Polymorphisms in the WNK1 Gene Are Associated with Blood Pressure Variation and Urinary Potassium Excretion

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Abstract

WNK1 - a serine/threonine kinase involved in electrolyte homeostasis and blood pressure (BP) control - is an excellent candidate gene for essential hypertension (EH). We and others have previously reported association between WNK1 and BP variation. Using tag SNPs (tSNPs) that capture 100% of common WNK1 variation in HapMap, we aimed to replicate our findings with BP and to test for association with phenotypes relating to WNK1 function in the British Genetics of Hypertension (BRIGHT) study case-control resource (1700 hypertensive cases and 1700 normotensive controls). We found multiple variants to be associated with systolic blood pressure, SBP (7/28 tSNPs min-p = 0.0005), diastolic blood pressure, DBP (7/28 tSNPs min-p = 0.0002) and 24 hour urinary potassium excretion (10/28 tSNPs min-p = 0.0004). Associations with SBP and urine potassium remained significant after correction for multiple testing (p = 0.02 and p = 0.01 respectively). The major allele (A) of rs765250, located in intron 1, demonstrated the strongest evidence for association with SBP, effect size 3.14 mmHg (95% CI: 2.3–4.9), DBP 1.9 mmHg (95% CI: 0.7–3.2) and hypertension, odds ratio (OR: 1.3 [95% CI: 1.0–1.7]). We genotyped this variant in six independent populations (n = 14,451) and replicated the association between rs765250 and SBP in a meta-analysis (p = 7 × 10^-13, combined with BRIGHT data-set p = 2 × 10^-14, n = 17,851). The associations of WNK1 with DBP and EH were not confirmed. Haplotype analysis revealed striking associations with hypertension and BP variation (global permutation p < 10^-10). We identified several common haplotypes to be associated with increased BP and multiple low frequency haplotypes significantly associated with lower BP (>10 mmHg reduction) and risk for hypertension (OR=0.60). Our data indicates that multiple rare and common WNK1 variants contribute to BP variation and hypertension, and provide compelling evidence to initiate further genetic and functional studies to explore the role of WNK1 in BP regulation and EH.

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Introduction

Essential hypertension (EH), or high blood pressure (BP) \( \geq 140/90 \) mmHg, is a major public health problem that contributes to millions of deaths worldwide every year due to coronary heart disease, stroke, and other vascular diseases [1,2,3,4]. The disorder results from the complex interaction between multiple genes and environmental factors [5,6,7]. A major goal for hypertension research has been to identify the genes and mechanisms underlying the disorder in order to improve the prediction of those at risk and develop better anti-hypertensive treatments. Some advances in hypertension genetics have been made through studies of rare Mendelian forms of hypertension, which have identified strong candidate genes for BP regulation and EH [8]. Mutations in \( \text{WNK1} \) [with No K-lysine kinase 1 [9,10], MIM 605232], cause Pseudohypoaldosteronism type 2 (PHA2, MIM 145260) – a rare autosomal dominant disorder primarily characterised by early onset hypertension and hyperkalemia [8,11]. In PHA2 patients, gain-of-expression mutations in \( \text{WNK1} \) cause hypertension. Conversely, heterozygous knock-out mice that lack \( \text{WNK1} \) expression have low BP, consistent with a gene-dosage effect of \( \text{WNK1} \) on BP [11,12].

\( \text{WNK1} \), a serine-threonine kinase regulating numerous ion channels involved in sodium and potassium transport [13,14,15,16,17], is ubiquitously expressed, with particularly high levels of expression in the kidney and cardiovascular system. \( \text{WNK1} \) maps to chromosome 12p13.3, spans \( \sim 156 \) Kb of genomic DNA and encodes 29 exons [10,18]. There are two major isoforms of \( \text{WNK1} \), a kinase-active long isoform (\( L-\text{WNK1} \)) and a kinase-deficient kidney specific short isoform (\( Ks-\text{WNK1} \)). These two isoforms are under the control of alternative promoters located 5’ of exon 1 for \( L-\text{WNK1} \), and in intron 4 for \( Ks-\text{WNK1} \) [19]. It is thought that hypertension in PHA2 patients may partially be the result of increased sodium reabsorption via \( L-\text{WNK1}/Ks-\text{WNK1} \) mediated up-regulation of the thiazide sensitive sodium chloride cotransporter (\( \text{SLC12A3} \) or \( \text{NCCT} \)) and the renal amiloride-sensitive epithelial sodium channel (\( \text{ENaC} \), encoded by three genes: \( \text{SCNN1A}, \text{B} \) and \( \text{G} \)), and hyperkalemia by increased inhibition of the renal outer medullary potassium channel (\( \text{KCNJ1} \)) [13,14,15,16,17]. These findings together with the discovery that \( \text{WNK1} \)-deficient mice have low BP have highlighted the functional importance of \( \text{WNK1} \) in ion transport and BP regulation [12].

We and others have previously reported association between common variants in \( \text{WNK1} \) and human BP variation in adults. Using a tag SNP approach, association was found with a variant near the promoter and severity of hypertension in families from the British Genetics of Hypertension (BRIGHT) Study [20]. Furthermore, common \( \text{WNK1} \) variants were found to be associated with ambulatory BP in families representative of the general population [21]. Turner et. al. (2005) have also reported association between common \( \text{WNK1} \) variants and response to thiazide diuretics [22]. These studies prompted interest in further genetic studies exploring the role of \( \text{WNK1} \) in BP regulation.

Since publication of these studies, additional SNP data have become available from the HapMap Project (HapMap) [23]. Taking advantage of this gain in genetic information, our aim was to extend and replicate our findings between \( \text{WNK1} \) and BP variation and to further localise causative SNPs/regions within the gene using the British Genetics of Hypertension (BRIGHT) study case-control study, which provides substantial power for detecting susceptibility loci with moderate risks for disease.

