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Suppression of co-stimulation by human cytomegalovirus promotes evasion of cellular immune defenses

Short Title: HCMV regulates CD58 to control T and NK cells

Eddie C.Y. Wang†, Mariana Pjechova1,2, Katie Nightingale3, Virginia-Maria Vlahava1, Mihil Patel1, Eva Ruckova1,2, Simone Forbes1, Luis Nobre3, Robin Antrobus3, Dawn Roberts1, Ceri A. Fielding1, Sepehr Seirafian1, James Davies1, Isa Murrell1, Betty Lau4, Gavin S. Wilkie4, Nicolás M. Suárez4, Richard J. Stanton1, Borivoj Vojtesek2, Andrew Davison4, Paul J. Lehner3, Michael P. Weekes3, Gavin W.G. Wilkinson1*, Peter Tomasec††

1Cardiff University School of Medicine, Division of Infection and Immunity, Henry Wellcome Building, Heath Park, Cardiff CF14 4XN, UK
2Regional Centre for Applied Molecular Oncology (RECAMO), Masaryk Memorial Cancer Institute, Zluty Kopec 7, 65653 Brno, Czech Republic
3Cambridge Institute for Medical Research, University of Cambridge, Hills Road, Cambridge CB2 0XY, UK
4MRC-University of Glasgow Centre for Virus Research, Sir Michael Stoker Building, 464 Bearsden Road, Glasgow G61 1QH, UK

*these authors contributed equally to this study

Corresponding Author:
Eddie C. Y. Wang, Cardiff University School of Medicine, Division of Infection and Immunity, Henry Wellcome Building 1F-07, Heath Park, Cardiff CF14 4XN, UK; +44-2920687035; WangEC@cf.ac.uk

† deceased
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Abstract
CD58 is an adhesion molecule that is known to play a critical role in co-stimulation of effector cells and is intrinsic to immune synapse structure. Herein, we describe the first virally-encoded gene that inhibits CD58 surface expression. Human cytomegalovirus (HCMV) UL148 was necessary and sufficient to promote intracellular retention of CD58 during HCMV infection. Blocking studies with antagonistic anti-CD58 mAb and an HCMV UL148 deletion mutant (HCMVΔUL148) with restored CD58 expression demonstrated that the CD2/CD58 axis was essential for the recognition of HCMV-infected targets by CD8+ HCMV-specific cytotoxic T lymphocytes (CTLs). Further, challenge of peripheral blood mononuclear cells ex vivo with HCMVΔUL148 increased both CTL and natural killer (NK) cell degranulation against HCMV-infected cells, including NK-driven antibody-dependent cellular cytotoxicity, showing that UL148 is a modulator of the function of multiple effector cell subsets. Our data stress the impact of HCMV immune evasion functions on shaping the immune response, highlighting the capacity for their potential use in modulating immunity during the development of anti-HCMV vaccines and HCMV-based vaccine vectors.

Significance statement
HCMV is the major infectious cause of developmental disorders in babies due to its capacity to cross the placenta. HCMV is also a major pathogen in transplant recipients and HIV-AIDS patients. Despite inducing the strongest immune responses observed for any human pathogen, HCMV evades host defenses and persists for life. We report a novel viral stealth strategy. HCMV UL148 reduces surface expression of a key cell adhesion molecule (CD58), impairing the ability of NK and T cells to be activated by HCMV-infected cells. This is the first description of a viral gene targeting this pathway. Our findings highlight a role for CD58 in recognition of HCMV-infected cells and may be relevant for development of future antiviral therapies.
Introduction

Human cytomegalovirus (HCMV; species Human betaherpesvirus 5), is the major viral cause of congenital birth defects and an important pathogen capable of causing severe disease in immunocompromised and immune-naïve individuals. HCMV is noted for inducing the most potent cellular immune responses observed for any human pathogen. These responses, including expansions of cytotoxic T-lymphocytes (CTLs) and natural killer (NK) cells of a specific phenotype, are large and are maintained for life (reviewed in (1, 2)). HCMV, however, is not cleared by the host following primary infection. The mechanisms that underpin the ability of the virus to endure in the presence of such immunity has been the target of intense study, with the hope that the knowledge gained will inform the generation of anti-HCMV vaccines and also vaccine design, for which the maintenance of induced immune responses is paramount. Indeed, HCMV is being pursued as a vaccine vector in its own right as redirection of anti-CMV immunity can clear pathogens otherwise capable of persisting in their host (3, 4). The study of HCMV-encoded genes and proteins has revealed many strategies designed to avoid innate and adaptive immunity, which have defined a number of basic immune pathways essential to CTL and NK activity. These include at least four functions that inhibit HLA-I expression and ten that impair NK cell activation (reviewed in (5, 6)).

CTLs and NK cells use the supramolecular adhesion complex (SMAC) at the immune synapse (IS) to interact with their targets. The SMAC is a tightly packed intercellular complex of adhesion molecules and receptor/ligand pairs where antigen presentation and signaling take place, which regulate the secretion of cytotoxic granules and cytokines from the effector cells (7-9). There are many descriptions of HCMV acting to prevent expression of activating ligands and promote the expression of inhibitory receptors on the target cell surface, but reports of immune evasion mechanisms able to impede formation of IS structure have been limited to remodeling of the target cell actin cytoskeleton (10).

