A highly photostable and versatile two-photon fluorescent probe for the detection of a wide range of intracellular nitric oxide concentrations in macrophages and endothelial cells

Carla Arnau del Valle a, Lewis Williams a, Paul Thomas b, Robert Johnson c, Sathuwarman Raveenthiraraj c, Derek Warren c, Anastasia Sobolewski c, María Paz Muñoz a, Francisco Galindo d, María J. Marín a,⁎

a School of Chemistry, University of East Anglia, Norwich Research Park, Norwich NR4 7TJ, UK
b Faculty of Sciences, University of East Anglia, Norwich Research Park, Norwich NR4 7TJ, UK
c School of Pharmacy, University of East Anglia, Norwich Research Park, Norwich NR4 7TJ, UK
d Departamento de Química Inorgánica y Orgánica, Universitat Jaume I, Av. Sos Baynat s/n, Castellón de la Plana 12071, Spain

ARTICLE INFO

Keywords:
Near-infrared
Two-photon microscopy
Nitric oxide detection
Macrophages cells
Endothelial cells

ABSTRACT

Nitric oxide (NO) is involved in many biological processes affecting the cardiovascular, nervous and immune systems. Intracellular NO can be monitored using fluorescent probes in combination with fluorescence imaging techniques. Most of the currently available NO fluorescent molecular probes are excited via one-photon excitation using UV or Vis light, which results in poor penetration and high photodamage to living tissues. Here, we report a two-photon fluorescent molecular probe, DANPY-NO, able to detect NO in live cells. The probe consists of an o-phenylenediamine linked to a naphthalimide core; and operates via photoinduced electron transfer. DANPY-NO exhibits good sensitivity (LOD of 77.8 nM) and high selectivity towards NO, and is stable over a broad range of pHs. The probe targeted acidic organelles within macrophages and endothelial cells, and demonstrated enhanced photostability over a commercially available NO probe. DANPY-NO was used to selectively detect endogenous NO in RAW264.7 Y macrophages, THP-1 human leukemic cells, primary mouse (bone marrow-derived) macrophages and endothelial cells. The probe was also able to detect exogenous NO in endothelial cells and distinguish between increasing concentrations of NO. The NO detection was evidenced using confocal laser scanning and two-photon microscopes, and flow cytometry. Further evidence was obtained by recording the changes in the intracellular fluorescence emission spectrum of the probe. Importantly, the probe displayed negligible toxicity to the analysed biological samples. The excellent sensitivity, selectivity, stability and versatility of DANPY-NO confirm its potential for in vitro and in vivo imaging of NO.

1. Introduction

Nitric oxide (NO) is a short-lived gaseous free radical involved in many physiological processes of the cardiovascular, nervous and immune systems [1,2]. In mammalian cells, NO is synthesised from L-arginine by nitric oxide synthase (NOS) enzymes including neuronal NOS (nNOS), inducible NOS (iNOS) and endothelial NOS (eNOS) [3,4]. These enzymes are responsible for the endogenous biosynthesis of NO, and differ in structure, function and localization [5]. nNOS and eNOS are calcium-dependent and produce low concentrations of NO (nNOS) for short periods of time; in contrast, iNOS is calcium-independent and produces a continuous production of NO [6]. NO generated by nNOS in the brain functions as a neurotransmitter, having roles in smooth muscle control, behaviour,
employs two NIR photons as the excitation source exhibiting advantages lower-energy excitation using NIR leads to negligible autofluorescence and higher photostability of the probe [35]. Two-photon microscopy as it allows for deeper penetration of biological tissues, and causes probes reported for the detection of intracellular NO [23] naphthalimide has been previously used for the construction of based on Nile Red, that permitted the \( \text{(NIR)} \) excitation overcomes some of the drawbacks of the UV industriousness, electron spin resonance spectroscopy, chemiluminescence, or colourimetry [18–21]. Despite the advances reported to date, the direct detection of NO has proven challenging due to its low concentration, short lifetime, and high reactivity with several reactive oxygen species (ROS) [6]. The use of fluorescent molecular probes in conjunction with microscopic techniques such as confocal laser scanning microscopy (LSM) has shown advantages over other techniques for the detection of intracellular NO. These advantages include spatial and temporal resolution and high sensitivity [19,22]. Consequently, considerable efforts have been made to further the availability and applicability of these fluorescent NO molecular probes, being aromatic \( o \)-diamino- \( \text{(o-dia-}\)mino)-fluorophore derivatives the most popular organic fluorescent probes reported for the detection of intracellular NO [23–27].

