Photodynamic Therapy as a New Treatment for Chronic Rhinosinusitis - A Systematic Review

Anika Kaura¹, Rishi Shukla², Abigail Lamyman², Robert Almeyda³, Mark Draper⁴, Pablo Martinez-Devesa⁵, Ali Qureishi⁶

¹Ear Institute, University College London, London, UK
²Department of ENT Surgery, John Radcliffe Hospital, Headley Way, Headington, Oxford, UK
³Department of ENT Surgery, Royal Berkshire Hospital, Craven Road, Reading, UK
⁴Department of ENT Surgery, Milton Keynes University Hospital, Standing Way, Milton Keynes, UK

Abstract

This review examines the latest evidence for photodynamic therapy (PDT) in treating chronic rhinosinusitis. MedLine, EMBASE and TRIP Database searches were conducted using the terms: “photodynamic” or “phototherapy” or “photo” and “sinusitis” or “rhinosinusitis,” date range January 2000 to May 2020. A total of 192 records were initially identified, after duplicates and exclusions, 9 full papers and 3 abstracts were included. All study types including in-vitro, animal and human studies were evaluated. Whilst there is in-vitro evidence for the efficacy of PDT’s bactericidal effect on drug resistant bacteria and biofilm viability, there are few clinical studies. PDT is a promising area of research, but larger, focused studies looking at the safety, delivery, efficacy, and patient selection are required before it can be considered a viable treatment for CRS.

Keywords: Phototherapy, photodynamic therapy, paranasal sinus diseases, sinusitis, alternative therapies

Introduction

Chronic rhinosinusitis has a significant impact on patient quality of life and productivity, affecting 5-12% of the general population (1). It is classified as chronic when the core symptoms persist beyond 12 weeks; is difficult to treat or is recalcitrant when symptoms persist despite appropriate medical and surgical treatment. Recent work has focused on defining CRS as either primary or secondary based on the type of the inflammatory disease or endotype. The overall aim is to allow for tailored treatment regimens focusing either on a primarily infective or an inflammatory (type 2) cause. The previous classification of CRS with nasal polyps (CRSwNP) and without nasal polyps (CRSsNP) is no longer favored, as polyps can present in both CRS patients with and without polyps (3). The classification for primary and secondary CRS is summarized in Figure 1 which has been adapted from Grayson et al. (2). The pathophysiology of CRS is complex; we must consider the various possible predisposing and causative factors.

Allergy and Asthma

Allergy and asthma in the context of CRS have been extensively studied. The presence of inhalant allergy has been found to be significantly higher in both CRS patients with and without polyps (3). The prevalence of allergy appears to vary with phenotype; central compartment atopic disease and allergic fungal rhinosinusitis have stronger associations than CRS with and without polyp groups. There is a strong association between asthma and CRS, especially in those with both CRS and allergic rhinitis (4). A United Kingdom study showed the prevalence of asthma to be 44.9% in CRSwNP, 21.2% in CRSsNP, compared to the 9.95% in controls (3).

Environmental Irritants

It is hypothesized that the pollutants can affect ciliary function including mucociliary clearance (5), predisposing to repeated infection and inflammation. It has been demonstrated that air pollutants are correlated with CRS symptom severity, with CRSsNP being the most affected cohort (6). Studies have shown that occupational exposure to dust from metals, textiles and paper are associated with CRS (6-8).

Fungi

Like bacteria, fungi can form biofilms and secrete toxins, but their role in CRS pathogenesis is yet to be fully elucidated. The role of fungi, like Aspergillus, in subtypes where fungal balls occur and in allergic fungal rhinosinusitis is clearer (9). Ex-vivo studies have shown that Aspergillus niger stimulation resulted in increased pro-inflammatory cytokines like IL-6 (10). Shin et al. (11) demonstrated that CRS patients showed exaggerated humoral
and cellular responses to common airborne fungi, particularly Alternaria, and that this could explain the chronicity of inflammation seen in CRS.

**Virus**  
It is hypothesized that viral infection could initiate or exacerbate CRS; coronavirus was the most commonly found virus in patients with CRS (12). In a study from Iran with 76 CRS patients undergoing endoscopic sinus surgery, 33% were found to have at least one of rhinovirus and respiratory syncytial virus (13). Ex-vivo studies have demonstrated that rhinovirus infection can be associated with exacerbation of CRS, including increased susceptibility to secondary microbial infection and impairment of mucociliary clearance (14, 15).

**Bacteria and Biofilms**  
Part of the pathophysiology of CRS is thought to be secondary to bacteria like *Staphylococcus aureus*, where colonization results in the disruption of the normal mucosal barrier and the promotion of immune dysregulation, further amplified by antibiotic drug resistance and biofilm formation (16, 17). In addition, the overgrowth of “bad” bacteria displaces those bacteria that are considered part of a healthy microbiome, resulting in microbial dysbiosis (16).

Patients with CRS have been found to lack normal sinus mucociliary defense mechanisms and to be colonized by multi-drug resistant bacteria that form biofilms within sinus cavities and continue to cause chronic inflammation (18-20). Methicillin-resistant *S. aureus* (MRSA) and multidrug resistant *Pseudomonas aeruginosa* are found in the clinical isolates of CRS patients and are a cause of antibiotic treatment failures (18, 20). Traditional treatment with steroids, oral antibiotics, nasal douches, and sinus surgery are less effective in this population. For these patients and those with other types of CRS, alternative therapies must be evaluated, as supported by the recommendations from EPOS 2020 (21).

