Psychiatric disorders frequently co-occur with HIV, as preceding conditions or consequent to HIV infection. This potentially compromises HIV diagnosis and antiretroviral (ARV) treatment adherence. We provide a brief guide to the diagnosis and treatment of common mental disorders in people living with HIV/AIDS, including: prescribing psychotropics in HIV; neuropsychiatric side-effects of ARVs and other medications commonly prescribed in HIV; and the diagnosis and treatment of depression, anxiety, psychosis, agitation, sleep disturbance, pain, and mania. Psychotropic treatments recommended were drawn primarily from those available in the public sector of South Africa.

Psychotropics in HIV: basic principles

A few simple points should be kept in mind when considering the management of mental disorders in HIV-infected individuals. Firstly, patients with HIV infection are generally very sensitive to medication side-effects as they often metabolise drugs more slowly, have less lean body mass and have compromised blood-brain barrier functioning. Although most patients ultimately tolerate standard doses of most medications, it is advisable to start at low doses and escalate dosing slowly over time.1,2 Furthermore, PLWHAs often receive multiple medications (ARVs, antibiotics, tuberculosis (TB) medications, etc.). Consequently, healthcare providers need to avoid prescribing complex regimens (e.g. daily dose instead of twice daily, where possible), anticipate drug interactions, and consider possible mood, behavioural and cognitive effects of medications such as ARVs.

The fact that treating HIV infection and related conditions is essential for optimal psychiatric care is often under-appreciated. Close collaboration between psychiatrists, physicians, nursing staff and all members of the multi-disciplinary healthcare team is crucial. It is important to make the distinction between primary and secondary psychiatric symptoms (e.g.
optimal HAART regimen for patients with alternative ARVs should be considered. The management of symptoms. Furthermore, where possible, patients should be informed of potential side-effects and closely monitored for any emergence or exacerbation thereof. Although EFV is not absolutely contra-indicated in patients with a history of severe mental illness, patients should be informed of potential side-effects and closely monitored for any emergence or exacerbation of symptoms. Furthermore, where possible, alternative ARVs should be considered. The optimal HAART regimen for patients with severe symptoms, high levels of distress and those caused by other medications, delirium, central nervous system (CNS) infections, etc., as results of standard psychiatric treatment may be inadequate if these are not addressed.\(^1\)

**ARV neuropsychiatric side-effects**

The introduction of HAART has transformed HIV infection from a death sentence to a chronic treatable illness. Unfortunately, most ARVs have neuropsychiatric side-effects (Table 1),\(^1\) most commonly insomnia and headache, with efavirenz (EFV) being the agent most often implicated. These symptoms usually emerge shortly (within 3 months) after commencing ARVs and abate on withdrawal thereof. Although EFV is not absolutely contra-indicated in patients with a history of severe mental illness, patients should be informed of potential side-effects and closely monitored for any emergence or exacerbation of symptoms. Furthermore, where possible, alternative ARVs should be considered. The optimal HAART regimen for patients with CNS disease remains to be established. It is unclear whether ARV regimens with better CNS penetration are superior to others,\(^1\) but there is consensus that optimal peripheral viral suppression is necessary.\(^4\)

**Depression**

Depressive disorder is common in HIV-positive individuals, with a prevalence of 11.1% for major depressive disorder and 29.9% for mild depression in SA clinics.\(^1\) It has been suggested that depression is often under-diagnosed\(^1\) and insufficiently managed.\(^2\) The following screening questions may prove helpful in identifying patients requiring further treatment or referral: 
- ‘During the past month, have you often been bothered by feeling down, depressed or hopeless?’
- ‘During the past month, have you often been bothered by little interest or pleasure in doing things?’
- ‘Is this something with which you would like help?’

It is also important to ask patients about suicidal thoughts, self-esteem, feelings of guilt or worthlessness, and outlook. Although the *Diagnostic and Statistical Manual of Mental Disorders* (Fourth Edition) (DSM-IV) includes problems with sleep, energy and appetite as diagnostic criteria for depression, these may be HIV-related.

