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A multi-stage genome-wide association study of bladder cancer identifies multiple susceptibility loci

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Abstract

We conducted a multi-stage, genome-wide association study (GWAS) of bladder cancer with a primary scan of 589,299 single nucleotide polymorphisms (SNPs) in 3,532 cases and 5,120 controls of European descent (5 studies) followed by a replication strategy, which included 8,381 cases and 48,275 controls (16 studies). In a combined analysis, we identified three new regions associated with bladder cancer on chromosomes 22q13.1, 19q12 and 2q37.1; rs1014971, $(P=8\times10^{-12})$ maps to a non-genic region of chromosome 22q13.1; rs8102137 $(P=2\times10^{-11})$ on 19q12 maps to *CCNE1*; and rs11892031 $(P=1\times10^{-7})$ maps to the *UGT1A* cluster on 2q37.1. We confirmed four previous GWAS associations on chromosomes 3q28, 4p16.3, 8q24.21 and 8q24.3, validated previous candidate associations for the *GSTM1* deletion $(P=4\times10^{-11})$ and a tag SNP for *NAT2* acetylation status $(P=4\times10^{-11})$, as well as demonstrated smoking interactions with both regions. Our findings on common variants associated with bladder cancer risk should provide new insights into mechanisms of carcinogenesis.

Bladder cancer is the fourth most common incident cancer in men1 and its frequent recurrence requires regular screening and interventions. Cigarette smoking and occupational exposure to aromatic amines have been strongly linked to bladder cancer risk.1 A family history of bladder cancer is associated with an approximately two-fold increase in risk; however, multiple-cancer families are rare and no high-penetrance genes have been identified to date2⁻⁴. Large meta-analyses of candidate gene studies have provided support for associations between NAT2 slow acetylation phenotype5 (defined by *NAT2* haplotypes) and a common gene deletion of *GSTM16* with bladder cancer risk7⁻⁸. Further, gene-environment interactions have been shown for smoking and *NAT2* acetylation, with an increased risk in slow acetylators, apparent only among cigarette smokers7⁻⁸.

Previous genome-wide association studies (GWAS) in bladder cancer have identified common variants in four genomic regions on chromosomes 3q289 (*TP63*), 4p16.3 (*TMEM129, TACC3-FGFR3*)10, 8q24.219, and 8q24.311 (*PSCA*) that are associated with risk. Interestingly, the variants on 8q24.21 map to a region centromeric to *MYC* that has been identified in GWAS of breast, colorectal and prostate cancers, as well as chronic lymphocytic leukemia12⁻¹⁸. Also, in follow-up analyses, an association with bladder cancer risk has been suggested for variants near the *TERT-CLPTM1L* locus on chromosome 5p15.33, which has also been associated by GWAS with risk for basal cell carcinoma, cutaneous melanoma, lung, brain and pancreatic cancers19⁻²³. However, the previously reported association with bladder cancer did not achieve genome-wide significance.

We conducted a multi-stage GWAS involving 3,532 cases and 5,120 controls of selfdescribed European descent in stage I, and followed up the most notable signals in two stages of replication (stages IIa/b and III) totaling 8,381 cases and 48,275 controls (Figure 1 and Online Methods). Individuals with scan data in stage I were participants in two casecontrol studies carried out in Spain and the USA (Maine and Vermont component of the New England Bladder Cancer Study) and three prospective cohort studies in the USA and Finland (see Supplementary Table 1 online for details). Replication analyses in stage II were carried out using existing scan data from two earlier studies. First, we evaluated the 100 most significant SNPs (excluding previously reported loci and SNPs with pairwise $r^2>0.8$) in 969 cases and 957 controls from the Texas Bladder Cancer study in the USA (stage IIa)11. Five of these SNPs were further evaluated in a second scan of 1,274 cases and 1,832 controls in The Netherlands (stage IIb)9. Three of the five SNPs were included or tagged at a pair-wise $r^2>0.8$ in the Dutch scan, and risk associations were confirmed for all three. In stage III, the three SNPs plus a tagging SNP for the *NAT2* acetylation status were evaluated in 6,141 cases and 45,486 controls from 11 case-control and 3 prospective cohort studies in the USA and Europe (see Figure 1 and Supplementary Table 1).

After quality control analysis of genotypes, we combined the data sets in stage I resulting in 589,299 SNPs available for analysis (based on the common SNPs called from both the Illumina Human1M and Human 610-Quad) in 3,532 cases and 5,120 controls (**Online Methods**). A logistic regression model was fit for genotype trend effects (1 d.f.) adjusted for study center, age, sex, smoking status (current, former or never) and DNA source (blood/ buccal). The quantile-quantile (Q-Q) plot showed little evidence for inflation of the test statistics as compared to the expected distribution (corrected λ_{1000} subjects=1.021), which minimizes the likelihood of substantial hidden population substructure or differential genotype calling between cases and controls24 (**Online Methods and** Supplementary Figure 1). A Manhattan plot displays the results of the combined GWAS in stage I (Supplementary Figure 2).