A relatively new area of interest involves the use of light therapies. Light therapies include laser, near infra-red illumination, ultra-violet (UV) light therapy and photodynamic therapy (PDT). To date, PDT has been most effective in the treatment of cancer. It is often used to treat actinic keratosis and basal cell carcinoma. It has also been shown to be effective in early stage squamous cell carcinoma of the oropharynx, nasopharynx and larynx, although a clinical role for PDT in this context is yet to be defined (22).

In animal studies, PDT has proven effective in treating chronic wound infections by suppressing bacterial growth (23) and cellular responses to common airborne fungi, particularly Alternaria, and that this could explain the chronicity of inflammation seen in CRS.

**Main Points**  
- Photodynamic therapy involves applying a photoactive agent to a surface and then irradiating this area with light. It has been shown to have anti-inflammatory, anti-bacterial and anti-neoplastic effects.
- Photodynamic therapy aims to reduce the bacterial load causing chronic infection and immune dysregulation in chronic rhinosinusitis.
- Photodynamic therapy has been shown to be highly effective in-vitro and in-vivo against bacteria implicated in chronic rhinosinusitis.
- The clinical studies to date have shown promising results through improvements in both objective and subjective outcome measures, with no unacceptable adverse effects.
- Randomised control trials are required to fully assess the short and long term efficacy of this treatment modality.

**PDT** uses a photosensitizing agent solution to prime the surface area covered by biofilm, then light at a specific wavelength is administered to activate this solution. This triggers the formation of reactive oxygen species (ROS) which have anti-bacterial, anti-inflammatory and anti-neoplastic downstream activity (24). The ROS damage cell walls, allow translocation of further activated solution and damage inner organelles, resulting in apoptosis (25). This cell death mechanism is an entirely different pathway to that of antimicrobials and might offer an alternative option for combatting multi-drug resistant organisms (26, 27). Commonly used photosensitizers include ultra-methylene blue (selectively binds to microbial cell walls and biofilms) and aminolevulinic acid variants. In addition, in-vitro studies have shown that unlike antibiotics, bacteria do not develop resistance to repeated photodynamic therapy treatments (28). There is also much evidence that PDT can disrupt biofilm by further reducing bacteria viability and increase sensitivity to antibiotics (29).

There are numerous studies to support the bactericidal effect of photodynamic therapy on drug resistant and biofilm forming bacteria (30-37). Whilst this technique may have a role in treating CRS, the research in this area is only just evolving and there is a need to evaluate its potential in treating CRS. This review will focus on the studies that are relevant to PDT and the treatment of CRS.

At present, approved devices include the Sinuwave photodisinfection system licensed in Canada, and a study using this device in human studies has been conducted by Desrosiers et al. (38). Rhinolight is a device that emits UV and visible light and is predominantly being used in the treatment of allergic rhinitis in centers across Hungary and Germany (without the application of a photoactive agent). Its evidence base centers on allergic rhinitis and the ex-vivo study discussed here uses this device in conjunction with a photoactive agent (39, 40). The other studies that have been evaluated in this review use light emitting devices in lab-based settings, not commercially produced or licensed for use in humans.

This review article examines the current evidence base for use of photodynamic therapy in the treatment of chronic rhinosinusitis.

**Materials and Methods**  
MedLine, EMBASE and TRIP database searches were conducted in May 2020 using the following terms: “photodynamic” or “phototherapy” or “photo” and “sinusitis” or “rhinosinusitis”, date range January 2000 to May 2020.
Inclusion criteria:
- Original scientific contributions
- Studies that employ photodynamic therapy (agent that acts as a photosensitizer and irradiation with light which could be in the form of a laser, LED, etc.)
- Use bacterial isolates from the human sinus or CRS patients for in-vitro studies
- Must investigate the use of PDT in the context of CRS or bacteria that cause CRS

Exclusion criteria:
- Studies that use light therapy alone without a photosensitizer
- Studies that use planktonic or other research strains of bacteria (they may be referenced to provide supporting evidence but are not included in the main table of papers)
- Insufficient information/conference abstracts (Three conference abstracts, all by Desrosiers et al. (38, 41, 42), have been included as they provide preliminary information on the results in human studies)
- Reviews, books etc. that refer to other studies
- Studies that are not directly looking at CRS, but investigate other conditions like allergic rhinitis or acute rhinosinusitis

AK and RS independently assessed the title and abstracts of the identified articles to determine relevance. Any disagreement was resolved by discussion with senior authors. Database searching identified 192 records that met the search terms (Figure 2). After duplicates were removed, 161 abstracts were screened, of which 106 were then excluded. Fifty-five full text articles were assessed for eligibility against the inclusion and exclusion criteria. After 43 articles were excluded, nine full papers and three abstracts were selected to be included in the review. Some of the full text articles that were excluded (n=43) have been referred to in the paper where they support the evidence of the included articles, for example mechanisms of delivery of PDT.

