Dendrimer-Based Fluorescent Indicators: *In Vitro* and *In Vivo* Applications

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Abstract

**Background:** The development of fluorescent proteins and synthetic molecules whose fluorescence properties are controlled by the environment makes it possible to monitor physiological and pathological events in living systems with minimal perturbation. A large number of small organic dyes are available and routinely used to measure biologically relevant parameters. Unfortunately their application is hindered by a number of limitations stemming from the use of these small molecules in the biological environment.

**Principal Findings:** We present a novel dendrimer-based architecture leading to multifunctional sensing elements that can overcome many of these problems. Applications *in vitro*, in living cells and *in vivo* are reported. In particular, we image for the first time extracellular pH in the brain in a mouse epilepsy model.

**Conclusion:** We believe that the proposed architecture can represent a useful and novel tool in fluorescence imaging that can be widely applied in conjunction with a broad range of sensing dyes and experimental setups.

Introduction

Fluorescence sensing is an interdisciplinary field of research and technology with broad applications in life sciences, clinical research, diagnostics and control of industrial processes [1–2]. In particular, fluorescence imaging is exploited to unveil the dynamics of biologically relevant processes with high spatial and temporal resolution [3]. In this framework synthetic organic fluorophores with sensing abilities make it possible to measure *in vitro* and *in vivo* ion concentration [4–5], small molecules [6–7], environment polarity [8], membrane potential [9], and enzymatic activity [10–11]. They suffer from several unwanted properties, however, such as difficult targeting to the desired cell compartment or tissue, poor solubility, cell leakage. In addition they usually do not allow ratiometric imaging so that absolute measurements are not possible. In order to overcome these limitations we developed a nanostructure that can exploit the properties of interest of a given fluorescent indicator together with the added features stemming from other active molecules present on a dendrimer scaffold (e.g., targeting or calibration signals). We chose dendrimers as scaffolds owing to their many desirable properties. Dendrimers are hyperbranched nano-sized polymers that combine the advantageous features of nanoparticles (ideal size as *in vivo* carriers, multivalency), of polymeric materials (low cost, tunable properties, biocompatibility), and of small molecules (monodispersity and therefore detailed control on their properties) [12–15]. Some recent promising reports already presented some nanosized scaffolds functionalized with optical probes [14–16]. Yet a general (i.e. useful for a wide range of analytes, biological specimen and experimental setups) and effective architecture for optical nanosensors is still lacking, thus limiting their appeal for life scientists and their commercial interest.

In this work we propose a novel dendrimer-PEG copolymer-based architecture for fluorescence sensing. Molecular dynamics simulations and spectroscopy allow a complete structural comprehension of dendritic sensor structure and its relationship with dyes photophysical properties. The proposed structure showed a marked enhancement of the performances of the conjugated sensing fluorophores in comparison with previous reported dendritic structures [14].

Sensors for different biological relevant ions are reported; these sensors show good performances under different experimental setups and demonstrated their usefulness *in vitro*, in living cells and *in vivo*. We show that the novel properties conferred by the dendritic scaffold allow measurements that would be difficult or impossible with the free dyes. In particular we report the first extracellular pH imaging in the brain in a mouse epilepsy model.

Importantly, the molecular architecture shown here can be applied for a wide range of sensing dyes so that it represents a model for a family of dendrimer-based sensors of interest for other biologically relevant ions and small molecules.
Results and Discussion

Sensors synthesis and *in vitro* evaluation

The structures proposed here are schematically shown in Fig. 1a. Our indicators comprise a second generation polyamidoamine (PAMAM) dendrimer whose sixteen peripheral groups are functionalized with two sets of fluorophores on the periphery, namely sensing dyes and analyte-insensitive dyes for ratiometric correction. The conjugation of several copies of fluorophores on the same scaffold leads to signal enhancement. Polyethyleneglycol (PEG) spacers were introduced in order to tune the dendrimer-fluorophore and fluorophore-fluorophore distances in order to reduce quenching and other undesirable interactions often occurring in labeled dendrimers [17]. Sensors were obtained with three high-yield synthetic steps from commercially available reagents. A complete description of synthesis of the materials is reported in Figure S1. Characterization of the synthesized materials is proposed in Fig. S2, Fig. S3, Fig. S4 and Fig. S5.

