Isidore Rigoutsos

List of Publications by Year in descending order

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19636 12933 22,458 167 61 131 citations h-index g-index papers 173 173 173 32363 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	A Pattern-Based Method for the Identification of MicroRNA Binding Sites and Their Corresponding Heteroduplexes. Cell, 2006, 126, 1203-1217.	13.5	1,827
2	The Draft Genome of Ciona intestinalis: Insights into Chordate and Vertebrate Origins. Science, 2002, 298, 2157-2167.	6.0	1,539
3	Integrated Genomic Characterization of Pancreatic Ductal Adenocarcinoma. Cancer Cell, 2017, 32, 185-203.e13.	7.7	1,428
4	MicroRNAs to Nanog, Oct4 and Sox2 coding regions modulate embryonic stem cell differentiation. Nature, 2008, 455, 1124-1128.	13.7	1,288
5	Metagenomic and functional analysis of hindgut microbiota of a wood-feeding higher termite. Nature, 2007, 450, 560-565.	13.7	1,181
6	Dynamic changes in the human methylome during differentiation. Genome Research, 2010, 20, 320-331.	2.4	930
7	Coding-Independent Regulation of the Tumor Suppressor PTEN by Competing Endogenous mRNAs. Cell, 2011, 147, 344-357.	13.5	926
8	The complex transcriptional landscape of the anucleate human platelet. BMC Genomics, 2013, 14, 1.	1.2	913
9	The Expression of MicroRNA miR-107 Decreases Early in Alzheimer's Disease and May Accelerate Disease Progression through Regulation of \hat{l}^2 -Site Amyloid Precursor Protein-Cleaving Enzyme 1. Journal of Neuroscience, 2008, 28, 1213-1223.	1.7	745
10	The miR-17/92 cluster: a comprehensive update on its genomics, genetics, functions and increasingly important and numerous roles in health and disease. Cell Death and Differentiation, 2013, 20, 1603-1614.	5.0	722
11	Metagenomic analysis of two enhanced biological phosphorus removal (EBPR) sludge communities. Nature Biotechnology, 2006, 24, 1263-1269.	9.4	634
12	Accurate phylogenetic classification of variable-length DNA fragments. Nature Methods, 2007, 4, 63-72.	9.0	524
13	The amphioxus genome illuminates vertebrate origins and cephalochordate biology. Genome Research, 2008, 18, 1100-1111.	2.4	456
14	Combinatorial pattern discovery in biological sequences: The TEIRESIAS algorithm [published erratum appears in Bioinformatics 1998;14(2):229]. Bioinformatics, 1998, 14, 55-67.	1.8	451
15	Geometric hashing: an overview. IEEE Computational Science and Engineering, 1997, 4, 10-21.	0.6	413
16	Analysis of 13 cell types reveals evidence for the expression of numerous novel primate- and tissue-specific microRNAs. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E1106-15.	3.3	376
17	Use of simulated data sets to evaluate the fidelity of metagenomic processing methods. Nature Methods, 2007, 4, 495-500.	9.0	322
18	Sex hormone-dependent tRNA halves enhance cell proliferation in breast and prostate cancers. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E3816-25.	3.3	295

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19	Genomic analysis of immunity in a Urochordate and the emergence of the vertebrate immune system: "waiting for Godot― Immunogenetics, 2003, 55, 570-581.	1.2	278
20	High-resolution metagenomics targets specific functional types in complex microbial communities. Nature Biotechnology, 2008, 26, 1029-1034.	9.4	254
21	The microRNAs within the DLK1-DIO3 genomic region: involvement in disease pathogenesis. Cellular and Molecular Life Sciences, 2013, 70, 795-814.	2.4	246
22	An algorithm for point clustering and grid generation. IEEE Transactions on Systems, Man, and Cybernetics, 1991, 21, 1278-1286.	0.9	241
