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Mendelian randomisation analysis of circulating adipokines and C-reactive protein on breast cancer risk

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Circulating adipokines and C-reactive protein (CRP) have been linked to breast cancer risk in observational epidemiological studies. The causal nature of these associations is unclear because of the susceptibility of conventional observational designs to residual confounding, reverse causation and other forms of bias. Mendelian randomisation (MR) uses genetic variants as proxies for risk factors to strengthen causal inference in observational settings. We performed a MR analysis to evaluate the causal relevance of six previously reported circulating adipokines [adiponectin, hepatocyte growth factor (HGF), interleukin-6, leptin receptor, plasminogen activator inhibitor-1 and resistin] and CRP in risk of overall and oestrogen receptor-stratified breast cancer in up to 122,977 cases and 105,974 controls of European ancestry. Genetic instruments were constructed from single-nucleotide polymorphisms robustly (p < 5 × 10⁻⁸) associated with risk factors in genome-wide association studies. Colocalisation was performed as a sensitivity analysis to examine whether findings reflected shared causal variants or genomic confounding. In MR analyses, there was evidence for an association of HGF with oestrogen receptor-negative cancer (odds ratio per standard deviation increase: 1.17, 95% confidence interval: 1.01–1.35, p = 0.035) but little evidence for associations of other adipokines or CRP with overall or oestrogen receptor-stratified breast cancer. Colocalisation analysis suggested that the association of HGF with oestrogen receptor-negative breast cancer was unlikely to reflect a causal association. Collectively, these findings do not support an important aetiological role of various adipokines or CRP in overall or oestrogen receptor-specific breast cancer risk.

Introduction
Elevated body mass index is an important modifiable risk factor for breast cancer.1 With the discovery of adipose tissue as a functioning endocrine organ, much attention has focused on a putative role of adipokines – cytokines and hormones released by adipose tissue – as molecular mediators linking excess adiposity to breast cancer.2–4 Two adipokines in particular – leptin and adiponectin – have been strongly implicated in breast cancer aetiology. In vitro studies indicate that leptin may promote human breast cancer cell proliferation and adiponectin may exert antiproliferative effects5 and meta-analyses of observational studies support their opposing roles in breast cancer risk.4,6,7 Likewise, several other adipokines including hepatocyte growth factor (HGF), interleukin-6 (IL-6), plasminogen activator inhibitor-1 (PAI-1) and resistin have been linked to breast cancer risk in observational studies, albeit less consistently.8–10 Pre-diagnostic C-reactive protein (CRP), a systemic marker of inflammation that is synthesised in part by adipose tissue,11 has also been associated with breast cancer risk in prospective observational studies.12 Collectively, these observational findings suggest that pharmacological targeting of adipokines or CRP could be an effective strategy for breast cancer prevention among overweight or obese women. However, the causal nature of these risk factors in breast cancer risk, and thus their suitability as intervention targets, is unclear. This is because of the uncertain relevance of in vitro studies to humans, and the susceptibility of conventional observational analyses to residual confounding and reverse causation, all of which undermines causal inference.13,14

Additional Supporting Information may be found in the online version of this article.

Key words: adipokines, C-reactive protein, breast cancer, Mendelian randomisation

Abbreviations: BCAC: Breast Cancer Association Consortium; CRP: C-reactive protein; ER−: oestrogen receptor-negative; ER+: oestrogen receptor-positive; GWAS: genome-wide association study; HGF: hepatocyte growth factor; IL-6: interleukin-6; InSIDE: Instrument Strength Independent of Direct Effect; IV: instrumental variable; IVW: inverse-variance weighted; MB: megabase; MR: Mendelian randomisation; OR: odds ratio; PAI-1: plasminogen activator inhibitor-1; SD: standard deviation; SNP: single-nucleotide polymorphism

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Mendelian randomisation (MR) uses genetic variants as instruments (‘proxies’) for risk factors (including those that are potentially modifiable) to generate more reliable evidence on the causal effects of these factors on disease outcomes and so inform potential intervention strategies.\(^{15,16}\) The use of genetic variants as instruments minimises confounding and precludes reverse causation as germline genotype is largely independent of lifestyle and environmental factors and is fixed at conception. The statistical power and precision of MR analysis can be increased by employing a ‘two-sample MR’ framework in which summary genetic association data from independent samples representing genetic variant–exposure and genetic variant–outcome associations are synthesised in order to estimate causal effects.\(^{17}\)

More formally, MR is a form of instrumental variable (IV) analysis that can generate unbiased causal estimates of effects of risk factors on disease and health-related outcomes if the following conditions are met: (i) the instrument is robustly associated with the exposure of interest, (ii) there is no confounding of the instrument–outcome relationship and (iii) there is no pathway through which an instrument influences an outcome except through the exposure (‘exclusion restriction criterion’).

