Prof. TOLGA CAN

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International Researcher IDs

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Publons / Web Of Science ResearcherID: D-1395-2010

ScopusID: 6602569668 Yoksis Researcher ID: 30749

Biography

Tolga Can received his BS degree in computer engineering from Middle East Technical University, Turkey, in 1998 and his PhD degree in computer science from the University of California at Santa Barbara in 2004. He is currently a professor in the Computer Engineering Department, Middle East Technical University, Turkey. His main research interests include bioinformatics and scientific visualization. He has worked on protein structure visualization and alignment and on construction and analysis of large-scale protein-protein interaction networks.

Education Information

Doctorate, University of California, Santa Barbara, Bilgisayar Bilimleri, United States Of America 2000 - 2004 Undergraduate, Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, Turkey 1994 - 1998

Research Areas

Computer Sciences, bioinformatics, Engineering and Technology

Academic Titles / Tasks

Professor, Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, 2017 - Continues

Associate Professor, Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, 2011 - 2017

Assistant Professor, Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, 2007 - 2011

Lecturer, Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, 2006 - 2007 Research Assistant, University of California, Santa Barbara, College Of Engineering, Computer Science, 2000 - 2005

Academic and Administrative Experience

Middle East Technical University, Faculty of Engineering, Department of Computer Engineering, 2007 - 2010

Advising Theses

CAN T., Construction and analysis of tissue/disease specific protein-proteininteraction networks by integrating large scale transcriptomedata with genome scale protein-protein interaction networks, Doctorate, A.BURÇAK(Student), 2022

CAN T., Mini autonomous car architecture for urban driving scenarios, Postgraduate, G.Karabulut(Student), 2019

CAN T., Genome- and tissue-wide analysis of alternative polyadenylation events using clustering and feature learning methods, Postgraduate, P.Yılmazer(Student), 2019

CAN T., TARI Z. S., Shape models based on elliptic pdes, associated energies, and their applications in 2d and 3d, Doctorate, A.GENÇTAV(Student), 2018

CAN T., Tools and techniques for assessing functional relevance of genomic loci, Doctorate, B.OTLU(Student), 2017

CAN T., A layout algorithm for visualization of graph alignments, Postgraduate, A.AKARSU(Student), 2017

CAN T., A multiplex primer design algorithm for target amplification of continuous genomic regions, Doctorate, A.RAŞİT(Student), 2016

CAN T., ERSON BENSAN A. E., Alternative polyadenylation dependent 3'-UTR changes in triple negative breast cancers, Doctorate, T.TUNCER(Student), 2015

CAN T., AYDIN SON Y., A multi-layered graphical model of the relation among SNPs, genes, and pathways based on subgraph search, Postgraduate, G.ERSOY(Student), 2015

CAN T., ÖZCENGİZ G., Pattern search in pathogenic bacterial proteins for localization and secretory systems, Doctorate, O.ÖZCAN(Student), 2015

CAN T., Determination of the effect of polyadenylation SLR values on microarray data classification, Postgraduate, Ü.ASLAN(Student), 2014

CAN T., Identification of interaction sites of G protein-coupled receptors using machine learning techniques, Doctorate, M.EMRE(Student), 2014

CAN T., A database query based solution for chemical compound and drug name recognition, Postgraduate,

CAN T., Integer linear programming based solutions for construction of biological networks, Doctorate,

Ö.EREN(Student), 2014
CAN T., ŞENER C., Modeling of split step parabolic wave equation using the graphics processing unit, Postgraduate,

CAN T., ŞENER C., Modeling of split step parabolic wave equation using the graphics processing unit, Postgraduate S.SEKMEN(Student), 2014

CAN T., ŞENER C., GPU accelerated radio wave propagation modeling using ray tracing, Postgraduate, A.ZUBAROĞLU(Student), 2014

CAN T., Machine learning methods for using network based information in microRNA target prediction, Doctorate, M.SUALP(Student), 2013