In order to better understand the effects of the structure on the fluorescence properties of our sensors we synthesized three different architectures and compared them by optical spectroscopy and molecular dynamics (MD) simulations. Representative MD snapshots are shown in Fig. 1b. These architectures comprise

- Carboxy-Fluorescein (green) and Tetramethyl-Rhodamine (red) and differ for the presence or absence of PEG spacers. They consist of: i) dendrimer without PEG spacers (1); ii) dendrimer with fluorescein loaded onto 2 kDa PEG spacers, but rhodamine directly conjugated to the scaffold (2) and iii) dendrimer with both dyes loaded on PEG spacers (3). Fluorescein and rhodamine fluorescence quantum yields are reported in Fig. 1b and reveal that the presence of PEG spacers leads to an enhancement of sensor brightness of almost one order of magnitude. MD analysis shows that PEG spacers create a shell around the dendrimer core and play an important role in the structure. In fact they influence dye-to-dye distances, solvent accessibility and freedom of motion. Radial distribution functions and dye-to-dye distance were estimated by MD (Figure S6 and S7) and confirm that structure (3) is flexible and does keep a suitable distance between dyes, while in the absence of PEG spacers the structure is rigid and fluorophores are too closely packed. These observations are in good agreement with the measured quantum yield and suggest that structure (3) is the preferred architecture for the present dendrimer-based sensors. All the measurements in the followings are performed with this structure.

In order to explore the general applicability of this architecture we tested three different indicators for different biologically...
relevant ions. We conjugated carboxyfluorescein [18], methoxy-
quinoline [19] and PBFI [20] to our dendritic architecture
obtaining nanosensors for pH, Cl\(^-\) and K\(^+\), respectively. These
commercially available fluorescent indicators were loaded on the
nanostructure together with tetramethyl-rhodamine as internal
reference in order to perform ratiometric measurements (Figure S8).
The indicators were titrated in \textit{vivo} with the corresponding
analyte and the titration curves are reported in Fig. 1c. As can be
clearly observed all the sensing dyes (reported in green, blue and
yellow for pH, Cl\(^-\) and K\(^+\) respectively) readily respond to their
analyte while the red signal of the rhodamine is not affected in the
physiological range. As a consequence the ratio of the fluorescence
signal of the two dyes is not dependent on indicator concentration
and is determined only by analyte concentration. This insensitivity
to concentration in this ratiometric imaging is critical for practical
applications in the biological environment. Notably the sensing
dyes retain their photophysical properties upon conjugation to the
dendritic scaffold, in particular spectra (Figure S8) and affinity
constants are not significantly changed following conjugation.

**Living cells analysis**

The pH sensors were also tested in living Chinese Hamster
Ovary (CHO) cells. We focused our attention on pH sensing since
H\(^+\) ions are relevant in many physiological mechanisms and many
pH-sensing dyes were developed in order to monitor H\(^+\)
intracellular concentration. These measurements allowed us also to
demonstrate nanostructure delivery across the cell membrane
by electroporation. We recently proposed an optimized protocol
for dendrimer electroporation in living cells together with
strategies for intracellular targeting following cytoplasmic delivery
[14]. Figure 2a shows representative confocal images of the two
channels (fluorescein and rhodamine) and the corresponding
ratiometric map obtained dividing pixel-by-pixel the two images.
Importantly the measured pH value is stable upon prolonged
imaging (Figures S9 and S10), a very important feature in order to
reliably monitor time evolution. Notably no nanostructure
accumulation in specific organelles occurs. Often organic
fluorophores display significant protein binding owing to their
hydrophobicity with consequent change of Kd and cell compart-
mentalization; the hydrophilic PEG shell of the sensor avoids this
degradation of properties by protecting sensing dyes from undesired interactions. Moreover the scaffold prevents leakage of
the indicators out of the cell, a common drawback of membrane-
permeable fluorophores [21]. This dendritic structure is retained
inside living CHO cells up to four days (Figure S11) and no
leakage was observed during all time-intervals tested. These
measurements suggest the use of dendrimer-based structures for
long-term imaging and cell-tracking applications. We moreover
demonstrate the ability of our sensor to respond to intracellular pH
changes. CHO cells were clamped to desired pH values with nigericin and ratiometric imaging was performed yielding the
 calibration curve reported in Fig. 2b. Figure 2c shows the ratio
maps obtained for cells ranging from pH = 5 to pH = 7.5. These
images demonstrate that the indicator does respond to the
intracellular changes of pH with a change of the green-to-red
ratio. Therefore the ability of the free fluorescein to measure
changes around the physiological pH is retained but the
nanostructure yields (i) an accurate ratiometric correction, (ii)
localization the dyes in the cytoplasm with no compartmentaliza-
tion, and (iii) avoids cell leakage.

