**In vivo** fluorescence kinetics and localisation of aluminium phthalocyanine disulphonate in an autologous tumour model

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**Summary**

Sulphonated phthalocyanines are studied as photosensitisers for photodynamic therapy of cancer. Their strong fluorescence and tumour-localising properties make them also potentially useful for detection of cancer by fluorescence. For this purpose, we have studied the fluorescence kinetics and localisation of aluminium phthalocyanine disulphonate (AlPcS₃) in 4-nitroquinoline 1-oxide (4NQO)-induced dysplasia and invasive cancer of the oral mucosa of the hard palate in Wistar albino rats. Twenty-two rats were divided into six groups. Five groups were subjected to a 4NQO application period of 8, 12, 16, 20 or 26 weeks and one was a control group. The dysplasia varied from slight to severe and was correlated with the duration of the application period. All animals received a dose of 1 μmol kg⁻¹ AlPcS₃ i.v. Fluorescence images were recorded via a specially designed ‘palatoscope’ with excitation at 460±20 nm for autofluorescence, 610±15 nm for AlPcS₃ fluorescence and detection of emission at 675±15 nm. After subtraction of the two images the specific AlPcS₃ fluorescence remained. AlPcS₃-mediated fluorescence increased significantly when the severity of dysplasia increased (P < 0.04). Also the phenomenon of strong fluorescent spots on the fluorescence images was observed. This always occurred within the first 10 h after injection of AlPcS₃. Histological analysis showed a local alteration to the mucosa in 67% of these spots, which was either invasive cancer (29%) or inflammation (38%). These results suggest two different mechanisms of AlPcS₃ uptake in tissue, one associated with the presence of generalised dysplasia and another associated with local changes of the epithelial/connective tissue, which is not necessarily specific for tumours.

**Keywords:** oral squamous cell carcinoma; photodetection; phthalocyanine; fluorescence detection; endoscope; 4-nitroquinoline 1-oxide

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Photosensitive drugs can be used for therapy and detection of cancer. The therapeutic modality is called photodynamic therapy (PDT) and the detection modality is generally referred to as photodagnosis or photodetection (PD). After administration, the ideal drug for PDT or PD accumulates preferentially in premalignant or malignant tissue. When illuminated with light of suitable wavelength and dose, the sensitisers can be excited to a singlet state, which may decay to an excited triplet state via ‘intersystem crossing’. Subsequently, a cascade of events occurs, whereby the energy of the excited photosensitiser is used to create singlet oxygen from its triplet ground state, as well as free radicals, which induce local tissue necrosis (Henderson and Dougherty, 1992). Combined with selective illumination, tumour destruction with limited damage to normal tissue is possible. If more photosensitisers are retained in tumour than normal tissue, their fluorescence can be used for tumour localisation or detection.

The sensitisers mostly used in clinical applications is a derivative of haematoporphyrin (HpD) and commercially available as Photofrin. This drug is far from ideal because it induces skin photosensitivity, which can last up to 8 weeks after administration. Also the fluorescent component of HpD or Photofrin is porphyrin in the monomeric form whereas the porphyrin dimers and oligomers are the photodynamically active components, which makes prediction of therapeutic effects more complex (Kessel, 1982; Dougherty, 1987). Therefore other drugs such as sulphonated metallophthalocyanines (MPcS₃, n = 1–4) are under investigation for use as photosensitisers in PDT (Rosenthal, 1991; Van Lier and Spikes, 1989). These dyes have several advantageous characteristics over HpD such as chemical stability, a high absorption of deeply penetrating red light and a relatively low induced skin photosensitivity (Tralau et al., 1989). A high skin photosensitivity induced by metallophthalocyanines has been reported only for caesium phthalocyanine sulphonate (CePcS) (Brasseur et al., 1987).

In general, the photochemical properties of the phthalocyanines are determined by the central metal ion. Zinc and aluminium have been proposed as suitable central metals for phthalocyanines used for PDT because of the high triplet yield and fluorescence yield (Ambroz et al., 1991; Berg et al., 1989). The insolubility of the bare phthalocyanine molecule in saline hampers its use in biological systems, but this problem can be overcome by sulphonation. The degree of sulphonation and the position of the sulphonate groups is of importance for the behaviour of the phthalocyanine molecule (hydrophilic, amphiphi1ic or lipophilic) in biological systems. Sulphonation influences the amount of uptake and the localisation in the cell (Paquette et al., 1988; Peng et al., 1991a; Chan et al., 1990). Mono-sulphonated phthalocyanines seem to be less attractive as tumour localisers but yield substantial PDT-induced necrosis whereas tetra-sulphonated phthalocyanines are good tumour localisers but yield limited tissue damage (Van Leengoed, 1993; Berg et al., 1989).

