A putative causal relationship between genetically determined female body shape and posttraumatic stress disorder

Citation
Polimanti, R., A. B. Amstadter, M. B. Stein, L. M. Almli, D. G. Baker, L. J. Bierut, B. Bradley, et al. 2017. "A putative causal relationship between genetically determined female body shape and posttraumatic stress disorder." Genome Medicine 9 (1): 99. doi:10.1186/s13073-017-0491-4. http://dx.doi.org/10.1186/s13073-017-0491-4.

Published Version
doi:10.1186/s13073-017-0491-4

Permanent link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:34651976

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA

Share Your Story
The Harvard community has made this article openly available. Please share how this access benefits you. Submit a story.

Accessibility
A putative causal relationship between genetically determined female body shape and posttraumatic stress disorder

Renato Polimanti 1*, Ananda B. Amstadter 2, Murray B. Stein 3,4,5, Lynn M. Almli 6, Dewleen G. Baker 3,5, Laura J. Bierut 7, Bekh Bradley 6,8, Lindsay A. Farrer 9, Eric O. Johnson 10, Anthony King 11, Henry R. Kranzler 12, Adam X. Maihofer 3, John P. Rice 6, Andrea L. Roberts 13, Nancy L. Saccone 14, Hongyu Zhao 15, Israel Liberzon 11,16, Kerry J. Ressler 17,18, Caroline M. Nievergelt 3,5, Karestan C. Koenen 19,20,21, and Joel Gelernter 1,22 for The Psychiatric Genomics Consortium Posttraumatic Stress Disorder Workgroup

Abstract

Background: The nature and underlying mechanisms of the observed increased vulnerability to posttraumatic stress disorder (PTSD) in women are unclear.

Methods: We investigated the genetic overlap of PTSD with anthropometric traits and reproductive behaviors and functions in women. The analysis was conducted using female-specific summary statistics from large genome-wide association studies (GWAS) and a cohort of 3577 European American women (966 PTSD cases and 2611 trauma-exposed controls). We applied a high-resolution polygenic score approach and Mendelian randomization analysis to investigate genetic correlations and causal relationships.

Results: We observed an inverse association of PTSD with genetically determined anthropometric traits related to body shape, independent of body mass index (BMI). The top association was related to BMI-adjusted waist circumference (WCadj; R = −0.079, P < 0.001, Q = 0.011). We estimated a relative decrease of 64.6% (95% confidence interval = 27.5–82.7) in the risk of PTSD per 1-SD increase in WCadj. MR-Egger regression intercept analysis showed no evidence of pleiotropic effects in this association (P pleiotropy = 0.979). We also observed associations of genetically determined WCadj with age at first sexual intercourse and number of sexual partners (P = 0.013 and P < 0.001, respectively).

Conclusions: There is a putative causal relationship between genetically determined female body shape and PTSD, which could be mediated by evolutionary mechanisms involved in human sexual behaviors.

Keywords: Trauma, Genetics, Women, Anthropometric traits

Background

Posttraumatic stress disorder (PTSD) is a mental illness that affects susceptible individuals who have experienced, witnessed, or been confronted with an event involving actual or threatened dangers [1]. Individuals affected by PTSD present symptoms that, according to the DSM-5, include four diagnostic clusters (re-experiencing, avoidance, negative cognitions and mood, and arousal) [2]. At least one in nine American women meet lifetime diagnostic criteria for PTSD and women’s risk of PTSD is twice that of men [3]. Among women, PTSD is also more likely to be chronic and associated with greater functional impairment and distinct neurobiological profiles [4, 5]. Evidence suggests that women are more vulnerable than men to develop PTSD when exposed to similar traumas [6].

Women’s higher risk for PTSD likely arises from biologic and social/environmental factors. For example, sex differences in trauma exposure may partially explain the differential risk for PTSD. Women are more likely to be
exposed to sexual violence, which has a high conditional risk of PTSD [7]. Alternatively, the pattern of results from twin studies suggests that PTSD may be more heritable in women than in men [8, 9] and results from a recent genome-wide investigation also demonstrate higher molecular heritability in women [10]. Sex hormones, particularly estradiol and progesterone, have been implicated as mediators of this risk, suggesting that their effects on biological, behavioral, and cognitive pathways augment the vulnerability to PTSD [11]. A recent epigenome-wide association study supported this possible mechanism; it reported an estrogen-dependent association of HDAC4 with fear in female mice and women with PTSD [12].

Genetic data can be used to investigate this complex network of multiple factors. Polygenic risk score (PRS) analyses and Mendelian randomization (MR) studies are less biased by confounders than observational studies and can be used to investigate the predisposition to complex traits, identifying shared molecular pathways, pleiotropy, and causal relationships [13, 14]. Here, we used summary statistics from female-specific genome-wide association studies (GWAS) of traits that may be related to disease pathogenesis—i.e., anthropometric characteristics [15], reproductive physiology and behaviors [16], and educational attainment [17]—to examine the mechanisms underlying women’s vulnerability to PTSD. Although these traits were previously indicated as related to PTSD in women, testing a wide range of genetic correlations can provide useful etiological insights and help prioritize likely causal relationships [18]. A greater understanding of the biological and external social processes involved in PTSD will help to address the critically important interplay between two important public health issues: PTSD and women’s health.

