Mitochondrial Proteomics of the Retinal Pigment Epithelium at Progressive Stages of Age-Related Macular Degeneration

Curtis L. Nordgaard,1 Pabalu P. Karunadharma,1,2 Xiao Feng,1 Timothy W. Olsen,1 and Deborah A. Ferrington1,2

PURPOSE. Age-related macular degeneration (AMD) is the leading cause of vision loss in individuals over the age of 65. Histopathological changes become evident in the retinal pigment epithelium (RPE), a monolayer that provides metabolic support for the overlying photoreceptors, even at the earliest stages of AMD that precede vision loss. In a previous global RPE proteome analysis, changes were identified in the content of several mitochondrial proteins associated with AMD. In this study, the subproteome of mitochondria isolated from human donor RPE graded with the Minnesota Grading System (MGS) was analyzed.

METHODS. Human donor eye bank eyes were categorized into one of four progressive stages (MGS 1–4) based on the clinical features of AMD. After dissection of the RPE, mitochondrial proteins were isolated and separated by two-dimensional gel electrophoresis based on their charge and mass. Protein spot densities were compared between the four MGS stages. Peptides from spots that changed significantly with MGS stage were extracted and analyzed by using mass spectrometry to identify the protein.

RESULTS. Western blot analyses verified that mitochondria were consistently enriched between MGS stages. The densities of eight spots increased or decreased significantly as a function of MGS stage. These spots were identified as the α, β, and δ-ATP synthase subunits, subunit VIb of the cytochrome c oxidase complex, mitoflin, mtHsp70, and the mitochondrial translation factor Tu.

CONCLUSIONS. The results are consistent with the hypothesis that mitochondrial dysfunction is associated with AMD and further suggest specific pathophysiological mechanisms involving altered mitochondrial translation, import of nuclear-encoded proteins, and ATP synthase activity. (Invest Ophthalmol Vis Sci. 2008;49:2848–2855) DOI:10.1167/iows.07-1352

Age-related macular degeneration (AMD) is a leading cause of blindness among older adults in developed nations.1,2 Early clinical features of AMD include alterations in the retinal pigment epithelium (RPE), a monolayer between the photoreceptors and choroid that supports retinal function and homeostasis. The quantity and extent of lipoproteinaceous deposits (drusen) that form between the RPE and choroid correlate with progressive stages of AMD. A significant number of patients with the early features of AMD progress to advanced stages with impaired central visual acuity, characterized by either central geographic atrophy (aAMD) or subretinal choroidal neovascularization with exudation (eAMD).3 The personal and public costs of AMD, coupled with the aging of the U.S. population, create an urgent need to improve AMD prevention and treatment strategies over the next decade.4,5 Further development of rational therapeutic interventions for AMD requires a greater understanding of basic AMD disease mechanisms.

Several lines of evidence indicate a role for mitochondria in the pathogenesis of AMD. First, mitochondria are the major source of superoxide anion in the cell.6 The superoxide anion can generate highly toxic hydroxyl radicals and hydrogen peroxide that damage the cell by reacting with proteins, DNA, and lipids. Oxidative stress appears to play an important role in AMD, since human donor eyes affected by AMD contain increased levels of protein adducts resulting from the oxidative modification of carbohydrates and lipids,7,8 and higher levels of antioxidant enzymes.9,10 Second, mitochondrial (mt)DNA is more susceptible than nuclear DNA to damage from oxidation and blue light,11–15 and mtDNA damage in the retina and RPE accumulates with age.14,15 Such damage may indirectly impair the function of mtDNA-encoded subunits of the electron transport chain and cause increased superoxide anion production, leading to further mtDNA damage and superoxide anion production in a self-perpetuating, destructive cycle.16,17 Third, aging and cigarette smoking are two strong risk factors for AMD that are also associated with mitochondrial dysfunction,18–28 suggesting that aging and smoking may contribute to AMD through their effects on mitochondrial function. Finally, in two recent studies, direct evidence of mitochondrial alterations in AMD has been found.21,22 A morphologic analysis of human donor eyes affected by AMD found an accelerated decrease in the number of mitochondria and cross-sectional area relative to normal age-related changes.21 In addition, our previous proteomic analysis of the global human RPE proteome in AMD identified changes in the content of several mitochondrial proteins including mitochondrial heat shock proteins 60 and 70, ATP synthase-β, and the voltage-dependent anion channel.22 To characterize the mitochondrial changes associated with AMD better, we analyzed the RPE mitochondrial subproteome from human donor eyes categorized with the Minnesota Grading System (MGS).
METHODS