Data from the first stage confirm the associations reported with tag SNPs in the four previously identified genomic regions on chromosomes 3q28 (rs710521)9, 8q24.21 (rs9642880)9, 8q24.3 (rs2294008)11 and 4p16.3 (rs798766)10 as well as a suggested region in 5p15.33 (rs401681; a neighboring SNP, rs2736098, was also reported but data were not available in our study)19 (Table 1 **and** Supplementary Figure 3). Consistent with prior reports9·10, rs9642880 on 8q24.21 and rs798766 on 4p16.3 were most strongly associated with tumors of low grade/low risk of progression (Supplementary Table 2). A stronger association with low grade/low risk disease was also suggested for rs401681 on 5p15.33 (Supplementary Table 2). In addition, we used a copy number variation TaqMan assay7 to assess the presence of *GSTM1* on 1p13.3 to genotype stage I samples, and confirmed an association with increased bladder cancer risk (Table 1).

In a combined analysis based on case/control counts by genotype and study, we estimated odds ratios (ORs) using logistic regression analyses adjusted for study center. Meta-analyses of estimated ORs adjusted for age, sex, smoking status and DNA source produced comparable point estimates (Supplementary Table 3). Our combined analysis of stages I, II and III identified three novel genomic regions on chromosomes 22q13.1, 19q12 and 2q37.1 that were associated with bladder cancer risk below the threshold for genome-wide significance ($P < 5 \times 10^{-7}$)25 (Table 2 and Supplementary Figure 4 for study and stage specific estimates, Figure 2). We also confirmed a signal below genome-wide significance for rs1495741, which tags the NAT2 acetylator status26 previously reported as a bladder cancer susceptibility locus on 8p227³. The new SNP is located approximately 10kb of the 3' end of the gene.

The locus on chromosome 22q13.1, rs1014971 (P_{trend} =8.4×10⁻¹²; OR per C allele =0.88, 95%CI 0.85-0.91)), was primarily associated with high-risk tumors (Supplementary Table 2). The locus is located in a non-genic region, approximately 25 kb centromeric of the

catalytic polypeptide-like 3A (*APOBEC3A*) and 64 kb telomeric of the chromobox homolog 6 (*CBX6*). *APOBEC3A* is an apolipoprotein B mRNA editing enzyme that belongs to the cytidine deaminase gene family, which can play a role in the initiation of tumorigenesis by deamination of cytosine (C) to uracil (U)27. *CBX6* is a component of the chromatin – associated polycomb complex involved in transcriptional repression.

In the combined analysis, we observed an association with rs8102137 on chromosome 19q12 (P_{trend} =1.7×10⁻¹¹; OR per C allele =1.13, 95%CI 1.09-1.17), which maps to the cyclin E1 gene (*CCNE1*). *CCNE1* is a key member of the cyclin/cyclin-dependent kinase (Cdk)/retinoblastoma protein (pRB) pathway which determines the rates of cell cycle transition from G1 to S phase, and is commonly altered in bladder cancer and other tumors28. Cyclin E1 expression in bladder cancer has been associated with high grade or muscle invasive tumors and poor clinical outcome29. Consistently, rs8102137 was most strongly associated with risk of high grade/high risk tumors (Supplementary Table 2).

A third locus is marked by rs11892031 ($P=1.0\times10^{-7}$; OR per C allele =0.84, 95%CI 0.79-0.89) on chromosome 2q37.1 and resides in an intronic region of the UDP-glucuronosyltransferase (UGT) 1A gene locus, which encodes the UGT1A family of proteins. Glucuronidation by UGTs facilitates solubility and removal of substrates such as endo- and xenobiotics (including carcinogens in tobacco smoke) via bile or urine30. Genetic variation in *UGT1A* has been associated with predisposition to severe gastrointestinal toxicity of the anticancer drug irinotecan31. The *UGT1A* locus is represented by at least nine highly homologous transcripts, collectively known as UGTs, generated by alternative splicing. Tissue-specific loss or decreased expression of UGTs has been associated with several gastrointestinal cancers and bladder cancer32⁻34, as well as experimentally induced bladder cancer in animal models35.

Previously, a promising signal in the *CLPTM1L-TERT* locus on chromosome 5p15.33 was reported in a region in which common variants have been associated with multiple cancers in recent GWAS19⁻23. In addition, rare mutations in *TERT* have been linked to dyskeratosis congenita (a bone marrow failure syndrome), idiopathic pulmonary fibrosis, acute myelogenous leukemia and chronic lymphocytic leukemia36⁻39. In the first stage of this GWAS, we observed a moderately significant effect for rs401681 (P= 2.9 × 10⁻³), which was at genome-wide significance when combined with the Rafnar et al. data (P= 5.0 × 10⁻⁷; OR per C allele 1.11, 95% CI 1.07-1.16) (Table 1, Supplementary Figure 3).

