Application of two-photon flash photolysis to reveal intercellular communication and intracellular Ca\textsuperscript{2+} movements

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Abstract. Two-photon excitation makes it possible to excite molecules in volumes of much less than 1 fl. In two-photon flash photolysis (TPFP) this property is used to release effector molecules from caged precursors with high three-dimensional resolution. We describe and examine the benefits of using TPFP in model solutions and in a number of cell systems to study their spatial and temporal properties. Using TPFP of caged fluorescein, we determined the free diffusion coefficient of fluorescein \((D = 4 \times 10^{-6} \text{ cm}^2/\text{s})\) at 20°C, which is in close agreement with published values. TPFP of caged fluorescein in lens tissue \textit{in situ} revealed spatial nonuniformities in intercellular fiber cell coupling by gap junctions. At the lens periphery, intercellular transport was predominantly directed along rows of cells, but was nearly isotropic further from the periphery. To test an algorithm aiming to reconstruct the Ca\textsuperscript{2+} release flux underlying physiological Ca\textsuperscript{2+} signals in heart muscle cells, TPFP of DM-Nitrophen was utilized to generate artificial microscopic Ca\textsuperscript{2+} signals with known underlying Ca\textsuperscript{2+} release flux. In an experiment with mouse oocytes, the recently developed Ca\textsuperscript{2+} cage dimethoxyxymethoxyethyl-ethylene glycol-bis-(β-aminoethyl ether)-N,N,N',N' tetraacetic acid-4 (DMNPE-4) was released in the oocyte cytosol and inside a nucleolus. Analysis of the resulting fluorescence changes suggested that the effective diffusion coefficient within the nucleolus was half of that in the cytosol. These experiments demonstrate the utility of TPFP as a novel tool for the optical study of biomedical systems. © 2003 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.1582468]

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1 Introduction

Conversion of precursors by photolytic chemistry (“flash photolysis”) is a relatively new tool for biomedical research. By synthesis of suitable precursors (“caged compounds”) it is possible to optically probe cell and tissue function with high spatiotemporal resolution.\textsuperscript{1,2} Using conventional UV illumination, the photoactivation of caged precursors can be accurately targeted within the plane of focus of the optical system, but significant uncaging also occurs in the cone of illumination above and below the focal plane. With the introduction of two-photon excitation\textsuperscript{3} it has now become possible to confine the flash photolysis to a diffraction-limited volume in all three dimensions (<1 fl).\textsuperscript{4,5} The unique ability of two-photon flash photolysis (TPFP) to produce a three-dimensionally resolved chemical source has allowed entirely new types of biological experiments. For example, the highly localized release of neurotransmitters by TPFP has been used to map the distribution of membrane receptors on neurons and other cells at high resolution.\textsuperscript{5,7} In cardiac muscle, TPFP of caged Ca\textsuperscript{2+} has recently been utilized to probe the subcellular Ca\textsuperscript{2+} dynamics in intact cells.\textsuperscript{8–10}

Conventional photoactivation of caged fluorochromes (using one-photon photolysis) has been used to measure diffusion coefficients and protein turnover in intact cells.\textsuperscript{11–13} TPFP of caged fluorochromes should provide the improved spatial resolution of two-photon excitation. In general, intercellular dye transport has been studied by introducing dye via patch pipettes (e.g., Refs. 14–16) or loaded from the bathing medium (e.g., Refs. 17 and 18). However, TPFP of caged dyes could provide a source whose dimensions are closer to a point, which would simplify the analysis and even open the possibility of studying transport in microscopic regions of the cell.

In this paper we use the high resolution provided by TPFP of caged fluorochromes to measure diffusion coefficients in solution and optically probe transport properties between fiber cells of the mammalian lens. In a novel application of TPFP of caged Ca\textsuperscript{2+} we demonstrate how this method can help reveal the temporal properties of microscopic Ca\textsuperscript{2+} release events (“Ca\textsuperscript{2+} sparks”) in cardiac muscle cells and can be used to probe spatial transport properties in different subcel-
lular compartments of mouse oocytes. These experiments demonstrate that TPFP makes it possible to study a variety of cellular processes at high spatial and temporal resolution.

