Biomarkers, Genomics, Proteomics, and Gene Regulation

Abnormalities in Mitochondrial Structure in Cells from Patients with Bipolar Disorder

Anne M. Cataldo,*†‡ Donna L. McPhie,*†
Nicholas T. Lange,*‡§ Steven Punzell,‡
Sarah Elmiligy,†‡ Nancy Z. Ye,†‡
Michael P. Froimowitz,*§ Linda C. Hassinger,†
Emily B. Menesale,†‡ Laura W. Sargent,‡
David J. Logan,§ Anne E. Carpenter,§
and Bruce M. Cohen*‡

From the Department of Psychiatry,* Harvard Medical School, Boston; the Laboratory of Molecular Neuropathology,† the Neurostatistics Laboratory,§ and the Molecular Pharmacology Laboratory,‡ Mailman Research Center and Frazier Research Institute, McLean Hospital, Belmont; and the Broad Institute of Harvard and MIT,¶ Cambridge Massachusetts

Postmortem, genetic, brain imaging, and peripheral cell studies all support decreased mitochondrial activity as a factor in the manifestation of Bipolar Disorder (BD). Because abnormal mitochondrial morphology is often linked to altered energy metabolism, we investigated whether changes in mitochondrial structure were present in brain and peripheral cells of patients with BD. Mitochondria from patients with BD exhibited size and distributional abnormalities compared with psychiatrically-healthy age-matched controls. Specifically, in brain, individual mitochondria profiles had significantly smaller areas, on average, in BD samples (P = 0.03). In peripheral cells, mitochondria in BD samples were concentrated proportionately more within the perinuclear region than in distal processes (P = 0.0008). These mitochondrial changes did not appear to be correlated with exposure to lithium. Also, these abnormalities in brain and peripheral cells were independent of substantial changes in the actin or tubulin cytoskeleton with which mitochondria interact. The observed changes in mitochondrial size and distribution may be linked to energy deficits and, therefore, may have consequences for cell plasticity, resilience, and survival in patients with BD, especially in brain, which has a high-energy requirement. The findings may have implications for diagnosis, if they are specific to BD, and for treatment, if they provide clues to the underlying pathophysiology of BD. (Am J Pathol 2010, 177:575–585; DOI: 10.2353/ajpath.2010.081068)
of mitochondrial DNA or of nuclear-encoded genes linked to mitochondrially based oxidative phosphorylation (OXPHOS).

Despite the various findings suggesting mitochondrial dysfunction in BD and separate evidence that altered mitochondrial morphology is crucially linked to altered energy metabolism, mitochondrial structure and distribution have not previously been studied in BD. For these reasons, we examined the size (shape) and distribution of mitochondria in BD using postmortem human brain and primary peripheral cells, both skin fibroblasts and blood lymphocytes, from individuals with BD and healthy comparison subjects.

Materials and Methods

Tissue and Primary Cell Lines

Postmortem tissue was obtained from prefrontal cortex (PFC) from ten individuals with BD Type 1, (ages 30–80 years) and an equal number of age-matched controls with no known history of psychiatric illness (Table 1). All tissue was procured from the Harvard Brain Tissue Resource Center (HBTRC), McLean Hospital, Belmont, MA and had previously been fixed in 10% formalin under standard procedures listed on the HBTRC website.

Fibroblast lines, comprising 8 BD and 8 age-matched psychiatrically-well comparison (control) subjects (ages 20–65 years) (Table 2), and transformed lymphocyte (lymphoblast) lines, comprising 6 BD and 6 age-matched controls, were obtained from the Coriell Institute of Medical Research, Camden, NJ. Fibroblast cell lines were grown in Minimal Essential Media plus 15% fetal bovine serum (FBS) and 1% glutamine, penicillin, and streptomycin in 5% CO₂. Lymphocytes were grown in RPMI plus 15% FBS and 1% glutamine, penicillin, and streptomycin in 5% CO₂. All cell culture media was from Invitrogen (Invitrogen Corporation, Carlsbad, CA). Media was changed every other day. Cell lines were used between passages 3 and 14.

