Research Paper

Multiple Myeloma Tumor Cells are Selectively Killed by Pharmacologically-dosed Ascorbic Acid

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Abstract

High-dose chemotherapies to treat multiple myeloma (MM) can be life-threatening due to toxicities to normal cells and there is a need to target only tumor cells and/or lower standard drug dosage without losing efficacy. We show that pharmacologically-dosed ascorbic acid (PAA), in the presence of iron, leads to the formation of highly reactive oxygen species (ROS) resulting in cell death. PAA selectively kills CD138+ MM tumor cells derived from MM and smoldering MM (SMM) but not from monoclonal gammopathy undetermined significance (MGUS) patients. PAA alone or in combination with melphalan inhibits tumor formation in MM xenograft mice. This study shows PAA efficacy on primary cancer cells and cell lines in vitro and in vivo.

1. Introduction

Multiple myeloma (MM) is a plasma cell neoplasm. Four active classes of drugs including glucocorticoids, DNA alkylators (melphalan), proteasome inhibitors (bortezomib and carfilzomib) and immunomodulatory agents (thalidomide, lenalidomide, and pomalidomide), combined with or without autologous stem cell transplantation (ASCT) have led to complete remissions (CRs) in the large majority of newly diagnosed patients with MM (Alexanian et al., 2013; Gay et al., 2013; Liu et al., 2013; Bergsagel, 2014). These treatments have greatly improved patient progression-free and overall survival. However, there are at least three major problems limiting the administration of these agents: 1. All these drugs target both tumor and non-tumor cells; 2. Increased hematologic toxicity has been identified by combining alkylators with either immunomodulatory drugs (IMIDs) (Bergsagel, 2014); and 3. High doses of the DNA alkylating agent, such as melphalan, have strong cytotoxicity on gut epithelial cells and hematopoietic stem cells (Shaw et al., 2014). One way to deal with non-selective toxicity of high dose melphalan is to combine it with another agent which very specifically targets tumor cells and therefore decreasing melphalan dosing without loss of efficacy.

In the 1970s, Cameron and Pauling reported that high doses of vitamin C increased survival of patients with cancer (Cameron and Pauling, 1976, 1978). Recently, reports have shown that pharmacologically-dosed ascorbic acid (PAA) 50–100 g (Chen et al., 2007, 2005). In a previous study, we have reported that the labile iron pool (LIP) is significantly elevated in MM cells (Gu et al., 2015), suggesting that PAA treatment should target MM cells quite selectively. The higher LIP is the direct result of the low expression of catalytic metal ions like iron, PAA administered intravenously exerts pro-oxidant effects leading to the formation of highly reactive oxygen species (ROS), resulting in cell death (Yun et al., 2015; Ma et al., 2014). In the presence of catalytic metal ions like iron, PAA administered intravenously exerts pro-oxidant effects leading to the formation of highly reactive oxygen species (ROS), resulting in cell death (Yun et al., 2015; Ma et al., 2014; Du et al., 2012; Chen et al., 2007, 2005). In a previous study, we have reported that the labile iron pool (LIP) is significantly elevated in MM cells (Gu et al., 2015), suggesting that PAA treatment should target MM cells quite selectively. The higher LIP is the direct result of the low expression of the only known mammalian cellular iron exporter, Ferroportin 1 (Fpn1), in MM as demonstrated by our group (Gu et al., 2015). These findings led us to the hypothesis that PAA might specifically target MM cells with high iron content and may also act synergistically in combination with commonly used MM therapies.
2. Materials and Methods

2.1. Patients Samples

Peripheral-blood samples or bone marrow aspirates were obtained from patients with monoclonal gammopathy of undetermined significance (MGUS), smoldering multiple myeloma (SMM), and multiple myeloma (MM). Written informed consent was obtained from all participants. The de-identified clinical specimens in this study were approved by the institutional review board at the University of Iowa (HawLdRB protocol 201302833).

2.2. Gene Expression

Gene expression profiling (GEP) has been described previously (Zhan et al., 2006; Shaughnessy et al., 2007). The GEP access number of normal plasma cell (NPC), MGUS, and primary myeloma samples is GSE2658.

