A Viral Immunoevasin Controls Innate Immunity by Targeting the Prototypical Natural Killer Cell Receptor Family

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Cell

A Viral Immunoevasin Controls Innate Immunity by **Receptor Family** Receptor Family

Graphical Abstract

Highlights

- \bullet Physiological ligand for the prototypical NK1.1 antigens is an MCMV immunoevasin, m12
- m12 decoy inhibits NK cells via NKR-P1B yet activates via NKR-P1A/C allomorphs
- Crystal structure of mouse NKR-P1B bound to m12 reveals a ''polar claw'' docking mode
- Co-evolutionary viral m12 and host NKR-P1B/C polymorphisms impact NK responses in vivo

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In Brief

A viral decoy is the long-sought ligand for a key immunoregulatory receptor family on natural killer cells.

A Viral Immunoevasin Controls Innate Immunity by Targeting the Prototypical Natural Killer Cell Receptor Family

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SUMMARY

Natural killer (NK) cells play a key role in innate immunity by detecting alterations in self and non-self ligands via paired NK cell receptors (NKRs). Despite identification of numerous NKR-ligand interactions, physiological ligands for the prototypical NK1.1 orphan receptor remain elusive. Here, we identify a viral ligand for the inhibitory and activating NKR-P1 (NK1.1) receptors. This murine cytomegalovirus (MCMV)-encoded protein, m12, restrains NK cell effector function by directly engaging the inhibitory NKR-P1B receptor. However, m12 also interacts with the activating NKR-P1A/C receptors to counterbalance m12 decoy function. Structural analyses reveal that m12 sequesters a large NKR-P1 surface area via a ''polar claw'' mechanism. Polymorphisms in, and ablation of, the viral m12 protein and host NKR-P1B/C alleles impact NK cell responses in vivo. Thus, we identify the long-sought foreign ligand for this key immunoregulatory NKR family and reveal how it controls the evolutionary balance of immune recognition during host-pathogen interplay.

Natural killer (NK) cells are a subset of innate lymphoid cells (ILC) capable of recognizing an array of pathological target cells. As

efficient sentinels responsible for the clearance of cancerous, virus-infected, transplanted, antibody-opsonized, and stressed cells, they discriminate between healthy-self, altered-self, and non-self targets through the integration of balanced signals delivered via multiple families of paired germline-encoded NK cell receptors (NKR) [\(Raulet and Vance, 2006](#page-15-0)).

Several NKR families have been described, many of which are grouped together in large genomic regions, such as the natural killer gene complex (NKC). In rodents, most NKR are C-type lectin-like proteins encoded within the NKC. These include the Ly49 (*Klra1*) receptors, the CD94/NKG2 (*Klrd1/Klrc1*) receptors, the NKG2D (*Klrk1*) receptor, and the NKR-P1 (*Klrb1*) receptors, which interact with genetically-linked C-type lectin-related (*Clr/Clec2*) proteins ([Kirkham and Carlyle, 2014; Raulet and](#page-15-1) [Vance, 2006\)](#page-15-1).

There exist at least five functional mouse NKR-P1 receptors, three stimulatory isoforms (NKR-P1A,C,F), and two inhibitory isoforms (NKR-P1B,G) [\(Carlyle et al., 2008; Kirkham and Carlyle,](#page-15-2) [2014](#page-15-2)). Originally described almost 40 years ago [\(Glimcher et al.,](#page-15-3) [1977](#page-15-3)), the NKR-P1 proteins were the first receptors to be identified as selectively expressed by NK cells. Yet to date, NKR-P1C, the prototypical NK1.1 antigen in B6 mice [\(Glimcher et al.,](#page-15-3) [1977; Koo and Peppard, 1984; Ryan et al., 1992\)](#page-15-3), and NKR-P1A remain orphan-activating receptors with unknown physiological ligands. On the other hand, the paired NKR-P1F and NKR-P1G receptors recognize overlapping ''self'' Clr (*Clec2*) ligands to balance signals during NK cell education and effector responses, while the inhibitory NKR-P1B receptor recognizes the broadly expressed self Clr-b (*Clec2d*) ligand ([Carlyle et al., 2004; Chen et al.,](#page-14-0) [2011; Iizuka et al., 2003; Kveberg et al., 2009\)](#page-14-0). To date, Clr-b has

¹²Lead Contact

Figure 1. MCMV Encodes a Clr-b-Independent NKR-P1B Immunoevasin, the m02 Family Member, m12

Mouse NIH 3T3 fibroblasts were infected with MCMV^{Smith} overnight and then used as stimulators for BWZ reporter cells expressing a CD3Z-NKR-P1B¹²⁹ fusion receptor (BWZ.P1B¹²⁹).

(A) Cell-surface expression of Clr-b on mock, MCMV-infected, or UV-treated MCMV-infected cells.

(B and C) BWZ.P1B¹²⁹ reporter cell assays using the above stimulators in the absence or presence of blocking Clr-b mAb (10 µg/ml).

(D and E) Clr-b-deficient NIH 3T3.AClr-b cells, targeted by CRISPR/Cas9 genome editing, analyzed by flow cytometry and BWZ.P1B¹²⁹ reporter assays following MCMV infection.

(legend continued on next page)

been shown to be involved in ''missing-self'' recognition under diverse pathological states [\(Kirkham and Carlyle, 2014](#page-15-1)), including cancer [\(Carlyle et al., 2004\)](#page-14-0), genotoxic and cellular stress [\(Fine](#page-15-4) [et al., 2010](#page-15-4)), hematopoietic transplantation ([Chen et al., 2015\)](#page-15-5), im-mune escape of primary lymphoma cells ([Rahim et al., 2015](#page-15-6)), and cytomegalovirus and poxvirus infection ([Aguilar et al., 2015; Voigt](#page-14-1) [et al., 2007; Williams et al., 2012](#page-14-1)), while the remainder of the self Clr ligands remain less well characterized.

Among pathogens recognized by NK cells, cytomegaloviruses (CMV) demonstrate co-evolution with their natural hosts, likely due to cycles of natural selection for viral versus host fitness. Since early responses to CMV infection are largely NK cell mediated, these viruses have evolved a diverse array of immunoevasins that antagonize NK cell function. For example, the MCMV m157 decoy immunoevasin binds directly to inhibitory Ly49C/I receptors from certain mouse strains, while the stimulatory Ly49H receptor has been evolutionarily adapted to directly recognize m157 as a foreign antigen, in turn dominantly establishing MCMV resistance in B6 mice [\(Arase et al., 2002](#page-14-2)).

Here, we identify the MCMV m02 family member, m12, as a functional decoy ligand that directly engages the mouse NKR-P1B inhibitory receptor and inhibits NK cell cytotoxicity. Importantly, we also demonstrate that m12 directly interacts with two stimulatory orphan receptors, the prototypical NK1.1 antigen, NKR-P1C, and NKR-P1A. These findings demonstrate an important role for the NKR-P1 receptor family in NK cell-mediated immunity to infection and identify m12 as the first physiological NKR-P1C^{B6} ligand, almost four decades since its discovery as the prototypical NK1.1 antigen ([Glimcher et al., 1977](#page-15-3)).

RESULTS

The murine NKR-P1B ligand, Clr-b (Clec2d), is rapidly lost in response to cellular infection by diverse viruses [\(Aguilar et al.,](#page-14-1) 2015; Voigt et al., 2007; Williams et al., 2012). However, MCMV-infected fibroblasts also induce a Clr-b-independent NKR-P1B ligand ([Aguilar et al., 2015\)](#page-14-1). To establish whether this was due to a viral immunoevasin, we infected cells with different MCMV strains and used them as stimulators for BWZ.CD3Z/NKR-P1B¹²⁹ (BWZ.P1B¹²⁹) reporter cell assays ([Mesci and Carlyle,](#page-15-7) [2007](#page-15-7)). As shown in [Figure 1,](#page-3-0) mock-treated NIH 3T3 cells or cells exposed to UV-irradiated MCMV strongly stimulated BWZ.P1B¹²⁹ reporter cells, a stimulus that was blocked using Clr-b mAb, while weaker stimulation was observed upon live MCMV^{Smith} infection, which could not be blocked using Clr-b mAb ([Figures 1](#page-3-0)A–1C). Thus, live MCMV infection is required to both downregulate host Clr-b and induce an alternative Clr-b-independent ligand.

To confirm that this signal was Clr-b independent, we generated Clr-b-deficient NIH 3T3 cells (NIH 3T3. AClrb) by CRISPR/

Cas9-mediated genome editing. Only MCMV-infected, but not mock-treated, NIH 3T3. Δ Clrb cells stimulated BWZ.P1B¹²⁹ reporter cells ([Figures 1D](#page-3-0) and 1E). Employing another approach, we compared MCMV infection of primary adult ear fibroblasts (AEF) from B6 WT and B6.*Clr-b–/–* mice ([Chen et al., 2015](#page-15-5)); here, BWZ.P1B¹²⁹ reporter cells responded only to infected B6.*Clr-b–/–* AEF, whereas WT AEF behaved similarly to NIH 3T3 cells (Figures S1A and S1B). Notably, similar results were observed upon MCMV infection of primary *Clrb–/–/*b*2m–/–* AEF cells (Figure S1C) and rat fibroblasts (data not shown). Thus, a C lr-b/ β_2 m-independent NKR-P1B ligand is induced upon MCMV infection of mouse and xenogeneic rat fibroblasts.

Since NKR-P1B is inhibitory, these results suggest that MCMV may encode a decoy immunoevasin. To test this, we analyzed distinct MCMV strains and large genomic deletion mutants to determine whether any isolates lacked the NKR-P1B reporter signal upon infection. The MCMV^{Smith} and MCMV^{K181} strains both generated strong BWZ.P1B¹²⁹ reporter signals; however, none of the mutants tested nor the parental BAC-generated MCMV^{MW97} strain elicited significant BWZ.P1B¹²⁹ reporter re-sponses ([Figures 1](#page-3-0)F and 1G). This suggested that MCMV^{MW97} might possess a polymorphism or loss-of-function mutation that abrogated BWZ.P1B¹²⁹ reporter cell recognition.