2 Materials and Methods

2.1 Stationary Spot Two-Photon Flash Photolysis

Illumination for two-photon excitation was provided by a mode-locked titanium:sapphire (Ti:S) laser tuned to a wavelength of \(\sim 700\) and \(\sim 750\) nm for photolysis of caged Ca\(^{2+}\) and fluorescein, respectively. After passing it through a Pockels cell (Conoptics, Danbury, Connecticut), the beam was coupled into a modified confocal microscope (Zeiss LSM 410, Jena, Germany)\(^{19}\) to form a stationary spot parfocal with the confocal image plane but bypassing the scanner system. The photolysis volume (0.5 \(\mu\)m in plane and 1.4 \(\mu\)m axially\(^{19}\)) was determined by TPFP of caged fluorescein-dextran immobilized in a polyacrylamide gel. Fluorescence changes in response to stationary spot photolysis were monitored using the 488-nm line of an argon ion laser (Uniphase, San Jose, California) and the conventional confocal mode of the microscope. The pinhole was set to a diameter of \(\sim 1.5\) Airy units, resulting in a point spread function (PSF) with a full width at half maximum (FWHM) diameter of 400 nm in the plane and 900 nm in the axial direction. Accurate axial alignment between the photolysis spot (near-infrared light from the Ti:S laser) and the recording plane (488-nm light from the Ar\(^+\) laser) was ensured by adjustment of a lens in the Ti:S beam path to correct longitudinal chromatic aberration.\(^9\) Flash intensities were kept low (<1 mW at the sample) to prevent photobleaching of fluo-3 (or released fluorescein) and cage depletion at the site of photolysis (confirmed by the square dependence of released Ca\(^{2+}\)/fluorescein on flash intensity).

1. Uncaging of fluorescein in model solutions. Solutions for flash photolysis were prepared in a simple buffer containing 140 mM KCl and 20 mM N-2-hydroxyethyl piperazine-N’-2 ethane sulfonic acid (HEPES), pH 7.4. In this solution CMNB-caged fluorescein (Molecular Probes, Eugene, Oregon) was dissolved to a final concentration of 1 mM. The photolysis spot was focused \(\sim 50\) \(\mu\)m above the coverslip to avoid edge effects.

2. Uncaging of fluorescein in rat lenses. Lenses were dissected from adult rats into phosphate-buffered saline (PBS) at 37°C, transferred to an intracellular medium containing (in mM) MgCl\(_2\) 1, ethyleneglycol-bis-(\(\beta\)-aminoethylether)-N,N,N’,N’-tetraacetic acid (EGTA) 0.5, NaCl 10, Na₂ adenosine triphosphate (ATP) 2, KCl 20, K-glucosone 120, HEPES 10, pH 7.3, 300 mOsm/kg and cut through the equator using a fresh scalpel blade. Lens hemispheres were inspected for structural integrity using a dissecting microscope and transferred to a perfusion chamber sealed at the bottom with a coverslip so that the cut face of the lens rested on the coverslip. The chamber contained a solution of 1 mM CMNB-caged fluorescein (Molecular Probes) in an intracellular medium. The chamber was mounted on the two-photon/confocal microscope and caged fluorescein was released by stationary spot TPFP and monitored as described above. For x-y scanning the pixel dwell time was \(\sim 25\) \(\mu\)s and an average of two frames were recorded; during line scans the pixel dwell time was \(\sim 15\) \(\mu\)s and an average of four scan lines were recorded. Following recording of the fluorescence time course, the argon ion laser was used to image the local fiber cell structure in brightfield mode.

3. Fluorescent labeling of lens sections. Equatorial sections of adult rat lenses 10 \(\mu\)m thick were obtained as described previously.\(^20\) The sections were treated on slides for 1 h with blocking solution (3% w/v bovine serum albumin, BSA, 3% v/v fetal calf serum in PBS), washed three times for 5 min in PBS and treated for 2 h with either undiluted anti-MP70 (connexin\(\times 50\)) mouse monoclonal antiserum or with rabbit anti-connexin\(\times 46\) antibody (Alpha Diagnostic International, San Antonio, Texas) diluted 1:200 in blocking solution. The slides were washed three times for 5 min in PBS and treated for 1.5 h in the dark with antimouse (or antirabbit) Alexa Fluor 488 antibody, diluted 1:200 in blocking solution. After being washed three times for 5 min in PBS, the slides were labeled for 1.5 h in the dark with Alexa Fluor 350-conjugated wheat germ agglutinin (WGA, Molecular Probes; 40 \(\mu\)g/ml in PBS) to label fiber cell membranes.\(^21\) Finally, after washing them three times for 5 min in PBS, the slides were mounted in 10 \(\mu\)l Citifluor AF1 and stored at 4°C.

4. Generation of artificial model sparks. Artificial Ca\(^{2+}\) release events resembling Ca\(^{2+}\) sparks\(^22\) but with a known underlying release wave form were generated using TPFP of the Ca\(^{2+}\) caged DM-Nitrophen.\(^23\) Solutions for flash photolysis were prepared in a simple buffer (140 mM KCl and 20 mM HEPES, pH 7.4). To this solution 0.8 mM DM-Nitrophen (Calbiochem, Alexandria, Australia) and 100 mM Fluo-3 (Molecular Probes) were added. The free Ca\(^{2+}\) concentration was adjusted to 100 nM by adding CaCl\(_2\). Between 30 and 480 responses were averaged to improve the signal-to-noise ratio of line scan images of “artificial sparks.” Owing to the two-photon nature of the excitation process, the photolysis rate is expected to be proportional to the square of the flash intensity. To confirm the squared flash time course (for comparison with the reconstructed release time course), we also measured two-photon excited fluorescence in a dilute solution (100 \(\mu\)M) of fluorescein in response to flashes that were used to generate model sparks.

