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EPIDEMIOLOGICAL SCIENCE

Genome-wide association study identifies *RNF123* locus as associated with chronic widespread musculoskeletal pain

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ABSTRACT

Background and objectives Chronic widespread musculoskeletal pain (CWP) is a symptom of fibromyalgia and a complex trait with poorly understood pathogenesis. CWP is heritable (48%–54%), but its genetic architecture is unknown and candidate gene studies have produced inconsistent results. We conducted a genome-wide association study to get insight into the genetic background of CWP.

Methods Northern Europeans from UK Biobank comprising 6914 cases reporting pain all over the body lasting >3 months and 242 929 controls were studied. Replication of three independent genomewide significant single nucleotide polymorphisms was attempted in six independent European cohorts (n=43 080; cases=14 177). Genetic correlations with risk factors, tissue specificity and colocalisation were examined.

Results Three genome-wide significant loci were identified (*rs1491985*, *rs10490825*, *rs165599*) residing within the genes *Ring Finger Protein 123* (*RNF123*), *ATPase secretory pathway Ca²⁺ transporting 1* (*ATP2C1*) and *catechol-O-methyltransferase* (*COMT*). The *RNF123* locus was replicated (meta-analysis p=0.0002), the *ATP2C1* locus showed suggestive association (p=0.0227) and the *COMT* locus was not replicated. Partial genetic correlation between CWP and depressive symptoms, body mass index, age of first birth and years of schooling were identified. Tissue specificity and colocalisation analysis highlight the relevance of skeletal muscle in CWP.

Conclusions We report a novel association of *RNF123* locus and a suggestive association of *ATP2C1* locus with CWP. Both loci are consistent with a role of calcium regulation in CWP. The association with *COMT*, one of the most studied genes in chronic pain field, was not confirmed in the replication analysis.

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Chronic widespread musculoskeletal pain (CWP) is a common complex trait influenced by genetic and environmental factors, most of which have yet to be determined.¹ CWP and fibromyalgia syndrome are sometimes used interchangeably, although the latter is generally more severe and includes other features such as sleep disturbance, fatigue and depression.² It is thought to represent a subgroup at the more

Key messages

What is already known about this subject?

- Chronic widespread musculoskeletal pain (CWP) is a primary diagnostic feature of fibromyalgia.
- CWP is moderately heritable, but precise genes involved in the pathogenesis of CWP are yet to be identified.

What does this study add?

- This is the largest genetic study conducted on CWP to date and identified novel genetic risk loci (*Ring Finger Protein 123* and *ATPase* secretory pathway Ca²⁺ transporting 1).
- The genetic signal points to peripheral pain mechanisms in CWP, and shows genetic correlation with other traits, including body mass index and depression.

How might this impact on clinical practice or future developments?

The findings add to aetiological basis of CWP.

severe end of the spectrum of CWP.³ The prevalence of CWP is 10.6% in the world population and 14.2% in the UK population.^{4.5} It is associated with high societal cost.⁶ CWP is responsible for excess mortality,⁷ which is thought to be attributable to cardiovascular disease, respiratory disease and cancer. Females are more affected by CWP than males,⁴ and the prevalence rises with age.⁵ In addition to age and sex, a number of exposures have been proposed as risk factors for CWP,^{8 9} but only increased body mass index (BMI) has been consistently reported across studies, including longitudinal studies.^{10–12}

Broad-sense heritability estimates for CWP range between 48% and 54%, indicating a substantial genetic contribution.¹³ To date, the candidate gene approach has been extensively applied to identify genetic factors in CWP,¹⁴ but few agnostic studies have been published.¹⁵ The only genome-wide association study (GWAS) meta-analysis combining 14 studies identified a locus lying on chromosome 5 intergenic to *CCT5* and *FAM173B*.¹⁵ *CCT5* has previously been implicated in neuropathy¹⁶ and



eular¹²²⁷



Figure 1 Overview of study design.

