Title: Advanced glycation endproducts increase proliferation, migration and invasion of the breast cancer cell line MDA-MB-231

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Abstract

Diabetic patients have increased likelihood of developing breast cancer. Advanced glycation endproducts (AGEs) underlie the pathogenesis of diabetic complications but their impact on breast cancer cells is not understood. This study aims to determine the effects of methylglyoxal-derived bovine serum albumin AGEs (MG-BSA-AGEs) on the invasive MDA-MB-231 breast cancer cell line. By performing cell counting, using wound-healing assay, invasion assay and zymography analysis, we found that MG-BSA-AGEs increased MDA-MB-231 cell proliferation, migration and invasion through Matrigel\textsuperscript{TM} associated with an enhancement of matrix metalloproteinase (MMP)-9 activities, in a dose-dependent manner. Using Western blot and flow cytometry analyses, we demonstrated that MG-BSA-AGEs increased expression of the receptor for AGEs (RAGE) and phosphorylation of key signalling protein extracellular signal-regulated kinase (ERK)-1/2. Furthermore, in MG-BSA-AGE-treated cells, phospho-protein micro-array analysis revealed enhancement of phosphorylation of the ribosomal protein 70 serine S6 kinase beta 1 (p70S6K1), which is known to be involved in protein synthesis, the signal transducer and activator of transcription (STAT)-3 and the mitogen-activated protein kinase (MAPK) p38, which are involved in cell survival. Blockade of MG-BSA-AGE/RAGE interactions using a neutralizing anti-RAGE antibody inhibited MG-BSA-AGE-induced MDA-MB-231 cell processes, including the activation of signalling pathways. Throughout the study, non-modified BSA had a negligible effect. In conclusion, AGEs might contribute to breast cancer development and progression partially through the regulation of MMP-9 activity and RAGE signal activation. The up-regulation of RAGE and the concomitant increased phosphorylation of p70S6K1 induced by AGEs may represent promising targets for drug therapy to treat diabetic patients with breast cancer.

Key Words: Advanced glycation endproducts, diabetes, breast cancer, signalling pathway, methylglyoxal.

Abbreviations: AGE, advanced glycation endproduct; AP, activator protein; BSA, bovine serum albumin; DMEM, Dulbecco's Modified Eagle's medium; EDTA, ethylenediaminetetraacetic acid; ERK, extracellular-signal regulated kinase; FBS,
foetal bovine serum; JNK, c-jun kinase; RAGE, receptor for advanced glycation endproducts; MAPK, mitogen-activated protein kinase; MG, methylglyoxal; MG-BSA-AGES, methylglyoxal-derived bovine serum albumin AGEs; MMP, matrix metalloproteinase; NF-κB, nuclear factor kappa B; PI3K, phosphoinositol-3 kinase; PKC, protein kinase C; PMSF, phenylmethylsulfonyl fluoride; p70S6K1, ribosomal protein 70 serine S6 kinase beta 1; ROS, reactive oxygen species; SD, standard deviation; SDS-PAGE, sodium dodecyl sulphate polyacrylamide gel electrophoresis; SPM, serum-poor medium; STAT, signal transducer and activator of transcription.

1. Introduction

Diabetes and cancer are major health problems and prevalent diseases with a globally increasing incidence. Generally, cancer has an increased metabolism associated with elevated glycolytic rates which enhance the formation of advanced glycation endproducts (AGEs) [1]. AGEs are a group of heterogeneous macromolecules formed during a non-enzymatic reaction called the Maillard reaction [2]. This Maillard reaction is responsible for the intermolecular cross-linking between the reactive carbonyl group of sugars and the nucleophilic amino group of proteins, lipids or nucleic acids [3, 4]. Although this glycation reaction is a non-specific biochemical reaction, protein modification results in altered enzyme activity [5], immunogenicity [6], decreases ligand binding [7], and cross-linking of extracellular matrix proteins, which cause stiffening [8]. Increased protein glycation has been associated with several age-related [9] and chronic inflammatory diseases such as cardiovascular diseases [10], Alzheimer’s disease [11], stroke, retinopathy [12], nephropathy [13], neuropathy and cancer [14]. In addition, AGE formation is often accompanied by increased free radical activity, which can damage cell membranes and cause gene mutations resulting in malignant cell transformation [15].

The cellular effects of AGEs are mainly mediated through the receptor for advanced glycation endproducts (RAGE). RAGE is a 45-kDa transmembrane multi-ligand signal transduction receptor belonging to the immunoglobulin superfamily, which is highly expressed during embryonic development, especially in the brain, and then its expression decreases in adult tissues [16]. In pathological conditions, RAGE expression is induced by high glucose [17], reactive oxygen species (ROS), external stress, hypoxia [18], pro-inflammatory mediators and by AGE itself [17]. RAGE is
also over-expressed on activated immune cells [19], vascular cells [20] and cancer cells [21]. AGE-RAGE interactions trigger a diverse array of signalling pathways [22], including mitogen activated protein kinase (MAPK) such as extracellular-signal regulated kinase (ERK)-1/2 [20], p38 [23] and c-jun kinase (JNK), that induce the activation of transcription factors such as nuclear factor kappa B (NF-κB) [24], activator protein (AP)1 and signal transducers and activators of transcription (STAT)-3 [25]. This is accompanied by ROS production [26] and by subsequent transcription and changes in gene expression, which result in tumour growth through increased cell proliferation [27] and tumour metastasis through stimulation of cell migration and invasion [14]. In addition, matrix metalloproteinases (MMPs) and especially gelatinases (i.e. MMP-2 and MMP-9) are known to promote cancer progression with their central role during the invasion process as they degrade the connective tissue mainly composed of type I collagen (or gelatine for denatured collagen) and the basement membrane mainly composed of type IV collagen [28]. AGE-RAGE interactions also induce the activation of phosphoinositol-3 kinase (PI3K) [27], oncogenic Ras, protein kinase C (PKC) and members of Rho/GTPase (Cdc42 and Rac-1) signalling pathways, which lead to cell survival, stress responses and apoptosis, release of growth factors and pro-inflammatory cytokines, and motility with changes in cell shape, respectively [29].

