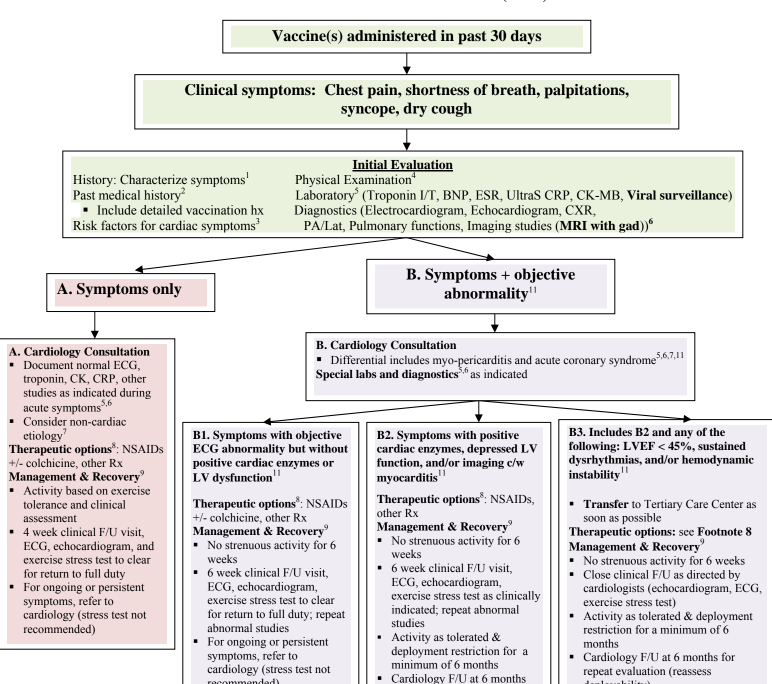
DoD Clinical Guidelines for Post-Smallpox Vaccine Associated Myopericarditis Vaccine Healthcare Centers Network (VHC)



Refer **all** cases to VHC Network for case review, entry into DoD Smallpox Vaccine Myopericarditis Registry, filing of VAERS report and natural history surveillance. With referral include: Patient and provider contact information, Echocardiograms, ECG, cardiac isoenzyme results, & copies of pertinent records.

Consultation: Call the DoD Vaccine Clinical Call Center at 866-210-6469 to request VHC and/or military cardiology clinical consultation.

recommended)

deployability)

