Antibody-drug conjugate targeting CD46 eliminates multiple myeloma cells

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Multiple myeloma is incurable by standard approaches because of inevitable relapse and development of treatment resistance in all patients. In our prior work, we identified a panel of macropinocytosing human monoclonal antibodies against CD46, a negative regulator of the innate immune system, and constructed antibody-drug conjugates (ADCs). In this report, we show that an anti-CD46 ADC (CD46-ADC) potently inhibited proliferation in myeloma cell lines with little effect on normal cells. CD46-ADC also potently eliminated myeloma growth in orthotopic xenograft models. In primary myeloma cells derived from bone marrow aspirates, CD46-ADC induced apoptosis and cell death, but did not affect the viability of nontumor mononuclear cells. It is of clinical interest that the CD46 gene resides on chromosome 1q, which undergoes genomic amplification in the majority of relapsed myeloma patients. We found that the cell surface expression level of CD46 was markedly higher in patient myeloma cells with 1q gain than in those with normal 1q copy number. Thus, genomic amplification of CD46 may serve as a surrogate for target amplification that could allow patient stratification for tailored CD46-targeted therapy. Overall, these findings indicate that CD46 is a promising target for antibody-based treatment of multiple myeloma, especially in patients with gain of chromosome 1q.

Introduction

The treatment of multiple myeloma (MM) has greatly improved in recent years with FDA approval of agents in the immunomodulatory imide drug (IMiD) and proteasome inhibitor drug classes. Nevertheless, myeloma remains incurable, and patients inevitably develop treatment-refractory disease. Furthermore, high-risk cytogenetic subgroups, including those with deletion of chromosome 17p or gain of chromosome 1q21, progress more rapidly through approved agents and have shortened overall survival (1, 2). Therefore, patients with relapsed/refractory (R/R) disease or with poor cytogenetic profiles are in dire need of novel therapies.

Antibody-based therapies have potential to fill this clinical need. Naked antibodies have recently shown increased promise with demonstration of single-agent activities of the anti-CD38 antibodies daratumumab and SAR650984 (3, 4). In addition, the anti–signaling lymphocyte activation molecule family member 7 (anti-SLAMF7) antibody elotuzumab was recently shown to improve outcome in combination with lenalidomide and dexamethasone in a randomized phase III trial (5). Antibody-drug conjugates (ADCs) have potential to further improve on the clinical efficacy of naked antibodies via targeted delivery of highly cytotoxic payloads directly to malignant plasma cells (6–8). ADCs have recently seen proof-of-concept clinical success in Hodgkin lymphoma (brentuximab vedotin) and human epidermal growth factor receptor 2–positive (HER2-positive) breast cancer (ado-trastuzumab emtansine) (9, 10). Because of the considerable potential for clinical benefit, novel ADCs should be evaluated in MM (11).

Our research objective is to identify a novel ADC for MM treatment, with an emphasis toward patients with R/R disease. We previously developed a novel antibody discovery method based on a phage antibody library selection on tissue using laser capture microdissection (12). By this method, antibodies were identified that bind to tumor cells residing in their natural microenvironment (12). The platform was pioneered on prostate cancer tissue. One novel antibody that showed excellent in vivo targeting properties (13) has been identified to target the CD46 antigen (also known as membrane cofactor protein, MCP; Y. Su and B. Liu, unpublished observations). CD46 is a multifunctional protein that has a role in complement inhibition, which may explain its overexpression on malignant cells (14), and cellular entry by pathogens including measles virus (15, 16). The latter quality has led to CD46 targeting in viral immunotherapy with the Edmonston strain of measles virus (17). In normal tissue, CD46 appears to have a low level of expression outside placenta and prostate (14).

The CD46 gene is located on the long arm of chromosome 1 (1q32.2), 50 Mbp from a clinically used FISH probe that may pro-
Results

CD46 antigen is highly expressed in myeloma cell lines. To evaluate whether CD46 was overexpressed in MM, we studied its cell surface expression by FACS on cell lines. CD46 was highly expressed on the cell surface of all MM cell lines tested (Figure 1A and Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI85856DS1). We next sought to quantify the CD46 antigen number per cell (referred to henceforth as antigen density), using methods described previously (25). The mean antigen density on MM cell lines RPMI8226 and MM1.S ranged from 454,668 to 470,991 for CD46, compared with 314,953 to 344,865 for CD38, a commonly used marker for MM (Figure 1B). It has previously been reported that extracellular CD46 antigen is shed from the cell surface of solid tumor cell lines (26). To assess whether MM cells shed CD46 antigen, we conducted Western blotting of cell lysates and supernatants from RPMI8226 cells. In either the presence or the absence of CD46 antibody stimulation, we found no appreciable shedding of CD46 antigen from MM cells into the culture supernatant (Supplemental Figure 2).

CD46 is upregulated in the context of the BM microenvironment. Myeloma is a disease in which the BM microenvironment promotes MM cell survival and chemotherapy resistance (27). The majority of MM patients have disease that is primarily localized to the BM. To assess whether the CD46 expression level in MM cells is impacted by this microenvironment, MM1.S cells were cocultured with the BM stromal cell line HS5. Analysis of mRNA expression by RNA sequencing (RNA-seq) showed the CD46 mRNA level increased in MM1.S when cocultured with HS5, compared with monocultures. (Supplemental Figure 1). RPKM, reads per kilobase of transcript per million mapped reads. CD46 and CD38 antigen densities in RPMI8226 and MM1.S, incubated with or without HS5-conditioned media (CM) for 3 days (data represent mean ± SEM, n = 3–5). Two-tailed Student’s t test, *P < 0.05, **P < 0.01.

Figure 1. CD46 is highly expressed on MM cells and is further increased in the setting of BM microenvironment. (A) CD46 expression in INA-6, RPMI8226, MM1.S, and MM1.R measured by FACS (solid lines), compared with nonbinding control (Ctrl = dashed lines, representative data, n = 3). (B) CD46 antigen density estimation compared with CD38 on RPMI8226 and MM1.S (data represent mean ± SEM, n = 3). (C) Coculture of MM1.S with HS5 BM stromal cells increases the expression of CD46 mRNA (n = 2). RPKM, reads per kilobase of transcript per million mapped reads. (D) CD46 and CD38 antigen densities in RPMI8226 and MM1.S, incubated with or without HS5-conditioned media (CM) for 3 days (data represent mean ± SEM, n = 3–5). Two-tailed Student’s t test, *P < 0.05, **P < 0.01.

