Dysfunctional MnSOD leads to redox dysregulation and activation of prosurvival AKT signaling in uterine leiomyomas

Vania Vidimar,1 David Gius,2 Debabrata Chakravarti,1 Serdar E. Bulun,1 Jian-Jun Wei,1,3 J. Julie Kim1*

AKT signaling promotes cell growth and survival and is often dysregulated via multiple mechanisms in different types of cancer, including uterine leiomyomas (ULMs). ULMs are highly prevalent fibrotic tumors that arise from the smooth muscular layer of the uterus, the myometrium (MM). ULMs pose a major public health issue because they can cause severe morbidity and poor pregnancy outcomes. We investigate the mechanisms driving ULM growth and survival via aberrant activation of AKT. We demonstrate that an acetylation-mediated impairment of the manganese superoxide dismutase (MnSOD) activity is prevalent in ULM cells compared to the normal-matched MM from the same patients. This impairment increases the levels of superoxide and oxidative stress, which activate AKT via oxidative inactivation of the phosphatase and tensin homolog deleted on chromosome 10 (PTEN). Redox activation of AKT promotes ULM cell survival under conditions of moderate but persistent oxidative stress that are compatible with ULM's prooxidative microenvironment. Moreover, because of impaired MnSOD activity, ULM cells are sensitive to high levels of reactive oxygen species (ROS) and superoxide-generating compounds, resulting in decreased ULM cell viability. On the contrary, MM cells with functional MnSOD are more resistant to high levels of oxidants. This study demonstrates a causative role of acetylation-mediated MnSOD dysfunction in activating prosurvival AKT signaling in ULMs. The specific AKT and redox states of ULM cells provide a potential novel therapeutic rationale to selectively target ULM cells because of their defective ROS-scavenging system.

INTRODUCTION

Abnormal cellular growth is a common neoplastic event in smooth muscle tissues, giving rise to tumors that escape normal growth control mechanisms and survive despite restrictive conditions (1). Uterine leiomyomas (ULMs), also called uterine fibroids, are benign smooth muscle tumors arising from the myometrium (MM) that grow to sizes that can exceed 10 cm in diameter and cause extreme morbidity in women, including vaginal bleeding, anemia, and poor pregnancy outcomes (2, 3). ULMs occur in approximately 70% of reproductive-age women and are the leading cause of hysterectomy in the United States (4, 5), resulting in a financial burden of $5.9 billion to $34.4 billion per year in health-related costs (6).

The pathophysiology of ULM has been relatively understudied. Previous findings from our group and others revealed that ULMs, unlike parental MM tissues, are characterized by abnormal activation of the AKT pathway (7, 8), a widely recognized critical regulator of tumor initiation and maintenance (9, 10). The biological relevance of the AKT pathway in ULM was demonstrated in our previous studies when, upon inhibition of AKT, multiple downstream effects, including decreased cell proliferation and viability, increased autophagy and death, and disruption of mitochondrial structures were observed in ULM cells (8, 11). However, the causes of AKT overactivation in ULM are unclear. In other systems, the AKT pathway was shown to be highly sensitive to variations of the intracellular redox state (12–14).

ULMs are hypoxic in nature, and, as in many disease states, low oxygen content and oxidative stress are closely associated (15). In ULMs, oxidative stress biomarkers, such as NADPH (reduced form of nicotinamide adenine dinucleotide phosphate) oxidase 4 (NOX4) (16) and 8-hydroxy-2'-deoxyguanosine (8-OH-dG) (17), have been shown to be increased compared to MM. Oxidative stress, which is characterized by elevated levels of reactive oxygen species (ROS), causes cellular damage. However, low to moderate levels of ROS play a critical role in fine-tuning normal cellular functions, such as proliferation, differentiation, and metabolic adaptation, because they are important second messengers that ensure cell homeostasis and mediate adaptation to stress (18–20). In ULM cells, it has been shown that ROS are important mediators of platelet-derived growth factor– and epidermal growth factor–dependent activation of mitogen-activated protein kinase signaling, thus promoting cell proliferation (21). Mitochondria are the major source and target of ROS, and to preserve their integrity under conditions of oxidative stress, cells have evolved sophisticated mechanisms to limit excessive ROS (22). In the mitochondria, the manganese superoxide dismutase (MnSOD; also known as SOD2) rapidly reacts with the superoxide anion (O2−), converting it into hydrogen peroxide (H2O2), which, in turn, diffuses across membranes and acts as a signaling molecule or is eliminated by catalase and other peroxidases in the form of water (23, 24). MnSOD is the primary ROS-scavenging enzyme of the cell and the only antioxidant enzyme indispensable for maintaining normal cell development and function (22, 23), and besides being a protective enzyme, it is also a regulator of intracellular redox signaling by controlling the O2−/H2O2 ratio. In cellular systems where the antioxidant machinery is impaired, a shift toward O2− has shown to create a favorable environment for cell transformation (25, 26). Recent mechanistic insights have demonstrated that the function of MnSOD is controlled by acetylation of specific and evolutionarily conserved lysine residue(s), suggesting that regulation of its activity via acetylation may play a key role in mitochondrial function (27). Hyperacetylation of MnSOD leads to loss of its enzymatic activity, resulting in increased O2− levels (28, 29).

