Identification of seven novel loci associated with amino acid levels using single
variant and gene-based tests in 8,545 Finnish men from the METSIM study


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ABSTRACT

Comprehensive metabolite profiling captures many highly heritable traits, including amino acid levels, which are potentially sensitive biomarkers for disease pathogenesis. To better understand the contribution of genetic variation to amino acid levels, we performed single variant and gene-based tests of association between nine serum amino acids (alanine, glutamine, glycine, histidine, isoleucine, leucine, phenylalanine, tyrosine, and valine) and 16.6 million genotyped and imputed variants in 8,545 non-diabetic Finnish men from the METabolic Syndrome In Men (METSIM) study. We identified five novel loci associated with amino acid levels ($P<5\times10^{-8}$): LOC157273/PPP1R3B with glycine (rs9987289, $P=2.3\times10^{-26}$); ZFHX3 (chr16:73326579, minor allele frequency (MAF)=0.42%, $P=3.6\times10^{-9}$), LIPC (rs10468017, $P=1.5\times10^{-8}$), and WWOX (rs9937914, $P=3.8\times10^{-8}$) with alanine; and TRIB1 with tyrosine (rs28601761, $P=8.8\times10^{-9}$). Gene-based tests identified two novel genes harboring missense variants of MAF<1% that show aggregate association with amino acid levels: PYCR1 with glycine ($P_{\text{gene}}=1.5\times10^{-6}$) and BCAT2 with valine ($P_{\text{gene}}=7.4\times10^{-7}$); neither gene was implicated by single variant association tests. These findings are among the first applications of gene-based tests to identify new loci for amino acid levels. In addition to the seven novel gene associations, we identified five independent signals at established amino acid loci, including two rare variant signals at GLDC (rs138640017, MAF=0.95%, $P_{\text{conditional}}=5.8\times10^{-40}$) with glycine levels and HAL (rs141635447, MAF=0.46%, $P_{\text{conditional}}=9.4\times10^{-11}$) with histidine levels. Examination of all single variant association results in our data revealed a strong inverse relationship between effect size and MAF ($P_{\text{trend}}<0.001$). These novel signals provide further insight
into the molecular mechanisms of amino acid metabolism and potentially, their perturbations in disease.

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INTRODUCTION

Amino acid levels are highly heritable biomarkers of human disease (1) that have been implicated in a range of clinical syndromes including type 2 diabetes/insulin resistance (2–4), liver disease (5), and Alzheimer’s disease (6). Previous studies have together identified >200 common variant signals associated with amino acid levels (7–20). However, the contribution of genetic variation to amino acid level trait variance, and the role of rare genetic variation in particular, is not fully understood.

One method of assessing rare variant associations is through aggregation of multiple rare variants into a single test (21). One such approach groups rare, protein-altering variants into one test for association for each gene (21). This method has been used successfully to identify gene-based associations with HAL for histidine levels and with PAH for phenylalanine levels (17). Notably, this result occurred in the absence of any single variant reaching genome-wide significance in either HAL or PAH, highlighting the importance of gene-based tests in identifying novel genetic loci for complex traits.

In this study, we performed genome-wide single variant and gene-based association analysis in 8,545 non-diabetic Finnish men from the METabolic Syndrome In Men (METSIM) study to identify genetic associations with serum amino acid levels. We identified seven novel amino acid loci – five from single variant tests (of which two signals replicated in the Northern Finnish Birth Cohort 1966 (NFBC1966) dataset) and two from gene-based associations. We also performed analyses conditioned on all previously known amino acid genome-wide association studies (GWAS) signals and identified five
additional novel and independent signals in known amino acid loci, of which three replicated in the NFBC1966 data. In total, we identified five novel and replicated loci-amino acid associations, and two novel gene-based associations. These results help clarify the role of the specific variants and genes in amino acid homeostasis.
RESULTS

GWAS for nine amino acids levels

To identify genetic variants associated with the nine amino acid traits measured in the METSIM study (alanine, glutamine, glycine, histidine, isoleucine, leucine, phenylalanine, tyrosine, and valine; see Supplemental Figure 1-3 and Supplemental Table 1), we analyzed 16.6M genotyped and imputed variants in 8,545 non-diabetic Finnish men of mean age 57 years and mean BMI 27 kg/m² (see Supplemental Table 1).

We identified 2,428 unique variants associated with at least one amino acid trait \((P < 5 \times 10^{-8})\), and a total of 2,580 variant-trait associations (see Supplemental Table 2, Supplemental Figures 4 and 5). Of the 2,580 variant-trait associations, the majority were with glycine (1,403 variants), followed by tyrosine (560), glutamine (164), alanine (95), leucine (89), isoleucine (87), phenylalanine (67), valine (62), and histidine (53). We present a summary of the variants and their distributions in independent loci in Supplemental Table 3.

