Let’s Talk TB:
A Supplement to GP CLINICS
Adverse Drug events With Anti Tuberculosis Therapy
What Every GP Should Know

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Standards of TB Care in India

• Pulmonary TB patients treatment:
  – Intensive Phase, First Line: Isoniazid (INH), Rifampicin (RIF), Pyrazinamide (PZA) and Ethambutol (EMB)
    • First 2 months
  – Continuation Phase: INH, RIF and EMB
    • Next 4 months

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Standards of TB Care in India

• Course of treatment can be complicated by occurrence of adverse events necessitating treatment interruptions and modified regimens

• Lack of adherence to drugs and treatment interruptions are the driving factors for emergence of drug resistance during the treatment

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Adverse drug events (ADE) during anti-tuberculosis therapy (ATT)

• 7% of all drug related adverse events
• 30% of fulminant hepatitis
First Line ATT

• This review is limited to the common adverse drug events on first line ATT
  – Extensively used by general practitioners
• Patients requiring second line drugs are best managed by pulmonary or infectious diseases physicians
  – Adverse events during second-line therapy are not discussed in this review

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# Common Adverse Events (First Line ATT)

## Table 1. Common Adverse Events with First Line ATT Drugs

<table>
<thead>
<tr>
<th>Drug</th>
<th>Common adverse events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid (INH)</td>
<td>Asymptomatic transient elevation of transaminases (20%), hepatitis, peripheral neuropathy, fever, skin rash, seizures, psychosis</td>
</tr>
<tr>
<td>Rifampicin (RIF)</td>
<td>Reddish orange color of urine and tears, Pruritus, GI intolerance, Isolated hyperbilirubinemia, hepatitis, pancytopenia, flu like syndrome, acute kidney injury</td>
</tr>
<tr>
<td>Pyrazinamide (PZA)</td>
<td>Nausea, vomiting, hepatitis, arthralgia, hyperuricemia, skin rash</td>
</tr>
<tr>
<td>Ethambutol (EMB)</td>
<td>Optic neuritis (1-5%), peripheral neuropathy, skin rash</td>
</tr>
<tr>
<td>Streptomycin (SM)</td>
<td>Ototoxicity, nephrotoxicity, skin rash</td>
</tr>
</tbody>
</table>

Drug Induced Liver Injury

- Drug induced liver injury is the most severe of the ADE
- Occurrence of ATT induced hepatitis is estimated to be 5-33%
- Depending on the definitions used to diagnose hepatitis
- Most widely used definition of drug induced hepatitis is serum aminotransferase level
  - >5 times the upper limit of normal [ULN] without symptoms
  OR
  - >3 times the ULN with symptoms of hepatotoxicity (nausea, vomiting, or pain abdomen)
Drug Induced Liver Injury

• ATT may be associated with asymptomatic elevation of transaminases in about 20% of patients

• It may also result in acute hepatitis, even sub-acute to fulminant hepatitis
  – May be fatal

• The frequency of hepatitis
  – With rifampicin is 0.6-2.7%
  – With Isoniazid is 0.6%

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First Line ATT Drugs

- PZA is the most hepatotoxic
  - 15% of patients experiencing hepatic adverse events when higher dose of PZA is used
- The risk of hepatitis is more with combination of INH with RIF or PZA

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Types of Hepatotoxicity and Clinical Features

• Hepatic adaptation
• Drug induced acute hepatitis
• Granulomatous hepatitis
• Cholestasis

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Hepatic adaptation

- Transient asymptomatic elevation of alanine aminotransferase (ALT) may be seen as a physiological response to drug exposure (Cytochrome P450 enzyme induction)
Drug induced acute hepatitis

• Associated with hepatocyte necrosis and elevation of hepatic transaminases with or without jaundice
• Patients may be asymptomatic or may present with nausea, vomiting, abdominal pain and jaundice
• Occasionally they may report constitutional symptoms including fever
• INH and Rifampicin can produce hepatotoxicity by this mechanism

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Granulomatous hepatitis

• e.g. Pyrazinamide can produce this and it is a hypersensitivity reaction to the inciting drug with granuloma formation

• Patients may present with fever, lethargy, body ache, rash, lymphadenopathy and hepatosplenomegaly

• Biochemical examination will reveal elevation of serum transaminases along with alkaline phosphatase
Cholestasis

• Usually associated with asymptomatic elevation of alkaline phosphatase along with bilirubin because of failure of bilirubin transport
  – e.g. seen with rifampicin
Risk Factors for Hepatotoxicity

- Age >35 years
- Children
- Female gender
- Recent child birth (<3 months post-partum)
- Alcohol abuse
- Abnormal baseline liver enzymes
- Slow acetylator status- associated with INH associated hepatotoxicity
- Malnutrition/hypoalbuminemia