Vide a surrogate marker for CD46. FISH is used clinically in MM for risk stratification, and amplification of 1q occurs commonly in MM (1, 18–20). Amplification of 1q21 (amp1q21) is considered a high-risk feature that becomes more prevalent at relapse (1, 2, 21). A recent report also suggests that amp1q21 remains a poor prognostic factor in the era of current protease inhibitor and IMiD therapies (22). Thus, antibody-based targeting of cell surface antigens such as CD46 that undergo genomic copy number gain on chromosome 1q is an attractive possibility in myeloma.

Our selection method preferentially identifies human monoclonal antibodies that are internalized via macropinocytosis, a tumor-selective pathway for cellular entry (23), and thus are well suited for ADC development (24). Therefore, we used these antibodies to develop a novel anti-CD46 ADC with the microtubule inhibitor monomethyl auristatin F (MMAF) (hereafter referred to as CD46-ADC). Herein we describe the preclinical characterization of CD46-ADC in MM using cell lines, orthometastatic xenograft models, and patient samples. We found high CD46 expression on MM cell lines and patient samples, and the expression level is further upregulated upon coculture with bone marrow (BM) stromal cells. CD46-ADC showed potent and specific myeloma cell killing activity in vitro and in vivo and on primary MM cells ex vivo. Thus, CD46-ADC has potential to be an efficacious treatment for MM. Finally, we evaluated the hypothesis that patients with disease demonstrating amp1q21 by FISH also coamplify the CD46 gene and upregulate antigen expression on the MM cell surface. The results support the use of 1q21 FISH as a biomarker for translation of CD46-targeting agents for use in MM.
Generation of anti-CD46 antibody and internalization by myeloma cells. We generated a panel of human monoclonal antibodies binding to domains 1 and 2 of human CD46 by phage and yeast antibody display (28) and identified a lead antibody, 23AG2, that binds to the target antigen specifically with high affinity. The equilibrium dissociation constant ($K_D$) of the antibody on a recombinant human CD46 protein fragment was 2.99 nM on the BLItz system (Supplemental Figure 3). Measured on living MM cells, $K_D$ values were 1.19 nM for RPMI8226 and 2.24 nM for MM1.S, respectively (Supplemental Figure 4). To determine whether the anti-CD46 antibody (23AG2) is internalized by MM cells, we incubated the antibody with MM1.R, analyzed internalization by FACS for CD46. For comparison, CD38 was also studied in parallel. MM1.S and MM1.R cells showed upregulation of CD46 when cultured with HS5 and BM61 BM stromal cells after 96-hour incubation ($n = 3$), suggesting that a factor from BM stromal cells may increase CD46 expression by MM cells (Figure 1D). In contrast, CD38 showed a variable response upon incubation of MM1.S and MM1.R cells with stromal cell–conditioned media ($P = 0.15$ and 0.8, respectively) (Figure 1D). The nature of the factor(s) and whether it is BM specific remains to be determined.

Figure 2. Potent and selective activity of CD46-ADC on MM cells compared with BM stromal cells, with potentiation of ADC effect in the context of MM-stromal interactions. (A) Confocal immunocytochemistry of MM1.R after 4 (left) and 18 (right) hours of incubation with anti-CD46 antibody (red). Late lysosomes shown with anti-LAMP1 antibody (green), and partial colocalization shown in merged panel (yellow). (B) Dose response for CD46-ADC inhibition of viability of MM cells compared with HS5 and BM61 BM stromal cells after 96-hour incubation ($n = 3$). NR: EC$_{50}$ not reached due to lack of killing at the highest concentration tested. (C) Lack of effect of nonbinding control ADC until approximately 100 nM ($n = 3$). (D) Annexin V and PI staining of MM cell line INA-6 for 0–10 nM CD46-ADC, with apoptosis and death by 48 hours (representative data, $n = 3$). (E) Sensitivity of MM1.S cell line to CD46-ADC is increased in the presence of HS5, BM61, or HS27A BM stromal cells. EC$_{50}$ was 2.25 nM on MM1.S alone and 0.77 nM, 0.92 nM, and 1.05 nM for MM1.S in the presence of HS5, HS27A, and BM61, respectively (data represent mean ± SEM, $n = 3$).
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The induction of apoptosis and death was detectable in MM cells tested for annexin V and propidium iodide (PI) after 48 hours of ADC treatment (Figure 2D). Next, we examined the effect of BM microenvironment interactions on the efficacy of CD46-ADC. Cocultures of MM1.S cells with BM61, HS5, or HS27A BM stromal cells enhanced the potency of CD46-ADC (Figure 2E), consistent with our observations of increased CD46 expression in coculture and correlation of potency to CD46 levels described above.

We next sought to validate that CD46-ADC cytotoxicity is dependent on binding to CD46 for specific tumor cell killing. First, competitive cell binding of the anti-CD46 antibody was measured and detected in the presence of increasing amounts of the recombinant CD46-Fc fusion protein (Supplemental Figure 8A). In addition, the CD46-Fc blocked the cytotoxic effect of CD46-ADC (Supplemental Figure 8B). Next, the effect of CD46 knockdown on myeloma cell cytotoxicity of CD46-ADC was tested. The MM cell line H929 was infected with lentivirus coexpressing green fluorescent protein (GFP) and shRNA against CD46. By quantita-

Figure 3. In vivo CD46-ADC antimyeloma activity in the RPMI8226-Luc disseminated xenograft model. (A) Study treatment scheme. RPMI8226-Luc cells were injected i.v. and established for 10 days. Starting on the tenth day (treatment day 1), a total of 4 injections of PBS, control nonbinding ADC (5 mg/kg), naked CD46 antibody (5 mg/kg), CD46-ADC (5 mg/kg), or bortezomib (1 mg/kg) were given twice per week (n = 5 mice per group). (B) Disease was monitored by bioluminescence imaging (BLI) (top views, dorsal; bottom views, ventral). BLI measurement in photons per second per cm² per steradian (p/s/cm²/sr) was translated to color to indicate disease activity in the mice by the legend shown at far right. Tx, treatment; mAb, naked antibody. (C) Kaplan-Meier survival curves of NSG xenografts transplanted with RPMI-Luc and treated with CD46-ADC or controls.
to reverse growth (Figure 3B). It should be noted that this bortezomib schedule was chosen for comparison with the ADC, not to simulate clinical use, which would be continuous. The survival of mice treated with CD46-ADC was significantly improved over that of control groups (hazard ratio [HR] = 0.151 between control-ADC and CD46-ADC; \( P = 0.004 \)), with most of the treated animals living until study was discontinued at day 200 (Figure 3C).

