Due to the overuse of antibiotics in medicine and food production, and their targeted mechanism of action, an increasing rate in spreading of antibiotic resistance genes has been noticed. This results in inefficient therapy outcomes and higher mortality all over the world. *Pseudomonas aeruginosa* (carbapenem-resistant) is considered one of the top three critical species according to the World Health Organization’s priority pathogens list. This means that new drugs and/or treatments are needed to tackle infections caused by this bacterium. In this context search for new/alternative approaches that would overcome resistance to classical antimicrobials is of prime importance.

The use of antimicrobial photodynamic inactivation (aPDI) and antimicrobial peptides (AMPs) is an efficient strategy to treat localized infections caused by multidrug-resistant *P. aeruginosa*. In this study, we have treated *P. aeruginosa* cells photodynamically in the presence and in the absence of AMP (CAMEL or pexiganan). The conditions for aPDI were as follows: rose bengal (RB) as a photosensitizing agent at 1–10 µM concentration, and subsequent irradiation with 514 nm-LED at 23 mW/cm² irradiance. The analysis of cell number after the treatment has shown that the combined action of RB-mediated aPDI and cationic AMPs reduced the number of viable cells below the limit of detection (<1log⁰CFU/ml). This was in contrast to no reduction or partial reduction after aPDI or AMP applied separately. Students t-test was applied to test the statistical significance of the results. Noteworthy, our treatment proved to be effective against all 35 clinical isolates of *P. aeruginosa* tested within this study, including those characterized as multiresistant. Moreover, we demonstrated that such treatment is safe and does not violate the growth dynamics of human keratinocytes (77.3–97.64% survival depending on the concentration of the studied compounds or their mixtures).

**Keywords:** antibiotic resistance, *Pseudomonas aeruginosa*, photoinactivation, CAMEL, pexiganan, XDR, MDR, antimicrobial peptides
INTRODUCTION

Since the discovery of penicillin in 1929, followed by a great success in the control of bacterial infections, a golden era of antibiotics has begun. This successful era lasted for many years, during which whole-cell screens, search for broad-spectrum natural and synthetic antibacterial drugs, were provided to the market and to the clinics. Later on, due to the increase in bacterial resistance to conventional drugs, the search turned into more targeted molecules, which, however, did not bring much success in recent years. Nowadays, we are facing the problem of global resistance and failure in antimicrobial drug discovery (Brown and Wright, 2016). Of particular concern are infections caused by the ESKAPE pathogens, which include Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa, and Enterobacter spp. (Rice, 2008).

Pseudomonas aeruginosa is a chief opportunistic pathogen that can cause nosocomial infections in susceptible persons in medical institutions. This bacterium can spread via human-to-human direct distribution, and also via water systems (up to 50%) in hospital wards (Blanc et al., 2004). In the hospitals, it was isolated from various medical devices, sanitary installations, but also from flowerpots (D’Agata, 2014). P. aeruginosa is responsible for the complicated infections, particularly in people with compromised immunity, e.g., oncological patients, people after transplantation, elderly people, that are frequently hospitalized. This bacterium causes skin and soft tissue infections, which can be fatal for people with burns and after surgeries. Mortality among P. aeruginosa-infected patients is estimated at 20%, but it can reach as high as 50%, e.g., in the case of placenta infection (Lautenbach et al., 2010; Ceniceros et al., 2016). The most dangerous population among P. aeruginosa isolates constitute those producing metallo-β-lactamases, conferring resistance to all penicillins, cephalosporins, and carbapenems. The latter has been known as the last resort drugs in the treatment of Gram-negative infections (Potron et al., 2015). Much attention has nowadays been paid to the development of strategies that can lower the use of antibiotics and slow down the spread of the resistance phenomenon. Being in-line with this trend, photodynamic inactivation of multiresistant pathogens has emerged as a promising alternative to antibiotics.

Antimicrobial photodynamic inactivation (aPDI), also known as photoantimicrobial chemotherapy (PACT), relies on the action of three elements: a small-molecular-weight chemical compound (photosensitizer, PS), light, and oxygen. Light irradiation activates PS, which leads to the generation of singlet oxygen (energy transfer) and/or oxygen radicals (electron transfer). All the reactive oxygen species (ROS) generated during aPDI are responsible for cytotoxic effect toward bacterial cells due to inactivation of proteins, lipids, and nucleic acids. Because of multtargeted action of ROS, acquiring resistance to this form of antibacterial treatment is highly unlikely and has not been experimentally confirmed so far (Giuliani et al., 2010; Tavares et al., 2010; Wainwright et al., 2017). Another advantage of PDI includes double selectivity based on the local delivery of a PS and light, that both need to act concomitantly to produce ROS. Such a local delivery of a PS and light allows avoiding systemic exposure and potential adverse effects of the treatment. Practically, every living microorganism can be inactivated by means of aPDI. Often the presence of bacterial cells is not sufficient to trigger disease, and the damage to host cells is caused by various virulence factors produced by the pathogen. aPDI has been shown to efficiently reduce virulence factors which seems to be a rational approach to control infection (Fila et al., 2017). It was found, however, that the efficacy of photoinactivation of Gram-negative species is less efficient as compared to Gram-positive ones, due to the presence of an outer membrane, which constitutes a natural barrier limiting a simple diffusion of a PS (Bertoloni et al., 1990). This means, that in vivo, a high concentration of a PS and irradiance have to be delivered to obtain satisfactory clinical outcome (Hashimoto et al., 2012; Fila et al., 2016). Thus in the case of in vivo treatment, the danger of cyto- and/or phototoxicity exists toward host eukaryotic tissues, resulting from higher light doses and/or higher PS concentration applied to photoinactivate Gram-negative bacteria. These might include damage to biomolecules leading to the breakdown of cell structure, and damage to organelles, as well as initiation of necrotic or apoptotic pathways. Various approaches have been reported to literature to overcome the problem of lower PDI efficacy toward Gram-negative bacteria as compared to Gram-positive ones, e.g., addition of positive charge to a PS (Hamblin et al., 2002; Tegos et al., 2006). Also, polymyxin B addition to anionic or neutral porphyrins enabled to carry out photoinactivation of Gram-negative species (Nitzan et al., 1992).

Another approach to diminish the negative effects of high doses of PS and light is to decrease the effective concentration of PS by combining its action with antimicrobial peptides (AMPs). AMPs have been studied in the context of combating drug-resistant pathogens as an encouraging alternative to classical antimicrobials. They are part of the innate immune system and act against diverse-spectrum microorganisms. AMPs are generally amphipathic and have a length of up to 50 amino acids. Their net positive charge and a high proportion of hydrophobic residues mediate binding to anionic groups on the bacterial cell surface and facilitate interaction with membranes to produce lethal pores (Zasloff, 2002). The bacterial surface has a net negative charge due to the presence of lipoteichoic acids and lipopolysaccharides as opposed to sphingomyelin, phospholipids, and cholesterol in the membrane of eukaryotic cells. Such differences between bacterial and eukaryotic membrane composition are the basis of selectivity for AMPs activity.