Light Microscopic Probes

Fibroblasts were plated at a density of 2.5 × 10⁴ cells per coverslip. MitoTracker-M7510, 125 nmol/L, (Invitrogen Corporation, Carlsbad CA), was used to visualize mitochondria according to manufacturer’s instructions. Cytoskeletal integrity was examined by immunocytochemistry (see below) using an antibody to tubulin I and II (Sigma-Aldrich Corp., St. Louis, MO and Cell Signaling, Danvers, MA) or with fluorescent phalloidin (Invitrogen Corporation, Carlsbad, CA). Cultured cells labeled with fluorescent M7510 were fixed with 4% paraformaldehyde at 37°C, washed two times with PBS, and treated with phalloidin (1:8000) for 20 minutes at room temperature for visualization of the actin cytoskeleton (actin filaments) and then mounted in gel mount that contained either 4',6-diamidino-2-phenylindole (DAPI; 200 nmol/L) or bis-benzimide (Hoechst 33258; 5 µg/ml) (both from Sigma-Aldrich Corp., St. Louis, MO) for identification of the nucleus and as a marker of nuclear changes indicative of apoptosis.

Immunocytochemistry of Cytochrome C in Fibroblasts

Cells were grown on glass coverslips and fixed as detailed above. Samples were washed three times with...
PBS, washed in dilution buffer (PBS + 2% BSA + 1% normal serum + 0.1% Triton-X100), and incubated with 0.5% SDS in PBS to facilitate antigen retrieval. After blocking in 10% serum, the cells were incubated in primary antibody to cytochrome C (1:2000) (Mitosciences, Eugene OR) overnight at room temperature with gentle rocking, then washed three times with dilution buffer, followed by incubation in fluorescent-conjugated secondary antibody for 2–3 hours at room temperature. To visualize the tubulin cytoskeleton, the cells were then incubated overnight with an antibody to tubulin directly conjugated to Alexa 555 (Cell Signaling, Danvers MA). Coverslips were then mounted and examined using a Zeiss epifluorescent microscope.

**Electron Microscopy**

Cells and vibratome-sectioned brain tissue were processed following standard EM protocols. All chemicals were from Ted Pella (Reading, PA). Briefly, tissue and cells were postfixed in 1% OsO4, dehydrated in a series of alcohols up to absolute alcohol and propylene oxide and embedded in Epon. Blocks were sectioned, stained, and examined on a JEOL 1200EX microscope.

**Measurement of Mitochondrial Area in EM Images**

A two-dimensional analysis using Image Pro Analyzer 6.3 (MediaCybernetics, Bethesda, MD) was used to quantify mitochondrial area from 10 BD and 10 control postmortem brain samples. Briefly, mitochondrial areas of 300 mitochondrial profiles were measured from digital images of cells from postmortem PFC at ×10,000 magnification. Ultrastructural images from thirty fibroblasts were obtained at random (magnification ×10,000) from each of seven healthy controls and eight BD cell lines (in total, 210 control fibroblasts and 240 BD fibroblasts). For each line of fibroblasts, 150 mitochondrial profiles in digital images from electron micrographs were measured with Image Pro 6.3 as above. Additionally the number of ring and cup shaped mitochondrial profiles were counted for each fibroblast sample. (Cup shaped was defined as being at least one half of a ring shape.) In both of the above data sets, the person doing the measurement was blind to the diagnostic condition of the sample.

**Analysis of Cytochrome C Immunohistochemistry of Human Prefrontal Cortex**

Vibratome sections of fixed PFC tissue from a subset of our brain samples (six BD and six Control subjects) were boiled in TBS for 3 minutes to facilitate antigen exposure and then rinsed three times in TBS. Endogenous peroxidase activity was blocked by incubating sections in 0.3% H2O2 in methanol for 30 minutes, and then sections were washed three times in dilution buffer. Sections were then blocked in 20% goat serum and incubated in anti cytochrome C antibody (Calbiochem, Gibbstown NJ) at a dilution of 15:1000 in dilution buffer. After washes, sections were incubated in biotinylated secondary antibody

**Table 2. Demographic Information for Fibroblast Lines Used**

<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Location biopsy</th>
<th>Family</th>
<th>Medications</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>M</td>
<td>Not clinically affected</td>
<td>Leg</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>61</td>
<td>M</td>
<td>Not clinically affected</td>
<td>N/A</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>30</td>
<td>F</td>
<td>Not clinically affected</td>
<td>The mesial aspect of the mid-upper left arm</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>23</td>
<td>M</td>
<td>Not clinically affected</td>
<td>N/A</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>27</td>
<td>M</td>
<td>Not clinically affected</td>
<td>N/A</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>29</td>
<td>M</td>
<td>Not clinically affected</td>
<td>N/A</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>25</td>
<td>F</td>
<td>Not clinically affected</td>
<td>Left forearm</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>52</td>
<td>F</td>
<td>Not clinically affected</td>
<td>The mesial aspect of the mid-upper left arm</td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

