Invasive Fungal Infection in War Wounds (CPG: 28)
Provide guidance on the recognition and comprehensive management of invasive fungal infection (IFI) in war wounds.

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BACKGROUND

Clinically significant infections, including invasive fungal wound infections (IFIs), have occurred in our wounded warrior patient population since the beginning of the current war in Afghanistan. During 2009-2010, a significant increase in the incidence of IFIs was observed among military personnel with wounds sustained in Afghanistan, corresponding to a greater frequency of improvised explosive device blast injuries sustained while on foot patrol in Helmand and Kandahar provinces.\(^1\)\(^-\)\(^3\) Of particular clinical concern was an apparent association between patient outcome and the presence of angioinvasive molds (e.g., order Mucorales, Aspergillus species, and Fusarium species). In general, IFIs are devastating infections associated with increased mortality, morbidity (e.g., amputation), and prolonged hospitalization for survivors.\(^2\)\(^,\)\(^4\)\(^-\)\(^12\) In civilian literature, mortality rates have been reported as high as 38%.\(^13\)\(^-\)\(^17\) Among the military population, the crude mortality rate was as high as 8% during the first two years of the outbreak.\(^6\)

Following recognition of the high number of IFI cases, the Joint Trauma System, in collaboration with the Trauma Infectious Disease Outcomes Study (TIDOS), launched an outbreak investigation. Review of the findings demonstrated that the most common mechanistic and clinical factors associated with IFI included dismounted blast injury, above knee traumatic amputations, extensive perineal/pelvic injury (observed trend, but not statistically significant), and massive packed red blood cell transfusion (≥20 units in the first 24 hours).\(^1\)\(^,\)\(^2\) Importantly, all IFI patients had a suspicious wound (i.e., unhealthy appearance), defined as recurrent tissue necrosis following at least two surgical debridements. (See Appendix A: Examples of Suspicious Wounds.)

The morbidity associated with IFI in war wounds, which may include significant tissue loss, necessitates early treatment of patients identified as high risk. Patients frequently require surgical amputations and/or amputation revisions, which include extending to more proximal levels (e.g., transtibial to transfemoral or transfemoral to proximal transfemoral, hip disarticulation, or hemipelvectomy).\(^18\) Although prevention strategies have not been clearly identified, early and aggressive debridement of devitalized tissue and removal of debris are universally accepted as the most important interventions. The treatment of IFI is based on three main principles: debridement of infected tissue, minimization of immunosuppression (e.g., avoidance of malnutrition or excessive blood product transfusion), and utilization of empiric dual antifungal medications (e.g., amphotericin B and a broad-spectrum triazole agent) when there is a strong suspicion of an IFI.\(^6\) The role of topical antifungal therapy in the prevention of IFI is not clear, but topical therapies have not been demonstrated to have adverse local or systemic effects.

EVALUATION AND TREATMENT

The most important aspect of evaluation and treatment of war wounds is the recognition of unhealthy or suspicious wounds followed by early, aggressive, and repetitive surgical debridement of all devitalized tissue and organic material.

After initial debridement, risk factors for invasive fungal infection will be assessed. Identified risk factors include:

- Dismounted blast injury.
- Above knee immediate traumatic amputation, or progressive transition from below knee to through knee to above knee amputation.
- Extensive perineal, genitourinary, and/or rectal injury.
- Massive transfusion > 20 units packed red blood cells within 24 hours of injury.

Additionally, a web-based clinical decision support tool to assist healthcare providers in assessing the probability of developing an IFI was jointly developed by the Surgical Critical Care Initiative (SC2I) and the TIDOS project...
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both based at the Uniformed Services University of Health Sciences (http://www.sc2i.org/ificdss). The tool was designed for early clinical evaluation in theater as well as upon arrival at the Role 4/5 facility (Military Medical Center).