Tissue Procurement and Grading

Briefly, globes were cooled in situ and stored at 4°C after postmortem enucleation until processed as in previous studies. The globes were processed according to the MGS, with the exception that both globes were photographed, dissected, and evaluated. After removing the vitreous, the neurosensory retina was gently peeled back and cut at the optic nerve head. The RPE was then carefully hydrodissected from Bruch’s membrane in balanced saline solution with gentle blunt mechanical debridement. Tissues used in the present study were dissected fresh and stored at ~80°C. There was no evidence during the clinical examination that the observed changes were associated with the cause of death. All research procedures adhered to the Declaration of Helsinki and had Institutional Review Board exemption from the human subjects committee at the University of Minnesota.

Mitochondrial Enrichment

The RPE from each pair of globes was combined (for globes assigned the same MGS grade) and processed collectively. The protein yield from the macular RPE (central 6 mm) was insufficient for two-dimensional (2-D) electrophoresis, and therefore the peripheral RPE was used exclusively. Previous proteomic analyses have demonstrated that AMD is associated with biochemical changes in the peripheral RPE22 and many biochemical changes detected in the macula are also detected in the periphery.23 These results support the use of peripheral RPE in the present study.

RPE tissues were homogenized in a buffer containing 0.5% Nonidet P-40, 1 mM phenylmethylsulphonyl fluoride, 20 mM i-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (pH 7.5), 10 mM KCl, 1.5 mM MgCl₂, 250 mM sucrose, 1 mM ethylenediamine tetraacetic acid, and 1 mM ethylene glycol tetracetic acid. After 150 μL of buffer was added to the RPE from a single pair of globes, the tissue was subjected to two freeze–thaw cycles before the tissue suspension was gently passed six times through a 26-gauge needle. The lysate was cleared of nuclei and intact cells by centrifugation for 15 minutes at 600g. Supernatants were removed and an additional 75 μL of buffer was added to the pellet, followed by a second homogenization and centrifugation. The second supernatant was combined with the first, and a 20-μL aliquot (homogenate) was removed for subsequent analyses. Mitochondria were then pelleted from the cleared cell lysate by centrifugation for 15 minutes at 13,000g. The supernatant, containing less dense organelles, membrane fragments, and soluble proteins (cytosol), was reserved for analysis of the preparation. The mitochondria-enriched pellet was washed once with buffer before resuspension in 8 M urea, 2% 3-(cholamidopropyl)dimethylammonio-1-propanesulfonate, and 0.5% amidoxulobetaine-14 (Calbiochem, San Diego, CA) to yield the mitochondrial fraction. To increase the solubility of membrane proteins, the mitochondrial fraction was subjected to two freeze–thaw cycles and incubated for 30 minutes in a water bath sonicator. Protein content was determined by using the bicinchoninic acid assay (Pierce, Rockford, IL).

2-D Gel Electrophoresis

2-D gel electrophoresis was performed as outlined22 and followed by staining with silver (Bio-Rad, Hercules, CA). Preliminary analyses determined that loading 100 μg of mitochondrial protein yielded the greatest number of spots with a linear response (data not shown), and this protein load was used to rehydrate 3 to 11 nonlinear IPG strips for first-dimension focusing. Samples in the same MGS stage were combined as necessary to yield 100 μg of mitochondrial protein.

2-D Gel Spot Quantitation and Analysis

A power analysis was performed to determine the change (x-fold) detectable given the study’s sample size and variability. This analysis determined that with a sample size of seven gels, a 2.5-fold change could be detected on average with 90% power. Two statistical models, reflecting two possible patterns of disease-related change (linear and stage-specific), were used to test for significant spot quantity changes. The probability for each spot is indicated for a given model, with an uncorrected significance level of P < 0.05 and P < 0.025 after the Bonferroni correction.

