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OPEN GWAS of allometric body-shape indices in UK Biobank identifies loci suggesting associations with morphogenesis, organogenesis, adrenal cell renewal and cancer

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Genetic studies have examined body-shape measures adjusted for body mass index (BMI), while allometric indices are additionally adjusted for height. We performed the first genome-wide association study of A Body Shape Index (ABSI), Hip Index (HI) and the new Waist-to-Hip Index and compared these with traditional indices, using data from the UK Biobank Resource for 219,872 women and 186,825 men with white British ancestry and Bayesian linear mixed-models (BOLT-LMM). One to two thirds of the loci identified for allometric body-shape indices were novel. Most prominent was rs72959041 variant in RSPO3 gene, expressed in visceral adipose tissue and regulating adrenal cell renewal. Highly ranked were genes related to morphogenesis and organogenesis, previously additionally linked to cancer development and progression. Genetic associations were fewer in men compared to women. Prominent region-specific associations showed variants in loci VEGFA and HMGA1 for ABSI and KLF14 for HI in women, and C5orf67 and HOXC4/5 for ABSI and RSPO3, VEGFA and SLC30A10 for HI in men. Although more variants were associated with waist and hip circumference adjusted for BMI compared to ABSI and HI, associations with height had previously been reported for many of the additional variants, illustrating the importance of adjusting correctly for height.

Abbreviations

ABSI	A body shape index
ABSI _{UKB}	A body shape index calibrated for UK Biobank participants
ADAMTS	A disintegrin and metalloprotease domains with thrombospondins motifs
AKR1C2	Aldo-keto reductase family 1 member C2
BMI	Body mass index
BOLT-LMM	Bayesian mixed-model association method
C5orf67	Chromosome 5 open reading frame 67
CADD	Combined Annotation Dependent Depletion
CASC20	Cancer susceptibility 20
CMIP	C-Maf inducing protein
COBLL1	Cordon-bleu WH2 repeat protein like 1
DLEU1	Deleted in lymphocytic leukemia 1
eQTL	Expression Quantitative Trait Loci
ERa	Oestrogen receptor a
ERI1	Exoribonuclease 1

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EYA FAM101A FGFR4 FUMA GO GWAS HC NUC DMU	Eyes absent homolog Filamin-interacting protein (readthrough region with <i>ZNF664</i>) Fibroblast growth factor receptor 4 Functional Mapping and Annotation Gene ontology Genome-wide association study Hip circumference
HC _{adj} BMI HI	Hip circumference adjusted for body mass index Hip index
HI _{UKB} HMGA1	Hip index calibrated for UK Biobank participants High mobility group AT-hook 1
HPA	Hypothalamic-pituitary-adrenal axis
KLF14	Kruppel-like factor 14
LD MAF	Linkage disequilibrium Minor allele frequency
MAGMA	Multi-marker Analysis of GenoMic Annotation
MC1R	Melanocortin 1 receptor
NEU1	Neutraminidase 1
NHANES	National Health and Nutrition Examination Survey
PEMP	Palmitoylated erythrocyte membrane protein
PLXND1	Plexin D1
SLC30A10	Solute carrier family 30 member 10
SNP	Single nucleotide polymorphism
SSPN	Sarcospan
TBX15	T-box transcription factor 15
TFAP4	Transcription factor AP-4
VEGFA	Vascular endothelial growth factor-A
VEP	Ensembl Variant Effect Predictor
WC	Waist circumference
WC _{adj} BMI	Waist circumference adjusted for body mass index
WHI	Waist-to-hip index
WHI _{ukb} WHR	Waist-to-hip index calibrated for UK Biobank participants Waist-to-hip ratio
WHR _{adj} BMI	Waist-to-hip ratio adjusted for body mass index
WNT	Wingless-type
XKR6	XK related 6
ZMIZ1	Zinc-finger MIZ-type containing 1
ZNF664	Zinc finger protein 664

The cardiometabolic complications of obesity are influenced by body shape, showing a positive association with abdominal size and an inverse association with gluteofemoral size¹. The waist-to-hip ratio (WHR) and body mass index (BMI) are used correspondingly as an index of body shape and an index of general obesity². The WHR, however, is moderately correlated with BMI and cannot differentiate abdominal from gluteofemoral size. Although waist (WC) and hip circumference (HC) are measures of specific body regions, they are both strongly correlated with BMI. To account for the correlation with BMI, genome-wide association studies (GWAS) have used traditionally residuals from linear models regressing each of WHR, WC, or HC on BMI, thus creating body-shape indices independent from BMI^{3,4}. Nevertheless, the adjustment of WC or HC for BMI introduces a positive correlation with height, which is stronger than the association of WC or HC with height⁴.

An alternative approach to creating body-shape indices independent from body size and general obesity has been implemented in the development of A Body Shape Index (ABSI) and Hip Index (HI)^{5,6}. Similarly to BMI, ABSI and HI are based on the principle of allometry, which accounts for the expansion of the dimensions of individual body parts relative to the total body size with log-linear rather than linear models⁷. Similar to the scaling of log-transformed weight to log-transformed height used for the development of BMI, log-transformed WC and HC have each been scaled to log-transformed weight and log-transformed height to develop ABSI and HI^{5,6}. This scaling accounts for the expansion of body circumferences proportional to body size, as reflected in height, and additionally accounts for the proportional expansion of body circumferences with general adiposity, as reflected in body weight. Consequently, the allometric body-shape indices are independent by design from height, as well as from BMI. Strong associations of ABSI with mortality and cardio-metabolic risk factors have been reported^{5,8} and we have demonstrated that ABSI achieves better mortality risk stratification than alternative body-shape indices, which are correlated with BMI⁹.

There are, however, no studies to date examining the genetic associations of ABSI and HI, no allometric counterpart of WHR and no insight into the influence of the correlation of traditional body-shape indices with height. The aims of our study, therefore, were to perform the first GWAS of allometric body-shape indices and to compare allometric and traditional body-shape indices with respect to their genetic associations. Our GWAS provides novel information for unbiased genetic associations of body-shape indices.

Results

We used data from the UK Biobank for 219,872 women and 186,825 men with white British ancestry (Supplementary Table S1). For allometric body-shape indices, we used ABSI, HI and a waist-to-hip index (WHI) calibrated for UK Biobank participants (ABSI_{UKB}, HI_{UKB} and WHI_{UKB}, see details in Methods). For traditional body-shape indices, we used WC, HC and WHR adjusted for BMI in linear models (WC_{adj}BMI, HC_{adj}BMI and WHR_{adj}BMI). We examined women and men separately, as pronounced sexual dimorphisms have been reported for the genetic associations of traditional body-shape indices¹⁰.

Genetic variants associated with allometric body-shape indices. We determined independent significant single nucleotide polymorphisms (SNPs) and consolidated them in independent genomic risk loci, represented by a locus lead SNP, with Functional Mapping and Annotation (FUMA) (see details in Methods).

A larger number of independent genomic risk loci were associated with WHI_{UKB} (282 in women, 97 in men) compared to ABSI_{UKB} (200 in women, 65 in men) and HI_{UKB} (171 in women, 75 in men) (Table 1). The highest-ranked lead SNPs associated with WHI_{UKB} (*rs72959041* in women and *rs577721086* in men) were both in the *RSPO3* locus and in very strong linkage disequilibrium ($r^2 > 0.99$). In addition, *rs72959041* was the second highest-ranked lead SNP for ABSI_{UKB} in women and was the highest-ranked lead SNP for HI_{UKB} in both sexes (Supplementary Table S2). This variant stood out with some of the largest effect sizes, with a positive sign of the regression coefficients for ABSI_{UKB} and WHI_{UKB} and a negative sign for HI_{UKB}, reflecting the phenotype of the minor allele. Most of the independent variants associated with WHI_{UKB} similarly showed regression coefficients with opposite signs for ABSI_{UKB} or HI_{UKB}. In women, the most outstanding examples of loci associated exclusively with ABSI_{UKB} were *VEGFA* and *HMGA1*, while the most prominent locus associated exclusively with HI_{UKB} were *AC022431.2* (*C5orf67*), *RP11-115J16.1*, *RP5-859D4.3* (*CASC20*) and the region including loci *HOXC4*, *RP11-834C11.14* and *HOXC5*, while the most prominent loci associated exclusively with HI_{UKB} were *RSPO3*, *VEGFA* and *RP11-95P13.2* (*SLC30A10*) (Supplementary Fig. S2).

Approximately two thirds of the independent significant SNPs but less than one fifth of the corresponding genomic risk loci associated in our study with WHI_{UKB} were novel, i.e. they have not been reported in the NHGRI-EBI GWAS Catalog¹¹ in association with WHR or WHR_{adj}BMI (Table 1, Supplementary Table S3). Nevertheless, over two thirds of the independent significant SNPs and one third of the corresponding genomic risk loci associated with ABSI_{UKB} were novel, i.e. not previously reported in association with WC or WC_{adj}BMI. Further, most of the independent significant SNPs and over two thirds of the corresponding genomic risk loci associated with HI_{UKB} were novel, i.e. not previously reported in association with HC or HC_{adj}BMI. It was notable, however, that many of the novel highest-ranked lead SNPs associated with allometric body-shape indices were in strong linkage disequilibrium (LD at $r^2 \ge 0.6$) with variants previously reported in association with the corresponding traditional body-shape index, which in our study showed lower significance (Fig. 1). Exceptions were the novel highly ranked lead SNPs associated with HI_{UKB} in men, *rs998584* in the *VEGFA* locus and *rs6066114* in the *EYA2* locus, which did not include in their clumps variants previously reported in association with HC or HC_{adj}BMI, while for *rs113733630* in the *TFAP4* locus, the previously reported variant was not in the same LD block (Supplementary Table S2).

The deleteriousness Combined Annotation Dependent Depletion (CADD) score was above the recommended cut-off 12.37¹² for 8% to 20% of the locus lead SNPs associated with allometric body-shape indices (Fig. 2). Nevertheless, almost half of all locus lead SNPs were in strong LD with a variant with genome-wide significance showing higher CADD than the cut-off.

