Asthma and chronic obstructive pulmonary disease (COPD) are inflammatory airway diseases that are characterized by different patterns of airway remodeling (1). Nevertheless, the decrease in lung function that characterizes both diseases is associated with an increased mass of bronchial smooth muscle (BSM) (2, 3), which is likely to be the most important abnormality responsible for the airway narrowing observed in response to bronchoconstricting stimuli (4). The mechanisms underlying such remodeling of smooth muscle remain largely unknown. On the one hand, in smooth muscle from asthmatic patients, excessive in vitro proliferation of BSM cells has been demonstrated (5–7). In addition, decreased apoptosis of BSM cells has also been demonstrated, although this was in a rat model of experimental asthma (7). On the other hand, in COPD, smooth muscle remodeling appears limited to airways distal to the fourth generation (3, 8). Although an increase in TGF-β1 production by BSM cells has been proposed, such localization and the complete mechanism remain unexplained (9). Whatever its cause, BSM remodeling is poorly sensitive to current therapeutics in both asthma and COPD.

Mitochondria play a major role in both cell proliferation and apoptosis (10, 11). In cancer, for instance, targeting of mitochondrial function and mitochondrial antiapoptotic protein bcl-2 has been used to either suppress the proliferation of tumor cells (10) or induce cell apoptosis in solid tumors (12). Mitochondria are also involved in other diseases, such as neuron-degenerative
diseases (11). However, their role in asthma or COPD remains to be investigated. We hypothesized that mitochondrial activation in BSM from asthmatic or COPD patients could contribute to smooth muscle remodeling. To investigate this hypothesis, we have compared mitochondrial mass, activity, and biogenesis in BSM obtained from asthmatics, COPD patients, and normal controls. We describe that both the mitochondrial mass and oxygen consumption were higher in the BSM from asthmatic subjects than in that from both COPD and controls. This feature, which is specific to asthma, is related to an enhanced mitochondrial biogenesis as a consequence of an increase in extracellular calcium influx upon activation of asthmatic BSM cells. We also demonstrate a specific mitochondria-dependent pathway for asthmatic BSM cell proliferation. Targeting such a pathway may thus represent a new approach for the treatment of airway remodeling in asthma.

RESULTS

Clinical populations

The clinical characteristics of all subjects are shown in Table I. All of the 14 severe persistent asthmatics were lifelong non-smokers and received stable treatments, including oral or inhaled corticosteroids and β2 agonists. 10 of them were atopic. The 17 moderate to severe COPD patients were either current or former smokers, and 9 of them received stable treatments, including oral or inhaled corticosteroids and/or β2 agonists. None of the asthmatic or COPD patients experienced a recent (<3 mo) exacerbation of the disease. The mean duration of the disease in asthmatic and COPD patients was 26 ± 4.6

and 18 ± 2.8 yr, respectively. Of the 19 control subjects who received no treatment, 8 of them were lifelong nonsmokers, whereas 11 were former smokers.

Table I. Clinical and functional characteristics of subjects

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Patients with asthma</th>
<th>Patients with COPD</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of patients</td>
<td>14</td>
<td>17</td>
<td>19</td>
</tr>
<tr>
<td>Age (yr)</td>
<td>42.3 ± 5.7</td>
<td>59.8 ± 3.0</td>
<td>60.4 ± 2.5</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>26.9 ± 1.1</td>
<td>24.0 ± 0.9</td>
<td>27.4 ± 1.1</td>
</tr>
<tr>
<td>Smoking history</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pack years</td>
<td>0 ± 0</td>
<td>46.3 ± 4.8</td>
<td>19.5 ± 5.7</td>
</tr>
<tr>
<td>Current (no. of patients)</td>
<td>0</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Former (no. of patients)</td>
<td>0</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Years since quitting</td>
<td>-</td>
<td>11.4 ± 4.8</td>
<td>13.1 ± 3.8</td>
</tr>
<tr>
<td>Treatments</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LABA (no. of patients)</td>
<td>11</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>ICS (no. of patients)</td>
<td>10</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>OCS (no. of patients)</td>
<td>6</td>
<td>3</td>
<td>0</td>
</tr>
</tbody>
</table>
| FEV₁
| Liters               | 2.2 ± 0.2            | 1.8 ± 0.1          | 2.9 ± 0.1 |
| Percentage of predicted value | 83.1 ± 6.3           | 60.2 ± 2.5         | 101.7 ± 3.2 |
| FEV₁; FVC ratio (% of FVC) | 71.9 ± 3.7           | 56.3 ± 2.2         | 81.8 ± 1.7 |
| FEF 25–75
| Liters sec⁻¹         | 1.9 ± 0.3            | 0.9 ± 0.1          | 2.9 ± 0.1 |
| Percentage of predicted value | 54.6 ± 7.8           | 28.6 ± 3.1         | 90.2 ± 4.0 |

Data are the mean ± the SEM. BMI, body mass index; LABA, long-acting β2 agonist; ICS, inhaled corticosteroid; OCS, oral corticosteroid; FEV₁, forced expiratory volume in one second; FVC, forced vital capacity; FEF 25–75, forced expiratory flow between 25 and 75% of FVC.
Mitochondrial mass and activity are increased only in the BSM of asthmatics

