Multidrug Resistance Conferred by Novel DNA Polymerase Mutations in Human Cytomegalovirus Isolates

Gillian M. Scott, 1,3,4* Adriana Weinberg, 5 William D. Rawlinson, 1,2,3,4 and Sunwen Chou 6

Virology Research, POWH and UNSW Research Laboratories, 1 and Department of Microbiology, SEALS, 2 Prince of Wales Hospital, Randwick, NSW, Australia; School of Medical Sciences, Faculty of Medicine, 3 and School of Biotechnology and Biomolecular Sciences, Faculty of Science, 4 University of New South Wales, Sydney, Australia; Infectious Diseases, Departments of Pediatrics and Medicine, University of Colorado Health Sciences Center, Denver, Colorado; 7 and Medical and Research Services, Veterans Affairs Medical Center, and Division of Infectious Diseases, Oregon Health and Science University, Portland, Oregon 6

Received 24 May 2006/Returned for modification 13 July 2006/Accepted 5 October 2006

The emergence of antiviral-resistant cytomegalovirus (CMV) strains is a continuing clinical problem, with increased numbers of immunocompromised patients given longer-duration antiviral prophylaxis. Two previously unrecognized CMV DNA polymerase mutations (N408K and A834P) identified separately and together in at-risk lung and kidney transplant recipients and a third mutation (L737M) identified in a liver transplant recipient were characterized by marker transfer to antiviral-sensitive laboratory strains AD169 and Towne. Subsequent phenotypic analyses of recombinant strains demonstrated the ability of mutation N408K to confer ganciclovir (GCV) and cidofovir (CDV) resistance and of mutation A834P to confer GCV, foscarnet, and CDV resistance. Mutation L737M did not confer resistance to any of the antiviral agents tested. A recombinant strain containing both N408K and A834P demonstrated increased GCV and CDV resistance compared to the levels of resistance of the virus containing only the A834P mutation. The addition of mutation N408K in combination with A834P also partially reconstituted the replication impairment of recombinant virus containing only A834P. This suggests that perturbation of both DNA polymerization (A834P) and exonuclease (N408K) activities contributes to antiviral resistance and altered replication kinetics in these mutant strains. The identification of these multidrug-resistant CMV strains in at-risk seronegative recipients of organs from seropositive donors suggests that improved prophylactic and treatment strategies are required. The additive effect of multiple mutations on antiviral susceptibility suggests that increasing antiviral-resistant phenotypes can result from different virus-antiviral interactions.

Immunocompromised patients with life-threatening illnesses caused by human cytomegalovirus (CMV) can be treated with the antiviral agents ganciclovir (GCV), valganciclovir (val-GCV), foscarnet (FOS), or cidofovir (CDV) (36, 41). The use of oral valaciclovir, GCV, or valGCV has also made the delivery of prophylactic anti-CMV therapy easier and has improved the outcomes for highly immunosuppressed patients (24, 34, 37). CMV strains resistant to antiviral agents arise under the pressure of antiviral selectivity, with very low levels of resistant genotypes detected in patients prior to antiviral treatment (20, 33). Therefore, the lengthy periods of antiviral administration required for effective inhibition of this latent virus provide the selective pressure for the emergence of antiviral-resistant CMV strains, with resistant CMV strains detected in 5 to 30% of immunocompromised patients treated for more than 2 months, depending on the patient group studied (3, 28, 31). In some patients, particularly severely immunocompromised children, antiviral-resistant CMV strains can emerge rapidly (within 6 weeks) after the start of antiviral treatment (18, 42). Other risk factors for resistant CMV infections include transplantation from a seropositive donor to a seronegative recipient (D+/R−), the use of prolonged low-dose oral antiviral prophylaxis or intermittent intravenous treatment, and the administration of highly potent immunosuppression (19, 27, 31).

CMV antiviral resistance develops via mutations in the UL97 protein kinase (responsible for GCV phosphorylation) or the CMV UL54 DNA polymerase (for reviews, see references 5 and 23). Mutations of the CMV UL97 protein kinase that confer GCV resistance have been well characterized and are localized at codons 460 (protein kinase functional domain VIb) and 520; in addition, single base changes and deletions within codons 590 to 607 have been found to confer GCV resistance (1, 6, 10, 13, 25, 43). This simplifies identification of the CMV UL97 protein kinase mutations associated with antiviral resistance by targeted PCR sequencing or restriction fragment length polymorphism analysis (6, 35, 39). Conversely, CMV DNA polymerase mutations associated with antiviral resistance occur throughout the functional domain region spanning codons 301 (DNA polymerase exonuclease I region) to 989 (domain VI) (4, 7–9, 11, 14, 15, 42, 44, 46). Furthermore, previously unrecognized mutations of the CMV DNA polymerase continue to be identified in CMV strains isolated from patients receiving antivirals (17, 39, 46).

