Proteomic data show an increase in autoantibodies and alpha-fetoprotein and a decrease in apolipoprotein A-II with time in sera from senescence-accelerated mice

S.J. Guo¹,³, C.H. Qi¹, W.X. Zhou¹, Y.X. Zhang¹, X.M. Zhang², J. Wang² and H.X. Wang²

¹Beijing Institute of Pharmacology and Toxicology, Beijing, China
²National Center of Biomedical Analysis, Beijing, China
³Shanghai Key Laboratory of Hypertension, Ruijin Hospital, Shanghai Jiaotong University School of Medicine, Shanghai, China

Abstract

We evaluated changes in levels by comparing serum proteins in senescence-accelerated mouse-prone 8 (SAMP8) mice at 2, 6, 12, and 15 months of age (SAMP8-2 m, -6 m, -12 m, -15 m) to age-matched SAM-resistant 1 (SAMR1) mice. Mice were sacrificed, and blood was analyzed by 2-dimensional electrophoresis combined with mass spectrometry. Five protein spots were present in all SAMP8 serum samples, but only appeared in SAMR1 samples at 15 months of age except for spot 3, which also showed a slight expression in SAMR1-12 m sera. Two proteins decreased in the sera from SAMP8-2 m, -6 m, and -12 m mice, and divided into 2 spots each in SAMP8-15 m sera. Thus, the total number of altered spots in SAMP8 sera was 7; of these, 4 were identified as Ig kappa chain V region (M-T413), chain A of an activity suppressing Fab fragment to cytochrome P450 aromatase (32C2_A), alpha-fetoprotein, and apolipoprotein A-II. M-T413 is a monoclonal CD4 antibody, which inhibits T cell proliferation. We found that M-T413 RNA level was significantly enhanced in splenocytes from SAMP8-2 m mice. This agreed with serum M-T413 protein alterations and a strikingly lower blood CD4⁺ T cell count in SAMP8 mice when compared to the age-matched SAMR1 mice, with the latter negatively correlating with serum M-T413 protein volume. Age-related changes in serum proteins favored an increase in autoantibodies and alpha-fetoprotein and a decrease of apolipoprotein A-II, which occurred in SAMP8 mice at 2 months of age and onwards. These proteins may serve as candidate biomarkers for early aging.

Key words: SAMP8; Serum proteins; 2-DE; Mass spectrometry; M-T413

Introduction

Alzheimer's disease (AD), as the prevalent form of progressive dementia, is representative of diseases related to impaired cognition: its early diagnosis so far remains challenging, although some symptomatic treatments and new drugs are emerging to delay and/or reverse its progress (1). It is therefore essential to identify the mechanisms of poor cognition by using biomarkers to facilitate the early diagnosis of neurodegenerative diseases characterized by deficits in learning and memory.

Mounting evidence has indicated that impaired cognition was slight but significant as early as at 2 months of age, and became more obvious with aging in senescence-accelerated mouse-prone 8 (SAMP8) mice compared to SAM-resistant 1 (SAMR1) mice, which exhibit normal aging processes (2,3). Moreover, SAMP8 mice also exhibited many of the biochemical findings of AD, such as decreased glucose metabolism and alterations in amyloid precursor protein, apolipoprotein E, bcl-2, presenilin-2, tau, and some oxidative markers (4-6). Taken together, these studies demonstrate that the SAMP8 mouse might be an ideal animal model for the investigation of the mechanisms underlying age-related learning and memory deficits in AD within the framework of gene and protein levels (7). Accordingly, a series of advanced genomic and proteomic investigations have been conducted in the brain, liver, and spleen tissue of SAMP8 mice and have shown that signal transduction change, impaired neuroprotection, abnormal fatty acid metabolism, depressed energy production, and enhanced oxidative stress-induced molecular damage were reportedly involved in learning and memory deficits (8-15). However, considering clinical practicability, it is undoubtedly optimal...
to identify the biomarkers contributing to these deficits in serum. The SAMP8 mouse has been employed since 1986 to assess age-related neurobiological features and to understand their pathological mechanisms. However, data on serum changes in accelerated senescence mice are relatively limited. For example, SAMP8 mice displayed an age-related decrease in testosterone and estradiol (16,17). In addition, dozens of abnormal metabolites were found in SAMP8 mouse sera. Among these, the most important metabolite responsible for the strain separation was a lack of inosine, which meant that the protective functions of anti-inflammation, immunomodulation and neuroprotection might be attenuated in SAMP8 mice (18). In addition, regarding serum proteins, decreased immunoglobulin E (IgE) (19) and increased anti-elastin antibodies (20) occurred in SAMP8 mouse serum, with the latter supporting a relationship between autoantibody and aging in SAMP8 mice for the very first time. To identify more age-relevant proteins, systematic views on serum protein changes during the aging process are warranted.