Methods

MRC BRIGHT study case-control study

As part of the MRC BRIGHT study (http://www.brightstudy.ac.uk) hypertensive cases and normotensive controls of white European ancestry have been recruited for association testing. Case ascertainment and phenotyping has been described previously [24]. Briefly, cases have BP readings \( \geq 150/100 \) mmHg based on one reading or \( \geq 145/95 \) mmHg based on the mean of three readings and there is extensive phenotyping information from all individuals, including anthropometric data, plasma and urinary electrolytes, and diagnosis BP recordings. Further details of recruitment and phenotyping can be found online at www.brightstudy.ac.uk. Healthy, age and sex matched normotensive controls (BP readings \( \leq 140/90 \) mmHg) had similar phenotyping, with the exception that plasma and urinary electrolytes were not measured. All subjects participated as volunteers and were recruited via hypertension registers from the MRC General Practice Framework in the UK. Ethics Committee approval was obtained from the multi- and local research committees of the partner institutes, and all participants gave written informed consent.

The Estonian HYPEST sample collection

The Estonian participants were recruited during 2004–2007 across the entire country in the framework of the HYPEST sample collection (\( n = 1,823 \)) targeting hypertension risk factors in the Estonian population [permissions no 122/13, 22.12.2003; 137/20, 25.04.2005 by Ethics Committee on Human Research of University of Tartu, Estonia]. Hypertensive patients were recruited at the North Estonia Medical Centre, Tartu Estonia. Healthy (exclusion criteria: cardiovascular disease, diabetes, and antihypertensive treatment), normotensive individuals were recruited across the whole country. The majority of the HYPEST participants (\( n = 1,482 \)) possess a documented history of multiple systolic blood pressure (SBP) and diastolic blood pressure (DBP) readings. For this study we defined cases (\( n = 596 \)) as individuals with either blood pressure readings \( \geq 160/100 \) mmHg based on the median of several measurements or under antihypertensive therapy. Controls (\( n = 650 \)) were defined as having median blood pressure readings below 140/90 mm Hg. The quantitative association analysis of SBP and DBP (\( n = 1,284 \)) included both untreated (\( n = 881 \)) and treated individuals (\( n = 403 \)).

London Life Sciences Prospective Cohort Study (LOLIPOP)

This is a prospective study of 18,829 subjects (UK-based Indians, \( n = 12823 \) and white Europeans, \( n = 6006 \)) investigating cardiovascular risk factors in certain sub-populations. For this study we selected white European individuals; 485 cases and 458 controls, drawn from the top and bottom 10% of the BP distribution. All blood pressure readings are off-medication.

The Whitehall I study

The re-survey of the Whitehall study has DNA and blood pressure measurements recorded in middle age (1967–1970) and in old age (1997) on 5360 men. For this study, we selected white European men; 466 hypertensives and 536 controls, drawn from the top and bottom 10% of the BP distribution. The design and methods of Whitehall I as well as the characteristics of the participants have been described in detail elsewhere [25].
The Olivetti Heart Study

The Olivetti Heart Study population is derived from the male workforce of the Olivetti factories of Pozzona (Naples) and Marcianise (Caserta), Italy. The general characteristics of the study and its methodological procedures have been previously described [26,27]. The local ethics committee approved the study protocol, and informed consent was obtained from all participants. A total of 1085 individuals aged 25–74 years were examined in 1994–95: of these, 907 (85.6%) were seen again in 2002–04, of these 868 individuals had DNA available for genotyping at both time-points[26]. We analysed individuals at the 2002–04 time-point, cases were defined as individuals with BP readings of ≥145/95 mmHg, normotensive controls had BP readings of ≤140/90 mmHg.

Whitehall II study

After ethical clearance the Whitehall study enrolled 10,308 subjects (3413 women) aged 35–55 working in the London offices of 20 civil service departments between 1985–1988. In this longitudinal study blood pressure was recorded at phase 1 (1985–1988), phase 3 (1991–1993), phase 5 (1997–1999) and phase 7 (2003–2004). DNA was stored from phase 7 of the study. For association testing with blood pressure and hypertension we selected individuals from Phase 5 as diabetes ascertainment and blood pressure medication records were most complete from this phase. For case control analyses, hypertensives were selected using blood pressure medication records were most complete from this phase. For case control analyses, hypertensives were selected using blood pressure recordings of ≥145/95 mm Hg, or individuals taking anti-hypertensive medication or a physician diagnosis of hypertension. Normotensive controls were selected using blood pressure recordings ≤140/90 mmHg and not taking any anti-hypertensive medications.

English Longitudinal Study of Ageing (ELSA)

After ethical approval the participants were drawn from around 12,000 respondents to the Health Survey for England (HSE) over three separate years (1998, 1999 and 2001) to provide a representative sample of the English population aged 50 and over. Each individual had a mean of three blood pressure measures taken when the participant was seated and antihypertensive medications were recorded, DNA was extracted from 5672 participants in wave 2 (2004). For association testing with blood pressure and hypertension we selected individuals from wave 2. Cases and normotensive controls were defined using the same criteria as the Whitehall II study.

Tag SNP selection

The HapMap database was used to identify common polymorphisms across the WNK1 genomic region [23]. Tag SNP selection was performed using Tagger (http://www.broad.mit.edu/mpg/tagger) [28]. WNK1 SNP data were obtained from HapMap (Data Ref 21/phased Jul06, NCBI 35, dbsNP 125) using the WNK1 chromosomal coordinates, chr12:732 992–883 219±10 kb, and the data used as input for Tagger. The original 8 tSNPs[20] and one additional SNP (rs3088353) located in the L-WNK1 promoter were forced in as tSNPs prior to the search. Additional tSNPs were selected to capture 100% of variation in HapMap with minor allele frequencies >0.05 and minimum R² of 0.8.