In this study, we characterized three novel mutations identified in CMV strains isolated from patients with clinically resistant CMV disease. Two of these, N408K (which occurs in DNA polymerase domain IV) and A834P (which occurs in

---

"Corresponding author. Mailing address: Virology Research, POWH and UNSW Research Laboratories, Level 3 Clinical Sciences Building, Prince of Wales Hospital, Avoca Street, Randwick, NSW 2031, Australia. Phone: 61-2 9382 9096. Fax: 61-2 9382 8533. E-mail: Gillian.Scott@sesi.health.nsw.gov.au.

"Published ahead of print on 16 October 2006.

"Within codons 590 to 607 have been found to confer GCV resistance (1, 6, 10, 13, 25, 43). This simplifies identification of the CMV UL97 protein kinase mutations associated with antiviral resistance by targeted PCR sequencing or restriction fragment length polymorphism analysis (6, 35, 39). Conversely, CMV DNA polymerase mutations associated with antiviral resistance occur throughout the functional domain region spanning codons 301 (DNA polymerase exonuclease I region) to 989 (domain VI) (4, 7–9, 11, 14, 15, 42, 44, 46). Furthermore, previously unrecognized mutations of the CMV DNA polymerase continue to be identified in CMV strains isolated from patients receiving antivirals (17, 39, 46).

In this study, we characterized three novel mutations identified in CMV strains isolated from patients with clinically resistant CMV disease. Two of these, N408K (which occurs in DNA polymerase domain IV) and A834P (which occurs in..."
DNA polymerase domain III), were identified in separate D+/R− lung transplant recipients from Australia and in a kidney transplant recipient from the United States. None of the patients responded to GCV, FOS, or CDV therapy (30, 39). The third mutation, L737M (DNA polymerase domain II), was identified in a liver transplant recipient receiving GCV (39). The phenotypic antiviral susceptibilities of recombinant CMV strains containing these mutations were assessed in vitro by two different assays, and the potential impacts of these mutations on DNA polymerase structure and function are discussed. These findings have implications for the management of D+/R− immunocompromised transplant recipients at high risk of developing multidrug-resistant CMV infections.

MATERIALS AND METHODS

Specimens, plasmid vectors, and viruses. Previously reported DNA polymerase mutations (N408K, L737M, and A834P) were identified in EDTA-anticoagulated blood specimens from patients with clinical evidence of infection with antiviral-resistant CMV (30, 39). These specimens were obtained with patient consent, and testing was carried out following the ethical guidelines of the South Eastern Sydney and Illawarra Area Health Service. CMV laboratory strains AD169 and Towne were obtained from the American Type Culture Collection (ATCC, strains ATCC VR538 and ATCC VR977, respectively). Previously described pBluescript plasmid vectors containing either the AD169- or Towne-derived CMV UL54 DNA polymerase genes with engineered Pmel restriction sites were used to transfer mutations to recombinant virus (7, 12). The construction of antiviral-sensitive AD169-derived and Towne-derived recombinant viruses used for marker transfer has been reported previously (7, 12). This includes the AD169-derived recombinant viruses T2211, which contains the secreted alkaline phosphatase (SEAP) reporter gene, a SwaI restriction site as well as with medium without an antiviral. The plaque numbers were counted, and the IC50 values were calculated for each assay after 7 days of incubation at 37°C with 5% CO2, the supernatant was removed and virus was incubated for 90-min absorption period, the supernatant was removed and replaced with 1 ml of medium containing 0.8% carboxymethyl cellulose as well as with medium without an antiviral. The plaque numbers were counted, and the IC50 values were calculated for each assay after 7 days of incubation. Each assay was carried out multiple times (7 to 23 times), and the averages and standard deviations were calculated. The GCV, FOS, and CDV susceptibilities of strain AD169-derived antiviral-sensitive strains T2211, T2233, and T2241 and recombinants T2293 (containing N408K), T2291 (containing A834P), and T2311 (containing N408K and A834P) and Towne-derived recombinant viruses T2296 (containing L737M) and T2287 (containing A834P) were also determined by a plaque reduction assay (PRA) with human embryonic fibroblasts (29). Each of these assays was carried out in triplicate, and the averages and standard deviations were calculated. Recombinant viruses with a greater than twofold increase in IC50 compared to those for the wild-type sensitive control strains were considered antiviral resistant, as this has been shown to be reproducible in previous studies of recombinant viruses (7, 13).

