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Multi-ethnic genome-wide association study for atrial fibrillation

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Short title: GWAS for Atrial Fibrillation

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Johnson. Dr. Kirchhof receives research support from European Union, British Heart Foundation, Leducq Foundation, Medical Research Council (UK), and German Centre for Cardiovascular Research, from several drug and device companies active in atrial fibrillation, and has received honoraria from several such companies. Dr. Kirchhof is also listed as inventor on two patents held by University of Birmingham (Atrial Fibrillation Therapy WO 2015140571, Markers for Atrial Fibrillation WO 2016012783). Kirsten Leineweber is an employee of Bayer AG. The genotyping of participants in the Broad AF Study and the expression analysis of left atrial tissue samples were supported by a grant from Bayer AG to the Broad Institute. The remaining authors have no disclosures.

Atrial fibrillation (AF) affects over 33 million individuals worldwide, ¹ is a common cause of stroke, ² and has a complex heritability. ³ We conducted the largest meta-analysis of genome-wide association studies for AF to date, consisting of over half a million individuals including 65,446 with AF. We performed combined and ancestry-specific meta-analyses as well as conditional and joint analyses. In total, we identified 97 loci significantly associated with AF including 67 of which were novel in a combined-ancestry analysis, and 3 in a European specific analysis. We sought to identify AF-associated genes at the GWAS loci by performing RNA-sequencing and expression quantitative trait loci (eQTL) analyses in 101 left atrial samples, the most relevant tissue for AF. We also performed transcriptome-wide analyses that identified 57 AF-associated genes, 42 of which overlap with GWAS loci. The identified loci implicate genes enriched within cardiac developmental, electrophysiological, and contractile or structural pathways. These results extend our understanding of the biological pathways underlying AF and may facilitate the development of therapeutics for AF.

Atrial fibrillation (AF) is the most common heart rhythm disorder, and is a leading cause of heart failure and stroke.² Prior genome-wide association studies (GWAS) have identified at least 30 loci associated with AF.^{4–9} We conducted a large-scale analysis with over half a million participants, including 65,446 with AF, from more than 50 studies. Our AF sample was composed of 84.2% European, 12.5% Japanese, 2% African American, and 1.3% Brazilian and Hispanic populations (**Supplementary Table S1**). We used the Haplotype Reference Consortium (HRC) reference panel to impute variants from SNP array data for 75% of the samples (**Figure 1**). In the remainder, we included HRC overlapping variants from 1000 Genomes imputed data, or from a combined reference panel. We analyzed 8,328,530 common variants (minor allele frequency (MAF) >5%), 2,884,670 low frequency variants (1%> MAF ≥5%), and 936,779 rare variants (MAF ≤1%).

The combined-ancestry meta-analysis revealed 94 AF-associated loci, 67 of which were novel at genome-wide significance (P-value (P) < 1×10^{-8}). This conservative threshold accounts for testing independent variants with MAF $\geq 0.1\%$ using a Bonferroni correction, while use of a more commonly utilized threshold of 5×10^{-8} resulted in the identification of an additional 10 loci (**Supplementary Table S2**). The majority of sentinel variants (N=92) were common (MAF >5%), with relative risks ranging from 1.04 to 1.55. Two low-frequency sentinel variants were identified within the genes *C1orf185* and *UBE4B* (**Figure 2**, **Table 1**, **Supplementary Table S3**, **Supplementary Figure S1**).

We then conducted a gene set enrichment analysis with the results from the combined-ancestry meta-analysis using MAGENTA. We identified 55 enriched gene sets or pathways that largely fall into cardiac developmental, electrophysiological, and cardiomyocyte contractile or structural functional groups (**Supplementary Table S4**). In total, 48 of the 67 novel loci contain one or more genes within 500kb of the sentinel variant that were part of an enriched gene set or pathway (**Supplementary Figure S2**).

Next, we performed ancestry-specific meta-analyses. Among individuals of European ancestry, we identified 3 additional loci associated with AF, each of which had a sub-threshold association ($P < 1 \times 10^{-6}$) in the combined-ancestry meta-analysis. These loci were located close to or within the genes *CDK6*, *EPHA3*, and *GOSR2* (**Supplementary Table S5**, **Supplementary Figure S3-4**). The region most significantly associated with AF in Europeans, Japanese, and African Americans (**Supplementary Figure S5-6**) was on chromosome 4q25, upstream of the gene *PITX2* (**Supplementary Figure S7**). We did not observe significant heterogeneity of effect estimates across ancestries for most associations, suggesting that top genetic susceptibility signals for AF have a relatively constant effect across ancestries (**Table 1**, **Supplementary Table S3**, **Supplementary Figure S8**). The proportion of heritability explained by the loci

from the European ancestry analysis was 42%, compared to previously reported 25%¹⁰ (**Supplementary Table S6**).

In conditional and joint analyses of the European ancestry results, we found 11 loci with multiple, independent AF-associated signals. At a locus centered on a cluster of sodium channel genes, we identified 3 regions that independently associate with AF within SCN10A, SCN5A and a third signal between both genes. At the previously described TBX5 locus, we detected a novel independent signal close to TBX3. Pairwise linkage disequilibrium (LD) estimates between independent variants at both loci were extremely low ($r^2 < 0.03$; Supplementary Table S7).

For 13 AF loci, the sentinel variant or a proxy (r²>0.6) was a missense variant. A missense variant (rs11057401) in *CCDC92* was predicted to be damaging by 4 of 5 *in silico* prediction algorithms (Supplementary Table S8); and was previously associated with coronary artery disease. Since most AF-associated variants reside in non-coding regions we sought to determine if the sentinel variants or their proxies (r²>0.6) fell within regulatory regions in heart tissues based on chromatin states from the Roadmap Epigenomics Consortium. At 64 out of 67 novel loci, variants were located within regulatory elements (Supplementary Table S9). Compared to 1000 Genomes control loci, AF-associated loci were significantly enriched within regulatory elements (Supplementary Figure S9).

