Malay Kumar Basu

Curriculum Vitæ

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Summary

Highly skilled bioinformatician and computational biologist, with more than 20 years of experience in the field, looking for a new exciting opportunity.

Academic Appointments

2012–Present **Assistant Professor**, *Department of Pathology*, University of Alabama, Birmingham, Birmingham, AL, USA.

Academic research, hiring, mentoring, teaching. An award-winning teacher and the chief architect of clinical genomics pipeline and led a team of software engineers and bioinformaticians.

2013–Present **Director**, *Genifx Genome Informatics Facility*, University of Alabama, Birmingham, Birmingham, Al LISA

ham, AL, USA.

Genifx genome analysis facility at UAB (http://genifx.uab.edu) is a UAB Health Science Foundation funded project to create a next-generation sequenced (NGS) data storage and analysis facility at UAB. Designed, created, and now maintains CHOIR/Genifx compute cluster at Department of Pathology,

2012–Present **Associate Scientist**, *Comprehensive Cancer Center*, University of Alabama, Birmingham, Birmingham, AL, USA.

Active researcher in cancer center.

2015-Present Senior Scientist, Informatics Institute, University of Alabama, Birmingham, Birmingham, AL,

USA.

Member, educational committee designing graduate and undergraduate curricula.

2014–Present **Assistant Professor**, Department of Clinical and Diagnostic Sciences, School of Health Profes-

sions, University of Alabama, Birmingham, Birmingham, AL, USA.

Advisor to the master of bioinformatics program.

2020-Present Scientist, Center for Clinical and Translational Science (CCTS), University of Alabama, Birm-

ingham, Birmingham, AL, USA.

Develop university-wide bioinformatics collaboration.

2010–2012 **Assistant Professor**, *J. Craig Venter Institute*, Rockville, USA.

Academic research, hiring, mentoring and leading a team of about 5 software engineers for develop-

ing annotation pipeline.

2009–2010 **Staff Scientist**, *J. Craig Venter Institute*, Rockville, USA.

Academic research, hiring and leading a team of 5 software engineers for developing genome annotation aims lines.

tation pipelines.

Dec 2008–Feb Senior Bioinformatics Engineer, J. Craig Venter Institute, Rockville, USA.

2009 Wrote several high-performance software for genome annotation.

2003–2008 **Postdoctoral Fellow**, National Center for Biotechnology Information (NCBI), National Insti-

tutes of Health (NIH), Bethesda, USA.

Research in the field of computational biology and bioinformatics under the supervision of Dr. Eu-

gene V. Koonin.

2002–2003 **Research Associate**, Center for Cellular and Molecular Biology (CCMB), Hyderabad, India.

Performed independent research in Bioinformatics. Single-handedly designed and built an eight node Linux cluster; installed hardware, performed software installation; helped in training and recruiting personnel for maintenance of the cluster.

Education

Degrees

- 2020–2021 **MBA**, *Collat Business School*, University of Alabama, Birmingham, USA. Anticipated 2021.
- 1995–2002 **PhD (Life Science)**, *Center for Cellular and Molecular Biology*, Hyderabad, India. Degree awarded in 2003.
- 1993–1995 **MTech (Biotechnology)**, *Jadavpur University*, Kolkata, India.
- 1990–1992 **MSc (Zoology with specialization in Cytology and Genetics)**, *University of Calcutta*, Kolkata, India.
- 1987–1990 **BSc(Honours in Zoology)**, *University of Calcutta*, Kolkata, India.