Regarding in-vitro studies, the papers included here are all relevant to investigating the effect of photodynamic therapy on biofilm forming bacteria, although not all use strains of bacteria from CRS patients or from the sinus cavity. The studies that use other isolates have not been included in the main table of results. According to the American Society of Microbiology (2010), a new approach has to prove an efficacy of 3 log reduction of colony forming units (CFU) before being able to use the term “antibacterial” (43).

Results
The studies included six in-vitro, one ex-vivo, one animal case report, three human case series and one randomized control trial (RCT), these are summarized in Table 1.

Pre-Clinical Studies (Laboratory)

In-vitro Studies
Preliminary studies by Zhao et al. (44) demonstrated antimicrobial properties of 5-aminolevulinic acid (ALA) mediated PDT on biofilm forming strains of S.aureus and S.epidermidis. The application of PDT appreciably reduced bacterial growth of S.aureus and S.epidermidis isolated from CRS patients. This was measured as a significant reduction in bacterial growth (measured as log reduction in colony forming units) when compared with control groups of the same planktonic strains of bacteria. In the biofilm S.aureus experiment, the mean log colony forming units (logCFU) was $8.68 \pm 0.05$ (control group), $6.90 \pm 0.96$ (experiment group) ($t=3.68, p<0.05$); and in biofilm S.epidermidis experiment the data was $8.67 \pm 0.05$ (control group), $7.29 \pm 0.61$ (experiment group, $t=5.07, p<0.01$). They present clear methodology for how biofilm was cultured and how conditions were controlled, including how the optimal photosensitizer concentration and light intensity were found. The CRS patients included here had received medical therapy prior to endoscopic sinus surgery (ESS), secretions from the middle meatus were sampled in order to obtain S.aureus and S.epidermidis strains for this in-vitro experiment. No further details regarding the type and length of medical treatment received by this cohort of CRS patients were included in the manuscript.

This study is supported by research undertaken by Biel et al. (45) using drug resistant strains of P.aeruginosa and MRSA. They demonstrated that PDT reduced the CRS polymicrobial biofilm by $>99.9\%$ after a single treatment. For the $300\mu g/mL$ concentration of the photoactive agent methylene blue (MB) there was a 6.5 log reduction of antibiotic-resistant multi-species bacterial biofilms after a single PDT treatment. When they used a higher MB concentration and lower light parameters, they achieved greater than seven logs of bacteria kill using two PDT light treatments. They showed clear methodology for each of the experimental groups and used an objective automated tool for counting colonies; however, the authors do not offer limitations or a critical appraisal of the study.

More recently, this group created an anatomically correct maxillary sinus model in order to conduct the same experiment using MB and 670 nm non-thermal activating light (46). Again, they showed a 99.9% reduction in biofilm for both P.aeruginosa and MRSA strains after one treatment as measured by log reduction in CFU. The model was created from human CT scans, and dimensions were taken as the average of 10 random male and female maxillary sinuses. The mixed species biofilm inoculum was pipetted into sterile silicone models and then allowed to shake for 24 hours. It is not clear how these results using silicone models and artificially grown biofilm can be extrapolated to how biofilm forms on human ciliated respiratory mucosa of the maxillary sinus, and the authors have recognized this as a limitation. The have shown methodological rigor by using 11 different treatment combinations to assess the effects of different concentrations of ethylenediaminetetraacetic acid (EDTA), EtOH and methylene blue. They were able to show that low concentration of EDTA added to MB results in improved PDT efficacy of killing biofilm forming bacteria.

