Switching to Rezolsta® (darunavir+cobicistat) or Evotaz® (atazanavir+cobicistat)

- Rezolsta and Evotaz are once daily fixed dose combinations (FDCs) of darunavir (DRV) 800 mg with cobicistat 150 mg, and atazanavir (ATV) 300 mg with cobicistat 150 mg respectively, and are indicated in combination with other antiretroviral medicinal products for the treatment of HIV-1 infection in adults aged 18 years or older.*

- There are clinical differences between ritonavir (r) and cobicistat boosted protease inhibitors (PIs) which need to be considered prior to switch:
  - Cobicistat’s drug interactions differ in that, unlike ritonavir, cobicistat does not induce glucuronidation (UGT1A1) or some CYP enzymes. Consequently, switching from PI/r to PI/cobicistat may increase levels of some drugs metabolised via these routes and route monitoring and/or dose modification (see table). Alternatively, consider remaining on ritonavir-boostered PI.
  - Cobicistat decreases estimated creatinine clearance by average 10 ml/min due to inhibition of tubular secretion of creatinine. This does not affect the actual glomerular filtration rate (GFR).

- Although bioequivalent to DRV+r 800+100 mg, DRV Cmin reduces by ~25-30% with cobicistat, but remains above the IC50 for DRV wild type virus. This may be relevant for some cohorts.

- The guidance below addresses key differences when switching from DRV+r or ATV+r to the FDCs and should be used in addition to the Product Labels and www.hiv-druginteractions.org.

### Darunavir-ritonavir to Rezolsta

**Rezolsta is NOT RECOMMENDED** if:

- Patient requires DRV 600 mg twice daily.
- Taking any of the following, which are not recommend with Rezolsta but may be used with DRV+r (with dose adjustment to twice daily for some drugs):
  - ARVs:
    - Efavirenz
    - Etravirine
    - Nevirapine
  - Anti-convulsants:
    - Carbamazepine
    - Phenoobarbital
    - Phenytoin
  - Other:
    - Bosantan
    - OBV/PTV/r ± D5V (AbbVie "2D/3D")**

- There are any DRV resistance associated mutations.
- eGFR <70 ml/min when co-administered with medicines that require dose adjustment based on renal function (e.g. tenofovir DF, lamivudine or emtricitabine).
- Patients aged <18 years where the safety and efficacy of Rezolsta has not been established.

**Rezolsta should be used with CAUTION** when:

- There are concerns about lower DRV exposure (compared to DRV+r) such as:
  - Protease inhibitor monotherapy.
  - Pregnancy, including if actively planning to conceive.
  - Where DRV is part of a regimen for patients with HIV encephalopathy or in patients with CSF HIV RNA escape.
  - Taking any other medicines, particularly those listed in the table. Conduct full medicines review.
  - Any combinations other than 2 NRTIs + Rezolsta as these have not been studied.

**Specific counselling points for Rezolsta:**

- The recommended dosing regimen is one tablet of Rezolsta taken once daily with food.
- Use all Prezista (DRV) and ritonavir prior to switch to Rezolsta.
- In approximately 2 years there may be a requirement to switch away from the fixed dose combination when a generic darunavir becomes available.

**Patient factors for Rezolsta:**

- The Rezolsta tablet is larger than other ARVs and may not be acceptable to some. Ensure the patient has seen the tablet prior to leaving clinic.

**Monitoring considerations for both Rezolsta and Evotaz:**

- Consider additional monitoring post switch for those with identified "cautions" or other clinical indications.
- Serum creatinine is expected to increase with a resulting reduction in eGFR of ~10 ml/min. Typically this plateaus after 4 weeks of cobicistat-based ART. If a further change in eGFR is observed, or other renal markers change, this should prompt review.
- Advise GP/primary care physician and other health professionals of the change including interpretation of eGFR and impact on non-ARVs.

