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Impact of the Adipokine Adiponectin and the Hepatokine Fetuin-A on the Development of Type 2 Diabetes: Prospective Cohort- and Cross-Sectional Phenotyping Studies

Norbert Stefan1,2,3, Qi Sun4,5, Andreas Fritsche1,2,3, Jürgen Machann1,6, Fritz Schick6, Felicia Gerst1,2,3, Charlotte Jeppesen7,3, Hans-Georg Joost8,3, Frank B. Hu4,5,9, Heiner Boeing10, Susanne Ullrich1,2,3, Hans-Ulrich Häring1,2,3, Matthias B. Schulze7,3

1 Department of Internal Medicine IV, University Hospital Tübingen, Tübingen, Germany, 2 Institute of Diabetes Research and Metabolic Diseases, Member of the German Center for Diabetes Research (DZD), Tübingen, Germany, 3 Deutsches Zentrum für Diabetesforschung (DZD), Neuherberg, München, Germany, 4 Department of Nutrition, Harvard School of Public Health, Boston, Massachusetts, United States of America, 5 Channing Division of Network Medicine, Department of Medicine, Brigham and Women’s Hospital and Harvard Medical School, Boston, Massachusetts, United States of America, 6 Section on Experimental Radiology, University Hospital Tübingen, Tübingen, Germany, 7 Department of Molecular Epidemiology, German Institute of Human Nutrition Potsdam-Rehbruecke, Nuthetal, Germany, 8 Department of Pharmacology, German Institute of Human Nutrition Potsdam-Rehbruecke, Nuthetal, Germany, 9 Department of Epidemiology, Harvard School of Public Health, Boston, Massachusetts, United States of America, 10 Department of Epidemiology, German Institute of Human Nutrition Potsdam-Rehbruecke, Nuthetal, Germany

Abstract

Background: Among adipokines and hepatokines, adiponectin and fetuin-A were consistently found to predict the incidence of type 2 diabetes, both by regulating insulin sensitivity.

Objective: To determine to what extent circulating adiponectin and fetuin-A are independently associated with incident type 2 diabetes in humans, and the major mechanisms involved.

Methods: Relationships with incident diabetes were tested in two cohort studies: within the European Prospective Investigation into Cancer and Nutrition (EPIC)-Potsdam study (628 cases) and the Nurses’ Health Study (NHS; 470 cases). Relationships with body fat compartments, insulin sensitivity and insulin secretion were studied in the Tübingen Lifestyle Intervention Program (TULIP; N = 358).

Results: Circulating adiponectin and fetuin-A, independently of several confounders and of each other, associated with risk of diabetes in EPIC-Potsdam (RR for 1 SD: adiponectin: 0.45 [95% CI 0.37–0.54], fetuin-A: 1.18 [1.05–1.32]) and the NHS (0.51 [0.42–0.62], 1.35 [1.16–1.58]). Obesity measures considerably attenuated the association of adiponectin, but not of fetuin-A. Subjects with low adiponectin and concomitantly high fetuin-A had the highest risk. Whereas both proteins were independently (both p < 1.8 × 10−7) associated with insulin sensitivity, circulating fetuin-A (r = −0.37, p = 0.0004), but not adiponectin, associated with insulin secretion in subjects with impaired glucose tolerance.

Conclusions: We provide novel information that adiponectin and fetuin-A independently of each other associate with the diabetes risk. Furthermore, we suggest that they are involved in the development of type 2 diabetes via different mechanisms, possibly by mediating effects of their source tissues, expanded adipose tissue and nonalcoholic fatty liver.


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* E-mail: norbert.stefan@med.uni-tuebingen.de
Introduction

Among several pathways involved in the pathogenesis of the epidemically spreading disease type 2 diabetes, an altered secretory pattern of the expanded and inflamed adipose tissue is thought to be important for the regulation of insulin sensitivity and subclinical inflammation in various tissues [1]. In this respect adiponectin has gained much attention in the past years because the circulating levels of this adipokine are not only markers of type 2 diabetes risk, but because adiponectin is strongly involved in its progression [2]. In analogy to dysregulated adipose tissue [2–4], there is increasing evidence that nonalcoholic fatty liver disease (NAFLD), which predictive of metabolic diseases [5–10], is also associated with an altered secretory pattern of proteins, which can be referred to as hepatokines, and which are both markers of the disease, and are involved in its pathophysiology [11]. Among them fetuin-A gained much attention during the recent years because of its association with type 2 diabetes and cardiovascular disease risk [12–17], and its important role in the pathogenesis of insulin resistance and subclinical inflammation [18–23].