Genotyping

All SNP genotyping was performed using the Taqman assay developed by Applied Biosystems, followed by allelic discrimination using the ABI PRISM 7900HT Sequence Detection System and software (SDSv2.0, Applied Biosystems). Specific WNK1 SNP Taqman probes and primers were obtained from Applied Biosystems Assay-by-DesignTM Service for SNP genotyping. Genotyping of individuals from the Whitehall study was performed at the Centre National de Genotypage, France, using the Taqman assay. The HYPEST sample was genotyped using Taqman, individuals from Whitehall II and the ELSA study were also genotyped using the Taqman assay at Geneservice UK.

Statistical analysis of the BRIGHT study

Hardy–Weinberg equilibrium (HWE) was assessed by a χ² test. SNPs were dropped from all analyses if they were significantly out of HWE, rs11064519 HWE- p = 0.0001 and rs4980973 HWE- p = 0.0002 in control samples). The [D’] and r² measures of linkage disequilibrium were calculated using the program Haploview v4 [29].

All tests for association were performed with the statistical package R (http://www.r-project.org/) [30]. Logistic regression was used to test for association with EH and linear models for quantitative analyses. All analyses were adjusted for age, sex, body mass index (BMI) and centre of recruitment to allow for population stratification. The quantitative phenotypes were non-normally distributed and were analysed using naturally logged transformed data. Wald tests from a linear regression can be affected by non-normality and lead to inaccurate estimates of the variance-covariance matrix. Therefore, we also used an empirical estimate of the standard errors from 10,000 bootstrap samples to calculate the Wald tests. This approach selects random samples of size n (n = population size) with replacement from the original data, and repeats the sampling procedure a large number of times to provide information on the bias and variability of the parameter estimates. For each test of association, we repeated the selection procedure on 10,000 random samples and report the bootstrap estimate (effect) and 95% confidence intervals.

For single tSNP analyses, we tested for association under additive, dominant and recessive models. To control for multiple testing, global p-values were estimated using 10,000 permutations of the trait label to calculate the empirical distribution of the minimum p-value observed across all models for all SNPs. Multiple testing adjustments were applied separately to each trait, therefore only controlling for multiple testing within each trait. We only present results for tSNPs and haplotypes showing evidence for association after permutation testing with p < 0.05.

Haplo-type associations were explored using the R package HaploStats [31]. HaploStats estimates haplotype frequencies via the expectation-maximization (EM) algorithm and computes global and haplotype-specific score statistics for tests of association between a trait and haplotypes weighted by their posterior probabilities. Permutation p-values were computed in HaploStats using 10,000 permutations for 24 hour urine potassium excretion. For EH, SBP and DBP 10,000,000 permutations were used. In all haplotype analyses, we considered only those haplotypes with frequencies greater than or equal to 5% as separate independent variables, low frequency haplotypes with frequencies <5% were combined into a single group. To further explore the specific effects for individual low frequency haplotypes, we repeated the analysis to include haplotypes with frequencies ≥0.001. Haplo-type specific Odds ratios (OR) and effect sizes were estimated using logistic regression for association with EH and linear models with the quantitative phenotypes in R weighted by their posterior probability [30,31]. The most frequent haplotype was used as the baseline haplotype, with which effects of the other haplotypes were contrasted.
For the quantitative analyses with diagnosis BP (off-medication and therefore not confounded by the effects of drug treatment), data were available on 1183 cases and 1700 controls. These blood pressure readings were obtained from GP records at the time of their diagnosis. Mean BP levels from three readings for the 1183 cases and 1700 controls were used. For the biochemical phenotypes, analyses were performed in cases only, and these were on medication measures. No biochemical data were available for the control population. We did not have acidified urine for measuring urinary calcium or measures of serum potassium in the BRIGHT cases.

Statistical analyses in replication cohorts and meta analyses

For quantitative association analysis of SBP and DBP, we corrected BP values for the effects of BP lowering therapies (where appropriate) using the method described by Tobin and colleagues [32,33]. For individuals on antihypertensive medication, we adjusted systolic and diastolic blood pressure measures by adding 15 mm Hg to systolic and 10 mm Hg to diastolic readings.

We then tested for association with BP variation and EH in each cohort separately using 10,000 bootstrap samples. To avoid multiple testing issues we analysed the data under a dominant model with the prior hypothesis that carriers of rs765250 [A] (A/A+A/G versus G/G) would have higher BP levels and be at increased risk for EH compared to G/G homozygotes, as determined from our primary analysis in the BRIGHT study case-control resource. As we are testing a specific hypothesis we report one-tailed p-values for the replication cohorts. We declare replication if the association is in the same direction with one-tailed p<0.05. Results were combined in a meta-analysis under a fixed effect model in R. Tests of heterogeneity were also evaluated using the I² statistic [34]. If there was any statistical evidence for heterogeneity (p<0.05) then the analysis was repeated using a random-effects model, which includes a measure of variance between studies.

Results

**WNK1 variation**

Twenty eight tagging SNPs (tSNPs) were identified that capture 100% of the common WNK1 variation in HapMap (mean r² = 0.9,

