Human cytomegalovirus elicits fetal γδ T cell responses in utero

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The fetus and infant are highly susceptible to viral infections. Several viruses, including human cytomegalovirus (CMV), cause more severe disease in early life compared with later life. It is generally accepted that this is a result of the immaturity of the immune system. γδ T cells are unconventional T cells that can react rapidly upon activation and show major histocompatibility complex–unrestricted activity. We show that upon CMV infection in utero, fetal γδ T cells expand and become differentiated. The expansion was restricted to Vγ9–negative γδ T cells, irrespective of their Vδ chain expression. Differentiated γδ T cells expressed high levels of IFN-γ, transcription factors T-bet and eomes, natural killer receptors, and cytotoxic mediators. CMV infection induced a striking enrichment of a public Vγ8Vδ1–TCR, containing the germline–encoded complementary-determining-region–3 (CDR3) δ1–CALGELGDDKLIF/CDR3γ8–CATWDTTGWFKLIF. Public Vγ8Vδ1–TCR–expressing cell clones produced IFN-γ upon coincubation with CMV–infected target cells in a TCR/CD3–dependent manner and showed antiviral activity. Differentiated γδ T cells and public Vγ8Vδ1–TCR were detected as early as after 21 wk of gestation. Our results indicate that functional fetal γδ T cell responses can be generated during development in utero and suggest that this T cell subset could participate in antiviral defense in early life.
Studies in several species have shown an important role for $\gamma\delta$ T cells in protection against infection, in tumor surveillance, in immunoregulation, and in tissue repair (Hayday, 2000; Wang et al., 2001; Holtmeier and Kabelitz, 2005; Pennington et al., 2005; Toulon et al., 2009). In general, they show a rapid and robust response before the development of the adaptive immunity mediated by conventional T cells. In comparison with $\alpha\beta$ T cells, $\gamma\delta$ T cells are not abundant in the peripheral blood but are highly enriched in tissues like the gut epithelium (Hayday, 2000; Holtmeier and Kabelitz, 2005). The majority of $\gamma\delta$ T cells in human adult peripheral blood use the TCR V region pair V$\gamma9$V$\delta2$ (note that according to an alternative nomenclature the V$\gamma9$ chain is also termed V$\gamma2$ [Holtmeier and Kabelitz, 2005]). This subset has been shown to react specifically toward nonpeptide low molecular weight phosphorylated metabolites (so-called phosophoantigens) and has been the subject of several clinical trials (Wilhelm et al., 2003; Diel et al., 2007; Kabelitz et al., 2007).

Probably in all species, $\gamma\delta$ T cells are the first T cells to develop (Hayday, 2000). In contrast to adult peripheral blood $\gamma\delta$ T cells, human neonatal cord blood $\gamma\delta$ T cells express diverse V$\gamma$ and V$\delta$ chains paired in a variety of combinations (Morita et al., 1994). Thus the adult-like V$\gamma9$V$\delta2$ subpopulation only represents a small fraction of the neonatal $\gamma\delta$ T cells (Parker et al., 1990; Morita et al., 1994; Cairo et al., 2008). Further illustrating the differences between adult and neonatal $\gamma\delta$ T cells, is the demonstration that in vitro exposure toward the same pathogen (Escherichia coli or Pseudomonas aeruginosa) results in expansion of V$\delta2^+$ $\gamma\delta$ T cells in adult peripheral blood but of V$\delta1^+$ $\gamma\delta$ T cells in cord blood (Kersten et al., 1996). In mice, $\gamma\delta$ T cells are important for the protection against an intestinal parasite infection in early life but not in adult life (Ramsburg et al., 2003), and during human T cell ontogeny $\gamma\delta$ T cells mature before $\alpha\beta$ T cells (De Rosa et al., 2004). However, so far it is not known whether pathogens in early life can activate human $\gamma\delta$ T cells. To gain insight into the ability of $\gamma\delta$ T cells to mount responses to viruses during fetal life, we studied the changes occurring in the $\gamma\delta$ T cell compartment during congenital CMV infection.

RESULTS

CMV infection in utero induces expansion of fetal $\gamma\delta$ T cells in newborns

To address whether human fetal $\gamma\delta$ T cells are responsive to CMV infection in utero, we first compared the percentage of $\gamma\delta$ T cells among all T cells in cord blood samples derived from 19 CMV-infected newborns versus 22 control CMV-uninfected newborns. In CMV-infected newborns, the percentage of $\gamma\delta$ T cells was significantly higher than in CMV-uninfected newborns (Fig. 1 A). To exclude the possibility that this higher percentage of $\gamma\delta$ T cells was the result of a decreased number of $\alpha\beta$ T cells, we determined the absolute number of $\gamma\delta$ T cells per microliter of blood. Indeed, significantly more $\gamma\delta$ T cells were present per microliter of cord blood in CMV-infected newborns in comparison with controls (Fig. 1 B). The higher number of $\gamma\delta$ T cells correlated with a higher percentage of $\gamma\delta$ T cells expressing the proliferation marker Ki-67 in CMV-infected newborns (Fig. 1 C).

The expansion of $\gamma\delta$ T cells in CMV-infected newborns is restricted to V$\gamma9^-$ cells, irrespective of the usage of the V$\delta$ chain

To further define specific subsets of $\gamma\delta$ T cells in cord blood of CMV-infected newborns, flow cytometry analysis was performed with antibodies specific against V$\gamma9$, V$\delta1$, V$\delta2$, and V$\delta3$. In combination with the pan-$\gamma\delta$ TCR antibody, the V$\gamma9$ antibody can make distinction between V$\gamma9^+$ and V$\gamma9^-$...
yδ T cells (Fig. 2 A). Vγ9 is the only member of the VγII family; thus the Vγ9− cells express Vγ chains of the VγI family (Hayday, 2000). The combination of Vδ1, Vδ2, and Vδ3 antibodies stained the vast majority (~90%; unpublished data) of the cord blood yδ T cells. This approach allows us to identify six yδ T cell subpopulations in cord blood: Vγ9+Vδ1+, Vγ9−Vδ1+, Vγ9∗Vδ2+, Vγ9−Vδ2+, Vγ9+Vδ3+, and Vγ9−Vδ3+. We detected higher percentages of yδ T cells negative for Vγ9, including Vγ9−Vδ1+, Vγ9−Vδ2+, and Vγ9−Vδ3+ yδ T cells in CMV-infected newborns compared with uninfected newborns (Fig. 2, A and B). On a selected number of CMV-uninfected and CMV-infected newborns, we performed a more detailed analysis of the γ chain usage. In CMV-uninfected newborns, there was a slight preference for Vγ4 and Vγ9, whereas upon CMV infection the VγI family members Vγ4 and Vγ8 were highly expanded (Fig. S1).

