## Hidemasa Kondo

List of Publications by Year in descending order

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218381 223531 2,188 52 26 46 h-index citations g-index papers 53 53 53 2317 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Discovery of Hyperactive Antifreeze Protein from Phylogenetically Distant Beetles Questions Its Evolutionary Origin. International Journal of Molecular Sciences, 2021, 22, 3637.	1.8	8
2	Characterization of microbial antifreeze protein with intermediate activity suggests that a bound-water network is essential for hyperactivity. Scientific Reports, 2021, 11, 5971.	1.6	15
3	Subzero Nonfreezing Hypothermia with Insect Antifreeze Protein Dramatically Improves Survival Rate of Mammalian Cells. International Journal of Molecular Sciences, 2021, 22, 12680.	1.8	8
4	An Ice-Binding Protein from an Antarctic Ascomycete Is Fine-Tuned to Bind to Specific Water Molecules Located in the Ice Prism Planes. Biomolecules, 2020, 10, 759.	1.8	8
5	Fish-Derived Antifreeze Proteins and Antifreeze Glycoprotein Exhibit a Different Ice-Binding Property with Increasing Concentration. Biomolecules, 2020, 10, 423.	1.8	17
6	Calcium-Binding Generates the Semi-Clathrate Waters on a Type II Antifreeze Protein to Adsorb onto an Ice Crystal Surface. Biomolecules, 2019, 9, 162.	1.8	15
7	Ice recrystallization is strongly inhibited when antifreeze proteins bind to multiple ice planes. Scientific Reports, 2019, 9, 2212.	1.6	43
8	Iceâ€binding proteins from the fungus <i>Antarctomyces psychrotrophicus</i> possibly originate from two different bacteria through horizontal gene transfer. FEBS Journal, 2019, 286, 946-962.	2,2	25
9	Multiple binding modes of a moderate ice-binding protein from a polar microalga. Physical Chemistry Chemical Physics, 2018, 20, 25295-25303.	1.3	14
10	Polypentagonal ice-like water networks emerge solely in an activity-improved variant of ice-binding protein. Proceedings of the National Academy of Sciences of the United States of America, 2018, 115, 5456-5461.	3.3	32
11	Concentration-dependent oligomerization of an alpha-helical antifreeze polypeptide makes it hyperactive. Scientific Reports, 2017, 7, 42501.	1.6	35
12	Hydrophobic ice-binding sites confer hyperactivity of an antifreeze protein from a snow mold fungus. Biochemical Journal, 2016, 473, 4011-4026.	1.7	49
13	Title is missing!. Kagaku To Seibutsu, 2014, 52, 10-12.	0.0	O
14	Hyperactive antifreeze protein from an <scp>A</scp> ntarctic sea ice bacterium <i><scp>C</scp>olwellia</i> sp. has a compound iceâ€binding site without repetitive sequences. FEBS Journal, 2014, 281, 3576-3590.	2.2	64
15	Annealing condition influences thermal hysteresis of fungal type ice-binding proteins. Cryobiology, 2014, 68, 159-161.	0.3	17
16	Cold adaptation of fungi obtained from soil and lake sediment in the Skarvsnes ice-free area, Antarctica. FEMS Microbiology Letters, 2013, 346, 121-130.	0.7	64
17	Ice-binding site of snow mold fungus antifreeze protein deviates from structural regularity and high conservation. Proceedings of the National Academy of Sciences of the United States of America, 2012, 109, 9360-9365.	3.3	92
18	ä¸å‡ã,¿ãƒ³ãƒ'ã,¯è³ª 機能ã•応甓. Kagaku To Seibutsu, 2010, 48, 381-388.	0.0	2

