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Elevated circulating branched chain amino acids are an early event in pancreatic adenocarcinoma development

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AUTHOR CONTRIBUTIONS

All authors participated in the analysis and interpretation of data. CW, CBC, PK, CY, YB, MKT and BMW performed the statistical analyses of the human data. CBC, MKT, SST, TJW, REG and BMW evaluated the platform for analysis of plasma metabolites in large cohort studies. CBC, JRM, AS and KP performed metabolite profiling experiments. PK, YB, MKT, SST, SO, MJS, ELG, ZRQ, DAR, JM, HDS, JMG, BBC, SL JWW, JEM, MNP and CSF assisted in data acquisition, management, and interpretation from the four cohort studies. JRM and MET conducted all mouse experiments with assistance from BPF, SMD, TP, AY TLD and ACK. CW, CBC, JRM, CSF, MGVH and BMW designed the study and drafted the manuscript with input from all authors.

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Abstract

Most patients with pancreatic ductal adenocarcinoma (PDAC) are diagnosed with advanced disease and survive less than 12 months¹. PDAC has been linked with obesity and glucose intolerance²⁻⁴, but whether changes in circulating metabolites are associated with early cancer progression is unknown. To better understand metabolic derangements associated with early disease, we profiled metabolites in prediagnostic plasma from pancreatic cancer cases and matched controls from four prospective cohort studies. We find that elevated plasma levels of branched chain amino acids (BCAAs) are associated with a greater than 2–fold increased risk of future pancreatic cancer diagnosis. This elevated risk was independent of known predisposing factors, with the strongest association observed among subjects with samples collected 2 to 5 years prior to diagnosis when occult disease is likely present. We show that plasma BCAAs are also elevated in mice with early stage pancreatic cancers driven by mutant *Kras* expression, and that breakdown of tissue protein accounts for the increase in plasma BCAAs that accompanies early stage disease. Together, these findings suggest that increased whole–body protein breakdown is an early event in development of PDAC.

PDAC is a leading cause of cancer–related death worldwide, and most patients have incurable disease at diagnosis¹. The best–characterized predisposing factors, current tobacco use and a first–degree relative with PDAC, both impart an approximate 1.8–fold increased risk for the disease^{5,6}. These risk factors, however, have thus far provided limited insight into the biology of early disease progression of sporadic tumors. Development and progression of PDAC is associated with altered systemic metabolism including obesity², glucose intolerance^{3,4}, and cancer-induced cachexia⁷. Nevertheless, no systematic examination of circulating metabolites has been performed to determine whether altered

metabolism may indicate subclinical pancreatic cancer or inform our understanding of early disease progression when interventions might improve patient outcomes.

Prior efforts to identify changes in circulating metabolites related to cancer have employed a cross–sectional design, comparing cancer–free controls to cases with blood samples collected at diagnosis⁸⁻¹⁰. This approach is problematic for discovery of changes related to early cancer progression, as consequences of advanced disease are likely to impact circulating metabolite profiles. This is particularly true for patients with pancreatic cancer, who commonly have significant anorexia, weight loss, and pancreatic insufficiency at the time of diagnosis¹. To investigate how altered metabolism might contribute to pancreatic malignancy, we profiled plasma metabolites in cases with PDAC and matched controls drawn from four prospective cohort studies with blood collected at least two years before cancer diagnosis (Supplementary Table 1). The median time between blood collection and PDAC diagnosis was 8.7 years.

In conditional logistic regression models, 15 plasma metabolites were associated with future diagnosis of PDAC to P<0.05; 3 metabolites, the branched chain amino acids (BCAAs) isoleucine, leucine and valine were significant to P 0.0006, the predefined significance threshold after correction for multiple–hypothesis testing (Fig. 1 and Supplementary Table 2). To evaluate the magnitude of risk for PDAC diagnosis, we divided participants into quintiles of increasing BCAA levels. Compared to the bottom quintile, subjects in the top quintile had at least a 2–fold increased risk of developing PDAC (Table 1, Supplementary Table 3 and Supplementary Fig. 1). As noted previously¹¹, circulating levels of the three BCAAs were highly correlated (Supplementary Table 4), reflecting their common pathways of metabolism¹² and leading to similar results for the sum total of BCAAs (Table 1 and Supplementary Table 3).

Circulating BCAAs are elevated in obese individuals and those with insulin resistance¹³. In study participants, plasma BCAAs were modestly correlated with markers of energy balance, obesity and glucose intolerance (Supplementary Table 4). To evaluate the independent effect of BCAAs on PDAC risk, we assessed models that included these markers and found that the associations of BCAAs with PDAC remained largely unchanged (Table 1). Circulating levels of BCAAs are also associated with future risk of diabetes^{11,14}. Since type 2 diabetes is a predisposing factor for PDAC¹⁵, we questioned whether the intermediate development of diabetes was underlying the BCAA association. Exclusion of subjects with diabetes at blood collection did not change our results (Table 1), indicating that we had not identified a signature of prevalent diabetes associated with later PDAC diagnosis. To determine whether circulating BCAAs were identifying a population at risk for diabetes, who are then at elevated risk of PDAC, we excluded subjects who developed diabetes between the time of blood collection and cancer diagnosis and found the results unchanged (Table 1). These data suggest that the association of BCAAs with future PDAC diagnosis is not dependent on intermediate development of diabetes.

To examine the contribution of circulating BCAAs to risk stratification models for PDAC, we evaluated area under the curve (AUC) of receiver–operating–characteristic (ROC) curves¹⁶ and net reclassification improvement (NRI)¹⁷ with low–risk and high–risk

categories. Compared to the base model, including circulating BCAAs led to a significant increase in AUC (Supplementary Table 5a and Supplementary Fig. 2), and a net 8.2% of cases moving to the high–risk category with a NRI of 5% (Supplementary Table 5b). Thus, in our population, inclusion of circulating BCAAs improved the ability of risk stratification models to identify future PDAC cases.

In stratified analyses, we noted no significant differences in the association of BCAAs with PDAC by cohort, sex, smoking status, BMI, or fasting status at blood collection (Supplementary Fig. 3, all *P*–interaction 0.14). To examine when circulating BCAAs were most associated with PDAC, we stratified cases and matched controls by time interval between blood collection and PDAC diagnosis (<2 years, 2–<5 years, 5–<10 years, 10 years). These analyses demonstrated particularly strong associations between 2 and 5 years prior to diagnosis (Fig. 2a and Supplementary Table 6).

