Comparison of Human Selenoprotein P Determinants in Serum between Our Original Methods and Commercially Available Kits

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Selenoprotein P (SeP) is a selenium (Se)-rich extracellular protein. SeP is identified as a hepatokine, causing insulin resistance in type 2 diabetes. Thus, the measurement of SeP in serum has received much attention, and several enzyme-linked immunosorbent assay (ELISA) kits for SeP determination are now commercially available. In the present study, we determined the serum SeP levels by our original ELISA and sol particle homogeneous immunoassay (SPIA) methods and also by commercially available kits, and these determinants were compared. We found a kit-dependent correlation of the determinants with our methods. These results suggest that the selection of kit is critical for comparison with our previous reports and for discussing the relationship between the serum SeP levels and disease condition.

Key words selenoprotein P; monoclonal antibody; enzyme-linked immunosorbent assay (ELISA); sol particle homogeneous immunoassay (SPIA); commercially available kit

Selenoprotein P (SeP; encoded by SELENOP) is a selenium (Se)-rich extracellular glycoprotein synthesized primarily by the liver.1) SeP contains the essential trace element Se as selenocysteine (Sec; an analog of cysteine having Se instead of sulfur), which is encoded by UGA codon in mRNA, formerly known as a stop codon.2) A stemloop RNA structure, called the Sec insertion sequence (SECIS), is essential for the incorporation of Sec during the biosynthesis of selenoproteins. The Sec insertion sequence (SECIS) is essential for the incorporation of Sec during the biosynthesis of selenoproteins. The SECIS sequence in the mRNA of selenoproteins is known as a stop codon.3) SeP functions as a Se transporter to maintain levels of selenoproteins, including antioxidant enzymes such as glutathione peroxidases (GPxs). SeP has multifunctional properties such as GPx-like enzyme activity and heavy metal binding activity.4) Thus, SeP plays a significant role in the antioxidative system in both a direct (GPx-like enzyme activity) and an indirect manner (Se transporter). We have previously reported that SeP serves as a hepatokine, promoting glucose intolerance and insulin resistance in type 2 diabetes.5) SeP is upregulated in the liver of type 2 diabetes patients, and high levels of SeP impair insulin signaling and glucose metabolism in both liver and muscle of mice.6) We have recently reported that SeP is involved in exercise resistance by suppressing the levels of exercise-induced reactive oxygen species (ROS) in skeletal muscle.7) Furthermore, it has been discovered that excess SeP decreases both pancreatic insulin levels and insulin secretion induced by high glucose.8) Collectively, previous studies suggest that excess SeP is a significant therapeutic target for type 2 diabetes.

Preparation of recombinant selenoprotein for biochemical experiments is still difficult because of the complicated translational mechanism of UGA codon as Sec, particularly for SeP, possessing 10 Sec residues per polypeptide.1,2) We have established a method for purifying SeP protein from human plasma,3) and purified SeP protein was used for the preparation of monoclonal antibodies (mAbs).4) SeP is composed of two domains: one Sec residue in the N-terminal region functions as the catalytic center for GPx-like enzyme activity and the other nine Sec residues in the C-terminal region functions as a Se transporter (Supplementary Fig. 1A).9) These domains are connected by a bridge containing histidine-rich regions, each of which has the typical heparin-binding motif XBBXB (B is a basic amino acid). The digestion of full-length SeP (FL-SeP) with plasma kallikrein results in the limited proteolysis (Arg-235–Gln-236 and Arg-242–Asp-243), which generates N-terminal (SeP-NF) and C-terminal fragments of SeP (SeP-CF) (Supplementary Fig. 1A).9) Domain structures of SeP are defined by using these SeP fragments. The rough epitope of prepared mAbs is also determined. Based on these results, we have previously developed an enzyme-linked immunosorbent assay (ELISA) specific for FL-SeP, and for both FL-SeP and SeP-NF, referred to as total SeP (Supplementary Figs. 1B, C). We further developed a sol particle homogeneous immunoassay (SPIA) for measuring the serum FL-SeP levels, which is applicable to various clinical analyzers.10)

Recently, measurement of human SeP has received much attention, and several ELISA kits for human SeP have become commercially available. In the present study, we compared the SeP determinants of our methods with those of three kinds of commercially available ELISA kits.