We estimated for each amino acid trait the phenotypic variation that genetic variants explained from 10.4% (histidine) to 28.5% (glycine) of variation (see Supplemental Table 4). Restricting analysis to genome-wide significant variant-trait associations \((P < 5 \times 10^{-8}\), see Supplemental Table 2), the proportion of phenotypic variation explained by significantly associated variants ranged from 1.3% (leucine) to 18.3% (glycine) (see Supplemental Table 4).
We attempted to validate genotypes at three rare imputed trait-associated variants with MAF<0.5% (see Supplemental Table 5). We confirmed two variants with no discordance between imputed and sequenced genotypes: rs141635447 (0/74 discordant) and chr16:73326579 (0/67). Variant chr3:125173967 showed a discordance rate of 39% (24/61), and was thus removed from subsequent analyses.

**Single variant analysis identifies novel associations at LOC157273/PPP1R3B, WWOX, LIPC, TRIB1, and ZFHX3**

Genome-wide single variant analyses identified five novel amino acid-associated loci at least 1 Mb away from the nearest known GWAS variant (see Table 1 and Supplemental Figure 6a-e). Of the five novel loci (see Table 1), two were located in the introns of LOC157273 (near PPP1R3B) and WWOX. At the LOC157273/PPP1R3B locus, intronic variant rs9987289-A was associated with decreased glycine levels (MAF=17.0%, $\beta$=-0.22, $P=2.3\times10^{-26}$, Supplemental Figure 6a). This variant was replicated in the NFBC1966 cohort ($\beta$=-0.15, $P=7.3\times10^{-4}$, see Table 1), and was associated with the risk of type 2 diabetes (Odds Ratio(OR)=1.05, $P=0.02$) and liver disease (OR=1.33, $P=4.7\times10^{-18}$, see Supplemental Table 6). Within the WWOX region, intronic variant rs9937914-G was associated with increased alanine levels (MAF=1.47%, $\beta$=0.36, $P=3.8\times10^{-6}$, Supplemental Figure 6b).

We identified two additional novel loci in regions previously highlighted by genome-wide association studies: first, in the region upstream of LIPC, a gene implicated in numerous lipid traits including high-density lipoprotein (HDL) cholesterol (22), phospholipids (20),
and the ratio of isoleucine and serum total cholesterol (serum-c) (10), rs10468017-T was associated with increased alanine levels (MAF=33.2%, $\beta=0.09$, $P=1.5\times10^{-8}$, Supplemental Figure 6c), and is in strong linkage disequilibrium (LD) with rs1532085 (LD $r^2=0.66$), a LIPC GWAS locus for HDL (23) and ratio of isoleucine and serum-c (10). Its association with increased alanine levels was confirmed in the NFBC1966 cohort ($\beta=0.08$, $P=7.7\times10^{-3}$, see Table 1). Second, rs28601761-G, for which we report an association with decreased tyrosine levels (MAF=42.2%, $\beta=-0.09$, $P=8.8\times10^{-9}$, Supplemental Figure 6d), is in strong LD with rs2954029 (LD $r^2=0.71$), a TRIB1 GWAS variant for low-density lipoprotein (LDL) cholesterol, triglycerides, and high-density lipoprotein (HDL) cholesterol levels (23, 24).

At the remaining novel locus, rare variant 16:73326579 was associated with increased alanine levels (MAF=0.42%, $\beta=0.76$, $P=3.6\times10^{-8}$). 16:73326579 is located within 300 kb of both HCCAT5 and ZFHX3, but is not in strong LD (LD $r^2<0.60$) with any coding variant observed in the GoT2D study (see Supplemental Figure 6e). We computed $P$ values ($P_{\text{ACT}}$) for the novel variants after correcting for the nine correlated amino acid traits (25). All of these five novel variants remained genome-wide significantly associated even after correcting for the nine amino acid traits ($P_{\text{ACT}} < 5.0 \times 10^{-8}$).

