The Antiretroviral Guide
A Tool for Providing Seamless Care and Assessing Antiretroviral Therapy in Hospitalized HIV+ Patients

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# A Tool for Providing Seamless Care and Assessing Antiretroviral Therapy in Hospitalized HIV+ Patients

## STEP 1 Admission Assessment

### Initial Patient Assessment

<table>
<thead>
<tr>
<th>Component</th>
<th>Comments</th>
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| **Medical History** | • Confirm admission diagnosis/HIV status  
• Depending on institutional practice, consider informing HIV clinic of admission for continuity of care.  
• Summary of previous and current medical conditions, including HBV, HCV, OIs, STIs, psychiatric, metabolic, etc.  
• Pregnancy or possibility of pregnancy  
• Vital signs, ROS, height, weight |
| **Social History**     | • Living arrangements  
• Income stability/job security  
• Social/family support  
• Alcohol/addictions/recreational drug use  
• Drug coverage plan (include ARV coverage, coverage for other medications) |
| **Laboratory Tests**   | • HIV-specific labs, including most recent CD4 count and HIV viral load  
• HAV, HBV, HCV status, toxoplasmosis serology, tuberculosis status if available  
• CBC, electrolytes  
• Organ function (assess overall stability)  
  - Renal (Scr, CrCl for renal drug dosing adjustments)  
  - Hepatic (ALT, AST, ALP, bilirubin, albumin, INR) |
| **BPMH/Medication Reconciliation** | • Allergies/intolerances  
  - Clarify the reaction, drug involved, date, and required treatment  
• Current ARV regimen; study drugs  
• Other prescription and non-prescription drugs, including inhalers, patches, topical medications, recent intra-articular injections (e.g. corticosteroids)  
• CAM/Herbal medications  
*Note: For all medications, clarify indication, drug, dose, frequency, formulation, route of administration and adherence*  
**Hospital Admission ARV Seamless Care Tips:**  
• If patient was taking ARVs prior to admission, was the patient adherent? Check with patient, outpatient refill history, community pharmacy, HIV program.  
• Check for any reasons why ARVs should be held in the hospital (recent non-adherence in the community, patient instability, significant drug toxicity on admission, significant illness in hospital, NPO, etc).  
• In NPO/critical care/severe nausea patients it might be necessary to stop all ARVs for the short-term depending on feeds and drug malabsorption issues.  
• Avoid use of partial ARV regimens to minimize the development of resistance (continue all drugs or stop all drugs together). If uncertain consult with HIV program.  
• Check if the patient is receiving therapy for HBV or HCV co-infection as these therapies should generally be continued during hospitalization.

ARV: antiretroviral; FDC: fixed dose combination; HAV: hepatitis A virus; HBV: hepatitis B virus; HCV: hepatitis C virus; NPO: nothing by mouth; OIs: opportunistic infections; ROS: review of systems; STIs: sexually transmitted infections
### Assess Antiretroviral (ARV) Therapy on Admission