There have been several studies on the subject of combining the action of aPDI with AMP through producing conjugates of a photosensitizer (mostly porphyrin derivatives, or xanthenes) with various peptides (Bourre et al., 2010; Yang et al., 2011; Liu et al., 2012; Dosselli et al., 2013, 2014; Johnson et al., 2013; Costley et al., 2017; Le Guern et al., 2017; Tsuchikama et al., 2017). However, the subject of a simple combination of two kinds of the molecules was underinvestigated. Such an approach is simpler and also cheaper as the conjugation step can be avoided. There has been only one elegant work done recently on the combination of three types of a photosensitizer,
nearly curcumin, methylene blue, and chlorin C6 with an AMP aurein 1.2, where free, non-conjugated molecules showed excellent activity (de Freitas et al., 2018). In the cited work, the authors focused on Gram-positive species, namely Enterococcus faecium, with an unknown antibiotic resistance pattern. In this work, we have been investigating Gram-negative species P. aeruginosa, which is currently considered a species of critical priority. As a photosensitizer for our analysis we have chosen rose bengal (RB) – 4,5,6,7-tetrachloro 2',4',5',7'-tetratriiodofluorescein, that is known to mainly generate singlet oxygen upon visible light exposure. RB possess a strong absorption band at 550 nm, and strong molar absorption coefficient (95,000 dm$^3$ mol$^{-1}$ cm$^{-1}$) (Ludvíková et al., 2016). RB properties have been extensively investigated in technology known as photochemical tissue bonding in tissue repair (Gu et al., 2011).

In our research, we have investigated the new potential therapy that could be applied in the treatment of local infections caused by multidrug-resistant P. aeruginosa. We examined the combined action of rose-bengal-based aPDI and two AMPs, namely pexiganan and CAMEL, showing significantly enhanced effect of the studied combination. Moreover, we investigated our approach toward the wide repertoire of clinical isolates of P. aeruginosa, including carbapenem-resistant strains, proving that the proposed approach is a viable antimicrobial strategy. Our extensive analysis revealed that such a new approach was not toxic to human skin keratinocytes, nor did it affect their growth dynamics.

MATERIALS AND METHODS

Bacterial Strains and Growth Conditions

The reference P. aeruginosa ATCC® 10145 strain was used in the study. The 35 clinical P. aeruginosa isolates, were provided by Julianna Kurlenda and isolated from patients hospitalized in the Provincial Hospital in Koszalin and the Provincial Hospital in Gdansk, and Sebastian Mucha from Independent Public Clinical Hospital No. 1 of Silesian Medical University in Katowice (Table 1). Bacterial cultures were grown aerobically in LB medium (Carl Roth, Karlsruhe, Germany) at 37°C with shaking (150 rpm) or on LB agar (Sigma-Aldrich, Munich, Germany). Antibiotic susceptibility for piperacillin (PIP); ticarcillin-clavulanic acid (TIM); piperacillin-tazobactam (TZP); cefepime (FEP); ceftazidime (CAZ); imipenem (IPM); meropenem (MEM); doripenem (DOR); gentamicin (GEN); amikacin (AMK); netilmicin (NET); tobramycin (TOB); ciprofloxacin (CIP); levofloxacin (LVX); colistin (CST); polymyxin B (PMB); fosfomycin (FOF); aztreonam (AZT) was determined by the Vitek 2 system, AST-N331 cards (bioMérieux, Carpenne, France). The Vitek 2 minimum inhibitory concentration (MIC) results were interpreted using an Advanced Expert System according to Clinical and Laboratory Standards Institute (CLSI) recommendations (CLSI, 2012). Categorization of resistant, intermediate resistant, susceptible P. aeruginosa was determined according to the guidelines of the European Centre for Disease Prevention and Control (ECDC) and the Center for Disease Control and Prevention (CDC). The description of strains as multidrug-resistant (MDR) was defined as acquired non-susceptibility to at least one agent in three or more antimicrobial categories, extensively drug-resistant (XDR) was defined as non-susceptibility to at least one agent in all but two or fewer antimicrobial categories (i.e., bacterial isolates remain susceptible to only one or two categories) (Magiorakos et al., 2012).

Chemicals

The photosensitizer used in the study was rose bengal (RB, 4,5,6,7-tetrachloro-2',4',5',7'-tetratriiodofluorescein, SigmaAldrich, Munich, Germany). The prepared stock solution of 10 mM RB was diluted with a sterile double distilled water and kept at 4°C for up to a month. All the reagents used for the synthesis were of pure analytical grade purchased from Sigma-Aldrich, Germany.

Peptides Synthesis

Peptides: CAMEL (CAM; KWKLFKIGAVLKNH$_2$) and pexiganan (PEX, GIGKFLKKAKFGBKVKILKKK-NH$_2$) were synthesized by solid-phase method using Fmoc (9-fluorenlymethyloxycarbonyl) chemistry on a Polystyrene Amide-RAM resin as described previously (Sikora et al., 2018). The peptides were purified by Reverse-Phase High-Performance Liquid Chromatography (RP-HPLC), and then freeze-dried. The mobile phase was water and acetonitrile containing 0.1% TFA (v/v). UV detection at 214 nm was used. The identity of the compounds was confirmed using mass spectrometry (ESI-MS). The compounds were analyzed by HPLC in the following conditions: 20–80% acetonitrile (0.1% TFA), 10 min.

Molecular weights of peptides were as follows: CAM – 1770.3 g/mol; PEX –2477.2 g/mol.

Determination of CAM, PEX MIC/MBC Values

Minimum inhibitory concentration (MIC, the lowest concentration of AMP that inhibited bacterial growth) toward P. aeruginosa 10145 was assessed as follows: a total of a 100 μL of bacterial suspension in brain-heart infusion medium (BHI, bioMérieux, Carpenne, France) was placed in 96-well plate. The inoculum of bacteria was 10$^4$ CFU in each well. A 100 μL of twice diluted AMP CAM or PEX was added to wells (concentration range 0–512 μg/mL). The cells were incubated 24 h at 37°C. The MBC (minimum bactericidal concentration) values were assessed as a minimal concentration of AMP that caused 99.9% death of bacterial cells (Cockerill et al., 2012).

Photoinactivation Experiments

Bacterial strains were cultured in 5 mL of LB medium for 18–22 h at 37°C with agitation (250 rpm). Cells were diluted with a fresh broth to the density of 0.2 McFarland units (10$^7$ CFU/mL). A total of 100 μL of each culture was loaded into a 96-well plate and incubated in the dark at 37°C for 15 min, either with or without the addition of RB. The concentrations of RB used in the study were 1–100 μM. Accordingly, in the photoinactivation experiments with the use of AMPs, PEX, or CAM was added along with RB and incubated in the dark at 37°C for 15 min. To remove unbound compounds, cells were washed three times

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TABLE 1 | Drug resistance characteristics of Pseudomonas aeruginosa clinical isolates used in the study.