Conditional analyses identify independent signals at five known amino acid loci: GLDC, HAL, ALDH1L1, ADAMTS3, and GCSH

We curated 1,519 unique known amino acid associated variants, and then used them as covariates in the genome-wide conditional analyses (see Supplemental Table 7,
After conditional analyses, we observed 227 unique variant-trait associations ($P_{\text{conditional}}<5\times10^{-8}$) (see Supplemental Table 8), whose distributions in genes are presented in Supplemental Table 3. Among these, we identified five novel signals at established amino acid loci distinct from the previously published GWAS variants (see Table 1): GLDC p.Q996H, associated with increased glycine (rs138640017-G, MAF=0.95%, $\beta=1.35$, $P_{\text{conditional}}=5.8\times10^{-40}$); HAL p.G283V, associated with increased histidine levels (rs141635447-A, MAF=0.46%, $\beta=0.85$, $P_{\text{conditional}}=9.4\times10^{-11}$); rs6564825-G, in an intron of PKD1L2 and 38 kb downstream of GCSH, was associated with increased glycine levels (MAF=11.7%, $\beta=0.17$, $P_{\text{conditional}}=2.0\times10^{-10}$) and nominally but not coincidently associated with expression level of PKD1L2 (see Supplemental Table 9); rs190671241-G, an intergenic variant near ADAMTS3, associated with increased phenylalanine levels (MAF=1.70%, $\beta=0.36$, $P_{\text{conditional}}=2.2\times10^{-9}$); and rs112981908-G, an intronic variant of the ALDH1L1 gene associated with decreased glycine levels (MAF=11.9%, $\beta=-0.14$, $P_{\text{conditional}}=3.1\times10^{-10}$). At each locus, we observed low pairwise LD ($LD^2<0.10$) between the novel variant identified in the METSIM data and the previously published GWAS variant(s). Notably, of the five novel signals at established amino acid loci, three replicated in the NFBC1966 data: GLDC p.Q996H, associated with increased glycine (rs138640017-G, $\beta=0.94$, $P=2.7\times10^{-10}$); HAL p.G283V, associated with increased histidine levels (rs141635447-A, $\beta=1.04$, $P=1.7\times10^{-5}$); and ADAMTS3 upstream variant rs190671241-G, associated with increased phenylalanine levels ($\beta=0.39$, $P=1.6\times10^{-4}$). Functional work is necessary to determine whether the novel signals represent additional
functional variants in genes known to play a role in amino acid metabolism (e.g. \textit{GLDC} and \textit{HAL}), or whether they point to novel mechanisms.

\textbf{Single variant associations exhibit an inverse relationship between allele frequency and effect size}

To visualize the relationship between allele frequency and effect size of amino acid-associated variants, we plotted the absolute value of effect size estimates vs. MAF for all loci associated with amino acid traits in the METSIM study ($P<5 \times 10^{-8}$) (see Table 1) and fitted a fractional polynomial spline to the data (see Figure 1). These results demonstrated a strong, inverse relationship between MAF and effect size ($P_{\text{trend}}<0.001$), consistent with past findings for other traits (e.g. (26)). This relationship is largely driven by variants with MAF<5\%, five of which were newly identified in this study.

\textbf{Variant associations with amino acid ratios}

Prior studies found more genome-wide significant variant associations with amino acid ratios as compared to amino acid traits alone (8, 12). We identified 15,220 significant variant-ratio associations (3,822 unique variants) from unconditional analyses of the 36 possible ratios among the nine amino acids measured in the METSIM study. These results are presented in \textbf{Supplemental Table 10} as a reference for other investigators.

\textbf{Gene-based tests identify novel gene associations with \textit{BCAT2} and \textit{PYCR1}}

To determine the joint contribution of the protein-truncating and missense variants of MAF<1\% on amino acid traits, we performed gene-based tests (see \textbf{Supplemental Table}
and Supplemental Figure 7) and applied a significance threshold based on the number of genes tested (~20,000). Given the high correlation among amino acid traits (see Supplemental Figure 2), we did not correct for the number of amino acid traits as a Bonferroni-corrected significance threshold would be overly strict. We identified six gene-trait associations ($P_{\text{gene}} < 2.5 \times 10^{-6}$), including four genes previously identified through single variant association tests: $ALDH1L1$ (8) and $GLDC$ (16) associated with glycine levels, $HAL$ with histidine levels (16) (see Supplemental Table 12), and $DHODH$ (previously associated with alanine-to-tyrosine ratio) with alanine levels, as well as two novel associations between $PYCR1$ and glycine levels ($P_{\text{gene}}=1.5 \times 10^{-6}$), and $BCAT2$ and valine levels ($P_{\text{gene}}=7.4 \times 10^{-7}$, see Table 2).