there is increasing evidence that small fibre neuropathy underlies a subset of fibromyalgia.¹⁷

Genetic factors are known to be shared by chronic pain conditions.¹⁸¹⁹ One of the most extensively studied chronic pain-associated genes encodes catechol-O-methyltransferase (COMT), an enzyme which regulates the production of catecholamines that act as neurotransmitters in the central nervous system (CNS) pain tract. A non-synonymous change of A to G encoding a valine (Val) to methionine (Met) substitution at codon 158 (Val158Met; rs4680) reduces the enzymatic activity of COMT. This single nucleotide polymorphism (SNP) has been reported to be associated with CWP in a small study of 122 participants,²⁰ but a subsequent association study of 3017 participants did not confirm earlier findings.²¹ An inconclusive role of COMT was observed for temporomandibular disorders (TMD) as well.^{22 23} Further investigation is required to identify genetic variants underlying CWP, which will shed light on the pathophysiological mechanisms underlying the development of chronic pain and may reveal therapeutic targets.

MATERIALS AND METHODS

An overview of study design is presented in figure 1.

Participant selection

For the discovery analysis, we performed a GWAS of CWP using UK Biobank (UKB) comprising 249 843 participants of European descent (6914 CWP cases and 242 929 controls). Independent SNPs passing a threshold p<5.0E-08 were submitted for replication in 43 080 individuals of European ancestry (14 177 CWP cases and 28 903 controls) from six independent cohorts originating in the UK (TwinsUK and The English Longitudinal Study of Ageing (ELSA)), the Netherlands (The Rotterdam Study 1, 2 and 3 (RS-1, RS-2 and RS-3)) and Norway (The Nord-Trøndelag Health Survey (HUNT)). The UKB dataset was used under project #18219. Description of each study cohort is presented in online supplemental text.

Phenotype

In UKB, CWP cases were defined by combining self-reported diagnosis of pain all over the body lasting for >3 months; simultaneous pain in the knee, shoulder, hip and back lasting 3+ months and fibromyalgia. Controls comprised those who reported no pain in the last month or reported pain all over the body in the previous month that did not last for 3 months or reported only ≥ 3 months of non-musculoskeletal pain (head-ache, facial and abdominal pain). Those reporting a self-reported

 Table 1
 Sample characteristics stratified by case/control status for discovery and replication cohorts

	Cases	Controls	P value	
Discovery cohort (UK Bio	bank)			
Female	4470 (64.7%)	128 599 (47.1%)	<0.0001	
Male	2444 (35.3%)	114330 (52.9%)		
Age (mean±SD)	57.8±7.45	57.0±8.09	<0.0001	
BMI (mean±SD)	30.02±5.97	26.83±4.40	< 0.0001	
Replication cohorts				
TwinsUK				
Female	1041 (93.7%)	3116 (87.6%)	< 0.0001	
Male	70 (6.3%)	440 (12.4%)		
Age (mean±SD)	54.78±10.48	50.12±13.21	< 0.0001	
BMI (mean±SD)	27.39±5.11	25.74±4.57	< 0.0001	
HUNT				
Female	6315	5836	< 0.0001	
Male	4241	7403		
Age (mean±SD)	55.95±9.48	54.82±10.31	<0.0001	
BMI (mean±SD)	27.37±4.33	26.52±3.88	< 0.0001	
ELSA				
Female	1090 (64.9%)	2660 (50.2%)	< 0.001	
Male	589 (35.1%)	2644 (49.8%)		
Age (mean±SD)	68.10±9.49	66.55±9.98	< 0.0001	
BMI (mean±SD)	28.60±4.98	27.08±4.22	< 0.0001	
RS-1				
Female	422	1323	< 0.0001	
Male	110	1281		
Age (mean±SD)	64.49±5.30	64.60±5.24	0.6660	
BMI (mean±SD)	26.98±3.91	26.14±3.54	< 0.0001	
RS-2				
Female	106	745	< 0.0001	
Male	38	676		
Age (mean±SD)	61.59±4.59	61.93±4.72	0.2651	
BMI (mean±SD)	28.54±4.73	27.77±3.91	0.0363	
RS-3				
Female	128	1516	< 0.0001	
Male	27	1263		
Age (mean±SD)	56.28±5.77	56.32±5.46	0.0348	
BMI (mean+SD)	28 54+4 86)	27 71+4 62	0.0827	