2.3. Viability Assay

Pharmacological ascorbic acid (PAA) was kindly provided by Dr. Garry R. Buettner (University of Iowa). Dr. Buettner prepares PAA as previously described (Du et al., 2010). Briefly, L-ascorbic acid was from MACRON Fine Chemicals/Avantor Performance Materia (Center Valley, PA, USA). A stock solution of 1.0 M ascorbate in de-ionized water (pH adjusted to 7.0 with NaOH) was made under argon and stored in a volumetric flask with a tight-fitting stopper at 4°C. Ascorbate concentration was checked at 265 nm, ε = 14,500 M⁻¹ cm⁻¹ (Buettner, 1998). The solution can be kept for several weeks without significant loss of ascorbate due to the lack of oxygen. CD138⁺ MM cells and CD138⁻ non-MM cells were isolated from MGUS, SMM, and MM patient samples using anti-CD138 immunomagnetic beads (Miltenyl Biotec, Auburn, CA). Cells were cultured with or without PAA at the described concentration for 1 h. After incubation, the cells were washed and cultured up to 24 h. Cell counts and viable cell number were determined using Trypan Blue staining.

2.4. Xenograft Mouse Model

The animal study was performed according to the guidelines of the Institutional Animal Care and local veterinary office and ethics committee of the University of Iowa, USA under approved protocol (IACUC 5081482). NOD.CγR1 mice 6–8 weeks old (Jackson Laboratory, Bar Harbor, Maine) were injected intravenously with ARP1 MM cells (1 × 10⁶) expressing luciferase. After one-week injection of ARP1 cells, mice were treated with either PAA (4 mg/kg) injected intraperitoneal once a day, 5 days every week for 3 weeks. Melphalan (3 mg/kg) was injected intraperitoneal once a day, 2 days a week for 3 weeks (Sanchez et al., 2012). Carfilzomib (3 mg/kg) was injected by in vein once a day, 2 days every week for 3 weeks (Eda et al., 2014). Bortezomib (3 mg/kg) was injected intraperitoneal once a day, 2 days a week for 3 weeks. The mice were euthanized when a humane endpoint was reached.

2.5. In Vivo Imaging System

Xenogen IVIS-200, an in vivo imaging system (IVIS), was used to analyze tumor burden and was indicated by quantification of luciferase intensity of mice pre- and post-treatments.

2.6. Cell Culture

Human myeloma cell lines (ARP1, OCI-MY5 and their derivative cell lines) were cultured in RPMI 1640 (Invitrogen, Carlsbad, CA), supplemented with 10% heat-inactivated FBS (Invitrogen), penicillin (100 IU/ml), and streptomycin (100 µg/mL) in a humidified incubator at 37°C and 5% CO₂/95% air. To increase cellular iron concentration, ferric nitrilotriacatate (Fe-NTA) was added in the cell culture media.

2.7. Western Blotting

Cells were harvested and lysed with lysis buffer: 150 mM NaCl, 10 mM EDTA, 10 mM Tris, pH 7.4, 1% X-100 Triton. Cell lysates were subjected to SDS-PAGE, transferred onto a pure nitrocellulose membrane (BioRad) and blocked with 5% fat-free milk. Primary antibodies for immunoblotting included: anti-AIF1 (1:1000, Cell Signaling), anti-RIP (1:1000, Santa Cruz Biotechnology), anti-RIP3 (1:1000, Cell Signaling), anti-Caspase3 (1:1000, Cell Signaling), anti-Caspase 8 (1:1000, Cell Signaling), anti-Caspase 9 (1:1000, Cell Signaling), phosphorylated γ-H2AX (1:1000, Enzo Life Sciences), and β-actin (1:1000, Cell Signaling) as loading control. Membranes were incubated with horseradish peroxidase (HRP)-conjugated anti-mouse secondary antibody (1:10,000, Santa Cruz Biotechnology, cat#: sc-2005) or anti-rabbit secondary antibody (1:10,000, AnaSpec Inc., cat#: AS-28177) for 1 h and chemi-luminescence signals were detected by HRP substrate (EMD Millipore). Pan-caspase inhibitor Q-VD-OPh (Sigma-Aldrich, MO) was at final concentration of 25 µM.