DIAGNOSIS CRITERIA

Diagnostic criteria for an IFI are: presence of a traumatic wound(s), recurrent necrosis following at least two consecutive surgical debridements, and laboratory evidence of fungal infection (i.e., mold culture positivity and/or histopathology indicating tissue invasion). This last criteria is usually available at deployed Role 2 or Role 3 Military Treatment Facilities (MTFs), so clinical suspicion is key to early intervention.

TOPICAL TREATMENT

Initiate topical antifungal therapy on patients with at least three of the above risk factors. Topical antifungal therapy should be initiated with 0.025% Dakins. Begin with Dakins low pressure, high volume irrigation in the operating room (OR) after the first or second operative debridement—use in lieu of saline irrigations for patients meeting criteria. Cover wounds with Dakins-soaked Kerlix dressing. Alternatively, an instillation vacuum dressing may be created by placing one additional infusion catheter per suction port on the vacuum dressing sponge; hold suction for 5 min and instill 50 cc 0.025% Dakins, then clamp catheters and restart vacuum; repeat every 1-2 hours.

A standardized operative note for wound description to be used throughout the continuum of care for patients at increased risk for IFI is available. Utilization of this operative note may facilitate the early detection of sequential wound necrosis (i.e., the first sign of IFI) – Appendix B. Description of Bastion Classification of lower limb injuries is presented in Appendix C and is to be documented on the first page of the Operative Note.

DEBRIEEMENT AND ANTIFUNGAL THERAPY

1. For patients transferred to any Role 3 strategic evacuation hub, risk factors for IFI should be assessed and ongoing resuscitation requirements should be addressed as needed. The patient should undergo surgical examination, wound washout, and debridement (if indicated) within 12-18 hours of arrival. Dakins wound irrigations or Dakins-soaked Kerlix dressings as described above should be initiated/continued.

2. Topical antifungal treatment using 0.025% Dakins solution via instillation vacuum dressing should be continued throughout the evacuation phase- if possible. Flight teams should receive instruction on management of the instillation vacuum device prior to leaving the MTF. In the event of malfunction during flight, the instillation may be held while vacuum dressing is continued. The surgeon on call should be then be contacted to evaluate the dressing immediately on arrival to the next level of care.

3. Upon arrival to the Role 4 MTF (i.e., regional treatment facility outside of the combat zone, but prior to arrival in the United States), the patient should undergo operative exploration, wound washout, and debridement (as indicated) within 12-18 hours. Histopathology and microbiology specimens should obtained at Role 4 on all patients with at least three risk factors for IFI and any with clinical suspicion. Topical antifungal therapy with 0.025% Dakins solution should be continued if there is continued suspicion or three risk factors for IFI, preferably using an instillation vacuum dressing. If not available, 0.025% Dakins soaked kerlix dressing should be used.

4. Upon arrival to an MTF in the United States, the patient should undergo surgical exploration, wound washout, and debridement within 12-18 hours. Histopathology and microbiology specimens should be
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on all patients with at least three risk factors for IFI and/or who have an unhealthy wound appearance (e.g., tissue necrosis). Topical Dakins dressings may be discontinued at any level of care when the treating surgeon observes healthy granulation, or when histopathology and cultures are negative for fungal infection or colonization.

5. If **tissue necrosis** is observed in wounds following two consecutive debridements, not including the first two debridements in theater, **broad-spectrum antifungal and antibiotic medications should be started immediately** and Infectious Disease consultation obtained. Liposomal amphotericin B is the primary choice due to its effectiveness against mucormycosis and its reduced potential to induce nephrotoxicity.24 Although voriconazole is ineffective against mucormycosis, it has shown to be an active agent against molds that are resistant to amphotericin B (e.g., Aspergillus terreus and Scedosporium prolificans).25

6. In general, patients with IFI are severely injured, and are predominantly prescribed intravenous formulations of antifungal agents as there is concern for inadequate gastrointestinal antifungal absorption in the septic patient.