Mass Spectrometry (MS)

Peptide extraction from polyacrylamide gels, peptide mass fingerprinting, and sequencing by tandem MS/MS were performed as described22 with the following modifications. Peptide masses were submitted to the Swiss-Prot database (http://www.expasy.org; provided in the public domain by Swiss Institute of Bioinformatics, Geneva, Switzerland) of human proteins by using a search engine (Mascot; Matrix Science Inc., Boston, MA). The peptide mass fingerprinting searches used mass tolerance settings of 50 ppm, carbamidomethylation of cysteine and oxidation of methionine (processing artifacts) as fixed and variable modifications, respectively, and 0 missed cleavages by trypsin. Protein identifications were considered verified by sequencing if the MS/MS spectra submitted to the Mascot-Swiss Prot human database yielded a significant (P < 0.05) match, with peptide and fragment mass tolerances set to 0.8 Da, carbamidomethyl as a fixed modification, and oxidation of methionine set as a variable modification.

1-D Gel Electrophoresis and Western Blot Analysis

RPE fractions were separated on 16 × 16-cm polyacrylamide gels (12%, 1.5 mm thickness or 10%, 1 mm thickness) by the method of Laemmli and transferred as described,23 except that for the 10% gels, the membrane was stained with ponceau S red before blocking, imaged, and destained with Tris-buffered saline (pH 11). Samples were also analyzed on 16 × 16 × 1.5-mm, 8% to 16% gradient gels (Jule Inc., Milford, CT) and stained with flamingo pink fluorescent protein stain (Bio-Rad). Fluorescent gels were excited with a transilluminator (model DR88M; Clare Chemical Research, Dolores, CO) and imaged (Chemidoc system; Bio-Rad). The density of individual bands was then quantified (Quantity One software; Bio-Rad) and tested by one-way ANOVA.

For blotting, the linear response of each primary antibody was verified for the protein load used. Antibodies directed against ATP synthase-α (1:500, A21350), ATP synthase-β (1:500, A21351), and mitochondrial NADH dehydrogenase 6 (1:1500, A31857) were purchased from Invitrogen-Molecular Probes (Eugene, OR). Other antibodies were directed against Lamp-1 (1:250, 611042; BD Biosciences, San Jose CA), tyrosinase (1:500, 35-6000), and CD54 (1:1500, 07-5403; Zymed Laboratories, South San Francisco CA), catalase (1:1000, ab16751; Abcam, Cambridge MA), heat shock protein 70 (1:5000, SPA-812), manganese superoxide dismutase (1:2000, SOD-110; Assay Designs, Ann Arbor, MI), gilial fibrillary acidic protein (1:2000, D097-35; MBL, Woburn, MA), and the voltage-dependent ion channel (1:1000, 529532; EMD Biosciences, San Diego, CA). Standards were loaded for comparisons between gels, and reaction densities were normalized to either the homogenous reaction density or to the total lane density of ponceau staining. Alkaline phosphatase– conjugated secondary antibodies were used for colorimetric development as described and imaged with a densitometer (model GS-800; Bio-Rad). Fluorescent gels were excited with a transilluminator (model DR88M; Clare Chemical Research, Dolores, CO) and imaged (Chemidoc system; Bio-Rad). The density of individual bands was then quantified (Quantity One software; Bio-Rad) and tested by one-way ANOVA.

RESULTS

Human donor globes were graded with the MGS. Donor demographics are presented in Table 1. The average time from death to enucleation and death to tissue cryopreservation for all donors was 4.5 ± 1.8 hours and 16.9 ± 4.2 hours, respectively (mean ± SD). Age matching was not possible, given the
available donor tissues. However, the distribution of donor ages was similar for MGS stages 1 to 3, with an increase at stage 4 (Fig. 1). Stage 1 of the MGS represents the control group. Stage 2 is characterized by the presence of numerous small, hard drusen in the macula and/or RPE pigmentary abnormalities. Stage 3 globes contain one or more large, soft drusen, numerous intermediate drusen, or nonmacular geographic atrophy. Globes categorized as MGS stage 4 contain central geographic atrophy or evidence of subfoveal choroidal neovascularization. Stage 3 globes contain one or more large, soft drusen, hard drusen in the macula and/or RPE pigmentary abnormalities. Stage 4 globes contain central geographic atrophy or evidence of subfoveal choroidal neovascularization.

