Application of photodynamic therapy in gastrointestinal disorders: an outdated or re-emerging technique?

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Photodynamic therapy (PDT) is a promising therapeutic modality that involves the administration of a photosensitizer followed by local illumination with a specific wavelength of light in the presence of oxygen. PDT is minimally invasive, has high selectivity for cancer, and has good patient compliance due to the simplicity of the procedure; therefore, PDT is widely used as a palliative and salvage treatment in patients with various gastrointestinal malignancies. When used as a salvage treatment for locoregional failures after definitive chemoradiotherapy for esophageal cancer, favorable results have been reported. PDT in conjunction with biliary stenting is a promising palliative treatment for unresectable cholangiocarcinoma, and can be used as an advanced diagnostic and therapeutic strategy in peritoneal dissemination of gastric cancer. Recent clinical reports of PDT for treating non-resectable pancreatic cancer also show promising results. To widen the application of PDT, the integration of PDT with molecular imaging and nanotechnology is being extensively studied. Based on these new developments, PDT is likely to re-emerge as a valuable technique in the treatment of diverse gastrointestinal diseases.

Keywords: Photochemotherapy; Gastrointestinal diseases; Gastrointestinal neoplasms; Palliative care; Photosensitizing agents
ly GI neoplasia [3].

So the question arises: is PDT an outdated technique? Although the role of PDT as a curative treatment has diminished, it is still widely used as a palliative and salvage treatment because it is minimally invasive, has high selectivity for cancer, and has good patient compliance due to the simplicity of the procedure. Advanced PDT technologies that are easier to deliver to local lesions, incorporate nanoparticles, and involve new photosensitizers and light sources have been developed. One attribute of PDT is that the majority of photosensitizers currently being used are fluorescent; this enables imaging to occur during therapy. Recently, this attribute has been exploited significantly. With these new developments PDT, in combination with improvements in endoscopic equipment, may be used to treat diverse GI diseases.

In this review, we summarize previously reported uses of PDT in GI diseases and introduce more recent clinical applications.

**PRINCIPLES AND MECHANISMS**

PDT is a well-established clinical modality that uses light energy to destroy dysplastic and malignant lesions. It requires three core elements: light of a specific wavelength, oxygen molecules, and a photosensitizer. A photosensitizer is a molecule that can absorb light energy and transfer that energy to adjacent oxygen molecules, which initiate a photodynamic reaction by forming cytotoxic molecular species including reactive oxygen, namely singlet oxygen [4]. Because the photosensitizer is preferentially localized to target tumor cells, and areas of light illumination can be controlled, this photodynamic reaction primarily damages the neoplastic lesion and preserves the adjacent normal tissue.

The GI tract starts at the mouth and ends at the anus. Most GI cancers occur at the mucosa layer, the most exposed part of the GI wall. With the development of flexible endoscopy, light energy can be directly delivered to a malignant lesion in the GI tract. This has made PDT popular as a minimally invasive local treatment modality for GI cancer [5]. Combined with imaging for dosimetry and demarcation, the potential for its use is even greater [6].

**ESOPHAGEAL CANCER**

PDT has been used as an endoscopic ablative technique to eliminate Barrett’s esophagus (BE) and, to a limited degree, high-grade dysplasia. This technique has resulted in regression and, in some cases, complete reversal of BE [7-10]. Since then, some studies of patients with esophageal dysplasia and cancer have reported that complete remission (CR) can be achieved in superficial lesions using PDT (Table 1) [11-19]. Studies on the use of PDT for esophageal squamous cell carcinoma (SCC) confined to the mucosal layer have suggested that PDT can be used as an alternative to esophagectomy given the low risk of lymph node metastasis in these patients. The majority of the patients included in the study had serious co-morbidities precluding surgical resection. Previous studies showed that the overall survival (OS) of patients with mucosal esophageal SCC treated with PDT was similar to that of patients who underwent esophagectomy [20,21]. There were no major complications and treatment-related deaths. Compared with endoscopic resection, PDT has a low complication rate, is simpler to perform, and is not limited by the size or circumference of the lesion. Given these results, PDT appears to be an ideal alternative to surgical or endoscopic resection in patients who have serious comorbidities and large or multifocal superficial SCC that cannot be completely resected endoscopically.