We then sought to link risk variants to candidate genes by assessing their effect on gene expression levels. First, since AF often arises from the pulmonary veins and left atrium (LA), we performed RNA sequencing, genotyping, and eQTL analyses in 101 human left atrial samples without structural heart disease from the Myocardial Applied Genomics Network repository. Second, we identified eQTLs from right atrial (RA) and left ventricular (LV) cardiac tissue from the Genotype Tissue Expression (GTEx)

project. Finally, we performed a transcriptome-wide analysis using the MetaXcan¹² method, which infers the association between genetically predicted gene expression and disease risk.

We observed eQTLs to one or more genes at 17 novel loci. Of the 10 eQTLs detected in LA tissue 8 were also detected in RA or LV, with consistent directionality. For example, we observed that rs4484922 was an eQTL for CASQ2 in LA tissue only. Although we detected more AF loci with eQTLs in the RA or LV data, for many of these (n=8) the results pointed to multiple genes per locus (**Supplementary Table S10-12**). LA eQTL studies may facilitate the prioritization of candidate genes, but are currently limited by sample size.

For the transcriptome-wide analyses we used GTEx human atrial and ventricular expression data as a reference. We identified 57 genes significantly associated with AF. Of these, 42 genes were located at AF loci, whereas the remaining 15 were >500 kb from an AF sentinel variant (**Supplementary Table S13**, **Figure 3**). The probable candidate genes at each locus are summarized in **Supplementary Table S12**. For example, at the locus with lead variant rs4484922 we observed results from all downstream analyses pointing towards the nearest gene *CASQ2*, at rs12908437 towards the gene *IGFR1*, and at rs113819537 towards the gene *SSPN*. However, for many loci the evaluation of candidate genes remains challenging.

We then sought to assess the pleiotropic effects of identified AF risk variants. First, we queried the NHGRI-EBI GWAS Catalog to detect associations to other phenotypes (**Supplementary Table S14**). Second, using the UK Biobank, ¹³ we performed a phenome-wide association study (pheWAS) for 12 AF risk factors (**Supplementary Table S15**). As illustrated in **Figure 4**, distinct clusters of variants were associated with AF as well as height, BMI, and hypertension. The pleiotropic effects at rs880315 (*CASZ1*)

to blood pressure¹⁴ and hyptertension¹⁴, also observed in the UK Biobank (hypertension, $P = 2.56 \times 10^{-34}$), might imply a shared molecular pathway between hypertension and AF.

In sum, we identified a total of 97 distinct AF loci from 65,446 AF cases and more than 522,000 referents. In recent pre-publication results, Nielsen et al., reported 111 loci from 60,620 AF cases and more than 970,000 referents, including more than 18,000 AF cases from our prior report. We therefore performed a preliminary meta-analysis for the top loci in non-overlapping participants from these two large efforts with a resulting total of over 93,000 AF cases and more than 1 million referents. In aggregate, we identified at least 134 distinct AF-associated loci (Supplementary Table S16).

Four major themes emerge from the identified AF loci. First, two AF loci contain genes that are primary targets for current antiarrhythmic medications used to treat AF. The *SCN5A* gene encodes a sodium channel in the heart, and the target of sodium-channel-blockers such as flecainide and propafenone. Similarly, *KCNH2* encodes the alpha subunit of the potassium channel complex, the target of potassium-channel-inhibiting medications such as amiodarone, sotalol, and dofetilide. *SCN5A* and *KCNH2* have previously been implicated in AF through GWAS, and candidate gene analysis and family-based studies. The school of the potassium channel complex and family-based studies.

Second, transcriptional regulation appears to be a key feature of AF etiology. *TBX3* and the adjacent gene *TBX5* have been shown to regulate the development of the cardiac conduction system. ¹⁹ Similarly, the *NKX2-5* gene is an early cue for cardiac development and has been associated with congenital heart disease ²⁰ and heart rate ²¹ (**Supplementary Table S14**). Further, reduced function of the transcription factor *PITX2* has been associated with AF, shortening of the left atrial action potential, and with

modulation of sodium channel blocker therapy in the adult left atrium.^{22–24} A transcriptional coregulatory network governed by *TBX5* and *PITX2* has been shown to be critical for atrial development.²⁵

Third, the transcriptome-wide analyses revealed a number of compelling findings. Decreased expression of *PRRX1* associated with AF, a result consistent with findings where reduction of *PRRX1* in zebrafish and stem cell-derived cardiomyocytes was associated with action potential shortening.²⁶ Further, increased expression of *TBX5* and *KCNJ5* associated with AF, a finding consistent with gain-of-function mutations in *TBX5* reported in a family with Holt-Oram syndrome and a high penetrance of AF.²⁷ Similarly, *KCNJ5* encodes a potassium channel that underlies a component of the I_{KACh} current, a channel that is upregulated in AF. Thus, prior studies support both the role of *PRRX1*, *TBX5*, and *KCNJ5* in AF and the observed directionality.

Fourth, many of the novel loci implicate genes that underlie Mendelian forms of arrhythmia syndromes. Mutations in *CASQ2* lead to catecholaminergic polymorphic ventricular tachycardia. ^{28,29} Pathogenic variants in *PKP2* impair cardiomyocyte communication and structural integrity, and are a common cause of arrhythmogenic right ventricular cardiomyopathy. ^{30,31} Mutations in *GJA5*, *KCNH2*, *SCN5A*, *KCNJ2*, *MYH7*, *NKX2-5*, have been mapped in a variety of inherited arrhythmia, cardiomyopathy, or conduction system diseases. ³² Our observations highlight the pleiotropy of variation in genes specifying cardiac conduction, morphology, and function, and underscore the complex, polygenic nature of AF.