Antiviral susceptibility analysis of the AD169-derived mutants. The GCV, FOS, and CDV susceptibilities of strain AD169-derived antiviral-sensitive strains T2211, T2233, and T2241 and recombinants T2293 (containing N408K), T2291 (containing A834P), and T2311 (containing N408K and A834P) were determined by the SEAP-based assay (12). Cell-free virus stock was inoculated at a multiplicity of infection of 0.01 to 0.03 into HFF cultures in 24-well plates and grown in the absence or presence of twofold increasing concentrations of antiviral. After 7 days of culture, aliquots of the supernatant from each well were analyzed for SEAP activity as described previously (12), and 50% inhibitory concentrations (IC50s) were calculated by linear regression analysis (12). Each assay was carried out multiple times (7 to 23 times), and the averages and standard deviations were calculated. The GCV, FOS, and CDV susceptibilities of AD169 wild-type virus (ATCC), T2293 (containing N408K), T2291 (containing A834P), and T2311 (containing N408K and A834P) were also determined by a plaque reduction assay (PRA) with human embryonic fibroblasts (29). Each of these assays was carried out in triplicate, and the averages and standard deviations were calculated. Recombinant viruses with a greater than twofold increase in IC50 compared to those for the wild-type sensitive control strains were considered antiviral resistant, as this has been shown to be reproducible in previous studies of recombinant viruses (7, 13).

Antiviral susceptibility analysis of the Towne-derived mutants. The GCV, FOS, and CDV susceptibilities of wild-type Towne (ATCC) and Towne-derived strains T2296 (containing L737M) and T2287 (containing A834P) were determined by PRA (40). A cell-free virus stock (50 PFU) was inoculated onto 80 to 100% confluent human lung (MRC-5) fibroblasts in 48-well plates (two assays per plate), and the plates were centrifuged at 600 × g for 30 min. After 1 h of incubation at 37°C with 5% CO2, the supernatant was removed and virus was grown in duplicate in the presence of 0.625 to 160 μM GCV, 0.625 to 1,600 μM FOS, or 0.0625 to 16 μM CDV in medium containing 0.8% carboxymethyl cellulose as well as with medium without an antiviral. The plaque numbers were counted, and the IC50 values were calculated for each assay after 7 days of culture. Each assay was carried out multiple times (8 to 11 times), and the averages and standard deviations were calculated. Recombinant viruses with a greater than twofold increase in IC50 compared to those for the wild-type sensitive control strains were considered antiviral resistant (7).

In vitro replication assays. The growth properties of mutant recombinant viruses over multiple cycles of replication were assessed by inoculating isolates strains T2211 and T2241 (wild-type controls), T2291 (containing A834P), and T2311 (containing N408K and A834P) into 24-well HFF cultures at a multiplicity of infection of 0.01. These experiments were carried out on three separate dates with independent viral dilutions. Four wells were inoculated per strain and after a 90-min absorption period, the supernatant was removed and replaced with 1 ml of medium containing 0.8% carboxymethyl cellulose as well as with medium without an antiviral. The plaque numbers were counted, and the IC50 values were calculated for each assay after 7 days of incubation. Each assay was carried out multiple times (8 to 11 times), and the averages and standard deviations were calculated. Recombinant viruses with a greater than twofold increase in IC50 compared to those for the wild-type sensitive control strains were considered antiviral resistant (7).

TABLE 1. Antiviral susceptibilities of AD169-derived mutants assessed by SEAP-based assay