A morphological analysis of BSM was performed in the three groups of subjects (Fig. 1, A–C). BSM mass was increased in both asthmatic and COPD patients as compared with controls (Fig. 1 D; P = 0.01). Ultrastructure of BSM mitochondria was then compared in asthma and COPD to control subjects (Fig. 2, A–C). The number of mitochondrial sections was higher in the BSM of asthmatics than in that of both COPD and controls (Fig. 2 D; P < 0.001). The mean area per section was, however, unchanged (0.1 ± 0.007, 0.1 ± 0.016, and 0.1 ± 0.009 μm² for asthmatics, COPD, and controls, respectively; analysis of variance [ANOVA] P = 0.87). Collectively, these results favor the hypothesis that the increase in mitochondrial mass in the BSM of asthmatics is related to an increase in number rather than in individual size. Similar results were obtained with cultured growth-arrested BSM cells (Fig. S1, A–D, available at http://www.jem.org/cgi/content/full/jem .20070956/DC1; P = 0.02). There was a significant increase in the mitochondrial density in asthmatic BSM cells both ex vivo (Fig. 2 E; P = 0.01) and in vitro (Fig. S1 E; P = 0.01) as compared with both COPD and controls. Based on electronic microscopy, the mitochondrial density has been shown to reflect the mitochondrial activity (13, 14). To assess the smooth muscle specificity of these results, we also analyzed ultrastructural mitochondrial parameters in other cell types from the same bronchial specimens. There was no difference between the three groups, in terms of both number (P = 0.35 and P = 0.89) and density of mitochondria (P = 0.47 and P = 0.87) in endothelial and epithelial cells, respectively (unpublished data). The increased mass of asthmatic BSM mitochondria was further confirmed, in vitro, by an increase in the porin content compared with that of both COPD and controls (Fig. 2 F; P = 0.001). We also found within the asthmatic population that both duration of the disease and forced expiratory volume in one second (FEV₁)/forced vital capacity

Figure 2. Increased mitochondrial mass and activity in asthmatic BSM. Representative electronic microscopic images from bronchial sections were obtained from an asthmatic (A), a COPD (B), or a control subject (C) and observed at 26,000× magnification. Some smooth muscle mitochondria were visualized (arrows). Bars, 0.2 μm. The number (D) and the density (E) of mitochondria were assessed from electronic microscopic images (n = 4 for each population). Mitochondrial mass was assessed by the porin content using Western blot (F; n = 8 for asthmatics, n = 5 for COPD, and n = 7 for controls). Endogenous cellular oxygen consumption was evaluated by oxygraphy (G; n = 5 for asthmatics, n = 4 for COPD, and n = 4 for controls). BSM cells (BSMC) were obtained from asthmatic (black columns), COPD (gray columns), and control subjects (white columns). Data are the mean ± the SEM. *, P < 0.05 between populations using ANOVA with the use of Bonferroni’s test.
Mitochondrial biogenesis is increased in asthmatic BSM cells through a calcium-dependent pathway

We next analyzed the mitochondrial network by confocal microscopy (Fig. 3, A–C). Surprisingly, asthmatic BSM cells presented a typical aspect of intense mitochondrial biogenesis, as shown by the presence of several budding areas with intense dots, and a wider network with increased ramifications (Fig. 3 A). Because the mitochondrial transcription factor A (mtTFA) is the main factor involved in mitochondrial biogenesis, we measured its protein content and transcription level in the three groups of BSM. As compared with both controls and COPD, mtTFA was increased in asthmatic BSM cells (Fig. 3, D and E). Upstream mechanisms that would explain such an increased mitochondrial biogenesis were then examined. The two transcription factors nuclear respiratory factor-1 (NRF-1) and peroxisome proliferator–activated receptor γ coactivator–1α (PGC-1α) were both elevated in asthmatic BSM cells at both the protein and the transcription levels (Fig. 3, D and E). Among the various factors that could activate PGC-1α, we observed that the calcium/calmodulin–dependent protein kinase IV (CaMK-IV) was phosphorylated and thus activated in asthmatic BSM cells compared with controls and COPD (Fig. 4 A). We thus investigated whether calcium homeostasis was deregulated in asthmatic BSM cells using microspectrofluorimetry. The resting calcium concentration was consistent in asthmatic (125 ± 8 nM), COPD (127 ± 7 nM), or control BSM cells (133 ± 4 nM; P = 0.60). However, the calcium response to acetylcholine was significantly altered in asthmatic BSM cells (Fig. 4, B–D). Whereas the amplitude of the calcium rise was unchanged (275 ± 20, 284 ± 43, and 272 ± 27 nM in asthmatics, COPD, and controls, respectively; P = 0.95), the area under the curve was significantly increased in asthmatic BSM cells (Fig. 4 E). A similar result was obtained in asthmatic BSM cells stimulated with histamine, confirming that it was not agonist-specific (Fig. S3, A–D, available at http://www.jem.org/cgi/content/full/jem.20070956/DC1). An enhanced calcium influx in asthmatic BSM cells accounts for this result, as both removal of extracellular calcium using EGTA and blockade of calcium channels using methoxyverapamil (D600) abolished such abnormal calcium responses in asthmatic BSM cells (Fig. 4 E and Fig. S3 D). Blockade of calcium influx using D600 also inhibited the activation of CaMK-IV (Fig. 4 A) and the subsequent activation of mitochondrial biogenesis through...
PGC-1α, NRF-1, and mtTFA (Fig. 5, A–C), leading to the increase in the mitochondrial mass as assessed by the porin content (Fig. 5 D). Thus, these results indicate that an enhanced extracellular calcium influx, specific to asthmatic BSM cells, is the initial priming event leading to an increased mitochondrial biogenesis and mass.

