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Germline BAP1 mutations induce a Warburg effect

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Carriers of heterozygous germline BAP1 mutations (BAP1^{+/-}) develop cancer. We studied plasma from 16 BAP1^{+/-} individuals from 2 families carrying different germline BAP1 mutations and 30 BAP1 wild-type (BAP1^{WT}) controls from these same families. Plasma samples were analyzed by liquid chromatography time-of-flight mass spectrometry (LC-TOF-MS), ultra-performance liquid chromatography triple quadrupole mass spectrometry (UPLC-TQ-MS), and gas chromatography time-of-flight mass spectrometry (GC-TOF-MS). We found a clear separation in the metabolic profile between BAP1^{WT} and BAP1^{+/-} individuals. We confirmed the specificity of the data in vitro using 12 cell cultures of primary fibroblasts we derived from skin punch biopsies from 12/46 of these same individuals, 6 BAP1+/- carriers and 6 controls from both families. BAP1+/- fibroblasts displayed increased aerobic glycolysis and lactate secretion, and reduced mitochondrial respiration and ATP production compared with BAP1^{WT}. siRNAmediated downregulation of BAP1 in primary BAP1^{WT} fibroblasts and in primary human mesothelial cells, led to the same reduced mitochondrial respiration and increased aerobic glycolysis as we detected in primary fibroblasts from carriers of BAP1^{+/-} mutations. The plasma and cell culture results were highly reproducible and were specifically and only linked to BAP1 status and not to gender, age or family, or cell type, and required an intact BAP1 catalytic activity. Accordingly, we were able to build a metabolomic model capable of predicting BAP1 status with 100% accuracy using data from human plasma. Our data provide the first experimental evidence supporting the hypothesis that aerobic glycolysis, also known as the 'Warburg effect', does not necessarily occur as an adaptive process that is consequence of carcinogenesis, but rather that it may also predate malignancy by many years and facilitate carcinogenesis.

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We discovered that inherited heterozygous germline mutations in the BRCA-associated protein 1 (*BAP1*) gene cause a high rate of mesothelioma and uveal melanoma.¹ Our data, confirmed and expanded by others in several *BAP1*^{+/-} families across the world, revealed that germline *BAP1* mutations in addition to mesothelioma and uveal melanoma, also cause other malignancies.^{2–5} We named this condition the 'BAP1 cancer syndrome'.^{2,6} So far, all individuals affected by this novel cancer syndrome developed at least one and often multiple cancers during their lifetime.⁷ Intriguingly, most *BAP1*^{+/-} patients experience significantly longer survival compared with patients who develop these same malignancies sporadically.^{7–9}

BAP1 is a deubiquitylase that regulates multiple activities¹⁰⁻¹⁶ by forming multi-protein complexes.⁶ BAP1 stabilizes PGC-1 α and promotes gluconeogenesis.¹⁷ Dey's team found that the glucose and hexose metabolic pathways were repressed in liver-specific *Bap1* knockout mice.¹⁸ These findings^{17,18} suggest that BAP1 may influence cellular metabolism. Very recently, we discovered a novel BAP1 cytoplasmic activity that, in concert with its nuclear activities, allows BAP1 to modulate gene–environment interaction and

contributes to the high incidence of cancer in $BAP1^{+/-}$ carriers. 19

Adult 'normal' differentiated cells derive energy mostly via oxidative respiration (the tricarboxylic acid (TCA) cycle and oxidative phosphorylation). Alterations in the major cellular pathways involved in energy production - glycolysis and TCA cycle coupled with oxidative phosphorylation - are frequently found in cancer cells.²⁰ Warburg's findings²¹ that cancer cells obtain approximately the same amount of energy from fermentation (glycolysis to lactic acid) as from the TCA cycle is considered a hallmark of malignancy.^{20,22-25} This phenomenon referred as the 'Warburg effect' protects tumor cells from hypoxia and provides a rich source of precursors for the biosynthesis of nucleic acid, fatty acids, and phospholipids that sustain tumor cell growth. Moreover, the lactic acid secreted in the extracellular space promotes tumor cell growth and protects tumor cells from immune cells.^{20,22-24} As to the emergence of the Warburg effect, the prevalent hypothesis is that the hypoxia that characterizes the initial growth of premalignant lesions provides micro-environmental selection forces that facilitate the growth of cell phenotypes that can adapt to this harsh environment through resistance to hypoxia

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and micro-environmental acidosis.²⁶ Thus adaptation to hypoxia via a Warburg effect and subsequent selection of those cell clones capable of surviving acidosis leads over time to malignant cell populations with a powerful growth advantage.²² In addition to tumor cells, embryonic and adult stem cells may also produce energy via aerobic glycolysis.²⁰ Similarly, within the tumor microenvironment, activated immune cells (macrophages, dendritic cells, and T cells) can also use aerobic glycolysis that helps them survive the hypoxic tumor environment.^{27,28}

Here we show that cells from $BAP1^{+/-}$ carriers do not need to go through a selection process that favors the emergence of clones with a Warburg effect required for tumor growth: normal primary $BAP1^{+/-}$ cells constitutionally derive a large part of their energy through aerobic glycolysis.