5. Imaging of rat cardiac myocytes. Enzymatically isolated rat cardiac myocytes were prepared from 250-g Wistar rats as described elsewhere.\(^24\) For line scan imaging, isolated ventricular myocytes were incubated with 5 \(\mu\)M of the AM form of fluo-4 (Molecular Probes) for 25 min at room temperature (20 to 22°C). The cells were then placed in a solution containing 2 mM CaCl\(_2\) and 10 to 20 \(\mu\)M of nifedipine (to allow identification of well-separated spark sites). The labeled cells were placed in the chamber of a confocal microscope (Zeiss LSM 410, Jena, Germany) with a glass coverslip as its bottom. Trains of action potentials were evoked by field stimulation at a frequency of 0.33 Hz to repeatedly activate spark sites. Indicator fluorescence was excited using the 488-nm line of an argon ion laser and detected between 510 and 550 nm (HQ filter, Chroma Technology, Brattleboro, Vermont). Fifty confocal lines were acquired during each field stimulation. The field stimulus was triggered after the first ten lines and the resulting Ca\(^{2+}\) transients were recorded during the remaining forty line scans. Line scanning was performed at the maximum scan speed of 1.4 ms/line. The total exposure time was therefore 70 ms/stimulus and trains of up to 100 stimuli could be recorded without detectable cell damage.
For comparison with experimental data, the photorelease and diffusion of caged fluorescein and caged Ca^{2+} were simulated using the measured PSFs of photorelease and confocal recording. The model equations were implemented using FACSIMILE (see also Refs. 24 and 25). For the modeling of artificial sparks, a diffusion coefficient of DM-Nitrophen (D_{DM}) in water was estimated as $4 \times 10^{-6}$ cm$^2$/s based on the molecular weight of DM-Nitrophen. On and off rates for DM-Nitrophen Ca^{2+} binding were $3 \times 10^7$ M$^{-1}$s$^{-1}$ and $0.144$ s$^{-1}$, respectively. The on and off rates for Ca^{2+} binding to fluo-3 were $2 \times 10^8$ M$^{-1}$s$^{-1}$ and $150$ s$^{-1}$, respectively, resulting in a $K_d$ of $0.75$ µM (similar to values used previously). For the full parameter set used to model sparks in intact heart cells, see Ref. 24.

7. Oocyte collection and microinjection. F1 hybrid mice of B6CB (C57Bl6JLac×CBA/CaLac) 21 to 28 days old were primed with an intraperitoneal injection of 7.5 IU pregnant mares' serum gonadotrophin (PMSG, Intervet, UK). At 44 to 52 h, their ovaries were punctured with a sterile needle and cumulus-enclosed oocytes were collected. The oocytes were mechanically stripped of their cumulus cells using a mouth pipette. The oocytes were microinjected as described previously with a solution containing fluo-3 pentapotassium salt (Molecular Probes) and the photolabile Cu^{2+} chelator dimethoxynitrophenyl-EGTA-4 (DMNPE-4) to final concentrations of 50 µM and 1 mM in the oocyte cytosol, respectively. Fluo-3 and DMNPE-4 were dissolved in 120 mM KCl, 20 mM HEPES, pH 7.5 for injection. To allow time for diffusion of dye within the oocytes, a period of at least 20 min from the time of microinjection was allowed and the oocytes were stored in M2 medium at 37°C for not more than 4 h. The injected oocytes had their zonae removed by a brief (10-s) incubation in acid Tyrode's solution. The oocytes were then immediately transferred to a heated chamber (Intracel) at 37°C containing 500 µl of M2 without bovine serum albumin. The oocytes were allowed to attach to the glass base in BSA-free M2 before the addition of 500 µl of M2 containing a fraction (14 mg/ml) V BSA.

8. Data processing. Image processing and analysis were performed using custom routines written in the IDL programming language (Research Systems, Boulder, Colorado).

3 Results
3.1 TPFP of Caged Fluorescein
At the cellular level, solutes are moved by diffusion, and most classical approaches for measuring diffusion (such as radioisotope tracers) have insufficient spatial or temporal resolution to measure diffusion coefficients with submicron resolution. Newer optical methods, such as fluorescence recovery after photobleaching (FRAP) and wide-field flash photolysis have increased spatial resolution but suffer from problems arising from the wide-field nature of the probe beam. Thus out-of-focus effects can contaminate the in-plane recordings of fluorescence change. Two-photon excitation provides a three-dimensionally resolved source of excitation which, in conjunction with a suitable photolabile compound, can provide a microscopic "point source" with which to probe local diffusive properties. To explore this idea further, we carried out tests using TPFP of caged fluorescein. A line scan image of the fluorescence changes in response to a 20-ms flash that was focused in a model solution containing 1 mM CMNB-caged fluorescein is shown in Fig. 1(a). The fluorescence rose rapidly after the onset of the flash and spread to a full width at half maximum of ~3 µm at the end of the flash. After the flash ended, the fluorescence rapidly declined to the background value, and ~15 ms after the end of the flash the fluorescence profile was well described by a Gaussian distribution [Fig. 1(b)]. Therefore, from this point in time forward, the solution of the diffusion equation for a point

