Home Trial Services
Enhance your trial. Improve their experience.

Reduce the patient burden and improve recruitment, engagement, and retention with home trial visits.

Bringing protocol visits to your patients—whether at home or another convenient, non-traditional location—makes it easier for them to participate in your clinical trial. Firma’s home trial visits help speed recruitment and increase retention for just about any clinical trial, not just elderly, pediatric caregiver-intense, and orphan-disease or rare-disease populations.

GLOBAL NETWORK OF HIGHLY SKILLED CLINICIANS
Our network of licensed, highly skilled, and clinical trial experienced home trial service providers (HTP) are trained on your protocol and study requirements, with their training documented in our electronic Learning Management System (eLMS). Firma staff can conduct protocol-specified activities such as:

- Biological Sample Collection
- Blood Draws
- Vital Signs
- Changes in Health and Medication Assessments
- Study Drug Administration
- EKG Administration
- Questionnaires
- Patient and Caregiver Training
- Chaperone Services
- AND MORE

99% EXECUTION RATE WITHIN PROTOCOL TIME FRAME

15,000+ HOME TRIAL VISITS COMPLETED SINCE 2015

2,000+ HEALTHCARE PROVIDERS IN DATABASE

45+ COUNTRIES COVERED BY OUR SERVICE

firmaclinicalresearch.com
DRIVEN BY TRANSPARENCY AND COMMUNICATION
Firma believes in full transparency. We start your study off right by determining the reporting you and your sites require, including the targeted KPIs that will best measure ongoing performance.

When assigned, the staff members’ CVs, licensure, and training certifications are sent to the PI/site for review and approval prior to the conduct of any patient visit. This sets us apart from many organizations and results in a much more successful engagement with study sites and principal investigators.

Weekly status updates through a custom portal ensure you always know the status of the home trial visits throughout the trial, enabling you to track patient engagement and achieve operational efficiency.

CUSTOM STRATEGIES = SEAMLESS TRIAL INTEGRATION
Enhance your patient-centric approach in clinical trials with our flexible and creative solutions that increase high-quality recruitment, support, and retention through:

- Real-time status updates and KPIs for clients
- 24-hour turnaround on document QC through electronic source document implementation
- Technology investment to support efficient and high-quality conduct, including eLMS, HTMS, electronic source visit document collection, and DocuSign
- Corporate alignment to the EU General Data Protection Regulation (GDPR)

“Our trials are quite complex and we have relied heavily on Firma to draw patient labs and ship them overnight to us at study lab sites. Several patients are international, so it is not a small accomplishment to get the labs to us overnight and in time for appropriate safety and efficacy monitoring. We simply could not carry out our studies without the remarkable dedication and professionalism of Firma.”

— Principal Investigator, large academic medical center