1Division of Reproductive Science and Medicine, Department of Obstetrics and Gynecology, Northwestern University, Chicago, IL 60610, USA. 2Department of Radiation Oncology, Northwestern University, Chicago, IL 60610, USA. 3Department of Pathology, Northwestern University, Chicago, IL 60610, USA. *Corresponding author. Email: j-kim4@northwestern.edu

Given the key role of the AKT pathway in regulating mitochondrial dynamics and the evidence that AKT inhibition causes damage of mitochondria in ULM (8), we investigated the function of the primary ROS mitochondrial scavenger MnSOD and its possible involvement in activating AKT in ULM. Here, we demonstrate an interplay between inactivating MnSOD acetylation and activation of redox-sensitive AKT signaling in promoting ULM survival and a role for ROS in determining ULM cell fate.

RESULTS

An acetylation-mediated alteration of MnSOD activity determines the unique redox profile of ULM

Alterations in redox homeostasis are generally associated with changes in the expression or activity of antioxidant enzymes, such as mitochondrial MnSOD (30). It has previously been shown that acetylation of MnSOD at lysine (K) 122 (MnSOD K122-Ac) leads to inactivation of MnSOD, increasing mitochondrial O$_2^\text{•−}$ levels (29). To investigate whether MnSOD was intrinsically acetylated in ULM, a tissue microarray (TMA) containing a cohort of matched human MM and ULM tissue cores from 60 hysterectomy specimens was immunostained with an antibody to MnSOD K122-Ac (Fig. 1A). For each hysterectomy specimen, two tissue cores were derived from MM and three were derived from ULM. Immunoreactivities for MnSOD K122-Ac and total MnSOD were scored for intensity on a four-point scale by a pathologist, with 0 being negative, 1 being weak, 2 being moderate, and 3 being strong. The frequency distribution of each staining score (0 to 3) was calculated for MnSOD K122-Ac and total MnSOD antibodies, and data are presented in Fig. 1B. No significant change in immunoreactivity for total MnSOD was observed between MM and ULM specimens, whereas immunoreactivities for MnSOD K122-Ac were significantly higher in ULM compared to normal MM (P = 0.0009). Moreover, 53.8% of ULM tissue cores displayed the strongest immunointensity [score, 3 (strong)] for MnSOD K122-Ac compared to normal-matched MM (30.2%) (Fig. 1B). Sirtuin 3 (SIRT3) is known to interact with and deacetylate MnSOD, increasing its dismutating activity (28, 29). To investigate a possible role of SIRT3 in promoting MnSOD hyperacetylation in ULM, its expression was analyzed by staining the TMA with an antibody against SIRT3. As shown in fig. S1A, the γ− test revealed a significant difference between MM and ULM, in that SIRT3 expression was slightly increased in ULM tissues.

To assess whether MnSOD acetylation in ULM was associated with protein damage caused by increased reactive oxygen/nitrogen species, the TMA was immunostained with 3-nitrotyrosine (3-NO) antibody (Fig. 1A). 3-NO is a well-established biomarker of oxidative protein damage by peroxynitrite (ONOO$^−$), a highly reactive free radical derived from the reaction of intracellularly cogenerated O$_2^\text{•−}$ and nitric oxide (NO) (31). In ULM tissue cores, significant levels of 3-NO correlated with high levels of MnSOD K122-Ac (P < 0.0001). The strongest immunointensity for 3-NO (score, 3) was detected in 42.5% of ULM tissue against 17% of normal MM (Fig. 1B). Higher 3-NO expression was also associated with increased levels of inducible nitric oxide synthase (iNOS) in ULM (fig. S1B; P < 0.0001). iNOS generates large amounts of NO in response to a variety of stimuli, including inflammation, hypoxia, and steroid hormones, and its activity is often increased in gynecologic disorders (32, 33). The increases in 3-NO and iNOS expression in ULM suggest that high levels of NO and O$_2^\text{•−}$ are simultaneously generated in ULM, and these high levels are likely due to augmented iNOS and impaired MnSOD activity, respectively.

Next, MnSOD activity was assessed in patient-derived primary ULM cells and normal-matched MM cells. Of 13 cases, 10 demonstrated significantly decreased MnSOD activity in ULM cells compared to normal-matched MM cells (Fig. 1C). Together, these results suggest that ULMs are characterized by an acetylation-mediated impairment of MnSOD activity compared to healthy MM cells.

Mitochondrial ROS derived from dysfunctional acetylated MnSOD lead to AKT activation in ULM cells

ULM tissues displayed higher levels of MnSOD K122-Ac than normal MM (Fig. 1, A and B). We observed these differences in cultured patient-derived primary cells as well. Western blot analyses revealed that in most of the samples analyzed, ULM cells have higher levels of MnSOD K122-Ac compared to MM cells, and this corresponded to higher levels of phospho-AKT(S473) (pAKT). On the other hand, MM cells with lower MnSOD K122-Ac and higher total MnSOD content had lower levels of pAKT (Fig. 2A and fig. S2A). In the three patient-derived MM and ULM cells (pt#11 to pt#13 from Fig. 1C) where no changes in MnSOD activity were detected, we did not observe significant variations in pAKT levels (fig. S2B).