Bipolar disorder

<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Location biopsy</th>
<th>Family</th>
<th>Medications</th>
</tr>
</thead>
<tbody>
<tr>
<td>23</td>
<td>F</td>
<td>BD onset at age 18 alcoholic, drug abuse</td>
<td>Posterior iliac crest</td>
<td>3 affected in family</td>
<td>Lithium</td>
</tr>
<tr>
<td>20</td>
<td>M</td>
<td>BD onset at age 17 drug abuse</td>
<td>Posterior iliac crest</td>
<td>6 affected in family</td>
<td>Lithium</td>
</tr>
<tr>
<td>28</td>
<td>F</td>
<td>BD</td>
<td>N/A</td>
<td>5 affected in family</td>
<td>Lithium; Haldol; Cogentin and Thorazine</td>
</tr>
<tr>
<td>24</td>
<td>M</td>
<td>BD medical disorder with seizures; alcoholism at age 15 depression at age 23</td>
<td>N/A</td>
<td>Mother and brother have BD</td>
<td>N/A</td>
</tr>
<tr>
<td>22</td>
<td>M</td>
<td>BD onset at age 20</td>
<td>N/A</td>
<td>Father has BD</td>
<td>Lithium hospitalized twice</td>
</tr>
<tr>
<td>65</td>
<td>M</td>
<td>BD onset at age 18</td>
<td>N/A</td>
<td>6 affected in family</td>
<td>None</td>
</tr>
<tr>
<td>26</td>
<td>M</td>
<td>BD onset at age 13</td>
<td>N/A</td>
<td>4 affected in family</td>
<td>Lithium and haldol hospitalized</td>
</tr>
<tr>
<td>51</td>
<td>F</td>
<td>BD onset at age 31</td>
<td>N/A</td>
<td>2 similarly affected sons and 1 daughter</td>
<td>Lithium</td>
</tr>
</tbody>
</table>

BD indicates bipolar disorder; M, male; F, female; N/A, not available.
for 30 minutes and ABC (Vector Labs, Burlingame, CA) for 1 hour. Staining was visualized with DAB and then sections were mounted on slides, dehydrated, and cleared with xylene. Eighty images of DAB-positive cells were taken in bright field at $\times 1000$. Before line profile analysis images were processed with Adobe Photoshop using the autocolor and subsequent grayscale commands. Grayscale images were analyzed with line profile analysis using Image Pro Analyzer 6.3 by drawing a line through the longest axis of the cell. Analysis was blind to the diagnostic condition of the sample. Line profile transects were centered and scaled to have mean 0 and variance 1, interpolated across a common median length across all brains and cells and smoothed objectively to remove high-frequency noise.27 Transects were then averaged across all cells sampled from each brain separately, and the result was averaged within groups for between-group comparison and assessed by a Kolmogorov-Smirnov test.

**Morphological Analysis of Mitochondrial Distribution Using Digital Images**

We quantified and analyzed mitochondrial density differences between fibroblasts obtained antemortem from the BD and control groups plated at a density of $2.5 \times 10^4$ cells per coverslip. Five random fields of view at $\times 630$ magnification were obtained from 24 to 41 cells from each of $n = 8$ BD subjects (total BD cells analyzed = 231) and in 20 to 40 cells from each of $n = 8$ healthy age-matched control subjects (total healthy cells analyzed = 251). The number of cells per image varied across subjects from 2 to 11. For each cell in each digital image, inner and outer cytoplasmic boundaries were traced manually. The inner boundary coincided with the nuclear envelope as determined by DAPI or Hoechst staining, and the outer boundary coincided with the external cell membrane of the fibroblast as determined by phalloidin staining of actin. No overlap was allowed between outer boundaries for different cells. For each cell, a set of concentric, equally spaced, scaled copies of the nuclear envelope were then aligned with the centroid of the envelope to segment the entire cytoplasmic area into expanding annuli. We then quantified mitochondria within each annulus by recording pixel gray levels against a low and clearly separated background. We defined total mitochondrial intensity within each annulus as the sum of gray levels of mitochondrial material (M7510-positive) segmented from image background.28 We used a mixed effects analysis of covariance model for formal statistical inference.29,30 Our model included an overall intercept term, a BD versus control grouping factor, radial perinuclear distance, and a group by distance interaction. Radial distances were treated as nested within subjects, and mitochondrial density was re-expressed on the square root scale to conform more closely to the constant variance and Gaussian distribution assumptions of the general linear model. This morphological distribution will be referred to as the “bulls eye” method.

