Testing Fluorescence Lifetime Standards using Two-Photon Excitation and Time-Domain Instrumentation: Fluorescein, Quinine Sulfate and Green Fluorescent Protein

Arne S. Kristoffersen 1 · Svein R. Erga 2 · Børge Hamre 1 · Øyvind Frette 1

Received: 15 June 2018 / Accepted: 20 July 2018 / Published online: 26 July 2018 © The Author(s) 2018

Abstract
It is essential for everyone working with experimental science to be certain that their instruments produce reliable results, and for fluorescence lifetime experiments, information about fluorescence lifetime standards is crucial. A large part of the literature on lifetime standards dates back to the 1970s and 1980s, and the use of newer and faster measuring devices may deem these results unreliable. We have tested the three commonly used fluorophores fluorescein, quinine sulfate and green fluorescent protein for their suitability to serve as lifetime standards, especially to be used with two-photon excitation measurements in the time-domain. We measured absorption and emission spectra for the fluorophores to determine optimal wavelengths to use for excitation and detector settings. Fluorescence lifetimes were measured for different concentrations, ranging from $10^{-3}$ to $10^{-5}$ M, as well as for various solvents. Fluorescein was soluble in both ethanol, methanol and sulfuric acid, while quinine sulfate was only soluble in sulfuric acid. Green fluorescent protein was prepared in a commercial Tris-HCl, EDTA solution, and all three fluorophores produced stable lifetime results with low uncertainties. No significant variation with concentration was measured for any of the fluorophores, and all showed single-exponential decays. All lifetime measurements were carried out using two-photon excitation and lifetime data was obtained in the time-domain using time-correlated single-photon counting.

Keywords Fluorescence · Lifetime · Fluorescein · Quinine sulfate · Green fluorescent protein · Two-photon · Time-domain

Introduction

When a fluorophore has absorbed the necessary energy to become excited, a number of processes, e.g. fluorescence, intersystem crossing, resonance energy transfer and internal conversion, can occur - each with a certain probability, characterised by the decay rate constants $k_r$ (radiative) and $k_{nr}$ (non-radiative), and the fluorescence lifetime $\tau$ of a fluorophore is the average time between its excitation and its return to the ground state:

$$\tau = \frac{1}{k_r + k_{nr}}.$$

The fluorescence lifetime is also sometimes defined as the time where the fluorescence intensity of an excited fluorophore population has decreased to $1/e$ of its initial value, which is mathematically equivalent to the average time excited.

The fluorescence lifetime is an excellent tool for investigating the dynamics of excited states and molecular interactions. For instance, a change in the lifetime for two separate fluorophore states will indicate a change in one or more of the quenching parameters described by the non-radiative decay rate constant $k_{nr}$. Fluorophores with a reliable and well established fluorescence lifetime - a fluorescence lifetime standard - are useful for anyone performing measurements within this field, whether it is for calibration purposes, testing for systematic errors, or simply to verify and compare own lifetime measurements to the standard. It is also important to have information about the types of fluorophores that are suitable for a given instrumentation setup. Especially useful is absorption and emission spectra, as well as the lifetime itself, but also it...
can be important to have information about which solvents and concentrations that give the most stable and reliable results. In this paper, we have selected three fluorophores and tested their suitability to serve as fluorescence lifetime standards. In our experiments, fluorescence lifetime data were obtained using time-correlated-single-photon-counting (TCSPC). We used a laser producing pulses in the femtosecond regime, tuneable in the wavelength range 690-1040 nm, for two-photon excitation, and a detector with a range of 400-800 nm. Naturally, the fluorophores were selected to have their absorption within the range of these wavelengths, meaning that for two-photon excitation the absorption should be roughly in the range of 345-520 nm. Other selection criteria were that the fluorophores should exhibit a single-exponential decay, be readily commercially available and also have a fluorescence lifetime between 0.1-10 ns. The upper boundary for the lifetime is both to avoid a shortening from oxygen quenching, but also to avoid a pile-up effect since the repetition rate of the laser was 80 MHz, or 12.5 ns between each pulse. In addition to these criteria, the selection of fluorescein, quinine sulfate and green fluorescent protein (GFP) were also based on a quantitative literature search comprising the past five years. The present three fluorophores were amongst those with the highest number of hits in publications dealing with fluorescence and fluorescence lifetime research.

