

# **Study Startup Support**

## **Efficiencies to Help You Meet Critical Milestones**

As a key element of study startup, IRB review can have a substantial impact on meeting critical study milestones. Advarra<sup>®</sup> provides flexible solutions to help accelerate the initial IRB review process and make it more efficient.

### Shorten Timelines and Enhance Study Success



#### Site Identification and Feasibility Support

Use site performance metrics from completed studies to identify quality, high performing sites through Advarra's SitelQ service.



#### Institutional Site Identification

Request our list of over 3,500 institutional sites that have worked with Advarra and/or have master agreements with us.



#### **Rapid Site Activation**

Onboard sites in less than one day and shrink total IRB review and approval time with Advarra's IRB-Ready<sup>®</sup> service.



#### **Project Management**

Manage resources, streamline information and meet IRB deadlines with the help of a client advocate; as your primary point of contact, your advocate understands your organization's unique needs.



#### **eTMF** Automation

Replace inefficient and error-prone manual processes with Advarra Connect<sup>®</sup>, Advarra's process for secure, direct transfer of information from the Advarra Center for IRB Intelligence (CIRBI) Platform to your eTMF.





#### **Proactive Budget Management**

Monitor your study's financial needs with custom spend-to-date reports and email notifications when nearing your specific budget ceilings.



#### **Real-Time Study Information and Reports**

Stay up to date on the IRB review process with real-time data from the Advarra CIRBI Platform. No need to search through your emails — IRB communications are managed through your CIRBI account.



#### **Draft Review and Advisory Opinion**

Submit a draft protocol for full board review and increase the likelihood of a smooth, efficient review with your formal study submission.



#### Informed Consent Template

Establish an IRB-approved template with the standard regulatory requirements and your organizational requisites so that you'll only need to update study-specific information with each new submission.



#### **Consulting Services**

Advarra Consulting offers a wide range of consulting services including GxP auditing, quality assurance, pharmacovigilance and REMS and HRP/IRB support. Audits, mock inspections, training and more can help ensure study compliance and success.



#### **IBC Support**

For research involving recombinant DNA, synthetic nucleic acid molecules, genetic engineering or gene therapy, Advarra's institutional biosafety committee (IBC) services can provide expert guidance and support to help you comply with NIH Guidelines. With coordinated IRB and IBC reviews, you don't have to wait for one committee to complete its review before the other can begin.

