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# A DIFFUSION EQUATION BASED ALGORITHM FOR DETERMINATION OF THE OPTIMAL NUMBER OF FIBERS USED FOR BREAST CANCER TREATMENT PLANNING IN PHOTODYNAMIC THERAPY

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## Abstract

It is essential in interstitial Photodynamic therapy (iPDT) treatment planning to ensure a homogeneous distribution within a tumor volume using cylindrical diffusing fibers while keeping the surrounding tissue intact. Light distribution is simulated through two algorithms based on the diffusion equation assuming diffusers as light sources. The first algorithm analyzes the diffusion equation and studies the effects of different variables (optical properties, delivered power, diffuser length, and position). Next, optical properties of breast were applied to estimate the volume that receives accepted light dose from one diffuser. In the second algorithm, multiple diffusers were simulated in order to find the relation between the volume and the number of required diffusers which are needed to cover cubical or cylindrical volume with sufficient light dose. Throughout this study, real values of optical properties, clinical laser power, and treatment time were considered to evaluate sufficient light doses. This study is in agreement with previous works in that optical properties are the major factors influencing light distribution in iPDT. It is shown that for a homogeneous phantom mimicking breast cancer and cubical or cylindrical shape, the number of required fibers  $N$  equal  $W \times L$  or  $D^2$  respectively.

**Keywords:** iPDT, diffusion equation, cylindrical diffuser fiber, sufficient light dose, breast cancer.

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## АЛГОРИТМ ОПРЕДЕЛЕНИЯ ОПТИМАЛЬНОГО ЧИСЛА ВОЛОКОН ИСПОЛЬЗУЕМЫХ ПРИ ВНУТРИТКАНЕВОЙ ФОТОДИНАМИЧЕСКОЙ ТЕРАПИИ РАКА МОЛОЧНОЙ ЖЕЛЕЗЫ НА ОСНОВАНИИ ДИФфуЗИОННОГО УРАВНЕНИЯ

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## Резюме

При планировании внутритканевой фотодинамической терапии (iPDT) с использованием цилиндрических диффузных волокон важно обеспечить однородное распределение света по всему объему опухоли, сохранив при этом целостность окружающей ткани. Авторы данной статьи смоделировали распределение света с помощью двух алгоритмов, основанных на уравнении диффузии, в которых в качестве источников света используются цилиндрические диффузоры. Первый алгоритм анализирует уравнение диффузии и изучает влияние различных переменных (оптических свойств источника, применяемой мощности, длины диффузора и его положения). Затем были использованы параметры оптических свойств молочной железы для оценки объема, который рассчитывает световую дозу от одного диффузора. Во втором алгоритме было смоделировано несколько рассеивателей для нахождения соотношения между объемом и количеством рассеивателей, необходимых для покрытия кубического или цилиндрического объема достаточной световой дозой. На протяжении всего этого исследования рассматривались реальные значения оптических свойств, клинической мощности лазера и времени лечения для оценки достаточных световых доз. Это исследование согласуется с предыдущими работами в том, что оптические свойства являются основными факторами, влияющими на распределение света

при iPDT. Показано, что, для однородного фантома, имитирующего рак молочной железы, кубической или цилиндрической формы, количество требуемых волокон  $N$  равно  $W \times L$  или  $D^2$ , соответственно.

**Ключевые слова:** внутритканевая фотодинамическая терапия, уравнение диффузии, волокно с цилиндрическим диффузором, достаточная световая доза, рак молочной железы.

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## Introduction

Photodynamic therapy (PDT) combines a light sensitive compound (photosensitizer) with light to generate reactive oxygen species, which, in sufficient quantities, lead to tissue destruction [1]. The development of photoactive compounds that absorb light at longer wavelengths, with greater penetration in tissue, has led to the development of treatments for larger, deep-seated tumors [2].

The diffusion approximation to modeling light transport works well for biological tissues at near-infrared PDT treatment wavelengths, where light scattering dominates over absorption. In addition, numerical solution of the diffusion equation, as used here, allow for the modeling of light transport in systems with time frames that are feasible for clinical application. Also, diffusion theory enables quick estimation of the fluence rate distribution in a homogeneous medium [3].

Recently, computer simulations suggested that cylindrical diffusing fibers (CDFs) are more effective than flat-cut fibers in delivering the therapeutic light in iPDT especially for bulky tumors [4, 5]. B. Farina et al. [6] simulated cylindrical diffuser as linear array of ideal point sources to extract fluence rate distribution in hollow organ. Baran and Foster [7] used GPU for simulation. They modeled the diffuser based on  $\text{TiO}_2$  particles. This model predicts a fluence distribution similar to that modeled by B Farina et al. They compared the radial degradation of fluence from the diffuser for both models and demonstrated that a model based on a linear of point sources is sufficient for determining fluence distribution. Baran and Foster in Ref [5], used the same diffuser modeled by them to make comparison between numbers of cylindrical diffusing fibers and flat cleaved ones.

A number of research groups have focused their efforts on optimizing the fiber placements. T.M. Baran et al. [5] demonstrated optimization of six CDFs positions in order to create treatment plans for iPDT of (6.1×7.5×7 cm) in brain. M.D. Altschuler et al. [8] used twelve CDFs in (2.5×3×5 cm). E. Oakley et al. [9] also demonstrated a treatment plan using finite element method and eleven CDFs to deliver an accepted light dose to (5×4×4 cm) in

head and neck cancer. Those studies did not indicate to the ability of single CDF to deliver accepted light doses around it. In the current study, benefits from analyzing the diffusion equation were taken to determine the whole volume with optical properties of breast cancer that has accepted light dose of all its points. This simulation study contains two codes which are regarded as important prior steps for practical procedure that we are planning to do in the laboratory. First code provides programming description of single CDF using diffusion equation. Four available CDFs in the laboratory were simulated in the second code. Results of this study clarifies interaction between external laser power, CDF length and medium optical properties and how it could be useful to determine the required number and full length of fiber tip to cover full volume with suitable fluence rate.

## Materials and Methods

### *Tissue model and software algorithm*

Following diagrammatic analysis of equation 2 assumes a cube ( $5^3 \text{ cm}^3$ ), divided into fractional ( $0.5^3 \text{ cm}^3$ ) cubes, fractional cubes could be smaller if requested. CDF position is the central cube ( $x, y, z = (2.5, 2.5, 5)$ ). While assuming that the decrement of CDF length starts from the top surface of the cube, the coordinate ( $z$ ) must be equal to the full length of the corresponding cube. Figure 1 represents the tissue model. The core of the PDT treatment planning simulation is the calculation of the light intensity (fluence rate) distribution in tissue generated by a CDF. Therefore, first code is made as flexible as to calculate the fluence rate  $\phi(r)$  as a function of each variable ( $h, l, s, \mu_{\text{eff}}, \mu_a$ ) independently in every point (pixel) in the cube. The second code allows choosing CDFs number (maximum is four). In addition to changing  $l, s, \mu_{\text{eff}}, \mu_a, t$ , volume and distance between pixels. Figure 2 displays the schematic of the codes. The light distribution, diffusion equation and tissue model were modeled in Matlab (2016a).