The goal of this paper is mainly to provide useful information about the present selection of fluorophores in terms of their suitability as fluorescence lifetime standards, and also gaining additional information about their properties with respect to existing literature, especially concerning two-photon excitation and lifetimes acquired in the time-domain. This instrumentation has become increasingly popular the last 10-15 years, especially in regards to working with living or fragile samples. The present study is also building on previous work with the same setup [21].

The first records of fluorescence lifetime measurements were made in the 1960s and 1970s [6, 7], however lifetime measurements from this time period were usually performed using pulse sampling oscilloscope techniques, which are not considered accurate by today’s standards. Since the late 1970s, with the introduction of high repetition rate lasers, several contributions have been made in determining lifetime standards [14, 20, 23, 33, 36]. However, many of the proposed standards have too long lifetimes to benefit today’s pico- and femtosecond instrumentation, and several were later found unsuitable due to double-exponential decays. Also, no systematic approach was done to evaluate the reliability of these lifetimes until 2007 [9], when comparisons of 13 different fluorophores were systematically examined by 9 separate laboratories in a very thorough and comprehensive work, although fluorescence was not induced by two-photon excitation in any of these measurements.

Fluorescein is available in a number of derivatives, and is widely used particularly because of a strong fluorescence signal, suitable absorption and emission wavelengths, as well as a number of other photophysical properties.

In 2014, a broad and extensive work on 20 fluorescein derivatives, which measured fluorescence lifetimes, quantum yields and emission maxima in ethanol and phosphate-buffered saline (PBS) was published [41]. There have also been several papers published in recent years, reporting on the basic fluorescent properties of various fluorescein dyes [8, 25, 27, 32, 40].

Quinine sulfate has been used as a fluorescence standard for more than 50 years [11, 13] and the photophysical properties of various quinine sulfate derivatives have also been extensively studied in the past [4, 17, 18, 26, 29]. However, fluorescence lifetime measurements are scarce after 1990.

In the 1990s, GFP from the jellyfish *Aequorea victoria* became one of the most widely studied and exploited molecules in biochemistry and cell biology [37], especially due to its unprecedented ability to serve as a marker of gene expression and protein targeting in intact cells and organisms [10, 31]. Fluorescence lifetime measurements have also been carried out [1, 30, 34, 35], however, both instrumentation and other measurement parameters differ quite significantly, and thereby also the reported fluorescence lifetimes.

A common denominator for all three fluorophores is that they are extensively used in a vast array of scientific areas, and that recent fluorescence lifetime measurements, especially using two-photon excitation and time-domain instrumentation is inadequately available in literature.

**Materials and Methods**

All three fluorophores were commercially available and purchased from Sigma Aldrich. Fluorescein for fluorescence, free acid (46955 SIGMA), Quinine sulfate, United States Pharmacopeia (USP) reference standard (1597005 USP), and GFP from jellyfish *Aequorea victoria*, recombinant expressed in *E. coli* (11814524001 ROCHE). The structural formulas of fluorescein and quinine sulfate are shown in Fig. 1. GFP is composed of 238 amino acid residues, and its structural formula is not shown here (see [37] for more information about the structure and composition of GFP). Ethanol and methanol of spectrophotometric grade, as well as 1 M sulfuric acid (H₂SO₄) were used as solvents for fluorescein, while quinine sulfate was only dissolved in H₂SO₄. The GFP was purchased and used in a solution form of
1 mg/ml in 5 mM Tris-HCl, 5 mM EDTA, pH 8.0. Fluorescein and quinine sulfate were prepared in concentrations of $10^{-3}$ M, $10^{-4}$ M and $10^{-5}$ M for all solvents. Samples were prepared by placing a droplet onto a microscope slide and covered by a 0.17 mm thick glass. All measurements were performed in room temperature (20°C).