23	Arginine methylation of Piwi proteins catalysed by dPRMT5 is required for Ago3 and Aub stability. Nature Cell Biology, 2009, 11 , $652-658$.	4.6	219
24	A linguistic model for the rational design of antimicrobial peptides. Nature, 2006, 443, 867-869.	13.7	214
25	Small RNA Sequencing across Diverse Biofluids Identifies Optimal Methods for exRNA Isolation. Cell, 2019, 177, 446-462.e16.	13.5	214
26	MicroRNA-134 Modulates the Differentiation of Mouse Embryonic Stem Cells, Where It Causes Post-Transcriptional Attenuation of Nanog and LRH1. Stem Cells, 2008, 26, 17-29.	1.4	213
27	New Tricks for Animal MicroRNAs: Targeting of Amino Acid Coding Regions at Conserved and Nonconserved Sites. Cancer Research, 2009, 69, 3245-3248.	0.4	207
28	Interactive exploration of RNA22 microRNA target predictions. Bioinformatics, 2012, 28, 3322-3323.	1.8	193
29	Reevaluation of human cytomegalovirus coding potential. Proceedings of the National Academy of Sciences of the United States of America, 2003, 100, 13585-13590.	3.3	171
30	Knowledge about the presence or absence of miRNA isoforms (isomiRs) can successfully discriminate amongst 32 TCGA cancer types. Nucleic Acids Research, 2017, 45, 2973-2985.	6.5	158
31	Evolution of distinct EGF domains with specific functions. Protein Science, 2005, 14, 1091-1103.	3.1	155
32	MINTbase v2.0: a comprehensive database for tRNA-derived fragments that includes nuclear and mitochondrial fragments from all The Cancer Genome Atlas projects. Nucleic Acids Research, 2018, 46, D152-D159.	6.5	155
33	Data mining and clinical data repositories: Insights from a 667,000 patient data set. Computers in Biology and Medicine, 2006, 36, 1351-1377.	3.9	150
34	Dissecting tRNA-derived fragment complexities using personalized transcriptomes reveals novel fragment classes and unexpected dependencies. Oncotarget, 2015, 6, 24797-24822.	0.8	146
35	miR-34a contributes to megakaryocytic differentiation of K562 cells independently of p53. Blood, 2009, 114, 2181-2192.	0.6	142
36	Beyond the one-locus-one-miRNA paradigm: microRNA isoforms enable deeper insights into breast cancer heterogeneity. Nucleic Acids Research, 2015, 43, 9158-9175.	6.5	134

3

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37	MiR-103a-3p targets the 5′ UTR of <i>GPRC5A</i> in pancreatic cells. Rna, 2014, 20, 1431-1439.	1.6	129
38	MINTmap: fast and exhaustive profiling of nuclear and mitochondrial tRNA fragments from short RNA-seq data. Scientific Reports, 2017, 7, 41184.	1.6	123
39	Short blocks from the noncoding parts of the human genome have instances within nearly all known genes and relate to biological processes. Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 6605-6610.	3.3	111
40	On the expanding roles of tRNA fragments in modulating cell behavior. Nucleic Acids Research, 2020, 48, 9433-9448.	6.5	107
41	Non-α-helical elements modulate polytopic membrane protein architecture11Edited by G. Von Heijne. Journal of Molecular Biology, 2001, 306, 349-362.	2.0	105
42	IsomiR expression profiles in human lymphoblastoid cell lines exhibit population and gender dependencies. Oncotarget, 2014, 5, 8790-8802.	0.8	103
43	What's in the mix: phylogenetic classification of metagenome sequence samples. Current Opinion in Microbiology, 2007, 10, 499-503.	2.3	100
44	N-BLR, a primate-specific non-coding transcript leads to colorectal cancer invasion and migration. Genome Biology, 2017, 18, 98.	3.8	97
45	"Off-Spotter― very fast and exhaustive enumeration of genomic lookalikes for designing CRISPR/Cas guide RNAs. Biology Direct, 2015, 10, 4.	1.9	96
