ORIGINAL ARTICLE

Linking diet, physical activity, cardiorespiratory fitness and obesity to serum metabolite networks: findings from a population-based study

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OBJECTIVE: It is not yet resolved how lifestyle factors and intermediate phenotypes interrelate with metabolic pathways. We aimed to investigate the associations between diet, physical activity, cardiorespiratory fitness and obesity with serum metabolite networks in a population-based study.

METHODS: The present study included 2380 participants of a randomly drawn subcohort of the European Prospective Investigation into Cancer and Nutrition-Potsdam. Targeted metabolomics was used to measure 127 serum metabolites. Additional data were available including anthropometric measurements, dietary assessment including intake of whole-grain bread, coffee and cake and cookies by food frequency questionnaire, and objectively measured physical activity energy expenditure and cardiorespiratory fitness in a subsample of 100 participants. In a data-driven approach, Gaussian graphical modeling was used to draw metabolite networks and depict relevant associations between exposures and serum metabolites. In addition, the relationship of different exposure metabolite networks was estimated.

RESULTS: In the serum metabolite network, the different metabolite classes could be separated. There was a big group of phospholipids and acylcarnitines, a group of amino acids and C6-sugar. Amino acids were particularly positively associated with cardiorespiratory fitness and physical activity. C6-sugar and acylcarnitines were positively associated with obesity and inversely with intake of whole-grain bread. Phospholipids showed opposite associations with obesity and coffee intake. Metabolite networks of coffee intake and obesity were strongly inversely correlated (body mass index (BMI): \( r = -0.57 \) and waist circumference: \( r = -0.59 \)). A strong positive correlation was observed between metabolite networks of BMI and waist circumference (\( r = 0.99 \)), as well as the metabolite networks of cake and cookie intake with cardiorespiratory fitness and intake of whole-grain bread (\( r = 0.52 \) and \( r = 0.50 \); respectively).

CONCLUSIONS: Lifestyle factors and phenotypes seem to interrelate in various metabolic pathways. A possible protective effect of coffee could be mediated via counterbalance of pathways of obesity involving hepatic phospholipids. Experimental studies should validate the biological mechanisms.

International Journal of Obesity (2014) 38, 1388–1396; doi:10.1038/ijo.2014.39

Keywords: metabolomics; physical activity; diet; network analysis; systems epidemiology

INTRODUCTION

Targeted metabolomics, which simultaneously studies many well-defined substrates and products of metabolism, is a powerful approach to detect metabolic alterations that are linked to human behavior such as diet and physical activity.1–3 Furthermore, metabolite levels in body fluids may also reflect consequent phenotypes including cardiorespiratory fitness and obesity.4–7 Therefore, targeted metabolomics may particularly help unravel biological mechanisms and identify pathways and networks through which behavioral factors and phenotypes are linked to metabolism. Knowledge about these mechanisms may offer great potential in terms of prevention of chronic diseases, which are strongly linked to human behavior and phenotypes.8,9 Consequently, adequate dietary and lifestyle recommendations could be generated.

Within the European Prospective Investigation into Cancer and Nutrition (EPIC)-Potsdam study, we applied a targeted metabolomics platform that covers 127 serum metabolites, including amino acids, acylcarnitines, choline-containing phospholipids and hexose. In preceding analyses in the EPIC-Potsdam study, we have studied cross-sectional associations of diet, physical activity, cardiorespiratory fitness and obesity measures with these serum metabolites.10–13 In the present study, we attempted to provide a comprehensive picture of the associations of these human behaviors and consequent phenotypes with serum metabolite networks. To this end, we visualized these associations in our population-based cohort by applying an innovative network approach, and evaluated
relationships between different lifestyle and phenotype networks.