Following recent reports of a fine-scale population structure in UK Biobank¹³, we checked matches with the SNPs reported in association with birth location. Only *rs1805007* (*MC1R*), reported in association with height, was identified in our study as a candidate SNP and was included in the LD block of locus lead SNPs for WC_{adj}BMI and HC_{adj}BMI in women and men (lowest $P = 4.4*10^{-11}$). Although *rs9268556* (MHC region), reported in association with forced vital capacity, was additionally identified as a candidate SNP for WHI_{UKB}, ABSI_{UKB} and WHR_{adj}BMI in women and HC_{adj}BMI in men (lowest $P = 2.4*10^{-9}$), this was not included in the LD block of a locus lead SNP.

Sexual dimorphism in the genetic associations of allometric body-shape indices. The genetic association patterns of allometric body-shape indices differed considerably between women and men. The heritability was larger in women compared to men, with up to three times more independent significant SNPs identified in women (Table 1), when the excess of women vs men in the dataset was only approximately 20%. Some 30% of the independent significant SNPs for ABSI_{UKB} but 10% or less for HI_{UKB} showed sex differences in effect size at $p_{sex} < 5^{*}10^{-6}$ (Table 1). Several highly ranked variants, however, were particularly affected (Supplementary Fig. S3). Thus, prominent associations with WHI_{UKB} and ABSI_{UKB} in women but not in men showed variants in loci *COBLL1*, *RP11-95P13.1*, *ADAMTS9-AS2*, *CMIP* and *AC022431.2* (*C5orf67*) and variants in the region including loci *ZNF664 and FAM101*, while *KLF14* was the most prominent locus associated with HI_{UKB} in women but not in men ($p_{sex} < 5^{*}10^{-28}$ for all independent significant SNPs in these loci). In addition, although variants in *RSPO3* locus were the highest-ranked associated with WHI_{UKB} and HI_{UKB} in both sexes, variants in the *VEGFA* and *RSPO3* loci were associated with ABSI_{UKB} almost exclusively only in women, while variants in the *VEGFA* locus were associated with HI_{UKB} mainly in men (Supplementary Fig. S3).

	WHI UKB	WHR _{adj} BMI	ABSIUKB	WC _{adj} BMI	НІ _{UKB}	
Women						
Total number						
Genomic Risk Loci (SNP-based) ^a	282	276	200	270	171	329
Lead SNPs ^a	586	579	368	502	282	618
Independent Significant SNPs ^a	1,988	1,959	1,273	1,586	814	2,008
Candidate SNPs ^a	58,203	58,172	42,447	49,768	21,553	65,934
Genomic Risk Regions (gene-based) ^b	237	229	158	228	136	295
Significant Genes ^b	759	748	584	707	299	923
Reported for Index in GWAS Catalog $^\circ$						
Genomic Risk Loci (SNP-based)	236 (84)	237 (86)	119 (60) **	201 (74) **	31 (18) *	87 (26) *
Independent Significant SNPs	729 (37)	718 (37)	278 (22) ***	478 (30) ***	40 (4.9)	121 (6.0)
Genomic Risk Regions (gene-based)	181 (76)	177 (77)	100 (63)	157 (69)	26 (19)	72 (24)
Significant Genes	342 (45)	337 (45)	172 (29) *	262 (37) *	33 (11)	90 (9.8)
Reported for Height in GWAS Catalog ^d	T	r	r.	1		r
Locus Lead SNPs	41 (15)	42 (15)	39 (20) **	93 (34) **	18 (11) ***	165 (50) ***
Independent Significant SNPs	132 (6.6)	127 (6.5)	103 (8.1) ***	227 (14) ***	52 (6.4) ***	440 (22) ***
Lead Genes	92 (39)	89 (39)	71 (45)	117 (51)	63 (46) *	174 (59) *
Significant Genes	196 (26)	188 (25)	141 (24) *	228 (32) *	96 (32) *	373 (40) *
Reported for Cancer in GWAS Catalog ^d						
Locus Lead SNPs	19 (6.7)	20 (7.2)	20 (10)	23 (8.5)	9 (5.3)	30 (9.1)
Independent Significant SNPs	92 (4.6)	96 (4.9)	73 (5.7)	66 (4.2)	26 (3.2) *	104 (5.2) *
Lead Genes	66 (28)	66 (29)	53 (34)	66 (29)	41 (30)	80 (27)
Significant Genes	149 (20)	150 (20)	129 (22) *	120 (17) *	70 (23)	227 (25)
Sex Difference ^e						
Independent Significant SNPs	489 (25)	482 (25)	373 (29) ***	328 (21) ***	85 (10) ***	66 (3.3) ***
Difference from Alternative Index ^f						
Independent Significant SNPs	64 (3.2)	48 (2.5)	259 (20) *	399 (25) *	103 (13) ***	987 (49) ***
Correlation		[1		[
Allometric vs Traditional Index r _{allo trad}	0.995		0.923		0.785	
Women vs Men r _{sex}	0.163	0.161	0.144	0.185	0.143	0.255
Heritability h ² g ^h	0.258 (0.001)	0.253 (0.001)	0.211 (0.001)	0.244 (0.001)	0.178 (0.001)	0.273 (0.001)
Men						
Total number	07	00	05	400	76	000
Genomic Risk Loci (SNP-based) ^a Lead SNPs ^a	97	82	65	188	75	262
	146	122	81	273	93	435
Independent Significant SNPs ^a Candidate SNPs ^a	617	571	266	842	222	1,400
Genomic Risk Regions (gene-based) ^b	23,462	22,398	11,505	29,621	7,328	47,701
Significant Genes ^b	83	77	61	164	52	222
Reported for Index in GWAS Catalog °	303	312	183	421	106	724
Genomic Risk Loci (SNP-based)	88 (91)	76 (93)	42 (65)	141 (75)	18 (24)	80 (31)
Independent Significant SNPs	173 (28)	161 (28)	42 (03) 65 (24) *	283 (34) *	22 (9.9)	110 (7.9)
Genomic Risk Regions (gene-based)			31 (51) *	108 (66) *	17 (33)	
Significant Genes	60 (72) 106 (35)	58 (75) 104 (33)	45 (25) *	162 (38) *	22 (21) *	68 (31) 86 (12) *
Reported for Height in GWAS Catalog ^d	100 (33)	104 (33)	43 (23)	102 (30)	22 (21)	00 (12)
Locus Lead SNPs	25 (26)	16 (20)	22 (34) ***	119 (63) ***	14 (19) ***	162 (62) ***
Independent Significant SNPs	64 (10)	55 (9.6)	40 (15) ***	235 (28) ***	30 (14) ***	373 (27) ***
Lead Genes	40 (48)	35 (9.0)	25 (41) *	103 (63) *	25 (48)	
Significant Genes	76 (25)	35 (45) 80 (26)	47 (26) ***	182 (43) ***	25 (46) 38 (36)	140 (63) 297 (41)
Reported for Cancer in GWAS Catalog ^d	10(23)	00 (20)	47 (20)	102 (43)	30 (30)	237 (41)
Locus Lead SNPs	10 (10)	10 (12)	10 (15)	19 (10)	6 (8.0)	26 (9.9)
Independent Significant SNPs	56 (9.1)	57 (10)	28 (11) *	49 (5.8) *	12 (5.4)	93 (6.6)
Lead Genes	28 (34)	27 (35)	22 (36)	49 (3.8) 47 (29)	17 (33)	59 (27)
Significant Genes	99 (33)	103 (33)	50 (27)	88 (21)	25 (24)	170 (23)
Sex Difference ^e	00 (00)		30 (27)	30 (21)		
Independent Significant SNPs	35 (5.7)	29 (5.1)	10 (3.8)	32 (3.8)	16 (7.2) **	28 (2.0) **
Difference from Alternative Index ^f	00 (0.7)		10 (0.0)	02 (0.0)	10 (1.2)	20 (2.0)
Binoronoo nonn Altornative muex	73 (12)	65 (11)	58 (22) ***	397 (47) ***	44 (20) ***	877 (63) ***
Independent Significant SNPs			~~ (<i>LL</i>)		(20)	0.1 (00)
Independent Significant SNPs Correlation	• • • •					
Correlation	1		0.879		0.759	
	0.994	0.162 (0.002)	0.879	0.225 (0.002)	0.759	0.276 (0.002)

Table 1. Counts and overlaps of independent genetic variants and loci associated with allometric and traditional body-shape indices in women and men. ^aBased on FUMA; ^bBased on MAGMA; ^cNumber (percentage from total per index) reported in the NHGRI-EBI GWAS Catalog¹¹ (https://www.ebi.ac.uk/gwas/ home, accessed on 07/04/2021) in association with the corresponding traditional body-shape index (with or without adjustment for body mass index, BMI), i.e. the waist-to-hip ratio for WHI_{UKB} and WHR_{adj}BMI (catalogue sets EFO_0004343, EFO_0007788, EFO_0004302), waist circumference for $ABSI_{UKB}$ and $WC_{adj}BMI$ (EFO_0004342, EFO_0007789, EFO_0004302), hip circumference for HI_{UKB} and HC_{adi}BMI (EFO_0005093, EFO_0008039, EFO_0004302). To ensure novelty, a match was counted for an independent significant SNP if any of the candidate SNPs in the LD block was reported and for a genomic risk locus (or region) if any of the corresponding independent significant SNPs (or genes) was reported; ^dNumber (percentage from total per index), counting indirect matches only for SNPs in strong LD, i.e. candidate SNPs within the corresponding LD block for independent significant SNPs and for a genomic risk locus (or region) only a match of the independent significant SNP (or gene) promoted to a locus lead SNP (or lead gene) (EFO_0004339, EFO_0004302 for height; EFO_0000311 for cancer); *Number (percentage from total per index) showing sex difference (p_{difference}<5*10⁻⁶ for any candidate SNP within the corresponding LD block); ^fAs for ^ebut reflecting difference from the corresponding alternative body-shape index, i.e. $WHR_{adj}BMI$ for WHI_{UKB} , $WC_{adj}BMI$ for $ABSI_{UKB}$, $HC_{adj}BMI$ for HI_{UKB} and vice versa; ABSI_{UKB}-a body shape index calibrated for UK Biobank participants; HC_{adj}BMIhip circumference adjusted for BMI; HI_{UKB} hip index calibrated for UK Biobank participants; WC_{adj}BMIwaist circumference adjusted for BMI; WHI_{UKB} -waist-to-hip index calibrated for UK Biobank participants; $WHR_{adj}BMI$ -waist-to-hip ratio adjusted BMI; h^2_g -estimated (pseudo-) heritability, based on the genetic relationship matrix in BOLT-LMM (approximate standard error, 316/number of individuals), comparison between men and women; rsex-Spearman's rank correlation coefficient between regression coefficients in women and men across all examined genetic variants (used in the test for difference of effect size or heritability); r_{allo-trad}-correlation coefficient as for r_{sex} but between the corresponding allometric and traditional index; Fisher's exact test was used to compare percentages between allometric and traditional body-shape indices: *P < 0.05; ***P*<0.001; ****P*<0.0001.