Despite the large number of fluorescent NO molecular probes reported in the literature for the detection of NO in living biological systems [28–34], there are still limitations that need to be overcome such as the excitation wavelength required for the measurement (mainly UV–Vis) and the high photobleaching which, in turn, affects the sensitivity of the detection and the spatiotemporal resolution. Near-infrared (NIR) excitation overcomes some of the drawbacks of the UV–Vis light as it allows for deeper penetration of biological tissues, and causes minimal photodamage upon long-term irradiation. In addition, the lower-energy excitation using NIR leads to negligible autofluorescence and higher photostability of the probe [35]. Two-photon microscopy employs two NIR photons as the excitation source exhibiting advantages over one-photon microscopy [36].

Several examples of two-photon excitable NO molecular probes have been reported in the literature, most of them based on an aromatic \( o \)-diamino-fluorophore. For example, Kim and co-workers developed \( o \)-phenylenediamine-based two-photon fluorescent probes that were able to selectively detect endogenous NO produced by RAW264.7 macrophages and had potential for deep-tissue imaging of NO [37,38]. Further improvements in the ability to monitor NO deep inside tissue were achieved when using a far-red emissive two-photon fluorescent probe based on Nile Red, that permitted the \textit{in situ} tracking of NO in an inflammatory process [39]. Among other two-photon fluorophores, 1,8-naphthalimide has been previously used for the construction of \( o \)-phenylenediamine-based fluorescent probes for the detection of NO in solution [40,41] and in different biological environments [42–45]. For example, Lyso-NINO was reported as a selective and highly sensitive NO probe that was able to detect endogenous NO in RAW264.7 macrophages and exogenous NO released by a donor in MCF-7 breast cancer cells [42], with low cytotoxicity to the latest. Similarly, the selective and highly sensitive probe for NO, LyNP-NO, was able to detect NO produced exogenously and endogenously in C6 glial cells and endogenously in rat hippocampus [44]. Despite the advances in the development of two-photon NO probes in recent years, to the best of our knowledge, there are no examples in the literature of highly biocompatible fluorescent NO probes that can be used for the successful detection of NO produced by the different forms of NOS in a broad range of biological environments and using different analytical techniques.

Here, we present a highly photostable, selective and biocompatible two-photon fluorescent probe (DANPY-NO, Fig. 1) that can be used for the detection of NO produced by iNOS in different macrophages, including a mouse macrophage cell line, primary mouse macrophages and human leukemic cells, and by eNOS in endothelial cells. DANPY-NO is a photoinduced electron transfer (PET)-based NO probe consisting of \( o \)-phenylenediamine linked to a naphthalimide core and bearing a simple piperazine scaffold. DANPY-NO shows good sensitivity and excellent selectivity towards NO in solution, and its fluorescence stability at different, biologically-relevant, pH values confirms its suitable applicability in physiological environments. The NO probe furthermore exhibits good photostability as confirmed when compared to a commercially available NO probe, 4,5-diaminofluorescein diacetate (DAF-2 DA), excellent biocompatibility, and extended applicability in cellular environments. The versatility of DANPY-NO as NO probe was confirmed using confocal LSM (images and intracellular fluorescence emission spectra), multiphoton microscopy and flow cytometry.