Figure 2. The expansion of yδ T cells in CMV-infected newborns is restricted to Vγ9− yδ T cells, irrespective of the usage of the Vδ chain. (A) Expression of Vγ9 versus Vδ1, Vδ2, or Vδ3 by yδ T cells from CMV-infected newborn Pos13 (top) and CMV-uninfected newborn Neg4 (bottom). Numbers in dot plots are the percentages of total T cells. (B) Box-and-whisker graph (defined as in Fig. 1) of the percentages of Vγ9+Vδ1+, Vγ9−Vδ1+, Vγ9−Vδ2+, Vγ9+Vδ3+, and Vγ9−Vδ3+ yδ T cell subpopulations of yδ T cells expressed as a percentage of total T cells. CMV+, n = 10–18; CMV−, n = 14–22.
γδ T cells from CMV–infected newborns are activated and differentiated

Next, we evaluated whether the expansion of fetal γδ T cells after congenital CMV infection was accompanied by activation and/or differentiation of these cells. A significant proportion of γδ T cells from CMV–infected newborns expressed the activation marker HLA-DR, whereas expression was virtually absent in uninfected controls (Table I). Downregulation of CD27 and CD28 expression has been shown to be associated with advanced or late differentiation in CD8 αβ T cells upon CMV infection (Appay et al., 2002; Marchant et al., 2003; van Leeuwen et al., 2006). These markers have also been used to identify differentiated human γδ T cells (Morita et al., 2007). Although CD27−CD28− γδ T cells were absent from CMV–infected newborns, a large proportion of γδ T cells showed this phenotype in CMV–infected newborns (Table I). This differentiation was most pronounced in the Vγ9− γδ T cell subpopulation (unpublished data). Collectively, these data clearly show that upon congenital CMV infection, γδ T cells are activated, undergo cell division, and become differentiated.

Expression of NK receptors (NKR)s, cytotoxic mediators, and IFN−γ is highly increased in γδ T cells of CMV–infected newborns

To gain insight into the function of fetal γδ T cells in newborns with congenital CMV infection, we compared the gene expression profiles of γδ T cells derived from three CMV–infected newborns versus three CMV–uninfected newborns. 1,622 genes were increased and 654 decreased upon infection (using the selection criteria described in Materials and methods; M > 0.05, P < 0.05). More than 100 genes associated with cell cycle showed increased expression upon CMV infection (as analyzed with DAVID; not depicted), coinciding with the expansion data (Fig. 1).

| Marker                  | CMV+               | CMV−               | p (CMV+ versus CMV−) |
|------------------------|--------------------|--------------------|----------------------|
| HLA-DR                 | 6.34 (2.87–10.68)  | 0.58 (0.27–0.80)   | <0.0001              |
| CD27− CD28−            | 43.82 (31.97–48.98)| 0.22 (0.13–0.45)   | <0.0001              |
| CD94+                  | 33.21 (23.68–57.20)| 4.44 (3.61–5.56)   | <0.0001              |
| NK2A+                  | 10.99 (4.27–20.64) | 3.84 (2.33–5.11)   | 0.0464               |
| NK2C+                  | 28.00 (10.77–35.54)| 1.17 (0.91–3.68)   | <0.0001              |
| CD158a/h+              | 14.70 (6.61–30.53) | 1.56 (0.94–3.34)   | <0.0001              |
| CD158b/j+              | 46.60 (29.07–51.38)| 2.75 (2.56–3.32)   | <0.0001              |
| NKG2A+                 | 69.18 (64.42–76.76)| 52.38 (46.26–63.08)| 0.0001               |
| KLRG1+                 | 38.85 (27.80–45.41)| 24.70 (16.73–28.72)| 0.0009               |
| perforin+              | 61.50 (51.97–73.32)| 0.67 (0.48–1.77)   | <0.0001              |
| granzyme A+            | 69.82 (57.97–79.63)| 4.38 (2.82–6.28)   | <0.0001              |
| CX3CR1+                | 43.21 (25.08–57.96)| 1.11 (0.65–2.10)   | 0.0003               |

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Cytotoxic mediators. In general, cytotoxic lymphocytes can kill target cells by two main mechanisms: exocytosis of granule–associated molecules, such as granzymes, perforin, and granulysin, or binding to receptors with ligands of the TNF superfamily (e.g., FasL and TRAIL). Among the >47,000 transcripts analyzed, the two genes displaying the most increased expression upon CMV infection were members of the granzyme family: granzyme B and granzyme H (Fig. 3). In addition, other granzyme family members (granzyme A and granzyme M), perforin, granulysin, FasL, and TRAIL were increased (Table S1). In CMV–uninfected newborns, there was either no or very few γδ T cells (CD94/NKG2A/NKG2C, CD158a/h [KIR2DL1/KIR2DS1], and CD158b/j [KIR2DL2/KIR2DS2]) or a significant fraction of γδ T cells (NK2D2 and KLRG1) expressing NKR on their membrane, as determined by flow cytometry (Table I). In CMV–infected newborns, significantly more γδ T cells expressed all these NRKs (Table I).

Chemokines and chemokine receptors. The genes for the chemokines CCL3 (MIP-1α), CCL4 (MIP-1β), and CCL5 +. CMV+, p (CMV+ versus CMV−)= 8–18; CMV−, n = 9–22.
measured by the mean fluorescence intensity (MFI), was consistently much higher in CD27−CD28− γδ T cells than in CD27−CD28+ γδ T cells (median within CD27−CD28− γδ T cells, 318 MFI; median within CD27+CD28+ γδ T cells, 102 MFI; P = 0.0023). Upon a brief polyclonal stimulation in vitro, the majority of CD27−CD28− differentiated γδ T cells of CMV-infected newborns produced IFN-γ (Fig. 4 B), whereas the CD27+CD28+ γδ T cells produced significantly less IFN-γ (median within CD27−CD28− γδ T cells, 68%; median within CD27+CD28+ γδ T cells, 22%; P = 0.0006). T-bet and IFN-γ expression within γδ T cells from CMV-uninfected newborns were similar to the expression found within CD27−CD28− γδ T cells from CMV-infected newborns (unpublished data).