Because the AKT pathway is redox-sensitive and ULMs are characterized by a reduced capacity of detoxifying ROS due to aberrant MnSOD activity, we next assessed pAKT levels upon oxidative stimulation in ULM versus MM cells. Cells were treated with paraquat (PQ) or H$_2$O$_2$ for 6 hours. PQ is used to specifically generate O$_2^\text{•−}$ when analyzing oxidative stress (34). In ULM cells, PQ induced pAKT levels at all the concentrations used (Fig. 2B). Similarly, H$_2$O$_2$ increased pAKT levels at all doses (with a maximum increase at 50 μM) except at 500 μM (Fig. 2C). No changes in pAKT expression were detected in MM cells with PQ or H$_2$O$_2$.

To further investigate the role of dysfunctional MnSOD in activating the AKT pathway in ULMs, primary ULM cells from different patients were infected with a control lentivirus (lenti-CTR) or a mutant MnSOD K122-R lentivirus (lenti–MnSOD K122-R), and pAKT levels were analyzed. It has previously been demonstrated that substitution of lysine (K) 122 with arginine (R) results in a constitutively active dominant-positive form of MnSOD that mimics deacetylation (29). Infection of ULM cells with lenti–MnSOD K122-R led to a significant increase in both MnSOD activity and protein levels as well as a decrease in pAKT expression in ULM cells, supporting the idea that mitochondrial ROS derived from dysfunctional acetylated MnSOD are capable of activating AKT in ULM (Fig. 2D and fig. S3). This was further corroborated by the fact that pAKT levels were reduced following treatment with the MnSOD mimetic GC4419, a small molecule that selectively detoxifies O$_2^\text{•−}$ (Fig. 2E).

Because ULM and MM cells displayed a different ROS detoxification potential, we then investigated the sensitivity of those cells to oxidative stimuli. Exposure of ULM and MM cells to increasing doses of PQ or H$_2$O$_2$ for 24 hours showed lower cell viability in ULM than in MM cells (Fig. 2F), demonstrating an increased sensitivity to oxidative insults in ULM cells.

ULM cells feature aberrant MnSOD acetylation and AKT activation under both normoxic and hypoxic conditions

ULMs are solid tumors featuring a hypoxic environment unlike the normal uterine MM tissues (35, 36). We investigated whether low oxygen content may contribute to the aberrant acetylation of MnSOD and activation of AKT in ULM. First, we observed that ULM cells grown in hypoxia (2% O$_2$) had higher levels of mitochondrial O$_2^\text{•−}$ compared to...
the normoxic counterpart (20% O₂), as shown by increased MitoSOX staining (Fig. 3A). Also, Western blotting analysis revealed that hypoxia [verified by the stabilization of hypoxia-inducible factor–1α (HIF-1α)] further augmented the expression of pAKT. Under hypoxia, MnSOD K122-Ac levels were comparable to those of ULM cells grown in normoxia and around twofold higher compared to MM (Fig. 3B). Accordingly, we found that MnSOD activity in ULM cells grown in hypoxia was still lower than that in control MM cells and that the MnSOD mimetic GC4419 simultaneously increased the capacity of ULM cells to dismutate O₂•− and reduced pAKT levels under hypoxic conditions as shown before under normoxic conditions (Fig. 3, C and D).

**Inactive MnSOD promotes AKT activation via mitochondrial ROS–dependent PTEN oxidation**

ROS induce oxidative posttranslational modifications that can negatively affect the structure and function of many proteins, including the phosphatase and tensin homolog deleted on chromosome 10 (PTEN), a major negative regulator of AKT signaling (37). Reversible oxidation of PTEN
by H$_2$O$_2$ results in its inactivation and translocation to the nucleus (38, 39). To investigate whether increased pAKT expression in ULM was due to upstream inactivation of PTEN, we first measured PTEN activity in MM and ULM cells, assessing the conversion rate of its sub-
strate, phosphatidylinositol 3,4,5-trisphosphate (PIP3), to phosphatidy-
linositol 4,5-bisphosphate. PTEN activity was detectable at very low
levels in four of five patient-derived ULM cells, whereas MM cells
displayed significantly higher levels of PTEN activity (Fig. 4A).

In ULM, increased MnSOD K122-Ac and 3-NO staining indicated enhanced oxidative stress and O$_2$$_{-}$ formation. To determine whether endogenous ROS were capable of oxidizing and inactivating PTEN while increasing pAKT, basal levels of oxidized PTEN were analyzed in untreated MM and ULM cells. Oxidation of PTEN induces the appearance of a higher-mobility (oxidized) form of PTEN on an SDS–polyacrylamide gel electrophoresis (SDS-PAGE) gel under nonreducing conditions (40). Accumulation of the oxidized form of PTEN was observed at higher levels in protein extracts from ULM cells compared to MM cells (Fig. 4B). Moreover, treatment of ULM cells with increasing concentrations of PQ induced a further accumulation of the oxidized form of PTEN, usually peaking at 500 μM (Fig. 4C). No increase in oxidized PTEN was detectable in MM cells under the same conditions (fig. S4A). Consistently, immunofluorescent staining showed increased nuclear accumulation of PTEN in ULM cells following treatment with 100 μM PQ compared to untreated ULM cells (fig. S4, B to D). Quantitation of the nuclear/cytoplasmic ratio of PTEN cellular distribution further confirmed a significantly increased nuclear localization of PTEN upon PQ treatment (fig. S4E).