No missense variants within either $PYCR1$ or $BCAT2$ achieved genome-wide significance with any amino acid trait in the single variant association tests, highlighting the utility of gene-based test in novel gene discovery (see Table 2). Despite only suggestive association evidence, the effect of these variants on amino acid trait variance was considerable (the range of absolute $\beta$: 0.36-1.02): the carriers of the two missense variants within the gene $PCYR1$ exhibited lower mean glycine levels, while the carriers of the three variants within the $BCAT2$ gene showed higher mean valine levels, suggesting altering their protein sequences would affect the serum glycine and valine levels, respectively (see Figure 2).
Amino acids are highly heritable traits whose levels have been implicated in the pathogenesis of human complex diseases such as type 2 diabetes (2–4) and Alzheimer's disease (6). Here, we leveraged dense, experimentally-determined and imputed genotypes and report seven novel amino acid associations in the METSIM study that replicated in NFBC1966. Of these associations, two were identified from single variant testing in the METSIM study and replicated in the NFBC1966 data, and two other loci were identified from gene-based analyses in the METSIM study alone. One of these newly identified variants from single-variant analyses, LOC157273/PPP1R3B variant rs9987289-A, also confers increased risk of type 2 diabetes and liver disease. In addition, we fine-mapped known amino acid loci and identified and replicated distinct association signals at three of these loci. Phenotypic variance explained for these nine amino acids by known and novel associations ranged from 10.4% for histidine levels to 28.5% for glycine levels in our data. These results further elucidate the potential mechanisms through which amino acid levels are perturbed, and their potential relationship to disease.

Novel loci highlight a potential role for genes implicated in lipid metabolism and human diseases

Of the five novel amino acid loci highlighted in this study through single variant analyses, four have previously been implicated in lipid metabolism. First, the rs9987289-A signal near LOC157273/PPP1R3B, for which we report an association with decreased glycine levels, has previously been associated with decreased high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), and total cholesterol...
levels (23), as well as increased $PPP1R3B$ expression levels in human liver tissue (23). In addition, overexpression of $Ppp1r3b$ led to a significant decrease in HDL-C and total cholesterol in a mouse model (23). This variant is also associated with increased risk of type 2 diabetes (27) and liver disease (28). Second, the rs10468017-T signal upstream of $LIPC$ associated with increased alanine levels has previously been associated with increased HDL-C (22), altered levels of several circulating phospholipids (20), and ratio of isoleucine and serum-c (10). Third, the signal near $TRIB1$ associated with decreased tyrosine levels has previously been associated with decreased total cholesterol, LDL-C, and triglyceride levels (23). Finally, we reported an intronic $WWOX$ variant, rs9937914-G, associated with increased alanine levels, and human carriers of predicted loss-of-function variants in $WWOX$ were reported to have lower HDL-C (31) levels; in addition, mice lacking $Wwox$ exhibit decreased fasting cholesterol, triglyceride and glucose levels (32). These variant associations may represent a secondary effect of altered lipid levels on amino acid metabolism, as previously demonstrated in the setting of obesity (33) and insulin resistance (2). Further work will be required to determine the mechanisms through which these lipid-related loci affect amino acid levels and human complex diseases.

**Gene-based tests highlight two novel loci not identified from single variant testing**

$PYCR1$, which we identified as a gene implicated in glycine levels, encodes a mitochondrial protein involved in biosynthesis of proline and generation of oxidative potential through NADP+ production (34). $PYCR1$ was recently identified as the genetic cause of autosomal recessive cutis laxa type 2, highlighting the importance of normal $PYCR1$ function in neurodevelopment (35). Functional studies of fibroblasts from affected
individuals found increased sensitivity to oxidative stress (36). As redox reactions are critical to amino acid biosynthesis (37), our finding that PYCR1 missense variants result in decreased glycine levels may suggest reduced oxidative potential in vivo.

We also identified an association between BCAT2 variants and valine levels through gene-based testing. BCAT2 encodes a mitochondrial enzyme responsible for the first steps in the breakdown of branched chain amino acids (isoleucine, leucine, and valine) (37); thus the relationship of BCAT2 with valine levels is clear. Prior literature reports that the deletion of exon 2 in the mouse homologue of BCAT2 resulted in a phenotype similar to Maple Syrup Urine Disease (38), an autosomal recessive human inborn error of metabolism characterized by high levels of branched chain amino acids and resulting neurologic symptoms due to the inability to catabolize dietary branched chain amino acids. A human case study of an adult male with mild neurologic symptoms (headaches and mild memory loss) has been reported similar findings, with R170Q and E264K missense variants in BCAT2 resulting in higher-than-expected levels of leucine, isoleucine, and valine (39). Therefore, our finding of BCAT2 missense variants resulting in increased valine levels is supported by prior literature.

