N-Acetylcysteine Amide Protects Against Oxidative Stress–Induced Microparticle Release From Human Retinal Pigment Epithelial Cells

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PURPOSE. Oxidative stress is a major factor involved in retinal pigment epithelium (RPE) apoptosis that underlies AMD. Drusen, extracellular lipid- and protein-containing deposits, are strongly associated with the development of AMD. Cell-derived microparticles (MPs) are small membrane-bound vesicles shed from cells. The purpose of this study was to determine if oxidative stress drives MP release from RPE cells, to assess whether these MPs carry membrane complement regulatory proteins (mCRPs: CD46, CD55, and CD59), and to evaluate the effects of a thiol antioxidant on oxidative stress–induced MP release.

METHODS. Retinal pigment epithelium cells isolated from human donor eyes were cultured and treated with hydrogen peroxide (H₂O₂) to induce oxidative stress. Isolated MPs were fixed for transmission electron microscopy or processed for component analysis by flow cytometry, Western blot analysis, and confocal microscopy.

RESULTS. Transmission electron microscopy showed that MPs ranged in diameter from 100 to 1000 nm. H₂O₂ treatment led to time- and dose-dependent elevations in MPs with externalized phosphatidylserine and phosphatidylethanolamine, known markers of MPs. These increases were strongly correlated to RPE apoptosis. Oxidative stress significantly increased the release of mCRP-positive MPs, which were prevented by a thiol antioxidant, N-acetylcysteine amide (NACA).

CONCLUSIONS. This is the first evidence that oxidative stress induces cultured human RPE cells to release MPs that carry mCRPs on their surface. The levels of released MPs are strongly correlated with RPE apoptosis. N-acetylcysteine amide prevents oxidative stress–induced effects. Our findings indicate that oxidative stress reduces mCRPs on the RPE surface through releasing MPs.

Keywords: microparticles, N-acetylcysteine amide, membrane complement regulatory proteins, AMD

Age-related macular degeneration (AMD) is the leading cause of blindness in the elderly affecting tens of millions of people worldwide, and is the most common cause of vision loss in the elderly in the United States. The dry form of AMD accounts for 80% to 90% of all AMD cases.1–4 The hallmarks of dry AMD are the appearance of drusen, and apoptosis of the retinal pigment epithelium (RPE).5–9 Drusen are extracellular lipid- and protein-containing deposits that accumulate mainly between the RPE and Bruch’s membrane. Drusen are a hallmark of aging and early AMD, as well as a risk factor for developing late AMD.10–14 Although understanding of the mechanisms of AMD has increased, there is neither a cure nor means of prevention for AMD. Therefore, there is a critical need to identify new mechanisms for AMD to develop preventive and therapeutic strategies for this age-related blinding disease.

The deposition of drusen between Bruch’s membrane and the RPE layer impairs the RPE cell layer through obstruction of nutrient flow to and from the choroid. It is believed that drusen are formed from cellular materials originating from RPE cells. The suggested mechanisms for drusen formation include cellular budding during apoptosis and exosomal trafficking during autophagy.15 An additional possible mechanism that has yet to be explored is through the generation of microparticles (MPs) by RPE cells.

Cell-derived MPs are small membrane-bound vesicles that range in diameter from 100 to 1000 nm. They are released from activated, injured, and apoptotic parental cells into the extracellular space.17–21 Microparticles shed by other organ systems under pathologic conditions have been noted to contain parental cell components and are capable of potentiating the disease state.17,22–25

The retina is highly susceptible to oxidative stress, which is recognized as a major risk factor for AMD.13,26–27 Retinal pigment epithelium cells spend their life recycling the lipid-rich photoreceptor outer segments, which requires an abundance of antioxidants and repair systems to maintain homeostasis.13,26 Disruption of this balance may lead to RPE dysfunction, cell death, and inflammation.5,13,26–28

Complement activation is currently thought to play an important role in AMD development.14,29 Membrane complement regulatory proteins (mCRPs: CD46, CD55, and CD59) provide normal cells with a first line of defense against complement attack, and have been identified on human RPE cells in culture30–33 and in situ.34–36 Oxidative stress leads to...
the loss of mCRPs on RPE cells. However, mechanisms leading to the loss of mCRPs on RPE cells under oxidative stress or during early AMD remain unknown. Therefore, we hypothesized that oxidative stress–induced loss of mCRPs on RPE cells is through the release of MPs that carry mCRPs.

N-acetylcysteine amide (NACA; also termed AD4), the amide form of N-acetylcysteine (NAC), is a thiol antioxidant with enhanced properties of lipophilicity, membrane permeability, and antioxidant capacity when compared with NAC. Emerging evidence provides strong support for NACA as a protective agent against oxidative stress under numerous pathologic conditions in vitro and in vivo, including a mouse model of retinal degeneration.