In the present study, we combined 2-dimensional electrophoresis (2-DE) with robust mass-spectrometry-based proteomic approaches to compare differential expression patterns of serum proteins between SAMP8 and SAMR1 mice at 2, 6, 12, and 15 months of age. This is the first time that the pathophysiological development of the SAMP8 mouse serum proteome has been investigated by evaluating global changes that occur with aging. Our results provide some information for understanding the dysfunction in the senescence-accelerated mouse and identifying potential novel serum biomarkers for early aging.

Material and Methods

Ethics statement
Male SAMP8 mice at 2, 6, 12, and 15 months of age (SAMP8-2 m, -6 m, -12 m, -15 m) and age-matched SAMR1 mice were generously provided by Dr. Toshio Takeda, Kyoto University, and maintained at the Laboratory Animal Center of the Academy of Military Medical Sciences (Protocol: SYXX JUN 2007-004). The mice were treated according to the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health with the approval (Protocol: SYXX JUN 2007-008) of the Animal Care and Use Committee of the National Beijing Center for Drug Safety Evaluation and Research.

2-DE and analysis
Mice (n = 4-5) were sacrificed after collection of blood samples (0.5 mL/mouse) by removing the eyeball. The blood was allowed to clot for 30 min at room temperature. Sera were obtained by centrifugation at 112 g for 10 min at 4°C and preserved at -80°C until use.

Each serum sample was separated by 2-DE. After determining protein concentration by the Bradford assay using bovine serum albumin as standard, 150 μg protein (for the comparative analysis of protein spots) or 1.5 mg protein from the serum of 1 mouse (for protein identification by mass spectrometry) was diluted with a rehydration buffer [8 M urea; 2% (w/v) CHAPS; 20 mM DTT; 0.5% (v/v) Immobilized pH Gradient (IPG) buffer, pH 3-10, and 0.002% bromophenol blue] to 350 μL and was then applied to IPG strips (18 cm, pH 3-10 linear, GE Healthcare Bioscience, Sweden). Isoelectric focusing was performed with the IPGphor system (GE Healthcare Bioscience) according to the following programmed settings: 30 V for 6 h, 60 V for 6 h, 200 V for 1 h, 500 V for 1 h, 1000 V for 1 h, and 8000 V for 1 h at gradient type, and 8000 V until reaching 64 kVh. Accordingly, the IPG strip was equilibrated for 15 min in an equilibration buffer containing 6 M urea, 50 mM Tris-HCl, 30% (v/v) glycerol, 2% (w/v) SDS and 0.02% (w/v) bromophenol blue with 10 g/L DTT, and then equilibrated for another 15 min in the same buffer but with 25 g/L iodoacetamide replacing the DTT. The second dimension electrophoresis was performed on 12.5% SDS-polyacrylamide gels with a low molecular weight marker (GE Healthcare Bioscience). Gels were then stained with silver for further analysis and with Coomassie brilliant blue R-250 for mass spectrometry for protein identification. The silver-stained gels were scanned at a 300-dpi resolution and protein spots were analyzed with the ImageMaster Platinum™ software (GE Healthcare Bioscience) according to manufacturer recommendations. For each sample, we performed electrophoresis followed by silver staining three times. Spots with a P value ≤ 0.05 for the t-test and a 2.0-fold or greater alteration in abundance were accepted as significantly changed proteins. The differentially expressed protein spots were cut and digested using a method based on a previous study by Jin et al. (21).