| **Is it the correct therapy?**  | • Usually HIV is treated with 3 active drugs; however some patients may be on > 3 drugs in cases of resistance.  
|                              | • There is also ongoing research on the use of 2-drug combinations.  
|                              | • There are many new co-formulations with several drugs included called fixed dose combinations (FDCs) or single tablet regimens (STRs).  
|                              | • Ritonavir and cobicistat are not considered “active drugs” (they are pharmacokinetic boosters to increase concentrations of certain ARVs).  
| (See Antiretroviral Agents/Handy Resources) | |  
| **Is there adequate ARV stock/drug coverage?** | • Ensure there is a supply of ARVs- check with patient, hospital stock, dispensing outpatient or community pharmacy.  
|                              | • Ensure the patient has active ARV drug coverage and is aware of potential drug costs upon discharge (see Discharge Assessment/Handy Resources).  
| **Are the doses/formulations correct?**  | • In some cases ARV doses may differ from the product monograph. Verify with the outpatient/community pharmacy or HIV program if needed.  
| (See Antiretroviral Agents/Handy Resources) | • Ensure doses are adjusted for significant renal/hepatic dysfunction or dialysis.  
|                              | • Some FDCs should be avoided if the CrCL < 50 mL/min and need to be split up into single drug formulations. When uncertain, consult with the HIV program.  
|                              | • Ensure the formulation is correct. Most ARVs are available in tablets or capsules and there are a few liquids; only zidovudine is available IV.  
|                              | • Consult specialized information on liquids, crushing tablets, or opening capsules. (See Handy Resources)  
| **Is therapy effective?** | • Verify CD4 count and viral load. Ideally the CD4 count should be > 200 cells/µL (i.e. > 0.200 x 10^9/L) to prevent OIs, although some patients are not able to achieve this degree of immune reconstitution.  
|                              | • The HIV viral load should be undetectable/not quantifiable if the patient is responding well to therapy. If the viral load is > 200-250 cells/µL while on ARVs, a genotypic ARV resistance test (GART) might be indicated (consult with HIV/ID team).  
|                              | • Monitoring efficacy: When starting therapy the HIV viral load is measured after 4-8 weeks to assess the initial response to therapy. In general, the CD4 count and viral load are monitored every 3-6 mos, depending on the response to treatment and the stability of the patient.  
|                              | • If the CD4 count is < 200 cell/µL, OI prophylaxis may be required to prevent certain infections like Pneumocystis pneumonia (PCP or PJP) (< 200), toxoplasmosis (< 100, if toxo Ab +) and Mycobacterium avium complex (MAC) (< 50). (See Handy Resources - OI guidelines)  
| **Is therapy safe?**  | • Ensure the patient is tolerating the current ARV regimen.  
| (See Antiretroviral Agents/Handy Resources) | • Common problems include GI (nausea, anorexia, diarrhea) and metabolic toxicities (high lipids, diabetes).  
|                              | • More serious toxicities may include skin rashes (not always serious), renal failure, hepatic failure (less common), pancreatitis, and anemia.  
| **Are there any drug-drug interactions?**  | • Common drug interactions involve absorption (pH and chelation/complexation interactions); metabolic (CYP450 3A4/2D6 and P-gp inhibition and induction interactions); and additive toxicity (renal, cytopenias).  
| (See Antiretroviral Agents/Handy Resources) | |  
| **Are there any scheduling issues?**  | • Most ARVs are best tolerated/absorbed with food; try to accommodate patient preferences when scheduling ARVs.  
| (See Antiretroviral Agents/Handy Resources) | • It is important to give a once daily regimen all at the same time and to give pharmacokinetic boosters (ritonavir, cobicistat) at the same time as the drugs they are boosting (e.g. protease inhibitors).  
|                              | • BID regimens should be scheduled q12h.  
| **Can the patient adhere to therapy?**  | • Ensure the patient is able to adhere to therapy during the hospitalization and whether this can be continued after discharge. There may be a number of factors that can affect short and long-term adherence (NPO, inability to eat/swallow, severe nausea, day passes, social, housing, addictions, toxicities, formulations issues, etc).  
| (See Antiretroviral Agents/Handy Resources) | |
STEP 2 Assessment During Course of Hospitalization

- For patients on ARVs, review medication profile daily or when medication changes are made.
- Monitor for common errors that may occur when transitioning from units including drug omissions, drug dosing issues, drug interactions with concurrent therapies prescribed over the course of hospitalization, scheduling of medications with food, auto-stops on antimicrobials (including ARVs and OI treatment/prophylaxis), etc.
- Monitor laboratory tests for efficacy and toxicity if these tests are ordered during hospitalization. Efficacy: CD4 count and HIV viral load (every 3-6 mos). Toxicity: CBC/diff, renal/hepatic function, GI effects. Long-term effects drug-specific (e.g. ↑ lipids/glucose, ↓ bone mineral density [BMD]).
**STEP 3** Discharge Assessment

**Assess Discharge Prescriptions**
- Discharge ARVs should be ordered and forwarded to the dispensing pharmacy with adequate time for preparation.
- If changes were made during hospitalization, ensure that the changes were approved by an HIV healthcare provider.
- Ensure that authorization for drug coverage is completed prior to discharge (see Handy Resources).
- Ensure OI treatment/prophylaxis prophylaxis medications are ordered if indicated.
- Verify that all other medications are ordered as appropriate including prescription, OTC and PRN drugs.
- If still indicated, re-start medications that were held on admission or during the course of hospitalization.