46	The mRNA-binding protein HuR promotes hypoxia-induced chemoresistance through posttranscriptional regulation of the proto-oncogene PIM1 in pancreatic cancer cells. Oncogene, 2016, 35, 2529-2541.	2.6	96
47	HuR Posttranscriptionally Regulates WEE1: Implications for the DNA Damage Response in Pancreatic Cancer Cells. Cancer Research, 2014, 74, 1128-1140.	0.4	91
48	Race Disparities in the Contribution of miRNA Isoforms and tRNA-Derived Fragments to Triple-Negative Breast Cancer. Cancer Research, 2018, 78, 1140-1154.	0.4	90
49	MINTbase: a framework for the interactive exploration of mitochondrial and nuclear tRNA fragments. Bioinformatics, 2016, 32, 2481-2489.	1.8	89
50	Argonaute CLIP-Seq reveals miRNA targetome diversity across tissue types. Scientific Reports, 2014, 4, 5947.	1.6	88
51	MicroRNA 125a and its regulation of the p53 tumor suppressor gene. FEBS Letters, 2009, 583, 3725-3730.	1.3	87
52	YAMAT-seq: an efficient method for high-throughput sequencing of mature transfer RNAs. Nucleic Acids Research, 2017, 45, gkx005.	6.5	84
53	A new computational method for the detection of horizontal gene transfer events. Nucleic Acids Research, 2005, 33, 922-933.	6.5	82
54	Therapeutic potential of FLANC, a novel primate-specific long non-coding RNA in colorectal cancer. Gut, 2020, 69, 1818-1831.	6.1	80

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55	BLOSUM62 miscalculations improve search performance. Nature Biotechnology, 2008, 26, 274-275.	9.4	79
56	The human platelet: strong transcriptome correlations among individuals associate weakly with the platelet proteome. Biology Direct, 2014, 9, 3.	1.9	77
57	Alu and B1 Repeats Have Been Selectively Retained in the Upstream and Intronic Regions of Genes of Specific Functional Classes. PLoS Computational Biology, 2009, 5, e1000610.	1.5	74
58	tRNA Fragments Show Intertwining with mRNAs of Specific Repeat Content and Have Links to Disparities. Cancer Research, 2019, 79, 3034-3049.	0.4	72
59	The Emergence of Pattern Discovery Techniques in Computational Biology. Metabolic Engineering, 2000, 2, 159-177.	3.6	71
60	A generic motif discovery algorithm for sequential data. Bioinformatics, 2006, 22, 21-28.	1.8	66
61	Evaluation of annotation strategies using an entire genome sequence. Bioinformatics, 2003, 19, 717-726.	1.8	65
62	Molecular dynamics simulations of Ago silencing complexes reveal a large repertoire of admissible â€~seed-less' targets. Scientific Reports, 2012, 2, 569.	1.6	62
63	The emerging roles of GPRC5A in diseases. Oncoscience, 2014, 1, 765-776.	0.9	61
64	A sensitive, support-vector-machine method for the detection of horizontal gene transfers in viral, archaeal and bacterial genomes. Nucleic Acids Research, 2005, 33, 3699-3707.	6.5	59
65	Profiles of miRNA Isoforms and tRNA Fragments in Prostate Cancer. Scientific Reports, 2018, 8, 5314.	1.6	57
66	In Silico Pattern-Based Analysis of the Human Cytomegalovirus Genome. Journal of Virology, 2003, 77, 4326-4344.	1.5	54
67	Transcriptional organization of the Clostridium acetobutylicum genome. Nucleic Acids Research, 2004, 32, 1973-1981.	6. 5	53
68	HuR is a post-transcriptional regulator of core metabolic enzymes in pancreatic cancer. RNA Biology, 2013, 10, 1312-1323.	1.5	53
69	A Bayesian Approach to Model Matching with Geometric Hashing. Computer Vision and Image Understanding, 1995, 62, 11-26.	3.0	52
70	GPRC5A is a potential oncogene in pancreatic ductal adenocarcinoma cells that is upregulated by gemcitabine with help from HuR. Cell Death and Disease, 2016, 7, e2294-e2294.	2.7	50