Diplomas

- 2011 **Certificate in Machine Learning**, *Online course*, *Stanford University*, Taught by Andrew Ng.
- 2011 **Certificate in Artificial Intelligence**, *Online course, Stanford University*, Taught by Sebastian Thrun and Peter Norvig.
- 1994 **Certificate in Unix and C**, Computer Science and Engineering Department, Jadavpur University, Kolkata, India.
 - C programming in Unix environment both PC and mainframe.
- 1993 **Programming Techniques and System Design Methodologies**, *Regional Computer Centre*, Kolkata, India.
 - Details of software design and database programming is several languages on PC and mainframe—CYBER 180/84A.
- 1992 **Postgraduate Diploma in Ecology and Environment**, *Indian Institute of Ecology and Envi*ronment, New Delhi, India.
 - Environmental engineering and laws.
- 1992 **Contact Programme in Molecular Biology**, *Department of Zoology*, Banaras Hindu University, Varanasi, India.
 - Basic molecular biology laboratory techniques.

Publications (* corresponding author)

Research Articles

- [1] Swenson-Fields K.I., Ward C. J., Lopez M. E., Fross S., Heimes Dillon A. L., Meisenheimer J. D., Rabbani A. J., Wedlock E., **Basu M. K.**, Jansson K. P., Rowe P. S., Stubbs J. R., Wallace D. P., Vitek M. P., and Fields T. A. Caspase-1 and the inflammasome promote polycystic kidney disease progression. *Front Mol Biosci*, 9:971219, November 2022. doi: 10.3389/fmolb.2022.971219 PMID: 36523654.
- [2] Suster D. I., Craig Mackinnon A., DiStasio M., **Basu M. K.**, Pihan G, and Suster S. Atypical thymomas with squamoid and spindle cell features: Clinicopathologic, immunohistochemical and molecular genetic study of 120 cases with long-term follow-up. *Mod Pathol*, February 2022.
- [3] Spurlock B., Parker D., **Basu M. K.**, Hjelmeland A., GC S., Liu S., Siegal G. P., Gunter A., Moran A., and Mitra K. Fine-tuned repression of Drp1-driven mitochondrial fission primes a 'stem/progenitor-like state' to support neoplastic transformation. *eLife*, 10:e68394, September 2021.
- [4] *Basu M. K., Massicano F., Yu L., Halkidis K., Pillai V., Cao W., Zheng L., and *Zheng X. L. Exome Sequencing Identifies Abnormalities in Glycosylation and ANKRD36C in Patients with

