respond differently to TNF-α or whether the experimental readouts were such that they placed in evidence one or the other outcome. Chances are that we have not heard the last word on the connection between TNF-α and Tregs.

Acknowledgments
A.M. Bilate was a recipient of a postdoctoral fellowship from the Pew Latin American Fellows Program.

Address correspondence to: Juan J. Lafaille, Molecular Pathogenesis Program, Kimmel Center for Biology and Medicine of the Skirball Institute, New York University School of Medicine, 540 First Avenue, New York, New York 10016, USA. Phone: 212.263.1489; Fax: 212.263.1171; E-mail: juan.lafaille@med.nyu.edu.


Can we build it better? Using BAC genetics to engineer more effective cytomegalovirus vaccines

Mark R. Schleiss
Department of Pediatrics, University of Minnesota Medical School, Center for Infectious Diseases and Microbiology Translational Research, Minneapolis, Minnesota, USA.

The magnitude and durability of immunity to human cytomegalovirus (HCMV) following natural infection is compromised by the presence of immune modulation genes that appear to promote evasion of host clearance mechanisms. Since immunity to HCMV offers limited protection, rational design of effective vaccines has been challenging. In this issue of the JCI, Slavuljica and colleagues employ techniques to genetically modify the highly related mouse CMV (MCMV), in the process generating a virus that was rapidly cleared by NK cells. The virus functioned as a safe and highly effective vaccine. Demonstration of the ability to engineer a safe and highly effective live virus vaccine in a relevant rodent model of CMV infection may open the door to clinical trials of safer and more immunogenic HCMV vaccines.

The urgent need for a vaccine against human CMV
Human CMV (HCMV) is an important cause of disease in immunologically compromised individuals, including recipients of solid organ and hematopoietic stem cell transplants and patients with advanced HIV disease. At greatest risk for HCMV-associated injury, however, is the developing fetus. HCMV is the most common agent of congenital viral infection in the United States, and among all infectious diseases is the most common cause of childhood neurological disability, including deafness, in the developed world (1, 2). Since severe and symptomatic congenital HCMV infections can be associated with a lifetime of disability, the economic burden associated with this infection is striking. When the Institute of Medicine was commissioned to prioritize vaccine development for the new millennium based on, among other factors, quality-adjusted life years (a marker of economic benefit), a vaccine for HCMV was ranked “head-and-shoulders” above all other potential new vaccines with respect to overall cost-effectiveness (3).

Although the need for an HCMV vaccine is compelling, it is less clear to whom such a vaccine should be administered, and what the constituents of such a vaccine should be. The correlates of protective immunity remain undefined, both for the nonpregnant individual and for the developing fetus. Subunit vaccines, typically based on recombinant expression of key targets of humoral and cellular immune responses to HCMV infection, have been evaluated in clinical trials, as have live-attenuated vaccines (4). Until recently, clinical trials have yielded little information about the potential for protective efficacy, largely because most studies have focused on the
end points of safety and immunogenicity and not the end point of protection against HCMV infection. A breakthrough in the HCMV vaccine field came in 2009, when, for the first time, a vaccine was shown to be effective against acquiring primary infection (5). This vaccine was based on a purified, recombinant formulation of the major envelope glycoprotein B, administered with a proprietary adjuvant known as MF59. In a phase II, placebo-controlled study, 18 of 225 (8%) women who received gB/MF59 vaccine acquired a primary HCMV infection in the one-year study window, compared with 31 of 216 in the placebo group (14%), for an overall vaccine efficacy of 50% (5). These data represent a major advance in the field, since they are the first evidence having been “hijacked” from the host genome during the course of coevolution of virus and host. Several of these gene products interfere with host adaptive immune responses, with the apparent goal of enabling immune evasion; others appear to play a proinflammatory role, seemingly to promote inflammation that presumably could facilitate widespread dissemination of CMV in the infected host. These gene products, which include cytokines, chemokines, GPCR homologs, and inhibitors of MHC class I and class II antigen presentation, are summarized in Table 1.

