Histone deacetylase inhibitor panobinostat induces calcineurin degradation in multiple myeloma

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Introduction

Multiple myeloma (MM) is a relapsed and refractory disease, one that highlights the need for developing new molecular therapies for overcoming of drug resistance. Addition of panobinostat, a histone deacetylase (HDAC) inhibitor, to bortezomib and dexamethasone improved progression-free survival (PFS) in relapsed and refractory MM patients. Here, we demonstrate how calcineurin, when inhibited by immunosuppressive drugs like FK506, is involved in myeloma cell growth and targeted by panobinostat. mRNA expression of PPP3CA, a catalytic subunit of calcineurin, was high in advanced patients. Panobinostat degraded PPP3CA, a degradation that should have been induced by inhibition of the chaperone function of heat shock protein 90 (HSP90). Cotreatment with HDAC inhibitors and FK506 led to an enhanced ant.myeloma effect with a greater PPP3CA reduction compared with HDAC inhibitors alone both in vitro and in vivo. In addition, this combination treatment efficiently blocked osteoclast formation, which results in osteolytic lesions. The poor response and short PFS duration observed in the bortezomib-containing therapies of patients with high PPP3CA suggested its relevance to bortezomib resistance. Moreover, bortezomib and HDAC inhibitors synergistically suppressed MM cell viability through PPP3CA inhibition. Our findings underscore the usefulness of calcineurin-targeted therapy in MM patients, including patients who are resistant to bortezomib.
promotes proliferation and cell survival in diffuse large B cell lymphoma–derived cell lines (13, 14). When we investigated whether calcineurin activation would affect MM cell survival, we found that PPP3CA (protein phosphatase 3, catalytic subunit, α isozyme), its catalytic subunit, is involved in myeloma cell growth.

Heat shock protein 90 (HSP90) functions as a chaperone stabilizing its client proteins and, as a non-histone protein, is one of the targets of HDAC inhibition (15). The chaperone function is inhibited by the acetylation induced by HDAC inhibitors (16). Panobinostat induces hyperacetylation of HSP90 in acute myeloid leukemia cells and inhibits its chaperone function, thereby leading to the proteasomal degradation of client proteins such as CXCR4 and AML1/ETO9a, both of which are involved in leukemogenesis (17, 18). These results would suggest that the ability of HDAC inhibitors to block the chaperone function of HSP90 might be important for facilitating their anticancer effects. We have shown that HDAC inhibition leads to the protein degradation of PPP3CA. Furthermore, we have demonstrated that PPP3CA was the common target of bortezomib and HDAC inhibitors and that aberrantly enhanced PPP3CA expression promoted bortezomib resistance. Our study points to the new role of calcineurin in the pathogenesis of MM and thus supports the possibility of developing novel strategies to target PPP3CA in MM patients.

**Results**

**Aberrantly increased PPP3CA expression observed in MM cell lines and MM cells isolated from patients with advanced disease.** To study the potentially pathogenic role of PPP3CA in MM, we examined the expression levels of PPP3CA in MM patients. We compared PPP3CA expression of MM cells isolated from patients suffering different stages of MM. A previous study involved high-resolution genomic and mRNA expression profiling of MM cells, thereby revealing candidate oncogenes for MM (19). We examined the expression of 179 candidate genes in MM patients previously reported using the Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) database (20), comparing it among different stages of patients. The disease stages of each patient were determined using the Durie-Salmon staging classification system (21). Those genes highly expressed in most advanced stage III cases (21) are likely to be involved in MM progression. Among the candidate genes examined, PPP3CA was one of 8 genes whose mRNA expression was significantly higher in samples from stage III patients compared with those from stage I patients (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/jci.insight.85061DS1).

We analyzed PPP3CA expression in non-MM and MM cell lines and found high PPP3CA expression only in the latter (Figure 1A). We then examined our MM patient samples (Supplemental Table 1). We found higher expression of PPP3CA in stage III compared with stage I (Figure 1B). Among the clinical
parameters, increased serum lactate dehydrogenase (LDH) was significantly associated with poor overall survival and progression-free survival (PFS) rates (22). PPP3CA expression was significantly higher in patients with increased serum LDH levels than those with normal LDH (Figure 1C). These correlations could support the contention that high PPP3CA expression levels might be involved in the pathogenesis of MM. An important role in cell adhesion–mediated drug resistance (CAM-DR) to bortezomib and conventional chemotherapy in MM is played by \( \alpha_4 \) integrins (23). Expression of \( \alpha_4 \) integrins tended to be high in MM patients with elevated PPP3CA expression, and the possible relevance of high PPP3CA expression to drug resistance was also supposed (Figure 1D).

Panobinostat induces protein degradation of PPP3CA through HSP90 inhibition. Next, we investigated the effects of panobinostat on PPP3CA protein expression in MM cell lines and found it was reduced in panobinostat-treated cells. The degree of protein expression change estimated by quantitative analyses of bands is displayed as indicated. (A) U266 and KMS-11 were treated with 20 nM panobinostat for 48 h followed by Western blot analysis. Actin served as a loading control. Nine (U266) and 3 (KMS-11) biologically independent experiments were performed. To determine the expression of PPP3CA mRNA in treated cells for 24 h, we performed relative quantification real-time PCR (n = 6). Four (U266) and 2 (KMS-11) biologically independent experiments were performed. (B) PPP3CA expression in U266 treated with DMSO (vehicle control) or 600 nM 17-AAG for 24 h. Three biologically independent experiments were performed. (C) IP assays using IgG2b or anti-PPP3CA antibody in U266. TCL, total cell lysate. Five biologically independent experiments were performed. (D) Protein expression in U266 treated with 20 nM panobinostat, 5 nM lactacystin, or both panobinostat and lactacystin for 48 h. Two biologically independent experiments were performed. (E) Protein expression in U266 treated with 20 nM panobinostat, 2 nM romidepsin, or 2 µM ACY-1215 for 48 h. Two biologically independent experiments were performed. (F) IP assays using anti-PPP3CA antibody in U266 treated with 2 µM ACY-1215 for 16 h. Two biologically independent experiments were performed. (G) Protein expression in U266 treated with 2 µM ACY-1215, 5 nM lactacystin, or both ACY-1215 and lactacystin for 48 h. Two biologically independent experiments were performed. (H) HSP70 protein expression in U266 treated with 600 nM 17-AAG for 24 hours. Two technically independent experiments were performed. (I) HSP70 expression in U266 treated with 20 nM panobinostat, 10 µM FK506, or both panobinostat and FK506 as indicated for 36 h. Cotreatment with 0.75 µM ACY-1215 and 10 µM FK506 for 48 h was also performed. Two biologically independent experiments were performed.

