Genes Regulate Blood Pressure, but “Environments” Cause Hypertension

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INTRODUCTION

Hypertension is among the first traits studied using genome-wide association study (GWAS). At first, GWAS by the Welcome Trust Case Control Consortium in 2007 did not identify any genome-wide significant single nucleotide polymorphism (SNP) (The Wellcome Trust Case Control Consortium, 2007), however, it was still asserted that GWAS would open the door to find the “missing heritability” of hypertension. In the following years, GWASs of hypertension were still unsuccessful in identifying robust loci until 2009. After shifting attention to the underlying quantitative traits (systolic blood pressure and diastolic blood pressure) and application of GWA meta-analyses of multiple cohorts to enlarge the sample size, the Global Blood Pressure Genetics (Global BPgen) Consortium and the Cohorts for Heart and Aging Research in Genome Epidemiology (CHARGE) Consortium identified 13 genome-wide significant signals (Levy et al., 2009; Newton-Cheh et al., 2009), although these SNPs had very small effect sizes and accounted for <0.2% of the overall blood pressure variation in the study populations.

WILL GWAS UNLOCK THE GENETIC BASIS OF HYPERTENSION?

There was a heated debate in HYPERTENSION on December 2010. Dominiczak and Munroe described the successes of aforementioned two studies, and predicted a bright future for GWAS in hypertension (Dominiczak and Munroe, 2010). On the contrary, Kurtz contended that GWAS had failed, and would continue to fail, to delineate the genetic basis of hypertension. He suggested that efforts and dollars should be shifted to other strategies and technologies that may hold greater chance for advancing our understanding of the genetic etiology of hypertension (Kurtz, 2010). Despite different opinions, both pro and con sides shared some common ground: low-frequency/rare variants are important, and the sample size should be further enlarged.

In 2016, three large-scale GWASs of hypertension were published in NATURE GENETICS (Ehret et al., 2016; Liu et al., 2016; Surendran et al., 2016). After applying newly designed microarray chips and meta-analyses of multiethnic populations with unprecedented large sample sizes (>300,000), these studies identified several new common loci of modest effects, and provided insights into the impact of low-frequency/rare variants. Subsequently, other genome-wide analyses of blood pressure traits (systolic, diastolic, and pulse pressure) were carried out in people of European ancestry drawn from UK Biobank (UKB), the International Consortium of Blood Pressure Genome Wide Association Studies (ICBP), the Genetic Epidemiology Research on Adult Health and Aging (GERA) cohort, and other combined cohorts with very large sample size (Hoffmann et al., 2017; Warren et al., 2017; Evangelou et al., 2018). These studies identified...
| PMID        | Year | Disease/trait | Initial sample size | Replication sample size | Major finding                                                                                                                                 |
|-------------|------|---------------|---------------------|-------------------------|------------------------------------------------------------------------------------------------------------------------------------------------|
| 19430479    | 2009 | DBP, SBP, HT  | 29,136 European ancestry individuals | 34,433 European ancestry individuals | They identified 13 SNPs for SBP, 20 for DBP, and 10 for hypertension.                                                                                   |
| 19430483    | 2009 | DBP, SBP      | 34,433 European ancestry individuals | Up to 100,347 European ancestry individuals, up to 12,889 Indian Asian ancestry individuals | They identified association between systolic or diastolic blood pressure and common variants in eight regions near the CYP17A1, CYP1A2, FG5, SH2B3, MTHFR, c10orf107, ZNF652, and PLCD3 genes. |
| 21572416    | 2011 | BP            | 19,608 East Asian ancestry individuals | 30,765 East Asian ancestry individuals | They identified genome-wide significant associations with SBP or DBP, which included variants at four new PP loci (ST7L-CAPZA1, FIGN-GRB14, ENPEP, and NPR3) and a newly discovered variant near TBX3. |
| 21909110    | 2011 | PP            | 74,064 European ancestry individuals | 48,607 European ancestry individuals | They identified genetic variants at 12 new loci to be associated with blood pressure.                                                                   |
| 26390057    | 2015 | DBP, SBP, PP  | 31,516 East Asian ancestry individuals, 35,352 European ancestry individuals, 33,126 South Asian ancestry individuals | 87,205 individuals, 48,268 East Asian ancestry individuals, 68,456 European ancestry individuals, 16,328 South Asian ancestry individuals | They identified 30 new blood pressure- or hypertension-associated loci.                                                                                   |
| 27618447    | 2016 | DBP, SBP, PP, HT | Up to 165,276 European ancestry individuals, up to 27,487 South Asian ancestry individuals | Up to 125,713 European ancestry individuals, up to 2,641 South Asian ancestry individuals, 4,632 Hispanic individuals, 22,077 African American individuals | They identified 30 new blood pressure- or hypertension-associated loci in the general population, including 3 rare missense variants in RBM47, COL21A1 and RRAS with larger effects than common variants. |
| 27618448    | 2016 | DBP, SBP, PP, HT | 120,473 European ancestry individuals, 21,503 African American individuals, 4,586 Hispanic individuals | 154,543 European ancestry individuals, 26,183 South Asian ancestry individuals | They identified 31 new blood pressure-associated loci.                                                                                                   |
| 27618452    | 2016 | DBP, SBP      | Up to 201,529 European ancestry individuals | Up to 140,886 European ancestry individuals | They identified 66 blood pressure-associated loci, of which 17 were new; 19 harbored multiple distinct association signals.                              |
| 27841878    | 2016 | DBP, SBP      | 295,529 European ancestry individuals, 8,231 Latino individuals, 3,058 African American individuals, 2,029 African British individuals, 7,701 East Asian ancestry individuals, 2,735 South Asian ancestry individuals, 1,979 mixed and unknown ancestry individuals | NA | They identified 39 new loci among 75 genome-wide significant loci.                                                                                  |
| 28135244    | 2017 | DBP, SBP, PP  | 140,886 European ancestry individuals | 190,318 European ancestry individuals | They identified 107 blood pressure-associated loci.                                                                                                    |
| 29403010    | 2018 | DBP, SBP      | 136,615 Japanese ancestry individuals | NA | They identified 1,407 trait-associated loci, 679 of which were novel.                                                                                  |
| 30224653    | 2018 | DBP, SBP      | 757,601 European ancestry individuals | 249,262 European ancestry individuals | They identify 535 novel blood pressure loci.                                                                                                           |
| 30578418    | 2019 | DBP, SBP      | 365,998 European ancestry individuals, 63,490 African ancestry individuals, 22,802 Hispanic individuals, 4,792 Asian ancestry individuals, 2,695 Native American ancestry individuals | 299,024 European ancestry individuals, 17,277 individuals | They discovered 208 novel common blood pressure SNPs and 53 rare variants.                                                                               |