**Instrumentation** A Ti:Sapphire laser (Coherent Chameleon Ultra), generating femtosecond pulses (pulse width 140 fs) at an 80 MHz repetition rate (12.5 ns between each pulse) was used for two-photon excitation of the samples. The laser is tunable in the wavelength range 690-1040 nm, with an average output power of 4 W at 800 nm. An electro-optical modulator (EOM) controlled the intensity of the laser, which in turn was guided by mirrors into a confocal inverted microscope (Leica TCS SP5) and focused by a water immersion $63\times$ objective with a numerical aperture NA = 1.2 and a working distance of 0.22 mm. The sample was scanned at a line frequency of 400 Hz and fluorescence was detected by a built-in photomultiplier tube (PMT) detector with a range of 400-800 nm. Line, frame, and pixel clock signals were generated and synchronized by another PMT detector (Hamamatsu R3310-02) and linked via a time-correlated single-photon counting (TCSPC) imaging module (SPC-830, Becker-Hickl, Berlin, Germany) to generate fluorescence lifetime raw data. Absorption spectra were measured using a spectrophotometer (Shimadzu UV-2450) and data was collected using the software UVProbe (v.2.42). Fluorescence emission spectra were measured using the lambda scan feature in the Leica LAS software.

**Absorption and Emission Measurements** All samples were diluted to a concentration of $10^{-5}$ M prior to both absorption and emission spectrum measurements. Before beginning the absorption spectrum measurements, a sample of distilled water was used to determine a baseline measurement with a very low absorption. The samples were placed in a cuvette and inserted in the spectrophotometer, which scanned in 1 nm steps from 200-700 nm. The lambda scan performs a fluorescence intensity measurement step-wise through a pre-determined detector wavelength interval. Since two-photon excitation is a non-linear process [42], the absorption spectra were used to help find a suitable excitation wavelength. For fluorescein, we used an excitation wavelength of 900 nm and a detector range of 440-700 nm. For quinine sulfate, excitation was at 800 nm and detector range set to 400-600 nm. For GFP, excitation was also at 800 nm, while the detector range was set to 450-590 nm. For all emission measurements the detector bandwidth was set to 30 nm and step size to 3 nm to achieve both a strong fluorescence signal and a smooth spectral curve.

**Fluorescence Lifetime Measurements** The spatial resolution of the fluorescence lifetime measurements were $128\times128$ pixels in order to achieve a good time resolution of 256 channels. Fine-tuning of the distance between sample and objective was necessary for each measurement to find focus, easily identified using a live scan showing the fluorescence intensity in real time. Each measurement was started by first initiating a live mode in the Leica LAS software, making sure that the sample was in focus and then starting the TCSPC module measurement, with an acquisition time of at least 1 minute, or until at least 10000 photons were collected for the brightest pixel. The laser intensity was adjusted to achieve a sampling rate of around $10^9$ photons per second. A higher sampling rate increases the risk of the so-called pile-up effect [5], where more than one photon is collected between two laser pulses. To ensure statistical significance, each sample was measured three times at different locations within the sample, then a new droplet of the same fluorophore was measured in the same way two times, so that each fluorophore-solvent-concentration combination was measured nine times. Lifetime values and uncertainties were calculated as an average of all nine measurements.

**Data Analysis** After collecting the fluorescence lifetime data with the TCSPC module, the software SPCLimage (Becker-Hickl, Berlin, Germany) was used to calculate the decay matrix. The collected photons for each pixel are stored as a histogram, and a decay curve is fitted to this data to calculate the fluorescence lifetime. A $\chi^2$ value indicates how well the fit is (1 indicates a best fit), and the matrix contains the calculated lifetimes for all the pixels in the image. Both lifetime data and the intensity (number of photons collected per pixel) was exported and further dealt with in MatLab® to make plots, calculate mean lifetimes and standard deviations. Figure 2 shows the lifetime images.
Fig. 2  The upper left shows the fluorescence intensity for quinine sulfate dissolved in sulfuric acid at 128x128 pixels, the lifetime image next to it is color-coded from the curve in the top right, which shows the distribution of lifetimes for the whole image. The lower part of the figure shows the exponential decay from detected photons for a single pixel in the image indicate by the blue cross-hair. The green curve is the instrument response function, and the $\chi^2$ is 1.14, indicating a very good fit for a single-exponential decay.