CAN T., Prediction of polyadenylation sites by probe level analysis of microarray data, Postgraduate,

Y.İLGÜNER(Student), 2013

Ç.ATA(Student), 2014

CAN T., Analysis of 3'UTR shortening events in breast cancer, Postgraduate, O.BALOĞLU(Student), 2013

CAN T., AYDIN SON Y., A framework for gene co-expression network analysis of lung cancer, Postgraduate,

E.AKDEMİR(Student), 2013

CAN T., Mining fungal effector candidates in biotrophic plant pathogens : rusts and mildews, Postgraduate, S.UĞUR(Student), 2012

CAN T., SINEC: Large scale signaling network topology reconstruction using protein-protein interactions and RNAi data, Postgraduate, S.HASHEMİKHABİR(Student), 2012

CAN T., Parallelization of functional flow to predict protein functions, Postgraduate, E.AKKOYUN(Student), 2011

CAN T., Machine learning methods for promoter region prediction, Postgraduate, H.ARSLAN(Student), 2011

CAN T., Resonctructing signaling pathways from RNAi data using genetic algorithms, Postgraduate, E.SERDAR(Student), 2011

CAN T., Identification of functionally orthologous protein groups in different species based on protein network alignment, Postgraduate, Ö.NEBİL(Student), 2010

CAN T., A distributed graph mining framework based on mapreduce, Postgraduate, S.ALKAN(Student), 2010

CAN T., A 3D topological tracking system for augmented reality, Postgraduate, M.ERCAN(Student), 2010

CAN T., A new approach for better load balancing of visibility detection and target acquisition calculations, Postgraduate, A.YİĞİT(Student), 2010

CAN T., A script based modular game engine framework for augmented reality applications, Postgraduate, M.FURKAN(Student), 2009

CAN T., Multi-resolution visualization of large scale protein networks enriched with gene ontology annotations, Postgraduate, S.YAŞAR(Student), 2009

CAN T., Prediction of protein-protein interactions from sequence using evolutionary relations of proteins and species, Postgraduate, T.DOĞACAN(Student), 2009

CAN T., Text classification in Turkish marketing domain and context-sensitive ad distribution, Postgraduate, M.ENGİN(Student), 2009

CAN T., Coevolution based prediction of protein-protein interactions with reduced training data, Postgraduate, B.PAMUK(Student), 2009

CAN T., ATALAY M. V., A classification system for the problem of protein subcellular localization, Postgraduate, G.ALAY(Student), 2007

Published journal articles indexed by SCI, SSCI, and AHCI

I. Differential expression of mRNA 3'-end isoforms in cervical and ovarian cancers DÖKEN D. N., ÖZGÜL İ., KÖKSAL BIÇAKCI G., Gol K., CAN T., ERSON BENSAN A. E. Heliyon, vol.9, no.9, 2023 (SCI-Expanded)

II. Targeting HIF1-alpha/miR-326/ITGA5 axis potentiates chemotherapy response in triple-negative breast cancer

Assidicky R., Tokat U. M., Tarman I. O., Saatci O., Ersan P. G., Raza U., Ogul H., Riazalhosseini Y., CAN T., Sahin O. BREAST CANCER RESEARCH AND TREATMENT, vol.193, no.2, pp.331-348, 2022 (SCI-Expanded)

III. A C-term truncated EIF2Bγ protein encoded by an intronically polyadenylated isoform introduces unfavorable EIF2Bγ-EIF2γ interactions.

Circir A., Bicakci G. K., Savas B., Doken D. N., Henden O., Can T., Karaca E., Erson-Bensan A. E. Proteins, vol.90, no.3, pp.889-897, 2022 (SCI-Expanded)

IV. Identification of an mRNA isoform switch for HNRNPA1 in breast cancers

Erdem M., Ozgul İ., Dioken D. N., Gurcuoglu I., Guntekin Ergun S., Cetin-Atalay R., Can T., Erson-Bensan A. E. Scientific Reports, vol.11, no.1, 2021 (SCI-Expanded)

V. MotifGenie: A Python Application for Searching Transcription Factor Binding Sequences Using ChIP-Seq Datasets.

Oguztuzun C., Yasar P., Yavuz K., Muyan M., Can T.