**In vivo pH imaging**

Finally, we used the pH indicator to detect dynamic pH changes
in the brain extracellular space in anesthetized mice. The
application of fluorescence imaging \textit{in vivo} is of great interest since
it makes possible to study biology in intact, native conditions. This
is known to be a challenging process for several reason: i) high
scattering and thus high background signal, ii) the complex
environment can interfere with sensor operation, iii) delivery and
targeting of the fluorescent probe are difficult, iv) spatial
distribution of the sensor is frequently inhomogeneous, v) sensor-
concentration normalization is required [22]. In particular
fluorescence sensing in the extracellular space is of great interest
as the brain matrix plays an important role in the regulation of
neural activity [23]. Notably most of the available fluorescence
indicators are not suitable for this \textit{in vivo} application owing to the
very fast drainage of the small dyes from the matrix.

The present pH indicator was injected in the brain cortex using a
micropipette and ratiometric imaging was performed as shown in
Fig. 3a. We performed two-photon fluorescence imaging while
simultaneously acquiring electrophysiological data. A bright signal
in both green and red channels was observed. The sensor was
localized in the extracellular space while no signal was observed
from blood vessels or cellular bodies (Figure S12).

Probably owing to its size, the sensor was retained in the
extracellular matrix and the signal remained stable for hours. We
stress that free fluorescein would have been removed very rapidly
and any measurement would have been impossible. Similarly to
living cells a ratio image was computed dividing pixel-by-pixel the
two colors signals. In order to demonstrate the ability of our sensor
to detect pH changes \textit{in vivo} we induced hypercapnia to the mouse
by controlled inhalation of CO\(_2\) since this phenomenon is known
to induce acidification of blood and tissues [24]. A time lapse pH
imaging during hypercapnia was performed and results are
displayed in Fig. 3b. The green-to-red ratio decreased after CO\(_2\)
administration indicating a drop in the pH value and it rapidly
reverted to the physiological value when standard oxygenation was
restored.

No spatial gradients were detected. This is expected since the
acidification kinetics is very slow in comparison to proton
diffusion. Note that the green signal alone would be completely
meaningless owing to the heterogeneous distribution of the sensor
and to focal plane changes during the measurement (hypercapnia
causes a strong vasoconstriction and therefore a significant
movement of the brain [25]). Figure S13 reports fluorescein,
rhodamine and ratio traces for three different experiments. After
the same stimulation the green curves are markedly different but
thanks to the ratiometric correction a reproducible response is
obtained, confirming the need for the internal reference in such \textit{in vivo}
extracellular imaging. To the best of our knowledge this is the
first example of two-photon brain pH imaging in living animals.

We finally performed a combined measurement of pH imaging
and electrophysiology in an animal model of epilepsy. Interictal
epileptic activity was induced by administration of the GABAA
antagonist bicuculline and ratiometric imaging and electrophys-
iology recordings are reported in Fig. 3c. After drug administra-
tion the highly synchronous activity characteristic of interictal
activity appears on the field trace [26] while no significant changes
in pH were recorded. Hypercapnia was induced and the
disappearing of neuronal activity correlated with a pH decrease.
The recovery of physiological pH value measured after air
ventilation coincides with the restart of interictal activity. This
result is consistent with a recent publication that highlighted the
relationship between brain extracellular pH values and epileptic
activity [27]. We would like to stress that this is a first example of
the relevance of the present nanostructure as element of combined
measurement protocols that could target the investigation of the
molecular fingerprints of a number of diverse pathological conditions.

Conclusion
A dendrimer-based indicator nanoarchitecture was presented that can overcome some of the more severe problems characteristic of the synthetic indicators currently available. Exploiting the unique chemical features of dendrimers we synthesized a monodisperse polymeric scaffold that is able to carry multiple sensing dyes together with other functional molecules. This represents a new paradigm in fluorescence sensing as virtually any small organic dye can be conjugated to the nanoscaffold and imaged under standard microscopy setups. Notably, this is not an alternative to the use of fluorescent dyes but a new and effective way to exploit their sensing properties. Successful applications in vitro, in living cells, and in vivo were presented. Most of these measurements would be difficult or impossible with standard methods and demonstrate the practical advantage of the present approach. We believe that the nanosystems presented in this work represent the prototypes of a novel and useful family of indicators optimized for biological imaging with a broad range of potential applications.