2. Results and Discussion

The novel two-photon NO molecular probe, DANPY-NO (Fig. 1), was designed as a PET-based fluorescent probe consisting of a 1,8-naphthalimide two-photon fluorophore conjugated to an \( o \)-phenylenediamine unit and a carboxyl-piperazine fragment. The NIR-excitable fluorophore, 1,8-naphthalimide, was chosen due to its well-known, remarkable photophysical properties such as good photostability, high quantum yield and pH insensitivity. The piperazine moiety was incorporated due to its versatile structure that could serve as scaffold to further the biological potential of DANPY-NO. An acetyl-derivative was specifically chosen here to prevent the protonation of the piperazine fragment at biological pHs. In DANPY-NO, the fluorescence emission is

![Fig. 1. Chemical structure of the two-photon NO molecular probe (DANPY-NO) and its reaction with NO in the presence of oxygen forming the triazole derivative (DANPY-NO-T). The fluorescence of the naphthalimide core is quenched in DANPY-NO due to a PET process. The formation of DANPY-NO-T hinders the PET process restoring the fluorescence of the core.](image-url)
quenched due to the PET that occurs from the electron-donating o-phenylenediamine to the naphthalimide core. The o-phenylenediamine moiety reacts with NO, in the presence of oxygen, forming the electron-poor triazole derivative, DANPY-NO-T, and enhancing the fluorescence emission of the naphthalimide core. Other examples of PET-based sensors have been reported in the literature for the detection of, for example, pH [46-48], endogenous NADH [49] and singlet oxygen [50], among others.

2.1. Synthesis and Characterisation of DANPY-NO

The probe was successfully synthesised in three steps from 4-bromom-1,8-naphthalic anhydride and 2-nitro-p-phenylenediamine (Scheme S1). DANPY-NO and the two intermediates of the synthesis were characterised by $^1$H NMR, $^{13}$C NMR, infrared spectroscopy and high-resolution mass spectrometry (HRMS) (see Figs. S1–S6 and Materials and Methods section in the SI for detailed characterisation). By measuring its UV–Vis absorption, and fluorescence excitation, emission and quantum yield. The photophysical properties of the NO probe were studied by measuring its UV–Vis absorption, and fluorescence excitation, emission and quantum yield. DANPY-NO exhibited a maximum absorption band at 396 nm ($\varepsilon = 1.13 \times 10^4$ M$^{-1}$ cm$^{-1}$ in DMSO, Figs. S7a and S8) and a fluorescence emission peak centered at 535 nm in DMSO (Fig. S7c), both characteristic of 1,8-naphthalimide structures.

2.2. Sensitivity and Selectivity of DANPY-NO towards NO

The ability of DANPY-NO to detect NO was first evaluated and confirmed in aqueous solution. As observed in Fig. 2, the weak fluorescence emission intensity of DANPY-NO at 556 nm in aqueous solution (13% DMSO) increased by 647% upon addition of an excess of the NO donor diethylenetriamine NONOate (0.5 mM) confirming the ability of the NO probe to detect NO via formation of the corresponding triazole, DANPY-NO-T. Additionally, a remarkable 13-fold increase in the fluorescence emission intensity of DANPY-NO was only observed following addition of NO, confirming the excellent selectivity of the developed compound towards NO. Furthermore, DANPY-NO remained sensitive to NO in the presence of the different species tested (Fig. S13). These results further confirm the suitability of DANPY-NO for the detection of intracellular NO.

2.3. Characterisation of DANPY-NO in RAW264.7 Y NO$^-$ Cells

To investigate the uptake of DANPY-NO by live cells, RAW264.7 Y NO$^-$ macrophages were incubated overnight with the probe and confocal LSM images were acquired showing bright green fluorescence upon excitation at 405 nm (Fig. S14). To further confirm that the observed green fluorescence was due to the DANPY-NO internalised by the macrophages, intracellular fluorescence emission spectra were recorded. The fluorescence emission spectrum of DANPY-NO inside RAW264.7 Y NO$^-$ matches that of DANPY-NO recorded in Dulbecco’s Modified Eagle Medium (DMEM) cell culture medium (black and red, respectively, in Fig. 4a). Importantly, the fluorescence emission spectrum of untreated cells (green in Fig. 4a) did not exhibit the characteristic fluorescence emission band at ca. 520 nm typical of 1,8-naphthalimide derivatives.

