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Association of serum orosomucoid with 30-min plasma glucose and glucose excursion during oral glucose tolerance tests in non-obese young Japanese women

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ABSTRACT

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Dr Tsutomu Kazumi; kazumi@mukogawa-u.ac.jp **Objective** Inflammatory markers are elevated in insulin resistance (IR) and diabetes. We tested whether serum orosomucoid (ORM) is associated with postload glucose, β -cell dysfunction and IR inferred from plasma insulin kinetics during a 75 g oral glucose tolerance test (OGTT). **Research design and methods** 75 g OGTTs were performed with multiple postload glucose and insulin measurements over a 30–120 min period in 168 nonobese Japanese women (aged 18–24 years). OGTT responses, serum adiponectin and high-sensitivity C reactive protein (hsCRP) were cross-sectionally analyzed by analysis of variance and then Bonferroni's multiple comparison procedure. Stepwise multivariate linear regression analyses were used to identify most important determinants of ORM.

Results Of 168 women, 161 had normal glucose tolerance. Postload glucose levels and the area under the glucose curve (AUCg) increased in a stepwise fashion from the first through the third ORM tertile. In contrast, there was no or modest, if any, association with fat mass index, trunk/leg fat ratio, adiponectin, hsCRP, postload insulinemia, the Matsuda index and homeostasis model assessment IR. In multivariable models, which incorporated the insulinogenic index, the Matsuda index and HOMA-IR, 30 min glucose (standardized β : 0.517) and AUCg (standardized β : 0.495) explained 92.8% of ORM variations.

Conclusions Elevated circulating orosomucoid was associated with elevated 30 min glucose and glucose excursion in non-obese young Japanese women independently of adiposity, IR, insulin secretion, adiponectin and other investigated markers of inflammation. Although further research is needed, these results may suggest a clue to identify novel pathways that may have utility in monitoring dysglycemia within normal glucose tolerance.

INTRODUCTION

Subjects with impaired glucose tolerance (IGT) or impaired fasting glucose (IFG) have an increased risk for type 2 diabetes.^{1–3}

Significance of this study

What is already known about this subject?

Postload glucose and glucose excursion during an oral glucose tolerance test are related to insulin resistance and/or impaired insulin secretion, both of which are associated with obesity.

What are the new findings?

- Postload glucose and glucose excursion during an oral glucose tolerance test are related to serum orosomucoid in non-obese young Japanese women.
- The relationships are independent of fat mass and distribution, markers of insulin resistance/ sensitivity, insulin secretion, adiponectin and other investigated markers of inflammation.

How might these results change the focus of research or clinical practice?

- Since insulin resistance and impaired insulin secretion are known to be causative in the development of diabetes, this might suggest that orosomucoid could be related to the glucose levels.
- These results may suggest a clue to identify novel pathways that may have utility in monitoring glucose dysmetabolism within normal glucose tolerance, although further research is needed.

Although more than half of the subjects who develop type 2 diabetes have IGT or IFG at baseline,⁴ prospective epidemiological studies have demonstrated that about 40% of subjects have normal glucose tolerance (NGT) at baseline.^{1 3 5} Therefore, other than fasting and 2-hour postload glucose, additional information might help us identify a group of subjects who might benefit from early lifestyle intervention.

Inflammation is recognized as a major etiological determinant of multiple disease states including cardiovascular disease, type 2 diabetes and metabolic syndrome.⁶ Several prospective studies have shown that high baseline levels of C reactive protein (CRP) are associated with the increased risk for type 2 diabetes.^{7 8} Among inflammatory markers, orosomucoid (ORM) has been less studied in relation to metabolic disorders, although the acute phase reactant protein is one of the most abundant plasma proteins, accounting for about 1% of all plasma proteins.⁹ In addition, most of the studies described above were conducted in Western countries where the prevalence of obesity is high. Recently, Qin et al¹⁰ have demonstrated that ORM activates AMP-activated protein kinase (AMPK) in skeletal muscle, which led to the activation of glycogen synthase and subsequent glycogen storage. As AMPK is important for glucose uptake and glycogen storage in the myocytes¹¹ and skeletal muscle is responsible for the majority of glucose disposal,¹² we aimed to examine whether ORM is associated with postchallenge glycemia in non-obese Japanese women and whether these associations, if any, depend on fat mass and distribution, insulin resistance (IR) and insulin secretion.