The MCMV-Encoded m12 Immunoevasin Targets

The P1B mediated Inhibition
To take advantage of the discrepancy between the MCMV strains, we conducted unbiased whole-transcriptome RNAsequencing (RNA-seq) using MCMV-infected NIH 3T3 cells to identify differences between MCMV^{Smith} and MCMV^{MW97}. RNAseq reads were then mapped to the MCMV genome sequence using TopHat analysis and visualized using the IGV browser. Since numerous MCMV immunoevasin genes are located at the left or right genomic termini, such as the m02 and m145 families, we analyzed differences within these regions [\(Figures 1](#page-3-0)H and 1I). Within the m145 family, few differences in transcript expression or genetic polymorphisms were noted between the two MCMV strains (Tables S1 and S2). In contrast, the m02 family displayed significantly more differences in transcript profiles and singlenucleotide polymorphisms (SNP). To screen these alleles functionally, we independently cloned all of the MCMV^{Smith} m02 and m145 family members and some other known immunoevasins (in total, 32 gene products spanning \sim 15% of the MCMV genome) into a mammalian expression vector, and then we tested them individually using human 293T transfectants as stimulators for BWZ.P1B^{B6} reporter cells. While none of the m145 family gene products yielded positive results, a single gene product in the m02 family, m12, was capable of directly stimulating BWZ.P1B^{B6}, but not parental BWZ reporter cells [\(Figures 1](#page-3-0)J and 1K). Thus, m12 is an NKR-P1B decoy ligand.

⁽F and G) Cell-surface Clr-b expression and BWZ.P1B¹²⁹ reporter cell assays using NIH 3T3 cells infected with different MCMV strains (Smith, K181, and MW97.01). Data are representative of at least three independent experiments.

⁽H and I) RNA-seq analysis was performed using RNA from NIH 3T3 cells infected with either MCMV-GFP^{Smith} or MCMV^{MW97}, and the resulting reads were mapped onto the MCMV genome. Histograms from MCMV-GFP^{Smith} (top) and MCMV^{MW97} (bottom) reads mapped to the (H) m02 and (I) m145 regions. SNP are shown as different colored lines in histograms.

⁽J and K) BWZ.P1B^{B6} reporter cell assays using HEK293T stimulator cells transfected with members of the (J) m02 or (K) m145 families. Data are representative of at least two independent experiments.

m12 Is a Givcosvlated Tvpe I Transmembrane Protein

We next characterized the topology, localization, and functional expression of m12. While a transmembrane (TM) domain was identified near the C terminus of m12 (Figure S2A), SignalP did not detect an N-terminal signal peptide. However, bioinformatic analyses identified a second in-frame ATG codon that yielded a canonical signal peptide (when analyzed by SignalP) and translational start site (according to the NetStart server). To validate the second ATG translational initiation site, we generated several m12 constructs, including native m12 (m12 $ATG1$, m12 $ATG2$) and N-terminal FLAG-tagged m12 with either a native or preprotrypsin signal peptide (m12^{NSP-FLAG}, m12^{PSP-FLAG}). These constructs were then analyzed using 293T transient transfectants, flow cytometry, and BWZ.P1B¹²⁹ reporter cell assays (Figures S2B and S2C).

Notably, cell-surface m12^{FLAG} was detected on transfectants using ATG2 and both signal peptides (m12^{NSP-FLAG}, m12^{PSP-FLAG}). confirming a type I TM topology (Figure S2B). In addition, all of the m12 constructs stimulated BWZ.P1B^{B6} reporter cells, with m12^{ATG1} yielding the weakest signal (Figure S2C). Collectively, these results suggest that ATG2 is preferred, the native signal peptide is cleaved, and m12 is expressed as a functional type I TM cell-surface protein.

Figure 2. Recognition of m12 by the Stimulatory NKR-P1C/NK1.1 Orphan Receptor

(A) Phylogenetic tree of the NKR-P1 family amino acid sequences from the B6, 129/BALB, and FVB strains using a ClustalW alignment.

(B-G) HEK293T cells transfected with m12^{Smith} were analyzed using BWZ reporter cells bearing the (B) NKR-P1B^{B6}, (C) NKR-P1C^{B6}, (D) NKR-P1B¹²⁹, (E) NKR-P1C¹²⁹, (F) NKR-P1B^{FVB}, or (G) NKR-P1C^{FVB} alleles. Data are representative of at least four independent experiments.

(H) BWZ reporter cells bearing a chimeric m12/ CD3z fusion receptor (BWZ.m12) were used to analyze HEK293T stimulator cells transfected with NKR-P1 receptors from the B6, 129/BALB, or FVB strains. Data are cumulative from four independent experiments normalized to control values. Mean \pm SEM are shown, with significance determined by two-way ANOVA relative to vector control.

To assess m12 biochemically, we conducted immunoprecipitation and western blot analysis of 293T transfectants. Under non-reducing conditions, the m12 protein exists both as a \sim 35–40 kDa monomer and a \sim 60–75 kDa dimer (Figure S2D). Under reducing conditions, only a single band of \sim 40 kDa was observed. Enzymatic digestion using endoglycosidase-H (EndoH) revealed a \sim 40 kDa EndoH-resistant band and a \sim 28 kDa EndoH-sensitive band, while treatment with peptide-Nglycosidase-F (PNGaseF) demonstrated a single band at \sim 28 kDa, in accordance with a predicted size of 24.6 kDa. These

data suggest that m12 is a highly N-linked glycosylated type I TM protein.

Receptor, NKR-P1C^{B6}
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We next tested whether m12 was capable of interacting with paralogous host receptor isoforms and alleles. To this end, BWZ reporter cells bearing CD3₂-fusion NKR-P1A/B/C/F/G receptors were constructed from the FVB, B6, and 129/BALB-strain alleles. Notably, the NKR-P1B^{FVB} and NKR-P1C^{B6} alleles react with NK1.1 mAb, whereas the NKR-P1B^{129/BALB} alleles do not [\(Carlyle et al., 1999, 2006; Ryan et al., 1992\)](#page-15-8), and NKR-P1A/ B/C forms a separate clade from NKR-P1F/G [\(Figure 2A](#page-5-0)).

When BWZ.NKR-P1 reporter cells were stimulated using 293T transfectants of the m12^{Smith} allele, the stimulatory NKR-P1A, NKR-P1F, and inhibitory NKR-P1G receptors did not appreciably recognize m12^{Smith} (data not shown), while all inhibitory NKR-P1B receptor alleles recognized m12^{Smith} ([Figures 2](#page-5-0)B, 2D, and 2F). Remarkably, the m12^{Smith} immunoevasin also significantly interacted with the stimulatory NKR-P1C^{B6} receptor, demonstrating that m12^{Smith} is a much-anticipated natural ligand for the prototypical NK1.1 antigen [\(Glimcher et al., 1977;](#page-15-3) [Koo and Peppard, 1984; Ryan et al., 1992\)](#page-15-3) [\(Figure 2C](#page-5-0)).

Figure 3. Direct Binding of m12 Protein to NKR-P1 Receptors

(A and B) Surface plasmon resonance sensograms and equilibrium binding curves (where appropriate) are shown for the binding of (A) m12^{Smith} (0.025-50 µM) and (B) m04^{Smith} (100 µM) to various NKR-P1 or Ly49 receptors, as indicated. Equilibrium dissociation constants (K_d) were calculated from two independent measurements, with each experiment performed in duplicate. Error bars represent the SEM. NB, no binding; ND, not determined.

(C) Sensograms are shown for the binding of 50 µM m12^{Smith} to NKR-P1B^{B6} and NKR-P1C^{B6} either before (solid lines) or after (dashed lines) exposure of immobilized receptor to the NK1.1 antibody.

(D) The relative binding of NK1.1 mAb to immobilized NKR-P1B^{B6} and NKR-P1C^{B6}.

(E) m12^{Smith} tetramer binding to HEK293T cells transfected with NKR-P1 alleles from B6, 129, and FVB strains. Numbers correspond to mean fluorescence intensities of the GFP⁺ gate.

Interestingly, the NK1.1⁻ NKR-P1C^{FVB} allele also recognized m12^{Smith}, but the NK1.1⁻ NKR-P1C^{129/BALB} allele did not [\(Figures](#page-5-0) [2E](#page-5-0) and 2G), demonstrating that paralogous stimulatory recognition of $m12^{Smith}$ is not unique to B6 mice.

To confirm these results reciprocally, we also generated BWZ.m12^{Smith} reporter cells and used them to semiquantitatively assess interactions with all NKR-P1 alleles on 293T trans-fectants ([Figure 2](#page-5-0)H). Using this approach, BWZ.m12^{Smith} reporters recognized NKR-P1BB6,129, FVB, NKR-P1AB6,129/FVB, and $NKR-PIC^{B6,\overline{F}VB}$, but not NKR-P1 C^{129} ; however, the stimulatory NKR-P1A/C paralogs interacted with m12^{Smith} more weakly than the inhibitory NKR-P1B counterparts in this cellular context. This was not due to splice variants nor inefficient surface expression of the stimulatory isoforms, as reciprocal domain swaps of NKR-P1B and NKR-P1C stimulated m12^{Smith} reporters similarly to native isoforms (Figures S2E and S2F). Collectively, dual recognition of m12 by paralogous NKR-P1 receptors with opposing signals suggests that NKR-P1:m12 interactions are evolving under selection pressures driven by host-pathogen interactions.