To elucidate the localisation of DANPY-NO within the RAW264.7 Y NO$^-$ cells, colocalisation studies were performed using LysoTracker Red DND-99, a marker of acidic organelles. As it can be seen in Fig. 4b (see Fig. S15 for scatterplot), there is a clear overlap between the fluorescence emission from DANPY-NO (green) and the fluorescence emission from the LysoTracker Red DND-99 (red), confirmed by the yellow spots visible when the green and the red channels are overlaid with the differential interference contrast (DIC) image. The degree of colocalisation between DANPY-NO and the LysoTracker in the acidic organelles was confirmed by a Pearson’s correlation coefficient of 0.74 ± 0.02 (n = 3 images, ca. 20 cells), which is excellent considering that DANPY-NO does not have a pending specific targeting moiety like previously reported probes of this type [42–45].
2.4. Intracellular Application of DANPY-NO for the Detection of NO in Macrophages

The potential of DANPY-NO to monitor NO concentrations in different cellular environments was investigated starting with its application in macrophages. The ability of DANPY-NO to detect endogenous NO was studied in RAW264.7 \( \gamma \)NO cells stimulated overnight with lipopolysaccharide (LPS) and IFN-\( \gamma \) to produce elevated concentrations of NO. Following incubation, the cells were imaged using confocal LSM. Stimulated cells incubated with DANPY-NO exhibited higher green fluorescence emission intensity compared to those unstimulated cells incubated with the probe (Fig. 5). These results confirmed the successful detection of endogenous intracellular NO in live macrophages by DANPY-NO. It is also important to notice that non-treated cells, both stimulated and unstimulated, did not show any evident fluorescence emission (Fig. S16). To further confirm the enhancement of the fluorescence emission intensity of DANPY-NO in stimulated macrophages due to the formation of the triazole derivative, DANPY-NO-T, in the presence of NO, the intracellular fluorescence

Fig. 3. Normalised a) fluorescence emission spectra and b) fluorescence response at 556 nm of DANPY-NO (8.5 \( \mu \)M) in the presence of NO and other biologically-relevant species (100 \( \mu \)M aqueous solution with 14% DMSO). \( \lambda_{\text{exc}} = 396 \) nm. Each experiment was repeated in triplicate and the relative standard errors are indicated by the error bars.

Fig. 4. a) Normalised fluorescence emission spectrum of: DANPY-NO in DMEM cell culture medium recorded in the fluorescence spectrophotometer (black); DANPY-NO internalised by RAW264.7 \( \gamma \)NO cells recorded in the confocal LSM (red); and non-treated control cells recorded in the confocal LSM (green); \( \lambda_{\text{exc}} = 405 \) nm. b) Confocal LSM images of RAW264.7 \( \gamma \)NO cells treated with DANPY-NO and LysoTracker Red DND-99. Images collected upon excitation at i) \( \lambda_{\text{exc}} = 405 \) nm, \( \Delta \lambda_{\text{em}} = 500-580 \) nm and ii) \( \lambda_{\text{exc}} = 561 \) nm, \( \Delta \lambda_{\text{em}} = 580-625 \) nm; and iii) composite image of green, red and DIC channels. Scale bars = 10 \( \mu \)m. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