In this study, we investigated if oxidative stress drives MP release from RPE cells, assessed whether these released MPs carry mCRPs, and evaluated the effects of NACA, a thiol antioxidant, on oxidative stress–induced MP release.

**Materials and Methods**

**Materials**

Cell culture dishes, flasks, Falcon Primaria 24-well plates (Becton-Dickinson, Inc., Lincoln Park, NJ, USA), and fetal bovine serum (FBS) were purchased from Fisher Scientific (Pittsburgh, PA, USA). Phenol red-free Dulbecco’s modified Eagle’s medium/F12 (DMEM/F12), penicillin G, streptomycin sulfate, hydrogen peroxide (H2O2), and NACA were purchased from Sigma-Aldrich Corp. (St. Louis, MO, USA). Dulbecco’s phosphate-buffered saline (D-PBS), Hanks’ balanced salt solutions (HBSS), fungizone, and Dead Cell Apoptosis Kit with Annexin V Alexa Fluor 488 and propidium iodide (PI) were obtained from Life Technologies, Haematologic Technologies (Essex, CT, USA). Isolated MPs were grown in Falcon Primaria 24-well plates with phenol-free DMEM/F12 containing 20% FBS, penicillin G (100 U mL⁻¹), streptomycin sulfate (100 µg mL⁻¹), and amphotericin B (0.25 µg mL⁻¹) at 37°C in a humidified incubator under 5% CO2. Cells were then subcultured into cell culture dishes and grown in phenol red-free DMEM/F12 containing 10% FBS with the same antibiotics as described above. In all experiments, simultaneous, parallel assays were performed on fourth to ninth passaged RPE cells seeded at the same time and density from the same parent cultures. All experiments were repeated at least three times.

For H2O2 treatment, RPE cells were treated with 500 µM H2O2 for 2 to 24 hours or 50 to 2000 µM H2O2 for 16 hours in serum-free and phenol-free DMEM/F12. Cells without H2O2 treatment serve as controls. Prior to experiments, cells were switched to MP-free medium to avoid contamination from endogenous MPs present in the serum. In some cases, RPE cells were preincubulated with 1 mM NACA for 8 hours prior to H2O2 exposure.

**Microparticle Isolation and Characterization**

Microparticle-free media, prepared by filtration through 0.1-µm filters, were used for all experiments. In each experiment, MPs were simultaneously isolated from vehicle- and H2O2-treated RPE cells by differential centrifugation and microfiltration as follows. After cell conditioned media were collected, cells were incubated with 0.05% trypsin/0.53 mM EDTA for 5 minutes, after which trypsin was neutralized with MP-free complete media. Cell suspensions were collected and pooled with the conditioned media that were collected prior to trypsinization. After centrifuging for 5 minutes at 500g at 4°C, cells were collected for flow cytometry analysis. Supernatants were collected and centrifuged at 1500g for 15 minutes at 4°C to remove cell debris. Each of the supernatants was collected and passed through a 1.2-µm filter to remove any larger extracellular vesicles, such as apoptotic bodies. Supernatants were then centrifuged at 20,000g for 30 minutes at 4°C. The pellets were resuspended, washed in D-PBS, and centrifuged for a total of three times. Isolated MPs were then processed for transmission electron microscopy (TEM), flow cytometry, Western blot analysis, or confocal microscopy as described below.

**Transmission Electron Microscopy**

Isolated MPs were fixed with 4% paraformaldehyde for 1 hour, washed in D-PBS, and centrifuged at 20,000g for 30 minutes, after which the pellet was resuspended in water and negative stained with 1% uranyl acetate for 1 minute. Samples were imaged with an AMT camera (Advanced Microscopy Techniques, Woburn, MA, USA) on a Philips-CM 100 (Philips, Andover, MA, USA) or JEOL JEM 1400 TEM (JEOL, Peabody, MA, USA) at the University of Michigan Microscopy and Image Analysis Core Facility.

**Flow Cytometry**

Isolated MPs were stained with the following antibody-fluorophores in varying combinations with compensation and IgG controls used where necessary: annexin V-FITC, annexin V-PE, PI, CD46-APC, CD55-PE, CD59-APC. Milk fat globule-epidermal growth factor (EGF) factor 8 (MFG-E8)-FITC,
and duramycin-FITC (Supplementary Table S1). Controls for IgG1 and IgG2a conjugated to APC were used. In some cases, MPs were exposed to 16 μM, 100-fold excess compared with MFG-E8, crGD for 5 minutes prior to staining with MFG-E8-FITC. Annexin V and PI staining was performed at room temperature for 15 minutes per the manufacturer’s instructions while all other staining was performed on ice for 1 hour. Samples were run on a LSR II flow cytometer (BD Biosciences, San Jose, CA, USA; Becton Dickinson) equipped with 450, 488, and 633 nm lasers with a side-scatter threshold set to 750. Acquisition was performed with BD FACSDiva software. The injection port was wiped and water was run through the cytometer between samples to minimize cross-contamination of samples. FlowJo version 10 (FlowJo, LLC, Ashland, OR, USA) was used to analyze and quantify data.