BMI, body mass index; ELSA, The English Longitudinal Study of Ageing; HUNT, The Nord-Trøndelag Health Survey; RS-1, RS-2 and RS-3, The Rotterdam Study 1, 2 and 3; SD, Standard deviation .

diagnosis of rheumatoid arthritis, polymyalgia rheumatica, arthritis not otherwise specified, systemic lupus erythematosus, ankylosing spondylitis and myopathy were excluded from the study (online supplemental figure S1). Further phenotype details for UKB and replication cohorts are provided in online supplemental text.

Genotyping and imputation

Genotyping and imputation methods across cohorts are summarised in online supplemental table S1 (online supplemental text).

Statistical analysis and in silico follow-up

The details of statistical analysis, and in silico follow-up are described in online supplemental text. In brief, GWAS in the discovery sample was performed using linear mixed-effects model implemented in BOLT-LMM (V.2.3.2).²⁴ An additive



Figure 2 Manhattan plot of a genome-wide association analysis of chronic widespread musculoskeletal pain (CWP). Each circle in the plot represents a single nucleotide polymorphism (SNP), which was positioned following genomic build GRCh37. The y-axis shows the corresponding –log10 p values and the x-axis shows chromosome position along with SNPs. The horizontal red dotted line indicates genome-wide significance threshold at $p=5.0\times10^{-8}$. The horizontal blue dotted line indicates suggestive genome-wide significance threshold at $p=5.0\times10^{-7}$. Gene labels represent nearest genes to independent SNPs located at loci associated with $p<5.0\times10^{-7}$.

genetic model for SNP effect on CWP was adjusted for age, sex, genotyping platform and the first 10 genetic principal components provided by UKB. A sensitivity GWAS (controls: 223606 and CWP cases: 6914) was performed excluding participants with chronic non-musculoskeletal pain such as headache, facial and abdominal pain from the controls. Independent SNPs at GWAS significant loci were identified using Conditional and Joint²⁵ analysis and submitted for replication. Independent SNPs across all replication cohorts were meta-analysed using fixed-effects model with both sample size, and inverse-variance weighting implemented in METAL.²⁶ SNP heritability was estimated using BOLT-REML²⁴ and converted to liability scale. Linkage disequilibrium score regression (LDSR)²⁷ was used to estimate inflation in test statistics and genetic correlations. We also estimated partial genetic correlations.²⁸ We used Functional Mapping and Annotation (FUMA) webtool²⁹ for the annotation of functional consequences of CWP-associated SNPs, gene mapping, tissue specificity and gene-set enrichment. Differential expression of replicated independent SNP was assessed using the GTEx V.8 tissues.³⁰ Colocalisation of GWAS-independent SNPs in human skeletal muscle and dorsal root ganglion (DRG) tissues was assessed using publicly available data.^{30 31} Functional annotation of GWAS-replicated locus was performed using Open Targets Platform.³²

RESULTS

Details of the discovery and replication cohorts are presented in table 1. Cases were enriched for females compared with controls in all cohorts (p<0.001) and were on average older in the discovery, and in three replication cohorts (p<0.05). In all cohorts, BMI was significantly higher in cases than controls (p<0.0001) except for RS-3 where a similar but non-significant trend was observed (p=0.0827).

