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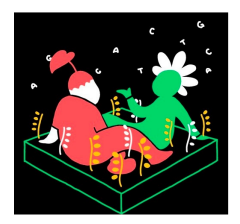
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Running4Research This Sunday, G2P is taking to the streets for the annual Boston Athletics Association 10K. Join us in supporting our team - your donation will help fund new and innovative projects that will accelerate the implementation of genomic medicine and the promise of precision health.



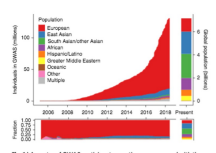


Fig. 1 | Ancestry of GWAS participants over time, as compared with the global population. Cumulative data, as reported by the GWAS catalog<sup>16</sup>. Individuals whose ancestry is 'not reported' are not shown.

PRESS In this podcast, Carrie Blout, MS, CGC, discusses the impact of whole genome sequencing, the rollout of a new <u>Preventative Genomics Clinic</u> at Brigham and Women's Hospital, and the future of genomic medicine.

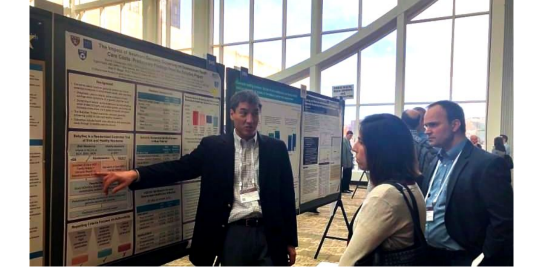
PERSPECTIVE Investigators from Broad and Mt. Sinai demonstrate that polygenic risk scores developed in one ancestry group cannot be accurately applied to another, making it even more critical to enroll diverse populations in genomic research.



THE FRANCA FUND With an aim to transform the practice of medicine so that everyone can access their genomic information to improve their health, we hosted leaders in science, business and industry, as well as prominent artists and fashion figures to discuss the Fund's vision, mission and goals.



OPINION <u>BabySeq</u> investigators found that 10-15% of enrolled babies have a potentially harmful genetic change that puts them at risk for a genetic condition, some of which could immediately impact the baby's parents. This left us faced with the decision of whether - and how - to return these results.



**TALKS** In April, genetic counselor Carrie Blout, MS, CGC, and econogenomics expert Kurt Christensen, PhD, presented at the ACMG Annual Clinical Genetics Meeting in Seattle, sharing data from our BabySeq newborn sequencing project.

## **Support Our Research!**



Consider making a tax-deductible gift to support our mission to accelerate the implementation of genomic medicine and the promise of precision health.

## Give today!

Donations support our current research projects in genomics and personalized medicine to predict and prevent conditions like cancer, heart disease and Alzheimer's, in addition to training junior faculty and students.