MATERIALS AND METHODS

Reagents Rat anti-human SeP mAbs (Clone AA3, AH5, and BD3) used in the present study were prepared as described previously.9) Human SeP was purified from human plasma, as described previously.8) Human frozen plasma was purchased from the Japanese Red Cross Kinki Block Blood Center (No. 2530012).

Subjects Blood samples were obtained from patients...
with suspected ischemic heart disease who were admitted to Kanazawa University Hospital from 2010 to 2011. Written informed consent for this study was provided from all patients. The experimental protocol was approved by the Ethical Committee of Kanazawa University (Approval No. 2152-1). The current study was conducted in accordance with the Declaration of Helsinki. A total of 21 serum samples were analyzed, and the subjects comprised eighteen males and three females (mean age $\pm$ standard deviation, 67.2 $\pm$ 10.0). Following an overnight fast, venous blood samples were obtained from each subject.

Measurement of FL-SeP and Total SeP by ELISA Measurement of FL-SeP and total SeP by ELISA was conducted, as described previously. Ninety-six-well microtiter plates were coated with 100 $\mu$L of 5 $\mu$g/mL rat anti-human SeP mAb AA3 (for FL-SeP) or BD3 (for total SeP) in 0.05M sodium bicarbonate buffer (pH 9.6) for 1h at room temperature. The wells were washed with phosphate buffered saline (PBS) containing 0.05% Tween 20 (wash buffer) and incubated at 37°C with PBS containing 0.1% bovine serum albumin (BSA) (Blocking buffer) for 1h. After washing, 50 $\mu$L of FL-SeP standard or serum sample diluted with PBS containing 0.05% Tween 20 and 0.1% BSA (PBS-Tween-BSA) was added to each well and incubated at 37°C for 1h. After washing, 50 $\mu$L of horseradish peroxidase (HRP)-conjugated rat anti-human SeP mAb AH5 (20 $\mu$g/mL) was added and incubated at 37°C for 1h. Finally, the plates were washed, and fifty microliters of 3,3',5,5'-tetramethylbenzidine (TMB, Merck, Darmstadt, Germany) was added to each well, and the enzyme-substrate reaction was allowed to proceed for 30 min in the dark. The reactions were stopped by the addition of 50 $\mu$L of 1 M sulfuric acid to each well. The absorbances were read at 450 nm on an OPTImax plate reader (Molecular Devices, Sunnyvale, CA, U.S.A.). Wells receiving all reagents apart from serum sample

![Graphs](image-url)
Measurement of FL-SeP by SPIA Measurement of FL-SeP by SPIA was conducted using a Model 7180 Hitachi automatic clinical analyzer (Hitachi, Tokyo, Japan), as described previously. Colloidal particles coated with rat anti-human SeP mAb were prepared as described previously. An aliquot of the sample (2 µL) was pipetted into a cuvette, followed by reaction buffer (120 µL). The composition of reaction buffer is as follows: 0.25 M Tris-buffer (pH 7.8) containing 50 mg/mL NaCl, 2 mg/mL ethylenediaminetetraacetic acid disodium salt (EDTA 2Na), 23 mg/mL polyethylene glycol 20000, 3.4 mg/mL gamma-butyrolactone, 3.5 mg/mL polyoxyethylene lauryl ether (Brij 35, Nacalai Tesque, Kyoto, Japan), and 2 mg/mL dodecyl(sulfonatophenoxy)benzenesulfonate (Pelex SS-H, Kao Corporation, Tokyo, Japan). After 5 min at 37°C, a mixture of colloidal gold particles coated with AH5 and AA3 (60 µL) was added and mixed. The reaction between the particles and FL-SeP in the samples resulted in the formation of agglutinates and a concomitant change in the absorbance signal. After the agglutination reaction started, the absorbance values at 505 nm (primary wavelengths) and 660 nm (secondary wavelengths) were measured for 5 min immediately. The differences between initial absorbance and the absorbance 5 min after the initiation of the reaction were calculated. By using the spline program of a Model 7180 Hitachi automatic clinical analyzer, a standard curve for FL-SeP was calculated.