A meta-analysis study has reported that breast cancer rates are increased by 23% for women who are diagnosed with type 2 diabetes [30]. Breast cancer is a heterogeneous disease of particular interest. It is classified into different subtypes based on expression of hormone (oestrogen and progesterone) receptors and on its capability to invade, which defines the degree of the malignancy. Much attention has been devoted to the most aggressive phenotype of breast cancer with the identification of the roles of AGEs and the increase of RAGE expression in patient’s tissues [31]. However, biological effects of AGEs-RAGE on breast cancer cells are poorly investigated. Here, in an attempt to understand the biological effects of AGEs/RAGE in breast cancer, we investigated the effects of methylglyoxal-derived bovine serum albumin AGEs (MG-BSA-AGEs) on an invasive and non-hormone-dependent breast cancer cell line MDA-MB-231, with regard to cell proliferation, migration and invasion. The signalling pathways underlying the MG-BSA-AGE-mediated effects were also
investigated. To determine the consequences of glycation on MDA-MB-231 cell functions, the biological effects of non-modified BSA were also determined.

2. Materials and Methods

2.1. Reagents

All reagents were obtained from Sigma-Aldrich (Dorset, UK) unless mentioned otherwise.

2.2. Preparation of MG-BSA-AGEs

Bovine serum albumin (BSA) fraction V (10 mg/mL) was incubated with 0.1 M methylglyoxal (MG) in 0.1 M sodium phosphate buffer containing 3 mM sodium azide (pH 7.4) and left at 37°C for 72 hours. This glycated BSA was dialysed against distilled water to remove unbound sugars. Dialysis was performed by stirring the samples at 4°C with daily changes of distilled water until equilibrium was reached. Non-modified BSA underwent the same preparation conditions but in the absence of MG. The resulting MG-BSA-AGEs were characterized using a fluorescence spectrophotometer (Luminescence spectrometer model LS 30 from Perkin Elmer LAS Ltd, Buckinghamshire, UK) with emission at 420 nm after excitation at 350 nm, which confirmed the higher intensity of MG-BSA-AGEs than that of non-modified BSA. Bacterial endotoxins were removed from non-modified BSA and MG-BSA-AGE solutions using detoxi-gel endotoxin-removing gel columns (Thermo Scientific, Rockford, USA). Endotoxin levels in MG-BSA-AGEs and non-modified BSA solutions were measured with an E-toxate kit based on the Limulus Amebocyte lysate assay and were found to be below the detection limit (< 0.125 EU/mL). Protein concentrations were determined using a Bradford-based assay (Bio-Rad Laboratories, Hertfordshire, UK) with BSA as a standard.

2.3. Culture of MDA-MB-231 breast cancer cells

MDA-MB-231 breast cancer cell line was obtained from American Type Culture Collection (Manassas, VA, USA) and cultured in complete medium composed of Dulbecco's Modified Eagle’s medium (DMEM) supplemented with 10% heat-inactivated foetal bovine serum (FBS), 2 mM glutamine and antibiotics (100 μg/mL
streptomycin, 100 IU/mL penicillin) at 37°C in a saturated air humidity/5 % CO₂ incubator.

2.4. Cell proliferation assay

MDA-MB-231 cells (2.5 × 10⁴/mL) were seeded in complete medium in 24-well plates (Nunc™, Fisher Scientific, Loughborough, UK). After 4 hours of incubation to allow the cells to attach to the bottom of the well, the medium was changed to a serum-poor medium (SPM) supplemented with 2.5% FBS in the presence or absence of different concentrations (25 – 200 µg/mL) of MG-BSA-AGEs or non-modified BSA. Each condition was performed in triplicate. After 72 hours of incubation, the cells were detached in 250 µL of 0.05% trypsin / 0.02% EDTA, and each cell suspension was diluted in 10 mL of isotonic solution and counted using a Beckman-Coulter counter (Buckinghamshire, UK). Cell viability was assessed using the trypan blue exclusion method, which stains dead cells; whereas viable cells actively exclude the dye. The percentage of cell viability was determined according to the following formula: % viability = (viable unstained cells) / (viable unstained cells + dead stained cells) × 100.

2.5. Cell migration assay

MDA-MB-231 cells (5 × 10⁴/mL) were seeded in complete medium on Thermanox® plastic coverslips (Nunc™) in 24-well plates. After 24 hours of incubation, the cells reached pre-confluence on the coverslips; the culture medium was changed to SPM for a further 24-hour incubation. At confluence, cells were washed three times with sterile PBS, and each monolayer was wounded on two sides with a sterile razor blade, which formed two wound edges per coverslip with cell denuded areas. Dislodged cells and cell debris were removed with sterile PBS. The coverslips with wounded cell monolayers were returned to the well containing SPM with or without different concentrations of MG-BSA-AGEs and non-modified BSA for 24 hours of incubation. Each condition was performed in duplicate. At the end of the experiment, the cells were rinsed with PBS, fixed with 100% ethanol for 5 minutes and then left to dry at room temperature. The cells were stained with 0.1% methylene blue for 2 minutes and washed abundantly with distilled water to reveal the degree of wound recovery. Four pictures from the wound edge of each side were taken to assess the cell migration by
counting the number of the migrated cells and by measuring the distance of the cell migration using image J software (http://rsbweb.nih.gov/ij/index.html).

2.6. Cell invasion Matrigel™ assay

Growth-factor reduced Matrigel™ (Becton Dickinson, Oxford, UK) was diluted (1:6) in serum-free medium then poured onto the porous membrane of a Transwell® 24-insert plate (Nunc™). The plate was incubated for 30 minutes to allow the gel to polymerise. MDA-MB-231 cells were seeded at 10⁴ cells /100 μL in SPM (in the upper chamber of the Transwell®) in the presence or absence of different concentrations of MG-BSA-AGEs or non-modified BSA (in the well corresponding to the lower chamber of the Transwell®). Each condition was performed in duplicate. After 24 hours of incubation, the non-migrated cells on the upper side of the porous membrane were removed using a cotton swab soaked with PBS. The cells that migrated across the porous membrane were fixed with 4% paraformaldehyde and then stained with 0.1% Giemsa stain for cell counting using a Zeiss optical microscope.

