# Stephen Y. Hwang

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Skills

Programming R (dplyr, tidyr, ggplot), Python, Perl, bash, Shiny, RMarkdown, knitr, Jupyter

Working knowledge of Pub/Sub, Dataproc, Dataflow, Cloud Composer (Airflow), Spark,

Hadoop, Dialogflow, AutoML

**Warehousing** SQL, BigQuery, Snowflake

Working knowledge of Redshift, MongoDB, Datastore, Hive, BaseX

Other technologies Google Data Studio, Tableau, Jira, Confluence, Git, Unix/Linux environments, AWS, REDCap

Working knowledge of Docker, Kubernetes

Professional Experience \_

### Senior Bioinformatics Scientist, RTI International, Research Triangle Park, NC

Jun 2015 - present

- Supervise multiple teams of analysts and developers as program manager and technical task lead
- Design and build extract, transform & load (ETL) pipelines on AWS & GCP to ingest and warehouse data
- Use R to implement data transformations (dplyr, tidyr) and to create novel visualizations (ggplot)
- Utilize Snowpipe to ingest Twitter JSON data into Snowflake
- Create intermediate tables from raw data to support ad-hoc analyses and reporting
- Design and create tools and pipelines to support analytics needs and facilitate analysis and insights
- Interface with technical and non-technical stakeholders to resolve complex business and technical issues

#### Data Engineer (Consultant), Animal Cancer DX, LLC, Raleigh, NC

Jun 2021 - present

- Designed and created an ETL pipeline to aggregate and stage experimental data on AWS S3, ingest data into RDS;
  application hosted on EC2 automatically generates and distributes reports with statistical analyses to stakeholders
- Designed and created a Shiny application to be hosted in a Docker container
- Designed and built data browser dashboard using Shiny and ggplot

#### Bioinformatics Data Scientist, Johns Hopkins Medical Institutions, Baltimore, MD

Aug 2013 - Jun 2015

- Completed 2-year data mining project in 10 weeks using R, SQL, and bash
- Designed and created an ETL batch process to stage public health datasets and ingest into a SQL database
- Used Python to implement data transformations (pandas) and create novel visualizations (matplotlib, ggplot)
- Designed and implemented analysis pipelines for large genomic datasets for exploratory analysis and publication
- Established unsupervised machine learning methods into quality control process

## $\textbf{Bioinformatics Analyst}, \ \mathsf{Johns\ Hopkins\ Medical\ Institutions}, \ \mathsf{Baltimore}, \ \mathsf{MD}$

Oct 2011 - Aug 2013

• Designed and created an ETL pipeline to ingest and transform public health data sources for analytic use, increasing the efficiency by 700% over the legacy workflow

#### Bioinformatics Researcher, Putonti Lab, Loyola University, Chicago, IL

Jan 2010 - Jan 2011

Created pipeline to aggregate and analyze genomic data extracted from NCBI repository

Education

## Johns Hopkins University

M.Sc. Bioinformatics

Baltimore, MD Aug 2013

**Loyola University** B.Sc. Bioinformatics

Chicago, IL May 2011

Publications

Gern, J. E., Jackson, D. J., Lemanske, R. F., Seroogy, C. M., Tachinardi, U., Craven, M., **Hwang, S. Y.**, ... Bacharier, L. B. (2019). The Children's Respiratory and Environmental Workgroup (CREW) Birth Cohort Consortium: Design, methods, and study population. *Respiratory Research*, 20(1), 115. doi:10.1186/s12931-019-1088-9

Vaz, M., **Hwang, S. Y.**, Kagiampaki, I., Phallen, J., Patil, A., O'Hagan, H. M., ... Baylin, S. B. (2017). <u>Chronic cigarette smoke-induced epigenomic changes precede sensitization of bronchial epithelial cells to single-step transformation by KRAS mutations</u>. *Cancer Cell*, 32, 360–376. doi:10.1016/j.ccell.2017.08.006