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CD28/B7-mediated costimulation is critical for early control of murine cytomegalovirus infection*

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Abstract

Control of acute murine cytomegalovirus (MCMV) infection is dependent upon both innate and adaptive immune responses, relying primarily upon natural killer (NK) and T-cell responses for control. Although CD28/B7 plays a clear role in T-cell responses in many antigen systems including some viral infections, the importance of costimulation during MCMV infection is unconfirmed. In addition, recent data suggest that CD28/B7 costimulation might also be important to Ly49H⁺ NK-cell expansion. We therefore hypothesized that CD28/B7 costimulation is critical to viral control after MCMV infection, and further that CD28/B7 costimulation plays a role in MCMV specific T and NK-cell responses. To test these hypotheses, we utilize C57BL/6 mice lacking costimulatory molecules B7-1 and B7-2 or CD28. After primary infection with MCMV, viral titers are significantly elevated in mice lacking CD28 or B7 compared with wild-type (wt) mice. Impaired viral control is associated with significant defects in peripheral T-cell responses to MCMV, which appear dependent upon CD28/B7 costimulation. Abnormal hepatic T-cell responses in CD28^{-/-} mice are preceded by impaired MCMV-specific Ly49H⁺ NK-cell responses. We conclude that CD28/B7 costimulation is not required for non-specific antiviral responses. We conclude that CD28-B7 costimulation is critical for early viral control during acute MCMV infection.

Keywords

Costimulation; Natural Killer cell; murine cytomegalovirus

Introduction

Control of acute cytomegalovirus (CMV) infection is dependent upon both innate and adaptive immune responses. It has been demonstrated that natural killer (NK) cell responses are critical to early viral control following acute CMV infection in some mouse strains (7,13,14,51,62). Viral clearance is further dependent upon intact T-cell responses, with CMV inducing specific cytotoxic T-cell (CTL) responses in infected hosts (16,48,52,54-56,66,67). Optimal CTL

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responses require professional antigen presenting cells (APC), and APC/T-cell interactions are thus critical to CTL differentiation in infected hosts (1,19,41). Although adoptive transfer of anti-CMV antibody is protective during acute infection (5,22,30,32,61), humoral responses to MCMV are slower to develop than cellular responses (9), and both B-cells and antibody appear dispensable during acute infection (29,74). Thus defects in either NK or T-cell responses have significant implications for viral control, which is clinically most obvious in patients with impaired NK or T-cell immunity (10,63,65).

T-cell responses to CMV occur through clonally restricted antigen receptors, resulting in proliferation and clonal expansion of CMV-specific cells (55). Generally, optimal activation of T-cells requires costimulation in addition to antigen specific signals (59). One such costimulatory mechanism is functionalized by activating receptor CD28 expressed on T-cell surfaces. Ligands for CD28, namely B7-1 and B7-2 (hereafter referred to as B7 molecules), are prototypic costimulatory molecules expressed primarily on antigen presenting cells (6,23, 24,34-36,59,76). Thus optimal activation of T-cells relies upon antigen presentation to the TCR, and is enhanced by costimulation via CD28/B7 ligation.

Although numerous studies demonstrate importance of costimulation in varying in-vitro antigen systems, there are few data published on costimulation during anti-viral responses, and even fewer utilizing in-vivo models. Of the few studies to date, CD28/B7 costimulation has been shown to have varying importance for T-cell regulation of other viruses [(18,44,49,70, 71), reviewed in (8)] and there are no studies evaluating the importance of costimulation in control of cytomegalovirus (CMV) infection. Despite this, there are circumstantial data suggesting that costimulation is important to control of murine CMV (MCMV). Included in MCMV's immune evasion repertoire are genes that interfere with expression of B7 molecules on antigen presenting cells. MCMV has been shown to down-regulate surface expression of both B7-1 (CD80) and B7-2 (CD86) costimulatory molecules in monocyte / macrophage and dendritic cells during infection (2,37,42). Given the known importance of these costimulatory proteins in development of adaptive T-cell responses in other systems, we hypothesized that if MCMV has evolved specific immune evasion mechanisms that down-modulate B7 molecule expression, then CD28/B7 costimulation must play a critical role in antiviral defense to infection with MCMV.

In addition to antigen specific T-cell responses, NK-cell subsets have recently been shown to expand in response to specific antigenic stimuli (21,26,58). One example is Ly49H⁺ NK-cell subset expansion in response to murine CMV (MCMV), which is triggered by activating receptor Ly49H binding to its recently described ligand MCMV protein m157 (4,12,21,25, 51,64,73). Because this NK subset expansion is similar to adaptive T-cell responses, it has been postulated that costimulation might also be important to NK-cell expansion and effector function (26,45). Murine NK-cells have been shown to express CD28, and current data suggest that CD28 activation is important for optimal NK proliferation by enhancing cytokine production in these cells (26,45). Additionally, NK-cell cytotoxicity is enhanced by either type of B7 molecule (26,39,75), although B7 stimulated NK cytolysis does not absolutely require CD28 (17,40). Taken together, these data suggest that costimulation could indeed play an important role in NK-cell subset expansion in response to MCMV infection.