Methods

Cohorts from the Psychiatric Genomics Consortium PTSD Workgroup

The individual data from the subjects included were made available by the members of the Psychiatric Genomics Consortium PTSD Workgroup. Details regarding participants, genotyping, quality control, imputation, and ancestry assignment were reported previously [10]. In the present study, we used data regarding female participants from seven cohorts: Collaborative Genetic Study of Nicotine Dependence [19]; Family Study of Cocaine Dependence [20]; Yale-Penn [21]; Grady Trauma Project [22]; Marine Resilience Study [23]; Nurses’ Health Study [24]; and Ohio National Guard [25]. Because the individuals in the GWAS that were used as sources for summary statistics for the traits of interest were of European descent, we restricted our analysis to women of that population. Quality control criteria for ancestry assignment and sex check are reported in our previous publication [10]. We considered imputed genotypes to maximize a consistent SNP (Single Nucleotide Polymorphism) panel between the training and testing sets. Imputed SNPs with high imputation quality (genotype call probability ≥ 0.8), minor allele frequency ≥ 1%, missingness per marker ≤ 5%, missingness per individual ≤ 5%, and Hardy-Weinberg equilibrium P > 10^{-4} were retained. After applying these quality control criteria, we retained information on 4,875,110 variants in a final sample of 3577 women (966 PTSD cases and 2611 trauma-exposed controls). To investigate the specificity of our findings for PTSD in women, we investigated a sample of 4628 men (501 PTSD cases and 4127 trauma-exposed controls) from the same cohorts of the Psychiatric Genomics Consortium PTSD Workgroup. The same quality control criteria were applied to the data from men. Principal component analysis of the final sample was conducted using PLINK 1.9 [26] after linkage disequilibrium (LD) pruning (R^2 < 0.2).

Polygenic risk score analysis

We conducted cross-phenotype PRS analyses using PRSice software [27] (available at http://prsice.info/). This method permits testing whether the polygenic components of inheritance have a substantial effect on multiple complex traits [28]. For polygenic profile scoring, we used summary statistics generated from female-specific GWAS of 17 traits (Table 1) conducted by the Genetic Investigation of ANthropometric Traits (GIANT) Consortium [29–32], the Reproductive Genetics (Reprogen) Consortium [33, 34], and the Social Science Genetic Association Consortium (SSGAC) [35, 36]. None of the PGC-PTSD cohorts was included in these previous GWAS and, thereby, no large overlap is expected among these samples. We considered multiple association P value thresholds (PT = 10^{-5}, 10^{-7}, 10^{-6}, 10^{-5}, 10^{-4}, 0.001, 0.05, 0.3, 0.5) for SNP inclusion, i.e. we conducted a high-resolution analysis. The PRS were calculated after using P value-informed clumping with a LD cut-off of R^2 = 0.3 within a 500-kb window, and excluding the Major Histocompatibility Complex region of the genome because of its complex LD structure. The PRS that were generated were fitted in regression models with adjustments for the top ten ancestry principal components to calculate Nagelkerke’s R as the figure of merit for prediction ability. We applied false discovery rate (FDR) multiple testing correction (Q < 0.05) to correct for the number of PRS and thresholds tested (a total of 170 tests). A gene ontology (GO) enrichment analysis was conducted based on the PRS results: the variants included in the significant PRS with direction concordant with PRS direction were considered in the enrichment analysis. Variants were mapped to the nearest genes and then the gene loci were entered.
into the enrichment analysis conducted using the PAN-
THER v11.1 Overrepresentation Test (release 20160715; 
Reference List: Homo Sapiens) [37]. FDR correction was 
applied to the enrichment results to account for multiple 
testing. The GO enrichment results were further investi-
gated using REVIGO [38]. Specifically, GO enrichments 
were used to make a graph-based visualization considering 
an allowed similarity of 0.7, UniProt as reference database, 
and SimRel method as the semantic similarity measure.

Two-sample Mendelian randomization analysis
The two-sample MR analysis was conducted using the R 
package MendelianRandomization (available at https://
cran.r-project.org/web/packages/MendelianRandomization /
index.html) and the GWAS summary statistics. This is a 
strategy to test cause–effect associations, in which evidence 
on the associations of genetic variants (i.e. instrumental variable) 
with the risk factor (i.e. exposure) and with the outcome comes from two sources [39]. We used 
SNPs associated with waist circumference adjusted for 
body mass index (BMI) (WC_{adj} P < 10^{-8} N = 31), which 
showed the most significant correlation with PTSD in our

### Table 1 GWAS summary statistics used for polygenic risk scoring

| Consortium | Trait Abbreviation | Women N | Variant N | Data link |
|-----------|-------------------|---------|-----------|-----------|
| Genetic Investigation of ANthropometric Traits (GIANT) | Body mass index | 128,698 | 2,494,584 | http://portals.broadinstitute.org/collaboration/giant/index.php/GIANT_consortium_data_files |
| | BMI (age ≤ 50 years) | 52,270 | 2,407,668 |
| | BMI (age ≥ 50 years) | 86,731 | 2,407,629 |
| | Height | 64,811 | 2,748,519 |
| | Hip circumference | 85,797 | 2,466,814 |
| | HIP adjusted for BMI | 84,787 | 2,466,814 |
| | Waist circumference | 90,543 | 2,444,344 |
| | WC adjusted for BMI | 89,759 | 2,473,027 |
| | Waist-hip ratio | 85,466 | 2,466,093 |
| | WHR adjusted for BMI | 84,222 | 2,467,768 |
| | WHR radial (age ≤ 50 years) | 34,523 | 2,366,527 |
| | WHR radial (age ≥ 50 years) | 56,150 | 2,366,567 |
| Reproductive Genetics (Reprogen) | Age at menarche | 182,416 | 2,441,881 | http://www.reprogen.org/data_download.html |
| | Age at menopause | 69,360 | 2,418,695 |
| Social Science Genetic Association Consortium (SSGAC) | Age at first birth | 158,856 | 2,470,136 | http://www.thessgac.org/data |
| | Education years | 122,427 | 7,643,300 |
| | Number of children ever born | 222,837 | 2,471,862 |
follow up a possible “evolutionary link” between female body shape and PTSD, we tested the association of genetically determined WC_{adj} with age at first sexual intercourse (AFS) and number of sexual partners (NSP). For AFS, we used female-specific summary statistics from a recent GWAS, including women of European descent only [42]. This previous GWAS was conducted considering a range of AFS, which also includes a small fraction of individuals who had their first sexual intercourse before 12 years of age. For NSP, we used UK Biobank data on a total of 58,356 unrelated women of European descent. We calculated the association of the top 31 SNPs associated with WC_{adj} (\(P < 10^{-8}\)) with NSP using a linear regression model. Before being entered into the analysis, NSP was adjusted for age and the top ten ancestry principal components and then normalized using appropriate Box–Cox power transformations.