Bioinformatics (Oxford, England), vol.37, pp.4238-4239, 2021 (SCI-Expanded)

VI. A prelude to the proximity interaction mapping of CXXC5

Ayaz G., Turan G., OLGUN Ç. E., Kars G., Karakaya B., Yavuz K., Demiralay O. D., CAN T., MUYAN M., Yasar P. SCIENTIFIC REPORTS, vol.11, no.1, 2021 (SCI-Expanded)

VII. A CpG island promoter drives the CXXC5 gene expression

Yasar P., Kars G., Yavuz K., Ayaz G., Oguztuzun C., Bilgen E., Suvaci Z., Cetinkol Ö., CAN T., MUYAN M. SCIENTIFIC REPORTS, vol.11, no.1, 2021 (SCI-Expanded)

VIII. JOA: Joint Overlap Analysis of multiple genomic interval sets

Otlu B., Can T.

BMC BIOINFORMATICS, vol.20, 2019 (SCI-Expanded)

IX. Alternative Polyadenylation Patterns for Novel Gene Discovery and Classification in Cancer BEGIK O., ÖYKEN M., ALICAN T. C., CAN T., Erson-Bensan A. E.

NEOPLASIA, vol.19, no.7, pp.574-582, 2017 (SCI-Expanded)

X. A multiplex primer design algorithm for target amplification of continuous genomic regions

OZTURK A. R., CAN T.

BMC BIOINFORMATICS, vol.18, 2017 (SCI-Expanded)

XI. Comparison of tissue/disease specific integrated networks using directed graphlet signatures Sonmez A. B., Can T.

BMC BIOINFORMATICS, vol.18, 2017 (SCI-Expanded)

XII. SUMONA: A supervised method for optimizing network alignment

TUNCAY E. G., Can T.

COMPUTATIONAL BIOLOGY AND CHEMISTRY, vol.63, pp.41-51, 2016 (SCI-Expanded)

XIII. Alternative Polyadenylation: Another Foe in Cancer

Erson-Bensan A. E., CAN T.

MOLECULAR CANCER RESEARCH, vol.14, no.6, pp.507-517, 2016 (SCI-Expanded)

XIV. 3 'UTR shortening and EGF signaling: implications for breast cancer

AKMAN H. B., ÖYKEN M., TUNCER T., CAN T., Erson-Bensan A. E.

HUMAN MOLECULAR GENETICS, vol.24, no.24, pp.6910-6920, 2015 (SCI-Expanded)

XV. Reconstruction of the temporal signaling network in Salmonella-infected human cells

Budak G., Ozsoy O. E., Son Y., Can T., Tunçbağ N.

FRONTIERS IN MICROBIOLOGY, vol.6, 2015 (SCI-Expanded)

XVI. The CHEMDNER corpus of chemicals and drugs and its annotation principles

Krallinger M., Rabal O., Leitner F., Vazquez M., Salgado D., Lu Z., Leaman R., Lu Y., Ji D., Lowe D. M., et al. JOURNAL OF CHEMINFORMATICS, vol.7, 2015 (SCI-Expanded)

XVII. Div-BLAST: Diversification of Sequence Search Results

Eser E., CAN T., Ferhatosmanoglu H.

PLOS ONE, vol.9, no.12, 2014 (SCI-Expanded)

XVIII. GPCRsort-Responding to the Next Generation Sequencing Data Challenge: Prediction of G Protein-Coupled Receptor Classes Using Only Structural Region Lengths

Sahın M. E., Can T., Son Ç. D.

OMICS-A JOURNAL OF INTEGRATIVE BIOLOGY, vol.18, pp.636-644, 2014 (SCI-Expanded)

XIX. Identification of Novel Reference Genes Based on MeSH Categories

Ersahin T., ÇARKACIOĞLU L., CAN T., Konu O., Atalay V., Cetin-Atalay R.

