Novel strategies in the use of lopinavir/ritonavir for the treatment of HIV infection in children

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Abstract: Lopinavir/ritonavir (LPV/r) is considered by many as the first choice protease inhibitor (PI) for children. This co-formulation avoids the need for children to take ritonavir separately to “boost” the levels of lopinavir. LPV/r has high virologic potency, an excellent toxicity profile and a high barrier to the development of viral resistance. However, LPV/r has poor tolerability of the oral suspension (due to the poor taste of ritonavir), difficult dosing requirements and metabolic side effects, especially hyperlipidemia. The new tablet low-dose formulation (100/25 mg) may allow more convenient antiretroviral treatment in children. Novel strategies of LPV/r in childhood could maximize its advantages. For example, infants infected with HIV despite single dose Nevirapine after birth need effective combination antiretroviral treatment. This can be given using a higher dose of LPV/r with therapeutic drug monitoring. Other novel uses include once daily LPV/r regimens in older children and adolescents and lower doses of LPV/r in certain populations, which may decrease hyperlipidemia. Heavily pre-treated children might benefit from a double PI/r regimen which includes LPV/r. The high potency of LPV/r needs to be balanced with convenient regimens, to enhance adherence and decrease toxicity whenever possible. The aim of this review is to discuss the rationale behind these novel strategies of LPV/r use in pediatric antiretroviral treatment as well as their results and limitations.

Keywords: human immunodeficiency virus, children, antiretroviral therapy, lopinavir, ritonavir

Introduction

Since the emergence of the HIV/AIDS pandemic children worldwide have suffered its devastating consequences. In 2007, UNAIDS estimated that 2.1 million children under 15 years old were living with HIV/AIDS, 420,000 children acquired the infection and 290,000 died of AIDS during this year.1,2

Without treatment, many HIV-infected children will die during the first year of life and half will not survive to their tenth birthday. Combination treatment (with 3 or more antiretroviral drugs) has dramatically changed the outlook for children with HIV infection, producing a marked decrease in mortality.3 This outstanding outcome has been particularly noticeable since the introduction of the protease inhibitors (PI) as part of Highly Active Antiretroviral Therapy (HAART) in 1997.4–6

The PIs are a class of antiretroviral drug that bind competitively to the viral protease enzyme that inhibits the cleavage of the gag-pol polyprotein. This results in production of immature non-infectious viral particles, and prevents subsequent cellular infection. Currently, 9 PIs are approved for the treatment of HIV-infection, 7 of which are approved...
Lopinavir/ritonavir in pediatric antiretroviral treatment

LPV/r was the sixth PI approved by the Food and Drug Administration (FDA). It is the only PI co-formulated as a fixed combination of lopinavir (LPV, initially known as ABT-378) and ritonavir (RTV). Ritonavir inhibits the hepatic metabolism of lopinavir increasing its plasma concentration. Ritonavir can be given separately to “boost” other PIs. However ritonavir suspension has an extremely poor taste and the capsule is large, making adherence in children extremely difficult. The co-formulation of RTV with LPV avoids some of these issues, although the suspension still has a poor taste.

LPV/r soft-gelatine capsule and oral solution were the first formulations approved for use in children. In March 2006, the capsule was replaced with a tablet formulation that used proprietary melt-extrusion technology. This offers several advantages over the capsule formulation, such as, lower pill burden, fewer gastrointestinal adverse effects and easier storage requirement (no need for refrigeration). In November 2007, the FDA approved a low strength tablet formulation (100/25 mg) for the use in children (available in the EU since 2008). The major disadvantage of the new tablet formulation is that the tablets can not be cut or crushed, as bioavailability is lost. This can make accurate dosing in children difficult. For example a child requiring 250/62.5 mg of LPV/r twice daily may either be given 200/50 mg in tablet form plus 50/12.5 mg as suspension twice daily, or 200/50 mg in tablet form in the morning and 300/75 mg in tablet form in the evening. Neither of these regimens is ideal and the pediatrician and caregivers will have to discuss which is most likely to achieve good adherence.

Currently several dosage forms of LPV/r are available for the treatment of pediatric HIV-infection (see Tables 1 and 2). A brief summary of the main pharmacokinetic and pharmacodynamic properties of LPV/r is shown in Table 3.