Throughout the paper, each observing point receives  $N$  rays.  $N$  represents the number of source points along the CDF. The points located at equal distances away from

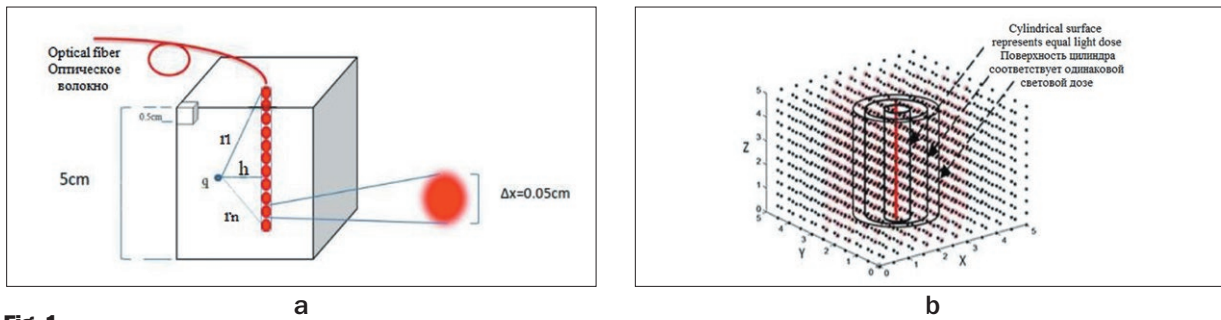


Fig. 1.

a – Cube, fractional cubes, CDF, point source and observing points are the components of the tissue model. Each point receives  $N$  rays.  $N$  is the number of total source points along the CDF (Dimensions are for clarification and they are not no scale);  
b – Cylindrical surfaces represent points that have equal fluence rate

Рис. 1.

а – Куб, дробные кубы, оптическое волокно с цилиндрическим диффузором (CDF), точечный источник и точки наблюдения являются компонентами модели ткани. Каждая точка получает  $N$  лучей.  $N$  – это общее количество точек-источников вдоль CDF (размеры даны для пояснения, и они не в масштабе);  
б – Цилиндрические поверхности представляют точки с одинаковой скоростью потока

CDF receive equal light dose and form a cylindrical surface around CDF (fig. 1b).

#### Laser treatment fiber

This study provides profiling of optical fiber which is made from RD-ML50 (Medlight S.A, Switzerland). Treatment fiber specifications are as follows: transmission of 630–760 nm, overall diameter 1 mm, illumination length of 50 mm, maximum (CW) power density (in air) of 0.5 W/cm, absolute maximum input power of 2.0 W (CW). Treatment regions: breast, prostate, brain, heart, lung and diaphragm. The diffuser length is 5 cm or less (in increments of 0.5 cm); density power, W/cm and energy, J/cm were specified for each fiber) [10].

#### Diffusion theory

Cylindrical diffuser fiber was modeled in two geometries. Two equations can be found in the literature that estimate the fluence rate emitted from a light source of a length  $l$  in heterogeneous/homogeneous media (equations 1 and 2). Equation 1 discretizes the diffusing part of the optical fiber as a sum of several point light sources [11–13]. Equation 2 considers the whole fiber as a finite line light source with 2D cylindrical light emission characteristics [14]. The first equation 1 is considered as complex equation to estimate the fluence rate, on the basis that the calculation of the fluence rate at a distance  $r$  from the fiber is the sum of each light source contribution. Whereas equation 2 computes the fluence rate values using the minimal distance  $r$  from the fiber.

$$\varphi(r) = \frac{3sl\mu'_s}{4\pi} \cdot \frac{1}{N-1} \cdot \sum_{i=1}^N \frac{e^{-\mu_{eff}r_i}}{r_i} \quad (1)$$

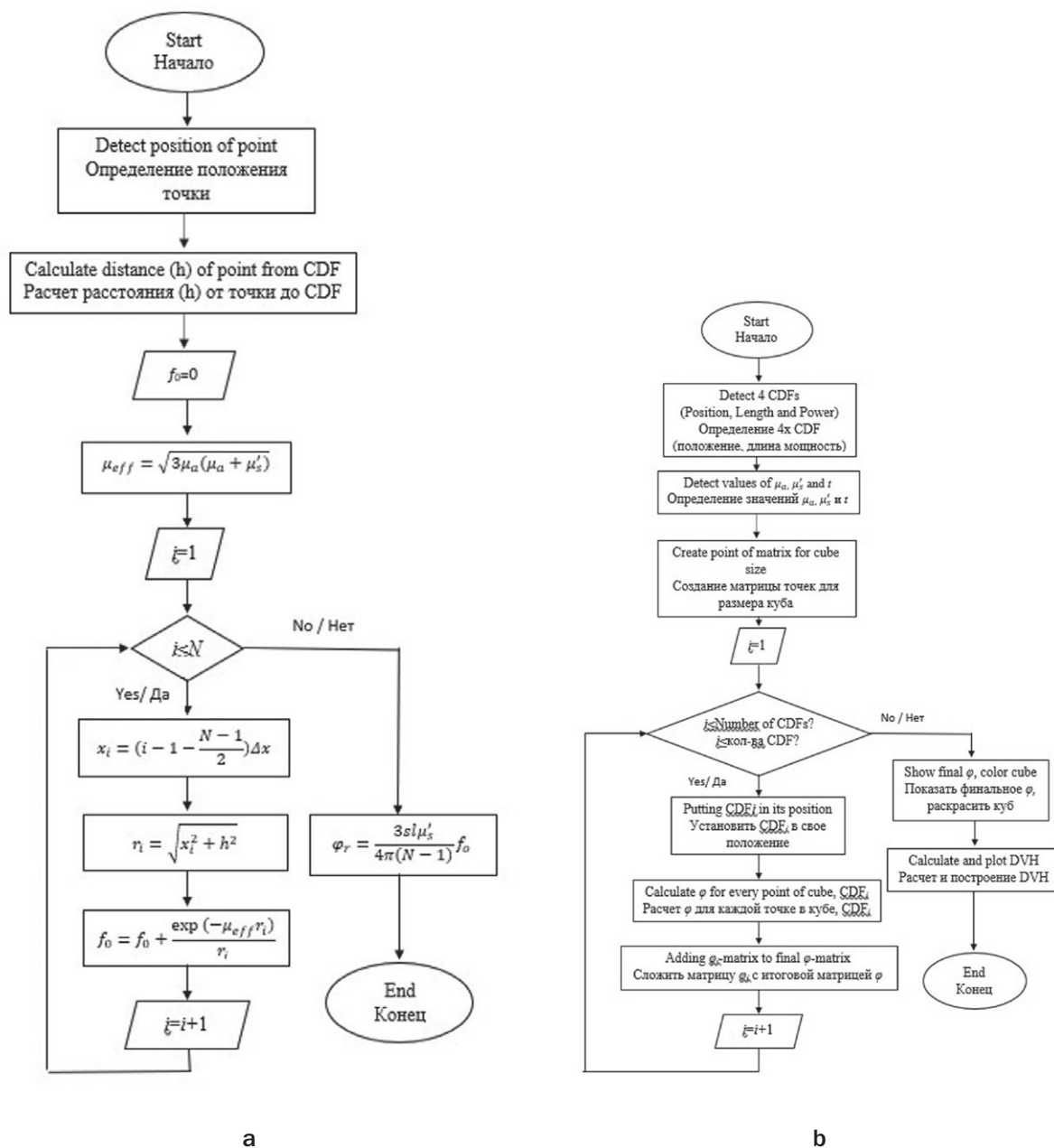
$$\varphi(r) = P \sqrt{\frac{2\delta}{\pi r}} \cdot \frac{e^{-r/\delta}}{2\pi \cdot \mu_a \cdot \delta^2} \quad (2)$$