If the patient answers ‘yes’ to one or more of the above questions, it is helpful to differentiate between mild-moderate and severe depression, to inform further management. In mild-moderate depression, patients usually experience transient or mild symptoms occasionally, have low levels of distress, and do not have suicidal thoughts or plans. Such symptoms can often be in relation to a recent diagnosis or the commencement of ARV treatment. These patients can be referred to a supportive adherence counsellor or can be considered for referral for psychotherapy where psychologists are available. In cases of severe depression, patients often have persistent, severe symptoms, high levels of distress and

<table>
<thead>
<tr>
<th>Table 1. ARV side-effects</th>
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<tr>
<td><strong>Class</strong></td>
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<td>Nucleoside reverse transcriptase inhibitors (NRTIs)</td>
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<tr>
<td>Non-nucleoside reverse transcriptase inhibitors (NNRTIs)</td>
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<td>Protease inhibitors (PIs)</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Antibacterials</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Other</td>
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\(\text{CNS = central nervous system.}\)

\(^{*}\text{Common: <10%; rare: <1%}.\)
suicidality. These patients should be referred to psychiatric services and/or treatment with antidepressant medication should be initiated.

**Antidepressants**

Using more than one antidepressant should be avoided, as the risk of serotonin syndrome may be increased in HIV-positive patients. The syndrome, which constitutes a medical emergency, presents with pyrexia, sweating, diarrhoea, hyperreflexia, myoclonus, loss of consciousness and seizures.

In general, the duration of treatment with antidepressant medication depends on whether or not the patient has experienced previous depressive episodes. For a first episode, medication should generally be continued for 6 - 12 months to prevent relapse. Treatment should be continued for 2 - 3 years in the event of a patient's second or third episode, and lifelong medication should be considered for >3 prior episodes.

St John's wort, a herbal product with antidepressant effects, may reduce the plasma concentrations and clinical effects of EFV, nevirapine (NVP) and lopinavir/ritonavir (LPV/r). Patients must therefore be informed that its concurrent use with these ARVs is contra-indicated.7

First-line agents include the serotonin selective re-uptake inhibitors (SSRIs), fluoxetine (for patients on first-line ART regimens) or citalopram (for patients on second-line regimens or receiving protease inhibitors (PIs)). In most public sector facilities, treatment with citalopram needs to be initiated by a psychiatrist. The side-effects, drug interactions and potential advantages of these SSRIs are outlined in Table 2.

Table 2. Side-effects, drug interactions and advantages of concurrent SSRI use with ART

<table>
<thead>
<tr>
<th>Agent</th>
<th>Dose</th>
<th>Side-effects</th>
<th>Drug interactions8,9</th>
<th>Notes</th>
</tr>
</thead>
</table>
| Fluoxetine | 20 - 60 mg/day | Nausea, dyspepsia, abdominal pain, anxiety (especially in first 10 days), headache, tremor, sexual dysfunction, hypomania, insomnia and agitation | • LPV/r: may increase fluoxetine levels – increased risk of serotonin syndrome | • Advantages: low cost, available at most centres  
• Agitation can be a big problem in the first few days: adequate explanation and reassurance can reduce impact  
• Safe in overdose |
| Citalopram | 20 - 60 mg/day | As for fluoxetine | • Not a potent inhibitor of most cytochrome-P450 enzymes: few drug interactions  
• Use with caution with NSAIDs/ warfarin | • Advantage over fluoxetine: starting dose can be halved (10 mg), and fewer drug interactions  
• Safe in overdose |

LPV/r = lopinavir/ritonavir; SSRI = serotonin selective re-uptake inhibitor; NSAIDs = non-steroidal anti-inflammatory drugs.

