PatientID: HDR91

Okitobba 06, 2023

Color Code

HR: High-Level Resistance
LR: Low-Level Resistance
IR: Intermediate Resistance

S: Susceptible

| DRUG.CLASS | DRUG | RESISTANCE.PROFILE | DRMS.above.20.percent.prevalence |
|------------|------|----------------------|----------------------------------|
| PI | ATV | S | |
| | DRV | ${f S}$ | |
| | FPV | \mathbf{S} | |
| | IDV | \mathbf{S} | |
| | LPV | \mathbf{S} | |
| | NFV | \mathbf{S} | |
| | SQV | \mathbf{S} | |
| | TPV | \mathbf{S} | |
| NRTI | ABC | $^{ m HR}$ | |
| | AZT | ${f S}$ | |
| | D4T | $_{ m HR}$ | |
| | DDI | $_{ m HR}$ | K65R;D67N;L74I;M184V |
| | FTC | $^{ m HR}$ | |
| | LMV | $^{ m HR}$ | |
| | TDF | IR | |
| NNRTI | DOR | PLR | |
| | EFV | $_{ m HR}$ | |
| | ETR | PLR | V106I;K103N |
| | NVP | $_{ m HR}$ | |
| | RPV | PLR | |

Appendix

Drug abbreviations in full

| DRUG.CLASS | ABBREVIATION | DRUG.NAME |
|------------|--------------|----------------|
| | ATV | Atazanavir |
| | DRV | Darunavir |
| | FPV | Fosamprenavir |
| PI | IDV | Indinavir |
| 11 | LPV | Lopinavir |
| | NFV | Nelfinavir |
| | SQV | Saquinavir |
| | TPV | Tipranavir |
| | ABC | Abacavir |
| | AZT | Azidothymidine |
| | DFT | Stavudine |
| NRTI | DDI | Didanosine |
| | FTC | Emtricitabine |
| | LMV | Lamivudine |
| | TDF | Tenofovir |
| | DOR | Doravirine |
| | EFV | Efavirenz |
| NNRTI | ETR | Etravirine |
| | NVP | Nevirapine |
| | RPV | Rilpivirine |
| | BIC | Bictegravir |
| | CAB | Cabotegravir |
| INSTI | DTG | Dolutegravir |
| | EVG | Elvitegravir |
| | RAL | Raltegravir |

Comments

| DRUG.CLASS | COMMENTS |
|------------|---|
| PI | |
| | D67N is a non-polymorphic TAM associated with low-level resistance to AZT. |
| | K65R confers intermediate reductions in susceptibility to TDF, ABC, and 3TC/FTC. It |
| NRTI | increases AZT susceptibility. In NRTI-experienced, INSTI-naive patients with K65R, |
| | TDF+3TC+DTG is usually highly effective and more effective than AZT/3TC/DTG. |
| | However, in patients receiving TDF+3TC+DTG, there is a risk of emergent DTG |
| | resistance that does not arise in NRTI-naive patients receiving TDF+3TC+DTG. |
| | L74V causes intermediate ABC resistance. L74I causes low-level ABC resistance. |
| | M184V/I cause high-level in vitro resistance to 3TC and FTC and low/intermediate |
| | resistance to ABC (3-fold reduced susceptibility). M184V/I are not contraindications to |
| | continued treatment with 3TC or FTC because they increase susceptibility to AZT and |
| | TDF and are associated with clinically significant reductions in HIV-1 replication. |
| NNRTI | K103N is a non-polymorphic mutation that confers high-level reductions in NVP and EFV |
| | susceptibility. It is the most commonly transmitted DRM. |
| | V106I occurs in 1% to 2% of viruses from untreated persons. It contributes to reduced |
| | NNRTI susceptibility only in combination with other NNRTI-resistance mutations. It is |
| | commonly selected in persons receiving DOR in combination with mutations at position |
| | 227. |

| INSTI | |
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