Figure 2. PPP3CA protein is degraded by HDAC (histone deacetylase) inhibitors in MM cells. The degree of protein expression change estimated by quantitative analyses of bands is displayed as indicated. (A) U266 and KMS-11 were treated with 20 nM panobinostat for 48 h followed by Western blot analysis. Actin served as a loading control. Nine (U266) and 3 (KMS-11) biologically independent experiments were performed. To determine the expression of PPP3CA mRNA in treated cells for 24 h, we performed relative quantification real-time PCR (n = 6). Four (U266) and 2 (KMS-11) biologically independent experiments were performed. (B) PPP3CA expression in U266 treated with DMSO (vehicle control) or 600 nM 17-AAG for 24 h. Three biologically independent experiments were performed. (C) IP assays using IgG2b or anti-PPP3CA antibody in U266. TCL, total cell lysate. Five biologically independent experiments were performed. (D) Protein expression in U266 treated with 20 nM panobinostat, 5 nM lactacystin, or both panobinostat and lactacystin for 48 h. Two biologically independent experiments were performed. (E) Protein expression in U266 treated with 20 nM panobinostat, 2 nM romidepsin, or 2 µM ACY-1215 for 48 h. Two biologically independent experiments were performed. (F) IP assays using anti-PPP3CA antibody in U266 treated with 2 µM ACY-1215 for 16 h. Two biologically independent experiments were performed. (G) Protein expression in U266 treated with 2 µM ACY-1215, 5 nM lactacystin, or both ACY-1215 and lactacystin for 48 h. Two biologically independent experiments were performed. (H) HSP70 protein expression in U266 treated with 600 nM 17-AAG for 24 hours. Two technically independent experiments were performed. (I) HSP70 expression in U266 treated with 20 nM panobinostat, 10 µM FK506, or both panobinostat and FK506 as indicated for 36 h. Cotreatment with 0.75 µM ACY-1215 and 10 µM FK506 for 48 h was also performed. Two biologically independent experiments were performed.
Figure 3. Calcineurin is indispensable for the maintenance of MM cell growth. The degree of protein expression change estimated by quantitative analyses of bands is displayed as indicated. (A) U266 was lentivirally transduced with control vector (sh-cont) or 3 different shRNAs against PPP3CA (KD #1, #2, and #3). Cell growth was determined by MTT assays (n = 5), and PPP3CA expression levels are displayed. Two biologically independent experiments were performed. (B) KMS-11 was lentivirally transduced with control vector or FLAG-PPP3CA (clone #1, #2, and #3). Cell growth (n = 5), and PPP3CA expression are displayed. Four (cell growth) and 2 (PPP3CA expression) biologically independent experiments were performed. (C) U266 was treated with 20 nM panobinostat, 10 µM FK506, or both panobinostat and FK506 as indicated for 36 h. Five biologically independent experiments were performed. (D) MTT assays in U266 and CD20-positive cells treated with 15 nM panobinostat, FK506, or both panobinostat and FK506 as indicated for 36 h (n = 5). Three biologically independent experiments were performed. (E) NOD/SCID (NOD/Shi/Jic-scid Jcl) mice bearing U266 cells were treated with vehicle (n = 6), panobinostat (n = 9), FK506 (n = 6), or both panobinostat and FK506 (n = 11). Treatment was continued for 15 days. Average of the ratio of the tumor volume on days 8 and 15 to that on day 1 is displayed for each condition. Error bars represent the standard errors. Difference between panobinostat (+)/FK506 (–) and panobinostat (+)/FK506 (+) on day 15 was analyzed using Scheffe test. *, significant. (F) The representative images of tumors of each group. (G) Protein expression in tumors of xenograft of mouse model. #1 and #2: vehicle treated. #3 and #4: panobinostat treated. Two biologically independent experiments were performed. (H) The representative immunohistochemical stainings by anti-cleaved caspase-3 are displayed (×40). The brown cells are positive for staining. Scale bar: 50 microns. Six biologically independent experiments were performed.
binostat-treated cells compared with mock-treated cells (Figure 2A and Supplemental Figure 2A). However, there was no reduction in PPP3CA mRNA expression (Figure 2A). Calcineurin is a client protein of HSP90, and PPP3CA expression was reduced by the HSP90 inhibitor 17-AAG (Figure 2B and refs. 24, 25). We confirmed the interaction between HSP90 and PPP3CA (Figure 2C). Lactacystin, a proteasome inhibitor, rescued the reduction of PPP3CA induced by panobinostat, thereby supporting the possibility that PPP3CA expression was reduced through protein degradation by panobinostat (Figure 2D).

Inhibition of HDAC6, which primarily deacetylates HSP90, leads to the hyperacetylation of HSP90 and the subsequent disruption of its chaperone function (15, 26, 27). Treatment with ACY-1215, a selective HDAC6 inhibitor (8), resulted in a stronger reduction in PPP3CA levels than did treatment with panobinostat or romidepsin — an inhibitor of HDAC1, -2, and -3 — though each agent alone induced similar levels of histone H3 acetylation (Figure 2E). It was shown that PPP3CA reduction by romidepsin occurred mainly through its transcriptional repression (Supplemental Figure 2B). We found that ACY-1215 treatment reduced the association between PPP3CA and HSP90 (Figure 2F). These results suggest that the inhibition of HDAC6 induces PPP3CA degradation by disrupting HSP90 chaperone activity. Lactacystin rescued the reduction of PPP3CA induced by ACY-1215 (Figure 2G).