- Immune-Mediated Thrombotic Thrombocytopenic Purpura. *Thromb Haemost*, November 2020. PMID: 33184803.
- [5] Meli A. P., Wang Y., de Kouchkovsky D. A., Kong Y., **Basu M. K.**, Ghosh S., and Rothlin C. V. IL-4-induced hysteresis in naïve T cell activation. *bioRxiv*, page 2020.08.31.275842, September 2020. doi: https://doi.org/10.1101/2020.08.31.275842.
- [6] Jin L., Chen Y., Crossman D. K., Datta A., Vu T., Mobley J. A., Basu M. K., Scarduzio M., Wang H., Chang C., and Datta P. K. STRAP regulates alternative splicing fidelity during lineage commitment of mouse embryonic stem cells. *Nat Commun*, 11(1):5941, 2020. PMID: 33230114.
- [7] Spurlock B., Gupta P., **Basu M. K.**, Mukherjee A., Hjelmeland A. B., Darley-Usmar V., Parker D., Foxall M. E., and Mitra K. New quantitative approach reveals heterogeneity in mitochondrial structure-function relations in tumor-initiating cells. *J Cell Sci*, 132(9), May 2019. PMID: 30910831.
- [8] Yu L., Tanwar D. K., Penha E. D. S., Wolf Y. I., Koonin E. V., and *Basu M. K. Grammar of protein domain architectures. *Proc Natl Acad Sci U S A*, 116(9):3636–3645, 2019. PMID: 30733291.
- [9] Ping Z., Soni A., Williams L. A., Pham H. P., **Basu M. K.**, and Zheng. X. L. Mutations in Coagulation Factor VIII are associated with more favorable outcome in patients with cutaneous melanoma. *TH Open*, 1(2):e113–e121, July 2017. PMID: 29152610.
- [10] Tanwar D. K., Parker D. J., Gupta P., Spurlock B., Alvarez R. D., *Basu M. K., and Mitra K. Crosstalk between the mitochondrial fission protein, Drp1, and the cell cycle is identified across various cancer types and can impact survival of epithelial ovarian cancer patients. *Oncotarget*, August 2016. PMID: 27509055.
- [11] Parker D. J., Iyer A., Shah S., Moran A., Hjelmeland A. B., **Basu M. K.**, Liu R., and Mitra K. A new mitochondrial pool of cyclin E, regulated by Drp1, is linked to cell-density-dependent cell proliferation. *J. Cell. Sci.*, 128(22):4171–4182, November 2015. PMID: 26446260.
- [12] Tian R., **Basu M. K.**, and Capriotti E. Computational methods and resources for the interpretation of genomic variants in cancer. *BMC Genomics*, 16 Suppl 8:S7, 2015. PMID: 26111056.
- [13] Tian R., **Basu M. K.**, and Capriotti E. ContrastRank: A new method for ranking putative cancer driver genes and classification of tumor samples. *Bioinformatics*, 30(17):i572–578, September 2014. PMID: 25161249.
- [14] Haft D. H., Selengut J. D., Richter R. A., Harkins D., **Basu M. K.**, and Beck E. TIGRFAMs and genome properties in 2013. *Nucleic Acids Res*, 41(D1):D387–D395, Jan 2013. PMID: 23197656.
- [15] *Basu M. K., Selengut J. D., and *Haft D. H. ProPhylo: partial phylogenetic profiling to guide protein family construction and assignment of biological process. *BMC Bioinform*, 12(1):434, November 2011. PMID: 22070167.
- [16] Haft D. H. and **Basu M. K.** Biological systems discovery in silico: radical S-adenosylmethionine protein families and their target peptides for posttranslational modification. *J. Bacteriol.*, 193(11):2745–2755, June 2011. PMID: 21478363.
- [17] Daphnia Genome Consortium. The ecoresponsive genome of *Daphnia pulex. Science*, 331(6017):555–561, Feb 2011.
- [18] Haft D. H., **Basu M. K.**, and Mitchell D. A. Expansion of ribosomally produced natural products: a nitrile hydratase- and nif11-related precursor family. *BMC Biology*, 8:70, 2010. PMID: 20500830.
- [19] Rogozin I. B., **Basu, M. K.**, Csürös M., and Koonin E. V. Analysis of rare genomic changes does not support the unikont-bikont phylogeny and suggests cyanobacterial symbiosis as

- the point of primary radiation of eukaryotes. *Genome Biol Evol*, 2009(0):99–113, June 2009. PMID: 20333181.
- *Basu M. K., Poliakov E., and Rogozin I. B. Domain mobility in proteins: functional and evolutionary implications. *Brief Bioinform.*, 10(3):205–16, 2009. PMID:19151098.
- [21] **Basu M. K.**, Rogozin I. B., Deusch O., Dagan T., Martin W., and Koonin E. V. Evolutionary dynamics of introns in plastid-derived genes in plants: saturation nearly reached but slow intron gain continues. *Mol Biol Evol*, 25:111–9, 2008. PMID: 17974547.
- [22] **Basu M. K.**, Rogozin I. B., and Koonin E. V. Primordial spliceosomal introns were probably U2-type. *Trends Genet*, 24(11):525–8, 2008. PMID:18824272.
- [23] **Basu M. K.**, Carmel L., Rogozin I. B., and Koonin E. V. Evolution of protein domain promiscuity in eukaryotes. *Genome Res.*, 18(3):449–61, 2008. PMID: 18230802.
- [24] **Basu M. K.**, Makalowski W., Rogozin I. B., and Koonin E. V. U12 intron positions are more strongly conserved between animals and plants than U2 intron positions. *Biol Direct*, 3(1):19, 2008. PMID: 18479526.
- [25] Babenko V. N., **Basu M. K.**, Kondrashov F. A., Rogozin I. B., and Koonin E. V. Signs of positive selection of somatic mutations in human cancers detected by EST sequence analysis. *BMC cancer*, 6:36, 2006. PMID: 16469093.
- [26] Rogozin I. B., **Basu M. K**, Jordan I. K., Pavlov Y. I., and Koonin E. V. APOBEC4, a new member of the AID/APOBEC family of polynucleotide (deoxy)cytidine deaminases predicted by computational analysis. *Cell cycle*, 4:1281–5, September 2005. PMID: 16082223.
- [27] **Basu M. K.** and Koonin E. V. Evolution of eukaryotic cysteine sulfinic acid reductase, sulfiredoxin (Srx), from bacterial chromosome partitioning protein ParB. *Cell cycle*, 4:947–52, July 2005. PMID: 15917647.
- [28] *Basu M. K. SeWeR: a customizable and integrated dynamic HTML interface to bioinformatics services. *Bioinformatics*, 17:577–8, June 2001. PMID: 11395442.
- [29] Ray M. K., Kumar G. S., Janiyani K., Kannan K., Jagtap P., **Basu M. K.**, and Shivaji S. Adaptation to low temperature and regulation of gene expression in Antarctic psychrotrophic bacteria. *J Biosci*, 23(4):423–35, 1998.