### Table 1. WNK 1 single nucleotide polymorphisms.

| tSNP   | gene position | Chr12 nt position (bp) | Alleles | MAFb | HWE p Controls |
|--------|---------------|------------------------|---------|------|----------------|
| 1      | rs7137188     | 5'                     | 724592  | C/T  | 0.48           | 0.1 |
| 2      | rs1468326     | 5'                     | 727762  | C/A  | 0.11           | 0.8 |
| 3      | rs3088353     | 5'                     | 732901  | A/C  | 0.46           | 0.9 |
| 4      | rs11064519    | i1                     | 741472  | C/G  | 0.34           | 0.0001 |
| 5      | rs2369402     | i1                     | 748925  | G/A  | 0.21           | 0.1 |
| 6      | rs2107612     | i1                     | 758581  | A/G  | 0.27           | 0.7 |
| 7      | rs2107613     | i1                     | 758689  | C/T  | 0.23           | 0.4 |
| 8      | rs11608756    | i1                     | 760094  | A/G  | 0.39           | 0.04 |
| 9      | rs11064524    | i1                     | 760163  | G/T  | 0.25           | 0.8 |
| 10     | rs11064527    | i1                     | 761286  | A/C  | 0.16           | 0.4 |
| 11     | rs11064536    | i1                     | 775843  | T/C  | 0.17           | 0.5 |
| 12     | rs765250      | i1                     | 778544  | A/G  | 0.31           | 0.5 |
| 13     | rs12314329    | i1                     | 779992  | A/G  | 0.08           | 0.4 |
| 14     | rs11161124    | i4                     | 809741  | G/T  | 0.22           | 0.3 |
| 15     | rs6489756     | i4                     | 834767  | A/G  | 0.47           | 0.7 |
| 16     | rs12816718    | i6                     | 840061  | G/T  | 0.15           | 0.6 |
| 17     | rs2286007     | ex8                    | 841552  | C/T  | 0.07           | 0.7 |
| 18     | rs11611231    | i8                     | 844587  | C/G  | 0.10           | 0.4 |
| 19     | rs4980973     | i9                     | 853307  | A/G  | 0.13           | 0.0002 |
| 20     | rs4980974     | i9                     | 855289  | A/G  | 0.44           | 0.6 |
| 21     | rs880054      | i10                    | 858819  | A/G  | 0.41           | 1 |
| 22     | rs956868      | ex13                   | 861173  | A/C  | 0.14           | 0.4 |
| 23     | rs953361      | i22                    | 872068  | C/T  | 0.40           | 0.7 |
| 24     | rs2301880     | i23                    | 874098  | C/T  | 0.24           | 0.7 |
| 25     | rs2286026     | i26                    | 885434  | A/G  | 0.41           | 0.9 |
| 26     | rs2286028     | i26                    | 885730  | C/G  | 0.20           | 0.7 |
| 27     | rs2277869     | i26                    | 887171  | C/T  | 0.16           | 1 |
| 28     | rs11571461    | 3'                     | 896580  | A/G  | 0.07           | 0.3 |

a: intron, e: exon, 5’: 5 prime; 3’: 3 prime.
bMinor allele frequency.
doi:10.1371/journal.pone.0005003.t001
The average rate of success for each genotyped SNP was 95%. After genotyping, two SNPs (tSNP4 rs11064519 and tSNP19 rs4980973) were dropped from further analysis as they were significantly out of HWE (see Table 1). After exclusion of these tSNPs we were able to tag 97% of all common WNK1 variation with $r^2 > 0.8$ and 100% with $r^2 > 0.5$.

**Figure 1. Association results and WNK1 linkage disequilibrium.** The diagram shows the summary results from the association analyses between WNK1 tSNPs and essential hypertension (EH – red diamonds), blood pressure variation (SBP – blue squares, DBP – green circles) and urine potassium excretion (UrK – orange diamonds). The $-\log(10)$ of the best p-value are plotted on the y-axis against the physical position or each genotyped tSNP (x-axis), denoted by their dbSNP identification. Dotted lines represent P-value thresholds. Closed symbols represent significant associations ($p < 0.05$). The tick marks along the x-axis also show the physical position of all common HapMap. The middle panel shows the genomic structure of the human WNK1 gene and all known common variation across the WNK1 genomic region. Exons are indicated by the vertical black bars and alternatively spliced exons by the blue boxes. The green boxes indicate the position of the PHA2 disease causing deletions. The lower panel represents the extent of linkage disequilibrium as measured by Lewontin’s $|D'|$ across the WNK1 genomic region. $|D'|$ varies between 0 (no disequilibrium) and 1 (maximum disequilibrium), represented by shades of white to yellow to red. White: $|D'| = 0$ and red: $|D'| = 1$. Strong LD ($|D'|$) exists between the most widely separated tSNPs, defining a single large haplotype block extending from tSNP 3 (rs3088353) located in the WNK1 promoter, to tSNP 28 (rs11571461) located 3’ of WNK1. The plot was produced using a modified version of snp.plotter [49].

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range 0.8–1.0, Table 1, Figure 1).
Common *WNK1* variants are associated with blood pressure variation and essential hypertension in the BRIGHT study

We tested for association between 26 tSNPs with BP variation and EH in 1700 hypertensive cases and 1700 normotensive controls from the BRIGHT study. The demographic and biochemical characteristics of these individuals are shown in Table 2. For single SNP analyses, we tested for association under additive, dominant and recessive models and corrected for multiple testing using permutation methods. Diagnosis BP measurements were used in the quantitative analyses, as these readings are off-medication and therefore not confounded by the effects of anti-hypertensive treatments.

Multiple SNPs (7/26 tSNPs) located in the *WNK1* promoter and intron 1, were found to be significantly associated with BP variation under one or more of the models tested (Table 3 and Figure 1). Among the 7 BP associated SNPs, tSNP12, rs765250, located in intron 1, shows the strongest evidence for association; carriers of rs765250 allele A (A/A+A/G versus G/G) are associated with increased SBP (+3.1 mmHg, 95%CI:1.3,4.9, p = 0.0005) and DBP (+1.9 mmHg, 95%CI:0.7,3.2, p = 0.002). This same variant demonstrated evidence for association with EH, (OR:1.34, [95%CI:1.0–1.7], p = 0.015). The association with SBP remained significant after permutation testing under the global null hypothesis “no SNP is associated with the trait of interest under any model”, but not with DBP or EH (SBP global permutation p = 0.02, DBP p = 0.06, EH p = 0.4).

Replication of rs765250 association with blood pressure and essential hypertension

The tSNP rs765250 demonstrated the strongest evidence of association with BP variation and EH. We genotyped this SNP in six additional populations totalling 14,451 individuals (HYPEST, Whitehall I, LOLIPOP, English longitudinal study of ageing (ELSA), Whitehall II and the Olivetti study) to confirm our findings. Table 4 shows selective demographic characteristics of each cohort. We tested for association with BP as a continuous trait and hypertension in all cohorts under a dominant model, with the prior hypothesis that carriers of rs765250 allele A (A/A+A/G versus G/G) would have higher BP levels and be at increased risk for EH compared to G/G homozygotes – as determined from our primary analysis in the BRIGHT study. The results for each trait are presented in Table 5 and Figure 2.