**ARV Dispensing/Coverage**
- Verify coverage of ARVs and other drugs.
- Consult with HIV program if drug coverage is an issue.
- Each province has different ARVs that are covered and drug coverage policies. (see Handy Resources)
- Other forms of drug coverage include:
  - Non-Insured Health Benefits (NIHB)
  - Interim Federal Health (IFH)
  - Private Insurance
  - Compassionate Access from pharmaceutical industry

**ARV Adherence**
- Address potential for non-adherence in outpatient setting.
- Reinforce important adherence and food requirements.
- Assess whether special adherence aids are required:
  - Medication schedule
  - Blister pack or daily observed therapy (DOT) at outpatient/community pharmacy
  - Consider giving DOT ARVs with daily opioids/methadone to increase adherence
  - Beepers, reminders, supports
  - Delivery of medications

**Outpatient Follow-up**
- Arrange for follow-up with HIV physician/care team.
- Arrange for follow-up with other health care providers such as the family physician.
- Communicate any changes in drug therapy to outpatient health care providers (e.g. physicians, HIV team, outpatient/community pharmacy).

### Antiretroviral Agents

<table>
<thead>
<tr>
<th>Drug/Trade Name</th>
<th>Formulations/Strengths</th>
<th>Usual Adult Dose/Food</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Individual Antiretrovirals</strong></td>
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<tr>
<td>NRTIs (Nucleoside Reverse Transcriptase Inhibitors)</td>
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<tr>
<td>abacavir (ABC)</td>
<td>Tab: 300 mg&lt;br&gt;Sol: 20 mg/mL</td>
<td>300 mg BID OR 600 mg daily&lt;br&gt;Take with or without food</td>
<td>• May ↑ risk of myocardial infarction&lt;br&gt;• Risk of HSR in individuals + for the HLA-B5701 gene; screen required before initiation; if + test, avoid abacavir&lt;br&gt;• Few drug interactions</td>
</tr>
<tr>
<td>Zidovudine (ZDV)</td>
<td>EC Cap: 125,200,250, 400 mg&lt;br&gt;Sol: 4 g/240 mL (SAP)</td>
<td>200 mg BID OR 400 mg daily&lt;br&gt;Take 90min ac or 2h pc</td>
<td>• GI intolerance&lt;br&gt;• Peripheral neuropathy, pancreatitis&lt;br&gt;• Few drug interactions</td>
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<tr>
<td>didanosine ( ddl)</td>
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<tr>
<td>Emtricitabine (FTC)</td>
<td>Cap: 200 mg (US) &lt;br&gt;Sol: 10 mg/mL (US)</td>
<td>200 mg daily&lt;br&gt;Take with or without food</td>
<td>• Well tolerated&lt;br&gt;• Few drug interactions&lt;br&gt;• Active against HBV&lt;br&gt;• Only available in Canada in a FDC</td>
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FDC: Fixed Dose Combination; HSR: hypersensitivity reaction
## 3. Antiretroviral Agents Continued