A further study by Zhang et al. (47) in 2017 investigated the effect of ALA mediated PDT on S.aureus biofilm, and the effect
<table>
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<tr>
<th>Title</th>
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<th>Cohort</th>
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<tr>
<td>In-vitro study of photodynamic therapy of antibiotic-resistant</td>
<td>Zhao et al. (44)</td>
<td>45 patients treated medically for CRS requiring FESS, mucus taken from middle meatus. 13 S.aureus and 16 S.epidermidis strains were identified. 5-aminolevulinic acid mediated PDT was applied to these strains as well as Planktonic strains.</td>
<td>Level 5 In-vitro</td>
<td>Mean IgCFU for experiment and control groups of planktonic and patient S.aureus and S. epidermidis strains.</td>
<td>In the biofilm S.aureus experiment, the mean IgCFU was 8.68±0.05 (control group), 6.90±0.96 (experiment group) (t=3.68, P&lt;0.05); and in biofilm S.epidermidis experiment the data was 8.67±0.05 (control group), 7.29±0.61 (experiment group, t=5.07, P&lt;0.01).</td>
<td>In-vitro study, only 2 major biofilm forming strains of bacteria used. Single outcome measure (IgCFU), clinical applicability requires evaluation.</td>
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<td>Antimicrobial photodynamic therapy treatment of chronic recurrent</td>
<td>Biel et al. (45)</td>
<td>Antibiotic resistant planktonic bacteria and fungi, and polymicrobial biofilms of Pseudomonas aeruginosa and MRSA were grown on silastic sheets and treated with a methylene blue photosensitizer and 670nm non-thermal activating light. Cultures of the planktonic microorganisms and biofilms were obtained before and after light treatment to determine efficacy of planktonic bacteria and biofilm reduction.</td>
<td>Level 5 In-vitro</td>
<td>Kill rate as measured by log CFU reduction in bacteria.</td>
<td>The CRS planktonic microorganism and biofilm study demonstrated that aPDT reduced the CRS polymicrobial biofilm by &gt;99.9% after a single treatment. For the 300 μg/mL MB concentration there was 6.5 log reduction of antibiotic-resistant multi-species bacterial biofilms after a single PDT treatment. Using a higher MB concentration (500 μg/mL) and lower light parameters achieved greater than 7 logs of bacteria kill using two PDT light treatments.</td>
<td>Multi-organism biofilm was treated with PDT rather than single organism biofilm.</td>
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<td>sinusitis biofilms</td>
<td>2011 USA</td>
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<td>5-aminolevulinic acid-mediated photodynamic therapy and its</td>
<td>Zhang et al. (47)</td>
<td>Study investigating the effect of 5-aminolevulinic acid with light emitting diode 633nm (ALA mediated PDT) on S.aureus. Biofilm forming MRSA and MSSA strains were isolated from CRS patients undergoing endoscopic sinus surgery. MRSA and MSSA were treated with ALA-PDT, compared with ALA-PDT combined with antibiotics (vancomycin, netilmicin or cefaclor). Control groups included no treatment, ALA alone, and light irradiation alone.</td>
<td>Level 5 In-vitro</td>
<td>Kill rate as measured by log CFU reduction in bacteria.</td>
<td>ALA-PDT was found to significantly inactivate S.aureus biofilm across all 15 strains, mean 5.75log10CFU/ml reduction in viable count; the effects were similar in the MSSA and MRSA groups. When ALA-PDT was combined with antibiotics (vancomycin, netilmicin or cefaclor) the bactericidal effect increased for at least 9 out of the 15 strains. They hypothesize that PDT breaks the sessile structure of biofilm leading to recovery of antibiotic sensitivity, although it is not clear why this happens in a strain dependent way.</td>
<td>The addition of antibiotics seems to have an additive effect to the PDT. Experimental groups were small, the conclusion that the effect in MRSA and MSSA strains is not significantly different may not be repeatable. The authors cannot explain why the additive effect of antibiotics is seen in some strains and not in others.</td>
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<td>strain-dependent combined effect with antibiotics on Staphylococcus</td>
<td>2017 China</td>
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<td>aureus biofilm</td>
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<td>Development and characterization of erythrosine nanoparticles with potential for treating sinusitis using photodynamic therapy</td>
<td>Garapati et al. (48) 2015 USA</td>
<td>Study of erythrosine mediated PDT in S. aureus cells using nanoparticle delivery system. In one arm of the experiment, cells were incubated with free erythrosine drug, and the other arm with erythrosine nanoparticles. The control arm was S. aureus only plus irradiation. Cells in all arms were irradiated at 6 time points with an LED (530nm).</td>
<td>Level 5 In-vitro</td>
<td>Kill rate as measured by log CFU reduction in bacteria. The uptake of erythrosine in S.aureus cells from nanoparticles and pure drug was approximately 14.83±0.15 and 0.60±0.19 g per mg of protein, respectively. This indicates the ability of bacteria cells to internalize erythrosine nanoparticles is better than the free drug. Photodynamic inactivation of erythrosine nanoparticles after 8, 16 and 24 h, was significantly higher compared to pure erythrosine. This could be attributed to the sustained released of erythrosine from nanoparticles.</td>
<td>This nanoparticle mediated method of PDT could potentially mean that reactive agent is only inserted into a sinus cavity once, and then the irradiation can be repeated at several time points to achieve maximum bactericidal effect. Further in-vivo studies are required to test the efficacy and safety of this method.</td>
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<td>The effect of antimicrobial photodynamic therapy on human ciliated respiratory mucosa</td>
<td>Biel et al. (52) 2012 USA</td>
<td>Study of aPDT treatment of EpiAirway™ (in-vitro airway tissue model that originates from normal, human-derived tracheal/bronchial epithelial cells) was performed. Treatment groups included a non-treatment control, laser light alone, photosensitizer alone, and therapeutic photosensitizer and light combination (aPDT).</td>
<td>Level 5 In-vitro</td>
<td>Histomorphological evaluation of the EpiAirway specimens. The EpiAirway™ histologic study demonstrated no histologic alteration of the respiratory cilia or mucosal epithelium in any of the treatment groups.</td>
<td>Unclear how accurately in-vitro airway tissue model reflects normal sinus ciliated respiratory epithelium.</td>
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<td>Photodynamic therapy of antibiotic-resistant biofilms in a maxillary sinus model</td>
<td>Biel et al. (46) 2013 USA</td>
<td>Antibiotic resistant polymicrobial biofilms of P. aeruginosa and MRSA were grown in an anatomically correct novel maxillary sinus model and treated with a methylene blue/EDTA photosensitizer and 670nm non-thermal activating light. Cultures of the biofilms were obtained before and after light treatment to determine efficacy of biofilm reduction.</td>
<td>Level 5 In-vitro</td>
<td>Kill rate was calculated as surviving CFU/ml in experimental conditions versus control (no light and no photosensitizer) and expressed as a log10 reduction from control for each individual organism. PDT reduced the CRS polymicrobial biofilm by &gt;99.99% after a single treatment. The best treatment results in biofilm reduction were achieved with PDT using the photosensitizer 1.25mM EDTA + 5% EtOH + 0.03%MB in the presence of 670nm light resulting in a 5 log10 (99.99%) reduction in P. aeruginosa biofilm and a 3.1 log10 (99.9%) reduction in MRSA biofilm after a single treatment. Low concentrations of EDTA added to MB results in improved PDT efficacy of multispecies biofilm bacterial kill.</td>
<td>In-vitro study using anatomically correct model based on CT scans. As the model is not lined by respiratory epithelium it is unclear as to what degree this reflects human physiology.</td>
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### Table 1. Summary of Studies of Photodynamic Therapy and Chronic Rhinosinusitis (Continue)