Experimental studies indicate years elapse between formation of the initial malignant clone and cancer diagnosis¹⁸, suggesting that occult PDAC might be present when we observe the strongest associations with elevated BCAAs. We therefore hypothesized elevated circulating BCAAs were a marker of early PDAC. To test this possibility, we conducted a prospective serial blood sampling study utilizing *LSL–Kras^{G12D/+};LSL–Trp53^{R172H/+};Pdx–1–Cre* (KPC) mice which develop PDAC with variable latency¹⁹. KPC mice progress through all histological stages of disease from normal pancreata to invasive adenocarcinoma with a median survival of approximately 21 weeks¹⁹ (Supplementary Fig. 4a). KPC mice initially displayed similar BCAA levels to littermate controls, but developed significant elevations from 15–17 weeks prior to sacrifice (Fig. 2b and Supplementary Fig. 4b). These data suggest circulating BCAA elevations accompany early PDAC.

LSL–Kras^{G12D/+}; Trp53^{flox/flox};Pdx–1–Cre (KP^{-/–}C) mice develop PDAC with more consistent kinetics, displaying precursor lesions (PanINs) with limited invasive cancer by 3–4 weeks of age (Fig. 2c) and a median lifespan of 10-12 weeks²⁰. At 3–4 weeks of age we observed no difference in weight or food consumption between KP^{-/–}C mice and littermate controls suggesting animals with early PDAC had not yet developed overt constitutional symptoms (Fig. 2d and Supplementary Fig. 4c). Consistent with findings in patients and KPC mice, circulating BCAAs were higher in KP^{-/–}C animals with subclinical PDAC when compared with controls (Fig. 2e), a pattern not observed for most other amino acids (Fig. 2f and Supplementary Fig. 4d-e).

We observed no significant differences in fasting blood glucose, response to glucose load, response to insulin challenge, or fasting insulin levels during intraperitoneal glucose and insulin tolerance tests in four–week–old KP^{-/-}C and control mice (Fig. 2g–h and Supplementary 4f-i). These findings argue that BCAA elevations are not reflective of hyperglycemia or insulin resistance and instead represent an early consequence of subclinical PDAC.

We examined whether other malignancies induced by the same genetic lesions caused elevated plasma BCAAs. Cre–recombinase introduction into lung or muscle of mice with the *Kras* and *Trp53* alleles from the KP^{–/–}C model leads to non–small cell lung cancer and

sarcoma, respectively²¹⁻²³. Neither model displayed the BCAA alterations seen with early PDAC (Supplementary Fig. 5). Re-implantation of cell lines derived from the KP^{-/-}C model into immunocompetent syngenic hosts also failed to cause BCAA elevations (Supplementary Fig. 6). These data argue BCAA elevations are associated with early stage autochthonous tumors arising in the pancreas and are not a general feature of *Kras*-driven cancer. They also suggest re–implanting cells from end–stage disease does not model the early disease state that produces BCAA elevations.

Chronic pancreatitis is a risk factor for PDAC²⁴, and pancreatic inflammation can promote PDAC development and progression in mice^{25,26}. Therefore, we examined whether BCAA elevations might be a cause or consequence of pancreatic inflammation in early disease. Mild, chronic pancreatitis in the absence of tumorigenesis failed to recapitulate BCAA elevations (Supplementary Fig. 7a-h), and prolonged increases in plasma BCAAs from dietary intervention did not cause pancreatic inflammation or pancreatitis (Supplementary Fig. 7i-o). Further studies are needed to understand the relationship between BCAAs and more severe pancreatitis.

Unlike other amino acids, the liver does not regulate plasma BCAAs levels^{27,28}; instead, levels are determined by dietary uptake, tissue utilization, and breakdown of muscle and other body protein stores^{27,29}. Therefore, plasma BCAAs may originate from short–term pools related to dietary uptake and disposal or long–term pools related to breakdown of tissue proteins. To interrogate the short–term pool, we fed four–week–old KP^{–/–}C and control mice an amino acid defined diet, in which 20% of leucine and valine were ¹³C–labeled. KP^{–/–}C and control mice consumed similar amounts of food when exposed to labeled diet for two hours, (Supplementary Fig. 8a) and we observed no difference in appearance and disappearance of plasma ¹³C–label (Fig. 3a and Supplementary Fig. 8b), arguing that gut uptake and peripheral disposal are similar in mice with or without PDAC.

To determine the contribution of long–term BCAAs pools to plasma levels, we exposed mice to labeled diet during a period of rapid growth in early life, and then switched them to unlabeled diet for three days to chase label from the short–term pool (Fig. 3b). Despite similar peripheral tissue protein labeling (Fig. 3c and Supplementary Fig. 8c), the fraction of labeled BCAAs in plasma was elevated in four–week–old KP^{-/-}C mice relative to littermate controls (Fig. 3d). Furthermore, by comparing plasma label under fed conditions, which is determined by both labeled long–term and unlabeled short–term pools, and fasted conditions, in which only labeled long–term pools contribute, we calculated that increased liberation of BCAAs from long–term body stores was solely responsible for BCAA elevations in KP^{-/-}C mice (Fig. 3d–e and Supplementary Fig. 8d). These data suggest that an early consequence of PDAC is enhanced breakdown of tissue proteins leading to plasma BCAA elevations. Consistent with this hypothesis, KP^{-/-}C mice with early PDAC have smaller fast–twitch muscles with sparing of slow–twitch and cardiac muscle (Fig. 3f and Supplementary Fig. 9). Notably, muscle atrophy associated with prolonged fasting and late–stage malignancy exhibits a similar pattern³⁰⁻³².

Increased muscle catabolism represents one aspect of cancer–associated cachexia, a wasting syndrome frequently affecting patients with advanced PDAC and contributing to worse

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outcomes³³⁻³⁶. Our findings, however, suggest that protein breakdown begins much earlier than previously appreciated and pre–dates onset of clinical cachexia. Inflammatory cytokines produced by immune and/or tumor cells have been implicated in cachexia^{31,37} and the low disease burden at the time of BCAA elevations suggests hormonal factors may be involved here as well. Liberation of tissue amino acids could support the elevated amino acid requirements of pancreatic cancer cells^{38,39}, with BCAAs and/or other amino acids derived from tissue breakdown contributing to disease progression. Because hepatic metabolism maintains relatively constant plasma levels of all amino acids except BCAAs^{27,29,40}, increased liberation of tissue amino acids are expected raise BCAA concentrations in blood. The association between elevated BCAAs and other metabolic disease states^{11,13,14,41} suggests that plasma BCAAs could be a general marker of increased protein turnover, and elevated BCAAs could contribute to the peri-diagnostic hyperglycemia commonly found in patients with PDAC⁴².

In participants from four large prospective cohorts, circulating BCAAs were associated with future development of PDAC. We observed similar BCAA elevations in two mouse models of PDAC, and demonstrated these elevations result from breakdown of peripheral protein stores. These findings provide new insight into how early disease impacts whole-body metabolism and suggest that muscle protein loss occurs much earlier in disease progression than previously appreciated.

