Nebulization as a tool for the delivery of photosensitizers in the photodynamic inactivation of respiratory diseases
GIULIA KASSAB

Nebulization as a tool for the delivery of photosensitizers in the photodynamic inactivation of respiratory diseases

Dissertation presented to the Graduate Program in Physics at the Instituto de Física de São Carlos, Universidade de São Paulo to obtain the degree of Master of Science.

Concentration area: Applied Physics
Option: Biomolecular Physics
Advisor: Prof. Dr. Vanderlei Salvador Bagnato

Corrected Version

(Original version available on the Program Unit)

São Carlos
2018
Kassab, Giulia


Dissertation (Master's degree - Graduate Program in Fisica Aplicada Biomolecular) -- Instituto de Fisica de São Carlos, Universidade de São Paulo - Brasil , 2018.

to my mom,

who always saw a scientist in me
ACKNOWLEDGEMENTS

I would like to express my gratitude to those that were part of this journey and were there for me over the past two years. To my advisor, Professor Vanderlei, who trusted me and allowed me to walk on my own two feet. To the funding agencies CAPES and FAPESP process #2017/12937-8, and the International Photodynamic Association (IPA), for the financial support. To Dr. Natalia and Dr. Mariana, who taught me so much, and in a certain way were my advisors, too. To the forty-five mice that were part of this study.

To the scientific contributions of: Professor Moacir, Dr. Eveline and the student Alana, from the DQ-UFSCar; Professor Vádila and her student Ana Elisa, from the DEQ-UFSCar; Professor Alexandra, from UNESP at Araraquara; Professor Cristina, Dr. Didi and Dr. Lilian, from CEPOF.

To all the support from the CEPOF office, the IFSC graduate service, the IFSC library, and the labs LAT and LIEPO. To Professor Ricardo, from the DGE-UFSCar, for providing the equipment. To the support from the colleagues at the Biophotonics lab, especially the ones in microbiology. To CEPOF, and IFSC as a whole.

To the everlasting room number five: Ila, Pan, Clara and Geisi. You were there when I was lost around the labs, and now you are my true friends. To other friendships that grew inside IFSC: Laís, Paula, Shirly and Eva. To Samara, who chose to take her first career steps beside me.

To all my family, in special: my mother Lucy and my father Francisco, who always encouraged me to dream big and work hard, and always pushed me into finding my own answers in a society where so many are taught to obey without question. To my brother Vitor, who I am so proud of. To my aunts Bruna and Angélica, who taught me what it means to be an independent, determined woman. To my Godmother Lydia, that never needed a blood bond to help me pursue my dreams.

To Lis, Mari and Mathias: you are my family in São Carlos and getting here would not have been possible without our talks, shared coffees and late-night sweets. To Luca, who did everything he could to help me. To my friends in São Paulo, especially Gui, Cesar and Rod, who never let the distance drive our friendship apart. Thank you!
“I believe in intuition and inspiration. (...) Imagination is more important than knowledge. For knowledge is limited, whereas imagination embraces the entire world, stimulating progress, giving birth to evolution. It is, strictly speaking, a real factor in scientific research.”

Albert Einstein,
in “Cosmic Religion: With Other Opinions and Aphorisms” (1931)
ABSTRACT


Pneumonia is one of the main causes of death worldwide, specially of the elderly and the children under 5 years old. The traditional antibiotic-based therapy faces a crisis due to the increase in resistance and a lack of new molecules approved. Recently, our research group demonstrated the photodynamic inactivation of streptococcal pneumonia in vivo, a technique to which the development of resistance is described to be unlikely. This study proposed to investigate the applicability of nebulization as a delivery method for photosensitizers, in the hope to advance the research of the photodynamic inactivation of bacterial pneumonia. First, the critical attributes for nebulization (droplet size and delivery rate), the extent of nebulization, and the stability of three photosensitizers were established, and they were all found to be compatible with the technique. Then, the delivery was validated in an animal model using the most promising compound. It was possible to activate it using extracorporeal infrared light without causing acute lung or liver damage. In conclusion, nebulization presented itself as a promising tool for the delivery of photosensitizers to the respiratory tract.

Keywords: Nebulization. Photosensitizers. Pulmonary delivery. Extracorporeal activation.
RESUMO

KASSAB, G. *Nebulização como uma ferramenta para a entrega de fotosensibilizadores na inativação fotodinâmica de doenças respiratórias*. 2018. 93 p. Dissertação (Mestrado em Ciências) - Instituto de Física de São Carlos, Universidade de São Paulo, São Carlos, 2018.

A pneumonia é uma das principais causas de morte no mundo, sobretudo de idosos e crianças menores de cinco anos. A terapia tradicional, baseada em antibióticos, enfrenta uma crise diante do aumento da resistência e do número reduzido de novas moléculas que são aprovadas. Recentemente, este grupo de pesquisa demonstrou a inativação fotodinâmica da pneumonia pneumocócica *in vivo*, uma técnica para a qual o surgimento de resistência é descrito como pouco provável. Este estudo se propôs a investigar a aplicabilidade da nebulização como método de entrega de fotosensibilizadores, na esperança de avançar a pesquisa da inativação fotodinâmica da pneumonia bacteriana. Inicialmente, os atributos críticos da nebulização (tamanho de gotícula e taxa de entrega), a extensão da dose nebulizada, e a estabilidade de três fotosensibilizadores foram estabelecidas. Todos eles se mostraram compatíveis com a técnica. Então, a entrega foi validada em um modelo animal, utilizando o composto mais promissor. Foi possível ativá-lo usando luz infravermelha extracorpórea sem que houvesse dano agudo pulmonar ou hepático. Em conclusão, a nebulização se mostrou uma ferramenta promissora na entrega de fotosensibilizadores ao trato respiratório.

LIST OF FIGURES

Figure 1 - Simplified Jablonski diagram. .................................................................29
Figure 2 - Chemical structure of Indocyanine Green (ICG). .................................30
Figure 3 - Chemical Structure of Photodithazine® (PDZ). ....................................31
Figure 4 - Basic monomeric structure of Photogem® (PTG). .................................31
Figure 5 - The tissue optical window. Hb: hemoglobin; HbO_2: oxygenated hemoglobin. Note how the relative absorbance varies logarithmically.........32
Figure 6 - Normalized absorbance of the three photosensitizers in water. .............33
Figure 7 - Deposition of microdroplets into the respiratory system according to their median diameter. For this model, the author considered the geometric standard distribution to be 2.2..........................................................35
Figure 8 - Hypothetical curve of the cumulative frequency for a measurement of particle size distribution from a nebulizer. The Volume Median Diameter (VMD) corresponds to a cumulative frequency of 50% of the droplets. D10 and D90 are the diameters for which the cumulative frequency is 10% and 90%, respectively..............................38
Figure 9 - Nebulizer chamber in jet nebulization. The air flow (blue arrows) comes from the pump through a tube (in green) and is pushed through the liquid creating droplets of various sizes. Larger droplets are retained and smaller droplets are carried out of the chamber towards the mouthpiece (orange arrow)..................................................................39
Figure 10 - Extent of nebulization for each solution of photosensitizer, being: ICG 50: Indocyanine green 50 μg/mL; ICG 500: Indocyanine green 500 μg/mL; PDZ 50: Photodithazine® 50 μg/mL; PDZ 500: Photodithazine 500 μg/mL; PTG 50: Photogem® 50 μg/mL; PTG 500: Photogem 500 μg/mL. ..........................................................................................42
Figure 11 - Rate of nebulization of the Omron® nebulizer for each of the concentrated solutions (500 μM) of photosensitizers. For PTG, there was no further droplet formation after 5 minutes. ...........................................43
Figure 12 - Concentration of photosensitizer in the vessel as the nebulization occurs. For PTG, there was no further droplet formation after 5 minutes. ....43
Figure 13 - Total wavelength chromatograms of ICG in three different conditions. There is no difference between the chromatograms of nebulized and freshly-prepared ICG. After 3 weeks in water, however, the main peak at 8.5 minutes disappears, and the peak at 6.9 minutes increases..............44
Figure 14 - Total ion chromatograms of ICG in three different conditions. Similarly to what happens in the wavelength chromatograms, here is no
difference between nebulized and freshly-prepared ICG, and after 3 weeks in water the main peak at 8.5 minutes disappears, and the peak at 6.9 minutes increases.

Figure 15 - Overlap of the absorbance spectra obtained from the two interest peaks in the wavelength chromatogram of ICG. The two spectra are quite similar, with the highest absorbance at around 780 nm. RT: retention time.

Figure 16 - Comparison between the mass spectra obtained from the two interest peaks in the wavelength chromatogram of ICG. The peak with the retention time (RT) of 8.5 minutes has a predominant m/z value of 753.3, and the peak at 6.9 min has predominant m/z values of 331.5 and 346.6.

Figure 17 - Custom-made illumination device for the photodynamic treatment of mice. It is composed of 18 laser diodes with maximum emission at 780 nm. The arrows indicate the laser beam propagation.

Figure 18 - Extracorporeal detection of fluorescence with the fiber optics setup. The red exes mark the spots where the measurements were taken.

Figure 19 - Isolation of the lung lobes for analysis. A knot was tied on the location indicated by the black line, so that the bronchoalveolar lavage would reach only the left lobe. The lobes on the right side (superior, middle, inferior and post-caval) were collected for histology.

Figure 20 - Fluorescence signal detected with the experimental setup of a solution of ICG 1.3 mM soon after its preparation, and after complete degradation of its main compound.

Figure 21 - Infrared images of the animals after receiving each of the treatments in experiment 1. The first row shows the left side, the second to the back, and the third the right side. The arrows indicate the spots where the fluorescence of ICG was observed.

Figure 22 - Infrared images of the animals that received each of the treatments from experiment 1, exposing the organs after the euthanasia. Organs in which fluorescence was evident are indicated with the black arrows.

Figure 23 - Relative fluorescence detected externally over time on a single animal that received nebulization of ICG 800 µM in experiment 2. There was no significant difference between the three sites, so the averages are shown.

Figure 24 - Relative fluorescence detected externally over time on animals that received nebulization with the increased concentration of ICG with (PDI) and without (ICG) exposure to infrared light (experiment 4). The horizontal bars indicate the duration of each of the treatments. The difference between treatments is statistically significant at 20 and 40 minutes.
Figure 25 - Relative fluorescence detected externally over time on each of the three positions for animals that received nebulization with the increased concentration of ICG with (PDI) and without (ICG) exposure to infrared light (experiment 4). There is no significant difference between positions for a given time and treatment combination. The difference between treatments is significant at 20 and 40 minutes for all positions. ......61

Figure 26 - Relative fluorescence detected externally at different times after the treatments from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light. There was no significant difference between the three sites, so the averages are shown................62

Figure 27 - Relative fluorescence detected directly at the exposed organs of interest at different times after the treatments from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light............64

Figure 28 - Relative fluorescence detected directly in the bladder, spleen and heart at different times after the treatments from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light........65

Figure 29 - Summary of the relative fluorescence detected directly in the organs over time after each treatment from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light....................66

Figure 30 - Correlation between the fluorescence detected externally on each side, and the fluorescence detected directly in the corresponding lung of the same animal. Data from the control, ICG and PDI groups from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light..................................................67

Figure 31 - Representative histological slides of the lungs from control and treated animals from experiment 3, with increasing magnification. Both groups showed healthy tissue, and there was no difference between them. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x). .................................................................68

Figure 32 - Representative histological slides of the lungs of each group in experiment 4, with increasing magnification. All animals had some level of lung injury, but the sick positive controls (C+) had higher scores than any other group. Note the patches of alveolar thickening in the 10-fold magnification pictures. The 100-fold magnification shows the contrast between the healthy tissue, with ample air pockets, and the dense injured tissue. The vascular congestion is more evident in the 400-fold magnification. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x))............................................................69

Figure 33 - Representative histological slides of the lungs from each group in experiment 5, with increasing magnification. All animals had some level of lung injury, and the groups were indistinguishable from one another. Note the patches of alveolar thickening in the 10-fold
magnification pictures. The degree of vascular congestion varied from animal to animal (see the 400-fold magnification), but there was no difference between the groups. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x).

Figure 34 - Representative histological slides from the livers of the groups in experiment 5, with increasing magnification. There was no difference between the groups. The scale bars correspond to 200 μm (100x) and 50 μm (400x).

Figure 35 - Percentage of macrophages in the bronchoalveolar lavage fluid from the animals at different times after either ICG alone or PDI. There was no significant difference between groups.