To test these hypotheses, we utilized C57/BL6 mice lacking costimulatory molecules B7-1 and B7-2 or CD28. After infection with MCMV, these mice demonstrate significant defects in viral control associated with impaired T and NK-cell responses. Impaired cellular responses to infection do not appear to be consequent to defective cytokine responses after infection in knockout mice. Thus our study for the first time confirms that CD28/B7 mediated costimulation is critical for early viral control during acute MCMV infection, and provides evidence that costimulation plays a role in adaptive cellular responses to MCMV.

Methods

Mice

Male C57BL/6_mice deficient of both B7-1 (CD80) and B7-2 (CD86) (referred to as B7^{-/-}) (11) or deficient of CD28 (referred to as CD28^{-/-})(26,60) were obtained from our colony. Age matched male C57BL/6 mice 8-10 weeks of age (Charles River) were used as controls. All animals were housed in a pathogen free environment adhering to the *Guide for the Care and Use of Laboratory Animals* prepared by the National Research Council (NIH Publication No. 86-23, revised 1985). After humane euthanasia, mice liver and spleen tissues were dissected aseptically, undergoing cellular isolation for flow cytometry analysis, viral titer analysis, or were frozen immediately in liquid nitrogen, then stored at -80°C for later analyses.

Viral infection

Primary CMV infection was achieved by intra-peritoneal (i.p.) injection of tissue culture passaged (once) salivary gland derived 5×10^4 PFU Smith strain MCMV (VR-1399/1981) obtained from ATCC (Rockville, MD). Prior to infection (day 0) cohorts of n=3 underwent analysis as controls. For infection experiments, n=6 each of wild type, B7^{-/-}, and CD28^{-/-} were infected, with cohorts of n=3 euthanized at days 2 and 6 post infection (pi).

Antibodies and flow cytometry

Fluorescent dye-conjugated antibodies specific for CD3 (PerCP-Cy5.5), NK1.1 (PE), Interferon- γ , and bromodeoxyuridine (BrdU, PRB-1) were all purchased from BD PharMingen (San Diego, CA). Cells isolated from livers or spleens of mice were stained with fluorescent dye conjugated mAbs and were analyzed by flow cytometry (FACScalibur, Becton Dickinson, Mountain View, CA)(26). Intracellular IFN- γ was analyzed in splenic NK cells (not further stimulated ex-vivo) using FITC-conjugated MAb to IFN- γ from BD PharMingen as previously described (27). Ly49H (APC) antibody was a kind gift from Wayne Yokoyama.

Isolation of hepatic mononuclear leukocytes

Hepatic mononuclear cells (MNCs) were isolated as described (26). Briefly, livers were perfused through the portal vein with 10 ml phosphate-buffered saline (PBS) to remove blood cells. Individual livers were ground with a 3-mL syringe insert and suspended in 10 ml medium. Single-cell suspensions were collected and centrifuged, and tissue debris was discarded. Cells were re-suspended in 2 ml 40% Percoll (Pharmacia, Uppsala, Sweden), layered on 2.5 ml 70% Percoll in 15 ml conical tubes. MNCs were recovered at the interface between 40% and 70% Percoll after centrifuge at 1600 rpm at room temperature for 25 minutes. Remaining red blood cells in MNCs were lysed by ammonium chloride solution. Hepatic MNCs were then subjected to flow cytometry analysis.

BrdU incorporation assay in-vivo

To determine cell proliferation in vivo, BrdU (Sigma, St Louis, MO; 1 mg/0.1 ml/mouse) was injected intraperitoneal into mice 3 hours prior to acquiring liver and spleen tissues. Splenocytes and hepatic MNCs were isolated, stained with fluorescent conjugated mAbs to CD3, NK1.1, and Ly49H followed by intracellular staining with FITC-labeled mAb to BrdU, and were analyzed by four-color flow cytometry as previously described (26). Subsets of CD3⁺NK1.1⁻, CD3-NK1.1⁺Ly49H⁺, and CD3⁻NK1.1⁺Ly49H⁻ cells were respectively gated to analyze BrdU incorporation. Ly49H monoclonal antibody was a gift from Dr. Wayne Yokoyama and biotinylated in our laboratory.

In vitro plaque assay

Spleen and liver tissues from day 0, 2, and 6 pi underwent in vitro plaque assay (IVPA) analysis to determine viral titer, as well as real time PCR to determine viral load. For IVPA, mouse embryo fibroblasts (MEF's) were grown to confluence in six well plates in DMEM (Gibco BRL). Following centrifugation (1000X G, 10 min) of 5ml homogenized tissue, 1 ml of supernatant was placed in each well. These plates were centrifuged at 1,000 G, and incubated at 37°C in 5% CO₂ for 3-4 hours. Plates were washed 3 times with PBS then covered with 3ml of 1% agar in DMEM. Following 6-7 days of incubation (37°C in 5% CO₂), plates were fixed in 10% formalin, stained with 1% crystal violet, and analyzed by low power phase contrast microscopy for plaque formation. All IVPA experiments were performed in duplicate, and viral titers are reported as mean \pm standard error of mean of both experiments.

IVPA techniques are time consuming and laborious, so we performed side-by-side comparisons of IVPA to real-time PCR to quantitate viral load in liver and spleen tissues. Linear regression analysis showed good correlation between IVPA and real time PCR results (R^2 0.89, p<0.001, data not shown). A drawback to real time PCR is that occasionally it was not sensitive enough to detect low viral copies, requiring the use of IVPA. Therefore when adequate copy numbers occur to allow detection, real-time quantitative PCR is an accurate surrogate for IVPA to quantify viral load in tissues.

