

Nathan J. Hillson

Lawrence Berkeley National Lab
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EDUCATION

Stanford University School of Medicine, Stanford CA
Postdoctoral Fellowship, Microbiology, 2009

Harvard Medical School, Boston MA
Ph.D., Biophysics, 2004

Rice University, Houston TX
B.A., Physics; Computational and Applied Mathematics, *cum laude*, 1999

Selected Course Work:

Macromolecular X-ray Crystallography; Chemical Biology; Simulation of Macromolecules; Single-molecule Biophysics; Molecular Approaches to Drug Action, Discovery, and Design; Proteins: Structure, Function and Catalysis; Genomics and Computational Biology; Molecular Biology and Genetics; Numerical Analysis; Partial Differential Equations; High Performance and Parallel Computing; Algorithms and Data Structures; Statistical Mechanics and Thermodynamics; Quantum Mechanics.

EXPERIENCE

DOE Agile BioFoundry, Emeryville CA **2016-present**
Title: *Lead Principal Investigator*
Responsibilities: To lead the development and demonstration of the Agile BioFoundry.

DOE Joint BioEnergy Institute (JBEI), Emeryville CA **2009-present**
Title: *Chief Information Officer* 2025-present
Director of Synthetic Biology Informatics 2014-present
Director of Synthetic Biology 2009-2014
Responsibilities: To develop and execute JBEI's Information Technology Roadmap. Lead and support initiatives involving automation, AI/ML, and analytics. Direct development and operation of laboratory automation, software engineering, and molecular/microbiological workflows.

DOE Joint Genome Institute (JGI), Berkeley CA **2012-present**
Title: *Program Lead, Synthetic Biology Informatics*
Responsibilities: To develop and demonstrate DNA screening biosecurity software and Synthetic Biology Institutional Review processes and software, and coordinate SynBio Informatics work at the JGI, JBEI, and ABF.

Lawrence Berkeley National Lab (LBNL), Berkeley CA **2009-present**
Title: *BioDesign Dept. Head, Biological Systems & Engin. Div.* 2015-present
Senior Staff Scientist, Biological Systems & Engin. Div. 2020-present
Staff Scientist, Biological Systems & Engin. Div. 2015-2020
Co-Deputy Director, Biological Systems & Engin. Div. 2015-2016
Staff Scientist, Physical Biosciences Division 2011-2015
Research Scientist, Physical Biosciences Division 2009-2011
Responsibilities: In addition to Agile BioFoundry, JBEI, and JGI responsibilities (see above), provide and participate in the strategic and operational leadership of the Biological Systems & Engineering Division and more broadly the LBNL Biosciences Area.

- Engineering Biology Research Center (EBRC), Emeryville CA** **2016-present**
Title: *Affiliate Investigator*
Responsibilities: Participation in road mapping activities as well as the Security Working Group.
- TeselaGen Biotechnologies, Inc., San Francisco CA** **2011-present**
Title: *Co-Founder and Chief Scientific Officer*
Responsibilities: To lead the design of TeselaGen's DNA design and assembly bioCAD/CAM platform, and provide software development quality assurance and control.
- Synberc, Emeryville CA** **2013-2016**
Title: *Affiliate Investigator*
Responsibilities: To further develop and demonstrate the ICE repository platform towards achieving a "Web of Registries", and to support and maintain Synberc's ICE repository instance.
- Stanford University School of Medicine, Stanford CA** **2004-2009**
Title: *Postdoctoral Research Fellow*
Research Advisor: *Lucy Shapiro, Department of Developmental Biology*
Research topics: Engineered the bacterium *Caulobacter crescentus* into a whole-cell Uranium biosensor. Developed a high-throughput microscopy screen to identify mutant strains of *Caulobacter* that mislocalize the polar histidine kinases DivJ/PlcC, or the cell-division protein ZapA. Protein-engineered a chimera of the essential histidine kinase CckA to generate a colorimetric reporter of CckA activity. Utilized the CckA chimera reporter system in a high-throughput screen to identify *Caulobacter* mutant strains that fail to activate CckA.
- Harvard Medical School, Boston MA** **2000-2004**
Title: *Graduate Thesis Student*
Research Advisor: *Prof. Christopher T. Walsh*
Department of Biological Chemistry and Molecular Pharmacology
Dissertation topics: Studied the enzymatic activity of VibF, a six domain non-ribosomal peptide synthetase utilized in the biosynthesis of vibriobactin, an iron chelating siderophore (virulence factor) of the cholera causing pathogen *Vibrio cholerae*. Mapped catalytic activity to individual domains of VibF. Demonstrated that VibF functionally operates as a dimer, via biochemical mutant domain activity regain and by ultracentrifugation studies. Utilized correlated-mutation computational analysis to assist the chimeric-engineering of glycosyltransferases to decorate the peptide scaffold of the antibiotic vancomycin with alternative sugar moieties.
- Harvard University, Cambridge MA** **2000**
Title: *Graduate Rotation Student*
Research Advisor: *Prof. Eugene Shakhnovich*
Department of Chemistry and Chemical Biology
Rotation topic: Adapted an existing all-atom protein folding model/program to study protein-peptide interactions in the MHC-II/10-mer peptide recognition system. Implemented the Perceptron learning device to cyclically refine the intra- and inter-molecular potentials to guarantee the lowest energy of the native (crystal structure) state of the protein-peptide complex when challenged with generated decoy structures.

