## LYMPHOID NEOPLASIA

# Immunoregulatory roles of versican proteolysis in the myeloma microenvironment

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#### **Key Points**

- Interplay between myeloma niche stromal cells and myeloid cells generates versikine, a novel damageassociated molecular pattern.
- Versikine may promote antigen-presenting cell maturation and CD8<sup>+</sup> T-cell activation/recruitment to the tumor bed.

Myeloma immunosurveillance remains incompletely understood. We have demonstrated proteolytic processing of the matrix proteoglycan, versican (VCAN), in myeloma tumors. Whereas intact VCAN exerts tolerogenic activities through Toll-like receptor 2 (TLR2) binding, the immunoregulatory consequences of VCAN proteolysis remain unknown. Here we show that human myeloma tumors displaying CD8<sup>+</sup> infiltration/aggregates underwent VCAN proteolysis at a site predicted to generate a glycosaminoglycan-bereft N-terminal fragment, *versikine*. Myeloma-associated macrophages (MAMs), rather than tumor cells, chiefly produced V1-VCAN, the precursor to versikine, whereas stromal cell–derived ADAMTS1 was the most robustly expressed VCAN-degrading protease. Purified versikine induced early expression of inflammatory cytokines interleukin 1 $\beta$  (IL-1 $\beta$ ) and IL-6 by human myeloma marrow-derived MAMs. We show that versikine signals through pathways both dependent and independent of Tpl2 kinase, a key regulator of nuclear factor  $\kappa$ B1-mediated MAPK activation in macrophages. Unlike intact VCAN, versikine-induced II-6 production was partially independent of TIr2. In a model

of macrophage-myeloma cell crosstalk, versikine induced components of "T-cell inflammation," including IRF8-dependent type I interferon transcriptional signatures and T-cell chemoattractant CCL2. Thus the interplay between stromal cells and myeloid cells in the myeloma microenvironment generates versikine, a novel bioactive damage-associated molecular pattern that may facilitate immune sensing of myeloma tumors and modulate the tolerogenic consequences of intact VCAN accumulation. Therapeutic versikine administration may potentiate T-cell-activating immunotherapies. (*Blood.* 2016;128(5):680-685)

## Introduction

Myeloma is a tumor of plasma cells, which are antibody-producing, terminally differentiated B lymphocytes.<sup>1</sup> Myeloma plasma cells typically live within the bone marrow microenvironment ("canonical" myeloma niche) but can often thrive in extramedullary sites and soft tissues ("noncanonical" niches).

We have hypothesized that infiltrating myeloid cells may exert crucial trophic and immunoregulatory functions in both "canonical" and "noncanonical" niches, in part through their regulation of extracellular matrix composition and remodeling.<sup>2</sup> We and others have previously demonstrated that versican (VCAN), a large, chondroitin-sulfate matrix proteoglycan, accumulates in myeloma lesions and have hypothesized that VCAN may contribute to myeloma niche immunoregulation.<sup>3,4</sup> VCAN has crucial, nonredundant significance in embryonic development<sup>5</sup> and emerging roles in cancer inflammation and metastasis.<sup>6-9</sup> VCAN promotes tolerogenic polarization of antigen-presenting cells

through Toll-like receptor 2 (TLR2).<sup>10</sup> VCAN is proteolytically cleaved by ADAMTS-type proteases in a highly regulated manner.<sup>5</sup> A cleavage product generated by proteolysis of the Glu<sup>441</sup>-Ala<sup>442</sup> bond within the VCAN V1 isoform, has been termed versikine.<sup>5</sup> Versikine has been shown to be bioactive (proapoptotic) during interdigital web regression in the mouse embryo<sup>11</sup>; however, the roles of VCAN proteolysis and/or versikine in tumor immunomodulation remain unknown.

## Study design

#### Versikine production, purification, and analysis

Methods for expression and purification of recombinant versikine from mammalian cells have been published.<sup>12,13</sup> We excluded endotoxin and

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Figure 1.

hyaluronan contamination as detailed in the supplemental Methods, available on the *Blood* Web site.

#### Primary bone marrow sample processing and cell culture

Bone marrow aspirates were collected with informed consent under a University of Wisconsin Institutional Review Board–approved protocol (HO07403) and processed as detailed in the supplemental Methods. Cell culture methods are detailed in the supplemental Methods.