Recent studies have indicated a beneficial role for PDT in salvage treatment for locoregional failures after definitive chemoradiotherapy (CRT) for esophageal cancer. When locally remnant lesions are found after CRT, salvage esophagectomy should be preferentially considered as the curative treatment option. However, the approach has been associated with higher rates of postoperative mortality and morbidity compared with planned esophagectomy [22]. Although EMR may be an alternative treatment in these cases, it is limited to superficial lesions and the procedure requires highly experienced endoscopists. On the other hand, PDT can be used for superficial cancers and for palliation of more advanced esophageal SCC. In a retrospective study of 37 consecutive local failure patients treated with salvage PDT using porfimer sodium, CR occurred in 22 patients (59.5%), the 5-year progression-free survival (PFS) rate was 20.7%, and the OS rate was 36.1% during a median...
### Table 1. Photodynamic therapy in esophageal cancer

<table>
<thead>
<tr>
<th>Indication</th>
<th>Study</th>
<th>Study subjects</th>
<th>Therapeutic target</th>
<th>Type of PS</th>
<th>PS drug delivery</th>
<th>Outcome</th>
</tr>
</thead>
</table>
No recurrence during follow-up (range, 4–51 months)
No severe complication                                         |
|                                         | Tanaka et al. (2011) [14]               | Human (n = 38) | 38 Superficial SCC (31 T1a, 7 T1b)     | Porfimer   | IV              | CR: 33/38 (87%)
No recurrence: 28/38 (74%)
5-Year OS rate: 76% after a median follow-up period of 64 months
No major complication and treatment related death               |
| Salvage treatment for local failures after CRT | Yano et al. (2005) [15]               | Human (n = 13) | 4 Local recurrence after achieving CR   | Porfimer   | IV              | CR: 8/13 (62%)
6/8 (75%): free of disease after median follow-up of 12 months
1-Year OS rate: 68.4%
No treatment-related deaths                                     |
|                                         | Yano et al. (2011) [12]               | Human (n = 37) | T1/T2/T3/T4 in 3/4/24/6 patients No/1 in 13/24 patients | Porfimer   | IV              | CR: 22/37 (59.5%)
5-Year PFS and OS: 20.7%, 36.1%
Complication: esophageal fistulae = 4 (10.8%), stenosis = 20 (54.1%), phototoxicity = 2 (5.4%) |
|                                         | Yano et al. (2012) [11]               | Human (n = 9)  | Local failure limited to T2 after CRT   | Talaporfin | IV              | CR: 5/9 (55.6%)
No dose limiting toxicity and phototoxicity                     |
|                                         | Hatogai et al. (2016) [16]             | Human (n = 113)| Local failure limited to T2 after CRT   | Porfimer   | IV              | CR: 66/113 (58.4%)
5-Year PFS and OS: 22.3%, 35.9%
Treatment-related death: 1.8%                                   |
Mean dysphagia-free interval: 66 days
35 Patients: required stent placement, mean interval to re-intervention of 58.5 days
Complication: perforation (2%), stricture (2%), candida esophagitis (2%), pleural effusion (4%), sunburn (6%
Procedure-related mortality: 1.8%
Median survival: 4.8 months                                      |
|                                         | Lindenmann et al. (2012) [18]          | Human (n = 171)| Advanced esophageal cancer              | Porfimer   | IV              | PDT as first palliation (n = 118)
Mean number of palliative treatment: 2.6
PDT was used in the first place, median survival was 50.9 months compared to 17.3 months if other options were used as initial modality (p = 0.012). |
|                                         | Yoon et al. (2012) [19]               | Human (n = 20) | Advanced obstructing esophageal cancer  | Porfimer   | IV              | Improvement of dysphagia score: 18/20 (90%)
6 Patients: required stent insertion following PDT at 63 days on average |

PS, photosensitizer; SCC, squamous cell carcinoma; IV, intravenous; CR, complete remission; CRT, chemoradiation therapy; PFS, progression-free survival; OS, overall survival; PDT, photodynamic therapy.
follow-up period of 55 months [12]. Another large-scale study of 113 patients who showed local failure limited to within T2 after CRT reported respective PFS and OS rates of 22.1% and 35.9%, but a 1.8% rate of treatment-related death [16]. Talaporfin sodium is a second-generation, rapidly cleared photosensitizer that is expected to have less phototoxicity compared with porfimer sodium. In a phase I study, five out of nine patients with local failure after CRT achieved CR following PDT [11]. It is particularly noteworthy that phototoxicity and other severe adverse events were not observed.

