

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: Ying, Kejun

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

POSITION TITLE: K00 Postdoctoral Fellow

EDUCATION/TRAINING *(Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)*

INSTITUTION AND LOCATION	DEGREE <i>(if applicable)</i>	Completion Date MM/YYYY	FIELD OF STUDY
Sun Yat-Sen University, Guangzhou, China	B.S.	06/2019	Life Science
Harvard University, Cambridge, MA	M.S.	05/2024	Computational Science Engineering
Harvard University, Cambridge, MA	Ph.D.	05/2025	Biological Science in Public Health
Stanford University, Stanford, CA	Postdoc	Present	Neurology (Tony Wyss-Coray’s Lab)
University of Washington, Seattle, WA	Postdoc	Present	Protein Design (David Baker’s Lab, co-mentorship)

A. Personal Statement

My research vision is to establish cellular aging as a tractable computational problem with engineered solutions by integrating causal AI models with evolutionary protein design. I aim to understand fundamental mechanisms of how cells maintain physiological function across the lifespan and translate these insights into precision therapeutics.

My training bridges computational and experimental approaches to aging biology. During my Ph.D. with Vadim Gladyshev at Harvard, I developed causality-enriched epigenetic clocks, foundation models for DNA methylation, and AI platforms for systematic intervention discovery. My postdoctoral work with Tony Wyss-Coray at Stanford on plasma proteomics and with David Baker at the University of Washington on protein design has prepared me to combine computational discovery with experimental validation and therapeutic engineering. This interdisciplinary training positions me to address fundamental questions in biochemistry and biophysics using modern computational and systems biology approaches.

UCSF’s Department of Biochemistry and Biophysics provides an ideal environment for my research program. The department’s strength in structural biology, computational biology, and systems approaches aligns perfectly with my vision. I envision collaborations with faculty in cryo-EM for structural characterization of engineered proteins, mass spectrometry experts for proteomic validation, and computational biologists for developing causal AI frameworks. The Mission Bay Campus’s state-of-the-art facilities, including advanced microscopy, cryo-EM, and high-throughput screening capabilities, would enable comprehensive investigation of cellular aging mechanisms and validation of designed therapeutics. I would benefit from UCSF’s collaborative culture and contribute by bringing expertise in causal inference, foundation models, and protein engineering while mentoring diverse students in quantitative approaches to fundamental biology.

My future research program will develop causal multimodal foundation models to identify aging mechanisms, build AI agent platforms for intervention discovery, and engineer therapeutic proteins informed by evolutionary

principles. This integrated approach will advance our understanding of cellular physiology while creating new tools for intervention in age-related cellular dysfunction.

B. Positions, Scientific Appointments, and Honors

Positions and Scientific Appointments

2025–Present	Postdoctoral Researcher, Tony Wyss-Coray's Lab & David Baker's Lab (co-mentorship), Stanford University & University of Washington
2025–Present	Co-Founder, Avinasi Labs, San Francisco, CA
2020–2025	Graduate Researcher, Vadim Gladyshev's Lab, Harvard Medical School, Boston, MA
2020	Graduate Researcher (Rotation), Eric Greer's Lab, Boston Children's Hospital, Boston, MA
2019	Graduate Researcher (Rotation), David Sinclair's Lab, Harvard Medical School, Boston, MA
2019	Graduate Researcher (Rotation), Brendan Manning's Lab, Harvard T. H. Chan School of Public Health, Boston, MA
2018–2019	Undergraduate Researcher, Zhou Songyang's Lab, Sun Yat-Sen University, Guangzhou, China
2018	Undergraduate Researcher, Xia Shen's Lab, University of Edinburgh, Edinburgh, UK
2018	Undergraduate Researcher, Matt Kaeberlein's Lab, University of Washington, Seattle, WA
2018	Undergraduate Researcher, Judith Campisi's Lab, Buck Institute for Research on Aging, Novato, CA
2017	Undergraduate Researcher, Danica Chen's Lab, University of California, Berkeley, Berkeley, CA
2015–2017	Undergraduate Researcher, Yikang Rong's Lab, Sun Yat-Sen University, Guangzhou, China

Other Experience and Professional Memberships

2023–Present	Core Member, Biomarkers of Aging Consortium
2024–Present	Organizer, Biomarker of Aging Challenge
2024–2025	President, Harvard Interdisciplinary Discussion on Disease and Health
2024	Organizing Committee Member, Biomarker of Aging Symposium 2024
2023	Organizing Committee Member, Biomarker of Aging Symposium 2023
2023–2024	Mentor, Yuanpei Young Scholars Program