Peptide mass fingerprinting by matrix-assisted laser desorption/ionization-time of-flight mass spectrometry (MALDI-TOF-MS)
An aliquot (0.5 μL) of tryptic peptides was mixed with 1.5 μL matrix consisting of a saturated solution of alpha-cyano-4-hydroxycinnamic acid in 50% (v/v) acetonitrile per 0.1% (v/v) trifluoroacetic acid. The mixture (1 μL) was immediately added to a MALDI plate and dried at room temperature. The dried spots were analyzed with a Bruker REFLEX III MALDI-TOF-MS (Bruker-Franzen, Germany) in positive ion mode at an accelerating voltage of 20 kV. The spectra were internally calibrated using trypsin autolysis products. The peptide mass fingerprints (PMF) were used to search through the SWISS-PROT and NCBI databases using the Mascot search engine (http://www.matrixscience.co.uk). The MS peak filtering included the following parameters: a mass range of 800 to 4000 Da with a tolerance of 0.1 Da; fixed and variable modifications due to carbamidomethylation (C) and oxidation (M),
respectively, and a maximum of 1 missed cleavage for trypsin digestion. The results were acceptable as a positive identification when score confidence was higher than 95%.

**Peptide sequencing by electrospray ionization nanoelectrospray tandem MS (ESI-MS/MS)**

ESI-MS/MS experiments were conducted using a Q-TOF 2 hybrid quadrupole TOF mass spectrometer (Micromass, UK) with a nanoflow Z-spray source. Peptide sequencing was performed using a palladium-coated borosilicate electrospray needle (Protana, Denmark) according to the method of Yan et al. (22). The MS was operated in the positive ion mode with a source temperature at 80°C, and a potential of 800 to 1000 V applied to the nanospray probe. The amino acid sequences of the peptides were deduced with the peptide sequencing program MasSeq and the database search was completed with the Mascot search engine using data processed through MaxEnt3 and MasSeq programs.

**Real-time quantitative polymerase chain reaction (RT-PCR)**

Splenocytes were isolated as described by Abe et al. (23). RNA from splenocytes was extracted using TRIzol reagents (Invitrogen Corporation, USA) according to manufacturer instructions. Quantification was carried out using absorbance at 260 nm and quality was confirmed by 1% agarose gel electrophoresis. RNA samples (n = 3) with an A260/A280 nm of 1.8-2.0 were stored at -80°C for further analysis.

For the M-T413 gene, the forward and reverse primers were 5′-GGC CCA TTC TCG AGC TTG TTT GTC CC-3′ and 5′-GTG GAC AGT GAG GCC AGG AT-3′, respectively; for the β-actin gene, the forward and reverse primers were 5′-CTG TTT ATT TCC AGC TTG GTC CC-3′ and 5′-GTG GAC AGT GAG GCC AGG AT-3′, respectively. These primers were designed using the Primer Premier 5.0 software and synthesized by Shenggong Corporation (China). Quantitative RT-PCR was carried out with a thermal cycler (Applied Biosystems, USA) by monitoring the amount of fluorescence increase due to the binding of SYBR green to double-stranded DNA. RT-PCR was performed on a final volume of 20 μL containing 1 μg RNA template, 1.6 μL of mixed primers, 400 μM of each dNTP, 0.4 μL AMV reverse-transcribed enzyme (5 U/μL), 400 0.4 μL Tfl DNA polymerase, 0.2 μL SYBR green, and 10 μL reaction buffer using an Access RT-PCR System (Promega Corporation, USA). Briefly, the RNA template was reverse-transcribed into cDNA templates for 45 min at 48°C and denatured for 2 min at 94°C, and the denatured cDNA templates were amplified by cycles of 94°C for 30 s, 64°C for 60 s, and 72°C for 120 s. Thirty-four cycles were performed to determine the linearity of the PCR amplification, and amplified β-actin cDNA served as a control for cDNA quality and quantification. Fluorescence measurements were recorded after each extension step. At the end of each PCR run, data were automatically analyzed by the system and amplification plots were generated. Data were calculated using the standard curve of the gene template, normalized with β-actin.

