

Enhancing the Clinical Trial Enterprise for Antibacterial Drug Development in the United States —November 18-19, 2019

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Disclosures

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Erin Duffy: CARB-X receives funding from the governments of the United States (BARDA, NIAID), United Kingdom (DHSC/GAMRIF), Germany (BMBF) and two charitable foundations, the Wellcome Trust and the Bill & Melinda Gates Foundation.

Scott Evans has grant funding from NIH. He has consultancies with AstraZeneca (Trial Executive Committee). He's received funding for editorial activities from DeGruyter and publishing royalties from Taylor & Francis. He's received travel expenses / honorariums for educational activities and meetings from the American Statistical Association, CIOMS, Huntington's Study Group, ACTION, FDA, Osaka University, National Cerebral and Cardiovascular Center of Japan, NIH, Society for Clinical Trials, American Society for Microbiology, Clinical Trials Transformation Initiative, University of Colorado, UCLA, International Society for Biopharmaceutical Statistics, Deming Conference on Applied Statistics, Antimicrobial Resistance and Stewardship Conference, World Antimicrobial Congress, and the CRT Conference. Dr. Evans serves on DMC or advisory committees for Takeda / Millennium, Pfizer, Roche, Novartis, Achaogen, Genentech, Amgen, GSK, Teva, Zeiss, Dexcom, Claret Medical, Vir, Arrebus, Five Prime, Shire, Gilead, Spark, Nuvelution, Tracon, Chiesi, WAVE, Advantagene, Braeburn, Cardinal Health, Covance, Lipocine, Microbiotix, Stryker, Rakuten, BENEFIT, AImmune, and the Austrian Breast & Colorectal Cancer Study Group (ABCSG)/Breast International Group (BIG) and the Alliance Foundation Trials (AFT).

Vance Fowler: Consultant: Pfizer, Novartis, Galderma, Novadigm, Durata, Debiopharm, Genentech, Achaogen, Affinium, Medicines Co., Cerexa, Tetrphase, Trius, MedImmune, Bayer, Theravance, Cubist, Basilea, Affinergy, Janssen, xBiotech, Contrafect, Regeneron, Basilea, Destiny, Amphlphi Biosciences. Integrated Biotherapeutics.

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-Royalties: UpToDate

-Patent pending: Sepsis diagnostics

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