FOOTNOTES: Last edited April 2009

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Footnote 1	Characterize symptoms, including chest pain type	Specify symptom location, character, onset, duration, intensity/severity, frequency, accompanying/associated symptoms, and alleviating/aggravating factors. All associated clinical symptoms should be detailed.		
		Categorize patient's chest pain type if present (choose one): 1. Pericarditis chest pain: Chest pain that is typical and made worse by supine position, improved with leaning forward, pleuritic, constant a. Detailed history is critical to case definition of suspect pericarditis – see case definitions, page 5 2. Myocarditis chest pain: angina-like, diffuse; not necessarily positional or pleuritic 3. Atypical chest pain: Pain, pressure, or discomfort in the chest, neck, or arms not clearly exceptional or not otherwise consistent with pain or discomfort of myocardial ischemic origin.		
		Reference:		
		http://www.guideline.gov/summary/summary.aspx?doc_id=6534		
Footnote 2	Assess past medical history	Detailed review of all systems, with attention to the following disorders: Lung disease Gastrointestinal disease Vascular disease (e.g., stroke, transient ischemic attack, peripheral arterial disease) Musculoskeletal disorders (e.g., impingement syndrome, thoracic outlet syndrome)		
		Reference: PMH study guide http://medinfo.ufl.edu/year1/bcs96/clist/history.html		
		Include vaccination history and adverse events (specify site of vaccination and lot number, if available)		
Footnote 3	Risk Factors for Cardiac Symptoms	 Personal History of angina, myocardial infarction (MI), congestive heart failure (CHF), percutaneous coronary intervention (e.g., balloon angioplasty, stent, atherectomy), coronary artery bypass graft (CABG), catheterization with stenosis ≥ 50% Age, sex, race/ethnicity (African American, Mexican American, American Indian, Native Hawaiian, some Asian American), diabetes, hypertension, 		
		smoking, dyslipidemia, family history of CAD (especially prior to age 55), obesity, physical inactivity, stress, and excessive alcohol consumption Reference: http://www.americanheart.org/presenter.jhtml?identifier=4726		
Footnote 4	Physical Examination	Perform a focused PE to include: gender and race/ethnicity, vital signs, ht, wt, detailed exam to include vaccination site, cardiac (jugular venous pressure if able), pulmonary, peripheral edema and lymphadenopathy. Reference: http://medicine.ucsd.edu/clinicalmed/introduction.htm		
Footnote 5	Laboratory studies	Report normal range as defined by individual hospital laboratory standards. Record units and normal range for laboratory. For troponin data, document 99th percentile cut-off for testing system used as well as name of testing system if available.		
Laboratory	Laboratory studies: All patients			
	Complete blood count	CBC at presentation, to include differential, with emphasis on eosinophil and lymphocyte count should be noted.		
	Cardiac enzymes	All Creatinine Kinase (CK), CK-MB, and troponin (I/T) values should be noted.		
	Inflammatory markers	Caution troponin positive based on 99 th percentile of testing system. All erythrocyte sed rate and C-reactive protein (CRP) (ultrasensitive, if available) values should be noted.		
Laboratory	studies as clinically indicated:			
	Immune complex screening	All C3, C4, CH50, Raji cell/C1q assay, and C3D values should be noted.		
	Brain natriuretic peptide	BNP at presentation to assess for heart failure		
	Viral surveillance	Smallpox related myopericarditis is a diagnosis of exclusion. No smallpox vaccine related cases have exhibited viral etiology to date. When considering other etiologies, viral surveillance is indicated.		

Serologies Consider ID consultation; PCR for vaccinia if available (consult CDC/VHC). A coxsackie A/B (enteroviruses), adenovirus, CMV, Parvovirus B19, influenz A/B, HHV-6, HSV-1, HIV, RSV, dengue, echovirus, encephalomyelitis, Epste Barr, Lyme, rhabdovirus, varicella, variola, yellow fever, hepatitis A/B/C IgM, core IgG values and titers during the evaluation should be noted; obtain specim for convalescent titers at 4 week interval. Other Cultures Consider ID consultation; all viral cultures (nasal wash, urine, feces) for adenovinfluenza viruses, parvovirus B19 or enteroviruses should be noted. Note all ANA, Anti-DS DNA, ENA, and similar values during the evaluation. Myocardial biopsy Auto-antibodies for myocardium; special studies, including PCR for vaccinia, parvovirus B19, Request specific assessment for eosinophils
Other Cultures Consider ID consultation; all viral cultures (nasal wash, urine, feces) for adenove influenza viruses, parvovirus B19 or enteroviruses should be noted. Collagen vascular screening Myocardial biopsy Auto-antibodies for myocardium; special studies, including PCR for vaccinia, parvovirus B19, Request specific assessment for eosinophils
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Consult VHC Network working group for updated information
Footnote Diagnostics
Diagnostics: All patients
Electrocardiogram (ECG) Note date, time, rate, rhythm, the presence of ectopy and abnormalities in wave intervals and segments
Typical ECG manifestations: Pericarditis: Acute
 Diffuse ST segment elevation, particularly leads I,II, III, aVF, aVL, a V5-V6 Diffuse PR segment depression
3. PR segment elevation in lead aVR Evolving 1. T-wave changes: notched, biphasic. Or low-voltage inversions.
Myocarditis: 1. Diffuse T-wave inversions without ST segment abnormality 2. Incomplete atrioventricular conduction blocks (usually transient) 3. Intraventricular conduction blocks (usually transient)
*When myocarditis and pericarditis occur together, ST segment abnormalities a may be evident. Reference:
Demangone, D. (2006) ECG manifestations: Noncoronary heart disease. <i>Emerg Medicine Clinics of North America</i> . (24) pp.113-131.
Chest X-ray PA and Lateral
Other diagnostics as clinically indicated:
Echocardiogram If only a range is estimated for ejection fraction (EF), note the midpoint of the r For pericardial effusions, record estimate of size and/or clinical significance" (s effusions may not be diagnostic.
Pulmonary functions With DLCO if indicated; diffusion capacity corrected for hemoglobin is a sensi measure of pulmonary interstitial disease and increased risk for hypoxia with activity.
Stress test Indicate whether an exercise tolerance, stress-echocardiogram, or nuclear/pharmacological stress test was performed during the hospital stay and result of the testing, if performed. Clinical correlation is recommended in the cases of a negative stress test result.
Cardiac catheterization If vessel occlusion identified, note the anatomical region affected and the degre stenosis present.
Holter & Event Monitor Consider for dysrhythmia evaluation
Imaging MRI with gadolinium; consider indium scan for detection of patchy inflammati If not available locally, contact VHC Network
Footnote 7 Consider acute coronary syndrome (myocardial infarction), aortic dissectio pneumothorax, pulmonary embolism, musculoskeletal pain, esophageal dis (gastroesophageal reflux, esophageal spasm), systemic autoimmune disease.
Footnote 8 Consult VHC Network Cardiology Working Group or Recent Literature for upon in treatment options.
Symptoms only (A) OR Non-steroidal anti-inflammatory therapy with or without colchicine (colchicine