<table>
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<tr>
<th>Title</th>
<th>Author Date Country</th>
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<tr>
<td>Ultraviolet light and photodynamic therapy induce apoptosis in nasal polyps</td>
<td>Nemeth et al. (40) 2012 Hungary</td>
<td>Ex-vivo study nasal polyp (NP) tissue was surgically collected from 21 consecutive patients with CRS associated with NP. The removed polyps were cut into pieces and tissue samples were irradiated in-vitro by different doses of combined ultraviolet and visible light (UV/VIS: 280-650nm) and by selective ultraviolet and visible light (sUV/VIS: 295-650nm). PDT was performed by pre-sensitizing tissue samples with 5-delta-aminolevulinic acid (DALA) then irradiated with visible light (VIS: 395-650nm), Tunel assay was applied to detect apoptosis of epithelial and inflammatory cells in irradiated and control nasal polyp tissue samples.</td>
<td>Level 5 Ex-vivo</td>
<td>Apoptosis rate of cells.</td>
<td>UV/VIS light significantly increased epithelial cell and subepithelial leukocyte apoptosis compared to control groups, PDT treatment showed the highest surface epithelial cell apoptosis rate as well as subepithelial leukocyte apoptosis rate compared to all other groups.</td>
<td>This study applies treatment to tissue in a non-physiologic environment and provides results only relating to the CRSwNP cohort.</td>
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<tr>
<td>Temporary regression of locally invasive polypoid rhinosinusitis in a dog after photodynamic therapy</td>
<td>Osaki et al. (51) 2012 Japan</td>
<td>Antivascular photodynamic therapy (PDT) using benzoporphyrin derivative monocoid ring A was applied in one dog with CRS and polyps, 8 months after initial presentation and after failed steroid treatment.</td>
<td>Level 5 Animal case report</td>
<td>CT findings, symptom recurrence.</td>
<td>Short term improvements in symptoms and scan findings up to 11 months post treatment.</td>
<td>In this single case report the subject needed multiple PDT therapies. The improvements appeared to be short lived. Frontal trepanations were used- a method not used by the other studies. Not evident how applicable the results are to human CRS treatment.</td>
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<tr>
<td>Title</td>
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<tr>
<td>Phototherapy for chronic rhinosinusitis</td>
<td>Krespi et al. (53)</td>
<td>A prospective randomized study with 23 symptomatic post-surgical CRS patients with positive cultures was conducted. Two groups (GR1 and GR2) were treated with NILI. GR1 was treated with a 940nm laser, while GR2 was treated with a topical photoactive agent, indocyanine-green, followed with 810nm laser. Saccharin test was performed 1 week following treatment.</td>
<td>Level 4 Case series</td>
<td>Nasal endoscopic scoring (NES), SNOT-20 scores and cultures-positivity / log reduction.</td>
<td>Significant improvement in SNOT scores in both groups. Of the 8 cultures in group 2, post treatment 2 were clear of bacteria and 2 showed significant log reduction. In group 2 there was a 50% reduction in mean NES score. Saccharin transit test for group 2 post treatment was normal in all cases. Two of the 23 patients experienced pain during treatment which subsided after reverting from continuous to pulsed mode.</td>
<td>Small group of test patients. Patients had varying degrees of pre-treatment surgery. NES scoring and saccharin tests were only performed in group 2. Saccharin transit test results suggest no adverse effect on ciliary movement. Little information provided regarding patient characteristics of the two groups, randomization, blinding and follow-up protocol. It appears the experiment was run in the first group which influenced how the second group experiments were conducted.</td>
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<tr>
<td>Sinuwave photodisinfection for the treatment of refractory chronic rhinosinusitis: a case series</td>
<td>Desrosiers et al. (38) 2013 Canada</td>
<td>Twenty-nine sinuses (13 frontal, 6 ethmoid, 10 maxillary) in nine patients with recalcitrant CRS persisting following technically successful FESS have been treated with the Sinuwave™ photodisinfection system.</td>
<td>Level 4 Case series</td>
<td>Short term follow-up has shown no delayed complications and somewhat surprisingly, resolution of disease in several patients.</td>
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<td>Conference abstract- full results not published yet.</td>
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<tr>
<td>Evaluation of the safety of antimicrobial photodynamic therapy (aPDT) for refractory chronic rhinosinusitis</td>
<td>Desrosiers et al. (42) 2016 Canada</td>
<td>Of the 44 trial patients, 31 were randomized to receive aPDT and a total of 43 treatments were delivered to 154 sinuses (52 frontal, 48 maxillary, 54 ethmoid).</td>
<td>Level 4 Case series-safety study</td>
<td>Pre and post treatment endoscopic visualization, CT imaging, ophthalmologic evaluation, olfactory testing.</td>
<td>aPDT of the paranasal sinuses can be safely performed in post endoscopic sinus surgery sinus cavities. No instances of ocular dysfunction or visual loss occurred. There was no trauma at the level of the surrounding sinus mucosa, and in several patients, there was resolution of disease.</td>
<td>Conference abstract- full results not published yet.</td>
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when combined with antibiotics. Ten methicillin sensitive *S. aureus* (MSSA) and 5 MRSA biofilm forming strains were isolated from CRS (with and without polyps) patients undergoing endoscopic surgery. The authors do not detail the clinical history or the previous treatments of these patients who were sampled.