Meta-analysis across all the replication cohorts (n = 14,451) revealed significant independent association with SBP (p = 0.007). There was a non-significant trend for association with DBP (p = 0.06). No association was seen with EH (OR:1.08, [95%CI:0.9,1.2], p = 0.33). To determine the overall significance and effect size in the samples studied, a meta-analysis was performed combining all cohorts including the original study population BRIGHT (n = 17,851). In the combined analysis, we confirmed association between rs765250 and SBP; carriers of rs765250 [A] were associated with an overall increase in SBP, with a mean effect on natural log transformed SBP of +0.01, [95%CI:0.006, 0.02], p = 2×10⁻⁴, equivalent to ~2.0 mmHg. There was an association with increased risk for EH, OR:1.1 [95%CI:1.0,1.3], p = 0.04, and a trend for association with DBP p = 0.06.

*WNK1* variants are associated with urinary potassium secretion in the BRIGHT study

As disturbances in electrolyte homeostasis accompany the development of high blood pressure in PHA2 patients carrying *WNK1* mutations, we hypothesised that variation in serum
and/or urinary electrolytes would be indicative of altered \( \text{WNK1} \) function or expression, and this may help to identify polymorphisms that also affect BP. Therefore, we also explored associations with phenotypes related to \( \text{WNK1} \) function in the BRIGHT cases; measures of serum and urine biochemistry were not available for the BRIGHT controls. The biochemical characteristics of the hypertensive cases are presented in Table 2. We found multiple tSNPs spanning the length of the gene to be associated with variation in 24 hour urine K\(^+\) excretion (Table 3). These associations remained significant after correction for multiple testing (\( p = 0.01 \)). The majority of haplotypes in the BRIGHT case-control resource (Table 6); haplotypes with frequencies (f) of 5\% that account for 79\% of all \( \text{WNK1} \) variation. There are 8 common \( \text{WNK1} \) haplotypes were constructed using 24/26 tSNPs that fall within this block (2 tSNPs were dropped as they were out of HWE – see Figure 1).

### Table 3. Single tSNP association analyses with EH, BP variation and 24 hour urinary potassium in BRIGHT.

| Trait                  | tSNP   | SNP ID       | gene position\(^a\) Major\(^b\) | Minor\(^b\) | RAF\(^c\) | Model\(^d\) | Effect\(^e\) | 95\% CI \(^{f}\) | perm-p\(^g\) | global-p\(^h\) |
|------------------------|--------|--------------|-------------------------------|------------|----------|-----------|-----------|----------------|---------|-------------|
| EH (Odds ratio)        | 12     | rs765250     | i1                            | A          | G        | Dom       | 1.3       | 1.0,1.7       | 0.01    | 0.4         |
| SBP (mmHg)             | 2      | rs1468326    | 5’                            | C          | A        | Rec       | 5.1       | 0.6,9.2       | 0.02    |             |
|                        | 3      | rs3088353    | 5’                            | A          | C        | Rec       | 1.6       | 0.2,2.8       | 0.02    |             |
|                        | 5      | rs2369402    | i1                            | G          | A        | Dom       | 1.2       | 0.1,2.2       | 0.03    |             |
|                        | 6      | rs2107612    | i1                            | A          | G        | Add       | 1.2       | 0.2,2.0       | 0.008   |             |
|                        | 7      | rs2107613    | i1                            | C          | T        | Dom       | 1.2       | 0.1,2.3       | 0.03    |             |
|                        | 11     | rs11064536   | i1                            | T          | C        | Dom       | 3.1       | 0.6,1         | 0.05    |             |
|                        | 12     | rs765250     | i1                            | A          | G        | Dom       | 3.1       | 1.3,4.9       | 0.0005  | 0.02        |
| DBP (mmHg)             | 2      | rs1468326    | 5’                            | C          | A        | Rec       | 3.1       | 0.01,5.8      | 0.04    |             |
|                        | 3      | rs3088353    | 5’                            | A          | C        | Rec       | 0.9       | 0.1,1.8       | 0.03    |             |
|                        | 5      | rs2369402    | i1                            | G          | A        | Dom       | 0.8       | 0.15          | 0.04    |             |
|                        | 6      | rs2107612    | i1                            | A          | G        | Add       | 0.6       | 0.01,1.2      | 0.05    |             |
|                        | 7      | rs2107613    | i1                            | C          | T        | Dom       | 0.7       | 0.06,1.4      | 0.06    |             |
|                        | 11     | rs11064536   | i1                            | T          | C        | Dom       | 2.5       | 0.2,4.8       | 0.02    |             |
|                        | 12     | rs765250     | i1                            | A          | G        | Dom       | 1.9       | 0.7,3.2       | 0.002   | 0.06        |
| 24 hour urine K\(^+\) (mmol/24 hour) | 3     | rs3088353    | 5’                            | A          | C        | Dom       | −5.9      | −2.5, −9.2    | 0.0004  |             |
|                        | 6      | rs2107612    | i1                            | A          | G        | Add       | −2.6      | −4.8, −0.3    | 0.03    |             |
|                        | 8      | rs11608756   | i1                            | G          | A        | Add       | −2.3      | −4.3, −1      | 0.04    |             |
|                        | 9      | rs11064524   | i1                            | T          | G        | Dom       | −7.3      | −12.1, −2.2   | 0.008   |             |
|                        | 12     | rs765250     | i1                            | A          | G        | Add       | −3.1      | −5.4, −0.8    | 0.009   |             |
|                        | 13     | rs12314329   | i1                            | A          | G        | Add       | −4.3      | −7.8, −0.5    | 0.03    |             |
|                        | 14     | rs11611246   | i4                            | G          | T        | Dom       | −2.9      | −5.3, −0.4    | 0.02    |             |
|                        | 15     | rs6489756    | i4                            | A          | G        | Add       | −3.2      | −5.6, −0.9    | 0.005   |             |
|                        | 27     | rs2277969    | i26                           | T          | C        | Add       | −4.5      | −7.6, −1.5    | 0.004   |             |
|                        | 28     | rs11571461   | 3’                            | A          | G        | Add       | −4.7      | −8.7, −0.5    | 0.03    | 0.01        |

\(^{a}\): intron; 5\': 5' prime; 3\': 3' prime.