Since the yield from initial density gradient separations was insufficient for proteomic analysis, differential centrifugation was used to produce a fraction enriched for mitochondria (mitochondrial fraction) and a fraction depleted of mitochondria (cytosolic fraction). It was assumed that (1) most of the proteins in the mitochondrial fraction would normally be expressed in the mitochondria, and (2) the enrichment procedure would not be affected by MGS stage. Therefore, any content changes would be associated with AMD rather than differences in the enrichment of organelles between stages. We tested these assumptions by Western blot analysis of the fractions. First, increasing loads of protein from all three fractions were probed with organelle-specific antibodies (Fig. 2A). All markers were detected in the homogenate. Lysosomal, peroxisomal, melanosomal, and cytosolic markers were clearly detected in the cytosolic but not mitochondrial fractions. Conversely, the outer (VDAC) and inner (ND6) membrane markers were clearly detected in the mitochondrial fraction but not the cytosolic fraction. The matrix protein MnSOD, however, was readily detected in all three fractions and indicates some leakage from the mitochondrial matrix and intermembrane space.

To determine whether the mitochondrial enrichment was effectively similar at each MGS stage, we probed a subset of samples with the same organelle markers. Since the mitochondrial fraction yield was not sufficient for both this analysis and the proteomic analysis, any changes in the enrichment process were measured indirectly by analyzing the homogenate and cytosolic fractions. It was assumed that a change in the mitochondrial fraction would correspond to a change in the cytosolic fraction, since greater organelle enrichment in the mitochondrial fraction would be accompanied by decreased organelle enrichment in the cytosolic fraction (e.g., more lysosomes in the mitochondrial fraction would mean fewer lysosomes in the cytosolic fraction). Each cytosolic reaction was normalized to its homogenate reaction to account for differences in total content between samples. No difference was detected between the four stages by one-way ANOVA (all \( P > 0.12 \); see Fig. 2B) for any of the organelle markers or mitochondrial markers tested. These results indicate that the mitochondrial preparation selectively enriched for mitochondria and that enrichment effectiveness was not significantly different between MGS stages.

Mitochondrial samples were also tested for retinal and choroidal protein contamination (data not shown) by probing for glial fibrillary acidic protein (GFAP), a retinal marker, and CD34, a choroidal marker expressed in the choriocapillaris. Little or no reactivity was detected for either marker. The mean ratio of GFAP signal for the mitochondrial fraction compared to a human retina positive control was 2.8% ± 3.0% (mean ± SD), and the ratio of CD34 signal compared to a human choroidal positive control was 1.1% ± 0.55%. The relationship between MGS stage and contaminant ratio was not significant for either

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**Figure 1.** Scatterplot comparison of human donor ages between MGS stages. **Horizontal bar:** mean age for each group. MGS stage 4 donors were significantly older than those in any other category (one-way ANOVA, \( P < 0.005 \) and the Tukey-Kramer post hoc test). The differences between the mean data for stages 1, 2, and 3 were not significant.

**Table 1.** Donor Demographics and Clinical Information

<table>
<thead>
<tr>
<th>MGS Grade</th>
<th>Sample Size (n)*</th>
<th>Mean Age (y ± SD)</th>
<th>Enucleation† (h ± SD)</th>
<th>Freeze‡ (h ± SD)</th>
<th>Cause of Death§</th>
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<tr>
<td>1</td>
<td>7/7/10/5</td>
<td>67 ± 10</td>
<td>4.2 ± 2.0</td>
<td>16.0 ± 3.9</td>
<td>Cancer (5), organ failure (4), sepsis (2), vascular accident or hemorrhage (4)</td>
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<td>2</td>
<td>7/6/6/6</td>
<td>70 ± 10</td>
<td>4.9 ± 1.7</td>
<td>17.4 ± 4.9</td>
<td>Cancer (4), myocardial infarct (1), organ failure (2), pneumonia (2), sepsis (1), vascular accident or hemorrhage (2)</td>
</tr>
<tr>
<td>3</td>
<td>7/8/6/7</td>
<td>73 ± 14</td>
<td>4.4 ± 1.7</td>
<td>18.2 ± 5.7</td>
<td>Cancer (5), cardiomyopathy (1), myocardial infarct (1), organ failure (2), pancreatitis (1), sepsis (2), surgical complications (1), vascular accident or hemorrhage (3)</td>
</tr>
<tr>
<td>4</td>
<td>5/8/3</td>
<td>86 ± 10</td>
<td>4.4 ± 1.9</td>
<td>16.0 ± 4.5</td>
<td>Asthma (1), cancer (1), myocardial infarct (1), organ failure (5), pneumonia (2), vascular accident or hemorrhage (1)</td>
</tr>
</tbody>
</table>