Invited book chapters and reviews

- [30] *Basu M. K. Bioinformatics over the Web: SeWeR, as you may think. In R P Grant, editor, Computational Genomics: Theory and Application. Horizon Press, UK, 2004.
- [31] **Basu M. K.** and Mishra R. K. Bioinformatics and the art of sequence analysis. In N C Gautam and M P Singh, editors, *Recent Advances in Biotechnology*. Shree Publishers, New Delhi, 2004.

Other publications

- [32] Koo H., **Basu, M. K.**, Crowley M., Aislabie J, and Bej A. K. Draft Genome Sequence of Pseudomonas sp. Strain Ant30-3, a Psychrotolerant Bacterium with Biodegradative Attribute Isolated from Antarctica. *Genome Announc.*, 2(3):e00522–14, June 2014. PMID: 24903870.
- [33] Haft D. H., **Basu. M. K.**, and Richter R. A. Braingrab: Capturing curator expertise as reusable annotation rules. *Nature Precedings*, 2009. doi:10.1038/npre.2009.3313.1.
- [34] Spurlock B., Parker D., **Basu M. K.**, Hjelmeland A., and Mitra K. Modulation of mitochondrial fission activity maintains ovarian tumor initiating cells dependent on mitochondrial energetics. *Free Radical Biology and Medicine*, 128:S74, November 2018.

[35] Massicano F., Staley E. M., Halkidis K., Kocher N. K., Williams L. A., Marques M. B., Guillory B. K., Cao W, **Basu M. K.**, and Zheng X. L. Exome Sequencing Identifies Glycosylation Defects As a Probable Cause of Immune Thrombotic Thrombocytopenic Purpura. *Blood*, 134(Supplement_1):217–217, November 2019.