Discovery genome-wide association study

Three genomic loci tagged by *rs1491985*, *rs10490825* and *rs165599* passed genome-wide significance threshold of p<5E-08 (figure 2). Observed inflation in test statistics (λ_{GC} =1.146, online supplemental figure S2) was due to polygenicity (LDSR intercept=1.002±0.0085, LDSR ratio=0.0118±0.0497) rather





Figure 3 Regional plots for three independent chronic widespread musculoskeletal pain associated single nucleotide polymorphisms (SNPs). Independent SNPs are coloured in purple. Other coloured circles indicate pairwise linkage disequilibrium (LD). The strength of LD (r^2) presented in the upper left corner of each plot.

than population stratification. SNP heritability of CWP was 0.05 ± 0.003 on the observed scale, and 0.33 ± 0.0004 on the liability scale meaning that the observed SNPs explain approximately 33% of the variance in CWP risk. Independent SNPs were located in the gene *Ring Finger Protein* 123 (*RNF123*) (chromosome 3, *rs1491985*, intronic variant, p=1.60E-08), *ATPase secretory pathway* Ca²⁺ transporting 1 (*ATP2C1*) (chromosome 3, *rs10490825*, intronic variant, p=1.30E-08) and COMT (chromosome 22, rs165599, 3'-untranslated region (3'-UTR) variant, p=2.50E-08), respectively (figure 3A–C; online supplemental table S2). Six additional loci near or within genes *HNRNPA1P46*, *LRRC3B*, *PDE6A*, *DPYSL2*, *ANXA11* and *AL138498.1* were identified at suggestive GWAS threshold of p<5E-07. Sensitivity GWAS excluding participants with chronic

A rs1491985

				Beta	Beta	
Study or Subgroup	Beta	SE	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI	
HUNT	-0.0107	0.0062	58.1%	-0.01 [-0.02, 0.00]		
ELSA	-0.0199	0.0092	26.4%	-0.02 [-0.04, -0.00]	-	
TwinsUK	-0.0348	0.0122	15.0%	-0.03 [-0.06, -0.01]	-	
RS-1	-0.032	0.0899	0.3%	-0.03 [-0.21, 0.14]		
RS-3	-0.2453	0.1381	0.1%	-0.25 [-0.52, 0.03]		
RS-2	-0.0165	0.1555	0.1%	-0.02 [-0.32, 0.29]		
Total (95% CI)			100.0%	-0.02 [-0.03, -0.01]	•	
Heterogeneity: Chi ² =	6.02, df =	5 (P = 0)				
Test for overall effect	: Z = 3.61	(P = 0.00)	-0.5 -0.25 0 0.25 0.5			

B rs10490825

				Beta	Beta
Study or Subgroup	Beta	SE	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% Cl
HUNT	0.014	0.0067	62.3%	0.01 [0.00, 0.03]	
ELSA	-0.0004	0.011	23.1%	-0.00 [-0.02, 0.02]	+
TwinsUK	0.0091	0.0141	14.1%	0.01 [-0.02, 0.04]	+
RS-1	0.1409	0.099	0.3%	0.14 [-0.05, 0.33]	
RS-3	0.0877	0.1685	0.1%	0.09 [-0.24, 0.42]	
RS-2	0.0058	0.1783	0.1%	0.01 [-0.34, 0.36]	
Total (95% CI)			100.0%	0.01 [0.00, 0.02]	
Heterogeneity: Chi ² =	3.21, df =	5 (P = 0)			
Test for overall effect	Z = 1.97	(P = 0.05)	5)		-0.5 -0.25 0 0.25 0.5

C rs165599

				Beta	Beta	
Study or Subgroup	Beta	SE	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% Cl	
HUNT	0.0026	0.0052	58.6%	0.00 [-0.01, 0.01]		
ELSA	0.0015	0.0077	26.7%	0.00 [-0.01, 0.02]	+	
TwinsUK	0.0066	0.0105	14.4%	0.01 [-0.01, 0.03]	+	
RS-3	-0.0905	0.1303	0.1%	-0.09 [-0.35, 0.16]		
RS-2	-0.0895	0.136	0.1%	-0.09 [-0.36, 0.18]		
RS-1	-0.0042	0.1395	0.1%	-0.00 [-0.28, 0.27]		
Total (95% CI)			100.0%	0.00 [-0.01, 0.01]	• • •	
Heterogeneity: $Chi^2 = 1.14$, $df = 5$ (P = 0.95); $I^2 = 0\%$						
Test for overall effect	: Z = 0.68	(P = 0.50)	-0.5 -0.25 0 0.25 0.5			

