



***Paving the way from
Trial to Treatment***

Who We Are



Frontier Clinical Research, LLC is a comprehensive Site Management Organization (SMO) that partners with local physicians to deliver high quality study conduct and drug/device development services.



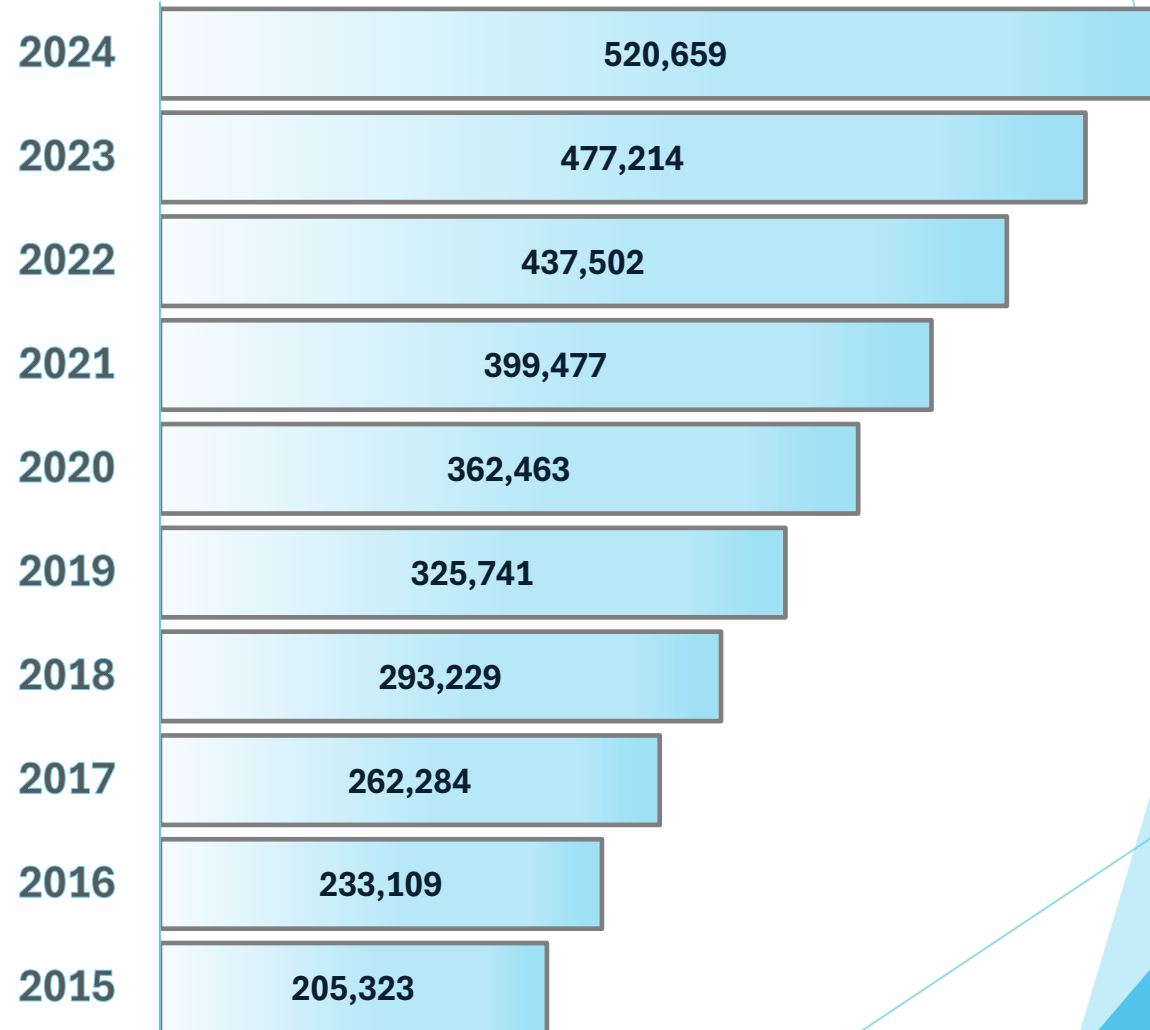
We have an enviable record of having provided services to many of the top pharmaceutical and biotechnology companies, both large and small.



Our clinical team has nearly 25 years of combined clinical research experience.

**In the last 10 years,
the number of
registered clinical
trials has more than
doubled!
(153% increase)**

Registered Clinical Trials by Year 2015-2024



How Can You Benefit From Research?