### Atazanavir-ritonavir to Evotaz

**Evotaz is NOT RECOMMENDED** if:

- Patient requires ATV 400 mg daily, with or without pharmacokinetic booster.
- Taking hormonal contraceptives, including those containing 30 µg of ethinylestradiol due to potential increase in estrogen exposure. Alternative forms of contraception (non-hormonal) should be considered.
- Taking any of the following, which are not recommend with Evotaz but may be used with ATV+r:
  - ARVs:
    - Efavirenz
    - Etravirine
    - Nevirapine
  - Anti-convulsants:
    - Carbamazepine
    - Phenoobarbital
    - Phenytoin
  - Other:
    - Bosantan
    - OBV/PTV/r ± D5V (AbbVie "2D/3D")**

- There are any ATV resistance associated mutations.
- eGFR <70 ml/min when co-administered with medicines that require dose adjustment based on renal function (e.g. tenofovir DF, lamivudine or emtricitabine).
- Patients aged <18 years of age where the safety and efficacy of Evotaz has not been established.

**Evotaz should be used with CAUTION** when:

- Pregnant, including if actively planning to conceive due to a lack of data.
- Taking any other medicines, particularly those listed in the table. Conduct full medicines review.
- Any combinations other than 2 NRTIs + Evotaz as these have not been studied.

**Specific counselling points for Evotaz:**

- The recommended dosing regimen is one tablet of Evotaz taken once daily with food.
- Use all Reyataz (ATV) and ritonavir prior to switch to Evotaz.
- In approximately 2 years there may be a requirement to switch away from the fixed dose combination when a generic atazanavir becomes available.
### Drug class

