BIOGRAPHICAL SKETCH

Provide the following information for the Senior/key personnel and other significant contributors. Follow this format for each person. **DO NOT EXCEED FIVE PAGES.**

NAME: David S. Perlin

eRA COMMONS USER NAME (credential, e.g., agency login): PERLIN

POSITION TITLE: Chief Scientific Officer and Professor

EDUCATION/TRAINING

INSTITUTION AND LOCATION	DEGREE (if applicable)	Completion Date MM/YYY Y	FIELD OF STUDY
Brandeis University, Waltham, MA	A.B.	1976	Biology
Cornell University, Ithaca, NY	Ph.D.	1980	Plant Physiology
Yale University School of Medicine, New Haven, CT	Postdoc.	1980-83	Biochemistry, Genetics
University of Rochester School of Medicine and Dentistry, Rochester, NY	Postdoc.	1983-85	Biochemistry

A. Personal Statement

My career spans over 30 years in translational infectious disease research with a strong emphasis on developing countermeasures to high-threat agents causing acute respiratory diseases. I have led five highly successful national drug accelerator programs sponsored by the NIH and/or DoD, which involved partnerships with industry. My work has focused on advancing novel anti-infective strategies, including the development of therapeutics for high-threat viral, bacterial, and fungal pathogens. Notably, I led two recent NIH Centers of Excellence in Translational Research (CETR) focused on overcoming antimicrobial resistance, where four antibiotic candidate programs have advanced to IND-enabling and de-risking stages. Most recently, I served as co-PI with Nobel Laureate Dr. Charles Rice (Rockefeller University) for the NIH AntiViral Drug Discovery (AViDD) program. Our consortium, the Metropolitan Antiviral Drug Accelerator (MAVDA), developed four advanced antiviral candidates including one that progressed to Phase II clinical trials.

My translational team—comprising seasoned researchers from both academia and industry—has supported the advancement of multiple small-molecule and biologic therapeutics, including four FDA-approved drugs and several others in late-stage clinical trials. For a decade, I also directed the Rutgers Regional Biocontainment Laboratory (RBL), a national center for research on highly transmissible high-threat pathogens, positioning me to contribute meaningfully to the biosafety and translational components of this program. My extensive publication record (>365 scientific papers, reviews, and chapters) further reflects my long-standing commitment to advancing science-driven therapeutic innovation.

Ongoing and recently completed projects that I would like to highlight include:

1U19Al1714010 Perlin (co-PI), Rice (co-PI) 05/17/2022 – 4/30/2025

NIH/NIAID

Metropolitan AntiViral Drug Accelerator (MAVDA)

1U19AI142731-01 Perlin (PI) 4/01/2019 – 3/31/2025

NIH/NIAID

Center to develop innovative therapeutics to multidrug resistant high-threat bacterial agents

2 R01 Al109025-11 Perlin (PI) 07/01/2023 - 06/30/2028

NIH/NIAID

Critical Factors Influencing Echinocandin Resistance in Candida glabrata.

3U19AI142731-02S1 Perlin (PI) 08/25/2020 – 04/30/2024

NIH/NIAID

A CETR-based partnership accelerator for rapid drug development targeting SARS-CoV-2 and pancoronaviruses

1 R01 Al138986-01 Perlin (PI) 05/01/2018 - 5/31/2024

NIH/NIAID

Novel bi-specific immunoprophylactics against multi-drug resistant Gram-negative bacterial infections