Figure 4 Forest plot for the association of (A) *rs1491985*, (B) *rs10490825*, and (C) *rs165599* with chronic widespread musculoskeletal pain. X-axis shows effect size measures are presented as beta value. The red square with horizontal black line represents the cohort-specific effect with a corresponding CI for the single nucleotide polymorphism (SNP) of interest. Size of the square indicates the weight of the study and reflects sample size. The vertical black line indicates 'line of no effect'. Overall effect is presented as a black diamond. Test statistics for each cohort, meta-analysis and heterogeneity are available on the left-hand side. The *rs1491985* and *rs10490825* were not present in The English Longitudinal Study of Ageing (ELSA); therefore *rs9870858* and *rs17329848* were used as proxy SNPs, respectively (online supplemental text).

non-musculoskeletal pain provided similar findings except that COMT locus now became suggestively significant (p=5.3E-08) (online supplemental figure S3).

Replication results and meta-analysis

Results are presented in online supplemental table S3, with meta-analysis of the six replication samples as shown in figure 4 (online supplemental tables S4, S5). Given the significance threshold for replication: 0.05/3=0.017, association between CWP and *rs1491985* was considered replicated (sample-size based p=0.0002; standard-error based p=0.0003). *Rs10490825* showed suggestive association with CWP (sample-size based p=0.0227; standard-error based p=0.0490) and demonstrated a consistent direction of effect in five of the six replication samples. *Rs165599* did not replicate (sample-size based p=0.7300; standard-error based p=0.5000) and the direction of effect was not consistent across cohorts: in three cohorts, allele A was protective, while in the other three it was the risk allele.

None of the three SNPs displayed statistically significant heterogeneity in the replication cohorts.

CWP shares genetic components with BMI, depression, age at first birth and years of schooling

Two hundred and nine traits from LD-hub (online supplemental text) were examined for genetic correlation with CWP. We selected traits for which the absolute value of the correlation coefficient (r_g) was >0.2, and for which the Bonferronicorrected p was <0.01/209=4.78E-05. Twenty-three traits fulfilled these criteria (online supplemental figure S4). The highest positive genetic correlation was observed for depressive symptoms (r_g =0.65) and the highest negative correlation was observed for college completion (r_g =-0.61). Many of the 23 genetically correlated traits were correlated with each other raising concerns about their independency of correlations with CWP. We therefore calculated partial genetic correlations



Figure 5 (A) Manhattan plot of the genome-wide genebased associationanalysis, (B) & (C) The circus plot displaying chromatininteractions (Ci) and expression quantitative trait loci (eQTLs) onchromosomes 3 and chromosomes 22, respectively, (D) Venn diagramshowing overlap of genes implicated by genome-wide gene-basedanalysis implemented in MAGMA, positional mapping (Pos Map), chromatin interaction mapping (Ci Map), and expression quantitativetrait locus mapping (eQTL Map). (A) The y-axis shows the -log10transformed two-tailed p-value of each gene from a linear model and the chromosomal position on the x-axis. The red dotted line indicates the Bonferroni-corrected threshold for genomewide significance of the gene-based test. (B, C) The most outer layer of the circus plotdisplaying Manhattan plot with -log₁₀ p-values forchronic widespread musculoskeletal pain associated independent singlenucleotide polymorphisms (SNPs). Each SNP is presented with rsID.Linkage diseguilibrium (LD) relationship between independent SNPs at he locus and their proxies are indicated with red ($r^2 > 0.8$) and orange ($r^2 > 0.6$). Grey SNPs indicate minimalLD with $r^2 \le 0.20$. The outer circle represents chromosome with genomic risk loci arehighlighted in blue. Either Ci- or eQTL mapped genes are displayed onthe inner circle. Ci- and eQTL mapped genes are presented in orangeor green color, respectively. Genes mapped with both approaches arecolored red.