<table>
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<tr>
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</table>
| lamivudine (3TC) | Tab: 100, 150, 300 mg; Sol: 10 mg/mL; Note: 100 mg tabs also for HBV infection (Hepovir) | 150 mg BID OR 300 mg daily Take with or without food | • Well tolerated  
• Few drug interactions  
• Active against HBV |
| stavudine (d4T) | Cap: 15, 20, 30, 40 mg; Sol: 1 mg/mL (SAP) | 300 mg daily Take with or without food | • Peripheral neuropathy, pancreatitis  
• Hyperlipidemia  
• Few drug interactions |
| tenofovir disoproxil fumarate (TDF) | Tab: 150, 200 (US); 300 mg; Pwdr: 40 mg/g (US) | 300 mg daily Take with or without food | • Nephrotoxicity; ↓ in bone mineral density (BMD)  
• Few drug interactions  
• Active against HBV |
| tenofovir alafenamide (TAF) | See FDC products | See FDC products | • TAF will largely replace TDF in most tenofovir formulations.  
• ↓ renal and bone toxicity with TAF vs. TDF  
• More drug interactions than TDF; avoid with potent P-gp inducers; dose adjust with P-gp inhibitors  
• Active against HBV |
| zidovudine (AZT, ZDV) | Cap: 100 mg; Tab: 300 mg (US); IV: 10 mg/mL; Syrup: 10 mg/mL | 300 mg BID OR 200 mg TID Take with or without food | • GI intolerance  
• Headache, insomnia  
• Bone marrow suppression, macrocytic anemia, neutropenia  
• Few drug interactions |

### NNRTIs (Non-Nucleoside Reverse Transcriptase Inhibitors)

<table>
<thead>
<tr>
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</table>
| efavirenz (EFV) | Cap: 50, 200 mg; Tab: 600 mg | 600 mg daily Take qHS on empty stomach or with low-fat snack to minimize CNS S/E | • CNS effects- vivid dreams, nightmares, insomnia, dizziness  
• Rash (usually self-limiting, unless high risk features)  
• Hyperlipidemia  
• Inducer of CYP3A4, 2B6  
• Avoid in pregnancy if possible |
| etravirine (ETV) | Tab: 25, 100, 200 mg | 200 mg BID OR 400 mg daily Take with food | • Nausea  
• Rash (usually self-limiting, unless high risk features)  
• Inducer of CYP3A4 (weak)  
• Inhibitor of CYP2C, 2C19 (weak-moderate) |
| nevirapine (NVP) | IR Tab: 200 mg; XR Tab: 400 mg; Syrup: 10 mg/mL (SAP) | IR: 200 mg daily x 14 days (lead-in) then 200 mg BID OR 400 mg daily XR: 400 mg daily (after 14 day lead-in) Take with or without food | • Rash (may be more serious with hepatitis, check for high risk features)  
• Avoid starting in men with CD4>400 and women with CD4>250 due to ↑ risk of hepatitis  
• Inducer of CYP3A, 2B6 |
### 3. Antiretroviral Agents Continued