**Confocal Microscopy**

Ten microliters of MFG-E8-FITC stained and washed samples for flow cytometry, prior to being diluted for flow cytometry, were pipetted onto a standard slide, coverslipped, and sealed with nail polish. Samples were imaged on a Leica SP5 confocal microscope (Leica Microsystems CMS GmbH, Wetzlar, Hesse, Germany) using a ×63 oil immersion lens, ×10 digital magnification, and a 488-nm laser.

**Cell Death Detection**

**Flow Cytometry of Cell Death.** Retinal pigment epithelial apoptosis and necrosis were evaluated by Dead Cell Apoptosis Kit with Annexin V Alexa Fluor 488 and PI (Life Technologies) by flow cytometry, using the same setup mentioned above, according to procedures outlined by the manufacturer. FlowJo version 10 was used to analyze and quantify data.

**TUNEL Assay.** Retinal pigment epithelial cells grown on sterile coverslips were treated with 0 to 2000 μM H2O2 for 16 hours. The coverslips were washed and stained with PI (0.15 mM) for 15 minutes at room temperature. After three washes, coverslips were fixed and subjected to TUNEL assay using the cell death detection kit (In Situ Cell Death Detection Kit, Cat#: 11664817910; Roche Applied Science, Indianapolis, IN, USA) according to the manufacturer’s protocol. Finally, the coverslips were washed three times with PBS, mounted on slides using VECTASHIELD antifade mounting medium with 4',6-diamidino-2-phenylindole (DAPI; Vector Laboratories, Burlingame, CA, USA). Cells were viewed with an epifluorescence microscope (model E800; Nikon, Melville, NY, USA). Digital images were collected with a cooled charge-coupled device (CCD) camera and the allied software (ACT; Nikon). Percentage of TUNEL-positive and PI-positive cells was quantified with ImageJ software using the cell counter plug-in. Representative images were obtained by merging single images of DAPI (blue), PI (red), and TUNEL (green) from the same field of view using ImageJ software.

**Cellular GSH Measurement**

Glutathione (GSH), also known as γ-glutamyl cysteinyglycine, is an important tripeptide thiol antioxidant. Its intracellular concentrations were measured by a Glutathione Cell-Based Detection Kit (Blue Fluorescence; Cat#:640360; Cayman Chemical Company, Ann Arbor, MI, USA) following procedures outlined by the manufacturer. Briefly, a cell-permeable dye (monochlorobimane) was used to react with cellular GSH to generate a highly blue fluorescent product that was measured with a LSR II flow cytometer using a 405 nm laser and 450/50 bandpass filter. Data were quantified using FlowJo version 10.

**Western Blot Analysis**

Western blot analysis was performed using the technique described previously11 with modifications. Briefly, intact cells or isolated MPs were lysed with RIPA Lysis Buffer System (Cat#: sc-24948; Santa Cruz Biotechnology) containing 150 mM NaCl, 25 mM Tris- HCl (pH 7.4), 2 mM EDTA, 1.0% Triton X-100, 1.0% sodium deoxycholate, 0.1% SDS, 1 mM sodium orthovanadate, 2 mM phenylmethylsulfonyl fluoride (PMSF), and complete protease inhibitor cocktail. Lysed samples were centrifuged at 14,000g for 10 minutes at 4°C and the supernatants collected. The protein content of each sample was determined by a bicinchoninic acid (BCA) assay kit (Cat#: BCA1; Sigma-Aldrich Corp.), with bovine serum albumin used as a standard. Proteins of whole-cell lysates or of MP lysates were solubilized in Laemmli sample buffer (62.5 mM Tris [pH 6.8], 25% glycerol, 2% SDS, 0.01% bromophenol blue; Bio-Rad, Hercules, CA, USA) without β-mercaptoethanol (nonreducing conditions), heated at 95°C for 5 minutes, cooled on ice, and then applied to 4% to 20% linear gradient long-life TGX (Tris-Glycine eXtended) gels (Cat#: 4561094; Bio-Rad). After electrophoresis, proteins were transferred to a low fluorescence polyvinylidene difluoride (PVDF) membrane (Cat#: IPFL00010; EMD Millipore Immobilon PVDF Transfer Membranes; Billerica, MA, USA), and immunoblotted with primary antibody against CD46 (2 μg/mL), CD59 (2 μg/mL), CD63 (2 μg/mL), β-actin (1:3000 dilution), or glyceraldehyde-3-phosphate dehydrogenase (GAPDH; 1 μg/mL; Supplementary Table S2) overnight in the cold room with constant rocking. Primary antibodies were visualized with fluorescently conjugated secondary antibody and an 473-nm wavelength was used for excitation and a greater than or equal to 510-nm long-pass (LPR) filter was used for emission acquisition. Alexa Fluor Fluor 488-conjugated secondary antibody, a laser with 473-nm wavelength was used for excitation and a greater than or equal to 665-nm long-pass (LPR) filter was used for emission acquisition. Fluorescent Western blotting images were processed and merged with ImageJ software.