Where  $P$  is power of one point,  $D = \frac{1}{3(\mu_a + \mu'_s)}$ ,  $\delta$  is the optical penetration depth in m and is equal

to  $\sqrt{D/\mu_a}$ ,  $s$  is the power of the point source in mW/cm;  $\varphi(r)$  is the fluence rate in mW/cm; the quantity  $\mu_{eff} = \sqrt{3 \cdot \mu_a \cdot (\mu_a + \mu'_s)}$  is the effective attenuation coefficient in tissues. The differential  $\Delta x = l/(N-1)$  is the length of the elemental (discretized) source segment. The odd integer  $N$  is the number of points used in the summation over the source, with one point always placed in the middle of the CDF. The distance between the  $i_{th}$  point of the linear light source and the observing point is  $r_i = \sqrt{x_i^2 + h^2}$ , where  $x_i = [(i-1)-(N-1)/2] \cdot \Delta x$  is the cylindrical coordinate along the fiber from the center of the linear source.  $h$  is the distance between the observing point and the fiber axis as it is shown in fig. 1a. The numerical value of the summation should be independent of  $N$  (or  $\Delta x$ ) if  $N$  is large enough. Accurate results of the summation can be obtained if  $\Delta x = 0.05$  cm. In this study, summing  $N = 101$  is available [8, 15]. Obviously, equation 2 is simpler than 1. Depending on geometrical optics equation 1 could be regarded as more accurate method than the equation 2. That is why fluence rate is estimated using the equation 1 throughout this study. For simplicity, we use the light fluence (fluence rate  $\times$  exposure time) for the PDT dose throughout the paper. The illumination time, through this study, was not fixed (300 sec in the first algorithm and 150 sec in the second algorithm). Final light dose at each point is the summing of light doses that were received from all diffusers. Specifically, tissue necrosis occurs above a threshold light dose [16]. The higher threshold of target dose is determined by minimizing damage to surrounding tissue. To make this study closer to reality, thresholds light dose are assumed to be 20–50 J/cm<sup>2</sup> and 90–300 J/cm<sup>2</sup> with optical properties of breast and prostate cancer, respectively [8, 17].

#### Optical properties of the studied tissues and patient's prostate

Optical properties at 732 nm in human prostate before and after PDT are taken from Altschuler et al. [8] and listed in table 1. While Optical properties of heart, breast,



**Fig. 2.** The block diagram illustrating two software planning algorithms:

a – Algorithm code for analyzing diffusion theory and CDF specification;

b – Algorithm code to calculate total fluence light and plotting color cube and Dose Volume Histogram (DVH)

**Рис. 2.** Блок-схема, иллюстрирующая два алгоритма программы:

a – Алгоритм кода для анализа теории диффузии и спецификации цилиндрических диффузоров (CDF);

b – Алгоритм кода для расчета общей плотности света и построения цветового куба и гистограммы доза-объем (DVH)

Table 1

Optical properties at 732 nm in human prostate gland (before and after PDT) for patient number 3 and 5 [8]

Таблица 1

Оптические свойства предстательной железы (до и после ФДТ) для 732 нм пациентов 3 и 5 [8]

| Patient Number<br>Номер пациента | Before PDT<br>До ФДТ    |                          | After PDT<br>После ФДТ |                          |
|----------------------------------|-------------------------|--------------------------|------------------------|--------------------------|
|                                  | $\mu_a, \text{cm}^{-1}$ | $\mu_s', \text{cm}^{-1}$ | $\mu_a$                | $\mu_s', \text{cm}^{-1}$ |
| 3                                | 0.15                    | 22.0                     | 0.07                   | 33.4                     |
| 5                                | 0.21                    | 11.8                     | 0.13                   | 7.18                     |

Table 2

Optical properties of human heart, breast, lung, diaphragm tissues [11]

Таблица 2

Оптические свойства тканей сердца, молочной железы, легких, диафрагмы человека [11]

| Tissue<br>Ткань                                | $\lambda, \text{nm}$ | $\mu_a, \text{cm}^{-1}$ | Average<br>Усредненное | $\mu_s', \text{cm}^{-1}$ | Average<br>Усредненное |
|--|----------------------|-------------------------|------------------------|--------------------------|------------------------|
| Heart<br>Сердце                                | 661                  | [0.18–0.12]             | 0.15                   | [5.22–90.80]             | 48                     |
| Breast (normal)<br>Молочная железа<br>(норма)  | 660                  | [0.037–0.11]            | 0.074                  | [11.4–13.5]              | 12.45                  |
| Breast (tumor)<br>Молочная железа<br>(опухоль) | 690                  | [0.07–0.1]              | 0.085                  | [14.7–17.3]              | 16                     |
| Lung<br>Легкое                                 | 661                  | [0.49–0.88]             | 0.68                   | [21.14–22.52]            | 21.83                  |
| Diaphragm<br>Диафрагма                         | 661                  | [0.15–1.08]             | 0.62                   | [9.65–21.7]              | 15.7                   |

lung, diaphragm tissues are taken from Julia J.L. Sandell and T.C. Zhu review [11] and given in table 2.

## Results and discussion

The present study was independent of allowed illumination/dark time or oxygen consumption. Each factor in equation 1 has a different effect. Throughout this part increment in distance is 0.125 cm. Illumination treatment time is 5 min. Source power is 0.5 mW/cm. The following discussion presents four effects.