Fig. 1 Flash photolysis of CMNB-caged fluorescein. (a) Line scan image of the fluorescence increase caused by a two-photon flash lasting 20 ms. The flash time course is indicated beside the line scan image. Scale bars: 5 µm, 20 ms. (b) Profile through the fluorescence distribution at the center of the photolysis volume. The theoretical curves were calculated by evaluating Eq. (1) at $x=0$ using diffusion coefficients $D=4 \times 10^{-6}$ cm$^2$/s, $4 \times 10^{-6}$ cm$^2$/s and $4 \times 10^{-7}$ cm$^2$/s.
source in three dimensions should be a good approximation of the recorded transient [with an appropriate choice of parameters; compare also Eq. (3.5) in Ref. 30]:

$$F(x,t) - F_0 = \frac{C}{D(t+t_0)^{3/2}} \exp\left[-\frac{x^2}{4D(t+t_0)}\right],$$  \(1\)

where \(F_0\) is the background fluorescence, \(D\) is the fluorescein diffusion coefficient, \(C\) is a constant that characterizes the amplitude of the distribution, and \(t_0\) is a time offset. For a given trial choice of \(D\), \(C\) and \(t_0\) were adjusted so that the profile shown in Fig. 1(b) was fit by the resulting Gaussian distribution. Essentially \(t_0\) depends only on the width of the profile and \(C\) is determined by the peak amplitude at the center of the profile shown in Fig. 1(b) (a value of \(~580\) resulted in a good fit that was largely independent of the choice of \(D\)). Using this procedure, the best fit of the decay time course of fluorescence decay [Fig. 1(c)] was obtained for \(D = 4 \times 10^{-6}\) cm\(^2\)/s, a value close to the reported diffusion coefficient of fluorescein in aqueous solution (\(5.1 \times 10^{-6}\) cm\(^2\)/s at 25°C).\(^{31}\) This datum suggests that TPFP of caged fluorescein can be used to obtain a measure of macromolecular diffusion at a microscopic distance.

### 3.2 TPFP of Caged Fluorescein in the Mammalian Lens

Molecular mobility also plays an important role in intercellular communication. Communication between cells can occur via specialized membrane channels of which gap junction (GJ) channels form a superfamily.\(^{32}\) These GJs allow low molecular-weight solutes to pass between cells and also allow electrical coupling. In the mammalian lens, GJ-mediated intercellular communication is important for maintaining the high transparency of this avascular organ.\(^{31}\) The lens is composed of many closely packed fiber cells that are connected by GJ “plaques” formed by connexin proteins. Mammalian fiber cells express two connexin isoforms, connexin46 (Cx46) and connexin50 (Cx50).\(^{34,35}\) To visualize GJ plaques, a longitudinal section of lens tissue was labeled with antibodies against Cx50 and the fluorescein-labeled membrane marker wheat germ agglutinin. Large GJ plaques on the broad sides of fiber cells are clearly visible in a longitudinal section through fiber cells [Fig. 2(a)]. Two-photon imaging of the stained section revealed that smaller plaques were also present in the bands of fluorescence between fiber cells. These gray bands represent the membrane on the narrow sides of fiber cells seen in cross-section. The close-packed arrangement of fiber cells and the location of GJ plaques in the cell membrane can be more clearly seen in transverse sections through the equator of the lens [Fig. 2(b)]. In this view of a region close to the lens periphery, fiber cells are discernible in cross-section. Note that the fiber cells are accurately aligned in rows and connected by large GJ plaques (stained with an antibody to Cx46) on the broad sides of fiber cells within these rows. Also visible in this section are smaller bright spots representing GJ plaques on the narrow sides of fiber cells that connect fiber cells in adjacent rows. In a tissue region deeper in the lens, the distribution of GJ plaques (stained with the same Cx46 antibody) was much less regular [Fig. 2(c)]. In this region cells were not as regularly organized into rows and the Cx46 staining was more uniformly distributed around the cell periphery.

From these structural data, one might expect regional variations in functional cell-to-cell coupling. To test this idea, we developed a method to measure diffusion from an identified cell using TPFP of caged fluorescein which, as shown earlier, should have sufficient spatial resolution to measure diffusion from the very narrow, but elongated, fiber cells. (Note that the “hourglass” wide-field illumination pattern would cause variable degrees of photolysis over a number of fiber cells that curve through the focal plane. Since this illumination pattern is convolved with the complicated 3-D cell geometry, quantitative analysis of such an experiment would be problematic.) To achieve TPFP of a fluorescent tracer whose movement could be monitored by confocal microscopy, we added CMNB-caged fluorescein to the intracellular medium whose movement could be monitored by confocal microscopy. We used CMNB-caged fluorescein to the intracellular solution that was used to bathe bisected (equatorial section) rat lenses. After allowing time for diffusion of caged fluorescein into the cut fiber cells, TPFP was used to liberate fluorescein inside a target cell by placing the stationary spot from
a Ti:S laser $\sim 40 \mu m$ from the cut surface. When the photolysis spot was placed in a peripheral fiber cell [corresponding to the region shown in Fig. 2(b)], the fluorescein spread rapidly into adjacent cells within the same row. This can be seen in the steady-state image of fluorescence taken $\sim 10 s$ after continuous photorelease was initiated [Fig. 2(d)]. Upon closer inspection, a small increase in fluorescence was also detected in cells adjacent to the row containing the chosen target cell. This probably reflects transport through gap junctions in the narrow sides of fiber cells. In contrast, when TPFP was performed in a region deeper within the lens [corresponding to the region shown in Fig. 2(e)], the pattern of fluorescence observed after $\sim 10 s$ did not indicate strong directed transport, as observed at the periphery. Rather, the fluorescence distribution was approximately isotropic around the target cell in which the fluorescein was released [Fig. 2(e)]. A smaller increase in fluorescence was also observed in some of the more distant cells. These results suggest that functional intercellular coupling is correlated with the local distribution of GJ plaques.