Figure 36 - Comparison of selected models of acute lung injury. A and B: normal mouse lungs. The alveolar walls are very thin, and the majority of the alveoli contain no cells (magnification in A, 100x; B, 400x). C and D: lungs from a mouse euthanized 18 h after intratracheal instillation of 5 ng/g lipopolysaccharide. Note the patchy nature of the injury (C, 100x) and the presence of inflammatory infiltrates and vascular congestion (D, 400x). E and F: lungs from a mouse euthanized 21 days after the administration of intratracheal bleomycin. Note the presence of fibrotic areas (arrows) (E, 200x; F, 400x). G and H: lungs from a mouse euthanized 12 h after aerosolization of Escherichia coli, 1 x 10⁸ CFU/mL. Note diffuse thickening of the alveolar spaces and intra-alveolar neutrophilic infiltrates (G, 200x; H, 400x). HE stain.

Figure 37 - Histopathology of mice treated with 5 mg/kg of talaporfin sodium (Laserphyrin®) and 50 J/cm² of light (664 nm laser) including the tumor and adjacent normal liver tissue. The regional boundary line between the laser radiation area and the outside was clear (a). The bottom row of pictures shows: the tumor (b); ischemic necrosis and focally living hepatocytes around Glisson in the peripheral area of the irradiation (c); and the slight denatured hepatocytes around the irradiation area (d). HE stain.
Table 1 - Parameters of the calibration curves to determine the concentration of photosensitizer before and after nebulization. .......................................................... 40

Table 2 - Specific parameters for the HPLC-MS-DAD analysis of each photosensitizer. .................................................................................................................. 41

Table 3 - VMD and span of the droplets formed by the Omron NE-C801 jet nebulizer for each of the tested solutions. ................................................................. 41

Table 4 – Groups of experiment 4. The exes mark the treatments received by each group. .............................................................................................................. 54
### LIST OF ABBREVIATIONS AND ACRONYMS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANOVA</td>
<td>Analysis of Variance</td>
</tr>
<tr>
<td>APC</td>
<td>Allophycocyanin</td>
</tr>
<tr>
<td>A.u.</td>
<td>Absorbance units</td>
</tr>
<tr>
<td>BALF</td>
<td>Bronchoalveolar lavage</td>
</tr>
<tr>
<td>C-</td>
<td>Negative control</td>
</tr>
<tr>
<td>C+</td>
<td>Positive control</td>
</tr>
<tr>
<td>CAP</td>
<td>Community acquired pneumonia</td>
</tr>
<tr>
<td>CD16</td>
<td>Cluster of differentiation 16</td>
</tr>
<tr>
<td>CD32</td>
<td>Cluster of differentiation 32</td>
</tr>
<tr>
<td>CD45</td>
<td>Cluster of differentiation 45</td>
</tr>
<tr>
<td>CFU</td>
<td>Colony-forming units</td>
</tr>
<tr>
<td>cps</td>
<td>Counts per second</td>
</tr>
<tr>
<td>CQAs</td>
<td>Critical quality attributes</td>
</tr>
<tr>
<td>D&lt;sub&gt;10&lt;/sub&gt;</td>
<td>Diameter value for a cumulative frequency of 10%</td>
</tr>
<tr>
<td>D&lt;sub&gt;90&lt;/sub&gt;</td>
<td>Diameter value for a cumulative frequency of 90%</td>
</tr>
<tr>
<td>DAD</td>
<td>Diode-array detection</td>
</tr>
<tr>
<td>FSC</td>
<td>Forward scattering</td>
</tr>
<tr>
<td>HAP</td>
<td>Hospital acquired pneumonia</td>
</tr>
<tr>
<td>Hb</td>
<td>Hemoglobin</td>
</tr>
<tr>
<td>HbO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Oxygenated hemoglobin</td>
</tr>
<tr>
<td>HE</td>
<td>Hematoxylin and eosin</td>
</tr>
<tr>
<td>HPLC</td>
<td>High performance liquid chromatography</td>
</tr>
<tr>
<td>ICG</td>
<td>Indocyanine green</td>
</tr>
<tr>
<td>LED</td>
<td>Light-emitting diode</td>
</tr>
<tr>
<td>MMAD</td>
<td>Mass median aerodynamic diameter</td>
</tr>
<tr>
<td>MRSA</td>
<td>Methicillin-resistant <em>Staphylococcus aureus</em></td>
</tr>
<tr>
<td>MS</td>
<td>Mass spectrometry</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffer solution</td>
</tr>
<tr>
<td>PCVs</td>
<td>Pneumococcal conjugate vaccines</td>
</tr>
<tr>
<td>PDI</td>
<td>Photodynamic inactivation</td>
</tr>
<tr>
<td>PDT</td>
<td>Photodynamic therapy</td>
</tr>
<tr>
<td>PDZ</td>
<td>Photodithazine®</td>
</tr>
<tr>
<td>PPV</td>
<td>Pneumococcal polysaccharide vaccine</td>
</tr>
<tr>
<td>PTG</td>
<td>Photogem®</td>
</tr>
<tr>
<td>RI</td>
<td>Refractive index</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>rpm</td>
<td>Rotations per minute</td>
</tr>
<tr>
<td>RT</td>
<td>Retention time</td>
</tr>
<tr>
<td>SSC</td>
<td>Side scattering</td>
</tr>
<tr>
<td>TIC</td>
<td>Total ion chromatogram</td>
</tr>
<tr>
<td>TWC</td>
<td>Total wavelength chromatogram</td>
</tr>
<tr>
<td>UV-Vis</td>
<td>Ultraviolet and visible light</td>
</tr>
<tr>
<td>VAP</td>
<td>Ventilator-associated pneumonia</td>
</tr>
<tr>
<td>VMD</td>
<td>Volume median diameter</td>
</tr>
</tbody>
</table>
LIST OF SYMBOLS

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{1}\text{O}_2$</td>
<td>Singlet oxygen</td>
</tr>
<tr>
<td>$^{3}\text{O}_2$</td>
<td>Triplet oxygen</td>
</tr>
<tr>
<td>$h\nu$</td>
<td>Light</td>
</tr>
<tr>
<td>$\phi_T$</td>
<td>Triplet state quantum yield</td>
</tr>
<tr>
<td>$\phi_F$</td>
<td>Fluorescence quantum yield</td>
</tr>
<tr>
<td>$\lambda_{max}$</td>
<td>Wavelength of maximum absorbance</td>
</tr>
<tr>
<td>$m/z$</td>
<td>Mass-to-charge ratio</td>
</tr>
<tr>
<td>$+ve$</td>
<td>Positive</td>
</tr>
<tr>
<td>$-ve$</td>
<td>Negative</td>
</tr>
<tr>
<td>$-$</td>
<td>Absent</td>
</tr>
<tr>
<td>$+$</td>
<td>Mildly present</td>
</tr>
<tr>
<td>$++$</td>
<td>Moderately present</td>
</tr>
<tr>
<td>$+++ $</td>
<td>Extensively present</td>
</tr>
</tbody>
</table>
CONTENTS

1 INTRODUCTION ................................................................................................................................. 25
1.1 Objectives ........................................................................................................................................ 26

2 LITERATURE REVIEW .......................................................................................................................... 27
2.1 Current challenges in the treatment of bacterial pneumonia ............................................................... 27
2.2 Photodynamic inactivation of microorganisms .................................................................................... 28
2.2.1 Mechanism .................................................................................................................................... 29
2.2.2 Photosensitizers ............................................................................................................................. 30
2.2.3 Light sources and light penetration ............................................................................................... 32
2.2.4 Photodynamic Inactivation of S. pneumoniae using indocyanine green and infrared light .......... 33

2.3 Respiratory delivery via nebulization ................................................................................................. 33
2.3.1 Types of Nebulizers ...................................................................................................................... 34
2.3.2 Characteristics of an ideal drug for nebulization .......................................................................... 34

3 EFFICACY OF THE NEBULIZATION OF DIFFERENT PHOTOSENSITIZERS .... 37
3.1 Purpose ............................................................................................................................................. 37
3.2 Methodology ................................................................................................................................... 37
3.2.1 Materials ...................................................................................................................................... 37
3.2.2 Determination of the volume median diameter (VMD) ................................................................. 38
3.2.3 Rate and extent of nebulization ................................................................................................... 39
3.2.4 Verification of Stability ................................................................................................................. 40
3.2.5 Statistical Analysis ..................................................................................................................... 41
3.3 Results ............................................................................................................................................... 41
3.3.1 Volume Median Diameter (VMD) ................................................................................................ 41
3.3.2 Rate and Extent of Nebulization .................................................................................................. 42
3.3.3 Stability of the Photosensitizers during Nebulization ................................................................. 44
3.4 Discussion ......................................................................................................................................... 47
3.5 Conclusion ......................................................................................................................................... 49

4 NEBULIZATION AND EXTRACORPOREAL ACTIVATION OF INDOCYANINE GREEN IN VIVO ......................................................................................................................... 51
4.1 Purpose ............................................................................................................................................ 51
4.2 Methodology ................................................................................................................................... 51
4.2.1 Animal Model ............................................................................................................................... 51
4.2.2 Materials and Devices .................................................................................................................. 52
4.2.3 Experimental Procedures and Data Collection ............................................................. 53
4.2.3.1 Experiment 1: Infrared imaging detection of nebulized vs instilled ICG .................. 53
4.2.3.2 Experiment 2: Extracorporeal monitoring of photosensitizer delivery over time....... 53
4.2.3.3 Experiment 3: Tissue damage and immune response to the treatment ................. 54
4.2.3.4 Experiment 4: Tissue damage and treatment monitoring with an increased dose of ICG ................................................................. 54
4.2.3.5 Experiment 5: Body distribution of lung-delivered ICG with and without exposure to light and time-dependency of the tissue and immune response ........................................... 55
4.2.4 Fluorescence Data Processing ..................................................................................... 56
4.2.5 Flow Cytometry ........................................................................................................ 56
4.2.6 Histological Analysis ................................................................................................. 57
4.3 Results .............................................................................................................................. 58
4.3.1 Selective Fluorescence of ICG with excitation at 780 nm ........................................... 58
4.3.2 Delivery of ICG via Nebulization versus Instillation ................................................. 58
4.3.3 Extracorporeal Detection and Activation of the Nebulized ICG in the Lungs ........... 60
4.3.4 Body Distribution of the Photosensitizer ................................................................ 62
4.3.5 Tissue Damage and Immune Response ..................................................................... 68
4.4 Discussion ......................................................................................................................... 72
4.5 Conclusion ...................................................................................................................... 80
5 CONCLUSIONS .................................................................................................................. 81
REFERENCES ....................................................................................................................... 83
APPENDIX A – Chromatograms of the photosensitizer solutions before and after nebulization ......................................................................................................................... 91
ANNEX A – Certificate of Approval from the Ethics Committee ........................................ 93
1 INTRODUCTION

Pneumonia is one of the main causes of death worldwide, specially of the elderly and the children under 5 years of age.\textsuperscript{1-2} Since it is mainly caused by bacteria, the treatment usually includes antibiotics.\textsuperscript{2} However, the antibiotic-based therapy faces a crisis due to the increase in resistance and a lack of new molecules.\textsuperscript{3} A promising alternative to this approach is photodynamic inactivation (PDI), the microbiological application of photodynamic therapy (PDT), to which the development of resistance is described to be unlikely.\textsuperscript{4} PDI is based on the combination of a photosensitizer, light at a certain wavelength, and molecular oxygen, to generate reactive oxygen species (ROS) that cause cell death.\textsuperscript{4}

Since 2013, our research group studies the applicability of PDI in the treatment of streptococcal pneumonia, using the photosensitizer indocyanine green (ICG) and extracorporeal activation with infrared light. The initial in vitro studies showed that it was possible to eliminate S. pneumoniae using doses of ICG and light that were unharmful to macrophages.\textsuperscript{5} Moreover, there was evidence that the PDI could potentialize the antimicrobial activity of such macrophages during the infection.\textsuperscript{5}

In the \textit{in vivo} studies, a single session of PDI was enough to reduce the bacterial burden and increase the survival rate of previously infected mice.\textsuperscript{6} However, ICG was delivered to the mice using instillation, which has the benefits of being local, but is uncomfortable and not well accepted in the clinical practice.\textsuperscript{6} Within our program of lung decontamination, efficient photosensitizer delivery to the airways is a key point, so this study proposed to investigate the applicability of nebulization as a method of delivering photosensitizers to the respiratory tract for the PDI of respiratory pathogens.