CD58 (LFA-3) on target cells acts to promote cell-to-cell adhesion and IS formation, and to provide a co-stimulatory signal through its receptor, CD2, on effectors (11-23). Recent studies have highlighted the importance of CD2 engagement for co-stimulation of CD4+ T-cells in HCMV infection (24) and adaptive NK cells (25, 26), and have identified the CD2/CD58 axis as the primary co-stimulatory pathway for CD28-CD8+ CTLs (19, 27, 28). Cell surface expression of CD58 has been reported to be either upregulated (29, 30) or downregulated (31) by HCMV infection depending on the HCMV strain used. The aim of
our study was to dissect this phenomenon, thereby determining the functional relevance of regulating CD58 expression on HCMV-infected cells. We describe the identification of the first viral-encoded gene responsible for downregulating CD58 and detail the broad impact of this novel function on both CTL and NK cell recognition in the context of HCMV infection.
Results

HCMV UL148 is a novel viral function down-regulating cell surface expression of CD58

HCMV strain AD169 has previously been reported to upregulate CD58 (29, 30), whereas proteomic analysis with low passage HCMV strain Merlin has suggested the opposite (31). We confirmed these effects directly by flow cytometry (Fig.1). AD169 has suffered a spontaneous deletion in its genome during in vitro culture, involving a 15kb sequence designated the UL/b’ region (32). We therefore investigated whether the function responsible for CD58 downregulation resided within UL/b’. Surface expression of CD58 was analyzed in cells infected with a complete library of HCMV strain Merlin UL/b’ single gene deletion mutants (loss of function screen, Fig.2A) and an adenovirus vector library over-expressing each UL/b’ gene individually (gain of function screen, Fig.2B, Fig.S1). Both screens identified HCMV UL148 as the gene responsible, with loss of UL148 from HCMV strain Merlin (referred to as HCMV∆UL148) (Fig.S2) resulting in CD58 upregulation (Fig.2A, C), and ectopic expression of UL148 resulting in CD58 downregulation (Fig.2B, D).

To screen for additional cell surface targets of UL148, we used plasma membrane profiling (PMP) of cells infected with HCMV, comparing Merlin to HCMV∆UL148. Filtering for proteins with Ig-, MHC, Cadherin, C-type lectin and TNF InterPro functional domains (31, 33) was used to extend interrogation of UL148, focusing the analysis on immune receptors and ligands directly involved in NK or CTL functions (Fig.2E, Dataset S1 ‘Summary’). This proteomic analysis identified CD58 as the only cell surface molecule targeted by UL148 that fell within these categories (Fig.2E).

CD58 is retained within the cell by UL148

Expression of CD58 was then studied during the course of HCMV infection. CD58 was gradually downregulated from the cell surface (Fig.1), whereas expression increased in whole cell lysates (Fig.3A). A faster migrating EndoH-sensitive CD58 glycoform accumulated in HCMV-infected cells and contrasted with the EndoH-resistant form detected in cells infected with HCMV∆UL148 (Fig.3A & B). This result is consistent with UL148 retaining CD58 as an immature precursor in the ER prior to processing through the Golgi complex. This model was further supported by co-immunoprecipitation of CD58 with UL148 from infected cells visualized either using immunoprecipitation with V5-tagged UL148 and Western blotting with anti-CD58 (Fig.3C) or in a global proteomic analysis.
using stable isotope labeling by amino acids in cell culture and immunoprecipitation (SILAC-IP) (Fig.S3). In conclusion, UL148 was both necessary and sufficient to mediate cell surface down-regulation and intracellular retention of CD58.

**UL148 is a potent modulator of CTL function**

The functional impact of CD58 regulation on CTL recognition was investigated in the context of HCMV infection by using HLA-A2-restricted CD8\(^+\) CTL lines generated to HCMV-IE1 VLEETSMVL (VLE) and HCMV-pp65 NLVPMVATV (NLV) peptides. These lines were tested by using CD107a degranulation assays and intracellular cytokine staining to detect cytokine production against autologous fibroblasts (uninfected or infected with Merlin or HCMVΔUL148) pulsed with a range of peptide concentrations. Absence of peptide led to minimal CTL activation. Peptide pulsing of uninfected cells resulted in a large increase in degranulation, which was significantly impaired by infection with HCMV strain Merlin. Deletion of UL148 resulted in recovery of degranulation in both CTL lines against HCMV-infected fibroblasts, including some experiments in which activation by HCMVΔUL148 was equivalent to that observed by peptide-pulsed uninfected cells (Fig.4A). A similar effect was observed for cytokine production in a third CTL line (Fig.S4). This occurred even though both Merlin and HCMVΔUL148 down-regulated HLA class-I evenly by more than tenfold (Fig.4B), indicating that these effects were independent of signals supplied through TCR recognition of HLA-I at that peptide dose. UL148 function could therefore compensate for more than tenfold differences in HLA-I expression. Significant differences in degranulation between Merlin and HCMVΔUL148 were observed over a narrow peptide range, with close to a doubling of the proportion of activated CD8\(^+\) CTLs occurring at 1μg/ml peptide, corresponding to a molar concentration of ~1μM. Doubling or decreasing the peptide dose by 20-fold overcame these differences (Fig.4C).