To assess whether acetylated MnSOD was responsible for PTEN oxidation due to increased O$_2$$_{-}$ and oxidative stress, ULM cells were treated with PQ together with the MnSOD mimetic agent GC4419. Treatment of ULM cells with the combination of 100 μM PQ and 10 μM GC4419 reduced both the appearance of the oxidized form of PTEN and pAKT levels induced by PQ alone (Fig. 4D). Together, these results suggest that dysfunctional acetylated MnSOD sustains AKT activation through mitochondrial ROS–dependent oxidation of PTEN in ULM.
Fig. 3. ULM cells are characterized by aberrant MnSOD acetylation and AKT activation under normoxic and hypoxic conditions. (A) MitoSOX Red was used to assess mitochondrial superoxide levels in ULM cells under normoxic (20% O₂) and hypoxic (2% O₂) conditions after 48 hours of incubation time. MitoSOX fluorescence was quantified by analyzing the fluorescence intensity of more than 10 cells for each condition using ImageJ software (***P < 0.0001; one-way ANOVA, n = 3 independent experiments ± SD), RFU, relative fluorescence unit. (B) ULM cells were grown under normoxia and hypoxia for 48 hours, and levels of MnSOD K122-Ac and pAKT(S473) were analyzed. MM cells grown in normoxia were used as control. Hypoxia was verified by detection of HIF-1α and AKT levels were analyzed by Western blotting, and densitometric analysis of three independent experiments is shown as means ± SD (***P < 0.0001; one-way ANOVA, n = 3).

MK-2206 effectively triggers death of ULM cells through simultaneous AKT inhibition and increase of superoxide

The potential involvement of the AKT pathway in regulating ULM growth and survival has previously been reported by Sefton et al. (8) and Hoekstra et al. (11). MK-2206 is a selective allosteric pan-AKT inhibitor that leads to multiple downstream effects depending on the dose and/or duration of the treatment. Treatment of ULM cells with increasing concentrations of MK-2206 effectively decreased pAKT protein levels at all the concentrations used (fig. S5A). In agreement with previous findings, 25 μM MK-2206 reduced viability of ULM cells (fig. S5B) despite the fact that there is no increase in cleaved caspases (8). Down-regulation of all AKT isoforms by specific small interfering RNA (siRNA) transfection did not affect ULM cell viability (fig. S5C). Given the recent evidence that the MK-2206 compound promotes ROS generation (41, 42) and disrupts mitochondrial structures (8), we sought to determine whether MK-2206 specifically induced O₂•−, causing ULM cell death. We observed that treatment of ULM cells with increasing concentrations of MK-2206 (from 1 to 10 to 25 μM) resulted in a dose-dependent increase in O₂•− levels (Fig. 5, A and B), unlike silencing of AKT, which did not alter O₂•− levels (fig. S5D). High doses of MK-2206 increased superoxide generation in MM cells as well (fig. S5E). Reduced glutathione (GSH) is another ROS-scavenging molecule that is oxidized [to GSSG (oxidized GSH)] and is one of the first lines of defense against oxidative stress in mammalian cells (43). We found that MK-2206 decreased the levels of the reduced form of GSH in a dose-dependent manner (Fig. 5C), which is suggestive of GSH oxidation by ROS, because its levels were inversely proportional to MK-2206–induced O₂•− levels (Fig. 5, A and B). The GSH synthesis inhibitor buthionine sulfoximine (BSO) and the antioxidant N-acetyl-cysteine (NAC) were used as negative and positive controls, respectively. Treatment of ULM cells with the MnSOD mimetic GC4419 inhibited MK-2206–induced increase of O₂•− in ULM cells (Fig. 5, A and B). Moreover, we found that MK-2206–induced cytotoxicity was completely reversed by GC4419 in ULM and MM cells. Although viability of MM cells decreased with 25 μM MK-2206, the decrease observed in ULM cells was greater (Fig. 5D). These results suggest that the AKT inhibitor MK-2206 promotes superoxide-mediated toxicity in the background of inactive AKT.

AKT serves as a survival factor in the prooxidative environment of ULM

To further demonstrate the idea that ROS are cytotoxic in ULM when AKT is inactive or, in other words, the idea that AKT serves as a survival factor for ULM under high oxidative stress, AKT was silenced using siRNA (fig. S6A) in ULM cells that were consequently treated with increasing concentrations of H₂O₂. H₂O₂ led to a progressive decrease of ULM cell viability compared to control siRNA (siCTR) cells treated with the same doses of H₂O₂ (Fig. 6A). Moreover, viability of MM cells under the same conditions was unaffected (Fig. 6B and fig. S6B). Together, these results suggest that AKT serves as a survival factor in the prooxidative environment occurring in ULM.