	Software Publications
SaucePan	Protein clustering software meta-package. Implements a novel language-modeling algorithm to identify orthologs. Internally used in JCVI.
ProPhylo	A high-performance parallel Perl framework for genome-scale phylogenetic profile comparison and the associated databases. Available at https://github.com/malaybasu/ProPhylo.
NCBIWeb	Perl modules to automate NCBI web server.
AnnotationRules	Functional annotation of protein using rules. Used in annotation pipeline at JCVI.
SeWeR	A very popular and widely used interface for bioinformatics services. SeWeR has been translated into several languages, and written using JavaScript and DHTML. Available at http://www.bioinformatics.org/sewer.
Pastel	A Perl framework for generating Scalable Vector Graphics (SVG) and animation. The API closely resembles Java Graphics2D API with state-of-the-art computational geometry support. Available at http://www.bioinformatics.org/pastel.
BioSVG	The first application of SVG in bioinformatics. A Perl framework for generating high-resolution vector graphics for biological data. Available at http://www.bioinformatics.org/biosvg.
Savvy	A CGI-based plasmid drawing software that generates print-quality, editable plasmid maps in SVG format. Available at http://www.bioinformatics.org/savvy.
ABI.pm	A module for parsing ABI chromatogram files. Available from CPAN (http://search.cpan.org/~malay/ABI-0.01).
SeqToolBox	My personal sequence analysis toolbox. Used in many projects that I wrote. Available at https://github.com/malaybasu/SeqToolBox
Font::TTFMetrics	A Perl module for parsing True Type Font file. Available from CPAN (http://search.cpan.org/~malay/Font-TTFMetrics-0.1/).
pastel-ttf2svg.pl	Converts TTF file to SVG font file. Available from CPAN. (http://www.cpan.org/pub/CPAN/authors/id/M/MA/MALAY/pastel-ttf2svg-0.04.zip).
Clinical genomics pipeline	I was the chief architect of the clinical genomics pipeline of at UAB, Dept. of Pathology.
	Awards and honors

Awards and honors

2020 Excellence in Teaching Award by Graduate school, University of Alab
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2017–2019 Elected member of UAB faculty senate.

2017 Nominated for Dean's excellence award in teaching, School of Medicine, University of Alabama, Birmingham.

Nominated for Dean's excellence in teaching award, School of Medicine, University of Alabama, Birmingham.

2012-Present Adjunct faculty F1000 prime.

2003–2008 NIH intramural fellowship.

2002 Travel award from O'Reilly publishing.

2002 Travel award from Council for Scientific and Industrial Research, India.

1997–2001 Senior research fellowship from Council for Scientific Industrial Research, India.

1995–1997 Junior research fellowship from Council for Scientific Industrial Research, India.

1993–1995	Fellowship from Department of Biotechnology, Govt. of India.
	Awards and honors from lab
2019	Felipe Massicano won departmental travel award to attend American Society of Hematology annual meeting held in Orlando, FL.
2019	Felipe Massicano won Abstract Achievement Award at the American Society of Hematology annual meeting held in Orlando, FL.
2018	Amrita Lakhanpal, high school student. Her project was awarded Intel Excellence in Computer Science award, 2018.
	Editorial boards
2021—-Present	Editor, BMC Genomics Data (Springer).
2020-Present	Reviewing board Member, Cancers (MDPI, IF: 6.126).
2015—-2020	Editorial board Member, Heliyon (Elsevier).
2014—-Present	Appointed editorial board member; Bioinformatics section, BioMed Research International (Impact: 2.88).
2013–Present	Appointed editorial Board member; Evolutionary Biology Section, BioMed Research International
2011–Present	Reviewing board member, Frontiers in Bioinformatics and Computational Biology.
	Grant review
2021	Israel Science Foundation (ISF), Israel.
2017	National Science Foundation.
2017	Selected for NIH Early Career Reviewer (ECR) program.
2015	National Science Foundation (NSF) career award.
	Councils and committees
2019	Member, organizing committee, Midsouth Bioinformatics Conference (MCBIOS), 2019.
2019	Poster judge, Midsouth Bioinformatics and Computational Biology society conference, 2019
2019	Faculty Advisory Committee of Bryan Guillory.
2018	Member, faculty search committee, Genomic Diagnostic and Bioinformatics division, Department of Pahtology.
2018	Judge, Pathology research day symposium.
2017—-Present	Member, Bioinformatics Educational Committee, Informatics Institute, UAB.
2017—-Present	Member, Informatics Gateway Committee, a research evaluation committee created by CCTS and Informatics institute for facilitating bioinformatics collaborations.
2017—-Present	Member departmental IRR review committee