Nebulization is widely used in the medical practice, since it allows the drug to reach the target site directly, reducing the required dose and limiting the side effects.\textsuperscript{7} Nebulized antibiotics have been part of a successful regimen for the management of cystic fibrosis, and have shown promising results in the treatment of ventilator-associated pneumonia.\textsuperscript{8} Plus, the equipment is available at hospitals, and already part of the usual management of pneumonia in children.\textsuperscript{9} However, the only publication that we were able to find so far that investigates nebulization for the photodynamic inactivation of pathogens is a pilot study from 2011, in which the properties of a nebulized methylene blue solution were assessed in an \textit{ex vivo} model.\textsuperscript{10}

This motivated the study of the compatibility between nebulization and other photosensitizers, in particular those that had potential to be activated externally. Three
photosensitizers were chosen based on their absorbance spectra and activity against microorganisms: ICG, the chlorin derivative Photodithazine® (PDZ), and the porphyrin derivative Photogem® (PTG).

1.1 Objectives

The primary objective of this study was to determine the efficacy of nebulization as a delivery method for photosensitizers. For that, it proposed to:

- Assess the critical attributes for nebulization (droplet size and delivery rate), the extent of nebulization, and the stability of the photosensitizers ICG, PDZ and PTG during the process;
- Validate the nebulization delivery of ICG to the lungs, measure its body distribution, and evaluate the toxicity and the immune response to a simulated PDI treatment in a murine model.
2 LITERATURE REVIEW

2.1 Current challenges in the treatment of bacterial pneumonia

Lower respiratory infections are the fourth largest cause of death worldwide, and the main cause of death in low-income countries. The most frequent lower respiratory infections are acute bronchitis and bronchiolitis, influenza, and pneumonia. Pneumonia is the main infectious cause of death in Europe and the United States. It is also the worldwide main cause of death of children younger than 5 years old. In Brazil, pneumonia is the number one cause of hospitalization. Although the number of hospitalizations has decreased over the past decades, the in-hospital mortality increased, what is mainly explained by the aging of the population and the occurrence of cases of pneumonia that are more difficult to treat.

The European Respiratory Society defines pneumonia as an acute illness of the lower respiratory tract that includes cough and at least one other symptom: new focal chest signs, new lung shadowing shown by radiography, otherwise unexplained fever for more than 4 days, or otherwise unexplained tachypnea/dyspnea. Community Acquired Pneumonia (CAP) is contracted from contact with the infection in day-to-day life. It is predominantly bacterial in origin, being Streptococcus pneumoniae its most prevalent pathogen. Other important agents are Haemophilus influenza, Pseudomonas aeruginosa, Chlamyphila pneumoniae, Mycoplasma pneumoniae, Legionella pneumophila, and Coxiella burnetii. About 30% of cases are coinfections with viruses. However, in the vast majority of CAP cases, there is no investigation of the etiological agent. In such situations, the treatment is based on the most prevalent microorganisms of that locality. The main classes of antibiotics employed for CAP in Brazil are macrolides, beta-lactams and quinolones.

Hospital Acquired Pneumonia (HAP), also called nosocomial pneumonia, is the one that develops after at least 48 hours after the patients admission. Its reported mortality rate ranges from 20 to 50%, the highest amongst nosocomial infections. Ventilator-associated pneumonia (VAP) is the one contracted at least 48-72 hours after endotracheal intubation. The most relevant agents of HAP and VAP are also bacteria, like Staphylococcus aureus, Pseudomonas aeruginosa, Escherichia coli, and Klebsiella, Acinetobacter, and Enterobacter species. Knowledge of the etiological agents is of great importance in the treatment of these infections, since patients that receive the wrong initial therapy have a high risk of mortality and morbidity. However, the delay in starting the treatment also leads to a poor prognostic.
A great concern in HAP and VAP cases is the present of methicillin-resistant *Staphylococcus aureus* (MRSA), which is associated with elevated mortality rates and treatment costs.\textsuperscript{19} Traditionally, the first-choice drug for MRSA infections is vancomycin, that due to its low penetration in the lungs and high renal toxicity, leads to a failure rate the can reach 70\%.

Even with new drugs like linezolid, tigecycline and ceftaroline, persists the difficulty in increasing the success rate of treatments, and the worry with the development of resistance.\textsuperscript{19-20} Linezolid, for example, was approved for clinical use in 2000, and cases of resistance in patients were reported as early as 2002.\textsuperscript{21} In a study from 2014, the occurrence of non-susceptibility to this antibiotic remained relatively low, but a number of different resistance mechanisms had already been observed by then.\textsuperscript{21}

Another approach to hinder the burden of pneumonia is vaccination. Two types of vaccines are currently available for *S. pneumoniae*, the main agent in CAP: the pneumococcal polysaccharide vaccine (PPV) has been recommended for adults since the mid-1980’s, but it lacks efficacy in neonates and infants;\textsuperscript{22} the pneumococcal conjugate vaccines (PCVs), designed to overcome that, were first approved in 2000.\textsuperscript{23} However, pneumococcal vaccination faces two main challenges: first, each vaccine is only effective against the serotypes contained in it; second, the reduction of the said serotypes increases the colonization of other serotypes that are not covered by the vaccines, and of other pathogen species like *S. aureus* and *H. influenzae*.\textsuperscript{22} Thus, new vaccines need to be developed continuously, similarly to what happens to antibiotics.\textsuperscript{22}

### 2.2 Photodynamic inactivation of microorganisms

The photodynamic inactivation (PDI) of microorganisms was first described in 1900 by Oscar Raab.\textsuperscript{24} It is based in the use of a photosensitizer that accumulates preferably in the pathogens, and that is activated by light at a specific wavelength, and in the presence of molecular oxygen generates reactive species that are toxic to the target.\textsuperscript{3} PDI is particularly interesting for fighting infections because there is no evidence of cross-resistance with antibiotics, and the restriction of the light exposure offers a further degree of selectivity.\textsuperscript{24-25} More than that, since PDI can damage a variety of molecules, unlike the target-specific
antibiotics, the development of resistance of previously-susceptible strains to this treatment is described as highly unlikely.\textsuperscript{4}

### 2.2.1 Mechanism

The photodynamic process starts with an electron transfer to a higher energy orbital after the absorption of the light.\textsuperscript{27} The excited molecule may dissipate this energy emitting fluorescence, or go through an intersystem crossing and go to a triplet state.\textsuperscript{27} From there, the energy can be dissipated non-radioactively, or generate toxicity through either of two mechanisms: in the type I reaction, the photosensitizer reacts with adjacent organic molecules and forms reactive oxygen species (ROS); in the type II reaction, there is an energy transfer to the molecular oxygen, which is a triplet in its basal state ($^3\text{O}_2$), and it goes to a singlet state ($^1\text{O}_2$).\textsuperscript{27} Both reactions contribute to the phototoxic effect, and both singlet oxygen and the ROS have short half-life times, limiting their diffusion and therefore the effect of the PDI to the site of light exposure.\textsuperscript{28,29} The Jablonski diagram illustrates this mechanism (fig.1.).

![Jablonski Diagram](image)

(1) Absorption; (2) Fluorescence; (3) Intersystem crossing; (4) Type II reaction (energy transfer); (5) Type I reaction.

Figure 1 - Simplified Jablonski diagram.
Source: By the author.
2.2.2 Photosensitizers

Figure 2 - Chemical structure of Indocyanine Green (ICG).
Source: By the author.

Indocyanine Green (fig. 2) is a water-soluble dye that emits fluorescence when exposed to infrared light. Its absorption peak in human plasma is 805 nm. The fluorescence quantum yield of ICG is \( \phi_F = 0.027 \pm 0.005 \), and its triplet state quantum yield is \( \phi_T > 0.01 \). ICG is used in the diagnosis of cancers, the determination of the cardiac debt and liver function, and in ophthalmic angiography. It is not ideal for the photodynamic therapy of cancers because of its short half-life time in the blood. However, studies employing ICG for the elimination of microorganisms through PDI have had promising results in vitro and in vivo.

Photodithazine® (fig. 3) is the N-methyl glucosamine salt of chlorin e₆ and therefore is a chlorophyll derivative. The glucosamine moiety provides better water solubility to the chlorin e₆, and it is eliminated in acid pH. PDZ absorbs greatly in the red region, with \( \lambda_{\text{max}} = 662 \text{ nm} \), and penetrates well into biological tissue. It also has a greater photodynamic efficiency than well-established photosensitizers like Photofrin®. Clinical trials of this drug for the treatment of tumors showed low dark toxicity, high selectivity and phototoxicity, and fast clearance. PDZ was shown to be effective against a multispecies biofilm containing Candida albicans, Candida glabrata and Streptococcus mutans in vitro after irradiation with LED light at 660 nm.
Photogem® (fig. 4) is a hematoporphyrin derivative, composed of monomers and oligomers with very similar structure and distribution to Photofrin II. Hematoporphyrin derivatives are the most frequently used photosensitizers in the treatment of tumors, and PTG is of particular interest because it has been approved for clinical use in Brazil. Its absorbance of light is most intense at 369 nm (Soret band), but it also has smaller peaks at 507, 540, 570 and 620 nm (Q bands). PTG was shown to be effective against Candida species in vitro and in vivo.
2.2.3 Light sources and light penetration

For PDI to be effective, light needs to be able to reach the pathogens in sufficient dose, despite the dispersion and absorption of the adjacent layers of tissue. Therefore, the choice of wavelength should take into account the depth of light penetration, the absorbance spectrum of the photosensitizer, and the location of the infectious agents.

![Image of tissue optical window](image)

Figure 5 - The tissue optical window. Hb: hemoglobin; HbO\(_2\): oxygenated hemoglobin. Note how the relative absorbance varies logarithmically.
Source: HUANG et al.\(^{49}\)

It is desirable to have the light excitation at around 650 to 1000 nm, because light at this range penetrates deeper into biological tissue, since it is less absorbed by water, melanin and hemoglobin (fig. 5).\(^{49}\) Because of that, this range is known as the “optical window” or the “therapeutic window” for photodynamic therapy.\(^{49-50}\) However, most photosensitizer do not absorb wavelengths greater than 800 nm, so the usual excitation sources have peaks that range from 650-850 nm.\(^{51-52}\)

Figure 6 shows the absorbance spectra of the three photosensitizers used in this study when dissolved in water. All of them have some absorbance in the optical window, although only ICG has its peak absorbance within this range.
2.2.4 Photodynamic Inactivation of S. pneumoniae using indocyanine green and infrared light

The application of ICG and extracorporeal infrared light to the treatment of bacterial pneumonia has been studied by our research group since 2013. In a study published in the Journal of Biophotonics in 2017, the in vitro inactivation of S. pneumoniae was found to be effective using concentrations of ICG as low as 5 μM when combined with a 780 nm laser device or 10 μM when using an 850 nm LED. In these conditions, the treatment was safe for RAW 264.7 macrophages, and seemed to enhance their ability to fight the bacteria.\(^5\)

The in vivo investigation found a reduction in the bacterial burden and an increase in the survival rate of SKH-1 hairless mice infected with S. pneumoniae after a single PDI session using ICG 100 μM and 120 J/cm\(^2\) of light at 780 nm.\(^6\) In this study, the light exposure did not seem to be harmful to the animals. Additionally, the ICG alone was no different form the control, suggesting that the activation with light was essential to the observed effects. Therefore, both the in vitro and in vivo findings suggested that the combination of ICG and infrared light has a great potential to treat pneumococcal pneumonia.