Direct Interactions between the NKR-P1A/B/C and m12 Direct Interactions between the NKR-P1A/B/C and m12

Next, we expressed soluble m12 and NKR-P1 receptor ectodomains and assessed their interaction by surface plasmon resonance (SPR). Here, $m12^{Smith}$ bound with similar affinity to the inhibitory NKR-P1B^{B6} (K_d = 5.8 μ M) and stimulatory NKR-P1C^{B6} ($K_d = 4.1 \mu M$) receptors ([Figure 3](#page-6-0)A). As observed using reporter cell assays, m12 recognition was found to be highly allele sensitive, with m12^{Smith} binding to NKR-P1B¹²⁹ and NKR-P1B^{FVB} with affinities 8-fold ($K_d = 0.7 \mu$ M) and 40-fold ($K_d =$ 0.15 μ M) higher than observed for NKR-P1B^{B6} ([Figure 3](#page-6-0)A). As controls, m12Smith did not bind to the structurally related Ly49AB6 receptor and m04 did not bind appreciably to any NKR-P1 alleles tested [\(Figure 3B](#page-6-0)). Notably, binding of m12^{Smith} to NKR-P1 C^{B6} , but not NKR-P1 B^{B6} , could be blocked using NK1.1 mAb ([Figure 3C](#page-6-0)), which only recognizes NKR-P1C^{B6} (and NKR-P1 B^{FVB}) but not NKR-P1 B^{B6} [\(Figure 3D](#page-6-0)) [\(Carlyle](#page-15-8) [et al., 1999, 2006; Ryan et al., 1992\)](#page-15-8).

We further generated m12^{Smith} tetramers and used these to stain transient 293T.NKR-P1 transfectants using an IRES-GFP reporter vector. These m12 tetramers were found to bind directly to cells expressing NKR-P1A^{B6,129,FVB}, NKR-P1B^{B6,129,FVB}, and NKR-P1CB6,FVB, but not NKR-P1C¹²⁹, NKR-P1F, or NKR-P1G [\(Figure 3E](#page-6-0)). Notably, m12^{Smith} tetramer staining of the activating NKR-P1A/C receptors was significantly lower than for the inhibitory NKR-P1B receptors at similar IRES-GFP reporter levels, indicating a weaker avidity of m12^{Smith} for the cellular NKR-P1A/C isoforms. In addition, the m12^{Smith} tetramers bound to fresh ex vivo B6-strain lymphocyte subsets, which mainly consisted of the NKR-P1B⁺ subset of NK cells, with significantly lower staining observed for the remaining NK cells, NKT cells,

and T cells, likely due to the weaker m12:NKR-P1A/C interactions at the cell surface [\(Figure 3F](#page-6-0)). Interestingly, differential results were observed for ex vivo lymphocytes from a number of distinct mouse strains (B6.*Clr-b–/–*, 129, BALB/c, FVB; Figure S3A), suggesting that differential susceptibility to MCMV m12 variants might be expected.

Reciprocally, we also generated NKR-P1B tetramers from the B6, 129, and FVB strains and used these to stain m12-expressing cells. Here, NKR-P1B^{B6,129,FVB} tetramers variably stained transient 293T.m12 and 293T.Clr-b transfectants, stable BWZ.m12 transductants, and NIH 3T3 cells infected with MCMV variants, demonstrating their utility in the absence of m12 mAb (Figures S3B–S3C and data not shown). Importantly, differences in NKR-P1 tetramer staining levels were observed using distinct NKR-P1 receptors across several m12 variants from distinct MCMV strains; these differences likely reflect their respective avidities for m12 variants at the cell surface (as they follow a similar affinity hierarchy to SPR results, where $FVB > 129 > B6$ for m12^{Smith}; see also below). Collectively, the above results demonstrate that there exists allelic and isoform specificity among the direct m12:NKR-P1A/B/C protein-protein interactions.

Structure of m12³ Complexed to NKR-P1B
We next determined the structure of NKR-P1B^{B6} bound to m12^{Smith} (Figure 4: Figure S4; Table S3). NKR-P1B adopted a classical C-type lectin-like domain (CTLD) fold and possessed a compact β 3- β 4 loop rather than the extended conformation observed in murine NKR-P1A ([Kolenko et al., 2011](#page-15-9)) (Figure S4A). Overall, m12 adopts an eight-stranded β -sandwich fold comprising one large (AIDEF) and one small (BGHC) β sheet, similar to that described for the m04 immunoevasin (Figures S4B–S4D) [\(Berry et al., 2014; Sgourakis et al., 2014](#page-14-3)). Like m04, the m12 I strand possesses two cysteine residues (Cys168 and Cys173) that form intra-molecular disulfide bonds with the CD loop (Cys88) and A strand (Cys58), respectively. However, m12 also contains an additional disulfide bridge between Cys40- Cys95 that serves to tether the relatively long N-terminal extension to the CD loop (Figure S4). Moreover, m12 possesses an additional α helix between the B and C strands. Thus, m12 possesses unique structural features that are directly involved in NKR-P1B receptor binding.

Topologically, m12 lies across the top of the NKR-P1B CTLD, such that the m12 β sheets lie parallel to the interface [\(Figure 4](#page-8-0)A). Here, m12 straddles NKR-P1B in a claw-like manner, whereby a relatively flat ''palm'' engages the planar top surface of the NKR-P1B CTLD, while several "fingers" grasp around the periphery [\(Figures 4B](#page-8-0)–4C). This binding mode results in an extensive interface that buries a total solvent-accessible surface area of 2,180 $\rm \AA^2$, which is considerably larger than the 1,680 $\rm \AA^2$ observed upon binding of the KACL endogenous ligand to the human NKp65 receptor [\(Li et al., 2013](#page-15-10)) (Figures S4E and S4F).

⁽F) m12^{Smith} tetramer binding to ex vivo splenocytes from B6 mice gated as indicated: NK cells (NKp46⁺CD3⁺); NKT cells (DX5⁺CD3⁺ or CD1d(αGalCer) tetramer⁺CD3⁺); conventional T cells (DX5⁻CD3⁺); $\gamma \delta$ T cells ($\gamma \delta$ TCR⁺CD3⁺). Numbers above gates indicate percentage of positive cells; numbers at top right corner represent the median fluorescence intensity of cells positive for 2D12 or m12 tetramer (see also Figure S3A). Specific staining, shaded histograms; fluorescence minus one (FMO) control, dashed lines. Data shown are representative of at least two independent experiments.

The interface is characterized by high shape complementarity and is dominated by hydrophilic interactions, including 14 H bonds and 8 salt-bridges ([Figure 4B](#page-8-0) and Table S4). The m12 palm encompasses residues from the A and B strands as well as the interconnecting loop (Ser60-Leu66) that form an extended surface that simultaneously engages the β 2- β 3 and β 4- β 5 loops as well as the β 4 strand of NKR-P1B. Here, the interaction is stabilized by a number of H bonds that are primarily derived from the m12 and NKR-P1B main chains ([Figures 4](#page-8-0)C–4D). In contrast to the flat, featureless palm, the m12 fingers are comprised of multiple large, primarily polar residues that form highly specific interactions around the edges of the docking site ([Figures 4C](#page-8-0)

Figure 4. Structure of the m12:NKR-P1B **Complex**

(A) Overview of $m12^{Smith}$ (blue) bound to NKR-P1B B6 (salmon) in cartoon representation. Sugar moieties are shown as sticks.

(B) Positive (blue) and negative (red) electrostatic potentials are shown on the surface of m12 that binds to NKR-P1B.

(C–G) Top view of the m12/NKR-P1B interface. m12 residues that form the ''palm'' (cyan sticks) and ''fingers'' (blue sticks) are shown relative to the NKR-P1B surface. Selected interactions between NKR-P1B and the m12 ''palm'' (D) and ''fingers'' (E–G). H-bonds and salt-bridges are shown as black and green dashed lines respectively.

and 4E–4G). For example, the m12 N terminus contains a highly charged $_{48}ERR_{50}$ motif that flanks the NKR-P1B CTLD ([Figure 4E](#page-8-0)). In addition, the loop region connecting the NKR-P1B β 1 sheet to the α 1 helix is similarly flanked by a triad of residues (Lys69, Asn73, Val76) that emanate from the unique m12 α 1 helix [\(Figure 4](#page-8-0)F). Further polar interactions are centered on the base of the m12 I strand, where Asp165 and Lys167 extend downward in order to engage Glu200, Asn206, Arg207, and Asn150 [\(Figure 4](#page-8-0)G). Overall, the breadth and the polar claw style of the interactions with NKR-P1B underlie the capacity of m12 to target this receptor family.

m12^{Smith/MW97} Impacts NKR-P1B

Immunoevasin Function Despite differential stimulation of BWZ.P1B^{129,B6} reporter cells by NIH 3T3 fibroblasts infected using MCMV^{Smith} versus MCMV^{MW97} ([Figure 1](#page-3-0)G), quantitation of m12 transcripts by qRT-PCR revealed similar mRNA levels between the two viruses [\(Fig](#page-9-0)[ure 5](#page-9-0)A). However, a single G271A SNP was observed in m12 $^{MWB\bar{7}}$ that resulted in a non-conservative E91K charge substitution. To determine whether this allelic m12 SNP affected immunoevasin function,

we compared BWZ.P1B^{B6} reporter stimulation by m12^{MW97} versus m12^{Smith} in 293T transfectants; indeed, m12^{MW97} was recognized more weakly than m12^{Smith} and host Clr-b [\(Fig](#page-9-0)[ure 5B](#page-9-0)). This suggests that m12^{Smith} is a superior NKR-P1B immunoevasin versus m12MW97, at least in B6 mice. Importantly, this finding was confirmed and extended using NKR-P1B tetramers, whereby all three allelic NKR-P1B^{B6,129,FVB} tetramers stained transient 293T.m12^{Smith} transfectants better than 293T.m12MW97 transfectants (Figure S3B).

We further validated these results using cytotoxicity assays. Here, we generated stable retroviral transductant YAC-1 target cells expressing native m12Smith, m12MW97, or host Clr-b

Figure 5. An Allelic MCMV m12 Polymorphism Controls NKR-P1B-Mediated Recognition and Inhibition of Cytotoxicity

(A) Quantitative real-time RT-PCR analysis of host *Clec2d* and MCMV *m12* transcript expression during infection of NIH 3T3 cells with MCMV^{Smith} or MCMV^{MW97}

(B) BWZ.P1B^{B6} reporter cell analysis of HEK293T stimulators transfected with host Clr-b, m12Smith, m12^{MW97}, or empty vector.

(C and D) YAC-1 cells transduced with $m12^{Smith}$, m12MW97, Clr-b, or empty vector were used as stimulators for BWZ.P1BB6 or BWZ.P1CB6 reporter cells.