**CD58 co-stimulatory function occurs only in HCMV-infected cells**

The specificity of the UL148 effect on CD58 was explored by using a monoclonal antibody (mAb) that inhibited CD2/CD58 interaction. Application of anti-CD58 mAb resulted in significant blocking of CTL activity towards cells infected with HCMVΔUL148 over a range of peptide concentrations, in some cases reducing it to the levels observed when using Merlin-infected cells as targets (Fig.4D). We performed a more detailed analysis that combined all blocking experiments by normalizing data to the isotype control in each experiment. These combined data showed that blocking of CD58 produced an effect that was observable even against Merlin-infected targets, that became significantly different from the blocking of activation by HCMVΔUL148 at the peak peptide concentration.
Irrespective of higher levels of surface CD58 compared to HCMV-infected targets, CTL activation measured by CD107 degranulation against uninfected cells could not be reduced by anti-CD58 mAb treatment, regardless of the peptide concentration used (Fig.4E,F). This differential effect of peptide loading and targets implies that HCMV-specific CD8+ CTLs are exquisitely sensitive to the context in which they receive activation signals, in that CD58-mediated co-stimulation became relevant only when these CTL were faced with an HCMV-infected target.

**UL148 significantly alters the ex vivo PBMC response to HCMV-infected cells**

The above data were generated using in vitro expanded T-cell lines. To test function in a more physiologically relevant setting, we challenged peripheral blood mononuclear cells (PBMC) ex vivo with fibroblasts infected with either Merlin or HCMVΔUL148 in the absence of exogenous peptide, comparing the degranulation responses of key effector cell subsets to the two viruses. In autologous assays and in the absence of peptide, CD3+CD8+ T-cells significantly increased their activation in response to HCMVΔUL148 in 3 of 9 subjects (Fig.4G). The CD2/CD58 axis has also been reported to be important in the activation of ‘adaptive’ NK cells defined through expression of CD57 and NKG2C, the expansion of which is associated with previous HCMV infection (25, 26). Responses of ex vivo NK cells were tested in the presence of Cytotect (purified IgG from HCMV-seropositive subjects) or IgG from HCMV-seronegative individuals, included as a negative control to measure antibody-dependent cellular cytotoxicity (ADCC) as well as standard NK cell function. NK cell function was measured against both allogeneic human fetal foreskin fibroblasts (HFFFs) and autologous HCMV-infected skin fibroblasts. The greatest effect of removing UL148 was observed in an allogeneic ADCC setting with smaller, but significant increases in NK cell activation in the absence of Cytotect (Fig.5A,B, Fig.S4). In an autologous setting, removing UL148 significantly increased the recognition of HCMV-infected targets only in the presence of Cytotect (Fig.5C). Further analysis to phenotype the responsive subset indicated it resided in CD57+ NK cells (Fig.5D), with different subjects showing significantly different responses to ΔUL148 in one, other or both of CD57+NKG2C- and CD57+NKG2C+ NK populations (Fig.5E). CD57+NKG2C+ populations were not analyzed because they represented less than 3% of total CD3-CD56+ NK cells in all but one subject.
Discussion

HCMV has become a paradigm for viral immune evasion, with the study of the activities of its genes and proteins unveiling many aspects of immune function. We have now identified HCMV UL148 as the first recognized virally encoded downregulator of the cell adhesion molecule CD58, the intracellular retention of which reduces \textit{ex vivo} activation of both CTLs and NK cells. This function is compatible with UL148 being an ER-resident type 1 transmembrane glycoprotein containing an ER retention motif (RRR, at residues 314-316) (34, 35). The CD58/CD2 axis may become particularly important when infected target cells exhibit sub-optimal activation signals, for example due to the action of multiple HCMV-encoded immune evasins.

To date, UL148 has only been assigned one other viral function. In the HCMV strain TB40/E, UL148 disruption alters the ratio of glycoprotein H/glycoprotein L (gH/gL) complexes involved in virus entry, resulting in increased infectivity of epithelial cells, in part due to a direct interaction between UL148 and those complexes, and most likely in the ER (34). Our SILAC-IP analysis of proteins binding UL148 during infection with HCMV strain Merlin did not demonstrate a specific interaction with gH or gL (Fig.S3), suggesting underlying complexity in UL148 interactions associated with the HCMV strains used and their cellular tropisms. In this regard, there is also the possibility that the host proteins targeted by UL148 may differ depending on the cell type infected by HCMV, with our data derived from fibroblasts. The rhesus cytomegalovirus (RhCMV)-encoded ortholog of UL148 (Rh159) also has effects on virus tropism, although in this case disruption of the gene renders the virus unable to spread in epithelial cells (36). Further Rh159 exhibits immune regulatory functions, impairing the surface expression of the NKG2D ligands, MICA, MICB, ULBP1 and ULBP2 (37). Interestingly, although both Rh159 and UL148 act by binding to, and retaining intracellularly their target proteins, we and others have shown that UL148 does not bind any NKG2D ligands (37). In HCMV, these ligands are targeted by UL16, UL142, US9, US18 and US20 (5, 38). It will be interesting to determine whether there is a common theme of CMV-encoded ER-resident proteins that impact on both immune evasion and cell tropism.