2.4. Intracellular Application of DANPY-NO for the Detection of NO in Macrophages

The potential of DANPY-NO to monitor NO concentrations in different cellular environments was investigated starting with its application in macrophages. The ability of DANPY-NO to detect endogenous NO was studied in RAW264.7 \( \gamma \)NO macrophages stimulated overnight with lipopolysaccharide (LPS) and IFN-\( \gamma \) to produce elevated concentrations of NO. Following incubation, the cells were imaged using confocal LSM. Stimulated cells incubated with DANPY-NO exhibited higher green fluorescence emission intensity compared to those unstimulated cells incubated with the probe (Fig. 5). These results confirmed the successful detection of endogenous intracellular NO in live macrophages by DANPY-NO. It is also important to notice that non-treated cells, both stimulated and unstimulated, did not show any evident fluorescence emission (Fig. S16). To further confirm the enhancement of the fluorescence emission intensity of DANPY-NO in stimulated macrophages due to the formation of the triazole derivative, DANPY-NO-T, in the presence of NO, the intracellular fluorescence
emission spectra of the investigated samples were recorded. As expected, the fluorescence emission intensity in stimulated cells was higher compared to the intensity of unstimulated cells incubated with DANPY-NO, and negligible fluorescence was observed for unstimulated and stimulated control cells (Fig. 5b). Colocalisation experiments with LysoTracker Red DND-99 confirmed the overlap between the green fluorescence observed in stimulated cells incubated with DANPY-NO (fluorescence of DANPY-NO-T) and the red fluorescence of the marker of acidic organelles (Pearson’s correlation coefficient of 0.71 ± 0.03 – n = 3 images, ca. 20 cells; Fig. S17). The successful performance of DANPY-NO in the detection of endogenous NO produced by stimulated RAW264.7Y NO– cells was also confirmed by flow cytometry (Fig. S18). Cells stimulated with LPS and INF-γ to produce NO and incubated with DANPY-NO exhibited a higher mean-fluorescence intensity compared to unstimulated cells incubated with DANPY-NO.

To prove the selective detection of NO by our probe, RAW264.7Y NO– cells were treated with an inhibitor of the NOS, N(o)-nitro-L-arginine methyl ester (L-NAME), to block the production of NO in stimulated RAW264.7Y NO– cells, as confirmed with a Griess assay (Fig. S19). Cells pre-treated with L-NAME displayed a reduced fluorescence emission compared to the stimulated cells that did not receive a pre-treatment with L-NAME (Fig. S20) confirming the selectivity of DANPY-NO towards NO in RAW264.7Y NO– cells.

An important feature to be considered when designing an intracellular fluorescent probe is the photobleaching of the probe over the irradiation time. To study the photostability of DANPY-NO over time, RAW264.7Y NO– cells incubated with DANPY-NO were imaged at different exposure times following continuous irradiation (λexc = 405 nm). The photostability of our probe was compared to the photostability of DAF-2 DA (5 μM), a commercially available probe commonly used for the detection of intracellular NO (λexc = 488 nm). The fluorescence emission intensity of DANPY-NO exhibited a 16 ± 5% decrease after 5 min of continuous irradiation compared to a 38 ± 10% reduction observed for DAF-2 DA following the same irradiation time (Fig. 6 and Fig. S21). These results confirm the remarkable photostability of DANPY-NO which could be beneficial for long-term monitoring of biological processes involving NO generation, e.g. monitoring of NO released following bacterial infection.

Due to the excellent performance of DANPY-NO in sensing NO in RAW264.7Y NO– macrophages, the next step was to evaluate its potential in primary mouse cells. Bone marrow-derived macrophages (BMDMs) were used as a model of primary cells. BMDMs were able to produce concentrations of nitrates of 30.4 ± 0.7 μM upon overnight stimulation with LPS (50 ng/mL) and INF-γ (100 ng/mL) compared to the negligible concentration of nitrates for unstimulated cells (0.6 ± 0.2 μM), as proven by the Griess assay. DANPY-NO was able to successfully detect NO endogenously produced by stimulated primary BMDMs (Fig. S22). Confocal LSM images of fixed cells stimulated and incubated with DANPY-NO exhibited higher fluorescence emission than the unstimulated cells incubated with DANPY-NO (Fig. S22a). BMDMs stimulated or unstimulated and without DANPY-NO loaded were also imaged as controls and exhibited no fluorescence. These results were further supported with the intracellular fluorescence emission spectra of the four different investigated samples, which clearly confirmed the ability of DANPY-NO to detect intracellular NO production in primary cells upon stimulation (Fig. S22b). Colocalisation experiments of DANPY-NO with LysoTracker Red confirmed the presence of the NO probe in the acidic organelles of the BMDMs (Pearson’s correlation coefficient of 0.84 ± 0.10 – n = 3 images, ca. 10 cells; Fig. S23).