Data availability
Data supporting the findings of this study are available within the article and its Additional files. GWAS Summary level data used to calculate PRS in this study were obtained from links reported in Table 1.

Results
We used summary statistics generated from female-specific GWAS of 17 traits (Table 1) to yield polygenic profile scoring and tested the association of the derived PRS with PTSD considering multiple PT and using samples from the Psychiatric Genomics Consortium PTSD Workgroup (966 PTSD cases and 2611 trauma-exposed controls). Six PRS showed correlations with PTSD that survived FDR multiple testing correction (\(Q < 0.05\); Table 2; Additional file 1). Although LD score regression is designed to analyze large sample sizes [18], using this method we nevertheless replicated the directions of five correlations and two of them survived multiple testing correction (Additional file 2). The strongest PRS result was observed for WC_{adj}; women with high genetically predicted WC_{adj} have low PTSD risk (PT: \(10^{-8}\); \(R = -0.079, P < 0.001, Q = 0.011\)). Additional file 3 shows the distribution of WC_{adj} PRS in PTSD cases and controls. The same variants included in the WC_{adj} PRS also drove four of the other associations observed: waist–hip ratio (WHR, PT: \(10^{-7}\); \(R = -0.064, P = 0.001, Q = 0.023\)), WHR adjusted for BMI (WHR_{adj}_{BMI}, PT: \(10^{-7}\); \(R = -0.059, P = 0.001, Q = 0.030\)), WHR_{adj} in women aged < 50 years (WHR_{adj:<50}, PT: \(10^{-8}\); \(R = -0.076, P = 1.25 \times 10^{-4}, Q = 0.011\)), and WHR_{adj} in women aged > 50 years (WHR_{adj:>50}, PT: \(10^{-8}\); \(R = -0.065, P = 0.001, Q = 0.023\)). Additional file 4 reports the strong correlations of WC_{adj} PRS with these other anthropometric traits. The association between WC_{adj} PRS and PTSD appears to be specific to women; no correlation was observed in men (Additional file 5).

Among the significant PRS, the finding related to Age at First Birth (AFB) PRS is the only one that appears to be independent from the WC_{adj} signal because its top correlation is due to the cumulative effects of a large number of variants (AFB top-PT = 0.3, SNP N = 64,579) and not to the effect of a small number of strongly associated loci (WC_{adj} top-PT = \(10^{-8}\), N = 31). We observed that women with a high genetically predicted AFB have low PTSD risk (\(R = -0.064, P = 0.001, Q = 0.023\)). Additional file 6 shows the distribution of AFB PRS in PTSD cases and controls. No association between AFB PRS and PTSD was observed in men (Additional file 7). Since this genetic overlap appears to be related to the additive effect of a large number of variants (N = 64,579), we investigated which molecular mechanisms are shared between PTSD and AFB. The GO enrichment analysis based on AFB results highlighted several potential molecular mechanisms involved (\(Q < 0.05\); Additional file 8, respectively). Because we observed a large number of significant GO enrichments (\(Q < 0.05\) related to the AFB–PTSD result (N = 40), we conducted a similarity-based network analysis among the significant GO enrichments. This analysis highlighted a GO cluster (Additional file 9) related to multiple signaling mechanisms (e.g., calcium-mediated signaling and MAPK cascade).

To investigate further the WC_{adj}–PTSD genetic correlation, we applied a MR approach to assess the causative mechanisms of this association. We used an inverse-variance-weighted method to calculate an overall estimate of the causal effect considering the coefficients related to the association of these variants with PTSD and