Materials and Methods

Materials
Carboxyfluorescein-NHS, Tetramethylrhodamine-NHS, PBFI were purchased from invitrogen. 6-Methoxyquinoline-N-sulfonyl acid was prepared according to the published procedures [19]. BOC-PEO₂k-NHS was purchased from Laysan bio. Ethylenediamine-core G₂ dendrimer and all the other reagents were obtained from Sigma Aldrich.

Synthesis and characterization
The synthesis of dendrimer-based sensors is schematically reported on Fig. S1. The general procedures are reported in the followings. In order to pegylate dendrimers 100 nmol of G₂ PAMAM were reacted with 20 molar equivalents (1.2 eq to dendrimer amines) of BOC-PEG₂k-NHS in DMSO at room temperature overnight. The solution was diluted with water and extensively dialyzed against water in order to remove the unreacted PEG and lyophilized. BOC deprotection was achieved dissolving the pegylated products in a 0.5% TFA solution in water and stirring for 48 h at room temperature. TFA and solvent were removed by freeze-drying. PEG conjugation and BOC deprotection were verified by NMR spectroscopy. Fluorescent labeling of dendrimer or PEG amines was achieved by means of N-hydroxysuccinimide (NHS) activated esters. When commercially available NHS-activated dyes were used. NHS-dyes were dissolved in DMSO together with the amines-bearing macromolecules and incubated under stirring overnight at room temperature. Notably in these conditions the reaction is quantitative and allows the control of the average number of dye per dendrimer. Otherwise non-activated dyes were conjugated with the following procedure. NHS (100 molar equivalents), EDC (100 molar equivalents), triethylamine (500 molar equivalents) and the sensing dye (100 molar equivalents) were added to rhodamine-labeled dendrimers (typically 50 nmol) in anhydrous DMSO and the solution was heated to 50°C overnight. The solution was cooled, diluted with water and extensively dialyzed against water.

Computational methods
All of the molecular dynamics (MD) simulations and data analysis were performed with the AMBER 11 suite of programs [28]. The molecular models for the sensors were constructed as composed by a central (CEN) unit and the repetitive branching units (REP) which constitute respectively the core and the scaffold of the G₂ PAMAM dendrimer, by the peripheral green (GRE) and red (RED) dye units and by the 2 kDa PEG chains composed by 44 monomers (PEG). All of the non-standard residues that compose the sensors were parametrized following a well-validated procedure for dendritic molecules [29]. All of the sensors were immersed in a periodic box extending 14 Å from the solute containing explicit TIP3P water molecules and the suitable amount of Na⁺ and Cl⁻ ions to ensure the neutrality of the system and the experimental salt concentration of 150 mM NaCl.
The MD simulation strategy was the same adopted in precedence by our group for the simulation of different dendrimers [30–31]. The MD runs lasted for 100 nanoseconds under NPT periodic boundary condition at 300 K and 1 atm of pressure. Such a long simulation time was necessary in order to allow the complete folding of the sensor molecules and to guarantee that an equilibrated configuration was successfully and reliably reached. The root mean square deviation (RMSD) data were obtained from the MD trajectories in order to verify that all of the systems converged to the equilibrium with good stability. The MD runs were carried out using the sander and pmemd.cuda modules within the AMBER 11 suite of programs working on NVIDIA Tesla 2050 GPU cards. All of the structural analyses were obtained by processing the equilibrated MD trajectories with the ptraj module within AMBER 11.

In vitro fluorescence measurements
Excitation and emission spectra were recorded by means of a Cary Eclipse fluorometer (Varian, Palo Alto, CA). The temperature of the cell compartment was controlled by a built-in Peltier cooler (Varian). Excitation and emission band-pass of 5 nm was employed. Typically, 500 μL samples were used in quartz cuvettes (Hellma, Milan, Italy). Titration of dendrimer-based sensors fluorescence was performed on 1 μM samples dissolved in water. Sensors were titrated with the corresponding analyte and emission spectra of the sensing dye and Rhodamine were obtained for each point.