ONLINE METHODS

HUMAN STUDIES

Study Population—Our study population included pancreatic cancer cases and controls from four prospective cohort studies: Health Professionals Follow–Up Study (HPFS), Nurses' Health Study (NHS), Physicians' Health Study I (PHS), and Women's Health Initiative–Observational Study (WHI–OS). HPFS was initiated in 1986 when 51,529 U.S. men 40–75 years working in health professions completed a mailed biennial questionnaire. NHS was established in 1976 when 121,700 female nurses aged 30–55 years completed a mailed biennial questionnaire. PHS is a completed trial initiated in 1982 of aspirin and β – carotene among 22,071 male physicians, aged 40–84 years. After trial completion, study participants were followed as an observational cohort. WHI–OS consists of 93,676 postmenopausal women, aged 50–79 years, enrolled from 1994–1998 at 40 U.S. clinical centers. Participants completed a baseline clinic visit and annual mailed questionnaires. The study was approved by Human Research Committee at Brigham and Women's Hospital (Boston, MA) and participants provided consent.

We included incident pancreatic adenocarcinoma cases diagnosed after blood collection through 2010 with available plasma and no prior history of cancer. Cases were identified by self– report or follow–up of deaths. Deaths were ascertained from next–of–kin, postal service, or National Death Index; this method captures >98% of deaths⁴³. Medical records were reviewed by physicians blinded to exposure data to confirm pancreatic cancer diagnoses. Similar to prior studies in these cohorts^{4,44-46} and based on a predefined analysis plan, we included only cases diagnosed 2 years after blood collection, as the weight loss and insulin resistance which develop due to pancreatic cancer manifest in the 2 years before

diagnosis^{42,47}. For each case, we randomly selected two controls, matching on cohort (also matches on sex), year of birth (+/–5 years), smoking status (never, past, current, missing), fasting status (<8 hours, 8 hours), and month/year of blood collection (+/–3 months in HPFS, +/– 3 months in NHS, +/–6 months in PHS, and exact matching in WHI). Controls were alive without cancer at the case's diagnosis date and provided a blood sample. Covariate data were obtained from baseline questionnaires in PHS and WHI and questionnaires prior to blood collection in HPFS and NHS, as described previously^{4,45}. In HPFS, NHS, and PHS, cancer stage among cases was directly classified based on medical record review as: (1) local disease amenable to surgical resection; (2) locally advanced disease that is unresectable, but without distant metastases, or (3) distant metastatic disease. In WHI, medical records were coded using Surveillance Epidemiology End Results summary staging, which classifies tumors as localized, regional, or distant. These stages were then classified in the same manner as in HPFS, NHS and PHS, as local, locally advanced and metastatic disease, respectively.

The initial dataset included 454 cases and 908 controls. Seven controls had insufficient plasma for metabolite profiling. One case and control were excluded due to missing data for >10% of metabolites.

Plasma Samples—Blood samples in EDTA tubes were collected from 18,225 men in HPFS from 1993–1995, 14,916 men in PHS from 1982–1984, and 93,676 women in WHI–OS from 1994–1998, and in Heparin tubes for 32,826 women in NHS from 1989–1990. Comparisons of participant characteristics in the blood collection cohort and the full cohort are provided for each study in Supplementary Table 7. Blood samples in HPFS and NHS were collected by participants, mailed overnight on cold–packs, and then spun to collect and store plasma (delayed processing), while PHS and WHI participants' whole blood was separately immediately into plasma and stored. An overview of procedures for collection and storage of samples from each cohort is provided below and summarized in Supplementary Table 8.

Health Professionals Follow–up Study: Upon arrival at the blood lab, vials were centrifuged in order to separate the various component parts. Cryo storage tubes were labeled with the appropriate study members ID number and the separated blood components were pipetted into them. This process produced 5 tubes of plasma, 2 tubes of white blood cells, and 1 tube of red blood cells for each cohort member. The tubes were then stored in liquid nitrogen freezers. A bulk tank, holding up to 3,000 gallons of LN2 automatically feeds each individual freezer whenever the freezer's sensors indicate that coolant is required.

Nurses' Health Study: Blood samples were separated into components (plasma, white blood cells and red blood cells) and pipetted into 8 cryotubes with 5 tubes of plasma, 2 tubes of white blood cells and 1 tube of red blood cells. Samples were immediately frozen in vapor phase liquid nitrogen freezers. The NHS Blood Lab stores all biologic samples associated with the Blood Study in-house in a large liquid–nitrogen (LN2) freezer farm. The cryotubes are stored in the vapor phase of LN2 freezers; the highest freezer temperature is – 130°C near the top of the freezer and the lowest temperature is –196°C at the bottom near

the liquid nitrogen. All freezers are alarmed and monitored continuously either by NHS laboratory staff or a central security desk (nights and weekends).

Physicians Health Study: Blood collection kits were sent to all participants with instructions to have blood drawn into the EDTA tubes that were provided. Two tubes were centrifuged for plasma, and a third tube was for whole blood. The specimens were received in the laboratory on chill packs within 24 hours of being drawn. Upon receipt, the samples were refrigerated and re–aliquotted into nine 1.2 mL tubes (three whole blood and six plasma), all frozen at –82°C.

Women's Health Initiative: Blood samples were collected on all WHI–OS participants at a baseline clinic visit in the fasting state. Blood samples were maintained at 4°C for up to one hour until plasma or serum was separated from cells. Centrifuged aliquots were put into – 70°C freezers within two hours of collection. Samples were shipped frozen by overnight delivery to a central facility and kept within –70°C freezers.

Plasma samples were grouped based on cohort, so that all cases and controls from a single cohort study underwent metabolite profiling as a batch. Sample triplets (pancreatic cancer case, matched control #1, and matched control #2) were distributed randomly within the batch, and the order of the case and two matched controls within each triplet was also randomly designated. Therefore, the case and its two controls were always run in the same batch and were always directly adjacent to each other in the analytic run, thereby limiting variability in platform performance across matched case–control triplets.

For participants from all four cohorts, plasma samples were thawed once to aliquot them from large volume vials into the smaller volumes needed for shipment to the Broad Institute of the Massachusetts Institute of Technology and Harvard University (Cambridge, MA). The samples were refrozen at the Broad Institute and then thawed a second time to perform metabolite profiling. Therefore, for all cases and controls, plasma samples had been thawed twice at the time of metabolite profiling.

We previously measured hemoglobin A1c (HbA1c) in 389 cases and 757 controls in the laboratory of Dr. Nader Rifai (Children's Hospital, Boston, MA) using reagents from Roche Diagnostics (Indianapolis, IN). We measured plasma insulin in 386 cases and 743 controls, plasma proinsulin in 388 cases and 746 controls, and plasma C-peptide in 408 cases and 785 controls using reagents from Diagnostic Systems Laboratory (Webster, TX) and Millipore Corporation (Billerica, MA). Randomly inserted samples from quality control (QC) plasma pools had mean intra–assay coefficients of variance (CVs) of 2.0% for HbA1c, 5.4% for insulin, 3.1% for proinsulin, and 4.9% for C–peptide⁴.