* Some protein samples from the same stage were combined.  † Length of time interval from death to enucleation.  ‡ Length of time interval from death to freezing of dissected tissue.  § Number of donors is indicated in parentheses. Organ failure includes respiratory failure and cardiac failure. Vascular accident or hemorrhage includes cerebral vascular accident.
Mitochondrial Proteomics

FIGURE 2. Verification of mitochondrial enrichment by Western blot analysis. (A) Western blot analysis of RPE fractions (homogenate, cytosolic, and mitochondrial) probed with antibodies specific for organelle markers. Blots indicate that the fractionation procedure resulted in selective depletion of mitochondria from the cytosolic fraction and selective enrichment of mitochondria in the mitochondrial fraction. Protein loads ranged from 5 to 35 μg of protein in 5 μg increments. +, the HeLa cell lysate–positive control. Only a faint Hsp70 (Cyto) reaction was detected in the mitochondrial fraction after considerably over-exposing the blot, as determined by comparing the positive control reaction intensities. Lyso, lysosomes; melan, melanosomes; tyrosinase antibody; perox, peroxisomes; catalase antibody; cyto, cytosol; Hsp70 antibody; OM, outer mitochondrial membrane; VDAC antibody; IM, inner mitochondrial membrane; complex I subunit 6 (ND6) antibody; and mtx, mitochondrial matrix, MnSOD antibody. (B) Semiquantitative comparison of subfractionation efficacy between MGS stages. Western blot measurements of organelle markers were compared for multiple samples from each stage. Each cytosol immune reaction was normalized to its corresponding homogenate immune reaction to account for differences in total content between samples. Bars, mean ± SE, n = 6 for each group. No significant differences were detected for any marker by one-way ANOVA, suggesting that the subfractionation of cellular contents into cytosolic (i.e., nonmitochondrial) and mitochondrial fractions was not affected by MGS stage.

GFAP or CD34 (P = 0.33 and P = 0.51, respectively). These results indicate that the levels of contaminating protein from the retina and choroid are nearly undetectable and do not change as a function of MGS stage.

Having established that the preparation enriches for mitochondria and that the preparation effectiveness does not differ significantly between MGS stages, we analyzed the mitochondrial fraction by 2-D gel electrophoresis (Fig. 3). Preliminary observations indicated that a protein load of 100 μg of protein would yield the greatest number of spots within the linear range of the silver stain dye (data not shown). Automated spot detection and matching were verified manually, and individual spot densities were normalized to the total protein density to account for staining or protein load differences. An average of 440 spots was resolved per gel and 222 consistently resolved spots were tested for two disease-related patterns of change: a linear, progressive change determined by linear regression and any difference between stages determined by one-way ANOVA. Eight spots changed significantly according to one or both of these models (Fig. 4, P < 0.05; after Bonferroni correction, P < 0.025) and were excised for identification by MS analysis. Initial protein identification by MALDI-TOF (matrix-assisted desorption ionization–time of flight) MS was verified by MS/MS sequencing (Table 2, Fig. 4). We deviated from this protocol for one spot, ATP synthase-δ, which initially produced a nonsignificant match of three peptides by MALDI-TOF MS. Because of its small size and low content of lysine and arginine residues (trypsin cleavage sites), a maximum of only six peptides would have been measured from ATP synthase-δ under ideal conditions. Sequence data for the two most intense peptide peaks in the MALDI-TOF spectrum matched exclusively to ATP synthase-δ. The match was considered valid since we detected three of six possible peptides, sequences from the two most intense peaks matched to ATP synthase-δ, and the observed mass and pl closely matched the predicted values.