New strategies on the use of LPV/r in pediatric antiretroviral treatment

The limitations observed with the long-term use of LPV/r in children have impelled the scientific community to explore...
new ways to benefit from the high antiviral potency of LPV/r minimizing its disadvantages.

Use of LPV/r in infants

The risk of disease progression in HIV is inversely correlated with the age of the child, with the youngest children at greatest risk of rapid progression.\(^{16,17}\) Currently, both European and American guidelines recommend that antiretroviral treatment should be started in every HIV infected infant under 12 months of age regardless of immune status or viral load.\(^{18}\) This recommendation follows a 4-fold reduction in HIV progression/mortality among infants starting HAART at less than 3 months of age compared to later, in both a large cohort meta-analysis and a randomized controlled trial.\(^{19,20}\)

This is particularly relevant in resource-limited settings where most children with HIV/AIDS live. The mortality rate of HIV-infected children in developing countries is 45% to 59% by 2 years of age compared with 10% to 20% in the EU and the US but early ART among infants between 6 to 12 weeks with CD4% ≥ 25 has been associated with a reduction in mortality of 76%.\(^{19,21,22}\)

Babies may be infected with HIV despite attempts to prevent mother-to-child transmission. In resource-limited settings these infants will often have been given the NNRTI nevirapine (NVP). Those infants who become infected despite nevirapine use have a high risk of NNRTI resistance, raising concerns about the efficacy of NNRTI-based regimens within the first year of life in NVP-exposed infants. The efficacy of highly active agents other than NVP to treat very young infants, such as PIs, needed to be assessed.\(^{23}\)

LPV/r could thus be considered as one of the first-line agent choices for early antiretroviral treatment because of its liquid formulation.\(^{24}\)

Nelfinavir was the first PI used extensively in children. High doses were needed to achieve effective drug levels in infants and there was great intersubject variation. Poor long-term viral suppression was reported with nelfinavir given in the first 3 months of life (11 out of 16 infants experienced virological failure and 30% developed resistance).\(^{25}\)

LPV/r was initially approved in Europe for children older than 2 years and it has become the first choice PI in children.\(^{26}\) However young infants have a higher apparent clearance

### Table 2

<table>
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<tr>
<th>Pediatric dosing scheme for LPV/r(^{19,48})</th>
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<tr>
<td>Children aged 14 days to 6 months</td>
<td>16/4 mg/kg or 300/75 mg/m(^2) twice daily</td>
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<tr>
<td>Children aged 6 months to 12 years</td>
<td></td>
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<tr>
<td>• Weight: 7 to 15 kg</td>
<td>12/3 mg/kg or 230/57.5 mg/m(^2) twice daily</td>
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<tr>
<td>• Weight: 15 to 40 kg</td>
<td>10/2.5 mg/kg or 230/57.5 mg/m(^2) twice daily (max dose of 400/100 mg twice daily)</td>
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<tr>
<td>Children aged 6 months to 18 years with co-administration of EFV, NVP, NFV or (fos)amprenavir in either naïve or treatment-experienced patients</td>
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<tr>
<td>• Weight: &lt; 15 kg</td>
<td>13/3.25 mg/kg twice daily</td>
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<tr>
<td>• Weight: &gt; 15 kg</td>
<td>11/2.75 mg/kg twice daily without exceeding the adult dose</td>
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**Abbreviations:** LPV/r, lopinavir/ritonavir; EFV, efavirenz; NVP, nevirapine; NFV, nelfinavir.