It is intriguing that blocking the CD2/CD58 interaction with an anti-CD58 monoclonal antibody did not inhibit effector activation by uninfected peptide-pulsed target cells, but did inhibit activation by HCMV-infected cells (Fig. 4D-F). A degree of inhibition of Merlin-infected targets was expected since CD58 was still present albeit at a reduced level.
compared to HCMVΔUL148-infected cells (Fig.4B). The absence of any effect against uninfected targets (which had equivalent levels of CD58 to HCMVΔUL148-infected cells) is surprising. The multiple immune evasion mechanisms employed by HCMV, such as impairment of TCR signaling via HLA-I downregulation or via inhibitory receptors such as LIR-1 binding HCMV UL18 (39), alter the balance of activating and inhibitory signals received by effectors. Our data are consistent with the concept that there is much more complexity in the way CD8+ CTLs are activated by virally infected targets. HCMV-specific CTLs may be ‘re-tuned’ to activate when exposed to infected targets in response to virus-encoded modulation of multiple activating and inhibitory ligands.

Some speculative evidence for this idea may be gleaned from the phenotype of HCMV-specific CTLs in vivo. HCMV drives massive, stable CD8+ CTL expansions, but it is interesting to note that the detailed phenotype and responsiveness of these cells is unusual compared to those described in classical models of T-cell differentiation (reviewed recently in (1)). For example, continuous in vitro stimulation by anti-CD3/anti-CD28 beads induces hallmarks of exhaustion in CD8+ T-cells (IL-7Rlo; and high levels of the immune cell inhibitor, programmed cell death protein 1 (PD1) (40)) that are reduced by co-stimulation through anti-CD2 signals (28). In contrast, although HCMV-specific CD8+ CTLs have been reported as showing an exhausted/senescent and/or terminally differentiated phenotype, they exhibit proliferative capacity, the ability to change co-stimulatory and chemokine receptor-phenotype (41, 42) and do not show all the functionally associated classical markers of exhaustion such as PD1. They are generally PD1lo (39), whereas HCMV increases PD-L1 expression on infected cells (31), which would be consistent with HCMV-specific CTLs adapting to receive fewer inhibitory signals through the PD1/PD-L1 axis.

It is possible that immune adaptation occurs in all effector cells facing HCMV-infected targets, and that this leads to the unusual effector phenotypes observed in HCMV-seropositive subjects. With regard to NK cells, an ‘adaptive’ NK subset that is CD57+NKG2C+FceR1- is expanded in HCMV-infected individuals and involved in ADCC (43, 44). In vitro expanded NKG2C+ NK cells exhibit higher levels of CD2 expression (45), which could aid activation in the face of lower levels of CD58 on target cells. We found that CD57+ NK cells exhibited enhanced ADCC in response to targets infected with HCMVΔUL148 compared to Merlin, with different subjects showing an impact on either or both NKG2C− and NKG2C+ NK cells within the CD57+ population (Fig.5). This is not unexpected, as NKG2C null individuals show CD57+ NK expansions following HCMV
infection (2), which exhibit a requirement for CD2 co-stimulation in their responses (26). The data are consistent with expanded effector subsets in HCMV-seropositive individuals showing greater responsiveness to an activation pathway being inhibited by HCMV.

Beyond CTL and NK cell recognition, it is notable that HCMV infection specifically induces cell surface expression of intercellular adhesion molecule 1 (ICAM1), which is reported to have an involvement equivalent to that of CD58 in formation of the SMAC (31, 46, 47). In the context of differential expression of such adhesion molecules. It is also tempting to speculate that ICAM1 may play an essential separate function in HCMV biology, and the specific downregulation of CD58 might provide an elegant mechanism for compensating for ICAM1 induction on infected cells. Indeed, ICAM1 is important in facilitating endothelial transmigration and permeability, and its induction could enhance dissemination of virus through host tissues (48) or play a role in direct cell-to-cell transfer of virus (49).

Finally, our data highlights that significant ex vivo ADCC does occur against HCMV-infected cells, even with HCMV strain Merlin, which encodes many NK cell immune evasion mechanisms and multiple Fc binding proteins (50). The impact of deleting UL148 on ADCC and general immune responses suggest that HCMV might be manipulated to drive the activation of multiple different effector cell types, making the emerging field of HCMV-based vector design all the more important for vaccine development.
Materials and Methods

Cells
Human fetal foreskin fibroblasts immortalized with human telomerase (HF-TERT), HF-TERTs transfected with the coxsackie-adenovirus receptor (HF-CAR), and TERT-immortalized donor dermal fibroblasts have been described previously (10). Cells were maintained in DMEM/10% fetal calf serum at 37°C/5% CO₂.

Viruses
HCMV strain Merlin RCMV1111/KM192298 (RL13-, UL128-) (51), AD169 varUK/BK000394, and Merlin recombinants containing single gene deletions in U₇/b’ were generated as described previously (51) (SI1). Tagged HCMV recombinants were generated as described previously (38, 52) (SI1). Recombinant adenovirus vectors expressing individual HCMV UL/b’ genes were generated as described previously (53) and validated for expression (54).