Given uncertainty surrounding the role of various previously reported adipokines and CRP in breast cancer aetiology, we performed two-sample MR analyses to evaluate the causal relevance of circulating adiponectin, HGF, IL-6, leptin receptor, PAI-1, resistin and CRP in overall and oestrogen receptor-stratified breast cancer risk.

**Materials and Methods**

**Study population**

Summary genome-wide association study (GWAS) statistics were obtained from genome-wide meta-analyses on 122,977 breast cancer cases and 105,974 controls of European ancestry.\(^{18}\) Summary statistics were also obtained from analyses on 69,501 oestrogen receptor-positive (ER+) and 21,468 oestrogen receptor-negative (ER−) breast cancer cases (105,974 controls) of European ancestry.\(^{18}\) Genotype data for a subset of the overall sample (61,282 cases and 45,494 controls) were obtained by direct genotyping using an Illumina Custom Infinium array (OncoArray) consisting of approximately 530,000 single-nucleotide polymorphisms (SNPs) or by imputation with reference to the 10,000 Genomes Project Phase 3 reference panel.\(^{19}\) All SNPs with a call rate of <95%, evidence of violation of Hardy–Weinberg equilibrium (\(p < 1.0 \times 10^{-7}\) in controls or \(p < 1.0 \times 10^{-12}\) in cases) and SNPs with concordance of <98% among 5,280 duplicate sample pairs were removed. An additional 1,128 SNPs where the cluster plot was judged to be inadequate were removed. In imputation, the following SNPs were additionally removed: those with a minor allele frequency of <1%, a call rate of <98%, and those that could not be linked to the 1,000 Genomes Project or differed significantly in frequency from this panel. In total, 469,364 of the 533,631 SNPs that were manufactured on the OncoArray were used in imputation. Results for OncoArray analyses were combined with those from the previous Illumina iSelect Collaborative Oncological Gene-Environment Study (iCOGS) genotyping project (46,785 cases; 42,892 controls) along with 11 other breast cancer GWAS (14,910 cases; 17,588 controls), using a fixed-effects meta-analysis. The OncoArray, iCOGS and individual GWAS were adjusted for principal components of ancestry and OncoArray and iCOGS analyses were further adjusted for country and study, respectively. All participating studies had the approval of their appropriate ethics review board and all participants provided informed consent.

**Instrument construction**

Genetic instruments to proxy risk factors were constructed either using a ‘monogenic’ approach (restricted to cis-acting variants, i.e. located ≤1 MB of the transcription start site of the protein-coding gene) or a ‘polygenic’ approach (combining cis- and trans-acting variants, independent of genomic position), depending on the number of genome-wide significant (\(p < 5 \times 10^{-8}\)) variants available to proxy each respective risk factor. When few SNPs are available as instruments, the restriction of an instrument to cis-variants can help to minimise horizontal pleiotropy (an instrument influencing an outcome through one or more biological pathways independent of that of the exposure), a violation of the exclusion restriction criterion, as cis-variants are more likely to have direct effects on protein levels than trans-variants (i.e. those >1 MB of the transcription start site of the protein-coding gene).\(^{16}\) Genetic instruments to proxy HGF, IL-6, leptin receptor and resistin were constructed by obtaining cis-acting SNPs robustly associated with these markers (\(p < 5 \times 10^{-8}\)) in GWAS of individuals of European ancestry that were replicated in independent samples. For risk factors with ≥3 independent (\(r^2 < 0.01\)) cis- or trans-SNPs available as proxies in GWAS of individuals of European ancestry (adiponectin, CRP, PAI-1), these SNPs were combined
Table 1. Characteristics of studies used to construct instrumental variables

| Trait       | Author                | Sample size   | Mean age  | % Female |
|-------------|-----------------------|---------------|-----------|----------|
| Adiponectin | Dastani et al.20      | 30,708–38,276 | NA        | NA       |
| CRP         | Ligthart et al.22     | 204,402       | NA        | NA       |
| HGF         | Sun et al.24          | 3,301         | 43.7      | 48.9     |
| IL-6        | Swerdlow et al.25     | 133,449       | 59.0      | 49.0     |
| Leptin receptor | Sun et al.24    | 3,301         | 43.7      | 48.9     |
| PAI-1       | Huang et al.21        | 30,395        | 54.3      | 50.0     |
| Resistin    | Sun et al.24          | 3,301         | 43.7      | 48.9     |