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<tr>
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<th>Usual Adult Dose/Food</th>
<th>Comments</th>
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<tbody>
<tr>
<td><strong>rilpivirine (RPV)</strong></td>
<td>Tab: 25 mg</td>
<td>25 mg daily</td>
<td>- Headache, dizziness, insomnia, vivid dreams, depression (mild-moderate)</td>
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<tr>
<td>Edurant</td>
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<td>50 mg daily with rifabutin</td>
<td>- Do not administer with PPIs (CI)</td>
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<tr>
<td>FDC: Complera, Odefsey</td>
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<td>Take with a meal (400 kcal minimum)</td>
<td>- Spacing required with H2RAs and/or antacids (↑ pH decreases RPV absorption)</td>
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<td>- Do not administer with a liquid nutritional drink (↓ RPV absorption)</td>
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<td>- Inducers/inhibitors of CYP3A may affect RPV concentrations</td>
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<td>- Avoid initiation if viral load &gt; 100,000 c/mL or CD4 &lt; 200 cells/µL</td>
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<tr>
<td><strong>PIs (Protease Inhibitors)</strong></td>
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<td></td>
<td>- Headache, dizziness, insomnia, vivid dreams, depression (mild-moderate)</td>
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<tr>
<td><strong>atazanavir (ATV)</strong></td>
<td>Cap: 100 mg (US), 150, 200, 300 mg Pwdr: 50 mg/1.5 g dispersible oral powder packet (US)</td>
<td>400 mg daily (unboosted) OR 300 mg daily with RTV 100 mg (boosted) Take with food</td>
<td>- Lower risk for metabolic S/E than other PIs</td>
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<tr>
<td>Reyataz</td>
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<td>- Avoid/space from antacids, H2RAs, and/or PPIs (ATV absorption)</td>
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<tr>
<td>FDC: Evotaz</td>
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<td>- Inhibitor of CYP3A, UGT1A1</td>
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<td></td>
<td>- Use with PK booster recommended; may also use unboosted</td>
</tr>
<tr>
<td><strong>darunavir (DRV)</strong></td>
<td>Tab: 75, 150, 400, 600, 800 mg Susp: 100 mg/mL*</td>
<td>DRV 600 mg + RTV 100 mg BID OR DRV 800 mg + RTV 100 mg daily (naïve subjects) Take with food</td>
<td>- GI intolerance</td>
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<tr>
<td>Prezista</td>
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<td></td>
<td>- Lower risk for metabolic S/E than other PIs</td>
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<tr>
<td>FDC: Prezobix</td>
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<td></td>
<td>- Inhibitor of CYP 3A4</td>
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<td></td>
<td></td>
<td>- Use with PK booster required</td>
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<td></td>
<td>- DRV/RTV BID dosing often used in more experienced patients with underlying DRV resistance (see product monograph)</td>
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<tr>
<td><strong>fosamprenavir (fAPV)</strong></td>
<td>Tab: 700 mg Susp: 50 mg/mL</td>
<td>IAPV 1400 mg BID (unboosted) OR IAPV 700 mg + RTV 100 mg BID (boosted) OR IAPV 1400 mg + RTV 100-200 mg daily (boosted) Take tabs with or without food; Susp ac</td>
<td>- GI intolerance</td>
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<tr>
<td>Telzir / Lexiva (US)</td>
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<td>- Rash (usually self-limiting, unless high risk features)</td>
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<td></td>
<td>- Metabolic S/E</td>
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<td></td>
<td></td>
<td></td>
<td>- Inhibitor of CYP 3A4</td>
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<td></td>
<td></td>
<td></td>
<td>- Use with RTV PK booster recommended</td>
</tr>
<tr>
<td><strong>lopinavir (LPV)</strong></td>
<td>Tab: 250,625 mg Pwdr: 50 mg/g (US)</td>
<td>1250 mg BID OR 750 mg TID (unboosted) Take with food</td>
<td>- GI intolerance (diarrhea- treat with fiber, calcium supplements)</td>
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<tr>
<td>(see Kaletra under Fixed-Dose Combination (FDC) Products)</td>
<td></td>
<td></td>
<td>- Metabolic S/E, lipodystrophy</td>
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<tr>
<td><strong>nelfinavir (NFV)</strong></td>
<td>Tab: 250 mg Tab: 125 mg Pwdr: 50 mg/g (US)</td>
<td>1250 mg BID OR 750 mg TID (unboosted) Take with food</td>
<td>- Inhibitor of CYP3A4</td>
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<tr>
<td>Viracept</td>
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<td>- Only non-boostable PI</td>
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<td></td>
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<td>- High variability in absorption</td>
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3. ANTIRETROVIRAL AGENTS CONTINUED