Gene-level associations with allometric body-shape indices. We examined gene-level associations with Multi-marker Analysis of GenoMic Annotation (MAGMA) employed in FUMA, which uses the association statistics of all SNPs included in a gene (see Methods for details).

Similar to SNP-based analyses, a larger number of non-overlapping genomic risk regions were associated with WHI_{UKB} (237 in women, 83 in men) compared to $ABSI_{UKB}$ (158 in women, 61 in men) and HI_{UKB} (136 in women, 52 in men) and the proportions of novel gene-based regions were comparable to the proportions of novel SNP-based loci (Table 1). There were, however, differences in the lead genes (Fig. 3). The highest-ranked region for WHI_{UKB} and ABSI_{UKB} in women was *FAM101A* and for WHI_{UKB} and HI_{UKB} in men was *DLEU1*. Prominent novel genes with their corresponding genomic risk regions were *XKR6* for ABSI_{UKB} and *ERI1* for WHI_{UKB} and ABSI_{UKB} in men, *EYA2* for HI_{UKB} in both women and men, *SLC30A10* and *SSPN* for HI_{UKB} in women and *PEMP* and *AKR1C2* for HI_{UKB} in men (Supplementary Table S4). Although few independent significant SNPs had previously been reported in association with cancer (or were in strong LD with previously reported SNPs, Supplementary Fig. S4), one third of the lead genes had previously been reported in association with cancer (Table 1), including the prominent lead genes *DLEU1 and ZMIZ1* and a notable cluster of significant genes in the region of *NEU1* on chromosome 6 (Supplementary Fig. S5).

Competitive gene-set analysis showed associations of allometric body-shape indices with gene sets related to embryonal morphogenesis and organogenesis, regulation of DNA binding, gene expression, biosynthetic processes and cell differentiation, transcription factor complexes, extracellular matrix component, circulatory system development, skeletal system development, ossification and chondrocyte differentiation and several cancers, with only a single gene set related to fat cell differentiation, which was associated with $ABSI_{UKB}$ in women (Fig. 4). A larger number of gene sets showed significant associations with WHI_{UKB} and $ABSI_{UKB}$ in women (42 and 29, correspondingly) compared to men (16 and 2) but a similar number were associated with HI_{UKB} (14 in women, 12 in men), although with only three overlapping.

Associations with expression Quantitative Trait Loci (eQTL) were significant in both women and men for adipose tissue (subcutaneous and visceral omentum), arteries (tibial, coronary and aorta) and, unexpectedly, for female reproductive organs (breast mammary tissue, uterus, ectocervix, endocervix and fallopian tube) and, in women only, also for ovary and vagina (Fig. 5).

Comparison between allometric and traditional body-shape indices. On the one hand, WHI_{UKB} and WHR_{adj}BMI were very strongly phenotypically correlated with each other and both were uncorrelated with height or BMI (Supplementary Table S5). Correspondingly, the ranking of independent significant SNPs associated with them was in excellent agreement, especially in women (Fig. 6a,d, Supplementary Fig. S6), with only some 5% showing differences in effect size at $p_{difference} < 5 \times 10^{-6}$ (Table 1, Supplementary Fig. S7 for women, Supplementary Fig. S9 for women, Supplementary Fig. S10 for men).

On the other hand, while the allometric indices $ABSI_{UKB}$ and HI_{UKB} were phenotypically uncorrelated with height and BMI, the traditional indices $WC_{adj}BMI$ and $HC_{adj}BMI$ were uncorrelated only with BMI. The adjustment for BMI was apparently introducing a moderate positive correlation with height, as height was only weakly

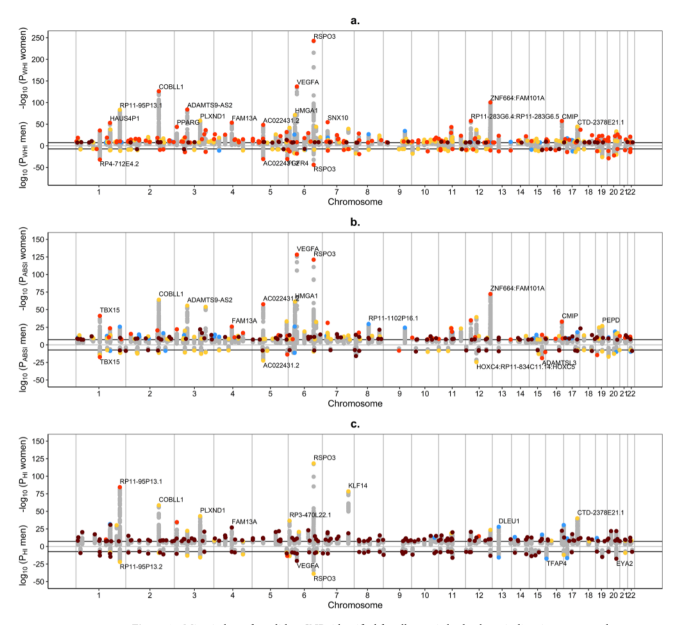


Figure 1. Miami plots of candidate SNPs identified for allometric body-shape indices in women and men. (a) GWAS of waist-to-hip index calibrated for UK Biobank participants (WHI_{UKB}); (b) GWAS of a body shape index calibrated for UK Biobank participants (ABSI_{UKR}); (c) GWAS of hip index calibrated for UK Biobank participants (HI_{UKB}); P-P-values were derived from BOLT-LMM infinitesimal models; SNP-single nucleotide polymorphism; horizontal lines correspond to the genome-wide significance cut-off $P = 5*10^{-8}$. Genomic risk loci with their corresponding locus lead SNPs were identified with FUMA v1.3.6a. All candidate SNPs are shown in grey, locus lead SNPs are colour-coded as follows: (grey circle) candidate SNPs; (dark red circle) novel genomic risk locus identified in the current study, with no previously reported candidate SNPs; (orange circle) genomic risk locus with a previously reported locus lead SNP; (yellow circle) genomic risk locus with a previously reported SNP in strong linkage disequilibrium (LD) with the locus lead SNP at $r^2 \ge 0.6$; (cyan circle) genomic risk locus with previously reported other candidate SNP. SNPs identified for allometric body-shape indices in the current study were matched against SNPs reported in the NHGRI-EBI GWAS Catalog¹¹ (https:// www.ebi.ac.uk/gwas/home, accessed on 07/04/2021) in association with the corresponding traditional bodyshape index (with or without adjustment for body mass index, BMI), i.e. the waist-to-hip ratio for WH_{IIKR} (catalogue sets EFO_0004343, EFO_0007788, EFO_0004302); waist circumference for ABSI_{UKB} (EFO_0004342, EFO_0007789, EFO_0004302); hip circumference for HI_{UKB} (EFO_0005093, EFO_0008039, EFO_0004302).

correlated with unadjusted WC and HC (Supplementary Table S5). Notably, a larger number of independent significant SNPs were associated with WC_{adj}BMI and HC_{adj}BMI, with half or more of them showing significant differences in effect size compared to their allometric counterparts $ABSI_{UKB}$ and HI_{UKB} (Table 1, Supplementary

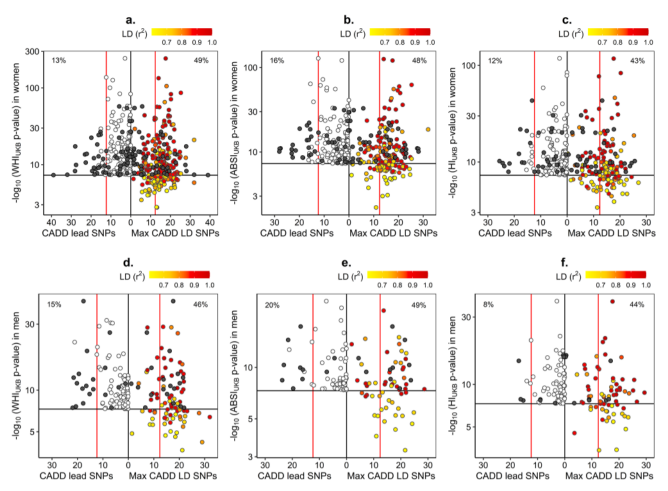


Figure 2. Deleteriousness (CADD) score of locus lead SNPs identified for allometric body-shape indices. (a) waist-to-hip index (WHI_{UKB}) calibrated for UK Biobank women (n = 282 genomic risk loci with the corresponding locus lead SNPs); (b) a body shape index (ABSI_{UKB}) calibrated for UK Biobank women (n = 200); (c) hip index (HI_{UKB}) calibrated for UK Biobank women (n = 171); (d) WHI_{UKB} for UK Biobank men (n = 97); (e) ABSI_{UKB} for UK Biobank men (n = 65); (f) HI_{UKB} for UK Biobank men (n = 75). CADD-Combined Annotation Dependent Depletion; LD-linkage disequilibrium; SNP-single nucleotide polymorphism; red vertical lines-recommended cut-off 12.37 for CADD (the higher the more deleterious)¹²; horizontal line-genome-wide significance cut-off ($P = 5*10^{-8}$); left-hand side-CADD for the locus lead SNP of each genomic risk locus, with the proportion above the cut-off; right-hand side-candidate SNPs in strong LD with the locus lead SNP (colour-coded according to $r^2 \ge 0.6$), showing the maximum CADD within the LD block, plotted with the corresponding significance on the y-axis; $P_{WHI/ABSI/HI}$ -P-values for body-shape indices, derived from BOLT-LMM infinitesimal models; (black circle)-marks the locus lead SNP when this is showing the maximum CADD within the corresponding LD block; percentages (top corners)-percentage above the cut-offs for both, CADD and genome-wide significance (all differences between left-hand side and right-hand side proportions were significant at P < 0.0001 when compared with Fisher's exact test, except for ABSI_{UKB} in men (P = 0.0008).