7. **When voriconazole is administered intravenously**, it requires a solubilizing excipient (i.e., sulfobutyl ether β-cyclodextrin), which may accumulate in patients with impaired renal function. A black box warning has been issued due to adverse effects of the accumulating solute in an animal model. Nevertheless, the effects of elevated sulfobutyl ether β-cyclodextrin are unknown in humans.26 Clinical experience to date has not shown permanent renal impairment with this off-label use of voriconazole in the wounded military population.27

8. Posaconazole is another triazole agent that has been found to have a 60-70% response rate as a salvage regimen against mucormycosis when prescribed orally.28,29 Recently, an intravenous formulation was approved and has shown to be useful.30 **Dual administration of liposomal amphotericin B and a broad-spectrum triazole (i.e., clinical experience has been primarily with voriconazole) is recommended as the first-line antifungal agents as many of the wounds incurred by combat casualties grow more than one mold.**31 Furthermore, broad-spectrum antibiotics covering both gram-positive and gram-negative organisms (e.g., Vancomycin and Meropenem) are prescribed as fungal-infected wounds frequently have bacterial growth as well.

9. **Particular attention should be given to aggressive debridement of non-viable tissue at each debridement procedure.** The extent of necrosis and appearance of the wound before and after completion of the operation should be documented in the operative note. **Appendix B** shows a standardized operative note for wound description to be used for patients at increased risk for IFI. Whenever a significant amount of necrotic tissue is debrided, repeat debridement should be performed in 24 hours or less.

10. **Topical antibacterial and antifungal beads may be considered in cases of proven or strongly suspected IFI,** and may be used in conjunction with vacuum/instillation dressings. The beads should be made with liposomal amphotericin B-500 mg, voriconazole-200 mg, tobramycin-1.2 gm, and vancomycin-1 gm.

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**Tissue Biopsy in OR**

**Biopsy should be done at the time of wound exploration (after initial surgical debridement)** once the casualty has been evacuated from the theater of conflict (in theater if patient evacuation is delayed) and repeated on subsequent explorations if there are persistent fevers and wound necrosis raising suspicion for IFI.

1. Tissue samples should be obtained from each lower extremity in patients with bilateral lower extremity amputations. Compromised muscle and adipose tissue should both be sampled.
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2. Other sites sampled should be at the discretion of the operative surgeon.

3. At least one specimen should be taken from the junction of viable and necrotic tissue (the last piece of borderline-viable tissue removed).

4. For each site sampled, two tissue samples will be collected fresh in two separate sterile specimen containers.
   - One specimen (1 cm³) for histopathological examination
   - One specimen (1 cm³) for fungal and bacterial culture

**OR STAFF RESPONSIBILITIES**

The histopathology specimen must leave the OR as a fresh specimen.

1. Order histopathology and cultures (aerobic, anaerobic, and fungal). Special studies are not routinely done, but may be requested (e.g., mycobacterial and viral).

2. Clearly label specimens as “blast biopsy protocol”. Labels should also contain the following information:
   - Site (e.g., left lower extremity)
   - Patient’s name, DOB, and hospital identification number

3. Directly contact the histopathology lab during working hours and the on-call pathologist after hours and on weekends to let them know they will receive a histopathology specimen shortly. Deliver the histopathology specimen to the Pathology Lab as soon as possible.

**PATHOLOGY STAFF RESPONSIBILITIES**

Pathology staff will coordinate processing as rapidly as possible (≤ 24 hours).

1. Histopathological specimen will be stained with hematoxylin and eosin (H&E) and Gomori Methenamine Silver (GMS)/Periodic Acid-Schiff (PAS) stains and evaluated for fungal elements.

2. Microbiological specimen will be cultured for aerobes, anaerobes, and fungi.

3. Mycobacterial and/or viral cultures will not be done routinely under this protocol, but may be done with special request.