**Only asthmatic BSM cell proliferation is mitochondria dependent**

To further assess the specific role of mitochondria in asthmatic BSM proliferation contributing to airway remodeling, we next compared BSM cell growth in the presence or absence of anaerobic glycolysis. For this purpose, BSM cell proliferation curves were plotted in the three groups of subjects using either glucose or galactose in the culture medium (Fig. 6, A and B). In the presence of glucose, which allows ATP to be produced by both aerobic and anaerobic glycolysis, asthmatic BSM cell growth was significantly increased compared with that of COPD and control subjects (Fig. 6 A; P = 0.02), with a concomitant decrease in the doubling time (Fig. 6 C; P = 0.02). When galactose, which only allows cells to produce ATP by mitochondrial oxidative phosphorylations, was substituted for glucose, the doubling time of the cell growth from both COPD (P = 0.02) and control subjects (P = 0.02) significantly increased, whereas that of asthmatic cells remained constant (Fig. 6, B and C). Because an increase in cell growth can be related to a decreased apoptosis and/or an increased proliferation, we analyzed Annexin V binding and BrdU incorporation, respectively. Taking into account the percentage of Annexin V–positive cells, spontaneous apoptosis was not altered in asthmatic BSM cells as compared with that of COPD and controls (Fig. S4, available at http://www.jem.org/cgi/content/full/jem.20070956/DC1). However, BrdU incorporation increased in BSM cells from asthmatics as compared with that in both COPD and controls in the presence of glucose (Fig. 6 D; P = 0.01). Incubation for 11 d in the absence of glucose significantly inhibited BrdU incorporation in BSM cells from controls (P = 0.02) and COPD patients (P = 0.02), but not in that from asthmatics (Fig. 6 D; P = 0.99).

Collectively, these results demonstrate that the proliferation of asthmatic BSM cells mainly uses mitochondrial-dependent...
Increased mitochondrial mass explains increased asthmatic BSM cell proliferation

To determine whether the increased mitochondrial biogenesis found in asthmatic BSM cells is a cause or a consequence of the asthmatic BSM cell increased proliferation, three alternative approaches were used. First, mitochondria-deficient BSM cells were generated using culture with ethidium bromide, as previously described (10, 15). After 16 d of culture, all of the asthmatic BSM cells died, whereas those from both COPD patients and controls were still alive up to 30 d in the ethidium bromide medium (unpublished data). 6 d of incubation with ethidium bromide significantly decreased the amount of porin in the three groups of subjects (Fig. 7). Fig. 6 C demonstrates that ethidium bromide significantly increased the doubling time of asthmatic BSM cells (P < 0.001), but did not change that of both COPD (P = 0.15) and control cells (P = 0.09). Similarly, proliferation of mitochondria-deficient asthmatic BSM cells decreased (P < 0.001), whereas that of COPD and controls remained unchanged (Fig. S5, available at http://www.jem.org/cgi/content/full/jem.20070956/DC1).

However, ethidium bromide also decreased BrdU incorporation of BSM cells from the three groups of subjects (Fig. 6 D). In a second alternative approach, we stimulated mitochondrial biogenesis using cyclic GMP for 6 d, as previously described (16). The amount of mitochondria increased in such stimulated BSM cells from both COPD and controls (Fig. 7; P = 0.02 for both). Because the amount of mitochondria was already up-regulated in BSM cells from asthmatics, cyclic GMP failed to additionally increase the porin content (P = 0.2). Cyclic GMP significantly decreased the doubling time (P = 0.01), and increased the BrdU incorporation (P = 0.02) and the proliferation of BSM cells from COPD patients (P = 0.01) (Fig. 6, C and D, and Fig. S5). Similarly, cyclic GMP also decreased the doubling time (P < 0.001) and increased the BrdU incorporation (P < 0.001) and the proliferation of BSM cells from controls (P < 0.001; Fig. 6, C and D, and Fig. S5). However, cyclic GMP had no effect in asthmatic BSM cell growth and proliferation. Finally, in a last approach, we analyzed the effect of altering calcium homeostasis on the proliferation of BSM cells. D600, which blocks calcium influx (Fig. 4 E) and the resulting increase in mitochondrial biogenesis and content only in asthmatic BSM cells (Fig. 5), also significantly inhibited the increased BrdU incorporation (P < 0.005 between experimental conditions versus glucose within a population using paired Student’s t tests).

oxidative phosphorylations, whereas that of COPD and control subjects mainly uses mitochondrial-independent anaerobic glycolysis.

Increased mitochondrial mass explains increased asthmatic BSM cell proliferation

To determine whether the increased mitochondrial biogenesis found in asthmatic BSM cells is a cause or a consequence of the asthmatic BSM cell increased proliferation, three alternative approaches were used. First, mitochondria-deficient BSM cells were generated using culture with ethidium bromide, as previously described (10, 15). After 16 d of culture, all of the asthmatic BSM cells died, whereas those from both COPD patients and controls were still alive up to 30 d in the ethidium bromide medium (unpublished data). 6 d of incubation with ethidium bromide significantly decreased the amount of porin in the three groups of subjects (Fig. 7). Fig. 6 C demonstrates that ethidium bromide significantly increased the doubling time of asthmatic BSM cells (P < 0.001), but did not change that of both COPD (P = 0.15) and control cells (P = 0.09). Similarly, proliferation of mitochondria-deficient asthmatic BSM cells decreased (P < 0.001), whereas that of COPD and controls remained unchanged (Fig. S5, available at http://www.jem.org/cgi/content/full/jem.20070956/DC1).

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incorporation in BSM cells from asthmatics, thus confirming that this enhanced cellular calcium influx represents the initial priming event (Fig. 8).

DISCUSSION

This study indicates that whereas both asthma and COPD are characterized by BSM remodeling, a specific mitochondria-dependent pathway is required for BSM proliferation only in asthma. This pathway is initiated by an altered calcium homeostasis, upon the activation of asthmatic BSM cells (Fig. S6, available at http://www.jem.org/cgi/content/full/jem.20070956/DC1). Proliferation of BSM in both health and COPD is, at the very least, less mitochondria dependent. Thus, these results suggest that mitochondria may represent a specific new therapeutic target in airway remodeling in asthma.