Recently, it was recently found that tetra-sulphonated zinc phthalocyanines showed a doubling of PDT-induced tumour necrosis by changing the illumination wavelength to 692 nm, according to an observed red shift in the absorption spectrum (Griffiths et al., 1994). Among these sulphonated metallophthalocyanines, aluminum phthalocyanine disulphonate (AlPcS₃) seems an interesting compound. In vitro studies showed uptake of AlPcS₃ in cells and a substantial cytotoxicity (Peng et al., 1991b; Chan et al., 1991). In vivo studies on the effect of the central metal ion and degree of sulphonation show that AlPcS₃ displays a high tumour fluorescence and an adequate tumour necrosis after illumination (Van Leengoed, 1993). AlPcS₃ seems a possible alternative for HpD as a sensitisser for clinical PDT owing to its tumour-localising and photodynamic properties and is therefore interesting for further investigations.

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Most in vivo fluorescence studies were performed with animal models in which xenografts were used to mimic the clinical situation. It is known that differences exist between the actual clinical situation and the tumour xenografts such as the presence of a fibrous layer surrounding the implanted material or other host responses (Fodstat, 1988). A tumour model that has a human counterpart will have the advantage of being clinically comparable. It has been shown that porphyrins accumulate in chemically induced premalignant and malignant tissues in animals (Crean et al., 1993; Mang et al., 1993). The tumour model used in the present study is based on the induction of squamous cell carcinomas with 4-nitroquinoline 1-oxide (4NQO) in the hard palate of the rat and closely resembles the clinical and histological appearance of human squamous cell carcinoma (Nauta et al., 1995; Prime et al., 1986). When 4NQO is applied three times a week, well-differentiated squamous cell carcinomas will develop within 26 weeks. During the application period tumours are preceded by dysplasia of the oral epithelium that varies from slight to severe, and is correlated with the duration of the application period. The whole mucosal area between the molars is dysplastic and tumours arise locally, sometimes in multiple spots. When the 4NQO application is continued more tumours arise and the existing tumours expand.

To detect sensitiser fluorescence in vivo several systems have been developed that include endoscopes connected to devices like intensified CCD cameras or photomultipliers (Profio et al., 1983; Andersson et al., 1987; Brodbeck et al., 1987; Rogers et al., 1990). Promising results have been established with endoscopic detection of tumours in clinical settings with HpD or Photofrin (Kato et al., 1992; Monnier et al., 1990), although it has been noted that most papers are case reports and that the real value of sensitiser-based PD needs to be established (Bown, 1993). We have developed an endoscope-based imaging system for the detection of sensitisers in the palate of the rat. The aims of this experiment were to study the fluorescence kinetics of AlPcS$_2$ in the 4NQO-palate tumour model and the ability of AlPcS$_2$ to localise in non-invasive epithelial disorders and squamous cell carcinoma of the mucosa of the hard palate.

Materials and methods

Photosensitiser

Aluminium phthalocyanine disulphonate was obtained from Porphyrin Products (Logan, UT, USA). The AlPcS$_2$ was prepared via the direct sulphonation method. After receiving the AlPcS$_2$ the purity was analysed by high-performance liquid chromatography in a gradient from 20% to 90% methanol. The fraction consisted of >90% pure AlPcS$_2$ of one isomer. For injection the drug was first dissolved in 0.1 M sodium hydroxide (pH 12). This solvent was diluted to an injectable volume with phosphate-buffered saline (PBS). The pH was adjusted by adding an amount of 0.1 M hydrochloric acid equal to the amount of sodium hydroxide.

Imaging system, AlPcS$_2$, excitation and fluorescence detection

A schematic drawing and description of the palatoscopy developed for the purpose of illumination and acquisition of fluorescence images is presented in Figure 1. The region of interest was the mucosa of the hard palate between the molars of the rat. With this endoscope a fairly homogenous beam was obtained, which illuminated an area of approximately 1 cm in diameter. From the centre of the beam the light intensity gradually diminished to not less than 90% at the outer part of the beam. The imaging system projected a full-screen view of the hard palate, including the molars and allowed detailed analysis of the mucosa. The images were digitised by a personal computer-based framegrabber and averaged over 16 frames (Van Leengoed, 1993). The detection range of the charge coupled device (CCD) camera was between 0 and 30 nW cm$^{-2}$ and fluorescence was detected in the linear part of the range between 10 and 25 nW cm$^{-2}$. The images were analysed by imaging software (IAS) using a pixel measurement program that allows measurement of grey-scale values of fields of interest and subtraction of different recorded images.