Contributing to groundbreaking medical advancement



Provide cutting-edge therapies previously unavailable to patient populations



Supplement income from Research Activities and space leasing



Research Staff with track record of clinical expertise provided by Frontier



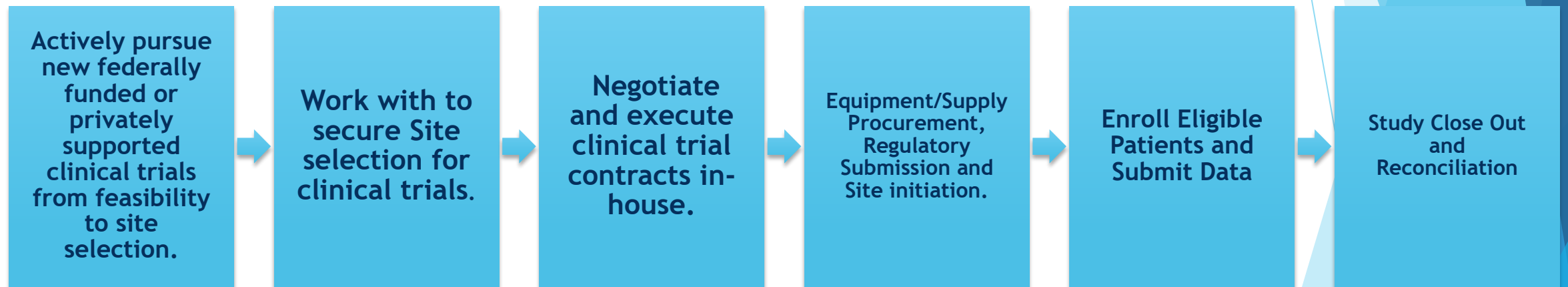
Minimal administrative burden to Clinic Staff and Physicians



No upfront cost to the Investigator

The Clinical Trial Process

Frontier Clinical Research, LLC, provides clinical trial expertise through the entire clinical trial:



Capabilities

Frontier Clinical Research has a highly qualified and well-trained staff with a track record of meeting and/or exceeding enrollment, adhering to protocols and completing studies on time. We have quick turn-around times for initial documents (CDA, CTA, and budget), Central IRB approval, recruiting and start-up. Our thorough and accurate documentation has resulted in low query rates and favorable audits. We have extensive outpatient Phase II-IV trial experience in areas including proof-of-principle, devices, PK/Absorption, safety, efficacy and tolerability.

Top Enrolling Sites for the Following Indications:

Anemia
Asthma
COVID-19
Diabetes

Influenza
Migraines
RSV
Vaccines

C. Diff
Constipation
Colorectal Cancer
Esophagitis

Frontier has conducted studies for many of the top pharma companies worldwide



* Partner-in-Care

Approved Study Medications & Devices

The research being conducted today becomes tomorrow's life-changing and life-saving treatments. The data collected from the patients enrolled in Frontier's research study projects have led to the success of the following FDA drug & device approvals:

- ▶ Alder: *AIMOVIG* - Treatment of Migraines
- ▶ Astellas: *VEOZAH* - Treatment of VMS
- ▶ Astra Zeneca: *TUDORZA* - Treatment of COPD
- ▶ Biocryst: *PERAMIVIR* - Treatment of Influenza
- ▶ DermaSensor - Skin Cancer Screening Device
- ▶ Eli Lilly: *FOUNDAYO* - Weight Loss Treatment
- ▶ GlaxoSmithKlein: *BLUJEPA* - Treatment of Urinary Tract Infection
- ▶ Janssen: *TREMFYA* - Treatment of Ulcerative Colitis
- ▶ Novo: *TRESIBA* - Treatment of Diabetes
- ▶ Shionogi: *XOFLUZA* - Treatment of Influenza
- ▶ Sanofi: *SOLQUA* - Treatment of Type 2 Diabetes
- ▶ Synergy: *TRULANCE* - Treatment of Idiopathic Constipation or IBS-c
- ▶ Vibrant Gastro: *VIBRANT* - Treatment of Constipation

Our Principal Investigators

Marcy Goise, MD
Family Medicine

Tiffany Pluto, DO
Family Medicine

Marc Happe, DO
Gastroenterology

Ward Paine, MD
Internal Medicine

Jeffry Pilney, MD
Family Medicine

Kevin Blankenship, MD
Emergency Medicine

Our Sub-Investigators

Charles Calabrese, DO
Gastroenterology

Alyssa Lorenze, MD
Gastroenterology

Bchara Janadri, MD
Pediatrics

Wilbur Sine, MD
Psychiatry/Family
Medicine

Erica Murray, PA-C
Emergency
Medicine

Rod Ali Hojat, MD
OB/GYN

Robin Goodwin, FNP
Family Medicine

Challenges to clinical trials

- ▶ Takes an average of 31 weeks / 7 months from Site Identification to Site initiation
- ▶ 11% of sites selected are never activated due to their inability to enroll subjects
- ▶ Poor site selection can increase costs by 20% causing delay of study
- ▶ 80% of clinical trials fail to meet enrollment deadlines
- ▶ 54% of new investigators are never used again due to poor enrollment or poor data quality

Clinical research development Partnering for success

- ▶ In an era of decreased margins, sponsors are looking to partner with research sites who can quickly and efficiently **recruit patients** and **produce quality data**.
- ▶ Aligning your sites enrollment with the top 30% of clinical research sites cannot be understated. It is the single most important factor in determining the success of your clinical research program.

Questions

- ▶ What questions do you have regarding research?
- ▶ What questions do you have regarding Frontier Clinical Research?

Please reach out to Frontier's Business Development Manager, Tancredi Calabrese

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We look forward to partnering with your clinical practice and building its clinical research capabilities to their utmost potential.