2.7. Gelatine zymography

Matrix metalloproteinase (MMP) activities in the cell-culture media that had been used for the cell invasion assay (as aforementioned) were measured. The medium was collected and centrifuged (600 g for 15 minutes at 4°C). Protein concentration was determined using the Bradford protein assay (Bio-Rad), and the samples (100 μg of protein) were mixed with an equal volume of non-reducing sample buffer. The samples were incubated at room temperature for 10 minutes, after which they were subjected to electrophoresis on 7.5% sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) containing 1 mg/mL gelatine as a substrate. The gels were washed in re-naturation buffer (2.5% Triton X-100) for 30 minutes at room temperature to remove SDS and to re-nature the MMPs in the gels. The gels were then rinsed in activation buffer (50 mM Tris-HCl, 0.2 M NaCl, 5 mM CaCl₂ and 0.02% Brij 35) for 30 minutes to activate the MMPs. Gels were incubated overnight at 37°C with fresh activation buffer and stained with 0.5% Coomassie blue R-250 for 2 hours at room temperature. After destaining the gels, MMP gelatinase activity was detected as a white band on a dark background and quantified by densitometry using Image J software.
2.8. Preparation of cell lysates and Western blot analysis

MDA-MB-231 cells (6 × 10^5/2 mL) were seeded in complete medium in 6-well plates (Nunc™). After 48 hours of incubation, the medium was renewed with SPM for a further 24-hour incubation, and MG-BSA-AGEs or non-modified BSA was added for 5, 10, 30 and 120 minutes of incubation. After washing in cold PBS, all the intracellular proteins were extracted after lysing the cells with 120 µL/well of ice-cold radioimmunoprecipitation buffer (pH 7.5) containing 25 mM Tris-HCl, 150 mM NaCl, 0.5% sodium deoxycholate, 0.5% SDS, 1 mM EDTA, 1 mM sodium orthovanadate, 1 mM phenylmethylsulfonyl fluoride (PMSF), 1% Triton X-100 and 1 µM leupeptin.

For a time-course study of RAGE expression, the cells were treated with or without MG-BSA-AGEs or non-modified BSA for 10 minutes and 24-72 hours of incubation. The cells were then lysed with 80 µL/well of ice-cold buffer containing 10 mM Tris HCl (pH 7.4), 50 mM NaCl, 5 mM EDTA, 1% Triton X-100, 0.05% SDS, 50 mM sodium fluoride, 100 µM sodium orthovanadate, 10 mM beta-glycerophosphate, 10 mM sodium pyrophosphate, 100 µg/mL PMSF, 3 mM benzamidine, 1 mM dithiothreitol, 10 µM leupeptin, 5 µM pepstatin A and a cocktail of protease inhibitors. Protein lysates (100 µg) were mixed with 2× Laemmlli sample buffer, denatured by boiling in a water bath for 15 minutes and then centrifuged.

To remove the cell debris, the cell lysates were centrifuged at 20,000 g for 30 minutes at 4°C. The amount of proteins for each sample was determined by Bio-Rad protein assay and was fixed at 20-50 µg for equal protein loading. An equal volume of protein samples and 2× sample buffer were mixed in Eppendorf® tubes and then placed in the boiling water for 15 minutes. Samples were separated along with pre-stained molecular weight markers by 12% SDS-PAGE. Proteins were electroblotted onto nitrocellulose membranes and the membranes were blocked for 1 hour at room temperature in Tris-buffered saline (TBS)-Tween (pH 7.4) containing 1% BSA. Membranes were stained with the following primary antibodies diluted in the blocking buffer, overnight at 4°C on a rotating shaker: mouse monoclonal antibodies to phospho-extracellular signal-regulated kinase (p-ERK1/2, Tyr204 of ERK1, sc-7976; 1:1,000 dilution), rabbit polyclonal antibodies to total ERK1/2 [C-16] (t-ERK1/2, sc-93; 1:1,000 dilution) and mouse monoclonal antibodies to RAGE [E-1]...
(sc-74473; 1:1,000 dilution) provided by Santa Cruz Biotechnology (Heidelberg, Germany) and mouse monoclonal antibodies to GAPDH [6C5] (ab8245; 1:2,000 dilution) and to p70S6K1 [6B2] (ab119252; 1:1,000) and rabbit monoclonal antibodies to phospho-p70S6K1 (p-p70S6K1, Tyr412, ab78413; 1:1,000) by Abcam (Cambridge, UK). After washing five times for 10 minutes in TBS-Tween at room temperature, the membranes were stained with either rabbit anti-mouse or goat anti-rabbit horseradish peroxidase-conjugated secondary antibodies diluted in TBS-Tween containing 5% de-fatted milk (1:1,000 dilution) for 1 hour at room temperature with continuous mixing. After a further 5 washes in TBS-Tween, proteins were visualised using ECL chemiluminescent detection (Amersham Biosciences, Buckinghamshire, UK) and analysed using GeneSnap software with Gene tool image analyser (Syngene, Cambridge, UK).

2.9. RNA extraction and real-time RT-PCR
MDA-MB-231 cells (6 × 10⁵/2 mL) were seeded in complete medium in 6-well plates. After 4 hours of incubation, the medium was renewed with SPM in the presence or absence of 100 µg/mL MG-BSA-AGEs for 24-72 hours of incubation. Whole RNA extraction was performed using the RNeasy mini Kit (Qiagen Inc, Valencia, CA) in order to quantitatively monitor by real-time PCR the mRNA expression of human Rage (accession number AB036432) related to internal control house-keeping gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH, accession number DQ403057). Complementary DNA (cDNA) was produced from total RNA extracts via reverse transcription using Transcriptor first-strand cDNA synthesis kit (Roche Molecular Systems, Pleasanton, CA), and the reaction was carried out in Tetrad2 Thermal Cycler (Bio-Rad). The primer pair sequences used from Invitrogen (Life technologies, Paisley, UK) were: 5'-GGC TGG TGT TCC CAA TAA GG-3' and 3'-TCA CAG GTC AGG GTT ACG GTT C-5' for human Rage as described previously [21] and 5'-TGA TGA CAT CAA GAA GGT GAA G-3' and 3'-TCC TTG GAG GCC ATG TGG GCC AT-5' for human GAPDH as designed using the primer3 software. The real-time PCR was performed using a QuantiTect Reverse Transcription kit containing PCR SyberGreen Master Mix (Applied Biosystems, Grand Island, NY) and performed on 7500 real-time PCR system (Applied Biosystems). For each analysis, a negative control was prepared using all the reagents...
except the cDNA template. All the reactions were run in triplicate, and the fold-change of expression was calculated according to delta delta Ct (ΔΔCt) method as follows: fold-change = 2(ΔΔCt) with ΔΔCt = ΔCt (RAGE$_{treated}$ – GAPDH$_{treated}$) – ΔCt (RAGE$_{control}$ – GAPDH$_{control}$).