1 R01 Al141183-01 Perlin (PI) 12/01/2018 - 11/30/2023

NIH/NIAID

Novel bi-specific immunotherapeutic against high-threat Gram-negative pathogens

U19 Al109713-01 Perlin (PI) 3/01/2014 – 2/28/2019

NIH/NIAID

Center to develop therapeutic countermeasures to high-threat bacterial agents

Citations

- a. Meyer C, Garzia A, Miller MW, Huggins DJ, Myers RW, Hoffmann HH, Ashbrook AW, Jannath SY, Liverton N, Kargman S, Zimmerman M, Nelson AM, Sharma V, Dolgov E, Cangialosi J, Penalva-Lopez S, Alvarez N, Chang CW, Oswal N, Gonzalez I, Rasheed R, Goldgirsh K, Davis JA, Ramos-Espiritu L, Menezes MR, Larson C, Nitsche J, Ganichkin O, Alwaseem H, Molina H, Steinbacher S, Glickman JF, Perlin DS, Rice CM, Meinke PT, Tuschl T. Small-molecule inhibition of SARS-CoV-2 NSP14 RNA cap methyltransferase. Nature 2024.
- b. Alvarez N, Adam GC, Howe JA, Sharma V, Zimmerman MD, Dolgov E, Rasheed R, Nizar F, Sahay K, Nelson AM, Park S, Zhou X, Burlein C, Fay JF, Iwamoto DV, Bahnck-Teets CM, Getty KL, Lin Goh S, Salhab I, Smith K, Boyce CW, Cabalu TD, Murgolo N, Fox NG, Mayhood TW, Shurtleff VW, Layton ME, Parish CA, McCauley JA, Olsen DB, **Perlin DS**. 2024. Novel Pan-Coronavirus 3CL Protease Inhibitor MK-7845: Biological and Pharmacological Profiling. Viruses. 2024 Jul 18;16(7):1158. doi: 10.3390/v16071158.
- c. Shurtleff VW, Layton ME, Parish CA, Perkins JJ, Schreier JD, Wang Y, Adam GC, Alvarez N, SBahmanjah S, Bahnck-Teets CM, Boyce CW, Burlein C, Cabalu TD, Campbell BT, Carroll SS, Chang W, de Lera Ruiz M, Dolgov E, Fay JF, Fox NG, Shih Lin Goh SL, Hartingh TJ, Hurzy DM, Kelly III MJ, Klein DJ, KlinglereF-M, Krishnamurthy H, Kudalkar SN, Mayhood TW, McKenna PM, Murray EM, Nahas D, Nawrat CC, Park S, Qian D, Roeckera AJ, Sharma c, Shipe WD, Su J, Taggart RV, Truong Q, Wu Y, Zhou X, Zhuang N, **Perlin DS**, Olsen DB, Howe JA, and McCauley JA 2024 Invention of MK-7845, a SARS-CoV-2 3CL protease inhibitor employing a novel difluorinated glutamine mimic. J Med Chem. 2024 Mar 14;67(5):3935-3958. doi: 10.1021/acs.jmedchem.3c02248.
- d. Chang CW, Oswal N, Murugan M, Goldgirsh K, Tsao W, Park S, Perlin DS. A novel cellular tool for screening human pan-coronavirus antivirals. Antiviral Res. 2025 Jun 10;240:106212. doi: 10.1016/j.antiviral.2025.106212. PMID: 40505777

B. Positions, Scientific Appointments, and Honors

Positions and Scientific Appointments

2022-	Deputy Director Consortium Integration and Strategic Initiatives for Research,
	Lombardi Comprehensive Cancer Center Consortium
2019-	Professor, Dept. of Microbiol & Immunol, Georgetown Univ Sch. of Med, Washington,
D.C. 2019-	Professor, Dept. of Medical Sciences, Hackensack Meridian School of Medicine
2019-	CSO, Hackensack Meridian Health Center for Discovery and Innovation, Nutley, NJ
2002-2018	Professor, Department of Microbiology, Biochemistry and Molecular Genetics, New Jersey Medical School, Rutgers Biomedical and Health Sciences, Newark, NJ
2009-2018	Director, Rutgers Regional Biocontainment Laboratory
2006-2018	Executive Director, Public Health Research Institute, NJ Med Sch-UMDNJ,
	Newark, NJ

2005-2006 President, Public Health Research Institute, Newark, NJ

1992-2002 1989-1991 1985-1988 Honors	Adjunct Associate Professor, New York University School of Medicine, New York, NY 1992-2018 Member, Public Health Research Institute, New York, NY Associate Member, Public Health Research Institute, New York, NY Assistant Member, Public Health Research Institute, New York, NY
2025 2023 2022 2021, 2024 2020, 2021	Fellow, Infectious Diseases Society of America Dr. Sol J. Barer Award for Vision, Innovation and Leadership Notable Health Care Leader- Crain's New York Business Named top 25 national innovator by Modern Healthcare EJI, honored as top Scientist in New Jersey

2018 Fellow, American Academy of Microbiology

2017- 2021 Chair, Audit Committee, American Society for Microbiology (ASM)

NJBIZ, Healthcare Hero award for COVID-19 response

2017 Named inaugural Editor-in-Chief for Journal of Fungi

2012- Executive Committee, Board of Directors, Aaron Diamond AIDS Research Center

2009-2012 Distinguished Visiting Professor, University of Manchester, United Kingdom

Named top 10 Health Care Influencer in New Jersey-RO1-NJ

2005 Fellow, The New York Academy of Sciences

Contributions to Science

2021

2019. 2020

1. Drug discovery targeting high-threat bacteria, fungi and viruses.

Multidrug-resistance plagues global and U.S. healthcare and with few new antibiotics making it to market from a diminished pipeline, there is an unmet medical need for new therapeutics to treat drug-resistant infections. The Perlin Lab has been involved in developing new targets and novel chemical scaffolds against high-threat fungal and bacterial pathogens. They are developing both narrow- and broad-spectrum agents against high-threat multi-drug resistant bacterial and fungal pathogens commonly associated with systemic infections among immunosuppressed patients. I lead an NIH designated Center of Excellence in Translational Research (CETR) and an Antiviral Drug Discovery accelerator (AViDD) that is developing a new generation of antibiotics against high-threat bacteria and viruses like SARS- CoV-2. These programs are collaborative public-private partnerships that accelerate the discovery and development of novel antibiotics and antivirals by joining together academic and industry researchers and providing critical core resources to turn highly promising early concept molecules into potential therapeutics suitable for clinical evaluation.