**Statistical Analysis**

Data are presented as mean ± SD, and analyzed using GraphPad Prism version 5 (GraphPad Software, Inc., San Diego, CA, USA). Student’s t-test or 1-way ANOVA with Sidak’s multiple comparisons test was used to compare groups. Correlation between PS-positive MPs versus RPE apoptosis was determined using Pearson’s correlation test. Serum-free samples were used as negative controls, termed control in all plots, and subtracted out from the H2O2-treated samples when normalized to control is stated. P values less than 0.05 were considered as statistically significant.

**RESULTS**

**Characterization of RPE-Derived Microparticles**

It is well established that H2O2 triggers RPE cell death by induction of the apoptotic signaling cascade through oxidative stress. To evaluate whether oxidative stress induces MP release from RPE cells, we treated cultured human RPE cells derived from a 52-year-old or a 61-year-old male Caucasian donor with 500 μM H2O2 or vehicle (control) for 16 hours with or without NACA (a thiol antioxidant). Microparticles released from RPE cells were characterized by TEM, the gold standard for vesicle...
size determination, and by flow cytometry, the most common method for analysis of MPs. Figure 1A shows representative TEM images and Figure 1B shows a quantitative histogram displaying the distribution of MP sizes in all four groups. As shown in Figure 1, MPs released from RPE cells are heterogeneous and range in diameter from 100 to 1000 nm with no statistical difference in size among the four groups.

To characterize MPs by flow cytometry, we first used fluorescent beads of known sizes (0.5 and 1 μm) and confirmed that a LSR II flow cytometer detected events within the size range of MPs (Fig. 2A). A gate encompassing the bead populations was set and maintained across all experiments. The majority of events in MP samples isolated from both control and H2O2-treated cells with or without NACA fell within the gate in a similar pattern to the distribution of the one-half micron beads, suggesting that we detected the MPs visualized with TEM. Total microparticles based on light-scattering measurements were significantly increased in H2O2-treated samples. This increase was reversed by pretreatment of RPE cells with NACA, a thiol antioxidant (Fig. 2B; P < 0.05).

A hallmark of MPs is phosphatidylserine (PS) externalization, which results from the loss of asymmetry of the cell membrane during apoptosis. Samples obtained from both control and H2O2-treated RPE cells demonstrated positive annexin V staining, suggesting that PS is exposed on the surface of MPs released from RPE, we stained isolated MPs with annexin V(630 nm). MFG-E8 has been suggested to bind PS more effectively than annexin V(630 nm). Therefore, we used fluorescently (FITC) labeled MFG-E8 to stain PS on MPs (Fig. 3A; lower panels). As shown in Fig. 3A, MFG-E8 was superior to annexin V for the staining of PS on MPs, especially for the MPs released from control RPE cells. Fig. 3B shows summarized data by using two different PS-binding proteins, annexin V, and MFG-E8, to label MPs isolated from control and H2O2 (500 μM) for 16 hours- and treated RPE cells. Compared with annexin V, MFG-E8 detected more PS-positive MPs from control RPE cells (4-fold increase; P < 0.001; Fig. 3B). Despite this, H2O2 treatment still significantly increased the number of PS-positive MPs detected by either annexin V (6-fold increase, P < 0.001) or MFG-E8 (1.78-fold increase, P < 0.002). Due to the fact that MFG-E8 can also bind to integral receptors (ζ3β1 or δβ1), we used a cyclic RGD peptide to block ζ3β1 or δβ1 integrin receptors possibly present on the MPs. Preincubation with cyclic RGD did not affect the percentage of PS-positive MPs, as measured by MFG-E8-FITC (Fig. 3C), suggesting that the increase in MFG-E8 binding to the surface of MPs was due to increased affinity for PS and not an additional binding site.

In addition to characterization of MPs by TEM and flow cytometry, we performed Western blot analysis to test whether CD63, an exosomal marker, was present on the MPs. As shown in Supplementary Figure S1, CD63 was detected in whole-cell lysates, but not in the lysates prepared from MPs, therefore ruling out contamination with exosomes.