To study the time course of the increase in fluorescence in response to the uncaging of fluorescein at higher temporal resolution, a confocal line was scanned repeatedly across several cells in a region similar to that shown in Fig. 2(e). The resulting line scan image showed that the fluorescence increased rapidly in the cell in which the photolysis spot was located [Fig. 3(a)] and approached a steady level with a half time to peak of $\sim 200 ms$. The fluorescence in the cell immediately above this cell also increased quite rapidly (with a half time of $\sim 800 ms$) and ultimately reached a level of $\sim 50\%$ of that in cell 1. The fluorescence in the cells adjacent to these two cells approached a saturating level significantly more slowly (with a half time of $>2 s$) and the fluorescence increased to $<10\%$ of the maximal level reached in the target cell [Fig. 3(b)]. It is interesting that the maximal fluorescence level recorded in one of the cells (cell 4) adjacent to the bright target cell was smaller than that recorded in cell 3, which was next to the comparatively dimmer cell 2. These observations suggest that the coupling strength (i.e., permeability) may vary significantly from cell to cell.

### 3.3 TPFP of Caged $Ca^{2+}$ to Measure $Ca^{2+}$ Release Fluxes

The previous results demonstrate the ability of TPFP of caged compounds to measure microscopic transport. Such measurements could also provide a method by which to gauge the strength of a subcellular chemical source since the rate of release is determined by the illumination intensity and time course. Within muscle cells, microscopic $Ca^{2+}$ signaling events have been observed (called $Ca^{2+}$ sparks) that are elementary events underlying muscle activation. However, determination of the $Ca^{2+}$ flux underlying $Ca^{2+}$ sparks (which is due to $Ca^{2+}$ release channel gating) has previously relied on untested mathematical algorithms for flux reconstruction from confocal fluorescence images (see Ref. 24). An alternative approach is to parameter fit a suitable model to experimental data. To test this approach, we used TPFP to provide a controlled chemical source of $Ca^{2+}$ with spatiotemporal properties comparable in size to the source underlying $Ca^{2+}$ sparks. The model included $Ca^{2+}$ release, $Ca^{2+}$ and indicator diffusion, and $Ca^{2+}$ binding to $Ca^{2+}$ buffers (as well as optical blurring). The distribution of a $Ca^{2+}$ release flux in the model ($q_{Ca}$) was described by a function with six adjustable parameters:

$$\begin{align*}
q_{Ca}(r,t) &= \frac{I_0}{2F(\sqrt{\pi}\sigma)^3} \exp\left(-\frac{r^2}{\sigma^2}\right)f(t),
\end{align*}$$

where

$$f(t) = \frac{1}{1 + \exp\left(-\frac{t-t_1}{\tau_1}\right)} \frac{1}{1 + \exp\left(-\frac{t-t_2}{\tau_2}\right)}.$$

and $F$ is Faraday’s constant, $I_0$ is related to the amplitude of release, $\sigma$ describes the spatial extent of the source, and $t_1$, $t_2$, $\tau_1$, and $\tau_2$ characterize the time course of release. By appropriate choice of the temporal parameters, this relationship can be used to describe approximately rectangular shaped or exponentially rising and falling release time courses. In the fitting process the parameters $I_0$, $I_1$, $t_1$, $t_2$, $\tau_1$, and $\tau_2$ varied to achieve an optimal fit of the model to the recorded fluorescence data. The diffusion coefficients of the indicator dye, $Ca^{2+}$ and the $Ca^{2+}$ binding characteristics were set to...
values reported in the literature (see also Ref. 24). If the uncaging flash intensity is sufficiently small (so that local cage depletion is negligible), the square of the flash time course should be directly proportional to the Ca²⁺ release flux from the cage. A line scan image of an artificial spark generated by TPFP of the Ca²⁺ cage DM-Nitrophen in a small volume of test solution (see Sec. 2) is shown in Fig. 4(a). In this experiment, the flash time course had a rectangular shape and lasted for 10 ms. The fitting procedure resulted in a line scan image that was very similar to the "artificial spark" [Fig. 4(b)] both in time course [Fig. 4(c)] and spatial extent [Fig. 4(d)]. The model Ca²⁺ release flux was in good agreement with the flash photolysis time course [Fig. 4(e)], suggesting that our fitting method could provide an adequate description of the underlying Ca²⁺ release flux. Similar experiments with exponentially decaying flashes also showed good agreement between the reconstructed release time course and the square of the flash time course. Applying this algorithm to an average (n = 11 from one site) Ca²⁺ spark recorded from an isolated rat ventricular myocyte showed that Ca²⁺ release rises rapidly and decays approximately exponentially with a time constant of ~15 ms [Fig. 4(f)].