| Table 2 | Top genetic correlations between PTSD and PRS tested |
|---------|-----------------------------------------------|
| PRS     | PT   | SNP N | R        | P value | Q value |
| WC_{adj} | \(10^{-8}\) | 31   | -0.079  | 6.77 \times 10^{-5} | 0.011 |
| WHR_{adj:<50} | \(10^{-8}\) | 19   | -0.076  | 1.25 \times 10^{-4} | 0.011 |
| WHR_{adj:>50} | \(10^{-8}\) | 19   | -0.065  | 0.001   | 0.023  |
| AFB      | 0.3   | 64,579 | -0.064  | 0.001   | 0.023  |
| WHR      | \(10^{-7}\) | 32   | -0.064  | 0.001   | 0.023  |
| WHR_{adj} | \(10^{-7}\) | 51   | -0.059  | 0.003   | 0.030  |
| Menarche | \(10^{-3}\) | 1510 | 0.049   | 0.013   | 0.074  |
| BMI_{<50} | \(10^{-7}\) | 26   | 0.048   | 0.015   | 0.077  |
| HIP      | \(10^{-8}\) | 94   | 0.045   | 0.024   | 0.108  |
| Height   | 0.1   | 25,369 | 0.042   | 0.035   | 0.144  |
| WC       | \(10^{-8}\) | 20   | 0.034   | 0.082   | 0.266  |
| BMI_{<50} | \(10^{-5}\) | 104  | 0.033   | 0.099   | 0.306  |
| BMI      | \(10^{-4}\) | 316  | 0.032   | 0.103   | 0.312  |
| N3B      | \(10^{-6}\) | 2    | -0.029  | 0.141   | 0.386  |
| H3P_{adj} | 0.05  | 13,338 | 0.029   | 0.147   | 0.390  |
| EDU      | \(10^{-8}\) | 17   | -0.021  | 0.294   | 0.641  |
| Menopause| \(10^{-5}\) | 163  | 0.018   | 0.360   | 0.686  |
WC_adj (Fig. 1; Additional file 10). This analysis showed a significant association of effect sizes of WC_adj-associated SNPs with effect sizes estimated in our PTSD sample (OR = 0.35; 95% confidence interval [CI] = 0.17–0.73; \( P = 0.008 \)). This translated to a relative decrease of 64.6\% (95% CI = 27.5–82.7) in the risk of PTSD per 1-SD increase in WC_adj. This result was consistent considering multiple MR approaches (Additional file 11). Furthermore, the consistency between PRS and MR analyses confirms that there is no sample overlap that is biasing our results. Indeed, these two approaches are susceptible to biases from sample overlaps, but the bias would be in opposite directions: risk of false-positive results in PRS and risk of false-negative results in two-sample MR [43]. Considering the MR-Egger regression intercept, we also observe that the markers did not show evidence of pleiotropic effects (\( P_{\text{pleiotropy}} = 0.979 \)), suggesting that the association of these variants with PTSD is mediated by their effect on body shape. To investigate further the potential causal relationship between PTSD and female body shape, we tested whether the PTSD PRS correlated with WC_adj. A causal relationship between two traits should show asymmetry in the effect sizes of associated variants [41]. We observed asymmetry between PTSD and WC_adj; PTSD-associated variants showed no effect on WC_adj (\( P > 0.10 \); Additional file 12) but strongly predicted BMI (\( P = 5.68 \times 10^{-7} \)) and other BMI-related anthropometric traits (Additional file 13). These results support a causal relationship between WC_adj and PTSD.

We tested whether genetically determined female body shape correlated with sexual behaviors in women using female-specific genetic information regarding AFS and NSP (Additional file 10). Considering a penalized robust MR-Egger regression, we observed associations of WC_adj with AFS (Beta = −0.25, SE = 0.10, \( P = 0.013 \); Fig. 2) and NSP (Beta = 0.16, SE = 0.04, \( P < 0.001 \); Fig. 2) with significant evidence of pleiotropy in both results (\( P_{\text{pleiotropy}} = 0.017 \) and < 0.001, respectively), suggesting that the effects of these variants on sexual behaviors could be due to multiple mechanisms.

**Discussion**

We found associations of the genetic risk for PTSD in women with genetically determined female body shape and reproductive behaviors. These findings provide novel insight into the mechanisms that may underlie the genetic vulnerability to PTSD in women.

Our strongest result reflects an inverse relationship between genetically determined female body shape and the genetic risk for PTSD. We observed that SNPs associated with anthropometric traits related to body shape independently from BMI (i.e. WC_adj, WHR, WHR_adj, WHR_adj<50).

---

**Fig. 1** SNP-exposure (WC_adj associations) and SNP-outcome (PTSD associations) coefficients used in the MR analysis. Error bars (95% CIs) are reported for each association. The solid line represents the inverse-variance-weighted estimate.
WHR_{adj} > 50) are linked to the risk for PTSD. No analogous association was present in our male cohort. Human body shape is the result of strong evolutionary pressures, including sexual selection, with evident differences in selection between women and men [44]. The dissimilarities between sexes are largely due to adaptation mechanisms related to female reproductive function [44]. To facilitate childbirth, the pelvis is proportionately larger in women than in men. There is also variability in fat accumulation and distribution that are related to women’s need for energy to be used during pregnancy and nursing: women have a higher percentage of body fat; men accumulate fat in the abdominal region, while women accumulate fat in hips, buttocks, thighs, and the lower abdomen [45]. Consistent with the presence of different regulatory mechanisms related to human body shape, GWAS from the GIANT consortium identified several loci with sex-specific effects and most of them have larger effects in women than in men [30, 32]. Although these sex differences and their regulatory mechanisms are well-known, to our knowledge no large study has investigated the role of genetically determined body shape in women’s mental health. In our study, we observed that a genetically determined evolution-related body shape (i.e. low WC_{adj}, WHR, and WHR_{adj}) was associated with increased risk of PTSD. A small study reported an association between observational (i.e. phenotypically observed) WHR and PTSD concordant with our finding [46]. Our data are most consistent with a causal relationship between WC_{adj} and PTSD in women. Considering MR-Egger regression intercept, we did not observe a pleiotropic effect between WC_{adj} and PTSD. Conversely, there was an asymmetry in the effect sizes of associated variants: WC_{adj}-associated variants affect the risk of PTSD, but PTSD-associated variants do not affect female body shape. Previous evolutionary studies reported that body shape in women is one of the traits related to reproductive attractiveness that has been selected during human evolution [47]. We note that female reproductive attractiveness, which is determined by evolutionary selection pressures, is distinct from socio-cultural norms about female attractiveness that are known to differ over time and place [48]. Our data can be interpreted as supporting that specific anthropometric characteristics may be related to PTSD via their role in increasing the risk of traumatic experiences related to undesired sexual behavior. Further evidence of an apparent causal relationship between WC_{adj} and experiences related to sexual function is provided by the analysis of AFS and NSP. Genetically determined WC_{adj} is associated with AFS and NSP; this provides additional evidence of the relationship between the female body shape and behaviors related to sexual interactions. As indicated by the significant evidence of pleiotropy, the relationship between female body shape and sexual behavior is likely mediated by multiple factors, including behavioral mechanisms and physiological processes related to reproductive function. Unfortunately, we cannot investigate further the relationship between female body shape and traumatic specific experiences, such as sexual assault, because trauma type information is not currently available in the PGC PTSD dataset. We hope to be able to do it in the future. In particular, we believe that longitudinal studies that include detailed life course assessments of traumatic experience will be needed to better understand the role of female body shape in risk of sexual assault or other traumatic event exposure.