2.3 Respiratory delivery via nebulization

Although vapors and aerosols have been employed in the treatment of respiratory diseases for many centuries, the term “nebulizer” only appeared in the end of the 19th century,
when the first mechanical pumps for air flow generation were created.\(^\text{53}\) The technology kept developing throughout the decades, and modern nebulizers are capable of delivering more than 60% of the nominal dose to the respiratory system.\(^\text{54-55}\) Nowadays, nebulization is used in the medical practice to deliver bronchodilators, antibiotics, mucolytic drugs and local anesthetics.\(^\text{7}\) It allows the drug to reach the target site directly, reducing the required dose and limiting the side effects.\(^\text{7}\)

### 2.3.1 Types of Nebulizers

Three types of equipment are currently available in the clinical practice: jet, ultrasonic, and vibrating-mesh nebulizers.\(^\text{55}\) Jet nebulizers are the most commonly used type.\(^\text{56}\) They contain Venturi orifices and deflectors, that combined with a high air flux, create a negative pressure on the liquid surface.\(^\text{53,55}\) Then, because of the superficial tension, microdroplets are created.\(^\text{55}\) Ultrasonic nebulizers are based on a piezoelectric transductor that vibrates at a high frequency in the base of the recipient, creating the microdroplets of liquid.\(^\text{55}\) In the case of vibrating-mesh nebulization, the liquid is pushed through micrometric opening, creating the aerosol.\(^\text{55}\)

The choice between nebulization method depends on the stability of the active compounds and the formulations to each of these processes.\(^\text{55}\) Although ultrasonic nebulizers are less noisy than the jet ones, they are more expensive and there is an increase in temperature inside the nebulization chamber that makes them incompatible with thermolabile formulations.\(^\text{57}\) Ideally, prescriptions for nebulization should include both the drug and the equipment.\(^\text{58}\)

### 2.3.2 Characteristics of an ideal drug for nebulization

The deposition of the droplets in the respiratory tract is dependent on their diameter, so this must be taken into account in the development of a nebulization method.\(^\text{8,53}\) Lung deposition seems to be optimal when droplet diameter is between 2-5 µm, whereas larger particles will have a larger deposition in the oropharynx (fig.7).\(^\text{59}\)
Figure 7 - Deposition of microdroplets into the respiratory system according to their median diameter. For this model, the author considered the geometric standard distribution to be 2.2.

Source: CLARK.59

The droplet size distribution and the drug output (or delivery rate) are considered to be the critical quality attributes (CQAs) of nebulized drugs.56 For a given nebulizer, the CQAs vary according to aspects of the solution or suspension, mainly viscosity, surface tension, and concentration.56 Changes in the equipment, for example increasing its air pressure, also have an influence in the CQAs.60

Other important aspect to be assessed in the development of drugs for nebulization is the stability of the active compounds, that may not be compatible with some of the processes of droplet formation.60 Additionally, the residual volume (and therefore the extent of the dose released) changes according to the nebulization method, the equipment, and the drug formulation.63-64

In the context of fighting infections, aerolized antibiotics have been part of a successful regimen for the management of cystic fibrosis, and have shown promising results in the treatment of non-cystic fibrosis bronchiectasis and ventilator-associated pneumonia.8 However, a single publication was found when searching for nebulization and photodynamic inactivation of pathogens: a pilot study from 2011, in which the properties of a nebulized
methylene blue solution were assessed.\textsuperscript{10} The first photodynamic treatment of a respiratory infection was focused on papillomatosis and went through clinical trials on the late 1980s.\textsuperscript{63} Surprisingly, although the larynx was the target tissue in this and other similar studies, the photosensitizers were administered intravenously.\textsuperscript{63-64}
3 EFFICACY OF THE NEBULIZATION OF DIFFERENT PHOTOSENSITIZERS

3.1 Purpose

Recently, our research group demonstrated the photodynamic inactivation of streptococcal pneumonia in vivo, using instillation to deliver the drug, which is uncomfortable and not well accepted in the clinical practice. Within our program of lung decontamination, efficient photosensitizer delivery to the airways is a key point. Since nebulization has clear advantages when compared to instillation and intravenous delivery, this study proposed to investigate the applicability of the jet nebulization as a method of delivering photosensitizers to the respiratory tract for the PDI of respiratory pathogens.

Three photosensitizers were chosen based on their activity against microorganisms: indocyanine green is a fluorescent dye used in diagnostics, and it is active against S. pneumoniae both in vitro and in vivo, Photodithazine® is a chlorin e₆ derivative that successfully reduced the viability of a multispecies biofilm, and Photogem® is a hematoporphyrin derivative with activity against Candida species. The critical attributes (droplet size and delivery rate), the extent of nebulization, and the stability of the photosensitizers to the process were assessed to ensure that they were compatible with the jet nebulization.

3.2 Methodology

3.2.1 Materials

All solutions of ICG (Indocyanine Green, Ophtalmos, Brazil), PDZ (Photodithazine®, Fotoditazin, Russia) and PTG (Photogem®, Photogem, Russia) were freshly prepared in distilled water at the concentrations of each experiment. The stock solution of Photodithazine® has 5 mg/mL, while the Indocyanine Green and the Photogem® are provided in powder. Exceptionally for the in vivo experiments, the chosen solvent was sterile water for injection, to avoid contamination. The commercially available Omron NE-C801 jet nebulizer (Omron, Kyoto, Japan) was used for all experiments, with different nebulizing chambers for each photosensitizer, and for the controls phosphate buffered saline (PBS) and distilled water.
3.2.2 Determination of the volume median diameter (VMD)

Figure 8 - Hypothetical curve of the cumulative frequency for a measurement of particle size distribution from a nebulizer. The Volume Median Diameter (VMD) corresponds to a cumulative frequency of 50% of the droplets. D10 and D90 are the diameters for which the cumulative frequency is 10% and 90%, respectively. 

Source: By the author.

The size of the particles released by a nebulizer usually follows a normal distribution. The volume median diameter (VMD) represents the diameter correspondent to a cumulative frequency of 50% of the particles, or in other words, half of the droplets are smaller or equal to the VMD, and half of them are larger. The measure of spread for the distribution of diameters is the Span, and it is calculated by the RT Sizer software as $\text{Span} = (D_{90} - D_{10})/\text{VMD}$, where $D_{90}$ and $D_{10}$ are the diameters for which 90% and 10% of the particles are smaller or equal to, respectively. Figure 8 shows a cumulative frequency curve for a hypothetical measure of particle size distribution, in which the VMD, $D_{90}$ and $D_{10}$ are indicated.

The droplet size was determined using an optical measurement, with the Spraytec system (Malvern Instruments model RTS5134) and the software RT Sizer. In this equipment, a 3mW Helio-Neon laser generator produces laser light at a wavelength of 632.8 nm. The laser beam is expanded by a lens and passes through the spray where a part of the light energy is diffracted at different angles depending on the size distribution of the droplets in the spray. The diffraction angle is inversely proportional to the size of the droplet. The lens used in the experimental tests was a 100 mm focal length lens that provided a particle size range, based on volume median diameter of the particle size distribution, from 2.5 to 125 µm. It is worth
mentioning that actual range of the instrument is wider than listed to accurately measure droplets both above and below the volume median diameter.

The nebulizer was placed in the middle of the optical path, and around 100 measurements were collected for each sample. For the solutions of ICG, PDZ and PTG at 500 μg/mL, the “opaque” configuration was selected, with a refraction index (RI) of 1.5. For water and PBS, the “water” configuration was chosen, with RI = 1.33. Water and PBS were chosen as controls to help explain whether the VMD results found were intrinsic to the equipment, or variable according to the characteristics of each solution.

3.2.3 Rate and extent of nebulization

![Nebulizer chamber in jet nebulization](image)

Figure 9 - Nebulizer chamber in jet nebulization. The air flow (blue arrows) comes from the pump through a tube (in green) and is pushed through the liquid creating droplets of various sizes. Larger droplets are retained and smaller droplets are carried out of the chamber towards the mouthpiece (orange arrow).

Source: By the author.

In jet nebulization, the solution or suspension of drug in placed in a nebulizer chamber, like the one pictured in figure 9. The design of the chamber helps the retention of larger droplets, that drain back to the main liquid, while the smaller droplets are carried out of
the equipment (orange arrow). For that reason, an indirect measurement was used to quantify the nebulization, by measuring the change in mass of the nebulizer chamber and the concentration of the remaining solutions.

The photosensitizers were prepared in two concentrations: the same used for the VMD determination (500 μg/mL), and a lower one (50 μg/mL) for further comparison. The chamber was weighted before and after the addition of 2.0 mL of each solution. For the concentration of 50 μg/mL, the nebulizer was on for 6 minutes, and afterwards the chamber was weighted again, and the concentration of the remaining liquid determined. For 500 μg/mL, the nebulization was interrupted after each minute passed, the chamber was weighted, a small aliquot (30 µL) was removed and diluted to determine concentration, and the chamber was weighted once more. All measurements were performed in triplicate.

A calibration curve was prepared for each photosensitizer using UV-vis spectrometry (Varian Cary® 50, Varian Medical Systems, United States), based on their peak absorbance in water, as shown in table 1. The reason why the concentration range in the curves is so smaller than the ones used in the nebulizer is to respect the detection limits of the equipment and ensure linearity. Also, diluting the sample before measurement was a way to reduce the amount of solution that had to be removed for each measurement.

Table 1 - Parameters of the calibration curves to determine the concentration of photosensitizer before and after nebulization.

<table>
<thead>
<tr>
<th>Photosensitizer</th>
<th>ICG</th>
<th>PDZ</th>
<th>PTG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak Absorbance (nm)</td>
<td>780</td>
<td>405</td>
<td>368</td>
</tr>
<tr>
<td>Concentration Range (μg/mL)</td>
<td>4.0 – 7.0</td>
<td>3.0 – 7.0</td>
<td>4.0 – 8.0</td>
</tr>
<tr>
<td>Linearity (R²)</td>
<td>0.9944</td>
<td>0.9977</td>
<td>0.9899</td>
</tr>
</tbody>
</table>

Source: By the author.

### 3.2.4 Verification of Stability

The composition of the solutions prior to and after nebulization was determined by high performance liquid chromatography (HPLC) coupled with mass spectrometry (MS) and diode-array detection (DAD) of absorbance spectra. The nebulizer was placed inside of a clean plastic chamber and the nebulization was performed for 6 minutes with 2.0 mL of each photosensitizer at 1.0 mg/mL. The droplets were allowed to set before the chamber was open. Still, it was only possible to collect a few microliters of each “after” sample, and that is the reason why the initial concentration was higher than in other experiments.

The aliquots from before and after the procedure were diluted in the same mixture as
the initial mobile phase to a final concentration of 100 μg/mL prior to analysis. For ICG, a solution stocked for three weeks at 4 °C was also analyzed. A different method of analysis was developed for each photosensitizer according to published literature, in the hope to separate potential degradation products.\textsuperscript{66-68} The Agilent 1200 HPLC system (Agilent Technologies, United States) and a reverse-phase column (NST 18 80A, 150 mm x 4,6 mm x 5 μm) were used. After separation, the mobile phase was split between the API 200 triple quadrupole mass spectrometer and the Agilent 1200 diode array for detection. Analysis was performed with the software Analyst 1.5.1. The particularities of each method are summarized in Table 2.

Table 2 - Specific parameters for the HPLC-MS-DAD analysis of each photosensitizer.

<table>
<thead>
<tr>
<th>Method</th>
<th>ICG</th>
<th>PDZ</th>
<th>PTG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aqueous Phase (A)</td>
<td>0.1 M Ammonium Acetate in water (pH 5.3)</td>
<td>0.1% formic acid</td>
<td>0.1% formic acid</td>
</tr>
<tr>
<td>Organic Phase (B)</td>
<td>Acetonitrile</td>
<td>Acetonitrile</td>
<td>Acetonitrile</td>
</tr>
<tr>
<td>Gradient</td>
<td>30% to 90% B in 12 minutes</td>
<td>45% to 100% B in 20 minutes</td>
<td>35% to 100% B in 20 minutes</td>
</tr>
<tr>
<td>Total Run Time</td>
<td>25 minutes</td>
<td>40 minutes</td>
<td>36 minutes</td>
</tr>
<tr>
<td>Ion Detection</td>
<td>m/z = 50 – 900 (+ve)</td>
<td>m/z = 50 – 900 (+ve)</td>
<td>m/z = 50 – 1300 (-ve)</td>
</tr>
<tr>
<td>Absorbance Detection</td>
<td>200 – 900 nm</td>
<td>200 – 900 nm</td>
<td>200 – 900 nm</td>
</tr>
<tr>
<td>Injection Volume</td>
<td>20 μL</td>
<td>20 μL</td>
<td>20 μL</td>
</tr>
</tbody>
</table>

Source: By the author.

3.2.5 Statistical Analysis

For all experiments, measurements were taken in triplicate. The analysis of variance (one-way or two-way ANOVA) was the test of choice whenever applicable, with a post-hoc Tukey test. The groups were considered different with a level of significance of 0.05. All calculations were performed using the software Origin 9.0 (OriginLab, Northampton, USA).

3.3 Results

3.3.1 Volume Median Diameter (VMD)

The VMD and span values for the photosensitizer solutions, water and PBS are summarized in Error! Autoreferência de indicador não válida.. The obtained VMD values are
close to 9 µm and do not seem to vary greatly depending on the composition of the solution. The obtained span values are all smaller than 0.3, indicating a small dispersion of particle diameters.

Table 3 - VMD and span of the droplets formed by the Omron NE-C801 jet nebulizer for each of the tested solutions.