### Table 3

<table>
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<th>Pharmacology and pharmacokinetics of LPV/r(^{19,48,49})</th>
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<td><strong>Absorption</strong></td>
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<td>Absorption of LPV/r liquid formulation is affected by the presence of food (the AUC and the C(<em>{max}) of LPV increased by 130% and 56% respectively when given with a high-fat meal compared with a fasting state). LPV/r tablets may be taken with or without food as long as the tablets are swallowed whole, without being chewed, crushed or broken. C(</em>{max}) of LPV of 9.8 ± 3.7 µg/mL 4 hours after the intake of the drug have been reported in adults after multiple dosing with 400/100 mg twice daily during 3–4 weeks. Minimum concentration within a dosing interval (12 h) was 5.5 ± 2.7 µg/mL Minimum effective concentration in treatment naïve adults has been established at &gt;1 µg/mL. AUC during a 12 hour dosing interval was 92.6 ± 36.7 µg·h/mL. The absolute bioavailability of LPV/r has not been established in humans.</td>
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| **Distribution** |  |
| LPV is approximately 98%–99% bound to plasma proteins (alpha-1-acid glycoprotein and albumin transport LPV). LPV/r accumulates intracellularly. Intracellular/plasma concentration of 1.18 has been reported. LPV is lipid soluble therefore penetrates the cerebrospinal fluid where a significant reduction of HIV viral load has been shown. |

| **Metabolism** |  |
| LPV is extensively metabolized by the hepatic cytochrome P450 system, almost exclusively by the CYP34 and CYP34A3 isoenzymes. RTV is a potent CYP34 inhibitor and consequently increases plasma levels of LPV when the two drugs are co-administered. RTV has been shown to induce metabolic enzymes, resulting in the induction of its own metabolism. |

| **Elimination** |  |
| After administration of LPV/r, approximately 10.4% ± 2.3% and 82.6% ± 2.5% of the administered dose can be found in urine and feces respectively after 8 days. Unchanged LPV accounted for nearly 2.2 and 19.8% of the administered dose in urine and feces, respectively. The apparent oral clearance of LPV is 5.98 ± 5.75 L/h. |

**Abbreviations:** LPV/r, lopinavir/ritonavir; AUC, area under the curve; C\(_{max}\), maximum plasma concentration; LPV, lopinavir; RTV, ritonavir.
of drugs and altered absorption. Extrapolating the dosage schedule followed in older children could lead to potential toxicity and/or lower exposure in young infants to ART, with incomplete virological suppression and the subsequent risk of viral resistance.27

The different absorption and distribution of LPV/r in young children compared with older children was initially observed by Verweel et al in a retrospective cohort study of 23 children who underwent a 12-hour pharmacokinetic sampling for LPV. Children under 2 years of age had a significantly lower Cmin and Cmax compared to children older than 2 years of age receiving LPV/r 230/57.5 mg/m2 twice daily. A dose increase by 37% resulted in an adequate LPV trough concentration in children younger than 2 years. Therefore a higher dose of 300/75 mg/m2 twice a day in children less than 2 years old was suggested.28 Chadwick et al investigated the pharmacokinetics, safety and efficacy of LPV/r plus two NRTIs in infants aged between 6 weeks and 6 months in a prospective trial including 21 infants. Infants received LPV/r 300/75 mg/m2 twice daily and were followed for 24 weeks. LPV/r clearance was slightly higher than that observed in older children, but the median AUC-time curve 0–12 h was similar to that seen in older children receiving 230/57.5 mg/m2 of LPV/r. The trough levels stabilized after the first 2 weeks of the study, which according to the authors, could be explained by improved absorption of the drug, better technique of administration, dietary changes or variation in RTV oral clearance.24

A median decrease in HIV viral load (VL) of 3.13 log10 copies/mL was observed at week 24, but only 38% had an undetectable VL (<400 copies/mL) at weeks 16 and 24. The virological response at week 16 was not associated with the LPV/r exposure at week 2, but viral suppression improved over time during the study. One infant interrupted treatment within the first 2 weeks of the study because of vomiting. 14.3% of the infants experienced a grade 3 or higher adverse events (transient neutropenia) but all were asymptomatic and transient. The authors concluded that a twice-daily dose of 300/75 mg/m2 of LPV/r in infants under 6 months gave a similar exposure to that reported in older children with favorable clinical and virological efficacy.24