| Strain | Source of isolation | PIP | TIM | TZP | FEP | CAZ | IPM | MEM | DOR | GEN | AMK | NET | TOB | CIP | LVX | CST | PMB | FOF | AZT | Resistance status |
|--------|---------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|------------------|
| 1      | 10145               | ND  | R   | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 2      | 3042/s              | Urine | R   | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 3      | 3318/p              | Ear |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 4      | 2969/s              | Wound |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 5      | 1561/o              | Bronchi | R   | R   | R   | R   |     |     |     |     |     |     |     |     |     |     |     |     |     | R    | MDR  |
| 6      | 3404/p              | Urine | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 7      | 3117/s              | Abdomen |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 8      | 3146/s              | Wound |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 9      | 4190/p A            | Wound | R   | R   | R   |     | IR  |     |     |     |     |     |     |     |     |     |     |     |     | R    | MDR  |
| 10     | 1959/o              | Bronchi | R   | R   | R   | R   | R   |     |     |     |     |     |     |     |     |     |     |     |     | R    | XDR  |
| 11     | 3109/o              | Urine | R   | R   | R   | R   | R   | R   |     |     |     |     |     |     |     |     |     |     |     | R    | MDR  |
| 12     | 1286/o              | Urine, blood | R   | R   | R   | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 13     | 55/K                | Bronchi | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     |     | R    | XDR  |
| 14     | 978/K               | Pleura | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     |     |     | R    | XDR  |
| 15     | K3                  | Blood | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     |     |     |     |     | R    | MDR  |
| 16     | K4                  | Blood | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     |     |     |     |     | R    | XDR  |
| 17     | 792/K               | Wound | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | IR  |     |     |     |     |     | R    | MDR  |
| 18     | 393/K               | Urine | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     | IR  |     |     |     |     | R    | XDR  |
| 19     | 133/K               | Urine |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 20     | 23/K                | Urine |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | IR   |     |
| 21     | 887/K               | Wound | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     | R   | R   |     |     |     | XDR  |
| 22     | 169/K               | Nose | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | IR  |     |     |     |     |     | XDR  |
| 23     | 3752/sz             | Ear |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 24     | 4908/p              | Urine | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 25     | 3007/sz             | Uterus |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 26     | 15/p1               | Tracheostomy tube |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 27     | 149/p               | Urine |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 28     | 153/s               | Urine |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 29     | 547/p               | Ear |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 30     | 1885/K              | Bronchi | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     | R    | XDR  |
| 31     | 556/K               | Wound | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   | R   |     | R    | XDR  |
| 32     | 1651/p              | Urine |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |
| 33     | 1304/s              | Urine | R   | R   | ND  | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    | MDR  |
| 34     | 1804/p              | Urine | R   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    | IR   |
| 35     | 2284/p              | Ear |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    | IR   |
| 36     | 143/p               | Wound |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     | R    |

Antimicrobial susceptibility testing included following antimicrobials: PIP, piperacillin; TIM, ticarcillin-clavulanic acid; TZP, piperacillin-tazobactam; FEP, ceftazidime; CAZ, ceftazidime; IPM, imipenem; MEM, meropenem; DOR, doripenem; GEN, gentamicin; AMK, amikacin; NET, netilmicin; TOB, tobramycin; CIP, ciprofloxacin; LVX, levofloxacin; CST, colistin; PMB, polymyxin B; FOF, fosfomycin; AZT, aztreonam. R, resistant; IR, intermediate resistant; no letter – sensitive.
with 1 mL of sterile PBS. Samples were irradiated with a total fluence of 15, 30, and 60 J/cm² (duration of irradiation 668, 1335, and 2668 s, respectively). For the irradiation procedure, LED lamps (SecureMedia, Poland), designed and produced for the Laboratory of Molecular Diagnostics, emitting green light were used (λ<sub>max</sub> 514 nm, FWHM = 33 nm: the width of a spectrum curve measured between those points on the y-axis which are half the maximum amplitude) (Ogonowska et al., 2018). The irradiance was 23 mW/cm². The distance from a LED to an illuminated sample was 10 cm, and the power density measurement represents the value at the distance of 10 cm. Aliquots incubated in the dark with and without RB served as dark controls. Additionally, during each experiment light-only treatment was performed without the addition of RB. After irradiation, 10 µL aliquots were taken to perform 10-fold serial dilutions in PBS, ranging from 10<sup>-1</sup> to 10<sup>-4</sup>. Ten microliter aliquots of each dilution were plated on LA plates (Carl Roth, Karlsruhe, Germany). Cells were incubated overnight at room temperature, and then, for 2–4 h at 37°C, the colonies formed were counted, and the results were analyzed statistically. Each experiment was performed in triplicate.

**Photosensitizer Uptake**

Overnight cultures prepared as those for the photoinactivation experiments were diluted to 0.2 McFarland units. A total of 100 µL of culture was placed in Eppendorf tube and washed three times with PBS. After the last centrifugation step, cells were diluted in 100 µL of PBS. Then, RB was added to a final concentration of 1, 2, 5, and 10 µM and incubated for 15 min in the dark at 37°C. Next, cell suspensions were centrifuged (5 min, 3300 rcf) and supernatants were placed in a 96-well plate. Absorption was measured at λ = 565 nm. The cells treated similarly but without incubation with RB served as background control. The concentrations of RB accumulated were read based on calibration curves obtained from the measurements of free RB.

**Fluorescence Microscopy**

Overnight cultures of *P. aeruginosa* were diluted 10-times (c.a. 3.1 McFarland units). A total of 100 µL of culture was placed in Eppendorf tube and washed with PBS 3 times, following 5 min centrifugation at 3300 rcf. Next, CAM, PEX and/or RB were added to a final concentration of 50 µM or 100 µM. Cells were incubated in the dark at 37°C for 5–30 min. The excess of the compounds was centrifuged (5 min, 3300 rcf) and the pellet was resuspended in 50 µL PBS. Four µL of bacterial suspension was placed on microscope glass. The observation was carried out under a fluorescence microscope (Olympus BX51, Hamburg, Germany) with the F-View Soft Imaging System digital camera. Excitation/emission spectra were λ<sub>ex/em</sub> = 510–550 nm/> 570 nm. At least three observations were made on three independent days.

**Photohemolysis Assay**

A 2%-solution of sheep erythrocytes was prepared in PBS by diluting a 1 mL stock solution (100%) in 50 ml PBS. Erythrocytes were centrifuged (5 min, 3300 rcf), and resuspended in 50 mL of the RPMI-1640 medium (Sigma-Aldrich, Hamburg, Germany). A 400 µL of a 2% solution of erythrocytes were placed in Eppendorf tube, together with CAM, PEX and/or RB to a final concentration of 5 µM each. Cells were incubated in the dark for 5 or 15 min at 37°C. Then, cells were washed twice with PBS and suspended in a final volume of 400 µL. A 200 µL was transferred into 96-well plate and kept in the dark, the second half of the solution was irradiated (fluence: 30 J/cm², irradiance: 23 mW/cm²). The plates were centrifuged (6 min, 3100 rcf), and supernatants were transferred to a new 96-well plate. The absorbance of released hemoglobin was measured at λ = 470 nm. A 10% SDS-treated erythrocytes were used as a control of complete hemolysis (100% hemolysis). The data were presented according to the equation:

\[
\% \text{integrity} = 100\% - \% \text{hemolysis}
\]

Three independent experiments were performed and presented values are the mean ± SD.