conditionally independent of each other. Using hierarchical clustering of genetic correlations we identified seven clusters (online supplemental figure S5A), with seven traits selected to represent each cluster (BMI, triglycerides, depressive symptoms, coronary artery disease, smoking, age of first birth and years of schooling) to quantify partial genetic correlation with CWP. We found depressive symptoms ($r_g=0.59$), BMI ($r_g=0.20$), age of first birth ($r_g=-0.26$) and years of schooling ($r_g=-0.17$) independently correlated with CWP (online supplemental figure S5B and table S6).

Tissue-specific expression of CWP mapped gene sets

The results of functional consequences of GWAS-independent SNPs and their proxies are presented in online supplemental figure S6 (online supplemental text). Four different gene mapping strategies were implemented in FUMA (genome-wide gene-based association analysis, positional, expression quantitative trait locus (eQTL) and chromatin interaction mapping) linking annotated SNPs to 89 genes of which *MST1*, *GMPPB*, *APEH*, *RNF123*, *ARVCF*, *AMIGO3*, *IP6K1*, *TANGO2* and *TRAIP* were identified using all four methods (figure 5A–D).³³ Mapped genes were investigated for tissue-specific gene expression and gene-set enrichment. In 54 specific GTEx tissues types, differentially expressed gene sets enriched for skeletal muscle, several brain tissues, heart, whole blood, pancreas and transverse colon (figure 6A, online supplemental table S7). In 30 general GTEx

tissue types, differentially expressed gene sets enriched for skeletal muscle, pancreas, heart, blood and brain (figure 6B, online supplemental table S8). In both sets of GTEx tissues, overall enrichment for differentially expressed gene sets containing *RNF123* and *ATP2C1* genes were stronger for skeletal muscle than other tissues. *RNF123* was found to be highly expressed in skeletal muscle compared with other tissue types (figure 6C). None of the hallmark gene sets available in the molecular signature database was identified in the analysis.

Putative causal genes in RNF123 locus

Colocalisation analysis identified a 93% probability of shared eQTL variant rs6809879, which controls Cadherin Related Family Member 4 (CDHR4) expression in the skeletal muscle and CWP association signal near the RNF123 locus (online supplemental table S9, online supplemental figure S7A). Additionally, significant colocalisation was found for rs13093525, which controls APEH expression in DRG at exon level (72% probability of shared variant with RNF123 locus). Both rs6809879 and rs13093525 were in complete LD with independent SNP rs1491985 (R²=1) (online supplemental table S10, online supplemental figure S7B). No evidence of skeletal muscle or DRG eQTL colocalisation was observed for ATP2C1 and COMT loci. Functional annotation of RNF123 locus identified nine genes (SLC25A20, NDUFAF3, DAG1, HYAL1, GMPPB, TRAIP, RHOA, CACNA2D2 and IMPDH2) specific to musculoskeletal system diseases, of which CACNA2D2, NDUFAF3 and IMPDH2 enriched as druggable targets (online supplemental figure S8).

DISCUSSION

CWP is a prevalent condition with moderate heritability and serves as a cardinal diagnostic feature of fibromyalgia. Therefore, our findings are of importance for better understanding the genetic basis of fibromyalgia. We report here the largest GWAS of CWP to date using 249 843 participants from the UKB, identifying 3 genome-wide significant loci implicating *RNF123*, *ATP2C1* and *COMT*. The association in *RNF123* was replicated, whereas *ATP2C1* showed a suggestive association, and the *COMT* locus did not replicate in 43 080 individuals from independent cohorts.