Next, a luciferase reporter–bearing MM1.S cell line was used in a second xenograft model of orthometastatic MM. Two different ADC doses and a single-dose regimen were investigated (Figure 4). Mice were treated once every 3–4 days at either 4 mg/kg or 0.8 mg/kg for a total of 4 injections. A third group was treated with a single dose of 4 mg/kg. Tumor burden continually increased in the control groups treated with vehicle and control ADC (Figure 4A). Naked CD46 antibody delayed increase in bioluminescence, but mice succumbed by day 53. The single dose showed elimination of bioluminescence by day 36, but all mice relapsed and succumbed by day 139 (Figure 4C). The 0.8-mg/kg (4-dose) regimen eliminated bioluminescence through day 43, but all succumbed to relapse (Figure 4B). A comparison group was treated with bortezomib, which delayed increase in the bioluminescence activity but failed to reverse growth (Figure 3B). It should be noted that this bortezomib schedule was chosen for comparison with the ADC, not to simulate clinical use, which would be continuous. The survival of mice treated with CD46-ADC was significantly improved over that of control groups (hazard ratio [HR] = 0.151 between control-ADC and CD46-ADC; \( P = 0.004 \)), with most of the treated animals living until study was discontinued at day 200 (Figure 3C).

Anti–CD46-ADC potently eliminates MM cell line xenografts in vivo. RPMI8226 cells expressing firefly luciferase (RPMI8226-Luc) were used to establish an orthometastatic xenograft model in NOD.Cg-Prkdc-scid Il2rg \( ^{tm1Wjl} \)/SzJ (NSG) mice, as previously described (30). A total of 4 injections of CD46-ADC at 5 mg/kg were given once every 3–4 days. CD46-ADC resulted in near-complete elimination of myeloma cell bioluminescent signal, whereas controls (vehicle, nonbinding ADC, naked antibody) did not (Figure 3, A and B). A comparison group was treated with bortezomib, which delayed increase in the bioluminescence activity but failed to reverse growth (Figure 3B). It should be noted that this bortezomib schedule was chosen for comparison with the ADC, not to simulate clinical use, which would be continuous. The survival of mice treated with CD46-ADC was significantly improved over that of control groups (hazard ratio [HR] = 0.151 between control-ADC and CD46-ADC; \( P = 0.004 \)), with most of the treated animals living until study was discontinued at day 200 (Figure 3C).

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by day 173. The 4-mg/kg (4-dose) regimen eliminated bioluminescent activity throughout the duration of the study, and all mice survived to study discontinuation at day 212. Together, these studies showed potent in vivo activity of CD46-ADC. This effect is dose dependent, with significant activity at a very low dose (0.8 mg/kg) (HR = 0.215 compared with control-ADC; \(P = 0.004\)) and apparent curative potential at a moderate dose of 4 mg/kg.

**Establishing a clinical biomarker for CD46 expression in MM.** The CD46 gene is located on 1q32, in proximity to a clinically used FISH probe at 1q21. We thus postulated that patients with ampl1q21 may also amplify 1q32 and therefore carry increased CD46 expression. To explore this we mined data sets from previous gene expression analyses in MM (31–33). By Affymetrix array, CD46 transcript increased 3-fold in MM cells compared with normal plasma cells (\(P = 6.375 \times 10^{-9}\)) and was also significantly increased in a sequential manner between monoclonal gammopathy of unclear significance (MGUS) and smoldering myeloma (asymptomatic proliferative plasma cell states) (Supplemental Figure 9, A–C). Furthermore, data annotated by 1q21 FISH status (33) demonstrated coamplification of CD46 by array comparative genomic hybridization (Supplemental Figure 9D) and a concomitant 2.8-fold increase in CD46 transcript expression (\(P = 0.002\)) compared with MM samples that were 1q21 normal by FISH (Supplemental Figure 9E).

Analysis of additional data derived from the CoMMpass Study (Interim Analysis 6) also confirms that about 30% of newly diagnosed patients demonstrate focal amplification of the myeloid cell leukemia-1 (MCL1) gene located at 1q21 (Figure 5A). In addition to 1q21, the entirety of the 1q arm is amplified for a similar fraction of patients (Figure 5A, gray track). Furthermore, coamplification analysis for individual samples demonstrates that 85.4% of patients carrying MCL1 amplification also amplified CD46, and the majority of the 1q arm (purple line). Thus, there is a high level of coamplification between a negative prognostic factor (amp1q21) and the gene for the CD46-ADC target. Additionally, comparison of the correlations of MCL1 with CD46 copy number and MCL1 with CKS1B (another gene located at 1q21) demonstrates that the 1q21 and 1q32 loci correlate similarly (Figure 5B and Supplemental Figure 11) with the overall population (Figure 5C). Thus, data from both published studies and the CoMMpass Study indicate that nearly all patients with amp1q21 also demonstrate CD46 gene amplification and increased CD46 mRNA expression.

**Cell surface CD46 is increased in myeloma samples with gain of 1q21 by FISH.** To validate 1q21 FISH as a clinical biomarker, we next measured CD46 cell surface expression in a cohort of patients at diagnosis or at relapse with a pure monoclonal population of MM
cells (Supplemental Table 1). An initial cohort of 10 MM patient samples was evaluated, 7 of which had amp1q21 and 3 of which did not (Supplemental Table 1, samples 1–10). CD46 was coexpressed on the MM cell surface with well-known myeloma antigens CD38 and CD138 (Figure 6A), but expressed at low levels on nonplasma cells (NPCs) that have negative/low CD38 and CD138 expression and represent a heterogeneous mixture of normal mononuclear cells (MNCs) (Supplemental Figure 11, A and B). The average mean fluorescence intensity (MFI) (anti-CD46) of CD138-positive/CD38-positive cells by FACS was 152,049 (SEM 22,767) with amp1q21, significantly higher compared with 37,113 (SEM 9,926) in patients with normal 1q21 (P = 0.014, 2-tailed t-test; Supplemental Figure 11C). Thus, CD46 was overexpressed on MM cells from all patients and further amplified in patients with amp1q21, with low expression on nonmalignant NPCs.