The technique of dual wavelength excitation (Baumgartner et al., 1987) was used to detect the AlPcS$_2$ fluorescence. The AlPcS$_2$ as purchased showed a typical aluminium phthalocyanine absorbance spectrum in the monomeric form with a minimal absorbance between 400 nm and 550 nm followed by a small peak at 590 nm to 615 nm and a large absorbance peak at 672 nm (Figure 2). Since the monomeric form is responsible for phthalocyanine fluorescence (Wagner et al., 1987) it was expected that AlPcS$_2$ fluorescence in vivo could be excited at a wavelength around 610 nm. In pilot experiments we found that autofluorescence images of the palate excited at 460±20 nm combined with a high-pass dichroic mirror (DM) of 550 nm and at 610±15 nm combined with a high-pass DM transmitting light above 650 nm did not differ very much in fluorescence intensity and pattern when the fluorescence of both excitation wavelengths was detected at 675±15 nm. After subtraction of the images of the two wavelengths, less than 10% of the original value remained (Figure 3). Also the autofluorescence images obtained at 460±20 nm did not alter after drug injection (Figure 3), whereas when excited at 610±15 nm AlPcS$_2$ fluorescence was easily detected at a dose of 1 µmol kg$^{-1}$. Satisfactory fluorescence intensities in recording images were obtained with light generated by a halogen-lamp (Ha) with an irradiance of 0.2 mW cm$^{-2}$ after passing through the excitation filter. The power of the lamp was checked regularly during the experiments but no adjustments were necessary.

Experimental procedure and assessment criteria

Squamous cell carcinomas and dysplasia were induced by an application of 4NQO three times a week. The rats were briefly anaesthetised by a mixture of nitrous oxide—oxygen—halothane and painted with 4NQO on the mucosa of the hard palate. During the application period the rats were housed under standard housing conditions. For this experiment 22 Wistar albino rats divided into five groups were used. Each
group was subjected to a different 4NQO application period, namely 8, 12, 16, 20 and 26 weeks. Two untreated animals served as controls.

Ideally, the light beam should be perpendicularly incident on the whole palatal surface. However, this is only partly possible owing to the anatomical curvature of the palate.

Also accurate repositioning of the animal is required. This was achieved by anaesthetising the animals with a mixture of oxygen–nitrous oxide–ethrane, and placing them in a stereotactic frame which was fixed on an XY-table. The endoscope was fixed to the table on a photographic standard that was adjustable in height.

After recording autofluorescence images all rats were injected with 1 μmol kg⁻¹ AlPcS₂ via a tail vein. Fluorescence images were recorded at 2, 4, 6, 8, 10, 24, 32, 52, 72, 100 h after injection. The rats were sacrificed with an intracardial overdose of pentobarbital after the last fluorescence image was recorded. The palate with the surrounding hard and soft tissues, including the skull, were removed in one piece. The palates were fixed in 4% formalin, decalcified in 4% formic acid with 0.34 M trisodium citrate dihydrate for approximately 4 weeks. The degree of decalcification was checked by X-ray analysis. Punch biopsies of 3 mm in diameter (Biopsy Punch, Stiefel, Germany) were taken at locations that displayed as strong fluorescent spots (hotspots) in the fluorescence image or that were clinically suspect for squamous cell carcinomas but did not fluoresce. The palates and biopsies were cut transversely and processed for standard haematoxylin- and eosin-stained histological sections. The slides of the biopsies and the adjacent epithelium in the palatal mucosa were examined by light microscopy and the epithelial dysplasia was assessed according to the epithelial atypia index (EAI) grading list by Smith–Pindborg by an independent observer (PGJN), without knowledge of 4NQO treatment time or of the outcome of the fluorescence measurements (Smith and Pindborg, 1969).

The average grey-scale value of the total area between the molars at every fluorescence recording after subtraction of autofluorescence was measured for all rats. The grey-scale values of the fluorescent hotspots were measured and compared with the total area measurements.