### Distance perpendicular to the fiber axis ( $h$ )

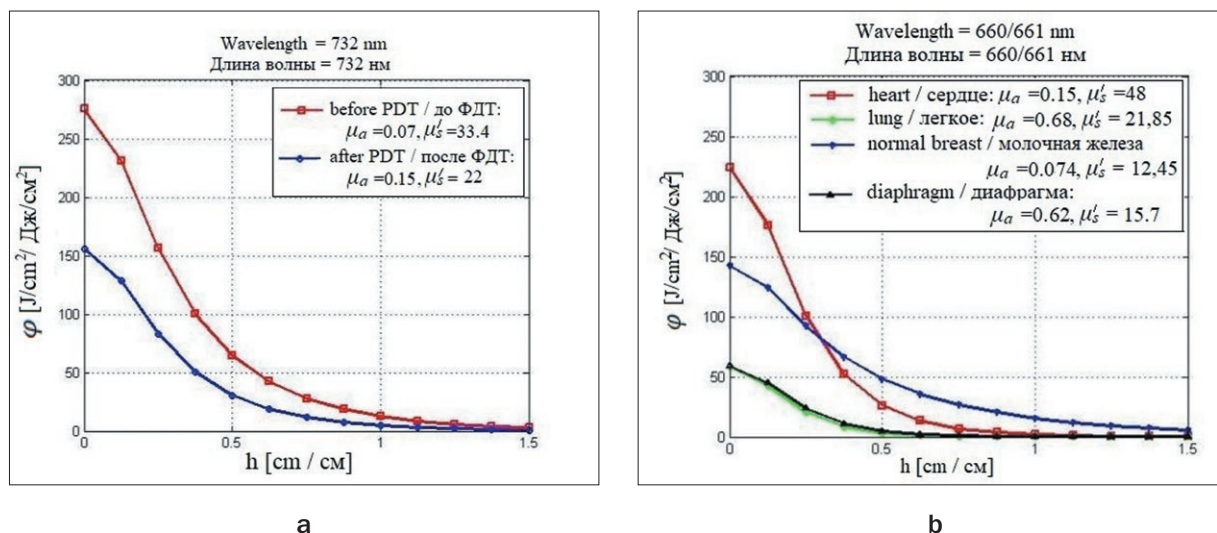
Figure 3a shows the variation of calculated light fluence distribution for different distances in the same prostate gland (optical properties are taken from table 1). The average optical properties were very different between the upper and lower line. The light fluence rate at 0.5 cm far away from the CDF varied from 64.59 J/cm<sup>2</sup> (before PDT) to 30.5 J/cm<sup>2</sup> (after PDT). Significant change of optical properties was observed before and after PDT treatment in the same patient number 3. Figure 3b shows the variation of calculated light fluence distribution for different distances in heart, breast, lung and diaphragm (optical properties are mentioned in the table 2). As a result, closed values of fluence rate were observed when

optical properties approximately the same (of lung and diaphragm). Absorption coefficient of breast is the smallest, so it allows higher distribution light ( $\varphi=50 \text{ J/cm}^2$  at 0.5 cm away from CDF). From figure 3b; the values of  $\varphi$  are 50 J/cm<sup>2</sup> at 0.375 cm for heart, and 50 J/cm<sup>2</sup> at 0.5 cm for breast. Thus, practically two CDFs are required to deliver 100 J/cm<sup>2</sup> for points at those distances. In contrast, for lung or diaphragm two CDFs deliver accepted fluence rate 90 J/cm<sup>2</sup> at 0.125 cm only. More explanation is provided in following absorption and scattering coefficients effects.

### Absorption and scattering coefficients effects

Remarkable advances were made in the last two decades to determine the *in vivo* optical properties in humans in a variety of organs. The absorption coefficient varies largely over the visible spectrum, while the scattering coefficient of tissue decreases monotonically as the wavelength increases [18]. The presence of chromophores affects the absorption coefficient. Although the actual range of the *in vivo* optical properties is tissue type dependent, generally varied in the ranges  $\mu_a=0.03\text{--}1.6 \text{ cm}^{-1}$  and  $\mu_s'=1.2\text{--}40 \text{ cm}^{-1}$  [11]. Since equation 1 is a nonlinear equation of two parameters  $\mu_a$  and  $\mu_s'$ , it was very difficult to separate effects of  $\mu_s'$  and  $\mu_a$  from





**Fig. 3.** Fluence rate curve plotted at different distances:

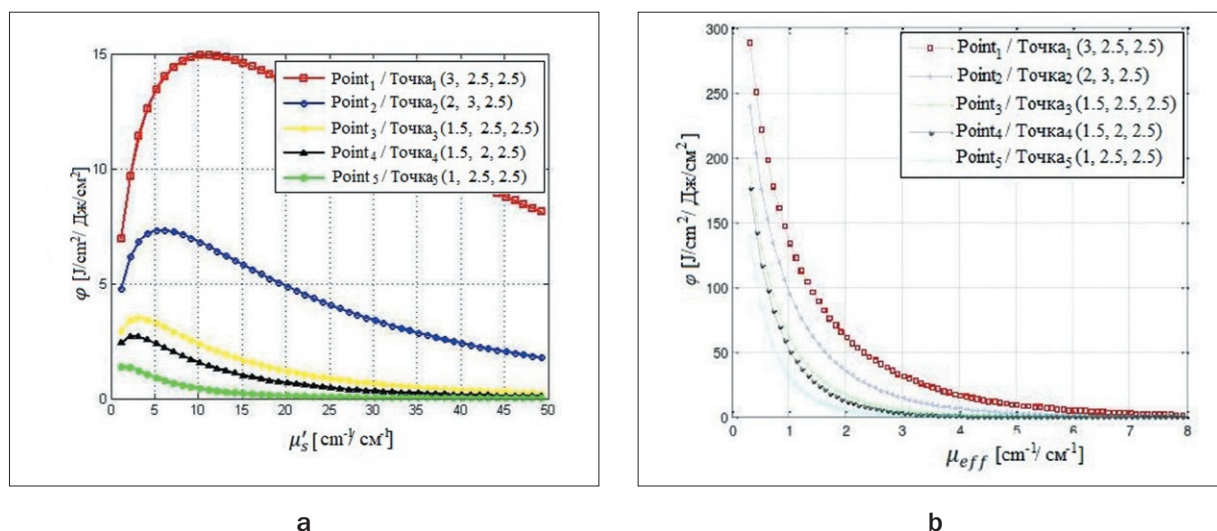
a – in prostate gland;

b – for other organs

**Рис. 3.** Кривая светового потока на разных расстояниях:

a – в предстательной железе;

b – для других органов



**Fig. 4.** The dependence of light fluence rate at different points from:

a –  $\mu'_s$ ;