DISCUSSION

Oxidative stress plays a role in the pathophysiology of several gynecologic diseases, including ULM. Previous studies showed that multiple oxidative stress biomarkers, such as NOX4 (16), 8-OH-dG (17), and malonaldehyde (44), as well as serum oxidative stress markers (45), are increased in ULM. Moreover, unlike immortalized MM cells, activity of both catalase and extracellular SOD3 is significantly reduced in...
immortalized ULM cells (46). To date, our study is the most comprehensive study to investigate the redox regulatory system in exclusively patient-derived ULM and MM specimens and, thus, significantly furthers our understanding of the redox biology of ULM. We have demonstrated that patient-derived primary ULM cells are inherently characterized by an acetylation-mediated impairment of MnSOD activity compared to healthy MM cells. In ULM, inactivating acetylation of MnSOD increases the levels of mitochondrial ROS, leading to a state of persistent oxidative stress that activates AKT through oxidative inactivation of PTEN. Redox-sensitive AKT activation in turn promotes cell survival in the prooxidative ULM microenvironment (Fig. 6C). Therefore, our results point to a potential key signaling role of mitochondrial ROS and mild oxidative stress in the biology of ULM. Because of the dysfunctional MnSOD activity, ULM cells also feature a defective response to high ROS concentrations. In contrast to MM cells, whose viability is less affected by exposure to oxidants, ULM cells that are deficient in detoxifying ROS are very sensitive to high doses of O$_2^-$. In contrast to MM cells, whose viability is less affected by exposure to oxidants, ULM cells that are deficient in detoxifying ROS are very sensitive to high doses of O$_2^-$. 

Several reports suggest that moderate increases in the levels of ROS, especially O$_2^-$, can affect numerous aspects of tumor initiation and progression (47, 48). Oxidative stress can result from defects in the cellular antioxidant response, of which MnSOD is a major component (30, 49). Tao et al. (29) demonstrated that the enzymatic function of MnSOD is regulated by lysine acetylation. They found that, in Sirt3$^{-/-}$ mouse embryonic fibroblasts, increased acetylation of MnSOD at lysine 122 (MnSOD K122-Ac) resulted in the inactivation of MnSOD and the subsequent increment of O$_2^-$ levels, promoting a tumor-permissive environment. We found that MnSOD K122-Ac was highly expressed in ~60% of ULM human tissues compared to the matched MM tissues analyzed, and this coincided with a decreased activity of MnSOD in ~70% of the patient-derived ULM cells compared to the normal-matched MM cells. We also showed that, in ULM, 3-NO and iNOS were accumulated in a similar fashion to the K122-acetylated inactive form of MnSOD (~60%). 3-NO is a well-known biomarker of oxidative stress (31) that is formed from the reaction between tyrosine residues of proteins and ONOO$, a damaging oxidant whose production depends on the availability of mitochondrial O$_2^-$ and NO. Therefore, high levels of MnSOD K122-Ac, 3-NO, and iNOS suggested a correlation between inactivating MnSOD acetylation and establishment of a prooxidative milieu in ULM that could be ascribed to increased mitochondrial O$_2^-$ levels. Up-regulation of NOX4 was also found in ULM compared to MM (16). It has previously been reported that mitochondrial O$_2^-$ derived from MnSOD deficiency increases the activity of extra-mitochondrial NADPH oxidase, the major source of cytosolic O$_2^-$, via a feed-forward mechanism (50–52). This leads to a vicious cycle of ROS-induced ROS release between mitochondrial and cytosolic sources of O$_2^-$ that synergistically fosters oxidative stress and may contribute to the pathogenesis of many tumor types, including ULM. Our study demonstrates the importance of acetylated MnSOD in a pathological context. The reasons for increased acetylated MnSOD in ULM are unknown. To date, no acetylates that acetylate MnSOD have been identified (53). It is known that mitochondrial SIRT3 plays a role in MnSOD deacetylation (28, 29), and members of the sirtuin deacetylase family were shown to be dysregulated in cancer (54). We did not detect lower SIRT3 expression in ULM, but rather an increase, suggesting that augmented acetylation of MnSOD is not likely due to a reduced deacetylation of MnSOD by SIRT3 but due to other unknown mechanisms. The increased acetylation of MnSOD may occur in instances of metabolic reprogramming. Cancer cells often exhibit an altered metabolism that is characterized by a shift from oxidative phosphorylation to
glycolysis, also known as the Warburg effect, and is associated with cancer cell growth and survival (55). The link between metabolic reprogramming and protein acetylation has been previously documented and includes the lactate-mediated increase of intracellular pH and the glycolytic increase in the production of acetyl–coenzyme A, which is the substrate used by acetyltransferases to acetylate proteins (56–58). ULMs appear to have features associated with the Warburg effect, such as increased hypoxia, acetylation, and augmented ROS. Whether metabolic reprogramming has occurred in ULM and whether this increases acetylation of MnSOD or other key proteins will require further investigation.

It has been shown that ULMs are locally hypoxic with abnormal vasculature, and it is thought that low oxygen levels may contribute to ULM pathogenesis and growth (35, 36). We found that, under hypoxic conditions, ULM cells exhibited increased levels of pAKT compared to the normoxic counterpart as well as higher levels of acetylated MnSOD and lower MnSOD activity than normal MM cells. Moreover, rescuing MnSOD activity prevents pAKT accumulation in hypoxia as similarly shown in normoxia. These data show that a low-oxygen environment further promotes oxidative stress and AKT dysregulation in ULM.