2.8. Statistical Analysis

GEP data were analyzed by one-way ANOVA test using log2 transformed Affymetrix Signals and presented by boxplot. The comparisons of tumor burden were analyzed either by student t-test (2 groups) or by one-way ANOVA test (≥2 groups). Kaplan-Meier test was performed for survival with the use of SPSS 16.0 software (SPSS, Chicago, IL). Two-tailed p value at an alpha level of 0.05 was considered to indicate statistical significance. Graphs were generated using Prism 6 software.

2.9. Electron Microscopy

Electron microscopy was performed by the Central Microscopy Research Facility personnel at the University of Iowa. Images were captured on JEOL JEM 1230.

3. Results

3.1. Pharmacological Ascorbic Acid Selectively Kills Primary Myeloma Cells

We analyzed the response to PAA of both CD138⁺ primary MM cells (high cytosolic iron) and CD138⁻ non-MM bone marrow (BM) cells obtained from 13 patients. The 13 patients included 2 pre-cancer of monoclonal gammopathy of undetermined significance (MGUS), 2 smoldering MM (SMM) and 9 MM patients. Patient demographic, disease characteristics and therapy are listed in Supplementary Table 1 and Supplementary Fig. 1. The survival of CD138⁺ cells in vitro was significantly decreased following PAA treatment in all 9 MM (Fig. 1A, grey bars, p < 0.01). In contrast, no significant change of cell viability was observed in CD138⁻ BM cells from the same patients (Fig. 1A, black bars). The same effect of PAA was also observed in the SMM patients (Fig. 1B). However, almost no response to PAA was detected in CD138⁻ cells from the 2 MGUS patients (Fig. 1C). We predicted that this would be the case because MGUS patients have much lower cytosolic iron compared to MM patients (Supplementary Fig. 2) as the consequence of lower expression of transferrin receptor 1 (TIR1), the cellular iron receptor-mediated importer (Supplementary Fig. 2A), and higher expression of Ferroportin 1 (Fpn1), the iron exporter (Supplementary Fig. 2B).
3.2. Pharmacological Ascorbic Acid Decreases Melphalan Doses in Myeloma Treatment

To investigate whether PAA may be effective in killing MM cells alone and when combined with currently used MM therapies. ARP1 MM cells expressing luciferase were injected intravenously into NOD.Cy-Rag1 mice. Seven combinations (control, PAA, melphalan, carfilzomib, melphalan + PAA, carfilzomib + PAA and bortezomib) were tested in vivo (Sanchez et al., 2012; Eda et al., 2014). Tumor progression was monitored by an in vivo imaging system (IVIS). Compared to the control group, all treatments inhibited MM cell growth significantly (p < 0.05) (Fig. 1D). Within the single agent treatments, melphalan only showed a higher decrease in tumor burden when compared to PAA treatment and other single agents tested. Also, the combination of melphalan plus PAA showed greater tumor burden reduction than each drug alone, suggesting a synergistic activity between these two drugs (p < 0.05). Under the experimental conditions described in the Fig. 1, no additive effect was found when PAA was combined with carfilzomib (p > 0.05). Bortezomib was not given in combination with PAA because ascorbic acid directly inactivates bortezomib by forming a tight and reversible complex through its vicinal diol group (Perrone et al., 2009; Harvey et al., 2009).

The above results showing that a synergistic effect was observed in MM treatment by combination of PAA with melphalan encouraged us to further determine if PAA addition would allow a decrease in melphalan dosing without losing efficacy. Therefore, mice were treated with 3 different doses of melphalan (1, 3, and 5 mg/kg) plus PAA. Tumor burden at three weeks of treatment showed that a single agent melphalan at the lowest dose was able to inhibit tumor growth better than PAA alone (Fig. 1E & F). Further, the presence of tumor at the highest dose of melphalan was detected only after four weeks confirming that the high dose of melphalan had greater anti-tumor effect. In contrast, no difference in outcome was observed when melphalan was combined with PAA even at the lowest dose. Reduction of mouse weight was not observed suggesting lack of toxicity (Supplementary Fig. 3). Tumor burden was almost undetectable in mice treated with any of the three combinational therapies (Fig. 1F). Survival curves confirmed that high doses of single agent melphalan (3 and 5 mg/kg) extended MM mouse survival (Fig. 1G) better than PAA alone. However, the combination of PAA with low-dose melphalan (1 mg/kg) extended MM mouse survival significantly compared with low-dose melphalan alone (Fig. 1G; p < 0.05). Importantly, no survival differences were observed between low and high doses of melphalan when given in combination with PAA (Fig. 1G and Supplementary Fig. 4).