In conclusion, we conducted the largest AF meta-analysis to date and report a more than three-fold increase in the number of loci associated with this common arrhythmia. Our results lay the groundwork for functional evaluations of genes implicated by AF risk loci. Our findings also broaden our

for AF.		

understanding of biological pathways involved in AF and may facilitate the development of therapeutics

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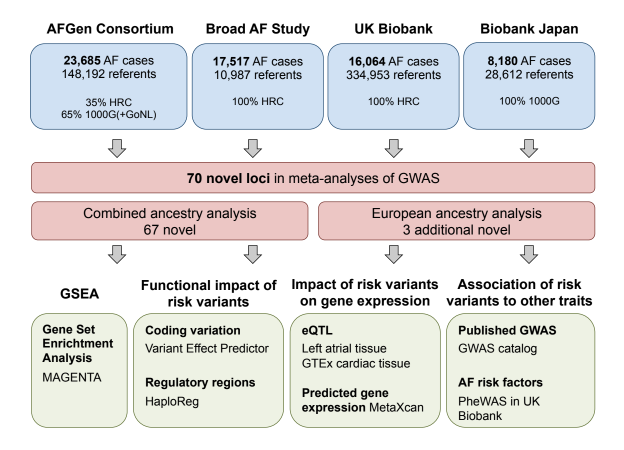
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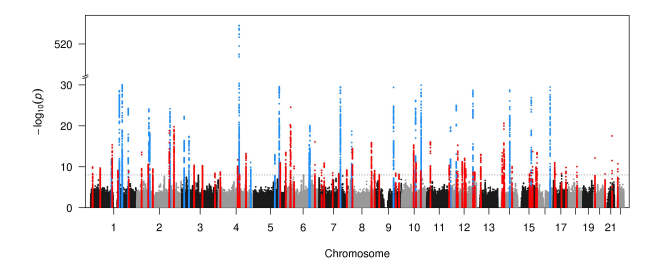
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Figure 1. Study and analysis flowchart



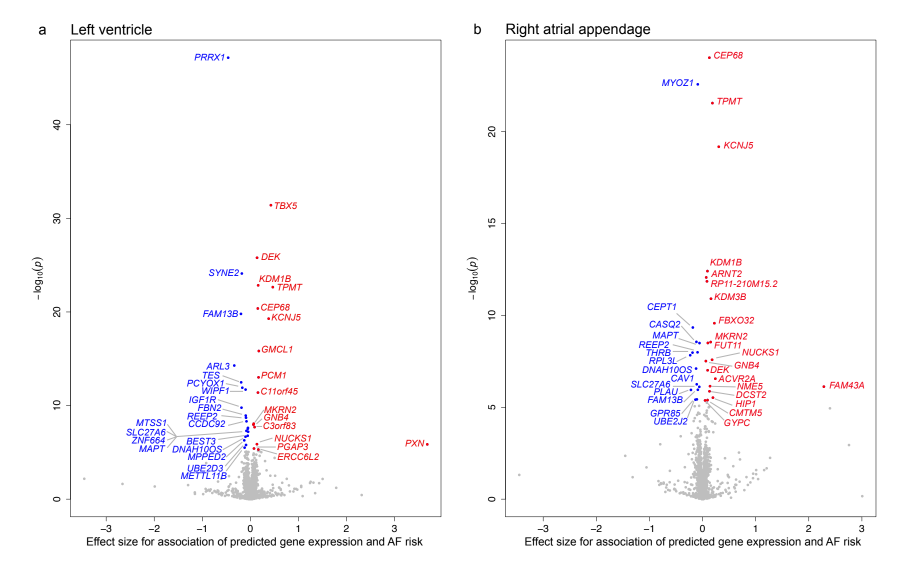
Top, overview of the participating studies, number of AF cases and referents, and the percent of samples imputed with each reference panel. Middle, summary of the primary analyses and the newly discovered loci for AF. Bottom, overview of the secondary analyses to evaluate AF risk variants and loci.

Figure 2. Manhattan plot of combined-ancestry meta-analysis



The plot shows 67 novel (red) and 27 known (blue) genetic loci associated with AF at a significance level of $P < 1 \times 10^{-8}$ (dashed line), for the combined-ancestry meta-analysis. The y-axis has a break between – log10(P) of 30 and 510 to emphasize the novel loci.

Figure 3. Volcano plot of MetaXcan results from human heart tissues.



The plots show the results from MetaXcan based on left ventricle (a, N=190) and right atrial appendage (b, N=159) tissue from GTEx. Each plotted point represents the association results from MetaXcan for an individual gene. The x-axis shows the effect size for MetaXcan associations of predicted gene expression and AF risk for each tested gene. The y-axis shows the $-\log 10(P)$ for the MetaXcan associations per gene. Genes with positive effect (red) showed an association of increased predicted gene expression with AF risk. Genes with negative effect (blue) showed an association of decreased predicted gene expression with AF risk. The highlighted genes are significant after Bonferroni correction for all tested genes and tissues with a $P < 5.36 \times 10^{-6}$. The result for one gene for right atrial appendage (b) is not shown (*SNX4*, Effect = 6.94, P = 0.2).

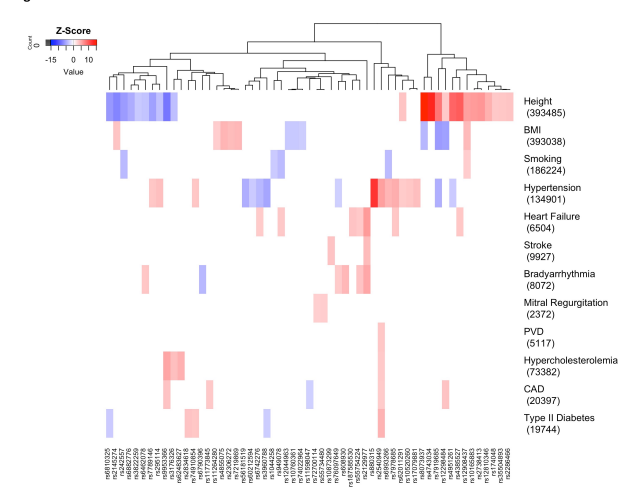


Figure 4. Cross-trait associations of AF risk variants with AF risk factors in the UK Biobank.

The heatmap shows associations of novel and known sentinel variants at AF risk loci from the combined-ancestry meta-analysis. Shown are variants and phenotypes with significant associations after multiple testing, correcting for 12 phenotypes via Bonferroni with P < 4.17x10⁻³. Listed next to the phenotypes on the y-axis are the number of cases for binary traits or total sample size for quantitative traits.

