# Keisuke YANAGISAWA

Assistant Professor, School of Computing, Tokyo Institute of Technology



<b>Current Status:</b>	Assistant Professor
Address:	2-12-1 W8-76, Ookayama, Meguro-ku, Tokyo, 152-8550 Japan
Phone:	+81-3-5734-3645 (Office)
Email:	yanagisawa [at] c.titech.ac.jp
URL:	https://keisuke-yanagisawa.github.io/
Date of Birth:	13 May 1991
Place of Birth:	Tokyo, Japan
Citizenship:	Japan

## **Education:**

04/2016-03/2019	Department of Computer Science, School of Computing,
	Tokyo Institute of Technology, D.Eng.
04/2014-03/2016	Department of Computer Science, Graduate School of Information
	Science and Engineering, Tokyo Institute of Technology, M.Eng.
04/2010-03/2014	Department of Computer Science, Faculty of Engineering,
	Tokyo Institute of Technology, B.Eng.

# **Employment:**

03/2020-present	Assistant Professor, Tokyo Institute of Technology
11/2019-03/2020	Part-time Lecturer, Tokyo Institute of Technology
04/2019-03/2020	JSPS Research Fellow (PD), Japan Society for the Promotion of Science
04/2017-03/2019	JSPS Research Fellow (DC2), Japan Society for the Promotion
	of Science

## Grants:

FY2022-FY2024	"Development of chemical substructure-based virtual screening method
	for huge compound library", JSPS KAKENHI (grant-in-aid for
	Scientific Research (B)), 22H03684, 17 030 000 yen
	(PI: Yutaka Akiyama)

- FY2020-FY2022 "Comprehensive prediction of cryptic binding sites by multi-task deep learning", JSPS KAKENHI (grant-in-aid for young researchers), 20K19917, PI, \$40 000
- FY2020 "Computer-aided lead optimization with cosolvent molecular dynamics (CMD)", Grant for young researcher, School of Computing, Tokyo Institute of Technology, PI, \$4 700
- FY2019 "Improvement of cosolvent MD which enables the systematic search of binding sites and the novel screening way of drug candidates", JSPS
  KAKENHI (grant-in-aid for JSPS fellows), 19J00878, PI, \$13 000
- FY2017-FY2018 "Development of divide-and-conquer based docking method using common partial structures of hundreds of millions of compounds", JSPS KAKENHI (grant-in-aid for JSPS fellows), 17J06897, PI, \$19 000

### Awards:

03/2021	2019 Education Award of Excellence,
	Tokyo Institute of Technology (as a member out of 17 members)
02/2020	Seiichi Tejima Doctoral Dissertation Award,
	Tokyo Institute of Technology.
12/2017	Grand Prize (Schrodinger K.K. Prize),
	4th Computer-Aided Drug Discovery Contest.
07/2015	Student Encouragement Prize,
	2nd Computer-Aided Drug Discovery Contest.
06/2015	2014 SIGBIO Best Student Presentation Award, IPSJ SIGBIO.
07/2014	Student Encouragement Prize,
	1st Computer-Aided Drug Discovery Contest.
03/2014	2013 Tokyo Institute of Technology Academic Excellence Awards,
	Tokyo Institute of Technology.

09/2009 Bronze Prize, 2009 High School Chemistry Grand Prix.

## **Current Research Interests:**

Molecular Dynamics, Computational Biology, Bioinformatics, Structural Biology, Protein-Ligand Docking, Protein-Ligand Interaction, Computational Drug Discovery, Virtual Screening, Structure-based Virtual Screening, Machine Learning

