INTRODUCTION

The Human Biomedical Research Act (HBRA) came into force on 1 November 2017. As clinical research regulated by the new HBRA regulations makes up approximately 95% of all regulated research in KK Women’s and Children’s Hospital (KKH), KK Research Centre (KKRC) seeks to reduce the risks of research non-compliance through research internal audits.

OBJECTIVES

The audit framework aims to increase oversight of clinical research in KKH through a broad-based study selection using a simple dynamic algorithm based on risk evaluation. We hope that the framework will ensure regulatory compliance and lead to an overall enhancement of research quality at KKH.

METHODOLOGY

Phase 1 – Preparation Work
- Familiarised with HBRA regulations
- Obtained reference from Health Sciences Authority Good Clinical Practice Inspection Framework

Phase 2 – Identification of Areas to Augment Research Compliance and Quality
- Collaborated with SingHealth Office of Research, Integrity and Compliance (ORIC)
- Identified areas where compliance and quality can be augmented

Phase 3 – Development of Research Quality Audit Framework
- KKRC defined and established audit framework scope, study selection process and audit conduct workflow

Phase 4 – Seeking Medical Board’s Approval for Implementation
- Obtained approval by KKH Chairman, Medical Board in December 2018

RESULTS

- The KKH Research Quality Audit Framework was launched in January 2019 and a hospital policy has been developed.
- Research studies for audit were selected based on a risk-based approach:
  - Level 1: Institutional Risk
    - Based on level of research activities in each division within KKH
    - (At least 1 study from each division will be selected annually to maintain oversight)
  - Level 2: Study Protocol Risk
    - Based on MOH Audit Risk Assessment
      - (Risk scoring of research protocols including minor, pregnant women, target recruitment number, length of research study, complexity of research protocols, number of research sites, etc.)
- Audit visits have been conducted for 8 HBRA regulated research till May 2019 and it is on an ongoing basis.
- Areas for improvement identified during the audits were communicated with the research study teams through audit reports and also sharing with the research community (e.g. Clinical Research Coordinator Network).

CONCLUSION

Risks of research regulatory non-compliances have been reduced through the corrective and preventive actions from the findings of the internal audits as well as communication with the KKH research community.