2019	Poster judge, Midsouth Bioinformatics and Computational Biology society conference, 2019
2019	Faculty Advisory Committee of Bryan Guillory.
2018	Member, faculty search committee, Genomic Diagnostic and Bioinformatics division, Department of Pahtology.
2018	Judge, Pathology research day symposium.
2017—-Present	Member, Bioinformatics Educational Committee, Informatics Institute, UAB.
2017—-Present	Member, Informatics Gateway Committee, a research evaluation committee created by CCTS and Informatics institute for facilitating bioinformatics collaborations.
2017—-Present	Member, departmental IRB review committee.
2017—-2019	Elected member of UAB faculty senate.
2017—-2019	Member, graduate curriculum committee, UAB faculty senate.
2017–2018	Chair, Bioinformatics Power Talks steering committee, a joint initiative between Informatics Insitute and CB2 computational biology and bioinformatics group at UAB.
2015–Present	Thesis advisory committees of, Joseph Palmer, John Schoelz, Sean Wilkinson
2014—Present	Advisor to the Molecular Tumor Board, UAB.
2012—-2014	Faculty Advisory Council member for representing the following divisions of Department of Pathology: Informatics, Neuropathology, Forensic Pathology.

- 2012—-Present Founder, Computational Biology and Bioinformatics activity (CB2; http://uab.edu/cb2) at UAB.
 - 2006–2007 Institute selected member of Fellows Committee (FELCOM) at NIH.
 - 2007 Chief judge of Bioinformatics and Computational Biology section of Fellows Award of Research Excellence (FARE) at NIH.
 - 2006 Judge, RECOMB satellite workshop on comparative genomics, Montréal, Canada.

Invited talks and workshops

- 2020-04-06-09 Bioinformatics workshop at Yale School of Medicine.
 - 2020-04-06 Department of Immunology, Yale University, Connecticut, USA. (Postponed for lockdown).
 - 2020-03-13 Annual Translational and Transformative Informatics Symposium (ATTIS), 2020. "Reading the book of life: the grammar of genes".
 - 2019-10-07 Department of Microbiology and Immunology, University of Mississippi Medical Center, University of Mississippi, Oxford, Mississippi, USA. Visited various department and delivered a talk entitled: "Reading the book of life: The language of the genes".
 - 2019-03-30 Midsouth Bioinformatics and Computational Biology Society Conference, 2019. "Reading the book of life: Language of Genomes".
 - 2018-04-25 Translational Bioinformatics Symposium, UAB. "DBGES: A Novel Gene Expression signature for developing a mitochondria based targeted therapy".
 - 2018-01-29 UAB department of medicine, Hematology and Oncology conference: "Cancer genome data-mining to identify a novel gene-expression signature for ovarian cancer prognosis".
 - 2017-05-03 UAB 1st Annual Translational Bioinformatics Mini-Symposium. "A novel gene-expression signature for ovarian cancer prognosis".
 - 2018-01-29 UAB department of medicince, Hematology and Oncology conference: "Cancer genome data-mining to identify a novel gene-expression signature for ovarian cancer prognosis".
 - 2017-05-03 UAB 1st Annual Translational Bioinformatics Mini-Symposium. "A novel gene-expression signature for ovarian cancer prognosis".
 - 2016-09-21 UAB Department of Biology. "Language of the genes".
 - 2016-02-09 UAB Department of Pathology. Molecular and Cellular Pathology Seminar "Unraveling Novel Biological Paradigms from Large-scale Analysis of Cancer Genomes".
 - 2014-03-06 Invited speaker, Hudson Alpha Institute of Bioinformatics, AL.
 - 2013 Genetics and Genomics Seminar Series (Jan 2013): "Investigating Biological Systems Using Phylogenetic Profiling".
 - 2011 Invited faculty to teach in the training course "Molecular Methods for Characterization, conservation and utilization of biodiversity", held in Feb-March, 2011 in Hyderabad, India.
 - 2012 University of Alabama, Birmingham.
 - 2011 Center for DNA fingerprinting and Diagnostics, Hyderabad, India.
 - 2008 Memorial Sloan-Kettering Cancer Center, NY.
 - 2008 Institute of Genome Sciences, Baltimore, MD.
 - 2008 University of Maryland, College Park, MD.
 - Speaker in O'Reilly Bioinformatics Technology conference, 2002, held in Tucson, Arizona. I delivered a talk entitled, "DHTML and Scalable Vector Graphics in Bioinformatics".