71	TRNA-derived fragments as sex-dependent circulating candidate biomarkers for Parkinson's disease. Parkinsonism and Related Disorders, 2019, 65, 203-209.	1.1	49
72	Dictionary building via unsupervised hierarchical motif discovery in the sequence space of natural proteins., 1999, 37, 264-277.		45

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73	Decoy for microRNAs. Nature, 2010, 465, 1016-1017.	13.7	44
74	In silico structural and functional analysis of the human cytomegalovirus (HHV5) genome 1 1Edited by F. Cohen. Journal of Molecular Biology, 2001, 310, 1151-1166.	2.0	41
75	Assessment of isomiR Discrimination Using Commercial qPCR Methods. Non-coding RNA, 2017, 3, 18.	1.3	40
76	Consequential considerations when mapping tRNA fragments. BMC Bioinformatics, 2016, 17, 123.	1.2	38
77	Mitochondrial tRNA-lookalikes in nuclear chromosomes: Could they be functional?. RNA Biology, 2015, 12, 375-380.	1.5	37
78	IsomiRs and tRNAâ€derived fragments are associated with metastasis and patient survival in uveal melanoma. Pigment Cell and Melanoma Research, 2020, 33, 52-62.	1.5	37
79	HuR's post-transcriptional regulation of death receptor 5 in pancreatic cancer cells. Cancer Biology and Therapy, 2012, 13, 946-955.	1.5	36
80	Nuclear and mitochondrial tRNA-lookalikes in the human genome. Frontiers in Genetics, 2014, 5, 344.	1.1	36
81	Ribosomal RNA fragmentation into short RNAs (rRFs) is modulated in a sex- and population of origin-specific manner. BMC Biology, 2020, 18, 38.	1.7	36
82	The biogenesis pathway of tRNA-derived piRNAs in Bombyx germ cells. Nucleic Acids Research, 2017, 45, 9108-9120.	6.5	34
83	Unification of miRNA and isomiR research: the mirGFF3 format and the mirtop API. Bioinformatics, 2020, 36, 698-703.	1.8	33
84	Dictionary-driven prokaryotic gene finding. Nucleic Acids Research, 2002, 30, 2710-2725.	6.5	31
85	Finding patterns in three-dimensional graphs: algorithms and applications to scientific data mining. IEEE Transactions on Knowledge and Data Engineering, 2002, 14, 731-749.	4.0	30
86	Human and mouse introns are linked to the same processes and functions through each genome's most frequent non-conserved motifs. Nucleic Acids Research, 2008, 36, 3484-3493.	6.5	30
87	microRNAs in colon cancer: A roadmap for discovery. FEBS Letters, 2012, 586, 3000-3007.	1.3	30
88	Multi-focal control of mitochondrial gene expression by oncogenic MYC provides potential therapeutic targets in cancer. Oncotarget, 2016, 7, 72395-72414.	0.8	30
89	Evaluation of Methods for De Novo Genome Assembly from High-Throughput Sequencing Reads Reveals Dependencies That Affect the Quality of the Results. PLoS ONE, 2011, 6, e24182.	1.1	29
90	The Evil, microRNA-143, K-Ras axis in colon cancer. FEBS Letters, 2011, 585, 693-699.	1.3	27

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91	Integrative Multi-omic Analysis of Human Platelet eQTLs Reveals Alternative Start Site in Mitofusin 2. American Journal of Human Genetics, 2016, 98, 883-897.	2.6	27
92	Structural details (kinks and non-Â conformations) in transmembrane helices are intrahelically determined and can be predicted by sequence pattern descriptors. Nucleic Acids Research, 2003, 31, 4625-4631.	6.5	26
93	Post-transcriptional regulation of BRCA1 through its coding sequence by the miR-15/107 group of miRNAs. Frontiers in Genetics, 2015, 6, 242.	1.1	26
94	Dictionary-driven protein annotation. Nucleic Acids Research, 2002, 30, 3901-3916.	6.5	24
95	Metabolic and transcriptional patterns accompanying glutamine depletion and repletion in mouse hepatoma cells: a model for physiological regulatory networks. Physiological Genomics, 2004, 16, 247-255.	1.0	23