We next measured cell surface antigen density by FACS to quantify the expression difference between amp1q21 and normal 1q21 patients. Samples were analyzed for CD46 cell surface expression on MM cells and matching NPCs from patients with and without amp1q21. In a second cohort of 10 patients, unselected MNCs were analyzed (Supplemental Table 1, samples 11–20). In 5 patients with amp1q21 the mean CD46 antigen density on MM cells was 313,190 (SEM 68,849), but other nontumor cell populations all had low CD46 expression (antigen density range 11,593–23,764) (Figure 6E and Supplemental Table 2). In BM from normal donors, monocytes (mean 58,320, SEM 6,874) and granulocytes (mean 54,439, SEM 10,688) had the highest CD46 antigen density, whereas others again were relatively low (range 8,443–23,772) (Figure 6F and Supplemental Table 3). Similar results were obtained from peripheral blood (PB) samples from normal donors, showing that CD46 antigen density is highest on monocytes (mean 56,237, SEM 11,649) and granulocytes (mean 40,523, SEM 8,165), but otherwise low (range 3,698–8,256) (Figure 6G and Supplemental Table 4).

In summary, CD46 expression was high on MM cells from 100% of patients (n = 25, cumulatively) and further amplified in patients with amp1q21.

Figure 6. CD46 is overexpressed on cell surface of primary MM patient cells and further amplified in patients with amp1q21 compared with normal 1q.

(A) FACS plot showing that CD46 surface expression correlates with CD38 in CD138-selected cells to identify the MM population by FACS (representative data, n = 25). (B) Quantitative FACS results for CD46 antigen density from MM versus NPCs from patients with normal (Nml) 1q (n = 5). (C) Quantitative FACS results for CD46 antigen density from MM versus NPCs from patients with amp1q21 (1q+, n = 5). (D) CD46 antigen density is further increased in amp1q21 patients (n = 5) compared with patients with normal 1q (n = 5). (E) CD46 antigen density on various BM normal cell populations compared with MM cells from 7 additional patients. (F) CD46 antigen density on various BM cell populations from 3 normal donors. (G) CD46 antigen density on various peripheral blood cell populations from 3 normal donors. Data represent mean ± SEM. Two-tailed t-test, *P < 0.05, **P < 0.01.
patients with amp1q21. The overall CD46 expression on normal hematopoietic cells is low. Notably, monocytes and granulocytes expressed relatively higher levels of CD46 compared with other normal cell populations. Interestingly, benign plasma cells from normal donors also had low CD46 antigen density (mean 22,475, SEM 4,869; Figure 6F and Supplemental Table 3), suggesting that high CD46 on MM cells occurs with malignant transition.

Anti–CD46-ADC potently and selectively induces primary MM cell apoptosis and death. We first studied internalization of our anti-CD46 antibodies by primary MM cells. As shown in Figure 7A (left panel), the anti-CD46 antibody was internalized and partially colocalized with the late lysosomal marker LAMP1. In contrast, no internalization into CD138-negative MNCs was observed (Figure 7A, right). We next evaluated whether CD46-ADC was specifically cytotoxic to primary MM cells ex vivo. CD138 positively selected cells from a patient with MM were incubated with CD46 antibody for 18 hours and costained with anti-LAMP1 antibody (green) and Hoechst dye (blue). Representative cell illustrates the intracellular localization of CD46 antibody, partially colocalizing with the late lysosomal marker LAMP1 (left). CD138-negative cells were treated in the same fashion and showed minimal binding of CD46 antibody without discernible internalization (right). Images were taken using a digital confocal microscope (Fluoview, Olympus) at ×60 magnification. (B) CD46-ADC depletes the number of CD138-positive, CD138-positive MM cells more potently in patients with amp1q21. Mean values with SEM are shown for CD46-ADC treatment of 2 patient samples with amp1q21 (red lines) and 2 patient samples with normal 1q21 (blue lines), compared with cells treated with nonbinding control ADC (black). (C) CD46-ADC does not affect the number of NPCs up to a concentration of 100 nM (n = 4) (data represent mean ± SEM).

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Figure 7. Ex vivo evaluation of CD46-ADC in patient sample MM cells. (A) Internalization of CD46 antibody (red) into MM patient cells ex vivo. CD138 positively selected cells from a patient with MM were incubated with CD46 antibody for 18 hours and costained with anti-LAMP1 antibody (green) and Hoechst dye (blue). Representative cell illustrates the intracellular localization of CD46 antibody, partially colocalizing with the late lysosomal marker LAMP1 (left). CD138-negative cells were treated in the same fashion and showed minimal binding of CD46 antibody without discernible internalization (right). Images were taken using a digital confocal microscope (Fluoview, Olympus) at ×60 magnification. (B) CD46-ADC depletes the number of CD138-positive, CD138-positive MM cells more potently in patients with amp1q21. Mean values with SEM are shown for CD46-ADC treatment of 2 patient samples with amp1q21 (red lines) and 2 patient samples with normal 1q21 (blue lines), compared with cells treated with nonbinding control ADC (black). (C) CD46-ADC does not affect the number of NPCs up to a concentration of 100 nM (n = 4) (data represent mean ± SEM).

The overall CD46 expression on normal hematopoietic cells is low. Notably, monocytes and granulocytes expressed relatively higher levels of CD46 compared with other normal cell populations. Interestingly, benign plasma cells from normal donors also had low CD46 antigen density (mean 22,475, SEM 4,869; Figure 6F and Supplemental Table 3), suggesting that high CD46 on MM cells occurs with malignant transition.