In the present study we now asked two questions: first, to what extent are circulating levels of these proteins related to incident type 2 diabetes independently of each other? Second, because the circulating levels of these two proteins strongly reflect the dysregulation of their source tissues, adipose tissue and liver, can they be used to estimate the contribution of expanded and inflamed adipose tissue and NAFLD to the pathogenesis of insulin resistance and impaired beta cell function?

For this we investigated associations of circulating adiponectin and fetuin-A with incident type 2 diabetes by applying a head to head comparison of these proteins in two large cohort studies, the European Prospective Investigation into Cancer and Nutrition (EPIC)-Potsdam study and the Nurses’ Health Study (NHS). In addition, we studied the independent relationships of the circulating levels of these proteins with precisely measured body fat mass and distribution, liver fat content, insulin sensitivity and insulin secretion in subjects of the Tubingen Lifestyle Intervention Program (TULIP).

Subjects and Methods

EPIC-Potsdam study

The EPIC-Potsdam Study is part of the multi-centre prospective cohort study EPIC [24]. In Potsdam, Germany, 27,548 subjects (16,644 women and 10,904 men) were recruited from the general population between 1994 and 1998. The age range was 35–65 years in women and 40–65 years in men. The baseline examination included anthropometric measurements, a personal interview including questions about prevalent diseases, and a questionnaire about socio-demographic and lifestyle characteristics [13]. Follow-up questionnaires were sent out every 2 to 3 years to update information about socio-demographic and lifestyle characteristics in 1976 [15]. In 2000–2001, 18,717 NHS participants aged 33–79 years provided blood samples. Among these participants, a prospective, nested, case-control study was conducted to examine plasma biomarkers in relation to type 2 diabetes risk. After excluding women with self-reported prevalent diabetes, cardiovascular disease, or cancer at baseline, 470 cases of type 2 diabetes cases from the date of blood draw through June 2006 were prospectively identified and confirmed. Risk-set sampling was used to randomly select one control for each case from the rest of population who remained free of diabetes when the case was diagnosed; the probability of being selected as a control is proportional to the length of follow-up. Cases and controls were further matched for age at blood draw (61 year), date of blood draw (63 months), fasting status (fast for 8 h or not), and race (white or other races) [15]. The study protocol was approved by the institutional review board of the Brigham and Women’s Hospital and the Human Subjects Committee Review Board of Harvard School of Public Health.

TULIP

A total of 358 Caucasians, who participated in the Tubingen Lifestyle Intervention Program (TULIP) [24], were included in the present analyses because they fulfilled at least one of the following criteria: a family history of type 2 diabetes, a BMI $\geq$ 27 kg/m$^2$, previous diagnosis of impaired glucose tolerance or gestational diabetes. Informed written consent from subjects participating in the Tubingen studies was obtained and the Ethical Committee of the University of Tubingen, Germany had approved the protocols.

Anthropometrics and metabolic parameters were measured as previously described [13,25,26]. Glucose tolerance was determined according to the 1997 World Health Organization diagnostic criteria [27]. Insulin sensitivity from the OGTT was estimated as proposed by Matsuda and DeFronzo [28]. In a subgroup (N = 244) insulin sensitivity was also measured during a euglycemic, hyperinsulimemic clamp [26]. The insulinogenic index, a precise estimate of glucose-induced insulin secretion, was assessed from the OGTT as follows: (insulin at 30 min-insulin at 0 min)/glucose at 30 min-glucose at 0 min).

Measurement of adiponectin and fetuin-A

In the EPIC-Potsdam study and in the TULIP study adiponectin levels were determined with enzyme-linked immunosorbent assays (ELISA, Linco Research, Inc., St Charles, MO). Fetuin-A levels were measured using an immunoturbidimetric method (BioVendor Laboratory Medicine, Modreci, Czech Republic). In the NHS, both, adiponectin and fetuin-A levels were measured by enzyme immunoasays from R&D Systems (Minneapolis, MN).