Hierarchical clustering was performed on a variant level using the complete linkage method based on Euclidian distance. The coloring represents the Z-score oriented towards AF risk, red indicates increased value of trait or increased risk for disease in the same direction of AF risk, and blue indicates increased value of trait or increased risk for disease in the opposite direction of AF risk. Abbreviations, BMI, bodymass index, CAD, coronary artery disease, PVD, pulmonary vascular disease.

Table 1. Novel loci in combined-ancestry meta-analysis

Rsid	Chr	hg19	Risk/Ref Allele	RAF [%]	RR	95% CI	P	Nearest Gene(s)	Func	imp Qual	I ² _{HET}	P _{HET}
rs187585530	1	10167425	A/G	0.53	1.55	1.36-1.77	1.18x10 ⁻¹⁰	UBE4B	missense	0.81	0	1.000
rs880315	1	10796866	C/T	37	1.04	1.03-1.06	5.04x10 ⁻⁰⁹	CASZ1	intronic	0.97	40.7	0.150
rs146518726	1	51535039	A/G	3	1.18	1.12-1.24	2.05x10 ⁻¹⁰	C1orf185	intronic	0.96	0	1.000
rs4484922	1	116310818	G/C	68	1.07	1.05-1.08	4.57x10 ⁻¹⁶	CASQ2	intronic	0.98	0	0.689
rs79187193	1	147255831	G/A	95	1.12	1.08-1.16	8.07x10 ⁻¹⁰	GJA5	upstream	0.97	39.8	0.190
rs4951261	1	205717823	C/A	38	1.05	1.03-1.06	1.17x10 ⁻⁰⁹	NUCKS1	intronic	0.99	0	0.788
rs6546620	2	26159940	C/T	75	1.07	1.05-1.09	2.96x10 ⁻¹⁴	KIF3C	intronic	0.95	33	0.201
rs6742276	2	61768745	A/G	61	1.05	1.03-1.06	2.42x10 ⁻¹¹	XPO1	upstream	0.99	0	0.731
rs72926475	2	86594487	G/A	87	1.07	1.05-1.10	3.49x10 ⁻¹⁰	REEP1,KDM3A	intergenic	0.97	38.7	0.180
rs56181519	2	175555714	C/T	74	1.08	1.06-1.10	1.52x10 ⁻¹⁹	WIPF1,CHRNA1	intergenic	0.94	0	0.519
rs295114	2	201195602	C/T	60	1.07	1.05-1.09	1.76x10 ⁻²⁰	SPATS2L	intronic	1.00	21.9	0.275
rs2306272	3	66434643	C/T	32	1.05	1.04-1.07	4.54x10 ⁻¹¹	LRIG1	missense	0.99	30.6	0.218
rs17490701	3	111587879	G/A	86	1.07	1.05-1.10	5.43x10 ⁻¹¹	PHLDB2	intronic	0.97	46.8	0.111
rs4855075	3	179170494	T/C	14	1.06	1.04-1.08	4.00x10 ⁻⁰⁹	GNB4	upstream	0.95	10.1	0.348
rs3822259	4	10118745	T/G	68	1.05	1.03-1.06	1.93x10 ⁻⁰⁹	WDR1	upstream	0.96	0	0.922
rs3960788	4	103915618	C/T	42	1.05	1.04-1.07	2.09x10 ⁻¹²	SLC9B1	intronic	0.98	35.7	0.183
rs55754224	4	114428714	T/C	25	1.05	1.03-1.07	9.25x10 ⁻⁰⁹	CAMK2D	intronic	0.99	0	0.511
rs10213171	4	148937537	G/C	8	1.11	1.08-1.14	6.09x10 ⁻¹⁴	ARHGAP10	intronic	0.96	0	0.584
rs174048	5	142650404	C/T	16	1.07	1.05-1.09	1.05x10 ⁻¹¹	ARHGAP26,NR3C1	intergenic	0.99	0	0.852
rs6882776	5	172664163	G/A	67	1.06	1.05-1.08	3.18x10 ⁻¹⁴	NKX2-5	upstream	0.95	0	0.858
rs73366713	6	16415751	G/A	86	1.11	1.09-1.14	5.80x10 ⁻²¹	ATXN1	intronic	0.94	0	0.879
rs34969716	6	18210109	A/G	31	1.09	1.07-1.11	2.91x10 ⁻²⁵	KDM1B	intronic	0.80	19.5	0.290
rs3176326	6	36647289	G/A	80	1.06	1.04-1.08	7.95x10 ⁻¹¹	CDKN1A	intronic	0.95	0	0.450
rs117984853	6	149399100	T/G	9	1.12	1.09-1.15	8.38x10 ⁻¹⁷	UST	downstream	0.83	56.5	0.100
rs55734480	7	14372009	A/G	27	1.05	1.03-1.07	7.34x10 ⁻¹⁰	DGKB	intronic	0.94	0	0.441
rs6462078	7	28413187	A/C	75	1.06	1.04-1.08	1.35x10 ⁻¹¹	CREB5	intronic	0.98	22.2	0.278
rs74910854	7	74110705	G/A	7	1.10	1.07-1.13	3.36x10 ⁻⁰⁹	GTF2I	intronic	0.74	24.4	0.265