If the patient has a co-morbid sleep disorder or chronic pain, tricyclic antidepressants (TCAs), such as amitriptyline or imipramine, should be considered (Table 3). These can be prescribed as monotherapy: 100 - 150 mg/day, or 25 - 50 mg/day as augmentation.

In patients with co-morbid anxiety and/or patients who have not responded to SSRIs, the use of venlafaxine (Efexor) – a serotonin and noradrenaline re-uptake inhibitor (SNRI) – could be considered. The drug is not available at all centres and needs to be initiated by a psychiatrist (at a dose of 37.5 mg/day, increased to a maximum of 225 mg/day). Treatment with venlafaxine should never be stopped abruptly, as discontinuation symptoms can occur. Potential side-effects include nausea, insomnia, dry mouth, somnolence, sweating, headache, nervousness, constipation, sexual dysfunction and elevation of blood pressure at higher doses. The drug should be avoided in patients at risk of arrhythmia.2

**Anxiety**

Anxiety is a normal human emotion and may be adaptive in many circumstances. However, when it is present for prolonged periods of time, is excessive in relation to the person's current life stressors, or interferes with daily functioning, an anxiety disorder may be present. Anxiety symptoms often mimic common mental conditions and may occur as part of depression, or alone. It is important to exclude and treat physical causes that can resemble the physical symptoms of anxiety, such as thyroid disease, cardiac disease and seizures. The DSM-IV
distinguishes between the following anxiety disorders:

- **generalised anxiety disorder (GAD):** excessive and pervasive worry and tension about a variety of events and activities in daily life, associated with somatic symptoms
- **panic disorder:** recurrent, unexpected sudden attacks of overwhelming anxiety
- **phobias:** excessive fears of specific objects (e.g. spiders) or situations (e.g. flying or social situations)
- **post-traumatic stress disorder (PTSD):** distressing dreams or flashbacks, nervousness, poor sleep and avoiding reminders following a life-threatening or traumatic event
- **obsessive-compulsive disorder (OCD):** repetitive, uncontrollable thoughts or images that are disturbing; or an inability to cease performing rituals or repetitive actions.

If symptoms of the above disorders are present for ≥1 month, where possible, we advise considering pharmacotherapy and referral to a psychologist. SSRIs are considered first-line treatment. Importantly, people with anxiety disorders may be particularly prone to adverse effects and tolerate high initial doses poorly.

### Table 4. Benzodiazepine use with ART

<table>
<thead>
<tr>
<th>Benzodiazepines (short-term prescribing only)</th>
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</table>
| Long-acting (half-life >20 hours) | Diazepam (Pax, Valium): 2 - 30 mg (oral or IV; never IM) up to 3 times daily  
Clonazepam (Rivotril): 0.5 - 2 mg twice daily | |
| Intermediate-acting (12 - 24 hours) | Lorazepam (Ativan, Tranqipam): 1 - 12 mg (oral, IM or sublingual) up to 3 times daily  
Alprazolam (Alzam, Xanor): 0.25 - 4 mg (see interactions below) twice daily | |
| Short-acting (6 - 12 hours) | Oxazepam 10 - 30 mg twice daily | |
| Ultra-short-acting (<6 hours) | Midazolam (Dormicum): 7.5 - 15 mg (see interactions below) usually stat dose, but may be used up to 3 times daily | |

**Benzodiazepine interactions with ARVs**

- **Diazepam**
  - Use with caution with EFV and LPV/r: may need dose adjustment because of increased sedation, confusion and respiratory depression

- **Clonazepam**
  - EFV: possible increase or decrease in clonazepam levels; avoid combination  
  - NVP: possible decrease in clonazepam concentrations and symptoms of withdrawal  
  - RTV: likely to increase levels of clonazepam – use with caution

- **Alprazolam**
  - EFV: may increase levels of alprazolam – avoid  
  - RTV: increases alprazolam effect when RTV is started; after 10 days no significant interaction  
  - NVP: may reduce alprazolam effect

