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2012-03-07

REC Reference nr 230408-011
IRB nr 00006240

DR MOA BENEDICT
DEPARTMENT OF FAMILY MEDICINE
CR DE WET BUILDING
UFS

Dear Dr Benedict

ECUFS NR 27/2012
DR MOA BENEDICT

DEPT OF FAMILY MEDICINE


**PROJECT TITLE: KNOWLEDGE OF MIDWIVES AND DOCTORS WORKING IN MATERNITY
UNITS OF FREE STATE DISTRICT HOSPITALS ON THE MANAGEMENT OF POSTPARTUM
HAEMORRHAGE.**

- You are hereby kindly informed that the Ethics Committee approved the above project at the meeting held on 6 March 2012.

(Prof WJ Steinberg did not take part in the discussion of this study)

- Committee guidance documents: Declaration of Helsinki, ICH, GCP and MRC Guidelines on Bio Medical Research. Clinical Trial Guidelines 2000 Department of Health RSA; Ethics in Health Research: Principles Structure and Processes Department of Health RSA 2004; Guidelines for Good Practice in the Conduct of Clinical Trials with Human Participants in South Africa, Second Edition (2006); the Constitution of the Ethics Committee of the Faculty of Health Sciences and the Guidelines of the SA Medicines Control Council as well as Laws and Regulations with regard to the Control of Medicines.
- Any amendment, extension or other modifications to the protocol must be submitted to the Ethics Committee for approval.
- The Committee must be informed of any serious adverse event and/or termination of the study.
- A progress report should be submitted within one year of approval of long term studies and a final report at completion of both short term and long term studies.
- Kindly refer to the ECUFS reference number in correspondence to the Ethics Committee secretariat.

Yours faithfully


For CHAIR: ETHICS COMMITTEE
cc Prof WJ Steinberg