Fig. S7, Supplementary Fig. S8), except for $WC_{adj}BMI$ and $ABSI_{UKB}$ in women, which showed more similarity (Fig. 6b). The main differences between $WC_{adj}BMI$ and $ABSI_{UKB}$ (Fig. 6b,e) and between $HC_{adj}BMI$ and HI_{UKB} (Fig. 6c,f) concerned independent variants which were either previously reported in association with height or were in strong LD with variants reported in association with height (Supplementary Fig. S9, Supplementary Fig. S10). Half or more of the locus lead SNPs for $WC_{adj}BMI$ in men and for $HC_{adj}BMI$ in men and women had previously been associated with height, compared to less than one third for $ABSI_{UKB}$ and HI_{UKB} (Table 1). More comparable proportions of lead genes (with gene-level significance) had previously been reported in association with height (less than half for allometric indices, Supplementary Fig. S11, and more than half for traditional indices), but the absolute number of lead genes identified for traditional indices was considerably larger (Table 1).

Discussion

Our study presents the first GWAS of allometric body-shape indices, performed separately in women and men with white British ancestry, and a comparison with traditional body-shape indices. One third of the genomic risk loci associated with $ABSI_{UKB}$ and over two thirds of the genomic risk loci associated with HI_{UKB} in our study were novel. Genetic associations were sexually dimorphic, with fewer independent variants identified in men compared to women. The highest-ranked independent variant for WHI_{UKB} , $ABSI_{UKB}$ in women and HI_{UKB} was

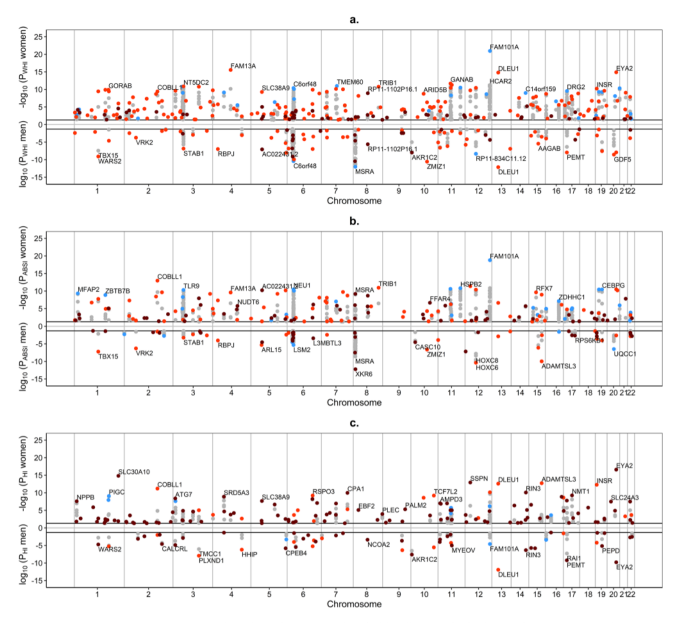


Figure 3. Miami plots of significant genes identified for allometric body-shape indices in women and men. (a) waist-to-hip index calibrated for UK Biobank participants (WHI_{UKB}); (b) a body shape index calibrated for UK Biobank participants (ABSI_{UKB}); (c) hip index calibrated for UK Biobank participants (HI_{UKB}); P-P-values were derived from MAGMA v1.08 employed in FUMA v1.3.6a and were adjusted with Bonferroni correction for 19,088 protein-coding genes; SNP-single nucleotide polymorphism; horizontal lines-correspond to P = 0.05after Bonferroni correction. Significant genes within 250 kb window were consolidated in genomic risk regions represented by a lead gene. All significant genes are shown in grey, lead genes are colour-coded as follows: (grey circle)-significant genes; (dark red circle)-novel genomic risk region identified in the current study, with no previously reported significant gene; (orange circle)-genomic risk region with a previously reported lead gene; (cyan circle)-genomic risk region including a previously reported significant gene (not the lead gene). Genes identified for allometric body-shape indices in the current study were matched against genes reported in the NHGRI-EBI GWAS Catalog¹¹ (https://www.ebi.ac.uk/gwas/home, accessed on 07/04/2021) in association with the corresponding traditional body-shape index (with or without adjustment for body mass index, BMI), i.e. the waist-to-hip ratio for WHI_{LKR} (catalogue sets EFO_0004343, EFO_0007788, EFO_0004302); waist circumference for ABSI_{UKB} (EFO_0004342, EFO_0007789, EFO_0004302); hip circumference for HI_{UKB} (EFO_0005093, EFO_0008039, EFO_0004302).

rs72959041 in the *RSPO3* locus. WHI_{UKB}, combining WC and HC, showed higher sensitivity to detect genetic associations compared to the regional indices ABSI_{UKB} and HI_{UKB}, based individually on WC or HC. The genetic association patterns of WHI_{UKB} and WHR_{adi}BMI were very similar. ABSI_{UKB} and HI_{UKB} showed fewer genetic