Taken together, our data have demonstrated that the small particles released and prepared under our experimental conditions are MPs (100–1000 nm), but not exosomes (30–100 nm).

Oxidative Stress Induces Microparticle Release in a Dose- and Time-Dependent Manner

To further investigate the role of oxidative stress in MP release from RPE cells we performed time- and dose-dependent studies. Treatment with H2O2 causes both dose- and time-dependent increases in the percent of PS-positive MPs, as measured by MFG-E8-FITC, with peaks at 200 μM and 16 hours (Figs. 4A, 4B).

Another aminophospholipid suggested to be present on the surface of MPs is phosphatidylethanolamine (PE), which can be targeted by duramycin. To see whether PE is externalized at the surface of MPs released from RPE, we stained isolated MPs from both control and H2O2-treated RPE cells with duramycin-FITC. Similar to PS, H2O2 induces PE-positive MP release in a dose- and time-dependent manner with peaks at 500 μM and 16 hours (Figs. 4C, 4D).

As H2O2 is able to drive both cellular apoptosis and necrosis, we performed TUNEL assay to confirm that MPs are being released primarily from apoptotic RPE cells (Supplementary Fig. S2). At 16 hours, serum free (control) and 50 μM H2O2 showed no signs of TUNEL- (green) or PI- (red) positive cells. Minor apoptosis (10%) was detected at 100 and 200 μM H2O2 with no necrosis. At 500 μM H2O2 there was significant apoptosis (80%) with minor necrosis (7%). There were increasing levels of necrosis in the 1000 μM H2O2 (35%) and 2000 μM H2O2 (50%) treatment groups. As 500 μM H2O2 for 16 hours constantly released higher MPs, we used 500 μM H2O2 for 16 hours for most of the data presented in this study.
Microparticle Release Correlates With RPE Apoptosis

In other organ systems, cellular apoptosis leads to the release of MPs. Furthermore, oxidative stress induces RPE apoptosis. To test whether MP release correlates with RPE apoptosis, we evaluated both RPE apoptosis, by annexin V and PI double staining, and released PS-positive MPs, by MFG-E8 staining, over time. As shown in Fig. 5, released PS-positive MPs strongly correlated with RPE apoptosis ($r = 0.9326$, $P < 0.05$).

Oxidative Stress Alters the Abundance of Membrane Complement Regulatory Proteins

Under normal physiological conditions, RPE cells express at least three mCRPs: CD46, CD55, and CD59. Following induction of oxidative stress in RPE cells, membrane abundance of these proteins is reduced. We assessed whether MPs may be one of the mechanisms by which mCRPs are shed from RPE cells under oxidative stress. Flow cytometric analysis showed that exposure of human RPE cells (derived from a 52-year-old male Caucasian donor) to 500 µM H$_2$O$_2$ for 16 hours significantly increased the percent of MPs staining positive for mCRPs: CD46 (9-fold, $P < 0.0001$), CD55 (6-fold, $P < 0.002$), and CD59 (5-fold, $P < 0.0001$), compared with controls (Fig. 6A). In addition, RPE cells with staining positive for mCRPs were significantly reduced (Fig. 6B), suggesting the loss of mCRPs from RPE cell surface is associated with the generation of mCRP-positive MPs. Similar results were obtained in experiments performed in a human RPE cell line, adult retinal pigment epithelium (ARPE)-19 cells (Supplementary Fig. S3), and cultured human RPE cells derived from another donor (a 61-year-old male Caucasian donor; Supplementary Fig. S4).

$N$-Acetylcysteine Amide Attenuates Oxidative Stress–Induced Effects on Microparticles and on RPE Cells

$N$-acetylcysteine amide can protect ARPE-19 cells against oxidative stress–induced death and decrease in cellular GSH.
To determine if NACA can protect cultured human RPE cells against H$_2$O$_2$-induced MP release, cell death, and decrease in cellular GSH levels, we pretreated RPE cells with the thiol antioxidant NACA and analyzed cell apoptosis, the released MPs, and cellular GSH levels by flow cytometry. Pretreatment of RPE cells with NACA significantly attenuated 500 μM H$_2$O$_2$-induced increase in PS-positive MPs (Fig. 7A), which was further confirmed by confocal microscopy (Fig. 7B). In addition, NACA pretreatment prevented the elevation of mCRP-positive MPs in 500 μM H$_2$O$_2$-treated samples (Figs. 7C–E; P < 0.0001, compared with H$_2$O$_2$ without NACA). Furthermore, the observed NACA effect on MPs was associated with significant attenuation of RPE cell apoptosis (Fig. 8A; P < 0.0001) and with restoration of cellular GSH to control levels (Fig. 8B; P < 0.0001).

N-acetylcysteine amide also significantly prevented RPE cells from loss of mCRPs induced by 500 μM H$_2$O$_2$ (Figs. 8C–E; P < 0.0001).