**Figure 2** Absorbance spectrum of AlPcS₂ in methanol. The arrows are positioned at the wavelength areas used for excitation of autofluorescence at 460 nm and AlPcS₂ fluorescence at 610 nm.

**Figure 3** Four digitised fluorescence images of a palate of the same 4NQO-treated rat with on each image on the left and right the molars (M) and the region of interest between the molars. (a) An autofluorescence image of the palate excited with 460 nm and (b) the autofluorescence image excited with 610 nm, both before injection of AlPcS₂. When these images are subtracted almost no autofluorescence remained (not shown). (c) The 610 nm and (d) the 460 nm fluorescence image of the same rat 6 h after injection of 1 μmol Kg⁻¹ AlPcS₂. The 460 nm image is unaltered and after subtraction of these two images clearly only AlPcS₂-mediated fluorescence remains (not shown).
Results

Fluorescence intensities and EAI of palates

The images of normal rat palate had a typical AlPcS\textsubscript{2} fluorescence pattern. A small band along the molars fluoresces strongly and the intensities decrease in the middle of the palate. This was also seen in the 4NQO-treated rats and this pattern slightly interfered with the diagnosis of hotspots because it made the interpretation of the images more complex (Figure 3). We were able to reduce the autofluorescence signal to a negligible level of approximately 5 pixels for all groups (0 h) by subtracting the 610 nm and 460 nm images. The grey-scale levels of AlPcS\textsubscript{2} fluorescence after subtraction were for the normal rats (with the lowest AlPcS\textsubscript{2} fluorescence of all groups) approximately 80 pixels. Therefore analysis of AlPcS\textsubscript{2} localisation was not hindered by autofluorescence. Two rats that were treated for 8 and 26 weeks respectively with 4NQO were lost during the experimental procedure. The fluorescence intensity of AlPcS\textsubscript{2} increased with increasing 4NQO treatment. Figure 4a shows the course of the detected AlPcS\textsubscript{2} fluorescence of the mucosa between the molars of a rat treated for 26 weeks with 4NQO. As hotspots were only present between 2 and 10 h after injection it was decided to restrict statistical analysis to that period. No hotspots were observed 24 h after injection. The area under the curve as presented in Figure 4a, between 2 and 10 h should give a good impression of the fluorescence kinetics of tumour tissue. The area was approximated by means of a weighted sum of the fluorescence measurements at times 2, 4, 6, 8 and 10 h respectively using ‘weights’ 1, 2, 2, and 1. Figure 4b shows the approximated areas vs the number of weeks of 4NQO application for every rat. An extension of the Wilcoxon rank-sum test (Cuzick, 1985; Stepniewska and Altman, 1992) was used to confirm a positive trend between the area under the curve (between 2 and 10 h for all rats) and the number of weeks of 4NQO application (P<0.04). Figure 4c shows the EAI indices vs the 4NQO application period. As was expected, a highly significant relationship is present between the 4NQO application period and the EAI (extension of Wilcoxon rank-sum test; P<0.001). In Figure 4d the approximated areas vs the EAI is plotted. The EAI was designed for assessing the severity of the dysplasia and has a maximum score of 44, but invasive squamous cell carcinoma (SCC) was not included in the grading system. We did not exclude SCC from the data analysis since it is an important feature of the model. It was previously observed that the dysplasia induced in this model will eventually lead to invasive SCC. For these reasons we assigned the number 50 for all cases of SCC. The size of this number is arbitrary, however; varying this number between for instance 50 and 100 does not affect the outcome of the analysis as the essence of the test is analysis of rank-order. The approximated areas under the curve vs the EAI showed a positive trend (extension of Wilcoxon rank-sum test; P<0.05).

This increase in fluorescence cannot be explained by variations in the individual amount of AlPcS\textsubscript{2} administered owing to differences in weight. The rats in each group were treated for different periods with a possibility of allowing the longest treatment group (26 weeks) to gain more weight. In

![Figure 4](image-url)

**Figure 4** (a) A typical example of the fluorescence intensities plotted vs the time (h) of one rat. For statistical analysis the area under the curve between 2 and 10 h (shaded area) should give a good impression of the fluorescence kinetics as we observed hotspots only in this time segment. Each data point represents the average of 16 frames gained at one time point. (b) The calculated areas under the curve for each rat are plotted against the weeks of 4NQO application. There was a significant trend (P<0.04) indicating an increasing fluorescence signal when the application period is longer. (c) The EA indices plotted against the weeks of 4NQO application (P<0.001). (d) The areas under the curve for each rat plotted against the EAI (P<0.05). We attributed the number 50 to squamous cell carcinoma (SCC). The EAI was designed for dysplasia with a maximum score of 44. In b–d the line indicates the trend of the data.
Figure 5 the average weight per group is plotted. Initially, the weight increased with prolonged treatment due to ageing, but the body weight of the rats in the group that were treated for 26 weeks was lower than the 20 weeks or 16 weeks groups, although the 26 weeks group had the highest fluorescence intensities. These rats probably lost weight as a result of their malignancy.