96	Aging-associated changes in microRNA expression profile of internal anal sphincter smooth muscle: Role of microRNA-133a. American Journal of Physiology - Renal Physiology, 2016, 311, G964-G973.	1.6	23
97	Accurate Profiling and Quantification of tRNA Fragments from RNA-Seq Data: A Vade Mecum for MINTmap. Methods in Molecular Biology, 2018, 1680, 237-255.	0.4	21
98	Motif discovery without alignment or enumeration (extended abstract)., 1998,,.		20
99	Post-transcriptional Regulation of BRCA2 through Interactions with miR-19a and miR-19b. Frontiers in Genetics, 2016, 7, 143.	1.1	20
100	Inhibition of the Single Downstream Target BAG1 Activates the Latent Apoptotic Potential of MYC. Molecular and Cellular Biology, 2011, 31, 5037-5045.	1.1	18
101	An miRNA Expression Signature for the Human Colonic Stem Cell Niche Distinguishes Malignant from Normal Epithelia. Cancer Research, 2017, 77, 3778-3790.	0.4	18
102	An Approximation Algorithm for Alignment of Multiple Sequences using Motif Discovery. Journal of Combinatorial Optimization, 1999, 3, 247-275.	0.8	16
103	Insights into the Regulation of a Common Variant of HMGA2 Associated with Human Height During Embryonic Development. Stem Cell Reviews and Reports, 2009, 5, 328-333.	5.6	15
104	Implementation of geometric hashing on the Connection Machine. , 0, , .		14
105	OMiR: Identification of associations between OMIM diseases and microRNAs. Genomics, 2011, 97, 71-76.	1.3	14
106	Increasing cell density globally enhances the biogenesis of Piwi-interacting RNAs in Bombyx mori germ cells. Scientific Reports, 2017, 7, 4110.	1.6	14
107	Short RNA regulators: the past, the present, the future, and implications for precision medicine and health disparities. Current Opinion in Biotechnology, 2019, 58, 202-210.	3.3	14
108	Long-term exposure of human endothelial cells to metformin modulates miRNAs and isomiRs. Scientific Reports, 2020, 10, 21782.	1.6	14

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109	Searching in parallel for similar strings [biological sequences]. IEEE Computational Science and Engineering, $1994,1,60-75.$	0.6	13
110	Unraveling the role of microRNA/isomiR network in multiple primary melanoma pathogenesis. Cell Death and Disease, 2021, 12, 473.	2.7	13
111	Atypical transcription of microRNA gene fragments. Nucleic Acids Research, 2010, 38, 2775-2787.	6.5	12
112	DELPHI: A pattern-based method for detecting sequence similarity. IBM Journal of Research and Development, 2001, 45, 455-473.	3.2	11
113	IsoMiRmap: fast, deterministic and exhaustive mining of isomiRs from short RNA-seq datasets. Bioinformatics, 2021, 37, 1828-1838.	1.8	11
114	Threshold-seq: a tool for determining the threshold in short RNA-seq datasets. Bioinformatics, 2017, 33, 2034-2036.	1.8	10
115	Global secondary structure packing angle bias in proteins. Proteins: Structure, Function and Bioinformatics, 2003, 53, 252-261.	1.5	8
116	Short RNAs: How Big Is This Iceberg?. Current Biology, 2010, 20, R110-R113.	1.8	8
117	Exploration of CCA-added RNAs revealed the expression of mitochondrial non-coding RNAs regulated by CCA-adding enzyme. RNA Biology, 2019, 16, 1817-1825.	1.5	8
118	Sequence homology detection through large scale pattern discovery. , 1999, , .		7
119	Sensitive detection of sequence similarity using combinatorial pattern discovery: A challenging study of two distantly related protein families. Proteins: Structure, Function and Bioinformatics, 2005, 61, 926-937.	1.5	7
