7:30  Continental Breakfast

8:00  Framework for Effective Use of CED to Inform Decision Making
      Alan Garber MD, Stanford University

What are major considerations in determining whether specific medical services or research questions can be credibly evaluated using data collecting using clinical registries? In what situations might coverage with evidence development be expected to produce actionable data versus those where CED it is likely inadequate to answer critical questions?

9:30  Break

Clinical Registry Case Studies (Chair – Barbara McNeil, MD, PhD, Harvard Medical School)

9:45  Medicare ICD Registry  Stephen Hammill, MD, Mayo Clinic

10:45 National Emphysema Treatment Trial. Scott Ramsey, PhD, Fred Hutchinson

11:45 Lunch

Clinical Registry Case Studies (Chair – Marc Berger, MD, Eli Lilly)

12:45 ABMT for Breast Cancer, Wade Aubry, MD, UCSF

1:45  Drug Eluting Stents, Mark Hlatky, MD, Stanford University

2:45  Break

3:00  Lessons Learned and Implications for Policy

   Moderator: Robert Mechanic, MBA, The Health Industry Forum, Brandeis University
   Panel:  Troyen Brennan, MD, Aetna, Inc.
          Kathy Buto, MPA, Johnson & Johnson
          Robert Califf, MD, Duke University
          Sharon Levine, MD, The Permanente Medical Group
          Steve Phurrough, MD, MPA, CMS

4:30  Meeting adjourns