**Cell Profiler Analysis of Mitochondrial Distribution in Fibroblasts**

An independent morphometric distributional analysis was run using the software Cell Profiler31 (Broad Institute, Cambridge MA) on a subset of the BD and control fibroblast lines (4 BD and 4 Control lines) to determine the effect of lithium treatment on mitochondrial distribution. In Cell Profiler, three single images from five fields per line (24 to 80 cells per line), one from the nuclear channel stained with Hoechst, one from the phalloidin channel to mark cell edges, and one from the M7510 channel were loaded into the analysis software. To control for any difference in intensity of the raw M7510 images and to obtain an accurate measure of the distance from the middle of the nucleus to the pieces of the mitochondrial network images from this channel were pre-processed in Image J (National Institutes of Health, Bethesda MD) using the sharpen, find edges, and make binary commands as used in Koopman et al32 before loading images into Cell Profiler. The Cell Profiler Relate module was then used to measure the distance from the nucleus boundary as a metric of the extent of the mitochondrial network in each cell. This same analysis was also used to quantify the distribution of cytochrome C staining in fibroblasts.

**Lithium Treatment of Fibroblast Cultures**

Lithium carbonate (LiCO$_3$) (Sigma Aldrich, Allentown, PA), at levels chosen to match those typically seen in blood in treated patients, was added to fibroblast cultures plated at $0.8 \times 10^4$ cells per well. Media was changed every other day, and cells were treated with lithium for 5 days.

**TUNEL Staining for Detection of Apoptotic Changes**

TUNEL staining of fibroblast cultures plated at $2.5 \times 10^4$ cells per coverslip was done as in McPhie et al.26

**Measurement of ATP Levels in Fibroblasts**

Cells were plated at a density of $2.5 \times 10^4$ cells per well in 24-well plate. On the day after plating, media in the wells receiving serum starvation was changed to media containing 1.5% FBS and left for 24 hours before running the assay. Thirty minutes before the assay, 125 $\mu$mol/L H$_2$O$_2$ was added to the wells receiving serum starvation. ATP levels were measured with the ATPlite 1step kit (Perkin Elmer, Waltham MA) according to manufacturer’s instructions. Three triplicate samples (for a total of nine readings) of each fibroblast line were measured for baseline ATP levels and ATP levels after the stress of serum starvation and 125 $\mu$mol/L H$_2$O$_2$ treatment.
Cytochrome C Expression Levels and Intensity Correlation C (Colocalization) of M7510 Staining and Cytochrome C Immunocytochemistry

Fibroblasts were plated on coverslips and labeled with M7510 and fixed as above. The cells were then probed with a specific antibody to cytochrome C as above without the steps of visualizing β tubulin. Images taken at ×630 were analyzed with Cell Profiler’s Measure Object Intensity module, and integrated intensity levels of cytochrome C expression were compared between groups. Integrated intensity is defined as the sum of the pixel intensities within the mitochondrial network. A total of 295 control cells and 276 BD cells were analyzed. All images were taken with the same exposure time. The Measure Correlation module was used to determine the degree of overlap between the two mitochondrial labels.

Results

More Mitochondria of Smaller Area Are Observed in BD Brains

Previous studies have indicated mitochondrial differences in volume density and numbers of mitochondria in oligodendroglia from postmortem brain in Schizophrenia. In this study we first asked whether abnormalities in mitochondrial morphology and size were present in postmortem BD brain. Prefrontal cortex was chosen, as it is a documented site of abnormalities in brain structure and function in patients with BD. Ultrastructural examination of human brain tissue from healthy controls and BD brains showed that most mitochondria were small rounded structures of uniform size—0.2 to 1.5 μm in diameter with occasional rod-shaped profiles up to 2 μm in length. A smooth outer membrane with invaginated inner cristae bound each organelle and small mitochondrial profiles were seen interspersed with several larger profiles. Examples of tissue quality and mitochondrial profiles analyzed are illustrated in Figure 1, A and B. To quantify differences in mitochondrial size we measured mitochondrial area in electron micrographs from postmortem PFC of BD and control brains and binned the area measurements into equal quintiles. We found significantly smaller individual mitochondrial areas in the BD sample (P = 0.03; t test) (Figure 1C).

Because our sample size was small, we could not do a full analysis of drug effects on mitochondrial area. However, using a one-way analysis of variance to look at correlation with lithium treatment, one of the most common medications used for BD, no significant differences in mitochondrial area were seen between the four BD subjects on lithium versus the other BD subjects on other medications. Additionally, we did not see an association between mitochondrial area and age or postmortem delay (see supplemental Figure 1 at http://ajp.amjpathol.org).

