

STANFORD UNIVERSITY

Administrative Panel on Human Subjects in Medical Research

2017-2018
IRB #6: Roster

Palo Alto, CA 94306
Assurance #FWA00000935

The Panel is composed of 24 members

VOTING MEMBERS

Fifteen Members affiliated with Stanford University

OAKES, David D. (M.D.) (CHAIR)
Professor, Emeritus
Surgery

MEYER, Timothy W. (M.D.)
Professor
Medicine/Nephrology (VA)

AMYLON, Michael D. (M.D.)
Professor, Emeritus
Pediatrics

MOLVIN, Celia (CIP)
Nonscientific Member

DOHERTY, Anastasia (CIP)
Nonscientific Member

MUNDY, David C.H. (M.Div.)
Reverend
Nonscientific Member

GILL, Manjit (CIP)
Nonscientific Member

RECHT, Lawrence D. (M.D.)
Professor
Neurology and Neurological Sciences

GOEL, Ashima (M.S.)
Nonscientific Member

HOLZER, Alison (Ph.D.)
Clinical Research Manager
Pediatrics

SCANDLING, John D. (M.D.)
Professor
Medicine/Nephrology

LANE, Alfred (M.D.)
Professor
Dermatology and Pediatrics

STOCKDALE, Frank E. (M.D.)
Professor, Emeritus
Medicine/Oncology

MATTMANN, Debra C. (R.N., M.S.N.)
Nursing Administrator
GCRC-SHC

WILSON, Darrell M. (M.D.)
Professor
Pediatrics/Endocrinology

Three Outside Nonscientific Members Otherwise Unaffiliated with Stanford

PARKER, George W. (M.B.A.)
Controller (retired)

PETERHANS, Laura (M.A.)
Teacher (retired)

EIGENBROD, Richard A. (M.B.A.)
Business Consultant

NON-VOTING MEMBERS

Six Ex Officio members

BANGHART, Dawn (C.H.P.)
Sr. Health Physicist
Environmental Health and Safety (EH&S)
(Alternate for Lance Phillips)

PHILLIPS, Lance (M.S., C.H.P., C.S.P.)
Radiation Safety Officer
Environmental Health and Safety

CAPLUN, Elizabeth
Deputy Director
Office of the Vice Provost and
Dean of Research
(Alternate for Kathy McClelland)

SEGAL, Ellyn D. (Ph.D.)
Biosafety Officer
Environmental Health and Safety

MCCLELLAND, Kathy
Research Compliance Director
Office of the Vice Provost and
Dean of Research

THOMPSON, Kathleen
Director
Research Management Group
Office of the Dean of the School of
Medicine

ICH/GCP: Stanford University Administrative Panels on Human Subjects in Medical Research (IRB) are in compliance with Good Clinical Practices as consistent with U.S. Food and Drug Administration Code of Federal Regulations (21 CFR 50 and 56) and DHHS (45 CFR Part 46).

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