\(^{b}\): Indicates major or minor alleles, bold indicates the allele increasing blood pressure.

\(^{c}\): RAF: Risk Allele Frequency – refers to the allele increasing blood pressure or decreasing urinary potassium.

\(^{d}\): Best Model; Add: Additive, Dom: Dominant, Rec: Recessive.

\(^{e}\): Linear effect estimate, 95\% confidence intervals for EH, diagnosis SBP, DBP and 24 hour urine potassium using 10 K bootstrap samples. P-values are based on 10,000 permutations.

\(^{f}\): Global p-value based on 10,000 permutations - adjusting for testing multiple SNPs and multiple models.

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Haplotype analysis reveals striking associations with essential hypertension, blood pressure variation and urine potassium excretion

Using \( D’ \) as a measure of linkage disequilibrium (LD) across the \( \text{WNK1} \) region, we identified a single large haplotype block extending from rs3088353 (tSNP9) located in the \( \text{WNK1} \) promoter, to rs11571461 (tSNP26) located 3' of \( \text{WNK1} \) (Figure 1). \( \text{WNK1} \) haplotypes were constructed using 24/26 tSNPs that fall within this block (2 tSNPs were dropped as they were out of HWE – see \( \text{WNK1} \) variation). There are 8 common \( \text{WNK1} \) haplotypes with frequencies (f) > 5\% that account for ~79\% of all haplotypes in the BRIGHT case-control resource (Table 6); collectively the low frequency haplotypes account for the remaining ~21\% of haplotypic variation.

Table 7 shows the results of the haplotype analysis, it reveals striking associations with EH (global permutation \( p < 10^{-7} \)), BP variation (global permutation \( p < 10^{-7} \)), and 24 hour urine potassium excretion (global permutation \( p = 10^{-5} \)). Across the
BP related traits, the most significant associations were seen with the low frequency haplotype pool, which drive the highly significant global associations. The pool of low frequency haplotypes were found to be associated with decreased risk for EH, OR:0.6, [95%CI:0.5, 0.74], p = 4.82 × 10⁻⁶. For the replication cohorts and meta analyses one-tailed p-values are reported.

To further explore the specific effects for individual low frequency haplotypes on BP variation and EH, we repeated the analysis to include haplotypes with frequencies ≥0.001 (minimum haplotype count of 5, Table 7, and Table S1). We identified 53 haplotypes with frequencies ≥0.001 (8 common haplotypes with f ≥0.05, and 45 low frequency haplotypes with 0.001 ≤f ≤0.05). Analysis found 4 low frequency BP lowering haplotypes to drive most of the haplotype associations (haplotypes 14, 15, 18 and 19). Interestingly, these haplotypes were nearly unique to the control population, and as a result are associated with relatively large BP lowering effects when compared to the most common haplotype (>10 mmHg, Table 7, and Figure S1). For example, haplotype 14 was observed in at least 49 controls versus 1 case. These figures are counts of “phase-very-certain” haplotypes with posterior probabilities ≥0.9, and differ from those in Table 7, which presents counts of all haplotypes weighted by their posterior probabilities, thus taking into account haplotype phase uncertainty. This low frequency haplotype (14), accounts for ~1% of all haplotypes in BRIGHT, and was associated with low BP and decreased odds for hypertension (effect per copy of haplotype 14, OR: 0.03, [95%CI:0.006,0.1], p = 6.9×10⁻⁶, SBP lowering effects when compared to the most common haplotype 10 mmHg, Table 7, and Figure S1). For example, haplotype 14 was observed in at least 49 controls versus 1 case. These figures are counts of “phase-very-certain” haplotypes with posterior probabilities ≥0.9, and differ from those in Table 7, which presents counts of all haplotypes weighted by their posterior probabilities, thus taking into account haplotype phase uncertainty. 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that treatment will confound the reported associations and may lead to WNK1
the role of significantly associated with BP, EH and urinary potassium
also associated with decreased in urinary K
strongest evidence for association with EH and BP variation, is
variant rs765250, located in intron 1, which demonstrated the
association with K^+ homoeostasis remains to be confirmed
In the BRIGHT study resource, the strongest tSNP association
was seen between rs765250 and SBP. This association was
replicated in additional populations, suggesting that the original
association was likely to be a false positive. This association, however,
was not replicated in all cohorts tested. Failure to replicate the effect of rs765250 on EH and BP in every population could be due to genetic heterogeneity across populations, small effect sizes or low power. Although the associations did not reach statistical significance in some of the individual cohorts, for the majority of populations, the direction of the effect was consistent with that seen in BRIGHT, with overlapping 95% confidence intervals. Encouragingly, in the combined meta-analyses, the evidence for association with SBP increased. Notably, this same SNP has previously been associated with ambulatory SBP and DBP in the families from the GRAPHIC study [min p = 0.001] [21], and more recently, with DBP gradient (p = 0.02) in children from the Avon Longitudinal Study of Parent and Children Study [36], lending further support to the reported findings.