Harvard University, Boston MA 2000
Title: Graduate Rotation Student
Research Advisor: Prof. Christopher T. Walsh
Department of Biological Chemistry and Molecular Pharmacology
 Rotation topic: Studied the enzymatic activity of two previously uncharacterized proteins (YbtU and YbtT) implicated in the biosynthesis of yersiniabactin, an iron chelating siderophore necessary for the virulence of the plague causing pathogen *Yersinia pestis*. Attempted to measure cofactor binding and detect novel chemical species produced by YbtU, a protein predicted to be a putative reductase. Assayed for increased hydrolysis rates of a substrate analog mediated by YbtT, assigned as a putative external thioesterase.

Harvard University, Boston MA 1999
Title: Graduate Rotation Student
Research Advisor: Prof. Gerhard Wagner
Department of Biological Chemistry and Molecular Pharmacology
 Rotation topic: Used NMR spectroscopy to study the BID protein, a pro-apoptotic factor in the cell-death signaling pathway which becomes 100 times more potent after truncation by caspase-8. Conducted an experiment to collect a novel HSQC spectra of truncated BID. Analyzed differences in the spectra of truncated and full-length protein to identify residues whose chemical environments were changed due to the cleavage.

Los Alamos National Laboratories, Los Alamos NM 1996-1998
Title: Undergraduate Summer Research Associate
Research Advisor: Angel Garcia, Ph.D.
Theoretical Division (T-10): Theoretical Biology and Biophysics
 Area of study: Composed Monte Carlo code to study minimalist off-lattice protein folding models. Investigated pressure effects on folding thermodynamics and kinetics of the theoretical Thirumalai protein. Optimized numerical Lambda Repressor protein model to best fit experimental B-factor data.

TEACHING *Teaching fellow*, Biochemistry and Protein Structure, Harvard University 2001
 Taught weekly session of 15 undergraduates and led review sessions.
 Graded exams and weekly problem sets.

Teaching assistant, General Physics I and II, Rice University 1996-1999
 Taught weekly class recitation sessions and review sessions.

HONORS
 Genscript/Molecular Cloud: 2019 Distinguished Research Cloud Scientist Award
 Journal of Biological Engineering 2016 Publication of the Year
 NSF Bay Area Regional I-Corps Summer 2015, Trailblazer Award
 Berkeley Lab Director's Awards for Exceptional Achievement: Technology Transfer 2013
 Emerging Leaders in Biosecurity 2013 Fellow
 Synthetic Biology Leadership Accelerator Program 2012 Fellow
 Joint BioEnergy Institute 2012 Entrepreneur Award
 Damon Runyon Cancer Research Foundation Postdoctoral Research Fellow
 National Defense Science and Engineering Predoctoral Fellowship
 National Science Foundation Graduate Research Fellow
 Harvard University Distinction in Teaching Award
 Sigma Pi Sigma Physics Honor Society

PATENTS

Hillson, N.J. (2023) Scar-Less Multi-Part DNA Assembly Design Automation. U.S. Patent No. US202403112565A1. Issued September 19, 2024.