The CDR31 and CDR32 are highly restricted upon congenital CMV infection

To study the impact of CMV infection during fetal life on the TCR repertoire of γδ T cells, we assessed the degree of junctional diversity of the complementary-determining-region-3 (CDR3) of the Vδ1 (CDR31) and Vδ2 (CDR32) chains by spectratyping on 11 CMV-uninfected and 13 CMV-infected cord blood samples. CMV-uninfected cord blood samples showed polyclonal profiles for both CDR31 (as described previously; Beldjord et al., 1993) and CDR32. In contrast, the CDR31 and CDR32 repertoires became highly restricted in the vast majority of CMV-infected newborns (Fig. 5 A and Fig. S3). To quantify this restriction, we calculated an index of oligoclonality for CDR31 and CDR32 as described previously (Déchanet et al., 1999; Pitard et al., 2008). For both CDR3s, the index of oligoclonality
was significantly higher in CMV-infected newborns than in CMV-uninfected newborns (Fig. 5 B). Moreover, the restriction of CDR.3δ1 of CMV-infected newborns was enriched for the same length (11 aa) in all CMV-positive newborns (Fig. 5 A, arrows). This length was either absent or minimally present in CMV-uninfected newborns (Fig. 5 A). In contrast to CDR.3δ1, the length of the enriched CDR.3δ2 sequences in CMV-infected newborns varied from newborn to newborn (Fig. S3). It is of note that CMV-infected newborn Pos10 showed a polyclonal CDR.3δ1 repertoire corresponding with the absence of CD27−CD28− differentiated Vδ1+γδ T cells (Fig. 5 A). In contrast, Vδ2+γδ T cells of this newborn were well differentiated, corresponding to a restricted CDR.3δ2 repertoire (Fig. S3).

Public germline-encoded CDR.3δ1 and CDR.3γ8 sequences are highly enriched in CMV-infected newborns

Because the CDR.3δ1 of CMV-infected newborns was highly enriched at 11 aa, we wondered whether this region included the same or similar sequences. Strikingly, at amino acid level in all 12 sequenced CMV-infected newborns the CDR.3δ1 of 11 aa had exactly the same sequence: CALGELGDDKLIF, or ELGDD for short (Table S2; Fig. 5 A). At the nucleotide level, two variants were observed: the first D of ELGDD was either formed by the g of the diversity gene δ3 (Dδ3) and the c of the joining gene δ1 (Jδ1) or completely formed by the gat of the Dδ3 (Table S2, dark gray). Besides ELGDD itself, few longer variants were present in some CMV-infected newborns, which were enriched as well, containing one (Pos3) or two (Pos3 and Pos11) extra Ts after the first D of ELGDD (Table S2). In contrast to the other CDR.3δ1 sequences, the highly enriched ELGDD sequence did not contain P/N additions and was thus completely germline encoded (Table S2).

In comparison with CDR.3δ1, the degree of shared CDR.3δ2 sequences among the CMV-infected newborns was much less clear (Table S3). Among eight CMV-infected newborns, three exhibited enrichment of the same CDR.3δ2 sequence (Pos8, Pos9, and Pos10; Table S3). No other obvious similarities were found between different enriched CDR.3δ2 sequences of different CMV-infected newborns. Vδ1 almost always paired with Jδ1, whereas Vδ2 had a preference for Jδ3 (Table S2 and Table S3). Sequencing of CDR.3δ1 and CDR.3δ2 of CMV-uninfected newborns confirmed the polyclonal repertoire as found by spectratyping (Table S2 and Table S3).

We wondered whether the Vδ1 chain, containing the public CDR.3δ1, from CMV-infected newborns cells had a preference for pairing with specific Vγ chains. By costaining of Vδ1 and Vγ2/3/4, Vγ5/3, Vγ8, or Vγ9, we determined that within CMV-uninfected newborns Vδ1 had a preference

Figure 4. Differentiated (CD27−CD28−) γδ T cells from CMV-infected newborns express highly the transcription factor T-bet and produce high levels of IFN-γ. Flow cytometry plots for T-bet (A) and IFN-γ (B), gated on CD27−CD28− versus gated on CD27+CD28+ γδ T cells of a CMV-positive newborn (Pos12), representative of seven CMV-infected newborns. Cells were stimulated for 4 h with PMA/ionomycin before intracellular staining for IFN-γ. Unstimulated cells showed no IFN-γ staining.
Figure 5. The CDR3\delta1 and CDR3\delta2 repertoire of γδ T cells from CMV-infected newborns are oligoclonal, and CDR3\delta1 is highly enriched for a single sequence. (A) Spectratyping plots of the CDR3\delta1 of CMV-uninfected and CMV-infected newborns. Each box represents one donor. The numbers at the left top of each box represent the percentage of Vδ1+ γδ T cells, expressed as percentage of total T cells. The numbers at the right top of each box represent the percentage of CD27⁻ CD28⁻ cells of Vδ1+ γδ T cells. The arrows indicate the sequences at CDR3\delta1 of 11-aa size of the CMV-infected newborns that have been sequenced: CALGELGDDKLIF (Table S2). (B) Index of oligoclonality for CDR3\delta1 and CDR3\delta2, determined as described in Materials and methods. Lines indicate medians.
for pairing with Vγ2/3/4 (Fig. 6 A), whereas Vβ8 and Vβ3 had rather a preference for Vγ9 and Vγ5/3, respectively (not depicted). In contrast, in CMV-infected newborns with highly expanded γδ T cells, Vβ1 had a clear preference for pairing with Vγ8 (Fig. 6 A), whereas Vβ2 and Vβ3 had rather a preference for Vγ2/3/4 (not depicted). Because of this preferential pairing of Vβ1 with Vγ8, we performed spectratyping for CDR3γ8 on six CMV-uninfected and six CMV-infected newborns. The CDR3γ8 of CMV-uninfected newborns showed a polyclonal repertoire. In contrast, the CDR3γ8 repertoire became highly restricted in CMV-infected newborns, showing a high enrichment at a length of 11 aa in five out of six CMV-infected newborns. The sixth CMV-infected newborn (Pos13) had a high enrichment at 12 aa (Fig. 6 B). Sequencing revealed that the CDR3γ8 sequences at 11 aa contained all the same sequence: CATWDTTGWFKIF (DTTGW for short). Pos13 had an extra Y after D (Fig. 6 B; Table S4). This sequence was not detected in CMV-uninfected newborns (Table S4). As for the public CDR3γ8 sequence, the public CDR3γ8 was completely germline encoded (Table S4).

The public Vγ8Vδ1 TCR reacts against CMV-infected target cells
To verify whether the public Vγ8Vδ1 TCR reacts against CMV-infected target cells, we generated γδ T cell clones expressing the public TCR containing the CDR3δ1-ELGDD and CDR3γ8-DTTGW from CMV-infected newborns Pos4 (11 public clones) and Pos6 (21 public clones). All clones expressing CDR3δ1-ELGDD coexpressed CDR3γ8-DTTGW, whereas clones with a different CDR3δ1 expressed other CDR3γ’s (unpublished data), showing in a direct way the preferential pairing between CDR3δ1-ELGDD and CDR3γ8-DTTGW. A brief coincubation (6 h) of public

Figure 6. The Vδ1 chain on γδ T cells of CMV-infected newborns preferentially pairs with a public Vγ8 chain. (A) The percentage of Vδ1+ γδ T cells positive for Vγ2/3/4, Vγ5/3, Vγ8, or Vγ9 determined in three CMV-infected (Pos4, Pos6, and Pos13) and three CMV-uninfected newborns. (B) Spectratyping for CDR3γ8 of six CMV-uninfected newborns (top row) and six CMV-infected newborns (bottom row). The arrow indicates the public CDR3γ8 sequence CATWDTTGWFKIF of 11 aa (Table S3). The CDR3γ8 of Pos13 contains 1 aa more (Y).
Differentiation and oligoclonal expansion of fetal \( \gamma\delta \) T cells can occur early during gestation