Teaching Experience

2021–Present Co-director and designer of a new undergraduate bioinformatics course, "Introductory bioinformatics".

2018-Present Co-director and designer of a new bioinformatics course, "INFO 510: Programming with biological data". Course master of "CB2-101: Introduction to Scientific Computing". A highly rated (rating 2014-Present score 9.5/10) hands-on 48 hours training course, open to everyone, at UAB. Also a graduate course of 3 credit hours. http://cmb.path.uab.edu/training/cb2-101.html. Course master of "CB2-201: Bioinformatics and Computational Biology". A highly rated 2015-Present (9.5/10) 40hr hands-on training course. Also a 3 credit graduate course. http://cmb.path. uab.edu/traning/cb2-201.html. 2018 Taught in GBS779: Translational Research. Course master: Eddy Yang. Organizer and course master of "Bioinformatics Power Talks". This is a new university-wide 2017-2018 bioinformatics research and journal discussion club, jointly sponsored by Computational Biology and Bioinformatics (CB2) and Informatics Institute, UAB. This is also a registered GBS course with 1 credit. Course master of "Computational Genomics". Advanced bioinformatics course. GBS 787. 2014 No longer offerred. Taught and evaluated evolutionary genomics GBS 722. 2014-02-05 2013-2017 Course master of a highly rated Computational Biology and Bioinformatics (CB2) journal club. Also a graduate school course of 1 credit hour. 2013 Taught and evaluated "Molecular Evolution" GBS 722. 2013 Highly rated laboratory medicine course on "Genomics" (Rating 4.65/5.00). School of Health Professionals "Introduction to Bioinformatics." 2012 Mentoring Filepe Massicano, Postdoctoral fellow. 2018-2020 2019-2020 Christopher Coffee, Undergraduate student intern. 2016-2018 Lijia Yu, Visiting Fellow. Joining Cambridge University, UK to pursue Ph.D. Amrita Lakhanpal, high school students. Her project was awarded Intel Excellence in Com-2017-2018 puter Science Award. 2017 Margaret Bell. GBS Rotation student 2014-2016 Deepak Tanwar, Visiting Fellow then Research Assistant. Currently pursuing Ph.D. in ETH Zurich. Emannuel Penha, Visiting Fellow. Now faculty in Brazil. 2014-2016 2014-2015 Aseygul Bulut, Part-time student assistant. Paul Boothe, Medical student intern in Cancer Research Experience for Students (CaRES) 2013 program. 2013 Darshan Patel, Biotechnology Intern, School of Health Professions. Meghna Yadigiri, Summer Intern at J. Craig Venter Institute. 2011 Conferences Oral presentation Basu MK (2020) Reading the book of life: the language of proteins. ISCB-Latin America, 2020-10-29 SolBio BioNetMX.

Basu MK (2020) Reading the book of life: the language of proteins. Intelligent Systems in

2020-07-14

Molecular Biology (ISMB).

- 2020-07-13 Basu MK (2020) Exome sequencing identifies abnormalities in glycosylation and ANKRD36C defects as probable causes of immune-mediated thrombotic thrombocytopenic purpura (TTP). Intelligent Systems in Molecular Biology (ISMB).
- 2019-12-7 Massicano F, Staley E, Halkidis K, Kocher N, Williams LA, Marques MB, Guillory B, Cao W, Basu MK, Zheng XL (2019) Exome sequencing identifies glycosylation defects as a probable cause of immune thrombotic thrombocytopenic purpura, American Society of Hematology annual conference, Dec 7-10, 2019, Orlando, FL.