**Flow cytometry analysis**

CD4+ T cells in peripheral blood were analyzed with a FACScalibur flow cytometer equipped with the CellQuest software (BD Biosciences, USA). Blood samples (n = 5) were incubated with phycoerythrin-conjugated anti-CD4 antibodies (BD Biosciences) and fluorescein isothiocyanate-conjugated anti-CD3 (BD Biosciences) for 30 min at 4°C in the dark. A 0.1-mL blood sample was added to 2 mL BD FACS Lysing Solution (BD Biosciences) to lyse red blood cells. Following fixation with 1% formaldehyde, the percentage of CD4+ T cells was detected. The white blood cell count and lymphocyte size were examined by automatic cytometry. The formula for calculating the CD4+ T cell count was defined as the white blood cell count x percentage of lymphocyte x percentage of CD4+ T cells.

**Statistical analysis**

Data are reported as means ± SD. The Student paired t-test was employed to evaluate the differences in means between the 2 strains and Pearson’s correlation analysis was used to identify the relationship between the continuous variables of interest. A comparison was considered to be significant if the two-sided P value was less than 0.05.

**Results**

**Comparison of protein expression profiles between sera from SAMP8 and SAMR1 mice**

We first separated serum proteins from SAMP8-2 m, -6 m, -12 m, and -15 m and age-matched SAMR1 mice by 2-DE. A total of 315 spots were detected on the maps by silver staining, and the overall protein expression profiles with pH 3-10 and molecular masses of 10 to 90 kDa were very similar within each predetermined group after analysis by the ImageMaster 2-D Platinum software, indicating the stability and reproducibility of 2-DE in our test system.

Two representative 2-DE gel images from SAMP8 and SAMR1 mice are illustrated in Figure 1, showing that 7 differentially expressed proteins were detected when the two strains were compared. Specifically, spots 1-5 appeared in all SAMP8 mouse serum samples, but only appeared in SAMR1 mouse samples at the age of 15 months, with the exception of spot 3, which showed a slight expression in SAMR1-12 m mouse sera (Table 1 and Figures 1, 2A and 3A). In addition, spots 6 and 7 appeared in the sera from SAMP8-2 m, -6 m, and -12 m,
and divided into 2 spots each with mass and charge differing from those of the same spots in SAMP8-15 m sera (Figure 2B).

Identification of proteins by MALDI-TOF-MS with PMF and ESI-MS/MS with peptide sequences

After MALDI-TOF-MS and ESI-MS/MS analyses, spots 3 to 6 were identified as Ig kappa chain V region (M-T413), chain A of an activity suppressing Fab fragment to cytochrome P450 aromatase (32C2_A), alpha-fetoprotein, and apolipoprotein A-II (Apo A-II), respectively. However, because the database search yielded no peptides whose score was high enough to provide unambiguous results and because no qualified peptides could be detected by ESI-MS/MS sequencing, the remaining 3 protein spots unfortunately were not identified in the present study. Information about the 7 aforementioned protein spots is summarized in Table 1.