SUBJECTS AND METHODS

Study population
EPIC-Potsdam is part of the multicenter EPIC-study, a prospective cohort study conducted in 23 European centers, with the scope of investigating diet and cancer risk that also includes other risk factors and chronic diseases. EPIC-Potsdam is the largest German study center and recruited 27,548 adults of the Potsdam area aged mainly 35–65 years between 1994 and 1998.14 The EPIC-Potsdam study was approved by the ethics committee of the Medical Society of the State of Brandenburg. Participants provided written informed consent before they participated in the baseline examination, which included anthropometric and blood pressure measurements, a blood sample collection as well as a personal interview on medical history, a sociodemographic and lifestyle questionnaire and a self-administered food frequency questionnaire; further details are given below.15 All measurements were conducted by qualified staff following standardized procedures.15 During the blood sample collection, 30 ml of venous blood were drawn, rapidly processed and fractionated into serum, plasma, ‘buffy coat’ (leukocytes) and erythrocytes, which were stored in straws at −196 °C until metabolomic analysis.15

Of all the EPIC-Potsdam participants who had provided blood samples at baseline (n = 26,444), a subcohort of 2500 participants was randomly drawn for biomarker measurements.16 By randomly selecting the subcohort, it was assured that the results are representative of the full cohort. For the present analysis, the EPIC-Potsdam subcohort was used, and participants with implausible energy intakes (< 800 and >6000 kcal per day) or missing information on serum metabolite concentrations and covariates were excluded (n = 120). Baseline characteristics of the included participants have been published previously.10 In brief, participants were 61% women with mean age of 49.8 years and mean body mass index (BMI) of 26.1 kg m−2. Of the participants, 4.5% had prevalent type 2 diabetes, and those were more likely to have increased hexose levels.

In 2007, a subgroup of 208 EPIC-Potsdam participants took part in a validation study for physical activity assessment, which also included repeated objective measurement of physical activity and cardiorespiratory fitness and a blood sample collection at two time points 4 months apart. Note that this subgroup was independent of the previously described subcohort. Blood was drawn from participants who had fasted over night by qualitative analysis. Of the 208 participants, 50 men and 50 women was fractionated by centrifugation and stored in a freezer at −26 °C until metabolomic analysis.15

Measurement of serum metabolites
Metabolite concentrations were measured in 10 μl baseline serum samples of the EPIC-Potsdam subcohort and in serum samples of 100 participants of the validation study for physical activity assessment with the AbsoluteIDQ p150 kit (BIOCRATES, Innsbruck, Austria).24,25 The targeted metabolomics method simultaneously determined concentrations of 163 predefined metabolites including acylcarnitines (Cxy), amino acids, hexose (sum of six-carbon monosaccharides, including glucose, but without distinction of isomers) and choline-containing phospholipids (lyso-, diacyl- and acylalkyl-phosphatidylcholines). For the phosphatidylcholines, only the total number of carbon atoms and double bonds across two fatty-acid side chains could be determined. Lyso-phosphatidylcholines, sphingomyelins and acylcarnitines contained a single fatty-acid side chain that was detected. All samples were analyzed at the Genome Analysis Center (Helmholtz Zentrum München) between 2009 and 2010. Sample preparation and metabolite quantification of these cohort samples has been described previously in full detail.26,27 In brief, an automated robotic system (Hamilton ML Star, Bonaduz, Switzerland) conducted the following procedure: 10 μl of serum was pipetted onto filters with stable isolate-labeled internal standards in 96-well plates and dried in nitrogen stream. Amino acids were derivatized with 5% phenylisothiocyanat reagent and the plates were dried again. The other metabolites and internal standards were extracted using 5 ml ammonium acetate in methanol, and then centrifuged and filtered. Final extracts were diluted with mass spectrometry running solvent and analyzed using an API 4000 triple quadrupole mass spectrometer (AB Sciex, Darmstadt, Germany). Metabolites were quantified by multiple reaction monitoring in combination with internal standards, and metabolite concentrations were calculated with the MetQ software package (BIOCRATES). The metabolomics method has been validated by the manufacturer according to the Food and Drug Administration guideline ‘Guidance for Industry—Bioanalytical Method Validation, May 2011’, which implies proof of reproducibility in a certain error range. The limit of detection for the individual metabolites was set to three times the values of the buffer-only-containing samples.

Statistical analysis
Within the subcohort, the metabolite network was calculated using Gaussian graphical modeling.26 In brief, Gaussian graphical modeling is a data-driven method that uses the high degree of correlation between the high degree of correlation between the high degree of correlation between the high degree of correlation between lipid metabolites to construct metabolite networks. Each node in the network represents one metabolite and each edge between two nodes represents the dependency of two metabolites reflected by their partial correlation. Pearson’s partial correlation coefficients between each possible pair of metabolites were calculated with adjustment for all the other metabolites.
to elminate the indirect effects, and the covariates age and sex, in accordance with previous studies.28,29 As we used 127 metabolites, the metabolite network contained 127 nodes and could possibly contain \( \frac{127 \times (127 - 1)}{2} = 8001 \) edges. We chose a cutoff of >0.20 for the absolute value of the partial correlation coefficient between metabolites for the edges to be depicted. Therefore, the number of edges was reduced to 206. All calculations were done with SAS version 9.2 (SAS Institute Inc, Cary, NC, USA). Metabolite networks were then visualized with the yed graph editor (yWorks GmbH, Tuebingen; http://www.yworks.com).