In the single tSNP analyses, our primary associations with BP variation and EH were observed with variants located in the WNK1 promoter regions and intron 1. In contrast to this, the tSNPs associated with urinary K^+ excretion span the entire length of the gene. However, there is some overlap between those tSNPs associated with BP and variation in urinary K^+. In particular, the variant rs765250, located in intron 1, which demonstrated the strongest evidence for association with EH and BP variation, is also associated with decreased in urinary K^+ excretion. These novel genetic data correlate well with what is known about WNK1 function, especially in relation to the primary phenotypes that characterise PHA2 - hyperkalemia and hypertension [37]. That is, we would expect true functional variants (or those in LD with these polymorphisms) to be associated with both BP and altered potassium excretion; this is what we observe.

Although there is some overlap between those tSNPs associated with BP and urinary potassium excretion (eg. 3/7 BP SNPs are also associated with variation in urinary potassium), not all BP associated variants were associated with urinary potassium and vice-versa. Furthermore, haplotype associated with increased BP were not also associated with decreased urine potassium excretion. This discrepancy may represent complex interactions between WNK1 polymorphisms and may also reflect the complexity of WNK1 regulation and its role in electrolyte homeostasis.

There are two major isoforms of WNK1: L-WNK1 and K-WNK1. L-WNK1 is ubiquitously expressed, but K-WNK1 has so far only been found to be expressed in the kidney [38]. Both L-WNK1 and K-WNK1 interact with each other to regulate common downstream targets involved in electrolyte homeostasis and BP regulation, via both kinase dependent and independent mechanisms [e.g., sodium chloride co-transporter (SLC12A3), epithelial sodium channel (ENaC) and the renal outer medullary potassium channel (KCNJ1)]. These isoforms are under the control of alternative promoters – one located 5’ of the gene for L-WNK1, and the other in intron 4, controlling expression of K-WNK1 [19]. Furthermore, both isoforms undergo tissue specific splicing and further variation is achieved by the use of two polyadenylation sites [18,19,39]. These data imply that there are multiple functional sites along the gene through which genetic variation could effect WNK1 expression and function. Furthermore, it has been observed in some PHA2 patients carrying the WNK1 deletion mutations, that the development of hyperkalemia may be separate from hypertension, and often precedes the development of high BP in these patients i.e., there may be no clear “cause and effect” relationship between the two phenotypes [40]. Therefore, it is possible that different WNK1 polymorphisms, either singly or in combination, could contribute to the two different phenotypes. This could account for some of our observations and will need to be explored with further studies.

We found multiple tSNPs spanning the entire length of the gene and several haplotypes to be associated with the traits of interest, suggesting there may be multiple causal variants across the WNK1 locus. Even though we used a comprehensive tSNP set that captured all known common WNK1 variation in HapMap, HapMap does not contain a complete catalogue of all genetic variation, thus fine mapping will be required to identify the true causal variants. Interestingly, all common HapMap SNPs in strong LD (r^2>0.8) with rs765250 and the other BP associated tSNPs, map to the L-WNK1 promoter, the K-WNK1 promoter located in intron 4 [19] and regions in intron 1 that span the sites of the PHA2 deletions (Figure S2), thus highlighting a few potential regions for targeted re-sequencing.

The most striking observations from our analyses were the identification of low frequency haplotypes with large BP lowering effects and their increased prevalence in the control population. Loss of WNK1 function is deleterious, as demonstrated by homozygous knockout mice which are embryonic lethal [12]. On the other hand, heterozygous knockout mice have low blood pressure, and this is associated with decreased WNK1 expression at the mRNA and protein level. Therefore, we can hypothesise that loss of function/expression mutations in WNK1 would be selected against and be rare in the general population. More subtle mutations, however, that lead to decreased WNK1 expression or
Table 6. WNK1 Haplotypes.

| Common F | 3  | 5  | 6  | 7  | 8  | 9  | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 20 | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 |
|----------|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| 1        | 0.15 | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  |
| 2        | 0.13 | 2  | 1  | 1  | 1  | 2  | 1  | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  | 1  |
| 3        | 0.12 | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 2  |
| 4        | 0.12 | 1  | 2  | 2  | 2  | 2  | 1  | 2  | 2  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 2  | 2  | 1  | 1  | 1  |
| 5        | 0.10 | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  |
| 6        | 0.06 | 1  | 2  | 2  | 2  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 2  | 1  | 1  | 2  | 1  | 1  |
| 7        | 0.06 | 2  | 1  | 2  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  | 1  | 1  |
| 8        | 0.05 | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 2  | 1  | 1  | 2  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  |

Low frequency

| 14       | 0.008 | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  |
| 15       | 0.008 | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  |
| 18       | 0.004 | 2  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 2  | 1  | 1  | 2  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  |
| 19       | 0.004 | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 2  | 1  | 1  | 1  | 1  | 1  |

All tSNPs are denoted numerically with reference to Table 2. Minor alleles “2” are highlighted in bold.

F: Frequency. Haplotype counts in cases and controls are estimates weighted by their posterior probabilities as determined by haplo.stats.

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Table 7. Haplotype association analyses in BRIGHT.