Expression of M-T413 in SAMP8 splenocytes

Previous studies have demonstrated that decreased T cell immune function is closely related to age-associated cognitive impairment in SAMP8 mice (23-25). The cause of the decreased T cell immune function in SAMP8 mice remains an open question. In the present study, we identified a differentially expressed protein (M-T413) via joint PMF and peptide sequencing (Figure 3 and Table 1). M-T413 is a monoclonal CD4 antibody binding to the CD4 V1 domain and can inhibit T cell proliferation in a mixed lymphocyte response, thus acting to immunosuppress the CD4+ T cell response (26-28). In the present study, M-T413 was expressed in all SAMP8 mouse sera and in the sera from SAMR1-12 m and -15 m mice (Figure 2, Figure 3A and Table 1), exhibiting a close association with senescence. However, since an anti-M-T413 antibody is not commercially available, serum M-T413 protein could not be determined by ELISA or Western blot. However, as an Ig kappa chain V region, we hypothesized that M-T413 was produced by B lymphocytes mainly derived from the spleen. Therefore, the M-T413 mRNA level was assessed in splenocytes from SAMP8 and age-matched SAMR1 mice. M-T413 mRNA was significantly enhanced in splenocytes from SAMP8-2 m mice (Figure 3B) compared to that from age-matched SAMR1 mice, in agreement with M-T413 protein changes in sera from 2-month old mice (Figure 3A), which once again validated the correlation of M-T413 with early impaired cognition. However, inconsistent with the M-T413 protein existing in SAMR1-12 m mouse sera, we observed that the M-T413 RNA level was elevated in the SAMR1 mouse spleen from 6 months of age, and maintained at a higher level until 15 months of age (Figure 3B). It is likely that splenocytes are not the unique source of serum M-T413, which warrants further evaluation.

CD4+ T cell count in SAMP8 blood

Furthermore, another report by Abe et al. (23) indicating the decreased number of CD4+ T cells in the SAMP8-2 m mouse spleen with respect to the age-matched SAMR1 mouse spleen, combined with our observation of the elevated M-T413 RNA level, led us to speculate that M-T413 might be accountable for CD4+ T cell variability. This speculation was further strengthened by the fact that no material changes were observed in the
| Spot | Description | NCBI accession No. | Theoretic Mr (Da/p) | Observed Mr (Da/p) | Peptide count | Sequence coverage | PMF Score | Peptide count | ESI-MS/MS Score | Fold-change (SAMP8/SAMR1) |
|------|-------------|--------------------|---------------------|-------------------|---------------|------------------|------------|--------------|----------------|------------------|
| 1    | Unnamed protein product | 74200884 | 43660/5.01 | 29698/4.67 | 4 | 8% | 35 | N | N | +/– | 5.41 ± 1.19 |
| 2    | Leo1 protein | 52139032 | 35905/3.96 | 29698/4.83 | 5 | 20% | 36 | N | N | +/– | 6.18 ± 1.68 |
| 3    | Ig kappa chain V region (M-T413) | 110545 | 12214/5.18 | 30133/4.90 | 3 | 58% | 49 | 2 | 188 | +/– | 12.7 ± 0.9 |
| 4    | Chain A, structure of an activity-suppressing Fab fragment to cytochrome P450 aromatase (32C2_A) | 7766934 | 23788/6.23 | 30133/5.07 | 9 | 46% | 74 | 2 | 165 | +/– | 3.16 ± 0.34 |
| 5    | Alpha-fetoprotein | 59797643 | 69158/6.09 | 6.72/5.36 | 5 | 24% | 34 | 2 | 139 | +/– | 1.98 ± 0.67 |
| 6    | Apolipoprotein A-II | 114003 | 11319/5.25 | 12736/4.48 | 3 | 28% | 42 | 1 | 83 | -4.8 ± 1.1 | -4.5 ± 0.9 | -7.6 ± 2.3 | - |
| 7    | Unnamed protein product | 12844891 | 11859/8.59 | 12754/4.88 | 2 | 23% | 27 | N | N | -6.7 ± 1.9 | -2.4 | -4.6 ± 0.9 | - |

Mr = relative molecular mass; pI = isoelectric point; PMF = peptide mass fingerprints; SAMP8 = senescence-accelerated mouse-prone 8; SAMR1 = senescence-accelerated mouse-resistant 1. Peptide count means the number of peptides matched in MS analysis. The trends of protein changes are reported as upward (positive value) or downward (negative value) fold-change of spot volumes (means ± SD, n = 5). N = no significant result; - = not determined, +/– = the spot only appeared in SAMP8 sera.