Hillson, N.J. (2019) Scar-Less Multi-Part DNA Assembly Design Automation. U.S. Patent No. 11,749,379. Issued September 5, 2023.

Hillson, N.J. (2016) Scar-Less Multi-Part DNA Assembly Design Automation. U.S. Patent No. 10,373,703. Issued August 6, 2019.

Hillson, N.J. (2012) Scar-Less Multi-Part DNA Assembly Design Automation. U.S. Patent No. 9,361,427. Issued June 7, 2016.

Chang, C., Bharadwaj, R., Singh, A., Chandrasekaran, A., and **Hillson, N.J.** (2012) Microfluidic Platform for Synthetic Biology Applications. U.S. Patent US10131903B2. Issued November 20, 2018.

Hillson, N.J., Shapiro, L., Hu, P., and Andersen, G.L. (2008) Heavy Metal Biosensor. U.S. Patent No. 8,697,388. Issued April 15, 2014.

RESEARCH

Gaillard, W. R., Sustarich, J., Li, Y., Carruthers, D. N., Gupta, K., Liang, Y., Kuo, R., Tan, S., Yoder, S., Adams, P. D., Garcia Martin, H., **Hillson, N. J.**, & Singh, A. K. (2025). "High-Throughput Microfluidic Electroporation (HTME): A Scalable, 384-Well Platform for Multiplexed Cell Engineering". *Bioengineering*, 12(8), 788. <https://doi.org/10.3390/bioengineering12080788>

Kim, H., **Hillson, N.J.**, Cho, B.K. et al. (2025) "Abstraction hierarchy to define biofoundry workflows and operations for interoperable synthetic biology research and applications". *Nat Commun* 16, 6056. <https://doi.org/10.1038/s41467-025-61263-6>

Zournas, A., Incha, M.R., Radivojevic, T., ..., **Hillson, N.**, et al. (2025) "Machine learning-led semi-automated medium optimization reveals salt as key for flaviolin production in *Pseudomonas putida*". *Commun Biol* 8, 630. <https://doi.org/10.1038/s42003-025-08039-2>

Chris M Yeager, **Nathan J Hillson**, Katherine L Wozniak, Vivek K Mutalik, Christopher W Johnson, Kyle Pomraning, Philip Laible, Taraka Dale, Adam M Guss. (2025) "The tier system: a host development framework for bioengineering". *Current Opinion in Biotechnology*. <https://doi.org/10.1016/j.copbio.2025.103260>

Matthew J. Szarzanowicz, Lucas M. Waldburger, Michael Busche, Gina M. Geiselman, Liam D. Kirkpatrick, Alexander J. Kehl, Claudine Tahmin, Rita C. Kuo, Joshua McCauley, Hamreet Pannu, Ruoming Cui, Shuying Liu, **Nathan J. Hillson**, Jacob O. Brunkard, Jay D. Keasling, John M. Gladden, Mitchell G. Thompson, Patrick M. Shih. (2024) "Binary vector copy number engineering improves *Agrobacterium*-mediated transformation". *Nat Biotechnol*. <https://doi.org/10.1038/s41587-024-02462-2>

Shannon Green, **Nathan J. Hillson**, John Moulton, Harshini Mukundan, Daniel P. Regan, Deepti Tanjore. (2024) "A Biomanufacturing Plan to Confront Future Biological Threats: Expert Panel Policy Review and Near-Term Recommendations", edited by Francesco Femia and Christine Parthemore. The Janne E. Nolan Center on Strategic Weapons, an institute of the Council on Strategic Risks. <https://councilonstrategicrisks.org/2024/09/04/a-biomanufacturing-plan-to-confront-future-biological-threats/>