In this study, we paid special attention when comparing data from severe asthmatics to that of COPD patients because both diseases have been shown to present smooth muscle remodeling (2, 3, 17). Using a variety of different experimental approaches, we provide evidence that asthmatic BSM express a higher number of active mitochondria and a clear aspect of intense mitochondrial biogenesis. In our study, we assessed mitochondrial mass using various parameters, including the number of mitochondria by electron microscopy both ex vivo and in vitro, the mitochondrial network by confocal microscopy, and the porin content by Western blot. All of these methods provided consistent results. We also found significant correlations between in vitro BSM porin content, which is a relevant quantitative estimate of mitochondrial mass, and both the duration of the disease and the FEV 1 /FVC ratio within the asthmatic population. These correlations further support our hypothesis because, on the one hand, the longer the duration of the disease that is known to favor airway remodeling (18), the higher the mitochondrial mass, and on the other hand, the lower the FEV 1 /FVC ratio, which also reflects airway remodeling, the higher the mitochondrial mass. Moreover, to obtain a comprehensive assessment of mitochondrial content, we additionally measured the expression level of mtTFA by both quantitative RT-PCR and Western blot to assess mitochondrial respiratory chain content, as mitochondrial overall content and respiratory chain content are not always linked (19).

The increased number of mitochondria was limited to asthmatic BSM cells compared with endothelial and epithelial cells from asthmatic, COPD, or control subjects. However, two characteristics of such asthmatic populations deserve further comment. First, it is unlikely that asthma treatments interfered with the observed changes because, on the one hand, 9/17 COPD patients took treatments similar to that of asthmatics, and, on the other hand, no difference has been found between controls and COPD. Second, the younger mean age of the asthmatic population is unlikely to contribute to this increase in mitochondrial content because (a) this increase persisted when comparing asthmatics to a subgroup of nonsmoking controls whose mean age was similar, and (b) in skeletal muscle, if not any other muscle, age does not influence mitochondrial content (20). To the best of our knowledge, this is the first description of such mitochondrial characteristics in any type of smooth muscle cells with potential pathophysiological implications. It has been previously reported that in a variety of differentiated tissues, a mitochondrial dysfunction increases mitochondrial biogenesis, suggesting a cellular compensatory mechanism (21). However, there was no mitochondrial dysfunction in our study because the respiration of asthmatic BSM mitochondria was efficiently coupled. Similarly, artificial activation of mitochondrial biogenesis of mouse neonatal cardiac myocytes largely induced a coupled respiration (22).

Mitochondrial biogenesis is controlled by many factors. PGC-1α is a well-known master activator of mitochondrial biogenesis through the production of both NRF-1 and mtTFA in various cell types, including myoblast (23), fibroblast, or adipocytes (24). In this study, we demonstrated that such a cascade is activated in asthmatic BSM cells, as shown by the concomitant up-regulation of PGC-1α, NRF-1, and mtTFA. Moreover, cyclic GMP has been shown to activate PGC-1α and mitochondrial biogenesis in various cell lines, including U937, L6, and PC12 (16, 25). We have observed that cyclic GMP induces mitochondrial biogenesis in both control and COPD BSM cells, but not in asthmatic BSM cells. These findings suggest that mitochondrial biogenesis in asthmatic BSM cells may already be up-regulated. In this connection, cyclic GMP also improved cell proliferation of control and COPD BSM cells, but not those of asthmatics. It could be argued that cyclic GMP regulates many different genes in smooth muscle, such as vascular smooth muscle (26). However, cyclic GMP has been shown to inhibit, rather than enhance, the proliferation of vascular smooth muscle cells, as well as that of mesangial cells and various fibroblasts (26). In addition, direct improvement of mitochondrial biogenesis by transgenic overexpression of PGC-1α activates skeletal muscle atrophy (27) and cardiac muscle dysfunction (22). Conversely, cyclic GMP increases the proliferation of endothelial cells, but the role of mitochondria in this phenomenon has not been investigated so far (28).

PGC-1α activation can be calcium-dependent (29). In this study, we found that CaMK-IV was more phosphorylated in

Figure 8. Effect of methoxyverapamil (D600) on BSM cell proliferation. BSM cell proliferation was measured using BrdU incorporations. Cells were cultured in the absence (−) or presence (+) of 1 μM D600 for 48 h. BSM cells were obtained from asthmatic (black columns; n = 4), COPD (gray columns; n = 4), and control subjects (white columns; n = 4). Data are the mean ± the SEM. *, P < 0.05 between populations within an experimental condition using ANOVA with the use of Bonferroni’s test. †, P < 0.05 between the absence and the presence of D600 within a population using paired Student’s t tests.
asthmatic BSM cells than in both controls and COPD. Rises in calcium concentration have been previously shown to activate CaMK-IV in other cell types, including skeletal muscle cells (29) or osteoclasts (30). Interestingly, we also demonstrated that calcium homeostasis in asthmatic BSM cells was altered, thus providing a mechanistic explanation for the increased activation of calcium-dependent signaling enzymes such as CaMK-IV (31). A D600-sensitive calcium influx accounts for such asthma-induced alteration in calcium homeostasis. We also provide evidence that this calcium influx was the initial priming event because, when blocked, mitochondrial biogenesis and subsequent asthmatic BSM cell–increased proliferation was inhibited. Whereas blockade of such influx may be beneficial, clinical responses remain to be examined because previous studies using methoxyverapamil were focused on short-term effects for up to 4 wk and did not assess airway remodeling (32, 33).