ALA-PDT was found to significantly inactivate *S. aureus* biofilm across all 15 strains, mean 5.75log<sub>10</sub>CFU/mL reduction in viable count; the effects were similar in the MSSA and MRSA groups. Through the live/dead staining they were able to show that in the ALA-PDT group, the dead cells were predominantly distributed in the upper layer of the biofilm, this could be due to lower concentration of photosensitizer in the inner layer or inability of light to penetrate these regions. When ALA-PDT was combined with antibiotics (vancomycin, netilmicin or cefaclor) the bactericidal effect increased for at least 9 out of the 15 strains. They hypothesize that PDT can break the sessile structure of biofilm resulting in increased antibiotic sensitivity, the authors cannot explain why the added effect with antibiotics is seen in some strains and not others. Regarding the similar effect of ALA-PDT in MSSA and MRS biofilms, the authors recognize that small experimental groups were used and therefore these results may not be generalizable, and further studies are required.

Garapati et al. (48) investigated the role of erythrosine nanoparticles in PDT. They offer detailed and thorough methodology regarding the development of their erythrosine-loaded PLGA (biodegradable polymer) nanoparticle delivery system. They demonstrated that by using nanoparticles to deliver erythrosine inside MRSA cells (from human sinus), there was significantly better uptake of erythrosine than using the free drug (14.83 micrograms per mg of protein compared with 0.6). After 1 hour of incubation, the uptake of erythrosine was ~25 times higher in the presence of erythrosine nanoparticles compared to the free drug, MRSA cells with free erythrosine drug (group 1) and those with erythrosine nanoparticles (group 2) were irradiated at 6 time points with an LED (530 nm). The control groups were MRSA alone, and MRSA plus LED irradiation. There was no effect on MRSA viability in the control groups. At 0.5- and 2-hour time points, groups 1 and 2 produced similar rates of MRSA inactivation. At 8, 16 and 24 hours, photodynamic inactivation of MRSA was significantly higher for group 2. At 16 hours in group 1 there was 5-fold reduction in mean log10CFU/mL compared with complete loss of viability in group 2. They found erythrosine nanoparticles (and irradiation) were highly effective in killing MRSA cells and this could be attributed to the sustained release of erythrosine from nanoparticles. The paper deduces that the nanoparticle system of delivering photoactive reagent into the cells will have more effective bactericidal activity compared with neat reagent. They propose that erythrosine nanoparticles could be delivered into a sinus cavity by using a powder insufflation technique, and that potentially a patient could undergo repeat light therapy without the need for further delivery of the erythrosine, as it is being released continuously from the nanoparticles. They acknowledge that whilst they have shown success in the in-vitro setting, in-vivo studies are required to evaluate the safety and efficacy of the erythrosine nanoparticle delivery system.

**Other In-vitro Studies (Supporting Evidence)**

A more recent study by Parasuraman et al. (49), supports the hypothesis that PDT is more effective when the photosensitizer is delivered into bacterial cells using nanoparticles. They used toluidine
blue (TB) encapsulated in mesoporous silica nanoparticles and a red diode laser (670 nm), on *P. aeruginosa* and *S. aureus* (not isolated from the sinus). The study demonstrated significant benefit of using TB in nanoparticles compared with TB alone in terms of reactive oxygen species production, cell inactivation, cell viability and importantly biofilm formation. This is a comprehensive and well-designed study that looked at many outcomes to determine efficacy; these included detection of reactive oxygen species, cell viability, extracellular polymeric substances quantification, protein leakage, lipid peroxidation, biofilm inhibition and anti-biofilm efficiency (live and dead cells) using confocal laser scanning microscopy.

Gandara et al. (50) investigated the effect of toluidine blue (TB) and photodynamic inactivation using 635 nm laser, and the additive effects of using near-infrared treatment (980 nm laser) and proteinase K treatment. Applying TB as the photosensitizer to *S. aureus* biofilm (research strain), and then applying consecutive treatment with 980 nm and 635 nm lasers produced the largest reduction in biofilm viability (4.5-log viable count decrease), which was significantly more than the effect of TB alone or TB with only one of the lasers. This group proposed that enzymatic digestion of biofilm components using proteinase K could enhance the effect of PDT. When biofilm was treated with proteinase K before TB-PDT there was increased reduction in bacteria CFU counts compared with TB-PDT alone (4.3 vs. 5.46 logs CFU/mL, respectively). This study highlights the additive effect of multiple laser treatment and enzyme treatment in further reducing biofilm viability in an in-vitro setting. The authors...
recognize that the therapeutic window for near infrared diode lasers is narrow to avoid thermal damage to the host tissue, and this must be further investigated with regards application on human respiratory mucosa that is both safe and with high bactericidal efficacy. They highlight that in-vitro studies investigating the effects of PDT use wide ranging concentrations of photosensitizer and selected wavelengths of light and there it is difficult to compare efficiencies between various studies to obtain mechanistic conclusions.