Antibodies and other reagents
All reagents were obtained from Biolegend, except Aqua live/dead dye (ThermoFisher), CD3-PE-Cy7 (clone UCHT-1, Beckman Coulter), CD8-APC-H7 (clone SK1, BD Biosciences), CD56-PE (clone N901, Beckman Coulter), CD58 (Abcam) CD107a-FITC (clone H4A3, BD Biosciences), MHC class-I (clone W6/32, Serotec) NKG2C-PE (clone 134591, R&D systems), CD155 (5D1, (55)), actin (Sigma), V5-agarose (Abcam), V5 (Serotec), anti-mouse-AF647 (ThermoFisher), anti-mouse-HRP (BioRad), anti-rabbit-HRP (BioRad), mouse IgG, UL141 (56), EndoH (NEB), PNGaseF (NEB). Other reagents were CD8-APC (clone HIT8a), CD56-BV605 (clone HCD56), CD57-APC (clone HNK-1), CD58 (clone TS2/9), CD107a-PerCP-Cy5.5 (clone H4A3), TNFα-BV421 (clone Mab11), IFNγ-PE-Cy7 (4S.B3), MHC class-I-PE (clone W6/32).

Flow cytometry
For HCMV infections, adherent cells were harvested with TripLE Express (Thermofisher) or HyQTage™ (GE Healthcare), stained in PBS/1%BSA buffer at 4°C with relevant antibodies, fixed with 4% paraformaldehyde and analyzed on an Accuri C6 flow cytometer (BD Biosciences) and with Accuri C6 software. For CD107a degranulation assays, data were gathered on an 11-color Attune NxT flow cytometer (ThermoFisher) and analyzed using Attune NxT or FlowJo V10 software.
**Immunoblotting**

Cells were lysed and boiled in reducing denaturing Nu-PAGE lysis buffer, and protein samples were separated on Nu-PAGE gels, transferred onto nitrocellulose membrane (GE Life Sciences), stained with relevant antibodies and Supersignal West Pico chemiluminescent substrate and imaged on Hyperfilm-MP (GE Life Sciences). In co-immunoprecipitation experiments, cells were lysed in Triton X-100 lysis buffer, and protein complexes were captured with V5-agarose prior to SDS-PAGE and immunoblotting.

**NK and T-cell assays**

HCMV-specific CTL lines were grown from PBMCs stimulated with irradiated (6000 RADs) autologous, peptide-coated fibroblasts as described previously (10). Degranulation assays were performed as described previously (57), using effector:target ratios of 10:1. The targets were pulsed with peptide at various concentrations and the excess washed off, while for blocking studies, a final concentration of 10 µg/ml anti-CD58 mAb, TS2/9, was used. Statistical testing of data was carried out by using Graphpad Prism 5.0 for ANOVAs with Tukey’s multiple comparisons post-tests; \( p < 0.05 \) was considered significant. Assays detecting ex vivo PBMC responses were carried out as described previously (58), with adaptation to a CD107a degranulation readout as described (57) and using multicolor flow cytometry to identify responding CD3\(^+\)CD8\(^+\) T-cells, CD3\(^-\)CD56\(^+\) NK cells and populations defined by expression of CD57 and NKG2C. Cytotect (Biotest) or IgG purified from HCMV-seronegative subjects was incubated with targets for 10 mins at 37°C at a concentration of 100 µg/ml prior to addition of an equivalent volume of effectors.

**Proteomics**

Proteomics was performed as described previously (31). Briefly, plasma membrane glycoproteins were oxidized with sodium metaperiodate and then biotinylated with aminooxybiotin (Cambridge Bioscience) prior to lysis of cells with Triton X-100. Biotinylated glycoproteins were captured with streptavidin agarose, and then washed, denatured, alkylated and digested on beads with Trypsin (Gibco). Peptides were fractionated by using strong cation exchange. Enriched peptides were labeled with tandem mass tag (TMT) reagents, combined at a 1:1 ratio, and then pre-fractionated by offline high-pH-reversed-phase chromatography (Agilent). Mass spectrometry and data analysis were performed as described previously by using an Orbitrap Fusion. P-values were estimated using Benjamini-Hochberg-corrected significance A or Significance B values from Perseus version 1.2.0.16 (59). SILAC-IP experiments were conducted and analyzed as described previously (10).
Ethics statement

Healthy adult volunteers provided blood and dermal fibroblasts for this study after providing written informed consent. The study was approved by the Cardiff University School of Medicine Research Ethics Committee ref. nos: 10/20 and 16/52.

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References


**Figure Legends**

**Figure 1: HCMV infection down-regulates cell surface CD58**
Human fibroblast (HF-TERT) cells infected with (A) HCMV strain Merlin or (B) strains Merlin or AD169 (MOI=5) or (A, B) mock-infected, and analyzed at the indicated time points post-infection by flow cytometry for cell surface expression of CD58 and MHC class-I. clgG = isotype control IgG.