Torres-Tiji, Yasin; Sethuram, Harish; Gupta, Abhishek; McCauley, Joshua; Dutra-Molino, Joao-Vitor; Pathania, Ruchi; Saxton, Lisa; Kang, Kalisa; **Hillson, Nathan**; Mayfield, Stephen. (2024) "Bioinformatic prediction and high throughput in vivo screening to identify cis-regulatory elements for the development of algal synthetic promoters" *ACS Synthetic Biology* sb-2024-00199b <https://doi.org/10.1021/acssynbio.4c00199>

Berezin C-T, Aguilera LU, Billerbeck S, Bourne PE, Densmore D, Freemont P, ... **Hillson NJ**, ..., et al. (2023) "Ten simple rules for managing laboratory information". *PLoS Comput Biol* 19(12): e1011652. <https://doi.org/10.1371/journal.pcbi.1011652>

Nava AA, Fear AL, Lee N, Mellinger P, Lan G, McCauley J, Tan S, Kaplan N, Goyal G, Coates RC, Roberts J, Johnson Z, Hu R, Wu B, Ahn J, Kim WE, Wan Y, Yin K, **Hillson N**, Haushalter RW, and Keasling JD. (2023) "Automated Platform for the Plasmid Construction Process." *ACS Synth Biol*. Nov 10. doi: 10.1021/acssynbio.3c00292

Nathan J. Hillson. (2023) "A Procedural Framework for Benchmarking Biofoundry Capabilities" *ACS Synthetic Biology* DOI: 10.1021/acssynbio.3c00491

Douglas Densmore, **Nathan J. Hillson**, Eric Klavins, Chris Myers, Jean Peccoud, and Giovanni Stracquadanio. 2023. Introduction to the Special Issue on BioFoundries and Cloud Laboratories. *J. Emerg. Technol. Comput. Syst.* 19, 3, Article 26 (July 2023), 2 pages. <https://doi.org/10.1145/3609485>

Chen Y, Gin JW, Wang Y, de Raad M, Tan S, **Hillson NJ**, et al. (2023) "Alkaline-SDS cell lysis of microbes with acetone protein precipitation for proteomic sample preparation in 96-well plate format Alkaline-SDS cell lysis of microbes with acetone protein precipitation for proteomic sample preparation in 96-well plate format. *PLOS ONE* 18(7): e0288102. <https://doi.org/10.1371/journal.pone.0288102>

Zhao, Huimin, **Hillson, Nathan**, Kleese van Dam, Kerstin, and Tanjore, Deepti. (2023) "Artificial Intelligence and Machine Learning for Bioenergy Research: Opportunities and Challenges", DOE/SC-0211. U.S. Department of Energy Office of Science and Office of Energy Efficiency and Renewable Energy. <https://doi.org/10.2172/1968870>

Hillson, N., Harmon-Smith, M., Simirenko, L., Greene, D., and Brink, K. (2023). Biorisk Management Case Study: United States Department of Energy Joint Genome Institute. Stanford Digital Repository. <https://doi.org/10.25740/rr427mq4842>.

Hector G Martin, Tijana Radivojevic, Jeremy Zucker, Kristofer Bouchard, Jess Sustarich, Sean Peisert, Dan Arnold, **Nathan Hillson**, Gyorgy Babnigg, Jose M Marti, Christopher J Mungall, Gregg T Beckham, Lucas Waldburger, James Carothers, ShivShankar Sundaram, Deb Agarwal, Blake A Simmons, Tyler Backman, Deepanwita Banerjee, Deepti Tanjore, Lavanya Ramakrishnan, Anup Singh. (2023) "Perspectives for self-driving labs in synthetic biology", *Current Opinion in Biotechnology*, <https://doi.org/10.1016/j.copbio.2022.102881>

Kosuke Iwai, Maren Wehrs, Megan Garber, Jess Sustarich, Lauren Washburn, Zachary Costello, Peter W. Kim, David Ando, William R. Gaillard, **Nathan J. Hillson**, Paul D. Adams, Aindrila Mukhopadhyay, Hector Garcia Martin & Anup K. (2022) "Scalable and automated CRISPR-based strain engineering using droplet microfluidics" *Microsyst Nanoeng* 8, 31. <https://doi.org/10.1038/s41378-022-00357-3>