**Photo- and Cytotoxicity Assay Based on MTT**

HaCaT cells (CLS 300493) were seeded at the density of 1 × 10<sup>4</sup> cells/well the day before treatment in three biological replicates for each condition in two 96-well plates (for light and dark conditions, respectively). Cells were grown in a standard humidified incubator at 37°C and 5% CO₂ atmosphere in DMEM high glucose medium supplemented with 1mM sodium pyruvate, 1 mM non-essential amino-acids, 100 U/mL penicillin, 100 µg/mL streptomycin, 2 mM glutamine, 10% fetal calf serum (all reagents from Life Technologies/Thermo Scientific). The compounds were added to the final concentration of 5 µM or 10 µM RB, 5 µM or 10 µM of AMP (CAM or PEX), or mixtures thereof. The compounds were added directly to the medium, and further incubated for 15 min at 37°C. Afterward, the cells were washed twice with PBS and finally dissolved in 100 µL PBS. Next, cells were subjected to illumination with 514 nm-light (irradiance: 23 mW/cm², 22 min). Cell survival was measured after 24 h of incubation at 37°C by MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] assay. Briefly, a 10 µL of MTT solution (12 mM) was applied to each well and incubated for 4 h at 37°C. Cells were then lysed in DMSO, and the absorbance of the formazan was measured at 550 nm using a plate reader (Victor 1420 multilabel counter, Perkin Elmer). The results are presented as a fraction of untreated cells, and calculated as a mean of three independent biological replicates with the standard deviation of the mean.

**Analysis of Cell Culture Growth Dynamics Based on the xCELLigence System**

HaCaT cells (CLS 300493) were seeded the day before treatment, in 7 technical replicates for each condition, at a density of 1 × 10<sup>4</sup> per well (according to the manufacturer’s protocol) on E-plate (ACEA Biosciences Inc.). Cells were grown in a standard humidified incubator at 37°C and 5% CO₂ atmosphere, in xCELLigence RTCA instrument, (ACEA Biosciences Inc.) in
DMEM medium as indicated above. Next day, when cells were in their logarithmic phase of growth, the experiment was carried out. To do this, the plates were removed from the xCELLigence device, spent medium removed by aspiration and changed to a medium containing a studied compound (as indicated). After 15 min incubation at room temperature in darkness, the cells were washed twice with PBS and the fresh medium was added. Then, cells were exposed to 514 nm light (irradiance: 23 mW/cm², 22 min), and returned to an xCELLigence device for 70–120 h. The cell index was measured every 15 min and recorded automatically.

**Statistical Analysis**

The results of photodynamic inactivation, are presented as the average of at least three independent experiments. Statistical significance was assessed using Student's *t*-distribution method with the use of Excel software. Three biological replicates of each strain were included separately into the analysis. For comparison with the use of RB and 514 nm-light, we subjected treated clinical isolates groups, ANOVA (Welch test) and RIR Tukey post *hoc* testing were performed using a STATISTICA 10 software (StatSoft Inc., 2011, United States).

**RESULTS**

**Photoinactivation of *Pseudomonas aeruginosa* With RB**

In order to assess the initial conditions for photoinactivation of *P. aeruginosa* with the use of RB and 514 nm-light, we subjected the bacterial cells (*P. aeruginosa* 10145 strain) to incubation with increasing concentration of RB, and illuminated cells with various light doses. *P. aeruginosa* is not vulnerable to the action of RB-mediated photokilling at low concentrations of RB (40 µM) and low light doses (15–30 J/cm²). We observed that even at a very high concentration of 80 µM RB and a light dose of 30 J/cm², the decrease in the viable cell count was less than 2 log₁₀ units. Increasing the RB concentration to a 100 µM resulted in 3 log₁₀ units reduction in bacterial count. Accordingly, increasing the light dose to 60 J/cm² resulted in 6 log₁₀ units reduction in survival as compared to reference cells (cells not treated with light and RB) (Figure 1). It is worth mentioning that the light itself has the impact on survival (Amin et al., 2016) e.g., the 60 J/cm² treatment without RB results in 1 log₁₀ units reduction in bacterial count. In further experiments this light dose was excluded, to avoid the influence of light alone on the results of experiments.

**Photoinactivation of *P. aeruginosa* in the Presence of Antimicrobial Peptides Is Increased**

Next, we checked the influence of the AMP on the photoinactivation efficiency. To this end we treated the cells with RB and AMP together in a single reaction mixture, incubated for 15 min, and subjected to illumination procedure as described in the Section "Materials and Methods." We applied RB concentrations up to 10 µM, where no effect on the viability of bacterial cells could be observed. As for the concentrations of CAM and PEX, we chose two different concentrations below MBC value. MBC for PEX was 18 µM, and for CAM – 25 µM. As it can be seen (Figure 2), RB itself at concentrations up to 10 µM, does not have any significant impact on *P. aeruginosa* survival, neither in the presence nor in the absence of light. As for the PEX influence on the viability of cells, 5 µM concentration resulted in 2.9 logs reduction, both in the presence and in the absence of light, whereas 10 µM concentration was responsible for the significant decrease in cell number below the limit of detection (Figure 2A). The analysis of the combined effect of RB and PEX (5 µM RB and 5 µM PEX) revealed enhanced bactericidal action that resulted in the significant decrease in cell survival (below the limit of detection, > 6 logs). A similar trend was observed for the combination of RB with CAM. In this case, the action of 10 µM CAM resulted in 2.08 logs reduction in surviving cells number. When CAM was mixed with RB (both at equal concentrations of 10 µM), and subjected to illumination, the reduction in survival was below the detection limit (Figure 2B).

It is worth to notice that the observed significant reduction in the number of viable bacterial cells was connected to the presence of light. This further means that the observed killing was related to a combination of aPDI with AMPs, rather than the interaction of the RB itself with AMP.

**The Presence of Antimicrobial Peptide Affects the Accumulation of RB in *P. aeruginosa* Cells**

To investigate the basis of the observed enhanced effect of RB and PEX/CAM on bacterial cell survival we analyzed the process of RB accumulation in *P. aeruginosa* cells. Bacterial cells were incubated either with RB alone, or RB and one of the AMPs. After 15 min incubation, cells were pelleted, and the unbound RB remained in the supernatant. The concentration of RB that remained in supernatants was assessed based on
FIGURE 2 | Photoinactivation of Pseudomonas aeruginosa with RB in the presence of antimicrobial peptides. The cells were incubated with rose bengal alone (RB), antimicrobial peptide (PEX or CAM) alone, or with rose bengal and one of the antimicrobial peptides together (RB+PEX or RB+CAM). Results for PEX are shown in (A), for CAM in (B). The concentrations of compounds are indicated on the abscissa. In the case of RB and AMP mixtures, cells were either kept without light application (dark) or they were illuminated with a total light dose of 30 J/cm². Each result is a mean ± SE of log₁₀ CFU/ml. Significance at respective p-values is marked with asterisks (*p = 0.002 for PEX, and *p = 0.003 for CAM). Arrows indicate the lowest effective concentration of RB/AMP resulting in a reduction of bacterial cell number below the limit of detection.

the calibration curve and subtracted from RB concentration at which the cells were incubated. In general, when cells were incubated with RB only, the amount of accumulated RB was 5.1 µM (for 10 µM initial RB concentration). The accumulation increased almost twofold for 10 µM initial RB concentration in the presence of either PEX (RB accumulation = 9.2 µM) or CAM (RB accumulation = 8.7 µM) (Figure 3). Interestingly, a similar effect of increased RB accumulation was observed for PEX and for CAM, although the phototoxic effect of aPDI was more pronounced in the presence of RB/PEX than in the presence of RB/CAM (Figure 2). No significant difference in RB accumulation efficiency was observed with respect to time the cells were incubated with the studied compounds.