We showed that part of the antimaloma effect achieved by HDAC inhibitors is mediated by the inhibition of HSP90. Indeed, HSP90 inhibition, in combination with bortezomib, represents a promising strategy for treating MM (28). However, HSP90 inhibitors, including 17-AAG, are known to activate the heat shock transcription factor 1 (HSF1). HSF1-dependent transcriptional induction of HSP70 protects cancer cells from apoptosis (29). This inhibition of apoptosis is thought to reduce the efficacy of the antitumor effects conferred by HSP90 inhibitors. We confirmed that 17-AAG enhanced protein expression of HSP70 (Figure 2H). In contrast, only a small increase was seen in the expression of HSP70 in samples treated with HADC inhibitors alone or in combination with FK506 (Figure 2I). These results support the idea that calcineurin-targeting therapy might be more accurately compared with conventional HSP90 inhibitors, which can induce HSP70 and thereby dampen any antitumor effects.

Inhibition of calcineurin reduces the growth of MM cells in vitro and in vivo. To evaluate the role of PPP3CA in MM cell viability, we knocked down PPP3CA in MM cells and found that their growth was suppressed in a PPP3CA expression level–dependent manner (Figure 3A). In contrast, overexpression of PPP3CA enhanced the growth of MM cells (Figure 3B), thereby suggesting that PPP3CA expression levels might regulate MM cell growth. The calcineurin complex is composed of PPP3CA and calcineurin B, the regulatory subunit of calcineurin (10). FK506 inhibits the association between PPP3CA and calcineurin B (30, 31). To investigate the role of calcineurin B in PPP3CA expression, we examined PPP3CA protein expression in MM cells simultaneously treated with HDAC inhibitors and FK506. PPP3CA expression was more strongly reduced by a combination of panobinostat or ACY-1215 with FK506 compared with HDAC inhibitor alone (Figure 3C and Supplemental Figure 3A). FK506 alone had no suppressive effect on the growth of MM cells. By contrast, FK506 — when used in tandem with panobinostat or ACY-1215 — induced a further reduction in cell growth compared with an HDAC inhibitor alone (Figure 3D and Supplemental Figure 3B). (4;14) is one of the major chromosomal abnormalities linked to poor prognosis (32). We investigated combinatorial effect of panobinostat and FK506 on growth of t(4;14)-positive (KMS-11, KMS-18, and KMS-26) and t(4;14)-negative (U266 and KMS-12PE) cells. Growth of myeloma cells were reduced by panobinostat, and this reduction was enhanced by the addition of FK506 in both t(4;14)-positive and -negative myeloma cell lines (Supplemental Figure 3C and refs. 33–35). Of note, panobinostat did not exert an overt effect on cell growth in normal B cells, and FK506 did not augment its cytotoxic effect (Figure 3D). Next, we studied the pathogenic role of calcineurin in MM in vivo using a xenograft mouse model of myeloma cells. Tumor sizes tended to increase in control and FK506-treated mice. In contrast, it decreased in panobinostat-treated mice. This reduction in tumor size was further enhanced by combined treatment with panobinostat and FK506 for 15 or 22 days (Figure 3, E and F, and Supplemental Figure 4, A–C). Reduced PPP3CA expression, enhanced histone H3 acetylation, and cleavage of caspase-3 in vivo were confirmed in the xenograft samples of panobinostat mice (Figure 3G). Further, the addition of FK506 augmented panobinostat-induced apoptosis in vivo (Figure 3H). Taken together, the addition of FK506 enhances the antimaloma effect of panobinostat through PPP3CA reduction, thereby supporting the importance of calcineurin in the pathogenesis of MM.

The effects of calcineurin inhibition on apoptosis and proliferation of MM cells and the identification of functional targets of PPP3CA in MM. We studied whether induction of apoptosis and/or suppression of MM cell
Figure 4. Calcineurin inhibition enhances apoptosis and blocks proliferation, and PPP3CA regulates NF-κB signaling in MM cells. (A) Apoptotic protein expression of KMS-11 treated with 20 nM panobinostat, 10 μM FK506, or both panobinostat and FK506 for 36 h is displayed. Two biologically independent experiments were performed. (B) KMS-11 was treated with 20 nM panobinostat, 10 μM FK506, or both panobinostat and FK506 for 36 h. Induced apoptosis was evaluated by flow cytometry. Annexin V (+) and propidium iodide (PI) (-): early apoptotic cells. Annexin V (+) PI (+): late apoptotic cells. Two biologically independent experiments were performed. (C) BrdU assays in KMS-11 treated with 15 nM panobinostat, FK506, or both panobinostat and FK506 as indicated for 48 h (n = 6). Two biologically independent experiments were performed. (D) Cell growth and PPP3CA expression in U266 treated with 20 nM panobinostat, 10 μM cyclosporine A, or both panobinostat and cyclosporine A as indicated for 48 h (n = 5). Two biologically independent experiments were performed. (E) NFATc1 (nuclear factor of activated T cells, cytoplasmic, calcineurin-dependent 1) protein in U266 treated with 20 nM panobinostat, FK506, or both panobinostat and FK506 as indicated for 72 h. TATA-binding protein (TBP) served as a loading control. Four biologically independent experiments were performed. (F and G) Cytoplasmic (C) and nuclear (N) NF-κB p50 in U266 lentivirally transduced with control vector (sh-cont) or shRNA against PPP3CA (sh-PPP3CA) (F) and control vector (vector) or PPP3CA (FLAG-PPP3CA) (G). α-tubulin and TBP served as a loading control. N/C ratios of NF-κB p50 are also displayed. Two biologically independent experiments were performed. (H) C and N NF-κB p50 in U266 treated with DMSO or 10 μM FK506 for 48 h followed by Western blot analysis. Two technically independent experiments were performed.
proliferation would contribute to the anti-MM effects observed in the simultaneous treatment with HDAC inhibitors and FK506. Combined treatment with panobinostat and FK506 enhanced cleavage of caspase-7 and -3 while increasing the BCL2-family protein BimS expression compared with panobinostat alone (Figure 4A). The previous study showed that Bim gene KO mouse embryo fibroblasts displayed diminished HDAC inhibitor–mediated apoptosis (36). These results suggest the importance of enhanced expression of Bim in HDAC inhibitor–mediated apoptosis and the possibility of blocking the effect of combined treatment with panobinostat and FK506 by shRNA Bim knockdown. Flow cytometric analysis confirmed that FK506 enhanced MM apoptosis induced by panobinostat or ACY-1215 (Figure 4B and Supplemental Figure 5). In addition, the proliferation of MM cells was inhibited by panobinostat, an inhibition that was enhanced by the addition of FK506 (Figure 4C).
A, another calcineurin B inhibitor, also increased the reduction of cell viability and PPP3CA expression induced by panobinostat (Figure 4D).