120	Reply to Backes and Keller: Identification of novel tissue-specific and primate-specific human microRNAs. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E2851-E2851.	3.3	7
121	RNase \hat{I}^0 promotes robust piRNA production by generating $2\hat{a} \in {}^2$, $3\hat{a} \in {}^2$ -cyclic phosphate-containing precursors. Nature Communications, 2021, 12, 4498.	5.8	6
122	The transcriptional trajectories of pluripotency and differentiation comprise genes with antithetical architecture and repetitive-element content. BMC Biology, 2021, 19, 60.	1.7	5
123	Building dictionaries of 1D and 3D motifs by mining the Unaligned 1D sequences of 17 archaeal and bacterial genomes. Proceedings, 1999, , 223-33.	0.3	5
124	The web server of IBM's Bioinformatics and Pattern Discovery group. Nucleic Acids Research, 2003, 31, 3645-3650.	6.5	4
125	MicroRNA Target Prediction. Modecular Medicine and Medicinal, 2010, , 237-263.	0.4	4
126	Beyond mRNAs and Mirnas: Unraveling the Full-Spectrum of the Normal Human Platelet Transcriptome Through Next-Generation Sequencing. Blood, 2012, 120, 3298-3298.	0.6	3

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127	Well-behaved, tunable 3D-affine invariants. , 0, , .		2
128	The web server of IBM's Bioinformatics and Pattern Discovery group: 2004 update. Nucleic Acids Research, 2004, 32, W10-W15.	6.5	2
129	High-Throughput Sequencing of the Human Platelet Transcriptome. Blood, 2010, 116, 481-481.	0.6	2
130	Expression Profiling of a Panel of Apoptosis Related Micrornas in Patients with Acute Myeloid Leukemia. Blood, 2015, 126, 4971-4971.	0.6	2
131	MicroRNA-139 Expression Is Dispensable for the Generation of Influenza-Specific CD8+ T Cell Responses. Journal of Immunology, 2022, 208, 603-617.	0.4	2
132	Managing statistical behavior of large data sets in shared-nothing architectures. IEEE Transactions on Parallel and Distributed Systems, 1998, 9, 1073-1087.	4.0	1
133	Knowledge discovery in biological domains (tutorial AM-3). , 2000, , .		1
134	Anchoring millions of distinct reads on the human genome within seconds. , 2010, , .		1
135	The Complex Transcriptional Landscape of the Human Platelet. Blood, 2012, 120, 390-390.	0.6	1
136	Comments on: "A comprehensive repertoire of tRNA-derived fragments in prostate cancer― , 0, , .		1
137	Abstract A41: GPRC5A acts as a potent oncogene in pancreatic cancer. Cancer Research, 2016, 76, A41-A41.	0.4	1
138	Megakaryocyte-specific knockout of the Mir-99b/let7e/125a cluster lowers platelet count without altering platelet function. Blood Cells, Molecules, and Diseases, 2021, 92, 102624.	0.6	1
139	Report on BIOKDD04. SIGKDD Explorations: Newsletter of the Special Interest Group (SIG) on Knowledge Discovery & Data Mining, 2004, 6, 153-154.	3.2	0
140	Abstract A18: HuR, an RNA binding protein, is critical for the DNA damage response in pancreatic cancer cells, 2012, , .		0
141	Abstract 628: A novel chemoresistance mechanism: HuR post-trancriptionally regulates WEE1, the mitotic inhibitor, upon DNA damage in pancreatic adenocarcinoma cells , 2013, , .		0
142	Towards a Reference Human Platelet Transcriptome: Evaluation Of Inter-Individual Correlations and Its Relationship With a Platelet Proteome. Blood, 2013, 122, 2297-2297.	0.6	0
143	Visuo-Tactual Coarseness Estimation Tasks: Experiments and Modeling. , 1992, , 257-270.		0
144	Characterization of the Short RNA Transcriptome of the Anucleate Platelet. Blood, 2014, 124, 4990-4990.	0.6	0

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145	Abstract A05: The mRNA-binding protein HuR promotes hypoxia-induced chemoresistance through post-transcriptional regulation of the serine-threonine kinase PIM1., 2015,,.		0
