Background

The use of indocyanine green angiography (ICG-A) to assess tissue perfusion has been described in gastrointestinal surgery to assess bowel perfusion to guide resection or anastomosis as well as in reconstructive procedures to predict skin and soft tissue viability. We present a novel use of this technology in which the Stryker SPY-PHI system (a handheld indocyanine green fluorescence imaging system) was utilized to aid in tissue resection boundaries based on perfusion in a patient with acute necrotizing fasciitis.

Summary

A 68-year-old, morbidly obese female presented with several days of worsening perineal and left gluteal pain. Examination of her left labia majora and left gluteal fold was consistent with a necrotizing soft tissue infection. Computed tomography confirmed the diagnosis, demonstrating significant subcutaneous emphysema and abscess formation. She was taken emergently to the operating room for primary debridement. On postoperative day two she returned to the OR for wound washout and possible re-debridement. Grossly non-viable tissue was first excised; then fluorescence perfusion assessment was utilized to assess for non-viable and poorly perfused tissue that would have otherwise not been debrided based on visual examination. The patient subsequently recovered quickly without the need for further debridement. She was discharged with local wound care.

Conclusion

Given that the surgical tenet for necrotizing soft tissue infections is serial debridement, the use of fluorescence perfusion assessment as an adjunct for intraoperative debridement decisions may potentially minimize the need for repeated debridement, thus achieving quicker source control. To our knowledge, this is the first documented use of this technology in the management of necrotizing soft tissue infections.

Key Words

indocyanine green angiography; necrotizing fasciitis; NSTI

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Case Description

Necrotizing soft tissue infections (NSTI) are life-threatening conditions that require prompt surgical intervention. Due to the aggressive nature of NSTIs, delay in diagnosis and treatment increases morbidity and mortality, and clinical diagnosis based on history, physical examination, and high clinical suspicion remain crucial.\textsuperscript{1-3} In addition to antibiotics and resuscitation, the mainstay of NSTI management is early surgical debridement.\textsuperscript{4} This typically requires multiple returns to the operating room to debride grossly infected and non-viable tissue until complete source control is obtained.

The use of indocyanine green angiography (ICG-A) to assess tissue perfusion has been well-described in bowel surgery to evaluate bowel perfusion for resection or before anastomosis, but it is also utilized in reconstructive procedures to predict skin and soft tissue viability.\textsuperscript{5-7} Several plastic surgery reports have shown that the addition of ICG-A with clinical examination was a useful and safe technique in monitoring free flap compromise both intraoperatively and postoperatively, leading to improved flap success rates.\textsuperscript{8} After a review of the literature, this technology has not been utilized in the setting of NSTI debridement. We present a case of necrotizing fasciitis in which the Stryker SPY-PHI system (a handheld indocyanine green fluorescence imaging system) was utilized to aid in tissue resection boundaries based on perfusion.

A 68-year-old, morbidly obese female who presented to the emergency department with several days of worsening perineal and left gluteal pain while visiting from outside the country. Examination of her left labia majora and left gluteal fold was consistent with a necrotizing soft tissue infection. Computed tomography (CT) demonstrated significant subcutaneous emphysema and abscess formation, consistent with necrotizing fasciitis (Figure 1). Laboratory values were consistent with an elevated WBC of 21000, hyponatremia, and lactic acidosis. Given these findings, she was taken emergently to the operating room for primary debridement.

Twenty-four hours after initial debridement, she returned to the OR for wound washout and possible further debridement. Grossly non-viable tissue was first excised as normal. Then, fluorescence perfusion assessment was utilized to assess non-viable and poorly perfused tissue (Figure 2), revealing multiple areas of poorly perfused tissue that would have otherwise not been debrided based on visual examination. Image-guided excision was therefore undertaken. Given that this technology was not utilized to change the intent of the procedure, the known safety profile of ICG, and the emergent nature of this procedure, IRB approval was not obtained.