To explore the possibility that \( \gamma\delta \) T cells could develop a response toward CMV infection early during fetal life, we analyzed the \( \gamma\delta \) T cells from fetal cord blood samples collected between 20 and 29 wk of gestation (from 12 CMV-

clones with CMV-infected human embryonic lung fibroblasts induced IFN-\( \gamma \) production, which was blocked by the presence of a soluble anti-CD3 antibody (OKT3) showing the involvement of the public V\( \gamma8\delta1 \) TCR/CD3 complex in the recognition of CMV-infected target cells (Fig. 7 and Fig. S4). Control \( \gamma\delta \) T cell clones of CMV-uninfected newborns did not show CMV-induced IFN-\( \gamma \) production. To gain insight into the antiviral activity of the \( \gamma\delta \) T cell clones, we conducted additional experiments. Public clones killed infected target cells (Fig. 8 A) and inhibited CMV replication (between one and two \( \log_{10} \) inhibition; Fig. 8 B), whereas control V\( \gamma9\delta2 \) T clones had no or only a moderate effect (Fig. 8).

Figure 7. \( \gamma\delta \) T cell clones expressing the public V\( \gamma8\delta1 \) TCR display reactivity against CMV-infected cells via TCR/CD3. Clones were coincubated for 6 h with human embryonic fibroblasts not infected (white bars) or infected (gray bars) with CMV (TB40/E). During coincubation either a control IgG2a antibody (ctrl) or the anti-CD3 antibody OKT3 (anti-CD3) was present in soluble form. Results are shown for one public clone from CMV-infected newborn Pos4 (V\( \gamma8\delta1\)-Pos4) and for one public clone of CMV-infected newborn Pos6 (V\( \gamma8\delta1\)-Pos6) and are representative of five independent experiments involving 11 different public V\( \gamma8\delta1 \) clones.

Figure 8. Public V\( \gamma8\delta1 \) clones kill CMV-infected target cells and inhibit CMV replication in vitro. (A) CMV-infected (TB40/E) human embryonic fibroblasts were coincubated with either \( \gamma\delta \) T cell clones expressing the public V\( \gamma8\delta1 \) TCR (derived from CMV-infected newborns Pos4 and Pos6) or a control V\( \gamma9\delta2 \) clone (derived from a CMV-uninfected newborn) at the indicated effector to target ratios. After 4 h of coincubation, the level of DNA fragmentation in the target cells was quantified. Results are representative of three independent experiments. (B) Human embryonic fibroblasts were incubated with CMV for 2 h, washed, and incubated with medium alone, with a public V\( \gamma8\delta1 \) clone from CMV-infected newborn Pos6 or with a control V\( \gamma9\delta2 \) clone from a CMV-uninfected newborn. After 7 d, the quantity of infectious CMV from the supernatant was determined by a plaque assay (PFU, plaque forming units). Shown are the mean \( \pm \) SEM of quadruplicate determinations. Results are representative of two independent experiments.

negative and 13 CMV-positive fetuses). At these earlier gestation times, the \( \gamma\delta \) T cells were already clearly differentiated (down-regulation of CD27 and CD28) and showed high expression of perforin (Fig. 9 A), granzyme A, and NKR (not depicted). From four CMV-infected fetuses, we performed spectratyping and sequencing for CDR3\( \delta1 \) at time of delivery and at earlier gestation time (Fig. 9 B). We found that the CDR3\( \delta1\)-ELGDD sequence was already enriched at as early as 21 wk of gestation (Fig. 9 B and Table S2). Few other CDR3\( \delta1 \) sequences that were present at early gestation time were also present at time of delivery (Table S2, fetus Pos4 [14 aa] and fetus Pos13 [17 aa]). As observed at time of
delivery, CDR3δ2 spectratyping showed more variability between fetuses (unpublished data). Furthermore, CDR3δ2 appeared to vary with time within the same fetus (Table S3, compare CDR3δ2 sequencing data of Pos4 at 20 wk, 5 d and at 40 wk, 0 d of gestation). Thus, the enriched CDR3δ1-ELGDD sequence appeared early during gestation in CMV-infected fetuses and remained present with time. In contrast, the enriched sequences of the CDR3δ2 were variable from one fetus to the other and changed during gestation time. Furthermore, the CDR3γ sequence associated with the CDR3δ1-ELGDD sequence, namely CDR3γ8-DTTGW, was also already enriched at as early as 21 wk of gestation.

Figure 9. Differentiation and oligoclonal (CALGELGDDKLIF) expansion of fetal γδ T cells can occur early during gestation. (A) Expression of CD27/CD28 and perforin by γδ T cells from a representative CMV-uninfected (bottom) and a representative CMV-infected fetus (Pos4; top), both at the gestational age of 20–21 wk. Dot plots are presented and numbers indicate the percentages of γδ T cells negative for CD27 and CD28 and positive for perforin. CMV−, results are representative of 12 (CD27/CD28) and 7 (perforin) fetuses (gestation range: 20 wk, 3 d–29 wk, 2 d); CMV+, results are representative of 13 (CD27/CD28) and 5 (perforin) fetuses (gestation range: 20 wk, 5 d–29 wk, 2 d). (B) Spectratyping for CDR3δ1 of four CMV-infected fetuses for which we had blood samples both at the time of delivery and at earlier gestation times. The enrichment for the CDR3δ1 size of 11 aa consists of the sequence CALGELGDDKLIF (Table S2).
as early as 21 wk of gestation. Despite the recent description of selective impairments of γδ T cells in preterm infants (Gibbons et al., 2009), γδ T cell are able to develop robust responses toward CMV infection in utero at as early as 21 wk of gestation.