3.3. The Therapeutic Effect of Pharmacological Ascorbic Acid Depends on Cellular Iron and Reactive Oxygen Species

We subsequently asked whether PAA was selectively killing MM tumor cells by generating ROS, we treated OCI-MYS MM wild-type (WT) cells with N-acetyl cysteine (NAC) or catalase. Both catalase and NAC protect cells from oxidative damage. OCI-MYS cells pretreated with NAC and catalase became resistant to PAA even at high doses (Fig. 2A). Importantly, adding deferoxamine (DFO), an iron chelator, to OCI-MYS cells before PAA treatment was also sufficient to prevent PAA-induced cellular death (Fig. 2A) but bathocuproinedisulfonic acid disodium salt (BCS), a selective copper chelator, was not able to block PAA-induced cellular death (Supplementary Fig. 5A) suggesting that iron is essential for PAA to achieve its anti-cancer activity. DFO is a poorly membrane permeable iron chelator, however, it has been used to chelate intracellular iron in multiple studies including those ones that describe cells expressing Fpn1 (Asano et al., 2011; Paradkar et al., 2008; Al-Qenaei et al., 2014). We reasoned that high cytosolic iron would catalyze PAA auto-
oxidation leading to cell death. Because MM tumor cells have a higher labile iron pool (LIP) than non-tumor cells, we hypothesized that PAA's anti-cancer effect is dependent on LIP. We have previously shown that Fpn1, the only known mammalian iron exporter, is down-regulated in MM cells at the expression levels leading to higher LIP. We next sought to determine if higher Fpn1 levels in MM tumor cells could also block cell death mediated by PAA. We overexpressed and confirmed Fpn1 expression by qRT-PCR in OCI-MY5 cells (Supplementary Fig. 5B, OE-Fpn1). We noticed that 4 mM PAA was able to kill OCI-MY5 transfected with empty vector (EV) but not to overexpressing Fpn1 (OE-Fpn1) cells (Fig. 2B). Five-fold greater concentration of PAA (20 mM) was required to successfully kill OE-Fpn1 cells. Since the overexpression of Fpn1 in OCI-MY5 cells inhibits PAA anti-cancer activity, we next explored whether iron supplementation was able to restore sensitivity of Fpn1 OCI-MY5 cells (Fig. 2D). Consistent with our hypothesis, DFO, an iron chelator (Fig. 2C & D), abolished the ability of PAA to reduce cell viability in both EV and OE-Fpn1 OCI-MY5 cells pre-treated with iron. Interestingly, we noticed that EV OCI-MY5 cells in Fig. 2C treated with PAA alone showed a stronger sensitivity than those cells in Fig. 2B. We speculated that even if the PAA incubation is 1 h in both experiments the cells in Fig. 2C were kept in culture almost 24 h longer than cells in Fig. 2B for incubation with other reagents before PAA treatment. A possible explanation is that the longer incubation with culture media may slightly increase cellular iron (Goto et al., 1983). This was supported by the evidence that either addition of DFO (Fig. 2C) or overexpression of Fpn1 (Fig. 2D) rescued PAA-induced MM cell death even cell viability was different between control groups, because both DFO and overexpression of Fpn1 decrease cellular iron.

