Nonalcoholic Steatohepatitis (NASH)
Drug Development Solutions

ProSciento provides scientific and operational expertise in the design and management of all aspects of NASH clinical trials. Strengthened by deep therapeutic area knowledge and experience, ProSciento offers highly customized clinical R&D programs for clients globally, addressing the complexities of biomarker selection, access to eligible patients, and multinational clinical trial site access to specialized methodologies.

Distinct Challenges in NASH Clinical Drug Development

The development pathway for NASH drugs is complex despite recent advances in the field. Distinct challenges for NASH clinical research include:

- Few validated biomarkers and endpoints and uncertain regulatory pathways for drug approval
- Crowded development space with competition for clinical trial subjects and sites for NASH clinical trials
- Limited number of clinical trial sites with access to specialized methodologies and relevant NASH experience.

Science-Driven Strategies to Address Challenges in NASH Clinical Research

Targeted Biomarker Selection and Study Design

Strengthened by 15+ years in clinical R&D focused on metabolic diseases, ProSciento’s scientific services team has deep expertise in selecting and deploying proven biomarkers and complementary advanced methodologies for NASH clinical research. We assist clients in designing studies utilizing biomarker insights for specific metabolic pathways, including, but not limited to, glucose homeostasis, lipid metabolism, inflammation, fibrosis and liver dysfunction. The services and expertise ProSciento provides to clients are further bolstered by scientific partnerships, including a NASH advisory board bringing together renowned scientists from leading academic institutions, and technology partnerships, including innovators in advanced imaging and circulating biomarkers.

Patient Access and Enrichment Strategies

ProSciento provides unique, effective solutions to considerably improve patient access and enrichment in the enrollment phase of NASH clinical trials. One such solution is ProSciento’s patient pre-screening methodology to identify patients with steatosis or steatohepatitis prior to imaging techniques or biopsies. The noninvasive methodology utilizes vibration controlled transient elastography (VCTE) FibroScan, proprietary clinical algorithms, and state-of-the-art metabolomic testing. The outcome is a wider number of eligible patients and detailed enrichment data for the enrollment phase of a NASH clinical trial, and far fewer screen failures during expensive imaging procedures.
Science-Driven Clinical Trial Design and Management

A significant challenge in NASH clinical research is the limited number of sites with access to complex methods and tools, including MRI-PDFF, MRE and biopsy capabilities. ProSciento’s hub & spokes clinical trial management platform solves this challenge by directly establishing and managing site access to specialized methods. What is unique to ProSciento’s platform is the integration of its specialized method hubs and global network of clinical trial sites with operational and scientific acuity established from nearly a decade of experience in NASH and 15 years as a clinical research center of excellence for metabolic diseases.

Key Components Include:

HUBS – Centers with Specialized Methodologies
ProSciento has a global network of 45+ method hubs, which are specialized centers with capabilities in liver biopsies and advanced imaging, including MRI-PDFF, MRE and VCTE FibroScan.

SPOKES – Qualified Clinical Trial Sites Near Each Hub
ProSciento manages the relationship between its global network of 200+ clinical trial sites and its network of specialized method hubs. To ensure efficiencies for complex multi-site NASH studies, clinical trial sites are selected within geographic proximity of pre-qualified method hubs. This removes the burden from individual sites to procure access to specialized methods. This also broadens the number of sites that can participate in a study and increases access to patients to participate in clinical trials.

ProSciento’s Approach for Highly Effective NASH Clinical Research Programs

- **Study Design & Biomarker Utilization**: Science-driven selection and utilization of biomarkers and methods
- **NASH patient access**: Focused pre-screening and patient enrichment strategies
- **NASH hub & spokes**: Multi-site trial management, including site access to specialized methods
- **Study Conduct & Data Analysis**: Full-scope CRO services, leveraging 15+ years of metabolic experience

Contact us at bd@prosciento.com to discuss your NASH drug development program.