Chadwick and colleagues also reported the use of LPV/r-based regimens in infants less than 6 weeks of age in a prospective, phase I/II study. This included 10 infants with confirmed HIV-1 infection aged between 2 and 6 weeks who received 300/75 mg/m2 of LPV/r twice daily plus 2 NRTIs and were followed 24 weeks. The median LPV AUC of 36.6 µg/mL was significantly lower than that found in infants aged between 6 weeks and 6 months of age. The half-life of LPV was similar to that seen in older age, so the authors postulated that reduced bioavailability of LPV could be the main cause for the lower LPV exposure rather than enhanced LPV metabolism. Altered absorption of LPV/r could be due to the food intake of the infants enrolled, though there was no correlation between the time and volume of formula milk received and LPV exposure in this study. The authors suggested that the addition of other foods might enhance LPV bioavailability in this age group. This needs to be evaluated in larger trials. This is particularly important because most infants that might benefit from early ART also suffer concomitant food insecurity.27

The authors assessed that despite the lower peak and average LPV exposure observed, the LPV trough was similar to that reported in older infants and an excellent virological response was achieved. The long-term follow-up of these patients would help to determine the variations of LPV/r pharmacokinetics and its long-term efficacy.27

These initial studies suggest that LPV/r may be used in young infants, but if used therapeutic drug monitoring (TDM) should be always considered. Data from larger trials are needed to evaluate the impact that other factors (like genetics or race) might have on LPV/r exposure during the first months of life. An improvement in the palatability of the oral formulation of LPV/r is urgently needed if extensive use of this drug is recommended after the early diagnosis of the HIV infection. Otherwise, poor adherence with possible incomplete viral suppression and resistance might happen early in life and make future antiretroviral treatment even more complex.

Monotherapy with LPV/r

The dramatic success of HAART has also brought important complications such as lipodystrophy, dyslipidemia or lactic acidosis. Complex dosing schedules with a high daily pill burden hinders the benefits of HAART and compromises its long-term use. Therefore, strategies like monotherapy with highly potent ART such as a boosted PI (PI/r) seemed an appealing way to enhance adherence and decrease complications. Single-drug antiretroviral therapy may be less toxic, easier to use and less costly while effectively maintaining long-term virological suppression and preserving future treatment options.29

Adult studies – LPV/r monotherapy after initial HAART

The use of LPV/r monotherapy as an NRTI-sparing treatment simplification strategy in patients with sustained viral
suppression was evaluated by Moltó et al in a retrospective cohort of 51 adults; 95% of the patients who were followed until 48 weeks maintained viral suppression, with a sustained increase in CD4% cell counts and a significant decrease in triglyceride levels. The authors concluded that this treatment simplification approach was safe and effective in routine clinical practice, especially in those patients already receiving a LPV/r based regimen.30

Similar results were obtained in the OK04 study group in a randomized, open label, non-inferiority clinical trial in 205 patients with suppressed viral replication on LPV/r plus 2 NRTI. Patients were randomized to continue HAART or LPV/r monotherapy with the reintroduction of 2 NRTIs if virological rebound was observed. At week 48, the percentage of patients without therapeutic failure was 94% in the monotherapy arm versus 90% in the triple therapy group. The percentage of patients with HIV RNA ≤50 copies/mL at week 48 by intention-to-treat analysis was 85% in the monotherapy groups versus 90% in the HAART arm, without statistically significant differences. Episodes of low level viremia were more common in patients receiving LPV/r monotherapy although the long-term significance of these “blips” is unknown. This study showed that LPV/r monotherapy (with reintroduction of NRTIs if needed) was non-inferior to conventional HAART.31

LPV/r monotherapy in ART-naïve adult patients

The use of LPV/r as monotherapy as initial therapy in adult patients was evaluated in the MONARK study;32 an open-label, randomized, 96-week clinical trial which compare the efficacy of LPV/r monotherapy (n = 83) with LPV/r plus ZDV and 3TC (n = 53) as an initial treatment in naïve patients with VL <100,000 copies/mL. LPV/r monotherapy showed lower rates of virological suppression; in an intent-to-treat analysis 64% of the patients in the monotherapy group showed VL <50 copies/mL, whereas 75% of those on conventional HAART maintained viral suppression. In the on treatment analysis, there was a significant difference on the patients who achieved an undetectable VL in both groups (80% in LPV/r monotherapy vs 95% in the HAART group). Three patients in the monotherapy arm without virological suppression acquired new resistance mutations with modest impact on LPV susceptibility. This study did not support the use of LPV/r monotherapy in ART-naïve adult patients.