Finally, we generated mitochondria–deficient BSM cells by depletion of mitochondrial DNA with ethidium bromide, which is a potent inhibitor of mitochondrial DNA replication and transcription (10, 15). Mitochondria–deficient BSM cells from asthmatics were unable to proliferate, thereby confirming the importance of mitochondria in asthmatic BSM cell proliferation. Thus, the increased mitochondrial biogenesis observed in asthmatic BSM cells appears to be a cause rather than a consequence of the asthmatic BSM cell increased proliferation. However, it is well known that training can increase mitochondrial biogenesis in skeletal muscle (34). It is unlikely that a similar phenomenon appears to the BSM from asthmatics. In this study, none of the asthmatics presented recent exacerbations, and all of these patients were treated by relaxant β2 agonists.

In conclusion, this study reveals that asthmatic BSM is characterized by an increased mitochondrial biogenesis that, in turn, enhances cell proliferation. Mitochondria may thus represent a new target for the treatment of asthmatic smooth muscle remodeling. Further studies are required to assess whether drugs interacting with mitochondrial biogenesis, including drugs acting at the site of calcium homeostasis, can prevent and/or reverse BSM remodeling in asthma.

MATERIALS AND METHODS

Study populations. A total of 14 patients with severe persistent asthma, 17 moderate to severe COPD patients, and 19 normal controls were prospectively recruited from the Centre Hospitalier Universitaire of Bordeaux according to both the Global Initiative for Asthma (35) and the Global Initiative for Chronic Obstructive Lung Disease guidelines (36). All subjects gave their written informed consent to participate in the study, after the nature of the procedure had been fully explained. The study followed recommendations outlined in the Helsinki Declaration and received approval from the local ethics committee. Bronchial specimens from all subjects were obtained by either fiberoptic bronchoscopy or lobectomy, as previously described (37, 38). All experiments were performed on phenotypically confirmed smooth muscle cells between passages 2 and 4. Cells were seeded on glass coverslips for confocal microscopy and microspectrofluorimetry, on culture flasks for electron microscopy and protein extraction, or on culture plates for RNA extraction. We used cells transfected with the plasmid mitochondrion-targeted GFP (mito-GFP) (14) to obtain confocal images of the mitochondrial network. Such images were acquired with Fluoview laser scanning microscope (Nikon) and reconstituted in three-dimensional images using Imaris Software (Bitplane) (38). Immunoblotting was performed on cell protein extracts (38, 40) using primary antibodies directed against porin, mtTFA, NRF-1, and PGC-1α. After reverse transcription, real-time quantitative PCR was performed on a Rotor-Gene 2000 (Corbett Research) (37, 38, 40), using appropriate primers designed to target mtTFA, NRF-1, or PGC-1α. Endogenous cell oxygen consumption, as well as coupling degree, were assessed in a thermostatically controlled chamber equipped with a Clark oxygen electrode (Oxygraph System; Hansatech), as previously described (14).

Cell calcium was assessed by microspectrofluorimetry using Indo-1 probe, as previously described (40, 41). Cell proliferation was evaluated using both BrdU incorporation and cell counting. Cell apoptosis was finally studied using FITC–Annexin V and flow cytometry. A complete description of all methods is available in the Supplemental materials and methods.

Statistical analysis. The statistical analysis was performed with NCSS 2001 software. Comparison between the three groups was performed by means of ANOVA, with the use of Bonferroni’s test for multiple comparisons or paired Student’s t tests. Values are presented as the mean ± the SEM. A Pearson correlation matrix was built between in vitro and in vivo measurements. A P value <0.05 was considered statistically significant.

Online supplemental material. Fig. S1 provides ultrastructural characteristics of mitochondria from pellets of BSM cells. Fig. S2 demonstrates coupled endogenous BSM cell respiration. Fig. S3 provides BSM cell intracellular calcium responses to histamine. Fig. S4 shows spontaneous BSM cell apoptosis. Fig. S5 shows the effect of ethidium bromide and cyclic GMP on BSM cell proliferation. Fig. S6 illustrates the mechanisms of BSM proliferation in asthma. The online version of this article is available at http://www.jem.org/cgi/content/full/jem.20070956/DC1.

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The authors have no conflicting financial interests.

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REFERENCES


SUPPLEMENTAL MATERIALS AND METHODS

Study population
A total of 14 patients with asthma, 17 COPD patients, and 19 normal controls were prospectively recruited from the Centre Hospitalier Universitaire of Bordeaux.

Asthmatic patients
Asthmatic patients were enrolled using the following inclusion criteria. Patients >18 yr had to exhibit characteristic symptoms (i.e., wheezing and breathlessness), as well as bronchial hyperresponsiveness confirmed either by a significant improvement by >15% in the forced expiratory volume in 1 s (FEV₁) 10 min after the inhalation of 200 µg of salbutamol, or a provocative concentration of methacholine required to lower the FEV₁ by 20% (PC20) of <4 mg/ml according to the American Thoracic Society criteria (1). Patients must all be graded as severe persistent asthmatics according to the Global Initiative for Asthma guidelines based on both clinical features and daily medication regimen (2). All patients must also be lifelong nonsmokers with no recent exacerbation (<3 mo). Exclusion criteria were recent bacterial or viral infections (<3 mo).

COPD patients
COPD patients were enrolled using the following inclusion criteria. Patients >18 yr had to present chronic symptoms (i.e., cough and breathlessness) and poorly reversible airflow obstruction defined by both an improvement of <10% in the FEV₁ 10 min after the inhalation of 200 µg of salbutamol or ipratropium bromide, and a postbronchodilator FEV₁/forced vital capacity ratio <70% according to the Global Initiative for Chronic Obstructive Lung Disease (GOLD) guidelines (3). Patients must be graded as moderate to severe COPD according to the GOLD guidelines (3). Exclusion criteria were recent bacterial or viral infections (<3 mo).