b –  $\mu_{eff}$

**Рис. 4.** Зависимость интенсивности светового потока в разных точках от:

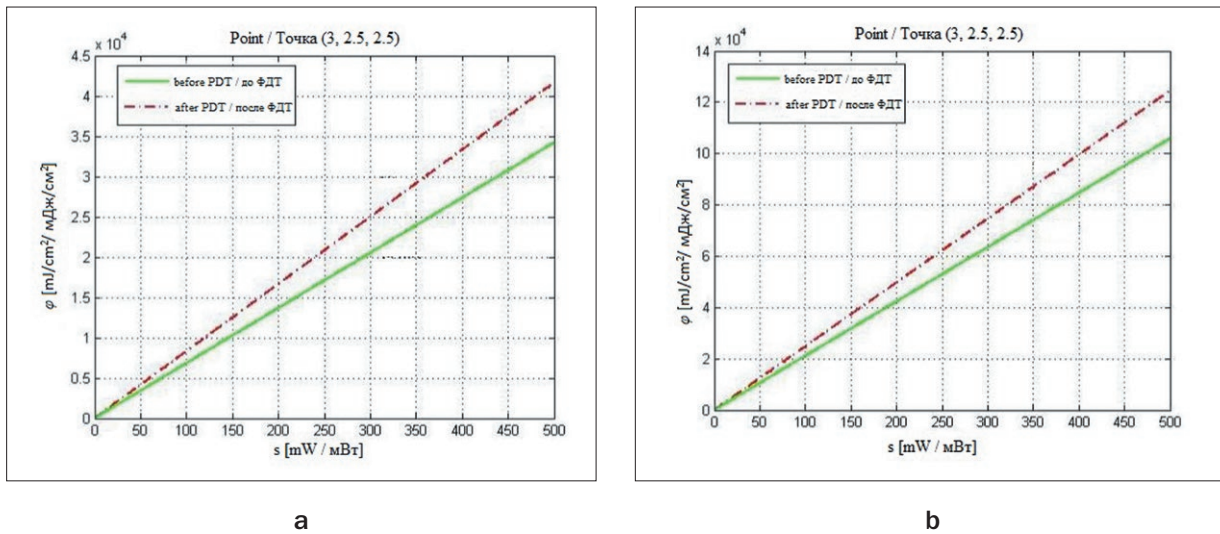
a –  $\mu'_s$ ;

b –  $\mu_{eff}$

each other. Now, the studied medium is supposed to be heterogeneous, and five observing points are considered. Therefore, to understand the effective attenuation coefficient effect, it is required to fix  $\mu'_s$  at 20.6 cm<sup>-1</sup> (is the average of its range) and  $\mu_{eff}$  is changing along 0.33–14.13 cm<sup>-1</sup>. We used these values to understand the expected standard behavior of optical properties, and to anticipate

the variation in tissue constitution that yields the tissue optical properties at any desired wavelength [19].

In the opposite case,  $\mu'_s$  is changed along its range and  $\mu_{eff}$  is fixed at value (7.23 cm<sup>-1</sup>); the average of its range. To explain  $\mu_{eff}$  and  $\mu'_s$  effects, five observing points were taken at the distances ( $h_1=0.5, h_2=0.7, h_3=1, h_4=1.12, h_5=1.5$ ) cm away from CDF. These five points represent



**Fig. 5.** Fluence rate as a function of power density  $s$ :

a – Before PDT:  $\mu_s' = 11.8 \text{ cm}^{-1}$ ,  $\mu_a = 0.21 \text{ cm}^{-1}$ ; after PDT:  $\mu_s' = 7.18 \text{ cm}^{-1}$ ,  $\mu_a = 0.13 \text{ cm}^{-1}$

b – Before PDT:  $\mu_s' = 22 \text{ cm}^{-1}$ ,  $\mu_a = 0.15 \text{ cm}^{-1}$ ; after PDT:  $\mu_s' = 33.4 \text{ cm}^{-1}$ ,  $\mu_a = 0.07 \text{ cm}^{-1}$

**Рис. 5.** Интенсивность оптического потока как функция плотности мощности  $s$ :

a – До ФДТ:  $\mu_s' = 11.8 \text{ см}^{-1}$ ,  $\mu_a = 0.21 \text{ см}^{-1}$ ; после ФДТ:  $\mu_s' = 7.18 \text{ см}^{-1}$ ,  $\mu_a = 0.13 \text{ см}^{-1}$

b – До ФДТ:  $\mu_s' = 22 \text{ см}^{-1}$ ,  $\mu_a = 0.15 \text{ см}^{-1}$ ; после ФДТ:  $\mu_s' = 33.4 \text{ см}^{-1}$ ,  $\mu_a = 0.07 \text{ см}^{-1}$

the nearest five cylindrical surfaces. Figure 4 shows the relationship between the light fluence rate and the optical properties.

Figure 4a shows that the fluence rate (for point 1) rapidly increases between 1.2 and  $10 \text{ cm}^{-1}$ , then approximately fixed between 10 and  $15 \text{ cm}^{-1}$ . As a result, reduced scattering coefficient does not cause increase in fluence rate after value  $10 \text{ cm}^{-1}$ . For point 2, decline in fluence rate occurs after  $7 \text{ cm}^{-1}$ . Although the difference in distance is only 2 mm between point 1 and 2, there is a large difference in fluence rate between them. For other points higher values of fluence rate are along  $1.2\text{--}5 \text{ cm}^{-1}$ . As a result, in experimental phantom, for higher distances, it is good to make  $\mu_s' \leq 10 \text{ cm}^{-1}$ . For larger distances (0.7, 1, 1.12, 1.5 cm), it is better to make  $\mu_s' \sim 5, 3.2, 3.2, 1.2 \text{ cm}^{-1}$ , respectively. Figure 4b shows the relationship between the light fluence rate and the effective attenuation coefficient as exponential decay curves. As shown in figure, it is clear that values corresponding to point 1 (0.5 cm) have higher fluence rate values, while lower lines represent other points respectively. For  $\phi(r)$  that was lower than  $90 \text{ J/cm}^2$ ;  $\mu_{\text{eff}}$  is  $1.53 \text{ cm}^{-1}$  for  $h_1$ . For other points  $\mu_{\text{eff}}$  was 1, 0.73, 0.6,  $0.5 \text{ cm}^{-1}$ , respectively. Two algorithms allow to change ranges and values of  $\mu_s'$  and  $\mu_{\text{eff}}$  to get accepted light dose and could be useful in preparing phantoms at laboratories. As a result, knowledge of optical properties of tissues is of great importance for interpretation and quantification of the diagnostic data. Fluence rate at virtual diffuser points depends on optical properties of studied medium. It was  $\sim 87.7 \text{ J/cm}^2$  with prostate optical properties  $\mu_s' = 14.3$ ,  $\mu_a = 0.3 \text{ cm}^{-1}$ . While it became