Recently Ghosn et al reported the 96-week follow-up of the MONARK study.33 By an intention-to-treat analysis, 47% of those initially randomized to LPV/r monotherapy had sustained viral suppression (<50 copies/mL). The occurrence of low-level viremia in some patients during follow-up, with the subsequent risk of drug resistance, has discouraged the use of LPV/r monotherapy in ART-naïve HIV-infected adults.

Pediatric studies

Antiretroviral treatment in children achieves less viral suppression than in adults. Therefore, the use of LPV/r monotherapy in naïve children has been avoided and the potential role of LPV/r monotherapy in ART-experienced children with viral suppression evaluated with caution. A prospective clinical trial of the use of LPV/r monotherapy as maintenance in Thai children (aged between 2 and 28 years) after VL suppression is ongoing (HIV Netherlands Australia Thailand Research Collaboration). Results will not be available for some time. A Paediatric European Network for the Treatment of AIDS (PENTA) trial of LPV/r monotherapy after HAART is also planned. Meanwhile, based on the current evidence in adults, LPV/r monotherapy should be discouraged in children, particularly during the first years of life.

LPV/r once-daily regimens

Once-daily administration regimens could increase convenience and adherence as LePrevost et al showed with abacavir (ABC) plus 3TC use in children.34 However, the efficacy of once-daily ART is highly dependent on the maintenance of inhibitory concentrations throughout the entire dosing interval.35

Adult studies

In treatment-naïve adults the administration of LPV/r in a single daily dose showed similar immunological and virological outcomes to the standard twice-daily regimen. Based on this experience, several studies have evaluated this new strategy with other drugs such as SQV/r in adult HIV-infected patients.36,37

Pediatric studies

In children, the change to a once-daily combination including boosted atazanavir in extensively ART experienced-children was associated with a significant risk of virological failure. Limited data are available about the use of other PIs, such as LPV/r, in a once-daily regimen. Pilot studies of once-daily LPV/r in children (dosed at 460/115 mg/m2) found similar pharmacokinetics to adult studies. However there was marked variation between individuals and studies; the observed median Cmin were just above the
minimal effective concentration and around 50% of patients showed trough levels below inhibitory levels. The new tablet formulation gave better drug levels and less variation (see Table 4).38

The use of LPV/r once daily is still not recommended in pediatric antiretroviral treatment guidelines because there is no evidence yet from large trials.39 A PENTA trial is planned to study this in more detail (PENTA 18).40 All the pediatric trials of once-daily LPV/r have not included children under 2 years old who have a more erratic LPV/r exposure. If LPV/r once daily is used in older children TDM should be carried out. Nevertheless, the advent of the tablet formulation of LPV/r, particularly the low-dose tablet, increases the feasibility of LPV/r once-diary regimens in children.

Double boosted PI regimens

When planning salvage regimens for children failing on NRTI/NNRTI-based HAART; pediatricians are often faced with multiple resistance mutations and limited options. Furthermore, response to salvage treatment containing a single PI might be suboptimal if little efficacy remains in the NRTI component or if progressive toxicity is associated with continued used of NRTIs.41

Dual ritonavir-boosted PI (PI/r) regimens represent an option for salvage or maintenance therapy for patients with reverse transcriptase mutations or intolerance.42 Double PI/r have the advantage of the boosting effect of RTV on plasma levels of both PIs and in vitro data have shown a synergistic effect of saquinavir (SQV) when combined with LPV/r.43

The efficacy of dual boosted PI combination has been assed in few studies among children. The combination of LPV/r and saquinavir gave good virological responses, but was associated with increases in cholesterol. This combination is used in Thailand as an alternative for second line treatment in PI-naïve children who failed NRTIs/NNRTIs regimens, but the high pill burden makes adherence difficult (see Table 5).

Double boosted PI/r regimens should not be used alone in naïve patients with high viral loads because they have been shown insufficient to suppress viral replication. Landman et al reported 61 naïve HIV-infected adults who received fosamprenavir/atazanavir/RTV or saquinavir/atazanavir/RTV; viral suppression was only achieved in 40% and 42% of patients respectively at week 16.44

However, the encouraging results in pre-treated children suggest that double PI/r regimens may be an effective and safe option for these children. However, they also carry a high pill burden, and thus adherence support is extremely important. The additional benefit that TDM could add to this strategy needs to be studied in more detail in children, because a dose reduction could reduce the frequency of metabolic side effects without compromising its efficacy. The new low-dose tablet formulation of LPV/r allows the possibility of double PI/r regimen with less toxicity and better acceptability.