### 3.4 TPFP of Caged Ca²⁺ in Mouse Oocytes

It is well known that cells contain microscopic intracellular compartments such as the nucleus, and Ca²⁺ signaling within the nucleus has been suggested to be important in the determination of gene expression. To examine whether previous measurements of Ca²⁺ transport in the cytoplasm are also applicable to nuclear Ca²⁺ transport, we used TPFP of caged Ca²⁺ to probe nuclear Ca²⁺ diffusion. For these experiments we used DMNPE-4 injected into mammalian oocytes. Although similar to DM-Nitrophen, DMNPE-4 has a lower affinity for Mg²⁺, which makes it more suitable for experiments in the presence of physiological Mg²⁺ levels. Flashes with a rectangular time course lasting 60 ms were used to photolytically liberate Ca²⁺ in the oocyte cytoplasm and inside a nucleolus [see Fig. 5(a)]. Comparison of the fluorescence changes in response to these flashes suggests that the fluorescence signal from the Ca²⁺ indicator fluo-3 spreads less quickly inside the nucleolus [Fig. 5(b)] than in the cytoplasm. This point can be seen more clearly in Fig. 5(c), which compares the spatial profiles of the fluorescence distribution in the cytoplasm and the nucleolus toward the end of the photolysis flash. The time course of the spatial spread of the fluorescence response in these two compartments is summarized in Fig. 5(d). To test the idea that the faster spread in the cytoplasm could be explained by a larger effective diffusion coefficient in the cytoplasm (compared with the nucleolus), calculations using a simplified model of diffusion were performed. The model considered the movement of a diffusible substance that was released for 60 ms from the measured photolysis volume (see Sec. 2) and its movement was characterized by an effective diffusion coefficient D_{eff}. Note that the model did not attempt to capture the full complexity of the Ca²⁺ movements, binding, and indicator diffusion in the oocyte because the details of the Ca²⁺ buffering and indicator binding properties are not well characterized in mouse oocytes. The evol

**Fig. 4** Reconstruction of microscopic Ca²⁺ release fluxes. (a) Line scan image of an artificial Ca²⁺ spark generated by TPFP of the Ca²⁺ cage DM-Nitrophen using a flash that lasted 10 ms. The image shows the increase in fluo-3 fluorescence in response to the uncaging flash. Scale bar: 2 μm. (b) The best-fit simulated line scan image that was calculated using a model of Ca²⁺ and fluo-3 movements and binding. (c) In this graph the time courses of normalized fluorescence recorded at various locations of the artificial spark (dashed lines) are compared with the corresponding time courses of the "best fit" model calculation (solid lines). (d) Comparison of fluorescence profiles extracted from experimental data (dashed) and fit (solid). (e) This image shows the distribution of Ca²⁺ release flux underlying the simulation that best fit the experimental data. The graph next to the image compares the estimated time course of release (gray) with the flash time course. (f) The fitting approach validated using TPFP [(a) to (e)] was applied to an average Ca²⁺ spark recorded in an intact heart cell loaded with the indicator fluo-4. The average spark is shown next to the fitted spark. The time course of Ca²⁺ release underlying the fitted spark is shown on the right. Scale bars: 2 μm, 10 ms.
solution of the FWHM of the calculated fluorescence distribution was determined for a number of effective diffusion coefficients and compared with the experimental data. The spread of fluorescence in the cytosol was well described by the model when an effective diffusion coefficient of $5 \times 10^{-6} \text{ cm}^2/\text{s}$ was used. The observed time course of fluorescence spread in the nucleolus, on the other hand, was well fitted by a curve obtained using $D_{\text{eff}} = 2.5 \times 10^{-6} \text{ cm}^2/\text{s}$ in the model simulation, suggesting that the effective diffusion coefficient in the nucleolus was only half of that in the cytosol.

4 Discussion

In this study we demonstrated the general utility of TPFP, which was used to reveal intercellular communication of lens fiber cells and Ca$^{2+}$ movements within cardiac myocytes and mouse oocytes. In all of these experiments the unique three-dimensional spatial localization provided by TPFP was critical.