-		0.45 (0.00)		0.44 (0.00)	0.11.0.00	0.44 (0.00)						
	0.21 (0.03)	0.15 (0.03)	0.17 (0.03)	0.11 (0.03)	0.11 (0.02)	0.11 (0.02)	go-bp: positive regulation of rna biosynthetic process (1520)					
	0.18 (0.03)	0.14 (0.03) 0.15 (0.03)	0.16 (0.02) 0.20 (0.03)	0.11 (0.02) 0.13 (0.03)	0.10 (0.02) 0.13 (0.03)	0.09 (0.02) 0.14 (0.03)	go-bp: positive regulation of gene expression (1842)					
	0.22 (0.03)	0.15 (0.03)		0.13 (0.03)		0.14 (0.03)	go-bp: positive regulation of transcription by rna polymerase ii (1127)					
	0.22 (0.04)	0.17 (0.03)		0.13 (0.03)	0.13 (0.03)		go-mf: regulatory region nucleic acid binding (894)					
	0.19 (0.03)		0.12 (0.03)	0.10 (0.03)	0.11 (0.03)	0.07 (0.03)	go-mf: sequence specific dna binding (1057)					
	0.23 (0.04)	0.16 (0.04)		0.15 (0.03)	0.12 (0.03)	0.11 (0.03)	go-mf: dna binding transcription factor activity (1615)					
	0.31 (0.05)	0.22 (0.05)	0.25 (0.05)	0.16 (0.04)	0.18 (0.04)	0.16 (0.04)	go-mf: double stranded dna binding (911)					
	0.24 (0.04)	0.17 (0.04)		0.16 (0.04)	0.14 (0.03)	0.11 (0.03)	go-mf: proximal promoter sequence specific dna binding (514) go-mf: sequence specific double stranded dna binding (824)					
	0.15 (0.03)	0.10 (0.03)		0.10 (0.02)	0.09 (0.02)	0.10 (0.02)	go-bp: positive regulation of biosynthetic process (1864)					
	0.25 (0.04)		0.18 (0.04)	0.18 (0.04)	0.13 (0.04)	0.12 (0.04)	go-mf: transcription factor binding (614)					
	0.59 (0.11)		0.24 (0.10)	0.30 (0.09)	0.23 (0.09)	0.24 (0.09)	curated: zhu cmv all dn (115)					
	0.71 (0.13)	0.60 (0.12)		0.29 (0.11)	0.20 (0.11)	0.16 (0.11)	curated: zhu cmv 24 hr dn (81)					
	0.23 (0.04)	0.20 (0.04)	0.17 (0.04)	0.11 (0.04)	0.07 (0.04)	0.13 (0.04)	go-bp: tube morphogenesis (786)					
	0.31 (0.06)	0.25 (0.05)		0.19 (0.05)	0.21 (0.05)	0.14 (0.05)	go-mf: dna binding transcription activator activity (399)					
	0.34 (0.06)	0.24 (0.06)	0.24 (0.06)	0.20 (0.06)	0.11 (0.05)	0.19 (0.05)	curated: kegg pathways in cancer (318)					
	0.32 (0.06)	0.27 (0.06)		0.07 (0.05)	0.08 (0.05)	0.14 (0.05)	curated: greshock cancer copy number up (323)					
	0.22 (0.04)	0.18 (0.04)		0.04 (0.04)	0.09 (0.03)	0.09 (0.03)	curated: dacosta uv response via ercc3 dn (838)					
100	0.20 (0.04)	0.18 (0.04)	0.14 (0.03)	0.10 (0.03)	0.08 (0.03)	0.09 (0.03)	go-bp: tube development (966)					
	3.67 (0.69)	3.27 (0.66)	1.64 (0.63)	2.12 (0.61)	1.53 (0.58)	1.72 (0.58)	curated: nikolsky breast cancer 7p15 amplicon (11)					
	0.28 (0.05)	0.29 (0.05)		0.30 (0.05)	0.18 (0.05)		go-bp: skeletal system development (498)					
	0.37 (0.07)		0.26 (0.06)	0.18 (0.06)	0.11 (0.06)	0.15 (0.06)	curated: wang mll targets (274)					
	1.11 (0.22)	1.01 (0.21)	0.65 (0.20)	0.50 (0.19)	0.28 (0.18)	0.71 (0.18)	curated: rozanov mmp14 targets subset (32)					
	0.31 (0.06)	0.28 (0.06)	0.20 (0.05)	0.25 (0.05)	0.19 (0.05)	0.17 (0.05)	go-bp: ossification (359)					
	0.15 (0.03)	0.10 (0.03)	0.08 (0.03)	0.08 (0.03)	0.08 (0.02)	0.11 (0.02)	go-bp: response to endogenous stimulus (1548)					
	0.32 (0.06)	0.31 (0.06)	0.02 (0.06)	0.03 (0.06)	-0.03 (0.05)	0.14 (0.05)	curated: schuetz breast cancer ductal invasive up (335)					
	0.19 (0.04)	0.20 (0.04)	0.13 (0.03)	0.13 (0.03)	0.10 (0.03)	0.11 (0.03)	go-bp: circulatory system development (1001)					
	0.15 (0.03)	0.13 (0.03)		0.11 (0.03)		0.10 (0.03)	go-bp: negative regulation of biosynthetic process (1435)					
	0.19 (0.04)		0.14 (0.03)	0.16 (0.03)		0.13 (0.03)	go-bp: animal organ morphogenesis (1005)					
	0.15 (0.03)	0.11 (0.03)	0.08 (0.03)	0.09 (0.03)	0.09 (0.03)		go-bp: cellular response to endogenous stimulus (1300)					
	0.19 (0.04)	0.17 (0.04)	0.17 (0.04)	0.12 (0.03)	0.12 (0.03)		go-bp: negative regulation of transcription by rna polymerase ii (778)					
	0.22 (0.05)	0.20 (0.04)		0.16 (0.04)	0.12 (0.04)	0.14 (0.04)	go-bp: cardiovascular system development (671)					
	0.24 (0.05)		0.17 (0.05)	0.17 (0.04)	0.14 (0.04)	0.09 (0.04)	go-bp: heart development (526)					
	0.41 (0.09)	0.34 (0.08)	0.31 (0.08)	0.26 (0.08)	0.19 (0.07)	0.32 (0.07)	go-bp: morphogenesis of a branching structure (191)					
	1.24 (0.27)	1.01 (0.26)	0.43 (0.24)	0.31 (0.23)	-0.13 (0.22)	0.65 (0.23)	curated: mccabe hoxc6 targets dn (21)					
1.12	0.36 (0.08)	0.30 (0.07)	0.10 (0.07)	0.08 (0.07)	0.12 (0.06)	0.10 (0.06)	curated: boquest stem cell up (246)					
-	0.35 (0.08)	0.26 (0.07)	0.23 (0.07)	0.22 (0.07)	0.17 (0.06)	0.17 (0.06)	go-bp: cell fate commitment (246)					
	0.37 (0.08)	0.25 (0.08)	0.21 (0.07)	0.19 (0.07)	0.15 (0.07)	0.13 (0.07)	curated: reactome sumoylation (179)					
	1.04 (0.23)	0.88 (0.22)	0.15 (0.21)	0.60 (0.20)	0.40 (0.19)		curated: reactome met activates ptk2 signaling (30)					
1000	0.18 (0.04) 0.36 (0.08)	0.13 (0.04) 0.29 (0.08)	0.12 (0.04) 0.18 (0.07)	0.16 (0.03) 0.11 (0.07)	0.11 (0.03) 0.13 (0.07)	0.13 (0.03) 0.17 (0.07)	go-bp: embryo development (952)					
	0.38 (0.08)	0.29 (0.08)	0.18 (0.07)	0.64 (0.15)	0.13 (0.07)	0.17 (0.07)	curated: onder cdh1 targets 2 up (236)					
	0.34 (0.08)	0.35 (0.08)	0.24 (0.07)	0.41 (0.07)	0.27 (0.07)	0.26 (0.07)	go-cc: extracellular matrix component (47) go-bp: skeletal system morphogenesis (235)					
	0.29 (0.07)	0.32 (0.07)	0.15 (0.06)	0.35 (0.06)	0.21 (0.06)	0.21 (0.06)	go-bp: connective tissue development (261)					
	0.46 (0.11)		0.27 (0.10)	0.53 (0.09)	0.29 (0.09)	0.31 (0.09)	go-bp: chondrocyte differentiation (116)					
	0.28 (0.17)	0.19 (0.16)	0.49 (0.15)	0.76 (0.15)	0.37 (0.14)	0.52 (0.14)	go-bp: regulation of chondrocyte differentiation (47)					
	0.27 (0.08)	0.35 (0.08)	0.20 (0.07)	0.36 (0.07)	0.21 (0.07)	0.20 (0.07)	go-bp: cartilage development (201)					
	0.64 (0.19)	0.48 (0.18)	0.31 (0.18)	0.82 (0.17)	0.51 (0.17)	0.40 (0.17)						
	0.23 (0.11)	0.31 (0.11)	0.01 (0.10)	0.46 (0.10)	0.38 (0.10)	0.25 (0.10)	go-bp: embryonic skeletal system development (124)					
	0.14 (0.03)	0.13 (0.03)	0.16 (0.03)	0.10 (0.03)	0.11 (0.03)	0.11 (0.03)						
1000	2.19 (0.57)	1.38 (0.55)	2.44 (0.52)	0.85 (0.49)	0.73 (0.47)	1.34 (0.48)						
	0.03 (0.32)	0.64 (0.30)	1.34 (0.29)	0.54 (0.26)	0.65 (0.25)	0.44 (0.25)	go-bp: heterochromatin assembly (10)					
	0.41 (0.13)	0.34 (0.13)	0.25 (0.12)	0.35 (0.12)	0.22 (0.11)	0.59 (0.11)	go-bp: embryonic placenta development (84)					
	-0.68 (0.68)	-0.66 (0.65)	0.56 (0.62)	2.21 (0.60)	0.47 (0.57)	2.87 (0.57)	go-mf: ketosteroid monooxygenase activity (6)					
	0.12 (0.04)	0.09 (0.04)	0.10 (0.04)	0.11 (0.03)	0.06 (0.03)	0.16 (0.03)	go-bp: negative regulation of developmental process (885)					
	0.73 (0.24)	0.60 (0.23)	0.64 (0.22)	0.59 (0.21)	0.33 (0.21)	0.99 (0.21)	go-bp: embryonic placenta morphogenesis (26)					
	0.10 (0.04)	0.07 (0.04)	0.06 (0.04)	0.11 (0.04)	0.07 (0.04)	0.17 (0.04)	go-bp: negative regulation of cell differentiation (664)					
	0.88 (0.28)	0.72 (0.26)	0.75 (0.25)	0.70 (0.24)	0.33 (0.23)	1.06 (0.23)	go-bp: labyrinthine layer morphogenesis (21)					
	0.30 (0.08)	0.34 (0.07)	0.16 (0.07)	0.14 (0.07)	0.08 (0.06)	0.21 (0.06)	go-bp: fat cell differentiation (208)					
	WHI	ABSI	HI	WHI	ABSI	HI						
		Women			Men		GO-MF GO-BP GO-CC Curated -log10 (Padj)					
		vomen			Men		GO-IVIE GO-DE GO-GO Cuiateu					

Figure 4. Gene sets associated with allometric body-shape indices. ABSI-a body shape index calibrated for UK Biobank participants (ABSI_{UKB}); Currated-curated gene set; GO-BP-gene ontology biological process; GO-CC-gene ontology cellular component; GO-MM-gene ontology molecular function; HI-hip index calibrated for UK Biobank participants (HI_{UKB}); WHI-waist-to-hip index calibrated for UK Biobank participants (WHI_{UKB}); WHI-waist-to-hip index calibrated for UK Biobank participants (WHI_{UKB}); #-reactome runx1 regulates transcription of genes involved in wnt signaling. The figure includes gene sets identified using competitive gene-set analysis with MAGMA v1.08 employed in FUMA v1.3.6a as significant for at least one of the allometric indices in women or men with adjusted $P_{adj} < 0.05$, incorporating Bonferroni correction for multiple comparisons for 15,485 gene sets. Gene sets with $P_{adj} < 0.05$ for WHI_{UKB} in women were sorted in descending order of P_{adj} for WHI_{UKB} in men, then the remaining gene sets with $P_{adj} < 0.05$ for WHI_{UKB} in women were sorted in descending order of P_{adj} for WHI_{UKB} in women and last, the remaining gene sets with $P_{adj} < 0.05$ for HI_{UKB} in women were sorted in descending order of P_{adj} for HI_{UKB} in women and last, the remaining gene set associated only with ABSI_{UKB} in women. The horizontal lines mark the start of the next sorting criterion. White cells are gene sets with $P_{adj} \ge 0.05$ for the corresponding body-shape index.

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Women				Men				-log10 (P _{adj})		
WHI	ABSI	HI	WHI	ABSI	HI			0 4 8 12		
0.050 (0.013)	0.029 (0.012)	0.023 (0.012)	0.019 (0.011)	0.018 (0.011)	0.017 (0.011)	Thyroid				
0.049 (0.012)	0.033 (0.012)	0.024 (0.011)	0.007 (0.011)	-0.011 (0.010)	0.032 (0.010)	Lung				
0.028 (0.009)	0.024 (0.008)	0.026 (0.008)	0.012 (0.008)	0.002 (0.007)	0.022 (0.007)					
0.074 (0.017)	0.050 (0.016)	0.069 (0.016)	0.039 (0.015)	0.022 (0.014)	0.038 (0.015)	Bladder				
0.081 (0.013)	0.070 (0.013)	0.037 (0.012)	0.025 (0.012)	0.034 (0.011)	0.032 (0.011)	Nerve Tibial				
0.039 (0.017)	0.055 (0.016)	0.044 (0.015)	0.037 (0.014)	0.030 (0.014)	0.026 (0.014)					
0.041 (0.016)	0.061 (0.016)	0.046 (0.015)	0.038 (0.014)	0.026 (0.014)	0.028 (0.014)					
0.053 (0.017)	0.072 (0.016)	0.054 (0.015)	0.052 (0.015)	0.037 (0.014)	0.036 (0.014)	비행 전에 집에 가지 않는 것이 같이 있는 것이 없는 것이 없다.	esophageal Junction			
0.063 (0.014)	0.063 (0.013)	0.041 (0.012)	0.037 (0.012)	0.034 (0.011)	0.042 (0.011)	Artery Aorta				
0.081 (0.016)	0.084 (0.015)	0.060 (0.014)	0.046 (0.014)	0.042 (0.013)	0.046 (0.013)	Artery Coronary Arteries				
0.069 (0.013)	0.079.(0.013)	0.051 (0.012)	0.048 (0.012)	0.042 (0.012)	0.039 (0.011)	Artery Tibial	55 5			
0.059 (0.014)	0.044 (0.012)	0.039 (0.013)	0.018 (0.013)	0.012 (0.012)	0.032 (0.012)	Vagina				
0.058 (0.013)	0.047 (0.012)	0.048 (0.012)	0.027 (0.011)	0.023 (0.011)	0.021 (0.011)	Ovary				
0.080 (0.016)	0.061 (0.015)	0.057 (0.014)	0.039 (0.013)	0.030 (0.013)	0.047 (0.013)	Fallopian Tube	Female reproductiv	ve system		
0 101 (0 015)	0.080 (0.015)	0.057 (0.014)	0.053 (0.014)	0.044 (0.013)	0.048 (0.013)	Cervix Ectocervix Cervix Endocervix				
0.107 (0.016)	0.084 (0.014)	0.068 (0.013) 0.068 (0.015)	0.055 (0.013) 0.056 (0.014)	0.048 (0.012) 0.044 (0.013)	0.038 (0.012) 0.048 (0.013)	Uterus Convix Estaconvix				
0.108 (0.017)	0.088 (0.016)	0.074 (0.015)	0.044 (0.015)	0.024 (0.014)	0.057 (0.014)	Breast Mammary 1	Issue			
0.085 (0.015)	0.071 (0.014)	0.074 (0.013)	0.045 (0.013)	0.016 (0.012)	0.053 (0.012)	Adipose Visceral Omentum				
0 1 12 (0.014)	0.098 (0.014)	0.076 (0.013)	0.044 (0.012)	0.028 (0.012)	0.058 (0.012)	Adipose tissue				