Since 2-D gel electrophoresis does not efficiently resolve membrane-associated proteins or high-molecular-weight proteins, mitochondrial samples were also separated using 1-D gradient gels (8%-16%) and stained with flamingo pink fluorescent protein stain. No additional changes were found after comparing densities of individual protein bands, suggesting that there were no major protein changes beyond those detected by the 2-D gel analysis (data not shown).

A significant number of the affected proteins were subunits of the mitochondrial ATP synthase complex that is responsible for generating ATP and maintaining the mitochondrial membrane potential. Hence, we sought to verify our original results by probing RPE homogenates with antibodies specific for ATP synthase-α and -β. There was no significant change in ATP synthase content when tested by one-way ANOVA (although...
there was a strong trend for the ATP synthase-β ANOVA, $P = 0.058$; data not shown). Since there was a poor correspondence between the results obtained by 2-D gel analysis of the mitochondrial fraction versus 1-D Western blot analysis of the homogenate, we re-evaluated the 2-D gels to gain a better understanding of the discrepancy. For example, it has been noted that 1-D Western blot and 2-D gel results are more likely to correlate for proteins that migrate as a single spot in 2-D gels than for those that migrate as multiple spots (e.g., as charge trains or isoelectric variants). The one ATP synthase-β spot and two ATP synthase-α spots that decreased at later MGS stages did appear to migrate as part of a charge train at the approximate theoretical pI for each subunit. Both subunits contain predicted and/or experimentally verified phosphorylation and acetylation sites that could account for the observed charge trains. Peptide mass fingerprint analysis identified three additional ATP synthase-β spots and three additional ATP synthase-α spots from the charge trains, although those spots were not affected by MGS stage (Fig. 5A and data not shown). This suggests that the 1-D Western blot measured both the ATP synthase spots that changed with MGS stage as well as those that did not. Consequently, 2-D gel analysis may be more sensitive for detecting changes in specific posttranslationally modified subsets of a total protein population.

For further understanding of the difference between ATP synthase content measured by 1-D Western blot and 2-D gel, we compared the sum of all identified spot densities for each subunit across MGS stages (Fig. 5B). Unlike the 1-D Western blot results, there was a significant effect of MGS stage on total ATP synthase-α, -β, and -δ content (all $P < 0.05$). This difference may reflect technical differences between the two techniques or differences in mitochondrial and homogenate ATP synthase content. For example, ATP synthase spots migrating outside of the charge train would have contributed to the Western blot measurements but would have gone undetected by the 2-D gel analysis. In addition, the ATP synthase-α and -β subunits are imported from their site of translation in the cytosol. Any decrease in their import could have distinct effects on measurement of the homogenate and mitochondrial ATP synthase content. It is also possible that components of the ATP synthase in RPE localize to the plasma membrane as noted for other cell types. If so, then extramitochondrial ATP synthase protein would be measured in the homogenate but not in the mitochondrial fraction.

**FIGURE 3.** Resolution of mitochondrial fraction proteins by 2-D gel electrophoresis. Representative gel demonstrating the resolution of 100 μg of human donor RPE mitochondrial fraction protein. Proteins were separated in the first dimension using a nonlinear pH gradient from 3 to 11 and in the second dimension using 13% SDS-polyacrylamide gel electrophoresis, then stained with silver. Spots that changed significantly by one-way ANOVA or linear regression analysis are boxed and numbered according to Table 2.

**FIGURE 4.** Protein spot densities and corresponding protein identity. Individual spot densities were normalized to the total spot density for each gel and compared by one-way ANOVA (A) and linear regression (L). The test that reached significance and its associated probability are indicated above the normalized spot densities, which are reported as the mean ± SE for each MGS stage. The number of measurements for each stage is indicated below the y-axis. Protein identifications (numbered according to Table 2) were obtained by MALDI-TOF mass spectrometry and verified by MS/MS peptide sequencing. Note the y-axis scale difference between the top and bottom panels.

**DISCUSSION**

We recently identified several mitochondrial protein changes in RPE isolated from donor eyes affected with AMD using a global proteomic analysis. In the present study, we sought to identify specifically the changes in the mitochondrial subpro-
teome by analyzing an enriched mitochondrial fraction from human donor RPE categorized using the MGS. Changes in the content of eight proteins crucial for mitochondrial translation, import of nuclear-encoded proteins, and metabolism were associated with AMD onset and progression. The use of proteomics to analyze human donor tissue has thus led to the identification of specific mitochondrial pathways affected by AMD.