Our investigation also identified an association between AFB and PTSD: genetically predicted young maternal AFB is associated with an increase of PTSD risk. This is consistent with epidemiologic observations of an inverse association between maternal AFB and psychiatric distress: mothers giving birth in their teens have an increased risk of a lifetime behavior disorder, PTSD, or an anxiety disorder compared to women who were older.
at first birth [49]. The AFB–PTSD correlation is due to a genome-wide genetic overlap that includes thousands of independent variants, which is different from the body shape result, which was influenced by a small number of variants. This could indicate that these two traits share some pathways involved in their predisposition. Our enrichment analysis identified several molecular processes. Among them, we observed a large GO cluster of signaling mechanisms that includes several that are known to be involved in the response to traumatic experience, such as calcium-mediated signaling and the MAPK cascade [50, 51]. Observational studies have reported that early trauma exposures are associated with an increased risk of teen pregnancy. In particular, childhood sexual abuse is associated with increased risk for subsequent adolescent pregnancy [52, 53]. In line with this scenario, we note that the AFB GWAS [35] was conducted without excluding birth events from non-consensual sex. For instance, although it accounts for a small fraction of the sample investigated, some of the cohorts included in this previous analysis also included births during childhood. Accordingly, a possible explanation of the AFB–PTSD correlation is that childhood abuse and sexual victimization can contribute to AFB and PTSD, and some of the trauma-response mechanisms involved may be shared among these traits. These mechanisms appear to be different from those related to the WC_{adj} results. The relationship between AFB and PTSD seems to be due to a large genetic overlap (i.e. pleiotropy); WC_{adj} appears to be involved in a causal mechanism related to women’s vulnerability to PTSD (i.e. no pleiotropy).

Conclusions
In summary, using a genetic approach, we found two associations with the risk of PTSD in women related to body shape and reproductive behaviors. Both mechanisms appear to be female-specific (i.e. there was no such association in men) and they suggest that sex differences in trauma type prevalence and molecular mechanisms of trauma response contribute to the greater vulnerability to PTSD observed in women. Our study demonstrates how genetic research can successfully contribute to the dissection of biological and non-biological mechanisms relevant to women’s health.

Additional files

Additional file 1: Genetic correlations between PTSD and PRS tested in women. (DOCX 19 kb)
Additional file 2: PTSD genetic correlations tested with LD score regression analysis. (DOCX 12 kb)
Additional file 3: Distribution of WC_{adj} PRS in PTSD cases and controls. (DOCX 333 kb)

Additional file 4: Genetic correlation of WC_{adj} PRS (SNP N = 31, PT = 10^{-4}) with the other anthropometric traits associated with PTSD in women. (DOCX 12 kb)
Additional file 5: Correlation between WC_{adj} PRS and PTSD in men. (DOCX 12 kb)
Additional file 6: Distribution of AFB PRS in PTSD cases and controls. (DOCX 361 kb)
Additional file 7: Correlation between AFB PRS and PTSD in men. (DOCX 12 kb)
Additional file 8: Gene Ontology (GO) enrichments (Q value < 0.05) for variants with AFB GWAS P < 0.3 and effect direction concordant with the AFB-PTSD correlation. (DOCX 14 kb)
Additional file 9: Similarity network based on GO enrichment results related to the AFB-PTSD correlation. Bubble color indicates the GO P value; bubble size indicates the frequency of the GO term in the Gene Ontology Annotation (UniProt-GOA) Database which annotates all UniProt entries with GO terms. Highly similar GO terms are linked by edges in the graph, where the line width indicates the degree of similarity. (DOCX 179 kb)
Additional file 10: SNP-exposure and SNP-outcome coefficients of the variants in included in the WC_{adj} PRS. (DOCX 15 kb)
Additional file 11: SNP-exposure (WC_{adj} associations) and SNP-outcome (PTSD associations) coefficients used in the MR analysis. The lines (see legend) represents the results observed considering different methods. (DOCX 383 kb)
Additional file 12: Correlation between PTSD PRS and WC_{adj}. (DOCX 12 kb)
Additional file 13: Top results of the reverse PRS analysis of PTSD with respect to the main anthropometric traits. (DOCX 12 kb)

Abbreviations
AFB: Age at first birth; AFS: Age at first sexual intercourse; BMI: Body mass index; FDR: False discovery rate; GO: Gene Ontology; GWAS: Genome-wide association study; MR: Mendelian randomization; NSP: Number of sexual partners; PRS: Polygenic risk score; PTSD: Posttraumatic stress disorder; SNP: single nucleotide polymorphism; WC_{adj}: BMI-adjusted waist circumference; WHR: Waist–hip ratio; WHR_{adj}: WHR adjusted for BMI; WHR_{adj,50}: WHR in women aged < 50 years; WHR_{adj,50}+: WHR in women aged > 50 years