In the next step, we considered different exposures. To reflect dietary behavior, we selected three foods that were previously found to be linked to serum metabolites and chronic disease risk in this population.10,11 This included low intake of whole-grain bread, as an established risk factor, as well as low intakes of coffee and cake and cookies as less-established risk factors. Physical activity was reflected by objectively measured physical activity energy expenditure. To mirror consequent phenotypes, we selected cardiorespiratory fitness and obesity with the single metabolites that we previously obtained.12,13 These were \( \beta \)-coefficient from linear mixed models regression analysis for physical activity and cardiorespiratory fitness, and Spearman’s partial correlation coefficients for diet and obesity measures. Physical activity and cardiorespiratory fitness data were additionally z-transformed to make the \( \beta \)-coefficients directly comparable. All of the association measures were multivariable adjusted for age, sex, alcohol intake, smoking, education and, if applicable, prevalent diseases and the other exposures. Note that the adjustment models differed slightly for different exposures as they were based on our previous studies.10–12 We colored the network based on the direction and strength of association between each exposure and the metabolites, and thus ended up with one metabolite network for each exposure.

To compare the metabolite networks of different exposures, we additionally calculated the correlation of the association measures between the different exposures and 127 serum metabolites, so essentially ‘the correlation of the correlation coefficients’. As an example to obtain the correlation of the metabolite networks of BMI and coffee, we calculated the correlation of the correlation coefficients between BMI and 127 serum metabolites with the correlation coefficients between coffee intake and 127 serum metabolites. As these were normally distributed for all exposures, we calculated Pearson’s correlation coefficients. Therefore, we obtained the correlation of metabolite networks of different exposures.

RESULTS

On the basis of their partial correlations, the 127 serum metabolites measured in 2380 EPIC-Potsdam participants were depicted in a metabolite network (Figure 1). In the metabolite network, there was a big cluster of phospholipids and acyl carnitines, one cluster of amino acids and C6-sugar. Within the network of the phospholipids, subgroups of sphingomyelins, lyso-phosphatidylcholines, diacyl-phosphatidylcholines and acyl-alkyl-phosphatidylcholines with different chain length and fatty-acid desaturation emerged. The center of the network was dominated by diacyl-phosphatidylcholines and their hydrolysis products (lyso-phosphatidylcholines). These were not only connected to acyl-acyl-phosphatidylcholines, sphingomyelins and acyl carnitines. Amino acids and C6-sugar were not connected to the phospholipids. Seven individual metabolites were not connected to any other metabolite, namely, hexose, ornithine, proline and C9-acyl carnitine, as well as diacyl-phosphatidylcholines C42:4, and acyl-alkyl-phosphatidylcholines C42:1 and C44:3. Free carnitine and propionyl-carnitine were linked as a pair. Although this network approach was purely data-driven, those phospholipids that are known to be only one reaction step apart were often strongly positively correlated, and thus, neighboring in the network, for example, sphingomyelins C18:0 and C18:1 (desaturase) and C16:1 and C18:1 (elongase) or amino acids glycine and serine (hydroxyl-methyl-transferase).

When looking at the association between diet, physical activity, cardiorespiratory fitness and obesity with the serum metabolite network, it was observed that the phenotypes (obesity and cardiorespiratory fitness) were more strongly linked to the metabolite network compared with the behavioral characteristics (diet and physical activity; Figures 2–4). The amino acids were particularly positively linked to cardiorespiratory fitness and physical activity, as well as intake of whole-grain bread and cake and cookies. Acylcarnitines were positively associated with obesity measures, and inversely associated with intake of whole-grain bread, coffee and cake and cookies. Sphingomyelins were particularly positively related to cardiorespiratory fitness and coffee intake, and inversely with the intake of whole-grain bread. Acyl-alkyl-phosphatidylcholines were positively linked to cardiorespiratory fitness and coffee intake, and inversely with obesity measures. Most lyso-phosphatidylcholines were positively associated with cardiorespiratory fitness, physical activity and coffee intake, and inversely with the intake of whole-grain bread and cake and cookies. Diacyl-phosphatidylcholines were positively associated with obesity measures and cardiorespiratory fitness, and inversely with intake of whole-grain bread, coffee and cake and cookies. Seven individual metabolites were not connected to any other metabolites with the correlation coefficients directly comparable.