**Figure 2: CD58 cell surface down-regulation is mediated by UL148**
(A) Human fibroblast (HF-TERT) cells infected with a library of HCMV strain Merlin deletion mutants, (MOI=5, 72h post infection) and (B) human fibroblast (HF-CAR) cells infected with a library of adenovirus vectors encoding HCMV strain Merlin UL/b’ genes, (MOI=5, 48h post-infection) as indicated were analyzed by flow cytometry for cell surface expression of CD58 and MHC class-I. MHC class-I down-regulation, which is a standard marker of HCMV infection, was used for quality control. Median fluorescence intensity (MFI) values relative to control cells (set to 1) are shown. (C, D) Representative plots from panels (A) and (B), respectively. (E) Scatterplot of cell surface proteins modulated by UL148 analyzed by plasma membrane profiling. The complete data spreadsheet is shown in SI3. Proteins that contained Ig-/ MHC/ Cadherin/ C-type lectin/ TNF InterPro functional domains were included in the scatterplot. Significance B was used to estimate p values (59). The complete data spreadsheet is shown in Dataset S1.

**Figure 3: UL148 retains immature CD58 intracellularly and interacts with CD58**
Human fibroblast (HF-TERT) cells were infected (MOI=5) with HCMV strain Merlin (HCMV) or a UL148 deletion mutant (ΔUL148) and lysates were analyzed by immunoblotting (A) at the indicated time points post-infection or (B) 72h post-infection. EndoH or PNGaseF glycosidases were used as indicated. CD155, which is known to be retained in the ER by HCMV UL141 and actin were used as controls. (C) HF-TERT cells were infected with HCMV strain Merlin (HCMV) or HCMV recombinants expressing V5-tagged UL148 (HCMVUL148.V5) or UL141 (HCMVUL141.V5) and whole cell lysates (WCL) were analyzed by immunoblotting or co-immunoprecipitation (V5-IP). The known interaction between CD155 and UL141 served as a control.

**Figure 4: UL148 modulates CTL function against HCMV-infected cells through CD58**
(A) Human dermal fibroblast (D007) cells were infected with HCMV strain Merlin (HCMV) or an UL148 deletion mutant (ΔUL148) (MOI=10, 72h). Cells were pulsed with VLE or NLV peptide at 1 µg/ml and used in standard CD107 degranulation assay as targets for VLE-
or NLV-specific T-cell lines generated from donor D007. Means + SEM of quadruplicate samples are shown. (B) Expression of CD58 and MHC class-I on D007 cells. (C) Summary of HCMV vs HCMVΔUL148 data from 5 separate experiments standardizing values to the mock-infected control at the peptide concentrations indicated. One-way ANOVA with Tukey (for >5 means) multiple comparison post-hoc tests showed significance differences between HCMV and HCMVΔUL148 at *** p<0.001. (D) Effect of anti-CD58 mAb or an isotype control added at the start of assays to a concentration of 10µg/ml at the indicated peptide concentrations. Means + SEM of quadruplicate samples are shown. (E) Summary data of 3 different blocking experiments following standardization to isotype control values. (F) Further blocking assay on uninfected fibroblasts at lower concentrations of peptide pulsing. One-way ANOVA with Tukey (for>5 means) multiple comparison post-hoc tests showed significance differences at ** p<0.01 and *** p<0.001. (G) Degranulation of CD3+CD8+ T-cells in PBMC of 9 HCMV-seropositive donors challenged with autologous fibroblasts infected with HCMV strain Merlin or HCMVΔUL148 in the absence of peptide (MOI=10, 72h). Points are means of triplicate samples. Paired t-Test showed the p value indicated. # marks the 3 of 9 donors showing significant differences (p<0.05) when comparing responses between Merlin and HCMVΔUL148 using one-way ANOVA with Tukey multiple comparison post-hoc tests with %change from the Merlin response indicated in brackets.

**Figure 5: Deletion of UL148 alters the response of NK cells**

Fibroblasts were infected with HCMV strain Merlin (HCMV) or a UL148 deletion mutant (ΔUL148) (MOI=10, 72h). Cytotect or HCMV-negative IgG was added to a final concentration of 50 µg/ml. The cells were used to stimulate PBMCs from healthy donors stimulated overnight with 1000 IU/ml IFNα. CD3−CD56+ NK cell responses to allogeneic fibroblast (HF-TERT) cells from (A) a representative donor and (B) ten donors comparing responses with control HCMV-negative IgG and Cytotect. CD3−CD56+ NK cell responses to autologous dermal fibroblasts are shown in (C) with a summary of nine donors comparing responses with control HCMV-negative IgG and Cytotect. Allogeneic ADCC responses were further split into (D) CD57− and CD57+ NK responses, and (E) CD57−NKG2C−, CD57+NKG2C+ and CD57+NKG2C− NK responses in 7 to 9 subjects. Data passed D’Agostino & Pearson omnibus normality testing. Points are mean of triplicate cultures. Paired t-Test showed the indicated significance p values. # marks donors showing significant differences (p<0.05) when comparing individual responses between Merlin and HCMVΔUL148 using one-way ANOVA with Tukey multiple comparison post-hoc tests. (2C-) indicates NKG2C− subjects, detected by flow cytometry.
Figure 1