Metabolite Profiling—Profiles of endogenous, polar metabolites were obtained using liquid chromatography–tandem mass spectrometry (LC–MS) at the Broad Institute of the Massachusetts Institute of Technology and Harvard University (Cambridge, MA). The LC–MS methods were designed to enable broad measurement of metabolic markers and intermediates, including metabolites from central metabolism and amino acid metabolism, using low plasma sample volumes⁴⁸. LC–MS parameters for targeted analyses, including

chromatographic retention times and mass spectrometry multiple reaction monitoring settings (declustering potentials, collision energies, and lens voltages), were determined using over 300 commercially available reference compounds. A subset of 133 polar metabolites were measurable in plasma using a combination of two distinct hydrophilic interaction liquid chromatography (HILIC) methods, one operated under acid mobile phase conditions with positive ion mode MS detection and the other under basic elution conditions with negative ion mode MS detection.

The acidic HILIC method using positive ionization mode MS analyses was similar to the method described in Wang et al¹¹. Briefly, the LC–MS system consisted of a 4000 QTRAP triple quadrupole mass spectrometer (AB SCIEX) coupled to an 1100 Series pump (Agilent) and an HTS PAL autosampler (Leap Technologies). Plasma samples (10µL) were extracted using nine volumes of 74.9:24.9:0.2 (v/v/v) acetonitrile/methanol/formic acid containing stable isotope– labeled internal standards (valine–d8, Isotec; and phenylalanine–d8, Cambridge Isotope Laboratories). The samples were centrifuged (10 min, 9,000 x g, 4°C) and the supernatants (10 µL) were injected onto an Atlantis HILIC column (150 x 2.1 mm, 3 µm particle size; Waters Inc.). The column was eluted isocratically at a flow rate of 250 µL/min with 5% mobile phase A (10 mM ammonium formate and 0.1% formic acid in water) for 1 minute followed by a linear gradient to 40% mobile phase B (acetonitrile with 0.1% formic acid) over 10 minutes. The ion spray voltage was 4.5 kV and the source temperature was 450°C.

A second method using basic HILIC separation and negative ionization mode MS detection was established on an LC–MS system consisting of an ACQUITY UPLC (Waters Inc.) coupled to a 5500 QTRAP triple quadrupole mass spectrometer (AB SCIEX). Plasma samples (30μ L) were extracted using 120 μ L of 80% methanol (VWR) containing the internal standards inosine–¹⁵N₄, thymine–d₄, and glycocholate–d₄ (Cambridge Isotope Laboratories). The samples were centrifuged (10 min, 9,000 x g, 4°C) and the supernatants were injected directly onto a Luna NH2 column (150 x 2.0 mm, 5 μ m particle size; Phenomenex) that was eluted at a flow rate of 400 μ L/min with initial conditions of 10% mobile phase A (20 mM ammonium acetate and 20 mM ammonium hydroxide (Sigma– Aldrich) in water (VWR)) and 90% mobile phase B (10 mM ammonium hydroxide in 75:25 v/v acetonitrile/methanol (VWR)) followed by a 10 min linear gradient to 100% mobile phase A. The ion spray voltage was –4.5 kV and the source temperature was 500°C.

Raw data were processed using MultiQuant 1.2 software (AB SCIEX) for automated LC– MS peak integration. All chromatographic peaks were manually reviewed for quality of integration and compared against a known standard for each metabolite to confirm compound identities. Internal standard peak areas were monitored for quality control, to assess system performance over time, and to identify any outlier samples requiring re– analysis. A pooled plasma reference sample was also analyzed after sets of 20 study samples as an additional quality control measure of analytical performance and to serve as reference for scaling raw LC–MS peak areas across sample batches. Metabolites with a signal to noise ratio <10 were considered unquantifiable. Metabolite signals were analyzed in relation to pancreatic cancer risk as LC–MS peak areas, which are proportional to metabolite concentration and appropriate for metabolite clustering and correlative analyses.

Of the 133 metabolites measured (Supplementary Fig. 10), 83 were included in the analyses of our nested pancreatic cancer case–control population (Supplementary Table 9). In pilot work⁴⁹, we determined that 32 metabolites had poor reproducibility in samples with delayed processing, so these metabolites were excluded as they could not be reliably measured in two of the participating cohorts. In the current study, three Heparin plasma pools (57 total QC samples) and three EDTA plasma pools (128 total QC samples) were randomly interspersed among participant samples as blinded QC samples. We calculated mean CVs for each metabolite across QC plasma pools and set an *a priori* threshold of 25% for satisfactory reproducibility. Based on this criterion, 13 metabolites with mean CV>25% were excluded from our analyses. Five metabolites had undetectable levels for >10% of participants and were also excluded. We evaluated plasma from ten volunteers with plasma collected simultaneously in Heparin and EDTA tubes. For the branched chain amino acids, Spearman correlation coefficients between Heparin and EDTA samples were 0.85 for isoleucine, 0.88 for leucine, and 0.95 for valine.

For metabolites meeting the threshold for statistical significance after multiple- hypothesis correction (isoleucine, leucine and valine), LC-MS peak areas were converted to absolute concentrations using stable isotope-labeled standards. Briefly, external calibration curves of MS response were determined using solutions of isotope–labeled ${}^{13}C_6$, ${}^{15}N$ – leucine, ¹³C₆, ¹⁵N-isoleucine (Cambridge Isotope Laboratories), and d₈-valine (Isotec). A 1 $\mu g/\mu L$ solution of each standard was prepared in water. 20 μL of each stock solution were added to 180 µL of reference pooled plasma and the resulting solution was then serially diluted using pooled plasma to generate a calibration curve. For multiple reaction monitoring MS analyses, the bond cleavage products and collision energy and declustering potential settings were the same as those used for the endogenous metabolites: natural leucine 132/86, CE=18 and DP=50; ¹³C₆, ¹⁵N- leucine 134/87, CE=18 and DP=50; natural isoleucine 132/86, CE=18 and DP=50; ¹³C₆, ¹⁵N- isoleucine 139/92, CE=18 and DP=50; natural valine 118/72, CE=18 and DP=25, and d_8 -valine 126/80, CE=18 and DP=25. Three separate plasma samples were prepared at each concentration and were analyzed using the acidic HILIC LC-MS method described above. The median concentrations of endogenous isoleucine, leucine and valine in the reference pooled plasma were calculated and the concentration of each metabolite in each study sample was determined from the response ratio relative to the nearest reference pooled plasma sample in the analysis queue.