(E and F) YAC-1 transductants as above were used as targets in ⁵¹Cr-release cytotoxicity assays with sorted NKp46⁺NKR-P1B⁺ or NKp46⁺NKR-P1B⁻NK-LAK effectors. Data are representative of at least two independent experiments. Mean \pm SEM values are shown, with pairwise significance determined by two-way ANOVA.

(G) Mapping of polymorphisms on the m12:NKR-P1B structure. Residues that differ between NKR-P1BB6 and NKR-P1B¹²⁹ are shown in pink, whereas those that differ between NKR-P1B¹²⁹ and NKR-P1B^{FVB} are shown in cyan. The E91K polymorphism identified in m12^{MW97} is highlighted in green, and the likely position of the A34V substitution found in m12^{C4A} is indicated in black. The m12 $_{48}ERR_{50}$ motif is denoted by a dashed oval.

hibited NKR-P1B⁺ NK-LAK cell cytoxicity [\(Figure 5](#page-9-0)E). Importantly, inhibition by m12^{Smith} and host Clr-b was mediated via $NKR-PIB^{B6}$, as it was not observed using NKR-P1B– NK-LAK effectors [\(Figure 5F](#page-9-0)). Notably, similar results were observed using sorted FVB-strain NKR-P1B⁺ and NKR-P1B⁻ NK-LAK effectors, partitioned using NK1.1 mAb (PK136), which recognizes NKR-P1B^{FVB} (but not NKR-P1C^{FVB}) ([Carlyle](#page-15-8) [et al., 1999](#page-15-8)) (Figures S5C and S5D). The

(indexed via IRES-GFP levels; Figure S5A). As observed using 293T transfectants, BWZ.NKR-P1 reporter analyses using YAC-1 stimulators show that BWZ.P1B^{B6} cells recognize m12^{Smith} more strongly than m12^{MW97} [\(Figure 5](#page-9-0)C). In contrast, BWZ.P1B^{FVB} cells recognize m12^{Smith} and m12^{MW97} similarly (Figure S5B), suggesting that the energetic profile of the interaction is allele specific. However, we could not detect significant stimulation of BWZ.P1C^{B6} reporter cells using YAC-1.m12^{Smith} stimulators ([Figure 5D](#page-9-0)), perhaps due to the lower m12 expression levels on these sorted stable transductants; notably, these data are consistent with reduced m12 tetramer staining of NKR-P1A/C relative to NKR-P1B ([Figure 3](#page-6-0)E).

Next, YAC-1 transductants were incubated as targets with sorted NKp46⁺NKR-P1B⁺ or control NKp46⁺NKR-P1B⁻B6-strain lymphokine-activated killer (NK-LAK) effectors in ⁵¹Cr-release cytotoxicity assays. As shown in [Figure 5,](#page-9-0) YAC-1.m12^{MW97} target cells were killed equivalently to control YAC-1(–) targets, while both YAC-1.m12^{Smith} and YAC-1.Clr-b target cells significantly insimilar recognition of m12^{Smith} and m12^{MW97} using BWZ.P1B^{FVB} reporters (Figure S5B) and NKR-P1BFVB tetramers (Figure S3B) but strong inhibition of cytotoxicity only by m12Smith suggests differential recognition of the m12 alleles by other stimulatory receptors, such as the NKR-P1A/C paralogs [\(Figure 2](#page-5-0)H; see also below).

The m12^{Smith}:NKR-P1B^{B6} structure provided an opportunity to understand the molecular basis underpinning the highly allelesensitive nature of this pivotal interaction. For example, NKR- $P1B^{129}$ differs from NKR-P1B^{B6} by 16 residues within the CTLD, but only 4 contact m12^{Smith} (S157R, Y158F, G201S, and S204T). Of these, the S157R and G201S polymorphisms would likely result in additional contacts with m12^{Smith} and would thereby account for the marked increase in affinity of NKR-P1B¹²⁹ compared to NKR-P1B^{B6} ([Figure 5G](#page-9-0)). In addition, the polymorphic m12 residue, E91, did not directly contact NKR- $P1B^{B6}$ but instead packed tightly against the m12 N terminus, thereby facilitating the interaction of the m12 $_{48}ERR_{50}$ motif

with NKR-P1B^{B6} [\(Figure 5](#page-9-0)G). Here, the m12^{MW97} E91K polymorphism would likely disrupt this packing and would thus indirectly abrogate the m12^{MW97}:NKR-P1B^{B6} interaction.

Collectively, these data demonstrate that an allelic m12MW97 SNP results in less-efficient NKR-P1B immunoevasin function, while the m12^{Smith} immunoevasin is capable of effective inhibition of both B6 and FVB NK-LAK effectors using YAC-1 targets. Moreover, the crystal structure provides a molecular basis for understanding the impact of host and viral allelic variation across the m12:NKR-P1B axis.

Wild MCMV Isolates Demonstrate Host-Driven Evolution
of the m12 Gene Product

The above findings support the existence of both host- and pathogen-driven evolutionary selection pressures, including the lossof-function m12^{MW97} SNP. Since NKR-P1B and NKR-P1C are polymorphic between mouse strains, we investigated whether additional m12 polymorphisms also exist in wild-derived MCMV isolates. Thus, we searched databases to compare m12 sequences from additional MCMV strains, including the Smith, K181, MW97.01, WP15B, C4A, C4B, C4C, C4D, G4, AA18d, N1, and NO7 isolates [\(Smith et al., 2013](#page-15-11)); notably, with

Figure 6. Wild-Derived Isolates of MCMV Show Host-Driven Evolution of the m12 Gene Product

(A) Phylogenetic tree of allelic m12 amino acid sequences derived from sequenced MCMV strains (Smith, K181, MW97.01, WP15B, C4A, C4B, C4C, C4D, G4, AA18d, NO7, and N1).

(B–G) Unique m12 alleles (Smith, MW97, C4A, and G4) were cloned into the pIRES2-EGFP vector, and HEK293T transfectants were used as stimulators for (B) BWZ.P1B^{B6}, (C) BWZ.P1C^{B6}, (D) BWZ.P1B¹²⁹, (E) BWZ.P1C¹²⁹, (F) BWZ.P1B^{FVB}, and (G) BWZ.P1C^{FVB} reporter cells. Data are representative of at least three independent experiments.

(H) BWZ.P1C^{FVB} reporter cells stimulated using 293T.Clr-b transfectants; cumulative mean \pm SEM from four independent experiments is shown. Significance was determined by two-way ANOVA relative to vector control.

the exception of K181 and WP15B, each of the other strains differed from the m12^{Smith} allele by at least one amino acid substitution (MW97.01, C4A/D/B), with extensive polymorphisms observed for some wild isolates (G4), including loss of signal peptide (AA18d), and premature truncations in the m04-like domain (C4C,N1,NO7) [\(Figure 6A](#page-10-0); Figure S6).

To assess these functionally, we cloned unique m12 gene products and then tested their recognition using BWZ.NKR-P1 reporters and 293T.m12 transfectant stimulators. We observed differences in recognition among all MCMV m12 alleles when tested against the different NKR-P1B and

NKR-P1C alleles (Figures $6B-6G$). The m12^{C4A} variant has an A34V substitution yet engaged all NKR-P1B alleles similarly to m12Smith [\(Figures 6](#page-10-0)B, 6D, and 6F); this was also true using NKR-P1B tetramers (Figure S3B). This result is not surprising, since the most N-terminal m12 residue visible in the structure (S39) is distal to the NKR-P1B binding site [\(Figure 5G](#page-9-0)). The m12^{MW97} variant, as observed previously, has an E91K substitution and engaged NKR-P1B^{B6} much less efficiently than m12^{Smith} in both reporter assays and using NKR-P1B tetramers, yet this difference was reduced using the NKR-P1B¹²⁹ allele and was absent using the NKR-P1B^{FVB} allele ([Figures 6B](#page-10-0), 6D, and 6F; Figure S3B). The highly diversified m12^{G4} variant was not recognized by NKR-P1 B^{B6} , interacted with NKR-P1 B^{129} similarly to m12^{MW97}, and interacted with NKR-P1B^{FVB} less well than other m12 variants [\(Figures 6B](#page-10-0), 6D, and 6F; Figure S3B).

When tested against the NKR-P1C paralogs, all m12 variants interacted with NKR-P1C B6 to some extent, although m12 G4 recognition was very weak ([Figure 6](#page-10-0)C). None of the m12 variants were found to interact with NKR-P1C¹²⁹ [\(Figure 6](#page-10-0)E), which may not be functionally expressed, due to the absence of a key cysteine residue (C122S) likely to be involved in intra-molecular stabilization. In contrast, NKR-P1C^{FVB} was stimulated strongly

(D and E) NIH 3T3 fibroblasts infected with the above MCMV variants were used as targets in ⁵¹Cr-release cytotoxicity assays with sorted (D) NKR-P1B⁺ or (E) NKR-P1B– B6-strain splenic NK-LAK effectors.

(F) Viral PFU titers in spleens of WT B6 and B6.*Nkrp1b–/–* mice infected in vivo using the above MCMV variants, analyzed 3 days post-infection. Data are pooled from two independent infections using $1x10^6$ PFU/mouse.

(legend continued on next page)

by m12 $^{\rm Smith}$ and m12 $^{\rm C4A}$ but very weakly by m12 $^{\rm MW97}$ and m12 $^{\rm G4}$ ([Figure 6G](#page-10-0)), in turn suggesting that m12^{Smith} immunoevasin function may be counterbalanced by host NKR-P1C^{FVB} activation relative to m12^{MW97} (see also below). Notably, NKR-P1B^{B6} and NKR-P1C^{FVB} only differ in a single m12 contact residue (S204T), thereby explaining their similar responses to the m12 variants [\(Figure 5G](#page-9-0)). Interestingly, Clr-b also weakly stimulated NKR-P1C^{FVB} reporters [\(Figures 6G](#page-10-0) and 6H), demonstrating that host Clr-b represents a novel activating self NKR-P1C ligand in the FVB strain. These data provide further evidence of the existence of host- and pathogen-driven evolution of the viral m12 decoy and reciprocal host adaptation of the NKR-P1B/C paralogs. These findings also validate the hypothesis that the NKR-P1B/C receptors have been co-evolving as a paired receptor clade to recognize similar host and viral ligands.