Abbreviations: CRP, C-reactive protein; HGF, hepatocyte growth factor; IL-6, interleukin-6; NA, not available in publication; PAI-1, plasminogen activator inhibitor-1.

into multi-allelic instruments to increase the variance in the risk factor explained by the instrument.20–22 As sensitivity analyses for adiponectin, CRP and PAI-1, effect estimates generated from multi-allelic instruments were compared to those obtained from instruments consisting of weakly correlated \( r^2 < 0.15 \) cis-variants to investigate horizontal pleiotropy in primary multi-allelic models.23 Across GWAS used to instrument various traits, sample sizes varied from 3,301 to 133,449 participants, mean ages of participants varied from 43.7 to 59.0 years, the proportion of females in the sample varied from 48.9 to 50.0%, and all participants were of European ancestry. Study-level information on sample size, mean age and percentage of female participants are presented in Table 1. Estimates of data set overlap across adipokine or CRP and breast cancer analyses are presented in Supporting Information Table S1. Complete summary genetic association data for all SNPs used to instrument each trait is presented in Supporting Information Table S2.

Statistical analysis

\( R^2 \) and \( F \)-statistics were calculated to assess the strength of instruments and to examine for weak instrument bias (i.e. reduced statistical power to reject the null hypothesis when an instrument explains a limited proportion of variance in an exposure), using previously reported methods.26 For instruments constructed using individual cis-variants (HGF, leptin receptor, resistin and sensitivity analyses for PAI-1), effect estimates were generated using the Wald ratio and standard errors were approximated using the delta method. For instruments constructed using \( \geq 3 \) independent variants (adiponectin, CRP, PAI-1), effect estimates were generated using inverse-variance weighted (IVW) random-effects models to account for over-dispersion in models.27 For instruments constructed using multiple weakly correlated cis-variants (IL-6 and sensitivity analyses for adiponectin, CRP), effect estimates were generated using IVW random-effects models with adjustment for correlations between variants.28 If underdispersion was present in random-effects models, the residual standard error was set to 1. For instruments constructed using multiple independent variants (adiponectin, CRP, PAI-1), MR-Egger (regression and intercept parameter) and weighted median estimation were used to evaluate for violations of the exclusion restriction criterion.27,29 MR-Egger regression can provide unbiased estimates of causal effects even when all IVs in an instrument are invalid provided that the instrument strength independent of direct effect (InSIDE) assumption is met (i.e. that there is no association between the strength of IV–exposure associations and the magnitude of horizontal pleiotropy). The MR-Egger intercept can provide a formal statistical test for directional pleiotropy (i.e. where the net horizontal pleiotropic effect across an instrument does not average to zero). The weighted median estimate can provide unbiased estimates of causal effects when at least 50% of the information in an instrument derives from valid IVs. This approach has two advantages over MR-Egger in that it provides improved precision as compared to the latter and does not rely on the InSIDE assumption. For adiponectin, CRP and PAI-1 analyses, we also visually examined for evidence of potential outliers which may be indicative of horizontal pleiotropy by generating scatter plots, forest plots and funnel plots. Leave-one-out permutation analysis was also performed for these three traits to examine whether any results were driven by an influential SNP within instruments.