Statistical Analysis—To compare baseline characteristics, we used conditional logistic regression conditioned on the matching factors and including the covariate of interest. Partial Spearman correlation coefficients were calculated for metabolites and covariates. Metabolites were log– transformed to improve normality and included as continuous variables in conditional logistic regression models conditioned on matching factors and adjusted for age at blood draw (years, continuous), fasting time (<4, 4–8, 8–12, 12 hours, missing) and race/ethnicity (White, Black, other, missing). Using a conservative Bonferroni correction for multiple–hypothesis testing⁵⁰, metabolites with *P* 0.0006 (0.05/83) were considered statistically significant.

To provide estimates of effect magnitude, significant metabolites were examined in conditional logistic regression models after categorization into quintiles based on log-

transformed metabolite levels in controls. Separate quintiles were generated for fasting (8 hours since last meal) and non–fasting (<8 hours since last meal) participants, given the possible effect of fasting time on metabolite levels. Quintiles were generated from the population of selected controls, which may not exactly reflect the characteristics of the full cohort population. Odds ratios (OR) and 95% confidence intervals (CI) were also calculated per standard deviation (SD) change in log–transformed metabolite levels. To control for possible confounding, we evaluated regression models adjusted for body–mass index (BMI), physical activity, history of diabetes mellitus, HbA1c, plasma insulin, plasma proinsulin, and plasma C–peptide. We also evaluated regression models after excluding subjects with diabetes by self–report or HbA1c 6.5% at blood collection (prevalent diabetes). We further evaluated models that excluded subjects who developed diabetes after blood collection but >2 years prior to cancer diagnosis (incident diabetes not thought to be recent onset from pancreatic cancer)^{42,47}.

Metabolite values were considered missing when an LC–MS peak was below the limit of detection. In the primary analysis, any case or control with missing data for a metabolite was excluded from the analysis of that metabolite. However, we also conducted sensitivity analyses, in which participants with missing values were assigned the lower limit of detection or half of the lower limit of detection, and our results were unchanged.

We assessed heterogeneity of metabolite associations with pancreatic cancer risk across cohorts using Cochran's Q-statistic⁵¹. We examined associations in predefined subgroups by sex, smoking status, BMI, and fasting status. Statistical interactions were assessed by entering into models the main effect terms and cross–product terms of metabolites and stratification variables, evaluating likelihood ratio tests. We also examined associations by time between blood collection and the case's cancer diagnosis. In these time–based analyses, one stratum included 40 pancreatic cancer cases with blood collected within 2 years of diagnosis and their matched controls. These cases and controls were not part of the primary analysis population, but were included in the stratified analyses by time to more fully delineate the associations were also examined for circulating BCAAs with previously explored risk factors for pancreatic cancer in our cohorts (Supplementary Table 10) and with cancer stage at diagnosis (Supplementary Table 11).

Since association of a marker with disease does not indicate the suitability of the marker to serve as a screening test for the disease, we examined two approaches to quantify the value of metabolites in a multi–factor risk discrimination tool for pancreatic cancer. Discrimination quantifies the ability of one or more disease markers to separate cases (individuals with the disease) from controls (individuals without the disease). We investigated the discrimination of risk models for predicting pancreatic cancer diagnosis in the ten years after measurement of circulating BCAAs, i.e. the 10–year risk of pancreatic cancer.

In the first approach, we investigated receiver–operating–characteristic (ROC) curve analysis and calculated of the area under the ROC curve (AUC), also known as the concordance (C) statistic^{16,52}. The base model included age at blood collection (continuous),

cohort (HPFS, NHS, PHS, WHI; which also accounts for sex), race/ethnicity (white, black, other/missing), smoking status (never, past, current, missing) and fasting time (<4, 4–8, 8– 12, 12 hours, missing). Three subsequent models mirrored the base model, but additionally included: (1) body-mass index, physical activity, and history of diabetes, (2) circulating BCAAs, or (3) body- mass index, physical activity, history of diabetes, and circulating BCAAs. Each point on the ROC curve shows the effect of a rule for turning a risk estimate into a prediction of the development of an event. The y axis of the ROC curve is the true positive rate or sensitivity (i.e., the proportion of individuals with pancreatic cancer who were correctly predicted to have the disease). The x axis shows the false positive rate, which is the complement of specificity (i.e., the proportion of individuals without pancreatic cancer who were incorrectly predicted to have pancreatic cancer). The area under the ROC curve, the AUC, measures how well the model discriminates between case subjects and control subjects. An ROC curve that corresponds to a random classification of case subjects and control subjects is a straight line with an AUC of 50%. An ROC curve that corresponds to perfect classification has an AUC of 100%. The improvement in AUC for a model containing a new marker is defined as the difference in AUCs calculated using a model with and without the new marker of interest⁵³.

For context, the Breast Cancer Risk Assessment Tool, commonly referred to as the Gail Model^{54,55}, estimates a woman's risk for breast cancer using clinically available information including current age, age at menarche, age of first live birth, number of first degree relatives with breast cancer, number of previous breast biopsies, breast biopsies that show atypical hyperplasia, and race/ethnicity. The Gail Model is used to counsel women on appropriate screening tests for breast cancer⁵⁶, the use of tamoxifen as a chemopreventative agent⁵⁷, and for determining sample size calculations in randomized clinical trials of prevention strategies⁵⁸. Several studies have evaluated the discrimination of the Gail model using ROC curve analysis and calculated the AUC to be 0.58 to 0.63^{52,59-61}. Follow–up studies have described an AUC of 0.62 to 0.66 when breast density is added as an additional predictor in the original Gail Model^{59,60,62}.

Although ROC curves are commonly used, they have a number of limitations and may underestimate the ability of a new marker to contribute to risk prediction when added to previously defined predictors⁶³⁻⁶⁶. Another approach to evaluating model discrimination is to evaluate the ability of a new marker to shift an individual's risk up or down between pre– defined risk categories. This is known as the prediction increment of a marker and has been codified in an approach known as net reclassification improvement (NRI)¹⁷. The NRI (sometimes referred to as the net reclassification index) constructs reclassification tables separately for participants with and without events, and quantifies the correct movement between categories of risk; namely, to higher risk categories for participants with events and to lower risk categories for those without events. Furthermore, incorrect movement in categories of risk (downwards for events and upwards for non–events) reduces the net correct reclassification of individuals within the study population.

The NRI calculation is represented by the following formula:

$$NRI = [P(up|D=1) - P(down|D=1)] - [P(up|D=0) - P(down|D=0)]$$

Upward movement (up) is defined as a change into a higher risk category based on the new model and downward movement (down) as a change into a lower risk category based on the new model, where P indicates probability and D denotes the event indicator (1, event; 0, non–event).

Using the NRI, we evaluated the ability of the prediction model including circulating BCAAs to appropriately reclassify individuals into risk groups compared to the base model. The base model was calculated using conditional logistic regression conditioned on matching factors and adjusted for race/ethnicity, body–mass index, physical activity and history of diabetes. The subsequent model included the covariates in the base model with the addition of circulating metabolites. As in prior studies⁶⁷⁻⁶⁹, we defined the high–risk group as those individuals with risk for pancreatic cancer at least 2–fold greater than an individual with average risk.