MCMV Mutants Reveal NKR-P1B-Dependent Inhibition

To establish an immunoevasin role for m12, we generated m12modified variants of the parental MCMV^{MW97} strain, including both m12-deficient (Δ m12) and m12^{Smith}-revertant (m12^{Smith}) mutants. In vitro characterization revealed that all three MCMV variants similarly downregulated host Clr-b upon infection of NIH 3T3 cells (Figure S7A).

Using infected NIH 3T3 cells directly, the m12^{Smith}-revertant virus stimulated BWZ.P1 B^{B6} reporters similarly to the original MCMV^{Smith} virus, while the Δ m12 mutant, parental MCMV^{MW97}, and mock-infected NIH 3T3 cells lacked significant stimulation (notably, BWZ.P1B^{B6} reporters routinely fail to recognize endogenous host Clr-b levels on NIH 3T3 cells; [Figure 7](#page-11-0)A). In contrast, using BWZ.P1B¹²⁹ reporters, strong stimulation was observed with MCMV^{Smith} virus, m12^{Smith}-revertant virus, and mockinfected NIH 3T3 cells, compared with weak stimulation using parental MCMV^{MW97}, and no stimulation using Δ m12 mutant MCMV ([Figure 7](#page-11-0)B). Finally, using BWZ.P1B^{FVB} reporter cells, stimulation was observed using mock-infected cells, m12^{Smith}revertant virus, MCMV^{Smith}, and MCMV^{MW97}, but not the Δ m12 mutant virus ([Figure 7](#page-11-0)C). The Clr-b-independent nature of these responses was confirmed using blocking Clr-b mAb (data not shown), as well as using mutant NIH 3T3. ACIrb cells infected with each of the MCMV strains (Figures S7B-S7D). Together, these data demonstrate that m12 is a non-redundant NKR-P1B decoy immunoevasin and that the *m12* G271A SNP (E91K polymorphism) identified between MCMV^{Smith} and MCMV^{MW97} is responsible for the differential stimulation of BWZ.P1B reporter cells.

Next, we conducted ⁵¹Cr-release cytotoxicity assays using sorted B6-strain NKR-P1B⁺ and NKR-P1B⁻ NK-LAK effectors and MCMV-infected NIH 3T3 targets. Notably, mock, $MCMV^{MW97}$, and $\Delta m12$ -infected NIH 3T3 targets were efficiently

lysed, whereas m12^{Smith}-infected targets displayed significantly lower cytotoxicity in response to NKR-P1B⁺ effectors ([Figure 7](#page-11-0)D). Importantly, inhibition by m12^{Smith} was NKR-P1B dependent, as it was not observed using NKR-P1B– NK-LAK ([Figure 7](#page-11-0)E). These data reveal that m12^{Smith}, but not m12^{MW97}, is directly capable of inhibiting B6-strain NK cytotoxicity in an NKR-P1B-dependent manner upon live MCMV infection.

To evaluate m12 immunoevasin function in vivo, the three variant MCMV^{MW97} strains were used to infect B6-strain WT, C lr-b^{-/-}, and Nkrp1b^{-/-} mice, then MCMV virulence was evaluated 3 days later using plaque assays and quantitative realtime PCR (qPCR) for viral genomic copy numbers in infected spleens. As expected, due to weak interaction between $m12^{MW97}$ and NKR-P1B^{B6}, no significant difference was observed between the parental MCMV^{MW97} and Δ m12 mutant viruses in B6 WT mice in vivo ([Figure 7F](#page-11-0)); remarkably, however, the m12^{Smith}-revertant virus displayed significantly higher splenic viral titers (PFU) and viral genomic copy numbers in B6 WT mice ([Figures 7](#page-11-0)F–7G). Importantly, the enhanced virulence of the m12^{Smith} allele in vivo was NKR-P1B dependent and Clr-b independent, as it was not observed using B6-strain Nkrp1b^{-/-} mice [\(Figures 7](#page-11-0)F-7G), yet Clr-b^{-/-} mice displayed a similar trend to WT B6 mice (data not shown). Notably, increased virulence of MCMV^{Smith} in vivo has been previously observed using WT B6 mice relative to B6.*Nkrp1b–/–* mice [\(Rahim et al.,](#page-15-12) [2016\)](#page-15-12). This finding is confirmed to be m12-dependent here using the three MCMV^{MW97}-strain variants, where the m12^{Smith}-revertant virus exhibits enhanced virulence versus parental MCMV^{MW97} and Δ m12 mutant MCMV, in WT B6, but not B6.*Nkrp1b–/–*, mice. These findings confirm a non-redundant NKR-P1B immunoevasin function for m12 in vivo.

To further elucidate the effects of the polymorphic host NKR-P1A/B/C-clade isoforms and alleles, we next performed in vivo experiments in the FVB, 129, and BALB inbred mouse strains using the three allelic MCMV variants. In the FVB strain, where strong interactions are observed between m12^{Smith} and both the inhibitory NKR-P1B^{FVB} and stimulatory NKR-P1C^{FVB} recep-tors [\(Figures 6F](#page-10-0)–6G), m12^{Smith} function may be counterbalanced by the paired receptors. However, the m12^{MW97} immunoevasin function should be intact, since a moderate interaction is observed for the inhibitory NKR-P1B^{FVB} receptor yet a much weaker interaction for the stimulatory NKR-P1C^{FVB} receptor [\(Figures 6F](#page-10-0)–6G). Interestingly, in FVB mice, no significant difference was observed between the m12^{Smith}-revertant and Δ m12 mutant viruses in vivo, while the parental MCMV^{MW97} virus maintained significantly elevated viral titers ([Figure 7H](#page-11-0)). Thus, the m12MW97 allele possesses superior immunoevasin function in the FVB strain, where the m12^{Smith} allele encounters opposing function from the activating NKR-P1C^{FVB} (and perhaps NKR-P1A^{FVB}) paralog.

⁽G) Quantitative real-time PCR analysis of MCMV ie1 genomic DNA copy numbers relative to host c-*myc* (*ie1/Myc*); all values from two independent experiments were normalized relative to values from WT B6 mice infected with parental WT virus (m12^{MW97}). Data are pooled from two independent infections using 1x10⁶ or 1x10⁵ PFU/mouse.

⁽H) Viral PFU titers in spleens of inbred FVB mice infected in vivo using the above MCMV variants and analyzed 3 days post-infection with 5x10⁵ PFU/mouse. (I–K) Viral splenic titers from (I) BALB/c, 129, (J) B6.*Ly49H–/–*, and (K) BALB.B6-Cmv1^r mice, respectively, infected in vivo using MCMV variants, analyzed 3 days post-infection with 2x10⁵ PFU/mouse. For (F–K), Mean±SEM are shown, each data point representing an independent mouse, and significance determined by one-way ANOVA (F–J) or unpaired student's t-test (K).

On the other hand, in BALB/c and 129-strain mice, small but significant differences were observed only between the parental $MCMV^{MW97}$ and $\Delta m12$ mutant viruses in vivo, with the m12^{Smith}revertant virus showing an intermediate response [\(Figure 7](#page-11-0)I). Interestingly, the published BALB/c and 129-strain NKR-P1A/ B/C alleles are identical. In keeping with this, we previously observed moderate interactions between the inhibitory NKR-P1BBALB/129 receptor and most m12 alleles ([Figure 6](#page-10-0)D), no interactions between the stimulatory NKR-P1C^{BALB/129} receptor and any m12 alleles ([Figure 6](#page-10-0)E), and a weak interaction for the stim-ulatory NKR-P1A^{BALB/129} receptor and m12^{Smith} [\(Figures 2H](#page-5-0) and [3E](#page-6-0)); thus, these in vivo results predict that the NKR-P1ABALB/129 receptor may have a higher affinity for m12^{MW97} versus m12^{Smith}, in turn resulting in lower viral titers. Notably, while NKR-P1BBALB/129 appears to be functional upon overexpression, m12^{Smith} tetramers fail to stain ex vivo NK cells from the BALB/c and 129 strains (Figure S3A), suggesting that both the NKR-P1B/C^{129/BALB} receptors may be non-functional in vivo, in turn resulting in reliance on NKR-P1A^{129/BALB} alone.

To further validate the NKR-P1B-decoy function of the m12 immunoevasin, we also performed in vivo infections using cohorts of B6.Ly49H^{-/-} mice, which lack the dominant Ly49H^{B6} activating receptor that recognizes m157, and BALB.B6Cmv1r congenic mice, which are NK1.1⁺ and retain the B6-strain *Nkrp1* genes. Here, the dominant decoy function of the m12^{Smith}-revertant significantly augmented viral PFU titers in both the B6.Ly49H^{-/-} ([Figure 7J](#page-11-0)) and BALB.B6^{Cmv1r} [\(Figure 7](#page-11-0)K) congenic host strains, relative to the Δ m12 mutant and parental m12MW97 viruses. These data confirm that the superior decoy function of the m12^{Smith} allele versus the m12^{MW97} allele is enhanced in the absence of Ly49H, segregates with the B6 strain NKR-P1 (*Klrb1*) region, and is unaffected by epistatic effects from the BALB genomic background.

Taken together, the above findings demonstrate that the immunoevasin function of m12 in vivo is both host and virus strain dependent, and it can be functionally counterbalanced by both inhibitory and stimulatory NKR-P1A/B/C-clade polymorphisms and paralogs. These findings further confirm that complex interactions between host NKR and MCMV immunoevasins ultimately determine the outcome of host-pathogen interactions in vivo.