- **Midazolam**
  - Do not co-administer with EFV, indinavir or LPV/r  
  - Use with caution with NVP

- **Lorazepam/oxazepam**
  - No clinically significant interaction expected

### Table 5. Approach to prescribing psychotropics for the agitated patient

<table>
<thead>
<tr>
<th>Step</th>
<th>Agent</th>
<th>Dose</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral (first step after non-pharmacological treatment)</td>
<td>Lorazepam</td>
<td>1 - 2 mg (maximum 12 mg/day)</td>
<td>Repeat after 45 min to a maximum of 12 mg/day (Daily)</td>
</tr>
<tr>
<td></td>
<td>Promethazine</td>
<td>20 - 25 mg</td>
<td></td>
</tr>
<tr>
<td>IM/IV (second step after oral measures have failed/are not possible)</td>
<td>Lorazepam</td>
<td>1 - 4 mg IM</td>
<td>Have flumazenil to hand in case of respiratory depression</td>
</tr>
<tr>
<td></td>
<td>Haloperidol</td>
<td>5 mg IM</td>
<td>Should be the last drug considered as incidence of acute dystonia is high</td>
</tr>
<tr>
<td></td>
<td>Diazepam</td>
<td>10 mg over 10 min IV (never IM)</td>
<td>Repeat after 10 min if insufficient effect (up to 3 times)</td>
</tr>
</tbody>
</table>

IVI = intravenous; IM = intramuscular.

IV = intravenous; IM = intramuscular.
Either fluoxetine (for patients on first-line ART regimens) or citalopram (for patients on second-line regimens or receiving PIs) could be commenced at half the usual starting dose. Citalopram needs to be psychiatrist-initiated, but has the advantage over fluoxetine of being a scored tablet, making it easier to start at half the usual dose (10 mg). Venlafaxine (psychiatrist-initiated) should be considered for patients who have not responded to SSRIs.

Benzodiazepines (Table 4) provide rapid symptomatic relief of anxiety, but because of their potential to cause physical dependence and withdrawal symptoms,11 and the potential for abuse, these drugs should be used at the lowest effective dose for the shortest period of time (<3 weeks). Benzodiazepines should be used in conjunction with SSRIs during treatment initiation. Caution should be exercised because of serious interactions with ARVs, particularly ritonavir (RTV), and especially with alprazolam, midazolam and triazolam. Lorazepam and oxazepam have the least number of interactions with ARVs. Benzodiazepines also cause or exacerbate cognitive impairment and are sedating; therefore, patients must be advised not to drive, operate machinery or drink alcohol concurrently with their use.

**Psychosis**

Psychosis can occur at any time during the course of HIV disease. A psychotic syndrome includes at least 2 of the following symptoms:

- delusions (fixed false beliefs)
- hallucinations (auditory and other)
- disorganised speech or thought
- disorganised behaviour.

Psychotic disorders include schizophrenia, substance-induced psychosis, and psychosis secondary to a general medical condition such as HIV. Reported rates of new-onset psychosis in HIV-positive patients range from 0.5% to 15%.12 It is essential to differentiate psychotic symptoms caused by delirium or encephalopathy, to identify and treat the underlying cause; although short-term symptomatic treatment may include low-dose antipsychotics. In delirium, the psychotic symptoms may occur in the context of fluctuating attention, sleep/wake disturbance and poor orientation.