METHODS

We cross-sectionally studied 168 young women (50 collegiate athletes and 118 untrained students of the Mukogawa Women's University), aged 18-24 years whose details have been reported elsewhere.^{13 14} Athletes were students of the Department of Health and Sports Sciences and were also members of volleyball club, basketball club or track club (middle-distance runners) of the University. They had been training regularly for 2 years or longer prior to the study, 5 hours a day and 6 days a week and participated regularly in competitive events in their respective sport specialties. Non-athletes were students of the Department of Food Sciences and Nutrition. Although they were not engaged in any regular sport activity, they had 9367±1971steps/day (mean±SD of 77 non-athletes, who used a pedometer for 14 days and mean steps a day were calculated in each participants). This may explain in part the result that the difference in insulin sensitivity was small between athletes and non-athletes as previously reported.¹³ Subjects with clinically diagnosed acute or chronic inflammatory diseases, endocrine, cardiovascular, hepatic, renal diseases, hormonal contraception and unusual dietary habits were excluded. Nobody reported to receive any medications or have regular supplements. The study was in accordance with the Helsinki Declaration. All subjects were recruited as volunteers and gave written consent after the experimental procedure had been explained.

After a 12 hours overnight fast, a standard 75 g OGTT with glucose and insulin measurements at 0 (fasting), 30 min, 1 hour and 2 hours was performed in the morning. Plasma glucose was determined by the hexokinase/glucose-6-phosphate dehydrogenase method (interassay coefficient of variation (CV) <2%). Serum insulin was

measured by an ELISA method with a narrow specificity excluding des-31, des-32 and intact proinsulin (interassay CV <6%). Area under the response curve of plasma glucose (AUCg) and serum insulin (AUCi) was calculated by the trapezoidal method. Homeostasis model assessment-IR (HOMA-IR), the Matsuda index and the insulinogenic index (IGI) were calculated as previously reported.^{15–17}

Leptin, adiponectin, high-sensitivity CRP (hsCRP) and plasminogen activator inhibitor-1 (PAI-1) were measured in fasted samples as previously reported.^{13 14} ORM concentrations were measured in fasted samples by an immunoturbidimetric method using a commercially available kit (N Antiserum to Human α 1-acid Glycoprotein, Siemens Healthcare Diagnostics, Tokyo, Japan) and an autoanalyzer (JCA-BM6010, JEOL, Tokyo, Japan). Intra-assay and interassay CV at 87 mg/dL was 1.4% and 1.7%, respectively.

Fat mass, bone mass and lean mass for arms, legs, trunk and the total body were measured using whole-body dual-energy X-ray absorptiometry (Hologic QDR-2000, software V.7.20D, Waltham, Massachusetts, USA) as previously reported.¹³ ¹⁴ The leg region included the entire hip, thigh and leg. General adiposity was assessed using height-adjusted and weight-adjusted body fat. The former, fat mass index (FMI), was calculated as body fat mass in kg divided by height in meter squared and the latter, per cent body fat, was calculated as body fat mass in kg divided by body weight in kg×100. Abdominal fat accumulation was assessed by the ratio of trunk fat-to-leg fat.¹⁸

Data were presented as mean±SD unless otherwise stated. Due to deviation from normal distribution, IGI and hsCRP were logarithmically transformed for analyses. Cross-sectional differences were tested using analysis of variance among ORM tertile and then Bonferroni's multiple comparison procedure. Associations of ORM were analyzed by Pearson's correlation. In order to identify most important determinants of ORM, stepwise multivariate linear regression analyses were performed. Independent variables included were all variables that showed significant difference among ORM tertile. Comparisons of ORM levels between athletes and non-athletes were made with t-test. A two-tailed value of p<0.05 was considered significant. Statistics were performed with SPSS system V.17.0 (SPSS, Chicago, Illinois, USA).