To confirm our flow cytometry findings presented in Figures 6 through 8, we performed fluorescent Western blot analysis. A single antibody blotting with either CD46 or CD59 specific antibody (Supplementary Table S2) detected a single band with molecular weight of approximately 60 kDa for CD46 or approximately 20 kDa for CD59 in cultured human RPE cells (data not shown), confirming the findings by others. $^{35}$ By using multiplexed fluorescent Western blot analysis with low autofluorescence Immobilon-FL PVDF membrane (Cat#: IPFL00010; EMD Millipore), we directly visualized and compared the expression levels of CD46, CD59, GAPDH, and β-actin in the same blots without stripping. Stripping and reprobing of the blots can cause erroneous results due to the effect of stripping on target proteins. $^{38}$ As shown in Figure 9, CD46 antibody detected a single band of approximately 60 kDa, and CD59 antibody detected a single band of approximately 20 kDa in whole-cell lysates and MP lysates. CD59 signal is much stronger than that of CD46 in the same multiplexed blots. Compared with equal amount of whole-cell lysates (lanes labeled with 5 μg), both CD46 and CD59 are enriched in MPs in control, and H$_2$O$_2$-treated samples with or without NACA. H$_2$O$_2$ treatment increases the amount of CD46 and CD59 in MPs and decreases the amount of CD46 and CD59 in RPE cells, compared with control (left panel). Pretreatment with NACA decreases the amount of CD46 and CD59 in MPs and increases the amount of CD46 and CD59 in RPE cells, compared with H$_2$O$_2$ treatment alone (right panel). Therefore, our Western blot analysis confirms our flow cytometry findings: H$_2$O$_2$ induces the loss of CD46 and CD59 from RPE cells with the concomitant gain of CD46 and CD59 in MPs, all of which are attenuated by NACA.

It’s well-known that GAPDH and β-actin serve as loading controls for whole-cell lysates. However, whether GAPDH and β-actin can serve as loading controls for MP lysates is unknown. As shown in Figure 9, as expected, both GAPDH (red) and β-actin (green) are detected in whole-cell lysates, but only GAPDH (red) is detectable in MPs. Compared with equal amount of whole-cell lysates (lanes labeled with 5 μg), GAPDH expression levels in MPs (MP lysates) are much lower than those in RPE cells (whole-cell lysates). Due to lack of known loading controls for MP lysates, normalization and quantification were not attempted.

It is important to note that our flow cytometry analyzed CD46 and CD59 protein expression on the surface of RPE cells and of MPs, while our Western blot analysis detected CD46 and CD59 protein expression in whole cells and whole MPs.

To test if a lower concentration of H$_2$O$_2$ could cause similar effects as those induced by 500 μM H$_2$O$_2$ described above, we used 200 μM H$_2$O$_2$ that induced 10% RPE apoptosis with no necrosis (Supplementary Fig. S2). Similar to 500 μM, 200 μM H$_2$O$_2$ also significantly increased PS-positive MP and CD59-positive MP release, both of which were reversed by NACA pretreatment (Supplementary Figs. S5A, S5D). However, unlike 500 μM H$_2$O$_2$, 200 μM H$_2$O$_2$ had no statistically significant effect on CD46- or CD55-positive MP release (Supplementary Figs. S5B, S5C).

**Discussion**

Here, we find that 500 μM H$_2$O$_2$ treatment produced a marked increase in released MPs by human RPE cells. This MP increase was accompanied by increases in RPE apoptosis and loss of mCRPs (CD46, CD55, and CD59) on the RPE cell surface. The released MPs exposed PS and PE, and carried mCRPs on their surface. Moreover, we demonstrate for the first time in any cell types that NACA, a thiol antioxidant, prevented the H$_2$O$_2$-enhanced release of mCRP-positive MPs, and loss of mCRPs on the RPE cell surface.
Oxidative stress from reactive oxygen species is a major factor involved in the RPE death that underlies AMD. Previously, we and others demonstrated that oxidative stress induces RPE apoptosis and degeneration.49–54 Now we show that oxidative stress also induces RPE to release MPs that carry mCRPs.