Control subjects
Control subjects were enrolled using the following inclusion criteria. Subjects >18 yr had to have normal lung function test and have underwent a fiberoptic fibroscopy or a thoracotomy because of hemoptysis or abnormal image on x ray. They must be asymptomatic without any treatment, and only those in whom fiberoptic investigation and bronchial mucosa were normal were selected as controls. Exclusion criteria were any comorbidity, such as asthma or COPD, and recent bacterial or viral infections (<3 mo).

All subjects gave their written informed consent to participate in the study after the nature of the procedure had been fully explained. The study followed recommendations outlined in the Helsinki Declaration and received approval from the local ethics committee.

Study procedures

Bronchial specimens.
Bronchial specimens were obtained by either fiberoptic bronchoscopy or lobectomy, as previously described (4, 5). Fiberoptic bronchoscopy was performed after the anesthesia of the upper airways was achieved with lidocaine 10% spray. The fiberoptic bronchoscope (FB-5V; Pentax) was introduced, and biopsies were taken from various bronchial carinae from the middle lobe.

Because the smooth muscle remodeling in both asthma and COPD occurred at two different bronchial localizations, we thus collected asthmatic bronchial specimens from the third to the fourth generation, those from COPD patients from the fourth to the sixth generation and those from control subjects were collected from the third to the sixth generation. Specimens were immediately transferred to the laboratory in a sterile container containing DME.

Optic microscopy and immunohistochemistry.
Bronchial specimens were fixed in 10% buffered formalin and processed in standard fashion for paraffin embedding. 3-µm-thick sections were cut and stained with hematoxylin–eosin–safran or processed for immunohistochemistry. After deparaffinization and rehydration through graded alcohols, endogenous peroxidase was inhibited using a solution of 0.1% sodium azide and 0.3% hydrogen peroxide for 30 min, followed by two rinses in PBS (Invitrogen). Nonspecific staining was blocked using 10% BSA (Dako) for 30 min. Sections were incubated for 2 h at room temperature with mouse anti–human smooth muscle actin (clone 1A4) or an unrelated mouse antibody (both from Sigma–Aldrich). After rinsing in PBS, biotinylated rabbit anti–mouse F(ab’)2 (Dako) was applied to the sections for 2 h, followed by the streptavidin–biotinylated horseradish peroxidase complex (Dako) for an additional 2 h. After rinsing in PBS, 1 mg/ml 3,3’–diaminobenzidine tetrahydrochloride (Dako), plus 0.02% hy-
dorgen peroxide for 6 min, were used as substrate to develop a peroxide-dependent brown color reaction at room temperature. Finally, the sections were rinsed and counterstained with Mayer’s hematoxylin. There was assessable BSM in the bronchial specimens from all 14 asthmatics, 17 COPD patients, and 19 controls. The total area of smooth muscle layer was assessed manually in a blinded fashion using ScanView software (Soft Imaging System) at a magnification of 200×. This smooth muscle area was normalized by the whole area of the corresponding tissues and presented as percentages of whole area.

**Electron microscopy.**
Specimens were fixed in 2.5% glutaraldehyde in cacodylate buffer, postfixed in 1% osmium tetroxide, dehydrated, and embedded in epon. For each specimen, semithin sections (1 µm thick) were cut and stained with alkaline toluidine blue. We selected the first semithin section large enough to span from the epithelium to the muscular layer. 10 ultrathin serial sections (60 nm thick) were then cut on diamond knives. Three of these latter sections were subsequently randomly selected and placed on grids. Staining was performed with uracile acetate and lead citrate. Grids were then scanned by transmission electron microscopy (Tecnai 12; Philips) and examined by a pathologist from left to right and from top to bottom to locate every whole nucleated BSM, epithelial, and endothelial cell. Each ultrathin section was examined in its entirety. To evaluate the number of mitochondria, computerized photographs and measurements were performed in a blinded fashion by using ScanView software at a standard magnification of 6,000×. Mitochondrial densities were performed using ImageJ 1.34s free software (National Institutes of Health).

**Cell cultures.**
Primary cultures of BSM cells were established from bronchial specimens. After a fine dissection under a microscope, smooth muscle explants were cultured in six-well plates in a humidified atmosphere at 37°C with 5% CO$_2$. Cells were maintained in sterile DME containing 10% (vol/vol) FCS and 4.5 g/liter glucose, supplemented with 2 mM t-glutamine, 100 U/ml penicillin, 100 µg/ml streptomycin, and 0.25 µg/ml amphotericine B (antimycotic/antibiotic solution; all from Invitrogen), 1 mM sodium pyruvate, and 1% (vol/vol) nonessential amino acid mixture (both from Sigma-Aldrich). The medium was changed every 48–72 h. After 6–8 wk, confluent cells were rinsed twice with HBSS and passaged with trypsin-EDTA (both from Invitrogen).

To assess purity of the cells, an immunocytochemical method was used, using an indirect immunofluorescence technique. Cells of varying passage number were growth arrested by incubating the cells with serum-free DME supplemented with 10 µg/ml insulin, 5.5 µg/ml transferrin, 5 ng/ml selenium, 0.5 µg/ml BSA, and 4.7 µg/ml linoleic and oleic acid (ITS solution; Sigma-Aldrich). After 48 h, cells were rinsed twice in PBS and fixed with cold methanol for 20 min. Nonspecific staining was blocked using PBS containing 3% BSA for 30 min. Monoclonal antibodies (mAb) diluted in PBS with 1% BSA, including anti-α-smooth muscle actin (Sigma-Aldrich), anti–smooth muscle myosin (Sigma-Aldrich), anti-cytokeratin 18 (Sigma-Aldrich), and anti-factor VIII (Dako) were incubated for 1 h. Control slides were treated similarly, using an unrelated antibody (mouse IgG; Sigma-Aldrich). After rinsing with PBS containing 0.05% Tween20 (Sigma-Aldrich), the cells were incubated for 1 h with FITC-conjugated anti–mouse immunoglobulins (Dako). Slides were mounted with a drop of fluorescent mounting medium (Dako) and observed under a fluorescence microscope (Nikon). Depending on the experiments, cells with a confirmed smooth muscle phenotype were seeded on coverslips, chamber slides, 75-cm² culture flasks, and 6-, 12-, or 96-well plates. Only cells at passage 2 to 4 were used for this study.