If angioinvasive fungal elements or fungal elements among necrotic debris are reported on histopathology, or if cultures are positive in the setting of recurrent necrosis, treatment with systemic antifungal medications should be initiated (or continued). Treatment will require close consultation with Infectious Disease; however, as a general guideline, stop systemic antifungal medications if the wound remains clean/viable for two weeks and if the patient remains clinically stable. If the patient has a fungal infection in more than one body region (e.g., extremity/pelvis, abdomen, and chest), long-term treatment may be indicated.

**NOTE:**

1. Fungus can take up to six weeks to grow in culture medium. Therefore, it is recommended that the cultures be checked frequently for two weeks; then once a week for four additional weeks before they are considered final. In addition, wounds without recurrent tissue necrosis may have mold colonization and not a true infection.32
2. Initial studies have shown that combat IFI wound cultures growing order Mucorales will have a second non-Mucorales fungus present 30% of the time. Aspergillus species is more difficult to grow than order Mucorales, but should be suspected and empirically treated initially as it has been shown to be virulent in this patient population.\(^\text{21}\) Therefore, dual use of a broad-spectrum triazole and liposomal amphotericin B is suggested for wounds infected with either or both of these fungi. If long-term treatment is required, the antifungal medications should be targeted based on culture results.

As aggressive surgical debridement of all necrotic and infected tissue remains the mainstay of treatment for IFI, surgical exploration and debridement should continue at least every 24 hours until cessation of necrosis occurs. Wound coverage and closure should not occur until after the wound is clean, contracting, and granulating.

**PERFORMANCE IMPROVEMENT (PI) MONITORING**

1. Correct identification and documentation of patients at increased risk for IFI (≥ 3 risk factors). Wound exploration/debridement within 12-18 hours of arrival at each facility for patients at increased risk for IFI.

2. Appropriate tissue sampling/biopsy including site and specimen selection, specimen handling, and notification of the pathology department.

3. Topical Dakins dressing initiated in theater for patients at increased risk for IFI.

4. Appropriate documentation of wound appearance at each level of care using the standardized operative note for patients at increased risk for IFI.

5. Antifungal medications started when there is tissue necrosis on two consecutive debridements, not including the first two debridements.

6. Systemic and topical antifungal treatments stopped when no evidence of IFI on histopathology or culture when wounds have remained viable for at least two weeks after previously presenting with necrosis on two consecutive surgical debridements.

**INTENT (EXPECTED OUTCOMES)**

- Casualties at risk for IFI will be identified early along the continuum of care.
- Documentation with specific attention to risk factors for IFI will be performed at each level of care.

**PERFORMANCE/ADHERENCE MEASURES**

1. Casualties with ≥3 IFI risk factors are taken to OR within 12-18 hours upon arrival at Role 3 or 4 MTFs.

2. Casualties with ≥3 IFI risk factors have Dakin’s solution applied to wounds.

3. Standardized operative note will be used at Role 3 facilities in theater and Role 4 facilities for casualties with ≥3 risk factors for IFI.

**DATA SOURCE**

- Patient Record
- Department of Defense Trauma Registry (DoDTR)
SYSTEM REPORTING & FREQUENCY

The above constitutes the minimum criteria for PI monitoring of this CPG. System reporting will be performed annually; additional PI monitoring and system reporting may be performed as needed.

The system review and data analysis will be performed by the JTS Director, JTS Program Manager, and the JTS Performance Improvement Branch.

REFERENCES


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APPENDIX A: EXAMPLES OF SUSPICIOUS WOUNDS

All photos are from different patients.

(A) Patient is 8 days out from injury. His wound is showing central necrosis. Debridement day prior revealed healthy appearing, bleeding tissue.

(B) Patient is also 8 days out from injury. Patchy necrosis is seen throughout his right-sided hemipelvectomy. Upon closer inspection, one notices a ‘burnt butterscotch’ appearance overlying tissue. Anecdotally, this is an additional IFI clinical indicator.