PLOS ONE, vol.9, no.3, 2014 (SCI-Expanded)

XX. A Divide and Conquer Approach for Construction of Large-Scale Signaling Networks from PPI and RNAi Data Using Linear Programming

Ozsoy O. E., Can T.

IEEE-ACM TRANSACTIONS ON COMPUTATIONAL BIOLOGY AND BIOINFORMATICS, vol.10, pp.869-883, 2013 (SCI-Expanded)

XXI. Estrogen-induced upregulation and 3 '-UTR shortening of CDC6

AKMAN B. H., CAN T., Erson-Bensan A. E.

NUCLEIC ACIDS RESEARCH, vol.40, no.21, pp.10679-10688, 2012 (SCI-Expanded)

XXII. Large-Scale Signaling Network Reconstruction

Hashemikhabir S., Ayaz E. S., Kavurucu Y., Can T., Kahveci T.

IEEE-ACM TRANSACTIONS ON COMPUTATIONAL BIOLOGY AND BIOINFORMATICS, vol.9, pp.1696-1708, 2012 (SCI-Expanded)

XXIII. Using network context as a filter for miRNA target prediction

Sualp M., Can T.

BIOSYSTEMS, vol.105, pp.201-209, 2011 (SCI-Expanded)

XXIV. METADATA MANAGEMENT AND SEMANTICS IN MICROARRAY REPOSITORIES

Kocabas F., CAN T., BAYKAL N.

BALKAN JOURNAL OF MEDICAL GENETICS, vol.14, no.2, pp.49-63, 2011 (SCI-Expanded)

XXV. Bi-k-bi clustering: mining large scale gene expression data using two-level biclustering

Carkacioglu L., Atalay R. C., KONU KARAKAYALI Ö., Atalay V., CAN T.

INTERNATIONAL JOURNAL OF DATA MINING AND BIOINFORMATICS, vol.4, no.6, pp.701-721, 2010 (SCI-

Expanded)

XXVI. RRW: repeated random walks on genome-scale protein networks for local cluster discovery

MACROPOL K., Can T., Singh A. K.

BMC BIOINFORMATICS, vol.10, 2009 (SCI-Expanded)

XXVII. Integration of topological measures for eliminating non-specific interactions in protein interaction

networks

BAYIR M. A., GUNEY T. D., Can T.

DISCRETE APPLIED MATHEMATICS, vol.157, no.10, pp.2416-2424, 2009 (SCI-Expanded)

XXVIII. Discovering functional interaction patterns in protein-protein interaction networks

Turanalp M. E., Can T.

BMC BIOINFORMATICS, vol.9, 2008 (SCI-Expanded)

XXIX. Efficient molecular surface generation using level-set methods

Can T., Chen C., Wang Y.

JOURNAL OF MOLECULAR GRAPHICS & MODELLING, vol.25, no.4, pp.442-454, 2006 (SCI-Expanded)

XXX. Integrating multi-attribute similarity networks for robust representation of the protein space

Camoglu O., Can T., Singh A. K.

BIOINFORMATICS, vol.22, no.13, pp.1585-1592, 2006 (SCI-Expanded)

Articles Published in Other Journals

I. Informatics Olympiads in Turkey Team Selection and Training

CAN T., SIĞIRCI İ. O., ABUL O., DEMİRCİ M. F.

OLYMPIADS IN INFORMATICS, vol.9, pp.225-232, 2015 (Peer-Reviewed Journal)

Books & Book Chapters

I. Introduction to Bioinformatics

CAN T.

in: miRNomics MicroRNA Biology and Computational Analysis, Yousef Malik, Allmer Jens, Editor, Springer Protocols, pp.51-71, 2013

Refereed Congress / Symposium Publications in Proceedings

I. Transcription Elongation in Connection to 3'UTR Ends

ÖZGÜL İ., CAN T., ERSON BENSAN A. E.