Spot 6 could be detected in 2 of 5 SAMP8 sera at 6 months age. <sup>a</sup>Spot 7 could be detected in 1 of 5 SAMP8 sera at 6 months age. <sup>b,c</sup>Fold-change (15-month-old SAMP8/SAMR1) of spot 6 and spot 7 could not be calculated because they were divided into 2 separate spots in 15-month-old SAMP8 sera.

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Mr = relative molecular mass; pI = isoelectric point; PMF = peptide mass fingerprints; SAMP8 = senescence-accelerated mouse-prone 8; SAMR1 = senescence-accelerated mouse-resistant 1. Peptide count means the number of peptides matched in MS analysis. The trends of protein changes are reported as upward (positive value) or downward (negative value) fold-change of spot volumes (means ± SD, n = 5). N = no significant result; - = not determined, +/– = the spot only appeared in SAMP8 sera.

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number of CD4$^+$ T cells in the SAMP8-5 m mouse spleen in the study by Abe et al. (23), or in the M-T413 RNA level observed in the splenocytes of SAMP8 mice at 6 months and onwards in the present study (Figure 3B). To further address this hypothesis, we determined CD4$^+$ T cell counts in blood samples from both SAMP8 and SAMR1 mice. As shown in Figure 3C, the CD4$^+$ T cell count in SAMP8 mouse blood was strikingly lower than that in the age-matched SAMR1 mice, which negatively correlated with serum M-T413 protein volume ($r = -0.821$, $P = 0.013$). Taken together, these results support our speculation that M-T413 might be one of the important reasons for reduced CD4$^+$ T cells in SAMP8 mice.

Discussion

In clinical practice, the evaluation of imaging findings has been incorporated into the diagnosis of AD along with traditional clinical criteria (29,30). Generally, an ideal biomarker for disease diagnosis should have relatively high sensitivity and specificity, and should have potential advantages for early detection. In view of the difficulties encountered in AD diagnosis, the screening of early stage biomarker candidates from an ideal animal model is highly desirable. Bearing this in mind, we explored the alterations of serum proteins in SAMP8 mice related to age, number of CD4$^+$ T cells in the SAMP8-5 m mouse spleen in the study by Abe et al. (23), or in the M-T413 RNA level observed in the splenocytes of SAMP8 mice at 6 months and onwards in the present study (Figure 3B). To further address this hypothesis, we determined CD4$^+$ T cell counts in blood samples from both SAMP8 and SAMR1 mice. As shown in Figure 3C, the CD4$^+$ T cell count in SAMP8 mouse blood was strikingly lower than that in the age-matched SAMR1 mice, which negatively correlated with serum M-T413 protein volume ($r = -0.821$, $P = 0.013$). Taken together, these results support our speculation that M-T413 might be one of the important reasons for reduced CD4$^+$ T cells in SAMP8 mice.

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which, as a non-transgenic strain with accelerated senescence, are accepted as a suitable model for assessing the pathophysiological features of AD (31-33).

Specific expression of spots 1 to 5 in SAMP8 and SAMR1 mouse serum at 15 months of age enabled us to conclude that these differential spots might have a close relationship both with accelerated senescence and normal aging.

Unlike the spots mentioned above, spots 6 and 7 did not exhibit changes similar to those that occurred in the normal aging process, suggesting that their alteration was only associated with accelerated senescence. However, spots 1 to 7 showed significant changes as early as by 2 months of age, and thus they might serve as candidate biomarkers of early deficits in cognition. Accordingly, we analyzed these "differential" spots using MALDI-TOF-MS and ESI-MS/MS after tryptic in-gel digestion.