For context, the Emerging Risk Factors Collaboration⁷⁰ has examined the integration of novel risk factors into risk prediction models for cardiovascular disease. In these studies, additional potential risk predictors were added to a model of known risk predictors for cardiovascular disease, including age, sex, smoking status, blood pressure, history of diabetes, and cholesterol. The net reclassification improvement was then calculated for three 10-year risk categories for cardiovascular disease. C-reactive protein is a marker of systemic inflammation, and elevated CRP has been associated with an increased risk for cardiovascular events in numerous studies⁷¹⁻⁷³. Circulating CRP is currently used to inform decisions in the clinic regarding screening and risk reduction strategies^{74,75} and to design clinical trials testing novel treatments to reduce cardiovascular events^{76,77} In an analysis of nearly 250,000 individuals⁷⁸, the addition of CRP to know cardiovascular disease risk factors was associated with a statistically significant improvement in the area under the ROC curve and a NRI of 1.52% for 10-year risk of cardiovascular disease. In contrast, additional analyses demonstrated a <1% improvement in the NRI for body-mass index, waist circumference, waist-to-hip ratio, plasma fibrinogen, and circulating apolipoproteins⁷⁸⁻⁸⁰, such that the clinical utility of these additional predictors remains unclear⁷⁵.

All analyses were performed with SAS 9.2 statistical package. All *P*-values were two-sided.

MOUSE STUDIES

All studies were approved by the MIT committee on animal care (IACUC).

Experimental Mice—All experimental groups were assigned based on genotype. All animals were numbered and experiments conducted blinded. After data collection, genotypes were revealed and animals assigned to groups for analysis.

<u>KPC</u>: Experimental KPC mice were male mice on a mixed background, heterozygous for the conditional lox–stop–lox *Kras^{G12D}* allele, heterozygous for the conditional lox–stop–lox

 $Trp53^{R172H}$ allele and expressed Cre–recombinase under control of the Pdx–1–promoter $(Tg(Ipf1-cre)I^{Tuv})^{19}$. Littermate controls lacked either the LSL– $Kras^{G12D}$ allele, the Cre allele, or both. Control mice were sacrificed at the same time as their tumor–bearing littermates.

<u>KP^{-/-}C:</u> Experimental KP^{-/-}C mice were male mice on a mixed background, heterozygous for the conditional lox–stop–lox $Kras^{G12D}$ allele, homozygous for loxP sites flanking exons 2–10 of Trp53 and expressed Cre–recombinase under control of the Pdx–1– promoter $(Tg(Ipf1-cre)I^{Tuv})^{20}$. Littermate control mice lacked either the Cre–recombinase allele, *LSL–Kras*^{G12D} allele, or both. Inbred C57B6/J male mice containing the same alleles were also examined where indicated, and cancer cell lines derived from these mice were used for syngenic re–implantation studies.

<u>Non-small cell lung cancer:</u> Six-month-old male mice on a pure 129 background, heterozygous for the conditional lox-stop-lox $Kras^{G12D}$ allele, homozygous for loxP sites flanking exons 2–10 of Trp53 were administered 2.5x10⁷ pfu of Cre-expressing adenovirus as previously described^{21,22}. High-titer adenovirus was obtained from the Gene Transfer Vector Core (University of Iowa).

<u>Hindlimb sarcoma:</u> Four–week–old male mice on a mixed background, heterozygous for the conditional lox–stop–lox *Kras^{G12D}* allele and homozygous for loxP sites flanking exons 2-10 of *Trp53* were administered 2.5×10^8 pfu of Cre–expressing adenovirus as previously described²³. High–titer adenovirus was obtained from the Gene Transfer Vector Core (University of Iowa).

<u>Re-implantation, Pancreatitis and BCAA diet studies:</u> Male C57B6/J mice aged 4–6 weeks at the start of the study were used for these experiments.

Diets-Standard Chow Diet: RMH 3000 (Prolab).

<u>Amino Acid Defined Diet:</u> 1x BCAA (TD.110839) and 2x BCAA (TD.110843) were designed in consultation with and subsequently obtained from Harlan Teklad.

Labeled Amino Acid Defined Diet: 20% ¹³C–leucine and 20% ¹³C–valine labeled diets were based on diet TD.110839 and produced by Cambridge Isotopes and Harlan Teklad.

Plasma for Metabolomics—Plasma was collected for each experiment at the time points indicated. Mice were anesthetized under 2% isofluorane–oxygen mixture and retro–orbitally bled approximately 4.5 hours after the onset of the light cycle. Blood was immediately placed in EDTA–pretreated tubes and centrifuged to separate plasma. Plasma was aliquoted and frozen at -80°C for further analysis. Fasting blood samples were harvested in the same manner first thing in the morning after a 16–hour overnight fast.

Food Consumption—Mice were housed individually for 48 hours and remaining food pellets weighed at zero, 24 and 48 hours. A two–day average was then calculated for each mouse. Body weight was determined on the second day. To assess consumption of BCAA

defined diets, mice were housed individually and fed diets for 5 days. Food was weighed after 2 days of feeding and again on day 5, and the average consumption per 24hrs over the 72hr period was calculated.

Blood glucose, plasma insulin, glucose tolerance test, and insulin tolerance

test—Four–week– old KP^{-/-}C mice were fasted overnight and blood glucose measured using a One Touch Ultra handheld glucometer. 25μ L of plasma from the same mice was harvested in heparinized tubes, aliquoted, and frozen at -80° C for further analysis. Plasma insulin levels were determined using an ultrasensitive mouse insulin ELISA kit (Crystal Chem, #90080). After measuring fasting parameters, a glucose tolerance test was performed in accordance with published protocols⁸¹. Briefly, conscious mice received an intraperitoneal injection of 2g/kg glucose at time zero. Blood glucose was subsequently measured at 15, 30, 60, 90 and 120 minutes post–injection as described above. For insulin tolerance test, four–week–old KP^{-/-}C mice were fasted for 6 hours during daytime hours. Following initial blood glucose measurement, conscious mice received an intraperitoneal injection of 0.75 IU/kg recombinant human insulin (Novolin, Novo Nordisk). Blood glucose was subsequently measured at 15, 30, 60 and 90 minutes post–injection as described above.

KP^{-/-}**C Cell Lines and Re–implantation Studies**—End–stage tumors were dissected from C57B6/J KP^{-/-}C mice and mechanically chopped prior to trypsin disaggregation, with tumor cells then propagated in DMEM with 10% FBS, 2mM glutamine and penicillin/ streptomycin. Cell lines were negative for mycoplasma. For subcutaneous re–implantation studies, recipient mice were anesthetized with inhaled 2% isofluorane–oxygen mixture, low passage cell lines (passage 5 for each line) were resuspended at 2.5 x 10⁵ cells/100µL in sterile PBS, and100µL of either PBS (control) or cell suspension was injected in the flank of syngeneic mice. For orthotopic re–implantation studies, recipient mice were anesthetized with inhaled 2% isofluorane–oxygen mixture, a vertical incision made in the abdomen at the left mid–calvicular line, the spleen mobilized, and 25µL of either PBS or PBS containing 1.0 x 10⁵ cells (passage 3 for each line) was injected into the tail of the pancreas.