### 2.10. Flow cytometry analysis

MDA-MB-231 cells (6 × 10$^5$/2 mL) were seeded in complete medium in 6-well plates. After 48 hours of incubation, the medium was renewed with SPM for a further 24-hour incubation, and then 100 μg/mL MG-BSA-AGEs or non-modified BSA were added for a 10-minute incubation. Each condition was performed in triplicate. The cells were washed with PBS then scraped to maintain intact the structure of the transmembrane RAGE protein. After centrifugation (300 g for 10 minutes), 10$^6$ cells were re-suspended in 20 μL of PBS then 20 μg/mL of mouse anti-RAGE (E-1) antibody or 2 μg of mouse IgG1 isotype control (ab91353 from Abcam) were added. The mixture was then kept on ice for 45 minutes. The excess antibody was removed by washing the cells twice with PBS followed by centrifugation. After the second centrifugation, the cells were re-suspended in 20 μL of PBS, and 20 μg/mL of monoclonal anti-mouse antibody-FITC (Invitrogen, Paisley, UK) were added. The mixture was then kept on ice for 30 minutes. The RAGE expression level was measured using FACScan analysis software from FACSCalibur Flow Cytometer (Becton Dickinson).

### 2.11. Kinexus phospho-protein array analysis

MDA-MB-231 cells (6 × 10$^5$/2 mL) were seeded in complete medium in 6-well plates. After 48 hours of incubation, the medium was changed to SPM. After a further 24-hour incubation, 100 μg/mL MG-BSA-AGEs or non-modified BSA were added, and the cells which were then incubated for 10 minutes at 37°C. To determine the protein expression profile of signalling pathways downstream of RAGE, a phospho-protein array analysis (Kinetworks PhosphoSite Screen, KPSS-1.3) was performed by Kinexus Bioinformatics (Vancouver, Canada). Protein samples (500 μg) from MG-BSA-AGE- or non-modified BSA-treated and untreated cells were extracted according to the manufacturer’s instructions. The samples were used for a multi-immunoblotting assay based on SDS-polyacrylamide mini-gel electrophoresis with
20-lane multi-immunoblotters using different primary antibodies. Most of the multi-immunoblotting assays were performed at least twice. Protein expressions were visualised by chemiluminescence and relative expression was determined by Kinexus and expressed in counts per minute.

2.12. RAGE neutralization
To investigate whether MG-BSA-AGEs act through RAGE, the cells were treated with an anti-RAGE antibody to neutralize all the RAGE receptors. Briefly, MDA-MB-231 cells were seeded in complete medium in 24-well plates according to the protocols of upstream applications previously described. The medium was renewed with SPM, and then 20 µg/mL mouse monoclonal anti-RAGE [E-1] antibody or 20 µg/mL IgG1, which was used as an isotype control, were added. After 1 hour of incubation, the cells were treated with 100 µg/mL MG-BSA-AGEs for 10 minutes or a 24-72-hour stimulation at 37°C followed by RNA or protein extraction for real-time RT-PCR and Western blotting. The cells were also eventually subjected to all functional cell-based assays mentioned previously. Each condition was performed in triplicate, and each experiment was repeated three times.

2.13. Statistical analysis
Results are expressed as the mean ± standard deviation (SD). Experimental points were gathered for a minimum of three independent experiments. An unpaired Student's t-test was used for the comparison of two groups. A value of P < 0.05 was considered significant.

3. Results
3.1. MG-BSA-AGEs increase MDA-MB-231 cell proliferation
We first evaluated the effects of increasing concentrations of MG-BSA-AGEs and non-modified BSA varying between 25 µg/mL and 200 µg/mL on MDA-MB-231 cell proliferation (see Materials and Methods). We found that MG-BSA-AGEs affected cell number, in a dose-dependent manner, increasing then decreasing cell proliferation as a bell-shaped curve. A peak of stimulation (2.0-fold increase, p < 0.01) was attained in the presence of 50 and 100 µg/mL MG-BSA-AGEs, whereas the highest concentration (200 µg/mL) did not alter the cell growth compared with the untreated
cells used as control (Figure 1A). To estimate the specific effect of MG-BSA-AGEs on cell proliferation, non-modified BSA was also tested. Compared with the control, only a low concentration of BSA (50 µg/mL) slightly increased cell proliferation (1.25-fold, p < 0.05) whereas no effect was observed with other concentrations (Figure 1A). Furthermore, for each condition, the percentage cell viability was assessed by trypan blue exclusion and revealed a high cell viability between 97% and 85% from the control to the highest concentration of MG-BSA-AGEs or non-modified BSA (Figure 1B).

3.2. MG-BSA-AGEs increase MDA-MB-231 cell migration
To assess the effect of MG-BSA-AGEs on the migration of the breast cancer cell line MDA-MB-231, the cells were subjected to the wound-healing assay followed by a 24-hour treatment in SPM (which reduced the cell growth) with different concentrations (25-200 µg/mL) of MG-BSA-AGEs or non-modified BSA (Figure 2). Representative photomicrographs of cell migration in untreated condition used as control (Figure 2A), or treated conditions either with 50 µg/ml MG-BSA-AGEs (Figure 2B) or non-modified BSA (Figure 2C) are shown. Compared with the control (Figures 2A and 2D) and to non-modified BSA (Figures 2C and 2D), MG-BSA-AGEs at concentrations of 25-100 µg/mL increased the number of migrated cells (2.0-fold, p < 0.05, Figure 2D). MG-BSA-AGEs significantly increased the distance of cell migration in a dose-dependent manner with a peak at 50 µg/mL (2.0-fold increase, p < 0.01), compared with the control (Figure 2D). No change in cell migration was observed with the highest concentration of MG-BSA-AGEs (Figure 2D). Although non-modified BSA had no effect on MDA-MB-231 cell migration at most concentrations, treatment with the highest concentration (200 µg/mL) resulted in a significant inhibition (40% decrease, p < 0.05) in the distance migrated as compared with the control (Figure 2D).

3.3. MG-BSA-AGEs increase MDA-MB-231 cell invasion
The invasion of the basement membrane is a critical step during tumour progression because the basement membrane represents the last barrier for the cancer cells to reach the circulatory system and to be disseminated via metastasis. The effect of MG-
BSA-AGEs on the MDA-MB-231 cell invasion was estimated by counting the stained migrated cells that passed across the porous membrane after invading the Matrigel™, a reconstituted basement membrane (Figure 3). Representative photomicrographs of cell invasion in untreated condition considered as control (Figure 3A) or treated either with MG-BSA-AGEs (Figure 3B) or non-modified BSA (Figures 3C) are shown. Compared with the control (Figure 3A), MG-BSA-AGEs at 50 μg/mL and 100 μg/mL (Figure 3C) significantly increased MDA-MB-231 cell invasion (2.0-fold, p < 0.01) (Figure 3D). By contrast, MG-BSA-AGEs at 25 μg/mL and 200 μg/mL did not induce any significant changes in cell invasion, compared with the control. At all concentrations used, no effect of non-modified BSA on MDA-MB-231 cell invasion was observed, compared with the control (Figure 3D).