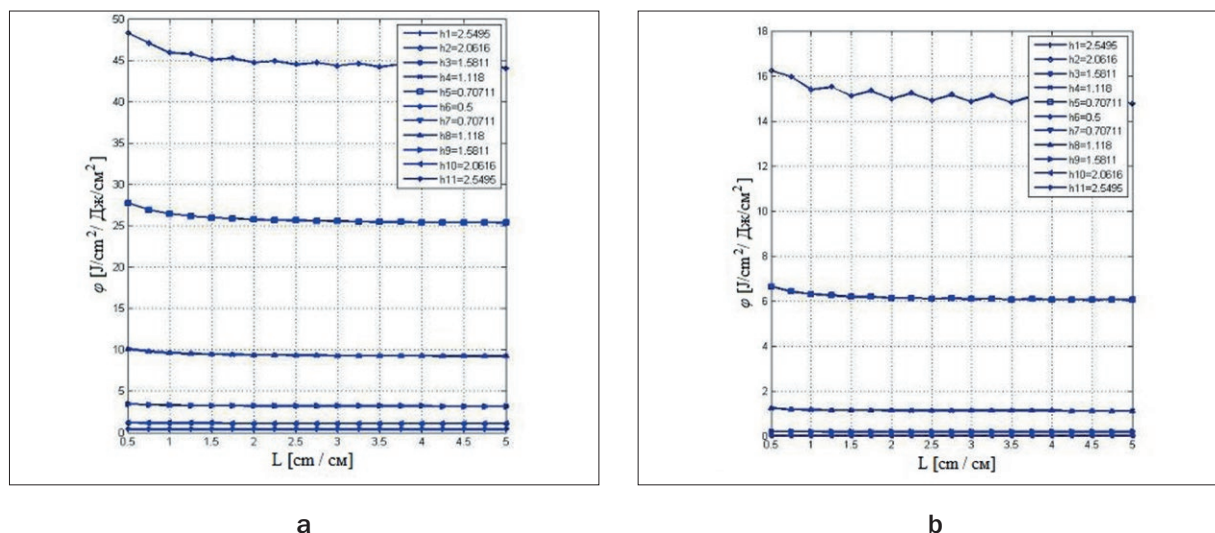
$199.5 \text{ J/cm}^2$  when  $\mu_a$  decreases by ten times. It is well known that absorption coefficient is included in  $\mu_{\text{eff}}$  expression. From our findings and for the optical properties of all biological tissues studied in ref. [11], it was found that the absorption coefficient  $\mu_a$  has a more significant influence on light fluence rate distribution than reduced scattering coefficient  $\mu_s'$ . It is noteworthy to say that our results are valid for wavelength range 600–800 nm at certain wavelengths of human tissues that was also mentioned in ref. [11].

#### Power density effects ( $s$ )

Source catheters are used for both light delivery and measurement of optical properties. Usually, the light dose is given in terms of external power density delivered by the light system. As it is noted in fiber specifications,  $s$  is  $0.5 \text{ mW/cm}$  at maximum allowed input power 2W. The power is considered to be  $0.5 \text{ mW/cm}$  through all previous study. Consequently, power of full diffuser length is 2.5 W. Maximum power density (in air) is  $500 \text{ mW/cm}$  (CW) as mentioned in the technical specifications of Medlight CDFs [10], so the power is varied between 0 and  $500 \text{ mW/cm}$  through the two codes. Optical properties of patients were taken from ref. [8]. As can be seen in figure 5a, optical properties of patient number 5 were  $\mu_s' = 11.8 \text{ cm}^{-1}$ ,  $\mu_a = 0.21 \text{ cm}^{-1}$  (before PDT). After PDT, the values changed to  $\mu_s' = 7.18 \text{ cm}^{-1}$ ,  $\mu_a = 0.13 \text{ cm}^{-1}$ . Fluence rate became larger after PDT [8].

Figure 5b shows linear relationship between  $s$  and  $\phi(r)$  that to deliver accepted fluence rate  $90 \text{ J/cm}^2$ , applied power should equal at least  $\sim 432 \text{ mW/cm}$ , while after PDT it decreased to  $369 \text{ mW/cm}$ . If  $s$  is limited at 250, flu-





**Fig. 6.** Fluence rate as function of diffuser length  $L$ .  $s = 0.5 \text{ mW/cm}$ ,  $\mu_s' = 14 \text{ cm}^{-1}$ ,  $t = 5 \text{ min}$ :

a –  $\mu_a = 0.09 \text{ cm}^{-1}$

b –  $\mu_a = 0.3 \text{ cm}^{-1}$

**Рис. 6.** Интенсивность оптического потока как функция длины диффузора  $L$ .  $s = 0.5 \text{ мВт/см}$ ,  $\mu_s' = 14 \text{ см}^{-1}$ ,  $t = 5 \text{ мин}$ .

a –  $\mu_a = 0.09 \text{ см}^{-1}$

b –  $\mu_a = 0.3 \text{ см}^{-1}$

ence rate is  $50 \text{ J/cm}^2$  (before) and  $61.5$  (after). It was clear that fluence rate became higher after the PDT. Results for patients 2, 4, 12 and 9 [8] confirm that  $\mu_a$  has a significant effect on light fluence rate distribution.

#### Cylindrical diffuser length effect

Returning to equation 1; the fluence rate clearly appears proportional to the tip length. Increasing the fluence rate is related to the density of number of source points over the diffuser ( $N$ ). As a result, the fluence rate at certain point is not affected by varying tip length. This result is illustrated by figure 6.

Figure 6 shows the relationship between  $\phi$  and  $L$ . The CDF was positioned at cube center, each line represented two opposite distances. Distances furthest from CDF were shown in small rectangle. As it was expected, light fluence rate was not affected by increasing length, for similar inputs and distances, but if  $\mu_a$  is changed to  $0.3 \text{ cm}^{-1}$ , then fluence rate will decrease to about  $15 \text{ J/cm}^2$  at  $h = 0.5 \text{ cm}$  while it is about  $45 \text{ J/cm}^2$  with  $\mu_a = 0.09 \text{ cm}^{-1}$ .

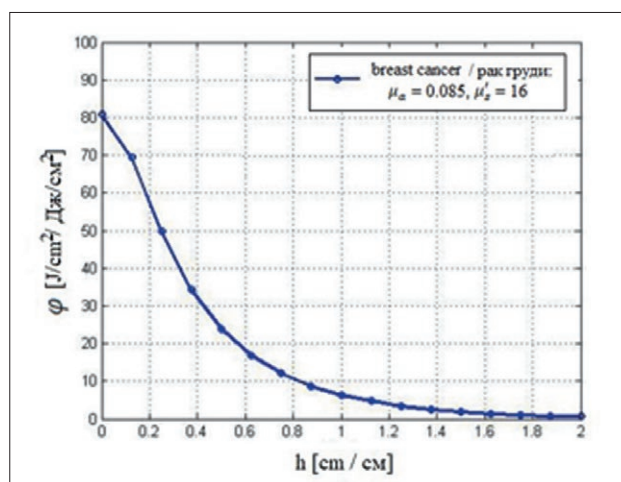
Clearly there is a big difference between  $\phi(h6)$  and  $\phi(h5)$ . As a result, there is major attenuation in fluence rate as distance increases.