DANPY-NO was further evaluated in human macrophages, known to produce lower NO concentrations than previously investigated macrophages. THP-1 cells were selected for these experiments since they are acute monocytic leukemic cells derived from the peripheral blood of an infant. THP-1 cells were first differentiated to macrophages using phorbol 12-myristate 13-acetate (PMA) treatment overnight. The concentration of nitrites produced by the differentiated THP-1 cells upon stimulation with LPS (5 μg/mL) for 48 h was found to be 2.55 ± 0.07 μM using the Griess assay. Confocal LSM images of live PMA-differentiated THP-1 cells incubated with DANPY-NO and stimulated using LPS for 48 h exhibited a bright emission intensity that contrasted the weak fluorescence observed in the images of unstimulated cells incubated with DANPY-NO (Fig. S24a). This apparent fluorescence emission enhancement of DANPY-NO when detecting NO in human macrophages was further confirmed when the intracellular fluorescence emission spectra were recorded (Fig. S24b). Finally, the ability of DANPY-NO to specifically detect NO in THP-1 cells was investigated using L-NAME. Confocal LSM images of THP-1 cells pretreated with L-NAME exhibited intensities that were comparable to those observed for THP-1 cells unstimulated and incubated with DANPY-NO. As expected, the fluorescence emission spectrum of THP-1 cells treated with L-NAME, stimulated and incubated with DANPY-NO (red in Fig. S24b) overlapped the spectrum of unstimulated control cells incubated with DANPY-NO and that of unstimulated control cells treated with L-NAME and incubated with DANPY-NO (black and blue, respectively, in Fig. S24b). Control cells without DANPY-NO loaded (unstimulated cells and stimulated cells with and without L-NAME treatment) showed negligible fluorescence intensity in the corresponding confocal images (Fig. S25), which are supported by the corresponding intracellular fluorescence emission spectra. The presence of DANPY-NO within the acidic organelles of the PMA-differentiated THP-1 cells was confirmed by
colocalisation with LysoTracker Red (Pearson’s correlation coefficient of 0.42 ± 0.06 – n = 3 images, ca. 10 cells; Fig. S26).

**DANPY-NO** was also able to monitor different concentrations of intracellular NO in THP-1 macrophages. Differentiated THP-1 macrophages were separately stimulated with two different concentrations of LPS (5 and 10 μg/mL). After overnight treatment, confocal LSM images and intracellular fluorescence emission spectra were recorded, which exhibited an increase in the fluorescence emission intensity of **DANPY-NO** concomitant with the increase in LPS concentration (Fig. 7; corresponding confocal LSM images shown in Fig. S27).

### 2.5. Intracellular Application of **DANPY-NO** for the Detection of NO in Endothelial Cells

To further explore the versatility of **DANPY-NO**, the potential of the probe to detect intracellular NO produced by eNOS was investigated in endothelial cells. NO is continuously produced by eNOS using L-arginine as substrate; and this production can be inhibited using L-NMMA or enhanced by treating the cells with a Ca<sup>2+</sup>-ionophore, since eNOS is a Ca<sup>2+</sup>-dependent enzyme. **DANPY-NO** was used to detect endogenous and exogenous NO in endothelial cells. To this aim, three types of samples containing **DANPY-NO** and their corresponding controls without the probe were prepared: 1) endothelial cells incubated with **DANPY-NO** overnight; 2) cells treated with Ca<sup>2+</sup>-ionophore A-23187 together with **DANPY-NO** overnight; 3) cells pre-treated with L-NMMA and incubated with **DANPY-NO** overnight. As confirmed by the Griess assay, higher NO levels were produced by cells containing Ca<sup>2+</sup>-ionophore A-23187 (0.295 ± 0.004 μM) and lower NO concentrations were observed in cells pretreated with L-NMMA (0.12 ± 0.02 μM), compared to untreated cells (0.21 ± 0.01 μM). The endothelial cells incubated with **DANPY-NO** and Ca<sup>2+</sup>-ionophore A-23187 showed an enhanced fluorescence emission intensity compared to those cells incubated only with **DANPY-NO** (Fig. S28a). In contrast, a reduction of the fluorescence emission intensity was observed for the cells pre-treated with L-NMMA (Fig. S28a, vii – ix). Confocal LSM images of control samples showed negligible fluorescence emission (Fig. S29). These results were confirmed by the intracellular fluorescence emission spectra recorded also in the confocal LSM (Fig. S28b) which exhibited the same trend as the confocal LSM images. These results demonstrated the ability of **DANPY-NO** to monitor different endogenous NO levels in endothelial cells.

