



Government Clinical Research Services

Committed to delivering fast, efficient, high-quality programs through the end-to-end use of technology & AI

Emmes is a full-service clinical research organization (CRO) dedicated to serving government, biopharmaceutical, non-profit, & academic partners in achieving their development & human health goals.

Founded nearly 50 years ago, we became one of the primary clinical research providers to the US government before expanding into publicprivate partnerships & commercial biopharma. Emmes has built industryleading capabilities in Vaccines & Infectious Diseases, Neuroscience, Ophthalmology, Cell & Gene Therapy & Rare Diseases in the following types of programs.

Statistical & Data Coordinating Centers (SDCCs)

Full Service Clinical Trials

Clinical Research Support Services (CRSS) Partnerships

Clinical Site Monitoring Centers (CSMCs)

Emmes spearheaded the industry's first truly unified electronic clinical platform technology, *Advantage eClinical Cloud*. The difficulty in dealing with numerous systems that do not sync well together led us to create a single application for patients, sites, CROs, & sponsors to conduct clinical research. The unified AI-driven platform delivers insights at scale from study start-up to filing, patient & site engagement, data management, biostatistics, & bioinformatics. Allowing the end user to put their focus back on the next generation of treatments.

Functional Area Expertise

Digital & Decentralized Clinical Trials (DCTs)

Study Design, Site Feasibility, & Protocol Development

Biostatistics, Data Science, & Bioinformatics

Pharmacovigilance & Medical Monitoring

Clinical Data Management

Global Regulatory Affairs

Clinical Site Management & Monitoring

Cloud Systems Design & Engineering

Key Statistics

1,000,000+

Enrolled Subjects

2,150+

Clinical Trials (CT) Completed

400+

Late-Stage CT in Public & Biopharma Sectors

32,000+

CT Sites Engaged Globally

75+

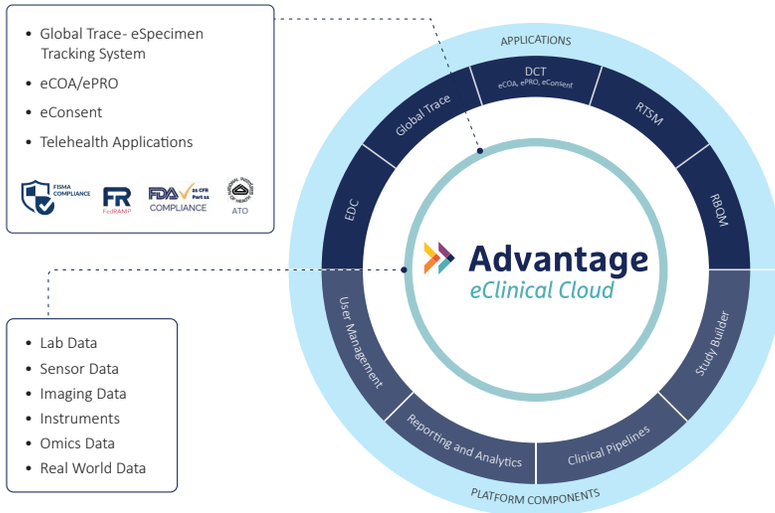
Countries Participating in Clinical Programs

450+

Manuscripts in Leading Peer- Reviewed Journals Since 2016



Advantage eClinical Cloud Unified Platform



One Unified Platform

Log in once to access and navigate across all applications

Single Database

Work with data across the platform without expensive messy integrations

Unified Study Build

Perform study builds in one place for all your applications

Streamlined Role Administration

Manage sites and users across all trials and applications

Active Government Partnerships

Partner with us on the next generation of clinical research

Vaccines & Infectious Diseases



650+ Vaccine & Infectious Disease Clinical Trials

NIAID | Division of Microbiology and Infectious Disease | Statistical Data Coordinating Center: Since 1996, we have supported more than 315+ domestic & international VID protocols as the SDCC for NIAID's DMID extramural program. More than 150 active clinical trials evaluating vaccines & treatment strategies for infectious pathogens, including COVID-19, Ebola, influenza, Zika, Chikungunya, e. coli, shigella, c. difficile, norovirus, & rotavirus, as well as emerging infectious diseases & bioterrorism threats

NIAID | Vaccine Research Center | Data Management, Warehousing, & Clinical Site Monitoring: Since 2001, we have been the clinical data management partner for more than 55 protocols for NIAID's VRC. Provide expert data management & data warehousing support for the development of new vaccines & mAb therapies for emerging infectious disease threats, including HIV, malaria, West Nile virus, Chikungunya virus, zika virus, Severe Acute Respiratory Syndrome (SARS), Ebola, Marburg, smallpox, RSV, & influenza (avian, seasonal, H1N1 & H5N1 pandemic influenza virus & universal influenza vaccine strategies)

Cell & Gene Therapy & Rare Diseases



125+ Cell & Gene Therapy Clinical Trials & 290+ Rare Projects

NHLBI | Blood & Marrow Transplantation Clinical Trials Network: Established in 2001 to conduct large multi-institutional clinical trials aimed at addressing issues in hematopoietic stem cell transplantation. Since then, over 10,000 patients have enrolled in 46 protocols from 150 global centers

NHLBI | CureSci: Emmes provides the management & coordination of the Sickle Cell Data Collection (SCDC) program, working cooperatively with NHLBI & patients, families, advocates, physicians, industry, & researchers

Neuroscience & Ophthalmology



180+ Neuroscience & 290+ Ophthalmology Clinical Trials

NIDA | Clinical Coordinating Center & Data & Statistics Center: Emmes has been the CCC for NIDA's Clinical Trials Network (CTN) since 2005, supporting 60+ protocols in substance use disorders. Emmes has also been the NIDA CTN DSC since 2009, supporting 50+ protocols to date

NIMH | NeuroAIDS Tissue Consortium (NNTC) Data Coordinating Center: Since 2003, Emmes has supported NIMH's unique multi-site repository that collects & distributes well-characterized antemortem & postmortem tissue specimens with associated clinical & serological data from HIV-infected individuals

NEI | National Eye Institute Support: Emmes has served as the NEI contractor since 2005 with 160+ protocols for their clinical trial program

Maternal & Pediatrics



230+ Maternal & Pediatric Projects

NICHD | Best Pharmaceuticals for Children Act Data Coordinating Center: Since 2009, Emmes supported more than 44 domestic protocols & as the DCC for the Pediatric Trials Network via the BPCA. Emmes supports the expeditious translation of quality pediatric research into scientifically supported Clinical Study Reports that allow the FDA to improve pediatric labeling

NAICS CODES

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BUSINESS SIZE

LARGE

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