Caerulein–induced Chronic Pancreatitis—Mice were treated with either USP–grade saline or 5µg caerulein (Sigma) via intraperitoneal injection daily, 5 days per week for 10 weeks as previously described²⁵. Blood was obtained and the mice sacrificed on the final day. Tissues were fixed in 10% formalin for subsequent histological analysis.

Plasma Markers of Pancreatitis—Plasma amylase and lipase measurements were performed by IDEXX BioResearch Laboratory (North Grafton, MA).

BCAA Diet Studies—Mice were fed either 1x or 2x BCAA diets for 10 weeks. Blood was obtained and mice sacrificed on the final day of the experiment. Tissues were fixed in 10% formalin for subsequent histological analysis.

Studies to determine source of BCAA elevations

<u>Acute uptake and disposal:</u> Following a 16– hour overnight fast, mice were fed 20% ¹³C– leucine and valine containing diet for two hours before removal of food, and food

consumption during this period quantified as described above. At the time points indicated by the red arrowheads in Figure 3a, $10-25\mu$ L of plasma was harvested from the tail vein of conscious mice in a heparinized tube and centrifuged to separate plasma. Plasma was aliquoted and frozen at -80° C for subsequent GC–MS analysis. Total ion counts from GC–MS analysis of leucine and valine were then normalized to norvaline internal standard and multiplied by fractional labeling to determine the amount of label present. This number for each animal was then normalized to that animal's food intake to control for inter– animal variation in labeled food consumption.

Long-term pool contribution of BCAA to plasma: Mice were exposed to 20% 13 C– leucine and valine labeled diets from 7 days to age to 24 days of age followed by three days of unlabeled diet (according to the protocol depicted in Figure 3b). Two cohorts of mice were used in this study. One cohort of mice was sacrificed on day 27 in the fed state and a second cohort of mice was sacrificed on day 28 after a 16–hour overnight fast (the points indicated by the red arrowheads in Figure 3b). At sacrifice, anesthetized mice were terminally bled and tissues harvested within 5 minutes, snap frozen in liquid nitrogen using Biosqueezer (BioSpec Products), and stored at -80° C for subsequent GC–MS analysis.

Total contributions from short and long-term pools were calculated according to the following equations:

Raw data and calculations are summarized in Supplementary Table 12.

Tissue and body weights—For measurement of tissue weights, mice were weighed prior to sacrifice, then gastrocnemius, tibialis anterior, soleus and heart were subsequently dissected and weighed. Muscle weights for each individual mouse were normalized to the body weight of that mouse.

LC–MS Plasma Amino Acid Measurements—Plasma amino acids were measured by LC–MS at the Koch Institute of the Massachusetts Institute of Technology (Cambridge, MA) using similar methods used for assessment of metabolites in human plasma. Raw data were analyzed as peak area tops using the open–access MAVEN software tool⁸².

GC–MS Assessment of {}^{13}C–leucine and valine Labeling—Plasma polar metabolites were extracted in ice–cold 4:1 methanol:water with norvaline internal standard (5µL plasma in 200µL extraction solution). Extracts were clarified by centrifugation and the supernatant evaporated under nitrogen and frozen at –80°C for subsequent derivitization. Dried polar metabolites were dissolved in 20µL of 2% methoxyamine hydrochloride in pyridine (Thermo) and held at 37°C for 1.5hr. After dissolution and reaction, tert–butyldimethylsilyl derivatization was initiated by adding 25µL *N*–methyl–*N*–(tert–

butyldimethylsilyl)trifluoroacetamide + 1% tert- butyldimethylchlorosilane (Sigma) and

incubating at 37°C for 1hr. The acid hydrolysis protocol was adapted from Antoniewicz and colleagues⁸³. Briefly, acid hydrolysis of tissue proteins was performed on snap frozen tissues by boiling 1–5mg tissue in 1mL 18% hydrochloric acid overnight at 100°C. 50µL supernatant was evaporated under nitrogen and frozen at -80°C for subsequent derivitization. Dried hydrolysates were re–dissolved in pyridine (10µL/1mg tissue) prior to tert–butyldimethylsilyl derivatization, which was initiated by adding *N*–methyl–*N*–(tert–butyldimethylsilyl)trifluoroacetamide + 1% tert–butyldimethylchlorosilane (12.5µL/1mg tissue, Sigma) and incubating at 37°C for 1h.

GC/MS analysis was performed using an Agilent 7890 GC equipped with a 30m DB– 35MS capillary column connected to an Agilent 5975B MS operating under electron impact ionization at 70eV. One microlitre of sample was injected in splitless mode at 270°C, using helium as the carrier gas at a flow rate of 1mlmin⁻¹. For measurement of amino acids, the GC oven temperature was held at 100°C for 3min and increased to 300°C at 3.5° Cmin⁻¹. The MS source and quadrupole were held at 230°C and 150°C, respectively, and the detector was run in scanning mode, recording ion abundance in the range of 100–605 *m/z*. MIDs were determined by integrating the appropriate ion fragments⁸³ listed in Supplementary Table 13 and corrected for natural isotope abundance using an algorithm adapted from Ferandez and colleagues⁸⁴.

Statistics—Appropriate statistical tests were performed where required. Two–sided unpaired student's *t*–tests were performed for all statistical analyses unless otherwise specified using Mircosoft Excel for Mac:2011 (Microsoft) or GraphPad Prism 6 (GraphPad Software). Two– sided repeated–measures ANOVA was performed to compare mean plasma glucose levels in the glucose tolerance test and insulin tolerance tests⁸⁵, using SAS 9.2 statistical package.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Plasma metabolites and risk of future pancreatic cancer diagnosis

P-values of the log-transformed, continuous metabolite levels from human plasma comparing pancreatic cancer cases and controls in conditional logistic regression models conditioned on matching factors and adjusted for age at blood draw (years, continuous), fasting time (<4, 4–8, 8–12, 12 hours, missing) and race/ethnicity (White, Black, other, missing). The dashed green line indicates the statistically significant P-value threshold after Bonferroni correction for multiple-hypothesis testing, P-trend 0.0006 (0.05/83). The dashed blue line indicates P- trend of 0.05.