- a. Wang Z, Koirala B, Hernandez Y, Zimmerman M, Park S, **Perlin DS**, Brady SF. 2022. A naturally inspired antibiotic to target multidrug-resistant pathogens. Nature. 2022 Jan 5. doi: 10.1038/s41586- 021- 04264-x
- b. Park S, Russo R, Westfall L, Shrestha R, Zimmerman M, Dartois V, Kurepina N, Kreiswirth B, Singleton E, Li SG, Mittal N, Ahn YM, Bilotta J, Connolly KL, Jerse AE, Freundlich JS, Perlin DS. A Novel Oral GyrB/ParE Dual Binding Inhibitor Effective against Multidrug-Resistant Neisseria gonorrhoeae and Other High-Threat Pathogens. Antimicrob Agents Chemother. 2022 Aug 16:e0041422. doi: 10.1128/aac.00414-22.
- c. Lovey A, Krel M, Borchardt A, Brady T, Cole JN, Do QQ, Fortier J, Hough G, Jiang W, Noncovich A, Tari L, Zhao Q, Balkovec JM, Zhao Y, **Perlin DS**. Development of novel immunoprophylactic agents against multidrug resistant Gram-negative bacterial infections. Antimicrob Agents Chemother. 2021 Aug 9:AAC0098521. doi: 10.1128/AAC.00985-21
- d. Hover BM, Kim SH, Katz M, Charlop-Powers Z, Owen JG, Ternei MA, Maniko J, Estrela AB, Molina H, Park S, **Perlin DS**, Brady SF. 2018. Culture-independent discovery of the malacidins as calcium- dependent antibiotics with activity against multidrug-resistant Gram-positive pathogens. Nat Microbiol. 3(4):415-422. doi: 10.1038/s41564-018-0110-1.

2. Drug resistance and tolerance mechanisms in fungi.

I have had a longstanding interest in antifungal drug resistance, which continues to be an emerging problem in medical mycology. We first reported the mechanism of clinical resistance to echinocandin class antifungal

drugs in 2005 and have provided a comprehensive molecular and clinical assessment of the resistance mechanism resulting in 150 papers and reviews. Our work has been instrumental in moving the field forward and has emphasized correlations between resistance mutations, genetics, enzyme kinetic inhibition, MIC, pharmacodynamics, resistance factors and clinical outcome. This multifactorial approach was critical to the development of revised CLSI breakpoints. My lab was established (Pfizer then Astellas) as a Global Clinical Reference Center for molecular evaluation of echinocandin resistant strains from patients failing therapy. We have evaluated the *FKS* mechanism in nearly one thousand clinical isolates. We have examined the relationship between resistance, virulence, and strain lineage, and we have used PK-PD studies to understand the importance of specific mutations and potential therapeutic response. Finally, in recent years, we have helped define underlying genetic and host factors that contribute to drug tolerance leading to emergence of echinocandin and multidrug resistance in *Candida species* including *Candida auris* and *Aspergillus* species.

- a. Arastehfar A, Daneshnia F, Cabrera N, Penalva-Lopez S, Sarathy J, Zimmerman M, Shor E, Perlin DS. Macrophage internalization creates a multidrug-tolerant fungal persister reservoir and facilitates the emergence of drug resistance. Nat Commun. 2023 Mar 2;14(1):1183. doi: 10.1038/s41467-023-36882-
- b. Arastehfar A, Daneshnia F, Hovhannisyan H, Fuentes D, Cabrera N, Quinteros C, Ilkit M, Ünal N, Hilmioğlu-Polat S, Jabeen K, Zaka S, Desai JV, Lass-Flörl C, Shor E, Gabaldon T, **Perlin DS.** Overlooked *Candida glabrata* petites are echinocandin tolerant, induce host inflammatory responses, and display poor *in vivo* fitness. mBio. 2023 Oct 31;14(5):e0118023. doi: 10.1128/mbio.01180-23.
- c. Garcia-Rubio R, Jimenez-Ortigosa C, DeGregorio L, Quinteros C, Shor E, **Perlin DS.** Multifactorial role of mitochondria in echinocandin tolerance revealed by transcriptome analysis of drug-tolerant cells. mBio. 2021 12(4):e0195921. doi: 10.1128/mBio.01959-21.
- d. Satish S, Jiménez-Ortigosa C, Zhao Y, Lee MH, Dolgov E, Krüger T, Park S, Denning DW, Kniemeyer O, Brakhage AA, **Perlin DS**. Stress-Induced Changes in the Lipid Microenvironment of β-(1,3)-d-Glucan Synthase Cause Clinically Important Echinocandin Resistance in Aspergillus fumigatus. MBio. 2019 10(3). pii: e00779-19. doi: 10.1128/mBio.00779-19.