DISCUSSION

The NK1.1 (NKR-P1; *Klrb1*) receptor family comprises both activating and inhibitory members, with a bona fide ligand of the activating NK1.1 antigen remaining unknown, despite being described 40 years ago [\(Glimcher et al., 1977\)](#page-15-3). We have identified m12 as a natural (viral) ligand for the prototypical NK1.1 antigen, the stimulatory NKR-P1C^{B6} orphan receptor [\(Glimcher](#page-15-3) [et al., 1977; Koo and Peppard, 1984; Ryan et al., 1992\)](#page-15-3). In addition, host Clr-b represents a second (self) NKR-P1C ligand in the FVB mouse strain. Moreover, we show that m12 functions as a decoy immunoevasin to subvert NK cell-mediated cytotoxicity by direct interaction with the inhibitory NKR-P1B receptor, which normally surveys infected cells for Clr-b downregulation during infection. The identification of the distantly Ig-related m02-family member, m12, as an NKR-P1A/B/C-clade ligand demonstrates sophisticated viral utilization of this protein fold to directly engage an NKR. Structurally, m12 clamped around the NKR-P1B receptor, in a polar claw mechanism, enveloping an extensive area of NKR-P1B that simultaneously provided insight into the specific targeting of the receptor by m12 and how allelic differences in this axis impacted upon recognition. Here, a number of the allelic polymorphisms in both the m12 immunoevasin and NKR-P1 receptors either mapped to the binding site or were in close proximity to it, thereby implying that both direct and indirect effects can impact the m12-NKR-P1 recognition axis.

The m12 footprint overlapped extensively with that observed for binding of the endogenous self ligand, KACL, to the closely related human NKp65 receptor [\(Li et al., 2013](#page-15-10)). Indeed, the utilization of existing binding sites on host receptors/ligands appears to be a common viral immune evasion strategy. Nonetheless, such structural mimicry does not necessarily underpin immunoevasin strategies, as structurally diverse immunoevasins, such as HCMV UL16 and MCMV m152, map to the same sites on ligands for the human and mouse activating NKG2D receptors, respectively (Mü[ller et al., 2010; Wang et al., 2012\)](#page-15-13). Further, some viral immunoevasins, including m157, can adopt unusual and unanticipated binding modes ([Berry et al., 2013\)](#page-14-4). Our work shows how the structurally divergent, viral Ig-like m12 immunoevasin utilizes binding mode mimicry to engage the inhibitory NKR-P1B receptor, which then underpins the ability of m12 to be reciprocally targeted by the structurally related activating NKR-P1A/C family members, in turn countering the m12 immunoregulatory properties.

The rodent *Klrb1* genes display considerable polymorphism, being subdivided into two antigenic and phylogenetic clusters in mice and rats: the NKR-P1A/B/C and NKR-P1F/G clades ([Car](#page-15-2)[lyle et al., 2008; Kirkham and Carlyle, 2014; Kveberg et al., 2009\)](#page-15-2). Strikingly, the NKR-P1B and NKR-P1C receptors appear to have undergone parallel divergence characteristic of paired recognition receptors, leading to the hypothesis that they are under selective pressure, within the constraints imposed by both endogenous self ligands and foreign viral ligands. Here, we provide compelling evidence of co-evolution at the host-pathogen interface, with host adaptation of the NKR-P1A and NKR-P1C (NK1.1) stimulatory receptor paralogs to directly recognize the m12 decoy, as well as the existence of wild-derived viral m12 immunoevasin alleles, including loss-of-function alleles, which may have evolved to avert direct NKR-P1A/C-mediated NK cell activation. We also provide strong evidence that m12 is a nonredundant NKR-P1B-targeting immunoevasin in vivo. However, the functional contribution of m12 to the outcome of host-pathogen interactions in vivo is ultimately determined by both the viral m12 and host NKR-P1A/B/C-clade polymorphisms.

For example, each m12 allele may contribute differentially to viral fitness in distinct mouse strains, and what may appear to be a superior immunoevasin in one mouse strain may be counterbalanced by opposing signals from both inhibitory NKR-P1B and activating NKR-P1A/C paralogs in another mouse strain. Our in-depth analyses employing eight inbred, mutant, or congenic mouse strains and three MCMV variants underscore how viral versus host fitness can be significantly impacted by polymorphisms in both the paired host receptors and viral immunoevasin.

The conserved use of decoy immunoevasins by MCMV (m12) and RCMV (RCTL) to subvert NKR-P1B-mediated NK cell immunosurveillance of pathological target cells highlights the role of the NKR-P1:Clr recognition axis in immunity to infectious disease and suggests that similar mechanisms may operate during HCMV infection of human cells ([Aguilar et al., 2015;](#page-14-1) [Voigt et al., 2007](#page-14-1)). Notably, the human NKR-P1A:LLT1 receptor-ligand pair are likely orthologous to some rodent NKR-P1:Clr interactions. NKR-P1A (CD161) is inhibitory and is expressed on $\sim 60\%$ of human NK cells [\(Lanier et al., 1994](#page-15-14)), resembling NKR-P1B in function and expression. LLT1 has more restricted expression than Clr-b and is induced in response to PRR agonists and infection [\(Aguilar et al., 2015;](#page-14-1) [Germain et al., 2011; Rosen et al., 2008](#page-14-1)). It remains to be determined whether HCMV may similarly target the human NKR-P1A receptor.

The NK1.1 antigen was the prototypical receptor identified to be selectively expressed by NK cells. Thus, while being used as a tool to study and purify NK cells for almost four decades, the physiological ligand for the activating NK1.1 antigen has remained a mystery. Here, we showcase m12 as a natural virally encoded ligand for this important NKR family. Moreover, NK1.1 is also found on some ILC and various T cell subsets, including invariant NKT cells, the ubiquitous mucosalassociated invariant T (MAIT) cells, CD4⁺ T helper cells, and activated CD8⁺ T cells. Thus, targeting of the NK1.1 antigen by a virally encoded ligand has important implications for both innate and adaptive immunity, virus-host co-evolution, and immunotherapy.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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Supplemental Information includes seven figures and five tables and can be found with this article online at <http://dx.doi.org/10.1016/j.cell.2017.03.002>.

O.A.A. designed, performed, and analyzed experiments and contributed to the manuscript writing. R.B. and Z.F. generated the purified m12 and NKR-P1 proteins and performed SPR analyses. R.B. solved, refined, and analyzed the structure and contributed to manuscript writing. M.M.A.R., M.T., B.P., M.M.T., and A.B.M. assisted with in vivo experiments. J.J.R. generated the m12 mutant and revertant viruses. T.N.H.L assisted with bioinformatic analysis. G.R.B. assisted with the structural analysis. C.L.K, A.M., B.P., and A.K. assisted with in vitro experiments. D.S.J.A., A.P.M., S.J., J.R., and J.R.C. contributed to the direction of the study. This work was done in the labs of A.P.M., S.J., J.R., and J.R.C. J.R.C. contributed to the design and interpretation of experiments, project management, and manuscript writing.

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STAR+METHODS

KEY RESOURCES TABLE

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Cell

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Animals

C57BL/6 (B6) mice were purchase from the Jackson Laboratory, FVB/N (FVB) and 129S6 (129) mice were purchased from Charles River. We have previously described the B6.Nkrp1b^{-/-} and B6.Clrb^{-/-} mice [\(Chen et al., 2015; Rahim et al., 2015\)](#page-15-5). B6.Ly49h^{-/-} and Balb.B6-Cmv1r mice have previously been described ([Fodil-Cornu et al., 2008; Scalzo et al., 1995](#page-15-16)). All animals were maintained according to approved protocols at the respective institutes: Sunnybrook Research Institute, University of Ottawa, and the University of Rijeka.

MCMV (Smith strain), MCMV-GFP, and MW97.01 (BAC-generated strain) have all been previously described ([Aguilar et al., 2015\)](#page-14-1). MCMV^{Smith} and MCMV-GFP were provided by Dr. Silvia Vidal (McGill University). The MCMV^{K181} strain was provided by Dr. Michael G. Brown (University of Virginia). The Δ m12 mutant virus and m12^{Smith}-revertant viruses were constructed using BAC technology that has previously been described [\(Wagner and Koszinowski, 2004](#page-15-18)).

Cells

UPITE
NIH 3T3, YAC-1, and HEK293T cells were purchased from the American Type and Culture Collection (ATCC). BWZ.36 cells were obtained from N. Shastri (UC Berkeley, CA, USA). B6 mouse embryonic fibroblasts (MEF) were obtained from T.W. Mak (University of Toronto, Canada). Cells were cultured in complete DMEM-HG, supplemented with 2 mM glutamine, 100 U/mL penicillin, 100 µg/ml streptomycin, 50 μg/mL gentamicin, 110 μg/mL sodium pyruvate, 50 μM 2-mercaptoethanol, 10 mM HEPES, and 10% FCS. Cells were maintained in incubator at 37° C, 5% CO₂.

Sf9 and Hi5 cells were cultured in Insect-XPRESS medium (Lonza) at 27°C and maintained with shaking at 120 rpm. HEK293S GnTi- adherent cells were cultured in DMEM supplemented with 10% FBS and 1% Pen/Strep/Glutamine (GIBCO) at 37°C in the presence of 5% CO2. For protein expression, FBS was reduced to 2%.

METHOD DETAILS

Virus infections

All viruses were passaged using MEF cells, and purified according to described protocols ([Aguilar et al., 2015](#page-14-1)). For in vitro infections, mouse fibroblasts were infected with MCMV at an MOI of 0.5 PFU/cell, centrifuged at 800xg for 30 min, and incubated at 37°C. For in vivo infections, mice were infected intraperitoneally with 0.2 -1x10⁶ PFU and sacrificed on day 3, tissues harvested, and the viral titers were assessed using plaque-forming assays ([Aguilar et al., 2015](#page-14-1)).

Flow cytometry and antibodies Flow cytometry was performed as previously described ([Chen et al., 2015](#page-15-5)). Clr-b mAb (4A6) has previously been described [\(Carlyle](#page-14-0) [et al., 2004](#page-14-0)). The NKR-P1B^{B6} mAb and hybridoma (2D12) were a kind gift from Drs. Koho Izuka and Wayne Yokoyama (lizuka et al., [2003\)](#page-15-15). All other mAb clones were purchased from eBioscience or Sigma-Aldrich and are has follows: NK1.1 (PK136); NKp46 (29A1.4); CD3ε (145-C11); FcγRII/III (2.4G2); CD49b/DX5 (clone DX5); γδTCR (clone GL3); FLAG (M2); streptavidin, SA-PE/APC (Thermo Fisher Scientific). The CD1d-tetramer loaded with PBS-57 (aGalCer analog) was a kind gift from Dr. Thierry Mallevaey. For tetramerization, purified biotinylated monomers (m12^{Smith}, NKR-P1B^{B6,129,FVB}) were incubated with SA-PE in a 4:1 molar ratio, aliquoted in 1/10th volumes over a 3 hr period at 4° C.