Fluence rate calculations were done using basic diffusion equation (eq. 1). As it is well known, the generating of matrixes in order to calculate fluence rate at each point is a cumbersome procedure. Conditions were determined to accept fluence rate. This condition could be changed as required for a given tissue, photosensitizer or wavelength. Our results agreed with prior works where optical properties have considerable effect on light distribution in any given medium [11]. However, in real

clinical producers, CDF length cannot be changed and practically, due to fixed distances between slots used for guiding PDT fibers inside the tissue which equal  $0.5 \text{ cm}$  on average [8], distance between tow CDFs could not be less than  $0.5 \text{ cm}$ .

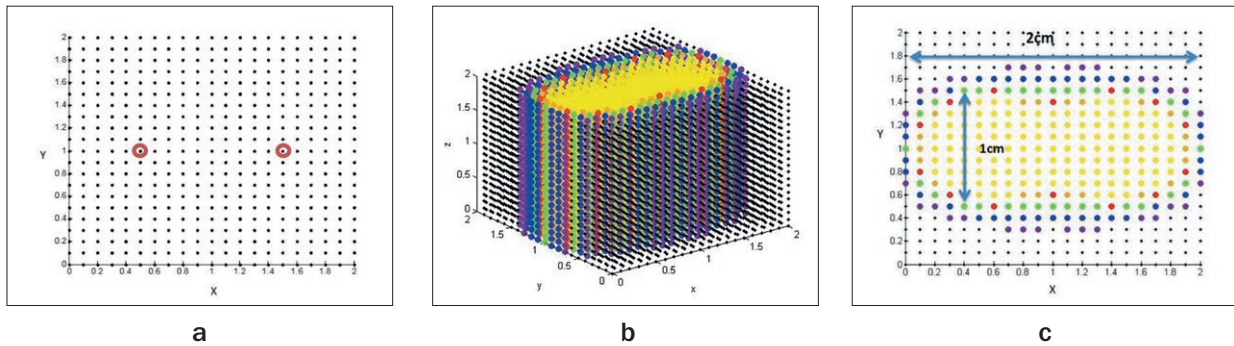
#### Volume versus number of CDFs

Actual input parameters had been taken into following consideration. Optical properties of breast cancer are  $\mu_a = 0.085 \text{ cm}^{-1}$ ,  $\mu_s' = 16 \text{ cm}^{-1}$  [11]. First algorithm is used to



**Fig. 7.** Fluence rate is as a function of distance for breast cancer tissue. Input parameters are:  $s = 50 \text{ mW/cm}$ ,  $t = 150 \text{ sec}$ . Light dose at  $h = 0.5 \text{ cm}$  is  $24 \text{ J/cm}^2$  for breast

**Рис. 7.** Интенсивность светового потока как функция расстояния для ткани рака молочной железы. Входные параметры:  $s = 50 \text{ мВт/см}$ ,  $t = 150 \text{ сек}$ . Световая доза при  $h = 0.5 \text{ см}$  равна  $24 \text{ Дж/см}^2$  для молочной железы



**Рис. 8.** Light distribution resulted from two CDFs with  $s = 50 \text{ mW/cm}$ ,  $t = 150 \text{ sec}$ ,  $\mu_a = 0.085 \text{ cm}^{-1}$ ,  $\mu_s' = 16 \text{ cm}^{-1}$ :

a – Top-down view shows positions of CDFs placed at  $(0.5, 1, 2)$ ,  $(1.5, 1, 2)$ ;

b – Yellow points have accepted dose and form volume  $1 \times 2 \times 2 \text{ cm}$ ;

c – Top-down view of cube after light distribution: light dose in red points is less than  $20 \text{ J/cm}^2$  and more than  $15 \text{ J/cm}^2$ , green points represent pixels which have dose that is less than  $15 \text{ J/cm}^2$  and more than  $10 \text{ J/cm}^2$ , blue/ violet points have the dose that is less than  $10 \text{ J/cm}^2$

**Рис. 8.** Распределение света, полученное при использовании двух CDF при  $s = 50 \text{ мВт/см}$ ,  $t = 150 \text{ с}$ ,  $\mu_a = 0,085 \text{ см}^{-1}$ ,  $\mu_s' = 16 \text{ см}^{-1}$ :

a – Положение CDF, размещенных в точках  $(0.5, 1, 2)$ ,  $(1.5, 1, 2)$  (вид сверху);

b – Желтым цветом выделены точки, получившие необходимую световую дозу. Их объем составляет  $1 \times 2 \times 2 \text{ см}$ .

c – Распределение света (вид сверху): доза света в красных точках составляет менее  $20 \text{ Дж/см}^2$  и более  $15 \text{ Дж/см}^2$ , зеленые точки представляют пиксели с дозой менее  $15 \text{ Дж/см}^2$  и более  $10 \text{ Дж/см}^2$ , синие/фиолетовые точки имеют дозу менее  $10 \text{ Дж/см}^2$

estimate fluence rate along  $2.5 \text{ cm}$  far away from single CDF (full length of tip is  $5 \text{ cm}$ ) is placed at center of cube ( $5 \times 5 \times 5 \text{ cm}$ ). Initial position of CDF is ( $x=2.5 \text{ cm}$ ,  $y=2.5 \text{ cm}$ ,  $z=5 \text{ cm}$ ); since  $x$ ,  $y$  can be changed along  $X$  and  $Y$  axes from  $0.5 \text{ cm}$  to  $5 \text{ cm}$ , while  $z$  must be set at the full length of CDF.

Figure 7 shows that sufficient light dose is reached for approximately  $0.25\text{--}0.5 \text{ cm}$  around CDF. Higher threshold of  $50 \text{ J/cm}^2$  is exceeded at  $h \leq 0.25 \text{ cm}$ . As a result, for optical properties of breast cancer, accepted light dose reaches to volume of one cylinder with full diameter of  $D = 1 \text{ cm}$ . Figure 8 is the result of second algorithm. Figure 8a shows two CDFs in a line. As can be seen from figure 8a, two CDFs can cover two centimeters in one direction of volume. While approximately only one centimeter in the vertical orientation could be covered with light dose of  $20\text{--}50 \text{ J/cm}^2$ .