Table 4 Summary of the pediatric studies on LPV/r once-daily regimens

<table>
<thead>
<tr>
<th>Author</th>
<th>Methods</th>
<th>Results</th>
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<tr>
<td>Rosso et al</td>
<td>21 HIV-infected ART-naive children</td>
<td>Median Cmin 1.59 mg/L in the once daily group vs 7.90 mg/L in the twice daily group. Cmin inhibitory for wild-type virus (&gt;1.0 mg/L) in 4 out of 7 children in the once daily group. No significant differences in the Cmin between groups</td>
</tr>
<tr>
<td>van der Lee et al</td>
<td>19 HIV-1 infected ART-experienced children with VL &lt; 50 copies/mL for at least 6 months</td>
<td>Median Cmin 2.88 ± 3.74 mg/L, median Cmax 10.77 ± 2.90 mg/L, median AUC0–12 h 149.8 ± 58.8 h*mg/L – comparable to adults receiving 800/200 mg of LPV/r once daily. Cmin inhibitory for wild-type virus (&gt;1.0 mg/L) in 47% children, less in younger children</td>
</tr>
<tr>
<td>la Porte et al</td>
<td>7 pretreated children aged 5 to 15 years.</td>
<td>Once daily&lt;br&gt;Median Cmin = 3.4 mg/L&lt;br&gt;Median Cmax = 13.5 mg/L&lt;br&gt;Median AUC0–24 h = 214.6 h*mg/L&lt;br&gt;Mean Cmin 3.1 ± 2.6 mg/mL, mean Cmax 14.8 ± 2.4 mg/L, mean AUC0–24 h 217.9 ± 44.9 mg/L.h LPV half-life = 5.8 ± 4.5 h; median time to maximum concentration = 5.8 h. Every child included in the study had an undetectable VL at week 24 of follow-up</td>
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<tr>
<td>van der Flier et al</td>
<td>15 HIV-1 infected children who had received at least 24 weeks of LPV/r treatment (with soft gel capsules) and had achieved virological suppression.</td>
<td>Twice daily&lt;br&gt;Median Cmin = 5.7 mg/L&lt;br&gt;Median Cmax = 9.8 mg/L&lt;br&gt;Median AUC0–12 h = 80.9 h*mg/L</td>
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Abbreviations: LPV/r, lopinavir/ritonavir; Cmin, minimum plasma concentration; Cmax, maximum plasma concentration; PK, pharmacokinetic; AUC0–24 h, 24 h area under the plasma concentration-time curve; AUC12 h, 12 h area under the plasma concentration-time curve.
Table 5 Summary of pediatric studies on double boosted PI regimens