Statistical analyses

In the EPIC Study and the NHS adiponectin and fetuin-A levels were categorized into quintiles based on subcohort or control participants. Hazard ratios as a measure of relative risk (RR) were computed using a weighted Cox proportional hazards model in EPIC-Potsdam, modified for the case-cohort design according to the Prentice method. Age was the underlying time variable in the counting processes, with entry defined as the subjects’ age at the time of recruitment and exit defined as age at the diagnosis of diabetes, or censoring. In NHS, odds ratios were calculated using
We computed RRs/ORs for each quintile of adiponectin and fetuin-A compared with the lowest quintile. The significance of linear trends across quintiles of adiponectin and fetuin-A was tested by assigning each participant the median value for the quintile and modeling this value as a continuous variable. Because this analysis indicated no departure from linearity, we also considered adiponectin and fetuin-A as continuous variables estimating the RR/OR associated with an increment of 1 SD.

We used information on covariates obtained from the baseline examination in multivariate analyses, namely sex, education, physical activity, smoking, and alcohol intake. Analyses in NHS were further adjusted for other matching factors beyond age (race, fasting status, time of blood drawing), as well as for body mass index (BMI) and waist circumference.

In the TULIP study, data are given as means ± SE. Data that were not normally distributed (e.g., liver fat, insulin sensitivity, body fat distribution; Shapiro-Wilk W test) were logarithmically transformed. A p-value ≤ 0.05 was considered statistically significant.

**Results**

**Association of circulating adiponectin and fetuin-A with diabetes incidence in the EPIC-Potsdam Study and the NHS**

The adjusted RRs/ORs of type 2 diabetes for quintiles of fetuin-A and adiponectin in the EPIC-Potsdam study and the NHS are shown in Table 1. Higher levels of fetuin-A were associated with an increased risk of diabetes, independent of adiponectin [RR/OR comparing extreme quintiles: EPIC-Potsdam: 1.23 [95% CI: 0.88–1.72], p for trend < 0.01; NHS: 2.05 [1.24–3.37]]. Contrary, higher adiponectin levels were associated with lower risk of diabetes in both cohorts (RR/OR comparing extreme quintiles: EPIC-Potsdam: 0.20 [0.13–0.30]; NHS: 0.18 [0.10–0.32]). Considering fetuin-A and adiponectin as continuous variables (per 1 SD increment), revealed a similar picture with consistent inverse associations for adiponectin and positive trends of higher fetuin-A with diabetes incidence.
associations for fetuin-A (Table 2). Adjustment for BMI and waist circumference attenuated the associations of adiponectin (by 29% in EPIC-Potsdam and 28% in NHS). However, the association of fetuin-A with diabetes risk was largely unaffected by adjustment for BMI and waist circumference (Table 2).

We then examined the joint effect of circulating adiponectin and fetuin-A by cross classifying participants by both variables using sex-specific medians as cut-offs. The RR/OR for the combination of a high fetuin-A- and a low adiponectin level compared with the opposite extreme was 3.54 (95% CI: 2.54–4.95) in EPIC-Potsdam and 4.61 (2.87–7.38) in NHS (Figure 1, A/B). Inclusion of BMI and waist circumference in the models attenuated the associations of circulating adiponec- 

**Table 2. Relative Risk of type 2 diabetes for plasma adiponectin and fetuin-A and attenuation of the risk by BMI and waist circumference.**

<table>
<thead>
<tr>
<th></th>
<th>Adiponectin</th>
<th>Fetuin-A</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>EPIC-Potsdam</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Model 1 *</td>
<td>0.35 Ref</td>
<td>1.21 Ref</td>
</tr>
<tr>
<td>(0.29, 0.41)</td>
<td>(1.09, 1.34)</td>
<td></td>
</tr>
<tr>
<td>Model 1 + BMI</td>
<td>0.43 +22.9</td>
<td>1.19 −1.7</td>
</tr>
<tr>
<td>(0.36, 0.52)</td>
<td>(1.06, 1.33)</td>
<td></td>
</tr>
<tr>
<td>Model 1 + BMI + waist circumference</td>
<td>0.45 +28.6</td>
<td>1.18 −2.5</td>
</tr>
<tr>
<td>(0.37, 0.54)</td>
<td>(1.05, 1.32)</td>
<td></td>
</tr>
<tr>
<td><strong>Nurses’ Health Study</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Model 1 †</td>
<td>0.40 Ref</td>
<td>1.32 Ref</td>
</tr>
<tr>
<td>(0.33–0.48)</td>
<td>(1.14–1.53)</td>
<td></td>
</tr>
<tr>
<td>Model 1 + BMI</td>
<td>0.47 +17.5</td>
<td>1.38 +4.5</td>
</tr>
<tr>
<td>(0.39–0.57)</td>
<td>(1.18–1.61)</td>
<td></td>
</tr>
<tr>
<td>Model 1 + BMI + waist circumference</td>
<td>0.51 +27.5</td>
<td>1.35 +2.3</td>
</tr>
<tr>
<td>(0.42–0.62)</td>
<td>(1.16–1.58)</td>
<td></td>
</tr>
</tbody>
</table>