Posters

- 2020-07-13 Basu MK (2020) Exome sequencing identifies abnormalities in glycosylation and ANKRD36C defects as probable causes of immune-mediated thrombotic thrombocytopenic purpura (TTP). Intelligent Systems in Molecular Biology (ISMB).
- 2019-12-7 Massicano F, Staley E, Halkidis K, Kocher N, Williams LA, Marques MB, Guillory B, Cao W, Basu MK, Zheng XL (2019) Exome sequencing identifies glycosylation defects as a probable cause of immune thrombotic thrombocytopenic purpura, American Society of Hematology annual conference, Dec 7-10, 2019, Orlando, FL.
- 2018-11-14 Spurlock B, Parker D, Basu MK, Hjelmeland A, Mitra K (2018) Modulation of mitochondrial fission activity maintains ovarian tumor initiating cells dependent on mitochondrial energetics. Presented at The Society for Redox Biology and Medicine's 25th Annual Conference (SfRBM 2018), November 14-17, 2018, Chicago, IL, USA.
- 2018-04-25 Yu L, Mitra K, Arend R, Basu MK (2018) DBGES-a predictive and prognostic gene-expression signature for ovarian and other cancers, UAB translational bioinformatics conference.
- 2017-07-09 Parker D, Spurlock B, Tanwar D, Basu MK, Mitra K (2017) Mitochondrial energetics regulated by mitochondrial fission modulates cell cycle towards maintaining stemness. GRC Cell Growth and Proliferation, July 9-14, 2017
- 2017-06-17 Yu L, Basu MK (2017) The language of protein domains, Pathology research retreat.
- 2016-11-05 UAB Comprehensive Cancer Center Symposium Mutation distribution and codon usage bias in oncogene and tumor-suppressor genes.
- 2016-09-09 UAB Core day: Genifx: Genome Informatics Facility at UAB.
- 2016-07-11 Intelligent Systems in Molecular Biology (ISMB), Orlando, FL "Position-dependent mutation distribution and codon usage bias in oncogene and tumor-suppressor genes".
- 2016-02-03 Parker D, Archana I, Basu MK, Mitra K (2016) Mitochondrial regulation of Cyclin E, AACR Precision Medicine Series: Cancer Cell Cycle.
- 2015-07-19 Parker D, Archana I, Basu MK, Mitra K. (2015) Mitochondrial fission protein Drp1 controls cell proliferation in a cell density dependent manner by regulating Cyclin E. Cell Symposia, Multifaceted Mitochondria, Chicago, IL, USA, July 19-21, 2015.
- 2014-11-18 UAB Core Day. "Genifx: Genome Informatics Facility @ UAB".
- 2014-10-06 UAB Comprehensive Cancer Center Symposium "ContrastRank: A New Method for Ranking Cancer Driver Genes and Tumor Samples Classification".
- 2013-11-05 UAB Comprehensive Cancer Center Symposium. Cancer Center Symposium. "The Role of Mitochondrial Fission and Fusion in Evolution of Cancer Genes", with Paul Boothe.

Skills

Bioinformatics

All aspects of classical bioinformatics, comparative genomics, evolutionary genomics, Next-generation sequence anlysis (NGS), RNASeq, whole-genome and exome analysis, single-cell data analysis, cancer genomics.

Data mining and Al

Traditional machine learning, regressions, SVM, Radom Forrest, clustering, PCA, PSLDA, SPLSDA, NMF, Deep learning using Keras, TensorFlow.

Computers

Software development

Algorithm design, object-oriented system design, design patterns, cross-platform and platform-dependent software development on GNU/Linux platform using different languages, network programming, database development, system programming, code maintenance and debugging.

Languages

C, C++, Java, Perl, Python, JavaScript, SQL, R, Octave, Matlab, ŁTEX.

Databases

MySQL, SyBase, SQLite, Barkley DB.

Cloud computing and containers

AWS, Google cloud, Docker

High Performance Computing

Extensive knowledge in hardware, purchasing, and setting up and running Linux cluster, parallel programming using MPI and PVM, and programming job schedulers like SGE and SLURM.

10/10

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