<table>
<thead>
<tr>
<th>Drug/Trade Name</th>
<th>Formulations/Strengths</th>
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<tr>
<td><strong>INSTIs (Integrase Strand Transfer Inhibitors)</strong></td>
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<tr>
<td>dolutegravir (DTG)</td>
<td>Tab: 50 mg Peds: 10, 25 mg tab*; 5 mg dispersible tab (all under study)*</td>
<td>50 mg daily (naïve subjects) OR 50 mg BID (experienced subjects or with certain CYP450 enzyme inducers) Take with or without food</td>
<td>• Well tolerated • GI intolerance, headache, insomnia • CK and/or transaminase elevation • Non-pathogenic ↑ SCr due to inhibition of renal tubular secretion (SCr: 10-15 µmol/L ↑) • Fewer drug interactions • Inducers/inhibitors of UGT1A1/CYP3A4 may alter DTG concentrations • Administer DTG 2h before or 6h after taking medications containing polyvalent cations (eg. Al, Ca, Fe, Mg, Zn) - (↓ DTG absorption); however may be taken with food at the same time as Ca and Fe</td>
</tr>
<tr>
<td>elvitegravir (EVG)</td>
<td>Tab: 85,150 mg*</td>
<td>Usual dose 150 mg daily with cobicistat 150 mg daily (boosted regimen) Take with food</td>
<td>• Well tolerated • GI intolerance, headache • CK and/or transaminase elevation • Non-pathogenic ↑ SCr due to inhibition of renal tubular secretion by cobicistat (SCr: 10-15 µmol/L ↑) • Modest inducer of CYP 2C9 • Cobicistat PK booster required • Administer EVG 2h apart from antacids or vitamin/mineral supplements containing polyvalent cations (eg. Al, Ca, Fe, Mg, Zn) - (↓ EVG absorption)</td>
</tr>
<tr>
<td>raltegravir (RAL)</td>
<td>Tab: 400 mg Chew Tab: 25,100 mg Pwdr: 20 mg/mL oral banana flavoured granular powder (100 mg/packet) (available in US; SAP in Canada)* 600 mg OD tab under study*</td>
<td>400 mg BID Take with or without food 1200 mg daily (2 x 600 mg OD tabs)-under study*</td>
<td>• Well tolerated • GI intolerance, headache, pyrexia • CK and/or transaminase elevation • Fewer drug interactions • Inducers/inhibitors of UGT1A1 may alter RAL concentrations • Concurrent or staggered administration not recommended with Al and/or Mg. May be given with antacids containing CaCO3. • Space from Fe, Zn by several hours (↓ RAL absorption) Note: 600 mg tabs may have different cation spacing recommendations once marketed</td>
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<tr>
<td><strong>CCR5 Receptor Antagonist</strong></td>
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<td>maraviroc (MVC)</td>
<td>Tab: 150, 300 mg</td>
<td>150-600 mg BID, depending on regimen and drug interactions Take with or without food</td>
<td>• Well-tolerated • GI intolerance, headache, orthostatic hypotension • Hepatotoxicity • Fewer drug interactions • Inducers/inhibitors of CYP3A4/P-gp may affect MVC concentrations (recent tropism screening test required; consult with HIV team regarding testing) • Only effective if virus has R5 tropism (screening test required)</td>
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### 3. ANTIRETROVIRAL AGENTS CONTINUED

#### Pharamcokinetic (PK) Boosters

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</table>
| ritonavir (RTV) | Tab: 100 mg, Sol: 80 mg/mL | 100-200 mg daily/BID as PK booster | - GI intolerance
- Hepatitis
- Metabolic S/E
- Many drug interactions
- Inducer of CYP 3A4, P-gp > 2D6
- Inducer of CYP 1A2, 2B6, 2C9, 2C19, UGT (clinically significant)
- Not used for ARV properties; used as a PK booster |
| Norvir | | Take with food | |
| FDC: Kaletra | | | |

| Cobicistat (cobi) | Tab: 150 mg* | 150 mg daily as a PK booster; use with daily EVG 150 mg, ATV 300 mg and DRV 800 mg | - Headache, insomnia, GI intolerance
- Non-pathogenic ↑ SCr due to inhibition of renal tubular secretion (SCr: 10-15 µmol/L ↑)
- Many drug interactions
- Inhibitor of CYP 3A4, P-gp > 2D6
- No ARV activity; used as a PK booster |
| Tybost* | | Take with food with other ARVs | |
| FDC: Stribild, Genvoya, Prezolobis, Evotaz | | | |