The risk associated with GSTM1 and NAT2 varied in strength across categories of cigarette smoking, whereas genotype risk associations by smoking categories were of similar magnitude for the eight susceptibility loci identified by GWAS (Supplementary Table 4). In a combined analysis, the risk association with GSTM1 deletion was strongest in never smokers (OR=1.75, 95%CI=1.44-2.13), and progressively weaker in former (OR=1.55, 95% CI=1.35-1.78) and current smokers (OR=1.25, 95% CI =1.07-1.46; Pinteraction = 0.008 for current vs. never smokers; Table 3). The stronger association of the GSTM1 deletion among non-smokers is a novel observation that was not evident in previous case-only metaanalyses7. rs1495741 located on the 3' end of NAT2 is a marker of the NAT2 phenotype associated with bladder cancer risk26. The rs1495741 GG genotype marking the slow acetylation phenotype, compared to the combined AG/AA genotypes corresponding to the intermediate/rapid acetylation phenotypes, showed a highly significant ($P=5.5\times10^{-7}$) association with increased bladder cancer risk that was limited to cigarette smokers $(OR=1.24, 95\% CI=1.16-1.32 P=4.3 \times 10^{-11}; P_{interaction}=6.3 \times 10^{-5})$ (Supplementary Figure 5) and Supplementary Table 3). This interaction is consistent with the role of NAT2 in the detoxification of bladder carcinogens such as aromatic amines from tobacco smoke.

Our three-stage study had adequate power to detect variants of moderate effect sizes over a range of common allele frequencies. For the newly discovered SNP markers, the power to detect the observed associations at a level of genome-wide significance was at 54%, 30%, 30% and 6% for rs104971, rs1495741, rs8102137 and rs11892031, respectively. In light of the limited power to discover SNPs with modest effect sizes, additional loci with similar effect sizes will likely be identified with larger scale GWAS. Based on a recent estimator40 that incorporates novel and previously reported loci together, we estimate that approximately two dozen additional bladder cancer susceptibility SNP markers of similar magnitude and frequencies might be discovered. Future studies should be powered with adequate sample size to detect additional variants.

With the exception of the *GSTM1* deletion, relative risk estimates for novel loci are based on associations using tag SNPs, which most likely underestimate the association with biologically important alleles. Accordingly, further studies are needed to define the functional variants and the clinical utility of risk models that combine genetic markers with epidemiologic risk factors for bladder cancer (i.e. smoking, occupational and environmental exposures, family history). Our combined analysis of 12,254 individuals with bladder cancer and 53,395 controls has uncovered three new genomic regions associated with bladder cancer risk. Fine-mapping studies of these three regions are needed to identify candidate variants for functional studies that should shed light into biological mechanisms for the associations reported through GWAS. This knowledge could establish the foundation for developing improved preventive, diagnostic and/or therapeutic approaches.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Appendix

Online Methods

Study Participants

Participants were drawn from 21 studies (Supplementary Table 1). For stage I, cases were defined as histologically confirmed primary carcinoma of the urinary bladder including carcinoma *in situ* (ICD-0-2 topography codes C67.0-C67.9 or ICD9 codes 188.1-188.9). Each participating study obtained informed consent from study participants and approval from its Institutional Review Board (IRB) for this study. For stage I only, participating studies obtained institutional certification permitting data sharing in accordance with the NIH Policy for Sharing of Data Obtained in NIH Supported or Conducted Genome-Wide Association Studies (GWAS).

Genotyping and Quality Control

For stage I, genome-wide genotyping was conducted using three chips, SBCS (HumanHap 1 Million), NEBC-ME/VT (Human Hap 610-Quad), ATBC, CPS-II and PLCO (cases)

(Human Hap 610) and controls from CGEMS/GEI for PLCO (Human Hap 550-r equivalent). DNA samples were selected for genotyping based on pre-genotyping quality control measures performed for GWAS at the Core Genotyping Facility of the NCI 4,089 blood samples and 2,813 buccal samples were analyzed. Repeat genotyping was performed on 38 blood samples (19 cases and 19 controls) and 10 buccal samples (2 cases and 8 controls) on Illumina 1M chips after suitable metrics identified performance issues. Cancer free controls (N=2003) were previously scanned in CGEMS18 and a lung cancer GWAS21.

Genotype clusters were estimated with samples by study with preliminary completion rates greater than 98% per individual study (namely SBCS, NEBC-ME/VT, PLCO, ATBC and CPS-II). Genotypes for the analytical build were based on study specific clustering. SNP assays with locus call rates lower than 90% were excluded.

SNPs with extreme departures from Hardy-Weinberg proportions ($P < 1 \times 10^{-7}$) were excluded from the association analysis due to the increased likelihood of spurious associations due to problematic assays or genotyping calling.42 Additional participants were excluded based on: 1) completion rates lower than 94-96% (n=203 samples); 2) heterozygosity of less than 22% or >35% (n=12); 3) inter-study unexpected duplicates (n=5); 4) phenotype exclusions (due to ineligibility or incomplete information) (n=94).

Assessment of population structure of study participants was performed with STRUCTURE43 by seeding the analysis with founder genotypes from three HapMap populations (Phase I and II build 26).44 A set of 12,898 SNPs with extremely low pair-wise correlation (r^2 <0.004) was selected for this analysis.45⁻47 A total of 55 participants (43 cases and 12 controls) were estimated to have less than 85% HapMap CEU admixture (Supplementary Figure 6). Principal component analysis (PCA) of scanned subjects (excluding inferred sib and half-sib pairs) was performed with GLU (a similar procedure to EIGENSTRAT)45^{,4}6 and did not reveal notable eigenvectors. Consequently, a study-specific indicator was used for the stage I analysis46.