Accordingly, we observed the enhanced effect of RB accumulation with the use of fluorescence microscopy. To this end, we incubated cells (overnight cultured) in the presence of 100 µM RB. We applied this concentration of RB to obtain sufficient staining of cells. As it can be easily followed, the fluorescence of the cells was not detected in the presence of RB alone, even if the incubation time was prolonged up to 30 min (Figure 4). Only upon simultaneous incubation of RB and PEX, the fluorescence of cells could be detected. Moreover, we observed that upon PEX addition the cells had a tendency to group themselves together, however, the observed aggregates should be further studied to indicate their nature. CAM was omitted from the experiments as in tests measuring quantitative accumulation, it presented similar to PEX accumulation profile. In summary, the obtained results indicated that accumulation of RB inside the cells is facilitated upon PEX presence.

The Integrity of Erythrocytes Is Affected Upon aPDI
To check biocompatibility of a compound often hemolysis assay is performed as a common measure of safety. Therefore, we checked how the treatment with RB and/or PEX affects the membrane of erythrocytes. The hemolytic behavior of PEX is similar to that of CAM so we omitted the latter from the experiment (Sikora et al., 2018). As it is clearly shown, in dark conditions membranes stay intact for the time of the experiment. Neither presence of RB, PEX nor the mixture of both
negatively affects erythrocytes membrane in the experimental conditions in the dark. However, upon light treatment, we observed that about 55% of erythrocytes were subjected to damage. And the process of lysis occurred only when RB and light treatment was performed (Figure 5). This indicates that RB and light treatment is phototoxic for erythrocytes. Interestingly, PEX itself did not damage the erythrocytes' membrane. This further means that PEX is relatively specific toward microorganisms.

**Cell Viability of HaCaT Cells Upon Photodynamic Treatment in the Presence of Antimicrobial Peptides Is Not Disturbed**

To assess the cyto- and phototoxic effect of rose bengal alone or in combination with AMPs on human skin cells the effect of phototreatment on human keratinocytes was measured. We tested the concentrations of the compounds that were used in the bacterial cells inactivation experiments and increased the dose to 10 µM of RB and AMPs. Based on the MTT assay results (Figure 6), the viability of the cells was not affected by the presence of RB alone, neither in dark (92.01 and 92.12% survival upon 5 or 10 µM) nor upon irradiation (90.08 and 89.56% survival upon 5 or 10 µM). Also, the combined concentrations of RB and PEX were not cyto- (97.64% survival) nor phototoxic (79.07% survival) even at higher 10 µM concentrations of compounds tested. After incubation of keratinocytes with CAM (10 µM) and upon irradiation, about 15.75% of cells were killed. Accordingly, when cells were incubated with a mixture of CAM and RB (each at 10 µM), 18.36% of cells were photoinactivated. This indicates that mostly CAM was responsible for the observed effect. Similar observations were made after analysis of dark toxicity. Again, CAM alone (10 µM) or in mixture with RB (10 µM) decreased the survival of keratinocytes by 17.3 and 23.73%, respectively. The light itself did not affect membrane integrity. Taken together the results indicate acceptable toxicity for the combined treatment of aPDI with PEX or CAM toward human skin keratinocytes. Although statistical significance was achieved for CAM treatment, the numerical values were not much different from these for PEX (Supplementary Table S1). Moreover, ~80% survival upon treatment is generally considered an acceptable level of toxicity as related to nearly 6 log₁₀ reduction in the viable bacterial count. It is also worth to emphasize that PEX itself does not cause HaCaT toxicity (neither with or without light).
interface. When the cells are grown on the surface, they alter the local ionic environment leading to an increase in electrode impedance. The more cells on the electrodes, the larger the change of impedance. The relative change in electrode impedance is represented by Cell Index (CI), which is a quantitative measure of cells attached to the electrodes. The higher the value of CI, the more cells are growing on the plate.

As the MTT assay measures the cell viability at the particular time point, we were interested in following the fate of cell growth during the longer time period. To study the eukaryotic cell growth dynamics after phototreatment, we applied the label-free automated monitoring of cell status using micro-electronic sensor plate (E-plate). The change of properties of the analyzed cells influences the passage of electrons and ions on a sensor surface thus providing an information about the biological status of cells. The physical parameter that is measured during such tests is electronic impedance, which is influenced by cell number, viability, morphology, the degree of adhesion. When cells are not present on a sensor E-plate, the impedance of the electrode depends on the ionic environment at the electrodes/solution interface. When the cells are grown on the surface, they alter the local ionic environment leading to an increase in electrode impedance. The more cells on the electrodes, the larger the change of impedance. The relative change in electrode impedance is represented by Cell Index (CI), which is a quantitative measure of cells attached to the electrodes. The higher the value of CI, the more cells are growing on the plate.

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mammalian cells, we were interested if it can be applied against various clinical isolates. As bacterial clinical isolates originating from a particular niche may substantially differ with respect to their properties, we evaluated the combined treatment of aPDI and AMP PEX on several clinical isolates. The analyzed clinical strains were isolated from diverse sources and expressed various antimicrobial resistance pattern (Table 1). Within the group of 35 analyzed clinical isolates, there was 14, which were characterized as multidrug- or extended drug-resistant P. aeruginosa. Interestingly for some of them, e.g., 156/o, 1885K, 556/k only combined treatment PEX and aPDI was sufficiently effective. All treatments efficiencies for individual test strains are presented in Table 2.

We investigated the efficacy of photodynamic inactivation alone, PEX treatment alone, and combined action of both approaches toward all 35 clinical isolates. The experiments were performed with light treatment or in the dark. We observed that in our experimental aPDI protocol photodynamic treatment itself had a very limited influence on cell survival (Figure 8 RB light). By applying this aPDI protocol, namely 30 J/cm², 5 µM RB we could not observe a significant bactericidal reduction in the number of surviving cells. According to the American Society for Microbiology statement every new approach has to prove an efficacy of min 3 log₁₀ reduction of CFU before it can be termed “antimicrobial.” Here, in our research, we indicated the reduction of ≥3 log₁₀ units, as antimicrobial reduction that is considered biologically relevant. When cells were treated with PEX only, either with or without the addition of light, the antimicrobial effect could be observed for 55% (20/36) of clinical isolates (Figure 8 PEX light, PEX dark). In the case of remaining 16 strains, the combination of PEX and RB was sufficient to obtain bactericidal reduction (i.e., reduction ≥3 log₁₀ units) even without the addition of light (Figure 8 PEX+RB dark) for seven isolates. As for the remaining 9 isolates out of the group of 16, the efficient bacterial cells killing (i.e., reduction ≥3 log₁₀ units) was only possible after combining PEX with aPDI. Of interest, in this group, the killing efficiency accounted for ≥5 log₁₀ units, with a single exception of isolate 4908/p (4.88 log₁₀ units reduction). This indicates that only a combination of AMP PEX and photodynamic treatment resulted in the inactivation of all analyzed clinical isolates (Figure 8 RB+PEX light). Worthy of notice is the fact that after combined treatment the reduction in cell survival was higher than 5 log₁₀ units in case of each strain (Table 2), again with a single exception already above mentioned.