In the next step, we considered different exposures. To reflect dietary behavior, we selected three foods that were previously found to be linked to serum metabolites and chronic disease risk in this population.10,11 This included low intake of whole-grain bread, as an established risk factor, as well as low intakes of coffee and cake and cookies as less-established risk factors. Physical activity was reflected by objectively measured physical activity energy expenditure. To mirror consequent phenotypes, we selected cardiorespiratory fitness and obesity with the single metabolites that we previously obtained.12,13 These were \( \beta \)-coefficient from linear mixed models regression analysis for physical activity and cardiorespiratory fitness, and Spearman’s partial correlation coefficients for diet and obesity measures. Physical activity and cardiorespiratory fitness data were additionally z-transformed to make the \( \beta \)-coefficients directly comparable. All of the association measures were multivariable adjusted for age, sex, alcohol intake, smoking, education and, if applicable, prevalent diseases and the other exposures. Note that the adjustment models differed slightly for different exposures as they were based on our previous studies.10–12 We colored the network based on the direction and strength of association between each exposure and the metabolites, and thus ended up with one metabolite network for each exposure.

To compare the metabolite networks of different exposures, we additionally calculated the correlation of the association measures between the different exposures and 127 serum metabolites, so essentially ‘the correlation of the correlation coefficients’. As an example to obtain the correlation of the metabolite networks of BMI and coffee, we calculated the correlation of the correlation coefficients between BMI and 127 serum metabolites with the correlation coefficients between coffee intake and 127 serum metabolites. As these were normally distributed for all exposures, we calculated Pearson’s correlation coefficients. Therefore, we obtained the correlation of metabolite networks of different exposures.

DISCUSSION

The present study used an innovative approach to generate multiple metabolite networks for different human behaviors and phenotypes, and compared these networks in the setting of a population study. Although the generation of the metabolite network was purely data driven, the metabolites were separated by their different biochemical classes. This was particularly achieved for amino acids, C6-sugar and acylcarnitines. The cluster of different phospholipids could not be completely differentiated, for example, there were still some direct connections between sphingomyelins and phosphatidylcholines. Biosynthesis of sphingomyelins includes the transfer of phosphocholine from phosphatidylcholines to ceramide,31 so there is a biological plausible link of these two groups. However, the direct link of these two different classes could also be a result of the limitations of the metabolomics measurements. The manufacturer of the metabolomics kit reported about possible interferences between individual metabolites.32 This could, for example, have affected hydroxy-sphingomyelin C14:1 and acyl-alkyl-phosphatidylcholine C30:1, two metabolites that were inversely correlated and neighboring in our metabolite network. Nevertheless, it was often the case that metabolites that are one reaction step apart in biological pathways were direct neighbors in our metabolite network. Krumsiak et al.33 previously showed in the Cooperative Health Research in the Region of Augsburg (KORA) cohort that the statistical method Gaussian graphical modeling could discriminate a pathway distance of one for most of these phospholipids with sensitivity and specificity ranging between 0.75 and 1.0. This previous study also showed that a pathway distance of two is indicated by strong negative correlations. This was also observed.
Partial correlation coefficient,
Mutually adjusted

Positive  Negative
---  0.2-0.4  ---
---  0.4-0.6  ---
---  0.6-0.8  ---

Figure 1. Serum metabolite network of the EPIC-Potsdam subcohort (n = 2380). Each node represents one metabolite and each edge between two nodes represents the partial correlation between two metabolites mutually adjusted for the other metabolites. Solid line represents positive correlation and dotted line negative correlation. The thickness of the line indicates the strength of the correlation.
in our study, for example, sphingomyelins C16:1 and C18:0 (desaturation and elongation), and diacyl-phosphatidylcholines C36:5 and C40:5 (two elongations). In summary, the metabolite network of our study population that was drawn with the data-driven method Gaussian graphical modeling could differentiate the metabolite classes and seemed to generally reflect biological plausible pathways.