Table 1
HCMV immune modulation genes

<table>
<thead>
<tr>
<th>CMV gene(s)</th>
<th>Putative immunomodulatory function</th>
</tr>
</thead>
<tbody>
<tr>
<td>UL16, UL18, UL142</td>
<td>Homologs of MHC class I molecules; promote NK cell evasion through binding of NK cell–inhibitory ligands, downregulation of NK cell–activating ligands</td>
</tr>
<tr>
<td>UL40, UL83, UL141</td>
<td>NK cell evasion genes; promote expression of non-classical HLA receptors; binding and inhibition of NK cell–activating proteins; downregulation of CD155 (activating receptor)</td>
</tr>
<tr>
<td>TRS1, IRS1</td>
<td>dsRNA-binding proteins; prevent activation of PKR following CMV infection; evasion of IFN response</td>
</tr>
<tr>
<td>UL111A</td>
<td>IL-10 cytokine homolog; possible downregulation of host inflammation to evade immune clearance</td>
</tr>
<tr>
<td>UL146, UL147</td>
<td>Homologs of C-X-C chemokines; possibly function to promote widespread dissemination of infection in host</td>
</tr>
<tr>
<td>UL128, UL130</td>
<td>Homologs of C-C chemokines; immune function unclear; ancillary proteins involved in viral entry into epithelial cells</td>
</tr>
<tr>
<td>UL33, UL78, US27, US28</td>
<td>Homologs of GPCRs; ligands may include multiple chemokines and cytokines; signaling molecules; may function as chemokine “sinks” to sequester immune activation proteins compatible with an immune evasion role</td>
</tr>
<tr>
<td>US2, US3, US6, US11</td>
<td>Genes interfering with MHC class I antigen processing; prevent development of CTL response to CMV through retention of MHC in the ER; facilitation of degradation of MHC; downregulation of MHC expression; immune evasion function</td>
</tr>
<tr>
<td>UL144</td>
<td>TNFR homolog; possible role modifying cytokine response to Th2 cell polarization; immune evasion</td>
</tr>
<tr>
<td>UL112</td>
<td>MicroRNA; functions at RNA not protein level; NK cell evasion</td>
</tr>
</tbody>
</table>

HCMV genes compromise immunity and complicate vaccine design
One of the most remarkable aspects of the biology of HCMV that has only recently begun to be elucidated is the role that multiple virus-encoded immune modulation genes play in the establishment (or lack thereof) of long-term immunity to this infection (6). Many of these immune modulation genes appear to be homologs of normal cellular immune effectors, presumably having been “hijacked” from the host genome during the course of coevolution of virus and host. Several of these gene products interfere with host adaptive immune responses, with the apparent goal of enabling immune evasion; others appear to play a proinflammatory role, seemingly to promote inflammation that presumably could facilitate widespread dissemination of CMV in the infected host. These gene products, which include cytokines, chemokines, GPCR homologs, and inhibitors of MHC class I and class II antigen presentation, are summarized in Table 1.

In addition to gene products that appear to modulate adaptive immune responses, HCMV also encodes proteins that impact innate responses, particularly the function of NK cells (Figure 1 and ref. 7). HCMV has evolved several genes that interact with NK cell–mediated pathways, modifying pathways of both inhibition and activation. One such protein, gpUL18, is encoded by the UL18 gene and appears to exert its evasive effect by binding to the NK cell inhibitory receptor leukocyte Ig-like receptor–1 (LIR-1) with a higher affinity than does the host MHC class I molecule (8, 9). Recently it has been shown that cells expressing another HCMV MHC class I homolog, the product of the UL142 gene, are protected from NK cell lysis (10). The NK cell evasion effect is related to the ability of gpUL142 to downregulate MHC class I peptides–related sequence A (MICA), which is a ligand for NK cell activation receptor NK group 2, member D (NKG2D) (11). A number of other CMV gene products, including those encoded by UL40, UL83, and UL141, also appear to function as NK cell evasins. The UL40 protein contains an epitope in its leader sequence that binds to HLA-E and is presented at the surface of infected cells. The gpUL40 peptide/HLA-E complex binds to CD94/NKG2A, resulting in NK cell inhibition (7). Remarkably, not all HCMV NK cell evasion genes function by encoding proteins: some gene products can promote evasion of innate immunity at the RNA level. Recently, a novel mechanism of NK cell evasion was attributed to an HCMV microRNA encoded by the HCMV UL112 gene (12). This effect appears to require interaction between the UL112 miRNA and a cellular miRNA (13).