Living cells studies
CHO cell lines were purchased from ATCC and cultured following manufacturer’s instructions. For live-cell microscopy cells were plated onto 35 mm glass-bottom dishes (WillCo-dish GWS-3522) and imaged at 37°C, 5% CO2. Electroporation was performed using Microporator (Digital Bio Technology, Korea) according to manufacturer instructions for CHO cells. After electroporation cells were centrifuged for 5 min at 1200 rpm in order to remove the excess of dendrimer in the medium. Cells were washed and fresh medium was added before imaging, usually 12–18 h after electroporation. Cell imaging was performed with a
Leica TCS SP2 inverted confocal microscope (Leica Microsystems) equipped with a 40×1.25 NA oil immersion objective (Leica Microsystems). Imaging was obtained illuminating the samples with the Ar and He-Ne lasers and with a 403 nm pulsed diode laser (M8905-01; Hamamatsu) at 50 MHz repetition rate. Fluorescence emission was collected with the microscope AOPS-based built-in detectors. Fluorescein and Rhodamine were imaged using the 488 nm (collection from 500 nm to 550 nm) and the 561 nm (collection from 575 nm to 700 nm) laser lines of the confocal system, respectively. To obtain a pH calibration of the sensor cells were clamped at the desired pH by using pH ionophores in a K+ enriched buffer [14].

In vivo imaging
All experimental procedures were authorized by the Italian Ministry of Health. Experiments were performed at 3–4 weeks of age on C57/Bl6j mice. Mice were anesthetized with urethane (1.6 gr/Kg) and ventilated with O₂ enriched air; a unilateral craniotomy above the visual cortex (centered 2 mm from the midline) was opened and the cortex surface constantly kept superfused with artificial cerebrospinal fluid (ACSF). A glass pipette containing an AgCl electrode was employed to inject in the cortex (150 μm depth, tip diameter 4 μm, pressure of 0.5 psi for 1–2 minutes) the dendrimer and to record extracellular field potentials. Extracellular field potentials were recorded employing an EXT-2F head-stage and extracellular amplifier (NPI) (gain of 1K, high pass filter 0.1 Hz low pass filter 300 Hz). In vivo two-photon imaging was performed with a Ultima Multiphoton Microscopy System (Prairie Technologies) equipped with a Mira 900 mode-locked laser source (Coherent). Fluorescein and rhodamine dyes were simultaneously excited with 820 nm 900 mode-locked laser source (Coherent). Fluorescein and rhodamine dyes were imaged using the 488 nm (collection from 500 nm to 550 nm) and the 561 nm (collection from 575 nm to 700 nm) laser lines of the confocal system, respectively. To obtain a pH calibration of the sensor cells were clamped at the desired pH by using pH ionophores in a K+ enriched buffer [14].

Supporting Information
Figure S1 Reactions scheme for the synthesis of the three sensor architectures. (a) Synthesis of structure 1: (i) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative. (b) Synthesis of structure 2: (i) , tetramethylrhodamine-NHS, DMSO, overnight rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, 98%. (iii) TFA 0.5% in water, 48 h rt 96%. (iii) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, quantitative. (b) Synthesis of structure 2: (i) , tetramethylrhodamine-NHS, DMSO, overnight rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, 98%. (iii) TFA 0.5% in water, 48 h rt 96%. (iii) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative. (b) Synthesis of structure 2: (i) , tetramethylrhodamine-NHS, DMSO, overnight rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, 98%. (iii) TFA 0.5% in water, 48 h rt 96%. (iii) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative. (b) Synthesis of structure 2: (i) , tetramethylrhodamine-NHS, DMSO, overnight rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, 98%. (iii) TFA 0.5% in water, 48 h rt 96%. (iii) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative. (b) Synthesis of structure 2: (i) , tetramethylrhodamine-NHS, DMSO, overnight rt, quantitative. (ii) BOC-PEG$_{2k}$-NHS, DMSO, overnight rt, 98%. (iii) TFA 0.5% in water, 48 h rt 96%. (iii) carboxyfluorescein-NHS, tetramethylrhodamine-NHS, DMSO rt, quantitative.

Figure S2 NMR spectra of G2-(Rh)$_{3}$-(PEG-BOC)$_{13}$. $^1$H-NMR spectra of G2-(Rh)$_{3}$-(PEG-BOC)$_{13}$. Structure scheme and peaks integration were reported.