Moleküler Biyoloji Kongresi MolBiyoKON18, İzmir, Turkey, 5 - 08 September 2018, pp.20

II. Deregulated APA and cancer specific APA isoforms

Begik O., Ercan M., Cingoz H., CAN T., ÖYKEN M., Erson-Bensan A. E.

Annual Meeting of the American-Association-for-Cancer-Research (AACR), Illinois, United States Of America, 14 - 18 April 2018, vol.78

III. Alternative Polyadenylation patterns for cancer classification

BEGIK O., ÖYKEN M., ÇINKILLI t., CAN T., ERSON BENSAN A. E.

HIBIT 2017, 28 - 30 June 2017

IV. Alternative Polyadenylation patterns for novel gene discovery and classification in cancer

BEĞİK O., ÖYKEN M., CAN T., ERSON BENSAN A. E.

Annual Meeting of the RNA Society, 30 May - 03 June 2017

V. APA isoform diversity in triple negative breast cancers

AKMAN TUNCER H. B., ÖYKEN M., AĞUŞ H. H., ERDEM M., YAVUZ E., CAN T., ERSON BENSAN A. E.

AACR Annual Meeting, USA, Washington, United States Of America, 1 - 05 April 2017, vol.77, pp.3374

VI. Identification of a potential cancer related lncRNA

BEĞİK O., cinkilli t., ÖYKEN M., CAN T., ERSON BENSAN A. E.

Functional Genomics to Systems Biology-EMBO, 12 - 15 November 2016

VII. Comparison of tissue disease specific integrated networks using directed graphlet signatures SÖNMEZ A. B., CAN T.

Proceedings of the 7th ACM International Conference on Bioinformatics, Computational Biology, and Health Informatics - BCB '16, Seattle, WA, USA, United States Of America, 2 - 05 October 2016

VIII. 3 UTR Length Changes In Connection With Estrogen Receptor Alpha In Breast Cancer

ÖYKEN M., CAN T., ERSON BENSAN A. E.

FEBS 2016, KUSADASI, Turkey, 3 - 08 September 2016

IX. 3 'UTR lengths change in connection with estrogen receptor-alpha in breast cancer

ÖYKEN M., CAN T., Erson-Bensan A. E.

41st FEBS Congress on Molecular and Systems Biology for a Better Life, Kusadasi, Turkey, 3 - 08 September 2016, vol.283, pp.207

X. Unsupervised identification of redundant domain entries in InterPro database using clustering techniques

RİFAİOĞLU A. S., DOĞAN T., CAN T.

6th ACM Conference on Bioinformatics, Computational Biology, and Health Informatics, BCB 2015, Georgia, United States Of America, 9 - 12 September 2015, pp.505-506

XI. Reconstruction of the Temporal Signaling Network in Salmonella Infected Human Cells

güngör b., EREN ÖZSOY Ö., AYDIN SON Y., CAN T., nurcan t.

23rd Annual International Conference on Intelligent Systems for Molecular Biology and the 14th European Conference on Computational Biology (ISMBECCB'2015), Dublin, Ireland, 10 - 14 July 2015

XII. Dbchem A database query based solution for the chemical compound and drug name recognition task

ATA C., CAN T.

Proceedings of the Fourth BioCreative Challenge Evaluation Workshop, BioCreative IV,, 1 - 03 October 2013, vol.2, pp.42-46

XIII. M4B: A novel method for designing and ordering of the genetic devices

Hashemikhabir S., Ersoy G., Oguz G., Yaldiz B., Tuncel Y., Budak G., Karaaslan S., AYDIN SON Y., CAN T. 2012 7th International Symposium on Health Informatics and Bioinformatics, HIBIT 2012, Cappadocia, Turkey, 19 - 22 April 2012, pp.123-127

XIV. Parallel SPICi

Hashemikhabir S., CAN T.