Toxicity evaluation of CD46-ADC in human CD46–expressing transgenic mice. We sought to study tolerability of CD46-ADC in vivo using a relevant animal model. The expression of murine CD46 is restricted to the inner acrosomal membrane of testes and the retina (34, 35). In addition, human CD46 and murine CD46 share weak homology, and our anti-CD46 antibody does not bind the mouse CD46 protein. For these reasons, the mouse xenograft experiments shown above provide limited data on potential toxicity of CD46-ADC. To circumvent this limitation, we studied in vivo tolerability of CD46-ADC using human CD46 transgenic mice (36). Following a single i.v. bolus injection of 6 mg/kg CD46-ADC, animals were monitored for body weight loss and sign of overt toxicity for 14 days. No significant body weight loss or overt sign of toxicity was observed (Figure 8A). At study discontinuation on day 14, necropsy study was performed. All organs appeared to be morphologically normal except for a slight increase in spleen size in CD46-ADC–treated animals. Histologic analysis of major organs showed no notable tissue damage (Figure 8B at ×20 and Supplemental Figure 14 at ×40 magnification). To assess whether CD46-ADC caused any notable effect on B cells in the spleen, we stained spleen sections with anti–mouse CD20 (Figure 8C). We measured diameters of CD20-positive regions in CD46-ADC–treated animals. Histologic analysis of major organs showed no notable tissue damage (Figure 8B at ×20 and Supplemental Figure 14 at ×40 magnification). To assess whether CD46-ADC caused any notable effect on B cells in the spleen, we stained spleen sections with anti–mouse CD20 (Figure 8C). We measured diameters of CD20-positive regions in CD46-ADC–treated animals. Histologic analysis of major organs showed no notable tissue damage (Figure 8B at ×20 and Supplemental Figure 14 at ×40 magnification). To assess whether CD46-ADC caused any notable effect on B cells in the spleen, we stained spleen sections with anti–mouse CD20 (Figure 8C).
expression level increases in the presence of a model of the BM microenvironment. Previous studies have characterized the CD46 gene as a transcriptional target of signal transducer and activator of transcription 3 (STAT3) (37). This finding is consistent with the notion that exposure of MM cells to marrow-derived cytokines, commonly associated with the activation of STAT3 activity, results in subsequent induction of CD46 transcript expression.

CD46 is an attractive cell surface target for ADC development because of its original identification from an unbiased screen-

Discussion

MM is incurable, and patients inevitably develop resistance to current therapies. New molecular targets and targeting agents have the potential to overcome limitations associated with current therapeutic strategies. Our study shows that CD46 is an attractive novel target for development of antibody therapy against MM. We have shown that CD46 is highly expressed on the surface of MM cell lines and primary MM cells (with relatively low expression on various normal BM and PB cell populations). In addition, the expression level increases in the presence of a model of the BM microenvironment. Previous studies have characterized the CD46 gene as a transcriptional target of signal transducer and activator of transcription 3 (STAT3) (37). This finding is consistent with the notion that exposure of MM cells to marrow-derived cytokines, commonly associated with the activation of STAT3 activity, results in subsequent induction of CD46 transcript expression.

CD46 is an attractive cell surface target for ADC development because of its original identification from an unbiased screen-
ing strategy that prioritized macropinocytosing antigens in their native conformations (12, 23). Our novel CD46-ADC potently and selectively eliminates MM cell lines in vitro and in vivo, and primary MM cells ex vivo. A unique distinction of CD46 is that the gene is located near a FISH probe already in routine clinical use (1). This 1q21 probe frequently undergoes genomic gain in patients with MM, especially in high-risk and R/R disease (2, 21). We have found that the genomic gain correlates well with CD46 antigen expression level and susceptibility to CD46-ADC. The discovery of a readily obtainable biomarker for target overexpression can prove extremely valuable for patient selection and stratification in clinical trials. Overall, our data suggest potential for CD46-ADC as a novel therapeutic for MM, with particular benefit predicted for patients with genomic amplification of the region where the CD46 gene resides (1q).

The finding of a putative drug target in high-risk or R/R MM with amp1q21 could be highly advantageous for the treatment of advanced disease. Putative oncogenes located on 1q include CKS1B and MCL1. In addition, CD46 overexpression itself is protective for cancer cell evasion of complement activation (38). MCL-1 is a prosurvival member of the B cell lymphoma 2 (BCL-2) family that is also overexpressed in high-risk MM patients and those with relapsed disease (39, 40). Importantly, MCL-1 upregulation has been described to evoke bortezomib resistance, which adds significant difficulty in treating advanced myeloma (41–43). These features of MM cells with 1q21 gain make CD46 an attractive target to combat drug resistance and oncogene accumulation in advanced disease.

If successful, there are several myeloma disease settings that may benefit from treatment with an efficacious ADC: (a) R/R MM with resistance to IMiDs and proteasome inhibitors, for which there is great clinical need for new therapies; (b) patients with high-risk disease features that tend to progress rapidly through current therapies — patients with amp1q21 at diagnosis fit into this category (2), making a particularly attractive group for treatment with CD46-ADC; and (c) addition of CD46-ADC to standard induction regimens for patients with newly diagnosed MM, with or without subsequent autologous stem cell transplant. Although standard approaches often achieve deep remission, they are not curative treatment for MM. The addition of a targeted cytotoxic therapy, such as an ADC that has a different mechanism of action, could potentially improve the odds of attaining cures that have thus far been elusive.

Current efforts on antibody targeting against MM are focused on a few lineage markers such as CD138 and CD38. Currently, 3 ADCs have been developed and entered clinical trials for MM. These include the anti-CD138 ADC indatuximab ravtansine (maytansinoid toxin DM4), the anti-CD56 ADC lornotuzumab mertansine (maytansinoid toxin DMI), and the anti-BCMA ADC GSK2857916 (MMAF toxin) (44–46). The linker/toxin combination for CD46-ADC is not used by any ADCs on a few lineage markers such as CD138 and CD38. Currently, 3 ADCs have been developed and entered clinical trials for MM. These include the anti-CD138 ADC indatuximab ravtansine (maytansinoid toxin DM4), the anti-CD56 ADC lornotuzumab mertansine (maytansinoid toxin DMI), and the anti-BCMA ADC GSK2857916 (MMAF toxin) (44–46). The linker/toxin combination used by CD46-ADC is most similar to the anti-BCMA ADC, except that the linker used is cleavable. The linker/toxin combination for CD46-ADC is not used by any ADCs currently in clinical trials, but has been described previously (29). A key differentiation between CD46 and the other targets of ADCs is that CD46 is a functional antigen and its expression is increased as the disease progresses. CD46 has a known role in negative regulation of complement, rendering cells dependent on its expression (14). In addition, the genomic region where CD46 resides is amplified in R/R MM (2, 21). This FISH biomarker for CD46 amplification is thus far unique in MM. Finally, the internalization mechanism of other MM antigens has not been characterized to assess whether they are optimally suited for ADC development (11). On the other hand, we have shown that our anti-CD46 antibodies enter cells through macropinocytosis (Y. Su and B. Liu, unpublished observations), an inherently tumor-selective pathway (23, 24), and our CD46-ADC showed high potency against MM cells, but selectively spared normal BM cells.