Figure S3 NMR spectra of G2-(PEG-BOC)$_{16}$. $^1$H-NMR spectra of G2-(PEG-BOC)$_{16}$. Structure scheme and peaks integration were reported.

Figure S4 NMR spectra of G2-(PEG-NH$_2$)$_{16}$. NMR spectra of G2-(PEG-NH$_2$)$_{16}$. 96% BOC deprotection was calculated from peak integrals.

Figure S5 DLS of dendrimer and dendrimer-PEG compounds. Dynamic light scattering analysis of G2 and G2-(PEG-NH$_2$)$_{16}$.

Figure S6 Radial distribution functions (RDF). Radial distribution functions (RDF) of the surface dyes (green and red) of sensors reported in Fig. 1b with respect to the center of mass of the sensor obtained from the MD trajectories. The RDF plots give indications about the presence of the dyes atoms in a certain zone of the system (spatial density). However, since these curves are calculated at each step of the simulation, and they are reported in the plots as averaged over the equilibrated phases of the MD trajectories, they give information also on the dynamics of the system – they provide indication on the time period in which a certain atom is present in a certain area in the space (dynamic density). In RDF plots, high and narrow peaks in a small area of these graphs mean not only high density of atoms in a certain zone, but also high localization and low mobility of these atoms. On the other hand, broad and low intensity peaks indicate low density and high vibrations. It is worth noting that the presence of PEG spacers enhance the mobility of the connected dyes (low peaks for the green dyes in 2 and for both red and green dyes in 3). When the surface groups are directly connected to the surface of the PAMAM dendrimer, however, they are prevented from moving freely (high peaks in 1 and, for red dyes, in 2) due to higher surface crowding and to the absence of the flexible linker (PEG).

Figure S7 Dye-Dye distances. a) Snapshot of dendrimer (3) taken from the equilibrated MD trajectories. Within the sensor, the G2 PAMAM dendrimer (CEN and REP) is represented as blue spheres, the PEG linear chains in grey and the centers of mass of the RED and GRE dyes as red and green spheres. Hydrogen atoms, water molecules and Cl$^-$ and Na$^+$ ions are not represented for clarity. The distances between dyes below are calculated according to the scheme in the figure – taken one peripheral group, the distance from the other dyes is calculated with reference on the center of masses. This is done for each peripheral dye and averaged over the total number of surface group – the same procedure is done also for the (1) and (2) sensors. The table below is filled accordingly. b) Number of dyes at a certain distance d between each surface group of the sensor. The distance d is calculated with respect to the center of masses of the surface groups and is expressed in nm. Data evidence that presence of the PEG spacers decreases strongly the crowding of the sensor surface (higher distance between the dyes).

Figure S8 Structures and spectra of the dendrimers-based sensors. Schematic structure (top) of dendrimer-based sensors for pH, chloride ions and potassium ions. Spectra of sensing dyes and rhodamine references upon analyte titration (bottom).

Figure S9 Photobleaching stability. Time lapse imaging of CHO electroporated with G2-(PEG-Rh)$_{3}$-(PEG-Fluo)$_{15}$. No changes in ratio were observed during prolonged imaging.

Figure S10 Ratiometric correction. Ratiometric imaging of cells with different fluorescence intensities reveals the independence of the ratio value from sensor concentration.
Figure S11  Long retention time in living cells. Prolonged time lapse of CHO cells electroporated with dendrimer-based architecture (3). Cells have been imaged for four days in order to demonstrate that the dendrimer is retained inside cells and no leakage occurs. The intracellular signal decrease over time as cells undergo several cell cycles and the fluorescent dendrimer is divided between the post-mitotic cells. No leakage was observed during this period.

(TIF)

Figure S12  Sensor localization in vivo. Representative images of sensor localization after intracranial injection at different depths in the visual cortex.

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Figure S13  Ratiometric correction in vivo. In vivo pH sensor response to hypercapnia. Different acquisitions series show different pathway in green and red signal but same ratio response thanks to the ratiometric correction.

(TIF)

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Author Contributions

Conceived and designed the experiments: LA GMIR FB. Performed the experiments: LA MB GP SS BS. Analyzed the data: LA MB SS. Contributed reagents/materials/analysis tools: LA. Wrote the paper: LA FB.