Cytochrome C Distribution in BD and Control Prefrontal Cortex

We examined the samples for distributional differences of mitochondria in brain between the two groups. Previous studies by Kung and Roberts indicated mitochondrial distributional differences in the neuropil of the striatum of schizophrenic subjects. To assess this parameter in our BD samples, we used cytochrome C immunocytochemistry as a marker of mitochondria in postmortem brain tissue from the PFC. The analysis chosen, line profile analysis of stained cells in PFC, showed no significant differences between groups in mitochondrial distribution. In this analysis the cytochrome C pixel intensity curves varied greatly in shape across subjects in the BD group and across subjects in the control group. Although the two groups of curves appeared different from one another, subsequent comparison of the group-averaged pixel intensity curves yielded no significant differences in overall shape or at curve extrema (P > 0.30).
BD Fibroblasts and Lymphocytes Exhibit Alterations in Mitochondrial Morphology

Although the identification of altered mitochondrial morphology (size) in the brain could be of value in studying various disease states, demonstrating consistent and reliable morphological abnormalities and a possible disease-specific relationship linked to mitochondrial function is difficult using postmortem brain tissue. This is due largely to variations in postmortem interval and fixation conditions. For these reasons, we selected two non-neuronal cell types for study—fibroblasts and lymphocytes—harvested antemortem from individuals with BD. Using the mitochondrial-selective fluorescent marker, M7510, we found that fibroblasts (Figure 2A) and lymphocytes (Figure 2C) from healthy controls displayed a highly organized and interconnected mitochondrial network that extended from the nuclear envelope to the plasma membrane. The network was composed of elements that ranged in shape from single profiles to interconnected stacks. A typical single mitochondrion was thin and straight with a relatively constant diameter with frequent branching. A few smaller sphere or cup-shaped profiles were observed. In contrast to control cells, the same cell types from patients with BD exhibited alterations in mitochondrial distribution and morphology. Mitochondria in the BD cell lines appeared as dense or bulky networks with perinuclear clustering, which was observed in greater than 70% of fibroblasts and lymphocytes in samples from BD patients of all ages (Figure 2, B and D). In addition to differences in the spatial distribution and clustering of the mitochondrial reticulum seen in the BD cells, we observed that mitochondria within the clusters appeared as thick, short segments or punctate, cup- or ring-shaped fluorescent profiles (see supplemental Figure 2, A–C at http://ajp.amjpathol.org). These clusters varied in size but were larger than those in control cells.

Spatial Distribution of Mitochondria in Primary Cells from BD Patients

We quantified the spatial variation in the perinuclear distribution of mitochondria using a segmentation algorithm. Cells from individuals with BD possessed higher mitochondrial densities at every radial distance from the nuclear envelope within the close, 15–70 µm, range relative to healthy controls (Figure 3, A–C). Also, our mixed effects ANCOVA model confirmed that for BD subjects the average mitochondrial density by radial perinuclear distance declined more steeply than that of the control group (BD group, \(-1.15/\mu m\); control group, \(-1.07/\mu m\), diagnosis by distance interaction \(P = 0.0008\)) (Figure 3C). We also measured cell area, cell length, cell width, and cell perimeter of the fibroblasts used in this analysis to verify that the observed mitochondrial distributional differences were not simply a result of BD cells being smaller, shorter, or fatter. No significant differences in any of these measurements were seen based on diagnosis after two sample \(t\) tests.

Morphometric Analysis of Fibroblast Mitochondria at the Ultrastructural Level

Mitochondrial area, length, width, and aspect ratio were quantified using a two-dimensional analysis in our fibroblast samples. Although mitochondria in the BD samples showed a trend toward smaller mean area, as was seen in brain, this trend did not reach significance \((P > 0.3)\). Quantification of the number of ring- and cup-shaped mitochondrial profiles showed that BD samples had more ring- and cup-shaped mitochondria (mean of 5.7 in 7 control lines and 8.5 in 8 BD lines; two-sample \(t\) test, \(P = 0.034\)). Examples of ring- and cup-shaped mitochondria are shown in supplemental Figure 2, D–H at http://ajp.amjpathol.org.
Lithium Effects on Mitochondrial Distribution in Fibroblasts

