Use of a carbocyanine dye as a marker of functional vasculature in murine tumours

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Summary An intravenously administered fluorescent carbocyanine dye, DiOC₃(3), has been evaluated for use in conjunction with Hoechst 33342 as a marker of murine tumour vasculature. DiOC₃(3) stains cells immediately adjacent to blood vessels and thus, like Hoechst 33342, outlines perfused tumour vasculature. The different fluorescence excitation and emission properties of DiOC₃(3) and Hoechst 33342 permit discrimination of the stains in the same tissue section. Mice tolerate a DiOC₃(3) dose of 1 mg kg⁻¹ i.v. with no ill effects. The dye has a distribution half-life in blood of 180s and staining of perivascular tumour cells is sufficiently stable to allow visualisation of vasculature for up to 30 min after DiOC₃(3) injection. However, DiOC₃(3) causes a 75% reduction in tumour blood flow as measured by laser Doppler techniques. Consequently, the compound appears to be most suitable as a second vascular marker, administered at some time after Hoechst 33342, to detect temporal and spatial fluctuations in tumour perfusion.

Quantitative histological study of microvascular perfusion in solid experimental tumours can be undertaken using blood-borne stains or tracer substances which label the functional tumour vasculature. Substances confined exclusively to the vascular compartment, such as Cr-labelled red blood cells and high molecular weight dextrans, have been used to measure functional tumour vascular space (Gullino & Grantham, 1964; Tannock & Steel, 1969). Most biological stains (for example, Evans blue, trypan blue or lissamine green) penetrate rapidly from blood vessels into the interstitial compartment following injection (Goldacre & Sylven, 1962). Some fluorescent dyes, such as fluorescein, behave similarly. Thus, unless animals are killed and tumours removed immediately after stain administration, no discrimination of tumour vasculature is obtained. Fluorescent or radio-labelled microspheres can also be used to label blood vessels but they are trapped in only a small fraction of tumour vessels seen in thin tissue sections.

However, identification of functional tumour vasculature can be achieved using intravenously administered dyes which avidly stain cells adjacent to the blood supply and thus penetrate slowly into tumour parenchyma. A DNA-binding, UV light-excited fluorescent stain, Hoechst 33342, has such properties and has been used to investigate vascular patterns within tumours (Reinhold & Visser, 1983), for cell selection from tumours using fluorescence-activated cell sorting (Chaplin et al., 1985; Olive et al., 1985), and as a vascular space marker in tumours (Smith et al., 1988).

Regions of tumour microvasculature may be subject to transient reductions in perfusion, flow stasis, or vascular collapse (Brown, 1979; Intaglia et al., 1977; Reinhold et al., 1977; Chaplin et al., 1986, 1987). Thus, stains such as Hoechst 33342, with a short half-life in blood, may not identify tumour vessels which are collapsed or otherwise non-functional during the few minutes following injection. Histological detection of such regions might be achieved by administration of a second vascular marker at some interval after Hoechst 33342 so that collapsed vessels, if they reopen, could be visualised. Conversely, cells near vessels which close during the interval between stain injections would be labelled only by Hoechst stain and not the second marker.

We have examined a fluorescent carbocyanine derivative, 3,3'-diethylxocarbocyanine (DiOC₃(3)), for use as a marker of tumour vasculature. DiOC₃(3) has been employed previously in photographic processing and as a membrane potential probe (Sims et al., 1974). The slow penetration rate of this dye has been used to select cells from different depths within multicell spheroids (Olive & Durand, 1987). DiOC₃(3) has a long alkyl side chain, increasing its lipophilicity and facilitating rapid entry into cells. This suggested to us that this dye might have in vivo staining characteristics similar to Hoechst 33342; that is, to stain cells immediately adjacent to blood vessels. Carbocyanine dyes are excited by visible light and can easily be discriminated from Hoechst 33342 when both stains are present in tissue sections examined by fluorescence microscopy. Thus, such dyes, used in conjunction with Hoechst 33342, could be employed in a double staining protocol to identify regions of transient perfusion in experimental tumours.