What is the clinical significance of the presence of this complex array of HCMV immune modulation genes? Key to the problem of HCMV immunity is the issue of reinfection. Mounting evidence indicates that people who have cleared primary viral infection and have latent, clinically quiescent infections can be reinfected with novel strains of HCMV. Such reinfestions can occur in women of child-bearing age, even in the face of long-standing immunity, and can lead to symptomatic transmission to the developing fetus, with its attendant morbidity (14–16). The role that immune evasion genes play in the context of in vivo infection was recently authenticated in rhesus macaques (17). In this study, macaques that had been naturally infected with the...
sus CMV (RhCMV, the rhesus macaque homolog of HCMV) were infected with a recombinant RhCMV virus expressing the SIV Gag protein. Despite their preexisting immunity, all inoculated macaques developed SIV Gag–specific T cell responses and shed the recombinant virus for up to three years after challenge. The ability of this virus to superinfect immune macaques, however, was abrogated by deletion of the RhCMV-encoded inhibitors of MHC class I antigen presentation. The authors pessimistically concluded that this ability to evade CTL-mediated immune memory “renders it unlikely” that an effective vaccine against HCMV is an “achievable goal” (17).

Building it better: engineering enhanced effectiveness in a CMV vaccine

In light of HCMV-mediated immune evasion mechanisms, what strategies could enable development of an effective vaccine? Does the presence of immune evasion genes make the effort to develop a live-attenuated vaccine hopeless? Or, do we have the technology to bioengineer a vaccine against HCMV that is “better” than natural immunity? The work of Slavuljica and colleagues reported in this issue of the *JCI* suggests that the answer, at least for a related CMV (mouse CMV [MCMV]), may be “yes” (18). In this study, MCMV was studied in a vaccine/challenge model. Like HMCV, MCMV encodes a wide array of immune evasion genes: some are highly homologous to those encoded by HCMV, and others are unique and species-specific. In particular, MCMV encodes a variety of gene products that interfere with NK cell–mediated clearance following infection, either by downregulating ligands for NKG2D or by encoding homologs of MHC class I molecules capable of binding NK cell inhibitory receptors. Examples include the m145-encoded protein, which decreases expression of the NKG2D ligand MULT-1 (19); the m155-encoded glycoprotein, which downregulates expression of the NKG2D ligand H60 (20); and the m152-encoded glycoprotein, which downregulates expression of proteins of the RAE-1 family of NKG2D ligands (21). The m138 protein functions by downmodulating multiple NKG2D ligands, including MULT-1 and H60 (22). MCMV mutants lacking these genes are attenuated in vivo, but if mice are depleted of NK cells prior to viral challenge, replication of MCMV can be restored.

Using this knowledge, how did Slavuljica and colleagues design a vaccine with the potential to confer more robust immune responses than natural infection (18)? The key was to modify MCMV in such a fashion as to override inhibitory signals delivered by self-MHC class I proteins or potentiated by viral proteins that downregulate NKG2D ligands (Figure 2). Implicit in this strategy is the hypothesis that not only could such a vaccine have an outstanding safety profile,
Recombinant MCMV overrides NK cell evasion genes, producing highly attenuated but immunogenic and protective vaccine. (A) Like HCMV, MCMV encodes a variety of gene products that interfere with NK cell–mediated viral clearance by downregulating activating ligands for NKG2D. Examples include the m145 protein, which decreases expression of MULT-1; the m155 glycoprotein, which downregulates H60; and the m152 glycoprotein, which downregulates proteins of the RAE-1 family. The m138 protein functions by downmodulating multiple NKG2D ligands, including MULT-1 and H60. NK cell evasion genes impair activation of NKG2D as indicated, resulting in viremia, disseminated infection, and end-organ disease. (B) Recombinant MCMV attenuated vaccine described by Slavuljica and colleagues (18). Recombinant virus was engineered to express the NKG2D activating ligand RAE-1γ at high levels. This modification overcomes other NK cell evasion genes, resulting in minimal replication in vivo, but eliciting strong cellular and humoral immune responses that provide protection against MCMV disease following challenge with virulent, salivary gland–adapted virus.
due to improved innate immune clearance following inoculation, but that it also could potentially be equally or more immunogenic than “natural” infection with wild-type MCMV, by virtue of increased NK cell activation leading to stronger adaptive immune responses. To engineer such a viral vaccine, Slavuljica and colleagues utilized the powerful mutagenesis techniques enabled by the availability of the MCMV genome cloned as an infectious BAC in Escherichia coli to insert a NKG2D ligand into the MCMV genome (18), an approach that has been exploited in designing other vaccine approaches in the MCMV model (23). The ligand employed in this study was a member of the RAE-1 family of NKG2D ligands, RAE-1γ (18). Insertion of the gene encoding RAE-1γ into the MCMV genome generated a recombinant virus with a profoundly attenuated phenotype in vivo, including in transgenic mice lacking type I interferon receptor and mice that were immune suppressed. Despite this attenuation, the recombinant vaccine efficiently primed and maintained adaptive immune responses, including CD8+ T cell responses, that were comparable to those induced by wild-type MCMV. Intriguingly, in a challenge infection model, mice immunized with the RAE-1γ-MCMV recombinant resisted challenge with virulent salivary gland–adapted virus, with improved survival compared with mice initially infected with wild-type MCMV, suggesting that expression of the NKG2D ligand RAE-1γ in the context of the recombinant MCMV provides an innate immune stimulus that results in enhanced effectiveness of the adaptive immune response compared with infection with wild-type MCMV. Adoptive transfer of CD8+ T cells from vaccinated mice to immunodepleted, infected mice resulted in protection against disease. In addition to the protective T cell effect, Slavuljica and colleagues identified an important role for antibody in a newborn pup challenge model mimicking neonatal HCMV disease (18). It was noted that pups born to vaccinated, but not control, mice were protected against MCMV infection and disseminated disease following challenge in the newborn period, indicating a role for transplacental antibody in prevention of disease.