Honors

2025	Semifinalist, Harvard President's Innovation Challenge, Health Care and Life Sciences Track
2024–2028	NIH/NIA F99/K00 Transition to Aging Research Award (Perfect Impact Score of 10)
2023	Best Poster Award, Inaugural Biomarker of Aging Symposium
2022	Best Poster Award, Gordon Research Conference, Systems Aging
2021	Hackathon Winner, Longevity Hackathon, VitaDAO
2016–2019	Yan-Sen Honor School Program, Sun Yat-Sen University
2016–2019	Yan-Sen Scholarship, Sun Yat-Sen University

C. Contribution to Science

1. **Development of causality-informed epigenetic biomarkers of aging:** A key focus of my research has been developing advanced biomarkers of aging based on DNA methylation patterns. I developed the first causality-informed DNA methylation clock for humans that distinguishes between damaging and adaptive aging changes (Ying et al., 2024a). Using Mendelian Randomization and machine learning models, I identified CpG sites causally linked to aging-related traits, leading to the development of two novel epigenetic clocks: DamAge and AdaptAge. I demonstrated that DamAge is associated with adverse conditions including increased mortality risk, while AdaptAge is linked to beneficial adaptations. This work provides new insights into the causal pathways involved in biological aging and creates more specific targets for potential interventions. The significance of this approach was recognized through an invited perspective on causal inference methods for epigenetic aging (Ying, 2024), and has led to a patent application for mapping CpG sites to quantify aging traits.

- a. **Ying, K.**, Liu, H., Tarkhov, A. E., Sadler, M. C., Lu, A. T., Moqri, M., Horvath, S., Kutalik, Z., Shen, X., & Gladyshev, V. N. (2024). Causality-enriched epigenetic age uncouples damage and adaptation. *Nature Aging (Featured on the February Cover)*, 1–16. PMID: 38243142; PMCID: PMC11070280
- b. **Ying, K.** (2024). Causal inference for epigenetic ageing. *Nature Reviews Genetics*, 26(1), 3–3. PMID: 39472743
- c. Tarkhov, A. E., Lindstrom-Vautrin, T., Zhang, S., **Ying, K.**, Moqri, M., Zhang, B., Tyshkovskiy, A., Levy, O., & Gladyshev, V. N. (2024). Nature of epigenetic aging from a single-cell perspective. *Nature Aging*, 4(6), 854–870. PMID: 38724733
- d. V. N. Gladyshev, **K. Ying**, "Mapping CpG sites to quantify aging traits" (2024). *WO2024039905A2* (Patent)

2. **Creation of comprehensive platforms for biological age profiling:** I have developed critical resources for the aging research community that facilitate systematic profiling and exploration of biological age. I created ClockBase (www.clockbase.org), a platform featuring biological age estimates based on multiple aging clock models applied to over 3,000 DNA methylation and gene expression datasets. Building on this foundation, I developed autonomous AI agent frameworks that systematically mine aging interventions from millions of molecular profiles (Ying et al., 2025a). I contributed to establishing unified frameworks for systematic curation and evaluation of aging biomarkers (Ying et al., 2025b). Additionally, I have developed MethylGPT, a foundation model for the DNA methylome (Ying et al., 2024b), advancing our ability to analyze epigenetic patterns in aging research. My work on biological age measurement has been recognized through invitations to present at the Keystone Symposia on Aging and other prominent conferences, and has resulted in a provisional patent application for high-dimensional measurement of biological age.

- a. **Ying, K.**, Tyshkovskiy, A., Moldakozhayev, A., Wang, H., Magalhaes, C. G. D., Iqbal, S., Garza, A. E., Tskhay, A., Poganik, J. R., Huang, K., Qu, Y., Glubokov, D., Jin, C., Lee, D., Liu, H., Leote, C., Trapp, A., Camillo, L. P. de L., Kerepesi, C., Moqri, M., Zhang, O., Jiang, K., Galkin, F., Zhavoronkov, A., Raamsdonk, J. M. V., Wang, M., Cong, L., Regev, A., Leskovec, J., Wyss-Coray, T., & Gladyshev, V. N. (2025). Autonomous AI agents discover aging interventions from millions of molecular profiles. *bioRxiv*, 2023.02.28.530532.
- b. **Ying, K.**, Paulson, S., Eames, A., Tyshkovskiy, A., Li, S., Eynon, N., Jacques, M., Grolaux, R., Seale, K., Jacques, E., Goeminne, L. J. E., Cipriano, A., Perez-Guevara, M., Emamifar, M., Casas Martínez, M., Kwon, D., Kosheleva, A., Snyder, M., Gobel, D., Herzog, C., McCartney, D. L., Marioni, R. E., Lasky-Su, J., Poganik, J. R., Moqri, M., & Gladyshev, V. N. (2025). A unified framework for systematic curation and evaluation of aging biomarkers. *Nature Aging*, 1–17.
- c. **Ying, K.**, Song, J., Cui, H., Zhang, Y., Li, S., Chen, X., Liu, H., Eames, A., McCartney, D. L., Marioni, R. E., Poganik, J. R., Moqri, M., Wang, B., & Gladyshev, V. N. (2024). MethylGPT: a foundation model for the DNA methylome. *bioRxiv*. PMID: 39574641; PMCID: PMC11580859
- d. **Ying, K.**, Tyshkovskiy, A., Chen, Q., Latorre-Crespo, E., Zhang, B., Liu, H., Matei-Dediu, B., Poganik, J. R., Moqri, M., Kirschner, K., Lasky-Su, J., & Gladyshev, V. N. (2024). High-dimensional Ageome Representations of Biological Aging across Functional Modules. *bioRxiv*. PMID: 39345525; PMCID: PMC11429788
- e. V. N. Gladyshev, **K. Ying**, "High-dimensional measurement of biological age" (2024). *Provisional Patent Application*