Chen Y, Kaplan Lease N, Gin JW, Ogorzalek TL, Adams PD, **Hillson NJ**, et al. (2022) "Modular automated bottom-up proteomic sample preparation for high-throughput Applications". *PLoS ONE* 17(2): e0264467. <https://doi.org/10.1371/journal.pone.0264467>

Rebecca Mackelprang, Katarzyna P. Adamala, Emily R. Aurand, James C. Diggans, Andrew D. Ellington, Samuel Weiss Evans, J. L. Clem Fortman, **Nathan J. Hillson**, Albert W. Hinman, Farren J. Isaacs, June I. Medford, Shadi Mamaghani, Tae Seok Moon, Megan J. Palmer, Jean Peccoud, Elizabeth A. Vitalis, India Hook-Barnard, and Douglas C. Friedman. (2022) "Making Security Viral: Shifting Engineering Biology Culture and Publishing". *ACS Synthetic Biology* 11 (2), 522-527. DOI: 10.1021/acssynbio.1c00324

Hector A. Plahar, Thomas N. Rich, Stephen D. Lane, William C. Morrell, Leanne Springthorpe, Oge Nnadi, Elena Aravina, Tiffany Dai, Michael J. Fero, **Nathan J. Hillson**, and Christopher J. Petzold. (2021) "BioParts—A Biological Parts Search Portal and Updates to the ICE Parts Registry Software Platform" *ACS Synthetic Biology* DOI: 10.1021/acssynbio.1c00263

Pomraning, K., Dai, Z., Munoz, N., Kim, Y., Gao, Y., Deng, S., Kim, J., Hofstad, B., Swita, M., Lemmon, T., Collett, J., Panisko, E., Webb-Robertson, B., Zucker, J., Nicora, C., De Paoli, H., Baker, S., Burnum-Johnson, K., **Hillson, N.**, and Magnuson, J. (2021) "Integration of Proteomics and Metabolomics Into the Design, Build, Test, Learn Cycle to Improve 3-Hydroxypropionic Acid Production in *Aspergillus pseudoterreus*. *Frontiers in Bioengineering and Biotechnology*. <https://doi.org/10.3389/fbioe.2021.603832>

Somtirtha Roy, Tijana Radivojevic, Mark Forrer, Jose Manuel Marti, Vamshi Jonnalagadda, Tyler Backman, William Morrell, Hector Plahar, Joonhoon Kim, **Nathan Hillson**, and Hector Garcia Martin. (2021) "Multiomics Data Collection, Visualization, and Utilization for Guiding Metabolic Engineering". *Frontiers in Bioengineering and Biotechnology* 9, 45. <https://doi.org/10.3389/fbioe.2021.612893>

Chris Lawson, Jose Manuel Martí, Tijana Radivojevic, Sai Vamshi R. Jonnalagadda, Reinhard Gentz, **Nathan J. Hillson**, Sean Peisert, Joonhoon Kim, Blake A. Simmons, Christopher J. Petzold, Steven W. Singer, Aindrila Mukhopadhyay, Deepti Tanjore, Josh Dunn, and Hector Garcia Martin. (2020) "Machine learning for metabolic engineering: A review" *Metabolic Engineering* <https://doi.org/10.1016/j.ymben.2020.10.005>

Garima Goyal, Nick Elsbree, Michael Fero, **Nathan J. Hillson**, and Gregory Linshiz. (2020) "Repurposing a microfluidic formulation device for automated DNA construction" *PLOS One* <https://doi.org/10.1371/journal.pone.0242157>

Gledon Doçi, Lukas Fuchs, Yash Kharbanda, Paul Schickling, Valentin Zulkower, **Nathan Hillson**, Ernst Oberortner, Neil Swainston, Johannes Kabisch. (2020) "DNA Scanner: a web application for comparing DNA synthesis feasibility, price, and turnaround time across vendors". *OUP Synthetic Biology*, ysaa011, <https://doi.org/10.1093/synbio/ysaa011>

