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Effects of protease inhibitors on the course of CMV retinitis in relation to CD4+ lymphocyte responses in HIV+ patients

Gerardus J van den Horn, Christina Meenken, Sven A Danner, Peter Reiss, Marc D de Smet

Abstract

Aim—To gain insight into the course of CMV retinitis (CMVR) in AIDS patients receiving protease inhibitors (PI), and to evaluate whether certain patterns of CD4 response are indicative of the clinical outcome and the risk of recurrence. Methods—15 consecutive AIDS patients receiving maintenance therapy for CMVR were included in a prospective observational cohort study at the university hospital between July and October 1996. Patients were evaluated for signs of CMVR activity and intraocular inflammation. CMVR recurrence was defined as the primary clinical endpoint. Follow up was performed until July 1997. No patient was lost to follow up. Clinical outcome was related to CD4+ lymphocyte counts, which were monitored every 6 weeks. Highly active antiretroviral treatment regimen including PI was started at study entry. Results—All recurrences (n=7) were in patients who failed to have a sustained increase in CD4 counts, whereas CMVR remained inactive during a follow up of 42–52 weeks in those who were able permanently to restore their CD4 values to 100×10^6/l or more (n=5). The remaining three patients died after 12, 16, and 50 weeks, respectively, without recurrences. All relapses of CMVR were seen after 6–16 weeks, and at CD4 counts well below 100×10^6/l. Conclusions—The beneficial effects of PI treatment correlate with the pattern of CD4 response. Sustained increases in CD4 counts achieved in the first 16 weeks of treatment are associated with a prolonged period of CMVR quiescence. Poor initial response is associated with a high risk of CMVR recurrence.

Cytomegalovirus retinitis (CMVR) is the most frequent opportunistic eye infection in patients with HIV infection. It is commonly seen in patients with very low CD4+ lymphocyte counts (below 50×10^6/l). Areas of necrotising retinitis, often with extensive intraretinal haemorrhages, are observed, while inflammatory reaction in the anterior segment and the vitreous cavity is usually mild or virtually absent. This lack of reaction is attributed to the inability to mount sufficient immunological responses. Therapy consists of lifelong anti-CMV regimes. In spite of therapy, recurrences are common.

Protease inhibitors (PI) may change this classic picture. Impressive results have been reported with triple drug combinations leading to declining plasma HIV-RNA levels below the detection limit of currently available assays in over 90% of individuals.1,2 Patients with opportunistic infections have sometimes shown prolonged remissions or improvement in the clinical manifestations of their diseases following initiation of PI treatment.3–5 Simultaneously, substantially increased CD4 counts have been observed. These observations reflect at least a partial restoration of immune function in patients receiving PI. However, improvement in immune function may also have disadvantageous effects: enhanced inflammatory activity has been observed shortly after installation of PI therapy.6

PI have also had a significant impact on CMVR. The overall incidence of CMVR has dramatically decreased. Recurrence free intervals have been prolonged in some individuals to the point where the role of maintenance therapy has been brought into question. However, CMVR has also been observed in patients on PI therapy at CD4 levels higher than previously reported.7 With the advent of PI, it has become necessary to re-evaluate our screening and therapeutic approaches with respect to CMVR.

To gain insight into the course of CMVR in patients receiving PI, we carried out a prospective observational study in AIDS patients on maintenance therapy for CMVR in whom PI treatment was initiated. By relating the course of CMVR to the CD4 response over time we attempt to evaluate whether certain patterns of CD4 response are indicative of the clinical outcome and the risk of recurrence.
Effects of protease inhibitors on the course of CMV retinitis in relation to CD4+ lymphocyte responses in HIV+ patients

Table 1  Patient characteristics before and during protease inhibitor (PI) treatment.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Duration of CMVR (weeks)</th>
<th>No of induction courses</th>
<th>Time to progression (weeks)</th>
<th>Recurrence (weeks)</th>
<th>Follow up (weeks)</th>
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<td>5</td>
<td>1</td>
<td>10</td>
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<td>Died at 12 weeks</td>
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<tr>
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<td>9</td>
<td>1</td>
<td>—</td>
<td>No</td>
<td>—</td>
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<tr>
<td>3</td>
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<td>4</td>
<td>16</td>
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<td>—</td>
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<tr>
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<td>40</td>
<td>1</td>
<td>—</td>
<td>No</td>
<td>Died at 50 weeks</td>
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<td>44</td>
<td>3</td>
<td>10</td>
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<td>—</td>
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<td>1</td>
<td>—</td>
<td>No</td>
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<td>No</td>
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<td>2</td>
<td>1</td>
<td>—</td>
<td>No</td>
<td>52</td>
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</tbody>
</table>