We estimated the inflation of the test statistic, λ , adjusted to a sample size of 1000 cases/ 1000 controls as per the method of de Bakker et. al: λ (corrected)= 1 + (λ -1) × [n_{case}^{-1} + n_{cont}^{-1}]/[2×10⁻³].48 The corrected estimated λ_{1000} is 1.021while the uncorrected λ is 1.086 (Supplementary Figure 1).

Twenty participant pairs were identified as potential relatives based on genotyping sharing in excess of theoretical expectations. A set of 4,546 SNPs were selected (with completion rates >95%, MAF>0.3 and r^2 <0.01 in the three HapMap populations) and used to run PREST49 to formally test for cryptic relatedness. 19 unexpected full-sib and 1 parent-child pairs were identified and excluded from PCA (but included in the association analysis). 243 expected duplicates (including 6 triplicates in ATBC) were evaluated and yielded a concordance rate of 99.99%.

The final participant count for stage I analysis was 3,532 cases and 5,120 controls (Supplementary Table 1). The number of SNPs available for association analysis in all studies but SBCS was 589,299. In the SBCS, genotyped with the Infinium HumanHap 1 M chip, after quality control metrics were applied, 1,002,634 SNPs were available and 571,643 overlapped exactly with the 610Quad/550k data.

TaqMan custom genotyping assays (ABI, Foster City, CA) were designed and optimized for 4 SNPs, including the tag SNP for *NAT2*. In an analysis of 1,107 samples from three studies, the comparison of the Illumina calls with the TaqMan assays showed an average concordance rate of 99.4% (range 99.2-99.8%); no shifts from wild type to homozygotes

were observed. The Illumina Infinium cluster plots for the four novel associations, rs1014971, rs8102137, rs11892031 and rs1495741 are shown in Supplementary Figure 7.

Association Analysis

Association analyses for stage I were conducted using logistic regression, adjusted for age (in five-year categories), sex, smoking (current, former or never), DNA source (buccal/ blood) and study. Each SNP genotype was coded as a count of minor alleles, with the exception of X-linked SNPs among men that were coded as 2 if the participant carried the minor allele and 0 if he carried the major allele.50 A score test with one degree of freedom was performed on all genetic parameters in each model to determine statistical significance. We assessed heterogeneity in genetic effects across studies using the I² statistic. For the inclusion of stage II and III data, we used genotype counts by case-control status and study, and conducted a fixed effects meta-analysis. We also conducted a meta-analysis based on estimates of allelic odds ratio adjusted by age, sex, smoking status, DNA source and study; the estimates did not materially differ from the fixed-effects meta-analysis (Supplementary Table 3).

Polytomous logistic regression was used to obtain estimates of effect for different tumor subtypes. Case-only analyses with tumor type as an outcome were used to test for differences in effect size across subtypes. Models for tumor grade constrained the effect size to increase linearly across levels. Genotype-smoking interactions were assessed using logistic regression for grouped data adjusted by study and including interaction terms. Forest plots by smoking, including summary estimates from fixed effects meta-analyses, are also shown for rs1495741.

Data analysis and management was performed with GLU (Genotyping Library and Utilities version 1.0), a suite of tools available as an open-source application for management, storage and analysis of GWAS data, and STATA.

Estimate of Recombination Hotspots

SequenceLDhot51 that uses an approximate marginal likelihood method52 was used to compute likelihood ratio (LR) statistics for a set of putative hotspots across the region of interest. We sequentially analyzed subsets of 100 controls of European descent (by pooling 5 controls from each study). We used Phasev2.1 to infer the haplotypes as well as background recombination rates. The analysis was repeated with five non-overlapping sets of 100 pooled controls.

Data Access

The CGEMS data portal provides access to individual level data for investigators from certified scientific institutions after approval of their submitted Data Access Request.

URLs:

CGEMS portal: http://cgems.cancer.gov/

CGF: http://cgf.nci.nih.gov/

GLU: http://code.google.com/p/glu-genetics/

EIGENSTRAT: http://genepath.med.harvard.edu/~reich/EIGENSTRAT.htm

SNP500Cancer: http://snp500cancer.nci.nih.gov/

STRUCTURE: http://pritch.bsd.uchicago.edu/structure.html

Tagzilla: http://tagzilla.nci.nih.gov/

References

- 1. Silverman, D.; SS, D.; LE, M.; N, R. Bladder Cancer. In: D, S.; Fraumeni, JF., Jr., editors. Cancer Epidemiology and Prevention. Oxford University Press; New York, NY: 2006. p. 1101-1127.
- 2. Kantor AF, Hartge P, Hoover RN, Fraumeni JF Jr. Familial and environmental interactions in bladder cancer risk. Int J Cancer. 1985; 35:703–6. [PubMed: 4008097]
- Murta-Nascimento C, et al. Risk of bladder cancer associated with family history of cancer: do lowpenetrance polymorphisms account for the increase in risk? Cancer Epidemiol Biomarkers Prev. 2007; 16:1595–600. [PubMed: 17684133]
- 4. Aben KK, et al. Segregation analysis of urothelial cell carcinoma. Eur J Cancer. 2006; 42:1428–33. [PubMed: 16737809]
- Lower GM Jr. et al. N-acetyltransferase phenotype and risk in urinary bladder cancer: approaches in molecular epidemiology. Preliminary results in Sweden and Denmark. Environ Health Perspect. 1979; 29:71–9. [PubMed: 510245]
- Bell DA, et al. Genetic risk and carcinogen exposure: a common inherited defect of the carcinogenmetabolism gene glutathione S-transferase M1 (GSTM1) that increases susceptibility to bladder cancer. J Natl Cancer Inst. 1993; 85:1159–64. [PubMed: 8320745]
- Garcia-Closas M, et al. NAT2 slow acetylation, GSTM1 null genotype, and risk of bladder cancer: results from the Spanish Bladder Cancer Study and meta-analyses. Lancet. 2005; 366:649–59. [PubMed: 16112301]
- Rothman N, Garcia-Closas M, Hein DW. Commentary: Reflections on G. M. Lower and colleagues' 1979 study associating slow acetylator phenotype with urinary bladder cancer: meta-analysis, historical refinements of the hypothesis, and lessons learned. Int J Epidemiol. 2007; 36:23–8. [PubMed: 17510073]
- 9. Kiemeney LA, et al. Sequence variant on 8q24 confers susceptibility to urinary bladder cancer. Nat Genet. 2008; 40:1307–12. [PubMed: 18794855]
- Kiemeney LA, et al. A sequence variant at 4p16.3 confers susceptibility to urinary bladder cancer. Nat Genet. 2010; 42:415–9. [PubMed: 20348956]
- 11. Wu X, et al. Genetic variation in the prostate stem cell antigen gene PSCA confers susceptibility to urinary bladder cancer. Nat Genet. 2009; 41:991–5. [PubMed: 19648920]
- 12. Eeles RA, et al. Identification of seven new prostate cancer susceptibility loci through a genomewide association study. Nat Genet. 2009; 41:1116–21. [PubMed: 19767753]
- Yeager M, et al. Identification of a new prostate cancer susceptibility locus on chromosome 8q24. Nat Genet. 2009; 41:1055–7. [PubMed: 19767755]
- 14. Crowther-Swanepoel D, et al. Common variants at 2q37.3, 8q24.21, 15q21.3 and 16q24.1 influence chronic lymphocytic leukemia risk. Nat Genet. 2010; 42:132–6. [PubMed: 20062064]
- 15. Tomlinson IP, et al. A genome-wide association study identifies colorectal cancer susceptibility loci on chromosomes 10p14 and 8q23.3. Nat Genet. 2008; 40:623–30. [PubMed: 18372905]
- Easton DF, et al. Genome-wide association study identifies novel breast cancer susceptibility loci. Nature. 2007; 447:1087–93. [PubMed: 17529967]
- 17. Zanke BW, et al. Genome-wide association scan identifies a colorectal cancer susceptibility locus on chromosome 8q24. Nat Genet. 2007; 39:989–94. [PubMed: 17618283]
- Yeager M, et al. Genome-wide association study of prostate cancer identifies a second risk locus at 8q24. Nat Genet. 2007; 39:645–9. [PubMed: 17401363]
- Rafnar T, et al. Sequence variants at the TERT-CLPTM1L locus associate with many cancer types. Nat Genet. 2009; 41:221–7. [PubMed: 19151717]
- 20. Stacey SN, et al. New common variants affecting susceptibility to basal cell carcinoma. Nat Genet. 2009