(C) Provides example of tissue necrosis in a previously healthy-appearing wound bed.

(D) Patient is recovering well, but still being treated with anti-microbial beads.
(E) Looking closely at the wound, one can see a ‘yellow-velvet’ covering to the wound. This is indicative of an Aspergillus infection.
### APPENDIX B: MD TRAUMA WOUND DEBRIDEMENT OP NOTE

<table>
<thead>
<tr>
<th>MD TRAUMA WOUND DEBRIDEMENT OP NOTE</th>
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<tbody>
<tr>
<td><strong>Date of Operation:</strong></td>
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<tr>
<td><strong>Pre-Operative Diagnosis:</strong></td>
</tr>
<tr>
<td><strong>Post-Operative Diagnosis:</strong></td>
</tr>
<tr>
<td><strong>Initial Bastion Amputation Class:</strong></td>
</tr>
<tr>
<td><strong>Surgeon(s):</strong></td>
</tr>
<tr>
<td><strong>Anesthesia:</strong></td>
</tr>
<tr>
<td><strong>EBL:</strong></td>
</tr>
<tr>
<td><strong>Fluids/Blood Products Administered:</strong></td>
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<tr>
<td><strong>OPERATIVE SITE #1: (specify)</strong></td>
</tr>
<tr>
<td><strong>Procedure</strong></td>
</tr>
<tr>
<td>[ ] Initial amputation (level) ______</td>
</tr>
<tr>
<td>[ ] Revision amputation (level) ______</td>
</tr>
<tr>
<td>[ ] Debridement/Washout Number_______</td>
</tr>
<tr>
<td>[ ] DPC</td>
</tr>
<tr>
<td>[ ] Exam/Dressing change under Anesthesia</td>
</tr>
<tr>
<td>[ ] Ex-Fix (initial)</td>
</tr>
<tr>
<td>[ ] Ex-Fix (revision)</td>
</tr>
<tr>
<td>[ ] ORIF</td>
</tr>
<tr>
<td>[ ] Orthopedic hardware removal</td>
</tr>
<tr>
<td>[ ] Other____________________________</td>
</tr>
<tr>
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<tr>
<td>[ ] None</td>
</tr>
<tr>
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<tr>
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<tr>
<td>[ ] Fungal culture</td>
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<tr>
<td>[ ] Histopathology</td>
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<tr>
<td>[ ] Other____________________________</td>
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### OPERATIVE SITE #2: (specify)

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<td>Total size of wound: _____ cm²</td>
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<td>Revision amputation (level) _____</td>
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<td>DPC</td>
<td>Approx 50% necrotic</td>
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<tr>
<td>Exam/Dressing change under Anesthesia</td>
<td>Approx 75% necrotic</td>
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<tr>
<td>Ex-Fix (initial)</td>
<td>Completely necrotic</td>
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<tr>
<td>Ex-Fix (revision)</td>
<td>Grossly purulent</td>
</tr>
<tr>
<td>ORIF</td>
<td>Gross mold</td>
</tr>
<tr>
<td>Orthopedic hardware removal</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>

**Samples Sent**
- None
- Aerobic culture
- Anaerobic culture
- Fungal culture
- Histopathology
- Other

**Adjunctive Therapy**
- Antimicrobial beads (type) _____
- Dakins soaked dressings
- Dakins Instill Device (started)
- Dakins Instill Device (Renewed)
- Negative pressure therapy
- Other