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Non-ARV Drug</th>
<th>Potential impact of switch on non-ARV</th>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analgesics</td>
<td>Methadone</td>
<td>Exposure ↑</td>
<td>Consider counselling patient on potential need for reduction in methadone dose. Advise methadone prescriber of potential for dose modification.</td>
</tr>
<tr>
<td></td>
<td>Diamorphine, morphine, hydromorphone, pethidine</td>
<td>Exposure ↑</td>
<td>Consider dose reduction and re-titration and/or monitor for signs of opiate toxicity.</td>
</tr>
<tr>
<td></td>
<td>Dihydromorphone</td>
<td>Unclear, may ↑ or remain unchanged</td>
<td>Counsel patient that may need to re-titrate dose.</td>
</tr>
<tr>
<td>Anti-microbials</td>
<td>Atovapoune, proguanil, sulfadiazine</td>
<td>Exposure ↑</td>
<td>Monitor for side effects.</td>
</tr>
<tr>
<td></td>
<td>Rifabutin</td>
<td>Unclear</td>
<td>Caution or avoid; Consult HIV/TB co-infection guidelines.</td>
</tr>
<tr>
<td>Anticoagulants</td>
<td>Warfarin, acenocoumarol, eltrombopag</td>
<td>Exposure ↑</td>
<td>Liaise with anti-coagulant prescriber prior to switch. For warfarin &amp; acenocoumarol check INR within one week of switch. For eltrombopag monitor impact on platelets.</td>
</tr>
<tr>
<td>Anti-convulsants</td>
<td>Lamotrigine, valproate</td>
<td>Exposure ↑</td>
<td>Consider dose reduction if currently exceeding maximum recommended dose; liaise with prescriber.</td>
</tr>
<tr>
<td></td>
<td>Carbamazepine, phenytoin</td>
<td>(↑PI &amp; ↓Cobicistat)</td>
<td>Not recommended with Rezolsta or Evotaz due to reduced PI and cobicistat exposure.</td>
</tr>
<tr>
<td>Anti-diabetics</td>
<td>Metformin</td>
<td>Exposure ↑</td>
<td>Careful patient monitoring and dose adjustment of metformin is recommended.</td>
</tr>
<tr>
<td></td>
<td>Gliclazide, glimepiride, glipizide, rosiglitazone, tolbutamide</td>
<td>Exposure ↑</td>
<td>Advise patient to check BMs frequently 1-2 weeks post switch and monitor for hypoglycaemia.</td>
</tr>
<tr>
<td></td>
<td>Nateglinide</td>
<td>Unclear, may ↑ or remain unchanged</td>
<td>Advise patient to check BMs frequently 1-2 weeks post switch and monitor for hypoglycaemia</td>
</tr>
<tr>
<td>Anti-hypertensives</td>
<td>Losartan, labetalol, irbesartan, torasemide</td>
<td>Exposure ↑</td>
<td>Counsel patient to monitor for hyotensive episodes and consider dose titration.</td>
</tr>
<tr>
<td>Anti-anginals</td>
<td>Voriconazole</td>
<td>Exposure ↑</td>
<td>Monitor for signs of toxicity such as vomiting, agitation, restlessless, dilated pupils and sinus tachycardia.</td>
</tr>
<tr>
<td>Anti-Parkinson’s agents</td>
<td>Apomorphine, rasagiline, ropinirole</td>
<td>Exposure ↑</td>
<td>Monitor for side effects and consider dose adjustment.</td>
</tr>
<tr>
<td>Beta-Blockers</td>
<td>Carvediol</td>
<td>Unclear, may ↑ or remain unchanged</td>
<td>Counsel patient to monitor for hypotensive episodes and bradycardia and consider dose titration.</td>
</tr>
<tr>
<td>Bronchodilators</td>
<td>Theophylline</td>
<td>Exposure ↑</td>
<td>Consider theophylline baseline (pre-switch) drug level and repeat levels. Monitor for signs of toxicity such as vomiting, agitation, restlessless, dilated pupils and sinus tachycardia.</td>
</tr>
<tr>
<td>Contraceptive &amp; HRT</td>
<td>Estrone-based contraceptives: estradiol, ethinylestradiol, norethisterone</td>
<td>Exposure ↑</td>
<td>Avoid coadministration as no dosing recommendations can be made on the use of Rezolsta or Evotaz (unlike ATV/r) with contraceptives. Alternative forms of (non-hormonal) contraception should be considered.</td>
</tr>
<tr>
<td>Cytotoxics</td>
<td>Anastrozole</td>
<td>Unclear, may ↑ or remain unchanged</td>
<td>Review risk versus benefit of change.</td>
</tr>
<tr>
<td></td>
<td>Dacarbazine, doloxifene, epirubicin, formestane, procarbazine</td>
<td>Exposure ↑</td>
<td>Careful monitoring for signs of toxicity.</td>
</tr>
<tr>
<td>Hepatitis C DAA</td>
<td>Ombitasvir/Paritaprevir/r ± DSV (AbbVie “2D/3D”)†</td>
<td>Exposure ↑</td>
<td>Rezolsta and Evotaz are contraindicated with ombitasvir/paritaprevir/r ± dasabuvir.</td>
</tr>
<tr>
<td>Immuno-suppressants</td>
<td>Mycophenolate</td>
<td>Exposure ↑</td>
<td>Review risk versus benefit of change.</td>
</tr>
<tr>
<td></td>
<td>Tacrolimus, ciclosporin, sirolimus</td>
<td>Unknown</td>
<td>An effect is unlikely, however, close monitoring of the immunosuppressant is recommended.</td>
</tr>
<tr>
<td>Mental health</td>
<td>Olanzapine</td>
<td>Exposure ↑</td>
<td>Consider dose reduction if currently exceeding maximum recommended dose. Write to mental health provider as may require monitoring and dose modification.</td>
</tr>
<tr>
<td></td>
<td>Bupropion, paroxetine, sertraline</td>
<td>Exposure ↑</td>
<td>Counsel patient about potential for side effects.</td>
</tr>
<tr>
<td></td>
<td>Duloxetine</td>
<td>Exposure ↑</td>
<td>Levels may significantly increase compared to PI/r. Write to prescriber and advise of need to monitor and dose reduction of duloxetine, especially if currently exceeding maximal dosing.</td>
</tr>
</tbody>
</table>

† 2D = OBV/PTV/r (Viekirax); 3D = OBV/PTV/r + DSV (Viekirax + Exviera)

There is a lack of data about the magnitude or clinical significance of any changes.