Lovelace Scientific Resources (LSR)

Established in 1987, LSR has conducted over 1000 multi-disciplinary clinical trials. We are passionate about our work and put patient safety and health at the forefront of our daily business. A significant strength of our research team is the participation of experienced Investigators who are physicians in community-based private practices or employed by LSR.

Customer expectations are met through experience and excellence in performing clinical trials. We obtain the highest quality data, provide effective patient recruitment and retention, and maintain flexibility and responsiveness to meet the needs of our customers.

**Site Locations**
- Albuquerque, NM
- Austin, TX
- Sarasota, FL
- Venice, FL

**LSR Capabilities**
- Own and Manage each Individual Site
- Conduct Multi-disciplinary Phase I-IV Trials

**Study Personnel**

**LSR Employees:**
- Physician Investigators (Employed by LSR and/or Contracted)
- Site Directors
- Registered Research Nurses and Certified Study Coordinators
- Regulatory Coordinators
- Research Assistants & Technicians
- Administrative Staff

**Central Services**
- Dedicated Patient Recruiters
- Research Database
- Contracts and Budget Department
- Marketing Services

**Frequently Asked Questions**

**Who are LSR’s customers?** LSR consistently works with pharmaceutical companies, CROs, biotechnical, and biomedical companies to bring new treatments, devices or procedures to the medical community and the general public.

**Description of LSR facilities?** LSR sites are in close proximity and affiliated with physician offices and medical centers. Physicians in these practice areas represent multidisciplinary specialties. LSR sites have the necessary equipment required to conduct simple-to-complex clinical trials, along with dedicated monitor space and access to ancillary services not available at the site.

**Who negotiates the clinical trial agreement?** Our Office of Research Contracts (ORC) is a central core service that manages Clinical Trial Agreements and budgets for all LSR locations.

**Who processes regulatory documents?** Dedicated regulatory coordinators manage initial submissions and maintenance of the regulatory documents. LSR typically utilizes Central IRBs but can work with a local IRB when required.

**How are patients recruited?** Physician and research site databases, community outreach and advertising. Dedicated recruiters identify patients to enhance study enrollment.

**Where are study records stored?** Study related documents are archived at the LSR corporate medical records facility.

To contact us please visit us online at [www.lsrtrials.com](http://www.lsrtrials.com), Google or Facebook!

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