Fluorescence intensities and EAI of biopsies
Images of 4NQO-treated rats showed some clear fluorescent hotspots varying in diameter between 1 and 4 mm (Figure 6). These spots occurred between 2 and 10 h after injection. No hotspots were seen after this time interval. The maximum level of fluorescence differs per spot as well as the time interval between administration of AlPcS2 and the peak levels. Some hotspots had their maximum at 2 h and some at 4 or 8 h after injection. This did not only occur among different rats but also on the palate of an individual rat. A total of 21 hotspots were seen in the complete group. The fluorescence intensities of the hotspots varied considerably. The average intensity of the hotspots was 135% (s.d ± 22%) of the fluorescence intensities of total palatal area measurements in a range of 108–215%. The intensity of the spot did not reveal information about a possible malignancy. This wide range indicates that the decision about whether something can be considered a fluorescent spot can only be made on visual information, guided by the fluorescence pattern on screen.

Six of the 21 hotspots proved to be invasive squamous cell carcinoma. In another eight biopsies inflammation was present. In most cases the inflammation was caused by included hairs or dietary fibres, and macrophages, lymphocytes and foreign body giant cells were present (Figure 7). Possibly the dysplastic mucosa is easily penetrated by such fibres. In seven of the fluorescent hotspots no specific alterations were found histologically. The EAI number did not differ from the EAI number of the surrounding palatal mucosa in these biopsies. By this biopsy method a sensitivity of 67% was achieved but a tumour specificity of only 29% when squamous cell carcinoma and inflammation were regarded as alterations to the mucosa. In three rats squamous cell carcinomas were found in areas where no hotspots were seen, resulting in 15% false negatives. The spots that represented squamous cell carcinoma were not always clinically suspect for invasive cancer (Figure 8).

Discussion
In this study we found that it is possible to localise squamous cell carcinomas induced by 4NQO with AlPcS2-mediated
fluorescence. AlPcS2 proved to be a sensitive probe for alterations to the mucosa but not for tumour specific, as only 29% of the biopsies were squamous cell carcinoma. However, when the grade of dysplasia increased, the fluorescence intensities of whole palates increased as well. The EAI (or the 4NQO application period) increased monotonically with the AlPcS2 fluorescence. There is an optimum time interval for the detection of hotspots in mucosa between 2 and 10 h after injection. After this interval AlPcS2 fluorescence could still be detected, up to 1 week after injection, but did not selectively localise in tumour tissue. The technique of dual wavelength excitation is suitable for fluorescence detection of AlPcS2. Tumours on the palate induced by 4NQO have a high production of keratin and this creates a high background fluorescence on the image. By choosing the excitation wavelengths properly it was relatively easy to reduce the (keratin) autofluorescence to an insignificant level. The photochemical properties of phthalocyanines are mainly determined by the macrocycle (Rosenthal et al., 1987). Therefore this technique can be applied independently of the number of sulphonate groups.

The AlPcS2 fluorescence detected in this experiment is composed of three levels: (1) non-specific fluorescence of AlPcS2 present in the vessels, connective tissue and normal epithelial tissue; (2) AlPcS2 fluorescence in dysplastic/neoplastic epithelial tissue; and (3) small areas with high fluorescence intensities displayed as hotspots on fluorescence images. Regarding the second fluorescence level, a significant relation was found in this experimental set-up between the fluorescence intensities of whole palates and an increasing severity of dysplasia. However, to be able to discriminate between normal tissue and dysplastic tissue by numerical analysis of the fluorescence image in a clinical setting a well-calibrated detection system is needed as well as knowledge of the uptake of AlPcS2 in different types of normal tissues.