<table>
<thead>
<tr>
<th>Solution</th>
<th>VMD (µm)</th>
<th>Span</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distilled Water</td>
<td>8.90</td>
<td>0.28</td>
</tr>
<tr>
<td>PBS</td>
<td>9.60</td>
<td>0.17</td>
</tr>
<tr>
<td>ICG 500</td>
<td>9.34</td>
<td>0.24</td>
</tr>
<tr>
<td>PDZ 500</td>
<td>9.00</td>
<td>0.28</td>
</tr>
<tr>
<td>PTG 500</td>
<td>9.19</td>
<td>0.25</td>
</tr>
</tbody>
</table>

Source: By the author.

3.3.2 Rate and Extent of Nebulization

Figure 10 shows the percentage of photosensitizers released six minutes after nebulization. In all cases, at least 65% of the dose was carried out of the equipment in the form of droplets. The 2-way ANOVA showed that the dose released was significantly larger for the higher concentrations of photosensitizers, and that less PDZ was nebulized compared to ICG and PTG.

Figure 10 - Extent of nebulization for each solution of photosensitizer, being: ICG 50: Indocyanine green 50 µg/mL; ICG 500: Indocyanine green 500 µg/mL; PDZ 50: Photodithazine® 50 µg/mL; PDZ 500: Photodithazine 500 µg/mL; PTG 50: Photogem® 50 µg/mL; PTG 500: Photogem 500 µg/mL.
Figure 11 - Rate of nebulization of the Omron® nebulizer for each of the concentrated solutions (500 \( \mu \text{M} \)) of photosensitizers. For PTG, there was no further droplet formation after 5 minutes.
Source: By the author.

Figure 12 - Concentration of photosensitizer in the vessel as the nebulization occurs. For PTG, there was no further droplet formation after 5 minutes.
Source: By the author.

It was possible to monitor the release of photosensitizer after each minute of
nebulization at 500 µg/ml (fig. 11). PTG reached a limit five minutes after and no nebulization was observed after that time, whereas ICG and PDZ continued to generate mist until the sixth minute. For all three solutions, the rate of delivery was high and stable initially, and its decline seemed to coincide with the limiting volume of solution in the vessel. The concentration of photosensitizer in the remaining solution was monitored as well (fig. 12). Although concentrations did not change drastically, they were all significantly higher at the end of the process, indicating the increase in concentration during the process.

3.3.3 Stability of the Photosensitizers during Nebulization

Figures 13 and 14 show the total ion chromatograms (TIC) and the total wavelength chromatograms (TWC) for the solution of ICG before and after nebulization, and after degradation in water. Figures 15 and 16 show the absorbance and mass spectra for the most important peaks found in the described chromatograms. ICG (m/z = 753.3) eluted with a retention time of 8.5 minutes, and there was an unidentified impurity present in both the fresh and the nebulized solutions (m/z = 331.5 and 346.6; RT= 6.9 min).

![Total wavelength chromatograms of ICG in three different conditions.](image-url)

**Figure 13** - Total wavelength chromatograms of ICG in three different conditions. There is no difference between the chromatograms of nebulized and freshly-prepared ICG. After 3 weeks in water, however, the main peak at 8.5 minutes disappears, and the peak at 6.9 minutes increases.
Comparing the peaks and their corresponding absorbance and mass spectra, the composition of the solution did not seem to change at all after the nebulization. The chromatogram of a solution that spent three weeks at 4 °C is shown to demonstrate the type of difference that would be observed in the case of a degraded sample. In this case, the peak correspondent to ICG is much smaller in the TIC than in the fresh solution (fig. 14), and entirely gone in the TWC (fig. 13). On the other hand, the peak at RT = 6.9 minutes has increased intensity in both chromatograms. It has the same mass and absorbance as the impurity observed at this retention time for the fresh ICG.
Figure 15 - Overlap of the absorbance spectra obtained from the two interest peaks in the wavelength chromatogram of ICG. The two spectra are quite similar, with the highest absorbance at around 780 nm. RT: retention time.

Source: By the author.

Figure 16 - Comparison between the mass spectra obtained from the two interest peaks in the wavelength chromatogram of ICG. The peak with the retention time (RT) of 8.5 minutes has a predominant m/z value of 753.3, and the peak at 6.9 min has predominant m/z values of 331.5 and 346.6.

Source: By the author.
No evidence of degradation was found for neither of the other photosensitizers (see Appendix A). PDZ eluted at 11.5 minutes (m/z = 597) and was the only compound detected in the TWC. Photogem® is a mixture of oligomers and it was not possible to correlate the observed peaks with specific molecular entities, but still the three main peaks in the TWC (at 6.1, 11.9 and 12.8 min) are present before and after the process of nebulization. The absorbance and mass spectra of all peaks remained unchanged.

3.4 Discussion

Measuring the size and distribution of droplet diameters helps to understand the dynamics and predict the lung deposition profile of nebulized drugs. The VMD is the dimension obtained by laser diffraction techniques, and was created in the attempt to be a more reliable measurement than the traditional mass median aerodynamic diameter (MMAD), obtained from other methods. In 1990, Rudolf and co-workers developed a semi-empirical model to describe deposition of aerosols in the human respiratory system. This model was further studied by Clark in 1995, that was able to demonstrate a strong correlation between the model and in vivo findings of other published results. According to this model, nebulizer clouds with VMD results of around 9 µm, like the ones in table 3, would have an average deposition fraction of over 80% of the inhaled dose, half of which would be deposited in the lungs. Although model suggests that an even greater deposition into the peripheral airways would be achieved with droplets of 2-5 µm, this would be very hard to achieve with a cheap and ubiquitous jet nebulizer. The high concentration of the photosensitizer solutions does not seem to affect greatly the size of the droplets, given that the obtained VMD results are between the ones for distilled water and PBS. Therefore, the droplets generated by the jet nebulizer are adequate for the delivery of photosensitizers to the lungs.

As for the delivered concentration, figure 10 shows that over 65% of the dose is nebulized for all three photosensitizers at 50 and 500 µg/mL. Total mass output in jet nebulization is dependent on equipment model and drug formulation, and it is incredibly variable, with efficiencies reported in the literature that range from 11% to 97%. However, it tends to be less dependent on the properties of the solution than for vibrating mesh or ultrasonic nebulizers. At the same concentration, we found no significant difference between photosensitizers. Yet, there seems to be a slight increase in the percentage of dose released for PDZ and PTG in the higher concentration compared to the lower.
The drug output is approximately constant for the first four minutes of nebulization, and it decreases over time, regardless of the photosensitizer (fig. 11). As it occurs for most jet nebulizers, the generation of droplets seems to be limited by the lowering volume of solution as the time goes by. As the initial output is higher for PTG, the volume limit is reached more quickly and nebulization stops after 5 minutes. For PDZ and ICG, the initial output is a little lower, and there is still mist formation for another minute. There does not seem to be any correlation between the VMD and the output rate.

During the experiment, we observed that the concentration of the compound in the remaining solution increased as the nebulization progressed (fig. 12). We proposed that the droplets formed carried less photosensitizer than the concentration of the solution, as it had been previously proposed by Diederik and co-workers in 2003. According to this model, as the concentration increased the droplets would have to have more compound in then, increasing the delivery over time. However, that was not observed in this experiment, since the volume limitation counteracts the rise in concentration, leading to the observed decrease in output from figure 11.

The output rate and the changes in concentration are important to determine the dose and nebulization time in the treatment design. Thus, it is important to notice that the results are given in terms of mass (µg/min and µg/mL) and not molarity. The outputs for PDZ (983.08 g/mol) and ICG (774.97 g/mol) are still comparable in terms of moles, but because PTG is a mixture of oligomers with high (however unpublished) molecular weight, it is unlikely that the increased mass output would reflect an increased molar output.

The concern about stability in the nebulization of many pharmaceuticals and the known degradation of ICG in water motivated a deep investigation of stability. To make sure that the process of droplet formation would not increase the degradation kinetics in any way, we studied the chemical composition of the three photosensitizer solutions with HPLC-MS-UV. The similarity between chromatograms was quite evident, but still the ionic mass and UV-Vis absorbance spectra of each of the peaks was compared to make sure that the composition was the same. All peaks had comparable intensities and corresponding spectra, so we concluded that the composition of the solutions had not changed because of the nebulization. The comparison of chromatograms and spectra of ICG is shown as an example, in figures 13 to 16.

Figures 13 and 14 also include a reference scenario in which such a change in composition occurs. The degradation of ICG in water is well described in the literature, which is why it is commercialized in powder form, to be dissolved immediately before use.
According to the manufacturer, solutions should be used within 10 hours of preparation. Therefore, after three weeks, we expected that the solution would no longer contain any photosensitizer. Surprisingly, the characteristic green color was sustained after this time, and it was found to be from the main degradation product detected (RT = 6.9 min). This compound and ICG itself have very similar absorbance spectra (fig. 15), what could mislead the analysis of stability through UV-Vis absorbance alone. A corresponding peak was also present in the freshly-prepared ICG solution. We could not determine if it was an impurity from the synthesis or a degradation product formed within the minutes before the analysis, and its absorbance and mass spectra did not correspond to any of the known products of ICG’s light-induced decomposition. Still, we found no evidence of an increase of such compound or decrease in ICG concentration after the nebulization process.

3.5 Conclusion

In summary, the findings of this chapter suggest that jet nebulization is an adequate method for the delivery of indocyanine green, Photodithazine® and Photogem® to the lungs. All three solutions are effectively carried out of the nebulizer, the microdroplets formed are of adequate size, and the compounds remain stable throughout the process. This is particularly promising considering the broad availability and reduced price of jet nebulization. Therefore, nebulization is a potential tool for the photodynamic treatment of not only pneumonia but other pulmonary disorders.
4 NEBULIZATION AND EXTRACORPOREAL ACTIVATION OF INDOCYANINE GREEN IN VIVO

4.2 Purpose

The nebulization of the three photosensitizers was equally successful, so indocyanine green was chosen as a model to demonstrate the delivery in vivo due to its prominent results in the treatment of bacterial pneumonia.\textsuperscript{5,6} Also, before studying the efficacy of the treatment, there was an interest in studying its effects in healthy organisms. Thus, this study aimed to assess the lung damage caused by the photodynamic action of nebulized ICG and infrared light in healthy mice.

Additionally, there was no published information about how the photosensitizer would be cleared from the lungs and distributed throughout the body. In this study, fluorescence imaging and spectroscopy were used to infer about the distribution of the photosensitizer delivered by nebulization within the lungs and to other internal organs in a murine model. Benefiting from the deep penetration of the infrared light in biological tissue, we observed the fluorescence of ICG extracorporeally and evaluated the applicability of its use as a monitoring tool for the proposed PDI treatment.

4.2 Methodology

4.2.1 Animal Model

Forty-five female BALB/c mice, from 8 to 10 weeks old, were obtained from CEMIB (Multidisciplinary Center for Biological Investigation on Laboratory Animal Science - Campinas, Brazil). They were anesthetized with a peritoneal injection of 5-10 mg/kg weight of xylazine (Anasedan, Ceva Santé Animale, Paulínia, Brazil) and 75-150 mg/kg weight of ketamine (Dopalen, Ceva Santé Animale, Paulínia, Brazil). The hair from the dorsal and lateral areas of the animals was removed using a depilatory cream (Veet\textregistered, Reckitt Benckiser, Cali, Colombia). Euthanasia was performed with an overdose of the same anesthetics. All animal experiments were approved by the Animal Ethics Committee of the São Carlos Institute of Physics, IFSC/USP – University of São Paulo (number 12/2016, approved on October 17\textsuperscript{th}, 2016).
4.2.2 Materials and Devices

Indocyanine Green (Ophtalmos, Sao Paulo, Brazil) was diluted in water for injection to the desired concentration immediately before use. The phosphate buffer saline (PBS) was prepared and sterilized to serve as the nebulization control. The Omron NE-C801 jet nebulizer was used, and a silicone tube was adapted to the mouthpiece of the nebulizer so that the aerosol would reach only the nose and mouth of the animals. The eyes were protected with damp gauze to prevent dryness. An illumination device was specially built for the treatment of mice, emitting monochromatic light at 780 nm with an irradiance of 60 mW/cm², and positioned in a way that the animal would be facing down and the light would reach the thorax from the back and the sides (fig. 17).

![Custom-made illumination device for the photodynamic treatment of mice. It is composed of 18 laser diodes with maximum emission at 780 nm. The arrows indicate the laser beam propagation. Source: GERALDE et al.]