Cells were stained in flow buffer (HBSS, 0.5% BSA and 0.03% NaN₃) on ice with primary monoclonal antibodies for 25-30 min, or secondary streptavidin conjugates for 15-20 min, washed between incubations and then analyzed using a FACSCalibur or FACSCanto II flow cytometer, or FACS sorted using a FACSAria (BD Biosciences). For staining of ex vivo splenocytes or NKp46⁺CD3⁻ lymphokine-activated killer (NK-LAK) effectors, cells were pre-incubated with 2.4G2 mAb for 10 min on ice prior to staining with antibodies. Cells were stained with tetramers by incubating on ice for 1 hr. Cells were analyzed by live gating for FSC/SSC, doublet excluded, exclusion of propidium iodide or DAPI for viability, and then according to cell surface markers. Flow cytometry files were analyzed using FlowJo software.

The ectodomains of the mouse NKR-P1 genes were PCR amplified from day 6 LAK cDNA (see Table S5 for primers) and cloned into a type-II MSCV vector to make chimeric receptors with intracellular CD3^r domains. MCMV m12/CD3^r chimeric reporters were constructed by cloning the ectodomain of m12 into a type-I MSCV vector ([Mesci and Carlyle, 2007](#page-15-7)). Retroviruses produced with these constructs were used to transduce BWZ.36 cells, and then sorted for IRES-GFP⁺ expression. All reporters were sorted to have

matched GFP expression. Reporter assays were done by co-culturing reporter cells (5x10⁴) with serially diluted amounts of stimulators in 96-well plates. Stimulators used were either MCMV-infected, mock-infected, or cells transfected with plasmids using Lipofectamine²⁰⁰⁰ (Thermo Fisher Scientific). Positive control cells were stimulated with 10 ng/mL PMA plus 0.5 μ M ionomycin. These cells were incubated overnight, washed with PBS, then resuspended in 150 µL of 1X CPRG buffer (90 mg/L chlorophenol-red-ß-Dgalactopyranoside (Roche), 9 mM MgCl2, 0.1% NP-40, in PBS), incubated at room temperature, then analyzed using a Varioskan microplate reader (Thermo Fisher Scientific), using OD 595-655.

^{- .} Cnromium release cytotoxicity assays
Splenic lymphokine-activated killer (LAK) effector cells were generated by harvesting spleens from mice, processing into single cell suspensions, ACK lysed (Thermo Fisher Scientific), then grown in 10% complete RPMI 1640 media supplemented with 2500 U/mL human rIL-2 (Proleukin; Novartis). On day 4, these cells were FACS sorted for NKp46⁺CD3⁻ NK cells subsetted based upon NKR-P1BB6/FVB (2D12 or NK1.1 mAb) expression, and used as effectors in ⁵¹Cr-release assays on day 6 or 7. Target cells were labeled with 50 μ Ci Na $_2$ ⁵¹CrO $_4$ (Perkin Elmer) in FCS for 1 hr at 37°C, washed, plated in 96-well V-bottom plates in serial dilutions with effector cells at corresponding ratios, and incubated at 37°C for 4 hr. Supernatants (100 μ l) were then transferred to scintillation plates (LumaPlate-96; Perkin Elmer), dried overnight, and counted using a Top Count NXT microplate Scintillation Counter (Packard Instrument Company). Percent specific lysis values were calculated relative to standard maximum release (2% Triton X-100) and spontaneous release (media) values.

Generation of NIH 3T3.ACIr-b cells using CRISPR-Cas9 technology

The pX458 vector was utilized in accordance with Dr. Zhang's protocol for generating targeting vectors ([www.genome-engineering.](http://www.genome-engineering.org/crispr/) [org/crispr/](http://www.genome-engineering.org/crispr/)**).** The sequence of oligos used for constructing *Clec2d* targeting vector were: sense: 5'-cac cgC TAC CTA TGC TTA GTC CCA C-3'; anti-sense: 5'-aaa cGT GGG ACT AAG CAT AGG TAG c-3'. This vector was transfected into NIH 3T3 cells that were then sorted for GFP⁺Clr-b⁻ expression. These cells were then subcloned using limiting dilution. Appropriate clones were selected lacking Clr-b cell surface staining and BWZ.NKR-P1B reporter stimulation.

recent expression and partications
Codon optimized genes encoding NKR-P1 receptors and m12 were synthesized by Integrated DNA Technologies. For NKR-P1B^{B6}, the unpaired cysteine residue at position 118 was replaced with a glycine. The ectodomains (residues 89-215) of NKR-P1B^{B6,129,FVB} and NKR-P1C^{B6} were cloned into a modified pHLSec vector upstream of a biotinylation site and a His tag (for SPR) or a His tag (for crystallization), and expressed via transient transfection in HEK293S cells as described previously ([Berry et al., 2013; Berry et al.,](#page-14-4) [2014\)](#page-14-4). For a single 6000 cm³ roller bottle, 600 µg plasmid DNA was mixed with 900 µL PEI (1 mg/ml) and 50 mL serum-free DMEM media. This mixture was added to HEK293S GnTi- cells after a 10 min incubation, followed by the subsequent addition of a further 200 mL DMEM and FCS to give a final concentration of 2%. Media was harvested after 3 and 6 days. DNA encoding m12^{Smith} (residues 29-208) was ligated into the pFastbac vector (ThermoFisher Scientific) upstream of a 6x Histidine tag and used to generate recombinant baculovirus according to the manufacturer's instructions. Specifically, the plasmid was transformed into DH10Bac *E.coli* cells in order to produce bacmid DNA. Recombinant baculoviruses were generated by transfection of Sf9 cells using Cellfectin 2. Protein was expressed in Hi5 insect cells by the addition of 2% P3 virus, and harvested after 48-72h. Cell culture media containing secreted m12 or NKR-P1 receptors was concentrated by tangential flow filtration and purified using nickel affinity and size exclusion chromatography using HisTrap and Superdex 200 columns 16/60 (GE Healthcare) respectively. For biotinylation, purified NKR-P1 receptors were buffer exchanged into 10 mM Tris and biotinylated overnight at 20°C prior to removal of free biotin via size exclusion chromatography. For SPR experiments, m12 protein was used immediately following purification. However, samples stored at 4°C at high concentration were prone to proteolytic degradation, resulting in a loss of \sim 4 kDa of mass from the carboxy-terminus, as judged by SDS-PAGE analysis. This truncated form of m12 could be selectively purified from the intact molecule by nickel affinity chromatography and bound to NKR-P1 receptors with the same affinity as the entire m12 ectodomain (data not shown). Accordingly, this truncated form of m12 was used for crystallization experiments. The ectodomains of Ly49A $B⁶$ and m04^{Smith} were produced in HEK293SGnTi- cells as described for NKR-P1 above and previously [\(Berry et al., 2013; Berry](#page-14-4) [et al., 2014](#page-14-4)).

SPR experiments were performed at 25°C on a Biacore 3000 using a running buffer consisting of 10 mM Tris pH8 containing 150 mM NaCl and 0.005% P20. Approximately 800 response units of biotinylated NKR-P1/Ly49 receptors were coupled on to streptavidin-coated chips according to the manufacturer's instructions (GE Healthcare). Remaining free streptavidin sites were blocked with D-biotin. m12 (0.025-50 μ M) or m04 (100 μ M) was injected over the immobilized receptors at a flow rate of 20 μ /min and the final response was calculated by subtraction of the response of an 'empty' flow cell (containing biotin blocked streptavidin). For blocking experiments, 50 µM m12^{Smith} was passed over immobilized NKR-P1B^{B6} and NKR-P1C^{B6} both before and after injection of 30 mL of 10 mg/ml NK1.1 (clone PK136-StemCell Technologies, Inc.). Equilibrium data were analyzed in GraphPad Prism.

For crystallization, the m12^{Smith} and NKR-P1B^{B6} ectodomains were mixed in a 1:1 molar ratio at a final protein concentration of 6.5 mg/ml. Crystals were obtained by the hanging drop vapor-diffusion method from a solution containing 41% PEG 400, 0.1 M Bis/Tris pH 6.3 and 0.4 M sodium iodide at 20°C. Crystals were flash cooled using liquid nitrogen and X-ray diffraction data were recorded on a Quantum-315 CCD detector at the MX2 beamline of the Australian Synchrotron. Data were integrated using iMOSFLM and scaled using SCALA from the CCP4 program suite. Details of the data processing statistics are given in Table S3.

Structure determination and refinement The structure was solved by molecular replacement using the Phaser program. The initial search model comprised a truncated form of the NKR-P1A CTLD (PDB ID: 3M9Z) that lacked the extended loop located between the β 3 and β 4 strands. The resulting unbiased electron density was not sufficient to build m12 de novo. However, molecular replacement using an extensively truncated fragment of m04 (PDB ID: 4PN6) comprising only the I, D, E' and F strands improved the quality of the electron density map to an extent that allowed the remainder of the m12 molecule to be built. The structure was refined using Buster. Model building was performed in Coot, and figures were prepared with Pymol [\(http://www.pymol.org](http://www.pymol.org)) and APBS. Details of the refinement statistics are provided in Table S4.

DNA/RNA Isolation, cDNA synthesis and PCR cloning Genomic DNA was isolated using Easy-DNA kit (Thermo Fisher Scientific). RNA was isolated using Total RNA isolation kit with gDNA removal (Norgen Biotek) and reverse transcription reactions were done using Superscript III cDNA synthesis kit (Thermo Fisher Scientific). Quantitative RT-PCR was performed on a CFX-96 PCR Detection System (Bio-Rad) using 20-50 ng of cDNA, and PerfeCTa qPCR mix (Quanta Biosciences). The primers used can be found in Table S5.