#### Fixed-Dose Combination (FDC) Antiretroviral Products

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<tr>
<td><strong>NRTI Backbones</strong></td>
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</table>
| Combivir | Zidovudine 300 mg Lamivudine 150 mg Tab | 1 tab BID Avoid if CrCl < 50 mL/min Take with or without food | - GI intolerance
- Headache, insomnia
- Bone marrow suppression, macrocytic anemia, neutropenia
- Few drug interactions |
| Descovy* | Tenofovir alafenamide (TAF) 10 and 25 mg Emtricitabine 200 mg Tab | 10/200 mg tab with RTV or cobicistat-boosted regimens 25/200 mg tab with other unboosted ARVs Avoid if CrCl < 50 mL/min Take with or without food | - ↑ renal and bone toxicity with TAF vs. TDF
- More drug interactions than TDF; avoid with potent P-gp inducers; dose adjust with P-gp inhibitors
- Active against HBV |
| Kivexa/Epzicom (US) | Abacavir 600 mg Lamivudine 300 mg Tab | 1 tab daily Avoid if CrCl < 50 mL/min Take with or without food | - May ↑ risk of myocardial infarction
- Risk of HSR in individuals + for the HLA-B5701 gene; screen required before initiation; if + test, avoid abacavir
- Few drug interactions |
| Trizivir | Zidovudine 300 mg Lamivudine 150 mg Abacavir 300 mg Tab | 1 tab BID Avoid if CrCl < 50 mL/min Take with food | - See Kivexa and Combivir comments |
| Truvada | Tenofovir (TDF) 300 mg Emtricitabine 200 mg Tab | 1 tab daily Adjustments required if CrCl ≤ 50 mL/min. Avoid if CrCl <30 mL/min or dialysis Take with or without food | - Nephrotoxicity; ↓ in bone mineral density (BMD)
- Few drug interactions
- Active against HBV |
### 3. ANTIRETROVIRAL AGENTS CONTINUED

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<th>Composition</th>
<th>Usual Adult Dose</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PI-based (with PK booster)</strong></td>
<td></td>
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</tr>
<tr>
<td>Evotaz*</td>
<td>Tenofovir 300 mg Atazanavir 300 mg Cobicistat 150 mg</td>
<td>1 tab daily</td>
<td>Avoid if CrCl &lt; 70 mL/min and also on TDF Take with food. See atazanavir and cobicistat comments.</td>
</tr>
<tr>
<td>Kaletra</td>
<td>Lopinavir/Ritonavir 100/25 mg (peds), 200/50 mg Tab</td>
<td>4 tabs (=800/200 mg) daily</td>
<td>Take with food (tabs, sol). GI intolerance, diarrhea. Higher risk for metabolic S/E than other PIs. Inhibitor of CYP 3A4; see RTV comments.</td>
</tr>
<tr>
<td>Prezcobix</td>
<td>Darunavir 800 mg Cobicistat 150 mg</td>
<td>1 tab daily</td>
<td>Avoid starting if CrCl &lt; 70 mL/min and also on TDF (e.g. Truvada, Viread). Take with food. See darunavir and cobicistat comments.</td>
</tr>
<tr>
<td><strong>INSTI-based Single Tablet Regimens (STRs)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genvoya*</td>
<td>Tenofovir alafenamide (TAF) 10 mg Emtricitabine 200 mg Elvitegravir (EVG) 150 mg Cobicistat 150 mg</td>
<td>1 tab daily</td>
<td>Avoid if CrCl &lt; 30 mL/min. Take with food. Administer 2h apart from antacids or vitamin/mineral supplements containing polyvalent cations (eg. Al, Ca, Fe, Mg, Zn) (EVG absorption). renal and bone toxicity with TAF vs. TDF. See Descovy, elvitegravir and cobicistat comments.</td>
</tr>
<tr>
<td>Stribild</td>
<td>Tenofovir (TDF) 300 mg Emtricitabine 200 mg Elvitegravir (EVG) 150 mg Cobicistat 150 mg</td>
<td>1 tab daily</td>
<td>Avoid starting if CrCl &lt; 70 mL/min. Discontinue if CrCl &lt; 50 mL/min. Take with food. Administer 2h apart from antacids or vitamin/mineral supplements containing polyvalent cations (eg. Al, Ca, Fe, Mg, Zn) (EVG absorption). renal and bone toxicity with TDF vs. TAF. See Truvada, elvitegravir and cobicistat comments.</td>
</tr>
<tr>
<td>Triumeq</td>
<td>Abacavir 600 mg Lamivudine 300 mg Dolutegravir (DTG) 50 mg</td>
<td>1 tab daily</td>
<td>Avoid if CrCl &lt; 50 mL/min. Take with or without food. Note: Additional 50 mg of dolutegravir should be given 12 hours after Triumeq if co-administered with certain CYP3A4 enzyme inducers. Administer 2h before or 6h after taking medications containing polyvalent cations (eg. Al, Ca, Fe, Mg, Zn) - (DTG absorption); however may be taken with food at the same time as Ca and Fe. See Kivexa and dolutegravir comments.</td>
</tr>
<tr>
<td><strong>NNRTI-based Single Tablet Regimens (STRs)</strong></td>
<td></td>
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<tr>
<td>Atripla</td>
<td>Tenofovir (TDF) 300 mg Emtricitabine 200 mg Efavirenz 600 mg</td>
<td>1 tab daily (hs)</td>
<td>Avoid if CrCl &lt; 50 mL/min. Take on an empty stomach. Take qHS on an empty stomach or with low-fat snack (to minimize CNS S/E of efavirenz). See Truvada and efavirenz comments.</td>
</tr>
<tr>
<td>Complera</td>
<td>Tenofovir (TDF) 300 mg Emtricitabine 200 mg Rilpivirine 25 mg</td>
<td>1 tab daily</td>
<td>Avoid if CrCl &lt; 50 mL/min. Take with a meal (400 Kcal). See Truvada and rilpivirine comments.</td>
</tr>
<tr>
<td>Odefsey*</td>
<td>Tenofovir alafenamide (TAF) 25 mg Emtricitabine 200 mg Rilpivirine 25 mg</td>
<td>1 tab daily</td>
<td>Avoid if CrCl &lt; 30 mL/min. Take with a meal (400 Kcal). See Descovy and rilpivirine comments.</td>
</tr>
</tbody>
</table>