Table 1 Differentially expressed metabolites between BAP1 ¹¹¹ and BA	P1*/-
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	Metabolomics profiling results		
	Plasma	Fibroblast cell cultures	
No. of total metabolites	412	495	
No. of known metabolites	246	226	
No. of metabolites with $VIP \ge 1.5$	28	21	

We found 412 plasma and 495 cells' chromatographic peaks corresponding to putative metabolites; of these, 246 plasma metabolites and 226 cell metabolites could be identified. Metabolites with VIP \geq 1 are listed in Supplementary Tables S2 and S3

Results

Individuals carrying heterozygous germline *BAP1* mutations can be reliably identified based on their metabolic profile. We analyzed the metabolite profiles of plasma samples from 46 members of two unrelated families carrying different germline *BAP1* mutations – the Wisconsin (W) and the Louisiana (L) families – that we have been following and treating for mesothelioma and other malignancies for > 10 years.¹ For details about individuals and samples, see Materials/Subjects and Methods section¹⁹ and Supplementary Table S1.

Plasma and fibroblast¹⁹ extracts were analyzed by liquid chromatography time-of-flight mass spectrometry (LC-TOF-MS), ultra-performance liquid chromatography triple guadrupole mass spectrometry (UPLC-TQ-MS), and gas chromatography time-of-flight mass spectrometry (GC-TOF-MS). A multivariate statistical analysis of the identified metabolites (Table 1), orthogonal partial least squares discriminant analysis (OPLS-DA), revealed a clear and statistically significant separation between BAP1^{WT} and BAP1^{+/-} individuals, both in plasma (OPLS-DA model: R2X = 0.208, R2Y (cum) = 0.951, Q2(cum) = 0.313) (Figure 1a) and in cell extracts (OPLS-DA model: R2X = 0.321, R2Y(cum) = 0.916, Q2(cum) = 0.215) (Figure 1b). The OPLS-DA model was not influenced by the year of collection of the samples (Figure 2a) and the age (Figure 2b) or gender (Figure 2c) of the individuals, suggesting that the differences observed in Figure 1a were specifically related to BAP1 status (Figure 2 and Supplementary Figure S1). To test whether the OPLS-DA model established with the metabolomics data of human subjects with known BAP1 status was able to



Figure 1 OPLS-DA analysis of metabolite profiles showed a global metabolic difference between $BAP1^{WT}$ and $BAP1^{+/-}$ individuals. (a) OPLS-DA score plot derived from, UPLC-TQ-MS and GC-TOF-MS spectral data of N = 61 plasma samples (37 $BAP1^{WT}$ (blue dots) and 24 $BAP1^{+/-}$ (red dots)), K = 412 variables, including identified metabolites (246) and unknowns. (b) OPLS-DA score plot of UPLC-TQ-MS and GC-TOF-MS spectral data of whole-cell extract samples from $BAP1^{WT}$ (N = 6) and $BAP1^{+/-}$ (N = 6) fibroblast cell cultures, K = 495 variables, including identified metabolites (226) and unknowns

clearly discriminate among samples from individuals with or without heterozygous germline *BAP1* mutations (Figure 2d), we analyzed blindly plasma samples from 76 individuals: $22 BAP1^{WT}$, $12 BAP1^{+/-}$, 1 individual of the W family with unknown *BAP1* status at the time the analysis was performed, and 41 healthy controls not related to the W and L

families. The OPLS-DA model based on plasma metabolomics data was able to clearly discriminate the 41 healthy controls as $BAP1^{WT}$ (Figure 2d). The BAP1 unknown sample (WI III-7) was classified among the $BAP1^{WT}$, a finding later confirmed by genomic testing (Figure 2d and Supplementary Figure S1).



Figure 2 Genotype of $BAP1^{WT}$ and $BAP1^{+/-}$ individuals can be predicted from plasma samples using an OPLS-DA analysis of their metabolite profile independently of year of collection, age, gender, or family of origin. (**a**-**c**) OPLS-DA analysis of metabolite profiles is not influenced by (**a**) the year of collection (2012 and 2014, displayed in the panel), (**b**) the age of the individuals, and (**c**) gender (M = male, F = female). (**d**) OPLS-DA analysis differentiates $BAP1^{+/-}$ carriers independently of family of origin. LC-TOF-MS and GC-TOF-MS were used for the metabolomic profiles of plasma samples from 22 $BAP1^{WT}$ subjects (blue dots), 12 $BAP1^{+/-}$ (red diamonds), 41 healthy controls unrelated to the W and L families (black triangles), and 1 individual of the W family with unknown BAP1 status at the time the analysis was performed – later confirmed to be $BAP1^{WT}$ (purple star)

Figure 3 Altered metabolic pathways for the most relevant distinguishing metabolites between $BAP1^{WT}$ and $BAP1^{+/-}$ cells. Glycolysis converts glucose into pyruvate. In the presence of oxygen, in the mitochondria, pyruvate is completely oxidized and the energy is stored as ATP (aerobic metabolism). In the absence of sufficient oxygen or in tumor cells because of the Warburg effect, the pyruvate is reduced to lactate (anaerobic metabolism). (**a**-**h**) Levels of glucose (**a**), glucose 6-P (**b**), glycerol (**c**), glycerol 3-P (**d**), pyruvate (**e**), citrate (**f**), fumarate (**g**), and malate (**h**) were measured in whole-cell extract samples from $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblast cell cultures. Bar plots show the mean \pm S.E.M. of the average metabolite intensity. Our analysis (see also Supplementary Table S3) revealed decreased levels of glucose 6-P (VIP = 2.42), glycerol 3-P (VIP = 1.59), pyruvate (VIP = 1.16), and citrate (VIP = 1.71) in $BAP1^{+/-}$ fibroblasts. A slight reduction in the levels of other TCA cycle intermediates – fumarate and malate – was also found. Glucose (VIP = 1.44) and glycerol (VIP = 1.09) levels were increased in $BAP1^{+/-}$ fibroblasts compared with $BAP1^{WT}$. *P < 0.05. (**i**) Basal lactate secretion is increased in $BAP1^{+/-}$ fibroblasts compared with $BAP1^{WT}$. The amount of lactate released (dotted red arrow) in the cell culture media was determined using a colorimetric assay; data shown as mean \pm S.E.M. of three independent experiments; *P < 0.05. ACO, aconitase; ADP, Adenosine tiphosphate; ALDO, aldolase; ATP, adenosine triphosphate; CS, citrate synthase; ENO, enolase; FH, fumarate hydratase; GPD, glycerol-3-phosphate dehydrogenase; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; GPI, glucose-6-phosphate isomerase; IDH, isocitrate dehydrogenase; LDH, lactate dehydrogenase; PKM, pyruvate kinase; OGDH, oxoglutarate (alpha-ketoglutarate) dehydrogenase; SUCL, succinate-CoA ligase; SDH, succinate dehydrogenase; TPI, triosephosphate isomerase