Limitations

Some limitations of this study should be considered. First, our analyses were limited by statistical power secondary to sample size. This was likely one of the contributing factors to our lack of replication of single variant signals in the NFBC1966 data, as the available NFBC1966 replication sample size was modest. Future meta-analyses of amino acid
associations are likely to clarify true signals from false positives. Second, our discovery and replication populations of Finnish men (and women for the replication study) limit the generalizability of our findings. However, this study design also provided the genetic homogeneity needed to identify Finnish-ancestry-specific rare variant associations with amino acid traits. Third, our results infer association between genetic markers and amino acid traits; however, elucidating the causal mechanism through which these variants affect amino acid levels will require further functional work.

Summary

These GWAS of nine amino acid traits in 8,545 participants of the METSIM study identified five novel single variant associations, including variant-trait associations near LOC157273/PPP1R3B, WWOX, TRIB1, LIPC, and ZFHX3, of which two were replicated in the NFBC1966 cohort and one (LOC157273/PPP1R3B) was also associated with the risk of type 2 diabetes and liver disease. In addition, we identified two novel gene-based signals driven by two and three potentially functional missense variants at PYCR1 and BCAT2, respectively. In BCAT2, we validated the association of one rare missense variant in the NFBC1966 study. Our use of a dense reference panel yielded 16.6M genotyped and imputed variants, allowing for high-resolution analyses and fine-mapping of independent genetic signals at GLDC, GCSH, ALDH1L1, ADAMTS3, and HAL; of which the signals at GLDC, HAL, and ADAMTS3 were replicated in the NFBC1966 data. Further work is needed to determine which of the variants identified in this study may affect gene function and the precise roles of the identified genes in amino acid metabolism. These analyses provide further insight into the molecular mechanisms of amino acid...
metabolism, and, given the importance of amino acid level perturbation in the
pathogenesis of numerous human diseases, may yield insights into a wide spectrum of
human complex disease.
MATERIALS AND METHODS

Study Participants

Of the 10,197 participants in the METSIM study, we analyzed the subset of 8,545 non-diabetic men of mean age 57.3±7.1 years and BMI 27.0±4.0 kg/m² with NMR amino acid trait measurements (see Supplemental Table 1). Institutional review boards at the University of Kuopio and Kuopio University approved the METSIM study. Written informed consent was obtained from each participant.

Amino Acid Trait Measurement

Blood samples from METSIM participants were obtained and stored in liquid nitrogen until measurement by NMR, as previously described (40). In brief, fasting serum samples collected at enrollment were stored at -80°C and thawed overnight in a refrigerator before sample preparation. A high-throughput serum NMR metabolomics platform was then used to quantify the levels of individual metabolites using a low-molecular weight metabolite data window (1H NMR spectra) used to identify amino acids (41). We then used iterative lineshape fitting with known chemical shifts to identify and quantify each specific metabolite (42).

We measured nine amino acid levels by NMR spectroscopy: alanine, glutamine, glycine, histidine, isoleucine, leucine, phenylalanine, tyrosine, and valine (see Supplemental Table 1). Visualization of the Pearson pairwise correlation matrix between the nine measured amino acid traits was generated using the corrplot package (https://cran.r-project.org/web/packages/corrplot/index.html) within R (see Supplemental Figure 2).
Genotyping and Imputation

METSIM participant samples were genotyped on the HumanOmniExpresss-12v1_C BeadChip (OmniExpress) and Infinium HumanExome-12 v1.0 BeadChip (Exome Chip) platforms. Quality controls included sample-level controls for sex and relatedness confirmation, sample duplication, and detection of sample genetic ancestry outliers using principal component analysis. Based on these quality control measures, we removed 14 samples with sex chromosome anomalies, 18 with evidence of participant duplication, 12 population outliers, and 9 samples with non-Mendelian inheritance inconsistencies. In addition, we removed one individual from each of seven monozygotic twin pairs.

We filtered variants with low mapping quality of probes to genome build GRCh37, low genotype completeness (<95% and <98% for the OmniExpress and ExomeChip, respectively), or Hardy-Weinberg equilibrium \( P<10^{-6} \).

We phased OmniExpress variants passing quality control with SHAPEIT v2 (43) and imputed them using minimac v2 (44). For imputation, we used a reference panel of 20.9M variants from the GoT2D study (including SNVs, indels, and large deletions) based on the whole genome sequence of 2,874 Europeans, including 1,004 Finnish individuals – the largest panel of Finnish genomes available (45). Following imputation, variants directly genotyped on the ExomeChip were added. In cases of common markers between imputed and genotyped variants, we used the directly genotyped call from the ExomeChip. The distribution of imputation quality and MAF for each imputed variants are presented in
Supplemental Figure 3. We carried forward 16,607,533 variants with high imputation quality (i.e. minimac RSQ ≥ 0.3) for further single variant association testing.