DNA Isolation

DNA were extracted from tissues using QIAamp Tissue Kit (QIAGEN GmbH, Hilden, Germany. DNA extracted from tissue homogenates were eluted in 100 μ l of distilled water and stored at -20°C until analysis. DNA were amplified in a total volume of 25 μ l with 200 nM of each primer and 1.0 U of Taq DNA polymerase (GIBCO BRL) added in 2.5 μ l of a PCR buffer (50 mM KCl, 20mM Tris-HCl (pH 8.4), and 1.5 mM MgCl₂).

Quantitative real time PCR

Primer sets and fluorescence resonance energy transfer type probes were designed and optimized for MCMV GB and β-actin genes. Plasmids homologous to the sequences amplified by our PCR primers were cloned for these genes, and using serial dilution of known plasmid concentrations, standard curves were constructed for quantitation. Regression equations had very high R^2 values (.95 β -actin and .99 GB), and were used for subsequent copy number calculations. Real time PCR results are expressed as copies of DNA/mRNA per copies of βactin. All samples were analyzed in duplicate, and results are expressed as mean copies \pm standard error of mean. Sequences for real time primers/probes designed are as follows: β -actin forward attgtgatggatcccggtga, reverse agctcatagctcttctcccag, probe cacccacactgtgcccatctac, CMV GB forward tgtactcgaagggagagct, reverse cgttcaccaccgaagacac, probe cgcctcgaacgtgttcagcctg. For cytokine analyses, primers for IL-1, TNF- α , and IFN- γ were obtained from Super Array. Sequences of other primers are as follows from 5' to 3': IL-2: forward, tcactcctcacagtgacctcaagt; reverse, agcgcttactttgtgctgtcct. IL-12a: Forward, tgtcttagccagtcccgaaacct; reverse, gtgaagcaggatgcagagcttcatt. IL-15: forward, ccatctcgtgctacttgtgtttcct; reverse: caggacgtgttgatgaacatttggac. IL-7a: forward, atccacctcacacgaggcacaagt; reverse, caaacacgaagcagtttgggacc. IFN-a: forward, tgtctgatgcagcaggtgg; reverse: aagacagggctctccagac. GAPDH: forward, aactttggcattgtggaagg; reverse, acacattgggggtaggaaca.

PCR reactions were carried out using a Smart-Cycler (Cepheid, Sunnyvale CA), using the following program: initial denaturation 4 min at 94°C, 35 cycles-denaturation 30s at 94°C, annealing 30s at 53°C, elongation 30s at 72°C, followed by final elongation 7 min at 72°C, then hold at 4°C.

Statistics

Statistical analyses utilized one-way ANOVA or one or two-tailed student-T test. P-values \leq 0.05 were considered significant for all testing. Means are expressed as mean \pm standard error. Statistical software used was GraphPad Prism (Version 4.03, GraphPad Software, San Diego, CA).

Results

CD28/B7 costimulation is critical for early viral control

To evaluate influences of costimulatory molecules B7 and CD28 on early viral control, wild type, B7^{-/-}, and CD28^{-/-} C57BL/6 mice (H-2^b) were infected with MCMV. Viral titers were measured 2 and 6 days after infection using both *in vitro* plaque assay (IVPA) and quantitative real time PCR for CMV DNA of tissue homogenates from spleen and liver.

In both liver and spleen, there were significantly elevated MCMV titers (3 fold) by day 2 in CD28^{-/-} mice, while viral titers in B7^{-/-} mice were not significantly different than wild type mice (Figure 1A&B). Previous studies have shown that six days after MCMV infection, MCMV specific NK subset expansion peaks, and anti-viral T-cell expansion is beginning (20, 21, 58). In the current study viral titers in both B7^{-/-} and CD28^{-/-} mice were significantly elevated (~2 logs) compared to wild type mice in both liver and spleen 6 days after infection (Figure 1). Interestingly, CD28^{-/-} mice had significantly higher viral titers (double) than B7^{-/-} mice in both tissues by day 6 pi (Figure 1A & B). In addition, hepatic titers were slightly but significantly higher than splenic titers at both time points (comparison not shown). Viral DNA loads from all tissues correlated very well with IVPA titers, further confirming IVPA results (R²=0.87; data not shown). We therefore conclude that both B7 and CD28 molecules play essential roles in early control of primary MCMV infection.

CD28 and B7 are required for T-cell expansion in response to MCMV

CD8 T-cells are known to be vitally important to viral control following MCMV infection (16,53,55,56). Because CD28/B7 costimulation is known to be variably important to T-cell responses in other models of viral infection (8,18,44,49), one possible explanation for observed differences in viral titers was that T-cells in these knockout mice do not respond normally to MCMV without costimulation. To test this hypothesis, we evaluated CD3⁺ NK1.1⁻ T-cell expansion in liver and splenic tissue following acute MCMV infection using flow cytometry measuring total numbers and cell division.

As previously published, wild type mice showed significant expansion of hepatic T-cell counts by day 6 after infection, which corresponded with increased BrdU incorporation (Figure 2A & B). In contrast, B7^{-/-} and CD28^{-/-} mice had significantly lower hepatic T-cell counts 6 days after infection (p=0.04, p=0.03, Figure 2A & B). Although this corresponded to impaired hepatic T-cell division, as suggested by impaired BrdU incorporation, which was significantly lower in B7^{-/-} and CD28^{-/-} mice than in wild type mice (p=0.01, p=0.002, Figure 2B), it is also possible that low hepatic T-cell counts could be consequent to impaired hepatic T-cell migration or T-cell death.