3.4. Pharmacological Ascorbic Acid Induces Both Necrosis and Apoptosis in Myeloma Cells

To determine the type of cell death induced by PAA, we performed transmission electron microscopy (TEM) experiments. Fig. 3A shows that in OCI-MY5 WT cells incubated with 4 mM PAA for 1 h and then left for another 2 h, PAA induced early necrosis (Fig. 3A, 60 min) followed by late apoptosis (Fig. 3A, 120 min). OCI-MY5 WT cells untreated appeared healthy and the mitochondria had visible cristae. However, once cells were treated with PAA, mitochondria started to swell and the cristae disappeared, but no remarkable chromatin condensation was identified (Fig. 3A, 60 min). In a later stage, chromatin condensation was seen in almost all cells, while mitochondrial membranes disappeared and most of cellular organelles were degraded (Fig. 3A, 120 min), consistent with apoptosis. Apoptosis can be induced by extrinsic stimuli through membrane death receptors or by intrinsic stimuli through mitochondrial signaling pathways (Hengartner, 2000; Kurokawa and Kornbluth, 2009). Our results further indicated that PAA induced mitochondria-mediated apoptosis with marked increase in caspases 3, 8, and 9 activity evidenced by increased cleavage of caspases (Fig. 3B). All three caspases were cleaved after 60 min post-PAA treatment. Consistently, PAA-induced MM cell death was partially blocked by adding a pan-caspase inhibitor Q-VD-OPh (p < 0.05; Supplementary Fig. 6). However, because necrosis was seen at earlier time points after PAA treatment we also hypothesized that extrinsic stimuli might be involved in PAA-mediated cell death and we tested the activation of receptor interacting protein kinase 1 and 3 (RIP1 and RIP3) (Nugues et al., 2014). Fig. 3B indicated that RIP1 and RIP3 were also cleaved.

3.5. Apoptosis-inducing Factor 1 Plays a Critical Role in Pharmacological Ascorbic Acid-induced Myeloma Cell Death

We subsequently tried to determine the molecular pathway by which PAA induced mitochondria-mediated apoptosis. Our hypothesis was that increased mitochondrial permeabilization was the trigger for the death signal transduction machinery. We focused our attention on apoptosis-inducing factor 1 (AIF1), because AIF1 induces cell death in caspase-dependent and caspase-independent manners (Nikolopoulou et al., 2013). We evaluate if PAA induced MM cell death depends on AIF1 at least partially. We generated OCI-MY5 cells with AIF1 knockdown (shRNA-AIF1) or overexpression (OE-AIF1). The
viability of AIF1-shRNA OCI-MY5 cells (Fig. 4A, top bar graph) was significantly higher than those cells expressing scrambled sequence after PAA treatment (Fig. 4A, top bar graph), while OE-AIF1 OCI-MY5 showed significantly less viability (Fig. 4A, bottom bar graph) than cells transfected with empty vector (EV) when treated with PAA (Fig. 4A, bottom bar graph). It is widely accepted that AIF1 must be cleaved and released from the mitochondria to the cytoplasm and then translocate to the nucleus to induce cell death (Fig. 4B) (Sevrioukova, 2011). We thus examined if PAA induced AIF1 cleavage. OCI-MY5 cells treated with PAA showed an increase in the AIF1 cleaved form by western blotting (Fig. 4C). Melphalan was not able to induce AIF1 cleavage in OCI-MY5 cells (Fig. 4C) probably because melphalan is an alkylating agent and produces a number of DNA adducts including DNA interstrand crosslinks (ICLs) considered to be the critical cytotoxic lesion (Spanswick et al., 2002) in an AIF1 independent mechanism. We hypothesized that the AIF1 cleavage was mediated by PAA reacting with LIP to form ROS. Thus, we incubated OCI-MY5 cells with or without DFO followed by PAA treatment. AIF1 was not cleaved after PAA incubation in OCI-MY5 cells pretreated with DFO confirming the crucial role of LIP in this process (Fig. 4D, white arrow). Earlier studies have shown that DFO is able to deplete LIP (Al-Qenaei et al., 2014). We also tested the level of phosphorylated γ-H2AX, a biomarker for DNA double-strand breaks, after PAA and melphalan treatment, and determined that PAA and high dose of melphalan induced γ-H2AX. However, a lower dose of melphalan with PAA was also able to induce γ-H2AX (Fig. 4C). These data support our earlier in vivo data (Fig. 1F) that combination of PAA and melphalan at lower dose inhibits tumor formation at the same level or greater than melphalan alone. Cellular localization of AIF1 was examined by immunolabeling electron microscope with and without PAA treatment in OCI-MY5 cells. This staining revealed that AIF1 localizes not only in the mitochondria, as seen in untreated cells (Fig. 4E, left panel), but also in cytoplasm and nuclei in PAA-treated OCI-MY5 cells (Fig. 4E, right panel). These results indicate that PAA by reacting with LIP and generating ROS induces mitochondria-mediated apoptosis in which AIF1 cleavage is important for cell death.