PDT has also been used for palliative treatment of patients with completely or partially obstructing esophageal cancer. Two randomized controlled trials were done to compare PDT with Nd:YAG thermal ablation for palliation of dysphagia in advanced esophageal cancer [23,24]. Both therapies had, overall, equal efficacy in improving dysphagia, but PDT yielded a more prolonged duration of response. Another retrospective analysis was done on 250 patients who received multimodal palliative treatment, including PDT, for inoperable esophageal cancer [18]. In this study, if PDT was used as an initial therapy, the median survival was 50.9 months compared to 17.3 months if other options (endoscopic dilatation, endoluminal brachytherapy, external radiation, chemotherapy, stenting, feeding tube, or palliative resection) were used initially.

CHOLANGIOCARCINOMA

Cholangiocarcinoma (CCC) is difficult to diagnose early and usually presents at an advanced stage, which may be unresectable. Biliary stenting, mainly through the use of a self-expanding metal stent, has become the mainstay of palliative treatment for unresectable CCC with malignant strictures [25]. However, preventing tumor ingrowth and maintaining stent patency are unresolved problems [26]. Palliative options available for patients with hilar CCC include PDT, which involves a single minimally invasive treatment that has a low complication rate and is well tolerated. PDT in conjunction with biliary stenting could be an attractive treatment because it reduces the local obstruction of tumor tissue by generating singlet oxygen and may result in long-term patency.

Several uncontrolled and controlled studies comparing stenting alone to stenting with PDT for unresectable CCC have been conducted (Table 2) [27-33]. These trials showed that PDT improved cholestasis and quality of life by diminishing the need for further procedures, such as stent revision and percutaneous biliary drainage. This results in a longer stent patency period because PDT may cause tumor “remodeling,” which can prolong or enhance the decompression effect. PDT was also associated with increased OS in patients with unresectable CCC. The adverse effects of PDT were predictable and relatively easy to treat. Neither study reported severe photosensitivity or procedure-related death. Thus, PDT seems to be a promising palliative treatment for unresectable CCC and should be considered part of the standard of care for these patients.

PANCREATIC CANCER

The survival statistics for pancreatic cancer (PanCa) remain dismal and the mainstay for PanCa treatment is surgery. For patients with non-resectable disease, the outlook is even more dismal. PDT can play a significant role in the management of patients with non-resectable disease. There have been two reports on the use of PDT for palliation of non-resectable PanCa [34,35]. While the two studies used different photosensitizing agents, both had promising results. The benchmark for tumor necrosis was met in both cases. In a study with verteporfin, one patient reverted to being resectable and underwent an R0 Whipple procedure [34]. Combined with other therapeutics and nanomedicine, PDT has great potential for PanCa treatment.