**DISCUSSION**

In this study, we demonstrate that CMV infection in utero leads to the oligoclonal expansion and differentiation of fetal γδ T cells, which express high levels of NKR and cytotoxic mediators and produce IFN-γ. Both activating (e.g., activating KIR, NKG2C, and NKG2D) and inhibitory (e.g., inhibitory KIR, NKG2A, and KLRG1) NKRs were highly expressed in γδ T cells derived from congenitally infected newborns. This would allow them to sense CMV-induced changes in infected target cells; HLA-E (ligand for KIR) expression is increased upon CMV infection, whereas classical MHC class I (ligands for KIR) expression is decreased (Wilkinson et al., 2008). In comparison with conventional T cells, it has been described that adult γδ T cells express high levels of NKR, like members of the C-type lectin and the KIR family (Battistini et al., 1997; De Libero, 1999; Pennington et al., 2005). We confirmed the expression of KLRG1 on γδ T cells in CMV-uninfected newborns (Eberl et al., 2005) and also showed that NKG2D is constitutively expressed. In contrast, unlike NK cells (Dalle et al., 2005), other NKR families (CD94/NKG2x and KIR family members) were not expressed or were expressed at low levels on γδ T cells from CMV-uninfected newborns. Thus the majority of NKR expression on adult γδ T cells is likely to be the consequence of infections after birth. CMV infection in utero induced the up-regulation of various cytotoxic mediators in fetal γδ T cells, including almost all members of the granzyme family, perforin, granulysin, FaSL, and TRAIL. Perforin and granulysin are membrane-disrupting molecules and most granzymes have been shown to be involved in killing of target cells, with most evidence for granzyme B (Lieberman, 2003; Chowdhury and Lieberman, 2008). In addition, other granzyme-mediated antiviral mechanisms have been recently described: granzyme A plays a proinflammatory role (Metkar et al., 2008), granzyme M targets α-tubulin (Bovenschen et al., 2008), and granzyme H cleaves La, a phosphoprotein involved in cellular and viral RNA metabolism (Romero et al., 2009). It is of note that granzyme H cleaves an adenovirus-encoded granzyme B inhibitor (Andrade et al., 2007). Analysis of the profile of cytokine genes expressed in fetal γδ T cells derived from CMV-infected newborns revealed the restricted high expression of IFN-γ. In parallel, we detected elevated levels of the T-box transcription factors T-bet and eomes, which are involved in the rapid and vigorous IFN-γ production by γδ T cells (Yin et al., 2002; Chen et al., 2007). In contrast, expression of other transcription factor genes like GATA3 (Th2) or ROR-γt (Th17) was not affected, coinciding with the absence of modulation of cytokine genes associated with these Th subsets. Only two chemokine receptors were significantly increased in the microarray analysis: CCR5 and CX3CR1 (fractalkine receptor). Fractalkine can be produced by endothelial cells in the context of CMV infection (Bolovan-Fritts et al., 2004), thus possibly attracting differentiated CX3CR1+ fetal γδ T cells to the site of infection. Together, our data indicate that fetal γδ T cells generated in utero during CMV infection are equipped with a range of antiviral effector mechanisms, including IFN-γ production and granule-mediated cytotoxicity. Indeed, γδ T cell clones generated from CMV-infected newborns killed CMV-infected cells and limited CMV replication in vitro. It is therefore likely that they participate in the limitation of the viral spread in the fetus. In kidney-transplanted patients with acute CMV infection, expansion of γδ T cells is associated with the clinical resolution, suggesting a protective role of the expanded γδ T cells (Lafarge et al., 2001; Halary et al., 2005).

We demonstrated that CMV infection during fetal life leads to the oligoclonal expansion of γδ T cells, which is characterized by highly restricted CDR31 and CDR32 repertoires and by the high enrichment of a public CDR31-CDR3γ8 sequence. Expanded γδ T cells were negative for Vγ9 and included Vδ1+, Vδ2+, and Vδ3+ cells. In contrast, in adult CMV-infected kidney-transplanted patients, expanded γδ T cells do not include Vδ2+ cells and there is no restriction of CDR3δ2 (Déchanet et al., 1999). Vγ9-Vδ2+ γδ T cells are very rare in the adult (Morita et al., 1994), providing a possible explanation of why Déchanet et al. (1999) did not detect any expansion of this subset in adults (Pitard et al., 2008).

TCR-δ chains have the highest potential diversity in the CDR3 loop (~10^16 combinations) among all antigen receptor chains (TCR-α, TCR-β, TCR-γ, TCR-δ, IgH, and IgL) because multiple D gene segments can join together, all D gene segments can be read in all three open reading frames, and N nucleotides can be inserted into the junctions of each of the joining segments (Chien and Konigshofer, 2007). Therefore, it was surprising to identify a high enrichment of exactly the same CDR31 sequence (i.e., public CDR3) in all fetuses with differentiated Vδ1+ γδ T cells upon congenital CMV infection (ELGDD). It has been suggested that much of the diversity of the CDR3 junctions of the δ chain may confer different affinities of the γδ TCR rather than the ability to recognize different ligands (Chien and Konigshofer, 2007). In addition, in adult CD8 αβ T cells, public CMV-reactive TCR sequences bind the MHC–peptide complexes with higher affinity than private MHC peptide–specific TCR sequences (Trautmann et al., 2005; Day et al., 2007). This suggests that the public CDR31-ELGDD is enriched by recognition of a CMV-induced ligand with high affinity. Our results show for the first time, to our knowledge, the expansion of a public γδ TCR CDR3 in the context of an infection. Furthermore, we demonstrated that the public CDR31 pairs with a public CDR3γ8 sequence (DTTGW), indicating that both the γ and δ chain are important for the recognition of the putative ligand. In addition, this public Vγ8Vδ1 TCR showed reactivity against CMV-infected...
target cells in vitro. It is of note that both the CDR3δ1-ELGD and CDR3γ8-DTTGW were germline encoded, as the CDR3δ involved was only formed by the V61 gene, one Dδ gene (Dδ3) and the Jδ1 gene, and the CDR3γ8 by the Vγ8 gene and the JγP1 gene, without any addition of P/N nucleotides. Similarly, the mouse T22/T10-binding CDR3δ does not contain N nucleotides (Adams et al., 2005; Chien and Konigshofer, 2007). This contrasts highly with αβ T cells, where the most critical amino acids in the CDR3α and CDR3β involved in the recognition of the MHC–peptide complex are encoded either completely or partially by N nucleotides (Davis et al., 1998; Chien and Konigshofer, 2007).

Despite the immaturity of the neonatal immune system and possible mechanisms of immunosuppression by regulatory T cells (Mold et al., 2008), CMV infection is efficient in stimulating vigorous responses of both γδ T cells and CD8 αβ T cells (Marchant et al., 2003) during fetal life. Studies in mice show that the protective role of γδ T cells in early life is not dependent on αβ T cells (Ramsburg et al., 2003). Conversely, in a model of West Nile virus infection, it has been shown that γδ T cells facilitate the CD8 αβ T cell response (Wang et al., 2006). In comparison with adult DC, fetal DC shows impaired functions (Goriely et al., 2004; Brandes et al., 2009), as shown in adult cells. Such mechanisms could contribute to the development of functional CD8 αβ T cell responses to CMV infection during fetal life (Marchant et al., 2003).

We conclude that human γδ T cells can mount a vigorous response to CMV infection during development in utero, providing an important mechanism by which the fetus can fight pathogens. Identification of the γδ TCR ligands induced upon CMV infection, like the putative ligand of the public Vγ8Vδ1 TCR, will likely be useful to design novel vaccination strategies against viral infection in early life.