DISCUSSION

We showed in our study that inactivation of the Gram-negative bacterium is efficient and can be exploited as potential antimicrobial therapy for treatment of local infections caused by multiresistant P. aeruginosa. By the combination of AMP with RB, otherwise inefficient toward Gram-negative bacteria, we obtained a 20-fold decrease in the concentration of RB used in the treatment to obtain min. 5 log₁₀ units reduction in cell survival (100 µM/60 J/cm² vs. 5 µM/30 J/cm²). This result should be considered as more than a 20-fold decrease as actually lower total fluence was used for the combined treatment as compared to the individual treatment of aPDI (i.e., without AMP). It is worth to emphasize that application of lower fluence (in this case 30 J/cm² vs. 60 J/cm²) results also in shortening of irradiation time (in this case 1335 s vs. 2668 s), which is of great clinical importance. The enhanced photokilling effect of the combined action of aPDI and AMP was due to the facilitated accumulation of RB in bacterial cells in the presence of AMP as compared to cells incubated only with RB (Figures 3, 4). Recently it was shown that in the presence of AMP aurein 1.2, the uptake of methylene blue (MB) was twice as much as compared to MB uptake alone. This doubled value of uptake was observed also in presented work for a combination of RB and CAM or RB and PEX (Figure 3). It is known from the literature data that the uptake process of a PS is not the only and critical determinant of efficient photodynamic inactivation. This further means that a PS does not have to be accumulated inside the cells to perform an efficient photodynamic action (Preuss et al., 2013). For example, chlorin-e6 uptake was significantly decreased in the presence of AMP, however, the synergistic action of aPDI and AMP was still observed (de Freitas et al., 2018). Such a characteristic of aPDI is an advantage as it means that bactericidal action can be started from the outside of the cell rather than from the inside where DNA could be affected and potentially mutagenized. We did not observe any difference in facilitating RB uptake between the two AMPs applied. Both CAM and PEX peptides had a similar impact on the accumulation of RB in P. aeruginosa cells. These two peptides have similar properties as for their structural features (helical peptides), and net positive charge (+6, and +10 for CAM and PEX, respectively), however, PEX was more potent in exhibiting antimicrobial action. PEX concentration reduced by 50% was sufficient to obtain a similar bactericidal effect as compared to CAM.

When considering the application of combined action of aPDI and AMPs, both components should be taken into account, namely PS and AMP. From the previously published research, it is known that not all combination of aPDI an AMP are robust and can be effective. Aurein 1.2 was effective when combined with aPDI, but only when MB or chlorin e6 were used as photosensitizers. In the case of another photosensitizer curcumin, the synergistic effect was not observed at all. This proves that the enhanced action of aPDI and AMP was PS-dependent (de Freitas et al., 2018). In our studies, RB proved to be very good PS for combined aPDI and AMP action, whereas it is commonly known that RB itself is not effective against Gram-negative species (Bezman et al., 1978; Schäfer et al., 2000; Demidova and Hamblin, 2005; Wen et al., 2017). This photosensitizer efficiently produces singlet oxygen, however, the outer membrane present in Gram-negatives prevents anionic RB from reaching the critical cellular structures like a cytoplasmic membrane. Thus, our hypothesis to employ cationic AMP increases outer membrane permeability and allows for the efficient accumulation of RB in cells proved valid. Our results from accumulation experiments confirmed the rationale of our hypothesis. It could be of further interest to explore if AMP facilitates RB loading into cells via direct interaction between the two, or AMP changes the properties of the cellular membrane, which further results in more efficient accumulation of RB.
TABLE 2 | The efficiency of RB-mediated photodynamic inactivation of *P. aeruginosa* clinical isolates in the presence and in the absence of PEX.

| Strain | Log<sub>10</sub> reduction in survival of bacterial cells after treatment<sup>a</sup>: | RB | PEX | RB+PEX | Light only | Antibiotic resistance status<sup>b</sup> |
|--------|-------------------------------------------------|-----|-----|--------|-----------|-----------------------------------|
|        | Light (J/cm<sup>2</sup>), Dark (µM)              | Light, Dark (µM), Light (J/cm<sup>2</sup>) | Light, Dark (µM), Light (J/cm<sup>2</sup>) | Light, Dark (µM), Light (J/cm<sup>2</sup>) | Light only (µM), Light (J/cm<sup>2</sup>) |
| 1      | 10145                                           | 0.21, 0.13 | 3.34, 2.70 | 5.69, 2.68 | 0.20 |
| 2      | 3042/s                                          | 0.44, 0.12 | 6.45, 6.45 | 6.45, 6.45 | 0.03 |
| 3      | 3318/p                                          | 0.48, 0.01 | 5.82, 5.82 | 5.82, 5.82 | 0.14 |
| 4      | 2969/o                                          | 0.35, 0.25 | 6.12, 4.49 | 6.12, 6.12 | 0.20 |
| 5      | 1561/o                                          | 1.11, 0.14 | 1.46, 1.33 | 5.37, 2.25 | 0.42 |
| 6      | 3404/p                                          | 0.81, 0.03 | 5.37, 5.37 | 5.37, 5.37 | 0.36 |
| 7      | 3117/s                                          | 0.35, 0.19 | 6.04, 6.04 | 6.04, 6.04 | 0.30 |
| 8      | 3146/s                                          | 0.63, 0.18 | 1.97, 1.48 | 5.90, 2.18 | 0.43 |
| 9      | 4190/p A                                        | 0.58, 0.41 | 3.44, 2.61 | 6.43, 3.07 | 0.54 |
| 10     | 1959/o                                          | 0.43, 0.21 | 0.98, 0.85 | 5.05, 5.05 | 0.12 |
| 11     | 3109/o                                          | 0.08, 0.11 | 5.13, 5.13 | 5.13, 5.13 | 0.01 |
| 12     | 1286/o                                          | 0.36, 0.24 | 0.56, 0.55 | 5.42, 2.70 | 0.19 |
| 13     | 55/K                                            | 0.44, 0.24 | 5.27, 5.27 | 5.27, 5.27 | 0.21 |
| 14     | 978/K                                           | 0.69, 0.08 | 5.75, 5.75 | 5.75, 5.75 | 0.49 |
| 15     | K3                                              | 1.10, 0.14 | 5.95, 5.95 | 5.95, 5.95 | 0.02 |
| 16     | K4                                              | 0.12, 0.05 | 6.04, 6.04 | 6.04, 6.04 | 0.07 |
| 17     | 792/K                                           | 1.16, 0.20 | 5.14, 5.14 | 5.14, 5.14 | 0.37 |
| 18     | 393/K                                           | 1.54, 0.19 | 5.17, 2.87 | 5.17, 2.77 | 0.31 |
| 19     | 133/K                                           | 0.10, 0.00 | 0.23, 0.31 | 5.02, 3.22 | 0.02 |
| 20     | 23/K                                            | 0.90, 0.23 | 0.67, 0.55 | 5.90, 5.90 | 0.49 |
| 21     | 887/K                                           | 0.68, 0.34 | 5.62, 5.62 | 5.62, 5.62 | 0.49 |
| 22     | 169/K                                           | 2.06, 0.02 | 2.67, 0.52 | 5.98, 5.98 | 0.07 |
| 23     | 3752/sz                                         | 1.12, 0.14 | 5.85, 5.85 | 5.85, 5.85 | 0.05 |
| 24     | 4908/p                                          | 0.02, 0.37 | 1.96, 1.63 | 4.88, 2.07 | 0.00 |
| 25     | 3907/sz                                         | 0.92, 0.07 | 3.09, 2.37 | 5.29, 3.56 | 0.86 |
| 26     | 15/p I                                          | 1.12, 0.32 | 5.64, 5.64 | 5.64, 5.64 | 0.16 |
| 27     | 149/p                                           | 0.78, 0.44 | 0.46, 0.47 | 5.97, 2.64 | 0.49 |
| 28     | 153/s                                           | 0.20, 0.10 | 0.49, 0.35 | 6.44, 3.38 | 0.20 |
| 29     | 547/p                                           | 0.37, 0.48 | 2.89, 1.64 | 5.81, 4.02 | 0.14 |
| 30     | 1885/K                                          | 0.79, 0.51 | 1.12, 0.83 | 6.05, 2.04 | 0.09 |
| 31     | 556/K                                           | 0.46, 0.29 | 0.67, 0.27 | 5.41, 1.93 | 0.25 |
| 32     | 1651/p                                          | 0.74, 0.22 | 0.46, 0.28 | 5.46, 2.42 | 0.03 |
| 33     | 1304/s                                          | 1.25, 0.33 | 5.70, 5.70 | 5.70, 5.70 | 0.65 |
| 34     | 1804/p                                          | 1.08, 0.01 | 5.26, 2.60 | 5.26, 2.37 | 0.25 |
| 35     | 2284/p                                          | 0.87, 0.41 | 1.04, 0.97 | 5.38, 2.14 | 0.09 |
| 36     | 143/p                                           | 0.93, 0.50 | 2.95, 2.27 | 5.83, 3.07 | 0.61 |