The NFATc1 is one of the PPP3CA substrates. The NFATc1 dephosphorylated by PPP3CA translocates from the cytoplasm to the nucleus, and its nuclear translocation is blocked by FK506 (37). We found that combined treatment with panobinostat and FK506 not only substantially suppressed nuclear NFATc1 expression, but also blocked calcineurin signaling (Figure 4E). FK506 alone suppressed nuclear NFATc1 expression. However, the viability of MM cells was not affected by FK506. Moreover, dephosphorylation of NFATc1 by PPP3CA does not appear to be related to those signals that promote MM survival (Figure 3D).

In addition to mediating NFATc1 activation, PPP3CA was shown to synergize with protein kinase C (PKC) in order to activate NF-κB in T cells (38). The latter is involved in the pathogenesis of MM (39). Panobinostat disrupts formation of the NF-κB/DNA binding complex, thereby inactivating NF-κB signaling (40). This inactivation may be due to downregulation of PPP3CA by panobinostat. In fact, PPP3CA knockdown reduced the nuclear/cytoplasmic ratio of NF-κB p50 subunit expression in MM cells (Figure 4F). Furthermore, its overexpression increased that ratio (Figure 4G). These results suggest that PPP3CA plays an important role in NF-κB signal activation in MM cells. Thus, it is possible that panobinostat reduces MM cell viability by suppressing PPP3CA–NF-κB signaling. On the other hand, FK506 did not affect the expression of the cytoplasmic or nuclear NF-κB p50 subunit in MM cells (Figure 4H).

**PPP3CA is a common target of panobinostat and bortezomib.** Next, we examined whether the PPP3CA reduction induced by panobinostat is relevant to the clinical effect that panobinostat adds to bortezomib therapy (1, 2). We examined a correlation between **PPP3CA** expression in CD138-positive cells and the clinical responses to bortezomib-containing therapies in newly diagnosed patients treated with VMP (bortezomib, melphalan, and prednisone regimen) or BD (bortezomib plus dexamethasone therapy) at 2 centers from 2012–2014 (Supplemental Table 1). **PPP3CA** expression was significantly higher in bortezomib-resistant patients versus those who were bortezomib sensitive (Figure 5A). Furthermore, PFS duration in patients with high **PPP3CA** expression was shorter than that of patients with low **PPP3CA** expression in bortezomib-containing therapy (Figure 5B). **PPP3CA** knockdown by lentiviral shRNA (Figure 5C, left) enhanced the sensitivity to bortezomib compared with controls (Figure 5C, right). Thus, **PPP3CA** expression levels correlated with bortezomib resistance. Bortezomib suppressed not only mRNA and protein expression of HDAC6 in a dose-dependent manner, but also **PPP3CA** protein expression (Figure 5D). Induction of protein acetylation by bortezomib was confirmed by hyperacetylation of histone H3 in MM cells treated with bortezomib (Figure 5D). These results indicate that bortezomib may suppress **PPP3CA** through HDAC6 inhibition. In fact, **PPP3CA** was transcriptionally suppressed by bortezomib (Figure 5D). Furthermore, cotreatment with panobinostat and bortezomib additively suppressed cell viability (Figure 5E). These results suggest that panobinostat might augment the antmyeloma effect of bortezomib by further reducing **PPP3CA** expression. Additive reduction of MM cell viability and **PPP3CA** expression was also observed in another combination treatment paring panobinostat with carfilzomib, another proteasome inhibitor (ref. 41 and Figure 5F). Suppression of cell viability and **PPP3CA** were enhanced by cotreatment with ACY-1215 and bortezomib (Supplemental Figure 6A). The antmyeloma effect of bortezomib was enhanced by cotreatment with FK506 (Supplemental Figure 6B). Combinatory treatment with panobinostat and bortezomib reduced cell growth both in vector and **FLAG-PPP3CA**–introduced cells compared with vehicle treatment. However, the growth ratio of **PPP3CA** overexpressed cells to that of control cells was comparable between vehicle and drug-treated cells. Thus, combination strategy did not block the growth enhancement induced by ectopic overexpression of **PPP3CA** (Supplemental Figure 7).

**Formation of osteoclasts is inhibited by panobinostat.** MM generates lytic bone lesions via the formation of osteoclasts (42). Osteoclast differentiation requires the induction of NFATc1 by receptor activator nuclear factor-κ-B ligand (RANKL). As FK506 inhibits osteoclast differentiation (43), we investigated the effect of panobinostat treatment on the latter. Panobinostat and FK506 each inhibited osteoclast differentiation, while the addition of both drugs strengthened the blockade of osteoclast formation compared with treatment with either agent alone (Figure 6A). **PPP3CA** protein levels, but not mRNA expression, decreased in differentiated osteoclasts treated with panobinostat (Figure 6B). NFATc1 activation is required for the induction of essential osteoclast transcription factors, such as microphthalmia-associated transcription factor (MITF) (44). In addition to the NFATc1-MITF pathway, MITF is upregulated by HSP70 (44). Although 17-AAG is a known potent cytotoxic drug, it is associated with side effects that cause bone loss due to increases in HSP70-MITF. In osteoclasts treated with panobinostat, MITF was reduced without any
increase in HSP70 (Figure 6C). These results suggest that treatment of MM with panobinostat would not cause osteoclast activation, as seen in 17-AAG treatment.