Acknowledgements
We thank study participants and research groups contributing to PGC-PTSD for sharing their data. Part of this research has been conducted using the UK Biobank Resource (application reference no. 19664). We acknowledge Naomi Wray of The University of Queensland (Brisbane, Australia) for insightful comments about the manuscript. We thank the members of the cited consortia of genome-wide association studies for making their data available. The members of the Psychiatric Genomics Consortium Posttraumatic Stress Disorder Workgroup are: Laramie E Duncan (Department of Psychiatry, Stanford University, Stanford, CA, USA, Broad Institute of MIT and Harvard, Stanley Center for Psychiatric Research, Boston, MA, USA), Andrew Ratanatharathorn (Department of Epidemiology, Gillings School of Global Public Health, University of North Carolina, Chapel Hill, Chapel Hill, NC, USA), Lynn M Almli (Department of Psychiatry and Behavioral Sciences, Emory University, Atlanta, GA, USA), Ananda B Amstadter (Department of Psychology, Virginia Commonwealth University, Richmond, VA, USA), Allison E Ashley-Koch (Department of Medicine, Duke Molecular Physiology Institute, Duke University Medical Center, Durham, NC, USA), Dewleen G Baker (Veterans Affairs San Diego Healthcare System and Veterans Affairs Center of Excellence for Stress and Mental Health, San Diego, CA, USA, Department of Psychiatry, University of California, San Diego, San Diego, CA, USA), Jean C Beckham (Veterans Affairs Durham Healthcare System, Durham, NC, USA, Department of Psychiatry and Behavioral Sciences, Duke University Medical Center, Durham, NC, USA), Laura J Bierut (Department of Psychiatry, Washington University School of Medicine, St Louis, MO, USA), Jonathan Bisson (Division of Psychological Medicine and Clinical Neurosciences, Cardiff University, Cardiff, UK), Bekh Bradley (Atlanta VA Medical Center, Atlanta, GA, USA, Department of
Funding
This study was supported by National Institutes of Health grants R01 MH106595 and U01 MH109532, and the VA National Center for PTSD Research.

Availability of data and materials
All data generated during this study are included in this published article and its Additional files.

Authors’ contributions
RP, MBS, and JG were involved in study design. ABA, LMA, DGB, LJB, BB, LAF, EOI, AK, HRR, AXM, JPR, ALR, NLS, HZ, IL, KJR, CMN, KCK, and JG were involved in cohort ascertainment, phenotypic characterization, genotyping, and/or recruitment. RP carried out statistical analysis. RP wrote the first draft of the manuscript and all authors contributed to the final version of the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
This study is based on the PGC-PTSD data freeze 1, which is an anonymized dataset. The cohorts contributing to the PGC-PTSD workforce included in the present analysis are: Collaborative Genetic Study of Nicotine Dependence; Family Study of Cocaine Dependence; Yale-Penn, Grady Trauma Project; Marine Resilience Study; Nurses’ Health Study; and Ohio National Guard. Each of the contributing studies has an institutional review board-approved protocol to investigate the genetics of PTSD and provided the data consistent with their approved protocols. All studies involved in the PGC-PTSD workforce comply with the Declaration of Helsinki. Additionally, we used GWAS summary statistics from the GIANT, Reprogen, and SSGAC consortia, which shared this information in accordance with the protocols approved for the cohorts participating in these studies.

Consent for publication
Not applicable.