*Relative Risk adjusted for age, sex, education (in or no training, vocational training, technical school, or technical college or university degree), occupational activity (light, moderate, heavy), sport activity (0, 0.1–4.0, >4.0 h/week), cycling (0, 0.1–2.4, 2.5–4.9, ≥5.0 h/week), smoking (never, past, current <20 cigarettes/d, current ≥20 cigarettes/d), and alcohol intake (0, 0.1–5.0, 5.1–10.0, 10.1–20.0, 20.1–40.0, >40.0 g/d).

† Odds Ratio adjusted for matching factors, including age at blood draw (yrs), race (white or not), fasting status (yes, no), and time of blood drawing, as well as smoking status (current smoker, past smoker, non-smoker), physical activity (in textiles), alcohol use (abstainer, <5.0 g/day, 5.0–14.9 g/day, ≥15.0 g/day), and education (registered nurse, bachelor, master and higher).

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Cross-sectional relationships in the TULIP

The 358 TULIP subjects had a mean age of 46 years and a mean BMI of 30 kgm⁻². To investigate relationships of circulating adiponectin and fetuin-A levels with metabolic traits we performed analyses in the total population and in subjects with normal glucose tolerance (NGT) and impaired glucose tolerance (IGT) separately (Table 1 in File S1).

**Relationships of circulating adiponectin and fetuin-A with body fat content and distribution and with insulin sensitivity.** In the total population circulating adiponectin, adjusted for age and gender, correlated negatively with BMI (r = -0.19, p = 0.0004) and with waist circumference (r = -0.26, p<0.0001). Negative correlations were also found with total body fat mass (r = -0.15, p = 0.008) and, more strongly, with visceral fat mass (r = -0.40, p<0.0001) and with liver fat content (r = -0.28, p<0.0001). Circulating fetuin-A correlated only very weakly with BMI (r = 0.11, p = 0.04) and not statistically significant with waist circumference (r = 0.10, p = 0.06), total body fat mass- (r = 0.07, p = 0.25), and visceral fat (r = 0.07, p = 0.20) mass. However, a positive correlation with liver fat content was found (r = 0.12, p = 0.04). In multivariate models including age and sex, both circulating adiponectin and fetuin-A were strongly and independently associated with insulin sensitivity estimated from the OGTT and measured during the clamp (Table 3, models 1). Inclusion of BMI and waist circumference in the models attenuated the associations of circulating adiponectin (change of the β; OGTT: −37%, clamp: −31%) on insulin sensitivity, but less so of circulating fetuin-A (OGTT: −17%, clamp: −18%) (Table 3, models 2). Similar relationships were found when these analyses were performed in subjects with and without NAFLD (OGTT: total N = 291, NAFLD = 93; clamp: total N = 203, NAFLD = 63). Here inclusion of BMI and waist circumference in the models attenuated the associations of circulating adiponectin with insulin sensitivity, particularly in subjects without NAFLD (change of the β; OGTT: −52%, clamp: −34%), while this association was only slightly attenuated or even became stronger in the smaller group of subjects with NAFLD (OGTT: −7%, clamp: +10%). The respective relationships of fetuin-A with insulin sensitivity were less strongly affected, both, in subjects without NAFLD (OGTT: −12%, clamp: −13%) and with (OGTT: −6%, clamp: −4%) NAFLD. After additional inclusion of liver fat content in the models 1, the associations of adiponectin and fetuin-A with insulin sensitivity were further attenuated and adiponectin, fetuin-A and liver fat content independently determined insulin sensitivity (Table 3, models 3). When we then divided non-obese subjects (N = 207) by the median insulin sensitivity estimated by the OGTT (13.78 arb.u.) we found circulating fetuin-A (OR for 1 SD: 0.37, 95% CI: 0.29–0.41, p = 0.0004) and with waist circumference (r = 0.19, p = 0.0004) and with liver fat content (r = 0.12, p=0.0001). Negative correlations were also found with total body fat mass (r = 0.07, p = 0.20) mass. However, a positive correlation with liver fat content was found (r = 0.12, p = 0.04). In multivariate models including age and sex, both circulating adiponectin and fetuin-A were strongly and independently associated with insulin sensitivity estimated from the OGTT and measured during the clamp (Table 3, models 1). Inclusion of BMI and waist circumference in the models attenuated the associations of circulating adiponectin (change of the β; OGTT: −37%, clamp: −31%) on insulin sensitivity, but less so of circulating fetuin-A (OGTT: −17%, clamp: −18%) (Table 3, models 2). Similar relationships were found when these analyses were performed in subjects with and without NAFLD (OGTT: total N = 291, NAFLD = 93; clamp: total N = 203, NAFLD = 63). Here inclusion of BMI and waist circumference in the models attenuated the associations of circulating adiponectin with insulin sensitivity, particularly in subjects without NAFLD (change of the β; OGTT: −52%, clamp: −34%), while this association was only slightly attenuated or even became stronger in the smaller group of subjects with NAFLD (OGTT: −7%, clamp: +10%). The respective relationships of fetuin-A with insulin sensitivity were less strongly affected, both, in subjects without NAFLD (OGTT: −12%, clamp: −13%) and with (OGTT: −6%, clamp: −4%) NAFLD. After additional inclusion of liver fat content in the models 1, the associations of adiponectin and fetuin-A with insulin sensitivity were further attenuated and adiponectin, fetuin-A and liver fat content independently determined insulin sensitivity (Table 3, models 3). When we then divided non-obese subjects (N = 207) by the median insulin sensitivity estimated by the OGTT (13.78 arb.u.) we found circulating fetuin-A (OR for 1 SD:
1.42 [95% CI 1.09–1.85]) but not circulating adiponectin (0.88 [0.77–1.01]) or hs-CRP levels (1.04 [0.98–1.11]) to predict the insulin resistant state, independently of age, sex, BMI and waist circumference. Similar relationships were found when NGT and IGT were analyzed separately (data not shown).