<sup>a</sup>The values were calculated by subtracting log<sub>10</sub> CFU/ml of treated samples from those of untreated controls (0 J/cm<sup>2</sup>; 0 µM RB, 0 µM PEX). The following concentration of compounds was used: 5 µM RB, 5 µM PEX, fluence 30 J/cm<sup>2</sup>. The initial number of cells was ∼10<sup>6</sup> CFU/ml. The presented values are mean of three independent experiments.

<sup>b</sup>MDR (multidrug resistant), XDR (extended drug resistant), see Section “Materials and Methods” for classification details.

While the process of PS accumulation is a light-independent phenomenon, i.e., no light is necessary for the process to occur, aPDI is from the definition a light-dependent process. The significant reduction in the number of viable bacterial cells was observed in the presence of light. This further means that the observed increased killing was related to a combination of aPDI (RB+light) with AMPs, rather than the interaction of the RB itself with AMP, at least for the particular concentrations tested. On the other hand, we could observe that application of RB together with PEX in dark resulted in enhanced bactericidal activity for about 30% of isolates. This observation points that RB itself facilitates PEX action, perhaps as a result of direct interaction. Further research is needed to explore the nature of the interaction between AMPs and RB.

The concept of improving the uptake of a PS in bacterial cells due to the action of the AMP, or more broadly cationic peptide, has been studied and described. Within the last few years, the results on the activity of porphyrin or xanthenes combined with cationic peptides against the model Gram-positive (*S. aureus*) and Gram-negative (mostly *Escherichia coli*, *E. coli*).
P. aeruginosa) species have been described (Bourre et al., 2010; Yang et al., 2011; Liu et al., 2012; Dosselli et al., 2013, 2014; Johnson et al., 2013; Costley et al., 2017; Le Guern et al., 2017, 2018). In all the referenced work, however, the authors presented the construction of conjugates composed of a PS and AMP rather than mixing separate components. The authors applied similar fluences as these presented in the current study, i.e., from 10 to 30 J/cm², and concentrations of 1 µM–1 mM to obtain at least 3–4 log₁₀ reduction in P. aeruginosa cell survival. Only in a single study the same photosensitizer as in the presented study, namely RB, was used. RB-(KLAKLAK)₂ conjugate activity was presented against P. aeruginosa, indicating 2.5 log₁₀ reduction upon 10 µM concentration (Costley et al., 2017). The same peptide was conjugated to, structurally similar to RB, eosin producing finally a conjugate that at 10 µM concentration resulted in >5 log₁₀ reduction in cell survival. However, in this case much higher fluence of 250 J/cm² was applied (Johnson et al., 2013). In our study, low light doses and low RB and AMPs concentrations resulted in the efficient killing of P. aeruginosa without conjugation of the two elements. In the only study concerning the evaluation of mixtures of PS (MB, chlorin e6, curcumin) and AMP (aurein 1.2), the applied combinations were not as much effective toward Gram-negative species (A. baumannii, E. coli). In contrast, our combinations were effective against Gram-negative species, and even toward diverse clinical isolates.

For any antimicrobial compound, no appreciable toxicity toward mammalian cells should be noted. For the mentioned eosin-(KLAKLAK)₂ 50% toxicity toward HaCaT cells was noticed (Johnson et al., 2013). This was in contrast to our results, where only limited up to 20% cytotoxicity was observed. Moreover, in our presented work, we showed for the first time that the growth dynamics of HaCaT cells after photodynamic treatment and/or combined treatment was not significantly disturbed. Our studies were the first to show, in contrast to the so far applied single time-point measurements, the dynamics of cell proliferation for several days. We could also observe a good agreement of real-time monitoring of growth dynamics with commonly applied MTT assay. In some cases, however, obtaining sufficient range of "therapeutic window," i.e., substantial reduction of bacterial cell number, while sparing host cells could not be obtained at all, although the obtained conjugates were very potent with respect to their antimicrobial properties (Le Guern et al., 2017). Summarizing, based on the results obtained by combining aPDI with AMPs a viable approach to obtain efficient killing of bacterial cells, and preserve no toxicity against human cells. Moreover, we believe that in particular cases, including the example presented within our research, simple mixing of PS and AMP is sufficient to obtain substantial antimicrobial activity, without the need of laborious conjugation process. At the same time, we have observed that RB+light treatment was hemolytic in our experimental in vitro conditions. We could see 55% hemolysis, which was a statistically significant value. The question remains about biological meaning of this observation in terms of the potential application of aPDI in clinics. From the definition, aPDI is a localized process, and the photosensitizer is applied topically, thus it actually has little impact on blood components degradation. Interestingly, there are several excellent works published by Kochevar group on the application of RB with green light to assess in vivo (Dutch belted rabbits model) the potential damage of retina and iris, and the treatment was assessed safe, rapid and effective (Zhu et al., 2016; Gallego-Munoz et al., 2017). Retina from RB+light-treated eyes seemed normal and choriocapillaris contained intact erythrocytes (Zhu et al., 2016).