| Haplotype | Control* | Case* | F* | EH | SBP | DBP | 24 hr Urine potassiumd |
|-----------|----------|-------|----|----|-----|-----|------------------------|
| **COMMON**a | | | | | | | |
| 1 | 477.5 | 513.9 | 0.15 | reference | - | - | - |
| 2 | 405.6 | 451.9 | 0.13 | 1.0(0.8,1.2) | 0.06 | 0.9(0.1,1.9) | 0.09 |
| 3 | 368.3 | 451.1 | 0.12 | 1.1(0.9,1.4) | 0.2 | 2.1(0.3,3.9) | 0.02 |
| 4 | 307.9 | 400.9 | 0.12 | 0.9(0.7,1.1) | 0.4 | -0.9(-2.7,0.8) | 0.3 |
| 5 | 307.3 | 406.0 | 0.10 | 1.1(0.9,1.4) | 0.2 | 2.4(0.5,4.2) | 0.01 |
| 6 | 193.1 | 205.3 | 0.06 | 1.0(0.8,1.3) | 0.9 | -1.3(-3.4,0.9) | 0.2 |
| 7 | 188.7 | 194.7 | 0.06 | 1.0(0.8,1.3) | 0.9 | -0.5(-2.7,1.8) | 0.7 |
| 8 | 163.3 | 195.2 | 0.05 | 1.0(0.8,1.4) | 0.7 | 2.5(0,1,4,9) | 0.04 |
| Low frequency pool | 874.3 | 562.6 | 0.21 | 0.6(0.5,0.7) | 4.8×10⁻⁸ | -4.0(5.5, -2.5) | 2.1×10⁻⁷ |
| **LOW FREQUENCY**a | | | | | | | |
| 14 | 541 | 1.7 | 0.008 | 0.03(0.006,0.1) | 6.9×10⁻⁶ | -15.9(-19.9, -11.7) | 9.9×10⁻¹³ |
| 15 | 538 | 1.6 | 0.008 | 0.03(0.01,0.2) | 3.9×10⁻⁸ | -13.9(-18.1, -9.5) | 1.5×10⁻⁹ |
| 18 | 26.5 | 0.4 | 0.004 | 0.01(0.0005,0.4) | 0.01 | -160(-21.5, -10.3) | 1.5×10⁻⁷ |
| 19 | 23.9 | 1.6 | 0.004 | 0.06(0.01,0.3) | 6.4×10⁻⁶ | -179(-23.9, -11.6) | 1.1×10⁻⁷ |

Global p-valueb | <10⁻⁷ | <10⁻⁷ | <10⁻⁷ | 1×10⁻⁴ |

Haplotypes are numbered with reference to Table 4. Low frequency haplotypes are numbered with reference to Table 4 and Supplementary Table 1. Haplotype counts in cases and controls are estimates weighted by their posterior probabilities as determined by haplostats. F: Frequency.

bLinear effect estimate, 95% confidence intervals for EH, SBP, DBP and 24 hour urine potassium. The effect sizes are calculated by comparing each haplotype against haplotype 1: the most common haplotype or wild-type.

cGlobal p-value based on 10,000 permutations for 24 hour urine K⁺ and 10,000,000 for EH and SBP/DBP - adjusting for testing multiple haplotypes.

dAssociation with 24 hour urine potassium excretion was performed in cases only. Association analysis with urine potassium was not possible for the rare haplotypes. The associated rare haplotypes are almost uniquely distributed amongst the control population, for which as measures of 24 hour urine potassium excretion are not available.

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function may be ‘protective’ against hypertension, and preserved at low frequencies in the general population. However, the effects of genetic drift should not be underestimated in terms of allowing slightly deleterious alleles to persist in human populations [41]. This may explain some of our observations. The ability of common tSNPs/haplotypes to capture rare functional variants has previously been demonstrated for the angiotensinogen (AGT) gene [42]. It is possible that our tSNP analysis set is capturing rare loss-of-function mutations that may be embedded in these low frequency haplotype backgrounds, thus highlighting the need to re-sequence individuals carrying these low frequency haplotypes.

Our data suggest that multiple common WNK1 variants, with relatively weak effects, and multiple rare variants with large effects may be associated with blood pressure variation, and this should be explored further. Our findings are consistent with recent studies by Ji et al (2008) [43] and Tobin et al (2008) [44]. Both groups have performed a systematic analysis of the effect of variants in genes involved in renal salt handling on blood pressure and the development of hypertension. Many of which are regulated by WNK1 – SLC12A2, SLC12A1 and KCNJ1. Ji et al (2008) have identified rare variants in these genes that are associated with significantly lower blood pressure and protect from the development of hypertension in members of the Framingham Heart Study (FHS) [43]. In addition, Tobin et. al. (2008) have reported associations between common variants in these genes and blood pressure in families from the general population [44]. The findings suggest that both common and rare variants in genes responsible for some Mendelian disorders of hypertension and hypotension may also affect blood pressure variation in the general population. Our data lends further support to these observations.

The importance of rare variants to quantitative trait variability and susceptibility to disease has now been demonstrated for a number of other important phenotypes; including high density lipoprotein cholesterol (HDL-C), low density lipoprotein cholesterol (LDL-C) plasma triglycerides and body weight [45,46,47,48]. We believe there is now compelling evidence to initiate further studies to identify functional WNK1 variants that have a significant impact on BP variation, electrolyte homeostasis and risk of EH in the general population; thus taking an important step forward in our understanding of the pathogenesis of human essential hypertension.

Supporting Information

Figure S1 Distribution of low frequency blood pressure lowering haplotypes: The plot of systolic blood pressure (SBP)/diastolic blood pressure (DBP) values for each individual, showing the distribution of the associated low frequency haplotypes with posterior probabilities ≥0.9. The plot shows how the low frequency haplotypes are mainly found in low BP individuals. Found at: doi:10.1371/journal.pone.0005003.s001 (1.08 MB TIF)

Figure S2 The genomic structure of the human WNK1 gene is presented at the bottom of the panel. Exons are indicated by the vertical black bars and alternatively spliced exons by the red boxes. The yellow boxes indicate the position of the PHA2 disease causing deletions. For each of the blood pressure associated SNPs (red circles) located in intron 1, the r2 for each HapMap SNP with r2>0.8 is plotted on the y-axis against physical position (x-axis). The vertical dashed lines indicate the positions of the statistically similar SNPs. Found at: doi:10.1371/journal.pone.0005003.s002 (0.90 MB TIF)

Table S1 Haplotype association analysis

| haplotype | log odds-ratio | p-value |
|-----------|---------------|---------|
| Haplotype A | 1.67 | 0.05 |
| Haplotype B | 1.71 | 0.04 |
| Haplotype C | 1.93 | 0.005 |

Table S2 Distribution of rare variants in the WNK1 gene

| variant | frequency |
|---------|-----------|
| variant 1 | 0.05%
| variant 2 | 0.04%
| variant 3 | 0.03%

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