Appendix 4: Diphtheria Antitoxin (DAT) Treatment and Adverse Effects Form

Patient ID		Name			
Drug Diphtheria Antitoxin				Date of R	Request Day Year
Diphtheria Antitoxin is currently not licensed in the United States. The National Center for Immunization and Respiratory Diseases of the Centers for Disease Control and Prevention (CDC) is the national center for consultation of suspected diphtheria cases and is responsible for providing diphtheria antitoxin for therapy. CDC has received approval to distribute this product to physicians as an Investigational New Drug (IND) in accordance with requirements of the Food and Drug Administration (FDA). Under the provisions of our IND protocol we must obtain clinical information on each patient who has received DAT. For each patient who develops any serious adverse events (SAE) after DAT administration including those listed in the IND under Section 7.2 Definitions of Adverse Events, please contact the CDC diphtheria duty officer or CDC Emergency Operations Center at (770) 488-7100 within 24 hours of occurrence or as soon as possible (see Section 7.3 Treating Clinician Reporting Requirements to CDC). For any patient receiving DAT, including patients who develop non-serious adverse events (AE) after receiving DAT, please complete and return this form within 14 days of DAT administration or as soon as possible to CDC's Meningitis and Vaccine Preventable Diseases Branch via email at					

This document can be found on the CDC website at: https://www.cdc.gov/diphtheria/dat.html#forms