GASTRIC CANCER

The outcome of gastric cancer patients has dramatically improved due to advancements in the accuracy of diagnostic endoscopy, and a decrease in the invasiveness of endoscopic therapies such as EMR and ESD. However, in patients with a background of intestinal metaplasia, it is still difficult to detect malignant lesions and evaluate the resection margin. Fluorescence navigation by photodynamic diagnosis (PDD) using 5-aminolevulinic
acid (ALA) can provide good visualization of gastric malignant lesions to determine the extent of the lesions. ALA is a naturally occurring amino acid and an early intermediate in the heme biosynthesis pathway. Orally administered ALA is mainly incorporated into malignant cells and metabolized to a photodynamically active protoporphyrin IX that emits a red fluorescence upon excitation with blue light [36]. The major advantage of ALA is that it is rapidly metabolized, significantly reducing the period of cutaneous photosensitivity. Several studies have demonstrated that ALA-mediated PDD is a promising tool for diagnosing minute lesions and peritoneal metastasis in gastric cancer. A Japanese study showed that laser-based endoscopy detected upper GI tumors as red fluorescent navigation (PDD-positive) in 23 of 27 lesions [37]. Interestingly, all but one of the intestinal type carcinomas were significantly PDD-positive, whereas signet ring cell carcinomas were PDD-negative. To evaluate the detection of peritoneal metastasis in gastric cancer, ALA-PDD was added to conventional diagnostic laparoscopy for 52 advanced gastric cancer patients [38]. Using ALA-PDD, peritoneal dissemination was detected in five of 24 patients with no macroscopic evidence of peritoneal metastases on white light. In another study, staging laparoscopy with ALA-mediated PDD was performed in 13 patients with serosa-invading gastric cancer [39]. When PDD was used, the peritoneal metastasis detection rate was significantly higher than with white light. These results show that ALA-PDD may be useful for preoperative staging of advanced gastric cancer.

Treatment of advanced gastric cancer, especially in serosa-positive cases, has focused on preventing peritoneal recurrence because the peritoneum is one of the most common recurrence sites after curative gastrectomy. Although several therapeutic options, including systemic adjuvant chemotherapy and intraperitoneal chemotherapy with or without hyperthermia and CRT, have been tried [40-42], there is no standard effective treatment to prevent peritoneal recurrence and significantly increase survival. PDT may be a good alternative treatment option because micrometastatic nodules on the peritoneal surface and intraperitoneal floating cancer cells are potentially suitable PDT targets. In a study using a peritoneal metastasis mice model, talaporfin, another second-generation photosensitizer, accumulated more

<table>
<thead>
<tr>
<th>Study</th>
<th>Study design</th>
<th>Study sample</th>
<th>Bisphosphonate</th>
<th>No. of PDT</th>
<th>Type of PS</th>
<th>Median survival</th>
<th>Adjvant therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ortner et al. (2003) [28]</td>
<td>RCT</td>
<td>20</td>
<td>PDT: II (0)/III (4)/IV (16)</td>
<td>Control: II (2)/III (2)/IV (15)</td>
<td>Porfimer sodium</td>
<td>PDT: 493 day</td>
<td>Control: (–)</td>
</tr>
<tr>
<td>Zoepf et al. (2005) [29]</td>
<td>RCT</td>
<td>16</td>
<td>PDT: II (1)/III (0)/IV (15)</td>
<td>Control: II (0)/III (0)/IV (16)</td>
<td>Photosan</td>
<td>PDT: 630 day</td>
<td>Control: (–)</td>
</tr>
<tr>
<td>Witzgall et al. (2006) [30]</td>
<td>Uncontrolled</td>
<td>68</td>
<td>PDT: I (1)/II (0)/III (14)/IV (53)</td>
<td>Control: I (5)/II (9)/III (12)/IV (30)</td>
<td>Porfimer sodium</td>
<td>PDT: 12 mon</td>
<td>Control: CTX, 6, RTX, 2</td>
</tr>
<tr>
<td>Kahaleh et al. (2008) [27]</td>
<td>Uncontrolled</td>
<td>19</td>
<td>PDT: I (2)/II (1)/III (7)/IV (9)</td>
<td>Control: I (1)/II (8)/III (10)/IV (10)</td>
<td>Porfimer sodium</td>
<td>PDT: 8 mon</td>
<td>Control: CTX, 8</td>
</tr>
<tr>
<td>Quynh et al. (2009) [31]</td>
<td>Prospective cohort</td>
<td>23</td>
<td>PDT: II (1)/III (10)/IV (12)</td>
<td>Control: II (1)/III (5)/IV (11)</td>
<td>Porfimer sodium</td>
<td>PDT: 25 day</td>
<td>Control: (–)</td>
</tr>
<tr>
<td>Cheon et al. (2012) [32]</td>
<td>Uncontrolled</td>
<td>17</td>
<td>PDT: I (6)/II (10)/IV (1)</td>
<td>Control: I (16)/II (29)/IV (26)</td>
<td>Porfimer sodium</td>
<td>PDT: 215 day</td>
<td>Control: (–)</td>
</tr>
<tr>
<td>Lee et al. (2012) [33]</td>
<td>Retrospective</td>
<td>15</td>
<td>PDT: II (6)/III (27)/IV (8)</td>
<td>Control: II (4)/III (7)/IV (4)</td>
<td>Porfimer sodium</td>
<td>PDT: 36 day</td>
<td>Control: (–)</td>
</tr>
<tr>
<td>Lee et al. (2013) [34]</td>
<td>Retrospective</td>
<td>18</td>
<td>PDT: I (4)/II (19)/IV (1)</td>
<td>Control: I (4)/II (19)/IV (1)</td>
<td>Porfimer sodium</td>
<td>PDT: 30 day</td>
<td>Control: (–)</td>
</tr>
</tbody>
</table>