#### Animal experiments

Animal experiments were carried out under an Institutional Animal Care and Use Committee–approved protocol (M2395).

#### **RNA** analyses

Cells were lysed in Trizol reagent and RNA extracted per standard methods. Reverse transcriptase polymerase chain reaction (RT-PCR) and RNA sequencing (RNA-seq) were performed as detailed in the supplemental Methods.

#### Immunohistochemistry

The University of Wisconsin myeloma tissue microarray has been previously reported.<sup>3</sup> A second myeloma tissue microarray was purchased from US Biomax (Rockville, MD; catalog no. T291b). Antibodies are listed in the supplemental Methods.

#### Immunoblot, cytokine detection, and cell-cycle analysis

Antibodies and reagents are listed in the supplemental Methods.

#### **Statistical Methods**

All in vitro experiments were performed in triplicate and statistical significance was assessed using 2-tailed Student *t* test or 2-way ANOVA analysis where appropriate. Multiple comparisons were made using the Tukey method. A *P* value <.05 was considered statistically significant.

## **Results and discussion**

To determine whether versikine acts as an endogenous damageassociated molecular pattern (DAMP), we exposed freshly explanted primary myeloma CD14<sup>+</sup> cells to recombinant human versikine (1  $\mu$ M). Versikine rapidly induced inflammatory cytokines interleukin-1 $\beta$ (IL-1 $\beta$ ) and IL-6 (Figure 1A-B). However, recombinant versikine had no discernible effects on cell-cycle progression of MM1.S myeloma cells (supplemental Figure 1) or primary CD138<sup>+</sup> myeloma cells (not shown).

VCAN, the precursor to versikine, has been variably reported to be expressed by carcinoma tumor cells or components of the carcinoma microenvironment; however, its cellular origin in the myeloma niche is unknown. We carried out RT-PCR using VCAN isoform-specific primers in paired myeloma CD138<sup>+</sup> plasma cells, CD14<sup>+</sup> monocytes/ macrophages, and CD14/CD138 double-negative fractions from 3 newly diagnosed patients. All 4 VCAN isoforms were predominantly expressed by CD14<sup>+</sup> monocytes/macrophages (supplemental Figure 2A). Known versicanases include ADAMTS-1, -4, -5, -9, -15, and -20.<sup>5</sup> We readily detected ADAMTS1, ADAMTS4, and ADAMTS5 messenger RNA (mRNA) expression in bone marrow mononuclear cell lysate whereas the rest were undetectable (data not shown). ADAMTS expression has been reported in bone marrow-derived mesenchymal stromal cells (BM-MSC).<sup>14</sup> We found that ADAMTS1 was expressed at higher levels by myeloma-derived BM-MSC compared with normal donor BM-MSC (supplemental Figure 2B). BM-MSC expressed much higher levels of ADAMTS1 message than either CD138<sup>+</sup> plasma cells or CD14<sup>+</sup> monocytes/macrophages (supplemental Figure 2C).

Inflammatory signaling in macrophages engages the MAP3K TPL2 (COT, MAP3K8).<sup>3,15</sup> *Tpl2* loss in murine bone marrow–derived macrophages (BMDM) abrogated II-1 $\beta$  production in response to versikine, but had little impact on II-6 production (Figure 1C). Interestingly, versikine induced II-12p40 but not II-10 (Figure 1D-E). Tpl2 negatively regulated II-12p40 production in response to versikine (Figure 1D), in a manner analogous to TLR agonists.<sup>16</sup> Whereas versikine induced II-12p40 in isolation, concurrent Fc $\gamma$  receptor ligation through addition of ovalbumin (OVA)/anti-OVA immune complexes promoted II-10 production and skewed macrophage polarization toward an immunoregulatory M2b phenotype (II-12<sup>lo</sup>-II-10<sup>hi</sup>)<sup>17</sup> (Figure 1E).

Intact VCAN is thought to signal through TLR2.<sup>10</sup> However,  $Tlr2^{-/-}$  BMDM showed only a partial II-6 production defect (approximately 50%) in response to recombinant versikine (Figure 1F).

Versikine stimulation of wild-type BMDM rapidly induced JNK, p38-MAPK, and AKT phosphorylation, but *Tpl2* loss only affected p38-MAPK phosphorylation (Figure 1G).