The fluorescence widefield images were taken using an infrared camera with excitation at 780 nm was used for imaging (more information about this camera can be found in the Master’s dissertation of Govone). The spectral detection of the fluorescence was performed with a fiber optics setup, under excitation at 780 nm and the collection of the spectra between 300 and 1000 nm. A filter (850 ± 40 nm bandpass) was included to reduce the signal from the excitation, while allowing the fluorescence of ICG to pass. This same setup was used to compare the fluorescence of a solution of ICG (1.3 mM) right after preparation, and after 3 weeks stored at 4°C.
4.2.3 Experimental Procedures and Data Collection

4.2.3.1 Experiment 1: Infrared imaging detection of nebulized vs instilled ICG

In this experiment, one mouse received 30 µL of ICG 200 µM diluted in PBS via instillation, one received 12 minutes of nebulization with the same solution, and another animal received 6 minutes of nebulization of ICG 800 µM in water for injection (n = 1). Infrared images were taken from each animal before the administration of ICG (control) and 40 minutes afterwards. They remained under anesthesia throughout the procedure and were placed into 4 positions for each time point: laid facing down (back), laid sideways with the left side up (left) and the right side up (right). After that, the animals were euthanized and a laparotomy was performed in order to obtain images that would show the trachea, lungs, stomach, heart, liver and intestines.

4.2.3.2 Experiment 2: Extracorporeal monitoring of photosensitizer delivery over time

In the same animal that received ICG 800 µM in the previous procedure, the fiber optics setup was used to collect fluorescence spectra from the sides of the thorax and the back (n = 1). The detector was placed on the skin above the intercostal space in three different sites (as shown in fig. 18) and three measurements were taken for each spot immediately before the nebulization (control) and after 10, 20, 30, and 40 minutes. The “back” refers to the central position, although the actual spot was a little to the left to avoid the dorsal spine.

Figure 18 - Extracorporeal detection of fluorescence with the fiber optics setup. The red exes mark the spots where the measurements were taken.
Source: By the author.
4.2.3.3 Experiment 3: Tissue damage and immune response to the treatment

Six animals were divided into either control or treated groups (n = 3). Treated animals received 6 minutes of nebulization ICG 800 µM in water for injection and 33 minutes and 11 seconds of light exposure at 60 mW/cm², that resulted in a fluence of 120 J/cm². Control animals received 6 minutes of nebulization with PBS and no light exposure. After 48 h, the animals were euthanized. Then, the lungs were collected and kept in 4% formaldehyde for the histological analysis.

4.2.3.4 Experiment 4: Tissue damage and treatment monitoring with an increased dose of ICG

In this experiment, 15 animals were divided into the five groups (n = 3): the negative controls (C-) did not receive any treatment or infection; the positive controls (C+) were infected with *S. pneumoniae* (as described in Geralde, 2017); one group received only the nebulized photosensitizer (ICG); one received the ICG and the light exposure, simulating the treatment (PDI); and finally, one group received the light exposure, and nebulized PBS instead of the photosensitizer (Light). The nebulization lasted for 10 minutes and had either ICG at a concentration of 1.30 mM in water for injection or PBS. For the PDI and Light groups, light exposure began halfway through the nebulization (5’) and lasted 33’11’’, resulting in a dose of 120 J/cm². The total treatment time was of about 38 minutes.

Table 4 – Groups of experiment 4. The exes mark the treatments received by each group.

<table>
<thead>
<tr>
<th>Group</th>
<th>Nebulization</th>
<th>PBS</th>
<th>Light exposure</th>
<th>Infection</th>
<th>Fluorescence detection</th>
<th>Histology</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>ICG</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>PDI</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Light</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>C+</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

Source: By the author.

In the treatment of the ICG and PDI groups, the extracorporeal detection of fluorescence (shown in fig. 18) was performed before the procedure (0 min), during and after the nebulization (5; 10 min), and then during and after the irradiation (20, 40 min). Measurements were taken in triplicate as previously described (fig. 1). After 48 hours,
euthanasia was performed and the right lobes of the lungs were collected and kept in a 4% formaldehyde buffered solution for the preparation of histological slides. The treatments given to each group are summarized in table 4.

4.2.3.5 Experiment 5: Body distribution of lung-delivered ICG with and without exposure to light and time-dependency of the tissue and immune response

For the photosensitizer distribution experiment, a total of 21 animals were divided into controls (n = 3, no treatment), ICG (nebulization only) and PDI (nebulization and light exposure). The treated animals were further divided into each of the timepoints: 1, 4 and 24 hours (n = 3). They received 6 minutes of nebulized ICG at a concentration of 800 µM, and the PDI group received 20 minutes of light exposure (72 J/cm²). Each animal was anesthetized after the corresponding time and the fluorescence of ICG was detected externally, as previously described.

Then, the animals were euthanized and the internal organs were exposed. The fiber was placed on the right and left sides of the lung, the stomach, liver, small and large intestines, bladder, spleen, and the heart of each animals, and measurements were registered in triplicate. To obtain the bronchoalveolar lavage fluid (BALF), a catheter was inserted into the trachea and used to deliver 0.4 mL of cold PBS to the left lobe of the lung, that was isolated from the other lobes with a suture knot on the right bronchus, as shown in figure 19. The PBS was recovered and kept in an ice bath for up to 2 hours until preparation for the flow cytometry. The liver and the right lobes of the lungs were collected for histological analysis and kept in an isotonic solution with 4% formaldehyde.
Figure 19 - Isolation of the lung lobes for analysis. A knot was tied on the location indicated by the black line, so that the bronchoalveolar lavage would reach only the left lobe. The lobes on the right side (superior, middle, inferior and post-caval) were collected for histology.
Source: By the author.

4.2.4 Fluorescence Data Processing

Using a custom-made C program, the fluorescence spectra obtained were preprocessed using spectral smoothing, with each point resulted from the average of its neighboring 25 points. The interval corresponding to the ICG fluorescence band (795-890 nm) was normalized by the amplitude of the excitation wavelength (780 nm) to account for changes in optical coupling, and then integrated to generate a single output. Then, using the software Origin 9.0 (OriginLab, Northampton, USA), all obtained values were divided by the median baseline for that location, that was either the measurement from the control group, or the “t = 0 min” timepoint. As a consequence, the results are shown in terms of relative fluorescence. They are represented by the mean and standard deviation obtained for each site and time point. The one-way ANOVA and a post-hoc Tukey test were used to compare the data sets, with a level of significance of 0.05.

4.2.5 Flow Cytometry

The BALF was centrifuged for 5 minutes at 1500 rpm, using the Centrifuge 5702 (Eppendorf, Hamburg, Germany). The supernatant was removed and the recovered cells were resuspended in 300 μL of PBS. Then, they were stained with a fluorescence-tagged anti-CD45
monoclonal antibody (APC Rat Anti-Mouse CD45, BD Biosciences, New Jersey, USA), using the protocol recommended by the seller. CD45 is a surface marker of all immune cells, so this staining procedure allows for their identification in flow cytometry. Non-specific binding sites were protected using the untagged CD16/CD32 monoclonal antibody.

Flow cytometry was carried out using the BD Accuri C6 Plus cytometer (Becton Dickinson, Franklin Lakes, United States) and the software BD Accuri C6 Plus 1.0.23.1 (Becton Dickinson, Franklin Lakes, United States) was used for data analysis. CD45-negative events were disregarded, and the classification of cell types was based on forward scattering (FSC, that correlates to size), side scattering (SSC, correlates to granularity), and autofluorescence.

4.2.6 Histological Analysis

The lung and liver samples were blocked into paraffin, sliced, stained with hematoxylin and eosin (HE) and mounted onto microscopy slides. Two slides were prepared from each sample. Each slide received a code and was analyzed using the Axio Observer Z1 microscope (Zeiss, Oberkochen, Germany), in search for signs of inflammation and tissue damage. In the lungs, such signs were: inflammatory infiltrates, hyaline membrane deposits, fibrosis, bronchiolar epithelial damage, hyperplasia, alveolar thickening and vascular congestion. Liver slices were searched for necrosis, inflammatory infiltrates, vascular congestion, microvacuolization and changes in color. Since the injuries were patchy, the extent of damaged tissue was compared semi-quantitatively (by assigning a symbol: -, +, ++ or +++). Only after that, the codes were matched with the corresponding experimental groups and the results were compared. Representative images of each group were obtained with the Pannoramic Desk digital slide scanner and the Pannoramic Viewer software (3DHISTECH Limited, Hungary).
4.3 Results

4.3.1 Selective Fluorescence of ICG with excitation at 780 nm

![Fluorescence signal detected with the experimental setup of a solution of ICG 1.3 mM soon after its preparation, and after complete degradation of its main compound. Source: By the author.]

ICG and its degradation product in water have very similar absorbance spectra (as shown in the previous chapter), but figure 20 shows that only the photosensitizer is able to emit fluorescence with excitation at 780 nm. It is also evident from fig. 20 that this experimental setup produces spectra with a lot of noise, what motivated the data processing described in item 4.2.4.

4.3.2 Delivery of ICG via Nebulization versus Instillation

The fluorescence from ICG when administered via instillation and nebulization was compared using infrared imaging. It was possible to see illuminated spots on the right and left sides of the animal 40 minutes after the instillation of the photosensitizer, that were not present after neither the nebulization of the lower now the higher concentration (fig. 21). After the organs were exposed (fig. 22), it became evident that these spots corresponded to a strong fluorescence signal coming from the stomach and the duodenum. The lungs, trachea and mouth of the animal that received instillation also exhibited intense fluorescence. After the
nebulization with the equivalent concentration (200 µM), the mouth and trachea showed signs of ICG fluorescence, but the lungs did not. As the concentration of ICG was increased to 800 µM, the signal in both the mouth and the trachea was also increased, and the fluorescence in the lungs became visible, although not as much as after instillation.

<table>
<thead>
<tr>
<th>Control</th>
<th>Instilled</th>
<th>Nebulized 200 µM</th>
<th>Nebulized 800 µM</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Left side" /></td>
<td><img src="image2" alt="Instilled left side" /></td>
<td><img src="image3" alt="Nebulized 200 µM left side" /></td>
<td><img src="image4" alt="Nebulized 800 µM left side" /></td>
</tr>
<tr>
<td><img src="image5" alt="Back" /></td>
<td><img src="image6" alt="Instilled back" /></td>
<td><img src="image7" alt="Nebulized 200 µM back" /></td>
<td><img src="image8" alt="Nebulized 800 µM back" /></td>
</tr>
<tr>
<td><img src="image9" alt="Right Side" /></td>
<td><img src="image10" alt="Instilled right side" /></td>
<td><img src="image11" alt="Nebulized 200 µM right side" /></td>
<td><img src="image12" alt="Nebulized 800 µM right side" /></td>
</tr>
</tbody>
</table>

Figure 21 - Infrared images of the animals after receiving each of the treatments in experiment 1. The first row shows the left side, the second to the back, and the third the right side. The arrows indicate the spots where the fluorescence of ICG was observed. 
Source: By the author.
Figure 22 - Infrared images of the animals that received each of the treatments from experiment 1, exposing the organs after the euthanasia. Organs in which fluorescence was evident are indicated with the black arrows.

Source: By the author.

4.3.3 Extracorporeal Detection and Activation of the Nebulized ICG in the Lungs

The fluorescence detected with the fiber optics system was significantly different after the delivery of ICG 800 µM with nebulization (fig. 23). The amount of fluorescence detected at each position at the same timepoint was not significantly different, so the average between the three positions in represented. The signal remained higher, between 1.5 and 3-fold compared to the baseline, throughout the whole interrogation time of the experiment.
Figure 24 - Relative fluorescence detected externally over time on animals that received nebulization with the increased concentration of ICG with (PDI) and without (ICG) exposure to infrared light (experiment 4). The horizontal bars indicate the duration of each of the treatments. The difference between treatments is statistically significant at 20 and 40 minutes.

Source: By the author.

Figure 25 - Relative fluorescence detected externally over time on each of the three positions for animals that received nebulization with the increased concentration of ICG with (PDI) and without (ICG) exposure to infrared light (experiment 4). There is no significant difference between positions for a given time and treatment combination. The difference between treatments is significant at 20 and 40 minutes for all positions.

Source: By the author.
Figure 24 shows the average fluorescence from the different positions, and the timeline of the treatment for experiment 4. The fluorescence increases in the animals that received only ICG, whereas after the light treatment the animals have the same fluorescence signal as they had before the nebulization of photosensitizer. Figure 25 shows the results achieved for each position individually, and there is no statistically significant difference between them.