4.1 Measurement of Macromolecular Diffusion by TPFP of Caged Fluorescein

A previous study demonstrated the use of caged fluorochromes to measure diffusion coefficients in solution using conventional UV uncaging. Here, we show that corresponding measurements can be conducted using TPFP of caged fluorochromes (albeit with the higher spatial resolution provided by TPFP). The diffusion coefficient for fluorescein obtained using TPFP ($4 \times 10^{-6} \text{ cm}^2/\text{s}$) is in reasonable agreement with a value reported in the literature ($5.1 \times 10^{-6} \text{ cm}^2/\text{s}$ at 25°C). The measurement in this study was performed in an air-conditioned room at a temperature of 20°C. Assuming a $Q_{10}$ of diffusion of 1.35, the value from Ref. 31 corresponds to $D = 4.4 \times 10^{-6} \text{ cm}^2/\text{s}$ at 20°C and is in good agreement with our result. In applying Eq. (1) to our data, we assumed that the fluorescein distribution was approximately spherical. Although the photolysis volume was not spherical (but rather an ellipsoidal shape, see Sec. 2) the decaying distribution will rapidly approach spherical symmetry, owing to diffusion. This was also confirmed by mathematical modeling (not shown), which indicated that deviations from spherical symmetry were <3% at times $>15$ ms after cessation of the photolysis flash. In fact, waiting for the diffusion profile from the photolysis spot to assume spherical symmetry is advantageous because it simplifies subsequent analysis. On the other hand, the source size is increased by this procedure, but the relative increase is not large compared with the size of the source produced by wide-field photolysis. Measurement of diffusion coefficients by TPFP provides an attractive alternative to other multiphoton techniques capable of quantifying diffusion [e.g., multiphoton fluorescence photobleaching recovery (MP-FPR) and fluorescence correlation spectroscopy (FCS)]. Compared with MP-FPR, it has the benefit that a bright signal is detected against a dark background (which should improve the signal-to-noise ratio) and data acquisition and analysis should be simpler than in equivalent FCS approaches. A potential limitation of our method is the commercial availability of appropriate caged compounds. Currently these are caged HPTS, caged fluorescein derivatives, and dextran-fluorescein. On the other hand, a succinimidyl ester of caged carboxyfluorescein is available that can be used to link the fluorochrome to amine groups on other macromolecules, extending the feasibility of our approach to a host of other applications.
4.2 TPFP of Caged Fluorescein to Probe Intercellular Coupling in Situ

The data presented in this study show that TPFP of caged fluorescein allows intercellular coupling to be measured in situ with high spatial resolution. Measurement of dye transfer is one of the primary tools used to study gap junction function and complements electrical measurements that provide information about ion movement. In isolated and cultured cell systems, the tracer dye is frequently introduced via a patch or intracellular pipette (e.g., Refs. 14–16). Alternatively, the dye can be introduced by scrape loading via the bathing medium.\textsuperscript{17,18,42} While this method can be used in whole tissues, the spatial resolution is comparatively low and generally limits this approach to studying average coupling properties in a whole tissue region.\textsuperscript{17} In comparison, the ability to produce a spatially confined source by TPFP provides not only local measurements of transport properties but also the ability to carry out multiple independent tests within the same preparation. Thus, it would be possible to measure the effect of an intervention on transport without having to physically wash out the tracer probe between interventions.

The data presented in Fig. 2 indicate that intercellular coupling changes qualitatively and quantitatively throughout the lens. While the coupling at the periphery was strongly directional, intercellular coupling deeper within the lens was approximately isotropic. The patterns of dye movement observed in a peripheral and a more central region within the lens were correlated with changes in the local distribution of gap junctions that were visualized by two-photon immunofluorescence microscopy. A more detailed analysis of our data indicates that there are also variations in intercellular coupling from cell to cell. This is suggested by the observation that the dye levels reached in cells adjacent to the target cell (which should be a function of local coupling strength) varied from cell to cell even though the membrane area in contact was comparable (cf. Fig. 2 and Fig. 3). Such variations may be due to differences in plaque density between fiber cells or different gap junction channel opening probabilities. Future experiments using the methods presented here should be able to examine these possibilities. Furthermore, it should be possible to extend the analysis of such dye transport data generated by TPFP-released fluorochromes to obtain quantitative estimates of local membrane permeabilities. Similar types of analysis have already been used in conventional dye transport studies, e.g., Refs. 17, 18, and 43. Using the technique described here, the analysis should benefit from the increased spatial resolution provided by TPFP.

An alternative approach to probe GJ permeability is to utilize fluorescence recovery after photobleaching (e.g., Ref. 43). After the fluorochrome is selectively bleached in a small region of tissue, the observed rate of fluorescence recovery should be a function of the intercellular coupling to cells that were not exposed to the bleaching illumination (which serve as a reservoir of dye). In principle, it should be possible to use two-photon excitation to selectively bleach the dye within a target cell and achieve local resolution similar to that with the uncaging approach. However, as noted earlier, the reduced contrast associated with FRAP methods decreases the signal-to-noise ratio. For example, in experiments using conventional one-photon illumination, the directed transport of tubulin was detected using caged fluorochromes,\textsuperscript{44} but could not be resolved in FRAP experiments.\textsuperscript{45}