Single Variant Analyses

We performed single variant association tests on imputed genotype dosages for all variants with a minor allele count ≥ 3. Association tests assumed an additive genotype model and accounted for cryptic relatedness among the Finnish population using the EMMAX linear mixed model approach (46), as implemented in EPACTS (http://genome.sph.umich.edu/wiki/EPACTS). We adjusted amino acid traits for age, age$^2$, and BMI, and then inverse normalized the residuals. We applied normalization of trait levels to control for type-I error caused by skewed distributions, although this normalization may reduce power to discover associated variants. We created association plots for the novel variants using LocusZoom (http://locuszoom.sph.umich.edu/locuszoom/). In addition, we computed $P$ values ($P_{ACT}$) for the novel variants after correcting for the nine correlated amino acid traits (25). We used a conventional significance threshold of $P < 5 \times 10^{-8}$ in single variant association testing.

Replication in the Northern Finnish Birth Cohort

The associations in the novel regions were replicated in silico in the Northern part of Finland: The 1966 cohort (the “Northern Finnish Birth Cohort”, or NFBC1966) (47). NFBC1966 is a prospective follow-up study of children from the two northernmost provinces of Finland born in 1966. All individuals still living in northern Finland or the
Helsinki area (n=8,463) were contacted and invited for clinical examination. A total of 6,007 participants attended the clinical examination at the participants’ age of 31 years. Among them, 5,402 samples were genotyped on Illumina HumanCNV370DUO Analysis BeadChip (48), and were then imputed to the Haplotype Reference Consortium (HRC) reference (49) and 1000 Genomes Project Phase 3 (50) on the Michigan Imputation Server (https://imputationserver.sph.umich.edu/index.html). The association for the novel region variants and rare variants were looked up in 2,591 samples. Given our focused hypothesis, we set a threshold for significance in replication as $P \leq 0.05$.

**Associations of Novel Amino Acid SNVs with End-organ Phenotypes**

We investigated the association of the novel amino acid regions variants with the risk of type 2 diabetes, Alzheimer’s diseases, and liver disease. For type 2 diabetes, we used publically available data in large-scale Europeans ($N = 159,208$) from the DIAGRAM consortium (http://www.diagram-consortium.org) (27). For Alzheimer’s disease, we examined associations in the ADGC consortium ($N = 54,162$) (51). Finally, for liver disease, we used association summary statistics data the GOLD consortium ($N = 7,176$) (28). We used proxy single nucleotide polymorphisms (SNPs) tightly linking with the novel variant if our variant was not available.

**Analysis of Amino Acid Trait Variance**

We estimated the phenotypic variance explained by genetic variants for inverse normalized amino acid traits as previously described through GCTA v1.26 (see Supplemental Figure 1) (52). We removed 1,153 close relatives through kinship cutoff
of 0.0075 in KING 1.4 (53), and then estimated the phenotypic variance in 7,392 unrelated samples (54). To account for the effect of population structure, we used the top ten principal components as covariates. In brief, we carried out a primary analysis that consisted of a simultaneous analysis of all 16.6M variants, and a secondary analysis considering only the 2,580 variants determined to be genome-wide significant for at least one amino acid trait (see Supplemental Table 2).

Validation of Imputed Rare Variants
We used TaqMan SNP genotyping (Thermo Fisher Scientific) or Sanger sequencing to validate genotypes at three trait-associated ($P<5\times10^{-8}$) and rare imputed variants (MAF<0.5\%) (see Supplemental Table 5). We examined all individuals predicted (on the basis of imputation) to be heterozygous carriers at any of the three sites, as well as additional non-carriers.

Genome-Wide Conditional Analyses
To identify additional independent genetic signals for amino acid traits at known GWAS loci, we conducted a comprehensive genome-wide conditional association analysis. We curated a database of prior published studies of genetic associations with amino acid and related traits to identify distinct variant associations in conditional analyses. To identify published studies, we screened a GWAS catalogue (http://www.ebi.ac.uk/gwas/), used SNIPPER (https://csg.sph.umich.edu/boehnke/snipper/) to query publicly available databases for published variants and loci, and performed literature review using PubMed (https://www.ncbi.nlm.nih.gov/pubmed/), and Google Scholar.
We focused on proteinogenic amino acid and related traits (e.g., citrate) in European populations (see Supplemental Table 7).

The curated list contained 2,615 variants (of which 1,519 were unique, with several variants having multiple trait associations) spanning >100 loci from 14 studies (see Supplemental Table 7). These associated variants were then filtered for pairwise LD ($r^2>0.95$) to 408 variants (see Supplemental Table 7). For the 2,580 amino acid associated variants identified in discovery analyses (see Supplemental Table 2), we performed a secondary analysis conditioning on the LD-pruned list of 408 independent genetic variants. A variant with $P$ value $<5\times10^{-8}$ was considered to be a novel secondary signal within known amino acid traits loci after conditioning on these 408 independent genetic variants.