When we divided participants by the medians of circulating adiponectin and fetuin-A levels, individuals with low adiponectin and high fetuin-A levels had the lowest insulin sensitivity compared to the other groups (Figure 1, C/D).

Relationships of circulating adiponectin and fetuin-A with insulin secretion. We next investigated whether circulating fetuin-A may be associated with glucose-induced insulin secretion in humans. In the total population circulating fetuin-A did not correlate with the insulinogenic index ($r = 0.01$, $p = 0.99$) when adjusted for age, gender and insulin sensitivity measured during the OGTT. However, this association depended on glucose tolerance status: while fetuin-A did not correlate with the insulinogenic index in subjects with NGT, a strong negative correlation between fetuin-A levels and the insulinogenic index was found in subjects with IGT ($r = -0.37$, $p = 0.0004$) (p for interaction = 0.024) (Figure 2, A/B). No significant relationships were found for circulating adiponectin with the adjusted insulinogenic index (all $p > 0.055$).

Discussion

During the last decade much effort has been made to identify important pathways involved in the natural history of type 2 diabetes. Thereby, several candidates were described, predominantly based on animal and on in-vitro studies [29–31]. However, often it was not possible to prove these pathways to be of high relevance for human metabolism. In human studies, on the other hand, several blood, genetic or phenotypic markers were found to predict incident type 2 diabetes [1–4]. Nevertheless, no precise mechanisms of action for several of these parameters are known and/or their predictive effect on the development of type 2 diabetes was either small or absent, which so far limits their potential in the prevention and the treatment of the disease.

Because these limitations largely do not apply to the adipokine adiponectin and the hepatokine fetuin-A, we here investigated to what extent circulating adiponectin and fetuin-A determine incident type 2 diabetes, independently of each other. Towards

Figure 1. Relative risk (in the European Prospective Investigation into Cancer and Nutrition (EPIC)-Potsdam Study; Panel A) and Odds Ratio (in the Nurses’ Health Study; Panel B) of type 2 diabetes for joint classifications of plasma adiponectin and fetuin-A. Groups with high/low adiponectin and fetuin-A levels were defined based on sex-specific medians. RRs were adjusted for age, sex, education, occupational activity, sport activity, cycling, smoking, alcohol intake, BMI, and waist circumference in EPIC-Potsdam and ORs for matching factors, including age at blood draw, race, fasting status, and time of blood drawing, as well as smoking status, physical activity, alcohol use, and education in the Nurses’ Health Study. Relationship of circulating adiponectin and fetuin-A with insulin sensitivity estimated from the oral glucose tolerance test (OGTT; N = 358, C) and measured during the euglycemic, hyperinsulinemic clamp (N = 244, D). Subjects were divided by the medians of circulating adiponectin and fetuin-A in groups with high and low levels. $p$ for trend after adjustment for age and sex.