Mood stabilizing drugs such as lithium are commonly used as treatments for BD, and all but one of our subjects with BD from whom fibroblasts were obtained were receiving lithium. Additionally, lithium treatment has been shown to have direct and indirect effects on mitochondria. Therefore, we asked whether lithium exposure could promote changes in mitochondrial structure similar to those seen in primary BD cells or, alternatively, whether exposure to lithium could rescue the distributional abnormalities in BD cells making them look more like controls. An age-matched subset of fibroblasts from both diagnostic groups (four BD and four Control) were treated acutely for 5 days with 1 mmol/L LiCO₃ (approximately the average serum level of lithium in patients). After distributional analysis we found that there was no significant effect of Li treatment in either group in a one-way analysis (or a group by Li treatment in a two-way analysis). There was, however, a significant difference in mitochondrial distribution between our two diagnostic groups (control versus BD; \( P < 0.05 \), two-sample \( t \) test) as observed in untreated cells (Figure 4, A versus C). The distributional analysis used for this part of the study, performed by Cell Profiler, provided an independent confirmation of disease-related differences observed in our bulls-eye analysis on a separate subset of cells from our BD and Control lines.

Cytochrome C Distributional Differences in BD Fibroblasts

To further characterize mitochondrial abnormalities seen in the BD fibroblasts, we asked whether the distribution of another component of mitochondria, cytochrome C, would also show differences in the BD lines. An age-matched subset of fibroblasts (four BD and four Control) was labeled with an antibody to cytochrome C. The immunocytochemical staining of this antibody under our staining protocol allowed us to visualize very specific staining of the mitochondrial network in the fibroblasts. After automated distributional analysis with Cell Profiler we saw a significant difference in cytochrome C distribution in the BD cell lines versus the lines from healthy subjects (two-sample \( t \) test, \( P < 0.0001 \)) (Figure 5). This result revealed that a separate component of the mitochondria also showed an altered distribution in the BD group, further confirming the Mitotracker M7510 result.

Cytochrome C Expression Levels and Intensity Correlation Analysis of M7510 Staining and Cytochrome C Immunocytochemistry

To determine whether the level of expression of cytochrome C was different between the two experimental groups, we quantified the staining intensity of an antibody specific for cytochrome C in images of BD and Control fibroblasts by an automated analysis. No significant differences in cytochrome C immunocytochemical staining intensity were found (20.61 ± 5.5 vs. 19.99 ± 6.1 Inten-
sity Units; two-sample \(t\) test NS; \(P = 0.63\). The values are reported as intensity units because microscopes are not calibrated to an absolute scale. Because similar localization (distributional) changes were seen with both the M7510, a probe that is sequestered in the mitochondria and binds to proteins that have thiol moieties, and a specific antibody to cytochrome C, we asked whether the two probes were labeling the same or different subsets of pieces of the mitochondrial network. Cell profiler automated intensity correlation analysis yielded a correlation coefficient of the two probes of 0.8 to 0.9, indicating that both probes are labeling the same pieces of the network.

**ATP Measurement of BD and Control Fibroblasts at Baseline and after Oxidative Challenge**

Abnormal mitochondrial shape and distribution may be associated with abnormal oxidative phosphorylation and production of ATP. Testing for this possibility, we observed no significant differences in ATP levels between the two groups either at baseline or after serum starvation and 125 μmol/L H₂O₂ challenge (see supplemental Figure 3 at http://ajp.amjpathol.org).

**Fibroblasts from Patients with BD Do Not Exhibit Substantial Cytoskeletal Alterations or Features of Apoptosis**

Mitochondria are known to move by closely associating with the cytoskeleton. We examined two key cytoskeletal proteins to see whether the mitochondrial distribution abnormalities might be a consequence of abnormalities in the cytoskeleton. Filamentous actin was examined by staining with phalloidin. Because mitochondria are directly associated with microtubules and microtubules have been demonstrated to have a direct role in mitochondrial biogenesis⁴⁰ we examined the microtubule cytoskeleton with an antibody to tubulin βI and βII. Using F-actin selective phalloidin (red label, Figure 6, A–D) or antibodies to β tubulin (red label, Figure 6, E–H), we found that in BD cells, abnormalities in mitochondrial shape and distribution (Figure 6, green label C, yellow-green label G) were not accompanied by substantial changes in the actin or tubulin cytoskeleton. Rather, BD cells showed a cytoskeletal pattern similar to that seen in healthy controls (Figure 6, B versus D, F versus H). In addition, DAPI labeling revealed no evidence of increased apoptosis or nuclear fragmentation in the BD cells. In addition, fluorescent staining of fibroblasts with the DNA-binding dyes, DAPI (A and C blue label) or bisbenzimide (not shown), revealed no evidence of increased apoptosis or nuclear fragmentation in the BD cells. Magnification: \(×1000\) (A–D), \(×630\) (E–H). Scale bar = 10 μm.