Discussion
In this study, we have shown the possible pathogenesis of PPP3CA, a catalytic subunit of calcineurin, in MM. In 4q22.3-4q24, where PPP3CA is located, gains in both copy number and gene amplification have been reported based on analyses of MM patient tumors (19). This suggests that high PPP3CA expression plays a pathogenic role in MM. Analyses of data from previous reports, and from our clinical samples, revealed that PPP3CA expression was higher in the tumors of advanced MM patients (stage III) than in those of nonadvanced patients (stage I). In addition, increased serum LDH in patients with high PPP3CA expression points to its potential relevance in poor prognosis. Furthermore, it has been suggested that PPP3CA maintains MM cell viability through activation of NF-kB signaling. Cotreatment of MM cells with panobinostat and FK506 synergistically reduced PPP3CA and MM cell viability. On the other hand, the cytotoxic effect exerted by panobinostat on CD20+ cells was subtle, while the addition of FK506 did not increase their viability. Moreover, development of T and B lineage cells was normal in PPP3CA-deficient mice, and panobinostat did not compromise donor lymphocyte reconstitution in a mouse BM transplantation model (45, 46). These results suggest that calcineurin-targeting therapy exerts an antimyeloma effect without inducing significant side effects in normal lymphoid systems. The 5TGM1/Kalwrij model of myeloma is an immune-competent model involving bone lesions and is considered suitable for analyses of the in vivo effects of antimyeloma drugs on T cell populations, as well as osteoclasts activity (47). However, treatment with FK506 significantly suppressed cellular immunity in immune competent mice (48). Transfer
B cells deficient in Blimp1 (the tumor suppressor gene) into mice lacking T cells (Rag1−/−) resulted in mutant cells expansion, but this transfer into WT mice did not in the previous study (49). Thus, it is supposed that B cell malignant cells will expand more in T cell–suppressed mice than control mice. These results suggest the possibility of enhanced growth of transplanted 5TGM1 cells by FK506 in a 5TGM1/Kalwrij model of myeloma. On the other hand, in our xenograft immunocompromised mouse model, tumor volume was comparable between panobinostat (−)/FK506(−)–treated and panobinostat (−)/FK506(+)–treated mice (Figure 3E and Supplemental Figure 4A). Furthermore, myeloma cells are responsible for producing osteoclastogenic activity (50), and FK506 is supposed to modify the osteoclast activity through enhanced growth of myeloma cells in the 5TGM1/Kalwrij model of myeloma. For this reason, the 5TGM1/Kalwrij model of myeloma may not be appropriate for analyzing the effects of combined HDAC inhibitors and FK506 on implanted tumors and osteoclasts activity. Recently, the activity of HDACs vis-à-vis nonhistone proteins such as HSP90 has attracted increasing attention as a key aspect of HDAC function (15). We discovered that HDAC inhibitors disturbed PPP3CA through inhibition of HSP90. ACY-1215 is a specific inhibitor of HDAC6, a class IIb HDAC (15). HDAC6 maintains HSP90 activation via its acetylation and prevents degradation of its client proteins. HDAC6-specific inhibition is not associated with disrupted epigenetic control of gene transcription, as has been observed in the usage of pan-HDAC inhibitors such as panobinostat (51). We investigated the effect of ACY-1215 on PPP3CA in order to elucidate the role HDAC6 plays in the protein stability of PPP3CA. Treatment of MM cells by ACY-1215 degraded PPP3CA with a reduction in cell viability. This degradation was mediated by the release of PPP3CA from HSP90. HDAC6-mediated acetylation of HSP90 might play an important role in the maintenance of PPP3CA stability and MM cell viability. HDAC1–3 downregulate their targets through transcriptional repression. Epigenetic recovery of tumor-suppressor genes such as p21 has been linked to the actions of HDAC inhibitors, including panobinostat (15, 52). However, some oncogenes such as CCND2 and Ki-67 are known to be suppressed by HDAC inhibitors (53). The reduction of PPP3CA mRNA by romidepsin suggested that PPP3CA could be the target of both protein degradation and transcriptional repression by HDAC inhibitors. Although FK506 by itself neither affects MM cell viability nor reduces PPP3CA expression, its use in cotreatment with HDAC inhibitors negatively impacts both. Calcineurin B has been shown to protect PPP3CA from protein degradation in in vitro experiments (54). This indicates that the ability of FK506 to additively inhibit any interaction between PPP3CA and calcineurin B promotes the degradation of unstable PPP3CA released from HSP90 by HDAC inhibitors. Thus, FK506 is supposed to enhance the antymeloma effects of HDAC inhibitors. High expression of PPP3CA will be important since PPP3CA mediates myeloma cell growth through PPP3CA–NF-xB signaling and could be the target of those HDAC inhibitors supported by FK506. The authors believe that this cotreatment would be clinically beneficial for a certain group of MM patients, as will be discussed in detail herein. Reduced intensity–conditioned allogeneic stem cell transplantation (RIST) is a tolerable and effective treatment for some MM patients who relapse after autologous stem cell transplantation (55). However, the reduced intensity of conditioning therapy may lead to the presence of residual disease. Although immunosuppressive drugs (e.g., FK506) are used to facilitate allograft survival and prevent graft-versus-host disease, they increase the risk of expanded residual disease and relapse. The enhanced antymeloma effect of combination therapy (panobinostat + immuno-suppressive drugs) may aid panobinostat-maintenance therapy in those patients receiving RIST. Approval of salvage therapy by panobinostat for relapsed and refractory patients indicates the possible effectiveness of such maintenance therapy. The results of our xenograft mouse model experiments have shown FK506's additive effect on the antymeloma effect exerted by panobinostat. In these experiments, panobinostat and FK506 were safely administered in vivo without a significant loss in body weight. The minimum dosage for oral administration of FK506 necessary to suppress immunity in vivo was 10 mg/kg body weight for 4 days (total 40 mg/kg for a week) in a previous study (48). In a clinical setting, a noncontinuous dosing schedule of panobinostat (e.g., 3 times per week) is necessary to reduce the risk of thrombocytopenia (52). To enhance these antymeloma effects, concurrent administration of panobinostat and FK506 should prove an effective therapy. Thus, 10 mg/kg of oral administration of FK506 (total 30 mg/kg for a week) is believed to be an appropriate and efficient treatment. Tumor volume differences between panobinostat (+)/FK506 (−)–treated and panobinostat (+)/FK506 (+)–treated mice were significant, as determined by Scheffe test analyses, both on days 8 and 15 (treatment for 15 days) and on days 8, 15, and 22 (treatment for 22 days) (Supplemental Figure 4, B and C). The ratio of tumor volume in panobinostat (+)/FK506 (+)–treated mice
versus panobinostat (+)/FK506 (–)-treated mice decreased on day 22 (0.459) compared with that on day 15 (0.623) (Supplemental Figure 4C). These results indicate that combinatory therapy for 22 days is more effective than that for 15 days. Thus, we believe that treatment should be carried out for at least 22 days.