rs62483627	7	106856002	A/G	24	1.05	1.03-1.07	5.17x10 ⁻⁰⁹	COG5	intronic	0.98	15.1	0.318
rs7789146	7	150661409	G/A	80	1.06	1.04-1.08	6.51x10 ⁻¹⁰	KCNH2	intronic	0.96	66	0.0193
rs7846485	8	21803735	C/A	87	1.09	1.07-1.12	3.71x10 ⁻¹⁵	XPO7	intronic	0.99	0	0.676
rs62521286	8	124551975	G/A	7	1.13	1.10-1.16	1.24x10 ⁻¹⁶	FBXO32	intronic	0.96	0	0.678
rs35006907	8	125859817	A/C	33	1.05	1.03-1.06	2.76x10 ⁻⁰⁹	MTSS1,LINC00964	regulatory reg.	0.97	0	0.542
rs6993266	8	141762659	A/G	54	1.05	1.03-1.06	9.73x10 ⁻¹⁰	PTK2	intronic	0.99	5.7	0.374
rs4977397	9	20235004	A/G	57	1.04	1.03-1.06	8.60x10 ⁻⁰⁹	SLC24A2,MLLT3	intergenic	0.95	38.3	0.166
rs4743034	9	109632353	A/G	23	1.05	1.03-1.07	3.98x10 ⁻⁰⁹	ZNF462	intronic	1.00	0	0.963
rs10760361	9	127178266	G/T	65	1.04	1.03-1.06	7.03x10 ⁻⁰⁹	PSMB7	upstream	0.97	0	0.680
rs7919685	10	65315800	G/T	53	1.06	1.04-1.07	5.00x10 ⁻¹⁶	REEP3	intronic	1.00	49.2	0.0965
rs11001667	10	77935345	G/A	22	1.06	1.05-1.08	1.06x10 ⁻¹¹	C10orf11	intronic	0.98	26.8	0.243
rs1044258	10	103605714	T/C	66	1.05	1.03-1.06	1.07x10 ⁻⁰⁹	C10orf76	3' UTR	0.98	14	0.325
rs1822273	11	20010513	G/A	27	1.07	1.05-1.09	8.99x10 ⁻¹⁷	NAV2	intronic	0.98	0	0.764
rs949078	11	121629007	C/T	27	1.05	1.04-1.07	4.77x10 ⁻¹¹	SORL1,MIR100HG	intergenic	0.97	0	0.600
rs113819537	12	26348429	C/G	74	1.05	1.03-1.07	2.23x10 ⁻⁰⁹	SSPN	upstream	0.98	0	0.597
rs12809354	12	32978437	C/T	15	1.08	1.06-1.11	5.48x10 ⁻¹⁶	PKP2	intronic	0.97	31.5	0.211
rs7978685	12	57103154	T/C	28	1.06	1.04-1.07	5.99x10 ⁻¹²	NACA	downstream	0.98	2.4	0.393
rs35349325	12	70097464	T/C	54	1.05	1.04-1.07	9.04x10 ⁻¹³	BEST3	upstream	0.96	0	0.863
rs11180703	12	76223817	G/A	56	1.05	1.03-1.06	3.58x10 ⁻¹⁰	KRR1,PHLDA1	intergenic	0.97	0	0.482
rs12810346	12	115091017	T/C	15	1.07	1.05-1.09	2.34x10 ⁻⁰⁹	TBX5-AS1,TBX3	intergenic	0.84	0	0.428
rs12298484	12	124418674	C/T	67	1.05	1.03-1.06	2.05x10 ⁻⁰⁹	DNAH10	intronic	1.00	0	0.973
rs9580438	13	23373406	C/T	32	1.06	1.04-1.07	1.01x10 ⁻¹³	LINCO0540,BASP1P1	intergenic	0.98	0	0.485
rs28631169	14	23888183	T/C	20	1.07	1.05-1.09	3.80x10 ⁻¹⁴	MYH7	intronic	0.97	14.5	0.319
rs2145587	14	32981484	A/G	28	1.08	1.06-1.10	2.32x10 ⁻²¹	AKAP6	intronic	0.94	0	0.888
rs73241997	14	35173775	T/C	16	1.07	1.05-1.10	1.10x10 ⁻¹³	SNX6,CFL2	intergenic	0.98	62.2	0.0318
rs10873299	14	77426711	A/G	38	1.05	1.03-1.07	9.62x10 ⁻¹¹	LRRC74,IRF2BPL	intergenic	0.96	4.4	0.381
rs62011291	15	63800013	G/A	23	1.05	1.04-1.07	6.14x10 ⁻⁰⁹	USP3	intronic	0.96	0	0.727
rs12591736	15	70454139	G/A	82	1.06	1.04-1.08	2.47x10 ⁻⁰⁹	TLE3,UACA	intergenic	0.92	0	0.966
rs12908004	15	80676925	G/A	16	1.08	1.06-1.10	1.95x10 ⁻¹⁴	LINCO0927,ARNT2	intronic	0.96	57.4	0.0520
rs12908437	15	99287375	T/C	39	1.05	1.03-1.06	1.25x10 ⁻¹⁰	IGF1R	intronic	0.98	0	0.818
rs2286466	16	2014283	G/A	81	1.07	1.05-1.09	3.53x10 ⁻¹⁴	RPS2	synonymous	0.92	0	0.882

rs8073937	17	7435040	G/A	37	1.05	1.04-1.07	1.02x10 ⁻¹¹	POLR2A,TNFSF12	intergenic	0.96	12.3	0.335
rs72811294	17	12618680	G/C	89	1.07	1.05-1.09	6.87x10 ⁻⁰⁹	MYOCD	intronic	0.95	32.3	0.206
rs242557	17	44019712	G/A	61	1.04	1.03-1.06	4.35x10 ⁻⁰⁹	MAPT	intronic	0.94	62.1	0.0319
rs7219869	17	68337185	G/C	44	1.05	1.03-1.06	1.49x10 ⁻¹⁰	KCNJ2,CASC17	intergenic	0.99	16.1	0.312
rs9953366	18	46474192	C/T	66	1.05	1.04-1.07	9.03x10 ⁻¹¹	SMAD7	intronic	0.93	0	0.565
rs2145274	20	6572014	A/C	91	1.11	1.08-1.14	6.97x10 ⁻¹³	CASC20,BMP2	regulatory reg.	0.96	19	0.295
rs7269123	20	61157939	C/T	58	1.05	1.03-1.06	5.59x10 ⁻⁰⁹	C20orf166	intronic	0.85	68.7	0.0123
rs2834618	21	36119111	T/G	90	1.12	1.09-1.14	2.93x10 ⁻¹⁸	LOC100506385	intronic	0.93	21.6	0.277
rs465276	22	18600583	G/A	61	1.05	1.04-1.07	1.84x10 ⁻¹¹	TUBA8	intronic	0.90	0	0.654

Abbreviations, Chr, chromosome, CI, confidence interval, Func, functional consequence (most severe consequence by variant effect predictor), HET,

heterogeneity, I², I-square, impQual, average imputation quality, P, P-value, RAF, risk allele frequency, reg, region, RR, relative risk.