RESULTS

Of 168 women studied, 2 athletes and 1 non-athlete had IFG, 4 non-athletes had IGT and 161 had NGT. Because serum ORM levels did not differ between athletes and non-athletes (126 ± 28 vs 127 ± 26 mg/dL, p=0.87), both groups were combined to be analyzed to obtain wider range of adiposity. Table 1 displays characteristics of women grouped by ORM tertile (<112, 113–134, \geq 135 µg/mL). As previously reported, ^{13 14} young women were normal weight, had normal waist circumference

Table 1 Features of 168 young Japanese women grouped by tertile of serum orosomucoid concentrations					
Tertile	First	Second	Third		
Range (mg/dL)	<112	113–134	≥135		
Orosomucoid (mg/dL)	100±10	125±6	155±20	abc	
Age (years)	20.3±1.1	20.3±1.0	20.6±1.1		
Fat mass index (kg/m ²)	5.6±1.6	5.3±1.8	5.2±1.8		
Waist circumference (cm)	73.3±5.5	73.5±5.5	72.9±5.9		
Percentage body fat (%)	26.6±6.1	25.5±6.1	25.2±6.3		
Trunk/leg fat ratio	1.24±0.24	1.24±0.25	1.21±0.24		
Leptin (ng/mL)	7.7±3.5	7.3±3.5	7.6±4.2		
Adiponectin (µg/mL)	13.4±5.1	12.2±4.5	11.6±4.1		
hsCRP (µg/dL)	44±97	23±33	32±48		
PAI-1 (ng/mL)	24±9	29±14	26±13		
Fasting glucose (mg/dL)	82±6	84±6	86±7	b	
30 min glucose (mg/dL)	97±13	123±8	148±20	abc	
1 hour glucose (mg/dL)	79±17	100±18	129±35	abc	
2 hours glucose (mg/dL)	80±12	91±17	104±24	abc	
Fasting insulin (µU/mL)	5.8±3.5	5.7±2.3	5.9±3.4		
30 min insulin (µU/mL)	56±35	53±25	52±33		
1 hour insulin (µU/mL)	32±17	43±20	52±31	ab	
2 hours insulin (μU/mL)	32±25	35±20	42±25		
∆glucoe30 (mg/dL)	15±14	40±8	62±18	abc	
∆insulin30 (µU/mL)	50±35	47±24	46±32		
AUCg (mg/dL, 2 hours)	168±18	203±19	245±38	abc	
AUCi (µU/mL, 2 hours)	70±32	78±29	87±43	b	
HOMA-IR	1.17±0.69	1.19±0.51	1.27±0.80		
Matsuda index	10.8±4.2	9.1±3.6	8.8±4.7	b	
IGI	10.2±20.9	1.2±0.6	1.2±3.4	ab	
Athletes (n, %)	17 30.4	18, 32.1	15, 26.8		

Mean \pm SD (n=56) in each tertile. Significantly different at p<0.05 or less.

a and b: first vs second and third tertile, respectively, c: second vs third tertile.

AUCg, area under the concentration curve of glucose; AUCi, area under the concentration curve of insulin; FMI, fat mass index; Δ glucose30 and Δ insulin30, incremental 30 min glucose and insulin concentrations, respectively; HOMA-IR, homeostasis model assessment of insulin resistance; hsCRP, high-sensitivity C reactive protein; IGI, insulinogenic index; PAI-1, plasminogen activator inhibitor-1.

and hence were insulin sensitive as indicated by normal fasting insulin levels.