In contrast to liver, overall splenic T-cell counts in B7^{-/-} and CD28^{-/-} mice were comparable to wild type mice (p=0.88, p=0.36, Figure 2A & B). Curiously, this was observed despite significant impairment of T-cell division in B7^{-/-} and CD28^{-/-} mice demonstrated by decreased BrdU incorporation (p=0.0007, p=0.05, Figure 2A & B). Despite maintenance of normal numeric T-cell responses, knockout mice have dramatically elevated splenic viral titers compared to wt mice 6 days after infection (Figure 1B). One possible explanation for these observations is that small changes in T-cell numbers driven by antigen-specific T-cell

proliferation could be obscured in the spleen as this is a primary lymphoid organ. Similar analyses of CD3⁺NK1.1⁺ cells (NKT) showed no significant expansion, and no significant differences between wild type and knockout mice (data not shown). From these studies we conclude that CD28/B7 costimulation is critical to T-cell division and subsequent peripheral expansion after infection with MCMV, and additionally that both B7 and CD28 influence accumulation of T-cells in the liver.

CD28, but not B7, is required for expansion of Ly49H⁺ NK-cells in liver

Robbins et al have recently shown that early T-cell responses and viral control of MCMV are dependent upon preceding NK-cell responses (57). Therefore we considered the possibility that diminished T-cell proliferative responses, and elevated viral titers in spleen and liver could also be additionally consequent to abnormal NK-cell responses in mutant mice. NK-cell responses to MCMV infection in normal mice begin at day 2 as non-specific responses, and peak 6 days after infection with subset expansion of Ly49H⁺ cells (20,21,58). We therefore hypothesized that impairment of early viral control observed in CD28^{-/-} and B7^{-/-} mice could be consequent to inadequate NK-cell expansion, and compared expansion between Ly49H⁺ and Ly49H⁻ NK-cells in livers after MCMV infection.

MNCs isolated from livers after MCMV infection were evaluated for Ly49H⁺ and Ly49H⁻ NK-cell expansion by flow cytometry as shown in Figure 3A, and summarized in Figure 3B. Confirming previously published findings, Figure <u>3B-a</u> shows significant expansion from baseline of Ly49H⁺ NK-cells in wt mice at day 2 of MCMV infection (p=0.01) that rise > 5 fold by day 6 of infection (p < 0.01)(20, 21, 58). This expansion is due to Ly49H⁺ NK-cell division, as confirmed by significant increases of BrdU incorporation in this subset at day 2 and 6 pi in wt mice (Figure 3B-b, p < 0.01 for both time points, compared to baseline). In contrast, non-specific Ly49H⁻ NK-cell expansion and division were modest (Figure 3B-c & d), which has also been consistently described by others (20, 21, 58).

As shown in Figure 3B, CD28 appears important for MCMV-specific NK subset expansion/ accumulation in liver tissue during MCMV infection. Unlike wt mice, CD28^{-/-} mice do not develop Ly49H⁺ NK-cell expansion/accumulation, showing no significant increase in $Ly49H^+$ NK counts above baseline (p > 0.05 for either time point, Figure 3B-a). In fact, CD28^{-/-} mice showed no significant difference between Ly49H⁺ (specific) and Ly49H⁻ (nonspecific) NK-cell counts 6 days after infection (p > 0.05, compare Figure 3B-a & c). Curiously, this deficit in Ly49H⁺ NK-cells occurred, despite significantly higher division rates in Ly49H⁺ NK-cells 2 days after infection in CD28^{-/-} mice (p < 0.01, Figure 3B-b). It is possible that this enhanced division was in part non-specific, as there was a concomitant similar, albeit smaller, elevation in Ly49H⁻ NK-cell division at this same time (p < 0.05, Figure 3B-d). Despite this, by 6 days after infection, Ly49H⁺ NK counts were significantly lower than wt controls, and Ly49H⁻ NK counts were not different than wt controls in liver tissues. Thus CD28 appears to be required for specific Ly49H⁺ NK-cell accumulation in the liver of infected mice, but does not appear to influence non-specific NK-cell responses to MCMV infection. It is unclear whether failed NK-cell expansion in CD28-/- mice livers reflects increased apoptosis in this population, and/or simply failure of normal recruitment of antigen specific NK-cells to the liver.

In contrast to CD28, B7 expression plays little or no role in expansion/accumulation of hepatic NK-cells after infection. Figure 3B-a shows that Ly49H⁺ NK-cells in B7^{-/-} mice significantly expand by day 6 pi compared to baseline (Figure 3B-a). Thus wt and B7^{-/-} mice showed no significant difference in Ly49H⁺ NK-cell counts 6 days after infection (p = 0.32), and BrdU incorporation in Ly49H⁺ NK-cells was also similar between wt and B7^{-/-} mice (Figure 3B-b).