#### **Publications:**

#### Peer-reviewed Journal Papers

- <u>Keisuke Yanagisawa</u>, Rikuto Kubota, Yasushi Yoshikawa, Masahito Ohue, Yutaka Akiyama. "Effective protein-ligand docking strategy via fragment reuse and a proof-of-concept implementation", ACS Omega, 7: 30265-30274, 2022/08.
- <u>Keisuke Yanagisawa</u>, Ryunosuke Yoshino, Genki Kudo, Takatsugu Hirokawa. "Inverse Mixed-Solvent Molecular Dynamics for Visualization of the Residue Interaction Profile of Molecular Probes", *International Journal of Molecular Sciences*, 23: 4749, 2022/04.
- Kazuki Takabatake, <u>Keisuke Yanagisawa</u>, Yutaka Akiyama. "Solving Generalized Polyomino Puzzles Using the Ising Model", *Entropy*, 24: 354, 2022/02.
- Jianan Li, <u>Keisuke Yanagisawa</u>, Yasushi Yoshikawa, Masahito Ohue, Yutaka Akiyama. "Plasma protein binding prediction focusing on residue-level features and circularity of cyclic peptides by deep learning", *Bioinformatics*, 38: 1110-1117, 2022/02.
- <u>Keisuke Yanagisawa</u>. "Virtual Screening Methods with a Protein Tertiary Structure for Drug Discovery", *JSBi Bioinformatics Review*, 2: 76-86, 2021/10. (in Japanese)
- Kazuki Takabatake, Kazuki Izawa, Motohiro Akikawa, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama. "Improved Large-Scale Homology Search by Two-step Seed Search Using Multiple Reduced Amino Acid Alphabets", *Genes*, 12, 1455, 2021.
- Masatake Sugita, Satoshi Sugiyama, Takuya Fujie, Yasushi Yoshikawa, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama. "Large-Scale Membrane Permeability Prediction of Cyclic Peptides Crossing a Lipid Bilayer Based on Enhanced Sampling Molecular Dynamics Simulations", *Journal of Chemical Information and Modeling*, 61, 3681–3695, 2021.
- Keisuke Yanagisawa, Yoshitaka Moriwaki, Tohru Terada, Kentaro Shimizu. "EXPRORER: Rational cosolvent set construction method for cosolvent molecular dynamics using large-scale computation", *Journal of Chemical Information and Modeling*, 61, 2744–2753, 2021.
- Masahiro Mochizuki, Shogo D. Suzuki, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama.
  "QEX: Target-specific druglikeness filter enhances ligand-based virtual screening", *Molecular Diversity*, 23, 11–18, 2019.
- Takashi Tajimi, Naoki Wakui, <u>Keisuke Yanagisawa</u>, Yasushi Yoshikawa, Masahito Ohue, Yutaka Akiyama: "Computational prediction of plasma protein binding of cyclic peptides from small molecule experimental data using sparse modeling techniques", *BMC Bioinformatics*, 19, 527, 2018. (14 pages)
- <u>Keisuke Yanagisawa</u>, Shunta Komine, Rikuto Kubota, Masahito Ohue, Yutaka Akiyama. "Optimization of memory use of fragment extension-based protein-ligand docking with an original fast minimum cost flow algorithm", *Computational Biology and Chemistry*, 74, 399–406, 2018.