In our study, we demonstrated that AKT activation occurs via oxidative inactivation of PTEN in ULM and suggested that this is driven by MnSOD deficiency. It is known that PTEN is sensitive to variations of the intracellular redox state (38, 39, 59). In our model, aberrant MnSOD activity leads to increased levels of mitochondrial superoxide that may directly inactivate PTEN if released to the cytoplasm. Multiple studies showed that superoxide can be released from the mitochondria toward the cytosol via voltage-dependent anion channels under conditions of elevated mitochondrial oxidative stress (60–62). However, because superoxide is a short-lived radical species, indirect mechanisms of superoxide-dependent PTEN oxidation cannot be excluded. For example, Dikalov’s group (51, 52) showed that reduced MnSOD activity leads to increased H$_2$O$_2$ release from the mitochondria, which can trigger PTEN oxidation. A similar mechanism may occur in ULM, resulting in acetylated MnSOD–dependent oxidation of PTEN and ultimately AKT activation. Moreover, although we believe that mitochondria-derived ROS are largely responsible for PTEN oxidation, other mechanisms, such as increased growth factor receptor signaling, cannot be ruled out in inactivating PTEN, as previously shown by other groups (63, 64).

Fig. 5. Treatment with the AKT inhibitor MK-2206 leads to superoxide generation in ULM cells. (A) Mitochondrial superoxide levels were assessed in ULM cells using MitoSOX Red. ULM cells were treated with vehicle (CTR), various concentrations of MK-2206 (MK; 1, 10, and 25 μM) and 10 μM GC4419 (GC) alone or with 25 μM MK-2206 (GC+MK25) for 6 hours in serum-free media. Representative pictures from three independent experiments are shown. (B) MitoSOX fluorescence was quantified by analyzing the fluorescence intensity of more than 10 cells for each condition using ImageJ software (****P < 0.0001 versus 0 μM MK-2205; ####P < 0.0001 versus 25 μM MK-2205; one-way ANOVA, n = 3 independent experiments ± SD). (C) GSH content was quantified using GSH-Glo reagent. ULM cells were treated at the indicated concentrations of MK-2206 for 24 hours. BSO (10 μM) and 50 μM NAC were used as negative and positive controls, respectively (*P < 0.05, ****P < 0.0001 versus vehicle (dimethyl sulfoxide); one-way ANOVA, n = 3 independent experiments ± SD). (D) MM and ULM cells were treated with 25 μM MK-2206 alone or in combination with 10 μM GC4419 for 24 hours. Cell viability was determined using WST-1 (**P < 0.01, ####P < 0.0001; one-way ANOVA, n = 3 independent experiments ± SD).
Our findings suggest that mitochondrial ROS play an important role as a survival factor in ULM through increased AKT activation. Although AKT is localized in the membrane and cytosol, nuclear localization (65) and localization in the mitochondria have been reported (66). Under oxidative stress conditions, cytosolic AKT can translocate into the mitochondria to preserve mitochondrial integrity and provide cytoprotection (66–68). It would be interesting to investigate whether AKT localizes to the mitochondria in ULM cells to preserve their integrity following oxidative stimulation.

Together, we have provided new insights into the pathophysiology of ULM that go beyond the comparative expression analysis of redox targets in MM versus ULM. We have demonstrated the mechanisms by which ULM cells have adapted to survive in otherwise suboptimal conditions that are very different from those of the normal tissue from which they arise, the MM. Using exclusively human tissues and patient-derived primary cells, we have found an acetylation-mediated impairment of the activity of a critical ROS-sensing protein, MnSOD, which promotes a mitochondrial ROS–mediated increase in oxidative stress. Dysfunctional MnSOD activity not only increases oxidative stress to levels that promote survival through activation of the AKT pathway but also renders ULM cells to be more vulnerable to further prooxidative stimulation. Not all fibroids exhibited high MnSOD K122-Ac and reduced MnSOD activity, and not all matched patient samples showed a correlation between MnSOD K122-Ac and pAKT levels. The biological reasons for these differences remain unknown, but future investigation on the influence of the most prominent mutations that occur in ULM (such as in the MED12, HMGA2, and FH genes) on the oxidative stress–mediated activation of AKT would be of interest. The differential response to ROS between ULM and MM cells has translational potential and paves the way for possible redox-modulating therapeutic strategies that can selectively target ULM while minimizing the effects on normal MM that has a functional ROS-scavenging system.

**MATERIALS AND METHODS**

**Collection of tissue samples and culture of primary cells**

Human ULM and MM tissues were collected from premenopausal women (age range, 30 to 52 years) undergoing hysterectomy or myomectomy at the Northwestern University Prentice Women’s Hospital (Chicago, Illinois), according to an International Review Board–approved protocol. At the moment of the surgery, subjects included in the study were not taking hormonal contraceptives or a gonadotropin-releasing hormone antagonist or agonist for at least 3 months. Informed consent was obtained from all the patients participating in the study. Primary ULM and MM cells were isolated and cultured as previously described (69) and passaged up to two times before they were discarded. Primary cells were cultivated in Dulbecco’s modified Eagle’s medium/nutrient Ham’s Mixture F-12 (DMEM-F12) 1:1 containing 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin at 37°C and 5% CO2 atmosphere. For experiments in hypoxia, cells were placed in a humidified incubator at 37°C under a 2% O2, 5% CO2, and balance N2 atmosphere for 48 hours.