To analyze the mitochondrial ultrastruture, BSM cells were growth arrested, as described in the previous section, for 6 d. 3 × 10⁶ cells were then collected and centrifuged at 400 g for 5 min. Cell pellets were fixed in 2.5% glutaraldehyde for 2 h at 4°C and processed for electron microscopy, as described in the previous section. Growth curves were built to assess the role of mitochondria in cell proliferation. For this purpose, 25,000 BSM cells were plated on 75-cm² flasks and cultured with 10% FCS-DME containing either 4.5 g/liter galactose or 4.5 g/liter glucose. In another set of experiments, we investigated the effects of 50 µg/liter ethidium bromide (Qbiogene) for up to 33 d, 1.3 g/liter cyclic cGMP (3 mM; Sigma-Aldrich) for up to 6 d, or 0.5 g/liter D600, 1 i.e., methoxysverapamil (1 µM; Sigma-Aldrich) for up to 2 d in 10% FCS-DME containing 4.5 g/liter glucose. For these latter experiments, medium was changed every 24 h, and, at the indicated time, BSM cells were harvested by trypsinization and counted in triplicate using a cytometer.

**Immunoblotting.**
Whole lysates from BSM cells were collected using 1% Triton X-100 lysis buffer in the presence of 2 mM Na orthovanadate, 1 mM EDTA, 50 µg/ml aprotinin, 100 µM leupeptin, 1 mM 1.4 Dithio-DL-Treitol, and 1 mM amino-ethyl-benzenesulfonyl fluoride hydrochloride (all from Sigma-Aldrich). Cellular extracts were reduced with mercaptoethanol, subjected to electrophoresis on a 10% acrylamide reducing gel, and transferred to PVDF membranes (Immobilon TM-P; Millipore). The immunoblots were then developed using mouse anti–human porin (MitoSciences), mouse anti–human β-actin (Sigma-Aldrich), rabbit anti–human phospho–CaMK-IV, rabbit anti–human PGC-1α, rabbit anti–human NRF-1, or rabbit anti–human mitochondrial transcription factor A (mtTFA; all purchased from Santa Cruz Biotechnology, Inc.) for 14 h at 4°C. For amplifica-
tion, biotinylated goat secondary antibody anti–mouse IgG (Bio–Rad Laboratories) or anti–rabbit IgG (Santa Cruz Biotechnology, Inc.) for 2 h at room temperature and a streptavidin–biotinylated horseradish peroxidase complex (Dako) were used. Immunoblots were revealed by enhanced chemiluminescence (Uptima). Blot images were acquired using BioCaptMW (Thermo Fisher Scientific), and band densities were quantified using ImageJ software.

**Endogenous cell oxygen consumption and determination of the coupling ratio.**

Endogenous cellular oxygen consumption was monitored on intact cells at 37°C in a 1–ml thermostatically controlled chamber (10⁶ cells/ml/run) equipped with a Clark oxygen electrode (Oxygraph System; Hansatech). The respiratory buffer was the glucose or the galactose culture medium described in Cell culture. The endogenous respiratory rates were expressed in ngatom O/min/10⁶ cells. To assess the mitochondrial coupling, respiratory rates were monitored using inhibitors of the phosphorylation system, i.e., 1 mM atracyloside, 1 mM carboxatractylsode, and 1 mM oligomycin in combination. Such inhibitors are considered as decouplers and can be used to evaluate the degree of mitochondrial ATP synthesis (6). Control experiments were performed using 1 mM cyanide as a blocker of the respiratory chain.

**Three-dimensional assessment of the mitochondrial network organization.**

To visualize the mitochondrial network, we targeted a GFP to the mitochondrial matrix space of the BSM cells. For this purpose, we used the mitochondrion-targeted GFP plasmid (mito–GFP), which was derived from the plasmid pEGFP-N1 (Clontech Laboratories, Inc.) containing the leader sequence of the E1α subunit of pyruvate dehydrogenase. 1 d before transfection, BSM cells were plated at 50% confluence in 10% FCS–DMEM. Cells were transfected with 1 µg of plasmid and 1.7% of the cationic lipid Lipofectamine 2000 (Invitrogen) in the absence of FCS. After 72 h, BSM cells were rinsed in PBS and fixed with 4% paraformaldehyde (PFA) for 20 min on ice. After drying, the slides were mounted with fluorescent mounting medium (Dako). Confocal differential interference contrast images were obtained using the FluoView laser scanning microscope (Nikon) and 60× oil-immersion objective. Z series sections were recorded in successive z axis serial sections at 0.5-µm intervals, and they were composed of optical sections in the x-y optical plane. Sections were reconstituted in three-dimensional images using Imaris Software (Bitplane).