MATERIALS AND METHODS

Study population. This study was approved by the Hôpital Erasme and Hôpital Saint-Pierre ethical committees. Women with suspected primary CMV infection were referred to the Fetal Medicine Units of the Hôpital Erasme or Hôpital Saint-Pierre. Diagnosis of primary maternal infection was based on anti-CMV IgG seroconversion or on the detection of high titers of anti-CMV–specific IgM, as described previously (Liesnard et al., 2000).

After maternal informed consent, 20–50 ml of cord blood was collected at birth (full term, >37 wk gestation). In some cases, fetal cord blood was collected at earlier gestation ages (<1 ml by cordocentesis). Diagnosis of congenital infection was based on the detection of CMV genome by PCR and/or by viral culture on amniotic fluid and/or on newborn urine collected during the first week of life. The study included 19 CMV-infected newborns and 22 uninfected control newborns as well as 13 infected and 12 uninfected fetuses. Symptomatic congenital infection was diagnosed in fetus Pos12 who had brain lesions at antenatal magnetic resonance imaging and an abnormal postnatal neurological development and in fetus Pos5 who had an abnormal postnatal neurological development.

Flow cytometry. The following antibodies were used: CD3–blue (clone SP34–2), γδ–PE (11F2), γδ–FITC (11F2), CD27–APC (L128), CD27–FITC (L128), CD94–APC (HP–3D9), CD158α–FITC (HP–3E4), CD158β–FITC (CH–L), HLA–DR–Cy7 (L243), NKGD2–APC (1D11), perfos–FITC (BG9), granzyme A–FITC (CB9), Ki–67–FITC (B56), and IFN–γ–FITC (25723.11; BD); V62–FITC (IMMU389), NKGD2–PE (Z199), CD3–ECD (UCHT1), and CD28–ECD (CD28.2; Beckman Coulter); V81–FITC (TS1; Thermo Fisher Scientific); NKGD2–APC (134591; R&D Systems); CXCR3–PE (2A9–1; MBL International); and T–bet–PE (4B10; eBioscience). Vγ5–PC5 (IMMU360) and Vδ3–FITC (PI1.5B) were derived from Beckman Coulter via custom design service. Vγ2/3–biotin and Vδ2/3–FITC (23D12), Vγ4–FITC, Vγ5/3–biotin (56.3), and Vγ8–biotin (R4.5.1) were provided by D. Wesch (Institute of Immunology, University of Kiel, Kiel, Germany; Kabelitz et al., 1994; Himz et al., 1997; Wesch et al., 1998). KLRG1–Alexa Fluor 488 (13F12) was provided by H. Pircher (University of Freiburg, Freiburg, Germany; Marcolino et al., 2004) and unlabeled Vδ3 antibody by E. Scotet (Institut National de la Santé et de la Recherche Médicale U601, Nantes, France; Peyrat et al., 1995). Staining was done on whole blood. Red blood cells were lysed using FACS Lysing solution (BD). The absolute number of γδ T cells in whole blood was determined using Trucount beads (BD). Intracellular staining for perforin–FITC, granzyme A–FITC, and Ki–67–FITC was performed with the Perm 2 kit (BD) and for T–bet–PE with the Fixperp staining buffer (eBioscience). For the detection of IFN–γ, PBMCs were stimulated for 4 h with 10 ng/ml PMA and 2 μM ionomycin in the presence of 2 μM monensin. Staining was done using the Cytofix/Cytoperm kit (BD).

Cells were run on the CyAn flow cytometer equipped with three lasers (405, 488, and 633 nm) and data were analyzed using Summit 4.3 (Dako).

Microarray analysis. PBMCs were isolated from cord blood by Lymphoprep gradient centrifugation (Axis–Shield). After depletion of remaining red blood cells and CD4+ cells by magnetic cell sorting (Miltenyi Biotech), CD3+γδ lymphocytes were sorted till high purity (>99%) with a MoFlo sorter (Dako). The γδ T cell yield varied from 80,000–300,000 cells per cord blood sample. Total RNA was isolated using the RNeasy Micro kit (QIAGEN) from sorted γδ T cells derived from three CMV-infected newborns and three CMV-uninfected newborns. RNA concentration was measured using the NanoDrop (Thermo Fisher Scientific) and RNA quality was assessed using the Bioanalyzer 2100 (Agilent Technologies). cRNA was amplified into biotin–labeled complementary RNA (cRNA) by one–round in vitro transcription using the Premier kit (Applied Biosystems). The cRNA was fragmented and hybridized on the Human Genome U133 Plus 2.0 GeneChip (Affymetrix). Staining and scanning was done on the Affymetrix platform. The procedures, from RNA quality control to generation of raw data (CEL files), were performed at DNAVision (Gosselies, Belgium). The raw data were analyzed using the Affy package of Limma (linear models for microarray data; www.biodoncult.org), including fitting a linear model (limfit) as described previously (Vermijlen et al., 2007). M– and A–values for each gene were generated. M (log2 of the fold change) is related to the degree of differential expression between the γδ T cells from CMV–infected newborns versus γδ T cells from CMV–uninfected newborns, whereas A is a measurement of the mean signal intensity. Genes were regarded as differentially expressed if the absolute M–value was >0.5 with a p–value <0.05. Genes with M–values >0.5 are enriched in the γδ T cells derived from CMV–infected newborns, whereas genes with M–values <−0.5 are enriched in the γδ T cells derived from CMV–uninfected newborns. The Database for Annotation, Visualization and Integrated Discovery (DAVID; http://david.abcc.ncifcrf.gov/) was used to assist in the discovery of functionally related groups of differentially expressed genes. Microarray data and procedures.
were deposited at Array Express (www.ebi.ac.uk/arrayexpress) under accession no. E-MEXP-2055.

**Spectratyping.** Total RNA was isolated from PBMC of cord blood of CMV-infected newborns and CMV-uninfected newborns, after which cDNA was generated using the First Strand cDNA synthesis kit (Fermentas). PCR (40 cycles) was performed with C8′-GTGAAATCTTCCAGCAGCAGAAGCAGCAGAAGGACAGCAGCAGAAGGCCTTAA-3′ and V8′ (5′-GCAAGACAGGGAAGAGCCTTAA-3′). Then a run-off reaction (one cycle) was performed using the fluorescently labeled C8-FAM primer (5′-ACGGATGGTTTGGTATGAGGCTGA-3′) and V8′ (5′-GAGYTTTGTTTCAGC-3′) and V8′ (5′-GCAAGACAGGGAAGAGCCTTAA-3′). The reaction products were run on a capillary sequencer (ABI3730xl or ABI3130xl analyzer) at DNAVision. The fluorescence intensity was analyzed using Peak Scanner 1.0 (Applied Biosystems). The index of oligoclonality was calculated as described previously (Déchanet et al., 1999; Pitard et al., 2008).