Neither CD28 nor B7 are required for expansion of splenic Ly49H⁺ NK-cells

To explore whether costimulation is required for splenic NK-cell expansion, we evaluated Ly49H⁺ and Ly49H⁻ NK-cells after infection by flow cytometry. In contrast to liver, and similar to our T-cell findings, there was no significant difference in splenic Ly49H⁺ NK-cell numbers between wild type, B7^{-/-}, or CD28^{-/-} mice at either 2 or 6 days after infection (p=0.56 and p=0.95, respectively, Figure 4A&B), although splenic expansion of Ly49H⁺ NK-cells was much more modest (~2 fold) than that seen in livers (>5 fold). In wt mice, MCMV specific Ly49H⁺ NK-cells showed a non-linear increase by day 6 (Figure 4B) that has been previously published (20, 21). The initial decline in splenic NK counts 2 days after infection has been previously postulated to be secondary to either redistribution of splenic NK-cells to more peripheral sites of infection, or to apoptosis or consumption of these cells in the spleen (20).

Figures 4B-b&c show effects of B7 and CD28 on splenic NK-cell division following MCMV infection. Both B7^{-/-} and CD28^{-/-} mice show small but statistically significant increases in BrdU incorporation in Ly49H⁺ NK-cells 2 days after infection when compared to wild type mice (p < 0.05). Nonetheless, this increased division corresponds to either no change or slight decreases in splenic Ly49H⁺ NK-cell numbers on day 2 (Figure 4B-a). Although the hypothesis that these cells might be redistributing to other sites of infection is sensible, there is no evidence that this occurs on day 2 pi in the liver, when liver counts of Ly49H⁺ NK-cells are not significantly increased in either B7^{-/-} or CD28^{-/-} mice (Figure 3B). Six days after infection, wild type, B7^{-/-}, and CD28^{-/-} mice show comparable BrdU incorporation that is significantly elevated from baseline. This cell division correlates with modest increases in Ly49H⁺ NK-cell numbers in splenic tissues by day 6 (Figure 4B-a&c). Importantly, when cell division for splenic Ly49H⁺ and Ly49H⁻ NK-cells are compared, there is identical BrdU incorporation in Ly49H⁻ NK-cells for wt and knockout mice (compare Figures 4B-b&d). Thus it seems that non-specific NK-cell expansion is intact in knockout mice, and that splenic NK expansion in response to MCMV occurs by a non-specific mechanism that does not require CD28 or B7 costimulation.

Effects of costimulatory molecules on early cytokine production

As an alternative hypothesis, it is possible that $B7^{-/-}$ or $CD28^{-/-}$ mice have some fundamental defect in early cytokine responses that could explain the observed defects in T and NK-cell expansion independent of costimulation. To test this hypothesis, we examined cytokine mRNA in liver and splenic tissues from wild-type and $CD28^{-/-}$ mice, including IFN- α , IFN- γ , IL-1, IL-2, IL-12, IL-15, IL-17, and TNF- α both 2 and 6 days pi. There were no significant differences between $CD28^{-/-}$ and wt mice for any cytokine transcripts 6 days pi in either tissue (data not shown). There were several differences noted at day 2 pi illustrated in Figure 5 and discussed below.

Several cytokines are known to be critical to T-cell responses to MCMV, including IFN- α and γ , IL-2, and IL-12 (43,57). There were no significant differences in IL-2 transcription in hepatic tissues 2 or 6 days after infection that might explain abnormal T-cells numbers and division in CD28^{-/-} mice (Figure 5). There were, however, elevated hepatic IFN- α transcripts observed in CD28^{-/-} mice 2 days after infection that became comparable to wt by day 6 pi (Figure 5). It has been previously shown that lack of early viral control by NK-cells leads to over-expression of IFN- α after MCMV infection, and that elevated IFN- α impairs T-cell responses to MCMV (57). Although there were decreased levels of IL-2 and IFN- γ transcripts observed in spleens of CD28^{-/-} mice 2 days pi compared to wt mice, these returned to normal by day 6 pi (not shown), and these deficits did not appear to influence overall splenic T-cell numbers after infection (Figure 2).

Further, there are several cytokines known to be critical in early NK responses to MCMV including IFN- α , IL-2, IL-12 and IL-15 (45,46), Importantly, there were no significant differences in these cytokine transcripts in liver between wild type and CD28^{-/-} mice to explain defective hepatic Ly49H+ NK-cell expansion observed in CD28^{-/-} mice (Figure 5)..IL-2, IL-12 and IL-15 levels were comparable in wt and CD28^{-/-} mice, and as previously mentioned, day 2 liver IFN- α mRNA levels were actually higher in CD28^{-/-} mice. "Although IL-2 levels were lower in spleens 2 days pi, this did not appear to influence splenic NK responses (Figure 4). Finally, there were no defects in non-specific Ly49H⁻ NK-cell expansion after infection in knockout mice when compared to wild type mice in either liver (Figure 3) or spleen (Figure 4), which functionally confirms that non-specific responses to MCMV are intact in these knockout mice.

Because interferon- γ is a dominant cytokine produced by NK-cells during MCMV infection (21), we examined whether costimulation influences early IFN- γ production by NK-cells in response to MCMV infection. As shown in Figure 6, numbers of Ly49H⁺ IFN- γ -producing NK-cells in spleens 2 days after infection are not impaired in knockout mice. One might anticipate early NK IFN- γ responses to be consequent to m157-Ly49H interaction, but this does not appear to be the case 2 days pi, because Ly49H⁻ NK cells showed the same fold increase in IFN- γ expression 2 days after infection (Figure 6B). These findings also suggest that decreased transcription of IFN- γ observed in whole spleen lysates (Figure 5) is occurring outside the NK-cell compartment. Although wt splenic Ly49H⁺ NK cells expressing IFN- γ are numerically somewhat lower than those previously published at this time point, we did not stimulate our cells ex-vivo to elicit this response. Unfortunately, hepatic NK-cells were too few to allow similar analysis of IFN- γ -producing cells by flow cytometry (all were used in Ly49H enumeration and BrdU experiments), but transcripts of IFN- γ 2 and 6 days pi were comparable in liver between wild type and CD28^{-/-} mice (Figure 5).

