Aaron R. Quinlan, Ph.D.

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Information Department of Public Health Sciences (F): 434.924.1312

Center for Public Health Genomics arq5x@virginia.edu

University of Virginia cphg.virginia.edu/quinlan

Research Chromosome stability and somatic genome evolution; Genomics software development; Cancer

Interests genomics; Population genomics; Genetics of complex disease

EDUCATION Boston College, Chestnut Hill, MA, USA Ph.D., Biology, 2008

College of William and Mary, Williamsburg, VA, USA

B.S., Computer Science, 1997

ACADEMIC University of Virginia, Charlottesville, VA, USA
EXPERIENCE Assistant Professor of Public Health Sciences 2011-

Center for Public Health Genomics

University of Virginia, Charlottesville, VA, USA

NRSA Postdoctoral Fellow (NHGRI) 2008-2011

HONORS AND Co-chair, NHLBI Exome Sequence Project Structural Variation Group. 2011-AWARDS Instructor for CSHL Advanced Sequencing Technologies Course. 2009-

> Fund for Excellence in Science and Technology Awardee, UVa. (1 of 5). 2011 Ruth L. Kirschstein (NRSA / F32) Postdoctoral Fellowship, NHGRI. 2009-2010

Presidential Fellowship, Boston College. 2004-2007

Editing and Ad hoc reviewer for Genome Research, Genome Biology, Bioinformatics, Review BMC Bioinformatics, Bioessays, and Genes

NIH: Special Emphasis Panel for Computational Analysis of ENCODE (U01) (16-May-2012)

Publications † denotes corresponding author

★ denotes joint first authors

- 16. Krumm N, Sudmant PH, Ko A, O'Roak BJ, NHLBI Exome Sequencing Project, **Quinlan AR**, Nickerson DA, Eichler EE. Copy number variation detection and genotyping from exome sequence data. Under review.
- 15. Layer R†, **Quinlan AR**†, Robins G, Skadron K. Binary Interval Search (BITS): A Massively Parallel Interval Intersection Algorithm. Under review.
- 14. **Quinlan AR** and Hall IM. Structural variation in mammalian genomes. Invited book chapter for Methods in Molecular Biology. In press.
- 13. Quinlan AR, Boland MJ, Leibowitz ML, Shumilina S, Pehrson SM, Baldwin KK, Hall IM. Paired-end DNA sequencing of induced pluripotent stem cell genomes reveals rare structural mutations and retroelement stability. *Cell Stem Cell*, 9:366-373, 2011.

- Keene KL, Quinlan AR, Hou X, Hall IM, Mychaleckyj, Onengut-Gumuscu S, Concannon P. Evidence for two independent associations with type 1 diabetes at the 12q13 locus. Genes and Immunity, 13:66-70, 2011.
- 11. **Quinlan AR**, Hall IM. The landscape of complex chromosomal rearrangements in the germline and somatic lineages. *Trends in Genetics*, 18:43-53, 2011.
- 10. Dale R, Pedersen B, Quinlan AR†. Pybedtools: a flexible Python library for manipulating genomic datasets and annotations. *Bioinformatics*, 24:3423-3424, 2011. packages.python.org/pybedtools/
- 9. Barnett D, Garrison E, **Quinlan AR**, Stromberg M, Marth G. BamTools: a C++ API and toolkit for analyzing and managing BAM files. *Bioinformatics*, 12:1691-1692, 2011. code.google.com/p/bamtools
- 8. 1000 Genomes Project Consortium.. A map of human genome variation from population-scale sequencing. *Nature* 7319:1061-73, 2010.
- 7. **Quinlan AR** and Hall IM. BEDTools: A flexible framework for comparing genomic features. *Bioinformatics*, 6:841-842, 2010. code.google.com/p/bedtools
- 6. Quinlan AR, Clark RA, Sokolova, S, Leibowitx ML, Zhang Y, Hurles ME, Mell JC, Hall IM. Genome-wide mapping and assembly of structural variant breakpoints in the mouse genome. *Genome Research*, 20:623-635, 2010. code.google.com/p/hydra-sv
- 5. Sackton, TB, Kulathinal RJ, Bergman CM, Quinlan AR, Dopman E, Marth GT, Hartl DL, Clark AG. Population Genomic Inferences from Sparse High-Throughput Sequencing of Two Populations of Drosophila melanogaster. *Genome Biol Evol*, 1:439-455, 2009.
- 4. Smith D, Quinlan AR⋆, Peckham HR, et al. Rapid whole-genome mutational profiling using next-generation sequencing technologies. Genome Research, 18:1638-1642, 2008.
- 3. Hillier LW, Marth GT, **Quinlan AR**, et al. Whole Genome Sequencing and SNP Discovery for C. elegans using massively parallel sequencing-by-synthesis. *Nature Methods*, 5:183-188, 2008.
- 2. Quinlan AR, Stewart D, Stromberg M, Marth GT. PyroBayes: Accurate quality scores for 454 Life Science pyrosequences. *Nature Methods*, 5:179-181, 2008.
- 1. Quinlan AR, Marth GT. Primer-site SNPs mask mutations. Nature Methods, 4:192, 2007.