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Recognition of Drug Induced Hepatitis
Recognition of Drug Induced Hepatitis

- It is important for doctors to suspect drug induced liver injury when patients on ATT present with nausea, vomiting, right hypochondriac pain, and jaundice

- Temporal pattern of disease evolution after exposure to drugs is important to diagnose drug induced adverse events
Recognition of Drug Induced Hepatitis

• Liver function tests must be requested on suspecting liver injury
  – Drug induced hepatitis is diagnosed when serum amino-transferase level >5 times the ULN without symptoms or >3 times the ULN with symptoms

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Recognition of Drug Induced Hepatitis

- Bilirubin may or may not be elevated
- Early detection of drug induced liver disease is critical to prevent severe or chronic liver disease
- Patients should be advised to report any untoward symptoms:
  - nausea, malaise, lethargy, right hypochondriac pain or new onset fever

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Management of Patients with Drug Induced Hepatotoxicity
Management of Patients with Drug Induced Hepatotoxicity

• Once hepatotoxicity is suspected, all hepatotoxic drugs must be stopped and promptly investigated as mentioned
  – Failure to discontinue a drug that is causing liver injury leads to poor outcome such as acute liver failure or chronic hepatitis
Management of Patients with Drug Induced Hepatotoxicity

• The specific risk factors for drug induced hepatotoxicity have to be carefully elicited

• In all patients with liver abnormality, history of hazardous intake of alcohol, other hepatotoxic drug ingestion must be enquired about and viral hepatitis must be ruled out
Management of Patients with Drug Induced Hepatotoxicity

• No hepatoprotective agent has been effective in ameliorating drug induced liver damage

• Non hepatotoxic ATT drugs which could be used are streptomycin, ethambutol and levofloxacin or moxifloxacin

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Management of Patients with Drug Induced Hepatotoxicity

• Patient should be observed for progress and the liver function tests [LFT] should be monitored once in 3 days

• Usually symptoms and laboratory abnormalities promptly improve within days or weeks once the inciting drugs are stopped

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Management of Patients with Drug Induced Hepatotoxicity

- When the ALT returns to less than 2 times the ULN, gradually drugs are reintroduced sequentially with rifampicin, INH and PZA in that order with a gap of 3-7 days between each drug and monitoring of LFT
  - If symptoms recur or ALT increases, the last drug added should be stopped

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Management of Patients with Drug Induced Hepatotoxicity

• In those patients who had experienced severe or prolonged hepatotoxicity, reintroduction of PZA may be avoided and the duration of ATT may be extended to 9 months

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Gastrointestinal Adverse Effects
Gastrointestinal Adverse Effects

• GI adverse events are usually minor
• Nausea, vomiting and abdominal discomfort, which may be self-limiting
• This may be due to mild gastritis and can be managed by the addition of proton pump inhibitors, anti-emetics, administering drugs after meals or by giving drugs at an interval
Gastrointestinal Adverse Effects

• Discontinuation of ATT is usually not required
• In all such patients with the above mentioned symptoms, LFT must be requested to rule out early liver toxicity
Dermatological Adverse Effects
Dermatological Adverse Effects

• Skin related ADE can occur with all anti TB drugs and is one of the common side effects in up to 6% of patients on ATT

• It can be in varying forms like maculopapular rash, erythema multiforme syndrome, acneiform eruptions, urticarial, lichenoid eruptions, the more severe exfoliative dermatitis and Steven Johnson Syndrome

• Exfoliative dermatitis also called erythroderma, a form of cutaneous hypersensitivity occurs after 6-8 weeks of therapy

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Dermatological Adverse Effects

• The propensity of the drugs to cause erythrodema is PZA >SM >EMB >RIF >INH with rates for PZA being 2.4%

• Steven Johnson syndrome is a life threatening mucocutaneous syndrome and it has been described with RIF, INH, PZA and thiacetazone

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Dermatological Adverse Effects

• In patients with itching/ minimal rash, ATT can be continued along with symptomatic treatment with anti-histamines

• If visible maculopapular rash, all the drugs have to be stopped and antihistamines have to be prescribed
  – Re-introduction of the ATT is to be done with INH followed by RIF, EMB and PZA with careful monitoring

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Neurological Adverse Effects
Neurological Adverse Effects

- CNS adverse events include seizures, psychosis, ototoxicity, and optic neuritis
- The drug most commonly implicated in psychosis is INH followed by cycloserine
- Seizures are a side effect of INH, fluoroquinolones, and cycloserine

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Neurological Adverse Effects

- Ototoxicity and neuromuscular blockade are adverse events of aminoglycosides, while ethambutol is known to cause optic neuritis.
- Ototoxicity and optic neuritis are dose dependent

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Neurological Adverse Effects

• It is important to ask patients about tinnitus, imbalance, vertigo or decreased hearing while the patients are on aminoglycosides.