	symptoms with chicative	addition to Conventional Therapy for south perioardities Desults of the calchising for		
	symptoms with objective findings, but with negative cardiac enzymes and no LV dysfunction (B1)	addition to Conventional Therapy for acute pericarditis: Results of the colchicine for acute pericarditis (COPE) trial. Imazio M, et al. <i>Circulation</i> 2005; 112:2012-16.)		
	Symptoms w/ positive cardiac enzymes or depressed LV function or imaging c/w myocarditis (B2)	Non-steroidal anti-inflammatory therapy. Other treatments to be considered in consultation with cardiology to include corticosteroid treatment. Consider biopsy for viral PCR, culture and assessment of inflammation (presence of eosinophils). Consider corticosteroids with evidence of eosinophilic inflammation and clinical deterioration		
	Progressive symptoms (LVEF < 45%, sustained dysrhythmias, hemodynamic instability) (B3)	 Conventional heart failure treatments (e.g., ACE inhibitors, nitrates, diuretics, select beta-blockers such as carvedilol or metoprolol succinate) Consider corticosteroids if no evidence of active infection and/or evidence of eosinophils in inflammatory infiltrate. Consider Vaccinia Immune Globulin (VIG)/IVIG only with expert consultant case review via VHC Network. 		
Footnote 9	Management and Recovery	Whenever possible, standardized follow up should occur at or be coordinated with Walter Reed Army Medical Center (WRAMC) or Brooke Army Medical Center (BAMC) in collaboration with VHC Network staff. Deployment restriction reference:		
		Maron et al. Task Force 4: HCM, Other Cardiomyopathies, and Marfan. <i>JACC</i> ;45 (8):1340–5.		
	Symptoms only (A) OR Symptoms with objective findings, but without positive cardiac enzymes or LV dysfunction (B1)	 Light physical activity at own pace for 4 weeks (A) No strenuous activity for 6 weeks (B1) Follow up in 4 weeks (A) to 6 weeks (B1) Asymptomatic at follow-up Repeat any previously abnormal studies Clinical evaluation to include stress test to assess exercise tolerance prior to clearance for return to duty Long-term follow-up will be completed by VHC Network Symptomatic and/or persistent/abnormal findings at follow-up Repeat any previously abnormal studies Clinical evaluation to include stress test (unless contraindicated) Repeat MRI if had previous enhancements or if symptomatic. Repeat at 12-18 months Consult cardiology for further recommendations Long-term follow-up will be completed by VHC Network 		
	Symptoms with positive cardiac enzymes or mild depressed LV function or imaging c/w myocarditis (B2) OR Progressive symptoms (LVEF < 45%, sustained dysrhythmias, hemodynamic instability) (B3)	 No strenuous activity for 6 weeks; deployment restriction for 6 months Clinical evaluation at 6 weeks and 6-12 months Asymptomatic at follow-up Repeat any previously abnormal studies at 6 weeks and 6-12 months Stress test at 6 weeks to assess exercise tolerance for rehabilitation; repeat at 6-12 months to assess exercise tolerance prior to clearance for deployment Long-term follow-up will be completed by VHC Network Symptomatic and/or persistent/abnormal findings at follow-up Clinical evaluation to include enzymes, ultra sensitive CRP, ECG, ECHO, stress test (unless contraindicated) Repeat MRI if had previous enhancements or if symptomatic. Repeat at 6 months to assess exercise tolerance prior to clearance for deployment. Clinical evaluation at 6 months to include repeat ECHO, stress test, and MRI If normal and asymptomatic, clear for deployment If normal MRI with continued symptoms, not cleared for deployment Continue cardiology follow-up at 6-12 month intervals until asymptomatic Long-term follow-up will be completed by VHC Network 		
Footnote 10	Disability Assessment	The majority of patients have recovered within 1 year. The natural history of this condition remains unknown. Careful functional assessment post-acute phase has not yielded definitive objective parameters. The long-term natural history of this condition (e.g., late onset arrhythmias, cardiomyopathy, recurrent myocarditis) has not been well defined. Development of new cardiac complications within 5 years following an episode of hypersensitivity myocarditis associated with immunization should be reported to the VHC Network clinical case management registry.		