Methods

Samples

Participants from more than 50 studies were included in this analysis. Participants were collected from both case-control studies for atrial fibrillation (AF) and population based studies. The majority of studies were part of the Atrial Fibrillation Genetics (AFGen) consortium and the Broad AF Study (Broad AF). Additional summary level results from the UK Biobank (UKBB) and the Biobank Japan (BBJ) were included (Figure 1). Cases include participants with paroxysmal or permanent atrial fibrillation, or atrial flutter, and referents were free of these diagnoses. Adjudication of atrial fibrillation for each study is described in the Supplementary Notes. Ascertainment of AF in the UK Biobank includes samples with one or more of the following codes 1) Non-cancer illness code, self-reported (1471, 1483), 2) Operation code (1524), 3) Diagnoses – main/secondary ICD10 (148, 148.0-4, 148.9), 4) Underlying (primary/secondary) cause of death: ICD10 (I48, I48.0-4, I48.9) 5) Diagnoses - main/secondary ICD9 (4273), 6) Operative procedures – main/secondary OPCS (K57.1, K62.1-4). ¹⁻³ Baseline characteristics for each study are reported in **Supplementary Table S17**. We analyzed: 55,114 cases and 482,295 referents of European ancestry, 1,307 cases and 7,660 referents of African American ancestry, 8,180 cases and 28,612 referents of Japanese ancestry, 568 cases and 1,096 referents from Brazil and 277 cases and 3,081 referents of Hispanic ethnicity. Samples from the UK Biobank, the Broad AF Study, and the following studies from the AFGen consortium: SiGN, EGCUT, PHB and the Vanderbilt Atrial Fibrillation Registry, were previously not included in primary AF GWAS discovery analyses. There is minimal sample overlap from the studies MGH AF, BBJ and AFLMU between this and previous analyses. Ethics approval for participation was obtained individually by each study.

Genotyping and Genotype Calling

Samples within the Broad AF Study were genotyped at the Broad Institute using the Infinium

PsychArray-24 v1.2 Bead Chip. They were genotyped in 19 batches, grouped by origin of the samples
and with a balanced case control mix on each array. Common variants (≥1% MAF) were called with
GenomeStudio v1.6.2.2 and Birdseed v1.33,⁴ while rare variants (<1% MAF) were called with zCall.⁵

Batch specific quality control (QC) was performed on each call-set including >95% sample call rate,
Hardy-Weinberg-Equilibrium (HWE) P > 1x10⁻⁶ and variant call-rate >97%. For common variants, a
consensus merge was performed between the call-sets from GenomeStudio and Birdseed. For each
genotype only concordant calls between the two algorithms were kept. The common variants from the
consensus call were then combined with the rare variants calls from the zCall algorithm. Samples from
all batches were joined prior to performing pre-imputation QC steps. Detailed procedures for
genotyping and genotype calling for the SiGN study,⁶ the UK Biobank,^{7,8} and the Biobank Japan⁶ are
described elsewhere. Details on genotyping and calling for all participating studies are listed in

Supplementary Table S18.

Imputation

Pre-imputation QC filtering of samples and variants was conducted based on recommended guidelines as described in **Supplementary Table S19.** QC steps were performed by each study and are described in **Supplementary Table S18.** Most studies with European ancestry samples performed imputation with the HRC reference v1.1¹⁰ panel on the Michigan Imputation Server.¹¹ Studies without available HRC imputation were included based on imputation to the 1000 Genomes Phase 1 integrated v3 panel (March 2012).¹² Participants of the SiGN study were imputed to a combined reference panel consisting of 1000 Genomes phase 1 plus Genome of the Netherlands.¹³ Studies from Brazil were imputed with the HRC reference v1.1 panel. Studies of Japanese ancestry or Hispanic ethnicity were imputed to the 1000G

Phase 1 integrated v3 panel (March 2012). Studies of African American ancestry were imputed to the HRC reference v1.1 panel or the 1000G Phase 1 integrated v3 panel (March 2012). Studies were advised to use the HRC preparation and checking tool (http://www.well.ox.ac.uk/~wrayner/tools/) prior to imputation. Prephasing and imputation methods for each study are described in **Supplementary Table S18**.

Primary statistical analyses

Genome-wide association testing on autosomal chromosomes was performed using an additive genetic effect model based on genotype probabilities. Each ancestry group was analyzed separately for each study. For the Broad AF Study, the primary statistical analysis was performed jointly on unrelated individuals, excluding one of each pair for related samples with PI HAT >0.2 as calculated in PLINK v1.90.^{14,15} Samples with sex mismatches and sample call rate <97% were excluded. Ancestry groups were defined with ADMIXTURE¹⁶ based on genotyped, independent, and high quality variants, using the supervised method with 1000Genomes phase 1 v3 samples as reference. A cutoff of 80% European ancestry was used to define the European subset and a cutoff of 60% African ancestry was used to define the African American subset. A Brazilian cohort within the Broad AF Study was analyzed separately. Principal components were calculated within each ancestry group with EIGENSTRAT¹⁷. For the UK Biobank, a European subset was selected within samples with self-reported white race (British, Irish, or other) and similar genetic ancestry. Genetic similarity was defined with the aberrant package in R¹⁸ based on principal components, following the same method as described for the UK Biobank. We excluded samples with sex mismatches, outliers in heterozygosity and missing rates, samples that carry sex chromosome configurations other than XX or XY, and samples that were excluded from the kinship inference procedure as flagged in the UK Biobank QC file. We further removed one sample for each pair of third degree or closer relatives (kinship coefficient >0.0442), preferentially keeping samples with AF

case status. Primary analyses for all other studies were performed at the study sites and the summary level data of the results were provided. Prevalent cases were analyzed in a logistic regression model and most incident cases were analyzed in a Cox proportional hazards model. Studies with both prevalent and incident cases analyzed these either separately using a logistic regression model or Cox proportional hazards model respectively, or jointly in a logistic regression model (Supplementary Table S18). Summary level results were filtered, keeping variants with imputation quality >0.3 and MAF * imputation quality * N events \geq 10. Post-imputation QC steps were also performed, which included a check of allele frequencies, inspection of Manhattan-plots, QQ-plots, PZ-plots, and the distribution of effect estimates and standard errors, calculation of genomic inflation (λ_{GC}), and consistent directionality for known AF risk variants. ¹⁹