Competing interests
Dr. Stein has in the last three years been a consultant for Actelion Pharmaceuticals, Healthcare Management Technologies, Janssen, Pfizer, Resilience Therapeutics, Tonix Pharmaceuticals, and Oxeia Pharmaceuticals. Dr. Kranzler has been an advisory board member, consultant, or CME speaker for Indivior and Lundbeck. He is also a member of the American Society of Clinical Psychopharmacology’s Alcohol Clinical Trials Initiative, which was supported in the last three years by AbbVie, Alkermes, Ethypharm, Indivior, Lilly, Lundbeck, Pfizer, Arbor, and Armarygala Neurosciences. Dr. Bierut is listed as an inventor on Issued US Patent 8,080,371, “Markers for Addiction” covering the use of certain SNPs in determining the diagnosis, prognosis, and treatment of addiction. Dr. Libenzon has been a consultant for ARMGO Pharmaceutical, Sunovion Pharmaceutical, and Trisman Pharma. The remaining authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1Department of Psychiatry, Yale University School of Medicine and VA CT Healthcare Center, 116A2, 950 Campbell Avenue, West Haven, CT 06516, USA. 2Department of Psychology, Virginia Commonwealth University, Richmond, VA, USA. 3Department of Psychiatry, University of California San Diego, La Jolla, CA, USA. 4Department of Family Medicine and Public Health, Psychiatry, Emory University, Atlanta, GA, USA; “Chia-Yen Chen” (The Analytic and Translational Genetics Unit, Massachusetts General Hospital, Boston, MA, USA, Psychiatric and Neurodevelopmental Genetics Unit, Center for Human, enetic Research, and Department of Psychiatry, Massachusetts General Hospital, Boston, MA, USA, Department of Psychiatry, Harvard University, Cambridge, MA, USA), Shareefa Dalvie (Division of Human Genetics, University of Cape Town, Cape Town, South Africa), Lindsay A Farrer (Biomedical Genetics, Boston University School of Medicine, Boston, MA, USA), Sandra Galea (Boston University School of Public Health, Boston, MA, USA), Melanie E Garrett (Department of Medicine, Duke Molecular Physiology Institute, Duke University Medical Center, Durham, NC, USA), Joel E Gerenser (Department of Psychiatry, Yale University School of Medicine, and VA CT Healthcare System, New Haven, CT, USA), Guia Guffanti (Department of Psychiatry, Harvard University, Cambridge, MA, USA, Department of Psychiatry, McLean Hospital, Belmont, MA, USA), Michael A Hauser (Department of Medicine, Duke Molecular Physiology Institute, Duke University Medical Center, Durham, NC, USA), Eric O Johnson (RTI International, Research Triangle Park, NC, USA), Ronald C Kesler (Department of Health Care Policy, Harvard Medical School, Boston, MA, USA), Nathan A Kimbrel (Veterans Affairs Durham Healthcare System, Durham, NC, USA, Department of Psychiatry and Behavioral Sciences, Duke University Medical Center, Durham, NC, USA), Anthony King (Department of Psychiatry, University of Michigan, Ann Arbor, MI, USA), Nastassja Koen (Department of Psychiatry and Mental Health, University of Cape Town, Cape Town, South Africa, MRC Unit on Anxiety & Stress Disorders, Groote Schuur Hospital, Cape Town, South Africa), Henny R Kranzler (Department of Psychiatry, University of Pennsylvania Perelman School of Medicine, Philadelphia, PA, USA), Rajendra A Morey (Department of Psychiatry and Behavioral Sciences, Duke University Medical Center, Durham, NC, USA, Durham VA Medical Center, Durham, NC, USA), Nicole R Nugent (Division of Behavioral Genetics, Department of Psychiatry, Rhode Island Hospital, Providence, RI, USA, Department of Psychiatry and Human Behavior, Alpert Medical School of Brown University, Providence, RI, USA), John P Rice (Department of Psychiatry, Washington University, St Louis, MO, USA), Stephen Ripke (Broad Institute of MIT and Harvard, Stanley Center for Psychiatric Research, Boston, MA, USA, The Analytic and Translational Genetics Unit, Massachusetts General Hospital, Boston, MA, USA, Department of Psychiatry and Psychotherapy, Charité, Campus Mitte, Berlin, Germany), Andrea L Roberts (Department of Social and Behavioral Sciences, Harvard T. H. Chan School of Public Health Cambridge, MA, USA), Nancy L Racine (Department of Genetics, Washington University, St Louis, MO, USA), Jordan W Smoller (Broad Institute of MIT and Harvard, Stanley Center for Psychiatric Research, Boston, MA, USA, Psychiatric and Neurodevelopmental Genetics Unit, Center for Human Genetic Research, and Department of Psychiatry, Massachusetts General Hospital, Boston, MA, USA), Daniel J Stein (Department of Psychiatry and Mental Health, University of Cape Town, Cape Town, South Africa, MRC Unit on Anxiety & Stress Disorders, Groote Schuur Hospital, Cape Town, South Africa), Murray B Stein (Department of Psychiatry, University of California, San Diego, La Jolla, CA, USA, Veterans Affairs San Diego Healthcare System, San Diego, CA, USA, Department of Family Medicine and Public Health, University of California, San Diego, La Jolla, CA, USA), Jennifer A Sumner (Center for Cardiovascular Behavioral Health, Columbia University Medical Center, New York, NY, USA), Monica Uddin (Department of Psychology and Carl R Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, IL, USA), Robert J Ursano (Center for the Study of Traumatic Stress, Department of Psychiatry, Uniformed Services University of the Health Sciences, Bethesda, MD, USA), Andrea E Wildman (Department of Molecular & Integrative Physiology and Carl R Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, IL, USA), Rachel Yehuda (James J Peters Bronx Veterans Affairs Hospital, Department of Psychiatry, Yeshiva University, New York, NY, USA, Department of Epidemiology, Columbia University School of Public Health, New York, NY, USA), Hongyu Zhao (Department of Biostatistics, Yale University, New Haven, CT, USA), Mark J Daly (Broad Institute of MIT and Harvard, Stanley Center for Psychiatric Research, Boston, MA, USA, The Analytic and Translational Genetics Unit, Massachusetts General Hospital, Boston, MA, USA), Israel Liberzon (Department of Psychiatry, University of Michigan, Ann Arbor, MI, USA, VA Ann Arbor Health System, Ann Arbor, MI, USA), Kerry J Ressler (Department of Psychiatry, Harvard University, Cambridge, MA, USA, Department of Psychiatry, McLean Hospital, Belmont, MA, USA), Caroline M Nievergelt (Veterans Affairs San Diego Healthcare System and Veterans Affairs Center of Excellence for Stress and Mental Health, San Diego, CA, USA, Department of Psychiatry, University of California, San Diego, San Diego, CA, USA), Karestan C Koenen (Broad Institute of MIT and Harvard, Center for Psychiatric Research, Boston, MA, USA, Psychiatric and Neurodevelopmental Genetics Unit, Center for Human Genetic Research, and Department of Psychiatry, Massachusetts General Hospital, Boston, MA, USA, Department of Epidemiology, Harvard T. H. Chan School of Public Health, Cambridge, MA, USA).
References

1. Pitman RK, Rasmusson AM, Koenen KC, Shin LM, Orr SP, Gilbertson MW, et al. Biological studies of post-traumatic stress disorder. Nat Rev Neurosci. 2012;13:769–87.

2. American Psychiatric Association. Trauma- and stressor-related disorders. In: Diagnostic and Statistical Manual of Mental Disorder. Washington, DC: American Psychiatric Association; 2013.

3. Kessler RC, Sonnega A, Bromet E, Hughes M, Nelson CB. Posttraumatic stress disorder. Psychol Bull. 2007;133:183–208.

4. Cortina LM, Kubiaik SP. Gender and posttraumatic stress: sexual violence as an explanation for women’s increased risk. J Abnorm Psychol. 2006;115:753–60.

5. Barban N, Jansen R, de Vlaming R, Vaez A, Mandemakers JJ, Tropf FC, et al. Novel genes identified in a high-density genome-wide association study for nicotine dependence. Hum Mol Genet. 2007;16:232–38.