Footnote 11	Case Definitions for Myocarditis and Pericarditis (MMWR 2003;52:492-6, www.edc.gov/mmwr/PDF/wk/mm5221.pdf)					
	Objective abnormalities					
Myo- carditis	Suspect (1) Symptoms (dyspnea, palpitations, or chest pain) (2) ECG abnormalities beyond normal variants, not documented previously (ST/T abnormality, paroxysmal supraventricular tachycardia, ventricular tachycardia, atrioventricular block, frequent atrial or ventricular ectopy) OR Focal or diffuse depressed LV function of uncertain age by an imaging study (3) Absence of evidence of any other likely cause	Probable (1) Meets criteria for suspected myocarditis (2) In addition, meets one of the following: Elevated levels of cardiac enzymes (Creatine Kinase-MB fraction, Troponin T or Troponin I), OR new onset of depressed LV function by imaging, OR abnormal imaging consistent with myocarditis (MRI with gadolinium, gallium-67 scanning, anti-myosin antibody scanning)	Confirmed Histopathologic evidence of myocarditis by endomyocardial biopsy or on autopsy.			
Peri- carditis	Suspect (1) Typical chest pain (made worse by supine position, improved with leaning forward, pleuritic, constant) (2) No evidence for alternative cause of such pain	Probable (1) Meets criteria for suspected pericarditis (2) Has one or more of the following: Pericardial rub on auscultation OR ECG with diffuse ST-segment elevations or PR depressions not previously documented OR echocardiogram revealing an abnormal pericardial effusion	Confirmed Histopathologic evidence of pericardial inflammation in pericardial tissue from surgery or autopsy			

Vaccine Healthcare Centers Network 202-782-0411

Web: www.vhcinfo.org
DoD Vaccine Clinical Call Center 1-866-210-6469