4. Discussion

High-dose vitamin C has been studied in multiple cancers and has shown controversial clinical effects (Cameron and Pauling, 1978, 1976; Creagan et al., 1979; Moertel et al., 1985). The contradictory clinical results can be at least partially explained by different routes of vitamin C administration applied, i.e., either orally or intravenously. Recent reports indicate that a certain ROS concentration is required for high-dose vitamin C to induce cytotoxicity in cancer cells. The generation of ascorbyl- and H2O2 radicals by PAA increases ROS stress in cancer cells (Du et al., 2012). These studies including preclinical and clinical were performed in solid tumors, such as glioblastoma (Herst et al., 2012), pancreatic cancer (Du et al., 2015), ovarian cancer (Ma et al., 2014), prostate cancer (Chen et al., 2012; Pollard et al., 2010), hepatoma (Verrax and Calderon, 2009), colon cancer (Pires et al., 2016), mesothelioma (Ranzato et al., 2011), breast cancer (Yun et al., 2015), bladder cancer (Gilloteaux et al., 2010), and neuroblastoma (Deubzer et al., 2011).
Fig. 4. Pharmacologic ascorbic acid induces AIF1 release from mitochondria. (A) Top bar graph represents OCI-MYS-Scr and -shRNA-AIF1 cells incubated with doxycycline to knockdown AIF1 for 2 days. Bottom bar graph represents EV OCI-MYS and OE-AIF1 OCI-MYS cells. All cells were treated with or without PAA at the specified concentrations. After 1 h treatment, PAA was washed away and cell viability was measured after 24 h. Knockdown and overexpression of AIF1 was confirmed by western blots. (B) Schematic representation of DFO inducing AIF1 cleavage, release and nuclear translocation in MM tumor cells. (C) OCI-MYS WT cells with or without PAA. After 1 h PAA was washed away and cells were incubated with melphalan (Mel, 0–80 μM) for 4 h then lysed. AIF1, β-actin and γ-H2AX levels were analyzed by western blots. (D) OCI-MYS WT cells were incubated with or without DFO (200 μM) for 3 h followed PAA (2 mM) treatment. After 1 h PAA was washed away and cells were lysed. AIF1 and β-actin levels were analyzed by western blots. White arrow represent AIF1 cleaved form. (E) Electron microscopy shows AIF1 immunolabeling stain of OCI-MYS WT cells treated without (left) or with (right) PAA (2 mM). N, M, C respectively represent nucleus, mitochondria and cytoplasm. Blue arrows indicate the nuclear membrane and red arrowheads indicate AIF1 gold beads in cytoplasm or mitochondria. Black arrowheads indicate AIF1 gold beads in nuclei.
treatment. Furthermore, AIF1 was originally discovered as an intermembrane space (IMS) component of mitochondria and characterized as a pro-apoptotic gene (Susin et al., 1999; Joza et al., 2001). Therefore, we focused on AIF1 as one of the pathways by which PAA-induced MM cell death. The pro-apoptotic AIF1 or truncated AIF1 (tAIF) is cleaved from the full-length AIF1 by calpains and/or cathepsins after a caspase-independent cell death insult (Joza et al., 2009; Modjtedhiedi et al., 2006; Sevrioukova, 2011; Artus et al., 2010). tAIF moves from the mitochondria to the cytosol and nucleus, where it initiates chromatosomely and caspase-dependent and caspase-independent cell death (Nikoletopoulou et al., 2013; Artus et al., 2010). Our data show that PAA increases AIF1 cleavage and translocation from mitochondria to cytoplasm and nucleus. Overexpression of AIF1 in MM cells increases while knock-down of AIF1 prevents PAA-induced MM cell death, indicating that AIF1 plays a critical role in mediating PAA-induced MM cell death. Because the mitochondrial apoptotic factors such as cytochrome c and Bcl-2 family proteins are also important for the activation of caspases, future work will have to determine if AIF1 is the major pathway related to PAA activity in cancer cells as well as the exact relationship with other mitochondrial apoptogenetic factors. In addition, the necrosis and apoptosis makers, such as RIP1/3 and caspases 3/8/9, are cleaved after PAA administration. It is therefore possible that PAA activates caspase 8 resulting in RIP1 cleavage and necrosis (Rajput et al., 2011) evidenced by strong caspase 8 cleavage after a short-term treatment with PAA.