6. Duncan LE, Ratanatharathorn A, Aiello AE, Almli LM, Amstadter AB, Ashley-Martin JA, et al. Interaction of the ADRB2 gene polymorphism with childhood trauma in predicting adult symptoms of posttraumatic stress disorder. JAMA Psychiat. 2014;71:1174–82.

7. Chang CC, Chow CC, Tellier LC, Vattikuti S, Purcell SM, Lee JJ. Second-generation PLINK rising to the challenge of larger and richer datasets. Gigascience. 2015;4:e7.

8. Euesden J, Lewis CM, O'Reilly PF. PRSice: Polygenic Risk Score software. Bioinformatics. 2015;31:1466–8.

9. International Schizophrenia Consortium, Purcell SM, Wray NR, Stone JL, Vischer PM, O’Donovan MC, et al. Common polygenic variation contributes to risk of schizophrenia and bipolar disorder. Nature. 2009;460:748–52.

10. Locke AE, Kahali B, Berndt SI, Justice AE, Pers TH, Day FR, et al. Genetic studies of body mass index yield new insights for obesity biology. Nature. 2014;514:86–92.

11. Kubiszyn LD, Bordellos P, Jun HI, Roberts AL, Cerda M, Bluestone N, et al. The weight of traumatic stress: a prospective study of posttraumatic stress disorder symptoms and weight status in women. JAMA Psychiat. 2014;71:44–51.

12. Yonkers KA, Smith MV, Foray A, Epperson CN, Costello D, Lin H, et al. Pregnant women with posttraumatic stress disorder and risk of preterm birth. JAMA Psychiat. 2014;71:987–904.

13. Boyraz G, Granda R, Baker CN, Tidwell LL, Waits JB. Posttraumatic stress, effort regulation, and academic outcomes among college students: a longitudinal study. J Couns Psychol. 2016;63:475–86.

14. Polimanti R, Gelernter J, Stein DJ. Genetically determined schizophrenia is not associated with impaired glucose homeostasis. Schizophrenia Res. 2017. http://doi.org/10.1016/j.schres.2017.10.033.
38. Supek F, Bosnjak M, Skunca N, Smuc T. REVIGO summarizes and visualizes long lists of gene ontology terms. PLoS One. 2011;6:e21800.
39. Burgess S, Scott RA, Timpson NJ, Davey Smith G, Thompson SG, EFC-InterAct Consortium. Using published data in Mendelian randomization: a blueprint for efficient identification of causal risk factors. Eur J Epidemiol. 2015;30:543–52.
40. Bowden J, Davey Smith G, Burgess S. Mendelian randomization with invalid instruments: effect estimation and bias detection through Egger regression. Int J Epidemiol. 2015;44:512–25.
41. Pickrell JK, Berisa T, Liu JZ, Segurel L, Tung JY, Hinds DA. Detection and interpretation of shared genetic influences on 42 human traits. Nat Genet. 2016;48:709–17.
42. Day FR, Helgason H, Chasman DI, Rose LM, Loh PR, Scott RA, et al. Physical and neurobehavioral determinants of reproductive onset and success. Nat Genet. 2016;48:617–23.
43. Burgess S, Davies NM, Thompson SG. Bias due to participant overlap in two-sample Mendelian randomization. Genet Epidemiol. 2016;40:597–608.
44. Polimanti R, Yang BZ, Zhao H, Gelernter J. Evidence of polygenic adaptation in the systems genetics of anthropometric traits. PLoS One. 2016;11:e0160654.
45. Fuente-Martin E, Argente-Arizon P, Ros P, Argente J, Chowen JA. Sex differences in adipose tissue: It is not only a question of quantity and distribution. Adipocyte. 2013;2:128–34.
46. Gandubert C, Scali J, Ancelin ML, Canièr I, Dupuy AM, Bagnolini G, et al. Biological and psychological predictors of posttraumatic stress disorder onset and chronicity: A one-year prospective study. Neuropsychobiology. 2016;73:61–7.
47. Jasienska G, Ziomkiewicz A, Ellison PT, Lipson SF, Thune I. Large breasts and narrow waists indicate high reproductive potential in women. Proc Biol Sci. 2004;271:1213–7.
48. Strahan EJ, Wilson AE, Cressman KE, Buote VM. Comparing to perfection: How cultural norms for appearance affect social comparisons and self-image. Body Image. 2006;3:211–27.
49. Tabet M, Flick LH, Cook CA, Xian H, Chang JJ. Age at first birth and psychiatric disorders in low-income pregnant women. J Womens Health (Larchmt). 2016;25:810–7.
50. Hauger RL, Olivares-Reyes JA, Dautzenberg FM, Lohr JB, Braun S, Oakley RH. Molecular and cell signaling targets for PTSD pathophysiology and pharmacotherapy. Neuropharmacology. 2012;62:705–14.
51. Singewald N, Schmuckermair C, Whittle N, Holmes A, Resler KJ. Pharmacology of cognitive enhancers for exposure-based therapy of fear, anxiety and trauma-related disorders. Pharmacol Ther. 2015;149:150–90.
52. Noll JG, Shenk CE, Putnam KT. Childhood sexual abuse and adolescent pregnancy: a meta-analytic update. J Pediatr Psychol. 2009;34:365–78.
53. Boyer D, Fine D. Sexual abuse as a factor in adolescent pregnancy and child maltreatment. Fam Plann Perspect. 1992;24:4–11.

Submit your next manuscript to BioMed Central and we will help you at every step:

- We accept pre-submission inquiries
- Our selector tool helps you to find the most relevant journal
- We provide round the clock customer support
- Convenient online submission
- Thorough peer review
- Inclusion in PubMed and all major indexing services
- Maximum visibility for your research

Submit your manuscript at www.biomedcentral.com/submit