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sensitivity may be a result of inflamed NAFLD [11,32–35].

in circulating fetuin-A and the resulting decrease in insulin

information from our and other studies suggest that the increase

can considerably depend on body fatness. Rather the available

data support that the adiponectin levels confer at least in part the effect of obesity on the type 2 diabetes risk. In contrast,

obesity, our data support that the adiponectin levels confer at least

attenuated after accounting for estimates of overall and visceral

adiponectinemia, but not of circulating fetuin-A, was considerably

was considerably attenuated after accounting for estimates of overall and visceral

adiponectinemia, but not of circulating fetuin-A, was considerably

Table 3. Determinants of insulin sensitivity in multivariate regression models in TULIP.

<table>
<thead>
<tr>
<th>Covariates</th>
<th>Insulin sensitivity_{OGTT}</th>
<th>Insulin sensitivity_{Clamp}</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate±SE</td>
<td>F-value</td>
</tr>
<tr>
<td>Models 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female sex</td>
<td>0.04±0.03</td>
<td>1.3</td>
</tr>
<tr>
<td>Age</td>
<td>−0.29±0.11</td>
<td>6.8</td>
</tr>
<tr>
<td>Adiponectin levels</td>
<td>0.38±0.07</td>
<td>30.4</td>
</tr>
<tr>
<td>Fetuin-A levels</td>
<td>−0.85±0.16</td>
<td>28.4</td>
</tr>
<tr>
<td>Models 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female sex</td>
<td>−0.01±0.04</td>
<td>0.24</td>
</tr>
<tr>
<td>Age</td>
<td>−0.17±0.10</td>
<td>2.6</td>
</tr>
<tr>
<td>Adiponectin levels</td>
<td>0.24±0.06</td>
<td>14.2</td>
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<tr>
<td>Fetuin-A levels</td>
<td>−0.70±0.14</td>
<td>23.8</td>
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<tr>
<td>BMI</td>
<td>−0.54±0.31</td>
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</tr>
<tr>
<td>Waist circumference</td>
<td>−1.16±0.43</td>
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</tr>
<tr>
<td>Models 3 *</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female sex</td>
<td>−0.004±0.04</td>
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</tr>
<tr>
<td>Age</td>
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<tr>
<td>Fetuin-A levels</td>
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<tr>
<td>Waist circumference</td>
<td>0.13±0.43</td>
<td>0.07</td>
</tr>
<tr>
<td>Liver fat content</td>
<td>−0.20±0.03</td>
<td>52.0</td>
</tr>
</tbody>
</table>

*N = 291 for insulin sensitivity_{OGTT} and N = 203 for insulin sensitivity_{Clamp}.

doi:10.1371/journal.pone.0092238.t003

this aim, we first chose an epidemiological approach and investigated the associations of circulating adiponectin and fetuin-A with incident type 2 diabetes by applying a head to head comparison of these proteins in two large cohort studies, the EPIC-Potsdam study and the NHS. In both studies we found that circulating adiponectin and fetuin-A were associated with risk of incident diabetes, independently of several confounders, and of each other. The consistency of the association suggests that it might be generalizable to healthy populations, at least to those with Caucasian origin. Because the strength of association of adiponectinemia, but not of circulating fetuin-A, was considerably attenuated after accounting for estimates of overall and visceral obesity, our data support that the adiponectin levels confer at least in part the effect of obesity on the type 2 diabetes risk. In contrast, the association of fetuin-A with diabetes risk does not appear to considerably depend on body fatness. Rather the available information from our and other studies suggest that the increase in circulating fetuin-A and the resulting decrease in insulin sensitivity may be a result of inflamed NAFLD [11,32–35].