Fero M.J., Craft J.K., Vu T., and **Hillson N.J.** (2020) "Combinatorial-Hierarchical DNA Library Design Using the TeselaGen DESIGN Module with j5". In: Chandran S., George K. (eds) *DNA Cloning and Assembly. Methods in Molecular Biology*, vol 2205. Humana, New York, NY. https://doi.org/10.1007/978-1-0716-0908-8_2

Ernst Oberortner, Robert Evans, Xianwei Meng, Sangeeta Nath, Hector Plahar, Lisa Simirenko, Angela Tarver, Samuel Deutsch, **Nathan J. Hillson**, and Jan-Fang Cheng. (2020) "An Integrated Computer-Aided Design and Manufacturing Workflow for Synthetic Biology". In: Chandran S., George K. (eds) DNA Cloning and Assembly. Methods in Molecular Biology, vol 2205. Humana, New York, NY. https://doi.org/10.1007/978-1-0716-0908-8_1

Sam Weiss Evans, Jacob Beal, Kavita Berger, Diederik A. Bleijs, Alessia Cagnetti, Francesca Ceroni, Gerald L. Epstein, Natália Garcia-Reyero, David R. Gillum, Graeme Harkess, **Nathan J. Hillson**, Petra A. M. Hogervorst, Jacob L. Jordan, Geneviève Lacroix, Rebecca Moritz, Seán S. ÓhÉigeartaigh, Megan J. Palmer, Mark W. J. van Passel. (2020) "Embrace experimentation in biosecurity governance" *Science* Vol. 368, Issue 6487, pp. 138-140 DOI: 10.1126/science.aba2932

Christopher B. Eiben, Tian Tian, Mitchell G. Thompson, Daniel Mendez-Perez, Nurgul Kaplan, Garima Goyal, Jennifer Chiniquy, **Nathan J. Hillson**, Taek Soon Lee, and Jay D. Keasling*. (2020) "Adenosine Triphosphate and Carbon Efficient Route to Second Generation Biofuel Isopentanol" *ACS Synth. Biol.* <https://doi.org/10.1021/acssynbio.9b00402>

Chiniquy J, Garber ME, Mukhopadhyay A, and **Hillson NJ***. (2020) "Fluorescent amplification for next generation sequencing (FA-NGS) library preparation". *BMC Genomics* 21, 85. <https://doi.org/10.1186/s12864-020-6481-8>

Sebastian Barthel, Sebastian Palluk, **Nathan J. Hillson**, Jay D. Keasling, and Daniel H. Arlow. (2020) "Enhancing Terminal Deoxynucleotidyl Transferase Activity on Substrates with 3' Terminal Structures for Enzymatic De Novo DNA Synthesis" *Genes* 11(1), 102 <https://www.mdpi.com/2073-4425/11/1/102>

Christopher B. Eiben, Tristan de Rond, Clayton Bloszies, Jennifer Gin, Jennifer Chiniquy, Edward E. K. Baidoo, Christopher J. Petzold, **Nathan J. Hillson**, Oliver Fiehn, Jay D. Keasling. (2019) "Mevalonate Pathway Promiscuity Enables Noncanonical Terpene Production", *ACS Synth. Biol.* <https://doi.org/10.1021/acssynbio.9b00230>

Chen Y, Guenther JM, Gin JW, Chan LJG, Costello Z, Ogorzalek TL, Tran HM, Blake-Hedges JM, Keasling JD, Adams PD, Garcia Martin H, **Hillson NJ**, Petzold CJ. (2019) "An automated 'cells-to-peptides' sample preparation workflow for high-throughput, quantitative proteomic assays of microbes". *J Proteome Res.* doi:10.1021/acs.jproteome.9b00455