Bortezomib plays a central role as a drug in MM therapies, both in transplantation-eligible and -ineligible patients. However, many patients develop a resistance to bortezomib during therapy, thereby impeding their long-term survival. Such resistance must be overcome if MM patients are to enjoy improved survival rates. Panobinostat, an FDA-approved drug, when used in combination with bortezomib is expected to overcome this resistance. Proteasome inhibition by bortezomib leads to the formation of aggresomes, which are dependent upon the interaction of HDAC6 with tubulin and the dynein complex (5). HDAC6 inhibition induces hyperacetylation of tubulin and blocks aggresome formation, which is accompanied by increased cellular stress and apoptosis (3). These results indicate that panobinostat enhances the antymyeloma effect of bortezomib by inhibiting aggresome formation. We found that high \( \text{PPP3CA} \) expression is indicative of a poor prognosis due to resistance to bortezomib. Thus, we believe that \( \text{PPP3CA} \) could serve as a target of therapy in bortezomib-resistant patients. In addition, we examined the expression of \( \text{PPP3CA} \) and several genes associated with bortezomib resistance (\( \text{PSMB5}, \text{CDK5}, \text{and} \ \text{CYP1A1} \)) in patients treated with bortezomib as well as dexamethasone in the millennium trial (56–59). We compared the expression of each gene between responsive (R) (minor response and better) and nonresponsive (NR) (no change and progressive disease) patients. There was no significant difference of expression in \( \text{CDK5} \) and \( \text{CYP1A1} \) between R and NR. However, \( \text{PSMB5} \) expression was substantially higher in NR compared with R, and a trend of higher expression of \( \text{PPP3CA} \) in NR compared with R was evident (Supplemental Table 2). The relevance of high \( \text{PPP3CA} \) expression vis-à-vis \( \text{PPP3CA} \) expression should be investigated by further analysis in a large cohort treated with a unified bortezomib-containing protocol. We have shown that bortezomib and panobinostat synergistically reduce MM cell viability through \( \text{PPP3CA} \) reduction. The antymyeloma effects of this combined therapy mediated inhibited aggresome formation and synergistic \( \text{PPP3CA} \) reduction. This is consistent with the clinical findings about the additive effect panobinostat provides to bortezomib and dexamethasone therapy (1, 2). \( \alpha_4 \) integrins are associated with CAM-DR, and the correlation between \( \text{PPP3CA} \) and \( \alpha_4 \) integrins expression in CD138-positive cells from MM patients was found. The reason for poor responses to VMP or BD treatments and short PFS durations in patients with high \( \text{PPP3CA} \) expression may be explained by the pathogenicity of high \( \text{PPP3CA} \) expression, as well as CAM-DR, resulting from high \( \alpha_4 \) integrin expression.

Bortezomib has been shown to not only exert a tumoricidal activity, but also to improve lytic bone lesions in MM patients. Moreover, it has been shown to suppress the formation of osteoclasts by blocking the NFATc1 induction caused by tumor necrosis factor receptor–associated factor 6 (TRAF6) (60). NFATc1

Figure 7. Proposed molecular mechanism of the clinical effects of calcineurin-targeting therapy in MM patients.

Schematic representation of the reduction in \( \text{PPP3CA} \) induced by panobinostat and bortezomib and the expected clinical effects of \( \text{PPP3CA} \) reduction in MM patients are displayed. CaM, calmodulin; CnB, calcineurin B; Ac, acetylation.
induction by calcineurin signaling also plays an important role in osteoclast formation (43). In this study, we have demonstrated that panobinostat disturbed osteoclast formation and suppressed PPP3CA. This finding is in accordance with a previous study in which panobinostat reduced bone density loss in a disseminated MM xenograft mouse model (61). Furthermore, addition of FK506 to panobinostat enhanced the blocking of osteoclast formation compared with treatments with either agent alone. This finding indicated that calcineurin-targeted therapy, using a combination of panobinostat and FK506, effectively inhibits lytic bone lesions, as well as MM cell viability. This might serve as a useful tool for halting the cycle induced by MM cells and osteoclasts.

In summary, we have revealed PPP3CA, a catalytic subunit of calcineurin, as a molecule related to both MM cell viability and osteoclast formation (Figure 7). PPP3CA is protected from protein degradation by binding to HSP90, and its degradation was induced by panobinostat and bortezomib through HDAC6 inhibition. Addition of FK506 enhanced PPP3CA reduction induced by panobinostat, and bortezomib has also transcriptionally suppressed PPP3CA. Furthermore, PPP3CA was related to bortezomib resistance. The development of new calcineurin-targeted therapies, which inhibit PPP3CA–NF-κB signaling by small molecules, is expected to profoundly improve the treatment of MM. This will overcome drug resistance and improve osteolytic lesions in a wide range of patients, including those receiving RIST who may be treated with panobinostat and FK506.

Methods

Gene expression analysis of candidate oncogenes in MM patients. mRNA expression data on candidate oncogenes from MM patient samples obtained during a previous study (20) were analyzed using the GEO (http://www.ncbi.nlm.nih.gov/geoprofiles/11574557). Statistical analyses examining the difference of expression of PSMB5, PPP3CA, CDK5, and CYP1A1 between R (complete response [n = 14], partial response [n = 76], and minor response [n = 23]) and NR (no change [n = 60] and progressive disease [n = 66]) patients who were treated with bortezomib (n = 169) or dexamethasone (n = 70) in millennium trial (59) were performed using GEO2R (National Center for Biotechnology Information, http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE9782, GPL96 platform).

Patient samples. CD138-positive cells were sorted from BM samples (Tokyo Women's Medical University and Nippon Medical School). The patient characteristics are described in Supplemental Table 1. Sensitivity to bortezomib-containing therapies was estimated in those patients who were treated with VMP (bortezomib, melphalan, and prednisone) or BD (bortezomib and dexamethasone).