We have investigated DiOC₃(3) for use as a marker of functional tumour vasculature and have examined pharmacokinetics, in vivo toxicity, stability of binding to tumour cells and vasoactive properties of the dye.

Materials and methods

Fluorescent dyes

The carbocyanine dye DiOC₃(3) was obtained from Molecular Probes Inc. (Eugene, OR, USA). The structure of symmetric carbocyanine dyes is shown in Figure 1 and can be described using the abbreviated notation DiYC₃(2n+1) (Sims et al., 1974). The lipophilicity of cyanine dyes increases with n in the above formula and the wavelengths of maximum absorption and emission increase with m. DiOC₃(3) has a molecular weight of 600 and exhibits green fluorescence when excited by blue light.

Carbocyanine dyes are poorly soluble in aqueous solution and DiOC₃(3) was dissolved in dimethyl sulphoxide (DMSO) and then diluted to 75% DMSO with phosphate-buffered saline before use. Hoechst 33342 (Sigma, St Louis, MO, USA) was dissolved in sterile phosphate-buffered saline and administered intravenously at a dose of 15 mg kg⁻¹.

Blood levels of DiOC₃(3) after i.v. injection were determined by fluorometric methods as described previously (Olive et al., 1985). Toxicity of DiOC₃(3) towards SCCVII tumour cells exposed to the stain in vivo was assessed as follows. Tumours were excised 5 min after simultaneous i.v. administration of Hoechst 33342 and DiOC₃(3) (1 mg kg⁻¹) and enzymatically disaggregated into a single cell suspension. Using a flow cytometer, cells were sorted into 10 fractions on the basis of the Hoechst 33342 diffusion gradient (Chaplin et al., 1985) and examined for clonogenicity by plating in 100-mm plastic tissue culture dishes kept in a
tumour with little or no contribution from skin or subcutaneous tissue. Red blood cell flux, number of moving RBCs and mean RBC velocity were continually monitored both before and after injection of DiOC$_3$(3), 1 mg kg$^{-1}$ i.v. The influence of intravenously injected DiOC$_3$(3) on the haematocrit of orbital sinus blood samples was also measured.

Results

Following intravenous administration, the blood concentration of DiOC$_3$(3) declined exponentially with a distribution half-life of 180 s (compared to 100 s for Hoechst 33342) and an elimination half-life of approximately 30 min (based on a two-compartment model of drug kinetics) (Figure 1). The apparent volume of distribution of DiOC$_3$(3) is very large, reflecting the lipophilic nature of the stain.

Mice tolerated an intravenous dose of 1 mg kg$^{-1}$ without obvious acute ill effects and were alive and well 6 weeks after drug administration. Doses greater than 5 mg kg$^{-1}$ resulted in death of the animal. The in vitro plating efficiency of SCCVII tumour cells exposed to DiOC$_3$(3) (1 mg kg$^{-1}$ i.v.) in vivo was not significantly reduced. Even cells located in the brightest Hoechst 33342 sort fractions, representing cells closest to tumour blood vessels and thus exposed to the highest concentration of DiOC$_3$(3), showed no reduction in clonogenicity (data not shown).

DiOC$_3$(3)-staining of cells immediately adjacent to blood

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**Figure 1** Blood levels of DiOC$_3$(3) in C$_5$H$_7$He mice following intravenous injection of 1 mg kg$^{-1}$. The mean $\pm$ standard deviation (s.d.) is shown and for those symbols without s.d. bars, the s.d. is less than the symbol size.

humidified incubator equilibrated with 5% CO$_2$, 5% O$_2$ and 90% N$_2$. Colonies were stained and counted 12 days later.