As an additional sensitivity analysis, colocalisation was performed to examine whether two traits showing evidence of an association in MR analyses share the same causal variant at a given locus. This is important to perform for analyses in which an instrument is restricted to either a single variant or a single gene region as findings may be more susceptible to bias through genetic confounding (i.e., exposure and outcomes are influenced by distinct causal variants that are in linkage disequilibrium with each other). The coloc R package uses approximate Bayes factor computation to generates posterior probabilities that associations between two traits represent each of the following configurations: (i) neither trait has a genetic association in the region \( H_0 \), (ii) only the first trait has a genetic association in the region \( H_1 \), (iii) only the second trait has a genetic association in the region \( H_2 \), (iv) both traits are associated but have different causal variants \( H_3 \) and (v) both traits are associated and share a single causal variant \( H_4 \).30 An assumption of coloc is that there is at most one causal variant per trait within the genomic region examined. Colocalisation analysis was performed for MR analyses showing evidence of association \( (p < 0.05) \) by generating...
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Table 2. Number of SNPs included in instrument, estimate of the proportion of variance in risk factor explained by the instrument ($R^2$) and $F$-statistic for each instrument, across all adipokines and CRP

| Risk factor       | Number of SNPs in instrument | $R^2$ | $F$-statistic |
|-------------------|------------------------------|-------|---------------|
| Adiponectin       | 8                            | 0.016 | 60.8          |
| CRP               | 45                           | 0.035 | 119.4         |
| HGF               | 1                            | 0.012 | 40.1          |
| IL-6              | 3                            | 0.002 | 51.4          |
| Leptin receptor   | 1                            | 0.54  | 3,872.7       |
| PAI-1             | 3                            | 0.0064| 65.5          |
| Resistin          | 1                            | 0.030 | 103.3         |

Abbreviations: CRP, C-reactive protein; HGF, hepatocyte growth factor; IL-6, interleukin-6; PAI-1, plasminogen activator inhibitor-1; SNPs, single-nucleotide polymorphisms.

$R^2$ indicates proportion of variance in risk factor explained by genetic instrument.

Results

For each risk factor, the number of SNPs included in the instrument and estimates of instrument strength ($R^2$ and $F$-statistics) are presented in Table 2. Across risk factors assessed, $F$-statistics ranged from 40.1 to 3,872.7, suggesting that analyses were unlikely to suffer from weak instrument bias.\(^3\) Conservative estimates of sample overlap across GWAS data sets used ranged from 0.0 to 7.3% (Supporting Information Table S1). Given minimal sample overlap and likely absence of weak instrument bias, the presence of overlap across some GWAS data sets is unlikely to have introduced bias into these analyses.

In MR analyses, there was little evidence to suggest associations of any of the adipokines or CRP with overall breast cancer risk (Table 3). In ER status-stratified analyses, there was evidence for an association of HGF with ER− breast cancer risk [odds ratio (OR) per standard deviation (SD) increase: 1.17, 95% confidence interval (CI): 1.01–1.35; $p = 0.035$]. This finding did not reach statistical significance using a Bonferroni-corrected threshold ($p < 0.007$). In sensitivity analyses, there was little evidence that this association colocalised (posterior probability $H_4 = 0.055$). Complete colocalisation analysis results are presented in Supporting Information Table S3. Regional Manhattan plots examining the association of all SNPs ±100 kb from the SNP used to instrument HGF for their association with this adipokine (Fig. 1) and with ER− breast cancer (Fig. 2) did not appear to support the presence of one or more independent causal variants for SNP-ER− breast cancer analyses.

In all other ER-stratified analyses, there was little evidence of association of other adipokines or CRP with breast cancer outcomes.

Findings for adiponectin, PAI-1 and CRP in sensitivity analyses using cis-SNP instruments for both overall and oestrogen receptor-stratified breast cancer were consistent with those from the primary analysis (Supporting Information Table S4). Findings were also consistent for these three risk factors when MR-Egger and weighted median estimation was employed to test for evidence of exclusion restriction.
Discussion

Our MR analyses in up to 122,977 breast cancer cases and 105,974 controls do not support an important aetiological role of six adipokines or CRP in breast cancer risk. In analyses stratified on oestrogen receptor status, there was some evidence for an association of HGF with risk of oestrogen receptor-negative breast cancer. However, this finding did not achieve statistical significance based on a Bonferroni-corrected threshold to account for multiple comparisons. Additionally, sensitivity analysis testing the probability that both traits share the same causal variant found little evidence for colocalisation.