<table>
<thead>
<tr>
<th>Virus genotype</th>
<th>DNA pol genotype</th>
<th>IC50 (μM)</th>
<th>Fold increase</th>
<th>IC50 (μM)</th>
<th>Fold increase</th>
<th>IC50 (μM)</th>
<th>Fold increase</th>
</tr>
</thead>
<tbody>
<tr>
<td>T2241</td>
<td>Wild type (AD169)</td>
<td>1.4 ± 0.5</td>
<td>42 ± 14</td>
<td>0.3 ± 0.1</td>
<td>0.2 ± 0.11</td>
<td>0.5 ± 0.1</td>
<td>0.25</td>
</tr>
<tr>
<td>T2211</td>
<td>Wild type + Pmel</td>
<td>1.3 ± 0.4</td>
<td>41 ± 12</td>
<td>1.2 ± 0.2</td>
<td>0.2 ± 0.11</td>
<td>0.5 ± 0.1</td>
<td>0.25</td>
</tr>
<tr>
<td>T2323</td>
<td>Wild type + Pmel</td>
<td>1.4 ± 0.4</td>
<td>45 ± 8</td>
<td>0.3 ± 0.1</td>
<td>0.2 ± 0.11</td>
<td>0.5 ± 0.1</td>
<td>0.25</td>
</tr>
<tr>
<td>Avg for wild types</td>
<td></td>
<td>1.33</td>
<td>42.7</td>
<td>0.25</td>
<td></td>
<td>18.8</td>
<td></td>
</tr>
<tr>
<td>T2293</td>
<td>N408K</td>
<td>5.6 ± 1.6</td>
<td>42 ± 14</td>
<td>5.2 ± 1.5</td>
<td>21.0</td>
<td>4.7 ± 2.3</td>
<td>18.8</td>
</tr>
<tr>
<td>T2291</td>
<td>A834P</td>
<td>7.1 ± 3.6</td>
<td>5.4</td>
<td>0.7 ± 0.4</td>
<td>3.0</td>
<td>0.7 ± 0.4</td>
<td>3.0</td>
</tr>
<tr>
<td>T2311</td>
<td>N408K + A834P</td>
<td>30.2 ± 8.8</td>
<td>22.7</td>
<td>308 ± 78</td>
<td>7.2</td>
<td>4.7 ± 2.3</td>
<td>18.8</td>
</tr>
</tbody>
</table>

a Mean values and standard deviations were calculated from 7 to 23 replicate experiments.

b Fold increase in IC50 compared to the average IC50 for wild-type sensitive strains.
minimum essential medium–3% fetal bovine serum. On days 1, 4, 5, 6, 7, and 8, 80-μl aliquots of the medium supernatant were removed and frozen for subsequent SEAP assay. At the end of the experiment, SEAP assays were performed with chemiluminescence, as described previously (12). The SEAP assay values observed over the time period were graphed, with points and error bars representing the means and standard deviations, respectively, of the four wells sampled per strain.

RESULTS

Antiviral susceptibilities of AD169-derived recombinant viruses. SEAP-based analysis and PRA consistently demonstrated the ability of mutation N408K to confer GCV and CDV resistance but not FOS resistance to AD169-derived recombinant virus T2293 (containing N408K) (Tables 1 and 2). SEAP-based analysis indicated that N408K conferred high-level resistance to CDV, with a 21.0-fold increase in the CDV IC_{50} of recombinant T2293 compared to the IC_{50} for the wild-type sensitive strains. The fold increase in CDV resistance (5.4-fold compared to those for the wild-type strains) was less pronounced when the increase was determined by PRA, which is probably related to the higher IC_{50} for the wild-type strains obtained by PRA compared to those obtained by SEAP-based analysis. The A834P mutation conferred GCV, FOS, and CDV resistance to recombinant virus T2291, as determined by SEAP-based analysis and PRA. Like N408K, the ability of mutation A834P to confer GCV resistance to T2291 was less pronounced when resistance was measured by PRA (2.2-fold increase in the GCV IC_{50} compared to that at the baseline) but was more clearly evident by SEAP-based analysis (5.4-fold increase in the GCV IC_{50} compared to that at the baseline).

The addition of N408K with A834P in recombinant T2311 produced 2.7- to 6.2-fold increases in GCV and CDV resistance compared to the levels of resistance of T2291 containing the single A834P mutation by both the SEAP assay and PRA (Tables 1 and 2). As expected, addition of N408K with A834P produced a negligible (1.1- to 1.7-fold) increase in FOS resistance, given the lack of FOS resistance produced by the N408K mutation in T2293 with a single mutation. The combined effect of the N408K and A834P mutations on antiviral resistance was less consistent when the resistance of the mutant with the single N408K mutation was compared to that of the mutant with dual mutations. These data suggest that N408K enhanced the GCV and CDV resistance of mutants containing A834P, but the effect was not consistently reciprocal.