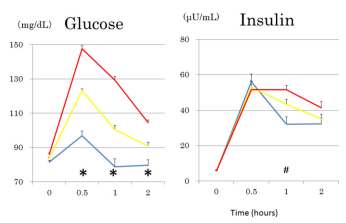
ORM levels were not associated with FMI, the percentage of body fat, trunk/leg fat ratio and serum leptin (table 1). In addition, ORM was not associated with serum hsCRP and PAI-1. Furthermore, no difference was found in serum adiponectin among three groups as well. In contrast, as ORM increased from the first through the third tertile (figure 1), postload glucose levels at 3 points of time increased in a stepwise fashion in both non-athletes and athletes, whereas postload insulin levels did not differ except for 1 hour insulin (table 1). AUCg and an incremental 30 min glucose concentration increased in a stepwise fashion, whereas an incremental 30 min insulin concentration did not differ (table 1). Women with the third (higher ORM) as compared with the first ORM tertile had lower IGI and the Matsuda index, and higher

fasting glucose, 1 hour insulin and AUCi. HOMA-IR and the percentage of athletes did not differ.

Multivariate stepwise linear regression analyses for ORM as a dependent variable was done, which included postload glucose values at 3 points of time, AUCg, IGI (log transformed), the Matsuda index and HOMA-IR as independent variables (table 2). AUCg and 30min glucose were independent determinants of ORM and explained 92.8% of variations of ORM. Subgroup analyses in athletes and non-athletes revealed similar results (table 2). Figure 2 shows the scatter plot between serum ORM and 30min glucose and AUCg.

DISCUSSION

The present study has demonstrated that increased ORM was associated with increased glucose excursion



Mean±SEs of glucose curves during 75 g oral Figure 1 glucose testing in young Japanese women divided into three groups according to tertile of serum orosomucoid: low, blue; median, yellow; high, red. *Under blue lines; glucose values at this time points are significantly different from each other among three tertile groups at p<0.05 or less. #Insulin values at 1 hour were significantly lower in the low as compared with the median and the high orosomucoid tertile. n=56 in each tertile. Range of orosomucoid (in mg/dL); low: <112, median: 113–134, high: ≥135.

(AUCg) and 30 min postload glucose in non-obese, young Japanese women. These associations were not related to sophisticated measures of general and abdominal adiposity (FMI, the percentage of body fat, trunk/ leg fat ratio) and serum leptin. Associations observed were not related to insulin secretion, insulin sensitivity/ resistance, adiponectin and other investigated markers of inflammation. Glucose excursion (AUCg) and 30 min postload glucose explained >90% of variations of ORM. The present results in young women confirmed positive associations of ORM with fasting and 2-hour glucose as previously reported in general population, and in overweight and obese people.¹⁹⁻²¹

Metabolomics now seeks to identify early biomarkers which are predictive of the development of dysglycemia and type 2 diabetes. A Finnish metabolic profiling study²² demonstrated that branched-chain and aromatic amino acids and orosomucoid referred to as al-acid glycoprotein, were predictors of both fasting and 2-hour glucose

 Table 2
 Multivariate stepwise linear regression analysis for
 orosomucoid as a dependent variable

glucose

AUCg

Non- athletes	Standardized β	Cumulative R ²	P values
AUCg	0.565	0.837	2.49E-22
30 min glucose	0.440	0.907	4.86E-16
Athletes			
30 min glucose	0.781	0.969	1.86E-20
AUCg	0.225	0.978	3.56E-05
Total			
30 min	0.517	0.844	3.39E-31

0.928

Independent variables included are the same in all models:

0.495

postload glucose levels at 3 points of time, AUCg, log IGI, HOMA-IR and Matsuda index.

AUCg, area under the concentration curve of glucose; HOMA-IR, homeostasis model assessment of insulin resistance; IGI, insulinogenic index.

at 6.5-year follow-up in a general population setting of middle-aged men and women.