Our findings are not consistent with some previous conventional observational analyses, notably those examining the association of adiponectin and leptin with breast cancer.6–9,12 A meta-analysis of 15 cohort and case–control studies reported a 34% risk reduction (95% CI: 13–50%) when comparing ‘highest’ to ‘lowest’ adiponectin levels, with moderate heterogeneity across studies ($I^2 = 53\%$).6 Likewise, a meta-analysis of 35 case–control studies reported a standardised mean difference in serum leptin levels of 0.46 ng/ml (95% CI: 0.31–0.60, $I^2 = 93.5\%$) when comparing breast cancer cases to controls.7 However, four prospective studies examining the relationship between prediagnostic leptin and breast cancer risk have shown conflicting results: two have supported positive associations (upper vs. lower quartile: OR 1.94, 95% CI: 1.37–2.75; upper vs lower tertile: OR 1.98, 95% CI: 1.20–3.29),32,33 whereas two found little evidence of association (upper vs lower quartile: OR 1.39, 95% CI: 0.93–2.09; upper vs. lower tertile: OR 0.83, 95% CI: 0.51–1.37).3,34 Divergent findings across studies could be attributable to residual confounding (due to unmeasured or imprecisely measured confounders) or reverse causation in some conventional observational analyses. For example, the ability to disentangle highly correlated measured adipokine levels from each other and from other metabolic perturbations associated with the obese phenotype (e.g. insulin resistance, chronic inflammation) may be limited when using conventional multivariable regression methods.35 Additionally, it is possible that previously reported associations of inflammatory markers like CRP with breast cancer could represent the effect of early stage or latent breast cancer on subsequent inflammation levels.36

Strengths of this analysis include the use of a MR approach to appraise the relationship of adipokines and CRP with breast cancer risk which should be less prone to confounding than conventional observational analyses and cannot be influenced by reverse causation. The use of a two-sample summary data MR framework afforded these analyses increased statistical power and precision by exploiting summary genetic data from several large genome-wide association studies. The incorporation of colocalisation analysis permitted evaluation of whether associations present were driven through genetic confounding or shared common causal variants. Finally,
through restricting summary genetic association data to that obtained in individuals of European descent and adjusted for principal components of ancestry, these analyses reduced the possibility of confounding through population substructures.

There are several limitations to these analyses. First, since analyses were performed using summary genetic data in aggregate, this precluded stratification according to menopausal status. Around 85% of samples included within the Breast Cancer Association Consortium (BCAC) data used were classified as postmenopausal at diagnosis, suggesting that findings were unlikely to be biased markedly by the presence of premenopausal or perimenopausal participants. Second, though attempts were made to circumvent potential violations of MR assumptions in our analyses through the use of cis-acting variants as primary instruments and in sensitivity analyses, we cannot rule out the possibility that false negative findings may have arisen through horizontally pleiotropic pathways biasing our findings toward the null. An assumption of the colocalisation package employed is that there is at most one causal variant per trait within the region examined. If both traits have a genetic association in the region examined, a low posterior probability generated for $H_4$ (probability that both traits share a single causal variant) could reflect the presence of multiple shared causal variants within that region. However, posterior probabilities generated in this analysis provided evidence that only HGF was likely to have an association in the HGF region examined, suggesting that this assumption was unlikely to be violated. Fourth, it is not known whether circulating levels of adipokines represent a good surrogate for tissue-specific levels in breast or adipose tissue. Fifth, we were unable to examine potential nonlinear associations of adipokines or CRP with breast cancer risk.

Beyond adipokines and inflammatory pathways, there are several other molecular mechanisms through which excess adiposity may influence subsequent risk of breast cancer. For example, recent MR evidence in the BCAC ($N = 98,842$ cases; $83,464$ controls) has suggested that elevated fasting insulin, a consequence of higher adiposity, increases breast cancer risk (OR per SD increase = 1.71, 95% CI: 1.26–2.31). Along with the recognised role of oestrogen levels in development of ER+ breast cancer, the hyperactivation of insulin-like growth factor pathways and oxidative stress have also been implicated as molecular mediators in breast carcinogenesis, both of which may potentially influence disease through dysregulation of mammalian target of rapamycin (mTOR) pathway signalling. Further examination of the putative role of these molecular pathways in carcinogenesis could help to identify potential pharmacological targets for the chemoprevention of breast cancer in high-risk groups and, ultimately, help to reduce the burden of breast cancer.

In conclusion, our MR analyses suggest that several adipokines and CRP are unlikely to causally influence breast cancer risk. Further exploration of potential molecular mediators linking adiposity to elevated breast cancer risk remains warranted.

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Conflict of Interest

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