In our study, we observed several associations between lifestyle factors and phenotypes with serum metabolite networks. As our study is of an observational design, we cannot prove biological mechanisms or demonstrate tissue origins of systemic metabolite changes. Nevertheless, our study may give some first ideas about possible mechanisms and pathways, which are discussed in the following paragraphs, but should be cautiously interpreted and confirmed in experimental studies.

We observed that obesity and cardiorespiratory fitness were more strongly linked to serum metabolite networks compared with diet and physical activity. Our results are in agreement with a
previous study, which reported that low cardiorespiratory fitness and high levels of visceral adipose tissue were strong predictors for impaired cardiometabolic health. This included altered lipoprotein-lipid profiles, deranged glucose-insulin homeostasis and increased inflammatory markers. However, our results have to be interpreted with caution as the different exposures were assessed with different methods. Particularly, diet was not measured objectively but self-reported in a food frequency questionnaire. Thus, it cannot be ruled out that imprecision of the dietary assessment may lead to underestimation of the true effect of diet. In contrast, both cardiorespiratory fitness and physical activity were measured objectively with heart rate and movement sensors and may be better comparable. Therefore, our results indicate that cardiorespiratory fitness may be stronger linked to metabolite networks than physical activity. Physical activity represents a lifestyle, whereas cardiorespiratory fitness can be considered a phenotype, which may not only be influenced by physical activity or inactivity but also by past physical activity.
other environmental and lifestyle factors and genetics. Therefore, cardiorespiratory fitness may reflect long-term exposure and be stronger linked to serum metabolites than physical activity.

On the level of metabolite classes, we observed that amino acids were particularly positively linked to cardiorespiratory fitness and physical activity. Morris et al. found that amino-acid excretion was reduced in participants with high fitness as indicated by lower urinary levels of amino acids. In contrast, in this previous study also plasma levels of many amino acids were lower in the high fitness group. With high energy expenditure, oxidation of branched chain amino acids is generally increased. However, as response to exercise, amino-acid biosynthesis as well as protein breakdown in skeletal muscle may increase. This could possibly increase the systemic pool of amino acids.

In the present study, C6-sugar was particularly positively linked to obesity measures. This may be a result of impaired glucose metabolism and insulin resistance in obese individuals. It has previously been reported that obesity correlated positively with levels of fasting plasma glucose. A recent intervention study could show that people with abdominal obesity were able to improve their levels of fasting plasma glucose with a combined physical activity and dietary intervention. This is in line with our findings that C6-sugar was further inversely correlated with cardiorespiratory fitness, physical activity and whole-grain bread.
intake. In our study population, we previously found that increased levels of C6-sugars were linked to a higher risk of type 2 diabetes.\(^3\) Thus, via this pathway, lifestyle modifications may offer potential for prevention of chronic diseases.

Acylcarnitines showed similar correlations with lifestyle factors and phenotypes as C6-sugar. Adams et al.\(^3\) reported increased levels of acylcarnitines in African Americans with type 2 diabetes, which may be a result of incomplete fatty-acid oxidation and via pro-inflammatory pathways trigger development of insulin resistance. In rats, fed with a 'cafeteria diet' with high-fat and low-fiber content, a serum accumulation of acylcarnitines was observed reflective of systemic mitochondrial dysfunction.\(^4\) This eventually led to tissue inflammation and development of obesity and metabolic syndrome. Thus, increased serum acylcarnitines may be reflective of inflammatory processes and oxidative stress involved in obesity.