The results found in the 4NQO tumour model need to be interpreted differently from xenograft tumour models when measuring tumour to normal tissue ratios, because no clinically visible borders exist between normal tissue, dysplasia and tumour. The 4NQO treatment will make large areas dysplastic so that an abrupt border of tumour–no tumour will not be present. It was possible to establish the outline of the strong fluorescent spots of the epithelium in the 4NQO model. The variation in time interval for the fluorescence intensities of the hotspots to reach their maximum values was remarkable. Some spots were present at 2 h and had completely disappeared at 8 h whereas others started to appear at 4–6 h. We found no difference between the grade of dysplasia of the biopsies that showed no invasive carcinomas and that of the adjacent mucosa. We expected a clear correlation between the presence of severe dysplasia or invasive tumour and fluorescent hotspots but most of the positive spots were associated with inflammation of the tissue. No conclusions regarding the nature of the tissue alterations could be drawn from the appearance of the spots on the image. The invasive squamous cell carcinomas and inflammation had no histological similarities except a loss of integrity of the basal membrane. Also the presence of cells in the case of inflammation have been associated with increasing fluorescence levels (Korbelik et al., 1991). The significantly increased fluorescence levels of the complete palate measurements, owing to increasing dysplasia, and the hotspots may have a different origin. The pathways to the cell are possibly passive diffusion or endocytosis of free phthalocyanine or
LDL receptor-mediated uptake of bound phthalocyanine (Jori, 1993; Ricchelli et al., 1991; Ben-Hur et al., 1987). Which of these is the most prominent or whether other unknown pathways are responsible for sensitiser uptake remains to be established (Hamblin and Newman, 1994). An increasing turnover rate and the accompanying high lipid metabolism of dysplastic cells possibly leads to a higher phthalocyanine uptake as seen in rats with a higher EAI number. However the phenomenon of the spot fluorescence, which occurred as a result of a high local accumulation of AlPcS₂, we interpret as follows: hotspots seem to arise as a result of the improved availability of AlPcS₂ to the epithelial cells owing to loss of biological barriers like the basal membrane. By (micro)infiltration of tumour cells into the stroma or mechanical destruction and inflammation, the process of the AlPcS₂ uptake can be much more efficient for epithelial cells in areas where the barrier is lost. Likewise a higher uptake of AlPc in tumours has been correlated with the presence of greater vessel permeability (Roberts and Hasan, 1993; Poon et al., 1992). Also the insertion of polyvinyl sponges induces a high accumulation of porphyrins in these sponges (Straight and Spikes, 1985). It has often been mentioned that the mechanism for the localisation of sensitisers in tumours is based upon a longer retention time of a sensitisiser in tumour tissue than in normal tissue. Based on the kinetics of the complete palate analysis and the spot localisations seen in this study we conclude that more AlPcS₂ is taken up by tumour or dysplastic epithelium than by normal epithelium but is also more rapidly cleared from the tumour than from the normal (underlying) tissue. Tumour uptake and clearance of sensitisers probably depends on the type of tumour, the mechanism of tumour induction (autologous or xenograft) and its anatomical location (Chan et al., 1989). Possibly, the results from this study are only accurate for epithelial disorders and other abnormalities (for instance sarcomas) at the palate may interact differently with AlPcS₂.

An important question is whether an optimum interval determined by fluorescence represents the best time interval for PDT treatment of tumours. The results of this study would suggest a time interval of 4–10 h after injection may yield optimal tumour necrosis while sparing the normal tissue as much as possible. However, we expect it to be difficult to induce necrosis solely in tumour tissue because of higher normal tissue levels of AlPcS₂. In most studies of PDT effects the time interval for illumination of AlPcS₂ was chosen at 24 h after injection but not all fluorescence kinetic studies show an optimal uptake of sensitiser at that point (Chatlani et al., 1991; Biolo et al., 1991b; Peng et al., 1991b; Barr et al., 1991; Pope et al., 1991; Van Leengoed, 1993). In fact, the maximum uptake in tumour tissue varies from 1 h to 48 h after injection, probably depending largely on the type of tumour and tumour–host tissues which are being studied.

In conclusion, AlPcS₂ can be used for photodiagnosis of premalignant and malignant disorders in epithelium and a fluorescence optimum can be expected between 2 and 10 h after injection. It would be interesting to study the possible influence of the time interval between injection and illumination on the selectivity of PDT-induced tumour necrosis.

Abbreviations
- hpD: haematoporphyrin derivative; AlPcS₂, aluminium phthalocyanine disulphonate; 4NQO, 4-nitroquinoline 1-oxide; EAI, epithelial atypia index; DM, dichroic mirror; BPF, bandpass filter.

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