RESEARCH SUPPORT

Project Title: New algorithms and tools for large-scale genomic analysis.

PI: Aaron Quinlan

Source: NIH/NHGRI (R21 ES020521-01)

Amount: \$437,112; Period funded: 19-Apr-2012 - 31-Mar-2016

Project Title: Identification of radiation sensitivity alleles by whole exome sequencing.

PI: Pat Concannon

Co-investigator: Aaron Quinlan

Source: NIH/NIEHS (R21 ES020521-01)

Amount: \$153,376; Period funded: 19-Aug-2011 - 31-Jul-2013

Project Title: New oncogenes and regions of genome instability in ovarian cancer.

PI: Aaron Quinlan

Source: University of Virginia Fund for Excellence in Science and Technology (FEST)

Amount: \$50,000; Period funded: 01-May-2011 - 30-Apr-2012

Project Title: Defining the genomic architecture of glioblastoma for improved therapy. PI: Aaron Quinlan

Source: University of Virginia Cancer Center Pilot Fund Amount: \$53,125; Period funded: 01-Apr-2011 - 31-Mar-2012

Project Title: Rates and patterns of recurrent structural variation in the mouse genome.

PI: Aaron Quinlan

Source: NIH/NHGRI (F32 HG005197-02) Period funded: 01-Aug-2009 - 31-Dec-2010

Invited Lectures

Towards a map of structural variation in the Exome Sequencing Project. NHLBI Exome Sequencing Project In-Person Meeting, March 28, 2012

Exploring the origin and extent of structural variation in human genomes.

Deans New Faculty Seminar Series, University of Virginia School of Medicine, Jan 19, 2012

ESP Structural Variation Project Group: goals, initial results, and future work.

NHLBI Exome Sequencing Project In-Person Meeting, June 9, 2011

Large-Scale Characterization of SV Breakpoints in Cancer.

Keystone Symposium on The Functional Impact of Structural Variation, Jan. 11, 2011

Efficient discovery of structural instability in repetitive regions of mammalian genomes.

Advances in Genome Biology and Technology, Feb. 2009

Approaches to rare allele discovery: More samples or more depth per sample? 1000 Genomes Analysis Meeting, Cold Spring Harbor Laboratories, May 2008.

ACADEMIC SERVICE

Member of the Univ. of Virginia Bioinformatics Core Advisory Committee.

01-Nov-2011 - Present.

Served on the Univ. of Virginia Bioinformatics Core Director Search Committee.

Summer 2011.

Teaching

Instructor for CSHL Advanced Sequencing Technologies Course.

2009 - Present

Guest Lecturer for graduate course in genomics (BIOCH 5080).

15-Feb-2011

THESIS COMMITTEES

Current

Michael Lindberg (Ph.D., Biochemistry and Molecular Genetics, qualified 2012)