RNF123 gene encodes E3 ubiquitin-protein ligase, has a role in cell cycle progression, metabolism of proteins and innate immunity.^{34 35} This gene is highly expressed in skeletal muscle than other tissues. Recent studies involving UKB samples also associated the locus with musculoskeletal pain.^{19 36} However, it is not clear how RNF123 may contribute to CWP. Using in silico follow-up, we identified CDHR4, APEH, SLC25A20, NDUFAF3, DAG1, HYAL1, GMPPB, TRAIP, RHOA, CACNA2D2 and IMPDH2 genes as putative causal candidates at the locus, of which CACNA2D2, NDUFAF3 and IMPDH2 can be targeted using known drugs.³⁷⁻³⁹ Notably, CACNA2D2 encodes the alpha-2/delta subunit of the voltage-dependent calcium channel complex, which is a receptor for gabapentinoids,⁴⁰ used by some in the management of fibromyalgia.^{41 42} Another prioritised gene CDHR4 belongs to cadherin superfamily has a role in calcium-ion binding to facilitate cadherin-mediated cell-cell interaction.43 44

Additionally, the *ATP2C1* locus demonstrated suggestive association in replication (p=0.0227). There was a consistent direction of effect for *ATP2C1* locus in six replication cohorts but not ELSA, where we used a proxy SNP, which had close to zero effect size (beta= -0.0004 ± 0.0110). This is the first study to implicate *ATP2C1* with musculoskeletal



Figure 6 (A) Differentially expressed gene (DEG) plots for chronic widespread musculoskeletal pain (CWP) in 54 tissue types from GTEX v8, (B) DEG plots for CWP in 30 general tissue types from GTEX v8 and (C) Differential expression of *RNF123* gene across tissue types from GTEX v8. (A, B) In both plots, the y-axis represents the —log10 transformed two-tailed p value of the hypergeometric test. Significantly enriched DEG sets (Bonferroni-corrected p value <0.05) are highlighted in red. (C) Y-axis represents transcripts per million (TPM) and x-axis represents the GTEx (V.8) tissues. The figure was adapted from GTEx portal (https://www.gtexportal.org/home/gene/ENSG00000164068).

pain using an agnostic approach. The ATP2C1 gene encodes for the ATP-powered magnesium-dependent calcium pump protein hSPCA1, which mediates Golgi uptake of cytosolic Ca(2+) and Mg(2+).⁴⁵ A loss of function mutation in the ATP2C1 leads to Hailey-Hailey disease (HHD), an autosomal dominant skin condition characterised by blistering and erosion of the epidermis.⁴⁶ Interestingly, HHD may be treated successfully with low-dose naltrexone, an opioid receptor antagonist, which has also been used in the management of fibromyalgia.^{47 48} A recent study showed that naltrexone is capable of restoring calcium homeostasis in natural killer cells of patients with chronic fatigue syndrome.⁴⁹ Additionally, the role of calcium regulation in pain processing is well known.⁵⁰⁻⁵² Taken together, our findings suggest a role in the regulation of calcium influencing CWP/fibromyalgia.

COMT is one of the most studied genes in human pain.⁵³ Almost 30 SNPs and 3 haploblocks of the COMT gene have been studied in acute clinical, experimental and chronic pain. Rs4680 of the COMT gene is extensively studied in many pain phenotypes such as pain sensitivity, TMD and fibromyalgia.⁵⁴ Across multiple ethnic populations, rs4680 was implicated with fibromyalgia.55 However, a meta-analysis of 8 case-control studies (589 fibromyalgia cases and 527 controls) did not confirm earlier association.⁵⁶ To date, the largest study that assessed the association between COMT haplotypes (rs4680, rs4818, rs4633 and rs6269) and fibromyalgia included 60 367 participants (2713 ICD-9 diagnosed fibromyalgia) and found no association.⁵⁷ They have also been refuted in other European CWP samples^{21 58} and a large candidate gene study of fibromyalgia.⁵⁹ However, we identified rs165599, located at 3'-UTR of COMT, associated with CWP in the discovery sample but not in the meta-analysis