Figure 5. eQTLs associated with allometric body-shape indices. ABSI-a body shape index calibrated for UK Biobank participants (ABSI_{UKB}); eQTLs-expression Quantitative Trait Loci; HI-hip index calibrated for UK Biobank participants (HI_{UKB}); WHI-waist-to-hip index calibrated for UK Biobank participants (WHI_{UKB}). The figure includes specific tissues from the Genotype-Tissue Expression (GTEx) v8.0 database, queried for associations with eQTLs with gene-property analysis, which were identified by MAGMA v1.08 employed in FUMA v1.3.6a as significant for at least one of the allometric indices in women or men, i.e. with adjusted $P_{adj} < 0.05$, incorporating Bonferroni correction for multiple comparisons for 54 tissues. White cells are tissues with $P_{adj} \ge 0.05$ for the corresponding body-shape index.

associations compared correspondingly to WC_{adj}BMI and HC_{adj}BMI, which were associated with a larger proportion of variants previously reported in association with height.

Our study highlights the importance of the *RSPO3* locus for body shape and fat distribution in individuals with white British ancestry and potentially in individuals with white ethnic background. RSPO3 protein, together with wingless-type WNT proteins, activate the canonical WNT/ β -catenin pathway, which plays a depo-specific role in regulating locally adipocyte hyperplasia, hypertrophy and browning¹⁴. *RSPO3* shows differential expression in adipose depots, with the highest expression in visceral, intermediate in abdominal subcutaneous and lowest in gluteal subcutaneous adipose tissue¹⁵. *RSPO3* additionally promotes angioblast specification and vascular development¹⁶, plays a key role throughout life in maintaining the structural zonation and the replenishment of damaged cells in the adrenal glands¹⁷, regulates the renewal and differentiation of stem cells¹⁸, and contributes to cancer development and progression^{19,20}. Genetic variants in the *RSPO3* locus associated with abdominal obesity have additionally been associated with dyslipidaemia²¹, thus relating the *RSPO3* gene to the metabolic syndrome. The highest-ranked *RSPO3* variants, *rs72959041* (in women) and *rs577721086* (in men), which are in strong linkage disequilibrium, are specific to European populations²² and have previously been reported in association with WHR_{adj}BMI^{3,4,23}. The active variant, however, may be *rs577721086*, as this is located in the attachment site of CCCTC-binding factor, which acts as a gene repressor, insulator or activator, but also regulates the chromatin structure and enables inter-chromosomal interactions²⁴.

Although the genetic associations of body-shape indices are interpreted traditionally from the perspective of the adipose tissue and insulin resistance²⁵, highly ranked variants associated with body shape are located in genes coding transcription factors, receptors and enzymes involved in morphogenesis, embryonal developmental, cell proliferation and cell survival, which have additionally been linked to various cancers. Thus, in addition to the RSPO3 proteins, HMGA1 and TFAP4 activate the WNT/ β -catenin pathway, stimulate cell migration and invasion and promote cancer progression^{26,27}. VEGFs, which are upregulated by hypoxia, are key factors for tumour-associated angiogenesis, tissue infiltration and metastasis²⁸. PLXND1, which is involved in angiogenesis and is upregulated by VEGFs, has a constitutively low expression in adult tissues but is overexpressed in cancer tissues and their vasculature²⁹. KLF14 is associated with insulin resistance, dyslipidaemia, type 2 diabetes and a female-specific shift of body fat from gynoid to abdominal stores, in agreement with the hip-specific association in women found in our study. In mice, adipocyte-specific deletion of Klf14 results in similar metabolic effects, while Klf14 knockout results in spontaneous tumorigenesis³⁰. TBX15 belongs to a family of transcription factors regulating differentiation, proliferation, tissue integrity and epithelial-mesenchymal transition, which are relevant to cancer development and metastasis³¹. Overexpression, mutations and epigenetic silencing of ADAMTS genes have been reported in different tumours³². SLC30A10 is a zinc transporter maintaining zinc homeostasis, which when dysregulated can result in cancer initiation and progression³³. FGFR4 plays a critical role in embryonic development, tissue repair, tumour angiogenesis and progression³⁴. EYA1-4 proteins can influence tumour progression through several mechanisms³⁵. Homeobox (HOX) genes code transcription factors with DNA-binding

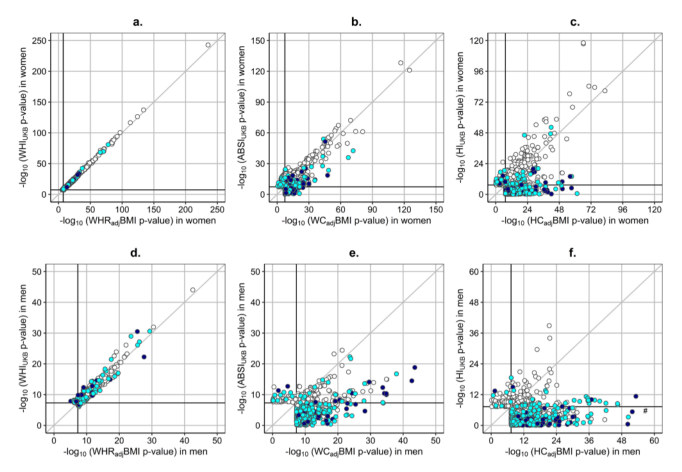


Figure 6. Significance ranking of independent significant SNPs: comparison between pairs of traditional and allometric body-shape indices. (a) independent significant SNPs (at $r^2 < 0.6$ within 1 Mb window) for WHI_{UKB} (n = 1,988) or WHR_{adi}BMI (n = 1,959) in women; (b) independent significant SNPs for ABSI_{UKB} (n = 1,273)or WC_{adi}BMI (n = 1,586) in women; (c) independent significant SNPs for HI_{UKB} (n = 814) or HC_{adi}BMI (n=2,008) in women; (d) independent significant SNPs for WHI_{UKB} (n=617) or WHR_{adi}BMI (n=571) in men; (e) independent significant SNPs for $ABSI_{UKB}$ (n = 266) or $WC_{adj}BMI$ (n = 842) in men; (f) independent significant SNPs for HI_{UKB} (n = 222) or $HC_{adj}BMI$ (1,400) in men. Association statistics p-values were derived from BOLT-LMM infinitesimal models. ABŚI_{UKB}-a body shape index calibrated for UK Biobank participants; BMI-body mass index; HC_{adi}BMI-hip circumference adjusted for BMI; HI_{UKB}-hip index calibrated for UK Biobank participants; SNP-single nucleotide polymorphism; WC_{adi}BMI-waist circumference adjusted for BMI; WHR_{adi}BMI-waist-to-hip ratio adjusted for BMI; WHI_{UKB}-waist-to-hip index calibrated for UK Biobank participants. Colour scale-colour-marked were only SNPs reported as associated with height in the NHGRI-EBI GWAS Catalog¹¹ (https://www.ebi.ac.uk/gwas/home, accessed on 07/04/2021), i.e. included in catalogue sets EFO_0004339 or EFO_0004302; (navy circle)-independent significant SNP reported in association with height; (cyan circle)-independent significant SNP in strong LD (at $r^2 \ge 0.6$) with a SNP reported in association with height.

activity, which regulate the formation of axial patterns and body shape during embryogenesis and have been implicated in cancer development³⁶. *COBLL1* is the ancestor of a family of proteins involved in morphogenesis and embryonal patterning in organisms with axial symmetry and is upregulated in castrate-resistant prostate cancer with poor prognosis³⁷. Upregulation of *DLEU1* can also promote tumorigenesis³⁸. A question, therefore, emerges whether excess abdominal fat mediates the development of cancers associated with abdominal obesity³⁹, or abdominal obesity and cancer are parallel outcomes with shared genetic predisposition. Although studies exploring a causal relationship between body shape and cancer are limited, Mendelian randomisation based on traditional body-shape indices has already provided evidence for causal positive associations of WHR or WHR_{adi}BMI with colorectal cancer^{40,41} and renal cell carcinoma⁴² but an inverse association with breast cancer⁴³.

Allometric body-shape indices showed sexually dimorphic genetic association patterns, like traditional bodyshape indices^{3,4,10,44}, with an overall lower heritability in men. There were, however, fewer genetic associations and with lower significance in men then would correspond proportionally to the slightly smaller sample size compared to women, as noted for traditional indices³. In contrast height, which is also phenotypically dimorphic between sexes, shows considerably larger heritability than body-shape indices and a closer agreement of the genetic association patterns between women and men⁴⁵. This raises the question, how women and men differ with respect to the regulatory factors determining fat distribution and body shape.