The fluorescence detected externally in experiment 5 was significantly after the nebulization in both ICG and PDI group for up to 24 hours (fig. 26). The animals that received only ICG had higher fluorescence signals after 4 and 24 hours than animals that also received light.

![Graph showing relative fluorescence over time for ICG and PDI treatments](image)

Figure 26 - Relative fluorescence detected externally at different times after the treatments from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light. There was no significant difference between the three sites, so the averages are shown.

Source: By the author.

### 4.3.4 Body Distribution of the Photosensitizer

The relative fluorescence detected in the organs of interest, where ICG was expected to be present, is shown in figure 27. Unfortunately, although some tendencies may be drawn from the graphs, the measurements from spot to spot showed huge variance, so it was not possible to find a statistically significant difference between most of the groups.
In the lungs, the signal increased after the nebulization, and remained higher than the one for the control throughout the 24 hours, even for the groups that were treated with light. In some cases, the fluorescence detected showed an increase of over 100-fold the baseline value. In both lungs, only at 24 hours the fluorescence from the ICG group is significantly higher than the one of PDI. In the stomach, the fluorescence signal peaks 4 hours after the nebulization, for both treatments, and reaches over a hundred times the baseline for the ICG group. At this time, the fluorescence signal for ICG is significantly higher than for PDI. At 24 hours, although some of the measurements were quite high, both ICG and PDI groups are indistinguishable from the control.
In the liver, the increase in fluorescence is already significant one hour after the nebulization, and it remains elevated for the 24 hours. There is no significant difference between the ICG and PDI groups. In the small intestines, the fluorescence is significantly increased for the ICG group at 4 and 24 hours. For the PDI group, none of the timepoints were significantly greater than the control. In the large intestines the only mean significantly greater than the control was for the PDI at 24 hours.
The relative fluorescence detected in the other organs, where ICG was not expected to be found, is shown in figure 28. In these organs, the increase in fluorescence was not as high as in the organs of interest, reaching an increase of only about 10-fold the baseline measurements. However, the results were much more consistent (note the smaller deviation bars compared to the graphs in fig. 27). In the bladder, the increase in fluorescence is significant for both ICG and PDI at 4 and 24 hours, and the results for the ICG group are significantly larger than PDI at both timepoints. The same behavior is observed in the measurements of the spleen and the heart, but in these organs the increase in the PDI group is already noticeable at 1 hour.

Figure 29 presents a summary of the same data shown in figures 27 and 28. It makes it possible to observe which organs showed the largest increase in fluorescence for each treatment and time. After one hour, the highest fluorescence was measured in the lungs and
liver, for both treatments. After 4 hours, the stomach shows the largest relative fluorescence, and after 24 hours, the highest increases in fluorescence were measured in the lungs (ICG) and the large intestine (PDI). However, the high variance between measurements in each organ limit the conclusions that can be drawn from it.

Figure 29 - Summary of the relative fluorescence detected directly in the organs over time after each treatment from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light.

Source: By the author.

An attempt to correlate the fluorescence detected in the lungs to the fluorescence detected externally is shown in figure 30. This graph combines all the data collected from controls, ICG and PDI groups. However, there was no linear correlation observed for neither the right or the left side.
Figure 30 - Correlation between the fluorescence detected externally on each side, and the fluorescence detected directly in the corresponding lung of the same animal. Data from the control, ICG and PDI groups from experiment 5: ICG 800 μM with (PDI) and without (ICG) exposure to 72 J/cm² of infrared light.
Source: By the author.
4.3.5 Tissue Damage and Immune Response

The histological analysis of experiment 3 showed that, 48 hours after the treatment, the animals from both the control and treated groups had no sign of lung damage (fig. 31). However, in experiments 4 and 5, all animals showed some level of alveolar thickening, vascular congestion and inflammatory infiltration in the lungs (figs. 32 and 33). Comparing the 10-fold magnification images, it is possible to see that only the latter two show patches of thickened alveolar tissue. Still, in the semi-quantitative comparison, the positive controls were distinguishably more injured than all the other groups. The lungs of the animals that received ICG or the simulated PDI treatment showed no difference from the untreated or the Light controls in neither experiment. There was no difference between the livers of the animals from experiment 5 either (fig. 34).

Figure 31 - Representative histological slides of the lungs from control and treated animals from experiment 3, with increasing magnification. Both groups showed healthy tissue, and there was no difference between them. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x).

Source: By the author.
Figure 32 - Representative histological slides of the lungs of each group in experiment 4, with increasing magnification. All animals had some level of lung injury, but the sick positive controls (C+) had higher scores than any other group. Note the patches of alveolar thickening in the 10-fold magnification pictures (blue arrows). The 100-fold magnification shows the contrast between the healthy tissue, with ample air pockets, and the dense injured tissue. The vascular congestion is more evident in the 400-fold magnification. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x).

Source: By the author.
Figure 33 - Representative histological slides of the lungs from each group in experiment 5, with increasing magnification. All animals had some level of lung injury, and the groups were indistinguishable from one another. Note the patches of alveolar thickening in the 10-fold magnification pictures (blue arrows). The degree of vascular congestion varied from animal to animal (see the 400-fold magnification), but there was no difference between the groups. The scale bars correspond to 2000 μm (10x), 200 μm (100x) and 50 μm (400x).

Source: By the author.
Figure 34 - Representative histological slides from the livers of the groups in experiment 5, with increasing magnification. There was no difference between the groups. The scale bars correspond to 200 μm (100x) and 50 μm (400x).

Source: By the author.
Finally, to further inquire about the immune response in the lungs, the composition of the BALF of the animals from experiment 5 was investigated using flow cytometry. The percentage of alveolar macrophages in the BALF after 1, 4 and 24 hours from nebulization with ICG 800 µM with (PDI) and without (ICG) 72 J/cm² of infrared light was compared against the controls (fig. 35). Although some of the animals had very different percentages of macrophages than others, there was no significant difference between the averages of each group.

![Figure 35: Percentage of macrophages in the bronchoalveolar lavage fluid from the animals at different times after either ICG alone or PDI. There was no significant difference between groups.](image)

Source: By the author.

### 4.4 Discussion

Since indocyanine green is a fluorescent dye, researchers have used its fluorescence signal to infer about its distribution in the body. In 2013, Zhang and co-workers used a fiber-based system to validate the development of a fluorescence molecular tomography method to study the pharmacokinetics of intravenously given ICG. This study proposed the use of a similar system to infer about the distribution of the photosensitizer delivered by nebulization within the lungs and to other organs of mice. However, in light of the findings of the previous chapter, that the main degradation product of ICG in water has a similar absorbance spectrum than ICG itself (see fig. 15), there was an interest in making sure that the fluorescence detection method would be selective to the active photosensitizer. Figure 20 shows the fluorescence signal detected for a fresh ICG solution and a completely degraded ICG solution.
using the experimental setup that was proposed for the animal studies. Although both solutions absorb greatly at 780 nm, only the fresh ICG emits fluorescence in the selected wavelength range. The degradation of ICG due to light exposure generates different compounds than the degradation in water, and no longer absorbs light at 780 nm, so there was no reason to assume such compounds would interfere with the readings. Thus, we proposed that the only corrections to be made were regarding the autofluorescence of the tissues and organs in which the measurements were made. For that, all the comparisons were made in terms of relative fluorescence (of the obtained values by the average baseline fluorescence of that given location).

The qualitative comparison using the infrared camera suggested that the distribution of ICG in the body is different when delivered via instillation and nebulization. The fluorescence detected externally in the animal that received instillation in figure 21 seems to come from the stomach, where the fluorescence is strongest for this animal in figure 22. The lack of that externally observable fluorescence in both the animals that received nebulization suggests that the amount of photosensitizer that initially goes to the stomach is much smaller using this method. This is important to notice because the ICG in the stomach could be activated by the light in the PDI treatment and cause unwanted effects.

However, for the same concentration, the nebulization was not able to generate the same fluorescence in the lungs as the instillation. The delivery of substances to the lungs of mice using aerosols can be quite challenging because of their small tidal volume, and specially using equipment that was designed for humans, the deposit in the upper respiratory tract seems inevitable. Still, by increasing the concentration of ICG in the nebulized solution, it was possible to see the fluorescence in the lungs. Moreover, ICG seems to be deposited in the mouth and throat after instillation as well as nebulization.

The main difference between the two methods, besides the accumulation in the stomach, is that much of the ICG is wasted in the nebulization, so the initial dose needs to be much higher in the latter. The instillation delivered about 6 nmol of ICG, while 1.6 μmol had to be put in the nebulizer in order to produce similar results. However, in humans, the efficiency of the nebulization is expected to be much higher (as discussed in the previous chapter).

Although it was not possible to see the difference in external fluorescence using the camera, the fiber optics setup showed that it significantly increased over time after nebulization with the higher concentration of ICG (fig. 23). This result was already expected, since the external fluorescence imaging presents a much lower signal. Using fluorescence imaging, the influence of the scattering and absorption is significantly higher when compared
to the one observed using the fluorescence spectroscopy. The main advantage on using an
external measurement is the possibility to monitor the kinetics of the photosensitizer
distribution without the interference of a surgical procedure to allow the fiberoptic probe
positioning at the organ surface.

This motivated another experiment that compared the relative fluorescence after
nebulization with and without exposure to the light, simulating what would happen in the PDI
treatment (figs. 24 and 25). This time, the concentration of ICG was increased even more, in
the hope that the difference in fluorescence would be even more evident.

The first thing to notice from this experiment is that, if it is possible to detect such a
difference, it is also possible to activate the photosensitizer deep into the tissue with that
wavelength. This is a confirmation of what was predicted by our previous studies with animal
models.\(^6,8^1\) Secondly, the fluorescence continues to increase for some time after the end of the
nebulization, and the peak in relative fluorescence was not higher for the increased
concentration of ICG. More research is needed to explain this effect, but it is possibly related
to the high concentration of the photosensitizer (0.8 and 1.3 mM), in which it is known to
aggregate and self-quench.\(^7^7\) Perhaps due to the distribution of ICG within the respiratory
system, the aggregates become undone and there are more molecules emitting fluorescence.
This is relevant because light should reach the lungs while the photosensitizer is around the
bacteria, to enhance effectiveness, and hopefully before it reaches the deeper tissue, to prevent
collateral effects. However, aggregated ICG may not be effective in the PDI treatment, so the
timeline of the treatment should be studied carefully.

Figure 25 compared the three different parts of the thorax to choose the best location for a
future monitoring of the treatment. However, they all had similar profiles and standard
deviations, so none of them was clearly better than the others. Also, in the group that was
exposed to light the fluorescence returned to baseline after 20 minutes, which probably
indicates the photobleaching of ICG as it goes through the photodynamic process. This helped
confirm that the activation of ICG was in fact occurring and led to the assumption that the
light dose delivered after this time was perhaps unnecessary.

To see if that was in fact the case, the light dose was reduced to 72 J/cm² in experiment 5.
This time, the external fluorescence of the animals was monitored for up to 24 hours, as figure
26 shows. However, the relative fluorescence is significantly higher after the animals received
the simulated treatment, which means that this light dose was not sufficient to bleach all of
the ICG. The external fluorescence continues to increase until 4 hours after the nebulization
and remains elevated after 24h. This corroborates with the proposal that ICG is initially
aggregated in the lungs, and as it slowly distributes into the tissue, its ability to fluoresce is restored.

Figures 26 also show that the relative fluorescence in the animals that only received ICG is much higher after 4 and 24 hours compared to the animals that had the simulated treatment (PDI). This difference probably accounts for the ICG that interacted with the light, generated the desired effect, and was eventually photobleached.

The fluorescence detected directly in the lungs was supposed to help quantify this difference. However, the results (shown in fig. 27) were incredibly variable, and as a consequence the Tukey test did not show a significant difference between most of the groups. It is possible that the increased variance was related to an uneven build-up of ICG in the lungs, that was not perceived in the external detection due to the diffusion of the light. Because of that, we decided not to conclude anything about the distribution of the photosensitizer in the lungs from this experiment alone.

The fluorescence detected in the other organs of interest (also in fig. 27) presence high variance as well. Although ICG is unstable in water, its interaction with proteins in the organism helps prevent the degradation reaction. That is why it is still possible to detect its fluorescence even after 24 hours. However, ICG administered intravenously has a reported half-life of only a few minutes. It is rapidly cleared from the blood, taken up by the liver, and excreted in the bile. Therefore, it was expected to be found in the liver at first, and then accumulate in the intestines. To some extent, this is what happens. Nevertheless, the fluorescence in the liver remains elevated until 24 hours, which once more suggests that ICG is slowly released from the lungs, circulates the blood and is cleared in the bile throughout the 24 hours.