- Takanori Hayashi, Yuri Matsuzaki, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama.
  "MEGADOCK-Web: an integrated database of high-throughput structure-based protein-protein interaction predictions", *BMC Bioinformatics*, 19, 62, 2018. (12 pages)
- <u>Keisuke Yanagisawa</u>, Shunta Komine, Shogo D. Suzuki, Masahito Ohue, Takashi Ishida, Yutaka Akiyama. "Spresso: An ultrafast compound pre-screening method based on compound decomposition", *Bioinformatics*, 33, 3836–3843, 2017.
- 14. Shuntaro Chiba, Takashi Ishida, Kazuyoshi Ikeda, Masahiro Mochizuki, Reiji Teramoto, Y-h. Taguchi, Mitsuo Iwadate, Hideaki Umeyama, Chandrasekaran Ramakrishnan, A. Mary Thangakani, D. Velmurugan, M. Michael Gromiha, Tatsuya Okuno, Koya Kato, Shintaro Minami, George Chikenji, Shogo D. Suzuki, <u>Keisuke Yanagisawa</u>, Woong-Hee Shin, Daisuke Kihara, Kazuki Z. Yamamoto, Yoshitaka Moriwaki, Nobuaki Yasuo, Ryunosuke Yoshino, Sergey Zozulya, Petro Borysko, Roman Stavniichuk, Teruki Honma, Takatsugu Hirokawa, Yutaka Akiyama, Masakazu Sekijima, "An iterative compound screening contest method for identifying target protein inhibitors using the tyrosine-protein kinase Yes", *Scientific Reports*, 7, 12038, 2017. (13 pages)
- 15. Shuntaro Chiba, Kazuyoshi Ikeda, Takashi Ishida, M. Michael Gromiha, Y-h. Taguchi, Mitsuo Iwadate, Hideaki Umeyama, Kun-Yi Hsin, Hiroaki Kitano, Kazuki Yamamoto, Nobuyoshi Sugaya, Koya Kato, Tatsuya Okuno, George Chikenji, Masahiro Mochizuki, Nobuaki Yasuo, Ryunosuke Yoshino, <u>Keisuke Yanagisawa</u>, Tomohiro Ban, Reiji Teramoto, Chandrasekaran Ramakrishnan, A. Mary Thangakani, D. Velmurugan, Philip Prathipati, Junichi Ito, Yuko Tsuchiya, Kenji Mizuguchi, Teruki Honma, Takatsugu Hirokawa, Yutaka Akiyama, Masakazu Sekijima. "Identification of potential inhibitors based on compound proposal contest: Tyrosine-protein kinase Yes as a target", *Scientific Reports*, 5, 17209, 2015. (13 pages)
- Keisuke Yanagisawa, Takashi Ishida, Yutaka Akiyama. "Drug Clearance Pathway Prediction Based on Semi-supervised Learning", *IPSJ Transactions on Bioinformatics*, 8, 21–27, 2015.

#### Peer-reviewed International Conference Papers

- Kazuki Takabatake, Kazuki Izawa, Motohiro Akikawa, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama. "Improved Homology Search for Metagenomic Analysis by Two-Step Seed Search with Reduced Amino Acid Alphabets", *The 10th International Conference on Bioinformatics and Biomedical Science (ICBBS2021)*, 2021.
- Kazuya Isawa, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama. "Antisense oligonucleotide activity analysis based on opening and binding energies to targets", In *Proceedings of the 27th International Conference on Parallel and Distributed Processing Techniques and Applications (PDPTA'21)*, 2021.

- Masahito Ohue, Ryota Ii, <u>Keisuke Yanagisawa</u>, Yutaka Akiyama. "Molecular activity prediction using graph convolutional deep neural network considering distance on a molecular graph", In Proceedings of the 25th International Conference on Parallel and Distributed Processing Techniques and Applications (PDPTA'19), 2019.
- 20. Takashi Tajimi, Naoki Wakui, <u>Keisuke Yanagisawa</u>, Yasushi Yoshikawa, Masahito Ohue, Yutaka Akiyama. "Computational prediction of plasma protein binding of cyclic peptides from small molecule experimental data using sparse modeling techniques", *The 29th International Conference on Genome Informatics (GIW 2018)*, 2018.
- <u>Keisuke Yanagisawa</u>, Shunta Komine, Rikuto Kubota, Masahito Ohue, Yutaka Akiyama. "Optimization of memory use of fragment extension-based protein-ligand docking with an original fast minimum cost flow algorithm", *The 16th Asia Pacific Bioinformatics Conference* (APBC2018), 2018.
- Takanori Hayashi, Yuri Matsuzaki, <u>Keisuke Yanagisawa</u>, Masahito Ohue, Yutaka Akiyama.
  "MEGADOCK-Web: an integrated database of high-throughput structure-based protein-protein interaction predictions", *The 16th Asia Pacific Bioinformatics Conference (APBC2018)*, 2018.
- <u>Keisuke Yanagisawa</u>, Shunta Komine, Shogo D. Suzuki, Masahito Ohue, Takashi Ishida, Yutaka Akiyama. "ESPRESSO: An ultrafast compound pre-screening method based on compound decomposition", *The 27th International Conference on Genome Informatics (GIW 2016)*, 2016.