Cell-derived extracellular vesicles are generally classified into three main types: MPs, exosomes, and apoptotic bodies. These groups are based primarily on their size and presumed biogenetic pathways. Microparticles (also termed ectosomes or microvesicles), formed by membrane blebbing, are a heterogeneous population of small vesicles of 100 to 1000 nm in diameter17–20; exosomes refer to smaller vesicles of 50 to 100 nm in diameter, generated by exocytosis of multivesicular bodies55; and apoptotic bodies are vesicles of 1000 to 5000 nm in diameter, and released as blebs from cells undergoing late apoptosis.56,57 They are released into the extracellular environment by various types of cells, including epithelial cells.58,59

Several research groups have investigated RPE-derived exosomes,16,53,60–64 and ARPE-19 blebs.65,66 However, a full manuscript investigating RPE-derived MPs has not published yet. In this study, we used both TEM and flow cytometry to characterize human RPE cells derived from two donors (52 years, 61 years) and ARPE-19 cells (19 years). To our knowledge, this is the first study to investigate MPs from RPE cells derived from older donors (52 years, 61 years), which could be more relevant to AMD. We show for the first time that MP sized vesicles are released from RPE cells under normal and oxidative stress conditions by TEM (Fig. 1), flow cytometry (Figs. 2–5, 6A, 7A, 7C–E), and confocal microscopy (Fig. 7B). Apoptotic bodies were eliminated by passing samples through a 1.2-μm filter. The centrifugal force (20,000 g) we used to isolate MPs is not sufficient to collect exosomes, which require a centrifugal force of 100,000g.17,66 Furthermore, exosomes are not detected by flow cytometry because their size is generally less than 100 nm,18 below the detection limit of our flow cytometer which could not detect 0.1-μm beads. Moreover, CD63, an exosomal marker, was not detected by Western blot analysis, indicating no contamination with exosomes in our MP population (Supplementary Fig. S1).

Two key events in early AMD are the appearance of drusen5–7 and RPE apoptosis.8,9 The importance of the RPE in drusen formation is evidenced by the fact that membranous debris, drusen-like deposits, and drusen build up on apical and basolateral sides of the RPE monolayer,5–7,67,68 and drusen are composed of materials found in RPE cells.5,15,28,69 The strong correlation between RPE apoptosis and PS-positive MP release was measured by annexin V-FITC and PI staining. Phosphatidylserine-positive MPs (%) were plotted on the x-axis, and the RPE apoptosis (%) was plotted on the y-axis for different time points (2–24 hours). Correlation coefficient (r) = 0.9326. Data are presented as mean ± SD (n = 3). P < 0.05.
sheddng (Fig. 5) suggests that MPs released from the apoptotic RPE cells may be early events in subclinical drusen, carrying with them the cellular components commonly found in drusen.

In this study, we found that the increase in PS-, PE-, and mCRP-positive MPs was accompanied by the increase in apoptosis and loss of mCRPs (CD46, CD55, or CD59) from 500 μM H$_2$O$_2$-treated RPE cells by flow cytometry, suggesting that the decreased mCRPs on the RPE surface result from increased shedding of mCRP-positive MPs. Our Western blot analysis confirms the flow cytometry findings for CD46 and CD59. Although we constantly found that CD55 was present on RPE cell surface and on MPs by flow cytometry (Figs. 6, 7D, 8D), we were unable to detect CD55 expression in lysates prepared from intact cells (whole-cell lysates) or lysates prepared from MPs by Western blot analysis, using two commercially available antibodies against CD55 (data not shown). One CD55 antibody we used was mouse anti-human CD55 monoclonal antibody (EPR66890, Cat#: ab133684; Lot#: GR93375-13; Abcam, Cambridge, MA, USA). Consistent with our results, Yang et al. did not detect CD55 expression in both native and cultured human RPE cells by Western blot analysis using a different commercially available antibody, although they did detect CD55 expression on RPE surface by flow cytometry. Thurman et al. reported that 1 mM H$_2$O$_2$ reduced the surface expression of CD55 (also known as decay accelerating factor [DAF]) and CD59 in ARPE-19 cells using flow cytometry, but no Western blot analysis of CD46, CD55, and CD59 was reported in their study. The reasons for not detecting CD55 protein expression by Western blot analysis are not clear. One possible explanation could be that CD55 antibodies commercially developed for Western blot analysis are not suitable for CD55 antigen in the RPE cells due to its unique posttranslational modifications. Additional investigations will be required to test this hypothesis in the future studies. Interestingly, Ebrahimi et al. found that cellular CD46 and CD59 proteins were decreased in ARPE-
19 cells treated with oxidized low-density lipoproteins and that CD59 and CD46 along with an exosomal marker CD63 were detected in culture supernatant. They suggested that the decreased levels of CD46 and CD59 were in part due to the release of exosomes and apoptotic particles. Two proteomic studies were performed on extracellular vesicles derived from ARPE-19 cells. One type of extracellular vesicles (ARPE-19 membrane blebs) was obtained by centrifuging conditioned medium for 15 minutes at 100,65 whereas another type of extracellular vesicles (exosomes) was obtained by centrifuging conditioned medium for 1 hour at 100,000g. Both proteomic studies did not demonstrate the detection of CD46, CD55, and CD59 on these vesicles. Although there are no standard procedures available to isolate different types of extracellular vesicles (mainly MPs, exosomes, and apoptotic bodies), the most common protocol for isolation of MPs consists of differential centrifugation with final centrifugal force of 20,000g. In this study, we followed this most common protocol to isolate and analyze MPs from cultured RPE cells derived from donor eyes as well as ARPE-19 cells. We detected PS- and PE-positive MPs, and found the presence of CD46, CD55, and CD59 on the surface of isolated MPs, suggesting the CD46 found in drusen during early AMD33,36 or CD59 detected in subretinal space could originate from RPE cells through the release of MPs.