The ability of **DANPY-NO** to detect exogenous NO in endothelial cells was studied using S-nitroso-N-acetylpenicillamine (SNAP) as NO donor. Endothelial cells were incubated overnight with **DANPY-NO** in the presence and absence of SNAP (220 μM). Following the corresponding incubation, confocal LSM images and spectra were acquired (Fig. S30). The confocal images exhibited a small difference in fluorescence emissions between cells incubated with **DANPY-NO** and cells incubated with SNAP, the latter being the brightest and thus confirming the presence of higher concentrations of NO. These results were supported with the fluorescence emission spectra that showed a higher intensity for cells treated with SNAP compared to cells incubated only with **DANPY-NO**. The fluorescence emission spectra of control cells that were not incubated with **DANPY-NO** did not show the characteristic emission intensity of the compound. **DANPY-NO** was partially colocalised with LysoTracker Red (Pearson’s correlation coefficient of 0.50 ± 0.10 – n = 3 images, ca. 5 cells; Fig. S31) in the endothelial cells.

### 2.6. Intracellular Characterisation of **DANPY-NO** under Two-Photon Excitation

Given the excellent performance of **DANPY-NO** as NO probe in solution and under one-photon excitation (405 nm), the ability of **DANPY-NO** to be excited by two-photon excitation (800 nm), and to be used to monitor NO production under these conditions were investigated using a multiphoton microscope (Fig. 8). RAW264.7 NO<sup>-</sup> macrophages, unstimulated or stimulated with LPS and IFN-γ, were incubated with **DANPY-NO** overnight. After this time, live cells were imaged using 800 nm irradiation with fluorescence collected between 500 and 550 nm. Unstimulated cells incubated with **DANPY-NO** exhibited a green fluorescence emission (Fig. 8) that was not observed in the control cells which had not been incubated with **DANPY-NO** (Fig. S32). Additionally, stimulated cells incubated with **DANPY-NO** showed a remarkable enhancement of the green fluorescence (Fig. 8) as expected from the previous results using confocal LSM. These results confirmed the capacity of **DANPY-NO** to be visualised under NIR light (800 nm) and its potential to detect intracellular NO under two-photon excitation using multiphoton microscopy.

### 2.7. Cytotoxicity Experiments of **DANPY-NO**

Importantly, the cytotoxicity of **DANPY-NO** towards all the cellular systems used was determined by CellTiter-Blue viability assay showing a great biocompatibility in all the cases (Fig. S33). The cytotoxicity of

![Fig. 7](image-url) Intracellularly recorded fluorescence emission spectra of THP-1 differentiated macrophages incubated with **DANPY-NO** (5 μM) and stimulated with different LPS concentrations (0, 5 and 10 μg/mL; black, blue and green spectra, respectively) overnight. Grey, red and yellow spectra correspond to cells without **DANPY-NO** and treated with 0, 5 and 10 μg/mL of LPS, respectively. λ<sub>exc</sub> = 405 nm. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

![Fig. 8](image-url) Multiphoton microscopy images of RAW264.7 NO<sup>-</sup> cells incubated overnight with **DANPY-NO** (5 μM): unstimulated cells (i – iii) and cells stimulated with LPS (0.7 μg/mL) and IFN-γ (17 μg/mL) (iv – vi). λ<sub>exc</sub> = 800 nm, Δλ<sub>em</sub> = 500–550 nm. Scale bars = 20 μm.
DANPY-NO towards RAW264.7 Y NO− macrophages showed a cell survival rate higher than 94% after overnight treatment with DANPY-NO (5 μM). In BMDMs, the cytotoxicity of DANPY-NO exhibited a cell viability of 88% when cells were incubated with the concentration of DANPY-NO used in the intracellular NO detection experiments (5 μM). Additionally, double the experimental concentration of DANPY-NO continued to exhibit low cytotoxicity to BMDMs. Cytotoxicity studies of DANPY-NO in PMA-differentiated THP-1 cells incubated for 48 h did not show a significant cell toxicity even at 10 μM (86% of cells were still viable). Finally, DANPY-NO also showed negligible cytotoxicity to endothelial cells in the range of concentrations used.