**Identification of tumour vasculature**

The ability of DiOC$_3$(3) to act as a marker of functional tumour vasculature was examined in SCCVII grown subcutaneously (s.c.) in C$_5$H$_7$He mice. The dye was administered via lateral tail vein injection at a dose of 1 mg kg$^{-1}$ in 50 $\mu$L volume. This dose provided optimal visualisation of tumour vasculature. Animals were killed by cervical dislocation at various intervals after dye injection and tumours were immediately excised and frozen at $-20^\circ$C in OCT embedding compound. Frozen sections 10 $\mu$m in thickness were rewarmed to room temperature, allowed to air dry, and examined using a microscope with 100 W mercury lamp illumination and an epifluorescence condenser using 430–490 nm excitation, a 510 nm dichroic mirror and a 520 nm long pass filter. Stain stability and bleaching were measured using an image intensified charge-injection device (CID) camera linked to a computer-based image processing system (Jaggi et al., 1988).

**Tumour blood flow measurements**

Relative changes in tumour blood flow following intravenous injection of DiOC$_3$(3) were measured using an infrared laser Doppler flowmeter (TSI Inc., St Paul, MN, USA). Electrical signals from the laser Doppler photodetector are proportional to the number of moving red blood cells (RBCs) and to mean RBC velocity. RBC flux (expressed as ml min$^{-1}$ 100 g tissue$^{-1}$) is the product of these two measurements. This instrument is capable of monitoring tissue microvascular flow continuously and noninvasively with a spatial resolution of approximately 1 mm$^3$ (Haumschild, 1986; Shepherd et al., 1987). Animals bearing s.c. implanted tumours were anaesthetised with ketamine (50 mg kg$^{-1}$ i.p.) and diazepam (10 mg kg$^{-1}$ i.p.), a 1–2 mm incision was made in the thinned skin directly overlying the tumour, and a 0.7 mm diameter laser Doppler needle probe was placed through this incision on to the tumour. Thus, flow measurements were recorded from a small, peripheral region of

![Figure 2](image-url)
The confidence number of stained tumour (r = 0.67) but tumours. Tumour 15 mm² was only in cells located a small distance from the vessel lumen.

The stability of DiOC₃(3) fluorescence in SCCVII tumours and the localisation of dye in relation to blood vessels were assessed using the fluorescence imaging system. In the first few minutes following DiOC₃(3) injection, the dye was seen only in cells located a small distance from the vessel lumen. With time, dye became detectable in cells more distant from the blood supply and fluorescence of the perivascular cells declined; that is, peak fluorescence intensity decreased and staining distance away from vessel increased. The peak-fluorescence/distance ratio is therefore a measure of dye localisation in relation to tumour blood vessels (Figure 4). Thirty to 60 min following DiOC₃(3) administration, the difference in fluorescence intensity between perivascular cells and cells more distant from the blood supply is essentially lost. Visualisation of tumour vasculature was most distinct if tumours were removed 5 min after dye injection. DiOC₃(3) fluorescence is relatively resistant to bleaching by continuous illumination of the tissue section with blue light using our optical system. In comparison, Hoechst 33342 fluorescence fades rapidly when excited by UV light.

DiOC₃(3) caused a reduction in blood flow in SCCVII tumours as measured using laser Doppler flowmetry (Figure 5). Baseline values for RBC flux varied between 4.8 and 8.0 ml min⁻¹ 100 g⁻¹ (mean 5.95 ± 1.23). DiOC₃(3) injection resulted in an initial transient reduction, recovery and then a gradual, significant decline in flux to levels which were approximately 25% of baseline values (mean 1.58 ± 0.73,

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**Figure 3** Quantitation of vascularity using DiOC₃(3). The number of stained regions per 1 mm² is plotted as a function of tumour weight for 29 subcutaneously implanted SCCVII tumours. The stain was administered i.v. to unrestrained, unanaesthetised mice. Counting was restricted to non-necrotic tumour regions. For each tumour, a total area of approximately 15 mm² was counted. The best fit linear regression line with 95% confidence limits is shown. The negative correlation is weak (r = 0.67) but statistically significant (P < 0.01).

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**Figure 4** Stability of DiOC₃(3) staining of SCCVII tumour cells in vivo. Peak fluorescence intensity declined and staining distance away from vessels increased with time after dye injection. Error bars represent standard error of the mean (s.e.m.) for 20 vessels per tumour. Below a fluorescence/distance ratio of about 5, counting blood vessels was not possible primarily because vessels near to each other could not be easily discriminated.