### Amino Acid Ratio Tests of Association

Prior investigations of genetic associations with amino acid trait variation have reported extensive findings with amino acid ratios (8, 12). While amino acid ratios were not the focus of our investigation, we included discovery analyses with 36 amino acid ratios listed in Supplemental Table 10.

### Gene-Based Tests of Association

We performed gene-based tests of association using SKAT-O (21) with EMMAX (46) to determine the joint contribution of protein-truncating (i.e. nonsense, frameshift, and essential splice variants) and missense variants with MAF<1% on amino acid traits, as
described in our previous study (55). For these analyses, we included only coding
variants directly genotyped on either the OmniExpress or Exome array. Missing
genotype data (proportion < 2%) were imputed with variant-specific mean genotype
since SKAT-O requires complete data (21). A total of 51,898 protein-truncating or
missense variants in 13,996 genes met these criteria (see Supplemental Table 11 for a
summary of variant distributions within genes). We considered a gene-based result
exome-wide significant at a p-value threshold of $2.5 \times 10^{-6}$ (0.05/20,000) to account for
the number of genes in these gene-based analyses.
Conflicts of Interest: A.J.K. and P.S. are shareholders of Brainshake Ltd., a company offering NMR-based metabolite profiling. A.J.K. and P.S. report employment relation for Brainshake Ltd. All other authors report no conflicts of interest.

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REFERENCES


FIGURE LEGENDS

Figure 1. Relationship between minor allele frequency and estimated beta coefficient (β) for loci associated with amino acid levels in the METSIM data. All novel amino acid loci (triangles in pink) are highlighted, in addition to novel signals at known amino acid loci (squares in green) identified through analyses conditioned on all known amino acid GWAS variants. The known amino acid signals are represented with blue circles. Gray dashed line represents a fractional polynomial spline fitted to the data points (P<0.001). β, on the y-axis, is the absolute value of the estimated regression coefficient for a given variant-trait association.

Figure 2. Plots show the trait values of rare variant carriers relative to the distribution of amino acid levels in all individuals. The tables in the right panel show gene-based tests of association with amino acid levels for genes PYCR1 and BCAT2. Histograms show the distribution of the inverse normalized residuals of the trait across all participants for the gene-based test of association at (a) PYCR1 with glycine levels and (b) BCAT2 with valine. The dashed gray line represents the mean inverse normalized residual of trait level for all individuals. The solid black line in each row represents the mean trait level for carriers of each variant. Triangles represent rare variant carriers. The locations of triangles relative to the distribution across all participants indicate the trait levels of rare variant carriers. No individuals were homozygous for the minor allele of any of the listed variants.
SUPPLEMENTAL DATA

Seven Supplemental Figures are available online as a single PDF file. Twelve Supplemental Tables are available in a single Excel file.
Table 1. Novel and known single variants associations with amino acid traits.

<table>
<thead>
<tr>
<th>Lead Variant</th>
<th>Trait</th>
<th>Chr:Pos ¹</th>
<th>Variant Annotation</th>
<th>METSIM</th>
<th>NFBC1966</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MAF (%)</td>
<td>β (SE)</td>
</tr>
<tr>
<td><strong>Novel Single Variant Associations with Amino Acid Traits</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rs9987289</td>
<td>Gly</td>
<td>8:9183358</td>
<td>LOC157273 intronic</td>
<td>17.0 (A)</td>
<td>-0.22 (0.02)</td>
</tr>
<tr>
<td>16:73326579</td>
<td>Ala</td>
<td>267 kb upstream of ZFHX3</td>
<td>0.42 (T)</td>
<td>0.76 (0.13)</td>
<td>0.41</td>
</tr>
<tr>
<td>rs10468017</td>
<td>Ala</td>
<td>45 kb upstream of LIPC</td>
<td>33.2 (T)</td>
<td>0.09 (0.02)</td>
<td>0.37</td>
</tr>
<tr>
<td>rs9937914</td>
<td>Ala</td>
<td>16:73422354</td>
<td>WWOX intronic</td>
<td>1.47 (G)</td>
<td>0.30 (0.02)</td>
</tr>
<tr>
<td>rs28601761</td>
<td>Tyr</td>
<td>49 kb downstream of TRIB1</td>
<td>42.2 (G)</td>
<td>-0.09 (0.02)</td>
<td>0.39</td>
</tr>
</tbody>
</table>