Sex steroid hormones play a central role in the regulation of fat distribution. Oestrogens counter fat accumulation in the abdominal area and favour fat accumulation in the gluteofemoral area⁴⁶. Oestrogen levels in blood⁴⁷ and the expression of ERa in the abdominal area⁴⁸ decrease after the menopause, which permits a functional androgen dominance and the development of android type obesity in post-menopausal women⁴⁹. Nevertheless, age-related differences in the genetic associations have been reported only for BMI and not for WHR_{adi}BMI¹⁰. In contrast, testosterone levels in women remain lower than in men at all ages⁵⁰ and are comparable between preand post-menopausal women, as ovarian androgen production is largely maintained long after the menopause⁵¹. The stress hormone cortisol, regulated by the hypothalamic-pituitary-adrenal (HPA) axis, is also closely involved in the regulation of fat distribution, with chronic cortisol excess resulting in the development of visceral adiposity and the metabolic syndrome⁵². Notably, the HPA axis shows a sexually dimorphic activity, with stronger responses to stimulation and suppression in women compared to men^{53,54} and increased responses to HPA stimulation in men after suppression of gonadal testosterone production⁵⁵. Abdominal obesity also influences HPA responsiveness, with larger WHR associated with a stronger response to HPA stimulation⁵⁶ and a weaker response to HPA suppression⁵³. Animal studies provide further evidence for a sexually dimorphic adrenal function, suggesting that androgens rather than oestrogens are important for adrenal regulation⁵⁷. Furthermore, absence of the androgen receptor in mice results in downregulation of the glucocorticoid receptor in the pituitary gland, adrenal hypertrophy and glucocorticoid overproduction⁵⁸. Thus, the higher HPA responsiveness in women, enabled by constitutively lower testosterone levels, may engage a more complex regulatory network and present more opportunities for genetic polymorphisms to act as rate-limiting steps in fat distribution, potentially explaining the pronounced sexual dimorphism in the heritability and the genetic association patterns of body-shape indices^{4,45}.

Despite the genetic and phenotypic differences in body-shape between women and men, the eQTL associations of genes associated with body-shape indices were in line with an involvement of adipose and vascular tissues, but also intriguingly of female reproductive organs in men as well as in women. This apparently paradoxical association in men may not be completely illogical, as female reproductive organs would be responsive to oestrogen-activated pathways and the formation of the Mullerian duct, from which the uterus and its adnexa are derived, is regulated by WNT signalling pathways⁵⁹, which may have pleiotropic roles.

We have previously demonstrated the need to adjust body-shape indices for BMI prior to using them in statistical models examining phenotypic associations⁹. We have now shown that it is also essential to account correctly for height, in order to avoid an "over-adjustment" arising from constraining the relationship between weight and height to a fixed proportion in BMI. In the case of $WC_{adj}BMI$ and $HC_{adj}BMI$, the adjustment for BMI does not reflect correctly the scaling of WC or HC with height and introduces phenotypic and genetic associations with height. When WC and HC are adjusted for height, in addition to BMI or weight, as in the allometric indices ABSI_{UKB} and HI_{UKB}, the relationship between weight and height is unconstrained and the correlation of WC and HC with height, as well as with BMI, is minimised. In the case of WHR_{adi}BMI, the phenotypic and genetic association patterns were very similar to the allometric counterpart WHI_{UKB}, despite the lack of additional adjustment for height, because the scaling coefficients for weight and height in individuals with white British ancestry were in proportion 1:2, similar to BMI. Nevertheless, given the large ethnic variability in body shape, this may not be universally applicable to other ethnicities. It would thus be advisable to use body-shape indices adjusted for height, as well as for BMI or weight, in order to minimise correlations with body size. Evaluating correctly the pleiotropic contributions is particularly important as many variants and genes associated with allometric body-shape indices are apparently related to growth and regulatory factors and a sizeable proportion of them have previously been reported in association with height. A mechanistic parallel with a pathway affecting either height or regional size when dysregulated in different periods of life could be drawn with growth hormone signalling. Growth hormone excess in adolescents is associated with increased height but an excess in adults, after epiphyseal closure, is associated with regional enlargement of the hands, feet and the face, i.e. acromegaly. Growth hormone signalling is further related to metabolic alterations and some cancers⁶⁰. A similar principle could be relevant to other growth or regulatory factors.

Our study benefited from a large sample size, anthropometric measurements performed by trained personnel according to standardised protocols and access to established bioinformatics pipelines. There were, however, several limitations. It should first be acknowledged, that the UK Biobank cohort is not representative of the UK population at the time of recruitment and is subject to a healthy volunteer bias which should be considered when interpreting the findings⁶¹. In addition, we lacked a validation cohort of similar ethnicity to the discovery dataset, although this would affect mainly the validity of associations with lower and borderline significance, while the highest-ranked genetic polymorphisms showed very strong and convincing associations. We also lacked datasets of comparable size including individuals with different ethnic backgrounds, as less than 5% of UK Biobank participants reported black or Asian ethnic background. Further, there were no reliable imaging measures of body composition (i.e. from dual-emission X-ray absorptiometry scans or magnetic resonance imaging) with comparable sample size. Furthermore, we did not perform a meta-analysis of waist and hip indices, since there are no other GWAS of ABSI and HI and we have already argued that WC_{adi}BMI and HC_{adi}BMI are not reliable body-shape indices. It would be useful, however, to consider a multi-trait meta-analysis approach⁶² when studies of other ethnicities are available. We did not specifically adjust our analyses for geographical location, which has been highlighted as a source of residual confounding of associations with BMI, bioelectric impedance fat mass measurements and height in UK Biobank¹³, as we minimised the number of adjustment variables to avoid introducing collider bias. There were, however, no matches of the reported variants related to geographical location with the main variants identified in our study, hence a confounding from fine-scale population structure is unlikely. Further, we could not perform mechanistic investigations linking genetic polymorphisms to adrenal function or cancer, which were beyond the scope of our study. Lack of Mendelian randomisation analysis, which may support a causal effect of the identified variants and genes on cancer and may further clarify the relationships in the network of mediators⁶³, is clearly a limitation. This, however, was also beyond the scope of the current study. Due to the large heterogeneity of cancers, associations would need to be consider separately for each individual cancer location and for each major histological type and the question of potential pleiotropy, which is very likely, would need to be addressed in detail. It should last be noted, that although examining $ABSI_{UKB}$ and HI_{UKB} enabled the identification of variants associated exclusively with the abdominal or the gluteofemoral regions, these associations may be determined by features other than fat accumulation, e.g. the status of lean mass in the gluteofemoral region. It would therefore be important to examine the association patterns of genetic variants identified for anthropometric indices with measures of body compositions, which would become available in the future for a larger part of UK Biobank participants.

In conclusion, in the first GWAS of allometric body-shape indices, we have identified novel genetic variants and genes and have obtained unbiased association statistics for individuals with white British ancestry, which would inform future studies of fat distribution, body shape and the disorders associated with them. The highestranked genes associated with body-shape indices have previously been associated in mechanistic studies with adrenal cell renewal, vascularisation and cancer development and progression, in addition to their functions in adipose tissue. The comparison of allometric and traditional body-shape indices demonstrated that adjustment of body-shape indices for height, as well as for BMI or weight, is warranted to avoid associations with height, which were more pronounced for $WC_{adj}BMI$ and $HC_{adj}BMI$. Differences between allometric and traditional body-shape indices with respect to their phenotypic and genetic associations were minimal when accounting correctly for height.

Methods

Study participants. UK Biobank is a prospective cohort with an ongoing follow-up, including 502,543 participants. Recruitment and data collection have previously been described^{64,65}. We excluded in total 95,846 participants. Exclusions were determined by a lack of genetic data or withdrawn consent (n = 15,229); outliers for heterozygosity or missing genotyping rate, or sex chromosome aneuploidy, or a mismatch between genetic and self-reported sex (n = 843); age below 40 or above 70 years when attending an assessment centre at baseline (n = 14); missing weight, height, waist or hip circumference measurements (n = 2,097); or pregnancy at baseline (n = 115). We further restricted the selection to participants with white British ancestry, defined by UK Biobank according to their genetic characteristics (excluded n = 77,548). The final dataset comprised 406,697 participants.

Genotyping and imputation. Genotyping, imputation and quality control were performed centrally by UK Biobank and have previously been described⁶⁵. Two genotyping arrays were used: Applied Biosystems UK BiLEVE Axiom Array (~ 50,000 participants) and a closely related Applied Biosystems UK Biobank Axiom Array (~ 450,000 participants).

Outcome measures and indices. We converted all anthropometric measurements to body-shape indices with adjustments either for BMI in linear models, or for weight and height in log-linear models. BMI was calculated by dividing weight (kg) by squared height (m).

For traditional body-shape indices, we used residuals of linear models regressing each of WC (cm), HC (cm) or WHR on BMI (WHR_{adj}BMI, WC_{adj}BMI, HC_{adj}BMI). For the corresponding allometric body-shape indices, we used ABSI and HI, and created a new waist-to-hip index (WHI), in order to complete the set. The published formulas for ABSI and HI have previously been derived for participants in the National Health and Nutrition Examination Survey (NHANES)^{5,6}:

ABSI = WC * Weight^{$$-2/3$$} * Height ^{$5/6$}

$$HI = HC * Weight^{-0.482} * Height^{0.310}$$

To avoid phenotypic correlations between anthropometric indices arising from differences in the anthropometric characteristics of UK Biobank and NHANES participants and to enable comparability with traditional residual methods, we calibrated ABSI and HI, as well as WHI, for UK Biobank participants (ABSI_{UKB}, HI_{UKB} and WHI_{UKB}). We derived the power coefficients from linear models regressing each of log-transformed WC (cm), HC (cm) or WHR on log-transformed weight (kg) and height (m) measured at baseline:

$$\log (Measure) \sim \beta * \log (Weight) + \gamma * \log (Height)$$

and generated allometric body-shape indices according to the general formula:

Index = Measure * Weight<sup>-
$$\beta$$</sup> * Height^{- γ}

where β and γ are the regression coefficients for weight and height (Supplementary Table S6). We included ABSI and HI calculated with the published regression coefficients from NHANES only for the phenotypic comparisons.

The absolute values of the power coefficients for weight and height for WHR_{UKB} were close to a ratio of 1:2 in both men and women, which corresponds to the relationship weight/height² in BMI (Supplementary Table S6). Therefore, in analogy to ABSI, for which the published coefficients were derived by rounding the power coefficients obtained in NHANES to simple fractions⁵, we generated for comparison a simplified version of WHI with the formula:

WHI = WHR *
$$[Weight(kg)/Height^{2}(cm)]^{-1/4}$$

As body-shape patterns show distinct differences between sexes¹⁰, we generated all indices and performed all statistical analyses separately for men and women. We used Blom's method for inverse normal transformation of anthropometric indices (package *RNOmni* in R), as in⁶⁶.