**Sequencing.** As described in Spectratyping, PCR (40 cycles) was performed on cDNA to amplify the sequences that contain the CDR3. CDR1, CDR2, or CDR3. PCR products were TA cloned according to the instructions of the manufacturer (Invitrogen). Sequencing was performed on recombinant plasmids purified from bacterial clones by cycle sequencing instructions of the manufacturer (Invitrogen). Sequencing was performed on 2, or CDR3

**CMV replication assay.** Confluent monolayers of human embryonic lung fibroblasts (HEL299) in flat-bottom 96-well plates were incubated with CMV (TB40/E) for 2 h (MOI 0.1), washed, and incubated with medium alone, with a public VγVδ1 clone or with a control VγVδ2 clone (150,000 cells per well). After 7 d, the quantity of infectious CMV from the supernatant was determined in quadruplicate by standard plaque assay titration (in plaque forming units).

**Statistical analysis.** Differences between CMV-infected newborns and CMV-uninfected newborns were determined using the nonparametric Mann–Whitney test using InStat software (GraphPad Software, Inc.). Differences were regarded as significant at P < 0.05.

**Online supplemental material.** Fig. S1 shows the Vγ chain expression of CMV-infected and CMV-uninfected newborns. Fig. S2 shows the association of the late differentiation phenotype of Vγ T cells with the expression of cytotoxic mediators and chemokine receptor CX3CR1. Fig. S3 shows the CDR3 repertoire. Table S1 provides an overview of differentially expressed genes in Vγ T cells from CMV-infected newborns versus CMV-uninfected newborns. Tables S2–S4 contain the sequencing data for CMV-infected and CMV-uninfected newborns. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20090348/DC1.

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**REFERENCES**

Adams, E.J., Y.H. Chien, and K.C. Garcia. 2005. Structure of a gammadelta T cell receptor in complex with the nonclassical MHC T22. *Science.* 308:227–231. doi:10.1126/science.1106885

Andrade, F., E. Fellows, D.E. Jenne, A. Rosen, and C.S. Young. 2007. Granulocyte H destroys the function of critical adenoviral proteins required for viral DNA replication and granulocyte B inhibition. *EMBO J.* 26:2148–2157. doi:10.1038/sj.emboj.7601650

Appay, V., P.R. Dunbar, M. Callan, P. Kleneman, G.M. Gillespie, L. Papagno, G.S. Ogg, A. King, F. Lechner, C.A. Spina, et al. 2002. Memory CD8+ T cells vary in differentiation phenotype in different persistent virus infections. *Nat. Med.* 8:379–385. doi:10.1038/nm0402-379

Battistini, L., G. Borsellino, G. Sawicki, F. Poccia, M. Salvetti, G. Ristori, and C.F. Brosnan. 1997. Phenotypic and cytokine analysis of human peripheral blood gamma delta T cells expressing NK cell receptors. *J. Immunol.* 159:3723–3730.
Beldjord, K., C. Beldjord, E. Macintyre, P. Even, and F. Sigaux. 1993. Peripheral selection of Vβ1 cells with restricted T cell receptor β gene junctional repertoire in the peripheral blood of healthy donors. J. Exp. Med. 178:121–127. doi:10.1084/jem.178.1.121

Bolovan-Fritts, C.A., R.N. Trout, and S.A. Spector. 2004. Human cyto- megalovirus-specific CD4+ T-cell cytokine response induces fractal kinase in endothelial cells. J. Virol. 78:13173–13181. doi:10.1128/JVI.78.23.13173-13181.2004

Bovenschen, N., P.J. de Koning, R. Quadir, R. Broekhuizen, J.M. Damen, C.J. Froelich, M. Slijper, and J.A. Kummer. 2008. NK cell protease granzyme M targets alpha-tubulin and disorganizes the microtubule network. J. Immunol. 180:8184–8191.

Brandes, K., M. Willingham, G. Bioley, N. Lévy, M. Eberl, M. Luo, R. Tampé, F. Lévy, P. Romero, and B. Moser. 2009. Cross-presenting human gammadelta T cells induce robust CD8+ alpha-beta T cell responses. Proc. Natl. Acad. Sci. USA. 106:2307–2312. doi:10.1073/pnas.0810059106

Brochet, X., M.P. Lefranc, and V. Giudicelli. 2008. IMGT/V-QUEST: the complete expressed human TCR V gamma repertoire by flow cytometry. J. Immunol. 178:2730–2736.

Chen, L., W. He, S.T. Kim, J. Tao, Y. Gao, H. Chi, A.M. Intlekofer, B. Harvey, S.L. Reiner, Z. Yin, et al. 2007. Epigenetic and transcriptional programs lead to default IFN-gamma production by gammadelta T cells. J. Immunol. 178:2730–2736.

Chien. 1998. Ligand recognition by alpha beta T cell receptors. Annu. Rev. Immunol. 16:57–82. doi:10.1146/annurev.immunol.16.1.57

Chien. 1998. Ligand recognition by alpha beta T cell receptors. Annu. Rev. Immunol. 16:57–82. doi:10.1146/annurev.immunol.16.1.57

Chondrology. 100:575–580. doi:10.1038/joas.2001.57

Dalle, J.H., J. Menezes, E. Wagner, M. Blagdon, J. Champagne, Chen, L., W. He, S.T. Kim, J. Tao, Y. Gao, H. Chi, A.M. Intlekofer, B. Harvey, S.L. Reiner, Z. Yin, et al. 2007. Epigenetic and transcriptional programs lead to default IFN-gamma production by gammadelta T cells. J. Immunol. 178:2730–2736.

Dalle, J.H., J. Menezes, E. Wagner, M. Blagdon, J. Champagne, Chen, L., W. He, S.T. Kim, J. Tao, Y. Gao, H. Chi, A.M. Intlekofer, B. Harvey, S.L. Reiner, Z. Yin, et al. 2007. Epigenetic and transcriptional programs lead to default IFN-gamma production by gammadelta T cells. J. Immunol. 178:2730–2736.

