Government and Public Health Services

Since 1990, PPD has provided clinical, regulatory and quality support to government and public health clients around the world. These research programs require a diversity of full medical product development and clinical support services, including full-service clinical trials, regulatory support contracts, clinical site monitoring contracts, statistical and data coordinating centers (SDCCs) and clinical research operations and management support (CROMS) contracts. Our government and public health services group (GPHS) is a small, nimble, project-centric team of more than 150 dedicated personnel that benefits from access to our global organization.

PPD PROVIDES:

• Programmatic resources for broad portfolios or full clinical program capabilities
• Multiple contracting and partnering models
• Deep experience in navigating and upholding all Federal contract requirements as a prime contract holder and subcontractor
• A DCAA-compliant accounting system
• Government program management expertise, including Earned Value Management

HOW OUR GOVERNMENT AND PUBLIC HEALTH EXPERTISE WILL ENSURE SUCCESS OF YOUR PIPELINE

1 Global Experienced and Passionate Workforce

- In the past 5 years:
  - 2,500+ trials in 100+ countries with 30,000+ dedicated team members

2 Patient and Caregiver Focus

- Patient advocacy groups support
  - Vaccine priority network of 1,000 sites
  - 300+ precontracted oncology research sites for rapid activation
  - Pediatric groups
  - Phase I units
  - Training program for capacity building at site and general population education
  - Up to 30% faster study startup and up to 10% faster study completion

3 Innovative Design and Data-driven Analysis

- Database of relevant patients
- Client Operations Dashboard

- Virtual trials
- Digital platforms
- Data visualization
- Time-saving data access
- Adaptive protocol design
- Risk-based monitoring

• 1,515+ government-funded clinical trials, including 1,067 infectious disease and vaccines trials, across NIH, CDC, BARDA, and the DoD

• Full-service support for NIAID/DAIDS-funded ACTIV-2 COVID-19 Outpatient Platform Trial

• Multi-service support of the NIAID/DCR-funded ACTIV-3/3b COVID-19 Inpatient Platform Trials

• 1 of 4 CROs providing global, full-service, and rapid response support for all parts of NIAID.

• Providing global, multi-service support for the CDC’s Tuberculosis Trials Consortium (TBTC)
Our industry-leading scientific and medical teams design, plan and implement full-scale global programs to help clients reach key milestones. They develop and provide counsel on feasibility and protocol optimization across a range of therapeutic areas to drive patient and site enrollment and provide expertise in the conduct and management of clinical studies.

This expertise results in an understanding of the current challenges in the landscape, as well as established strategies and processes to assure efficient and cost-effective drug, device, and evidence-based care development.