**Lentiviral transduction**

A lenti-CTR and a lenti–MnSOD K122-R were provided by D. Gius. ULM cells at 60% confluence were transiently infected with lenti-CTR and lenti–MnSOD K122-R (10^7 copies/ml for both) in the presence of polybreyne (8 μg/ml). After 2 days of infection, the culture medium...
of infected cells was replaced with fresh DMEM-F12 containing 10% FBS and 1% penicillin-streptomycin, and cells were cultured for an additional 2 days. Cells were then harvested, and protein extracts were prepared for both MnSOD activity assay and Western blot analysis.

**Chemicals**

GC4419 was provided by D. Gius and Galera Therapeutics Inc. MK-2206 was provided by Merck Sharp & Dohme Corp. and the National Cancer Institute, National Institutes of Health. PQ, H2O2, and NAC were purchased from Sigma-Aldrich, and BSO was purchased from Cayman Chemical.

**Immunohistochemical staining of TMAs**

Immunostaining of TMA was performed on an automated system by the Robert H. Lurie Comprehensive Cancer Center’s Pathology Core Facility. Anti-MnSOD (NB2P-20535; Novus Biologicals), anti–MnSOD K122-Ac (NCI-156, clone ID 33; Epitomics), anti–3-NO (06–284; EMD Millipore), INOS (PA3–030A; Thermo Fisher Scientific), and SIRT3 (ARP32389_P050; Aviva Systems Biology) antibodies were used at 1:200, 1:250, 1:500, 1:1000, and 1:250 dilutions, respectively. Primary antibodies were detected using a standard anti-rabbit secondary antibody followed by 3,3′-diaminobenzidine revelation (Dako).

**MnSOD activity assay**

MnSOD activity was measured using the Superoxide Dismutase Assay Kit (Cayman Chemical) following the manufacturer’s instructions for specific quantitation of mitochondrial MnSOD activity. Addition of 2 mM KCN (Sigma-Aldrich) (Cu–ZnSOD/SOD1 and FeSOD/SOD3 inhibitor) to the assay allowed specific detection of MnSOD activity. Absorbance was read at 450 nm using a Cytation 3 Cell Imaging Multi-Mode Reader (BioTek).

**SDS-PAGE and Western blotting**

Protein lysates were extracted from MM and ULM cells using M-PER mammalian protein extraction reagent (Thermo Fisher Scientific) with protease and phosphatase inhibitors (Sigma-Aldrich). Protein concentration was determined using Bradford reagent (Sigma-Aldrich). Equal amount of proteins were subjected to SDS-PAGE and subsequently transferred to polyvinylidene difluoride (PVDF) membranes. Immunoblotting was performed using the following primary antibodies: rabbit anti-pAKT(S473) (#9271; Cell Signaling Technology), rabbit antipan-AKT (#2932; Cell Signaling Technology), rabbit anti-AKT2 (#2964; Cell Signaling Technology), rabbit anti-AKT1 (AB2056; Cell Signaling Technology), rabbit anti-MnSOD (NB2P-20535; Novus Biologicals), rabbit anti–MnSOD K122-Ac (NCI-156 Clone ID 33; Epitomics), and mouse anti-actin (A1972; Sigma-Aldrich). Secondary antibodies were horse-radish peroxidase (HRP)–labeled anti-mouse or anti-rabbit (Bio-Rad). Chemiluminescence was detected by adding a chemiluminescent HRP substrate (Thermo Fisher Scientific) and measured with a Fujifilm LAS-3000 Imager.

**WST-1 viability assay**

ULM and/or MM cells were cultured in 96-well plates in complete DMEM-F12 1:1 medium until 80 to 90% confluence. At the end of treatments, WST-1 (BioVision) was added to each well (1:10 dilution) for 2 hours at 37°C, and absorbance was read at 420 nm with reference wavelength set at 650 nm. Data were analyzed following the manufacturer’s manual.

**PTEN activity assay**

MM and ULM cells were cultured in 100-mm dishes in complete DMEM-F12 1:1 medium until 80 to 90% confluence. Cells were lysed in ice-cold 25 mM tris-HCl (pH 8.0), 150 mM NaCl, 1% NP-40, 1 mM EDTA, and 5% glycerol lysis buffer. PTEN (500 μg) was immunoprecipitated from whole-cell lysates with 8 μl of rabbit anti-PTEN antibody (Cell Signaling Technologies) under agitation at 4°C overnight. The antigen-antibody complex was recovered with 30 μl of protein A Sepharose beads (Sigma-Aldrich) after 3 hours with agitation at 4°C, and then the beads were washed twice in lysis buffer, washed once in PTEN reaction buffer [25 mM tris-HCl (pH 7.4), 140 mM NaCl, 2.7 mM KCl, and 10 mM DTT], and resuspended in 80 μl of PTEN reaction buffer. PTEN phosphatase activity was measured using the Malachite Green Assay Kit (Echelon Biosciences) and PIP3 as substrate according to the manufacturer’s instructions.