Proteomic analysis of human tissue has many advantages, although there are some limitations. First, the postmortem analysis of pathologic human tissues is a highly valid approach to studying disease but may be limited by variations in the time interval from death to tissue collection and cryopreservation. Neither time interval, from death to enucleation nor death to the time of tissue freezing, varied significantly among the four MGS groups and consequently cannot account for any differences based on MGS stage. Second, although we identified several novel mitochondrial protein changes by 2-D gel analysis, some changes probably went undiscovered, because 2-D gels are less efficient at resolving membrane proteins and high-molecular-weight proteins. Two approaches were used to address these limitations: the use of an additional zwitterionic detergent (ASB-14) to increase protein solubility before first dimension focusing and a separate 1-D gel analysis to detect changes in SDS-soluble membrane proteins and high-molecular-weight proteins. No additional changes were detected by the 1-D gel analysis, suggesting that no major changes in protein content occurred beyond those detected by the 2-D gel analysis. Every proteomic strategy has particular strengths and

TABLE 2. Protein Identification by Mass Spectrometry

<table>
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<tr>
<th>Spot</th>
<th>Protein</th>
<th>Accession Number*</th>
<th>Theoretical Mass†/pI</th>
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<th>Coverage (%)</th>
<th>Error (ppm)</th>
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* Primary accession number from Swiss-Prot (http://www.expasy.org; provided in the public domain by the Swiss Institute of Bioinformatics, Geneva, Switzerland).
† Molecular mass in kilodaltons.
‡ Includes peptides sequenced by MALDI-TOF tandem MS/MS and ESI-tandem MS/MS. The number of significantly identical peptides is indicated.
§ Coverage of ATP synthase-δ was 53% of the total peptides expected within the measured mass range but was not sufficient for a significant MOWSE score. The match to ATP synthase-δ was significant after searching with the peptide sequences obtained by ESI-tandem MS/MS.

FIGURE 5. 2-D gel analysis of mitochondrial ATP synthase content. (A) Mass spectrometry analysis identified multiple ATP synthase-α and -β protein spots with altered isoelectric focusing points (isolectric variants, arrows). Only a subset of the isoelectric variants changed significantly with MGS stage (numbered according to Table 2), based on the initial 2-D gel analysis. Although the spots identified as ATP synthase-α and -β migrated as one of several ATP synthase isoelectric variants, ATP synthase δ appeared to migrate as a single spot. (B) Total mitochondrial ATP synthase-α, -β, and -δ content was estimated by summing the density of all identified spots (A, arrows). Mean densities ± SEM were normalized to the greatest density (MGS 2 for each subunit). The one-way ANOVA for each of the three ATP synthase subunits was significant (all P < 0.04).
weaknesses, and despite the limitations just described, the present 2-D gel analysis successfully compared the content of more than 200 protein spots across four stages of AMD. Given the restricted amount of RPE tissue provided by a single donor (and far less mitochondrial protein), our methodology was successful in identifying AMD-related mitochondrial protein changes.

Unlike other proteomic studies that seek to identify the proteins expressed in an organelle or subcellular structure, the present study sought to identify mitochondrial changes associated with AMD and hence did not require a perfectly pure mitochondrial preparation. The loss of some mitochondrial protein or retention of some nonmitochondrial protein should not affect the measurement of AMD-related changes if the loss or contamination does not vary between MGS stages, which was found for every contaminant examined. Similarly, the loss of some matrix contents regrettably decreased the number of mitochondrial proteins measured by the 2-D gel analysis but was not correlated with MGS stage and should not affect the validity of the results.