In addition to potentially driving or participating in drusen formation, our data indicate a possible effect of MP shedding on the retinal cells, the loss of mCRPs in the RPE layer may lead to enhanced complement attack and further cell death. Numerous studies point to deregulation of the complement

**FIGURE 8.** N-Acetylcysteine amide attenuates H2O2-induced apoptosis, decrease in cellular GSH levels, and loss of membrane complement regulatory proteins in RPE cells. Control and 500 μM H2O2-treated RPE cells with or without pretreatment of NACA were stained with annexin V-Alexa Fluor 488 and PI for apoptosis assay (A), stained with monochlorobimane (a cell-permeable dye) reacting with cellular GSH (B), or stained with fluorescently-labeled antibodies to CD46 (C), CD55 (D), or CD59 (E). Data are presented as mean ± SD (n = 3–5). **P < 0.01, ***P < 0.0001 compared with controls –NACA; #P < 0.0001 compared with H2O2-treated groups –NACA.

**FIGURE 9.** Fluorescent Western blot analysis of CD46 and CD59 proteins in RPE cells and RPE-derived MPs. Human RPE cells were treated vehicle (control; lanes labeled with C), 500 μM H2O2 (lanes labeled with H), or 500 μM H2O2 with NACA pretreatment (lanes labeled with N+H) for 16 hours. Cells and MPs were harvested and isolated after treatment. Intact cells or isolated MPs were lysed and proteins were measured. Proteins of whole-cell lysates or of MP lysates were subjected to Western blot analysis of membrane complement regulatory proteins (mCRPs) CD46 and CD59. The anti-CD46, anti-CD59 or anti-β-actin primary antibody was detected with Alexa Fluor 488-conjugated chicken anti-mouse IgG, and protein bands were pseudocolored green. The primary antibody against GAPDH was detected with Alexa Fluor 647-conjugated goat anti-rabbit IgG, and single bands were pseudocolored red. Shown is an overlay of green (CD46, CD59, and β-actin) and red (GAPDH) images. GAPDH and β-actin serve as loading controls for whole cell lysates. M indicates protein size marker.
cascade in AMD. Loss of mCRPs leads to RPE cells being vulnerable to complement attack. Here, we observed 500 µM H₂O₂-induced loss of CD46, CD55, and CD59 on the surface of cultured human RPE cells at two key points: activity of C₃ convertase and membrane attack complex (MAC) formation. CD46 and CD55 block CD59, while CD59 inhibits MAC assembly. Consistent with our results obtained by 500 µM H₂O₂, previous studies have shown that 1 mM H₂O₂ treatment reduced the surface expression of mCRPs on ARPE-19 cells by flow cytometry, and sensitized ARPE-19 cells to complement-sufficient serum-mediated complement activation and disruption of the barrier function. A lower concentration of H₂O₂ (200 µM) also induced PS-positive MP and CD59-negative MP releases that were prevented by NACA pretreatment (Supplementary Figs. S5A, S5D). Although some CD46- or CD55-positive MP release was detectable in control samples, 200 µM H₂O₂ had no significant effect on their release (Supplementary Figs. S5B, S5C). The mechanisms for this are unknown. Retinal pigment epithelial cells in vivo probably encounter a complex and varying mix of low concentrations of oxidative stress stimuli that might have additive and/or synergistic effects on MP release. CD46 and CD55 inhibit complement attack by blocking C₃ activation while CD59 prevents complement attack by the inhibition of the MAC. We speculate that there could be some sorting differences associated with these two different levels of protection and our data suggest that MP release may be a deliberate and coordinated process instead of a random release of membrane-bound vesicles.

In conclusion, this study demonstrates for the first time that oxidative stress induces human RPE cells to release PS-, PE-, and mCRP-positive MPs. The levels of released MPs are strongly correlated with RPE apoptosis. All of the effects induced by oxidative stress are prevented by NACA. Our data support the concept that RPE-derived MPs can carry components found in drusen, and suggest that oxidative stress could trigger or participate in drusen formation by releasing MPs from the RPE. Our study also describes a novel role for NACA in the prevention of mCRP-positive MP release, which might have broad therapeutic implications, particularly in diseases associated with loss of mCRPs on the cell surface, and with enhanced release of MPs.

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