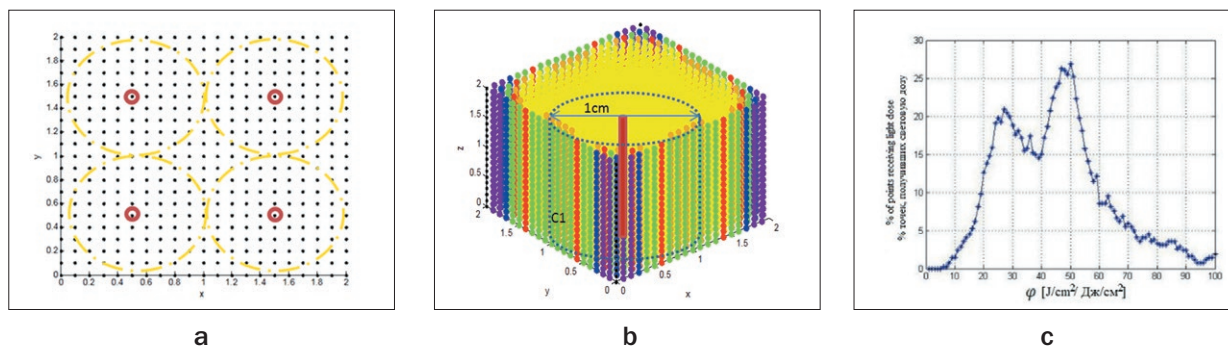
Length of CDFs must be near the height of the volume. For the breast cancer tissue, the covered volume equals approximately  $1 \times N \times L \text{ (cm)}$  as represented in figure 8c.  $N$  is the number of CDFs,  $L$  is the length of diffusers. The volume (with accepted light dose) is  $1 \times 2 \times 2 = 4 \text{ cm}^3$ . To cover full cube size which is  $2 \times 2 \times 2 \text{ (cm)}$ , four CDFs must be placed at  $(0.5, 0.5, 2)$ ,  $(0.5, 1.5, 2)$ ,  $(1.5, 1.5, 2)$ ,  $(1.5, 0.5, 2)$  (fig. 9a). The maximum covered volume with accepted light dose equals four times of cylinder  $c1$  (cylinder  $c1$  is indicated in figure 9b).

It can be concluded that for delivering sufficient light dose ( $20\text{--}50 \text{ J/cm}^2$ ) to any phantom which has cuboid or cylindrical shape and has optical properties of breast cancer, it is needed to place ( $W \times L$ ) or ( $D^2$ ) CDFs respectively

with distance ( $1 \text{ cm}$ ) between each two CDFs. Where,  $W$ ,  $L$  are the rectangular sides and  $D$  is the cylinder diameter. Practically, optimization theories are applied to optimize places of a lower number of CDFs [5, 8]. Finding an algorithm to choose suitable powers represents another practical solution to reduce required fibers.

#### Dose-volume histogram

A DVH, more properly called a cumulative dose volume histogram, expresses the percentage of an organ that has received more than a given dose [20]. It is useful to summarize the volumetric distribution of light doses. Second code allows plotting DVH (fig. 9c) that corresponds to figure 9b. For higher accuracy; distance between pixels has been reduced to  $0.05 \text{ cm}$ . Consequently, number of slices is 40 slices ( $2/0.05 \text{ cm}$ ); total considered points are  $40 \times 40 \times \text{number of slices}$ . Plotting the prism with finer voxels causes much longer time to get  $\phi_{total}$  matrix (final fluence rates resulting from four CDFs), while the time needed to calculate the DVH is only a second or so because no feasibility procedures are involved. Using DVH, it is possible to estimate percentage of points which accumulated sufficient light doses. With optical properties of breast cancer, sufficient light doses are attained in more than 95% of all cube points. Figure 9c shows DVH. Another case is studied with optical properties of brain cancer ( $\mu_a = 0.2 \text{ cm}^{-1}$ ,  $\mu_s' = 5 \text{ cm}^{-1}$ ) [5] (all other inputs are fixed). Results indicate that percentage of the covered volume with accepted light dose is reduced to 10% of the number of pixels with brain cancer ( $\mu_a = 0.2 \text{ cm}^{-1}$ ,  $\mu_s' = 5 \text{ cm}^{-1}$ ). Low level of percent coverage imposes using much more CDFs. T.M. Baran and



**Рис. 9.** Light distribution from four CDFs with  $s = 50 \text{ mW/cm}$ ,  $t = 150 \text{ sec}$ ,  $\mu_a = 0.085 \text{ cm}^{-1}$ ,  $\mu_s' = 16 \text{ cm}^{-1}$ :

a – Four CDFs placed at (0.5, 0.5, 2), (0.5, 1.5, 2), (1.5, 1.5, 2), (1.5, 0.5, 2);

b – The distributed light forms a volume that is approximately  $2 \times 2 \times 2 \text{ cm}$ . Colors are the same as in figure 8;

c – The Dose-volume histogram for the results of four CDFs with  $40 \times 40 \times 40$  pixels. The vertical axis is the percent of total pixels that receive light doses. Summing the number of points showed that more than 63000 pixels out of 64000 received sufficient light dose

**Рис. 9.** Распределение света, полученное при использовании четырех CDF при  $s = 50 \text{ мВт/см}$ ,  $t = 150 \text{ с}$ ,  $\mu_a = 0,085 \text{ см}^{-1}$ ,  $\mu_s' = 16 \text{ см}^{-1}$ :

a – Положение четырех CDF, размещенных в точках (0,5, 0,5, 2), (0,5, 1,5, 2), (1,5, 1,5, 2), (1,5, 0,5, 2);

b – Распределение света образует объем, который составляет примерно  $2 \times 2 \times 2 \text{ см}$ . Цвета такие же, как на рисунке 8.

c – Гистограмма «Объем-доза» для результатов четырех CDF с  $40 \times 40 \times 40$  пикселей. Вертикальная ось – процент от общего количества пикселей, которые получают световые дозы. Суммирование общего количества пикселей показало, что более 63000 пикселей из 64000 получили достаточную световую дозу

T.H. Foster et al. [7] used eight CDFs to deliver 2270–2350 J ( $333\text{--}1178 \text{ J/cm}^2$ ) to accumulate  $90 \text{ J/cm}^2$  in 90% of tumor volume that was  $5 \text{ cm}^3$  approximately. By raising the delivered power ( $s$ ) or time treatment the percentage of coverage rises as well. In this regard, consumption of oxygen and the kind of photosensitizer must be considered in *in vivo* studies according to the treated organ.

## Conclusion

This study is considered as the first inclusive description of cylindrical diffuser fibers using basic equation of diffusion theory (eq. 1). It was found that optical properties and distance away from CDF had the major effects on light distribution. For light absorbing media (i.e. prostate) which is virtually made of blood, high power is required in order to attain adequate light distribution

in the tissue. On the other hand, for highly scattering media such as breast tissue, increasing the number of CDF fibers with lower power is highly recommended to avoid thermal collateral damage. Moreover, the study provides a guideline for proper localization of optical detectors inside tissue for investigating fluorescence during PDT. In addition, this simulation was very accurate in calculating the light doses at each point because of small size of voxels. Using specific colors for each range of light dose allowed estimating the whole volume covered by the sufficient light dose without referring to the matrixes. This simulation would be flexible regarding changes in variables in equation 1 including time, distance between points and the size of volume structure.

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