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a, Graph of odds ratio (95% confidence interval) for future pancreatic cancer diagnosis among cohort cases and matched controls comparing highest versus lowest quintiles of circulating BCAAs stratified by time from blood collection to the case's cancer diagnosis. Odds ratio (95% confidence interval) from conditional logistic regression models conditioned on matching factors and adjusted for age at blood draw (years, continuous), fasting time (<4, 4–8, 8–12, 12 hours, missing) and race/ethnicity (White, Black, other, missing). Red horizontal line marks an odds ratio of 1.0. **b.** Graph of total plasma BCAAs in

KPC mice over time and in littermate controls. Each control data point is an average for one mouse over the course of the study (Supplementary Fig. 3b) and values for KPC mice living longer than 19 weeks are averaged for the >19 weeks time point. For weeks 15-17, n = 6KPC and n = 9 control, *t*-test, *P*=0.001. **c**, H&E staining of pancreatic tissue obtained from KP^{-/-}C mice and littermate controls at 3-4 weeks of age. Tissue are shown from a control mouse with histologically normal pancreas (left); a KP-/-C mouse with areas of PDAC adjacent to areas of normal pancreas (middle); and a KP-/-C mouse with areas of PDAC and pancreatic intraepithelial neoplasia (closed arrow heads) (right); scale bar = 50μ M. d, Mean (\pm SEM) body weights at 3-4 weeks of age for KP^{-/-}C mice and littermate controls (n = 7 $KP^{-/-}C$, n = 11 control mice). e, Mean (\pm SEM) total plasma branched chain amino acid levels from KP^{-/-}C mice and littermate controls at 3-4 weeks of age ($n = 10 \text{ KP}^{-/-}$ C, n = 14control mice, t-test, P=0.002). f, P-values for comparison of circulating amino acid levels in $KP^{-/-}C$ mice and littermate controls at 3-4 weeks of age, cys = cystine (n = 10 $KP^{-/-}C$, n = 14 control mice), The dashed red line indicates *P*-value of 0.05. g, top, glucose tolerance test in KP^{-/-}C mice and littermate controls at the time of weaning ($n = 7 \text{ KP}^{-/-}$ C, n = 11control mice) and bottom, insulin tolerance test in KP-/-C mice and littermate controls at four weeks of age (n = 7 KP^{-/-}C, n = 15 control mice). h, Mean (\pm SEM) fasting plasma insulin levels from KP^{-/-}C mice and littermate controls at four weeks of age (n = 7 KP^{-/-}C, n = 11 control mice).



Fig. 3. BCAA elevations are derived from a long-term pool of amino acids

a, Plasma levels (mean \pm SEM) of ¹³C–labeled leucine and valine normalized to food intake over time following a two–hour exposure to diets containing ¹³C–labeled leucine and valine. The time points correspond to the red arrowheads in the diagram. **b**, Diagram of experiment using labeled diets to investigate contributions to plasma BCAA levels from long–term pools. Two cohorts of mice were used for these experiments, one sacrificed in the fed state and a second sacrificed in the fasted state at the time points indicated by the red arrowheads. **c**, Fractional labeling of total amino acids in protein hydrolysate of gastrocnemius muscle

from fasted KP^{-/-}C mice and control littermates (n = 8 KP^{-/-}C, n = 6 control). **d**, Fractional labeling of plasma amino acids in KP^{-/-}C and control mice in the fed state (n = 3 KP^{-/-}C, n = 4 control). **e**, The calculated contribution of the short– and long–term BCAA pools to the BCAAs present in plasma . **f**, Mean (±SEM) gastrocnemius weight (left panel, *t*-test, P=0.01), a predominantly fast–twitch muscle, and heart weight (right panel) normalized to body weight (n = 6 KP^{-/-}C, n = 10 control).

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Model	Extreme Quintiles b	Per S.D.	Extreme Quintiles b	Per S.D.	Extreme Quintiles b	Per S.D.	Extreme Quintiles b	Per S.D.
Base model	2.11 (1.40-3.18)	1.30 (1.15–1.48)	2.08 (1.38–3.13)	1.31 (1.14–1.50)	2.00 (1.37–2.92)	1.23 (1.09–1.39)	2.13 (1.43–3.15)	1.30 (1.14–1.48)
+ BMI and physical activity	2.05 (1.34–3.12)	1.29 (1.14–1.48)	2.01 (1.32–3.06)	1.29 (1.12–1.49)	1.94 (1.31–2.86)	1.21 (1.07–1.38)	2.06 (1.37–3.09)	1.28 (1.12–1.47)
+ BMI, physical activity, and reported diabetes at blood collection	2.00 (1.31–3.05)	1.28 (1.13–1.46)	1.97 (1.29–2.99)	1.28 (1.11–1.48)	1.90 (1.28–2.81)	1.20 (1.06–1.37)	2.01 (1.34–3.03)	1.27 (1.11–1.46)
+ BMI, physical activity, reported diabetes, HbA1c, plasma insulin, proinsulin and C-peptide	1.86 (1.13–3.03)	1.24 (1.06–1.45)	1.81 (1.11–2.96)	1.24 (1.04–1.47)	1.67 (1.06–2.63)	1.14 (0.99–1.33)	1.89 (1.17–3.06)	1.22 (1.04–1.44)
Exclude subjects with reported diabetes or HbA1c 6.5% at blood collection	2.12 (1.37–3.27)	1.33 (1.16–1.52)	2.16 (1.39–3.35)	1.32 (1.14–1.54)	1.91 (1.28–2.85)	1.23 (1.08–1.41)	2.19 (1.44–3.34)	1.31 (1.14–1.51)
Exclude subjects with reported diabetes or HbA1c 6.5% at blood collection and those with reported diabetes after blood collection	2.18 (1.39–3.43)	1.32 (1.15–1.53)	2.21 (1.40–3.49)	1.33 (1.13–1.55)	1.94 (1.27–2.96)	1.22 (1.07–1.40)	2.25 (1.45–3.49)	1.31 (1.13–1.52)
Abbreviations: BCAAs, bran	ched chain amino acids;	BMI, body-mass in	dex; CI, confidence inter	val; HbA1c, hemog	lobin A1c; S.D., standar	d deviation		

^aOdds ratio (95% confidence interval) from conditional logistic regression models conditioned on matching factors and adjusted for age at blood draw (years, continuous), fasting time (<4, 4–8, 8–12, 12) hours, missing) and race/ethnicity (White, Black, other, missing). Subsequent models also adjusted for indicated measure of energy balance, hyperglycemia or insulin resistance: body-mass index (kg/m², continuous), physical activity (MET-hr/week, continuous), reported diabetes at blood collection (yes/no), Hemoglobin A1c (%, continuous), plasma insulin (µU/ml, continuous), plasma proinsulin (pM, continuous), and plasma C-peptide (ng/ml, continuous).

^bOdds ratios (95% CI) for the comparison of the fifth quintile to the first quintile (referent) for the circulating branched chain amino acids

 $^{\mathcal{C}}$ Sum of the concentrations of the three circulating branched chain amino acids, isoleucine, leucine and value