PDT, photodynamic therapy; PS, photosensitizer; RCT, randomized controlled trial; CTX, chemotherapy; RTX, radiation therapy.
in metastatic nodules than normal organs when injected intraperitoneally [43]. Talaporfin-mediated PDT also exerts considerable antitumor effects in vivo. The total weight of peritoneal metastatic nodules was lower in mice models with PDT delivered using nanoparticles loaded with indocyanine green. These mice also had significantly improved body weight loss and a higher survival rate [44]. These results show that PDT may be a promising diagnostic and therapeutic modality in peritoneal dissemination of gastric cancer.

**RECENT DEVELOPMENTS TO OVERCOME THE LIMITATIONS OF PDT**

As outlined above, PDT is a very attractive treatment modality for cancer because it has minimal toxicity and can be applied repeatedly without harming adjacent normal tissue [45]. PDT is currently widely used to treat skin diseases and easily accessible malignant and pre-malignant lesions.

To maximize the effect of PDT, further research is needed to enhance its three core elements: the photosensitizer, light, and oxygen. Fig. 1 shows recent developments to overcome the limitations of PDT.

The photosensitizer can cause hypersensitivity, allergy, toxicity, and unpredictable biodistribution profiles if administrated intravenously, especially in multiple doses. Unpredictable biodistribution can lead to the accumulation of photosensitizer in easily light-exposed uninvolved tissues such as the eyes and skin, and this can result in phototoxicity and prolonged photosensitivity when patients are exposed to ambient light. Local delivery of the photosensitizer may solve these problems. Local delivery may shorten treatment time, enhance therapeutic efficacy, and eliminate skin photosensitization completely. There are several approaches for the local delivery of PDT to target GI tumors; these include spraying via the working channel of the endoscope, intratumoral injection, and the use of photosensitizer-coated stents.

To enhance the clinical use of PDT, the second challenge is to improve the targeting of the drug-activating light to the target tissue. In PDT, superficial tissues can be easily illuminated using convenient light sources, but deep tissue tumors are more difficult for light to penetrate. For example, blue light (400 to 450 nm) penetrates approximately 1 mm into tissue, while orange light (590 to 620 nm) penetrates approximately 1.5 mm, and red light (620 to 750 nm) penetrates even further, up to 3 mm [46]. Newer light sources, such as light-emitting diodes (LEDs), are being explored to deliver light more effectively. LEDs can generate high-energy light of desired wavelengths and can be assembled in a range of geometries and sizes [47,48]. During PDT, monitoring tumor tissue oxygenation, by measuring the partial pressure of oxygen and the level of hypoxia markers, may be used to increase the concentration of singlet oxygen to the target tissue during light irradiation. In addition, fractionating light irradiation by controlling light/dark periods can promote tissue oxygen re-perfusion to compensate for oxygen depletion caused by photochemical reactions [49,50].

![Figure 1. Recent developments and future perspectives with regard to improving the efficacy of photodynamic therapy (PDT) to treat gastrointestinal tract cancers. EUS, endoscopic ultrasound; LED, light-emitting diode.](https://www.kjim.org)
NOVEL APPLICATIONS OF PDD AND PDT IN GI MALIGNANCY

Much research has been done to extend the application of photodynamic techniques to the early diagnosis and treatment of GI malignancy.