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Figure 4 Levels of ¹³C-metabolites in culture medium and cells after culturing *BAP1^{WT}* and *BAP1^{+/-}* cells with ¹³C-glucose for 24 h. *BAP1^{WT}* and *BAP1^{+/-}* fibroblasts were incubated in ¹³C-glucose medium. (a) ¹³C-glucose consumption rate measured in tissue culture medium from *BAP1^{WT}* and *BAP1^{+/-}* cells. (b–d) ¹³C incorporation into ¹³C-glucose (b), ¹³C-glucose 6-P (c), and ¹³C-citrate (d) measured in cell extracts from *BAP1^{WT}* and *BAP1^{+/-}* cells. (e) Extracellular concentration of ¹³C-lactate produced by *BAP1^{WT}* and *BAP1^{+/-}* fibroblasts after 24 h of cell culture using ¹³C-glucose medium. Bar plots show the mean \pm S.E.M.; **P* < 0.05

Variable importance in the projection (VIP) values were calculated to evaluate the contribution of individual metabolite to the OPLS-DA model. Large VIP values > 1.0 are the most relevant for explaining differences between $BAP1^{WT}$ and $BAP1^{+/-}$ groups. A total of 71 metabolites identified in plasma samples had VIP > 1, among which 28 had VIP values > 1.5 (Table 1 and Supplementary Table S2). A total of 111 metabolites identified in samples from cells in tissue culture had VIP > 1, among which 21 had VIP values > 1.5 (Table 1 and Supplementary Table S3). To determine how many metabolites were increased/decreased in the $BAP1^{+/-}$ samples, we calculated the fold change as the ratio $BAP1^{+/-}$ (BAP1^{WT} of the average metabolite amount (Supplementary Table S2).

Germline BAP1 mutations have significant effects on energy metabolism. Several of the altered metabolites between BAP1^{+/-} and BAP1^{WT} were glycolytic or TCA cycle intermediates (Figures 3a-h). Among the cell metabolites with a VIP \geq 1.5 (Supplementary Table S3 and Supplementary Figure S2), we found that glucose 6-phosphate (Figure 3b) was significantly decreased in $BAP1^{+/-}$ samples (VIP=2.42, P=0.00972). Glycerol 3-phosphate (Figure 3d) and citrate (Figure 3f) were also decreased in BAP1+/samples, although the group difference was not significant by Student's t-test (Supplementary Table S3). By extending the analysis to include the metabolites with VIP ≥ 1 (Supplementary Table S3), we found decreased levels of pyruvate (Figure 3e), while glucose (Figure 3a) and glycerol (Figure 3c) levels were increased in BAP1+/- samples. A small reduction was detected in fumarate (Figure 3g) and malate (Figure 3h) levels in BAP1+/- samples. Significantly increased lactate concentrations were detected in the conditioned cell culture medium of BAP1+/- fibroblast cell (+32.75% + 11.24) compared with BAP1^{WT} cultures (Figure 3i). Together, these findings suggested increased glycolysis and reduced aerobic mitochondrial respiration. However, the metabolomics data (Figure 3) might also fit the hypothesis of decreased hexokinase (HK) conversion (because the substrate metabolite, namely, glucose accumulates and the product metabolite, namely, glucose 6phosphate decreases) and decreased glycerol kinase (GK) activity (because the substrate metabolite, namely, glycerol

accumulates and the product metabolite, namely, glycerol 3-P decreases).

Metabolic rate analysis using ¹³C-glucose confirms increased glucose consumption and lactate secretion in BAP1+/- cells. Stable isotope (13C) tracking of central metabolic pathways provides a dynamic picture of metabolism and allows the experimental quantification of the integrated responses of metabolic networks.²⁹ We performed comparative profiling of the ¹³C-labeled metabolic pattern in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts. After growing fibroblasts in ¹³C-labeled glucose, we performed a metabolic rate analysis based on UPLC-QTOFMS tracing pattern of the stable isotope in metabolites in a steady state (Figure 4). Analysis of ¹³C-glucose levels in conditioned medium revealed a significantly increased consumption rate in BAP1+/- fibroblasts compared with BAP1^{WT} (Figure 4a). Analyses of cells extracts collected in parallel revealed increased levels of ¹³C-glucose in BAP1^{+/-} fibroblasts compared with BAP1^{WT} (Figure 4b). We also observed a reduction in intracellular ¹³C-glucose 6-P (Figure 4c) and ¹³C-citrate (Figure 4d) in BAP1^{+/-} fibroblasts, supporting the data shown in Figure 3 and Supplementary Table S3. Moreover, increased ¹³C-lactate concentrations were detected in the conditioned cell culture medium of BAP1+/- fibroblasts compared with $BAP1^{WT}$ (Figure 4e). Thus, at the steady state, $BAP1^{+/-}$ cells consumed more glucose (higher ¹³C-glucose consumption rate) and released more ¹³C-labeled lactate in the culture medium. At the same time, lower amounts of intracellular intermediates ¹³C-glucose 6-P and ¹³C-citrate were detected in BAP1^{+/-} cells. Mass distribution vectors³⁰ of ¹³C-labeled glucose 6-P, citrate and lactate (Supplementary Table S4) showed that: glucose 6-P was only observed in form M+6; citrate was observed in seven forms: unlabeled and M+1. M +2, M+3, M+4, M+5, and M+6; lactate was observed in four forms: unlabeled, M+1, M+2, and M+3. Fractional contributions of ¹³C-labeled glucose 6-P and citrate were lower, while fractional contribution of lactate was higher in $BAP1^{+/-}$ cells, confirming a faster glycolytic rate compared with BAP1^{WT} cells. These data indicate that $BAP1^{+/-}$ cells have an increased glycolytic rate compared with BAP1^{WT} cells. $BAP1^{+/-}$ cells take up more glucose because they rapidly metabolize it to lactate, which is then released extracellularly. A faster metabolic rate from glucose to extracellular lactate

