# Eric Edward Bryant, PhD

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## **Education & Experience**

2020-Pres. Amgen, Postdoctoral Research Fellow

Role: Developed and executed a research plan studying the genetics of mammalian

recombinant protein expression using CRISPR methods and transcriptomics.

Groups: Mammalian Expression & Cell Engineering / Integrated Protein Design

Managers: Irwin Chen & Rene Hubert

Keywords: High-throughput mammalian expression / NGS / RNAseq / Arrayed & pooled CRISPR

screening / Protein Engineering / Multispecifics / Biologics Discovery

2020 Spring Roche/Genentech (Contract), Clinical Genomics Data Science Consultant

Role: Evaluated variant calling from RNAseq while on short-term contract.

Department: Pharma Research & Early Development Informatics (pREDi)

Managers: Venus So & Vitalay Fomin

*Keywords:* Genome Analysis Toolkit (GATK) / RNAseq / Real World Data / High-performance

computing cluster (HPC) / Bioinformatics / CWL, Nextflow, Seven Bridges Genomics

2011-'18 PhD, Columbia University

Department: GSAS Biological Sciences

Thesis: Systems genetics of DNA damage tolerance —

cisplatin, RAD5 & CRISPR-mediated nonsense

Mentor: Rodney Rothstein
Co-Mentor: Alberto Ciccia

Committee: Songtao Jia, Elizabeth Miller & Matthew Weitzman

Keywords: Systems genetics / Genetic interaction networks / Landscape enrichment

analysis / Synthetic lethality / DNA replication, recombination & repair /

Chromosome mobility / CRISPR iSTOP / Bioinformatics / R statistical programming,

package and shiny app development

2009-'11 Gene Oracle, Research Assistant

*Role:* De-novo gene synthesis & degenerate codon library construction.

2006–'09 BS, University of California Los Angeles

Department: UCLA Microbiology, Immunology & Molecular Genetics

## Software examples

2019 screenmill R package: capture, annotate, quantify, review and analyze time-series

**colony growth.** This package was used for colony quantification and interaction analysis in Bryant et al. *NAR* 2019. A nice example analysis using screenmill can be found in Figure 1D. A

shiny app is included to enable manual review of processed images.

2017 iSTOP R package: design guides to introduce stop codons with CRISPR-mediated

**base editors.** This package was written to facilitate guide design for Billon et al. *Mol. Cell* 2017. iSTOP can be configured to generate any desired missense mutation using any hypothetical base editor. This package was used to design a base editing library for Cuella-Martin et al. *Cell* 2021.

### **Awards**

| 2018     | Departmental distinction for PhD dissertation defense                              |
|----------|--|
| 2017—'18 | TL1 NIH training grant, clinical and translational research                        |
| 2016—'17 | T32 NIH training grant, cancer biology   |
| 2013—'15 | T32 NIH training grant, biological sciences  |
| 2013     | James Howard McGregor award (student with unusual promise as a teacher of zoology) |

#### Lead contribution

2019-09-26 Rad5 dysregulation drives hyperactive recombination at replication forks resulting in cisplatin sensitivity and genome instability.

Bryant EE, Šunjevarić I, Berchowitz L, Rothstein R, Reid RJD.

Nucleic Acids Research. 2019 Sep 26;47(17):9144-9159

PMID: 31350889 - PMCID: PMC6753471 - DOI: 10.1093/nar/gkz631

2019-01-09 Systems genetics of DNA damage tolerance – Cisplatin, *RAD5* & CRISPR-mediated nonsense. **Bryant EE**.

Columbia University.

DOI: 10.7916/d8-k1do-kb09

2017-09-21 CRISPR-mediated base editing enables efficient disruption of eukaryotic genes through induction of STOP codons.

Billon P\*, Bryant EE\*, Joseph SA, Nambiar TS, Hayward SB, Rothstein R, Ciccia A.

Molecular Cell. 2017 Sep 21;67(6):1068—1079.e4

PMID: 28890334 - PMCID: PMC5610906 - DOI: 10.1016/j.molcel.2017.08.008

\*co-first authors

### Supporting contribution

2022-03-24 Temporal coordination between chromosome mobility and homologous recombination. Joseph F, Lee SJ, **Bryant EE**, Reid RJD, Šunjevarić I, Rothstein R. BioRxiv: 10.1101/2022.03.24.485580v1 – DOI: 10.1101/2022.03.24.485580

2021-02-18 Functional interrogation of DNA damage response variants with base editing screens.

Cuella-Martin R, Hayward SB, Fan X, Chen Xiao, Huang JW, Taglialatela A, Leuzzi G, Zhao J, Rabadan R, Lu C, Shen Y, Ciccia A.

Cell. 2021 Feb 18;184(4):1081-1097.e19.

PMID: 33606978 - PMCID: PMC8018281 - DOI: 10.1016/j.cell.2021.01.041

\* Acknowledged for bioinformatics support. I designed the CRISPR base editing guide library.

2020-08-26 Measuring chromosome pairing during homologous recombination in yeast.

Joseph F, Lee SJ, Bryant EE, Rothstein R.

*Methods in Molecular Biology*. 2021;2153:253—265.

PMID: 32840785 - DOI: 10.1007/978-1-0716-0644-5 18

2019-10-01 DNA damage triggers increased mobility of chromosomes in G1 phase cells.

Smith MJ, Bryant EE, Joseph FJ, Rothstein R.

Molecular Biology of the Cell. 2019 Oct 1;30(21):2620-2625

PMID: 31483739 - PMCID: PMC6761769 - DOI: 10.1091/mbc.E19-08-0469

2018-09-01 Increased chromosomal mobility after DNA damage is controlled by interactions between the recombination machinery and the checkpoint.

Smith MJ, **Bryant EE**, Rothstein R.

Genes & Development. 2018 Sep 1;32(17-18):1242—1251

PMID: 30181361 - PMCID: PMC6120718 - DOI: 10.1101/gad.317966.118

2016-10-01 A synthetic dosage lethal genetic interaction between CKS1B and PLK1 is conserved in yeast and

human cancer cells.

Reid RJD, Du X, Šunjevarić I, Rayannavar V, Dittmar J, Bryant EE, Maurer M, Rothstein R.

Genetics. 2016 Oct 1;204(2):807-819

PMID: 27558135 - PMCID: PMC5068864 - DOI: 10.1534/genetics.116.190231