

CURRICULUM VITAE

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Education

<u>Year</u>	<u>Degree</u>	<u>Field</u>	<u>Institution</u>
1989	B.A.	Biology	Division of Biology, Moscow State University, Moscow, USSR
1992	M.S.	Genetics	Department of Genetics, Moscow State University, Russia
1995	Ph.D.	Biochemistry	Institute of Molecular Genetics, Russian Academy of Sciences

Postdoctoral Training

<u>Dates</u>	<u>Specialty</u>	<u>Mentor</u>	<u>Place of Training</u>
1995-1997	Molecular Biology	Dr. A. Goldfarb	Public Health Research Institute, New York

Academic Appointments

1997-2003	Assistant Professor, New York University School of Medicine, New York, NY
2003-2005	Associate Professor (tenured), New York University School of Medicine
2005-2008	Professor, Dept. of Biochemistry, NYU School of Medicine
2008-	Julie Wilson Anderson Professor, NYU School of Medicine
2013-	Investigator, Howard Hughes Medical Institute

Awards/Honors

1997	The Helen Hay Whitney Foundation research fellowship (declined)
1998	The Searle Scholar Award
2001	The Irma T. Hirschl Career Scientist Award
2002	The Edward Mallinckrodt, Jr. Foundation Research Award
2002	Fogarty International Research Collaboration Award
2004	Keynote Lecture: The RNA Biochemistry Symposia. Blaubeuren, Germany
2004	United States-Israel Bi-national Science Foundation Award
2006	The Dynasty Foundation Award
2006	NIH Director's Pioneer Award
2008	Crain's "40 Under 40" Alumni
2008	Julie Wilson Anderson Professorship (endowment chair)
2009	The Vilcek Foundation Finalist Prize
2010	Keynote Lecture: The Harden Conference on Transcription. Cambridge, UK
2010	The Blavatnik Award, New York Academy of Science
2011	The Robertson Foundation Award
2012	Biogerontology Research Foundation Award
2012	Blavatnik Family Foundation Award
2013	Investigator, Howard Hughes Medical Institute

2013	Honorary Professor, Moscow Institute of Physics and Technology
2013	Master Scientist, NYU Langone Medical Center
2014	Dean's Honor Lecture, NYU School of Medicine
2016	The Neil Welker Memorial Award
2016	Engelhardt Memorial Lecture, IMB, Russian Academy of Science
2016	Elected Foreign Member, Russian Academy of Sciences
2017	Elected Fellow, American Academy of Arts and Sciences
2018	Glenn Award for Research in Biological Mechanisms of Aging

Research Statement

Our most significant contributions to science include:

1. Discovery and implications of RNA polymerase (RNAP) “backtracking” and “ratcheting”:

In 1997 we described back-and-forth sliding of RNAP along DNA and RNA. Our group then showed that this phenomenon, which we called “*backtracking*”, plays the key role in regulating gene expression (via pausing and termination), in transcriptional fidelity, in genome instability, and in coupling transcription to DNA repair. We were also the first to demonstrate that RNAP is a Brownian ratchet machine. Our findings explained in mechanistical details how RNAP translocates, how it responds to regulatory signals and factors, and how it terminates transcription.

2. Mechanisms of transcription termination and antitermination: We have uncovered the general mechanistical principles of the termination process in bacteria and formulated models explaining the intricate molecular pathways leading to both intrinsic and Rho-dependent transcription termination. We also studied how these processes are regulated in the cell and formulated the mechanistical models of factor-dependent (antitermination) and factor-independent (riboswitches) modes of regulation. Our studies uncovered novel functions of Rho, such as its role in silencing of horizontally transferred (and potentially toxic) genes and in preserving genomic integrity. We found that Rho functions as the global regulator of transcription acting at the 5'UTRs of hundreds of bacterial genes, and that sRNAs control Rho-dependent termination genome-wide, thus establishing sRNAs as transcription elongation factors.

2. Transcription coupled DNA repair (TCR): In 2014 we uncovered a general mechanism of TCR that relies on active RNAP backtracking. We found that in bacteria, UvrD binds RNAP during transcription elongation and, using its helicase activity, forces RNAP to slide backward along DNA. By inducing backtracking, UvrD exposes DNA lesions shielded by blocked RNAP, allowing the repair enzymes to gain access to sites of damage. The small molecule alarmone ppGpp and general elongation factor NusA contribute to UvrD-mediated TCR. Because backtracking is a shared feature of all cellular RNAP, this mechanism enables RNAPs to function as global DNA damage scanners in bacteria and eukaryotes.

3. Discovery and characterization of riboswitches: In 2002 we described the first ligand-sensing mRNAs that regulate biosynthetic genes in *B. subtilis*. Simultaneously, Ron Breaker reported similar findings in *E. coli*. Since then dozens of riboswitches have been described in bacteria and eukaryotes where they control numerous genes. We have shown that riboswitches can activate and suppress gene expression acting at the level of transcription termination (both intrinsic and Rho-dependent), translation initiation, and also modulating alternative splicing and mRNA stability (in plants).

4. Discovery of the eukaryotic RNA thermosensor: In 2006 we isolated a complex composed of the translation elongation factor eEF1A1 and a novel non-coding RNA (HSR1) that is required for activation of heat shock genes in mammals. We have accumulated evidence demonstrating that HSR1 serves as a *bona fide* molecular *thermosensor*, which is set for different temperatures in different organisms. We also showed that eEF1A1 orchestrates the whole process of heat

shock response, from transcription activation to mRNA stabilization, transport, and translation. These findings provide a new paradigm of cellular adaptation to stress, with far-reaching clinical implications.

5. Bacterial gasotransmitters: We have shown that endogenously produced gases NO and H₂S protect bacteria from oxidative stress, immune attack, and numerous antibiotics. These results support the emerging concept of antibiotic killing, which relies in part on oxidative damage, and establish NO- and H₂S-producing enzymes as promising new targets for antimicrobial therapy. We further showed that NO produced by bacteria inside *C. elegans* diffuses into animal's tissues where it activates a defined set of genes that protect nematodes from environmental stress and extend their lifespan.

Bibliography:

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2. Kashlev M., Nudler E., Goldfarb A., White T., Kutter E. (1993) Bacteriophage T4 Alc protein: a transcription termination factor sensing local modification of DNA. **Cell** **75**: 147-154.
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