Among the phospholipids, sphingomyelins, lyso- and acyl-alkyl-phosphatidylcholines were positively associated with coffee intake, and the latter class was inversely associated with obesity. It was previously observed that levels of acyl-alkyl-phosphatidylcholines were reduced in obese individuals.\(^5\) In addition, these phospholipid classes have been linked to lower risk of type 2 diabetes in this study population, whereas diacyl-phosphatidylcholines were linked to higher risk.\(^6\) In the present study, diacyl-phosphatidylcholines were positively associated with obesity and inversely with coffee intake. All of these phospholipids are primarily synthesized in the liver and secreted as part of blood lipoproteins.\(^7\) In a previous study, we could show that acyl-alkyl-phosphatidylcholines were positively correlated with high-density lipoprotein cholesterol, whereas diacyl-phosphatidylcholines were positively correlated with plasma triglycerides.\(^8\) Coffee intake has been associated with decreased risk of chronic diseases in prospective cohort studies, and particularly with lower risk of type 2 diabetes.\(^9\) However, the biological mechanisms are not yet understood. In the present study, it was additionally observed that the metabolite network of coffee was strongly inversely correlated with the metabolite network of obesity. Thus, our results indicate that the suggested protective effect of coffee could be mediated via counterbalance of pathways of obesity, in particular, on the level of phospholipids that are supplied from hepatic synthesis. Thus, our findings may point toward a crucial role of the liver. These assumptions may be supported by the fact that higher coffee intake has also been associated with lower risk of fibrosis in patients with non-alcoholic fatty liver disease.\(^10\) In addition, in an experimental study, coffee prevented induced liver cirrhosis in rats.\(^11\) Finally, coffee consumption has also been associated with lower risk of liver cancer in prospective cohort studies.\(^12\)

The metabolite network of cake and cookies was positively associated with the metabolite networks of cardiorespiratory fitness and intake of whole-grain bread. This finding was rather unexpected. Nevertheless, it is in line with the observation that cake and cookie intake was inversely associated with risk of chronic diseases in this study population.\(^13\) It is still to debate and see in other studies whether this observation has a biological background or is the result of some type of reporting bias, that is, that individuals with lower levels of cardiorespiratory fitness and low intake of whole-grain bread tend to under-report their cake and cookie intake compared with their counterparts. At this point, we can only conclude that the observation is interesting and that this issue should be investigated further.

The strengths of the present study include that we provided a comprehensive overview of the associations between phenotypes and lifestyle characteristics with serum metabolite networks. We conducted metabolomics analyses in a large population-based cohort measuring 127 metabolites in more than 2300 participants. In addition, we integrated data on objectively measured physical activity and cardiorespiratory fitness, obesity measures and questionnaire-based dietary assessment with metabolomics data using an innovative network approach. Study limitations include that we used a commercial metabolomics kit and could only investigate those metabolites that were included by the manufacturer. Therefore, we might have missed associations of other important metabolites, for example, other amino acids or lipid derivatives. It would be very interesting to study a broader range of metabolites with different targeted or untargeted metabolomics platforms in future studies. Furthermore, our main focus in the discussion was on the level of metabolite classes. Nevertheless, in some conditions, metabolite changes were different for individual metabolites within the same class, which is to be expected, as individual metabolites, for example, different amino acids, may carry very distinct metabolic functions. Another limitation of the study is that the assessment instruments of exposures were different. Therefore, they may be differentially prone to measurement error and, consequently, association measures between different exposures and serum metabolites may not be directly comparable. In addition, we used a cross-sectional design, which illustrates associations but not necessarily temporality and causality. However, knowledge is scarce about the association of these lifestyle factors and phenotypes with metabolite networks in a population setting.

In summary, we could visualize a metabolite network that reflected biological pathways with a data-driven approach based on 2380 EPIC-Potsdam participants. Our results support the concept that different lifestyle factors and phenotypes are reflected in metabolic pathways. Amino-acid metabolism seems to be particularly linked to cardiorespiratory fitness and physical activity, whereas levels of acylcarnitines and C6-sugar were linked to obesity and low intake of whole-grain bread. In addition, the metabolite networks of coffee and obesity were strongly inversely correlated. Therefore, a possible protective effect of coffee on chronic disease risk may be particularly mediated by hepatic phospholipids. Experimental studies need to further validate these findings.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

We specially thank Ellen Kohlsdorf and Wolfgang Bernigau for data management and Martin Floegel for his support with figure formatting. We also thank Julia Scarpa, Arsin Sabunchi and Werner Römisch-Margl for metabolomics measurements performed at the Helmholtz Centrum München, Genome Analysis Center, Metabolomics Core Facility. We thank all the EPIC-Potsdam study participants for their devoted participation in the study. This study was supported by the Federal Ministry of Science, Germany (Grant number 01 EA 9401), the European Union (Grant number 263759), the German Cancer Aid (Grant number 70-2201-Bo2), and by a grant from the Federal Ministry of Education and Research (BMBF) to the German Center for Diabetes Research (DZD eV).

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