We then focused on the relationship of both circulating proteins with anthropometrics and metabolic traits in precisely phenotype subjects of TULIP. We confirmed the strong correlations of adiponectinemia with measures of body fat mass and distribution in these subjects as well as the absence of such relationships for fetuin-A levels. Based on the known properties of adiponectin and fetuin-A to regulate insulin sensitivity, we confirmed that the circulating levels of these proteins were independently of each other associated with insulin sensitivity, estimated from the OGTT or measured by a euglycemic, hyperinsulinemic clamp. In agreement with the findings from the EPIC-Potsdam study and the NHS, the relationship of circulating adiponectin, but not of fetuin-A, was considerably attenuated after accounting for measurements of body fat content and distribution. Consequently we asked the question whether circulating fetuin-A may be a better predictor of insulin sensitivity than circulating adiponectin in subjects who are non-obese and could confirm this hypothesis in our study.

Having found strong independent associations of circulating adiponectin and fetuin-A, the two proteins that regulate insulin sensitivity, on the diabetes risk, we then asked whether they may differentially impact on insulin secretion, and thereby have distinct effects in the pathogenesis of type 2 diabetes. For adiponectin we have previously shown that this protein does not influence glucose-induced insulin secretion in humans [36]. In the present study we could show that fetuin-A levels are not associated with insulin secretion in our subjects. Based on the knowledge that subjects with IGT have an impaired beta cell function [37,38], we then tested the hypothesis that fetuin-A is particularly relevant specifically in this population that is at very high risk for the disease. Indeed, when we separated the individuals in those with NGT and IGT, a strong negative relationship of fetuin-A with disease. Indeed, when we separated the individuals in those with NGT and IGT, a strong negative relationship of fetuin-A with IGT have an impaired beta cell function [37,38], we then asked whether circulating fetuin-A may be a better predictor of insulin sensitivity than circulating adiponectin in subjects who are non-obese and could confirm this hypothesis in our study.

What is the relevance of our data for clinicians and researchers? Because the relative risk of incident diabetes was much higher for the combination of a high fetuin-A- and a low adiponectin level, than for the single circulating level of each protein, it may be
important for clinicians to measure both proteins when it comes to
the prediction of the risk of future type 2 diabetes. Whether fetuin-
A and adiponectin improve prediction of diabetes risk beyond
waist circumference and other classical risk factors remains,
however, uncertain. Furthermore, fetuin-A may become an
important determinant of insulin resistant states, particularly in
non-obese subjects where adiponectin and hs-CRP levels lost their
strong predictive power in our study.

For researchers we provide novel information that adiponectin
and fetuin-A are independently involved in the pathogenesis of
type 2 diabetes. Both proteins impact on the development of the
disease predominantly be the regulation of subclinical inflamma-
tion. Furthermore, we have support for the hypothesis that they
mediate the effects of their source tissues, expanded adipose tissue
and inflamed nonalcoholic fatty liver on glucose metabolism and
cardiovascular disease. In addition, we provide explorative
information about a putatively newly identified cross-talk of the
liver with the endocrine function of the pancreas. Whether there
are direct effects of fetuin-A on signalling cascades in beta cells or
whether fetuin-A induces a chronic pro-inflammatory process in
the human islets needs to be investigated in future studies.

Some possible limitations of our findings have to be considered.
The potential of residual confounding applies to our study as it
does to observational studies in general. We adjusted for a large
variety of known risk factors. Although fetuin-A and adiponectin
remained significantly associated with diabetes risk, we cannot rule
out that other unmeasured factors or that imprecision in the
measurement of covariates explain this observation. Also, we only
had a single blood drawing which might have introduced random
measurement errors in determining fetuin-A and adiponectin. The
lack of repeated measurements may have led to an understimation
of the observed associations. In conclusion, our findings support that
the adipokine adiponectin and the hepatokine fetuin-A are reliable
predictors of incident type 2 diabetes and insulin resistance, and that
they are strongly and independently of each other involved in their
pathogenesis. Moreover, we could provide indirect support that
adiponectin mediates, at least in part, the impact of dysregulated
and expanded visceral fat on metabolism, whereas increased fetuin-
A levels are largely a result of NAFLD. Finally, we provided novel
exploratory information that fetuin-A may play a role in the
pathogenesis of type 2 diabetes by affecting insulin secretion.

Supporting Information

Table S1 Characteristics of the TULIP study partici-
pants. (DOC)

Author Contributions

Conceived and designed the experiments: NS QS AF MBS FS FBH HB
HGJ HUH. Performed the experiments: NS QS AF JM CJ FG SU MBS.
Analyzed the data: NS QS AF JM CJ MBS. Wrote the paper: NS QS.

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