3. **Investigating aging and genetics of longevity in relation to immunity and disease:** My research has examined relationships between aging, genetics, and immunity, particularly in the context of diseases like COVID-19. I utilized Mendelian Randomization and multi-instrument analyses to investigate how genetic factors associated with longevity influence susceptibility to COVID-19 (Ying et al., 2021). I found that genetic variations supporting longer life are associated with lower risk of COVID-19 infection and hospitalization. I also identified key pathways in age-related COVID-19 severity, including Notch signaling and immune system development. In more recent work, I examined patterns of loss-of-function germline mutations in centenarians (Ying et al., 2024c), identifying potential longevity genes that may offer protection against age-related diseases. These findings provide new insights into the genetic basis of longevity and age-related disease susceptibility.

- a. **Ying, K.**, Zhai, R., Pyrkov, T. V., Shindyapina, A. V., Mariotti, M., Fedichev, P. O., Shen, X., & Gladyshev, V. N. (2021). Genetic and phenotypic analysis of the causal relationship between aging and COVID-19. *Communications Medicine*, 1(1), 35. PMID: 35602207; PMCID: PMC9053191
- b. **Ying, K.**, Castro, J. P., Shindyapina, A. V., Tyshkovskiy, A., Moqri, M., Goeminne, L. J. E., Milman, S., Zhang, Z. D., Barzilai, N., & Gladyshev, V. N. (2024). Depletion of loss-of-function germline mutations in centenarians reveals longevity genes. *Nature Communications*, 15(1), 9030. PMID: 39424787; PMCID: PMC11489729
- c. Yang, Z., Macdonald-Dunlop, E., Chen, J., Zhai, R., Li, T., Richmond, A., Klarić, L., Pirastu, N., Ning, Z., Zheng, C., Wang, Y., Huang, T., He, Y., Guo, H., **Ying, K.**, Gustafsson, S., ... Shen, X. (2022). Genetic Landscape of the ACE2 Coronavirus Receptor. *Circulation*, 145(18), 1398–1411. PMID: 35387486; PMCID: PMC9047645
- d. Castro, J. P., Shindyapina, A. V., Barbieri, A., **Ying, K.**, Strelkova, O. S., Paulo, J. A., Tyshkovskiy, A., Meinl, R., Kerepesi, C., Petrashen, A. P., Mariotti, M., Meer, M. V., Hu, Y., Karamyshev, A., Losyev, G., Galhardo, M., Logarinho, E., Indzhykulian, A. A., Gygi, S. P., ... Gladyshev, V. N. (2024). Age-associated clonal B cells drive B cell lymphoma in mice. *Nature Aging*, 4(10), 1403–1417. PMID: 39117982

4. **Exploring causal epigenetic alterations in aging and age-related conditions:** Understanding whether epigenetic changes causally affect lifespan and aging-related outcomes has been a central theme in my research. During my rotation in Dr. Eric Greer's lab, I studied RNA modifications in *C. elegans* using untargeted Mass spectrometry, contributing to research showing that 18S rRNA methylation can affect longevity and stress resistance across generations (Liberman et al., 2023). For my thesis work, I implemented genetic-based causal inference methods to identify DNA methylation sites affecting lifespan, differentiating damaging from adaptive epigenetic changes during aging. I also contributed to understanding DNAm clocks at the single-cell level through analysis of single-cell DNA methylation data (Zhang et al., 2022), helping annotate aging-related loci and elucidating the mechanisms underlying these biomarkers. The significance of my research in this area has been recognized with an NIH F99/K00 Transition to Aging Research award for predoctoral students.