Similar observations suggesting an enhanced immunogenicity and protective efficacy for a live-attenuated vaccine engineered to modify the impact of viral immune modulation genes have been made in vaccine studies in another rodent CMV model — the guinea pig CMV (GPCMV) (24). In this model, deletion of three MHC class I homologs, presumed NK cell evasins, was achieved using BAC technology. The resulting vaccine virus was highly attenuated in vivo, but vaccination was highly immunogenic and protective. Similar to the RAE-1γ vaccine generated by Slavuljica and colleagues (18), these results suggested that the safety and efficacy of a live-attenuated HCMV vaccine could be enhanced by modification of the virus/NK cell interaction: in the case of the MCMV vaccine, by engineering a NKG2D ligand into the viral genome; and in the case of the GPCMV vaccine, by removing putative class I “decoy” genes that mediate evasion of NK cell-mediated clearance (24).

Time for human clinical trials? What’s the next step?

In light of these encouraging results in the MCMV and GPCMV models (18, 24), it seems reasonable to proceed with phase 1 studies of similarly constructed live-attenuated HCMV vaccines. Unfortunately, regulatory bodies have expressed some reluctance in moving forward with studies of live-attenuated HCMV vaccines, because of perceived long-term theoretical health risks associated with such vaccinations (25). In this context, the latent infection observed with the vaccine designed by Slavuljica and colleagues (18), which they hypothesized may help "boost" long-term immune responses during episodes of abortive reactivation, may not be advantageous for an HCMV vaccine. Additional studies are warranted in rodent models to try to optimize both attenuation and immunogenicity of live-attenuated vaccines. However, clinical trials of candidate live-attenuated HCMV vaccines should not be delayed until exhaustive evaluations are completed in animal models. The intriguing data reported by Slavuljica and colleagues (18) demonstrating feasibility and efficacy of a rationally engineered vaccine in a relevant animal model should help drive translational studies of analogous approaches in human clinical trials. The exciting advent of molecular technologies to “reengineer” live virus vaccines may be applicable to other live virus vaccines used in clinical practice. These improvements in vaccine technology — aimed at not only improving efficacy but also ensuring safety — should in turn help increase confidence in the safety and vital importance of all vaccines in clinical practice.

Acknowledgments

This work is supported by NIH grants HD038416 and HD044864.

Address correspondence to: Mark R. Schleiss, Department of Pediatrics, University of Minnesota Medical School, Center for Infectious Diseases and Microbiology Translational Research, 2001 6th Street SE, Minneapolis, Minnesota 55455, USA. Phone: 612.624.1112; Fax: 612.624.8927; E-mail: schleiss@umn.edu.