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19	Comparison of functional properties of two fungal antifreeze proteins from ⟨i>Antarctomyces psychrotrophicus⟨ i> and ⟨i>Typhula ishikariensis⟨ i>. FEBS Journal, 2010, 277, 394-403.	2.2	76
20	Contribution of Asparagine Residues to the Stabilization of a Proteinaceous Antigen-Antibody Complex, HyHEL-10-Hen Egg White Lysozyme. Journal of Biological Chemistry, 2010, 285, 7686-7696.	1.6	18
21	Crystal structure and enhanced activity of a cutinaseâ€like enzyme from <i>Cryptococcus</i> sp. strain Sâ€2. Proteins: Structure, Function and Bioinformatics, 2009, 77, 710-717.	1.5	40
22	The crystal structure of a xyloglucanâ€specific endoâ€Î²â€1,4â€glucanase from <i>Geotrichum</i> sp. M128 xyloglucanase reveals a key amino acid residue for substrate specificity. FEBS Journal, 2009, 276, 5094-5100.	2.2	24
23	ONIOM Study of the Mechanism of the Enzymatic Hydrolysis of Biodegradable Plastics. Bulletin of the Chemical Society of Japan, 2009, 82, 338-346.	2.0	6
24	Critical contribution of VH–VL interaction to reshaping of an antibody: The case of humanization of antiâ€lysozyme antibody, HyHELâ€10. Protein Science, 2008, 17, 261-270.	3.1	48
25	Crystal Structure and Mutational Analysis of Ca2+-Independent Type II Antifreeze Protein from Longsnout Poacher, Brachyopsis rostratus. Journal of Molecular Biology, 2008, 382, 734-746.	2.0	66
26	Thermodynamic Consequences of Mutations in Vernier Zone Residues of a Humanized Anti-human Epidermal Growth Factor Receptor Murine Antibody, 528. Journal of Biological Chemistry, 2008, 283, 1156-1166.	1.6	86
27	Mass preparation and technological development of an antifreeze protein. Synthesiology, 2008, 1, 7-14.	0.2	23
28	ä¸å‡è›‹ç™½è³ªã®å§é‡ç²¾è£½ã;æ—°ãŸã³å¿œç"¨é—‹æ‹" ï¼å®Ÿç"¨åŒ—ã,'指å⁵ã™ã,‹è›‹ç™½è³ªç"究5 Syn	the <b>sia</b> logy	y, 2 <b>0</b> 08, 1, 7-1
29	Structural Consequences of Mutations in Interfacial Tyr Residues of a Protein Antigen-Antibody Complex. Journal of Biological Chemistry, 2007, 282, 6783-6791.	1.6	44
30	The Structural Basis for the Exo-mode of Action in GH74 Oligoxyloglucan Reducing End-specific Cellobiohydrolase. Journal of Molecular Biology, 2007, 370, 53-62.	2.0	52
31	A Human Antibody Fragment with High Affinity for Biodegradable Polymer Film. Bioconjugate Chemistry, 2007, 18, 645-651.	1.8	16
32	Relationship between the Size of the Bottleneck 15 $\tilde{A}$ from Iron in the Main Channel and the Reactivity of Catalase Corresponding to the Molecular Size of Substratesâ $\in$ <sub>i</sub> . Biochemistry, 2007, 46, 11-22.	1.2	54
33	Crystallization and preliminary X-ray crystallographic analysis of Ca2+-independent and Ca2+-dependent species of the type II antifreeze protein. Acta Crystallographica Section F: Structural Biology Communications, 2006, 62, 538-541.	0.7	7
34	Thermal Stabilization of Bacillus subtilis Family-11 Xylanase by Directed Evolution. Journal of Biological Chemistry, 2006, 281, 10236-10242.	1.6	106
35	Tetra-n-butylammonium bromide–water (1/38). Acta Crystallographica Section C: Crystal Structure Communications, 2005, 61, o65-o66.	0.4	214
36	Functions and Structures of Xyloglucan Hydrolases Belonging to Glycoside Hydrolase Family 74. Journal of Applied Glycoscience (1999), 2005, 52, 169-176.	0.3	3

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37	Tandem Repeat of a Seven-Bladed $\hat{l}^2$ -Propeller Domain in Oligoxyloglucan Reducing-End-Specific Cellobiohydrolase. Structure, 2004, 12, 1209-1217.	1.6	45
38	Can lipases hydrolyze a peptide bond?. Enzyme and Microbial Technology, 2003, 32, 655-657.	1.6	19
39	Crystallization and preliminary X-ray crystallographic study on a xyloglucan-specific exo- $\hat{l}^2$ -glycosidase, oligoxyloglucan reducing-end specific cellobiohydrolase. Acta Crystallographica Section D: Biological Crystallography, 2003, 59, 1838-1839.	2.5	6
40	Antifreeze proteins from snow mold fungi. Canadian Journal of Botany, 2003, 81, 1175-1181.	1.2	108
41	The Role of Hydrogen Bonding via Interfacial Water Molecules in Antigen-Antibody Complexation. Journal of Biological Chemistry, 2003, 278, 5410-5418.	1.6	60
42	Structural Consequences of Target Epitope-directed Functional Alteration of an Antibody. Journal of Biological Chemistry, 2003, 278, 24929-24936.	1.6	15
43	Alternate conformations observed in catalytic serine ofBacillus subtilislipase determined at 1.3â€Ã resolution. Acta Crystallographica Section D: Biological Crystallography, 2002, 58, 1168-1174.	2.5	67
44	NMR Analysis of Type III Antifreeze Protein Intramolecular Dimer. Journal of Biological Chemistry, 2001, 276, 1304-1310.	1.6	47
45	Structural Evidence for Entropic Contribution of Salt Bridge Formation to a Protein Antigen-Antibody Interaction. Journal of Biological Chemistry, 2001, 276, 23042-23050.	1.6	36
46	Crystal Structure of Anti-Hen Egg White Lysozyme Antibody (HyHEL-10) Fv-Antigen Complex. Journal of Biological Chemistry, 1999, 274, 27623-27631.	1.6	63
47	Highly efficient recovery of functional single-chain Fv fragments from inclusion bodies overexpressed in Escherichia coli by controlled introduction of oxidizing reagent—application to a human single-chain Fv fragment. Journal of Immunological Methods, 1998, 219, 119-129.	0.6	190
48	Crystallization and Preliminary X-Ray Studies of Two Serine Proteinase Inhibitors, BGIA and BGIT, from the Seeds of Bitter Gourd. Journal of Structural Biology, 1997, 120, 204-206.	1.3	2
49	Escherichia coli positive regulator OmpR has a large loop structure at the putative RNA polymerase interaction site. Nature Structural and Molecular Biology, 1997, 4, 28-31.	3.6	87
50	Crystallization and X-ray Studies of the DNA-binding Domain of OmpR Protein, a Positive Regulator Involved in Activation of Osmoregulatory Genes in Escherichia coli. Journal of Molecular Biology, 1994, 235, 780-782.	2.0	12
51	Identification of the fatty acid binding site on glutathione S-transferase P. Biochemical and Biophysical Research Communications, 1992, 189, 197-205.	1.0	16
52	The Absolute Stereochemistry of Okamurallene and Its Congeners, Halogenated C15Nonterpenoids from the Red AlgaLaurencia intricata. Chemistry Letters, 1991, 20, 33-34.	0.7	15