Antiviral susceptibility of the Towne-derived recombinant viruses. Mutation L737M did not confer GCV or FOS resistance to Towne-derived recombinant strain T2296 (Table 3). A twofold increase in CDV susceptibility compared to that of the sensitive wild-type Towne strain was demonstrated for T2296, but this virus was not considered CDV resistant, given that resistant viruses typically demonstrate a greater than twofold increase in antiviral sensitivity (7, 12). Towne-derived recombinant T2287, which contained the A834P mutation, showed resistance to all three drugs (GCV, FOS, and CDV) by PRA, similar to the findings presented above for the corresponding AD169-derived recombinant T2291.

Replication fitness of recombinant viruses. The amount of virus T2291 (containing A834P) measured by SEAP-based chemiluminescence after 4, 5, and 6 days in HFF culture was 4.2- to 6.5-fold less than that observed for the AD169-like wild-type strain T2211 (Fig. 1). A similar decrease was observed when T2291 was compared to the second wild-type control strain, T2241 (results not shown). To ensure that the smaller amount of growth was not due to relative underinoculation, strain T2291 was adjusted on two of the setup dates so that its SEAP levels at day 1 were slightly greater than those for the wild-type strains. These initial reductions in virus growth became less pronounced by days 7 and 8, at which point the amount of wild-type virus had reached a maximum plateau, with 2.9- and 1.3-fold differences between the amounts for wild-type virus T2211 and strain T2291, respectively. Early attenuation was also observed for dual recombinant virus T2311, with a 2.1- to 2.9-fold reduction in viral output after 4 to 6 days of culture compared to that of the wild-type strain. By day 8 the viral output of strain T2311, which contained dual...

### TABLE 2. Antiviral susceptibilities of AD169-derived mutants assessed by PRA*

<table>
<thead>
<tr>
<th>Virus</th>
<th>DNA pol genotype</th>
<th>GCV</th>
<th>FOS</th>
<th>CDV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>IC_{50} (μM)</td>
<td>Fold increase&lt;sup&gt;a&lt;/sup&gt;</td>
<td>IC_{50} (μM)</td>
</tr>
<tr>
<td>T2293</td>
<td>N408K</td>
<td>1.9 ± 0.5</td>
<td>5.3</td>
<td>78 ± 13</td>
</tr>
<tr>
<td>T2291</td>
<td>A834P</td>
<td>4.1 ± 2.0</td>
<td>2.2</td>
<td>431 ± 184</td>
</tr>
<tr>
<td>T2311</td>
<td>N408K + A834P</td>
<td>12.4 ± 1.8</td>
<td>6.5</td>
<td>732 ± 418</td>
</tr>
</tbody>
</table>

<sup>a</sup> Mean values and standard deviations were calculated from three replicate experiments.

### TABLE 3. Antiviral susceptibilities of Towne-derived mutants assessed by PRA

<table>
<thead>
<tr>
<th>Virus</th>
<th>DNA pol genotype</th>
<th>GCV</th>
<th>FOS</th>
<th>CDV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>IC_{50} (μM)</td>
<td>Fold increase&lt;sup&gt;a&lt;/sup&gt;</td>
<td>IC_{50} (μM)</td>
</tr>
<tr>
<td>Towne</td>
<td>Wild type</td>
<td>4.9 ± 0.5</td>
<td>1.3</td>
<td>62.8 ± 13.6</td>
</tr>
<tr>
<td>T2296</td>
<td>L737M</td>
<td>5.4 ± 1.9</td>
<td>1.1</td>
<td>63.4 ± 12.9</td>
</tr>
<tr>
<td>T2287</td>
<td>A834P</td>
<td>30.1 ± 10.2</td>
<td>6.1</td>
<td>486.3 ± 7.9</td>
</tr>
</tbody>
</table>

<sup>a</sup> Mean values and standard deviations were calculated from 8 to 11 replicate experiments.
mutations, was equivalent to the maximum plateau for wild-type strain T2211. Similar results were obtained for each of the three experiments carried out on different dates. The SEAP activity at day 5 for the wild-type controls (T2211 or T2241) was consistently greater than that for T2311, which was in turn greater than that for T2291, with a >2-standard-deviation difference in the SEAP activity of T2291 and those of the other strains. T2311 consistently demonstrated higher SEAP activity levels than T2291 whether the SEAP levels for T2311 on day 1 were slightly higher or lower than those for T2291 (results not shown). The addition of mutation N408K in dual recombinant virus T2311 therefore improved the replication fitness of virus T2291, which contained only A834P.