Interestingly, CD46 has been independently targeted for MM and other malignancies using oncolytic measles virus (17). An early report of 2 patients treated in phase I study in myeloma described substantial decreases in disease activity (47). It would be of interest to evaluate whether patients who tend to respond have amp1q21, as would be suggested by our study finding of higher CD46 expression levels in these patients. While promising, the approach needs to address immunogenicity of measles virus that limits repeat dosing. Human antibody-based approaches such as our CD46-ADC have less potential for immunogenicity. Nonetheless, the oncolytic measles virus approach has shown that CD46-targeted therapy can lead to tumor-specific effects (47, 48). Other than infusion-related reactions, oncolytic measles virus was well tolerated, showing that CD46 can be engaged in humans without triggering unacceptable toxicity (47, 48). It remains to be tested whether our CD46-ADC is safe in phase I trials.

We have found thus far that CD46-ADC was tolerable in mice, including transgenic mice that express the human CD46 gene. Our ex vivo study of CD46-ADC on primary cells showed potent cytotoxicity on myeloma cells, but not NPCs. Further study will determine potential differential effects of the ADC on subpopulations of NPCs. Because CD46 is expressed at low levels on all nucleated cells, there is potential for a tissue sink that could also impact the pharmacokinetics of CD46-ADC. We acknowledge this potential problem but believe it will be a minor issue as our quantitative measurement has shown that the CD46 antigen density on NPC subpopulations is low (typically ≤25,000 per cell). Ultimately, the pharmacokinetics and toxicity of CD46-ADC need to be investigated in nonhuman primates.

Mutations in CD46 are known to be a predisposing factor for about 10% of cases of atypical hemolytic uremic syndrome, with incomplete penetrance (49). CD46 expression is low on most normal cells, including endothelial cells (14, 50). A potential toxicity concern is that an antibody-based therapy targeting CD46 may block complement inhibition and potentially induce atypical hemolytic uremic syndrome. However, our ADC binds a conformational epitope residing in Sushi domains 1 and 2, outside of the interface where the complement pathway components bind (51). Consistent with this, we have observed that our CD46 antibody does not compete with complement binding and does not enhance complement activity in vitro assays (Y. Su and B. Liu, unpublished observations). Therefore the ADC is not expected to lead to excessive terminal complement cascade activation on normal cells.

In addition to MM, CD46 is overexpressed in prostate cancer, ovarian cancer, cervical cancer, breast cancer, endometrial cancer, lung cancer, and leukemia (38, 52, 53). Intriguingly, 1q gain is also
common in cancers from multiple tissue types (54, 55). We speculate that 1q gain may be the mechanism for CD46 upregulation in other cancers. Thus, CD46-ADC may have efficacy in other malignancies, particularly those that commonly have copy number gain of 1q.

In conclusion, we have identified a novel functional antigen, CD46, for ADC targeting of MM, with unique potential for high-risk and R/R disease that has genomic amplification at the CD46 gene locus and is in dire need of therapy. We have developed a novel CD46-ADC and shown that it is highly potent and selective in eliminating MM cells (cell lines and primary tumor cells) in preclinical models. We have found that CD46 genomic gain on chromosome 1q correlates with antigen amplification, and further identified a biomarker based on a clinical FISH test that can be used for patient stratification. Thus, our study could lead directly to the application of a novel ADC therapeutic for treating MM.

Methods

**MM patient and normal donor samples.** BM samples from MM patient samples 1–10 were separated into 2 fractions based on selection for MM patient and normal donor samples.

**Antibody generation.** We have previously identified a phage antibody binding to tumor cells in situ by laser capture microdissection-based selection using prostate cancer specimens (12). We later identified that CD46 is the target antigen and the original antibody binds to a conformational epitope residing in Sushi domains 1 and 2 (Y. Su and B. Liu, unpublished observations). A combined phage and yeast antibody display library selection approach (28) was used to identify additional anti-CD46 human antibodies. Briefly, the human CD46 gene fragment containing domains 1 and 2 was cloned into the pFUSE-Fc vector (Invivogen) to create a recombinant Fc fusion molecule that was produced in HEK293a cells, and purified by protein A chromatography.

**Cell surface antigen density determination.** Quantitative flow cytometry was performed to determine CD46 antigen density. Anti-CD46 and anti-CD38 antibodies were labeled with Alexa Fluor 647-conjugated streptavidin (Life Technologies). All samples were analyzed using an Accuri C6 (BD Biosciences) with a 96-well autosampler. Non-specific Fc receptor binding was minimized by preincubation with Clear Back reagent (Medical and Biological Laboratories Co.). To determine CD46 expression on patient samples, multicolor analysis was performed using FITC-conjugated anti-CD38 antibody (clone HIT2, Stem Cell Technologies) to identify MM cells in CD38-selected samples. For FACS of CD46 on various BM and PB cell populations, an extended antibody panel was used with anti-CD38 (clone HIT2) PerCP-Cy5.5, anti-CD19 (clone HIB19) BV510, anti-CD14 (clone M5E2) BV421, anti-CD11b (clone HL3) BV150, and anti-CD45 (clone HI30) BV515. A control IgG1 (YSC10) was created from a phage antibody picked randomly from the nonselected library, which does not bind to any cell surface antigen and hence is designated as a non-binding antibody in this article.

**MM cell lines.** MM cell lines bearing the firefly luciferase reporter gene were provided by Constantine Mitsiades (Dana-Farber Cancer Institute, Boston, Massachusetts, USA) (56). RPMI8226, MMLS, and MMR were purchased from American Type Culture Collection (ATCC). All MM cell lines were maintained in RPMI1640 with 100 U/ml penicillin, 100 μg/ml streptomycin, and 10% FBS. INA-6 was supplemented with IL-6 (2 ng/ml). Patient stroma culture, BM61, was derived from CD138-negative MNC fractions derived from BM aspirate.