Human THP-1 monocytic cells can be induced to generate macrophages that provide a functional model to study macrophage regulation, including polarization.<sup>18</sup> THP-1 macrophages transduced with versikine-plasmid or empty-vector control were cocultured with human myeloma MM1.S cells for 48 hours and RNA-seq analysis was performed on each cell type. Remarkably, only 23 genes were differentially expressed in MM1.S cells exposed to versikine-secreting macrophages and all were upregulated: among these, 13 genes defined a type I interferon (IFN) signature (Figure 1H). Interestingly, VCAN transcription was itself induced, suggesting a positive autoregulatory loop responding to cleaved VCAN. THP-1 macrophages expressing versikine demonstrated differential regulation (>2-fold) of 39 genes (4 downregulated, 35 upregulated): 13 upregulated genes defined a type I IFN signature (supplemental Table 1).

Figure 1. Versikine, a novel DAMP with immunoregulatory roles in the myeloma microenvironment. (A-B) Freshly explanted MAM from the human myeloma marrow cases indicated were exposed to 1  $\mu$ M versikine (Vkine) for 12 hours. Relative expression of IL-1 $\beta$  and IL-6 transcripts is shown. Black bars, vehicle; gray bars, versikine-treated; NS, not significant; veh, vehicle. (C-D) Wild-type (WT) and  $Tp/2^{-/-}$  BMDMs were treated with 1  $\mu$ M versikine, and cytokine concentrations were measured in the culture supernatant at 12 hours postexposure. (C) II-1 $\beta$  (left), II-6 (right); before II-1 $\beta$  assay, cells were treated with 5 mM ATP for 20 minutes. (D) II-10 (left); II-12p40 (right). (E) Versikine modulates macrophage polarization. BMDMs were exposed to vehicle, versikine alone, or versikine + OVA/anti-OVA immune complexes (IC), as previously described.<sup>17</sup> Versikine exposure resulted in M1-like phenotype (II-12<sup>h</sup>, II-10<sup>h</sup>) in the absence of concurrent Fc<sub>7</sub> ligation. Versikine + IC promoted macrophage polarization toward an M2b-like, immunoregulatory phenotype (II-12<sup>h</sup>, II-10<sup>h</sup>). (F) WT and  $Tp/2^{-/-}$  BMDMs were estimulated by versikine for 12 hours and II-6 protein was measured in the supernatant. (G) Signaling mediators induced by versikine stimulation of WT and  $Tp/2^{-/-}$  BMDMs. BMDMs were collected following stimulation with versikine at designed time-points (each number reflects minutes) and subjected to immunoblot analysis with the antibodies shown. (H) RNA-seq analysis of MM1.S myeloma cells exposed to versikine-transduced macrophages for 48 hours. Only 23 genes were eliferentially expressed and all were upregulated. Thirteen of 23 upregulated genes were ISGs (highlighted in gray). *VCAN* gene transcription changes are underlined. (I-J) Myeloma cell-macrophage cocultures were exposed to 0.5  $\mu$ M purified versikine for 4, 18, or 48 hours. Representative ISG transcription is shown for THP-1 (I) and MM1.S cells (J). Relative mRNA transcription is normalized to vehicle-only control at each time point. \*



**Figure 2. Versikine acts through IRF8 to promote transcription of ISGs.** (A) IRF transcription in MM1.S cells, following 48 hours of coculture with THP-1 macrophages in the presence of 0.5  $\mu$ M versikine (Vkine) or vehicle (veh). Expression is normalized to veh-only levels at 4 hours. (B) *IRF9* mRNA levels in MM1.S cells cocultured with THP-1 macrophages in the presence of 0.5  $\mu$ M versikine (gray bars) or vehicle (black bars) for indicated time lengths. (C) THP-1 cells expressing Vkine or an empty-vector (EV) control were transduced with control lentivirus (NT) or lentivirus expressing short hairpin (shRNA) targeting IRF8 (shIRF8) (see supplemental Figure 3 for validation of IRF8 knockdown at the protein level). Versikine-mediated induction of 3 ISGs shown was measured in the presence and absence of IRF8. (D) *EBI3* transcripts in patient-derived, freshly explanted MAM treated with 0.5  $\mu$ M versikine (gray bars) or vehicle (black bars) for indicated time lengths. (E) RT-PCR analysis for *EBI3* transcripts in patient-derived, freshly explanted MAM treated with 1  $\mu$ M versikine for 12 hours. Relative expression is normalized to vehicle-only control (= 1). (F) Staining of human myeloma bone marrow core biopsy consecutive sections with antibodies against neoepitope DPEAAE generated by V1-VCAN cleavage at Glu<sup>441</sup>-Ala<sup>442</sup> and T-cell marker CD8. DPEAAE constitutes the C terminus of versikine. Four patterns of staining were observed in 19 informative punches: Pattern (a): intense/moderate VCAN proteolysis-CD8 poor (single/doublet CD8<sup>+</sup> cells sparsely distributed within tumor). Pattern (c): weak/focal VCAN proteolysis-CD8 poor (single/doublet CD8<sup>+</sup> cells sparsely distributed within tumor). Pattern (d): absent VCAN proteolysis-CD8 poor (single/doublet CD8<sup>+</sup> cells sparsely distributed within tumor). (G) Proposed immunomodulatory roles of VCAN proteolysis. CD8 por (single-presenting cells, and tumor cells. Currently untested hypotheses are represented by broken lines. \**P* < .05, \*\**P* < .01, \*\*\**P* < .001.