Meta-analyses

Summary level results were meta-analyzed jointly with METAL using a fixed effect model with inverse-variance weighted approach, correcting for genomic control. ²⁰ Separate meta-analyses were conducted for each ancestry. The results for the Japanese ⁹ and Hispanic ¹ specific analyses have previously been reported and were not included as separate results. Variants were included if they were present in at least two studies and showed an average MAF \geq 0.1%. To correct for multiple testing, a genome-wide significance threshold of P < 1×10^{-8} was applied for each analysis. This threshold is based on a naive Bonferroni correction for independent variants with MAF \geq 0.1%, using an LD threshold of $r^2 <$ 0.8 to estimate the number of independent variants based on European ancestry LD. ²¹ As these meta-analyses are based on effect estimates and standard errors from both logistic regression and Cox proportional hazards regression, we report variant effects as relative risk, calculated as the exponential of effect estimates. For sentinel variants reaching genome-wide significance in the combined ancestry meta-analysis, we assessed if effect estimates were homogeneous across ancestries by calculating an I^2

statistic²² across ancestry specific meta-analyses. We account for multiple testing across 94 variants using a Bonferroni correction, resulting in a significance threshold of $P < 5.32 \times 10^{-4}$ for the heterogeneity test.

Broad AF LD reference and proxies

A linkage disequilibrium (LD) reference file was created including 26,796 European ancestry individuals from the Broad AF study. The LD reference was based on HRC imputed genotypes. Monomorphic variants and variants with imputation quality <0.1 were removed prior to conversion to hard calls. A genotype probability (GP) threshold filter of GP >0.8 was applied during hard call conversion. For multiallelic sites the more common alleles were kept. Variants were included in the final reference file if the variant call rate was >70%.

We identified proxies of sentinel variants as variants in LD of $r^2 > 0.6$ based on the Broad AF LD reference file, using PLINK v1.90. ^{14,15}

Meta-analysis of provisional loci

We meta-analyzed 111 variants from externally reported 23 provisional loci within predominantly non-overlapping samples from the Broad AF Study, BBJ, EGCUT, PHB, SiGN and the Vanderbilt AF Registry with METAL. The predominantly non-overlapping samples included a total of 32,957 AF cases and 83,546 referents, with minimal overlap from the studies MGH AF, BBJ and AFLMU. We subsequently meta-analyzed these results with the reported provisional results with METAL using a fixed effect model with inverse-variance weighted approach. We analyzed a total of 93,577 AF cases and 1,053,762 referents. We compared our discovery results with the provisional loci using the same significance cutoff of P < 5×10^{-8} for both results. Overlapping loci were identified, if the reported sentinel variants were

located within 500KB of each other. For overlapping loci with differing sentinel variants we calculated the LD between the sentinel variants, based on the Broad AF LD reference panel of European ancestry.

Variant consequence on protein coding sequence

The most severe consequence for variants was identified with the Ensembl Variant Effect Predictor version 89.7 using RefSeq as gene reference and the option "pick" to identify one consequence per variant with the default pick order. We queried sentinel variants and their proxies to identify tagged variants with HIGH and MODERATE impact including the following consequences: "transcript_ablation", "splice_acceptor_variant", "splice_donor_variant", "stop_gained", "frameshift_variant", "stop_lost", "start_lost", "transcript_amplification", "inframe_insertion", "inframe_deletion", "missense_variant" and "protein_altering_variant". We evaluated each identified consequence on the protein coding sequence with in silico prediction tools to assess potentially damaging effects. The evaluation included MutationTaster (disease causing automatic or disease causing), SIFT (damaging), LRT (deleterious), Polyphen2 prediction based on HumDiv and HumVar (probably damaging or possibly damaging).

Chromatin states

- 1) Chromatin state annotation. We identified chromatin states for sentinel variants and their proxies from the Roadmap Epigenomics Consortium 25-state model (2015)²⁹ using HaploReg v4.³⁰ We looked for chromatin states occurring in any included tissues as well as chromatin states occurring in heart tissue. Heart tissues include E065: Aorta, E083: Fetal Heart, E095: Left Ventricle, E104: Right Atrium and E105: Right Ventricle.
- **2) Regulatory region enrichment.** 1,000 sets of control loci were generated by matching SNPs to sentinel variants from the AF combined-ancestry analysis, with the SNPSnap³¹ tool. We used the European 1000 Genomes Phase 3 population to match via minor allele frequency, gene density, distance

to nearest gene and LD buddies using $r^2 > 0.6$ as LD cutoff and otherwise default settings. We excluded input SNPs and HLA SNPs from the matched SNPs. Loci were defined as SNPs and their proxies with $r^2 > 0.6$ based on LD from the European 1000 Genomes Phase 3 population. We identified SNPs in regulatory regions across all tissues and in cardiac tissues (E065, E095, E104, E105) based on the Roadmap Epigenomics Consortium 25-state model (2015)²⁹ using HaploReg v4.³⁰ Regulatory regions included the following states: 2_PromU, 3_PromD1, 4_PromD2, 9_TxReg, 10_TxEnh5, 11_TxEnh3, 12_TxEnhW, 13_EnhA1, 14_EnhA2, 15_EnhAF, 16_EnhW1, 17_EnhW2, 18_EnhAc, 19_DNase, 22_PromP and 23_PromBiv. We calculated the percent overlap of each annotation per locus, defined as number of SNPs per locus that fall in regulatory regions divided by total number of SNPs per locus. Statistical significance was calculated with a permutation test from the *perm* package in R.