**FACS analysis of cell surface CD46 expression.** To determine CD46 expression on MM cells, FACS was performed using biotin-labeled human anti-CD46 IgG1 followed by detection with Alexa Fluor 647-conjugated streptavidin (Life Technologies). All samples were analyzed using an Accuri C6 (BD Biosciences) with a 96-well autosampler. Non-specific Fc receptor binding was minimized by preincubation with Clear Back reagent (Medical and Biological Laboratories Co.). To determine CD46 expression on patient samples, multicolor analysis was performed using FITC-conjugated anti-CD38 antibody (clone AT1, Stem Cell Technologies) to identify MM cells in CD38-selected samples. For FACS of CD46 on various BM and PB cell populations, an extended antibody panel was used with anti-CD38 (clone HIT2) PerCP-Cy5.5, anti-CD138 (clone MI15) BV421, anti-CD45 (clone HI30) BV510, anti-CD19 (clone HIB19) BV150, and anti-CD45 (clone RPA-T4) PerCP-Cy5.5, anti-CD8 (clone RPA-T8) Pacific Blue, anti-CD3 (clone HIT3a) FITC, anti-Lineage FITC, anti-CD34 (clone 5B1) PE, anti-CD61 (clone VI-PL2) PE, anti-CD33 (clone HIM3-4) FITC, anti-CD14 (clone Hop9) PerCP-Cy5.5 (BD Biosciences), and Live/Dead-Near IR (Life Technologies/Thermo Fisher Scientific) on a FACSCanto II (BD Biosciences) flow cytometer.
PAGE. After semi-dry transfer to Immobilon-P membrane (Millipore), Western blotting was performed using anti-CD46 antibody H-294 (sc-9098, Santa Cruz Biotechnology) followed by anti-rabbit HRP (Jackson ImmunoResearch Laboratories) and detected by chemiluminescence with Pierce ECL Western Blotting Substrate (Pierce/Thermo Fisher Scientific) according to the manufacturer’s instructions. Images were captured using a C-DiGit blot scanner (LI-COR Biosciences).

**Antibody internalization by confocal microscopy.** Alexa Fluor 647-labeled anti-CD46 antibody was incubated with MM cell lines for 4 or 18 hours, washed with PBS, fixed with 4% PFA, permeabilized with PBS with 0.1% Triton X-100 and 1% BSA, and analyzed by confocal microscopy (Olympus FluoView). Nonbinding isotype antibody was studied in parallel as a control. For internalization by patient cells ex vivo, CD138-positive and CD138-negative (control) cells were incubated with Alexa Fluor 647-labeled anti-CD46 or nonbinding isotype control antibodies for 18 hours, processed, and analyzed as described above. Subcellular localization to lysosomes was assessed by costaining with LAMP1 antibody (clone D2D11, Cell Signaling Technology).

**ADC generation and characterization.** MMAF was conjugated to anti-CD46 IgG1 via an mcvcpab linker (29). To create the mcvcpab linker, N-ε-maleimidocaproyl-lysine succinimide ester (Pierce) was dissolved in anhydrous dimethylformamide (DMF) (final 0.14 mM). This solution was then added to valine-citrulline-p-aminobenzylalcohol (Consortis Biosystems) (final 0.14 mM). After brief agitation to dissolve all components, diisopropylethylamine (DIPEA) was added (final 0.41 mmol), incubated at room temperature for 1 hour, and precipitated with cold ethyl acetate (EtOAc) to form mcvcpab. Bis-nitrophenyl carbonate (0.41 mmol) was added along with DIPEA (0.41 mmol) to mcvcpab dissolved in DMF and incubated at room temperature for 4 hours followed by EtOAc precipitation as before to yield maleimidocaproyl-valine-citrulline-p-aminobenzylalcohol p-nitrophenol carbonate (mcvcpab-PNP), MMAF hydrochloride salt (Consortis Biosystems), N-hydroxysuccinimide, and DIPEA were added to DMF-dissolved mcvcpab-PNP and incubated at room temperature for 16 hours to form mcvcpabMMAF (29). CD46 IgG1 was reduced by tris(2-carboxyethyl)phosphine (TCEP) at 37°C for 2 hours, purified by Zeba spin column (Pierce/Thermo Fisher Scientific), and dialyzed against PBS to remove free MMAF and analyzed by HPLC using chromatography with Infinity 1200 LC System (Agilent). The drug-to-antibody ratio was estimated from area integration using OpenLab CDS software (Agilent).

**Apoptosis assay.** Induction of apoptosis and cell death in myeloma cell lines was evaluated using an annexin V–FITC Early Apoptosis Detection Kit (Cell Signaling Technology). Cell lines were seeded into 96-well plates at 4 × 10^4 cells per well and incubated with varying concentrations of CD46-ADC or nonbinding control ADC at 37°C for 48 hours. Apoptosis and cell death were assessed in triplicate by FACS for annexin V–FITC and PI. For primary samples, induction of apoptosis and cell death was also studied using the above approach, with the following modifications. Magnetic bead column-separated CD138-positive and -negative cell fractions were each plated at 4 × 10^4 cells per well and incubated with CD46-ADC or nonbinding ADC at 37°C. After 48 hours, cells were analyzed by FACS using annexin V–FITC, PI, and CD38 (AT1, Santa Cruz Biotechnology). Annexin V staining and PI staining were gated separately for the CD138-positive, CD38-positive MM cells and the CD138-negative, CD38-negative NPCs.

**Cell proliferation assays.** The firefly luciferase reporter–expressing lines were used to determine ADC potency in vitro. Cells were plated into 96-well plates at 2,000 per well and incubated for 96 hours with CD46-ADC. Following administration of luciferin, firefly luciferase activity was measured as an indicator of viability using a BioTek Synergy 2. The data were normalized against untreated control wells and EC_{50} estimated using GraphPad Prism version 6.0c.

For assessment of patient cell sensitivity to ADC, unselected MNC samples were plated at 4 × 10^4 cells per well in 96-well plates and treated with CD46-ADC or nonbinding ADC at 37°C for 48 hours. Cells were harvested after treatment, washed, and stained with phycoerythrin-conjugated anti-CD138 (BD Biosciences) and FITC-conjugated anti-CD38 antibodies. The numbers of CD138-positive, CD38-positive MM cells and CD138-negative, CD38-negative NPCs were gated and counted, with the curve constructed following normalization to the cell numbers in untreated wells. Treatments were performed in triplicate and plotted with SEM.