146	Abstract A71: Post-transcriptional regulation of the proto-oncogene PIM1 by the mRNA stability factor HuR: implications for pancreatic cancer therapeutic response and cell survival., 2015, , .		0
147	Abstract A78: Profiling of miRNA interactions in pancreatic adenocarcinoma by Argonaute CLIP-seq suggests a highly dynamic repertoire of targets. , 2015, , .		0
148	Abstract A34: Regulation of pancreatic cells by isomiRs. , 2015, , .		0
149	Abstract 2113: Targeting tumor-associated hypoxia to overcome chemoresistance in pancreatic ductal adenocarcinoma (PDA). , 2015, , .		0
150	Abstract 232: Characterization of the human miRNA-ome reveals numerous novel, tissue-specific miRNAs that are implicated in cancer biology. , 2015, , .		0
151	Abstract 3997: Expression profiling of a panel of apoptosis-associated microRNAs in acute myeloid leukemia identifies differentially expressed microRNAs that target epigenetic modifiers., 2015,,.		0
152	Abstract 231: MicroRNA isoforms come of age: Going beyond the one-locus-one-microRNA paradigm in cancer biology. , 2015, , .		0
153	Abstract A29: Gender-specific and race-specific regulatory noncoding RNAs are prevalent in healthy and in cancerous tissues. , 2016 , , .		0
154	Abstract PRO4: The molecular biology of nuclear and mitochondrial regulatory noncoding RNAs differs between white and African American patients with triple-negative breast cancer. , 2016, , .		0
155	Abstract A46: Tissue biology is shaped by noncoding RNAs that depend on gender, population, and race. , 2016, , .		0
156	Abstract 2665: SHOT-RNAs: A novel class of tRNA-derived functional RNAs expressed in hormone-dependent cancers. , 2016, , .		0
157	Abstract 1995: HuR dependent inhibition of PARG enhances PARP inhibitor therapy for DNA repair proficient and deficient pancreatic cancer cells. , 2016 , , .		0
158	Abstract P4-08-01: IsomiRs and tRNA fragments are race-dependent regulators in breast cancer. , 2017, , .		0
159	Abstract B43: Transcriptomic Heterogeneity of microRNA Isoforms and tRNA Fragments contributes to Race-based Differences in Breast and Prostate Cancers. , 2017, , .		0
160	Abstract 4423: Deep sequencing of lung cancer samples using different library preparation methods produces discordant short non-coding RNA profiles. , 2017, , .		0
161	Abstract 5438: Knowledge of which miRNA isoforms are expressed in a sample can successfully discriminate amongst the 32 TCGA cancer types. , 2017, , .		0
162	Abstract B64: Race and prostate cancer: miRNA isoforms and tRNA fragments could hold some of the answers. , 2018 , , .		0

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163	Abstract B63: Race and triple-negative breast cancer: Advances in noncoding RNA research together with a systems-biology-level analysis uncover key molecular differences. , 2018, , .		0
164	Abstract IA31: Novel nuclear and mitochondrial RNAs that are linked to key pathways and depend on sex, population origin, race, tissue, and disease. , 2020, , .		0
165	Abstract C010: The wiring between genes and short noncoding RNAs in cancer depends on race/ethnicity and sex. , 2020, , .		0
166	Abstract B077: Ribosomal RNAs are fragmented into short RNAs in a manner that depends on a person's sex, population origin, and race: implications for health disparities and personalized medicine. , 2020, , .		0
167	Abstract PO-127: The short non-coding RNAs known as "ribosomal RNA-derived fragments―(rRFs) are linked to race disparities in TNBC. , 2020, , .		0