6th International Symposium on Health Informatics and Bioinformatics (HIBIT), İzmir, Turkey, 2 - 05 May 2011, pp.86-90

XV. CONSTRUCTING SIGNALING PATHWAYS FROM RNAI DATA USING GENETIC ALGORITHMS

Ayaz E. S., CAN T.

6th International Symposium on Health Informatics and Bioinformatics (HIBIT), İzmir, Turkey, 2 - 05 May 2011, pp.95-98

XVI. The effect of representative training dataset selection on the classification performance of the promoter sequences

YAMAN A. G., CAN T.

6th International Symposium on Health Informatics and Bioinformatics, HIBIT 2011, İzmir, Turkey, 2 - 05 May 2011, pp.55-58

XVII. Coevolution based prediction of protein-protein interactions with reduced training data

Pamuk B., CAN T.

2010 5th International Symposium on Health Informatics and Bioinformatics, HIBIT 2010, Antalya, Turkey, 20 - 22 April 2010, pp.187-193

XVIII. Text Classification in the Turkish Marketing Domain for Context Sensitive Ad Distribution

Engin M., Can T.

24th International Symposium on Computer and Information Sciences, Güzelyurt, Cyprus (Kktc), 14 - 16 September 2009, pp.105-110

XIX. Human Skeletal and Muscle Deformation Animation Using Motion Capture Data

Bayer A. O., Sevinc A. M., CAN T.

16th International Conference in Central Europe on Computer Graphics, Visualization and Computer Vision, Plzen, Czech Republic, 4 - 07 February 2008, pp.167-168

XX. Analysis of protein-protein interaction networks using random walks

Can T., Orhan C., Singh A. K.

5th International Workshop on Bioinformatics, BIOKDD 2005 - In Conjunction with 11th ACM SIGKDD International Conference on Knowledge Discovery and Data Mining, KDD 2005, Chicago, IL, United States Of America, 21 August 2005, pp.61-68

XXI. CTSS: A robust and efficient method for protein structure alignment based on local geometrical and biological features

Can T, WANG Y.

2nd International Computational Systems Bioinformatics Conference, California, United States Of America, 11 - 14 August 2003, pp.169-179

Supported Projects

MUYAN M., DEMİRALAY Ö. D., CAN T., Project Supported by Higher Education Institutions, Genom Düzeyindeki Protein Bağlanma Verilerinin Biyoinformatik ve Deneysel Yöntemlerle Analizi, 2021 - 2022

ERSON BENSAN A. E., GÜLEZ B., BANERJEE S., ÖZKAN S., CAN T., Project Supported by Higher Education Institutions, Yaygın kanser tiplerinde transkriptom 3'UTR isoform çeşitliliğinin incelenmesi, 2017 - 2019

CAN T., Project Supported by Higher Education Institutions, G-Proteini Kenetli Reseptörlerin Etkileşimleri:hesaplamalı Yöntemlerin Ve Deneysel Çalışmaların Sentezlenmesi., 2010 - 2019

SON Ç. D., ÇELEBİ M., CAN T., Project Supported by Higher Education Institutions, G-proteine kenetli reseptörlerin etkileşim bölgelerinin işlemsel olarak 3D modellenmesi, 2017 - 2017

MUYAN M., ERSON BENSAN A. E., CAN T., Project Supported by Higher Education Institutions, Meme Kanseri Model Hücrelerinde Tüm Genom Düzeyinde RNA Ekspresyon Analizleri, 2015 - 2016

ERSON BENSAN A. E., ozdemir n., CAN T., Project Supported by Higher Education Institutions, CDC6 alternatif poliadenilasyonunun meme kanseri hücrelerinde incelenmesi, 2013 - 2015

Patent

CAN T., APADetect: Alternatif Poliadenilasyon Tespiti, Patent, CHAPTER G Physics, Standard Registration, 2016

Metrics

Publication: 53

Citation (WoS): 636 Citation (Scopus): 676 H-Index (WoS): 12 H-Index (Scopus): 13

Non Academic Experience

METU University of California, Santa Barbara University of California, Santa Barbara

METU