3.4. MG-BSA-AGEs enhance MMP-9 activity
Matrix metalloproteinases (MMP)-2 and MMP-9, also called gelatinases, play a central role in the degradation of the type IV collagen of the basement membrane, thus contributing to tumour invasion and metastasis. Compared with the MMP-9 activity in untreated MDA-MB-231 cell-condition medium (control) as assessed by gelatine substrate zymography, treatment with 50 and 100 μg/mL MG-BSA-AGEs significantly (p < 0.01) enhanced MMP-9 activity (1.5-fold and 2.1-fold, respectively, Figures 4A and 4B). A decrease in the activity of pro-MMP-9, which is the precursor and inactive form of MMP-9, was noticed when the cells were treated with 200 μg/mL MG-BSA-AGEs, compared with the control (Figures 4A and 4B). The activity of pro-MMP-2 was enhanced after treatment with MG-BSA-AGEs at all concentrations used; however, no active MMP-2 was observed (Figure 4).

3.5. MG-BSA-AGEs increase the phosphorylation of signalling proteins
To optimize incubation time corresponding to the maximal cell signalling induced by 100 μg/mL MG-BSA-AGEs (concentration previously shown to significantly increase all MDA-MB-231 cell functions), Western blotting was used to determine the expression levels of phospho-ERK1/2 (p-ERK1/2) in MDA-MB-231 cells treated with MG-BSA-AGEs for 5, 10, 30 and 120 minutes (Figure 5). MG-BSA-AGEs increased the phosphorylation of ERK1/2 in cells after 5 minutes of incubation (Figure 5). The maximum p-ERK1/2 expression level was reached at the 10-minute
incubation point. Expression levels of p-ERK1/2 slightly decreased after 30 minutes of incubation and nearly disappeared at 120-minute incubation point (Figure 5A). To confirm the effect of MG-BSA-AGEs on p-ERK1/2 expression level, MDA-MB-231 cells were treated with or without 100 μg/mL MG-BSA-AGEs or non-modified BSA for 10 minutes (Figure 5B). Compared with untreated cells (control), MG-BSA-AGE treatment increased the phosphorylation of ERK1/2 (1.8- and 2.8-fold increase for p-ERK1 and p-ERK2, respectively) while no change of ERK1/2 phosphorylation was observed in non-modified BSA-treated cells (Figure 5B).

A wider investigation of the downstream phosphorylated proteins of MG-BSA-AGEs/RAGE signalling pathways was proceeded as follows: the whole cell lysates of untreated MDA-MB-231 cells (control, Figure 5C1) or cells treated with 100 μg/mL MG-BSA-AGEs (Figure 5C2) and 100 μg/mL non-modified BSA (Figure 5C3) were analysed using the phospho-protein array analysis KPSS-1.3 for 35 phospho-proteins including p-ERK2 (Figure 5C). Like using Western blot analysis, quantification of the immuno-blots showed that MG-BSA-AGEs also resulted in a 3.0-fold increase of ERK2 phosphorylation, which reinforced the reproducibility and the validity of the data (Figure 5B). MG-BSA-AGEs increased the phosphorylation of the ribosomal protein serine S6 kinase (p70S6K)-beta 1 (25.4-fold increase), the signal transducer and activator of transcription (STAT)-3 (8.0-fold increase), MAPK p38 and glycogene synthase-serine kinase (GSK)-3α (5.6-fold and 3.2-fold, respectively), and MAPK/ERK protein-serine kinase 1/2 (MKK1/2, 4.6-fold increase), compared with untreated cells (Figure 5C4). Furthermore, the phosphorylation of c-Jun and its kinase JNK was increased (2.0-fold) in MG-BSA-AGE-treated cells as compared with untreated cells (control). The phosphorylation of the tumour-suppressor protein retinoblastoma-associated protein-1 (pRb) was also increased (2.6-fold) in MG-BSA-AGE-treated cells. Cell treatment with 100 μg/mL non-modified BSA had no significant effects on the phosphorylation of the proteins studied, compared with the control (Figure 5C).

3.6. MG-BSA-AGEs up-regulate RAGE expression
A time-course of expression levels of RAGE in MDA-MB-231 cells following treatment with 100 μg/mL MG-BSA-AGEs was established using Western blotting and real-time RT-PCR. MG-BSA-AGEs treatment led to up-regulation of RAGE
protein levels in a time-dependent manner (Figure 6A). Within 10 minutes, RAGE expression levels increased in MG-BSA-AGE-treated cells, compared with the basal level of RAGE expressed in untreated cells (Figure 6A). This data was further confirmed using flow cytometry analysis (Figure 6B). Western blot analysis revealed a peak of RAGE protein expression levels (3.0-fold increase, p < 0.0001) after 24 hours of incubation with MG-BSA-AGEs, whereas a peak of RAGE mRNA expression (5.0-fold increase, p < 0.001) was noticed after 48 hours of incubation (Figure 6C). After 72 hours of incubation with MG-BSA-AGEs, there was not significant difference in RAGE transcript and protein levels between MG-BSA-AGEs treated and untreated cells (Figure 6).

3.7. MG-BSA-AGEs stimulate MDA-MB-231 cells through RAGE

To check whether MG-BSA-AGEs stimulated MDA-MB-231 cells through their major receptor RAGE, the cells were pre-treated with either a neutralizing RAGE antibody to block all the RAGE receptors or with an isotype control IgG1 used as a negative control (Figure 7). RAGE neutralization process was applied to verify whether MG-BSA-AGEs induce the increased phosphorylation of signalling proteins such as ERK1/2 (the key protein of signalling pathways) and p70S6K1 (the most over-phosphorylated protein induced by MG-BSA-AGEs) through RAGE. The isotype control IgG1 did not affect ERK1/2 phosphorylation induced by MG-BSA-AGEs, compared with the basal expression level of p-ERK1/2 in untreated cells used as control (Figure 7A). However, 100 µg/mL BSA-AGEs did not enhance ERK1/2 phosphorylation in the cells pre-treated with neutralizing RAGE antibody (Figure 7A). In addition, the protein array analysis previously described was confirmed by a similar concomitant increase of p70S6K1 phosphorylation induced by MG-BSA-AGEs even in IgG1-treated cells, compared with the control (Figure 7B). The blockade of RAGE completely inhibited MG-BSA-AGE-induced p70S6K1 phosphorylation (Figure 7B). On the cell function aspects, the addition of MG-BSA-AGEs to IgG1-pretreated cells induced a significant increase in cell proliferation (1.8-fold, p < 0.001, Figure 7C), cell migration (2.0-fold, p < 0.05, Figure 7D) and cell invasion (1.6-fold, p < 0.05, Figure 7E) as compared with the untreated cells (control). However, the addition of MG-BSA-AGEs to cells pre-treated with anti-RAGE antibody did not change the cellular processes as compared with the control.
Altogether, MG-BSA-AGEs lost their biological effects after the blockade of RAGE, demonstrating that MG-BSA-AGEs act through RAGE.