Interpreting these cytokine results is somewhat complicated by differences in viral titers in knockout mice, as well as lack of cellular fraction specific analyses. Differences in IFN- γ expression in splenocyte lysates that were not observed in NK-cells highlight the possibility that splenocyte and hepatic lysates could mask subtle differences in cytokine production in individual cell subsets. Nonetheless, there appear to be no glaring defects in cytokine responses to infection in CD28 knockout mice, and taken together, these observations suggest that non-specific cytokine responses to MCMV are intact in CD28^{-/-} mice.

Discussion

This study confirms that costimulatory molecules B7 and CD28 are critical to host viral control during acute MCMV infection. Our viral titer data clearly demonstrate that both molecules are required for host immune control of virus replication following acute MCMV infection in both hepatic and splenic tissues. This influence in viral titers began as early as 2 days after infection, becoming more dramatic 6 days after infection in mice lacking B7^{-/-} or CD28^{-/-}. Early responses to MCMV (day 1-2) are known to be largely a consequence of APC release of cytokines, resulting in non-specific NK-cell expansion (3,19,47), while later viral control (day 6) is dependent upon antecedent non-specific and MCMV specific NK-cell responses, as well as subsequently developing T-cell responses (14,16,33,48,50,55,57). Because of the known importance of costimulation in adaptive immune responses, we further explored influences of costimulation on T and NK-cell responses in these knockout mice.

It is not entirely surprising that costimulation is critical to MCMV control, given recent data demonstrating the importance of B7 during herpes simplex virus and Epstein Barr virus infection (44,70,71). Further data supporting this hypothesis include MCMV targeting of DC responsiveness and costimulation as part of its immune evasion armamentarium (2,37,41,42).

B7 proteins enhance NK mediated killing, and conversely, B7-1 deficient antigen presenting cells display significant reductions in their ability to promote T-cell activation (23,39,75). Thus circumstantially, it seemed likely that costimulation via B7 and CD28 would be critical to MCMV control after acute infection. Despite these circumstantial data, there are other viruses, such as LCMV, that actually develop vigorous CD8 responses without costimulation (49). Thus ours are the first data that we are aware to definitively prove that costimulation is required for early control of CMV infection.

Antigen presentation and costimulation are known to be critical in development of normal Tcell responses in many antigen systems. Interestingly, MCMV has been shown to cause "immune paralysis" in DCs, which manifests as reduced release of IL-2 and IL-12, and impairment of T-cell activation (2,28). This impairment has recently been attributed to viral genes m138 and m147.5, which cause down-regulation of costimulatory molecules B7-1 (CD80) and B7-2 (CD86) (37,42) on DCs. Our studies confirm the importance of CD28/B7 costimulation in development of T-cell responses to MCMV in-vivo, as knockout mice showed impaired hepatic T-cell expansion and accumulation. This may be a direct consequence of diminished T-cell division, as there was significantly lower T-cell BrdU incorporation following MCMV infection in both B7^{-/-} and CD28^{-/-} mice compared to wild type mice. It is possible that poor expansion of T-cells in both B7^{-/-} and CD28^{-/-} mice is a consequence of early abnormalities in splenic IL-2 or IFN- γ transcription, but this will require further study. Overall, it is likely that this impaired T-cell response at least contributes to elevated viral titers seen 6 days after infection.

Much like APC/T-cell interactions, APC/NK interactions have also been shown to be critical in development of NK-cell responses to MCMV. Early during MCMV infection, APCs produce IFN- α/β and IL-12, causing IFN γ dependent non-specific NK expansion in-vivo (3,19,47). In addition to these non-specific NK responses, MCMV-specific Ly49H⁺ NK subset expansion occurs 4-8 days after infection, consequent to stimulatory receptor Ly49H⁺ activation (20), presumably activated by binding ligand m157 (12,64,73). This subset expansion has been reported to peak 6 days after infection in both liver and spleen tissue (3,20,21,58), and we reproduced these results in wild type mice. Recent work has shown that interaction between CD8 α^+ DC and NK-cells is important for expansion of this MCMV-specific Ly49H⁺ NK subset in response to acute infection (3). Altogether these observations by others are quite novel, demonstrating for the first time that NK-cells, long considered to be central to "innate" immunity, actually have "adaptive" characteristics.