Measurement of SeP by Using Commercially Available Kits The following kits were used for measurement of SeP in the present study: Selenotest ELISA (InVivo BioTech Services GmbH, Berlin, Germany), ELISA Kit for Selenoprotein P1, Plasma (Cloud-Clone, Houston, U.S.A.), and Human Selenoprotein P (SEPP1) ELISA kit (Kit C)—and the correlation between determinants was evaluated. The solid line represents the regression line. Formula, $r$, and $p$ values are shown in each figure.
Selenoprotein P (SEPP1) ELISA kit (CUSABIO, Wuhan, China). SeP content was measured according to the supplier’s protocol, and the standard material of each kit was used for the determinants.

**Statistical Analysis** Statistical analyses were performed using Excel software and Statcel 3. Correlation coefficients were assessed using Pearson’s correlation coefficient test and are shown in each figure. The regression line is also indicated. $p<0.05$ was considered to be significant.

**RESULTS AND DISCUSSION**

**Correlation of Serum SeP Levels between Our ELISA and SPIA Method** Twenty-one human serum samples with different concentrations of SeP were subjected to ELISA and SPIA for SeP using different mAbs. The rough epitope of mAbs used in the present study is as follows: AH5 and BD3, N-terminal region; AA3, C-terminal region (Supplementary mAbs used in the present study is AH5 and BD3, C-terminal region (Supplementary Fig. 1A). Therefore, the measurement system using AA3-AH5 is for FL-SeP, while BD3-AH5 is for total SeP, containing both FL-SeP and SeP-NF (Supplementary Figs. 1B, C). The significant correlation between AA3-AH5 ELISA and AA3-AH5 SPIA suggests that these mAbs are working well in both systems (Fig. 1A). In addition, we found good correlation between BD3-AH5 ELISA and AA3-AH5 SPIA (Fig. 1B), and between AA3-AH5 ELISA and BD3-AH5 ELISA (Fig. 1C), suggesting small variation of SeP-NF levels in serum samples used in the present study.

**Comparison between Serum SeP Determinants of Our Method and Those of Commercially Available Kits** The determinants of our methods using AA3-AH5 SPIA, AA3-AH5 ELISA, and BD3-AH5 ELISA were significantly correlated with those of Selenotest ELISA (Kit G) from InVivo BioTech Services GmbH Germany (Figs. 2A–C). In the case of Kit G, ELISA for SeP was developed by two kinds of anti-SeP mAbs, which were prepared by using recombinant mutant SeP protein (all Sec replaced with cysteine) as immunogen. Although the precise epitope of these mAbs has not been reported, the good correlation of the determinants in Kit G with those of our FL-SeP method suggests that Selenotest ELISA determines the FL-SeP.

Next, we further evaluated the correlation of determinants between our methods and ELISA Kit for Selenoprotein P1, Plasma (SEPP1) (Kit U) from Cloud-Clone U.S.A.; however, a significant correlation between these was not observed (Figs. 3A–C). In the case of Kit U, although precise preparation of Abs used was not described in the instructions, it appears that recombinant SeP protein from Tyr 60 to Ser 299 (Sec is not included in this sequence) might be used as an immunogen. The high values of SeP, from 20 to 60 µg/mL, in Kit U were obvious, and it is unclear whether unknown fragments of SeP are present in the human serum samples.

In the case of Human Selenoprotein P (SEPP1) ELISA kit (Kit C) from CUSABIO China, a significant correlation with our system was also not observed (Figs. 4A–C). Precise preparation of Abs used in Kit C was not described in the instructions, and it appears that recombinant mutant SeP protein (all Sec replaced with Ser) might be used as an immunogen. Similar to the determinants of Kit U, the high values of SeP, from 10 to 20 µg/mL, were observed. We also evaluated the correlation of determinants between Kit U and Kit C; however, a significant correlation between these was not observed (Supplementary Fig. 2).

In the present study, we compared the determinants of serum SeP of our ELISA and SPIA methods with those of commercially available kits. We found a correlation between our determinants and kit G, but not kit U or kit C. These results suggest that the choice of kit is critical for discussing about serum levels of human SeP including the reproducibility of our previous results.

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**Conflict of Interest** The authors declare no conflict of interest.

**Supplementary Materials** The online version of this article contains supplementary materials.

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