**Anti-psychotics**

Importantly, with regard to prescribing antipsychotics, HIV-positive patients may be more susceptible to extra-pyramidal side-effects (EPSEs), neuroleptic malignant syndrome and tardive dyskinesia. Antipsychotics should always be initiated at the lowest effective dose and for the shortest period of time necessary. Atypical or second-generation antipsychotics (SGAs), where available, are generally preferred over first-generation antipsychotics, because of the decreased risk of EPSEs. Risperidone is the most widely studied atypical antipsychotic (or SGA), and generally appears to be safe, although levels have been reported to increase with concurrent RTV use.11 An overlap in metabolic side-effect profiles, e.g. weight gain, dyslipidaemia and impaired glucose tolerance of the SGAs and ARVs (PIs and nucleoside reverse transcriptase inhibitors (NRTIs) in particular), complicates the risk-to-benefit equation.14,15 Beside the interactions between risperidone and RTV, antipsychotics do not generally significantly inhibit or induce P-450 enzymes and can safely be added to HAART regimens without causing toxicity or HAART failure. Theoretically, RTV may increase the serum levels of haloperidol; therefore, close monitoring of adverse effects is advised.

The use of clozapine in HIV-positive patients is not routinely recommended, although it may be helpful in otherwise medically stable patients with higher CD4 cell counts.13 It is not

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**Fig. 1. Decision tree for initiating appropriate anti-psychotic treatment for psychosis in HIV-infected individuals. EPS = extra-pyramidal side-effects; WCC = white cell count; LFTs = liver function tests.**

*Risperidone and quetiapine may be initiated ONLY by a psychiatrist.

†Depot medications are not contra-indicated. Consider a zuclopenthixol depot if clinically indicated; use a test dose of 50 - 100 mg, and repeat in 1 week. Thereafter, the depot can be administered monthly.
known whether HIV-positive patients receiving clozapine have a greater risk of agranulocytosis; therefore, extremely close monitoring of white cell count is recommended. Clozapine should only be initiated by a psychiatrist. The drug should be used with caution with LPV and RTV;\(^8,9\) as this could increase clozapine plasma concentrations, resulting in an increased risk of arrhythmias, seizures and haematological effects.

A guide to initiating appropriate treatment is provided in Fig. 1. Benzodiazepines can be added in the initial stages for agitation or aggressive or disruptive behaviour. Following diagnosis, a depot preparation can be given if adherence is likely to be a problem (or if the patient chooses it). A test dose should be considered.

Agitation

Aggressive or disruptive behaviour can occur in the context of psychiatric illness, physical illness, substance abuse or personality disorder. This can put staff and patients at risk. In general, it is important to attempt to examine the patient as thoroughly as possible before sedation to establish the underlying aggravating factors. It is also crucial to gain help from nursing staff, doctors and security staff where possible. The safety of the clinician, other patients and staff must be ensured. Non-pharmacological methods should be attempted first, e.g. talking down, distracting, and reassuring the patient. The patient must not be threatened, as this often escalates the situation. Oral treatment must be offered first and biperiden must be available if neuroleptics are introduced. Drug interactions between benzodiazepines and ARVs, and the approach to prescribing psychotropics in the agitated patient are summarised in Tables 4 and 5, respectively.

Sleep disturbance

A patient with insomnia may have difficulty with falling asleep, early-morning wakening and/or frequent waking during the night. Before treating insomnia with drugs (Table 6), it is important to consider and address underlying reversible causes such as depression, mania, pain, medication side-effects, substance abuse, and poor sleep hygiene. It must be ascertained whether the patient has realistic expectations of sleep and whether other medications are being given at appropriate times, e.g. stimulating drugs in the morning and sedating drugs at night. If medication is prescribed, then the lowest effective dose for short-term use only must be used, and patients must be advised of interactions with alcohol.

Table 6. Available classes of psychotropics for insomnia\(^*\)

<table>
<thead>
<tr>
<th>Agent</th>
<th>Dose</th>
<th>Side-effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Promethazine</td>
<td>10 - 25 mg</td>
<td>Can cause dry mouth and ‘hangover’ effect</td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>10 - 25 mg</td>
<td>Useful in patients with peripheral neuropathy</td>
</tr>
<tr>
<td>Oxazepam</td>
<td>15 - 30 mg</td>
<td>Risk of dependency; prescribe no more than 14 days’ supply, unless prescribed by a psychiatrist or neurologist</td>
</tr>
</tbody>
</table>

*Refer to the text for ARV drug interactions.