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Figure 5 $BAP1^{+/-}$ fibroblasts have increased glycolytic activity and decreased intracellular ATP. (a) ECAR was recorded over time in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts with the Seahorse XF96 extracellular flux analyzer. ECAR can be further dissected into different modules as depicted in the upper left scheme. The initial administration of a saturating concentration of glucose activates the glycolytic pathway producing NADH, protons, H₂O, and ATP. Protons induce a rapid increase in ECAR. The following inhibition of mitochondrial ATP synthesis with Oligomycin A shifts the energy production to glycolysis and induces an additional increase in ECAR. Finally, inhibition of glycolysis with 2-deoxyglucose (2-DG) causes a strong decrease in ECAR levels, confirming that the ECAR observed was generated by the glycolytic catavity of the cells. Representative experiments performed in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts are displayed on the right panel. (b) Levels of glycolysis and maximal glycolytic capacity in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts are displayed on the right panel. (b) Levels of glycolysis and maximal glycolytic capacity in $(C) BAP1^{WT}$ fibroblasts transfected with a pool of siRNAs targeting BAP1, and $(d) BAP1^{+/-}$ fibroblasts transduced with adenoviruses for BAP1 (AdBAP1), its catalytic inactive mutant BAP1(C91S) (AdBAP1(C91S)), or GFP as control (AdGFP). In panel (b), the WB shows BAP1 protein levels in $BAP1^{WT}$ and $siBAP1^{*/-}$ fibroblasts transduced with AdBAP1 (a pool of four different siRNAs targeting BAP1 (C91S), or control (AdGFP). (e) Total levels of further sire of the AdBAP1(C91S), or control (AdGFP). (e) Total levels of further sire of the adaption of the maximal glycolytic. (catavitic nective in the BAP1^{*/-} fibroblasts compared with AdBAP1. (catalytically inactive AdBAP1(C91S), or control (AdGFP). (e) Total levels of four different sire of the maximal glycolytic) is or other of the sire of the adaption of the maximal glycolytis and sire of the adaption of th

would quickly exhaust the glycolysis intermediates, explaining why $BAP1^{+/-}$ cells have lower levels of ¹³C-glucose 6-P in spite of the increased glucose consumption rate. Thus the changes observed did not appear related to decreased HK or GK activity, an interpretation further supported by the experiments described below.

BAP1^{+/-} cells rely largely on the glycolytic metabolism for energy production. To further characterize the glycolytic pathway in $BAP1^{+/-}$ cells, we measured the extracellular acidification rate (ECAR) in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblast cell cultures using the Seahorse XF96 Analyzer. When a saturating concentration of glucose (5 mM) is added, its catabolism to lactate through the glycolytic pathway produces ATP, protons, and thus a rapid increase in ECAR. We measured this glucose-induced response and reported it as the rate of glycolysis in basal conditions (Figures 5a and b). ECAR measurements (Figure 5b) revealed a significantly increased rate of glycolysis in *BAP1*^{+/-} fibroblast cell cultures (28.05 ± 1.76 mpH/min) compared with *BAP1*^{WT} (16.73 ± 1.52 mpH/min). The subsequent addition of the ATP synthase inhibitor – oligomycin A – inhibits mitochondrial ATP production and thus shifts the energy production to glycolysis. The consequent further increase in ECAR is a measure of the maximum glycolytic capacity of the cell. We found a significantly increased maximum glycolytic capacity in *BAP1*^{+/-} cells (35.88 ± 2.05 mpH/min) compared with *BAP1*^{WT} (22.44 ± 2.04 mpH/min) (Figure 5b).

Next, we silenced BAP1 in wild-type control fibroblasts using siRNAs (siBAP1) to test whether we were able to mimic the same increases in glycolysis and glycolytic capacity. ECAR measurements (Figure 5c) showed a significant increase in both parameters in BAP1-silenced cells (rate of glycolysis: $BAP1^{WT}$ 12.76±1.12 mpH/min, siBAP1 18.98±2.51 mpH/min; glycolytic capacity: $BAP1^{WT}$ 16.93±0.85 mpH/min, siBAP1 24.11±2.35 mpH/min). Similar results