**Ex-vivo Study**

Nemeth et al. (40) in 2012 performed an ex-vivo study on polyps taken from 21 consecutive CRS patients undergoing ESS. Patients were excluded if they used certain medication in the four weeks prior, including corticosteroids and leukotrienes. However, no details regarding the clinical history or prior medical or surgical treatment are described. Combinations of ultraviolet and visible light or photodynamic therapy using 5-deat-aminolevulenic acid were applied to the tissue samples using the Rhinolight device. The phototherapy group showed the highest rate of apoptosis of surface epithelial cells (80%) and subepithelial inflammatory cells (70%).

A major limitation of this study is that it is ex-vivo, where harvested polyps are experimented on, it is therefore not clear whether these results reflect in any way what occurs in-vivo in CRS patients with polyps and what the safety profile of the photosensitizing agent or the light irradiation would be. The au-
Animal Case Report  
A Japanese group have trialed the use of PDT in a dog with CRS and nasal polyps. Osaki et al. (51) describe the case of a dog with disease affecting the nasal cavity and frontal sinuses, which had failed steroid therapy. In this case, antivascular photodynamic therapy (PDT) using benzoporphyrin derivative monoacid ring A (BPD-MA) was administered intranasally via intravenous catheters fitted with cylindrical diffusers, and fibers with microlens inserted through small trephinations of the skin. After 15 minutes of instilling the photosensitizing agent, 690 nm laser light emitted by diode laser was applied via the fibers. After the first treatment there was resolution of symptoms and improvement of computerized tomography (CT) scan findings, however three months later symptoms recurred, and the dog was given a further course of treatment. The subject received four treatments over the space of 11 months. Whilst this case report showed improvement in clinical signs and symptoms after each PDT treatment using BPD-MA, these were clearly short lived. In terms of side-effects, post procedural facial swelling was reported after each treatment, lasting a few days each time and did not require any treatment. This study describes a new technique whereby PDT can be used to treat frontal sinus disease through percutaneous trephinations; although clinically it will be more pragmatic to deliver PDT endonasally via a frontal sinusotomy procedure. This mechanism for treatment would require careful evaluation in human subjects and it is yet unclear if any of the results or analysis would be applicable to the management of CRS in humans.

Safety Studies  
The safety of photodynamic therapy has been evaluated by Biel et al. (52) on an in-vitro tissue airway model. They tested PDT using methylene blue and Sinuwave technology on EpiAirway™; this histologic study demonstrated no histologic alteration of the respiratory cilia or mucosal epithelium in any of the treatment groups. It is not evident how representative EpiAirway™ is of in-vivo respiratory epithelium, so it remains unclear how much these results can be extrapolated.

Desrosiers et al. (41) evaluated the safety of using PDT in human subjects. A conference abstract only gives a summary of the results (full study has not been published). Forty-three PDT treatments were delivered to 154 sinuses. Outcomes included pre and post treatment endoscopic visualization, CT imaging, ophthalmologic evaluation, and olfactory testing using the University of Pennsylvania Smell Identification Test (UPSIT). There were no episodes of ocular dysfunction or mucosal damage. The most frequently reported side effect was transient mild pressure over the treated sinus.

Clinical Studies  
In a study by Krespi et al. (53), 23 patients were randomized to receiving laser (940 nm) alone (group 1) or laser (810 nm) and topical photosensitizing agent indocyanine-green (group 2). Post treatment saccharin transit tests for both groups were normal, suggesting no adverse effect on ciliary movement. In the PDT group, Sino-Nasal Outcome Test-20 (SNOT-20) scores dropped by 41% (p=0.0003) and nasal endoscopic scores (NES, based on severity of inflammation, ostial patency and crusting) halved (p=0.0005). In group 1, two of 13 patients were culture negative post treatment, and in group 2, two of 10 patients were culture negative. For the patients where cultures remained positive, there was no detail regarding the log reduction in bacterial growth, and the authors recognized that not measuring bacterial growth qualitatively was a limitation.

There were no serious adverse effects, some experienced minor discomfort during laser illumination, and two patients felt pain associated with heat that subsided when changing from a continuous to a pulsed method of light therapy.

The RoB2 Cochrane tool was used to assess risk of bias in this clinical study. A major limitation was they did not include a control arm. Patients with persistent CRS symptoms, with and without polyps, with at least one prior ESS surgery were recruited; the overall sample size was small. The paper does not offer further details regarding the clinical or surgical history of these patients, and therefore it is unclear whether the two groups were similarly matched in this regard. Patients were assigned to each group by consecutive recruitment randomly to one arm until the group was complete (n=13), then to the other arm. The severity of patient symptoms and endoscopic findings, along with response to treatment guided the number of treatments administered for each patient; it is not clear whether this followed a protocol or was at the discretion of the clinician. The mean follow-up time was 2.8 months, with a range of 2-6 months; it is not evident whether there was a standard follow up protocol; certainly, variable time after treatment could affect the reported outcomes and potentially the magnitude of change in either the SNOT-20 or NES scores.