Results and Discussion

Absorption and Emission  The main reason to measure absorption and emission spectra for the fluorophores was to be able to make well-founded decisions in choosing excitation wavelengths as well as setting detector intervals for fluorescence. All fluorophores were diluted to $10^{-5}$ M concentrations for both absorption and emission measurements, to avoid possible aggregating of molecules as well as still having a sufficient fluorescence signal strength. It should be noted that the emission spectra were not corrected in regards to detector sensitivity variation at different wavelengths, which may cause a slight shift in emission peaks. However, this was not crucial to our measurements since the detector range was relatively broad for all samples. Figures 3, 4 and 5 show the absorption and emission spectra of fluorescein dissolved in ethanol, methanol and sulfuric acid, respectively. The spectra for ethanol and methanol are very similar for both absorption and emission, while dissolved in sulfuric acid, the spectra are shifted towards the blue end of the visible spectrum by 20-30 nm. Also, there are two absorption peaks for ethanol and methanol, compared to one for the sulfuric acid solution. Multiple absorption (or emission) peaks are in general attributed to vibronic transitions, which describes simultaneous changes in electronic and vibrational energy levels of a molecule [12, 22].

Since there is a large range of different fluorescein derivates in use [41], the absorption and emission spectra also vary substantially. Our measurements are, however, in good accordance with recently published results of free acid type fluorescein [2, 15, 19], with two characteristic absorption peaks around 450 and 480 nm, and an emission peak around 520 nm for both ethanol and methanol solutions. The present work is to our knowledge the first published on fluorescein absorption and fluorescence...
**Fig. 3** Absorption and emission spectrum for a $10^{-5}$ M concentration of fluorescein in ethanol. $\lambda_{\text{abs}}$ indicates both peaks in the absorption spectrum. The blue curve represents absorption and the red curve represents emission. Both curves are normalised to unity.

**Fluorescein in ethanol**

$\lambda_{\text{abs}} = 457$ nm, 486 nm

$\lambda_{\text{em}} = 523$ nm

**Fig. 4** Absorption and emission spectrum for a $10^{-5}$ M concentration of fluorescein in methanol. $\lambda_{\text{abs}}$ indicates both peaks in the absorption spectrum. The blue curve represents absorption and the red curve represents emission. Both curves are normalised to unity.

**Fluorescein in methanol**

$\lambda_{\text{abs}} = 454$ nm, 482 nm

$\lambda_{\text{em}} = 519$ nm

**Fig. 5** Absorption and emission spectrum for a $10^{-5}$ M concentration of fluorescein in 1 M sulfuric acid. The blue curve represents absorption and the red curve represents emission. Both curves are normalised to unity.

**Fluorescein in sulfuric acid**

$\lambda_{\text{abs}} = 432$ nm

$\lambda_{\text{em}} = 492$ nm
properties in a sulfuric acid solution. We also note that fluorescein did not completely dissolve in distilled water, and therefore we do not include any measurement results using this solvent.

Quinine sulfate does not completely dissolve in water, ethanol or methanol, and is according to literature almost exclusively dissolved in sulfuric acid. Figure 6 shows the absorption and emission spectra for quinine sulfate dissolved in sulfuric acid. Absorption peaks at around 320 nm and 350 nm are in excellent agreement with both recent and earlier results [13, 16, 39]. We do, however, report a slightly red-shifted fluorescence emission maximum, at 481 nm, compared to typically around 450 nm found in literature [16, 39]. The emission peak in Fig. 5 is broader compared to those reported in literature, although the intensity at 450 nm is still quite high, at about 80% of the peak value.

The absorption and emission spectra for the (GFP) is shown in Fig. 7. Both the absorption peak at 397 nm and the emission at 507 nm are in excellent agreement with previous measurements of rGFP [10], and wild type [37], who reported 395 nm and 509 nm, and 395-397 nm and 504 nm, respectively, although they also reported a minor absorption peak at 540 nm and an emission shoulder at 540 nm, which were not pronounced in our results.

