Accelerate & Diversify Your Dermatology Program

With a robust database of prescreened participants across seven fully integrated sites, our team stands ready to meet your enrollment timelines and key study milestones.



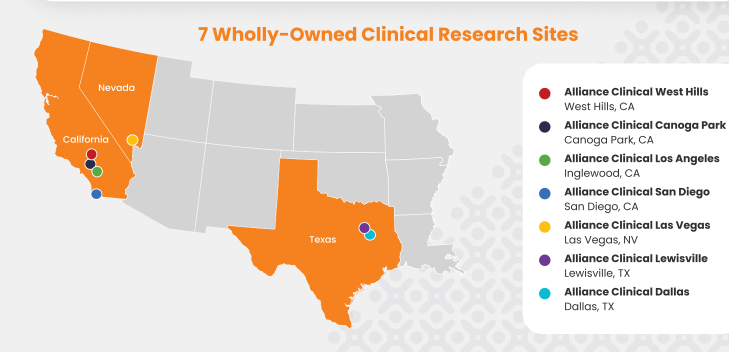
>250,000 database of consented, prescreened participants



85% of database from underrepresented populations

Proven Expertise Across a Broad Range of Dermatology Indications

Atopic Dermatitis	Rosacea	Alopecia Areata	Skin Barriers
Acne	Psoriatic Arthritis	Skin Hydration	Topicals





Three Keys to a Successful Dermatology Study

Rapid Study Startup. Audit-Ready Data. Diverse Participant Population.

Rapid Study Startup Enables Brisk Enrollment

We begin with a proprietary database of >250,000 prescreened participants. Harnessing technology, we then quickly identify and contact appropriate candidates across our sites, proactively scheduling them in anticipation of site activation. This accelerates time to FPI.

> In a recent atopic dermatitis study, our sites rapidly achieved their study caps ahead of schedule – with screen failure rates well below the industry average.

Comprehensive Focus on Quality Produces Audit-Ready Data

We use a streamlined visit management process that allows us to efficiently consent and treat participants with minimal downtime and maximum quality. We bolster that with:

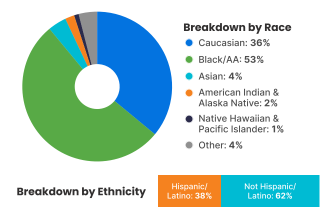
- · Overnight quality control on 100% of visits
- Automated eSource exception reporting
- Ongoing third-party internal audits

These tactics—implemented across all projects —enable us to catch and correct issues in near real time. The result: conclusive, audit-ready data for your clinical study.



Intentional Design Delivers Diverse Participant—and Staff—Population

Establishing purpose-built study sites in communities and cities with underrepresented residents, Alliance Clinical places a deliberate focus on helping ensure that innovative new therapies are effective across populations.



About Alliance Clinical Network

Alliance Clinical's fully integrated, wholly owned and operated clinical research sites help advance human health by delivering exceptional clinical trial data gathered from a proprietary database of >250,000 consented participants. Our centralized management harmonizes processes and systems to deliver consistent study execution across our seven sites—a solid foundation from which to meet your study needs.

To learn more, please contact us.



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