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4

De Libero, G. 1999. Control of gammadelta T cells by NK receptors. Microbes Infect. 1:263–267. doi:10.1016/S1286-4579(99)80043-4
Mold, J.E., J. Michaëlsson, T.D. Burt, M.O. Muench, K.P. Beckerman, Marchant, A., V. Appay, M. Van Der Sande, N. Dulphy, C. Liesnard, M. Marchant, A., and M. Goldman. 2005. T cell-mediated immune responses
Liesnard, C., C. Donner, F. Brancart, F. Gosselin, M.L. Delforge, and F. Lewis, D.B., and C.B. Wilson. 2001. Developmental immunology and role of
Levy, O. 2007. Innate immunity of the newborn: basic mechanisms and clini-
Maródi, L. 2006. Neonatal innate immunity to infectious agents.
Lieberman, J. 2003. The ABCs of granule-mediated cytotoxicity: new weap-
Peyrat, M.A., F. Davodeau, I. Houde, F. Romagné, A. Necker, C. Leget, J.P. Cervoni, N. Cerf-Bensussan, H. Vié, M. Bonneville, and M.M. Hallet. 1995. Repertoire analyses of human peripheral blood lymphocytes using a human V delta 3 region-specific monoclonal antibody. Characterization of dual T cell receptor (TCR) delta-chain expressers and alpha beta T cells expressing V delta 3 alpha C alpha-encoded TCR chains. J. Immunol. 155:3060–3067.
Pitard, V., D. Roumanes, X. Lafarge, L. Couzi, I. Garrigue, M.E. Lafon, P. Merville, J.F. Moreau, and J. Déchanet-Merville. 2008. Long-
Ramsburg, E., R. Tigelaar, J. Craft, and A. Hayday. 2003. Age-dependent requirement for γδ T cells in the primary but not secondary protective
Romero, V., D.E. Fellows, D.E. Jinne, and F. Andrade. 2009. Cleavage of La protein by granzyme H induces cytoplasmic translocation and interferes with La-mediated HCV-IRES translational activity. Cell Death Differ. 16:340–348. doi:10.1038/cdd.2008.165
Stagno, S. 2001. Cytomegalovirus. In Infectious Diseases of the Fetus and Newborn Infants. J.S. Remington and J.O. Klein, editors. W.B. Saunders Company, Philadelphia, PA. 389–424.
Tabi, Z., M. Moutafski, and L.K. Borysiewicz. 2001. Human cytomegalovirus pp65- and immediate early 1 antigen-specific HLA class I-restricted cytotoxic T cell responses induced by cross-presentation of viral antigens. J. Immunol. 166:5695–5703.
Toulon, A., L. Breton, K.R. Taylor, M. Tenenhausen, D. Bhavsar, C. Lanigan, R. Rudolph, J. Jameson, and W.L. Havran. 2009. A role for human skin-resident T cells in wound healing. J. Exp. Med. 206:743–750. doi:10.1084/jem.20081787
Trautmann, L., M. Rimbert, K. Echasserieux, S. Saulquin, B. Neveu, J. Déchanet, V. Cerundolo, and M. Bonneville. 2008. Selection of T cell clones expressing high-affinity public TCRs within Human cytomegalovirus-specific CD8 T cell responses. J. Immunol. 175:6123–6132.
van Leeuwen, E.M., G.J. de Blee, J.J. ten Berge, and R.A. van Lier. 2006. Human virus-specific CD8+ T cells: diversity specialists. Immunol. Rev. 211:225–235. doi:10.1111/j.0105-2896.2006.00379.x
Vanhecke, D., B. Verhaest, V. Debaker, G. Leclercq, J. Plum, and B. Vanderkerckhove. 1995. Differentiation to T helper cells in the thymus. Gradual acquisition of T helper cell function by CD3+CD4+ cells.
Vermijlen, D., D. Luo, C.J. Froelich, J.P. Medema, J.A. Kummer, E. Willems, F. Brset, and E. Wisse. 2002. Hepatic natural killer cells exclusively kill splenic/blood natural killer-resistant tumor cells by the perforin/granzyme pathway. J. Leukoc. Biol. 72:668–676.
Vermijlen, D., P. Ellis, C. Langford, A. Klein, R. Engel, K. Willimm, H. Joma, A.C. Hayday, and M. Eberl. 2007. Distinct cytokine-driven responses of activated blood gammadelta T cells: insights into unconventional T cell pleiotropy. J. Immunol. 178:4304–4314.
Wang, L., A. Kamath, H. Dax, L. Is, and J.F. Bukowski. 2001. Antibacterial effect of human V gamma 2V delta 2 T cells in vivo. J. Clin. Invest. 108:1349–1357.
Wang, T., Y. Gao, E. Scully, C.T. Davis, J.F. Anderson, T. Welte, M. Ledizet, R. Koski, J.A. Madni, A. Barrett, et al. 2006. Gamma delta T cells facilitate adaptive immunity against West Nile virus infection in mice. J. Immunol. 177:1825–1832.
Wesch, D., T. Hinz, and D. Kabelitz. 1998. Analysis of the TCR Vgamma repertoire in healthy donors and HIV-1-infected individuals. Int. Immunol. 10:1067–1075. doi:10.1093/immunol/10.8.1067
White, G.P., P.M. Watt, B.J. Holt, and P.G. Holt. 2002. Differential patterns of methylation of the IFN-gamma promoter at CpG and non-CpG sites underlie differences in IFN-gamma gene expression between human neonatal and adult CD45Ro- T cells. J. Immunol. 168:2820–2827.
Wilhelm, M., V. Kunzmann, S. Eckstein, P. Reimier, F. Weissinger, T. Ruediger, and H.P. Tony. 2003. Gammadelta T cells for immune therapy of patients with lymphoid malignancies. Blood. 102:200–206. doi:10.1182/blood-2002-12-3665
Williamson, D.J., D. Vermijlen, E.L. Wise, S.I. Clarke, R.E. Tigelaar, and A.C. Hayday. 2005. The integration of conventional and unconventional T cells that characterizes cell-mediated responses. Adv. Immunol. 87:27–59. doi:10.1016/S0091-6700(07)8002-6
Peyrat, M.A., F. Davodeau, I. Houde, F. Romagné, A. Necker, C. Leget, J.P. Cervoni, N. Cerf-Bensussan, H. Vié, M. Bonneville, and M.M. Hallet. 1995. Repertoire analyses of human peripheral blood lymphocytes using a human V delta 3 region-specific monoclonal antibody. Characterization of dual T cell receptor (TCR) delta-chain expressers and alpha beta T cells expressing V delta 3 alpha C alpha-encoded TCR chains. J. Immunol. 155:3060–3067.
Pitard, V., D. Roumanes, X. Lafarge, L. Couzi, I. Garrigue, M.E. Lafon, P. Merville, J.F. Moreau, and J. Déchanet-Merville. 2008. Long-term expansion of effector/memory Vdelta2-gammadelta T cells is a specific blood signature of CMV infection. Blood. 112:1317–1324. doi:10.1182/blood-2008-01-136713
Ramsburg, E., R. Tigelaar, J. Craft, and A. Hayday. 2003. Age-dependent requirement for γδ T cells in the primary but not secondary protective immune response against intestinal parasite. J. Exp. Med. 198:1403–1414. doi:10.1084/jem.200300680
Romero, V., D.E. Fellows, D.E. Jinne, and F. Andrade. 2009. Cleavage of La protein by granzyme H induces cytoplasmic translocation and interferes with La-mediated HCV-IRES translational activity. Cell Death Differ. 16:340–348. doi:10.1038/cdd.2008.165
Stagno, S. 2001. Cytomegalovirus. In Infectious Diseases of the Fetus and Newborn Infants. J.S. Remington and J.O. Klein, editors. W.B. Saunders Company, Philadelphia, PA. 389–424.