Eli Levy Karin April 2020

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Education		
2018 –	Postdoc. Advisor: Dr. Johannes Söding	Quantitative and Computational Biology, MPIBPC
2013 – 2017	Ph.D. Advisors: Prof. Tal Pupko and Prof. Itay Mayrose	Faculty of Life Sciences, TAU
	<u>Dissertation</u> : Statistical Techniques in Molecular Evolution: Improving insilico Sequence Simulations and Detecting Genotype-Phenotype Associations	
2012 – 2014	M.Sc in Theoretical and Mathematical Biology (direct Ph.D track)	Faculty of Life Sciences, TAU
	Grade average: 98	
2008 – 2011	B.Sc in Computer Science. Grade average: 90 (magna cum laude)	Faculty of Exact Sciences, TAU
	- Participated in "First Steps in Research for Excellent Students". Studied aspects of the human metabolic network at Prof. Eytan Ruppin's lab.	
	Grade: 97	
	<ul> <li>Participated in "Parallel Computation of Computer Vision Algorithms" workshop. Final project focused on <u>accelerating the calculation of optical flow fields</u>. Grade: 99</li> </ul>	
2005 – 2008	<b>B.Sc</b> in <b>Biology</b> . Grade average: 90	Faculty of Life Sciences, TAU

#### **Publications**

- \* denotes equal contribution / \* denotes co-correspondence
- Halabi K, <u>Levy Karin E+</u>, Guéguen L, & Mayrose I+. TraitRELAX A codon model for associating phenotypic traits with altered selective patterns of sequence evolution. *Submitted*. 2020.
- Levy Karin E, Mirdita M, & Söding J. MetaEuk sensitive, high-throughput gene discovery and annotation for large-scale eukaryotic metagenomics. *Microbiome*. 2020; *8:48*.
- Levy Karin E, Ashkenazy H, Hein J, & Pupko T. A simulation-based approach to statistical alignment. *Systematic Biology*. 2019; 2:252-266.
- Ashkenazy H, Sela I, <u>Levy Karin E</u>, Landan G, & Pupko T. Multiple sequence alignment averaging improves phylogeny reconstruction. *Systematic Biology*. 2019; 1:117-130.
- Mushegian A, <u>Levy Karin E</u>, & Pupko T. Sequence analysis of malacoherpesvirus proteins: pan-herpesvirus capsid module and replication enzymes with an ancient connection to "Megavirales". *Virology*. 2018; 513: 114-128.
- 9 Lavi B, <u>Levy Karin E</u>, Pupko T, & Hazkani-Covo E. The prevalence and evolutionary conservation of inverted repeats in proteobacteria. Genome Biology and Evolution. 2018; 3: 918–927.
- 8 <u>Levy Karin E\*</u>, Ashkenazy H\*, Wicke S, Pupko T, & Mayrose I. TraitRateProp: a web server for the detection of associations between phenotypic trait changes and specific sequence sites. *Nucleic Acids Research.* 2017; 45:W260-W264.
- Ashkenazy H\*, <u>Levy Karin E\*</u>, Mertens Z, Cartwright R, & Pupko T. SpartaABC: a web server to simulate sequences with indel parameters inferred using an approximate Bayesian computation algorithm. *Nucleic Acids Research*. 2017; 45:W453-W457.

- 6 <u>Levy Karin E</u>, Wicke S, Pupko T, & Mayrose I. An integrated model of phenotypic trait changes and site-specific sequence evolution. *Systematic Biology*. 2017; 6:917–933.
- Levy Karin E\*, Shkedy D\*, Ashkenazy H, Cartwright R, & Pupko T. Inferring rates and length-distributions of indels using approximate Bayesian computation. *Genome Biology and Evolution*. 2017; 9:1280-1294.
- 4 Preisner H, <u>Levy Karin E</u>, Poschmann G, Stühler K, Pupko T, & Gould S. The cytoskeleton of parabasalian parasites comprises proteins that share properties common to intermediate filament proteins. *Protist.* 2016; 167:526–543.
- 3 <u>Levy Karin E\*</u>, Rabin A\*, Ashkenazy H, Shkedy D, Avram O, Cartwright R, & Pupko T. Inferring indel parameters using a simulation-based approach. *Genome Biology and Evolution*. 2015; 7:3226-3238.
- Levy Karin E, Susko E, & Pupko T. Alignment errors strongly impact likelihood-based tests for comparing topologies. *Molecular Biology and Evolution*. 2014; 31:3057-3067.
- 1 Cohen O, Ashkenazy H, <u>Levy Karin E</u>, Burstein D, & Pupko T. CoPAP: co-evolution of presence-absence patterns. *Nucleic Acids Research*. 2013; 41:W232-W237.