**Detection of oxidized PTEN**

MM and ULM cells were cultured in 60-mm dishes in complete DMEM-F12 1:1 medium until 80 to 90% confluence. Untreated cells or cells treated with PQ at the indicated concentrations for 6 hours were washed twice with phosphate-buffered saline (PBS)/EDTA, scraped in PBS/EDTA, and collected by centrifugation. Cell pellets were washed with PBS and resuspended in 200 μl of lysis buffer [100 mM tris-HCl (pH 6.8), 2% SDS, and 40 mM N-ethylmaleimide] as described by Connor et al. (40). Equal amounts of protein were loaded on a non-reducing SDS-PAGE gel (8% polyacrylamide) and electroblotted onto a PVDF membrane, which was incubated first with anti-PTEN (Cell Signaling Technologies) and then with anti-rabbit secondary antibody conjugated to HRP (Bio-Rad). Chemiluminescence was detected by adding a chemiluminescent HRP substrate (Thermo Fisher Scientific) and measured with a Fujifilm LAS-3000 Imager.

**Immunofluorescence staining for oxidized PTEN**

ULM cells were grown on glass coverslips in a 12-well plate in complete DMEM-F12 1:1 medium until 70% confluence. After treatment with PQ at the indicated concentrations for 6 hours, cells were washed three times with PBS, fixed with 4% paraformaldehyde (PFA), permeabilized with 0.1% Triton X-100 in PBS, and blocked with 5% bovine serum albumin in PBS for 1 hour. Cells were then incubated with anti-PTEN antibody (1:200; Cell Signaling Technologies) overnight at 4°C. Anti-rabbit secondary antibody Alexa Fluor 488 (Thermo Fisher Scientific) was then added for 1 hour at room temperature. Coverslips were then mounted onto glass slides using ProLong Gold Antifade reagent with 4′,6-diamidino-2-phenylindole (DAPI) (Thermo Fisher Scientific) as counterstain to visualize the nuclei. Images were taken using a Leica DM5000 B microscope.

**PTEN nuclear/cytoplasmic ratio**

The nuclear and cytoplasmic distribution of PTEN was analyzed using ImageJ software and R. Details about the procedure can be found in the Supplementary Materials. The mean nuclear/cytoplasmic PTEN signal ratio of more than 10 cells per condition was measured using ImageJ software. Average ratios and SDs from independent experiments were plotted using GraphPad Prism software.

**Superoxide anion detection**

ULM cells were grown on glass coverslips in a 12-well plate in complete DMEM-F12 1:1 medium until 70% confluence. Cells were then washed twice with Hanks’ balanced salt solution (HBSS) Ca-Mg and preincubated...
with 5 μM MitoSOX Red (Thermo Fisher Scientific) in HBSS Ca-Mg for 20 min at 37°C in the dark. MitoSOX Red allows the selective visualization of O$_2^•^−$ generated in the mitochondria because it is rapidly oxidized by O$_2^•^−$ only. After treatments, the dye was removed, and the cells were washed three times with HBSS Ca-Mg and fixed with 4% PFA. Coverslips were then mounted onto glass slides using ProLong Gold Antifade reagent with DAPI (Thermo Fisher Scientific) as counterstaining to visualize the nuclei. Images were taken using a Leica DMS5000 B microscope.

**Relative fluorescence unit quantification**

The relative fluorescence units of individual cells were quantified using ImageJ software. Corrected total cell fluorescence units were determined using the following formula: integrated density of selected cell – (area of selected cell × mean integrated density of background readings).

**GSH measurement**

ULM cells were cultured in a 96-well white plate in complete DMEM-F12 1:1 medium until 80 to 90% confluence. At the end of treatments, GSH content was measured using the GSH-Glo Glutathione Assay (Promega Corporation) according to the supplier’s instructions. Luminescence was read using a luminometer plate reader (Cytation 3 Cell Imaging Multi-Mode Reader, BioTek).

**AKT knockdown, RNA isolation, and RT-PCR**

AKT1, AKT2, and AKT3 were silenced in ULM or MM cells by reverse transfection using siAKT1, siAKT2, and siAKT3 ON-TARGET plus SMARTpool (GE Dharmacon) and Lipofectamine RNAiMAX (Thermo Fisher Scientific) according to the manufacturer’s instructions. A nontargeting siCTR (GE Dharmacon) was used in parallel. Cells were harvested for immunoblotting or reverse transcription polymerase chain reaction (RT-PCR) after 72 hours from transfection.

RNA was isolated from ULM cells using TRIzol Reagent (Thermo Fisher Scientific) and reverse-transcribed with M-MLV Reverse Transcriptase (Thermo Fisher Scientific) following the manufacturer’s instructions. AKT3 TaqMan gene expression assay was purchased from Thermo Fisher Scientific on a QuantStudio12K Flex RT-PCR system (Thermo Fisher Scientific). 18S was used as housekeeping gene, and relative mRNA levels were calculated using the 2$^{-ΔΔCt}$ method. Each data point is the average of three replicates.

**Statistical analysis**

GraphPad Prism software was used for statistical analysis. According to the experimental design, $χ^2$ test, unpaired t test, paired t test, or one-way ANOVA was performed. Statistical analysis on fold change data was performed after log transformation of the data to obtain a more normalized distribution. Data from each patient were considered as an independent experiment.

**REFERENCES AND NOTES**


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Vania Vidimar, David Gius, Debra B. Chakravarti, Serdar E. Bulun, Jian-Jun Wei and J. Julie Kim

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