4. Discussion

Diabetes and cancer are a cause for concern because of the high incidence of these diseases in Western countries and, more recently, in developing countries. A growing body of epidemiological evidence suggests a molecular link between diabetes and breast cancer [31, 32]. In diabetic patients, the most circulating and abundant advanced glycation endproducts (AGEs)-derived protein is serum glycated albumin [33]. In patients with breast cancer, serum concentrations of AGEs were higher than in healthy controls [32]. To date, the biological effects of glycated albumin on human breast cancer cells have been poorly described.

Here, we have shown that MG-BSA-derived AGEs increase the proliferation, migration and invasion (associated with an enhancement of MMP-9 gelatinase activity) of the invasive and non-hormone-dependent cell line MDA-MB-231 in vitro. Furthermore, MG-BSA-AGEs increase expression of RAGE and phosphorylation of ERK1/2 and other key signalling proteins such as p70S6K1. In addition, the blockade of RAGE using a neutralizing RAGE antibody removes all these MG-BSA-AGE effects. By contrast, non-modified BSA showed little biological effects on MDA-MB-231 cells. These new findings suggest that AGEs might contribute to the development and progression of invasive breast cancer. RAGE over-expression and the phosphorylation of key proteins induced by AGEs might represent promising targets for drug therapy to treat diabetic patients with breast cancer.

MG-BSA-AGEs increased MDA-MB-231 cell proliferation, migration and invasion, in a dose-dependent manner as shown by a bell-shaped curve, while non-modified BSA had no biological effects, except a slight stimulatory effect on cell proliferation at 50 µg/mL and an inhibitory effect on cell migration at 200 µg/mL. This slight pro-mitogenic effect of non-modified BSA is correlated with the induction of ERK1/2 phosphorylation observed at short incubation time (10 minutes) at the same concentration (50 µg/mL) in our pilot studies (data not shown), whereas no increase in phosphorylation of ERK1/2 was induced by 100 µg/mL of non-modified BSA. The pro-mitogenic effect of native BSA was also reported by Chung and colleagues using tubular epithelial cells in similar conditions [34]. This slight positive effect might be
attributed to a traceable amount of AGEs reported to be observed in native albumin, the most common serum protein known to be readily glycated in vivo. Indeed, native albumin is modified by 0.3 mol/mol of sugars and this is likely to be by glucose, the major metabolic sugar [35]. However, modification by other sugars such as methylglyoxal cannot be ruled out and may contribute towards AGE formation on native albumin. In contrast, the inhibitory effect of high concentration of native BSA on cell migration may be caused by the inhibition of cell adhesion, which plays a critical role in cell motility, due to the high adsorption of BSA between the culture medium and the solid surface as it was previously reported [36]. Here, bell-shaped curves representing the MDA-MB-231 cell response to MG-BSA-AGE biological effects are indicative of a bivalent bridging mechanism describing the dimerization/oligomerization of specific cell-surface receptors after binding with the ligand [37]. Indeed, throughout this present study, the maximal cell response was obtained with 50 and 100 µg/mL MG-BSA-AGEs, which might correspond to the optimal oligomerization of the AGE receptor such as RAGE to induce the maximal signalling. Therefore, the loss of the cell response seen at the highest concentration (i.e. 200 µg/mL) of MG-BSA-AGEs may be due to impediment of RAGE oligomerization. Furthermore, using an epitope-defined monoclonal antibody that specifically recognises receptor oligomerization, Xu and colleagues demonstrated the importance of RAGE oligomerization for the formation of active signalling complexes [38]. In addition, a recent study provides insights into the factors that influence AGEs-RAGE binding including glycation agent like methylglyoxal and the RAGE structure [39]. Indurthi et al. (2012) used 1 and 5 mM methylglyoxal to modify BSA respectively; they demonstrated that higher concentration of methylglyoxal produced more AGEs, which showed increased binding affinity to RAGE. They suggested that higher glycation generate more potential binding sites for RAGE in the protein surface [39]. In the present study, we applied 100 mM methylglyoxal to generate MG-BSA-AGEs to ensure a high binding potential to RAGE. Using siRNA technology, RAGE has been shown to play a critical role in the growth of human breast cancer cell lines, including MDA-MB-231 [21]. Most studies on BSA-AGEs, including this investigation, compare the biological effects of MG-BSA-AGEs with native non-modified BSA, which in agreement with previous work show no biological effects or little effects at the concentrations or experimental
conditions used [40]. Indeed, native globular BSA even bound to the cell surface rarely triggers specific transduction signalling unlike BSA-AGEs [41, 42]. Therefore, the mode of action of MG-BSA-AGEs is dependent on pathways mediated via RAGE rather than BSA-mediated pathways.

In this present study, we also showed that the enhancement of MDA-MB-231 cell invasive capacity induced by MG-BSA-AGEs was associated with MMP-9 gelatinase activity without changing MMP-2 gelatinase activity. Non-modified BSA had no effect on both MMP activities. Invasion and metastasis are key cell events resulting in tumour progression and are the hallmarks of cancer malignancy. During metastasis, cancer cells degrade the matrix by production and secretion of MMPs. MMPs facilitate cell migration and invasion across the connective tissue and the basement membrane to reach the blood circulation which is the main route of tumour dissemination towards vascularized organs. A study has shown that incubating human lung adenocarcinoma cell with glyceraldehyde-AGEs for 48 hours, promoted cell migration and invasion across Matrigel™ with enhancement of MMP-2 activity but not MMP-2 mRNA [43]. Recently, Noh and colleagues reported MMP-2 from body fluid as a putative biomarker in metastatic breast cancer [44]. However, Hallett and colleagues targeted MMP-9 mRNA in a mouse model using anti-MMP-9 DNAzyme transfected into MDA-MB-231 cells, and they reported a decrease of breast cancer and confirmed the key role of MMP-9 in breast cancer metastasis [45]. Furthermore, using a murine macrophage cell line, it has been shown that AGEs have regulated MMP-9 production via RAGE, which is in agreement with our present work, and was associated with activation of ERK1/2, p38 MAPK and NF-κB [46]. Therefore, our results suggest that MG-BSA-AGEs might promote breast tumour progression by increasing cancer cell migration, invasion and MMP-9 activity.