- a. Liberman, N., Rothi, M. H., Gerashchenko, M. V., Zorbas, C., Boulias, K., MacWhinnie, F. G., **Ying, A. K.**, Flood Taylor, A., Al Haddad, J., Shibuya, H., Roach, L., Dong, A., Dellacona, S., Lafontaine, D. L. J., Gladyshev, V. N., & Greer, E. L. (2023). 18S rRNA methyltransferases DIMT1 and BUD23 drive intergenerational hormesis. *Molecular Cell*, 83(18), 3268–3282.e7. PMID: 37689068; PMCID: PMC11990152
- b. Li, T., Ning, Z., Yang, Z., Zhai, R., Zheng, C., Xu, W., Wang, Y., **Ying, K.**, Chen, Y., & Shen, X. (2021). Total genetic contribution assessment across the human genome. *Nature Communications*, 12(1), 2845. PMID: 33990588; PMCID: PMC8121943
- c. Zhang, B., Tarkhov, A. E., Ratzan, W., **Ying, K.**, Moqri, M., Poganik, J. R., Barre, B., Trapp, A., Zoller, J. A., Haghani, A., Horvath, S., Peshkin, L., & Gladyshev, V. N. (2022). *Epigenetic profiling and incidence of disrupted development point to gastrulation as aging ground zero in Xenopus laevis*. bioRxiv.
- d. Rothi, M. H., Sarkar, G. C., Al Haddad, J., Mitchell, W., **Ying, K.**, Pohl, N., Sotomayor-Mena, R. G., Nattale, J., Dellacona, S., Gladyshev, V. N., & Greer, E. L. (2024). The 18S rRNA Methyltransferase DIMT-1 Regulates Lifespan in the Germline Later in Life. *bioRxiv*. PMID: 38798397; PMCID: PMC11118296

5. **Leadership in aging biomarker standardization and community efforts:** I have contributed to establishing standards and community infrastructure in the aging biomarker field. As an organizing committee member for the Biomarker of Aging Symposium and organizer of the Biomarker of Aging Challenge (Ying et al., 2024d), I have helped develop frameworks for systematic evaluation of aging biomarkers. I contributed to establishing unified frameworks for curation and evaluation of aging biomarkers (Ying et al., 2025). As a member of the Biomarkers of Aging Consortium, I have co-authored papers addressing challenges in biomarker translation (Biomarkers of Aging Consortium et al., 2024). I have also developed computational tools to support aging research.

- a. **Ying, K.**, Paulson, S., Eames, A., Tyshkovskiy, A., Li, S., Eynon, N., Jacques, M., Grolaux, R., Seale,

- K., Jacques, E., Goeminne, L. J. E., Cipriano, A., Perez-Guevara, M., Emamifar, M., Casas Martínez, M., Kwon, D., Kosheleva, A., Snyder, M., Gobel, D., Herzog, C., McCartney, D. L., Marioni, R. E., Lasky-Su, J., Poganik, J. R., Moqri, M., & Gladyshev, V. N. (2025). A unified framework for systematic curation and evaluation of aging biomarkers. *Nature Aging*, 1–17.
- b. **Ying, K.**, Paulson, S., Reinhard, J., Camillo, L. P. L., Trauble, J., Jokiel, S., Biomarkers of Aging Consortium, Gobel, D., Herzog, C., Poganik, J. R., Moqri, M., & Gladyshev, V. N. (2024). An Open Competition for Biomarkers of Aging. *bioRxiv*. PMID: 39554132; PMCID: PMC11565782
- c. **Biomarkers of Aging Consortium**, Herzog, C. M. S., Goeminne, L. J. E., Poganik, J. R., Barzilai, N., Belsky, D. W., Betts-LaCroix, J., Chen, B. H., Chen, M., Cohen, A. A., Cummings, S. R., Fedichev, P. O., Ferrucci, L., Fleming, A., Fortney, K., Furman, D., Gorbunova, V., Higgins-Chen, A., Hood, L., Horvath, S., ... Gladyshev, V. N. (2024). Challenges and recommendations for the translation of biomarkers of aging. *Nature Aging*, 1–12. PMID: 39660064; PMCID: PMC11630784
- d. Moqri, M., Herzog, C., Poganik, J. R., **Biomarkers of Aging Consortium**, Justice, J., Belsky, D. W., Higgins-Chen, A., Moskalev, A., Fuellen, G., Cohen, A. A., Bautmans, I., Widschwendter, M., Ding, J., Fleming, A., Mannick, J., Han, J.-D. J., Zhavoronkov, A., Barzilai, N., Kaeberlein, M., ... Gladyshev, V. N. (2023). Biomarkers of aging for the identification and evaluation of longevity interventions. *Cell*, 186(18), 3758–3775. PMCID: PMC11090477

Complete List of Published Work:

<https://www.ncbi.nlm.nih.gov/myncbi/kejun.ying.1/bibliography/public/>