32C2_A, alpha-fetoprotein and Apo A-II

Besides the M-T413 protein mentioned in the Results section, 32C2_A and alpha-fetoprotein were also reported to be involved in senescence among differentially expressed proteins identified for the first time in the present study. In the present study, the expression of 32C2_A (spot 4), an autoantibody suppressing cytochrome P450 aromatase activity (34), was only detected in sera from all SAMP8 mice and SAMR1-15 m mice (Figure 2A). P450 aromatase, as a key enzyme in estrogen biosynthesis, can convert androgen to estrogen (35). We thus speculated that 32C2_A might inhibit estrogen biosynthesis by suppressing the activity of P450 aromatase. This was supported by our previous observation that the concentration of serum estradiol was lower in all SAMP8 mice compared to SAMR1 mice (17). In addition, elevations of M-T413, 32C2_A and anti-elastin antibodies (20) in SAMP8 mouse sera suggest that autoantibodies play a part in the impaired cognition of SAMP8 mice.

We also identified another protein, alpha-fetoprotein (spot 5), in serum samples from all SAMP8 mice and SAMR1-15 m mice (Figure 2A). Alpha-fetoprotein is a serum alpha-globulin synthesized by embryo hepatocytes and the yolk sac during the embryonic stage and is nearly undetectable in adult serum (36). It serves as a marker for the diagnosis of hepatocellular carcinoma owing to a raised synthesis and secretion when tumors occur (37). In the present study, a very novel finding was alpha-fetoprotein showed a close correlation with impaired cognition, so that the exact mechanism of its expression in sera of accelerating senescence mice remains worthy of further investigation.

Another protein involved in senile amyloidosis, Apo A-II (spot 6), was decreased in sera from SAMP8-2 m, -6 m, and -12 m mice, and divided into 2 spots in SAMP8-15 m mouse sera as shown in Figure 2B. There are 3 types of Apo A-II: type A Apo A-II, type B Apo A-II, and type C Apo A-II. In these 3 types, amino acid substitutions have been reported at a total of 4 different positions. An Apo A-II with methionine at position 26 was detected in SAMP8, SAMP3, SAMR2, and C57BL/6J mice with a moderate incidence of amyloid deposition; B Apo A-II with proline substituted at position 5 was mainly observed in SAMR1, SAMR4, SAMP6, and BALB/c mice with a low incidence of senile amyloidosis, and C Apo A-II was found in SAMP1, SAMP2, SAMP7, and SAMP9 mice with a severe and high incidence of amyloidosis (38). In view of this evidence, we conjectured that spot 6 might be B Apo A-II due to its higher abundance in SAMR1 mice, whereas its actual type requires further identification.

The limitations of the present study include 3 of the 7 proteins being unsuccessfully identified and no verification of M-T413 expression changes in SAMP8 mouse serum owing to the unavailability of an M-T413 antibody. Moreover, besides the 7 proteins mentioned above, the count and abundance of protein spots with an isoelectric point (pI) between 5.4-8.3 and relative molecular mass (Mr) of 5.3-6.7, and of protein spots with a pI between 4.9-8.0 and Mr of 2.8-3.2 also differed significantly between the 2 strains (Figure 1). These protein spots could not be further analyzed by MS because they were not separated into single spots.

Despite these limitations, our study still showed that the expression levels of some serum proteins, especially the autoantibodies, were significantly different between SAMP8 and SAMR1 mice from 2 months of age onwards. Our results may provide a reference point for the understanding of the basic pathophysiological mechanisms of learning and memory deficits, and these changed proteins may serve as candidate biomarkers of early impaired cognition. On this basis, future investigations can focus on elucidating the role and mechanisms of these proteins in age-related cognitive impairment and identify additional potential serum biomarkers using further proteomic methods such as narrow pH range IPG strips and the removal of high abundance serum proteins.

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