During MDA-MB-231 cell treatment with MG-BSA-AGEs, the cells displayed a maximum RAGE up-regulation after 24 hours of incubation. Recently, Shi and colleagues reported that AGEs up-regulate RAGE expression in various tissues, reaching a maximum after a 24-hour treatment, which facilitates the AGE-RAGE response by forming a positive feedback loop [41]. This suggests that RAGE up-regulation induced by MG-BSA-AGEs may contribute to the increased cellular response assessed as proliferation, migration and invasion. In addition, in endothelial progenitor cells, BSA-AGE-induced RAGE up-regulation has been demonstrated to
be ERK1/2 and p38 MAPK-dependent but not mediated by JNK MAPK [47]. We observed the peak up-regulation of RAGE mRNA expression level after 48 hours of treatment with MG-BSA-AGEs, whereas the protein expression level was lower, indicating a negative-feedback loop of the translational regulatory system of RAGE protein expression. To confirm this hypothesis, additional studies might be of interest on the key factors regulating the translational system of RAGE protein expression such as, microRNA and nuclear transcription factors p65/50 NF-κB and Sp-1/ERα, previously demonstrated to be involved in this regulatory process [48].

p-ERK1/2 is a phospho-protein widely known to play a central role in most cellular responses including cell proliferation, migration and invasion. We found that the optimal signal for MG-BSA-AGE-induced p-ERK1/2 expression in MDA-MB-231 cell line was obtained after 10-minute treatment. Similar early phosphorylation of ERK1/2 was reported to be induced by BSA-AGEs in various cell types [34]. We demonstrated that MG-BSA-AGEs enhanced phosphorylation of ERK1/2 and stimulated all cellular functions through RAGE, as these processes were attenuated after neutralization of the receptor using a specific monoclonal antibody. Thus, this result eliminates the involvement of other AGE receptors such as CD36, scavenger receptors class A type II, class B type I and AGE receptors 1, 2 and 3, which are mainly implicated in the detoxification of AGES rather than in signalling processes [49].

Although 100 μg/mL non-modified BSA did not change the phospho-proteome profile of MDA-MB-231 cells compared to untreated cells, at a similar concentration, MG-BSA-AGEs concomitantly increased the phosphorylation of p70S6K1, STAT-3, p38 MAPK, GSK-3 and MKK1/2. The p70S6K1 has been demonstrated to play a central role in tumour growth by regulating pro-angiogenic effects and protein synthesis of insulin [50, 51]. In addition, targeting p70S6K1 gene expression by micro-RNA 145 expression has been reported to inhibit tumour growth [52]. Constitutively active STAT proteins, which are critical in the regulation of cell cycle and cell growth, are found in various types of tumours including breast cancer [53]. Previous studies have reported p38 MAPK as a central kinase in a common intracellular signalling pathway that plays an essential role in human breast cancer cell migration and invasion by modulating the expression and activity of MMPs [54]. Furthermore, several lines of evidences suggest a link between p38 MAPK activation
and MMP-9 expression [55]. In some reports, the use of pharmacological inhibitors of GSK-3 and of p38 MAPK, which are proteins with increased phosphorylation observed in our study, in MDA-MB-231 cells and in a model mouse of breast cancer demonstrated the key requirement of these proteins in breast cancer development [56, 57]. Here, we also showed that MG-BSA-AGE-induced p70S6K1 over-phosphorylation was prevented after RAGE neutralization, which further removes all the biological effects of MG-BSA-AGEs-induced MDA-MB-231 cell proliferation, migration and invasion. Thus, altogether, targeting either the expression of phospho-proteins aforementioned such as p70S6K1 or targeting RAGE expression might prevent the progression and development of breast cancer in diabetic patients.

In conclusion, as a consequence of BSA glycation, this study demonstrated the stimulatory effects of MG-BSA-AGEs on cell proliferation, migration and invasion (with an enhancement of MMP-9 activity) through RAGE, in an invasive and non-hormone-dependent breast cancer cell line MDA-MB-231. Among the phospho-proteins involved in MG-BSA-AGE-induced signalling pathways, we also highlighted a concomitant increase in the phosphorylation of p70S6K1, which might be a new biomarker of the invasive subtype of the breast cancer and a promising target drug like RAGE to fight against breast cancer in diabetic patients.

Acknowledgements
We are grateful to the Ministry of Health, Saudi Arabia for providing a PhD scholarship to Hana Sharaf that enabled her to undertake this study. We would like to thank Dr Nasser Al-Shanti, School of Healthcare Science at Manchester Metropolitan University for his help with flow cytometry analysis and Tor Yip for his assistance with the illustrations.

References


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Figure legends

Figure 1: Effect of MG-BSA-AGEs and non-modified BSA on (A) the proliferation and (B) the viability of breast cancer MDA-MB-231 cells. Cells were treated for 72 hours of incubation with 25-200 μg/mL MG-BSA-AGEs or non-modified BSA. Cell number (A) and cell viability (B) were determined using an automatic cell counter and the trypan blue exclusion method, respectively. Control indicates non-treated cells. Results are presented as mean ± SD of three independent experiments. (*), (**), and (***) signify a statistically significant difference (p < 0.05, p < 0.01 and p < 0.001), compared with the control.

Figure 2: Effect of MG-BSA-AGEs and non-modified BSA on the migration of breast cancer MDA-MB-231 cells. Representative photomicrographs (×400 magnification) showing MDA-MB-231 cell migration measured from the wound edge
(indicated by the arrow) after 24 hours of incubation in (A) non-treated (control) or (B) 50 μg/mL MG-BSA-AGE-treated or (C) non-modified BSA-treated cells. Scale bar = 100 μm. (D) Quantification of the number of migrated cells (black bars) and the distance of migration (grey bars) of MDA-MB-231 cells treated in the aforementioned conditions. Results are presented as mean ± SD of three independent experiments. Results are presented as mean ± SD of three independent experiments. (*) and (**) signify a statistically significant difference (p < 0.05 and p < 0.01), compared with the control.