**Comments:**

### OPERATIVE SITE #3: (specify)

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Wound Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial amputation (level) _____</td>
<td>Total size of wound: _____ cm²</td>
</tr>
<tr>
<td>Revision amputation (level) _____</td>
<td>Clean</td>
</tr>
<tr>
<td>Debridement/Washout Number _____</td>
<td>Approx 25% necrotic</td>
</tr>
<tr>
<td>DPC</td>
<td>Approx 50% necrotic</td>
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<tr>
<td>Exam/Dressing change under Anesthesia</td>
<td>Approx 75% necrotic</td>
</tr>
<tr>
<td>Ex-Fix (initial)</td>
<td>Completely necrotic</td>
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<tr>
<td>Ex-Fix (revision)</td>
<td>Grossly purulent</td>
</tr>
<tr>
<td>ORIF</td>
<td>Gross mold</td>
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<tr>
<td>Orthopedic hardware removal</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>

**Samples Sent**
- None
- Aerobic culture
- Anaerobic culture
- Fungal culture
- Histopathology
- Other

**Adjunctive Therapy**
- Antimicrobial beads (type) _____
- Dakins soaked dressings
- Dakins Instill Device (started)
- Dakins Instill Device (Renewed)
- Negative pressure therapy
- Other

**Comments:**
APPENDIX C: BASTION CLASSIFICATION OF LOWER LIMB INJURY

<table>
<thead>
<tr>
<th>Class of limb injury</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>Injury confined to foot</td>
</tr>
<tr>
<td>2</td>
<td>Injury involving lower leg permitting effective below-knee tourniquet application</td>
</tr>
<tr>
<td>3</td>
<td>Injury involving proximal lower leg or thigh, permitting effective above-knee tourniquet application</td>
</tr>
<tr>
<td>4</td>
<td>Proximal thigh injury, preventing effective tourniquet application</td>
</tr>
<tr>
<td>5</td>
<td>Any injury with buttock involvement</td>
</tr>
</tbody>
</table>
APPENDIX D: ADDITIONAL INFORMATION REGARDING OFF-LABEL USES IN CPGS

PURPOSE

The purpose of this Appendix is to ensure an understanding of DoD policy and practice regarding inclusion in CPGs of “off-label” uses of U.S. Food and Drug Administration (FDA)–approved products. This applies to off-label uses with patients who are armed forces members.

BACKGROUND

Unapproved (i.e., “off-label”) uses of FDA-approved products are extremely common in American medicine and are usually not subject to any special regulations. However, under Federal law, in some circumstances, unapproved uses of approved drugs are subject to FDA regulations governing “investigational new drugs.” These circumstances include such uses as part of clinical trials, and in the military context, command required, unapproved uses. Some command requested unapproved uses may also be subject to special regulations.

ADDITIONAL INFORMATION REGARDING OFF-LABEL USES IN CPGS

The inclusion in CPGs of off-label uses is not a clinical trial, nor is it a command request or requirement. Further, it does not imply that the Military Health System requires that use by DoD health care practitioners or considers it to be the “standard of care.” Rather, the inclusion in CPGs of off-label uses is to inform the clinical judgment of the responsible health care practitioner by providing information regarding potential risks and benefits of treatment alternatives. The decision is for the clinical judgment of the responsible health care practitioner within the practitioner-patient relationship.

ADDITIONAL PROCEDURES

Balanced Discussion

Consistent with this purpose, CPG discussions of off-label uses specifically state that they are uses not approved by the FDA. Further, such discussions are balanced in the presentation of appropriate clinical study data, including any such data that suggest caution in the use of the product and specifically including any FDA-issued warnings.

Quality Assurance Monitoring

With respect to such off-label uses, DoD procedure is to maintain a regular system of quality assurance monitoring of outcomes and known potential adverse events. For this reason, the importance of accurate clinical records is underscored.

Information to Patients

Good clinical practice includes the provision of appropriate information to patients. Each CPG discussing an unusual off-label use will address the issue of information to patients. When practicable, consideration will be given to including in an appendix an appropriate information sheet for distribution to patients, whether before or after use of the product. Information to patients should address in plain language: a) that the use is not approved by the FDA; b) the reasons why a DoD health care practitioner would decide to use the product for this purpose; and c) the potential risks associated with such use.