- Landi MT, et al. A genome-wide association study of lung cancer identifies a region of chromosome 5p15 associated with risk for adenocarcinoma. Am J Hum Genet. 2009; 85:679–91. [PubMed: 19836008]
- 22. Shete S, et al. Genome-wide association study identifies five susceptibility loci for glioma. Nat Genet. 2009; 41:899–904. [PubMed: 19578367]
- Petersen GM, et al. A genome-wide association study identifies pancreatic cancer susceptibility loci on chromosomes 13q22.1, 1q32.1 and 5p15.33. Nat Genet. 2010; 42:224–8. [PubMed: 20101243]
- 24. Freedman ML, et al. Assessing the impact of population stratification on genetic association studies. Nat Genet. 2004; 36:388–93. [PubMed: 15052270]
- Genome-wide association study of 14,000 cases of seven common diseases and 3,000 shared controls. Nature. 2007; 447:661–78. [PubMed: 17554300]
- 26. García-Closas M, et al. A single nucleotide polymorphism identified in a genome-wide scan tags variation in the N-acetyltransferase 2 phenotype in populations of European background. Pharmacogenet Genomics. 2010
- 27. Conticello SG. The AID/APOBEC family of nucleic acid mutators. Genome Biol. 2008; 9:229. [PubMed: 18598372]
- Malumbres M, Barbacid M. To cycle or not to cycle: a critical decision in cancer. Nat Rev Cancer. 2001; 1:222–31. [PubMed: 11902577]
- 29. Richter J, et al. High-throughput tissue microarray analysis of cyclin E gene amplification and overexpression in urinary bladder cancer. Am J Pathol. 2000; 157:787–94. [PubMed: 10980118]
- Strassburg CP, Lankisch TO, Manns MP, Ehmer U. Family 1 uridine-5'-diphosphate glucuronosyltransferases (UGT1A): from Gilbert's syndrome to genetic organization and variability. Arch Toxicol. 2008; 82:415–33. [PubMed: 18491077]
- Ando Y, et al. Polymorphisms of UDP-glucuronosyltransferase gene and irinotecan toxicity: a pharmacogenetic analysis. Cancer Res. 2000; 60:6921–6. [PubMed: 11156391]
- Strassburg CP, Manns MP, Tukey RH. Differential down-regulation of the UDPglucuronosyltransferase 1A locus is an early event in human liver and biliary cancer. Cancer Res. 1997; 57:2979–85. [PubMed: 9230212]
- Strassburg CP, Nguyen N, Manns MP, Tukey RH. Polymorphic expression of the UDPglucuronosyltransferase UGT1A gene locus in human gastric epithelium. Mol Pharmacol. 1998; 54:647–54. [PubMed: 9765507]
- Giuliani L, et al. Can down-regulation of UDP-glucuronosyltransferases in the urinary bladder tissue impact the risk of chemical carcinogenesis? Int J Cancer. 2001; 91:141–3. [PubMed: 11149414]
- Iida K, et al. Suppression of AhR signaling pathway is associated with the down-regulation of UDP-glucuronosyltransferases during BBN-induced urinary bladder carcinogenesis in mice. J Biochem. 2009; 147:353–60. [PubMed: 19880377]
- Calado RT, Young NS. Telomere maintenance and human bone marrow failure. Blood. 2008; 111:4446–55. [PubMed: 18239083]
- Armanios MY, et al. Telomerase mutations in families with idiopathic pulmonary fibrosis. N Engl J Med. 2007; 356:1317–26. [PubMed: 17392301]
- Tsakiri KD, et al. Adult-onset pulmonary fibrosis caused by mutations in telomerase. Proc Natl Acad Sci U S A. 2007; 104:7552–7. [PubMed: 17460043]
- Calado RT, et al. Constitutional hypomorphic telomerase mutations in patients with acute myeloid leukemia. Proc Natl Acad Sci U S A. 2009; 106:1187–92. [PubMed: 19147845]
- 40. Park JH, et al. Estimation of effect size distribution from genome-wide association studies and implications for future discoveries. Nat Genet. 2010; 42:570–5. [PubMed: 20562874]
- 41. Hein DW. N-acetyltransferase 2 genetic polymorphism: effects of carcinogen and haplotype on urinary bladder cancer risk. Oncogene. 2006; 25:1649–58. [PubMed: 16550165]
- 42. Wigginton JE, Cutler DJ, Abecasis GR. A note on exact tests of Hardy-Weinberg equilibrium. Am J Hum Genet. 2005; 76:887–93. [PubMed: 15789306]

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- 43. Pritchard JK, Stephens M, Donnelly P. Inference of population structure using multilocus genotype data. Genetics. 2000; 155:945–59. [PubMed: 10835412]
- 44. Frazer KA, et al. A second generation human haplotype map of over 3.1 million SNPs. Nature. 2007; 449:851–61. [PubMed: 17943122]
- 45. Price AL, et al. Principal components analysis corrects for stratification in genome-wide association studies. Nat Genet. 2006; 38:904–9. [PubMed: 16862161]
- Patterson N, Price AL, Reich D. Population structure and eigenanalysis. PLoS Genet. 2006; 2:e190. [PubMed: 17194218]
- 47. Yu K, et al. Population substructure and control selection in genome-wide association studies. PLoS One. 2008; 3:e2551. [PubMed: 18596976]
- 48. de Bakker PI, et al. Practical aspects of imputation-driven meta-analysis of genome-wide association studies. Hum Mol Genet. 2008; 17:R122–8. [PubMed: 18852200]
- Sun L, Wilder K, McPeek MS. Enhanced pedigree error detection. Hum Hered. 2002; 54:99–110. [PubMed: 12566741]
- 50. Clayton D. Testing for association on the X chromosome. Biostatistics. 2008; 9:593–600. [PubMed: 18441336]
- 51. Fearnhead P. SequenceLDhot: detecting recombination hotspots. Bioinformatics. 2006; 22:3061–6. [PubMed: 17060358]
- Fearnhead P, Harding RM, Schneider JA, Myers S, Donnelly P. Application of coalescent methods to reveal fine-scale rate variation and recombination hotspots. Genetics. 2004; 167:2067–81. [PubMed: 15342541]



Figure 1. Study design of multi-stage GWAS of bladder cancer

See **Online Methods** and Supplementary Table 1 for details of study designs and sample sizes. *The tag SNP, rs1495741 located 3' of *NAT2*41 was genotyped in subjects in stage II and III studies as well as on the Illumina bead chips used in stage I. **Includes 338 additional cases from NBCS that were added to the final combined analyses.