**Fluorescence Lifetimes** Fluorescein fluorescence lifetimes were measured in concentrations of $10^{-3}$ M, $10^{-4}$ M and $10^{-5}$ M in solutions of ethanol, methanol and sulfuric acid. Quinine sulfate lifetimes were measured within the same concentration range, but only dissolved in sulfuric acid as it did not dissolve satisfactory in ethanol and methanol. We also attempted to dissolve both fluorescein and quinine sulfate in distilled water, however, none dissolved properly and since the following lifetime measurements were highly variable, the results are not included here. (GFP) fluorescence lifetimes were measured only for the original
The fluorescence lifetimes of quinine sulfate dissolved in sulfuric acid for 10⁻³ M, 10⁻⁴ M and 10⁻⁵ M concentrations are shown in Table 2. The lifetimes increase only slightly with dilution, although the uncertainty also increases by a similar value. All three conditions showed single-exponential decays, which is remarkable because several earlier works dating 25-35 years back deemed quinine sulfate unsuitable as lifetime standards because of exhibiting double exponential decays [26, 28]. This may well be the reason that fluorescence lifetime measurements of quinine are basically absent from literature since the early 1990s.

Table 2 Quinine sulfate fluorescence lifetimes [ns]

|        | 10⁻³ M | 10⁻⁴ M | 10⁻⁵ M |
|--------|--------|--------|--------|
| H₂SO₄  | 2.26 ± 0.10 | 2.28 ± 0.13 | 2.34 ± 0.18 |

The (GFP) fluorescence lifetime was only measured for the original solution of 1 mg/ml (GFP) in 5 mM Tris-HCl, 5 mM EDTA with a pH of 8.0. Our measurements gave a lifetime of 2.8 ns, as shown in Table 3. This agrees very well with earlier results, with 2.8 ns for a deprotonated species [34], and also very good with 2.7 ns reported for a GFP-Rac2 [38] and 2.9 ns for a recombinant GFP [35]. The latter, however, also reported double exponential decay, with the 2.9 ns component contributing 90% of the total fluorescence. All our measurements of GFP fluorescence lifetimes produced single-exponential decays.

Table 3 Green fluorescent protein fluorescence lifetime [ns]

|        | 5 mM EDTA, pH 8.0 |
|--------|------------------|
| 5 mM Tris – HCl | 2.79 ± 0.07 |