**Figure 3: Effect of MG-BSA-AGEs and non-modified BSA on the invasion of breast cancer MDA-MB-231 cells.** Representative photomicrographs (×100 magnification) of (A) untreated MDA-MB-231 cells (control), (B) cells treated with 100 μg/mL MG-BSA-AGEs and (C) non-modified BSA in the Matrigel™ invasion assay. Stimulated cells which had invaded the Matrigel™ (a reconstituted basement membrane) and migrated across the porous membrane were stained with Giemsa dye for counting. (D) Quantification of number of invaded MDA-MB-231 cells after treatment with 25-200 μg/mL MG-BSA-AGEs or non-modified BSA, compared with the control. Results are presented as mean ± SD of three independent experiments. (**) signifies a statistically significant difference (p < 0.01), compared with the control.

**Figure 4: Effect of MG-BSA-AGEs on MMP-9 and MMP-2 activities produced by breast cancer MDA-MB-231 cells during the Matrigel™ invasion assay.** (A) Representative gelatine zymographic analysis showing the activity of MMP-9 and MMP-2 from cell conditioned media collected after Matrigel invasion assay by MDA-MB-231 cells treated with 25-100 μg/mL MG-BSA-AGEs, compared with untreated cell-condition medium (control, c). The gels revealed gelatinolytic activities of pro- and active MMP-9 and pro-MMP-2. (B) Quantitative gelatinolytic activity of pro- and active MMP-9 and (C) pro-MMP-2 expressed as relative activity, which was calculated as a ratio of the control. Results are presented as mean ± SD of four independent experiments. (*), (**) and (***) signify a statistically significant difference (p < 0.05, p < 0.01 and p < 0.001), compared with the control.
Figure 5: Effect of MG-BSA-AGEs on the phosphorylation of signalling proteins in MDA-MB-231 cells. (A) Representative Western blot analysis showing phosphorylation of ERK1/2 (p-ERK1/2) induced by 100 μg/mL MG-BSA-AGEs after different incubation times varying from 5 to 120 minutes, compared with untreated cells (control). (B) Representative Western blot analysis (B1) showing the effect of 100 μg/mL MG-BSA-AGEs or non-modified BSA on the phosphorylation of ERK1/2 expression, compared with the control. (B2) Relative phosphorylation of ERK1/2 calculated as a ratio to the total ERK1/2 (t-ERK1/2), the loading control. Results are presented as mean ± SD of three independent experiments. (**) signifies a statistically significant difference (p < 0.01), compared with the control. (C) Multi-immunoblotting of 35 phospho-proteins expressed in untreated (C1) MDA-MB-231 cells, cells treated with 100 μg/mL MG-BSA-AGEs (C2) or cells treated with non-modified BSA (C3) after 10 minutes of incubation. (C4) The bar graph shows the relative expression of the relevant phospho-proteins expressed in counts per minute and calculated as a ratio to the untreated cells (control).

Figure 6: Time-course of RAGE protein and mRNA expression in breast cancer MDA-MB-231 cells treated with MG-BSA-AGEs. (A) Representative Western blot analysis (A1) showing the effect of 100 μg/mL MG-BSA-AGEs on RAGE expression in MDA-MB-231 cells after incubation for 10 minutes or 24, 48 or 72 hours, compared with untreated cells (control). Bar graph (A2) showing the relative expression of RAGE calculated as a ratio to GAPDH expression, the loading control. (B) Representative FACS analysis (B1) of the RAGE up-regulation induced by 100 μg/mL MG-BSA-AGEs after 10 minutes of incubation. IgG1 was used as an isotype control. Bar graph (B2) indicating the quantification of the percentage of RAGE-positive cells calculated as a ratio to isotype control. (C) Relative expression of RAGE mRNA as determined using real-time PCR analysis. Results are presented as mean ± SD of three independent experiments. (*) and (**) signify a statistically significant difference (p < 0.05 and p < 0.01), compared with the control.

Figure 7: RAGE mediates the effects of MG-BSA-AGEs in breast cancer MDA-MB-231 cells. Representative Western blot analysis showing the loss of MG-BSA-
AGE-induced phosphorylation of (A) ERK1/2 and (B) p70S6K1 after the blockade of RAGE by an anti-RAGE antibody, compared with untreated cells (control) and an isotype control IgG1. Bar graphs show the relative expression levels of (A2) p-ERK1/2 and (B2) p-p70S6K1 calculated as a ratio to the total ERK1/2 (t-ERK1/2) and total p70S6K1 (t-p70S6K1), the respective loading controls. Loss of MG-BSA-AGE-induced (C) cell proliferation, (D) migration and (E) invasion after RAGE blockade using a neutralizing blocking anti-RAGE antibody, compared with untreated cells (control) and an isotype control IgG1. Results are presented as mean ± SD of three independent experiments. (*) and (**) signify a statistically significant difference (p < 0.05 and p < 0.01), compared with the control.
Figure 1

A

Cell number ($\times 10^{6}$)

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<th>25</th>
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<tr>
<td>Control</td>
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<tr>
<td>MG-BSA-AGEs (μg/mL)</td>
<td>1±</td>
<td>2±</td>
<td>3±</td>
<td>2±</td>
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<tr>
<td>BSA (μg/mL)</td>
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B

Cell viability (%)

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<tr>
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<td>BSA (μg/mL)</td>
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Figure 2
Figure 3

A

B

C

D

Number of invaded cells

Control 25 50 100 200 25 50 100 200

MG-BSA-AGEs (µg/mL) BSA (µg/mL)

** **
Figure 4

A

B

C

MG-BSA-AGES (µg/mL)
Figure 5

A

Control  MG-BSA-AGEs (100 µg/mL)

RAGE  48h 72h 10kDa

GAPDH  26kDa

B

B1

Control IgG alone

B2

Control untreated cells  MG-BSA-AGE-treated cells

C

Relative expression of RAGE mRNA

Control  MG-BSA-AGEs (100µg/mL)

24 h incubation  48 h incubation  72 h incubation
Figure 7

A

MG-BSA-AGES (100 μg/mL)

+ - IgG1 (20 μg/mL)

C - + anti-RAGE Ab (20 μg/mL)

p-ERK1/2 44/42 kDa

t-ERK1/2 44/42 kDa

B

MG-BSA-AGES (100 μg/mL)

+ - IgG1 (20 μg/mL)

C - + anti-RAGE Ab (20 μg/mL)

p-p70S6K1 70 kDa

t-p70S6K1 70 kDa

C

Cell number (x 10^6)

Control IgG1 anti-RAGE Ab

MG-BSA-AGES (100 μg/mL)