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Figure 6 Mitochondrial respiratory function is impaired in $BAP1^{+/-}$ cells. (a) OCR was analyzed in real time using the Seahorse XF96 extracellular flux analyzer. When Oligomycin A is added, the ATP synthase complex is inhibited, therefore the respiratory chain-associated oxygen consumption is inhibited. Addition of the ATP synthesis uncoupler FCCP induces the maximal oxygen consumption by the respiratory chain. Addition of rotenone and antimycinA (Rot/AntA, complex I and III inhibitors, respectively) blocks the electron transfer as well as oxygen consumption by the respiratory chain. Displayed on the left is a representative experiment showing the different parameters that can be measured and used to determine mitochondrial respiratory function: basal respiration (OCR-BASAL), ATP production (OCR-ATP), maximal respiration (OCR-MMR), and spare respiratory capacity (OCR-SRC). Representative experiments performed in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts are shown on the right. (b) Reduced OCR-BASAL, OCR-ATP, OCR-MMR, and OCR-SRC in $BAP1^{+/-}$ fibroblasts compared with $BAP1^{WT}$, indicating an overall impaired mitochondria-linked aerobic respiration and ATP production in germline mutated $BAP1^{+/-}$ fibroblasts transfected with a pool of siRNAs targeting BAP1 and (d) $BAP1^{+/-}$ fibroblasts transduced with adenoviruses (Ad) for BAP1 (AdBAP1), its catalytic inactive mutant AdBAP1(C91S), or GFP as control (AdGFP). Bar plots show the mean \pm S.E.M. of three independent experiments. **P < 0.01, *P < 0.05

were obtained in primary human mesothelial (HM) cells, which we established in tissue culture from patients with benign pleural effusions treated with siBAP1 to reduce BAP1 expression levels (Supplementary Figures S3a and b).

We also transduced BAP1 in BAP1+/- fibroblasts with adenoviruses for wild-type BAP1 (AdBAP1) or its catalytic inactive mutant carrying the C91S point mutation (AdC91S)³¹ or a control adenovirus (AdGFP). Figure 5d shows that BAP1^{+/-} fibroblasts transduced with AdBAP1 had reduced glycolysis and glycolytic capacity; in contrast, AdC91S was ineffective at reducing glycolysis (rate of glycolysis in BAP1^{+/-} transduced with: AdGFP 24.94 ± 1.77 mpH/min, AdBAP1 6.92 ± 0.24 mpH/min, AdBAP1(C91S) 20.33 ± 2.92 mpH/min; glycolytic capacity: AdGFP 30.33 ± 1.71 mpH/min, AdBAP1 11.50 ± 0.43 mpH/min, AdC91S 25.08 ± 3.25 mpH/min). These data further indicate that BAP1 regulates glycolysis and that BAP1+/- cells rely largely on the aerobic glycolytic metabolism for energy production. Despite the enhanced glycolysis, we detected lower intracellular levels of total ATP in $BAP1^{+/-}$ compared with $BAP1^{WT}$ cells (Figure 5e). Conversion of glucose to lactate yields two ATP molecules; the subsequent TCA cycle generates ~36 molecules of ATP.32 The reduced intracellular levels of ATP in BAP1+/cells suggested that mitochondrial respiration was impaired, and therefore $BAP1^{+/-}$ cells were preferentially using glycolysis – rather than the TCA cycle – to produce ATP (Warburg effect).²¹

BAP1 mutations impair mitochondrial respiration. As citrate, a key TCA cycle intermediate, was decreased in carriers of germline BAP1 mutations (Figures 3f and 4d), we sought to determine whether the boost in the glycolytic pathway of energy production observed in $BAP1^{+/-}$ cells was associated with changes in mitochondrial respiration. First, we studied whether BAP1+/- cells exhibited normal mitochondrial morphology and mitochondrial membrane potential $(\Delta \Psi)$. Mitochondria were labeled in living cells by expressing green fluorescent protein specifically targeted to the mitochondria (mtGFP). mtGFP imaging (Supplementary Figure S4a) showed the normal three-dimensional interconnected tubular network characteristic of these organelles and no differences in mitochondrial number (Supplementary Figure S4b) or volume (Supplementary Figures S4c and d) between $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts. Also no differences were detected in the loading of the fluorescent dye tetramethyl rhodamine methyl ester (TMRM) (Supplementary Figure S4e), thus ruling out $\Delta \Psi$ differences between BAP1^{WT} and $BAP1^{+/-}$ fibroblasts.

Next, using the Seahorse XF96 Analyzer, we studied the oxygen consumption rate (OCR) in $BAP1^{WT}$ and $BAP1^{+/-}$

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fibroblast cell cultures. The OCR is indicative of (and proportional to) mitochondrial respiration and can be employed as an indicator of mitochondrial respiratory capacity and energy production to reveal defects in mitochondrial bioenergetics mechanisms. The consecutive addition of specific mitochondrial inhibitors to the cells - Oligomycin A, FCCP. and rotenone/antimvcin A – allows the measurements of distinct modules of oxygen consumption related to different mitochondrial processes (Figure 6a, left panel), Figure 6a (right panel) shows the OCR measurements over time in $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblast cell cultures. $BAP1^{+/-}$ fibroblasts exhibited significantly lower basal mitochondrial respiration (OCR-BASAL) (65.78 ± 4.38 pMoles/min) compared with $BAP1^{WT}$ (111.20 ± 6.92 pMoles/min), supporting lower ATP turnover and demand (Figure 6b). The addition of Oligomycin A inhibits ATP synthase (complex V of the respiratory chain) causing a decrease in OCR that correlates with the amount of oxygen consumption linked to mitochondrial ATP production (OCR-ATP). ATP-linked respiration was significantly decreased in $BAP1^{+/-}$ fibroblasts (48.86 ± 3.38 pMoles/min) compared with $BAP1^{WT}$ (81.48 ± 5.24 pMoles/min) (Figure 6b). The maximal ATP output of mitochondria can be determined by addition of FCCP, an uncoupling agent that collapses the proton gradient and disrupts the mitochondrial membrane potential, inducing maximal oxygen consumption and substrate oxidation by complex IV (OCR-MMR, maximal mitochondrial respiration). OCR-MMR was significantly lower in BAP1+/- fibroblasts $(139.73 \pm 11.61 \text{ pMoles/min})$ compared with BAP1^{WT} (227.97 ± 19.17 pMoles/min) (Figure 6b). The spare respiratory capacity (OCR-SRC) was determined by subtracting OCR-BASAL from FCCP-induced OCR-MMR. OCR-SRC measures the ability of the cells to respond to increased energy demand and was also significantly decreased in $BAP1^{+/-}$ fibroblasts (73.94 ± 7.51 pMoles/min) compared with $BAP1^{WT}$ (116.77 ± 14.00 pMoles/min) (Figure 6b). To summarize, in BAP1+/- fibroblast cell cultures we observed an overall reduction in all the OCR parameters that quantify mitochondrial respiration.