Jesus F. Barajas, Ryan P. McAndrew, Mitchell G. Thompson, Tyler W. H. Backman, Bo Pang, Tristan de Rond, Jose H. Pereira, Veronica T. Benites, Héctor García Martín, Edward E. K. Baidoo, **Nathan J. Hillson**, Paul D. Adams, and Jay D. Keasling. (2019) "Structural insights into dehydratase substrate selection for the borrelidin and fluvirucin polyketide synthases" *J Ind Microbiol Biotechnol* <https://doi.org/10.1007/s10295-019-02189-z>

Paul Opgenorth, Zak Costello, Takuya Okada, Garima Goyal, Yan Chen, Jennifer Gin, Veronica T. Benites, Markus de Raad, Trent R. Northen, Kai Deng, Samuel Deutsch, Edward E.K. Baidoo, Christopher J. Petzold, **Nathan J Hillson**, Hector Garcia Martin, and Harry R Beller. (2019) "Lessons from two Design-Build-Test-Learn cycles of dodecanol production in Escherichia coli aided by machine learning". *ACS Synth. Biol.* DOI: 10.1021/acssynbio.9b00020

Nathan Hillson, Mark Caddick, Yizhi Cai, Jose A. Carrasco, Matthew Wook Chang, Natalie C. Curach, David J. Bell, Rosalind Le Feuvre, Douglas C. Friedman, Xiongfei Fu, Nicholas D. Gold, Markus J. Herrgård, Maciej B. Holowko, James R. Johnson, Richard A. Johnson, Jay D. Keasling, Richard I. Kitney, Akihiko Kondo, Chenli Liu, Vincent J. J. Martin, Filippo Menolascina, Chiaki Ogino, Nicola J. Patron, Marilene Pavan, Chueh Loo Poh, Isak S. Pretorius, Susan J. Rosser, Nigel S. Scrutton, Marko Storch, Hille Tekotte, Evelyn Travník, Claudia E. Vickers, Wen Shan Yew, Yingjin Yuan, Huimin Zhao & Paul S. Freemont. (2019) "Building a global alliance of biofoundries". *Nature Communications*, 10:2040 <https://www.nature.com/articles/s41467-019-10079-2>

Garima Goyal, Zak Costello, Jorge Alonso Guitierrez, Aram Kang, Taek Soon Lee, Hector Garcia Martin, and **Nathan J. Hillson**. (2018) "Parallel Integration and Chromosomal Expansion of Metabolic Pathways" *ACS Synthetic Biology* DOI: 10.1021/acssynbio.8b00243

Thomas L. Ruegg, Jose H. Pereira, Joseph C. Chen, Andy DeGiovanni, Pavel Novichkov, Vivek K. Mutalik, Giovanni P. Tomaleri, Steven W. Singer, **Nathan J. Hillson**, Blake A. Simmons, Paul D. Adams & Michael P. Thelen. (2018) "Jungle Express is a versatile repressor system for tight transcriptional control". *Nature Communications* 9, 3617. <https://doi.org/10.1038/s41467-018-05857-3>

Sebastian Palluk, Daniel H. Arlow, Tristan de Rond, Rathin Bector, Justine S. Kang, Hratch M. Baghdassarian, Alisa N. Truong, Peter W. Kim, Anup K. Singh, **Nathan J. Hillson**, Jay D. Keasling. (2018) "De novo DNA synthesis using polymerase-tethered nucleotides" *Nature Biotech* doi:10.1038/nbt.4173

Jesus F. Barajas, Amin Zargar, Bo Pang, Veronica T. Benites, Jennifer Gin, Edward E. K. Baidoo, Christopher J. Petzold, **Nathan J. Hillson**, and Jay D. Keasling. (2018) "Biochemical Characterization of β -Amino Acid Incorporation in Fluvirucin B2 Biosynthesis". *ChemBioChem* 10.1002/cbic.201800169

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Nathan J. Hillson*, Hector A. Plahar, Jacob Beal*, and Ranjini Prithviraj. (2016) Improving Synthetic Biology Communication: Recommended Practices for Visual Depiction and Digital Submission of Genetic Designs. *ACS Synth. Biol.*, 5 (6), pp 449–451. DOI: 10.1021/acssynbio.6b00146

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