<table>
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<tr>
<th>Author</th>
<th>Methods</th>
<th>Results</th>
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<tr>
<td>Ananworanich et al54</td>
<td>HIV-NAT 017 group (1st study): 20 heavily pretreated Thai children: dose 50 mg/kg BD SQV plus 230/57.5 mg/m² twice daily LPV/r (50% also received 3TC)</td>
<td>HIV RNA was suppressed &lt;400 copies/mL in 80% of the children after 24 weeks of treatment. Median increase in CD4 counts = 6%</td>
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<tr>
<td>Kosalaraksa et al55</td>
<td>HIV-NAT 017 group (2nd study): 50 Thai children: dual boosted combination of SQV/LPV/r (as above) after 48 weeks of follow-up</td>
<td>Median rise in CD4% = 9%, median decrease in HIV RNA viral load = 2.8 log₁₀. Mean of both PIs exceeded therapeutic concentrations. 10% of patients had virologic failure associated with poor adherence, none selected major PI mutations. Median serum cholesterol and triglyceride increased significantly.</td>
</tr>
<tr>
<td>Bunupuradah et al56</td>
<td>HIV-NAT 017 group (3rd study): 96 week follow-up of the study conducted by Kosalarska et al (see above)</td>
<td>HIV RNA was suppressed below 400 copies/mL in 74% of the children after 96 weeks. 20% of patients had virologic failure. Median increase in CD4+ count = 558 cells/mm³. Total cholesterol and HDL increased significantly during the study, whereas triglycerides did not report any change in body shape. Both LPV and SQV Cₘₕ were high and stable during follow-up.</td>
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<tr>
<td>Robbins et al57</td>
<td>26 heavily experience-treated children and adolescents: Dose: either LPV/r 400/100 mg/m² twice daily without NNRTI or LPV/r 480/120 mg/m² twice daily with concomitant NNRTI. If the LPV inhibitory quotient [Cₘₕ/IC₅₀] was &lt;15, SQV was added</td>
<td>Median maximal decrease in viral load at week 8 = 1.57 log₁₀. However, the high dose was safe and well tolerated for up to 48 weeks. The significant initial increase of cholesterol did not worsen during the study and no significant gastrointestinal problems were observed even when SQV was added.</td>
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Abbreviations: LPV/r, lopinavir/ritonavir; SQV, saquinavir; IC₅₀, 50% inhibitory concentration; AUC₀–12, area under the plasma concentration-time curve; Cₘₕ, maximum plasma concentration; Cₘₖ, minimum plasma concentration; IC₅₀, minimum inhibitory concentration; NRTIs/NNRTIs, nucleoside reverse transcriptase inhibitor/non-nucleoside reverse transcriptase inhibitor; PI, protease inhibitor.

Low-dose LPV/r

The effect that different isoenzyme polymorphisms have on the pharmacokinetics of drugs such as efavirenz has raised the importance of genetic factors in determining the dosage of ART.45 Studies have looked at using LPV/r at reduced doses in certain ethnic groups.

Adult studies

Boyd et al showed a satisfactory response and adequate PK profile in Thai adults with a 50% reduction in the dose of indinavir/RTV. Lower doses of saquinavir/ritonavir (SQV/r) and LPV/r were also evaluated and a 70% dose of LPV/r is currently accepted for Thai adults.46

Pediatric studies

The desirable benefits of LPV/r lower doses such as cost reduction and decreased rates of metabolic side effects would be valuable in pediatric antiretroviral treatment. A pilot study of low-dose LPV/r conducted by Puthanakit et al included 24 ART-naïve HIV-infected Thai children aged between 2 and 18 years. Patients received 70% of the standard dose twice a day plus zidovudine (ZDV) and lamivudine (3TC). At week 48, there were no statistically significant differences in CD4% or viral load between children receiving the standard dose and those in the low-dose arm. The AUC₀–12 and the Cₘₕ were lower in the low-dose arm (by 29% and 31% respectively).47

This pilot study included a small number of patients, and a LPV/r dose-reduction is not yet recommended for PI-experienced children or infants. However the relevance of pharmacogenetic aspects on dosage schedule is proven and the need to adapt guidelines to different settings warranted. This study also suggests that TDM should be considered in all children on LPV/r in order to find out if a lower dose could be used. The advent of the new low-dose LPV/r tablets encourages the development of reduced dose studies among children due to its improved palatability and easier storage requirements compared with the liquid formulation.

Conclusions

The benefits achieved with LPV/r-based therapies in HIV-infected children outweigh the metabolic adverse effects observed with its prolonged used. Therefore, until other PIs develop child friendly formulations, LPV/r will be considered the first choice for PI in childhood. The new tablet low-dose...
formulation increases the possibility for more convenient antiretroviral treatment in children.

Novel strategies of LPV/r use could maximize its advantages during childhood; in infants a higher dosage with close TDM is a safe and effective option, especially in those exposed to single-dose NVP. Older children and adolescents could benefit from a once-daily LPV/r regimen and certain populations could be treated with lower doses if TDM can be guaranteed. Finally heavily pre-treated children might benefit from a double PI/r regimen which includes LPV/r.

All the previous approaches emphasize the lessons learnt with antiretroviral treatment in the past; high potency therapies need to be balanced with convenient regimens, to enhance adherence. No drug is effective if it is not taken and this is particularly challenging in children where both the patient and the caregiver needs to adhere to treatment.

Disclosures

The authors declare no conflicts of interest.

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