To support these findings, we measured OCR parameters in $BAP1^{WT}$ fibroblasts silenced for BAP1 (siBAP1). Similar to the $BAP1^{+/-}$ cells (Figure 6b), we found that in siBAP1 cells all of the four critical parameters of mitochondrial respiration were significantly decreased: OCR-BASAL ($BAP1^{WT}$ 95.89 ± 4.90 pMoles/min, siBAP1 68.44 ± 5.48 pMoles/min), OCR-ATP ($BAP1^{WT}$ 76.63 ± 4.55 pMoles/min, siBAP1 51.27 ± 6.75 pMoles/min), OCR-MMR ($BAP1^{WT}$ 109.27 ± 6.91 pMoles/min, siBAP1 84.36 ± 5.73 pMoles/min), and OCR-SCR ($BAP1^{WT}$ 22.87 ± 3.78 pMoles/min, siBAP1 10.91 ± 2.91 pMoles/min) (Figure 6c). Similar results were obtained in primary HM cells in which BAP1 was silenced (Supplementary Figures S3a and c).

Next, we tested whether the rescue of BAP1 protein levels would be sufficient to revert the impairment in mitochondrial respiration. $BAP1^{+/-}$ fibroblasts were transduced with AdGFP (control), AdBAP1, or AdBAP1(C91S). We found that the reintroduction of AdBAP1, but not of AdGFP or the catalytic inactive AdBAP1(C91S), was able to rescue mitochondrial respiration in $BAP1^{+/-}$ cells, as determined by measurements of OCR-BASAL (AdGFP 78.29±3.82 pMoles/min, AdBAP1

96.10 \pm 7.42 pMoles/min, AdBAP1(C91S) 76.61 \pm 5.14 pMoles/min), OCR-ATP (AdGFP 60.81 \pm 1.56 pMoles/min, AdBAP1 77.07 \pm 5.40 pMoles/min, AdBAP1(C91S) 63.46 \pm 4.96 pMoles/min), OCR-MRR (AdGFP 89.70 \pm 2.57 pMoles/min, AdBAP1 110.43 \pm 7.95 pMoles/min, AdBAP1(C91S) 85.86 \pm 3.72 pMoles/min), OCR-SCR (AdGFP 7.46 \pm 3.81 pMoles/min, AdBAP1 22.02 \pm 3.90 pMoles/min, and AdBAP1 (C91S) 6.53 \pm 5.26 pMoles/min) (Figure 6d).

Gene expression analysis of genes coding for enzymes participating in glycolysis, glycerol metabolism, the pentose phosphate pathway, glycogen metabolism, and TCA cycle revealed no significant transcriptional differences between $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts (Supplementary Table S5). Together our findings indicate that the metabolomic changes detected are regulated posttranscriptionally and linked specifically to BAP1 deubiquitylase activity.

Discussion

We discovered that normal primary cells carrying heterozygous germline *BAP1* mutations have increased aerobic glycolysis and impaired mitochondrial respiration. These metabolic changes in *BAP1*^{+/-} mutation carriers are so specific that, based on the metabolomics data from human plasma, we were able to predict *BAP1* status with 100% accuracy. The notion that genotype can be predicted by plasma metabolic analyses may seem implausible; however, the same results were reproducible regardless of the year of collection and gender or age of the individuals. Indeed, medium of fibroblast cell cultures derived from 12 of these same individuals retained many of the same metabolomics differences, underscoring the specificity of our findings.

We identified significant differences in glycolysis and TCA cycle metabolites between $BAP1^{+/-}$ and $BAP1^{WT}$ fibroblast cell cultures. $BAP1^{+/-}$ cells consumed more glucose and released more lactate in the culture medium, indicative of a faster glycolytic rate compared with BAP1^{WT}. These data indicate that $BAP1^{+/-}$ cells rely largely on the glycolytic metabolism for energy production. Indeed, in comparison with BAP1^{WT}, ECAR measurements in BAP1^{+/-} cells revealed a significantly increased rate of glycolysis, while we observed an overall reduction in all the OCR parameters that quantify mitochondrial respiration. ECAR was reduced and OCR restored in BAP1+/- cells transduced with WT BAP1 (AdBAP1) but not with a catalytic inactive BAP1 (AdBAP1 (C91S)). Moreover, similarly altered ECAR and OCR parameters were observed in BAP1^{WT} primary HM cells and fibroblasts silenced for BAP1, indicating that the effects observed are caused by the reduced BAP1 protein levels, independently of donor or cell type. Therefore, we anticipate that our results are also relevant to the numerous malignancies that carry somatic BAP1 mutations.33-44

The exact mechanism/s responsible for these metabolic changes is/are presently unknown. In a paper in press in *Nature*,¹⁹ we report that BAP1 modulates ER-to-mitochondria calcium (Ca²⁺) release.¹⁹ Intracellular Ca²⁺ modulates three rate-limiting enzymes of the TCA cycle.⁴⁵ Ongoing studies in our laboratory will determine whether the deregulation of intracellular Ca²⁺ signaling is also responsible for the distinctive metabolic signature of *BAP1* mutation carriers.