**DISCUSSION**

The identification of multidrug resistance-conferring DNA polymerase mutations N408K (GCV and CDV resistance) and A834P (GCV, FOS, and CDV resistance) separately and together in three CMV strains infecting immunosuppressed CMV-seronegative transplant recipients emphasizes the increased risk of antiviral-resistant CMV acquisition in these patients (21, 27, 30, 31, 39). Consistent with the known risk factors for the emergence of these antiviral-resistant CMV strains, these transplant recipients had received long-term GCV therapy, potentially at subclinical doses (3, 16, 20, 27). The subsequent persistence of these strains may be attributed to the ongoing but intermittent antiviral (GCV, FOS, or CDV) treatment that these patients then received (30, 39). Consistent extended high-dose antiviral prophylaxis or observational preemptive therapy based on improved diagnostic assays may reduce the frequency of antiviral-resistant CMV strains in high-risk D+/R− transplant recipients (21). Unfortunately, all current CMV drugs may result in dose-limiting toxicity, and less toxic but effective anti-CMV therapy needs to be developed.

Consistent with our findings for mutation N408K (Tables 1 and 2), mutations of DNA polymerase domain IV-exonuclease II (ExoII) domain typically confer various levels of resistance to GCV and CDV but not resistance to FOS (14). The levels of GCV and CDV resistance among strains with the N408K mutation were similar to the published levels of GCV and CDV resistance among strains with the N408D mutation (14). This asparagine residue is highly conserved among α-like DNA polymerases and is located within the ExoII domain, which binds to single-stranded DNA and the metal ions required for excision of mismatched bases (2). It is interesting that substitution of a neutral polar residue (asparagine) to a basic residue (lysine) or acidic residue (aspartic acid) at this important codon has similarly contributed to GCV and CDV resistance.

The domain III amino terminus of herpesviruses consists of a DNA template binding region (47), immediately followed by residues important for pyrophosphate binding and nucleotide incorporation (9, 15, 44). Mutation A834P is situated within a putative α-helical region (α helix Q) at the less well defined carboxy end of DNA polymerase domain III (32, 45), where two other CMV mutations (T838A and G841A) and a herpes simplex virus (HSV) mutation (R842S) associated with antiviral resistance have been identified (22, 26, 42). The HSV R842S mutation has little effect on the interaction of the HSV DNA polymerase with dGTP (26), even though the neurovirulence of HSV is attenuated by this mutation in mice (38). This evidence and the reduced level of CMV replication in vitro produced by the A834P mutation shown here suggest that mutations toward the domain III carboxy terminus of DNA polymerase alter DNA template and substrate interactions through conformational changes in the catalytic core (32, 45).

The close interactions of domain III and domain II residues in HSV- and RB69-derived DNA polymerase models support this hypothesis (26, 32, 45).

Qualitative antiviral resistance was consistently demonstrated for strains with the N408K and A834P mutations by two validated assays: the SEAP-based assay (which measures cell-free virus output to the medium) and PRA (which measures cell-associated virus replication) (12, 29). The different viral parameters measured by the two assays are likely to account for the quantitative differences in antiviral inhibitory concentrations indicated by the two assays, as suggested by the lower IC50 values demonstrated for even wild-type strain AD169 as measured by the SEAP-based assay compared with that measured by PRA (Tables 1 and 2). The greater dynamic range and increased number of replicates possible with the SEAP-based assay allow greater confidence in the mean values obtained. However, variations in results from laboratory to laboratory for standardized assays are well documented (29), requiring “in-house” comparisons to the relevant wild-type virus when qualitative antiviral resistance is determined by either method.

Further characterization of emerging DNA polymerase mutations that confer resistance to antiviral agents (such as N408K and A834P) and even those that result in antiviral-sensitive phenotypes (L737M) is necessary for the accurate and definitive detection of antiviral-resistant strains by rapid genotypic assays (35, 39). This and other studies are elucidating the total numbers of DNA polymerase mutations that confer antiviral resistance, but more investigations are required before this list is complete. As demonstrated here, investigation of novel antiviral resistance mutations increases our understanding of the functional regions of CMV DNA polymerase and...
will assist with the development of novel antiviral agents that inhibit CMV replication.

ACKNOWLEDGMENTS

This work was supported by project grant 300532 from the National Health and Medical Research Council of Australia and U.S. National Institutes of Health grant AI39938. We thank Gail Marousek, Laura Van Wechel, Heather Lichy, and Monique Nicolle for technical assistance.

REFERENCES


