

Office of Human Research Protection Programme (OHRPP) Post-Its:  
*Bringing you the latest updates on research policies, educational resources and event information*

## Education & Training

**Registration now OPEN!**

### **NHG PCR 001 Subject Recruitment & Informed Consent**

NHG PCR 001 Subject Recruitment and Informed Consent is an **enhanced** online course to equip Investigators, Clinical Research Coordinators, Research Assistants and other study team members with knowledge on:

Ethical and local regulatory requirements for subject recruitment and informed consent

How to plan, design and conduct the subject recruitment and consent process

Best practices and tools to maintain proper documentation

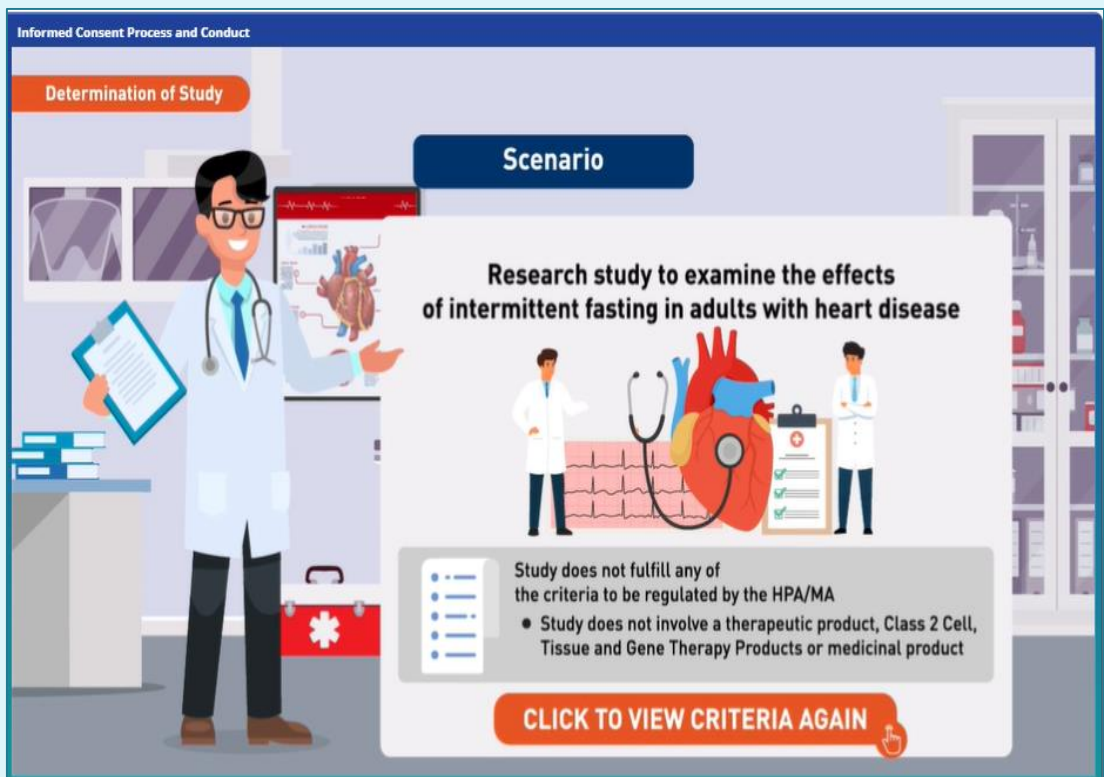
Common errors in the subject recruitment and informed consent process

### **What can you expect from NHG PCR 001 Subject Recruitment & Informed Consent course?**

**Interactive Guides on Subject Recruitment + Informed Consent Requirements For Research + Consent Scenarios**

Learn about **Remote Consent. What is it?** When can **Remote Consent** be used in research?

**Waiver of Consent (WOC)** How can your studies meet WOC criteria? How do you address IRB concerns?



**Informed Consent Process and Conduct**

**Determination of Study**

**Scenario**

Research study to examine the effects of intermittent fasting in adults with heart disease

Study does not fulfill any of the criteria to be regulated by the HPA/MA

- Study does not involve a therapeutic product, Class 2 Cell, Tissue and Gene Therapy Products or medicinal product

**CLICK TO VIEW CRITERIA AGAIN**

#### **TO REGISTER**

Visit <https://www.research.nhg.com.sg> > Training & Education > Register for Courses & Other Events > **NHG PCR 001**

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