The increased aerobic glycolysis - 'Warburg effect'20,46 displayed by BAP1+/- cells is frequently observed in cancer cells. In carriers of germline BAP1 mutations, the presence of a Warburg effect in both the cells that undergo malignant transformation and in the surrounding stromal cells, creates an environment that promotes carcinogenesis and tumor growth.^{20,46} Indeed, in all cell cultures from individuals carrying germline BAP1 mutations, we observed increased levels of extracellular lactate. Lactate is known to promote cancer growth,⁴⁷ stimulate angiogenesis,⁴⁸ and polarize tumor-associated macrophages into a pro-tumor M2 phenotype.⁴⁹ Alterations in cytokine levels and macrophage polarizations with a pronounced M2 phenotype were observed in the peritoneal cavity of asbestos-exposed $Bap1^{+/-}$ mice and were linked to an increased incidence of MM.⁵⁰ These findings support a model in which alterations in cellular metabolism due to reduced BAP1 levels may also induce immune system alterations associated with a microenvironment that favors malignant transformation.

In summary, our data indicate that the Warburg effect, in addition to being a hallmark of cancer cells,²⁵ is also found in normal cells from individuals carrying heterozygous germline *BAP1* mutations and may contribute to the high incidence of cancer observed among them.

Materials and Methods

Subjects. All participants (affected and unaffected family members) provided written informed consent according to the guidelines set forth by the Institutional Review Board of the University of Hawaii. Of the 14 individuals studied here from the W family, 7 were BAP1^{WT} and 7 were BAP1^{+/-} (all healthy at the time of sample collection; however, $2/7 BAP1^{+/-}$ individuals had been previously diagnosed with (a) MM and (b) with MM and Breast cancer, both in remission for >7 and >15 years, respectively). Of the 32 individuals from the L family, 23 were BAP1^{WT} and 9 were BAP1+/- (all healthy at the time of sample collection, except 1/9 BAP1+/individuals diagnosed with uveal melanoma and MM 9 years prior to sample collection, who had stable disease at the time of blood collection). A total of 61 plasma samples were collected over a period of 2 years, 37 from $BAP1^{WT}$ and 24 from BAP1+/- individuals. We also studied 12 fibroblast cell cultures we established from skin punch biopsies from 6 BAP1+/- carriers (3 from the W and 3 from the L family, all healthy at the time of sample collection, although 1 of them previously diagnosed with cancers 15 years prior, see above) and 6 BAP1^{WT} control individuals from the same families (3 from the W and 3 from the L family).19

Cell cultures. Human dermal skin fibroblasts were derived from explants of skin biopsies of *BAP1^{WT}* and *BAP1^{+/-}* W and L families' members.¹⁹ For details, see Supplementary Experimental Procedures.

Plasma sample preparation and analysis by LC-TOF-MS. Metabolites were extracted from plasma samples and analyzed by LC-TOF-MS.

An Agilent HPLC 1200 system (Agilent Corporation, Santa Clara, CA, USA) was used with chromatographic separations performed on a 4.6 × 150 mm, 5 μ m Agilent ZORBAX Eclipse XDB-C18 chromatography column. Mass spectral data were acquired using an Agilent model 6220 MSD TOF mass spectrometer equipped with a dual sprayer electrospray ionization source (Agilent Corporation). Agilent API-TOF Reference Mass Solution Kit was used to obtain accurate mass time-of-flight data in both positive and negative mode operation. During metabolite profiling, both plot and centroid data were acquired for each sample from 50 to 1000 Da over a 25 min analysis time. Data generated by LC-TOF-MS were semiquantitative data and are expressed in peak intensity. For details, see Supplementary Experimental Procedures.

Plasma sample preparation and analysis by UPLC-TQ-MS. Plasma samples were prepared as previously described with modifications.^{51–53} The supernatant was used for targeted metabolic profiling of 140 lipids and amino acids with an Acquity ultra performance liquid chromatography coupled to a Xevo TQ-S

mass spectrometer (UPLC-TQ-MS, Waters Corp., Milford, MA, USA).⁵³ Data generated by UPLC-TQ-MS were quantitative data and are expressed in μ M.

Plasma sample preparation and analysis by GC-TOF-MS. Metabolites extracted from plasma samples were analyzed using an Agilent 7890N gas chromatograph coupled with a Pegasus HT TOF mass spectrometer (Leco Corporation, St. Joseph, MI, USA). Electron impact ionization (70 eV) at full scan mode (*m*/*z* 40–600) was used, with an acquisition rate of 20 spectra/s in the TOF-MS setting. Data generated by GC-TOF-MS were semiquantitative data and are expressed in peak intensity. For details, see Supplementary Experimental Procedures.

Cell extract preparation and analysis by UPLC-TQ-MS and GC-TOF-MS. $BAP1^{WT}$ and $BAP1^{+/-}$ fibroblasts were grown as described above, and 10^7 cells were collected for the analysis.