### **Awards and Scholarships**

2018 – 2020	FEBS long-term postdoctoral fellowship	Federation of European
2018	EMBO long-term non-stipendiary postdoctoral fellowship	Biochemical Societies European Molecular Biology Organization
2017	Ernst Mayr award for best talk	Society of Systematic Biologists
2017	Travel scholarship	Society for Molecular Biology and Evolution
2017	Travel scholarship	Constantiner Institute for Molecular Genetics
2016	Travel scholarship	Manna Center, Faculty of Life Sciences, TAU
2015	Award for excellent achievements in teaching	Faculty of Life Sciences, TAU
2015	Dan David Prize for Ph.D students, Future – Bioinformatics category	Dan David Foundation
2015	Excellent Research Student Prize for the academic year 2014-2015	Edmond J. Safra Center, TAU
2014	Walter M. Fitch award finalist for Ph.D and post-doctoral students	Society for Molecular Biology and Evolution
2014 – 2017 2013	Ph.D fellowship Award for excellent achievements in teaching, research and studies	Edmond J. Safra Center TAU graduate school

#### Posters and Oral Presentations (international meetings)

<u>Levy Karin E</u> , Mirdita M, & Soeding J. MetaEuk – sensitive, high-throughput gene discovery and annotation for large-scale eukaryotic metagenomics. Microbiome COSI. <i>ISMB/ECCB</i> . July 21-25, 2019, Basel, Switzerland.	talk
<u>Levy Karin E</u> , Wicke S, Pupko T, & Mayrose I. An integrated model of phenotypic trait changes and site-specific sequence evolution. Mayr symposium. <i>Evolution</i> . June 23-27, 2017, Portland, Oregon, USA.	talk
<u>Levy Karin E</u> , Wicke S, Pupko T, & Mayrose I. An integrated model of phenotypic trait changes and site-specific sequence evolution. <i>SMBE</i> . July 2-6, 2017, Austin, Texas, USA.	poster

<u>Levy Karin E\*</u>, Rabin A\*, Ashkenazy H, Shkedy D, Avram O, Cartwright R, & Pupko T. Inferring indel parameters poster using a simulation-based approach. *SMBE*. July 12-16, 2015, Vienna, Austria.

<u>Levy Karin E</u>, Susko E, & Pupko T. Alignment errors strongly impact likelihood-based tests for comparing talk topologies. Fitch symposium. *SMBE*. June 8-12, 2014, San Juan, Puerto Rico, USA.

### Student supervision

Vladyslav	slav The EukBook project: Large-scale search for novel eukaryotic proteins	MSc thesis Oct19 – Apr20
Dembrovskyi	from public sequencing data	ivisc triesis Oct19 – Apr 20

# **Teaching Experience**

2013 – 2015	Introductory course in Math for Biology students (8888-42002)	Kahanoff Foundation,	
	I developed and taught a course aiming to better prepare new Biology undergrads for the first year of their studies.	Faculty of Life Sciences, TAU	
2012 – 2016	Perl Programming for Biology (0455-3065)	Faculty of Life Sciences, TAU	
(fall semesters)	I was responsible for the entire course for which I received an award from the Tel-Aviv University Graduate school.		
Oct 2010 – Jun	Computer Science for middle school students	Bialik Rogozin School, Tel- Aviv	
2011	As part of my studies for an additional B.Sc in Computer Science, I developed lesson plans and taught for a year on a weekly basis		

## **Academic Experience**

2015, 2016	Visiting scholar at the Molecular Evolution institute, Heinrich-Heine-Universität, Düsseldorf, Germany
2016	Visiting scholar at the IEB, Westfälische Wilhelms-Universität, Münster, Germany

### Referee Service (for scientific journals)

Referee Service (for Scientific Journals)	
Journal of Molecular Biology & Evolution	01.2019, 03.2020
Journal of Molecular Evolution	06.2016, 12.2017, 06.2018
Journal of Microbial Genomics	02.2016
Journal of BMC Evolutionary Biology	06.2015, 09.2016, 01.2018
Journal of <i>Bioinformatics</i>	Apr 2015

## **Employment**

2011 – 2012	Computational genomics scientist	Evogene Ltd.
Nov 2007 – Mar 2008	Molecular biology researcher	Prof. Gil Segal's lab, Faculty of Life Sciences TAU
2005 – 2006	Business intelligence researcher	Adkit Ltd.

# **Non-Academic Activity**

2011 – 2015	Volunteer at the open clinic	Physicians for Human Rights
2014 – 2015	Volunteer private tutor for math	
Feb 2014 – Jul 2014	Volunteer computers instructor	Migrant Worker & Refugee Community Education Center (CEC)
Oct 2013 – Jan 2014	Volunteer Hebrew teacher	CEC