

PROACTIVE DRUG DEVELOPMENT

Integrated CRO and CDMO Services

Saving up to **40%** in drug development time from lead candidate selection to clinical proof of concept, and beyond **IMAGINE** having your Investigational Brochure updated while your IND-enabling studies are in progress.

IMAGINE starting first-in-human trials sooner by timing small-scale drug formulations with clinical conduct.

IMAGINE the ability to plan your clinical program while your preclinical safety assessment is ongoing so that you can initiate your first-in-human trial as soon as you receive regulatory approval. **IMAGINE** timed bioanalytical analysis and safety data.

IMAGINE swift decision-making and reduced hand-offs with a dedicated, cross-functional program manager as your single point of contact throughout the lifecycle of your project.

At **Altasciences**, we do more than **IMAGINE**, we **anticipate** the needs of your project from the get-go.

HOW DOES PROACTIVE DRUG DEVELOPMENT WORK?

At Altasciences, we provide you with clear, customized roadmaps, supported by our real-time data generation, proprietary communication platforms, and central program management and scheduling.

Altasciences' **Proactive Drug Development Solution** is based on three core pillars:

- Tell Us Once™
- Ø.T.L.Ø.S.™
- A unique organizational structure

Together, these pillars translate into a results-driven exchange of information, up to 40% reduction in drug development timelines, and cost savings.

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TELL US ONCE™

ONE COMPANY ONE SYNCHRONIZED SOLUTION

Saving 25% to 40% in development time from lead candidate selection to clinical proof of concept

CDMO

comprehensive manufacturing and analytical solutions

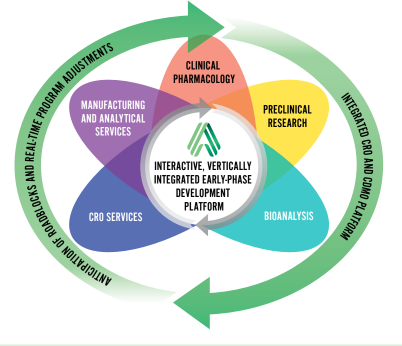


Tell Us Once™ is Altasciences' commitment to communication. Ask Albert is the proprietary database behind the commitment, that fuels integration across departments.

- Ask Albert—database collects and shares client-specific data, preferences, drug information, and results across internal teams, regardless of department or location.
- As a molecule advances, so does our clients' information.
- We eliminate the need for clients to repeat themselves at each stage of development or throughout a single study.

Sponsors only need to tell their story once, and we'll take care of the rest.





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We anticipate program-specific roadblocks and streamline an integrated approach to CRO and CDMO services under one program manager and one operational structure using a centralized scheduling system.

- Customized roadmap for our client's drug development program
- Dedicated cross-functional project manager
- Single centralized scheduling system
- Active timeline management
- Real-time responses to challenges that may arise, allowing proactive solutions to be applied.

Our unique structure—a grassroots level of integration with two executives leading all scientific and operational teams, eliminating the internal silos that can impact your timelines. This group of forward-thinkers and scientific experts become extensions of your team, dedicated to advancing your studies.

Scientific and operational teams are led by two executives:

- One executive covers the preclinical and clinical research.
- One executive is responsible for study design, reporting, bioanalysis, and manufacturing.



At Altasciences, we do more than **IMAGINE**. You can count on us to anticipate the needs of your project from the start, to seamlessly deliver timely, quality results.

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