Cell samples were prepared as previously described with modifications^{52,53} as described in Supplementary Experimental Procedures. The supernatant was used for targeted metabolic profiling of 140 lipids with a UPLC-TQ-MS (Waters Corp., Milford, MA, USA)⁵³ and, for untargeted metabolic profiling, with an Agilent 7890A gas chromatograph coupled to a Leco Pegasus time of flight mass spectrometer (GC-TOF-MS, Leco Corporation).⁵²

Data analysis. The acquired data files from LC-TOF-MS, UPLC-TQ-MS, and GC-TOF-MS were processed, combined, and analyzed using multivariate statistical tools to establish characteristic metabolic profiles associated with different genotypes. In total, 412 plasma and 495 cells' chromatographic peaks corresponding to putative metabolites were found by deconvolution after exclusion of peaks originating from internal standards, contamination, and artifacts. Of these putative metabolites, 246 plasma metabolites and 226 cell metabolites were identified by their mass spectra and corresponding retention index (Table 1; see Supplementary Information).

¹³C-glucose metabolic pattern analysis

Cell culture, labeling, and sample collection: Fibroblasts were seeded at 5×10^5 cells/T25 flask, in DMEM, $1 \times , 25$ mM Glucose (Corning, Corning, NY, USA, Cat. No. 10-017-CV) supplemented with 10% Dialyzed FBS (Sigma, Cat. No. F0392). Labeling was started 24 h after seeding: cells were washed with phosphate-buffered saline (PBS) and culture medium was replaced with DMEM and no glucose (Gibco-Thermo Fisher Scientific, Waltham, MA, USA, Cat. No. 11966-025) supplemented with 10% Dialyzed FBS and 25 mM D-Glucose U-¹³C6 (Cambridge Isotope Laboratories, Inc., Tewksbury, MA, USA, Cat. No. 110187-42-3). A sample of the labeling media was taken at time zero and stored as a reference for analysis. ¹³C-labeled samples of medium and cells were collected following a 24-h labeling period. Cells were rinsed with PBS, detached with trypsin, and subjected to centrifugation at 200 × g for 5 min, at 4 °C. Cell pellets were washed three times with cold PBS, and the dry pellets were stored at - 80 °C for subsequent analysis.

UPLC-QTOF-MS analysis: Cell culture medium samples were processed by adding 150 µl of acetonitrile to 50 µl of samples, while cell pellets ($3-6 \times 10^6$ cells) were extracted with 75% (v/v) methanol. After being vortexed for 10 min and centrifuged for 20 min at 16 100 × *g*, the supernatants (culture medium or cell extracts) were used for UPLC-QTOFMS analysis.

All analyses were performed on a Waters UPLC system (UPLC Acquity, Waters Corp., Manchester, UK) coupled with a quadrupole–time of flight mass spectrometer (Synapt G2, Waters Corp., Manchester, UK). Metabolites separation was achieved through a 2.1 × 100 mm 1.7 μ m Acquity amide and an Acquity HSS C18 column (100 × 2.1 mm i.d., 1.7 μ m; Waters Corp., Manchester, UK) equipped with ACQUITY UPLC VanGuard Pre-Column, separately, according to the published methods with modifications.⁵⁴ The column was maintained at 40 °C and a 5 μ l aliquot of sample was injected. The flow rate remained constant at 0.4 ml/min. UPLC-MS raw data obtained were analyzed using TargetLynx applications manager version 4.1 (Waters Corp., Manchester, UK). Quantification was achieved for each metabolite using linear regression analysis of the peak area of metabolite *versus* concentration.

Gene silencing with siRNA, adenovirus-mediated gene transfer and transfection. siRNA oligonucleotides targeting four different BAP1 mRNAs and one negative control siRNA were obtained from Qiagen (Germantown, MD, USA) (GeneSoluton siRNA: Hs_BAP1_1, Cat. No.: Sl00066696; Hs_BAP1_2, Cat. No.: Sl00066703; Hs_BAP1_3, Cat. No.: Sl00066710; Hs_BAP1_5, Cat. No.: Sl03036390; AllStars Negative control siRNA, Cat. No.: 1027280). Tranfection was performed with HiPerfect (Qiagen), using a 10 nM final concentration of siRNA in 10 $\%\,$ FBS medium for 24 h.

The adenoviruses expressing BAP1 and GFP were purchased from SignaGen Laboratories (Rockville, MD, USA) (Ad-BAP1, Cat. No. SL175127; Ad-GFP, Cat. No. SL100708). A customized adenovirus expressing Ad-Myc-BAP1(C91S) was produced by SignaGen Laboratories using their Ad.MAX System.

XF96 instrument setup and analysis. *BAP1^{WT}* and *BAP1^{+/-}* fibroblasts were seeded in XF96 Cell Culture Microplates. OCR and ECAR were measured simultaneously using the XF96 Analyzer (Seahorse Biosciences, North Billerica, MA, USA), and values were normalized to the number of cells per well using the crystal violet method. For details, see Supplementary Experimental Procedures.

Conflict of Interest

MC has a pending patent applications on BAP1. MC provides consultation for mesothelioma diagnosis at no cost to patients and colleagues and for a fee to lawyers. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed.

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Author contributions

Conceptualization: MC, WJ, PP, HIP, HY; methodology: AB, CG, GS, PM, VS, SaP, LP, SK; validation: CG, SiP, PP; formal analysis: AB, WJ, AN, SiP, LP; investigation: AB, SiP, LP, MS, GX; writing: AB, AN, MC; resources: PM, GS, HIP, WJ; visualization: AB, WJ, AN, SiP, LP; supervision: MC, GG, WJ, PP, HY; project administration: MC, HY; funding acquisition: MC, WJ, PP, HY.

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