Expression quantitative trait loci (eQTL)

Variants identified from GWAS were assessed for overlap with eQTLs from two sources:

1) Left atrial (LA) tissue from the Myocardial Applied Genomics Network (MAGNet) repository. We performed RNA sequencing (RNA-seq) on 101 left atrial tissue samples from the MAGNet repository (http://www.med.upenn.edu/magnet/) on the Illumina HiSeq 4000 platform at the Broad Institute Genomic Services. Left atrial tissue was obtained at the time of cardiac transplantation from normal donors with no evidence of structural heart disease. All left atrial samples were from individuals of European ancestry. A summary of the clinical characteristics for these samples is shown in Supplementary Table S20. Reads were aligned to the reference genome by STAR³² and assigned to genes based on the GENCODE gene annotation. ³³ Gene expression was measured in fragments per kilobase of transcript per million mapped reads (FPKM) and subsequently quantile-normalized and adjusted for age, sex, and the first 10 principal components. Genotyping was performed on the Illumina OmniExpressExome-8v1 array and imputed to the HRC reference panel. Principal components were

calculated with the smartpca program from EIGENSOFT¹⁷ and European ancestry was confirmed by assessing principal components in the samples combined with 1000 Genomes European samples.¹²
Associations between gene expression and genotypes were tested in a linear regression model with QTLtools,³⁴ in order to detect *cis*-eQTLs, defined as eQTLs within 1MB of the transcription start site of a gene. To account for multiple testing, a false discovery rate (FDR) was used to identify significant eQTLs with a FDR <5%.

2) Genotype-Tissue Expression (*GTEx*) **project.**³⁵ We queried the GTEx version 6p database for cis-eQTLs with significant associations to gene expression levels in the two available heart tissues: left ventricle and right atrial appendage.³⁶

Association between predicted gene expression and risk of atrial fibrillation

To investigate the association between predicted gene expression and AF disease risk, we employed the method MetaXcan.³⁷ MetaXcan extends the previous method PrediXcan³⁸ to predict the association between gene expression and a phenotype of interest, using summary association statistics. Gene expression prediction models were generated from eQTL datasets using Elastic-Net to identify the most predictive set of SNPs. Only models that significantly predict gene expression in the reference eQTL dataset (FDR <0.05) were considered. Pre-computed MetaXcan models for the two available heart tissues (left ventricle and right atrial appendage) in the genotype-tissue expression project version 6p (GTEx)³⁶ were used to predict the association between gene expression and risk of AF. Summary level statistics from the combined ancestry meta-analysis were used as input. 4859 genes were tested for left ventricle and 4467 genes were tested for right atrial appendage. Bonferroni correction was applied to account for the number of genes tested across both tissues, resulting in a significance threshold of P < 5.36x10⁻⁶, calculated as 0.05/(4859 + 4467).

Conditional and joint analyses

Conditional and joint analyses³⁹ of GWAS summary statistics were performed with Genome-wide Complex Trait Analysis (GCTA)⁴⁰ using a stepwise selection procedure to identify independently-associated variants on each chromosome. We used the Broad AF LD reference file for LD calculations.

Gene set enrichment analysis (GSEA)

A Meta-Analysis Gene-set Enrichment of variaNT Associations (MAGENTA) 2.4⁴¹ was performed with a combined gene set input database (GO_PANTHER_INGENUITY_KEGG_REACTOME_BIOCARTA) based on publicly available data. The analysis was conducted using the summary level results from the combined ancestry meta-analysis. 4046 gene sets were included and multiple testing was corrected via false discovery rate (FDR). Gene sets were manually assigned to one or more of the following functional groups: developmental, electrophysiological, contractile/structural, and other. Genes within 500 kilobases of a sentinel variant were identified based on the longest spanning transcribed region in the RefSeq gene reference. For each gene set, genes close to significant loci were listed. The selected genes were assigned to one or more functional groups based on their affiliation to gene sets. Functional groups from gene sets with a single label were preferentially assigned.

Association with other phenotypes

To determine if the sentinel AF risk variants had associations with other phenotypes, two sources of data were used:

1) GWAS catalog. We queried the NHGRI-EBI Catalog of published genome-wide association studies ^{42,43} (accessed 2017-08-31) to detect associations of AF risk variants with other phenotypes.

2) UK Biobank phenome-wide association study (PheWAS). A PheWAS was conducted in the UK Biobank in European ancestry individuals. Ancestry definition and sample QC exclusions were performed in the same manner as for the primary statistical analysis, as described above. We further removed one sample for each pair of second degree or closer relatives (kinship coefficient >0.0884), preferentially keeping the sample with case status or non-missing phenotype. We included the following phenotypes: height, body mass index (BMI), smoking, hypertension, heart failure, stroke, mitral regurgitation, bradyarrhythmia, peripheral vascular disease (PVD), hypercholesterolemia, coronary artery disease (CAD), and type II diabetes. Phenotype definitions are shown in Supplementary Table S21. Number of samples analyzed, as well as case and referents counts for each phenotype are listed in Supplementary Table S22. Binary phenotypes were analyzed with a logistic regression model and quantitative phenotypes with a linear regression model using imputed genotype dosages in PLINK 2.00.¹⁵ As covariates we included sex, age at first visit, genotyping array, and the first 10 principal components.

Proportion of heritability explained

We calculated SNP-heritability (h_g^2) of AF-associated loci with the REML algorithm in BOLT-LMM v2.2⁴⁴ in 120,286 unrelated samples of European ancestry from a subset of the UK Biobank dataset comprising a prior interim release as previously described in separate work from our group.² We defined loci based on a 1MB (+/- 500KB) window around 84 sentinel variants from the European ancestry meta-analysis. We transformed the h_g^2 estimates into liability scale (AF prevalence = 2.45% in UK Biobank). We then calculated the proportion of h_g^2 explained at AF loci by dividing the h_g^2 estimate of AF-associated loci by the total h_g^2 for AF, that was based on 811,488 LD-pruned and hard-called common variants (MAF \geq 1%).²

Data availability

The datasets generated during and/or analyzed during the current study are available from the

corresponding author on reasonable request. The summary level results of this study are available on the Cardiovascular Disease Knowledge Portal (http://www.broadcvdi.org/).

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