Cell lines. Human MM (U266, KMS-11, KMS-12PE, KMS-26, and RPMI8226) and non-MM (HEL, HL60, KG-1, THP-1, U937, BALL-1, NALM6, Jurkat, Molt-4, K562, Daudi, and Raji) cell lines were cultured in RPMI-1640 medium supplemented with 10 % FCS, penicillin, and streptomycin. HEK293T cells were cultured in DMEM supplemented with 10 % FCS, penicillin, and streptomycin. The source of cell lines is as follows: U266, RPMI8226, HEL, HL60, KG-1, THP-1, U937, Jurkat, Molt-4, K562, Daudi, Raji, and HEK293T (ATCC); KMS-11 and NALM6 (JCRB cell bank, National Institutes of Biomedical Innovation, Health and Nutrition); KMS-18 and KMS-26 (T. Otsuki, Kawasaki Medical School, Okayama, Japan); and BALL-1 (Riken BioResource Center).

Reagents. Panobinostat (LBH589)(SelleckChemicals), romidepsin (SelleckChemicals), ACY-1215 (Chemietek), bortezomib (Selleck Chemicals), FK506 monohydrate (Sigma-Aldrich), cyclosporine A (Sigma-Aldrich), 17-(allyamino)-17-demethoxygeldanamycin (17-AAG) (Sigma-Aldrich), lactacystin (Santa Cruz Biotechnology Inc.), and carfilzomib (Selleck Chemicals) were dissolved in DMSO and added to the culture medium at the indicated concentrations. Panobinostat for in vivo study was kindly provided by Novartis Pharma Inc. (Fabrikstrasse 28-Z1.28.1 Novartis Campus CH-4056). FK506 for in vivo study was provided by Astellas Pharma Inc.

Western blot analyses and immunoprecipitation. Total cell lysates were prepared as follows. Collected cells were washed with PBS and lysed in RIPA buffer (50 mM Tris-HCl [pH 7.4], 150 mM NaCl, 1 % NP-40, 0.1 % SDS, 0.1 % deoxycholate, 1 mM phenylmethylsulfonyl fluoride [PMSF], 2 μg/ml aprotinin, 2 μg/ml leupeptin, 1 μg/ml pepstatin A, and 1 % phosphatase inhibitor cocktail [Sigma-Aldrich]). Cell lysates were incubated with gentle rocking for 1 hour at 4°C and then centrifuged at 12,281 g for 20 minutes. The supernatant was analyzed as the whole cell lysate. Nuclear extracts were prepared as described previously (62). Extraction of nuclear and cytoplasmic protein fractions were performed using a Nuclear/Cytosol Fractionation Kit (BioVision, K266).
Western blot analyses were performed using anti-PPP3CA (Santa Cruz Biotechnology Inc., sc-9070 and sc-17808), anti–acetyl-Histone-H3 (Millipore, 06-599), anti-HDAC6 (Cell Signaling Technology, 7558), anti-NFATc1 (Santa Cruz Biotechnology Inc., sc-7294), anti–caspase-3 (Cell Signaling Technology, 9662), anti–cleaved caspase-3 (Cell Signaling Technology, 9661), anti–caspase-7 (Cell Signaling Technology, 9494), anti–cleaved caspase-7 (Cell Signaling Technology, 9491), anti-Bim (Cell Signaling Technology, 2819), anti–NF-xb p50 (Santa Cruz Biotechnology Inc., sc-7178), anti-MITF (Abcam, ab12039), anti-HSP70 (Enzo Life Sciences, ADI-SPA-812), anti-FLAG (Sigma-Aldrich, F3165), anti-actin (Sigma-Aldrich, A2066), anti–α-tubulin (Covance, MMS-489R), and anti-TATA binding protein (Abcam, ab818) antibodies. Immunoprecipitated proteins were recovered using Protein G-Sepharose 4B Conjugate (Invitrogen). The Luminata Forte HRP Western Substrate (Millipore) was used for chemiluminescent detection, and the images were analyzed with a LAS-1000 (FUJIFILM). Quantitative analyses of bands in analyzed images were performed using Multi Gauge (FUJIFILM). We also quantitated the degree of protein expression change from the blots in Figures 2 and 3. For the immunoprecipitation assays, total cell lysates were prepared as described previously (63), and anti-PPP3CA (Santa Cruz Biotechnology Inc., sc-17808), anti-HSP90α (Stressgen, ADI-SPA-840), and mouse IgG2b Isotype control (MBL, M077-3) antibodies were used.

RNA extraction, cDNA synthesis, and relative quantification real-time PCR. Total RNA was extracted from patient samples, MM cells, and osteoclasts using TRizol Reagent (Ambion). cDNA was synthesized from total RNA using a High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). cDNA, TaqMan Gene Expression Master Mix (Applied Biosystems), and TaqMan MGB primer-probe sets (Applied Biosystems) for the target and endogenous control gene were mixed and analyzed on an ABI 7500 Real-Time PCR System or a StepOnePlus Real-Time PCR System (Applied Biosystems). The TaqMan MGB primer-probe sets were as follows: human PPP3CA, Hs00174223_ml; human HDAC6, Hs00195869_ml; mouse PPP3CA, Mm01317678_ml; human β-actin, human ACTB (Beta Actin) Endogenous Control (Applied Biosystems); and mouse β-actin, mouse ACTB (Beta Actin) Endogenous Control (Applied Biosystems). Each patient’s mRNA amount of PPP3CA divided by that of U266 cells was designated as relative PPP3CA mRNA.

MTT assays. A total of 2 × 10^4 of the indicated MM cell lines were seeded in a 96-well plate and cultured for 48 hours as indicated. MTT assays were performed using the Cell Counting Kit-8 (DOJINDO) following the manufacturer’s instructions, and the plates were read using a Plate CHAMELEON (Hidex). Five independent clones were examined. CD20-positive cells were sorted using CD20 Micro Beads (Miltenyi, 130-091-104) as described in the manufacturer’s protocol. A total of 10 μg/ml pokeweed mitogen (PWH) (Sigma-Aldrich) was added to the culture medium to induce the proliferation of CD20-positive cells. In MTT assays of shRNA-transduced U266 or FLAG-vector–transduced KMS-11, a total of 1 × 10^4 (PWH) (Sigma-Aldrich) was added to the culture medium to induce the proliferation of CD20-positive cells. In MTT assays of shRNA-transduced U266 or FLAG-vector–transduced KMS-11, a total of 1 × 10^4 (PWH) (Sigma-Aldrich) was added to the culture medium to induce the proliferation of CD20-positive cells.