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**Figure 5.** Cytochrome C expression shows an altered distribution in BD fibroblasts. Distributional analysis of cytochrome C immunocytochemistry demonstrated an altered distribution of the mitochondria network in BD fibroblasts as compared with Control fibroblasts. (two-sample \(t\) test, \(P < 0.0001\)).

**Figure 6.** No substantial changes in the actin or tubulin cytoskeleton are seen that correlate with diagnosis in BD fibroblasts. Representative images of fibroblasts from patients with BD (C, D, G, and H) and healthy controls (A, B, E, and F) labeled with probes for mitochondria (M 7510 green A and C, and orange E and G), filamentous actin (red, phalloidin A–D), and tubulin β (red, E–H). The results confirm mitochondrial redistribution in the BD fibroblasts compared with controls. However, abnormalities in mitochondrial shape and distribution were not accompanied by changes in the actin cytoskeleton, observed by phalloidin, or antibodies to tubulin β I and II, when compared with healthy controls. In addition, fluorescent staining of fibroblasts with the DNA-binding dyes, DAPI (A and C blue label) or bisbenzimide (not shown), revealed no evidence of increased apoptosis or nuclear fragmentation in the BD cells. Magnification: \(×1000\) (A–D), \(×630\) (E–H). Scale bar = 10 μm.
Abnormalities of Other Membranous Organelles

Inherited or environmentally induced abnormalities of mitochondria might affect other subcellular organelles, as well. However, in contrast to the altered cellular arrangement of the mitochondria in the BD cells, other organelles such as the plasma membrane, ER, and Golgi apparatus appeared similar to those in control cells (data not shown).

Discussion

To our knowledge, this study is the first to demonstrate that cells (neurons) in the prefrontal cortices of postmortem brain from patients with BD and peripheral cells from living BD patients display abnormalities in the morphology (size and shape) and intracellular distribution of mitochondria. We observed two different mitochondrial abnormalities in BD samples. In the PFC of BD brain, there were significantly more mitochondria of smaller size than in age-matched controls. In peripheral cells from BD patients, we quantified an altered distribution of the mitochondrial network such that there was a significantly faster drop off of mitochondria as one measures outward from the nucleus toward the cell periphery. This result was consistent between two independent mitochondrial probes. There were also more ring- or cup-shaped mitochondrial profiles in the BD fibroblast samples. Although the number of ring- and cup-shaped mitochondria counted in our sample is modest when compared with the total number of mitochondria counted, the BD samples had significantly more of these atypically shaped mitochondria, which may reflect subtle changes in the critical network architecture of mitochondria in the cells. Mitochondrial size, shape, and distribution typically differ between tissues, and the same factors may not cause exactly the same mitochondrial changes in different tissues. However, the abnormalities of mitochondrial size observed in BD brain may also be features of mitochondria in peripheral cells in BD, in which we saw a trend to abnormalities of size. Larger sample sizes than those available would be needed to confirm this difference. Similarly, the abnormalities of mitochondrial distribution observed in peripheral cells might also be features of mitochondrial networks in brain cells in BD. Although the analysis chosen for the present study was not able to detect a distributional difference, a more comprehensive study on a larger number of samples than those available for the present study might define disease-related mitochondrial distributional differences in the brain.

Alterations in mitochondrial shape and distribution could be the underlying cause of energy dysfunction in BD, as observed in imaging studies, or may be secondary to other factors associated with BD. In either case, the mitochondrial abnormalities identified here may have implications for cell plasticity, resilience, and survival in patients with BD. They also may identify a pathognomonic marker for BD or related illnesses and might, with further study, provide clues for studies targeted at the repair of the mitochondrial dysfunctions observed in BD.