**RNA extraction, RT, and real-time quantitative PCR.**

BSM cells were washed twice with HBSS and lysed by the addition of 300 liters/well Trizol (Invitrogen) and 30 µl of chloroform (Sigma–Aldrich). The RNA was extracted from the aqueous phase after being centrifuged at 12,000 g for 15 min. RNA was precipitated in the presence of isopropanol (Sigma–Aldrich) at −20°C overnight. The pure RNA was recovered by centrifugation and washed with 80% ethanol (Sigma–Aldrich). The purity was assessed by electrophoresis (30 mA for 75 min) on 1.5% agarose gel (Bio–Rad Laboratories), followed by 20 min incubation in 10 µg/ml ethidium bromide (Sigma–Aldrich), showing 2 bands corresponding to the 28S and 18S fractions of ribosomal RNA. The concentration of RNA was measured spectrophotometrically by GeneQuant RNA/DNA calculator (GE Healthcare). The total RNA (1 µg) was reverse transcribed into cDNA using AMV RT (Promega), RNase inhibitor, and oligo d(T) as a primer at 42°C for 60 min, followed by heating at 94°C for 3 min.

Real-time quantitative PCR was performed on a Rotor–Gene 2000 (Corbett Research). In brief, triplicate PCR reactions were assembled in 0.1-ml strip tubes containing cDNA from 10 ng of total RNA, 0.2 µl 50X Titanium Taq DNA Polymerase, 1X Titanium Taq PCR Buffer (Clontech Laboratories, Inc.), 1 mM dNTP, 100 mM each of the appropriate primer, and 0.5X Sybr Green I (Invitrogen). The PCR was performed under the following conditions: denaturation at 95°C for 15 s, annealing temperature (64–72°C) depending on specific primers for 15 s, and extension at 70°C for 30 s for 30–40 cycles. Data collection was performed after each extension step, at a temperature at least 3.5°C lower than the melting temperature of the amplicon (generally between 80–85°C) to eliminate nonspecific fluorescence signal. PCR-negative controls were systematically made using water instead of cDNA or RNA sample without the RT step. All specific primers were designed using the primer analysis software (Oligo 6.6; Molecular Biology Insights) and were ordered from Sigma–Aldrich. Primers, sense and antisense, were as follows: for mtTFA (NM_013261), forward 5′-GAAGTCTGACTGCCGCTCCC-3′ and reverse 5′-ACTCCGCCCTATAAGCATC-3′; for PGC-1α (NM_013261), forward 5′-GTCAACCCCAAATCCTTTATT-3′ and reverse 5′-GCCGATCTTTGAACGTGAT-3′; for NRF-1 (NM_005011), forward 5′-AAGATACGACGCCAAACAC-3′ and reverse 5′-CCGGTGTTACACTGATAAGT-3′; for tyrosin 3-monoxygenase/tryptophan 5-monoxygenase activation protein, zeta polypeptide (YWHAZ) (NM_003406), forward 5′-GGGGATGTTGGAATTTTTATAAAG-3′ and reverse 5′-CGCAGGCAAAAACGGTTATGAG-3′; for hypoxanthine phosphoribosyltransferase 1 (HPRT-1) (NM_000194), forward 5′-GTTGAAAAGGACCABCAGAA-3′ and reverse 5′-AGTCAAGGCGATCTCACAACAA-3′; and for human acidic ribosomal phosphoproteins (PO) (NM_001002), forward 5′-CAACGGTTACACAGAGTC-3′ and reverse 5′-CTTCCTTTGGCTTCACCTTGA-3′. The efficiency of the PCR reactions was always >90%. The specificity of the amplified PCR products was examined with the melting curve analysis, and also in 2% agarose gel containing ethidium bromide. The RT-PCR expression of the target genes (i.e., mtTFA, PGC-1α, and NRF-1) was thus presented as an arbitrary unit and
normalized to endogenous references (geometric averaging of three internal control genes; i.e., YWHAZ, HPRT-1, and PO) according to geNorm software (7).

Microspectrofluorimetry.
Change in growth-arrested BSM cell intracellular calcium concentration ([Ca$^{2+}$]$_i$), was assessed using the Ca$^{2+}$-sensitive probe indo-1, as previously described (8, 9). In brief, cells were loaded with indo-1 (Calbiochem) and mounted in a perfusion chamber continuously perfused with phosphate salt solution containing either 2 mM Ca$^{2+}$ in the absence or presence of 1 μM D600, or 400 μM EGTA (Sigma-Aldrich) in the absence of extracellular calcium. BSM cells were stimulated with 10$^{-5}$ M acetylcholine or histamine (both from Sigma-Aldrich). Individual cell calcium levels were monitored continuously. Results were expressed as the mean ± the SEM of resting or peak of [Ca$^{2+}$]$_i$ rises (in millimoles) and area under the curves (nanomoles/second). Experiments were carried out at room temperature (22–25°C).

BrdU incorporation.
Growth-arrested BSM cells were cultured in 96-well plates for 24 h and stimulated with 10% FCS-DME containing glucose, galactose, ethidium bromide, or cyclic GMP for 24 h or containing D600 for 48 h, as described above. Control BSM cells remained in serum-deprived DME. DNA synthesis was measured using the BrdU kit according to the manufacturer’s instructions (Roche). In brief, BSM cells were incubated with 10 μM BrdU for 2 h at 37°C and fixed for 20 min at room temperature. Cells were denaturized for 30 min and incubated with the anti-BrdU antibody for 2 h at room temperature. Absorbance was measured at 370 nm in a microplate reader in triplicate. Results are expressed as a normalized ratio of BrdU incorporation, i.e., absorbance for test condition divided by absorbance for control serum-deprived condition.

Statistical analysis.
The statistical analysis was performed with NCSS 2001 software. Comparisons between the three groups were performed by means of ANOVA, with the use of Bonferroni’s t for multiple comparisons or paired Student’s t tests. Values are presented as the mean ± the SEM. A Pearson correlation matrix was built between in vitro and in vivo measurements. A P value <0.05 was considered statistically significant.

REFERENCES