Recombinant versikine induced upregulation of IFN-stimulated genes (ISGs) in THP-1 macrophages as well as in cocultured MM1.S myeloma cells (Figure 1I-J). IRF8 was dramatically induced at 18 and 48 hours in MM1.S cells, but induction of other IFN-regulatory factors (IRFs) was minimal (Figure 2A-B). Interestingly, IRF8 was upregulated in MM1.S cells only in the presence of versikine-producing

macrophages. Knockdown of IRF8 expression impaired ISG induction in THP-1 macrophages (Figure 2C). Irf8 expression in transplanted tumor cells has been shown to be inducible through an II-27–dependent mechanism.<sup>19</sup> We observed upregulation of the IL-27 subunit, EBI3, in tumor cells and in primary myeloma-associated macrophages (MAMs) treated with versikine (Figures 2D-E).

We stained myeloma bone marrow biopsy sections with antibodies against a VCAN neoepitope (DPEAAE<sup>441</sup>) generated by Glu<sup>441</sup>-Ala<sup>442</sup> cleavage of V1-VCAN.<sup>5</sup> DPEAAE<sup>441</sup> constitutes the C terminus of versikine. Consecutive sections were stained for T-cell marker CD8. We observed 4 staining patterns in 19 core biopsies (Figure 2F). Myeloma tumors displaying CD8<sup>+</sup> infiltration/aggregates<sup>20</sup> (n = 5 of 19) demonstrated intense/moderate VCAN proteolysis, as detected by the anti-DPEAAE antibody.

We hypothesized that the regulated degradation of VCAN by ADAMTS-type versicanases may modulate its tolerogenic potential by controlling VCAN bioavailability, disrupting its extracellular matrix networks and/or by generating novel bioactive fragments. We report that versikine, a product of VCAN proteolysis, possesses immunoregulatory activities that may promote innate myeloma sensing and "T-cell inflammation" through induction of type I IFN signatures, macrophage activation (IL-1β, IL-6, IL-12), T-cell chemotactic mediators (CCL2),<sup>21,22</sup> and IRF8 induction, the latter known to be critical for dendritic and myeloid-derived suppressor cell maturation and homeostasis, respectively,<sup>23,24</sup> as well as macrophage inflammatory responses<sup>25</sup> (Figure 2G). Taken together, our results suggest that versikine may antagonize the tolerogenic actions of intact VCAN and thus, may provide a novel antitumor strategy. The findings also suggest that, in addition to small leucine-rich proteoglycans previously shown to act as DAMPs,<sup>26</sup> fragments of large aggregating proteoglycans may have the capacity to stimulate innate immunity and provide a bridge to adaptive immunity.

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## Authorship

Contribution: C.H., S.F., J.J., S.X.C., J.L.J., S.P., and F.A. performed experiments; C.L., I.M., N.C., S.M., and P.H. provided reagents and/or advice on study design; S.S.A. provided crucial expertise, advice, and reagents throughout the study; F.A. was responsible overall for study design; C.H. and F.A. wrote the manuscript; and all coauthors reviewed, edited, and approved the manuscript.

Conflict-of-interest disclosure: C.H. and F.A. are listed as inventors in U.S. Provisional Patent Application No. 62/305 761 (March 9, 2016). The remaining authors declare no competing financial interests.

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