• As children may not be able to complain about decreased vision, or impairment of color perception (which is a classical symptoms of optic neuritis), ethambutol is best avoided in children.

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Neurological Adverse Effects

• Peripheral neuropathy, manifested by tingling and numbness in hands and feet is a side effect of INH (seen in up to 2% of patients)
  – Less commonly ethambutol and ethionamide

• The risk factors for peripheral neuropathy are malnutrition, HIV infection, diabetes mellitus, alcohol abuse and concomitant drugs having neuropathy potential such as Stavudine

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Neurological Adverse Effects

• It is best to prevent peripheral neuropathy by pyridoxine supplementation (40mg daily) to all patients with the neuropathy risk factors receiving INH.

• The involved drugs have to be stopped promptly on the first suspicion of seizures, psychosis, ototoxicity or visual toxicity and never administered again.

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Musculoskeletal Adverse Effects
Musculoskeletal Adverse Effects

• Mild arthralgia may be associated with PZA, and is associated with elevated uric acid
  – It can be managed symptomatically with NSAIDS

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Renal Adverse Events
Renal Adverse Events

• Acute kidney injury (acute renal failure) with normal urine output can occur with aminoglycosides
• Rarely, rifampicin can give rise to interstitial nephritis

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Drug Induced Haematological Adverse Events

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Drug Induced Haematological Adverse Events

• Immune mediated cytopenia of all three cell lines, including agranulocytosis have been described with ATT

• Thrombocytopenia is an uncommon but potentially life-threatening complication and is characterized by rapid destruction of platelets in susceptible patients due to immune mechanisms

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Drug Induced Haematological Adverse Events

• Patients usually present with purpuric rashes all over the body
  – Reported classically with RIF
  – PZA, EMB and INH are also implicated in various case reports

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Drug Induced Haematological Adverse Events

- The diagnosis is by resolution of symptoms and improvement of platelet count on stopping the drugs.
- Treatment is by discontinuation of the drugs, transfusion of platelets if platelet count is $< 20,000/\mu L$
- Patient has to be managed with 3 new anti tuberculosis drugs including one parenteral agent.

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Drug Induced Haematological Adverse Events

• Rifampicin is also implicated with life threatening drug induced hemolytic anemia due to immune mechanisms

• It can manifest with severe anemia requiring blood transfusion and jaundice due to indirect hyperbilirubinemia

• Re-challenge with the offending drug is contraindicated both in drug induced thrombocytopenia and hemolysis as minute quantity of the drug can trigger a very severe reaction

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Monitoring of Patients for ADE on ATT

• All patients should undergo baseline testing of serum bilirubin, transaminases, alkaline phosphatase, creatinine, and complete blood count

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Monitoring of Patients for ADE on ATT

• **In those with risk factors for hepatotoxicity:** every 2-4 weekly monitoring of liver function tests is recommended

• **In those without risk factors for hepatotoxicity:** routine LFT monitoring is not required unless symptomatic

• During each visit, patients should be asked about symptoms of possible ADE relevant to the regimen they are receiving

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CLINICAL HIGHLIGHTS

• The course of treatment of TB can be complicated by drug induced adverse events and GP’s knowledge about their occurrences and competency to diagnose them are critical for the successful outcome.

• Minor adverse reactions to TB medications include nausea, gastric intolerance, neuropathy, itching and joint pains that can be managed symptomatically and do not warrant stoppage of medications.

• Major adverse reactions include hepatitis, severe skin reactions, optic neuropathy thrombocytopenia, hemolysis, and renal failure. Symptoms and signs suggesting a major reaction should be treated by stoppage of drugs, and close monitoring.

• Drug induced liver injury is the most common serious adverse event which may present as nausea, vomiting, jaundice, right hypochondriac pain and requires stoppage of all hepatotoxic drugs with monitoring of liver functions and subsequent reintroduction of drugs sequentially.

• Patients with pre-existing liver disease, viral hepatitis, chronic alcohol consumption, pregnant women, or postpartum period, HIV infection and on concomitant hepatotoxic medications are particularly at high risk of ATT induced hepatitis and these risk factors must be carefully sought prior to the initiation of ATT and are to be monitored with LFT every 2-4 weekly on therapy.
• On reintroduction of ATT, it is important to avoid mono therapy to prevent emergence of drug resistant TB. Non hepatotoxic drugs like streptomycin, ethambutol and moxifloxacin can be given, when hepatotoxic drugs need to be stopped.

• Life threatening complications like immune thrombocytopenia, immune hemolytic anemia, Steven Johnson syndrome, and organ threatening complications such as ototoxicity, nephrotoxicity, and ocular toxicity are absolute contraindications to rechallenge with the suspected drug.

• All patients need to be counselled on how to identify ADEs, before initiation of ATT.
References:


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