Conclusions

To determine whether a fluorophore is suitable to use as a fluorescence lifetime standard, the two most important criteria is that it exhibits a single exponential decay and that it does not vary within at least two orders of magnitude in concentration. It is also important that the fluorophore has a strong fluorescence signal, and that it is completely soluble in a solvent. As a prerequisite, it should have absorption and emission peaks in the visible spectrum, as most lasers and detectors operate within this region, although this is of course dependent on what kind of fluorescence lifetime system one is using. All three tested fluorophores meet these criteria, since they were selected on this premise, although we were still able to rank them based on how well we think they are suitable as fluorescence lifetime standards. We found fluorescein to be the most suitable of the three fluorophores considered in this work, since the fluorescence lifetimes were stable with low uncertainties for all the tested solvents and concentrations. Fluorescein is also extensively used in a wide range of research fields, and is relatively affordable in large quantities. Quinine sulfate also yielded reliable lifetimes, however, its use was restricted with respect to solvent as it dissolved properly only in sulfuric acid. This makes it less suitable both for experimental work and as a lifetime standard compared to fluorescein, especially since sulfuric acid is a more hazardous chemical to work with than ethanol and methanol. (GFP) from the jellyfish Aequorea victoria, recombinant expressed in E. coli, did also produce very good fluorescence lifetime results, however, the suitability as a reference standard is not at the level of fluorescein because it is not readily available to dissolve in various solvents, and it is also more expensive compared to both fluorescein and quinine sulfate.
Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. Ahmed T, Shea K, Masters JR, Jones GE, Wells CM (2008) A pak4–link1 pathway drives prostate cancer cell migration downstream of hgf. Cell Signal 20(7):1320–1328
2. Al-Aqmar DM, Abdelkader H, Kana MTA (2015) Spectroscopic properties and amplified spontaneous emission of fluorescein laser dye in aqueous and water-soluble micellar media. J Mol Struct 1072:134–141
3. Bacsik BJ, Skoch J, Hickey GA, Allen R, Hyman BT (2003) Spectroscopic properties and amplified spontaneous emission of fluorescein laser dye in aqueous and water-soluble micellar media. J Mol Struct 1072:134–141
4. Barrow DA, Lentz BR (1984) Quinine as a fluorescence lifetime standard: conditions for effectively homogeneous decay. Chem Phys 76(2-3):163–167
5. Becker W, Bergmann A, Hink M, König K, Benndorf K, Biskup C (2004) Fluorescence lifetime imaging by time-correlated single-photon counting. Microsc Res Tech 63(1):58–66
6. Berlman I (1965) Handbook of fluorescence spectroscopy of aromatic molecules. Academic Press, New York
7. Birks JB (1970) Photophysics of aromatic molecules. Wiley-Interscience London, New York
8. Biswas S, Bhattacharya SC, Sen PK, Moulip SP (1999) Absorption and emission spectroscopic studies of fluorescein dye in alkanol, micellar and microemulsion media. J Photochem Photobiol A Chem 123(1-3):121–128
9. Boens N, Qin W, Basaric N, Hofkens J, Ameloot M, Pouget J, Lefèvre JP, Valeur B, Gratton E, VandeVen M et al (2007) Fluorescence lifetime standards for time and frequency domain fluorescence spectroscopy. Anal Chem 79(5):2137–2149
10. Chaffie M, Tu Y, Euskirchen G, Ward WW, Prasher DC (1994) Green fluorescent protein as a marker for gene expression. Science 263(5148):802–805
11. Drobnik J, Yeagers E (1966) On the use of quinine sulfate as a fluorescence standard. J Mol Spectrosc 19(1-4):454–455
12. Ferreira JA, Costa SM (2001) Solvent effects on the vibronic structure of s1? x0 transition of rhodamine 3b. J Mol Struct 565:35–38
13. Fletcher AN (1969) Quinine sulfate as a fluorescence quantum yield standard. Photochem Photobiol 9(5):439–444
14. Grynyczynski I, Nowaczyk K, Lakowicz J, Glovna D, Kawski A (1992) Picosecond fluorescence lifetime standards. In: FASEB J, vol 6, pp A314–A314. FEDERATION AMER SOC EXP BIOL 9650 ROCKVILLE PIKE, BETHESDA, MD 2084-3998
15. John J, Thomas L, George NA, Kurian A, George SD (2015) Tailoring of optical properties of fluorescein using green synthesized gold nanoparticles. Phys Chem Chem Phys 17(24):15,813–15,821
16. Joshi S, Pant DD (2012) Ground and excited state dipole moments of quinine sulfate dication: Solvatochromic shift of absorption and fluorescence spectra. J Mol Liq 172:125–129