17. Hansen SG, et al. Evasion of CD8+ T cells is criti
In situations of stress the heart beats faster and stronger. According to Marks and colleagues, this response is, to a large extent, the consequence of facilitated Ca\(^{2+}\) release from intracellular Ca\(^{2+}\) stores via ryanodine receptor 2 (RyR2), thought to be due to catecholamine-induced increases in RyR2 phosphorylation at serine 2808 (S2808). If catecholamine stimulation is sustained (for example, as occurs in heart failure), RyR2 becomes hyperphosphorylated and “leaky,” leading to arrhythmias and other pathology. This “leaky RyR2 hypothesis” is highly controversial. In this issue of the JCI, Marks and colleagues report on two new mouse lines with mutations in S2808 that provide strong evidence supporting their theory. Moreover, the experiments revealed an influence of redox modifications of RyR2 that may account for some discrepancies in the field.

The heart has a remarkable capacity to react to altered demand by changing the rate at which it beats and the force with which it contracts, thereby changing its output. Both the reduction of cardiac output in phases of rest and its increase in physical and emotional exercise (the fight or flight response) are essential for normal body homeostasis and long-term survival. It is not surprising therefore that cardiac rate and force are regulated at multiple levels, extrinsic and intrinsic to the heart, and in a highly complex and secured fashion. Stimulation of \(\beta_1\)-adrenergic receptors by the sympathetic neurotransmitter norepinephrine induces increased production of the second messenger cAMP, cAMP directly and indirectly (via activation of PKA) induces faster depolarization in sinoatrial node cells (the cells that generate the action potentials that trigger cardiac contraction) and thus acceleration of heart rate (i.e., it has a “positive chronotropic effect”) and stronger contraction (i.e., it has a “positive inotropic effect”) and faster relaxation (i.e., it has a “positive lusitropic effect”) in working myocytes. In chronic heart failure, one of the most frequent life-threatening diseases in Western societies, norepinephrine levels are chronically elevated, which leads to desensitization of the \(\beta_1\)-adrenergic signalling cascade and blunted responses. \(\beta\)-Blockers, introduced by Waaegstein and colleagues in the mid 1970s, protect the heart from chronic sympathetic stimulation and provide the largest prognostic benefit for patients with chronic heart failure.

Cardiac excitation-contraction coupling

The positive inotropic and lusitropic consequences of \(\beta_1\)-adrenergic receptor stimulation in cardiomyocytes are explained by changes in excitation-contraction coupling, i.e., the relationship between the cardiac action potential and myofilament activation (Figure 1 of this commentary). When the cell depolarizes during a cardiac action potential, L-type Ca\(^{2+}\) channels (LTCCs) open, allowing Ca\(^{2+}\) to enter the cell. This so-called trigger Ca\(^{2+}\) induces a much larger Ca\(^{2+}\) release from intracellular Ca\(^{2+}\) stores, known as the sarcoplasmic reticulum (SR), via large tetrameric ryanodine-sensitive channels, referred to as ryanodine receptor 2 (RyR2). The increase in Ca\(^{2+}\) concentration initiates a conformational change in the myofilaments and thereby contraction. Removal of Ca\(^{2+}\) from the cytosol via the sarcoplasmic/endoplasmic reticulum Ca\(^{2+}\) ATPase (SERCA) and the Na\(^{+}\)/Ca\(^{2+}\) exchanger (NCX) in the plasmalemma reverses the process. Importantly, the amount of Ca\(^{2+}\) entering the cell is, under normal conditions, exactly matched by the amount of Ca\(^{2+}\) leaving it via the NCX. \(\beta_1\)-Adrenergic receptors stimulate the system at numerous levels via PKA (Figure 1 of this commentary). Phosphorylation of LTCCs increases their open probability, allowing more Ca\(^{2+}\) to enter the cell. Phosphorylation of phospholamban (PLB), a small protein that when unphosphorylated inhibits SERCA, leads to disinhibition, i.e., increased reuptake of Ca\(^{2+}\) into the SR. This has at least two consequences: first, more Ca\(^{2+}\) in the SR and therefore more Ca\(^{2+}\) release during systole, which has a positive inotropic effect; and second, faster Ca\(^{2+}\) removal from the cytoplasm and thus faster

---

**Conflict of interest:** The author has declared that no conflict of interest exists.

**Citation for this article:** J Clin Invest. doi:10.1172/JCI45251.

---

**Is ryanodine receptor phosphorylation key to the fight or flight response and heart failure?**

Thomas Eschenhagen

Department of Experimental Pharmacology and Toxicology, Cardiovascular Research Center Hamburg, University Medical Center Hamburg Eppendorf, Hamburg, Germany.