4.3 TPFP of Caged Ca\textsuperscript{2+} to Measure Microscopic Ca\textsuperscript{2+} Release Fluxes

To advance our understanding of the microscopic basis of Ca\textsuperscript{2+} sparks (and similar local signals), it is desirable to obtain a direct measure of the Ca\textsuperscript{2+} release flux underlying the observed fluorescence signal. Several algorithms have been proposed in the literature to reconstruct the release flux from the fluorescence record.\textsuperscript{46,47} To overcome the noise limitations associated with such approaches, we have implemented a novel fitting approach in which a numerical model of the Ca\textsuperscript{2+} movements is fitted to the experimental data.\textsuperscript{24} This fitting approach is less noise sensitive than other methods (in which high-order numerical derivatives of noisy experimental data have to be calculated) and also provides measures of parameter sensitivity. A common shortcoming of all approaches (including our own) had been the lack of an experimental test that could verify the ability of these algorithms to provide a reasonable estimate of the Ca\textsuperscript{2+} release flux. To test our algorithm, we reconstructed the known time course of Ca\textsuperscript{2+} release produced by TPFP of DM-Nitrophen. This method not only produced events that were comparable in magnitude and time course to real Ca\textsuperscript{2+} sparks but also incorporated limitations of the recording systems in a way that would not easily be produced by purely synthetic data. By using low illumination intensities, we ensured that saturation effects were avoided so that the rate of uncaging was proportional to the square of the instantaneous illumination intensity. This point can be confirmed by increasing illumination power and examining the power dependence of total yield.\textsuperscript{1} To our knowledge, this is the first time that reconstruction algorithms have been so accurately tested.

Our results show that we can obtain a reasonable description of the time course of Ca\textsuperscript{2+} release on a time scale of milliseconds with a spatial resolution of \( \sim 1 \mu m \). From the reconstructed time course we can estimate that the total release current is \( >7 \) pA,\textsuperscript{24} which suggests that at least fifteen ryanodine receptors participate in generating Ca\textsuperscript{2+} sparks. Put another way, during a Ca\textsuperscript{2+} spark, \( \sim 200,000 \) Ca\textsuperscript{2+} ions are released. The kinetic features of Ca\textsuperscript{2+} release should be reproduced in models for cardiac excitation-contraction coupling and must reflect the intrinsic gating of intracellular Ca\textsuperscript{2+} release channels as well as the strength of the Ca\textsuperscript{2+} source. Our data and other studies\textsuperscript{8,9,39} suggest that TPFP of caged Ca\textsuperscript{2+} will be one of the key techniques to further clarify the microscopic basis and regulation of local Ca\textsuperscript{2+} signaling.

4.4 Probing Transport Properties in Mammalian Oocytes by TPFP of Caged Ca\textsuperscript{2+}

In experiments with mouse oocytes we have used DMNPE-4 to release Ca\textsuperscript{2+} at different points within the cell. DMNPE-4 (\( K_D \approx 19 \) nM for Ca\textsuperscript{2+} at pH 7.4) undergoes a \( \sim 21,000 \)-fold change in Ca\textsuperscript{2+} affinity upon photolysis and has a comparatively low affinity for Mg\textsuperscript{2+} (\( K_D \approx 7 \) mM).\textsuperscript{39} We achieved an approximately twofold increase in fluorescence (Fig. 4) during the peak of the transient at illumination powers that were significantly below intensities that caused visible photodam-
age. Assuming a Ca$_{2+}$ binding affinity of fluo-3 of $\sim 0.75$ $\mu$M inside the cell$^{24}$ and a resting [Ca$_{2+}$] of 100 nM, it can be calculated (see Ref. 22) that photolysis resulted in an increase of the free Ca$_{2+}$ concentration to $\sim 230$ nM within the cell. When the photolysis spot was placed inside a nucleolus, a fluorescence transient of similar amplitude was observed. However, the fluorescence spread less quickly than in the cytoplasm, which is presumably due to a reduced local diffusion coefficient. Although the spread of the fluorescence signal should be dominated by fluo-3 diffusion, the presence of a larger proportion of essentially immobile Ca$_{2+}$ buffers could also contribute by lowering the effective Ca$_{2+}$ diffusion coefficient.$^{48}$ Regardless of the origin of the changes in mobility between cytoplasm and nucleolus, it should be noted that the experimental detection of such subcellular heterogeneities between cellular compartments only becomes possible with the high spatial localization provided by TPFP. Such studies have potential applications in characterizing processes that are dependent on subcellular compartmentalization such as biochemical pathway kinetics and intracellular signaling.

5 Conclusions

Using TPFP we have conducted proof of principle initial experiments in a variety of cell systems that seek to clarify microscopic transport properties. We have shown that 3-D localized uncaging using TPFP provides a unique tool for studying significant biological problems. Using this method, we have shown spatial anisotropy in intercellular transport within the lens and tested a novel method for calculating Ca$_{2+}$ release fluxes. We also show that the microscopic transport properties within a small organelle (the nucleolus) can be different from those observed in the cytosol. Further developments of the methodology presented here should provide new insight into cellular processes at an unprecedented microscopic scale.

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