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Figure 2. Association results, recombination and linkage disequilibrium plots for four regions on chromosomes 22q13.1, 19q12, 2q37.1 and 8p22

Results of stage I (green circles), combined stages II and III (blue diamonds) and combined data from the three stages (red diamonds) with P-values for log-additive association results with recombination rates (cm/Mb) based on HapMap phase II data. Pairwise r² values based on control populations are displayed at the bottom for all SNPs included in the GWAS analysis. **Panel A** depicts chromosome 22q13.1 region (37,617,065 to 37,743,614). **Panel B** depicts the region of chromosome 19q12 (34,922,089 to 35,080,325). **Panel C** depicts the region of 2q37.1 (234,131,582 to 234,286,564). **Panel D** depicts the region of 8p22 (18,216,291 to 18,406,519). Genomic coordinates are based on NCBI Human Genome Build 36.3.

Table 1

Previously reported genetic variants associated with bladder cancer risk

Results of meta-analyses of allelic OR estimates for the markers reported to achieve genome-wide significance25. Studies in Kiemeney et al. 20089 Hampshire, LBCS, IBCS, TBCS, Sweden, EEBCS, Belgium, BBCS, ZBCS, MSKCC. Studies in Rafnar et al 200919 include NBCS, IBCS, LBCS, include: NBCS, LBCS, IBCS, TBCS, Sweden, Belgium, EEBCS, BBCS, ZBCS. Studies in Wu et al. 200911 include: NBCS, TXBCS1/2, New Sweden, TBCS, EEBCS, Belgium, ZBCS, BBCS.

Marker ^{a} , risk allele ^{b} , chr ^{c} , location ^{c} and gene ^{d}	Groups of studies	N ^e	Cases	Controls	Freq <i>f</i>	V V	llelic O 5%CI)	2 %	P value ^h
rs9642880 [T]	Previously reported ¹ 9	6	3,855	37,985	0.45	1.22	1.15	1.29	7.8E-12
Chr 8q24.21:	Stage I	5	3,525	5,108	0.45	1.21	1.13	1.29	4.6E-08
128787250	Combined	14	7,380	43,093		1.21	1.16	1.27	2.0E-18
MYC									
rs710521 [A]	Previously reported ¹ 9	6	3,855	37,985	0.73	1.19	1.12	1.27	1.1E-07
Chr 3q28:	Stage I	5	3,519	5,110	0.72	1.15	1.07	1.25	3.3E-04
191128627	Combined	14	7,374	43,095		1.18	1.12	1.24	1.8E-10
TP63									
rs2294008 [T]	Previously reported ¹ 11	13	6,667	39,590	0.46	1.15	1.10	1.20	2.0E-10
Chr 8q24.3:	Stage I	S	3,529	5,115	0.45	1.08	1.01	1.16	2.2E-02
143758933	Combined	18	10,196	44,705		1.13	1.09	1.17	4.4E-11
PSCA									
rs401681 [C]	Previously reported ¹ 19	6	4,147	34,988	0.54	1.12	1.06	1.18	5.1E-05
Chr 5p15.33:	Stage I	5	3,526	5,117	0.55	1.11	1.04	1.19	2.9E-03
1375087	Combined	14	7,673	40,105		1.11	1.07	1.16	5.0E-07
TERT-CLPTMIL									
rs798766 [T]	Previously reported ¹ 10	Π	4,580	45,269	0.19	1.24	1.17	1.32	9.5E-12
Chr 4p16.3:	Stage I	5	3,531	5,118	0.19	1.14	1.05	1.23	2.6E-03
1704037	Combined	16	8,111	50,387		1.20	1.14	1.26	3.9E-13
TMEM129 TACC3-FGFR3									
Deletion Assay	Previously reported $^{\dot{I}}$ 7,9	28	5,072	6,466	0.51	1.46	1.35	1.58	1.9E-21

Marker ^{a} , risk allele ^{b} , chr ^{c} , location ^{c} and gene ^{d}	Groups of studies	Ne	Cases	Controls	Freq <i>f</i>	¥ 5)	llelic Ol 5%CI)	24 %	P value ^h
Chr 1p13.3	Stage V	4	2,480	3,222	0.49	1.49	1.33	1.68	3.7E-11
GSTMI	Combined	32	7,552	9,688	0.53	1.47	1.38	1.57	5.0E-31
NCBI dbSNP identi	fier.								
, Risk allele shown ir	.0 נ								
Chromosome and N	CBI Human Genome Build 3	6.3 loca	tion.						
r Gene neighborhood	closest to the most notable SI	٨P.							
N: number of studie	s								
Risk allele frequency	y in control populations.								
Estimate assuming 1	multiplicative odds model, OF	ς, odds	ratio; CI,	95% confid	ence interv	/al.			
1 d.f. trend test.									
Summary estimates	differ slightly from previously	/ publis	hed becau	ise we used	a fixed eff	ects me	ta-analy	sis.	

