should be underlined that smaller dendrimers, i.e., under generation 4, do not form developed enough internal cavities to provide encapsulation, while dendrimers of generation 7 and above are too large, resulting in blocked pores that are less accessible. Generation 5 provides internal cavities large enough to host smaller molecules, such as APG4, and still open and exposed for solutes and ions to readily pass through.

Another important factor is surface functionalization. To prevent toxicity of the dendrimer-based nanosensor, the surface of the dendrimer must be electroneutral. It has been demonstrated that naked PAMAM dendrimers, which possess cationic amine terminals, induce the formation of transient holes in membranes of living cells. This increases toxicity and allows passage to solutes, including the dendrimer itself, across the plasma membrane. In the same study, a PEG functionalized-dendrimer was shown to have no effect over the same concentration range. It has also been shown that PEG-functionalized dendrimers, when inserted inside neurons, do not affect the integrity or electrical properties of the neurons.

While the dendrimer nanoparticle properties play an important role in the nanosensor design, the dye itself is especially important. Due to the fact that \( \text{Na}^{+} \) and \( \text{K}^{+} \) are both monovalent cations with very similar properties, dyes that bind \( \text{K}^{+} \) are likely to have a certain degree of affinity to \( \text{Na}^{+} \). Most ion-sensitive dyes are used to image “intracellular” ion dynamics, but due to differential ionic concentrations across the membrane, some dyes that are effective intracellularly will not be as useful in the extracellular setting. The classical \( \text{K}^{+} \) dye PBFI, for example, is more effective intracellularly, where there is a low concentration of \( \text{Na}^{+} \) and a high concentration of \( \text{K}^{+} \). Extracellularly, the respective ion concentrations are reversed, and PBFI loses sensitivity due to the confounding fluorescence introduced by the high \( \text{Na}^{+} \) concentration [Fig. 4C]. For this reason, it was important to assess the sensitivity of nanosensors based on various dyes in an extracellular-like solution. Figure 4C demonstrates that under extracellular conditions, APG4 is the most sensitive of the dyes tested.

The APG4 fluorescent nanosensor emits green light at \( \sim 540 \text{ nm} \), with a one-photon excitation maximum at \( \sim 490 \text{ nm} \). In two-photon mode, it has a broader excitation centered around 810 nm. As various organic dyes with red emission have similar two-photon excitation wavelengths around 800 nm, one could envisage using one common excitation wavelength while detecting both green and red emission simultaneously. This indeed proved useful for our design, enabling the improvement of the nanosensor via the incorporation of AF568, which can be excited with the same infrared wavelength.

After characterizing the nanosensor using sol-gels and cuvettes, we aimed to demonstrate its efficacy in living brain tissue. Iontophoresis in the tissue or superfusion of solutions are both possible methods for demonstrating differential fluorescence of the nanosensor in the tissue in response to a variable concentration of \( [\text{K}^{+}]_{o} \). However, it is more relevant to both physiology and pathology to assess whether the nanosensor is able to measure changes associated with \( [\text{K}^{+}]_{o} \), where the source of the \( \text{K}^{+} \) fluctuations is the tissue itself.

The experiment carried out in Fig. 6 demonstrates that the sensor is sensitive enough to detect such changes in \( [\text{K}^{+}]_{o} \) and correlates them to a well-established readout, the local field potential. To relate raw fluorescence to concentrations, one would need a calibration protocol for the nanosensor in living tissue. This would require the silencing of neuronal activity, as well as the blockade of a variety of \( \text{K}^{+} \) channels, using molecules such as 4-AP. Barium can also be used to block \( \text{K}^{+} \) channels, but the observed sensitivity of the nanosensor to barium may exclude it from employment in the calibration process, except at low concentrations (<100 \( \mu \text{M} \)). Alternatively, one could envisage extending the sol-gel approach depicted in Figs. 3 and 4 and combining it with the ratiometric nanosensor to calibrate the fluorescence signal ex vivo. This approach, once validated, would significantly accelerate the process of obtaining absolute extracellular \( \text{K}^{+} \) levels in situ.

\( \text{K}^{+} \) homeostasis has many contributing functional components taking part in the process, including Na-K-ATPase pumps, Na-K-Cl cotransporters (NKCC), inwardly rectifying potassium channels (K\(_{\text{i}}\)), voltage-gated potassium channels, and, in the case of the astrocytic synctium, gap junctions. This nanosensor is likely to facilitate future investigations of these mechanisms and their interactions. The spatiotemporal nature of the gathered data will allow for the investigation into the possible existence of \( \text{K}^{+} \) microdomains around activated neurons and the spatial extent of these domains.

In addition to validating the feasibility of the nanosensor for imaging in the ECS, this study highlights the range of possible extensions of the nanosensor strategy. For example, the ratiometric nanosensor design demonstrates the possibility of combining several optical readouts, laying the path for the design of multimodal optical nanosensors and introducing a broader range of possible applications.

**Disclosures**

The authors declare no financial or personal conflicts of interest.

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