| **Novel Single Variant Signals at Established Amino Acid Loci** |       |           |                    |        |          |      |   |            |            |           |        |       |       |
| rs138640017  | Gly   | 9:6533092 | GLDC Q996H | 0.95 (G) | 1.35 (0.08) | 3.35 | $3.5 \times 10^{-65}$ | rs140348140 | Gly (16) | $5.8 \times 10^{-40}$ | 1.02 | 0.94 (0.15) | $2.7 \times 10^{-10}$ |
| rs141635447  | His   | 12:96374381 | HAL G283V | 0.46 (A) | 0.85 (0.12) | 0.62 | $2.5 \times 10^{-13}$ | rs7954638 | His (16) | $9.4 \times 10^{-11}$ | 0.35 | 1.04 (0.24) | $1.7 \times 10^{-5}$ |
| rs6564825    | Gly   | 16:81153894 | PKD1L2 intronic | 11.7 (G) | 0.17 (0.02) | 0.57 | $3.1 \times 10^{-12}$ | rs74249229 | Gly (16) | $2.0 \times 10^{-10}$ | 10.4 | -0.01 (0.05) | 0.87 |
| rs190671241  | Phe   | 139 kb upstream of ADAMTS3 | 1.70 (G) | 0.36 (0.06) | 0.42 | $2.4 \times 10^{-9}$ | rs7954638 | His/Phe (8) | $2.2 \times 10^{-6}$ | 2.10 | 0.39 (0.10) | $1.6 \times 10^{-4}$ |
| rs112981908  | Gly   | 3:125858480 | ALDH1L1 intronic | 11.9 (G) | -0.14 (0.02) | 0.37 | $1.8 \times 10^{-8}$ | rs11047366 | Gly (16) | $3.1 \times 10^{-10}$ | e | e | e |

Abbreviations: β (SE) – estimated regression coefficient and standard error for the minor allele; Chr/Pos – variant chromosome and position based on hg19 build; MAF% (Allele) - minor allele frequency (in percent) with minor allele in parentheses; SNV – single nucleotide variant; Var% – trait variance explained by variant (in percent)

Amino acid abbreviations: Ala – alanine; Gly – glycine; His – histidine; Phe – phenylalanine; Tyr – tyrosine.

¹ Position based on hg19 build.

b Lead GWAS SNV within 1 Mb of the identified lead SNV.

c $P_{\text{conditional}}$ Values result from conditional analyses adjusting for known amino acid signals from previous studies (see Supplemental Table 7)

d These two variants were directly genotyped for validation and had 100% concordance with the imputed genotype (see text).

e This variant was not in the HRC panel used for analyses in the NFBC1966 dataset.
### Table 2. Novel gene-based associations with amino acid traits.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Gene</th>
<th>rsID</th>
<th>Chr:Pos a</th>
<th>Annotatio n</th>
<th>METSIM</th>
<th>NFBC1966</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MAF (%) (Allele)</td>
<td>Genotype Counts</td>
</tr>
<tr>
<td>Gly</td>
<td>PYCR1</td>
<td>rs3744807</td>
<td>17:79890818</td>
<td>G297R</td>
<td>0.13(T)</td>
<td>8532 / 22 / 0</td>
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<tr>
<td></td>
<td></td>
<td>rs142225075</td>
<td>17:79893020</td>
<td>L108V</td>
<td>0.10(C)</td>
<td>8528 / 17 / 0</td>
</tr>
<tr>
<td>Val</td>
<td>BCAT2</td>
<td>rs199999090</td>
<td>19:49299714</td>
<td>R331C</td>
<td>0.18(A)</td>
<td>8515 / 30 / 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rs117048185</td>
<td>19:49309776</td>
<td>Q60E</td>
<td>0.59(C)</td>
<td>8445 / 100 / 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rs201148940</td>
<td>19:49309937</td>
<td>H6R</td>
<td>0.02(C)</td>
<td>8542 / 3 / 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rs192923495</td>
<td>16:72057193</td>
<td>R317W</td>
<td>0.09(T)</td>
<td>8529 / 16 / 0</td>
</tr>
</tbody>
</table>

**Novel Gene-Based Associations with Amino Acid Traits**

- **Abbreviations:** β – estimated regression coefficient for the minor allele; Chr:Pos – variant chromosome and position based on hg19 build; MAF(%) – minor allele frequency (in percent) with minor allele in parentheses.
- **Amino acid abbreviations:** Ala – alanine; Gly – glycine; His – histidine; Val – valine.
- a Position based on hg19 build.
- b SNPs in *DHODH* previously associated with Alanine-to-Tyrosine ratio by Kettunen et al.(8).