

Experience a new perspective on research... Experience our research dedicated facility

New Phase Research & Development is a versatile clinical research institute located in Knoxville, Tennessee. With a focus on quality data, outstanding subject care, and active recruitment, New Phase has become a leader in successful clinical trials at a site level.

We currently operate as a fully dedicated clinical research facility in West Knoxville. With an expansive network of patient referrals and an in-house clinical research database, we are able to assess feasibility of potential trials as well as accurately predict recruitment potential and set realistic goals.

We invite you to explore New Phase as a potential partner and site for your next trial.

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Welcome!

Our dedicated clinical research facility is located in the heart of West Knoxville and is focused on the successful completion of clinical trials in an array of primary care indications. With a staff of experienced clinical research professionals including 2 research dedicated Principal Investigators, New Phase stands out above the rest in trial coordination and Principal Investigator involvement.

We do not focus on trial volume but instead we carefully select each trial for our facility and focus on enrollment and performance. Our full-time research team consists of dedicated professionals who work together to achieve one common goal; successful trials.

All team members are annually trained in GCP/ICH guidelines as well as attend bi-monthly workshops for in-depth training on various aspects of clinical research as well as new industry standards.



Principal Investigators

Dr. Evelyne Davidson, MDPrincipal Investigator



Dr. Natalie Clarke, MDPrincipal Investigator



New Phase is proud to have Dr. Evelyne Davidson as our research dedicated PI who oversees all trials at the West Knoxville facility.Dr. Davidson is a board eligible Internist with over 30 years of experience in patient care and management. Dr. Davidson has extensive experience in most primary care indications and is actively involved in the care of every clinical research subject. Dr. Davidson not only performs all protocol dictated examinations, but meets with each subject at every visit. Dr. Davidson is present in all informed consent discussions and personally evaluates all inclusion/exclusion criteria. Unlike most Principal Investigators, Dr. Davidson not only provides oversight, but also has an active role in the day to day coordination of each trial.

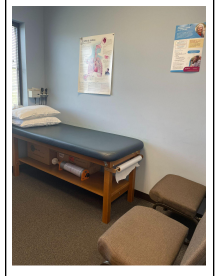
Dr. Clarke began working at New Phase Research & Development in March 2016. She graduated from Penn State University with a Bachelor of Science degree in Biology in 1996. After college, she participated in two years of research investigating the effects of various vasodilators on atherosclerotic arteries. From 1998-2002, she attended The Medical College of Georgia, earning her MD and completed her anesthesiology residency in 2006. Dr. Clarke moved from Georgia to Knoxville in 2010. While working at New Phase, Dr. Clarke has been involved in several studies including ones pertaining to diabetic neuropathy, IBS, COPD, asthma, rheumatoid arthritis, lupus and chronic low back pain.

Clinical Research Coordinators

Our dedicated research site is staffed with 6 full-time Clinical Research Coordinators and 1 Sub-Investigator, who work alongside Dr. Davidson and Dr. Clarke to move each trial in the right direction. With over 25 years of nursing and research experience, our Coordinators are some of the best in the industry.

The Coordination staff consists of one FNP-C, two CNAs, a BS in Health Science, and one LPN coordinator. Our Clinical Research Coordinators are assigned a maximum of three clinical trials each.

This allows each Coordinator to fully dedicate their time to each trial they undertake, while providing their subjects the time and follow-up they truly need.



Our staff has extensive experience with most EDC systems, including Medidata Rave, Oracle, Inform, and Datalabs. This experience, coupled with our site SOP of a 48hr data entry turnaround, has proven to be a highly successful combination in timely and accurate data entry.

Our Coordinators function in a team focused environment with 24hr access to the Principal Investigator and Clinical Research Director.

Our team atmosphere and open communication allows us to provide cohesive and consistent care to our subjects as well as accurate and detailed data collection.

Our Coordinators are all IV certified and IATA trained/certified. We have all necessary lab equipment including refrigerated centrifuge, 20F and -70F freezers, EKGs, and immediate access to dry ice.

Our Facility

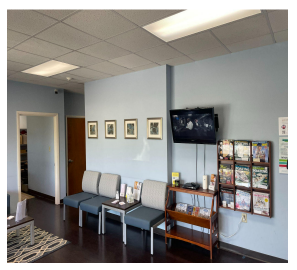
We have a secured double locked drug closet equipped with pin code access and deadbolt. We also maintain a drug storage refrigerator that is independently locked. Both the storage area and refrigerator are equipped with a digital glycol controlled min/max thermometer. These monitors are set to alarm staff of any deviations from the set range. Designated site staff maintain daily drug logs.



We maintain an active emergency action plan and are located approximately 6 miles

from the nearest hospital. We are equipped with a fully stocked crash cart that includes oxygen and an AED.

All staff members are AED and CPR trained. A private office is dedicated to onsite CRA visits. This office contains access to wireless internet, telephone, copier and fax machine.



Our subjects have the comfort of their own comfortable lounge area equipped with a LCD television. This provides extra comfort for extended subject visits in which we provide meals.

Our Patients

Our current subject database consists of over 10,000 subjects who are interested in, or have participated in clinical trials with New Phase Research & Development. We are able to search and create customized database lists to include diagnosis, current medications, and lab results. This allows us to accurately predict our enrollment for each potential protocol.

We utilize a variety of standard local advertising campaigns including print, digital targeting, radio, and television. We also attend local health expos, county fairs, and city events as a means to identify new subjects. Our relationships in the local medical community have been and still remain vital to our site success. We have developed very successful referral relationships with multiple specialties in Knoxville and the surrounding area. This allows us to reach our two groups of subjects we otherwise would not have direct access to.

We have a dedicated, full time team of recruiters who are specialized in patient enrollment. We also have an internal marketing and advertising team. This allows New Phase Research & Development to efficiently maximize enrollment and meet goals set by each sponsor.

Contact New Phase Research and Development
We are readily available to answer all questions, as well as provide
feasibility reports, for any potential protocols currently looking for sites.

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