3. Conclusions

In summary, we have reported a highly versatile PET-based two-photon fluorescent probe, DANPY-NO, for the detection of NO produced by iNOS and eNOS in a wide selection of cells and at a different range of concentrations. In solution, DANPY-NO was selective to NO over other potential biological interferences, exhibited a LOD of 77.8 nM, suitable for the detection of endogenous NO produced by biological samples, and its fluorescence was stable at different, biologically-relevant, pH values. The versatility of DANPY-NO for the detection of intracellular NO was confirmed using a range of techniques including confocal LSM (images and intracellular fluorescence emission spectra), multiphoton microscopy and flow cytometry. The probe was successfully used for the selective detection of intracellular endogenous NO produced by iNOS and eNOS in a wide range of cells including: mouse macrophages, human leukemic cells, primary mouse macrophages and endothelial cells; demonstrating an excellent biocompatibility with all the listed cellular samples. Furthermore, DANPY-NO exhibited higher photostability than a frequently used commercially available NO probe. Overall, DANPY-NO has proven to be an excellent candidate for the monitoring of intracellular NO produced at low and high concentrations; and its two-photon excitation nature makes it potentially valuable for both in vitro and in vivo detection of NO.

Author Contributions

C. A. V performed all the experimental work described in this paper and analysed the data, wrote the original draft of the manuscript and the ESI and contributed to the later reviewing and editing. L. W. provided guidance on the synthesis of DANPY-NO. P. T. provided training and supervision of the imaging techniques and contributed to reviewing and editing the manuscript. R. J. contributed to the endothelial cell culture and provided help with the intracellular experiments involving endothelial cells. S. R. isolated and supplied the BMDMs. D. W. provided guidance on the endothelial work, contributed to reviewing and editing the manuscript. A. S. provided guidance on the BMDMs work and supplied the THP-1 cells. M. P. M. provided training and supervision of the synthesis of DANPY-NO and contributed to writing, reviewing and editing the manuscript. F. G. contributed to the design of the project and reviewing the manuscript. M. J. M. supervised the entire research with editing the manuscript. F. G. contributed to the design of the project and synthesis of ESI and contributed to the later reviewing and editing. L. W. provided training and guidance on the photon excitation nature makes it potentially valuable for both in vitro and in vivo detection of NO.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgment

The authors acknowledge Dr. A. Goldson for training and guidance on the flow cytometer; Dr. P. Wilson for training and guidance on the multiphoton microscope. The authors would like to thank the Faculty of Sciences and School of Chemistry at the University of East Anglia and Mr. and Mrs. Whittaker oncology fellowship for financial support, and the EPSRC (Grant EP/S017909/1) that supported the purchase of the Edinburgh Instrument FSS fluorescence spectrometer used in this work.

Appendix A. Supplementary data

The SI contains the experimental section of this paper and the supporting figures including NMR spectra of DANPY-NO and its intermediates; electronic spectroscopic properties of DANPY-NO and sensitivity and selectivity characterisation; and supporting intracellular controls and further intracellular characterisation. Supplementary data to this article can be found online at https://doi.org/10.1016/j.jphotobiol.2022.112512.

References


Y. Yu, X. Zhang, Y. Dong, X. Luo, X. Qian, Y. Yang, Fusing the Nagano’s and the Anslyn’s chemistry for lyso-specific NO detection, Sensors Actuators B Chem. 346 (2021), 130562.

Y. Chen, Recent developments of fluorescent probes for detection and bioimaging of nitric oxide, Nitric Oxide 98 (2020) 1–19.