BP, blood pressure; DBP, diastolic blood pressure; SBP, systolic blood pressure; PP, pulmonary pressure; HT, hypertension.
hundreds of novel blood pressure loci that offer new biological insights into blood pressure regulation. Also, trans-ethnic genome-wide association study of blood pressure in up to 776,078 participants from the Million Veteran Program (MVP) and collaborating studies discovered 208 novel common blood pressure SNPs and 53 rare variants (Giri et al., 2019). Table 1 listed all high quality GWASs for hypertension or blood pressure published in NATURE GENETICS since 2007.

**GENE–ENVIRONMENT INTERPLAY**

Thirteen years GWAS for hypertension, from the beginning “failure” to the recent “success,” can we say GWASs have deciphered the genetic architecture of hypertension? The answer is obviously NO. This is because of the fact that environmental factors and gene-environment interactions are likely major contributors to the development of hypertension, however, they were largely ignored in present GWASs (Cooper, 2018).

As widely accepted, hypertension is a consequence of significant interaction between genetic and environmental factors. Here, the meaning of “environment” is extensive, including intrauterine, postnatal and evolutionary environments. It has long been known that intrauterine environmental factors (e.g., maternal nutritional perturbation, toxin exposure, and stress during pregnancy) may result in hypertension in adult life. Recent epidemiological and experimental studies further indicated that paternal environmental factors, before conception and during sperm development, were also linked to the development of hypertension in later life (Li et al., 2016). On the other hand, classic epidemiological studies have addressed many postnatal environmental risk factors for hypertension, including living environments (cold temperature, air pollution, and toxins), life styles (lack of physical activity, psychological stress, smoking, alcohol abuse, drug use, improper nutrition, excessive salt intake, and obesity), and other demographic differences in age, gender, race, socioeconomic status, etc. From an evolutionary perspective, hypertension can be viewed as a maladaptation disease caused by the discrepancy between today’s lifestyles and ancient adaptive genotypes. Physiologically, many risk factors for hypertension, such as enhanced salt and water avidity and vascular contractility, were adaptive traits associated with salt scarcity in the hot and humid climate of the ancestral African environment. As humans migrated out of Africa to cooler environments, the originally selected genes became maladaptive for new environments and turned into risk factors for hypertension (Young et al., 2005; Ji et al., 2016).

**DISCUSSION**

Due to the complexity of essential hypertension, previous GWAS didn’t find many robust variants. If more related factors are considered, and method like phenomics is used, future studies may help us to deeply understand this complex disease. As emphasized by Jeremy Berg, editor-in-chief of SCIENCE, genes alone do not determine our futures—environmental factors and chance also play important roles (Berg, 2016). Indeed, 13 years GWASs for hypertension really made a big step forward, however, if we leave behind another important factor, the environment, can also impede our progress. Incredibly, the environmental effects were ignored in almost all published GWASs of blood pressure traits or hypertension. Some studies were conducted in well-established cohorts like Global BPGen and CHARGE, and many environmental data should be available. Although analyzing the gene-environment interaction is still a great challenge, GWASs without consideration of environmental effects will definitely not go to address the key question regarding the genetic architecture of hypertension.

**AUTHOR CONTRIBUTIONS**

L-dJ and JX conceived the opinion. L-dJ, NT, Z-fX, and JX wrote the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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