![Graph](image_url)

**Figure 5** Effects of DiOC₃(3) (1 mg kg⁻¹ i.v.) on red blood cell (RBC) flux, number of moving RBCs (indicative of functional microvascular volume), and mean RBC velocity in subcutaneous SCCVII tumours as assessed by laser Doppler flowmetry. Error bars represent s.e.m. for five tumours.
When alone, 75% DMSO (75%) caused no changes in tumour blood flow. Intravenous administration of DiOC₃(3) did not alter blood haematocrit. In anaesthetised animals, DiOC₃(3) also caused a slight decrease (1.5°C) in body temperature.

Discussion

The fluorescent carbocyanine dye, DiOC₃(3), rapidly enters and stains cells immediately adjacent to the blood supply, thus identifying functional vasculature in murine tumours. Intravenous administration of either DiOC₃(3) or Hoechst 33342 results in similar staining patterns. In conjunction with Hoechst 33342, DiOC₃(3) appears useful as a vascular marker in double staining experiments designed to locate regions of transient perfusion in experimental tumours.

Carbocyanine dyes have been used extensively for measurement of cell membrane potential and DiOC₃(3) has been shown to stain proliferating cells to a greater extent than non-proliferating cells (Cohen et al., 1981). However, avid dye uptake by perivascular cells and slow penetration of DiOC₃(3) into tumour parenchyma dominate staining in vivo. DiOC₃(3) can be used for cell selection from multicell spheroids using fluorescence-activated cell sorting (Olive et al., 1986). On staining cells from solid tumours on the basis of DiOC₃(3) fluorescence has not, to date, been successful. The time required for mechanical and enzymatic dissociation of solid murine tumours into single cell suspensions and leakage of dye from cells damaged during disaggregation results in reduction of the large staining gradient initially present in intact tumours.

DiOC₃(3) is strongly fluorescent and if tumours are removed within 5 min of dye injection excellent contrast is obtained between stained and unstained tumour regions. For photomicrography or image processing, DiOC₃(3) (green fluorescence) is superior to Hoechst 33342 since CID cameras and most photographic film are maximally sensitive to green light. In addition, the dye is relatively resistant to bleaching thereby allowing prolonged illumination of the tissue section for vessel counting or fluorescence measurements. DiOC₃(3) can be successfully employed as a marker for functional tumour vasculature provided several limitations are kept in mind. Firstly, poor solubility requires the use of an organic solvent to maintain DiOC₃(3) in solution. In our studies, 50 μl of 75% DMSO solution was well tolerated by the animal and had no effect on vascular staining patterns or tumour blood flow. Secondly, although immediate tumour excision after DiOC₃(3) injection is not necessary for counting of stained regions, tumours should be removed rapidly, preferably within 5 min, since the staining gradient declines with time. Finally, the dye causes a significant and prolonged reduction in tumour blood flow as measured by laser Doppler methods. This vasoactive effect argues against the use of DiOC₃(3) as the first marker in a double staining regimen.

We have employed DiOC₃(3), in a double staining protocol together with Hoechst 33342, to detect regions of transient perfusion in murine tumours. Administration of DiOC₃(3) 20 min after Hoechst injection revealed regions of tumour vasculature with unmatched staining patterns indicative of transient vessel shutdown (Trotter et al., 1989). Approximately equal numbers of vessels opened (DiOC₃(3) staining only) and closed (Hoechst 33342 staining only) during the interval between stain injections (unpublished results). This suggests that administration of DiOC₃(3), and the resultant transient reduction in tumour blood flow, is not responsible for the staining mismatch observed.

In summary, the carbocyanine derivative DiOC₃(3) can be used to identify functional vasculature in experimental tumours. Like Hoechst 33342, the dye rapidly enters cells adjacent to the blood supply and penetrates slowly into tumour parenchyma allowing identification of the vasculature using fluorescence microscopy. The different fluorescence excitation and emission properties of DiOC₃(3) and Hoechst 33342 permit discrimination of the stains in tissue sections and thus the stains can be administered sequentially to detect regions of transient perfusion in murine tumours.

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