The stomach was an organ of interest as well, because of the comparison between nebulization and instillation. No fluorescence was seen in this organ with the camera 40 minutes after the nebulization (fig. 22), and figure 27 shows that after one hour, there was still no difference from the baseline. However, at 4 hours, there is a peak in the relative fluorescence for both ICG and PDI groups. It is possible that the photosensitizer that accumulated in the upper airways was cleared and swallowed, eventually reaching the stomach.

There was no particular reason for ICG to be found in the bladder, spleen, or heart. The fluorescence was monitored in these organs to infer about its presence in the blood and help understand the distribution as a whole. The fact that variability of results in figure 28 is much
smaller in these organs than the organs of interest suggests that the latter could be a consequence of how ICG distributes in the healthy organs, and not necessarily because of an animal-to-animal variability. These results also support the belief that part of the ICG delivered was degraded by the light exposure in the PDI groups, and that the photosensitizer is gradually released from the lungs to the rest of the body.

Figure 29 allows for the comparison of the relative fluorescence changes over time throughout the body. In the first hour, the highest values of relative fluorescence were found in the lungs and the liver. Since the extracorporeal activation covers the entire thorax, it could unintentionally activate ICG in the stomach, liver and heart. If the timeline of the treatment is similar to the ones presented in experiments 3 to 5, in which the illumination starts during or right after the nebulization and goes on for 20-33 minutes, the liver would be the organ most susceptible to collateral effects. On the other hand, treatment schedules with a longer waiting period could potentially harm the stomach.

Unfortunately, the high variance between the results limits a further discussion of the topic. Thus, detecting the fluorescence directly into the surface of the organs showed to be a limited technique to study the biodistribution of ICG. The difference in fluorescence from spot to spot suggest that the distribution, especially in the lungs, is not homogenous. Therefore, the presented results do not necessarily represent the organs as a whole. This method was initially proposed because of its simplicity. However, better results would probably be obtained from extracting the photosensitizer from the organs, which is complex in itself, and using more sophisticated techniques, like HPLC-MS.

This high variance also translated into a lack of correlation between the fluorescence detected in the lungs, and externally through the sides of the thorax (fig. 30). The external detection was much more consistent than the direct one, probably because the light diffused through the tissue and the uneven distribution of ICG from spot to spot did not matter as much. It was not possible to validate a real-time treatment monitoring proposal with this technique, but it does not mean that the external fluorescence does not correlate to the average ICG concentration in the lungs. The combination of extraction from the tissues and HPLC analysis is a possible method to overcome this problem, too.

Regarding the tissue damage of the treatment, experiment 3 provided the initial comparison between the simulated treatment and the controls (fig. 31). In experiment 4, the concentration of ICG and the nebulization time were increased to simulate a more intense treatment, and light-only, ICG-only and positive controls were included (fig. 32). In
experiment 5, samples were taken at different times in search for transient effects that are sometimes seen in other treatments (figs. 33 and 34). In a study from 2017, for example, mice that received an anti-infective peptide showed epithelial sloughing and bronchitis 4 hours after the treatment, but these signs were gone after 24 hours. Nonetheless, no histological difference was found between treated animals and healthy controls, regardless of the ICG and light dose, and the time passed after the procedure.

This suggests that, at least in the presented conditions, the PDI using ICG and light at 780 nm does not cause acute damage to the lungs or the liver. The lack of difference does not mean, however, that the treatment is not effective. In a previous study that used instilled ICG in the PDI of pneumonia, there was no histological difference in the lungs between groups, but a single treatment increased the survival rate in 30 days from 60% to 100%.

Figure 36 shows examples of lung histology pictures for healthy mice (36A and B) and different murine models of acute lung injury (36C through H). Comparing these with the obtained results of this study, it becomes evident that the animals from experiment 3 (fig. 31) were healthy, but the animals from experiments 4 and 5 (figs. 32 and 33, respectively) had some level of lung damage (+ or ++), characterized by patches of alveolar thickening and vascular congestion, consistent with the exposure to lipopolysaccharides (36C and D) or bacteria (36G and F). These animals were supposed to be healthy as well, but these findings suggest that they might have been exposed to lung pathogens unintentionally, prior to the experiment. Still, the extent of lung injury was significantly larger (+++) in the positive controls, that were infected with *S. pneumoniae* using a well-established protocol (see the bottom row of fig. 31).
Figure 36 - Comparison of selected models of acute lung injury. A and B: normal mouse lungs. The alveolar walls are very thin, and the majority of the alveoli contain no cells (magnification in A, 100x; B, 400x). C and D: lungs from a mouse euthanized 18 h after intratracheal instillation of 5 ng/g lipopolysaccharide. Note the patchy nature of the injury (C, 100x) and the presence of inflammatory infiltrates and vascular congestion (D, 400x). E and F: lungs from a mouse euthanized 21 days after the administration of intratracheal bleomycin. Note the presence of fibrotic areas (arrows) (E, 200x; F, 400x). G and H: lungs from a mouse euthanized 12 h after aerosolization of Escherichia coli, 1 x 10^8 CFU/mL. Note diffuse thickening of the alveolar spaces and intra-alveolar neutrophilic infiltrates (G, 200x; H, 400x). HE stain.

Source: Adapted from MATUTE-BELLO et al. 88
Figure 37 - Histopathology of mice treated with 5 mg/kg of talaporfin sodium (Laserphyrin®) and 50 J/cm² of light (664 nm laser) including the tumor and adjacent normal liver tissue. The regional boundary line between the laser radiation area and the outside was clear (a). The bottom row of pictures shows: the tumor (b); ischemic necrosis and focally living hepatocytes around Glisson in the peripheral area of the irradiation (c); and the slight denatured hepatocytes around the irradiation area (d). HE stain.

Source: KASUYA et al. 89

There was also no difference between the livers of the animals from experiment 5 (fig. 34). An example of liver damage caused by PDT is shown in figure 37. Because there was evidence of ICG in the liver 1 hour after the treatment (from fig. 27), there was a chance that there would already be some photosensitizer in this organ during the illumination, causing ischemic necrosis (like the one on fig. 37C). The absence of signs of necrosis suggests that the proposed treatment is safe for the liver.

The findings from the flow cytometry analysis of experiment 5 support the argument that there was no difference between treated and control animals (fig. 35). However, the high variability between subjects limits the drawing of conclusions from this experiment. The number of cells that were properly stained and detected by the equipment was small and varied greatly between samples (not shown). Thus, the BALF recovery and staining protocols should be improved for future experiments.

Furthermore, the study was overall limited by the small number of subjects in each group. The experiments 1 and 2 were proof-of-concept pilots, so a single animal was used for each group. However, for experiment 3 to 5, three animals in each group were probably not sufficient to overcome the high variability that is intrinsic to living creatures. Ethics plays a huge role in deciding how many animals should be used in an experiment, but future discussions of these findings would benefit from replicating the presented experiments.
4.5 Conclusion

The findings of this study suggest that nebulization is a viable method for the delivery of the photosensitizer indocyanine green to the respiratory tract in the photodynamic inactivation of pneumonia. In the murine model, both instilled and nebulized ICG are deposited in the upper and lower respiratory tract, but only instillation creates an immediate build-up of this photosensitizer in the lungs. However, the dose of ICG required using nebulization needs to be much higher than using instillation in order to yield similar results.

It is possible to verify the activation of nebulized ICG with extracorporeal light using fluorescence. The characteristic emission of this photosensitizer increases over time after delivery by nebulization and is reduced after illumination with infrared light. However, the return of the fluorescence signal to baseline does not seem to correspond to the complete consumption of the photosensitizer. ICG is slowly released from the lungs and is present in many organs for at least 24 hours after the nebulization, even if the animal goes through the illumination. This combination of nebulized ICG and extracorporeal light does not seem to be harmful to the lungs or livers of mice.

Although it is possible to infer about the presence of ICG in the body by detecting its fluorescence, this method was not considered ideal for the detection of ICG directly in the exposed organs, since the variation of results from spot to spot was impressively high. More research is required to fully understand the distribution and the effects of the photosensitizer in the lungs and other organs after the pulmonary administration.
5 CONCLUSIONS

This study proposed to investigate the applicability of nebulization as a delivery method for photosensitizers, in the hope to advance the research of the photodynamic inactivation of bacterial pneumonia. It showed that the photosensitizers indocyanine green, Photodithazine® and Photogem® are all compatible with the Omron NE C-801 jet nebulizer, and present attributes that suggest that the delivery to the lungs of patients will be efficient. It also showed conditions in which it was possible to: deliver indocyanine green using jet nebulization to a murine model; activate it using extracorporeal infrared light; and not cause acute lung or liver damage. However, questions remain to be answered regarding the body distribution of the photosensitizer in the organism after the pulmonary delivery, and the immune infiltration after the treatment. Still, nebulization presented itself as a promising tool for the delivery of photosensitizers to the respiratory tract.
REFERENCES


39 SILVA, R. C. Avaliação da eficiência fotodinâmica de fotosensibilizadores com aplicação em terapia fotodinâmica. 2007. 71 p. Dissertação (Mestrado em Ciências) - Instituto de Química de São Carlos, Universidade de São Paulo, São Carlos, 2007.


79 GOVONE, A. B. Desenvolvimento de um sistema de imagem de campo amplo de fluorescência para localização de linfonodo sentinel empregando indocianina verde. 2016. 88p. Dissertação (Mestrado em Ciências) - Instituto de Física de São Carlos, Universidade de São Paulo, São Carlos, 2016.


87 GERALDE, M. C. Avaliação in vivo da inativação fotodinâmica para tratamento de pneumonia. 2017. 110 p. Tese (Doutorado em Biotecnologia ) - Centro de Ciências Exatas e Tecnológicas, Universidade Federal de São Carlos, São Carlos, 2017.


APPENDIX A – Chromatograms of the photosensitizer solutions before and after nebulization

Total Mass Chromatogram: freshly prepared PDZ

Total Mass Chromatogram: nebulized PDZ

Total Wavelength Chromatogram: freshly prepared PDZ

Total Wavelength Chromatogram: nebulized PDZ
Total Mass Chromatogram: freshly prepared PTG

Total Mass Chromatogram: nebulized PTG

Total Wavelength Chromatogram: freshly prepared PTG

Total Wavelength Chromatogram: nebulized PTG
CERTIFICADO DA COMISSÃO DE ÉTICA NO USO DE ANIMAIS – CEUA/IFSC

Certificamos que o projeto intitulado “Descontaminação do sistema respiratório por inativação fotodinâmica: estudo in vivo”, protocolo nº 12/2016, sob a responsabilidade da pesquisadora Mariana Carreia Geralde e sob supervisão do Prof. Dr. Vanderlei Salvador Bagnato – que envolve a produção, manutenção e/ou utilização de animais pertencentes ao filo Chordata, subfilo Vertebrata (exeto humanos), para fins de pesquisa científica – encontra-se de acordo com os preceitos da Lei nº 11.794, de 8 de outubro de 2008, do Decreto nº 6.699, de 15 de julho de 2009, e com as normas editadas pelo Conselho Nacional de Controle da Experimentação Animal (CONCEA), e foi aprovado pela COMISSÃO DE ÉTICA NO USO DE ANIMAIS (CEUA) do Instituto de Física de São Carlos (IFSC), em reunião de 17/10/2016.

<table>
<thead>
<tr>
<th>Finalidade</th>
<th>Pesquisa Científica</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vigência da Autorização</td>
<td>17/10/2016 a 01/01/2019</td>
</tr>
<tr>
<td>Espécie/linhagem/raça</td>
<td>Camundongo Isogênico - BALB/c</td>
</tr>
<tr>
<td>Nº de animais</td>
<td>140</td>
</tr>
<tr>
<td>Peso/Idade</td>
<td>20-25mg / 8-10 semanas</td>
</tr>
<tr>
<td>Sexo</td>
<td>70 machos e 70 fêmeas</td>
</tr>
<tr>
<td>Origem</td>
<td>Faculdade de Medicina de Ribeirão Preto – USP CEMIB de Campinas/SP</td>
</tr>
</tbody>
</table>

São Carlos, 17 de outubro de 2016.

Prof. Dr. Fernando Fernandes Paiva
Presidente
Comissão de Ética no Uso de Animais – IFSC/USP