Because of these adaptive characteristics, and the obvious importance of APC/NK interactions, we postulated that similar to T-cells, costimulation might be important to expansion of MCMV-specific NK-cell subsets. Indeed, $CD28^{-/-}$ mice had significantly reduced hepatic Ly49H⁺ NK subset expansion, which correlated with significant impairment in Ly49H⁺ NK-cell division 6 days after infection. In contrast, B7 appeared dispensable to MCMV-specific NK subset expansion, as B7^{-/-} mice developed normal hepatic Ly9H⁺ NK counts and BrdU incorporation. Nonetheless it was conceivable that deficits in Ly49H⁺ NK subset expansion in CD28^{-/-} mice could be explained by some fundamental defect in innate responses in these mice. Importantly, there were no significant differences in IFN- α , IFN- γ , IL-12, or IL-15 transcripts after infection that might explain impaired NK subset expansion in CD28^{-/-} mice. Thus, although not definitively proven, our data suggest that CD28 costimulation enhances subset expansion of peripheral NK-cells. This provocative observation will require further study.

The mechanism by which CD28 might participate in hepatic NK-cell expansion is currently not known. MCMV-specific NK expansion is somewhat unique in that it does not depend upon MHC-I antigen presentation, but instead relies upon direct recognition of m157 (an MHC-I homolog) expressed on the surface of infected cells by activating receptor Ly49H⁺ (72). We

speculate that just as MHC/TCR signaling and antigen specific T-cell expansion is enhanced by CD28/B7 costimulation, m157/Ly49H signaling may be similarly enhanced, and therefore lack of CD28 stimulation leads to depressed NK responses seen in CD28^{-/-} mice. Unfortunately, this hypothesis doesn't fit neatly with our results, as B7^{-/-} mice do not show the same effect. This is in contrast to the utter dependence of peripheral T-cell responses upon both costimulatory molecules. At best, we can consider that peripheral NK-cell subset expansion is influenced by costimulatory molecule CD28, but appears independent of costimulatory ligand B7. If true, this also suggests that CD28 might recognize a different B7 family ligand in B7^{-/-} mice that facilitates MCMV-specific NK subset expansion.

Although costimulation appears to be important to splenic T-cell division in response to MCMV infection, decreased T-cell division observed in knockout mice did not negatively impact overall splenic T-cell counts. This could simply be because splenocytes responsive to MCMV comprise a small percentage of the splenocyte population, or that day 6 is too early to detect significant differences in T-cell counts. Alternatively, splenic expansion of T-cells after MCMV infection might not be solely reliant on increased cell division costimulated by B7 or CD28 molecules, but occur by some other mechanism, such as resistance to activation-induced T-cell death. In contrast, splenic NK-cell expansion appears to be altogether independent of costimulatory influence, occurring primarily by non-specific expansion, shown by equal expansion and division of both Ly49H⁺ and Ly49H⁻ NK-cells in wt and knockout mice. This is consistent with previously published hypotheses that splenic NK-cell expansion occurs by a mechanism different than that in the liver (68,69). Although this issue of splenic NK expansion has become somewhat controversial (38,68,69), it would appear that whatever the mechanism, splenic NK and T-cell expansion occurs independent of costimulation by B7 or CD28. It is interesting to note that development of relatively "normal" MCMV specific NKcell and T-cell numbers in splenic tissues was associated with better viral control in spleen tissues compared to liver (comparison not shown). Nonetheless, splenic viral titers were still much higher in both B7^{-/-} and CD28^{-/-} than wt mice suggesting that there is some important immune interaction missing in these knockout mice.

Interestingly, at almost every time point, CD28^{-/-} mice had significantly higher viral titers than B7^{-/-} mice. While this could be explained simply by CD28 being relatively more important to viral control, it is also possible that lack of CD28 molecules could have additional consequences. CD28^{-/-} mice still express B7 molecules, which can also bind CTLA-4 and PD-L1, known inhibitory receptors (15,23,31,40). It is therefore possible that absence of CD28 could cause unopposed B7/CTLA-4 or PD-L1 binding and signaling, down-regulating adaptive responses to MCMV. This could explain differences in viral titers between CD28^{-/-} and B7^{-/-} mice observed in livers as early as 2 days after infection and in both liver and spleen by day 6. Alternatively, because T-cell responses are in part dependent upon preceding NK-cell responses (57), it is also possible that B7^{-/-} mice, which develop normal NK but abnormal Tcell responses, have slightly better viral control than CD28^{-/-} mice, which have sequential impairment of both NK and T-cell responses. Robbins et al have shown that poor viral control by NK-cells leads to higher IFN- α levels after infection (57), which we observed in CD28^{-/-} mice on day 2. These hypotheses are not mutually exclusive, and are both consistent with previously published data, and likely explain differences in viral titers observed in our knockout mice.

In conclusion, costimulatory molecules B7 and CD28 are critical to viral control in mice during acute CMV infection, as absence of either molecule leads to significant elevations in viral titers. We feel that differences in viral titers observed in knockout mice are likely a direct consequence of impaired T-cell expansion, with more subtle contributions from impaired MCMV specific NK-cell expansion in CD28^{-/-} mice. Whether CD28/B7 costimulation is critical for NK subset expansion remains unproven, but this provocative observation certainly deserves further study.

These data thus support the hypothesis that down-regulation of costimulatory molecules by MCMV is an important immune evasion strategy during acute infection.

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Figure 1. Effect of costimulation upon viral titers during acute MCMV infection Wild type C57BL/6, B7^{-/-} or CD28^{-/-} knockout mice were infected with Smith MCMV (5 × 10^4 pfu) or receiving vehicle (PBS), and liver (A) and spleen (B) tissues were acquired 2 or 6 days after infection. Tissues were analyzed by plaque assay to determine viral titers, and results are reported in plaque forming units (PFU) per gm tissue. Each data bar represents n=3 mice. The data shown are representative of three comparable experiments.