A

24h 48h 72h

HCMV Merlin

cIgG

cell surface CD58

cell count

24h 48h 72h

HCMV Merlin

cIgG

cell surface MHC-I

cell count

B

72h

mock

HCMV Merlin

HCMV AD169

cIgG

cell surface CD58

cell count

72h

mock

HCMV AD169

HCMV Merlin

cIgG

cell surface MHC-I

cell count
Figure 2

A

ΔUL150/A
ΔUL148D
ΔUL148C
ΔUL148B
ΔUL148A
ΔUL148
ΔUL147A
ΔUL147
ΔUL146
ΔUL145
ΔUL144
ΔUL143
ΔUL142
ΔUL141
ΔUL139
ΔUL138
ΔUL136
ΔUL135
ΔUL133
ΔUL132
ΔUL131A
HCMV
mock

cell surface CD58

B

UL150/A
UL148D
UL148C
UL148B
UL148A
UL148
UL147A
UL147
UL146
UL145
UL144
UL143
UL142
UL141
UL139
UL138
UL136
UL135
UL133
UL132
UL131A
HCMV
mock

cell surface MHC-I

C

HCMV
mock

HCMV
ΔUL148

cell count

cell surface CD58

cell surface MHC-I

D

RAd

UL148

RAd

CTRL

RAd

UL148

RAd

CTRL

cell count

cell surface CD58

cell surface MHC-I

E

UL148 deletion down-regulates

UL148 deletion up-regulates

Log₁₀(signal : noise)

Fold change HCMVΔUL148 : HCMV

CD58
Figure 4

A VLE line, VLE (1 μg/ml) vs NLV line, NLV (1 μg/ml)

B HCMV mock vs HCMV ΔUL148 cell surface CD58

C 100% of mock %CD107+

D peptide c=0.04 μg/ml vs peptide c=0.2 μg/ml vs peptide c=1 μg/ml

E anti-CD58 % of isotype CTRL

F isotype CTRL vs anti-CD58 mock-infected targets

G p=0.049
Figure 5

A. NK vs allogeneic targets

B. Allogeneic targets

C. Autologous targets

D. NK vs allogeneic targets + Cytotect

E. NK vs allogeneic targets + Cytotect
Supplemental Information Legends

Figure S1: Details of the library of HCMV recombinants containing single gene deletions

Recombinant HCMVs (RCMVs) were generated by recombineering of the strain Merlin bacterial artificial chromosome BAC1111/KM192298 (RL13-, UL128-) (Stanton RJ, et al. (2010). J Clin Invest 120: 3191-3208). Whole-genome consensus sequences of passage 1 of each RCMV were derived by using the Illumina platform as described previously (Fielding CA, et al. (2014). PLoS Pathog 10: e1004058), and were deposited in GenBank. Deleted regions are shown between slashes in lower case, with the initiation codon underlined, and did not always include the whole protein-coding region. Except for the differences noted, the viruses were genetically identical to that recovered from the parental BAC (RCMV1111/KM192298).

* middle G tract near right end of US is 1 bp shorter
** contains an 1190 bp E. coli insertion element IS150 plus a 3 bp duplication at the right end inserted into US11. This insertion was considered irrelevant to this study, but the recombinant was not banked for further use and the sequence was not deposited in GenBank.

Two further recombinants (not listed) were also constructed and their whole-genome sequences verified. To insert a V5 tag after UL148 (RCMV2084), a recombineering cassette comprising KanR/rpsL/lacZ was amplified and inserted into the 5’ end of UL148 by using primers

<table>
<thead>
<tr>
<th>Forward primers</th>
<th>Reverse primers</th>
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<td>GCATTCTCGAGACGTCGGCGAAGGTTTCTGCTATGTCCGCGAGCGACGTGTA</td>
<td>TCAACACATTGCTGATTACAATGATGGCGGCTCTGCGTTTTGATAAACC</td>
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| CGAGGACGTCGGTTCGCCGAAGCCCTCCTGTGAGCGAAGATCATT | ACCTGGTTGTCGCAGCGGTAGCTGAGTTCTTTATGCTCCTT-3’. In a second round of recombineering, this cassette was removed and replaced by a V5 tag by using the complementary primers
| TGCTATGTCCGCAACGAGCTGACGGAGACGTGGTTTCGCCAAGCCTTTACGTA | GAATCAAGACCTAGGAGGCGGTTAGGGATTGGCTTTACCAGCGCT |
| And | (V5 tag and linker underlined). To insert a V5 tag after UL141 (RCMV2022), the KanR/rpsL/lacZ was amplified and inserted at the end of UL141 using primers
| GGCGATATGGGCTCCTGGTTTGTATAACCTACCTGGTGTCGCAGGCGTGGAGCGCT | GGTAAGCCATCCCTAACCAGCCTCTAGGTTCTTTGATACGTA
AGGGGACGAGCGGGCTATCGACGCCTACCGACTTACGATAGTTACC
CCGGTGTTAAAAAGATGAAGAGGCCTGTGACGGAAGATCACTTCG and
GCATATTTTAATCACACTATTCCACACTTTCAACACACTGCATTTTTTAACATCTTTTT
TTATTTATGCGTTCTCACTGAGGTTCTTATGGCTCTTG
(regions of homology to UL141 underlined). This cassette was then removed and replaced
with a V5 tag, using primers
GACGCCTACCAGACGATGATTTACCCGGGTGTAAAAAGATGAAGAGGGGCTCC
GGGGGGTCGGGTGGAAGTGGCGGTAAGCCAATCCCTAACCCGCT
And
ACACACTGCATTATTAAACACTCTTTTTTTTTTTTATCGTGTTCTCACGTAATC
AAGACCTAGGACCGGGTTAGGGATTGGCTTACCGCCTC (complementary
regions that overlapped to generate the V5 tag underlined).