Association testing. We obtained Pearson's coefficient for partial phenotypic correlation between anthropometric indices with function *pcor* (package *ppcor* in R), adjusting for age at baseline.

We used Bayesian linear mixed-model analysis BOLT-LMM v2.3 for genome-wide association testing^{67,68}, which incorporates in the statistical algorithm a correction for population stratification and thus accounts for the relatedness between UK Biobank participants. To estimate the parameters of the LMM, we used linkage disequilibrium (LD) scores from the 1000 Genomes EUR samples from individuals with European ancestry⁶⁹ and selected a coreset with high-quality genetic variants by restricting the list of variants released by UK Biobank after the centrally performed quality control to variants with missingness < 0.015, minor allele frequency MAF > 5% and Hardy–Weinberg exact test $P > 1*10^{-6}$ (Supplementary Table S7). We performed the main analyses with variants with MAF $\ge 1\%$ and imputation quality factor INFO > 0.1. We adjusted all models for age at baseline, age squared and a binary indicator of genotyping array. Supplementary Table S8 includes quality control parameters for BOLT-LMM. We obtained heritability estimates (h_g^2) from BOLT-LMM, based on the genetic relationship matrix. We report association statistics based on the BOLT-LMM infinitesimal models, which use a Gaussian single nucleotide polymorphism (SNP) effect prior.

Mapping, annotation and prioritisation of genetic variants. Individual genetic variants were defined with the chromosome, the genomic position in base pairs, the minor (alternative) allele and the major (reference) allele. For some SNPs there were more than one alternative alleles, hence the total number of genetic variants associated with a given body-shape index could be higher than the total number of SNPs. We used the SNP2GENE process of the web application Functional Mapping and Annotation (FUMA) v1.3.6a⁷⁰ to perform positional mapping, clumping and annotation of genetic variants according to the p-values of the association statistics obtained from the corresponding BOLT-LMM infinitesimal model for variants with high imputation quality (INFO>0.9). Independent significant SNP were defined as variants with genome-wide significance $(P \le 5^*10^{-8})$, which were in approximate linkage equilibrium with each other, at $r^2 < 0.6$ within a 1 Mb window. Variants with nominal significance (P < 0.05) in LD with an independent significant SNP ($r^2 \ge 0.6$ within a 1 Mb window) formed the corresponding LD block of candidate SNPs. To map LD, we used the "UKB release2b 10 k White British" panel in FUMA. Independent significant SNPs in LD with each other ($r^2 \ge 0.1$ within 1 Mb window) were consolidated in a clump, represented by a lead SNP with the lowest p-value. Lead SNPs with corresponding clump boundaries less than 250 kb apart were merged into a genetic risk locus, represented by the locus lead SNP with the lowest p-value. All candidate SNPs were mapped to genes within a maximum distance of 1 kb (based on their genomic position) and for functional consequences (based on Ensemble genes v92) with ANNOVAR employed in FUMA⁷¹. We used as a measure of pathogenicity of a given variant the deleteriousness score (Combined Annotation Dependent Depletion (CADD) score) calculated in FUMA and evaluated this against the recommended cut-off of 12.37¹². We additionally annotated the functionality of candidate SNPs with genome-wide significance with Ensembl Variant Effect Predictor (VEP) v90⁷². This maps each variant to the gene or location with the nearest protein-coding transcription start site within a window of 200 kb from the position of the examined variant.

Gene-based association analysis. For gene-level, gene-set and gene-property analysis, we used Multimarker Analysis of GenoMic Annotation (MAGMA) v.1.08⁷³, which is employed in FUMA v1.3.6a with default settings. The reference panel for LD and the Ensemble version for assigning SNPs to genes (within a symmetric window of 1 kb from both sides) were set as described for SNP annotation and prioritisation above.

Gene analysis was based on the summary SNP statistics of all variants with INFO > 0.9 (i.e. including the complete distribution). A SNP-wide mean model was used to calculate an association statistic and p-value for each gene. Significant genes were considered those with P < 0.05, after applying Bonferroni correction for multiple comparisons for 19,088 identified protein-coding genes. Significant genes with boundaries within 250 kb of each other were clumped in genomic risk regions, represented by a lead gene with the lowest *P*-value.

Competitive gene-set analysis was performed as a gene-level linear regression model to test whether the genes included in each gene-set (a binary indicator) showed stronger (positive) associations with the phenotype than other genes, generating a one-sided p-value. The model was conditioned by default on gene size, gene density (reflecting LD between SNPs in the gene), the inverse of the mean MAF in the gene (to account for potential power loss in very low MAF SNPs), and the log values of the three variables⁷³. FUMA v1.3.6a uses gene sets obtained from the Molecular Signatures Database (MSigDB) v7.0, including curated gene sets from online pathway databases and gene ontology (GO) terms. The significance of associations with gene sets was evaluated at P < 0.05, after Bonferroni correction for 15,485 examined gene sets.

The Genotype-Tissue Expression (GTEx) v8.0 database was used to perform gene-property analysis, as implemented in FUMA v1.3.6a, which examines associations with expression Quantitative Trait Loci (eQTL). This uses a similar linear regression model to gene-set analysis, but with the average \log_2 transformed gene expression values per tissue (a continuous variable), to test the (positive) relationship between highly expressed genes in a specific tissue and genetic associations represented by gene-level statistics, generating a two-sided p-value. The significance of associations with eQTLs was evaluated at P < 0.05, after Bonferroni correction for 54 individual tissue types. **Match against published reports.** We matched all candidate SNPs and the significant genes identified by MAGMA in FUMA against SNPs and genes previously reported in the NHGRI-EBI GWAS Catalog¹¹ (accessed on 07/04/2021, https://www.ebi.ac.uk/gwas/home) in association with anthropometric indices. Novelty for a genomic risk locus was concluded if there was no match of any candidate SNP included in the locus with a SNP reported in the NHGRI-EBI GWAS Catalog in association with the corresponding traditional index. Similarly, novelty for a genomic risk region was concluded if there was no match of any significant gene included in the region with a gene reported in the NHGRI-EBI GWAS Catalog in association with the corresponding traditional index. Match with reported associations with height or cancer were compared at the level of independent significant gene.

Difference analysis. To test for sex-dimorphic effects, we calculated a t-statistic as follows:

$$t_{sex} = \frac{\beta_{men} - \beta_{women}}{\sqrt{SE_{men}^2 + SE_{women}^2 - 2 * r_{sex} * SE_{men} * SE_{women}}}$$

where β are the regression coefficients for a given variant in men or women, SE are the corresponding standard errors and r_{sex} is the Spearman rank correlation coefficient between the regression coefficients in men and women for all examined genome-wide variants^{3,10}. We extracted the corresponding p-values (p_{sex}) from a t-distribution with function *pt* in R. We used a similar formula to compare heritability between sexes, replacing the regression coefficients with heritability estimates in men and women.

To test for difference in effect sizes between allometric and traditional indices, we calculated, separately in women and men, t-statistics and the corresponding $p_{\text{difference}}$ with a similar formula to the one used for sex dimorphisms:

$$t_{allo - trad} = \frac{\beta_{allo} - \beta_{trad}}{\sqrt{SE_{allo}^2 + SE_{trad}^2 - 2 * r_{allo - trad} * SE_{allo} * SE_{trad}}}$$

where β are the regression coefficients for a given variant for the corresponding allometric and traditional index in a pair (i.e. WHI_{UKB} and WHR_{adj}BMI, ABSI_{UKB} and WC_{adj}BMI, HI_{UKB} and HC_{adj}BMI), SE are the corresponding standard errors and r_{allo-trad} is the Spearman rank correlation coefficient between the regression coefficients for the allometric and traditional index for all examined genome-wide variants. To test for difference in effect sizes between the allometric waist and hip indices, the parameters for allometric and traditional indices were replaced with the corresponding parameters for ABSI_{UKB} and HI_{UKB} and r_{ABSI-HI} was calculated as above.

As differences in effect sizes were considered only for independent significant SNPs identified for at least one of the two compared sexes or indices and this would vary between comparisons, we evaluated differences relative to a single universal conservative cut-off $P < 5 \times 10^{-6}$, which is equivalent to a Bonferroni correction for 10,000 comparisons and reflects strong evidence for association.

We used R version 3.6.1. for the management of data and results⁷⁴.

Ethical approval and consent to participate. This research was conducted according to the principles expressed in the Declaration of Helsinki. The UK Biobank cohort has been approved by the North West Multicenter Research Ethics Committee, UK (Ref: 16/NW/0274). Written informed consent has been obtained from all study participants. The current study was approved by the UK Biobank access management board under application 41952. Participants who had withdrawn consent by the time of the analysis were excluded from the dataset.

Data availability

The data supporting the findings of the study are available *to bona fide* researchers upon approval of an application to the UK Biobank (https://www.ukbiobank.ac.uk/researchers/) and a material transfer agreement. The results from FUMA with additional annotations are included in Supplementary Table S9 (candidate SNPs) and Supplementary Table S10 (all other analyses) and the gene-level analysis from MAGMA and a list of the sets and traits from the NHGRI-EBI GWAS Catalog used for matching are included in Supplementary Table S10. Summary statistics from BOLT-LMM infinitesimal models for all candidate SNPs identified in our study are included in Supplementary Table S11. The SNP2GENE output will also be made available upon publication on the FUMA website (https://fuma.ctglab.nl/) and locus lead SNPs will be uploaded to the NHGRI-EBI GWAS Catalog (https://www.ebi.ac.uk/gwas/home).

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Author contributions

S.C., K.K.T. and E.R. conceived and designed the study. K.K.T. and E.E. provided statistical advice. S.C. led the research and performed the statistical analysis. S.C. had full access to all of the data in this study and takes responsibility for the integrity of the data and the accuracy of the data analysis. S.C. drafted the paper with contributions from E.E, K.K.T and E.R. All authors, i.e. S.C., E.E., E.R. and K.K.T. were involved in the interpretation of the results, the critical revisions of the paper, and the approval of the final version of the manuscript.

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Competing interests

The authors declare no competing interests.

Additional information

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