Figure 2. Costimulation in T-cell responses during acute MCMV infection

Wild type C57BL/6, B7^{-/-} or CD28^{-/-} knockout mice infected with Smith MCMV (5×10^4 pfu) or vehicle had liver and spleen tissues removed 2 or 6 days after infection. Three hours prior to organ removal, each mouse received i.p. 1.0 mg BrdU. Single splenocyte preparations or hepatic MNCs were prepared, stained with MAb to CD3 (PerCP-Cy5.5), NK1.1 (APC), and Ly49H (PE) followed by intracellular staining with mAb to BrdU (FITC) as described in Methods, and analyzed by flow cytometry. CD3⁺NK1.1⁻ T cells were gated and analyzed for BrdU and Ly49H expression (A). Note that a small portion of CD3+NK1.1-BrDU-T cells expressed Ly49H, consistently with our previous observation (unpublished data). **A**. Representative flow cytometric plots of liver and spleen from mice at day 2 or 6 pi. **B**. T-cell

numbers for each group at various time points pi from a representative experiment. Each data point represents mean \pm standard error for n=3 mice. The experiment was repeated twice with similar results.

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Figure 3. Effect of costimulation on LY49H⁺ NK-cell expansion in liver during acute MCMV infection

Wild type C57BL/6, B7^{-/-} or CD28^{-/-} knockout mice infected with Smith MCMV (5×10^4 pfu) had liver tissues removed 2 or 6 days after infection. Three hours prior to organ removal, each mouse received BrdU. Hepatic MNCs were prepared and NK-cells were enumerated and had division estimated by four color flow cytometry. The MNCs were stained with fluorescent dye-conjugated mAbs to CD3 (PerCP-Cy5.5), NK1.1 (PE) and Ly49H (APC) for cell surface markers, and intracellular staining with mAb to BrdU (FITC) as described in Methods. Percentages of each subset (CD3⁻ NK1.1⁺Ly49H⁺, CD3⁻NK1.1⁺Ly49H⁻,

CD3⁻NK1.1⁺Ly49H⁺BrdU⁺ or CD3⁻ NK1.1⁺Ly49H⁻BrdU⁺) were determined and then

multiplied by numbers of bulk MNCs. **A.** Representative flow cytometric plots at day 2 pi from CD3⁻NK1.1⁺ gated cells. **B.** Mean cell numbers of CD3⁻NK1.1⁺ gated Ly49H⁺ and Ly49H⁻ NK-cells and BrdU incorporation in hepatic MNC over time. Each data point represents mean \pm standard error for n=3 mice. This experiment was performed twice with comparable results.

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Figure 4. Effect of costimulation on splenic NK-cell expansion during acute MCMV infection Wild type C57BL/6, B7^{-/-} or CD28^{-/-} knockout mice infected with Smith MCMV (5×10^4 pfu) or vehicle had splenic tissues removed 2 or 6 days after infection. Three hours prior to organ removal, each mouse received i.p.1.0 mg BrdU. CD3⁻ NK1.1⁺ gated splenocytes were enumerated and had division estimated by flow cytometry, as described in Figure 3. **A**. Representative flow cytometric plots 2 days after infection. **B**. Ly49H⁺ and Ly49H⁻NK-cell numbers and BrdU incorporation in spleens over time. Each data point represents mean ± standard error for n=3 mice. This experiment was performed twice with comparable results.



Figure 5. Effect of costimulation on early cytokine production upon MCMV infection Wild type C57BL/6 or CD28^{-/-} knockout mice infected with Smith MCMV (5×10^4 pfu) had liver and splenic tissues evaluated 2 days after infection. Total RNA were prepared and subjected to real-time PCR analysis for IL-1, IL-2, IL-12, IL-15, IL-17, IFN- γ , TNF- α and IFN- α . There were no significant differences in IL-1, IL-17 or TNF- α transcripts between wild type and CD28^{-/-} mice either in liver or spleen (not shown). Data shown in are arbitrary transcripts units (relative to GAPDH) for IFN- α , IFN- γ , IL-2, IL-12 and IL-15 in liver or spleen 2 days pi. Significant differences are noted by p-values in this figure. Each data point represents mean \pm standard error for 3 - 4 mice.



Figure 6. Costimulation and Ly49H $^+$ NK-cell interferon γ (IFN- γ) responses during acute MCMV infection

Wild type C57BL/6, B7^{-/-} or CD28^{-/-} knockout mice infected with Smith MCMV (5×10^4 pfu) or receiving vehicle (PBS) had splenic tissues removed 2 days after infection. Single splenocytes were prepared, stained with MAb to CD3 (PerCP-Cy5.5), NK1.1 (PE) and Ly49H (APC) followed by intracellular staining with FITC-conjugated MAb to IFN- γ , and then analyzed by four-color flow cytometry. CD3⁻ NK1.1⁺ cells were gated as described in Figure 3, and further analyzed for Ly49H⁺ and IFN- γ expression. Ly49H⁺ NK-cells expressing IFN- γ were enumerated. **A**. Representative of flow cytometric plots for Ly49H⁺IFN- γ ⁺ NK-cells

from each group. **B.** Numbers of INF- γ -producing Ly49H⁺ and Ly49H⁻ NK-cells in each group. Each data point represents mean + standard error for n=3 mice.