Limitations of the results reported here include the modest number of tissue samples available. Our study also is limited in that only one brain region was examined with only a single analysis method. In addition, details of the medical history of the patients who provided the fibroblast samples, such as when in the disease the sample was taken, were not available. However, findings of alterations in mitochondria were consistent across three different tissues—brain, fibroblasts, and lymphocytes—from two different tissue collections. The size and distributional abnormalities observed were consistently seen in tissues from individuals with BD and were seen with a much lower frequency in psychiatrically healthy age-matched controls. Not all of the subjects were receiving the same medications, and the abnormality could not be replicated in control samples or rescued in BD samples by exposure of fibroblasts in vitro to the medication the subjects most commonly received (i.e., lithium). This suggests, but does not prove, that the effect is a corollary of illness rather than a consequence of treatment. Of course, even if the abnormality were drug related, it also would be of interest, as it might provide a clue to the mechanisms by which medications for BD act. Additionally, it is interesting that the mitochondrial abnormalities persist in vitro in cell lines and continue through many passages. A potential explanation for this is that the cell lines may carry genetic defects in nuclear or mitochondrial DNA similar to the mitochondrial DNA deletions seen by Kato et al in BD brains. Future work and nuclear and mitochondrial DNA sequencing may provide more information on this mechanism.

Evidence from a number of studies now shows that mitochondrial morphology is linked to energy metabolism and that changes in mitochondrial morphology reflect alterations in energy metabolism. Although the fibroblast lines in the present study did not show differences in ATP levels at baseline or with one level of oxidative challenge, a further or alternative stressor may bring out energy metabolism alterations. Alternatively, although an alteration in mitochondrial function would be consistent with our observations of altered mitochondrial location, it may also be that the only change present in the BD cells is a change in mitochondrial distribution. It has been suggested that mitochondrial distributional or localization changes alone may be the cause of psychiatric and other illnesses, in that the mitochondria in BD cells may not be able to deliver energy and other substrates to places in the cell that need it, thus leading to disease. Another point to be considered is that although we saw no differences in numbers of apoptotic cells in our samples, changes in mitochondrial distribution might render BD fibroblasts more sensitive or resistant to stress-induced apoptosis.

Mitochondria continually fuse and divide and the overall morphology of the mitochondrial network, which is prominent in eukaryotic cells, is modified by the balance of these two processes, which are regulated through the interactions of several well known mitochondrial GT-Pases. Deviation in either fission or fusion can result in cellular compromise leading to reduced cell function or even apoptosis and cell death.
Mutations in genes encoding mitochondrial fission or fusion proteins have been associated with a number of disorders of the nervous system and can directly affect mitochondrial function. Overexpression of fusion proteins such as MFN2 or the down-regulation of fission proteins such as DRP1, affects mitochondrial function by reducing mitochondrial respiration and ATP levels and affects mitochondrial morphology leading to decreased mitochondrial branching, spheroid mitochondrial tubules, and ring-shaped mitochondria and fragmentation. G72 G72 a candidate BD susceptibility gene has also been shown to affect mitochondrial fragmentation. Since one explanation for the increased numbers of smaller mitochondria seen in BD PFC is increased fragmentation, G72 or related proteins may play a role. There is also an ever growing set of studies indicating that dysregulation of mitochondrial fission and fusion may be important causative factors in the neurodegeneration seen in Alzheimer’s, Parkinson’s and Huntington’s disease.

Given the influence of fission and fusion events in mitochondrial structure, alterations in these processes might explain our findings of abnormal mitochondrial morphology and distribution in patients with BD and are appropriate targets for further study. Alternatively, a failure of mitochondrial transport due to a disruption in cytoskeletal proteins (i.e., actin, tubulin, or motor proteins such as kinesin) could contribute to altered mitochondrial clustering. Although our examination of the cytoskeleton in primary fibroblasts from patients with BD did not show any substantial changes in cytostructure, a more detailed study centered on cytoskeletal proteins may shed light on more subtle alterations between the groups.

Cumulatively, our findings indicate that changes in mitochondrial shape and distribution are a consistent feature of neurons and primary non-neuronal cells from individuals with BD. Furthermore, our data are consistent with evidence from MR and postmortem studies linking reduced energy production and mitochondrial dysfunction to BD. Because of the well-characterized link between mitochondrial structure and function, our current findings of mitochondrial structural changes in cells from patients with BD support the hypothesis that there are abnormalities of mitochondria in BD. Specifically, our findings support the possibility that alterations in mitochondrial structure and function may underlie abnormal brain development or activation in BD. As mitochondria are crucial for cell growth, function, and survival, they may also play a role in the neuronal and/or glial compromise or cell deficits seen in BD. The peripheral cell models of BD described here may prove valuable in further defining the fundamental relationships between mitochondrial structure and function in BD, examining how these interactions influence energy metabolism and elucidating molecular changes contributing to the expression of BD. In time, further study of the phenomena observed could lead to the identification of specific pathogenic and pathophysiological features of BD, useful for diagnosis and for the design of treatments targeted specifically to the abnormalities that underlie BD.

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