In the spotlight in the field of GI endoscopy is molecular imaging because it can be used to detect lesions at an early stage. Molecular imaging can be used to visualize the molecular fingerprint of target cells, and not just their gross morphology. Developments in molecular imaging are occurring with advances in new imaging techniques such as virtual chromoendoscopy and magnifying endoscopy. In molecular imaging, disease-specific biomarkers are targeted using fluorescently labeled exogenous molecular probes including antibodies, antibody fragments, peptides, nanoparticles, and activatable probes [51]. Combining a molecular probe with a photosensitizer can enhance tumor targeting and the detection of minute lesions by the endoscopist. Furthermore, it can enable optical biopsy and theragnosis, which combines simultaneous diagnosis and therapeutics. In our group, a peptide specific to colon cancer was isolated using phage display and then conjugated to hematoporphyrin (HPP), a promising second-generation photosensitizer [52]. Immunocytochemical staining using fluorescein isothiocyanate or HPP-conjugated peptides showed high binding affinity to colon cancer cell lines. The HPP-conjugated cancer-specific peptide showed an enhanced PDT effect, compared with HPP alone. These results show that targeted PDT using peptide probes may constitute promising candidate treatments for use in the development of colon cancer diagnostics and treatments.

Photosensitizer-embedded stents allow for repeatable intraluminal PDT, which leads to regression of surrounding GI malignant tissue and extends stent patency. A recent study of a polymeric photosensitizer (pulullan acetate-conjugated phorphoride A)-embedded self-expanding nonvascular metal stent, which permits repeatable PDT of CCC without systemic injection of the photosensitizer, reported that the stent sustained its PDT effects for at least 2 months in vitro [53]. Furthermore, in vivo testing using HCT-116 tumor-xenograft mouse models showed degradation of tumor cells and regression of xenograft tumors after laser exposure. PDT can improve the tissue penetration efficiency (PE%) of released drugs in non-vascular drug-eluting stents (DESs) by breaking the epithelial layer with singlet oxygen, a process called photochemical tissue penetration (PTP). We developed a PTP-DES with a photosensitizer (chlorin e6) incorporated in a gemcitabine-eluting polyurethane membrane [54]. This stent demonstrated a 2-fold increase in the PE% of gemcitabine, and tumor growth was effectively inhibited in an in vivo mouse model. Our results indicate that the PTP-DES system has great potential for improving the therapeutic effect of conventional DES.

Nanomedicine that combines PDT with molecular target therapy may be used to enhance the efficacy of anticancer treatments and improve outcomes for patients with highly treatment-resistant GI malignancy, while decreasing toxicity. Recently, we developed a novel photoactivatable multi-inhibitor nanoliposome (PMIL), which consists of a nanoliposome doped with a photosensitizer (benzoporphyrin derivative) in the lipid bilayer, and a nanoparticle encapsulating cabozantinib, a multikinase inhibitor [55]. This novel nanoscale drug not only causes photodynamic damage to tumor cells and microvessels but also initiates the release of cabozantinib inside the tumor. In a mouse model of PanCa, PMIL resulted in prolonged tumor reduction and metastasis suppression. These results open the door to the application of PDT in tumor-selective chemotherapy.

CONCLUSIONS

We reviewed a number of key studies that used PDT to treat GI malignancy. Drawbacks of PDT including suboptimal efficacy, phototoxicity, and high cost have curtailed the initial enthusiasm for this procedure in treating GI malignancy. PDT still has a role as a palliation and salvage treatment because it is a minimally invasive procedure. Much effort has been put into widening the application of PDT, and favorable results in areas such as early diagnosis, salvage treatment for local failures, and palliation of GI cancer, are shedding new light on the applications of PDD and PDT. Novel integration of PDT with existing technologies such as molecular imaging, DES, and nanotechnology may make PDT a feasi-
ble and desirable option for the treatment of GI cancers.

**Conflict of interest**

No potential conflict of interest relevant to this article was reported.

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