BrdU assays. A total of 5 × 10^3 KMS-11 cells were seeded in 96-well plates and cultured for 48 hours as indicated. Cell proliferation was analyzed using a Cell Proliferation ELISA, bromodeoxyuridine (BrdU) chemiluminescent Kit (Roche Diagnostics) following the manufacturer’s instructions. Plates were read on a Multi Gauge (FUJIFILM).
Plate CHAMELEON (Hidex), and the results were acquired by relative luminescence units (rlu) per second.

Flow cytometric analysis. To examine apoptosis, KMS-11 cells were stained with the ApoAlert Annexin V-FITC Apoptosis Kit (Clontech) following the manufacturer’s instructions, and cells were analyzed on an EPICS XL System II Flow Cytometer (Beckman Coulter). Data were analyzed using Kaluza 1.3 (Beckman Coulter).

Xenograft mouse model. Five- to 7-week-old NOD/Shi-iscid Jcl (NOD/SCID) female mice (CLEA Japan) were used for the in vivo assays. A total of $3 \times 10^7$ U266 cells in 100 μl of RPMI-1640 were mixed with the same volume of Matrigel (BD Biosciences), and the mixture was s.c. injected into the back of the mouse. Tumors were measured with calipers, and volumes were calculated using the formula $\alpha \times \beta \times 0.5$, where $\alpha$ is the shortest diameter and $\beta$ is the diameter perpendicular to $\alpha$, as described previously (64). When the diameter of the tumor reached 1.0 cm, treatment with vehicle, panobinostat, FK506, or both panobinostat and FK506 was initiated. Panobinostat (provided by Novartis Pharma Inc.) was dissolved in 5 % dextrose (Sigma-Aldrich), and 20 mg/kg panobinostat was i.p. injected into mice 3 times per week for 3 weeks. FK506 (20%; provided by Astellas Pharma. Inc.) was dissolved in water, and 10 mg/kg FK506 was administered orally to mice 3 times per week for 3 weeks. Tumor volume and mice body weight were assessed on alternate days. Mice treated with panobinostat showed body weight loss, and we set 20 mg/kg panobinostat as the maximum tolerated dose, with less than 15% body weight loss. After sacrifice, cleavage of caspase-3, PPP3CA expression, and acetylation of histone H3 in tumors from some mice were examined.

IHC. Tumors from the xenograft mouse model were put in optimal cutting temperature (OCT) compound (Sakura Finetek, 4583) and frozen. The frozen samples were processed as 6-μm sections with cryostat (Leica, CM1850). After fixation and permeabilization, the sections were incubated with anti–cleaved caspase-3 antibody overnight at 4°C centigrade and washed in PBS. The sections were then incubated with anti–rabbit secondary antibody (Vector, MP-7401) for 1 hour at room temperature and washed in PBS. A 3,3′-diaminobenzidine (DAB) reaction was performed using a DAB Peroxidase (HRP) Substrate Kit (Vector, SK-4100) following the manufacture’s instructions. Counterstaining by H&E was also performed.

Induction of mouse BM mononuclear cells into osteoclasts and tartrate-resistant acid phosphatase (TRAP) staining of osteoclasts. Induction of mouse BM mononuclear cells into osteoclasts and TRAP staining of osteoclasts were performed as previously described (65). Mouse BM cells were harvested from the femurs and tibias of female C57 BL/6J mice. They were seeded in 24-well plates and cultured in minimum essential medium supplemented with 10 % FCS, 10 ng/ml human macrophage colony-stimulating factor (R&D Systems), penicillin, and streptomycin for differentiation into macrophages. On days 3, 5, and 6, 50 ng/ml human RANKL (PeproTech) was added to the medium for the induction of macrophages into osteoclasts. We then counted TRAP-positive multinucleated cells (more than 3 nuclei).

Statistics. All data except for those pertaining to the xenograft mouse model are shown as the average ±SD. Differences between the 2 groups of samples were analyzed using a 1-tailed t test. For differences among the 3 groups of samples, ANOVA using a 1-way layout was performed. When the statistic model proved significant, the differences between combinations of the 2 groups were analyzed using a Tukey-Kramer test for multiple comparisons. PFS of the patients treated with bortezomib-containing therapies was defined as the duration from the time to start of therapy to that of disease relapse or progression. A Kaplan-Meier analysis was performed for the study of PFS. A P value less than 0.05 was considered significant. Statistics except for those about xenograft mouse model were analyzed with JMP ver.11 (SAS Institute). The time-dependent variations in tumor volume between panobinostat (+)/FK506(–)-treated and panobinostat (+)/FK506(+)–treated mice in a xenograft mouse model were analyzed with SAS ver. 9.4 (SAS Institute) using the GLM procedure. Tumor size difference between 2 groups on designated days were estimated by Scheffé test analyses.

Study approval. In the analyses of patients’ samples, all patients signed independent informed consent forms for the sampling and molecular and clinical analyses prior to participation in this study. All clinical investigation was conducted according to Declaration of Helsinki principles. This study was approved by the Institutional Ethics Committees of Tokyo Women’s Medical University and Nippon Medical School. All mouse protocols used in this study were approved by the Institutional Animal Care and Use Committee of Tokyo Women’s Medical University.

Author contributions
YI conceived the study, designed and performed the research, collected the patients’ samples, analyzed and...
interpreted data, and wrote and edited the manuscript. EO designed and performed the research, analyzed and interpreted data, and wrote and edited the manuscript. ST and SS performed the experiments inducing mouse BM mononuclear cells into osteoclasts. MI and HT collected the patients’ samples. YHW conducted the experiments of lentiviral production and transduction. AD performed the immunohistochemical experiments. JT, YM, and TM interpreted data, and wrote and edited the manuscript.

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