17. Joshi S, Pant DD (2014) Steady state and time-resolved fluorescence spectroscopy of quinine sulfate dication bound to sodium dodecylsulfate micelles: fluorescent complex formation. J Lumin 145:224–231
18. Joshi S, Varma T, Pant DD (2013) Steady state and time-resolved fluorescence spectroscopy of quinine sulfate dication in water and neutral micelles: effect of medium on the photophysics. Colloids Surf A Physicoeng Chem 425:59–67
19. Kim D, Scranton AB, Stansbury JW (2009) Analysis of association constant for ground-state dye-electro acceptor complex of photoinitiator systems and the association constant effect on the kinetics of visible-light-induced polymerizations. J Polym Sci A Polym Chem 47(5):1429–1439
20. Koester V, Dowben P (1978) Subnanosecond fluorescence lifetimes by time-correlated single photon counting using synchronously pumped dye laserexcitation. Biophys J 24(1):245–247
21. Kristofoffersen AS, Erga SR, Hamre B, Frette Ø (2014) Testing fluorescence lifetime standards using two-photon excitation and time-domain instrumentation: rhodamine b, coumarin 6 and lucifer yellow. J Fluoresc 24(4):1015–1024
22. Lakowicz JR (1999) Principles of fluorescence spectroscopy, 2nd. Academic/Plenum Publishers, Kluwer
23. Lampert RA, Chewter LA, Phillips D, O’Connor DV, Roberts AJ, Meech SR (1983) Standards for nanosecond fluorescence decay time measurements. Anal Chem 55(1):68–73
24. Magde D, Rojas GE, Seybold PG (1999) Solvent dependence of the fluorescence lifetimes of xanthene dyes. Photochem Photobiol 70(5):737–744
25. Magde D, Wong R, Seybold PG (2002) Fluorescence quantum yields and their relation to lifetimes of rhodamine 6g and fluorescein in nine solvents: improved absolute standards for quantum yields. Photochem Photobiol 75(4):327–334
26. O’Connor D, Meech S, Phillips D (1982) Complex fluorescence decay of quinine bisulphate in aqueous sulphuric acid solution. Chem Phys Lett 88(1):22–26
27. Orte A, Crovetto L, Talavera EM, Boens N, Alvarez-Pez JM (2005) Absorption and emission study of 2’,7’-difluorofluorescein and its excited-state buffer-mediated proton exchange reactions. J Phys Chem A 109(5):734–747
28. Pant D, Tripathi H, Pant D (1992) Time resolved fluorescence spectroscopy of quinine sulphate, quinidine and 6-methoxysquinoline: ph dependence. J Lumin 51(4):223–230
29. Pant D, Tripathi U, Joshi G, Tripathi H, Pant D (1990) Photophysics of doubly-charged quinine: steady state and time-dependent fluorescence. J Photochem Photobiol A Chem 51(3):313–325
30. Parsons M, Vojnovic B, Ameer-beg S (2004) Imaging protein–protein interactions in cell motility using fluorescence resonance energy transfer (fret)
31. Sheen J, Hwang S, Niwa Y, Kobayashi H, Galbraith DW (1995) Green fluorescent protein as a new vital marker in plant cells. Plant J 8(5):777–784
32. Sierakowiak D (2005) Spectral properties of fluorescein molecules in water with the addition of a colloidal suspension of silver. J Phys Chem B 109(30):14,382–14,386
33. Spears KG, Cramer LE, Hoffland LD (1978) Subnanosecond time-correlated photon counting with tunable lasers. Rev Sci Instrum 49(2):255–262
34. Striker G, Subramaniam V, Seidel CA, Volkmer A (1999) Photochromicity and fluorescence lifetimes of green fluorescent protein. J Phys Chem B 103(40):8612–8617
35. Suhling K, Siegel J, Phillips D, French PM, Levêque-Fort S, Webb SE, Davis DM (2002) Imaging the environment of green fluorescent protein. Biophys J 83(6):3589–3595
36. Thompson RB, Gratton E (1988) Phase fluorometric method for determination of fluorescence lifetimes. Anal Chem 60(7):670–674
37. Tsien RY (1998) The green fluorescent protein
38. Van Manen HJ, Verkuijlen P, Wittendorp P, Subramaniam V, Van den Berg TK, Roos D, Otto C (2008) Refractive index sensing of green fluorescent proteins in living cells using fluorescence lifetime imaging microscopy. Biophys J 94(8):L67–L69
39. Velapoldi RA, Tønnesen HH (2004) Corrected emission spectra and quantum yields for a series of fluorescent compounds in the visible spectral region. J Fluoresc 14(4):465–472
40. Zhang XF, Zhang J, Liu L (2014) Fluorescence properties of twenty fluorescein derivatives: lifetime, quantum yield, absorption and emission spectra. J Fluoresc 24(3):819–826
41. Zhang XF, Zhang J, Liu L (2010) Photophysics of halogenated fluoresceins: involvement of both intramolecular electron transfer and heavy atom effect in the deactivation of excited states. Photochem Photobiol 86(3):492–498
42. Zipfel WR, Williams RM, Webb WW (2003) Nonlinear magic: multiphoton microscopy in the biosciences. Nat Biotechnol 21(11):1369