It appears that experiments were performed consecutively for the two groups, rather than in parallel, so the results of the first study (group 1) appear to have influenced conducting the second study (group 2) experiments. Nasal endoscopic scoring was carried out in group 2 to “record the encouraging endoscopic results demonstrated in group 1.” In addition, saccharin transit tests were only carried out in group 2. It is not clear whether the nasal endoscopic scoring was carried out by the same clinician each time, or whether scores were independently checked, and could be considered subjective depending on the clinician’s experience; and if different clinicians carried out NES then inter-observer variability needs to be accounted for.
The study appears to have a design where experiments have been carried out consecutively and the first group 1 experiments have influenced how experiments and tests were carried out in the second group. The experiments should have been carried out in parallel using the same protocol. It is also unclear whether patients, clinicians or those carrying out the study were blinded to the type of treatment received.

Whilst there is a lot of missing information regarding the randomization process, the approach in blinding, how the similarity of the characteristics in the two groups were ensured, and how missing data was dealt with, we can conclude that there are considerable concerns regarding risk of bias, and this should be taken into account when interpreting the results and conclusions.

Canada has approved the use of Sinuwave technology for delivering photodynamic therapy in chronic rhinosinusitis. Desrosiers et al. (38) have pioneered the research in this field and after a small case series in 2013, this group conducted the first randomized control trial in 2016 (42). The results have only been published in conference abstract form and the full study results are yet to be published. Each treatment consisted of application of the photosensitizing agent to a previously operated sinus cavity and then illumination with a custom fiber-optic light diffusing balloon catheter. They recruited 23 CRS patients without polyps and 24 CRS patients with polyps to this study. Patients were randomized to PDT or to endoscopic irrigation with saline. Pre and post treatment measures included Sino-Nasal Outcome Test-22, endoscopic mucosal score, UPSIT smell test, conventional bacteriology, and Lund-Mackay endoscopic scores. The study showed PDT treatment improved symptoms and disease specific quality of life, the greatest improvement was in the CRS with polyps group receiving two treatments (endoscopic sinus score improvement of 47% at 6 months, p=0.007).

Discussion/Clinical and Research Consequences
Photodynamic therapy is a new technology that has been used in anti-cancer treatment and might have a role in the treatment of CRS. Studies suggest efficacy in both in-vitro and in-vivo settings, with safety studies so far demonstrating no unacceptable adverse effects.

Clinical studies have shown an improvement in objective and subjective outcome measures in patients with CRS. These studies have only used small sample sizes with follow up times of maximum six months and therefore do not demonstrate the potential long-term benefits or side-effects of PDT.

The studies discussed here use a variety of photosensitizing agents and light emitting devices (Sinuwave and Rhinolight) at different wavelengths. Further studies evaluating combinations of photosensitizing agents at different concentrations, along with each light emitting device at different wavelengths needs to be trialed in human subjects to understand the optimum setting for the eradication of antibiotic resistant biofilm forming bacteria. In addition, most of the clinical studies to date have compared groups of patients receiving different light therapies with no control arm. Studies with control arms receiving no therapy or other traditional medical therapy should be conducted.

The future of PDT will rely on further studies which accurately evaluate the long-term efficacy and sustainability of this intervention using both objective and subjective measures. The ideal study would be a large RCT with appropriately selected CRS patients (with primary and secondary CRS) followed up for more than one year. Evaluation methods should include quantitative measures of bacterial growth, CT scan findings, nasal endoscopic scoring, ciliary activity, olfactory assessment, and subjective measures including the SNOT-22 questionnaire.

Limitations
This review is limited by the lack of existing research, and the quality of included studies, especially clinical trials. Whilst photodynamic therapy is already being used successfully in other domains of medicine and there is in-vitro evidence for the efficacy of PDT, there is little evidence yet to fully support PDT as a viable treatment in CRS patients.

The in-vitro and ex-vivo studies evaluated here have used relatively small experimental groups and different photosensitizers and wavelengths of light. Therefore, it is difficult to compare the in-vitro studies to one another. It is yet unclear how the results of in-vitro studies that have used laboratory-grown biofilm from CRS isolates and/or silicone maxillary models can be extrapolated to the treatment of CRS in humans. Therefore in-vivo studies based on the in-vitro methods used are required to fully assess efficacy, durability, and safety. The clinical studies that have been conducted so far either have insufficient information regarding methodology or considerable risk of bias.

Conclusion
Research in CRS to date have shown that the factors in CRS pathophysiology include the sino-nasal microbiome, host immunity and mucosal barrier. Photodynamic therapy aims to reduce the bacterial load causing chronic infection and immune dysregulation. Preliminary data suggests that PDT is likely to be safe and has proven effective in-vitro. Further clinical research is required to evaluate the safety and efficacy of photodynamic therapy compared with traditional treatment, and how a healthier nasal microbiome can be restored.

The ultimate goal for research into PDT should be to demonstrate an ability to significantly improve the burden of CRS disease and identify the patients or the endotypes of the disease that are most likely to respond. For the treatment to reach clinical practice it will need to be proven as safe, effective, practical, and reproducible. Whilst this may take several years there is enough pre-clinical and early data for cautious optimism in this novel treatment modality.

Peer-review: Externally peer-reviewed.

Conflict of Interest: The authors have no conflicts of interest to declare.

Financial Disclosure: The authors declared that this study has received no financial support.

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