Figure 1. CT Imaging Revealed Significant Inflammation and Gas in Subcutaneous Tissues, Consistent With Necrotizing Fasciitis. Published with Permission

![Figure 1](https://example.com/figure1.png)

Figure 2. Visible Spectrum Light and ICG Fluorescence Imaging of Wound at Second Debridement. Published with Permission

![Figure 2](https://example.com/figure2.png)
The patient subsequently recovered quickly without the need for further debridement, with the remainder of her hospital stay requiring simple local wound dressing. Final wound cultures were consistent with a polymicrobial infection of *Escherichia coli* and *Bacteroides fragilis*, for which her antibiotics were appropriately narrowed. She was discharged on postoperative day nine with local wound care to her home country. By postoperative day 15, her wound continued to heal well with wet-to-dry dressing changes (Figure 3). At 4.5 months postop, the wound was completely healed (Figure 4).

**Discussion**

Given that the surgical tenet of necrotizing soft tissue infections is serial debridement procedures, the use of fluorescence perfusion assessment as an adjunct for intraoperative debridement decision making can help achieve quicker source control, thereby minimizing the need for repeated debridement. Given that many NSTI patients are hemodynamically unstable due to underlying sepsis and their need for serial debridement, patients are often subjected to prolonged intubation. The sequelae of long-standing intubation are well documented. Rapid source control and achievement of clinical stability will help reduce the possible complications of ventilator support. It would also help reduce the possible detrimental cumulative effects of general anesthesia.

Wound care, especially for larger wounds, may constantly have to be adjusted following every subsequent debridement. If the number of procedures was reduced, the final wound would be realized much sooner, and local wound care could be better established. Critical care length of stay would potentially be decreased, thereby helping offset the overall costs to both the patient and the hospital. According to Childers et al., the mean cost per minute of OR time ranges between $30 to $113. The cost of the Stryker SPY-PHI system is approximately $495 in direct costs to the patient, per use.
Indocyanine green has been proven to be a relatively safe entity, with no known metabolites. Once intravascularly injected, the binding of ICG to plasma proteins allows for its restriction in the vascular compartment, with eventual excretion into the bile. In fact, ICG has been utilized for decades in multiple areas of science and medicine, including cardiac surgery, surgical oncology, sentinel lymph node mapping in different cancers, hepatic surgery, and ophthalmology. Boni et al. reported on a series of 108 patients who underwent differing laparoscopic surgeries, all without any evidence of intraoperative or injection-related adverse effects. Depending on its use, the total dose of ICG ranges from 0.5 mg/kg of body weight in hepatic function testing to ≤2 mg/kg of body weight in cardiac output monitoring, though doses of up to 5 mg/kg appear to be safe. Rare cases of severe reactions, such as anaphylactic shock, have been reported, and close monitoring after ICG administration should be undertaken. Review of the literature indicates that there are multiple areas of interest being investigated in the use of ICG angiography.

As this is the first documented use of ICG in this setting, further studies would need to be conducted to compare its use to the standard serial debridentments. The patient’s clinical status is always taken into account when determining if further debridement is needed. It is unclear from this one case if her second debridement would have changed her clinical course. Good local wound care may have been all that was needed. Perhaps this technology would be better suited in a larger wound or in a patient who has undergone more debridentments without clinical improvement. Regardless, it may represent an adjunct to clinical examination when viability margins are uncertain.

**Conclusion**

To our knowledge, this is the first documented use of ICG angiography in the management of NSTI. While this technology is not specifically FDA-approved for this indication, we feel that the use of ICG fluorescence in necrotizing soft tissue infections is a new and novel way to improve and expedite patient care.

**Lessons Learned**

Indocyanine green angiography is an excellent tool to assess tissue viability in the setting of necrotizing soft tissue infection. It may aid in intraoperative decision-making for debridement of non-viable tissue otherwise not grossly appreciated, allowing for a more definitive surgery and fewer returns to the operating room for serial debridement.

**References**