**BM stromal cell coculture and conditioned media.** BM coculture experiments evaluating ADC activity in MML.S cells were performed using compartment-specific bioluminescence as previously described (56). Following ammonium-chloride-potassium lysis, CD138-positive cells were isolated from MNCs using the EasySep Human Whole Blood and BM CD138 Positive Selection Kit (Stem Cell Technologies). HS5 and HS27A were purchased from ATCC. BM61 cells were generated from cultures of CD138-negative BM MNCs from a myeloma patient sample. HS5, HS27A, or BM61 cells were grown in RPMI1640-supported cultures for later use in monoculture or coculture assays. Coculture assays were initiated using a 2:1 ratio of bone marrow stromal cells to MM cells.

**CD46 gene expression of MM cells in coculture was evaluated as follows:** MML.S cells were seeded in monoculture at a density of 5 × 10^5 cells/ml, or, for coculture, seeded at the same density over 70% confluent stromal cells. After 24 hours, cells were collected and treated with trypsin-free dissociation solution (Accumax), and CD138-positive cells were harvested after treatment, washed, and stained with phycoerythrin-conjugated anti-CD138 (BD Biosciences) and FITC-conjugated anti-CD38 antibodies. The numbers of CD138-positive, CD38-positive MM and CD138-negative, CD38-negative NPCs were gated and counted, with the curve constructed following normalization to the cell numbers in untreated wells. Treatments were performed in triplicate and plotted with SEM.

**BM stromal cell–conditioned medium was collected from HS5 cells cultured under serum-free conditions for 48 hours, as previously described (58).** MML.S and MML.R cell CD46 and CD38 antigen density was measured with or without the addition of conditioned media for 72 hours.

**CD46 knockdown.** CD46 was targeted for knockdown with shRNA oligo with sequence 5′-ATTGGAGAGAGCACGATTTAT-3′. The shRNA sequence was cloned into plKO.1-GFP vector, containing a U6 promoter to drive shRNA expression and an IRES-GFP as previously described (59). Lentivirus particles were produced in HEK293T cells by cotransfection of shRNA constructs with pPax2 and pMD2.G. H929 MM cell lines were incubated with lentivirus and assessed for infection by GFP expression and CD46 knockdown by FACS 7 days after infection.
In vivo animal study. For in vivo efficacy assessment of CD46-ADC, 5 × 10³ RPlM8226-Luc or MMLS-Luc cells were injected i.v. into NSG mice (4–6 weeks of age, male and female) (Jackson Laboratory) to create orthotumestatic MM xenograft models. Bioluminescence imaging (BLI) was used to monitor graft status (typically injected tumor cells established themselves in the BM and joint in 10 days). Four mouse groups were treated with CD46-ADC, control ADC (MMF-conjugated to a nonbinding human IgG1), naked anti-CD46 antibody, or vehicle control (PBS). Tumor status was assessed by BLI and results analyzed by Living Image (PerkinElmer). After treatment, mice were continuously monitored for survival endpoints over a period of at least 200 days. HR was determined by log-rank method and significance by Wilcoxon test. For tolerability assessment, transgenic mice expressing human CD46 under its native promoter (MCP) (36) were (used in strain C57BL/6, backcrossed into C57BL/6 background for over 8 generations). The following primers were used for genetic screening: hMCP-Tg1 (5′-ATTGTTGGGTCCCAGATCT-3′) and hMCP-Tg2 (5′-CGGAGAGGAGGTAGTACACG-3′). Eight-week-old male mice were used in the study. Both the CD46-ADC and the YSC10 nonbinding control ADC were tested. Animals were injected with an i.v. bolus of the testing agent at 6 mg/kg (n = 3) and monitored for body weight loss and other overt signs of stress for 14 days. At the end of the experiment, major organs were harvested, formalin-fixed, frozen in liquid nitrogen, cryosectioned by Cryostat (Leica Biosystems), and stained with H&E (ScyTek Laboratories). To assess the status of CD20-positive region in the spleen, we stained the spleen sections with goat anti-mouse CD20 antibody (clone M-20, Santa Cruz Biotechnology), and measured diameters of the CD20-positive regions following treatment (n = 74 for CD46-ADC-treated and 81 for control ADC-treated regions, respectively). P value was calculated by 2-tailed Student’s t test.

Data mining. CD46 mRNA expression and copy number data were mined from 3 published data sets (31–33). Data from the CoMMpass Study (Interim Analysis 6) were provided by the Multiple Myeloma Research Foundation (https://research.themmrf.org). A detailed explanation for these data mining procedures is provided in the Supplemental Methods.

Statistics. All data are presented as mean and SEM unless noted. Significance was determined using GraphPad Prism version 6.0c. Two-tailed Student’s t test was used for comparing 2 means. For comparing multiple groups, ANOVA was used with multiple comparison correction. Levels of significance are categorized as **P < 0.05, ***P < 0.01, ****P < 0.001, and *****P < 0.0001. Mouse model sample sizes were determined by preliminary in vivo experience with CD46-ADC and other ADCs published in the literature, rather than power calculation (46), and statistically significant results were observed.

Study approval. BM samples from MM patients were obtained from the UCSF and University of Colorado Anschutz Medical Campus hematologic malignancies tissue banks with approval from the UCSF and Western Institutional Review Boards, respectively. Informed consent was obtained from all who donated samples. Identifying patient information was replaced with sequentially assigned numbers, in accordance with Health Insurance Portability and Accountability Act guidelines. Mouse studies were approved by the UCSF Animal Care and Use Committee (AN092211-01) and Washington University in St. Louis Animal Studies Committee (20100272A1).

Author contributions

DWS, BTA, YS, CRB, and BL designed experiments. DWS, BTA, YS, CRB, AW, and BCH performed experiments. DAA, PW, and MAS aided in model system development. ACL, JLW, and TGM developed tissue banks and contributed patient samples. XW and JPA conducted the transgenic mice experiment. DWS, BTA, YS, CRB, and BL wrote the manuscript. BL conceived the overall project idea. All authors contributed to manuscript revision.

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