High oxidative stress and DNA damage activity are increased, while the anti-oxidant enzyme levels are decreased in MM patients (Mehdi et al., 2012). Several free radical drugs, such as As2O3 and ascorbic acid, have been used to treat MM, in which As2O3 generates ROS while ascorbic acid serves as an anti-oxidant agent. In MM preclinical and clinical studies, ascorbate was used as an adjunct drug and showed controversial results (Harvey et al., 2011; Perrone et al., 2009; Held et al., 2013; Sharma et al., 2012; Nakano et al., 2011; Takahashi, 2010; Sharma et al., 2009; Qazilbash et al., 2008). However, none of these tests used pharmacological doses of ascorbate and intravenous administration. It has been reported that ascorbate directly inactivates bortezomib activity by forming a tight but reversible complex through its vicinal diol moiety. In a pilot study, ascorbate was used as an adjunct drug and showed controversial results (Harvey et al., 2011; Perrone et al., 2009; Held et al., 2013; Sharma et al., 2012; Nakano et al., 2011; Takahashi, 2010; Sharma et al., 2009; Qazilbash et al., 2008). However, none of these tests used pharmacological doses of ascorbate and intravenous administration. It has been reported that ascorbate directly inactivates bortezomib activity by forming a tight but irreversible complex through its vicinal diol group (Perrone et al., 2009; Harvey et al., 2009). This dose of ascorbate in the combination with bortezomib is at a physiological level which has anti-oxidant effect. We did not perform the combination of PAA with bortezomib due to a possible chemical reaction described above. However, our pilot study also suggested that PAA could overcome drug resistance to bortezomib in MM cells (data not shown).

Our findings complement reported studies and further address the mechanism of action using clinical samples in which we observed that PAA killed tumor cells with high iron content, suggesting that iron might be the initiator of PAA cytotoxicity. In addition, combination of PAA with standard therapeutic drugs, such as melphalan, may significantly reduce the dose of melphalan needed. This is beneficial because high doses of melphalan are very toxic not only to tumor cells but also to normal tissues, such as hematopoietic stem cell and epithelial cells in the gut (Shaw et al., 2014; Bayraktar et al., 2013). The efficacy of high-dose melphalan by itself is clearly dose-dependent. Combined treatment of reduced dose melphalan with PAA achieved a significantly longer progression-free survival than the same dose of melphalan alone. These data also suggest that the bone marrow suppression induced by high-dose melphalan can be ameliorated by the combination of PAA with lower dose of melphalan because of the lack of toxicity of PAA on normal cells with low iron content. It is important to consider for future clinical studies that renal insufficiency could be a contraindication for usage of PAA in MM patients. Renal impairment is a common complication of MM (50%) and up to 5% require dialysis (Yadav et al., 2016a, b). The clinical toxicity is probably from oxalate, an end-product of ascorbate metabolism. Therefore, if creatinine clearance is <30 mL/min, high dose ascorbic acid should be not administered.

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Conflict of Interest

Authors declare no conflict of interest.

Author Contribution

J.X., H.X and X.Z performed experiments and analyzed the data; C.A. performed, analyzed and reviewed the electron microscopy experiments; K.L.C. and G.T. collected patients’ samples and clinical data and edited the manuscript; R.N. contributed in the electron microscopy experimental design; I.F. and F.Z. reviewed the data, wrote and edited the manuscript. All authors approved the manuscript.

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Appendix A. Supplementary Data

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