or any of the replication cohorts. This variant is not in LD with previously studied COMT SNPs rs4680, rs4818, rs4633 and rs6269, and was found not to be associated with chronic musculoskeletal pain including CWP neither when studied as a single SNP nor as a part of a haploblock.⁶⁰⁻⁶² Several explanations of our non-replication of COMT locus are possible. First, there was lower power pertaining to overall metaanalysis, which was estimated at 48% based on the effect size observed in the discovery sample (n=249843), replication sample size $(n=43\,080)$ and the number of tests conducted (n=3). Our meta-analysis did have 90% power to detect a relative risk as small as 1.04 but the estimated COMT effect was only 1.012 (beta=0.0027±0.004; OR=1.012, 95%) CI=0.97 to 1.05). However, our replication sample size was larger than many of the earlier studies that reported the association between COMT and CWP.^{20 63} Second, we observed a tendency towards non-significance for the COMT locus in the sensitivity GWAS due to the exclusion of participants with non-musculoskeletal pain from the control group suggesting that COMT predisposes to chronic pain in general. Finally, genetic factors underlying chronic pain and psychiatric comorbidity (e.g. depression and neuroticism) are known to be shared.⁶⁴ However, previous GWAS on chronic pain,^{28 65 66} depression⁶⁷ and neuroticism⁶⁸ have failed to detect an association with COMT. Thus, if there is a role of COMT in CWP, it is likely minimal.

Epidemiological studies have consistently reported higher BMI to be associated with an increased risk of CWP.^{10 11 69} Our analysis showed significantly higher BMI in CWP cases compared with controls (p<0.0001) in all cohorts except RS-3. In line with this, we observed a positive genetic overlap between BMI and CWP independent of genetic confounders. Similarly, genetically independent pairwise genetic correlation for depressive symptoms, age of first birth and years of schooling was seen with CWP. These findings indicate the presence of shared molecular pathways underlying these traits.

Functional analysis showed that FUMA mapped genes differentially expressed in skeletal muscle, several areas of the CNS, pancreas, whole blood and heart tissues. These findings suggest the involvement of nervous, musculoskeletal and neuroendocrine systems in CWP. These physiological systems have been implicated in fibromyalgia by previous studies.^{70–72} Evidence suggests that both peripheral and central pain mechanisms influence CWP.^{73 74} We observed overall stronger enrichment for differentially expressed gene sets in skeletal muscle than other GTEx tissues. Also, skeletal muscle and DRG eQTLs colocalise with the *RNF123* locus. These findings suggest a substantial involvement of peripheral pain mechanisms in CWP.

The study has limitations. The case definition of CWP depends on self-report together with exclusion of other conditions with symptoms leading to chronic pain.⁷⁵ A clinical diagnosis of CWP would have been infeasible in a sample this large. Also, we used common SNPs to estimate the heritability of CWP, so the contribution of other variants in the heritability estimated remains unknown. The phenotype definition used in this study to estimate SNP heritability has differed from the Kato *et al*¹³ study, where a modulated American College of Rheumatology⁷⁶ criteria based on self-report was used to estimate broad-sense heritability. However, using UKB samples, a study reported the SNP heritability of pain all over the body, regardless of chronicity, on the liability scale was 0.31 ± 0.072 .⁶⁴ We found a similar but slightly higher estimate for CWP (0.33 ± 0.0004), suggesting our definition is meaningful and CWP is a trait of high genetic influence. Finally, our findings cannot be generalisable to ancestry other than northern Europeans (online supplemental text).

In summary, this study identified a novel association for CWP in the *RNF123* locus and suggested the role of calcium regulation, by the involvement of the *CDHR4*, *CACNA2D2* and *ATP2C1* genes. The association of the *COMT* locus with CWP was not replicated, suggesting a small influence, if any. We found evidence that the epidemiological association of BMI and CWP is at least in part genetically mediated. Finally, our results suggest a profound role of peripheral mechanisms in the pathogenesis of CWP.

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