All the so far research concerning the discussed subject were performed on a very limited number of isolates, whereas it is evident that due to the huge genotypic and phenotypic variation of bacterial isolates, evaluating the results on many strains is of real value and actual necessity. Here, we evaluated 35 clinical isolates, which originate from various sources to verify if the combined protocol of aPDI and PEX treatment can be applied to a wide repertoire of strains. In our experimental conditions all strains could be eradicated, i.e., the survival of bacterial cells dropped below the limit of detection. To evaluate the robustness of our protocol against clinical strains, we applied conditions optimized for reference P. aeruginosa 10145 strain. This particular strain was vulnerable to the action of PEX and
for the combination of PEX and RB in the dark, and upon light activation for 5 \( \mu \)M PEX and 5 \( \mu \)M RB combination. Based on the results obtained for reference strain, we employed the same concentrations and fluence (30 J/cm\(^2\)) for the analysis of all the clinical isolates. As it appeared for almost half of the analyzed strains (47%) mere presence of PEX was sufficient to effectively reduce the number of bacterial cells >5 \( \log_{10} \) units. This means that PEX MBC values for these strains were much lower as compared to the reference 10145 strain. As a result, the concentration of PEX used for these experiments could be further reduced.

As aPDI practically from the definition is a localized treatment the most interesting strains are those isolated from local infections, e.g., wound, ear or bronchi. From the clinical point of view, the most important parameter of bacterial characterization is antibiotic resistance profile. In the previous work published by our group and others, it was shown that aPDI is equally effective toward antibiotic resistant and antibiotic susceptible strains (Maisch et al., 2014; Makdoumi and Backman, 2016). Here, the same applies to the combined treatment of aPDI and AMPs. Even strains that were considered extended drug-resistant strains (Maisch et al., 2014; Makdoumi and Backman, 2016). This means that PEX MBC values for these strains were much lower as compared to the reference 10145 strain. As a result, the concentration of PEX used for these experiments could be further reduced.

We chose RB as PS for our studies due to its photophysical properties, like the high production of singlet oxygen. Production of singlet oxygen in antimicrobial photodynamic therapy is highly appreciated as there are no effective mechanisms that could protect bacterial cells from the toxic singlet oxygen action. Moreover, RB is commonly used in various medical applications. This pink xanthene dye is used for the diagnosis of damage in corneal (Doughty, 2013), brucellosis (Ducrotoy and Bardosh, 2017). Rose Bengal in combination with the green light has been extensively studied with respect to photochemical tissue bonding and corneal transplants (Gu et al., 2011; Gallego-Munoz et al., 2017). This compound was also studied in anti-tumor immune responses in melanoma models (Liu et al., 2018). In terms of aPDI, RB was underappreciated due to its poor action against Gram-negative species. However, our results as well recent from other groups (Wen et al., 2017; Li et al., 2018) proved that RB is efficient photosensitizer also toward Gram-negative bacteria.

Methylene blue was more efficiently accumulated in Enterococcus faecalis cells upon the presence of the peptide aurein 1.2, thus delivering a higher local concentration of ROS produced upon light treatment. In the case of chlorin Ce6, however, direct interaction with AMP was postulated through hydrophobic interactions, which resulted in the prolonged half-life of the PS, thus providing an efficient enhancement of aPDI (de Freitas et al., 2018). At this stage, we are not able to identify the exact mechanism of enhanced bactericidal action of RB-based aPDI combined with PEX or CAM. We can only speculate about possible explanations, which can be ascribed to the improved action of either AMP or PS or interactions between the two, which might influence the structure or the function of each molecule. In our studied combination of RB+PEX/CAM the observed enhanced antimicrobial effect might be related to RB and AMP direct interaction, as we observed efficient killing of bacterial cells, not only in the presence of light but also in its absence (Table 2). RB seems to interact with PEX/CAM directly. The analysis of UV-Vis absorption spectra of RB alone and in combination with AMPs (less intense, red-shifted maximum absorption peak) might confirm such hypothesis (Supplementary Figure S1).

The overall picture seems to be more complex as a yet higher increase in antimicrobial efficacy of RB combined with AMPs is observed upon light irradiation. So overall the hypothesis explaining the enhanced action of aPDI and AMP studied by us can be proposed. Based on this hypothesis direct interaction of RB and AMPs occurs with no light share, which results in (i) higher accumulation of RB in bacterial cells, (ii) change of function and/or structure of PEX into a more potent membrane-disorganizing agent. In addition, upon light treatment, production of ROS occurs, which further increases the observed bactericidal effect.

When new approaches are introduced as potential medical applications, toxicity and biocompatibility issues should be investigated. In our experimental conditions, AMPs were not toxic at the concentrations tested (up to 10 \( \mu \)M). The combined action of AMP and aPDI showed, however, some toxicity (50% in hemolysis assay, 20% MTT assay). Such toxicity is acceptable, particularly if the treatment has local character. At the same time, the observation of HaCaT cell growth for a longer time revealed that cells after treatment took up growth and proliferation occurred. The dynamics was retarded in relation to untreated cells, but without any cytostatic effect observed. This indicates that the studied treatment has the potential to be exploited as a rational clinical application. More studies on complex in vivo models are required to eventually verify this hypothesis.

**CONCLUSION**

We proved that combination of RB-based antimicrobial photodynamic (aPDI) inactivation and AMPs is a viable alternative to classical antimicrobials, effective against multidrug-resistant and extended drug-resistant *P. aeruginosa*. This is the first study showing the significant effect of the combined action of aPDI and AMPs toward 35 clinical isolates with diverse antimicrobial resistance, resulting in equally effective photokilling. Specifically, AMP pexigan (PEX) was found to be a very potent molecule with evident selectivity toward Gram-negative bacterial cells. Clever and careful combination of photodynamic inactivation and AMPs opens up new possibilities for combating pathogens that cause localized infections. This proposed approach could help to design strategies that alleviate the increasingly common problem of antimicrobial resistance spreading.

**AUTHOR CONTRIBUTIONS**

JN conceived the study, analyzed the results and wrote the first draft of the manuscript, and performed some experiments.
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SUPPLEMENTARY MATERIAL

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FIGURE S1 | UV-Vis spectra of RB alone and RB in the presence of antimicrobial peptides CAMEL and pexiganan. Light absorption spectra were of analyzed compounds at concentration of 2 µM were measured in a wavelength range of 450–650 nm with 0.5 nm intervals, in quartz cuvettes (1 cm light path) containing appropriate solutions in 1 mL PBS, pH 6.8, using Beckman’s DU 650 spectrophotometer at room temperature (25°C).

TABLE S1 | Numerical values of HaCaT cell viability assay (MTT assay) results are presented in Figure 6.

absorbance spectra. Dr. Anna Woziwodzka (IFB UG&MUG) was acknowledged for statistical expertise. The study sponsor had no involvement in the study design, collection, analysis, and interpretation of data.
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.