3. Rapid detection of respiratory, GI tract and bloodstream infections, and associated resistance markers.

Early and appropriate antimicrobial therapy is critical to a favorable outcome for patients with respiratory and BSIs. Current diagnostic methods are inadequate and reducing the time from specimen collection to species identification and antimicrobial susceptibility is essential for improving patient outcome. For the past decade and one-half, my group has been involved in developing next-generation nucleic acid PCR- and RNA-based molecular beacon platforms for rapid identification of viral, bacterial and fungal pathogens, and associated drug resistance in high threat bacterial, viral and fungal pathogens.

- a. Kordalewska M, Perlin DS. 2022. Detection and Identification of Candida auris from Clinical Skin Swabs. Methods Mol Biol. 2022;2542:245-256. doi: 10.1007/978-1-0716-2549-1 18.
- b. Zhao Y, Lee A, Composto K, Cunningham MH, Mediavilla JR, Fennessey S, Corvelo A, Chow KF, Zody M, Chen L, Kreiswirth BN, Perlin DS. 2021 A novel diagnostic test to screen SARS-CoV-2 variants containing E484K and N501Y mutations. Emerg Microbes Infect. 2021 Dec;10(1):994-997. doi: 10.1080/22221751.2021.1929504.
- c. Zhao Y, Cunningham MH, Mediavilla JR, Park ^s, Fitzgerald S, Ahn HS, Li X, Zhan C, Hong T, Munk G, Chow KF, **Perlin DS**. 2021. An observational study of COVID-19 from a large healthcare system in Northern New Jersey: Diagnosis, clinical characteristics, and outcomes. Sci Rep Feb 23;11(1):4389. doi: 10.1038/s41598-021-83959-7.
- d. Kordalewska M, Zhao Y, Lockhart SR, Chowdhary A, Berrio I, **Perlin DS**. (2017) Rapid and accurate molecular identification of the emerging multidrug resistant pathogen *Candida auris*. J Clin Microbiol. May 24. pii: JCM.00630-17. doi: 10.1128/JCM.00630-17
- 4. Improving existing drug therapy. A key factor for successful therapy is whether a drug get to the site of infection at the desired level for efficacy. My research group has been using novel technology to image and quantify the level of drugs in life-threatening diseases resulting from intra-abdominal abscesses and

pulmonary lesions. This work provides insights into more effective therapy by increasing exposure levels and reducing the emergence of drug resistance resulting from suboptimal dosing. In addition to drug access, drug response is often limited factors such as an individual's metabolism, which effect whether a drug is present at the desired concentration over the course of therapy. Classically, therapeutic drug monitoring (TDM) has been used to assess drug levels in patients. This is often a laboratory-intensive process that can take several days. We are developing novel technology to rapidly assess drug levels of first-line antimicrobial agents in blood in real-time at the bedside.

- a. Lee A, Wang N, Carter CL, Zimmerman M, Dartois V, Shaw KJ, **Perlin DS**, Zhao Y. Therapeutic Potential of Fosmanogepix (APX001) for Intra-abdominal Candidiasis: from Lesion Penetration to Efficacy in a Mouse Model. Antimicrob Agents Chemother. 2021 Mar 18;65(4):e02476-20. doi: 10.1128/AAC.02476-20.
- b. Lee A, Prideaux B, Lee MH, Zimmerman M, Dolgov E, **Perlin DS**, Zhao Y. 2019. Tissue Distribution and Penetration of Isavuconazole at the Site of Infection in Experimental Invasive Aspergillosis in Mice with Underlying Chronic Granulomatous Disease. Antimicrob Agents Chemother 63(6). pii: e00524-19. doi: 10.1128/AAC.00524-19.
- c. Zhao Y, Prideaux B, Nagasaki Y, Lee MH, Chen PY, Blanc L, Ho H, Clancy CJ, Nguyen MH, Dartois V, **Perlin DS**. 2017 Unraveling Drug Penetration of Echinocandin Antifungals at the Site of Infection in an Intra- Abdominal Abscess Model. Antimicrob Agents Chemother. AAC.01009-17. doi: 10.1128/ AAC.01009-17.

Complete List of Published Work in MyBibliography:

(see: https://pubmed.ncbi.nlm.nih.gov/?term=perlin%2Bds&sort=date)