Converging evidence suggests that DNA damage occurs more readily in the mitochondrial than the nuclear genome, and that mtDNA damage could lead to RPE dysfunction. Blue light irradiation and treatment with hydrogen peroxide cause long-lasting mtDNA but not nuclear DNA mutations in cultured RPE cells.11–13 Several disorders caused by mtDNA deletions and mutations, such as maternally inherited deafness and diabetes (MIDD), MELAS (myopathy, encephalopathy, lactic acidosis, and stroke-like episode syndrome), specific forms of retinitis pigmentosa, and Leber’s hereditary optic neuropathy, can lead to retinopathy.31–33 Finally, mtDNA deletions in the RPE and retina increase with age and could contribute to the age-dependent onset of AMD.14,15 Because most of the mitochondrial genome encodes either oxidative phosphorylation subunits (OXPHOS; 13 protein-encoding genes) or tRNAs necessary for mitochondrial translation33 (22 tRNA genes), mtDNA mutations and deletions often affect these specific mitochondrial systems.34 Polypeptides encoded by mtDNA are synthesized by mitochondria-specific ribosomes, tRNAs, and translation factors. The mitochondrial translation factor Tu (Tufm) delivers aminoacylated tRNAs to the mitochondrial ribosome as part of a mitochondrial translation elongation complex.35 Because of its central role in mitochondrial translation, increased or decreased levels of Tufm have direct consequences for the synthesis of mtDNA-encoded proteins. In a yeast model, overexpression of Tufm rescued a mitochondrial phenotype caused by the MELAS tRNA mutation.36 Overexpression of Tufm in fibroblasts from patients with a mitochondrial translation deficiency also restored the translation of mtDNA-encoded OXPHOS subunits and the assembly of OXPHOS complexes.37 On the other hand, a single point mutation in the Tufm tRNA binding site led to a dramatic loss of mitochondrial translation in patient-derived fibroblasts.38 Although we did not measure mtDNA mutations or deletions directly, we did find a dramatic upregulation of Tufm at the earliest clinical stage of AMD (MGS stage 2). Tufm appears to be necessary for mitochondrial translation and to rescue translation defects, including those caused by tRNA mutations. These data indicate that changes in mitochondrial translation pathways are associated with an early AMD phenotype, possibly as a consequence of increased mtDNA damage in early AMD.

We also found AMD-related changes in the content of four OXPHOS subunits, including three from ATP synthase. The relatively consistent decrease in total ATP synthase-α, -β, and -δ (Fig. 5B) suggests a concerted loss of the ATP synthase complex between MGS stages 2 and 3. A subsequent loss of ATP synthase activity and decreased cellular ATP levels could be detrimental for RPE function and viability. Alternatively, a fraction of ATP synthase complexes exist as oligomeric supercomplexes that maintain mitochondrial morphology and the mitochondrial membrane potential.39–41 Because the F1 component of ATP synthase is believed to participate in its oligomerization,42 decreased content of the ATP synthase-α, -β, and -δ subunits may also affect ATP synthase supercomplex oligomerization, mitochondrial morphology, and the mitochondrial membrane potential. Although the consequences are presently unknown, loss of the ATP synthase F1 subunits could be a pathologic event occurring during early AMD.

The present results also indicate an association between AMD and deficient mitochondrial import of nuclear-encoded proteins. In both a previous global proteomic analysis of the RPE5,22 and the present mitochondrial subproteome analysis, we found a decrease in the content of the mitochondrial heat shock protein mtHsp70 associated with AMD. Decreased mtHsp70 content could be detrimental due to its pleiotropic functions in p53-mediated apoptosis, iron-sulfur cluster biogenesis, and mitochondrial calcium regulation.43–45 However, mtHsp70 is also required for the ATP-dependent import of nuclear-encoded proteins into the mitochondrial matrix.46 Decreased mtHsp70-dependent import could consequently affect many matrix-localized functions (including the TCA cycle and β-oxidation).

In summary, we performed a mitochondrial sub-proteome analysis of RPE categorized using the MGS. We replicated our previous observation that decreased content of mtHsp70 is associated with AMD and identified novel associations between AMD and both the decreased content of three ATP synthase subunits and increased content of a mitochondrial translation factor. These results are consistent with the hypothesis that mtDNA damage and mitochondrial dysfunction contribute to the pathogenesis of AMD. The identification of mitochondrial translation, import, and ATP synthase protein changes using proteomic analysis of clinically staged human donor tissue represents a powerful approach for identifying putative AMD disease mechanisms.

Acknowledgments

The authors thank the Minnesota Lions and Minnesota Lions Eye Bank personnel for their assistance in procuring eyes for this study, and the Mass Spectrometry Consortium for the Life Sciences (University of Minnesota) for technical assistance.

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