PCR cloning of m02 and m145 family members was done using gene-specific primers with appropriate restriction enzyme sites, cDNA from MCMV-GFP infected NIH 3T3, amplified with Q5 high-fidelity PCR system (New England Biolabs), and cloned into the pIRES2-EGFP vector (Clontech). The summary of primers can be found in Table S5. The m12^{C4A} gene was synthesized by conducting in situ mutagenesis using primers found in Table S5. The m12^{G4} gene was synthesized using GeneArt services from Thermo Fisher Scientific (Burlington, ON). Cloning of the Nkrp1 genes was mediated by PCR using gene specific primers (Table S5).

N-terminally FLAG-tagged constructs of m12 were generated using primers in (Table S5) and cloned into pIRES2-EGFP vector. These constructs were transfected into either HEK293T or NIH 3T3 cells and immunoprecipitations were done using anti-FLAG mAb bound to Protein A/G agarose beads in 1% NP-40 buffer. The isolates were run on SDS-PAGE and transferred to Immobilon PVDF membrane (EMD Millipore) and blotted for anti-FLAG-HRP conjugated antibody (Cell Signal Technologies) and developed with Immobilon ECL reagent. Blots were imaged using a MicroChemi 4.2.

RNA-Sequencing and Analysis Total RNA from either mock-infected, MCMV-GFP, or MW97.01-infected NIH 3T3 cells using mirVana kit (Thermo Fisher Scientific). These samples were quality checked using an Agilent Bioanalyzer 2100 and confirmed to have RNA integrity values of > 9 , then used to create a cDNA library for SOLiD RNA-Sequencing. For analysis of viral transcripts, raw data was mapped onto the MCMV^{Smith} reference genome (accession number GU305914) using TopHat algorithm and Cufflinks was applied to report RPKM values. Mapped reads were visualized using the Integrative Genomics Viewer (IGV version 2.3.34, Broad Institute, [https://www.broadinstitute.org/igv/\)](https://www.broadinstitute.org/igv/).

Mapped RNA-Seq reads were loaded visualized on the IGV Genome Browser using the MCMV^{Smith} reference genome (accession number GU305914). Membrane topology was determined using the TMpred server ([http://www.ch.embnet.org/software/](http://www.ch.embnet.org/software/TMPRED_form.html) [TMPRED_form.html](http://www.ch.embnet.org/software/TMPRED_form.html)). The SignalP 4.1 Server [\(http://www.cbs.dtu.dk/services/SignalP/\)](http://www.cbs.dtu.dk/services/SignalP/) was used for identification of predicted signal peptides. Detection of ORF start site was achieved using the NetStart 1.0 Prediction Server website ([http://www.cbs.dtu.](http://www.cbs.dtu.dk/services/NetStart/) [dk/services/NetStart/\)](http://www.cbs.dtu.dk/services/NetStart/).

QUANTITATION AND STATISTICAL ANALYSIS QUANTITATION AND STATISTICAL ANALYSIS

Data were analyzed using GraphPad Prism 7, using either a paired Student's two-tailed t test, or one-way/two-way ANOVA, where applicable. See figure legends for details. Statistical significance is defined as: $* p < 0.05; ** p < 0.01; ** p < 0.001$.

DATA AND SOFTWARE AVAILABILITY

Coordinates for the m12:NKR-P1B structure have been deposited in the protein data bank (PDB: 5TZN). The accession numbers for the FVB-strain sequences, Nkrp1 splice variants, and m12 variants reported in this paper are GenBank: KX443605–GenBank: KX443630.

Supplemental Figures

Figure S1. MCMV Encodes a Clr-b and β_2 m-Independent NKR-P1B Decoy Ligand, Related to [Figure 1](#page-3-0)

(A–C) Adult ear fibroblasts (AEF) from (A) WT B6, (B) B6.*Clr-b–/–*, and (C) B6.*Clr-b–/–*b2m–/– mice were infected with MCMVSmith and used as stimulators for BWZ.NKR-P1B¹²⁹ reporter cells in the absence or presence of blocking Clr-b mAb. Mean±SEM are shown in (A) and (B). Data representative of at least 2 independent experiments.

Figure S2. The MCMV m12 Gene Product Is a Glycosylated Type-I Transmembrane Protein Capable of Interacting with Numerous NKR-P1B/C Isoforms, Related to [Figures 1](#page-3-0) and [2](#page-5-0)

(A) Structural diagram of the m12 gene with 2 translational start sites (ATG1/2) and TGA stop codon labeled (top) and domain structure of the m12 protein (bottom). SP, signal peptide; EC, extracellular m04-like domain; TM, transmembrane domain; IC, intracellular domain.

(B) Cell surface expression of native and N-terminally FLAG-tagged m12 (NSP, native signal peptide; PSP, preprotrypsin signal peptide) constructs cloned into pIRES2-EGFP and transfected into HEK293T.

(C) Stimulation of BWZ.NKR-P1B^{B6} cells by different m12 constructs.

(D) Immunoprecipitation of m12ATG2/PSP-FLAG transfected into HEK293T cells, isolated under different conditions, and immunoblotted for FLAG expression; a dashed line indicates the merged white light image (molecular weight marker lane) with the ECL image (Western blot lanes). Data are representative of at least 3 independent experiments.

(E) Nkrp1 splice isoforms were cloned into pIRES2-EGFP using cDNA from B6, 129, and FVB splenic LAK, transfected into HEK293T cells, and used as stimulators for BWZ.m12Smith reporters.

(F) Intracellular (IC) and extracellular (EC) domain swaps of NKR-P1B^{B6} and NKR-P1C^{B6} were constructed and tested for their ability to stimulate BWZ.m12^{Smith} reporters upon transfection into HEK293T. Data are representative of at least 2 independent experiments.

Figure S3. Direct NKR-P1 and m12 Interactions Revealed by Tetramer Staining of Ex Vivo Immune Cell Subsets, Transfectants, and MCMV-Infected Cells, Related to [Figure 3](#page-6-0)

(A) m12^{Smith} tetramers were used to stain ex vivo splenic immune cell subsets from distinct mouse strains. Numbers above gates represent percentage of gated positive cells; numbers at top right corner represent MFI of positive cells for 2D12 and m12-tetramer staining to highlight a ~2-fold increase in NKR-P1B levels on

(B) NKR-P1B tetramer staining of HEK293T cells transfected with m12 variants. Numbers represent mean fluorescence intensity.

(C) NKR-P1B tetramer staining of MCMV-infected NIH 3T3 cells. Numbers represent percentage of gated positive cells.

B6.Clr-b^{-/-} NK cells relative to WT B6 NK cells (see [Figure 3](#page-6-0)F). Gated positive cells (shaded histograms) are shown compared to the corresponding B6 mouse FMO negative control samples (dashed lines).

Figure S4. Structure of NKR-P1B and m12 and Comparison to Known Structures and Docking Modes, Related to [Figure 4](#page-8-0)

(A) Comparison of the NKR-P1B C-type lectin-like domain with that of CLEC5A (PDB ID 2YHF) and NKR-P1A (PDB ID 3M9Z).

(B) Overview of the m12 ectodomain in cartoon representation. b-strands are colored blue; a-helices are red; and loop regions are shown in black. Cysteine residues that form intra-molecular disulfide bonds are shown as green sticks. The refined 2F*o-*F*^c* electron density map for the attached sugar moieties (blue sticks) is shown as a blue mesh and contoured at 3σ .

(C) Alignment of the m12^{Smith} (blue) and m04 G4 ectodomains (cyan) (PDB ID 4PN6).</sup>

(D) Schematic representation of the m12 ectodomain fold, colored as in (B). Amino acid residues involved in disulfide bonding and sugar attachment are labeled in green and black respectively.

(E) Side-by-side comparison of NKR-P1B (salmon) bound to m12 (blue) and NKp65 (red) bound to KACL (light blue) (PDB ID 4IOP). Structures are shown in the same orientation with respect to the receptor.

(F) Top view showing the footprint made by each ligand on its corresponding receptor. The molecular surface of the receptors is colored gray, with the residues that contact the ligand colored blue.

Figure S5. Generation and Characterization of YAC-1 Transductants, Related to [Figure 5](#page-9-0)

(A and B) YAC-1 cells were transduced with pMSCV2.2-IRES-EGFP retrovirus encoding m12^{Smith}, m12^{MW97}, Clr-b, or parental empty vector and sorted for matching GFP levels. Stable YAC-1 transductants were (A) analyzed by flow cytometry for Clr-b expression and (B) used as stimulator cells for BWZ.NKR-P1B^{FVB} reporter cells.

(C and D) Stable YAC-1 transductants were used as target cells in ⁵¹Cr-release cytotoxicity assays for (C) NKR-P1B⁺ or (D) NKR-P1B⁻ FVB-strain splenic NK-LAK effector cells. Mean±SEM are shown. Data are representative of at least 2 independent experiments.

Figure S6. Protein Alignments of m12 Variants from Sequenced MCMV Genomes, Related to [Figure 6](#page-10-0)

Amino acid sequences of m12 from laboratory MCMV strains (Smith, K181, and MW97.01) and wild-derived MCMV variants (WP15B, C4A, C4B, C4C, C4D, G4, AA18d, N1, and NO7) were aligned using ClustalW. The protein domains and secondary structure elements are labeled above the amino acid sequences; TM, transmembrane domain; IC, intracellular domain; blue arrows, B-sheets; red rectangles, α -helices. Dashed lines represent regions not visible in the crystal structure. m12 residues that make contacts to NKR-P1B^{B6} are highlighted with a pink background.

Figure S7. Parental MW97-Derived m12 Variants Retain Ability to Downregulate Clr-b and Differentially Stimulate NKR-P1B Reporters, Related to [Figure 7](#page-11-0)

(A–D) NIH 3T3 cells were infected with MCMV m12 variants (MOI = 0.5) and assessed for cell surface Clr-b expression 24 hr post-infection by flow cytometry. Numbers correspond to Clr-b MFI; shaded histograms, Clr-b staining; dashed lines, secondary reagent alone. NIH 3T3.DClr-b fibroblasts were infected with MCMV variants and used as stimulators for (B) BWZ.NKR-P1B^{B6}, (C) BWZ.NKR-P1B¹²⁹, and (D) BWZ.NKR-P1B^{FVB} reporter cells.