Patients and methods

Between July and October 1996, highly active retroviral treatment including PI was started in 15 consecutive AIDS patients receiving maintenance therapy for CMVR with either ganciclovir or foscarnet. CMVR was diagnosed on clinical grounds (that is, a characteristic funduscopy appearance and a favourable response to anti-CMV therapy) 2–54 weeks before onset of PI treatment. At the time of inclusion, several of the patients had required one or more reinduction courses because of CMVR progression (Table 1). CD4+ lymphocyte counts were determined at onset of PI therapy, and at 6–8 week intervals. Ophthalmological examination was performed every 2 weeks during the first 2 months, and every 4 weeks thereafter, and was focused on two particular aspects—inflammatory signs as an indicator of the patient’s ability to mount an inflammatory response and CMVR activity. The inflammatory reaction in the anterior and posterior segments was scored by slit lamp examination as either regular or excessive; CMVR activity was graded funduscopy as completely inactive, smouldering, or active. In case of active or smouldering lesions, fundus photographs were helpful to assess retinitis control. All assessments were carried out by two ophthalmologists experienced in AIDS related pathology. The clinical end point was defined by recurrence of CMVR. Recurrences were diagnosed if the borders of any lesion had advanced by more than half an optic disc diameter (750 µm), or if any new lesion appeared in a previously unaffected area, compared with baseline conditions. Time of initiation of PI was regarded as baseline.

Results

CD4 counts

All patients invariably had very low CD4 counts at study onset (mean CD4 25×10⁶/l, range 10–60). Figure 1 shows the individual CD4 responses to PI treatment of all CMVR patients. Three did not show any increase in their CD4 count after 8 weeks of treatment (non-responders). Seven patients had a rapid increase to values of 50×10⁶/l or more, followed by a return to nearly initial values over the following 6 weeks. A group of five patients showed a sustained increase in their CD4 counts, and reached values of 100×10⁶/l or more after 12–16 weeks of treatment. The elevated levels were maintained during the whole follow up period.

Vitritis

Six patients showed excessive inflammation in the vitreous cavity 6–8 weeks after onset of PI medication. They are indicated in Figure 1 by an asterisk. Four of them (nos 5, 6, 9, 11) had a concomitant rapid increase of their CD4 count to values above 100×10⁶/l. Patients experienced blurred vision and floaters, but none required a vitrectomy. Vitritis never interfered with funduscopy assessment.

CMVR

While using PI, seven patients needed reinduction because of CMVR reactivation. In two patients, CMVR remained smouldering, whereas it became completely inactive in six. CMVR activity in relation to the response to PI treatment is indicated in Figure 1 (bottom). All recurrences were in patients who failed to have sustained CD4 responses to PI treatment, whereas CMVR became inactive in those who were able to permanently restore their CD4 values to 100×10⁶/l or more. All relapses of CMVR were seen at CD4 counts well below 100×10⁶/l.

Discussion

This study demonstrates two important phenomena. Firstly, there exists a clear difference in clinical outcome between patients with and without a sustained CD4 response to PI treatment. During a follow up of 42–52 weeks none of the five patients with a sustained response relapsed, whereas seven of 10 patients without a sustained response needed reinduction therapy because of reactivation of CMVR. All recurrences occurred within 4 months. This
suggests that patients without sustained CD4 response do not benefit from PI with respect to CMVR.

The second phenomenon is that the manifestations of successful PI treatment in severely immunocompromised AIDS patients can be divided into two episodes: an early phase of approximately 6 weeks’ duration in which CD4 cells may rise substantially and in which the ability to mount an inflammatory response is enhanced. In this phase, inflammatory reactions such as vitritis may occur. This episode is followed by either a return of CD4 counts to baseline values or by further immune restoration during which CD4 counts remain elevated (sustained response). A return of CD4 counts to baseline conditions preceded a relapse of CMVR in five cases and seems to be indicative of imminent recurrence.

Shortly after initiation of PI treatment we observed vitritis in six patients. This phenomenon has not been previously associated with AIDS related CMVR. It probably reflects an increased inflammatory response to ocular CMV proteins resulting from increased immunocompetence due to PI treatment. It is associated in our data with an early increase of CD4 to values above 100×10⁶/l, although the presence of inflammation is not predictive of a sustained increase in CD4 count. In a recent paper Jacobson et al. report five cases of CMVR with CD4 counts around 200×10⁶/l at the time of CMVR diagnosis. These patients had started PI 4–7 weeks earlier while their CD4 count was less than 85×10⁶/l. Two of these patients also had vitritis. In our opinion, the apparent discrepancy between the diagnosis of CMVR and the relatively high CD4 counts in these patients is a consequence of the fact that in these patients a previously subclinical CMVR progressed to symptomatic disease, while CD4 counts rose simultaneously as a result of initiation of PI treatment. We believe that their observation that no recurrences of CMVR were seen during a follow up of 5–12 months is of greater importance. It is consistent with the lack of progression of CMVR as seen in our patients with sustained CD4 responses. It suggests that immune restoration can be achieved to an extent sufficient to prevent recurrence of CMVR. An important question, which should be addressed in future studies, is whether continued anti-CMV maintenance therapy is still mandatory in patients with a sustained increase in CD4. Another issue worth investigating is whether declining CD4 levels in patients who have previously experienced a sustained increase in CD4 should be regarded as indicative of imminent recurrence. Such studies should not only involve frequent ocular examinations but also include CD4 monitoring in association with other prognostic markers such as determination of CMV-DNA levels in the peripheral blood by polymerase chain reaction.