* May not be covered provincially; may be available via compassionate access (verify with HIV program/manufacturer) or Special Access Program (SAP) - Health Canada.
Handy Resources

Canadian Resources
Canadian HIV and HCV Services: http://hiv411.ca/
Programme National de Mentorat sur le HIV-SIDA (French): http://pnmvs.org/

ARV Drug Coverage
ARV Drug Access & Coverage in Canada:
Non-Insured Health Benefits (NIHB):

HIV Drug Information • HIV Patient Resources • Drug Interactions
Toronto General Hospital Site and HIV/HCV app: http://hivclinic.ca / http://app.hivclinic.ca
University of Montreal Site- HIV Medication Guide (in French also): www.hivmedicationguide.com
CATIE HIV/HCV Information Canadian Site: www.catie.ca
University of Liverpool Site (App)- HIV and HCV sites: www.hiv-druginteractions.org/ / www.hep-druginteractions.org/
HIV Insite (UCSF): http://hivinsite.ucsf.edu/insite?page=ar-00-02

HIV Drug Dosing in Renal or Hepatic impairment and Dialysis
Toronto General Hospital Site and HIV/HCV app:
HIV Insite (UCSF): http://hivinsite.ucsf.edu/InSite?page=md-rr-18

Crushing HIV Medications • ARV Liquid Formulations
Toronto General Hospital Site (see Crushing and Liquids) and HIV/HCV app:

Enteral ARV Administration

Opportunistic Infection (OI) Guidelines

HIV and Pregnancy
Maternik: http://www.catie.ca/en/resources/maternik
Oak Tree (BC) Perinatal Guidelines: http://www.bcwomen.ca/health-professionals/professional-resources/hiv-aids-resources/hiv-aids-clinical-guidelines
Saskatchewan Perinatal Protocols: https://www.skhiv.ca/pregnancy