Pain

Pain symptoms are common in HIV infection and may be caused by painful neuropathy, headaches, cancers and secondary infections. Pain disorders may be acute or chronic, with the latter often accompanied by depression, anxiety, and/or sleep disorders. In addition to analgesia use, psychotropics are frequently used to ameliorate pain symptoms:
- TCAs (e.g. amitriptyline at doses 25 - 75 mg at night): refer to the section on ‘Depression’ for further information. Other antidepressants should be considered if co-morbid depression is present. Duloxetine is an antidepressant registered for chronic pain but is not on State code in SA.
- Anti-convulsants: carbamazepine is not advised because of interactions with ARVs.;\(^*\) Gabapentin (Lyrica) is used in chronic pain, but is not freely available.

Bipolar affective disorder/mania

The essential characteristic of a bipolar mood disorder is one or more manic (or hypomanic) episodes with/without depressive episodes. Mania is a recognised presentation in HIV-infected individuals. A manic episode, which is severe enough to impair functioning or warrant hospitalisation, is characterised by abnormal and persistently elevated, expansive or irritable mood, with: grandiosity; decreased need for sleep; talkativeness; flight of ideas/accelerated thoughts; distractibility; and/or increased involvement in pleasurable activities with potentially negative consequences, e.g. excessive buying or sexual indiscretions.

Mood stabilisation

Management is described in Fig. 2. HIV-positive individuals may be more sensitive to the side-
effects of mood-stabilisers, especially in the case of neurocognitive dysfunction. Agents such as valproate, lamotrigine and lithium may be used cautiously, but carbamazepine (Tegevetol) should be avoided because of potential interactions with ARVs (RTV, NVP and EFV), and the risk of neutropenia.2,6,9

Valproate (Epilim) is generally considered the first-line treatment, but there is an additive risk of fatty liver with didanosine (ddI), abacavir (ABC), lamivudine (3TC), stavudine (d4T) and zidovudine (AZT). It is important to monitor the patient’s liver function and adjust the dose accordingly, and it is advisable to test pre-treatment hepatic transaminases (AST/ALT) and platelet levels. The potential for teratogenesis in women of child-bearing age remains a concern. Some drug interactions do occur with LPV/r and LPV. Valproate levels may be decreased with co-administration of RTV. An increase in the dose of valproate may be required. No significant interaction occurs with tenofovir (TDF), NVP or EFV.6,9

Lithium should be avoided in patients with dehydration and renal function impairment. The agent is not always well tolerated, and it may be advisable to limit its use to individuals with higher CD4 cell counts.6 Careful monitoring of lithium levels is needed, usually 5 days after any dose adjustment, then monthly and 3-6 monthly thereafter.

Lamotrigine (Lamictin) can also be considered and is mainly used for depression in bipolar disorder. In most State facilities its use needs to be initiated by a psychiatrist or neurologist and the dose needs to be increased gradually to avoid Stevens-Johnson syndrome. RTV decreases lamotrigine levels by about 50% due to induction of glucuronidation; therefore, an increased lamotrigine dosage may be required.

Additional treatments that can be used in manic episodes include antipsychotics (such as Risperidone and Quetiapine) and benzodiazepines.

**Conclusion**

Many patients with HIV/AIDS have co-occurring mental health conditions that affect ART adherence, quality of life, morbidity and mortality. Although close collaboration between physicians, psychiatrists and other members of the multidisciplinary healthcare team is the ideal, many clinicians work in settings where psychiatric support is not readily available. This article is intended to guide prescribing antipsychotics in these settings.

**Additional resources**

- Medicines Information Centre: http://www.mic.uct.ac.za; tel: +27 (0)21 406 6829
- HIV drug interactions: http://www.hiv-druginteractions.org
- Psychiatry services and resources in the Western Cape province: http://www.hivmentalhealth.co.za.

**References**