data to appear in "GSTM1 null and NAT2 Slow Acetylation Genotypes, Smoking Intensity, and Bladder Cancer Risk: Results from the New England Bladder Cancer Case-Control Study and Meta-Analyses" by Moore LE, Baris D, Figueroa J, Garcia-Closas M, Karagas M, Schwenn M, Johnson A, Lubin J, Hein DW, Dagnall C, Colt J, Kida M, Jones M, Schned A, Cherela S, Chanock S, Cantor K, Silverman D, Rothman N) Data from SBCS was excluded from stage I because it had been included in the previous meta-analyses published in Garcia-Closas et al. 20057... Data from NEBCS are reported separately (Unpublished

Table 2

Novel SNPs identified in a multi-stage GWAS of bladder cancer

Results of the meta-analysis of genotype counts included in combined stages I, II and III. The initial scan results were adjusted by age, gender, smoking status, study site and DNA source.

Marker ^{<i>a</i>} , minor allele ^{<i>b</i>} , chr ^{<i>c</i>} , location ^{<i>c</i>} and gene ^{<i>d</i>}	Studies included	Ne	Cases	Controls	Freqf	V V	Jlelic O 95%CI)	X %	P value ^h
rs1014971 [C]	Stage I	S	3,529	5,092	0.38	0.87	0.82	0.93	6.1E-05
Chr 22q13.1:	Stage II and III	15	8,277	47,517	0.37	0.89	0.85	0.93	3.0E-08
37662569	Combined	20	11,806	52,609	0.38	0.88	0.85	0.91	8.4E-12
CBX6, APOBEC3A									
rs8102137 [C]	Stage I	2	3,530	5,114	0.33	1.13	1.06	1.21	1.6E-04
Chr 19q12:	Stage II and III	15	8,261	47,708	0.32	1.13	1.08	1.18	2.6E-08
34988693	Combined	20	11,791	52,822	0.33	1.13	1.09	1.17	1.7E-11
CCNEI									
rs11892031 [C]	Stage I	5	3,524	5,108	0.08	0.79	0.70	0.89	6.9E-05
Chr 2q37.1:	Stage II and III	15	8,284	47,727	0.08	0.86	0.80	0.93	1.8E-04
234230022	Combined	20	11,808	52,835	0.08	0.84	0.79	0.89	1.0E-07
<i>UGT1A</i> cluster ^j									
rs1495741 [G]	Stage I	5	3525	5116	0.24	0.86	0.80	0.93	1.5E-04
Chr 8p22:	Stage II and III	15	8,279	47,744	0.22	0.87	0.83	0.92	6.8E-08
18317161	Combined	20	11,804	52,860	0.24	0.87	0.83	0.91	4.2E-11
NAT2									
⁴ NCBI dbSNP identifier.									
b Minor allele shown in []	<u> </u>								

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 $^{\mathcal{C}}$ Chromosome and NCBI Human Genome Build 36.3 location.

 $d_{\rm Gene}$ neighborhood closest to the most notable SNP.

 $e_{\rm N}$: number of studies

fMinor allele frequency in control populations.

 ${}^{\mathcal{E}}_{}$ Estimate assuming multiplicative odds model, OR, odds ratio; CI, 95% confidence interval.

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Table 3

Interaction of NAT2 tagSNP (rs1495741) and GSTMI deletion with cigarette smoking in bladder cancer risk

Results from logistic regression analyses of aggregated data adjusted by study. rs1495741 genotypes were classified according to an established approach based on the phenotype of NAT2 acetylators; AA/AG were collapsed to tag rapid/intermediate acetylation, and GG tags slow acetylation.26

	Cas	es	Contr	.ols					Р
<i>NAT2</i> (rs1495741)	AA/AG	99	AA/AG	66	OR	95%	CI	Ь	Interaction
All subjects	3,784	6,915	5,233	8,182	1.15	1.09	1.22	2.9E-07	
By smoking status									
Never smoker	760	1,202	1,679	2,758	0.95	0.85	1.06	3.3E-01	Ref.
Ever smoker	3,024	5,713	3,554	5,424	1.24	1.16	1.32	4.3E-11	6.3E-05
Former smoker	1,859	3,455	2,300	3,559	1.20	1.11	1.29	6.8E-06	6.7E-04
Current smoker	1,165	2,258	1,254	1,865	1.27	1.14	1.40	7.2E-06	1.6E-04
GSTMI Del	Present	Null	Present	Null					
All subjects	1,319	1,995	1,717	1,726	1.47	1.33	1.62	4.4E-14	
By smoking status									
Never smoker	210	346	519	510	1.71	1.38	2.12	6.9E-07	Ref.
Ever smoker	1,109	1,649	1,198	1,216	1.47	1.30	1.67	2.1E-09	1.1E-01
Former smoker	564	961	622	653	1.62	1.39	1.89	4.7E-10	6.9E-01
Current smoker	545	688	576	563	1.19	1.00	1.40	4.5E-02	8.1E-03