Figure S2: Transcriptome profiles of HCMV strain Merlin (RCMV1111) and ΔUL148
(RCMV2035)
(A) Whole-cell RNA was isolated from human fibroblast (HF-TERT) cells infected with
HCMV strain Merlin (RCMV1111) or a UL148 deletion mutant ΔUL148 (RCMV2035) at 72
h post-infection. Transcriptome profiles were derived from directional Illumina RNA-Seq
data generated from polyadenylated RNA. The genome is shown in five sections, with the
inverted repeats shaded grey. Protein-coding regions and non-coding RNAs are shown by
color- and white-shaded arrows, respectively, with gene nomenclature below, and introns
as narrow white bars connecting exons. The colours of protein-coding regions indicate
conservation among alpha-, beta- and gammaherpesviruses (core genes) or between
beta- and gammaherpesviruses (subcore genes), with certain noncore genes grouped into
related families. The yellow windows depict transcription profiles for rightward (magenta)
and leftward (cyan) transcripts. The extent of transcription is plotted as log₁₀ of the number
of reads per nucleotide (calculated by Bowtie2 assembly against the RCMV2035
sequence) per million viral reads, the overall number of reads for RCMV2035) being
normalised to that for RCMV1111. The number of viral reads (76 nucleotides) in the
RCMV1111 and RCMV2035 datasets was 11,857,464 (48% of the total) and 12,178,576
(52%), respectively. Among the regions containing zero reads (scored -2) is UL148 in
RCMV2035 (boxed). Except for UL148, the transcriptome profile of RCMV2035 was very
similar to that of RCMV1111. (B) Histogram showing the relative proportion (%) of
normalized sequence read counts mapping to individual HCMV protein-coding regions and
noncoding RNAs in RCMV2035 compared with RCMV1111. In this experiment, and
excluding UL148, transcription levels of the protein-coding regions and non-coding RNAs
were 105±16% of those in RCMV1111, ranging from 57% (UL146) to 183% (the region common to IRS1 and TRS1). Only UL148 was differentially expressed >2-fold.

**Figure S3: Identification of cellular proteins interacting with UL148 via SILAC-IP**

SILAC-labelled HFFF-TERT cells were infected (MOI=5, 72h) with unmodified HCMV strain Merlin (light-labeled cells), HCMVUL141.V5 (medium-labeled cells) or HCMVUL148.V5 (heavy-labeled cells) followed by immunoprecipitation of mixed lysates with anti-V5. Enriched proteins were digested with trypsin then subjected to mass spectrometry. To identify proteins that specifically interact with HCMV UL148, the following filters were employed: UL148:wt ratio of >2 and UL148:UL141 ratio of >2. UL141 served as an irrelevant control and did not interact with UL148 (heavy : light ratio = 1.3). Significance A was used to estimate p-values, which were adjusted for multiple hypothesis testing using the method of Benjamini-Hochberg (59).

**Figure S4: Effect of UL148 on cytokine production**

Autologous skin fibroblasts were infected with Merlin (HCMV) or a UL148 deletion mutant (∆UL148) (MOI =10, 72h) and then used as targets against a CD8+ T-cell line or PBMCs. CD107a degranulation and intracellular cytokine staining for IFNγ and TNFα were measured after 6h using standard procedures (Becton Dickinson). (A, B) Two separate experiments on a CD8+ T-cell line (D009-VTE) at a cognate peptide dose of 1 µg/ml. (C) PBMC data gating on CD57+CD3-CD56+ NK cells with targets in the presence of Cytotect. MFI = median fluorescence intensity for positive cells. Mean + SEM of quadruplicates shown. Two-way ANOVA comparing all means showed significance at *p<0.05, **p<0.01, ***p<0.001.

**Dataset S1: Mass spectrometric analysis comparing Merlin and Merlin∆UL148**

Excel spreadsheet, no figure legend provided.
<table>
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Supplemental Figure S2

(A) [Graphical representation of data]

(B) [Graphical representation of data]
Supplemental Figure S3

Fold enrichment HCMV UL148.V5 : wt

- CD59
- UL148
- LRCC8
- SLC25A10
- P4HB
- ATB8VA
- ZNF318
- TUBA1A
- YLPM1
- PIDA3
- US29
- TUBA1B
- TUBA1A
- ATP01B2
- TUBA1C
- SNAPE
- NCOA3

- p < 0.01
- p < 0.05
- p < 0.1
- p > 0.1
Supplemental Figure S4

A. **D009-VTE expt 1**

B. **D009-VTE expt 2**

C. **D007 - CD3-CD56+CD57+ NK cells vs allogeneic targets + Cytotect**