Since the prevailing insulin concentrations over the first 30 min of the OGTT, a direct estimate of insulin secretion during the OGTT, were not associated with ORM, insufficient insulin secretion was possibly not the primary factor. Studies from Asia have reported an independent association between elevated 30 min glucose and incident diabetes.^{23 24} However, this was associated with decreased β -cell function in those studies.^{23 24} An incremental 30 min glucose concentration may reflect the ability of endogenous insulin secretion to suppress hepatic glucose production.²⁵ It is suggested that hepatic IR during early phase insulin secretion may be related to early phase hyperglycemia.²⁵ Although there was no association between ORM and HOMA-IR, a more reflective surrogate of hepatic IR,¹⁶ the possibility cannot be ruled out that hepatic IR may be involved in ORM-early phase

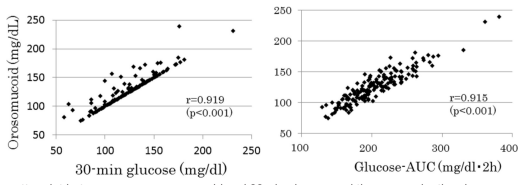


Figure 2 The scatter plot between serum orosomucoid and 30 min glucose and the area under the glucose curve (glucose-AUC) during a 75 g oral glucose tolerance test.

1.62E-29

hyperglycemia association because majority of circulating ORM is produced in the liver.²¹

In above-mentioned studies,^{19–22} serum ORM concentrations were considered a marker of low-grade chronic inflammation, which links obesity to IR and β -cell failure resulting in dysglycemia.²⁶ However, in the present study, there was no association of ORM with FMI, trunk/leg fat ratio, hsCRP and adiponectin, the last of which is known to stimulate glucose uptake through AMPK or peroxisome proliferator-activated receptor α activation.²⁷ Since IR and β -cell failure are known to be causative in the development of diabetes, this might suggest that ORM could be related to the glucose levels independently of the muscle insulin signaling: insulin-independent glucose uptake via skeletal muscle.

It is reported that contraction-stimulated, insulin-independent glucose transport in skeletal muscle may be mediated by AMPK.²⁸ ²⁹ Studies in mice have demonstrated that contraction-stimulated elevation of muscle ORM increases muscle glycogen and enhances muscle endurance,³⁰ which is mediated through activation of AMPK by ORM in skeletal muscle.¹⁰ Taken together, it seems reasonable to speculate that ORM per se may stimulate glucose uptake by myocytes and hence may lower blood glucose. Indeed, exogenous ORM improved glucose and insulin tolerance in diabetic db/db mice.²¹ However, in the present study, higher serum ORM was associated with higher, but not lower glycemia. It may be that sustained increase in circulating ORM might cause ORM resistance.²¹

Increased ORM may be a consequence of increased plasma glucose in the present study as it is reported that adipose ORM levels were elevated by high glucose³¹ and it has been suggested to modulate immune responses to protect adipose tissue from inflammation and metabolic dysfunction.²¹ However, this explanation appears to be unlikely in the present study because differentiated adipocytes were incubated with extremely high glucose (450 mg/dL)³¹ and because another study reported repressed adipose ORM expression in diabetic ob/ob mice, although ORM was expressed in adipose tissue at high levels.³² The biology underlying the association between high serum ORM concentrations and increased glucose excursion found in the present study is not known.

The strengths of the present study include homogeneous study population with scarce confounding factors, and accurate and reliable measures of body composition by dual-energy X-ray absorptiometry (DXA). Several limitations of this study warrant consideration. The cross-sectional design of the present study complicates the drawing of causal inferences, and a single measurement of biochemical variables may be susceptible to short-term variation, which would bias the results towards the null. We used crude measures of insulin sensitivity/IR and insulin secretion, which may be less accurate. Statistical power was not calculated. As we studied young Japanese women only, results may not be generalized to other gender, age populations, races or ethnicities.

In conclusion, elevated circulating orosomucoid was associated with elevated 30min glucose and glucose excursion in non-obese young Japanese women independently of general and abdominal adiposity, insulin secretion, insulin sensitivity/resistance, other investigated markers of inflammation and adiponectin. Further research is needed to determine the public health and clinical significance of these results. But these results may suggest a clue to identify novel pathways that may have utility in monitoring glucose dysmetabolism within NGT.

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