COMPOSITION
Kop® 50 Tablet: Each enteric coated tablet of Kop® 50 contains ketoprofen BP 50 mg.
Kop® 100 Tablet: Each enteric coated tablet of Kop® 100 contains ketoprofen BP 100 mg.
Kop® IM Injection: Each 2 ml Kop® Injection contains Ketoprofen BP 100 mg.
Kop® 2.5% Gel: Each 100 gm gel contains 2.50 gm Ketoprofen BP.
Kop® 100 SR Capsule: Each sustained release capsule contains 100 mg Ketoprofen BP.
Kop® 200 SR Capsule: Each sustained release capsule contains 200 mg Ketoprofen BP.

PHARMACOLOGY
Ketoprofen (Kop®) is a non-steroidal anti-inflammatory drug (NSAID) with analgesic and antipyretic actions. In addition to the inhibition of prostaglandin synthesis, it stabilizes lysosomal membranes in vitro and in vivo, inhibits leukotriene synthesis in vitro at high concentrations, and also exhibits antibradykinin activity in vivo. Ketoprofen (Kop®) produces analgesia by inhibiting the synthesis of prostaglandins peripherally and centrally. It has also been suggested that Ketoprofen (Kop®) causes the suppression of prostaglandin synthesis in the CNS (probably in the hypothalamus) leading to its antipyretic effect. Ketoprofen (Kop®) is rapidly and almost completely absorbed from the GI tract. It is approximately 99% bound to plasma protein, mainly albumin. Following single or multiple oral doses in healthy adults, the elimination half-life of the drug has averaged 1.1–4 hours. It is rapidly and extensively metabolized in the liver, principally via conjugation with glucuronic acid. Following a single oral dose of Ketoprofen in healthy adults, about 50–90% of the drug is excreted in urine and about 1–8% in faeces within 1–5 days; most urinary excretion occurs within 12–24 hours and most faecal excretion occurs within 24–48 hours. In case of IM injection, peak concentration of approximately 10mg/L is reached at about 0.5–0.75 hour after a 100 mg dose. The elimination half-life is approximately 1.88 hour.

INDICATION
Kop® is indicated in musculoskeletal and joint disorders such as ankylosing spondylitis, osteoarthritis, and rheumatoid arthritis, in periarticular disorders.
such as bursitis and tendinitis, in mild to moderate pain such as
dysmenorrhoea or postoperative pain, and in other painful and inflammatory
conditions such as acute gout or soft-tissue disorders.
Ketoprofen is used for symptomatic relief of mild to moderate pain, such as
postoperative (including that associated with dental surgery) postpartum
and orthopedic (including musculoskeletal strains or sprains) pain and
visceral pain associated with cancer.
Kop® IM injection is indicated for management of acute exacerbations of:
Rheumatoid arthritis, osteoarthritis, ankylosing spondylitis, periarticular
conditions such as fibrositis, bursitis, capsulitis, tendinitis, and tenosynovitis,
low back pain of musculoskeletal origin, sciatica, other musculoskeletal
conditions, acute gout and control of pain and inflammation following
orthopaedic surgery.
Kop® 2.5% Gel is an anti-inflammatory and analgesic preparation to be
applied topically to the painful area. It is indicated as a short-term treatment
for traumatic lesions (sprains, tendinitis, edema, bruises) and pain.

**DOSAGE AND ADMINISTRATION**
Kop® Tablet: Oral treatment with Ketoprofen is 50-100 mg daily, taken with
food to minimize gastrointestinal disturbance. For rheumatic disease, 100-
200 mg daily in 2-4 divided doses with food. For pain and dysmenorrhoea,
50 mg up to 3 times daily.
Kop® SR Capsule: 100-200 mg once daily, depending on patient’s weight
and on severity of symptoms
Kop® IM injection: Adults: 50 to 100 mg every four hours, repeated up to a
maximum of 200 mg in twenty-four hours for up to 3 days. Following a
satisfactory response oral therapy should be instituted with Kop® tablet. It is
recommended that the injection should not normally be continued for
longer than three days.
Elderly: As with other medications it is generally advisable in the elderly to
begin ketoprofen therapy at the lower end of the dose range and to maintain
such patients on the lowest effective dosage.
Paediatric dosage: Not established for intramuscular use.
It is not used as intravenous injection.
Kop® 2.5% Gel: Kop® 2.5% Gel is to be applied to the painful area twice daily.
The gel can be used with an occlusive dressing. Rub gently into the skin to ensure penetration.

For children: There is no contraindication to the use of this drug in children, but the appropriate dosage has not yet been established.

Dosage of Ketoprofen must be carefully adjusted according to individual requirements and response, using the lower possible effective dosage.

Dosage exceeding 300 mg daily has not been adequately studied and are not recommended.

ADVERSE EFFECT

Adverse reactions to Ketoprofen are usually mild and mainly involve the GI tract, particularly upper GI tract. Most Ketoprofen-induced adverse effects occur during the first month of treatment, and the frequency of adverse effects generally decreases with continued therapy.

Adverse reactions involving digestive system are dyspepsia, nausea, abdominal pain, diarrhoea, constipation, flatulence, anorexia, vomiting, stomatitis and that involving nervous system are headache, dizziness, malaise, depression, nervousness, dreams, etc. Other reactions are tinnitus, visual disturbance, rash, impairment of renal function, signs or symptoms of urinary-tract irritation.

CONTRAINDICATION AND PRECAUTION

Ketoprofen is contraindicated in patients with known hypersensitivity to the drug. Ketoprofen is contraindicated in patients in whom asthma, urticaria, or other sensitivity reaction is precipitated by aspirins or other NSAIDs, since severe, rarely fatal, anaphylactic reactions to Ketoprofen have been reported in these patients.

The risk of potentially serious adverse GI effects should be considered in patients receiving Ketoprofen, particularly in patients receiving chronic therapy with the drug. Ketoprofen should be used in patients with GI bleeding or active peptic ulceration only when the potential benefits justify the possible risks.

Ketoprofen should be used with caution in patients who may be adversely affected by a prolongation of bleeding time (e.g. patients receiving anticoagulant therapy), since the drug may inhibit platelet function.

Ketoprofen should be used with caution in patients with heart failure, hypertension, or other conditions associated with fluid retention, since
peripheral edema has been observed in some patients receiving the drug. Liver function should be monitored periodically during long-term Ketoprofen therapy.

Ketoprofen injection must not be given intravenously.

Ketoprofen gel should not be applied to patients who have allergy to ketoprofen, other anti-inflammatory agents and aspirin.

**DRUG INTERACTION**
As Ketoprofen may cause GI bleeding, inhibit platelet aggregation and prolong bleeding time, the drug should be used with caution and the patient should be carefully observed if the drug is used concomitantly with any anticoagulant or thrombolytic agent. Concomitant administration of Ketoprofen and hydrochlorothiazide has resulted in decreased urinary excretion of potassium and chloride compared with hydrochlorothiazide alone. Ketoprofen and salicylates appear to interact in a complex manner and they should not be used concomitantly. Concomitant use of Ketoprofen and probenecid is also not recommended. Ketoprofen should be avoided in patients receiving methotrexate.

**USE IN PREGNANCY AND LACTATION**
Embryopathic effects have not been recorded with Ketoprofen, but it is recommended to avoid medication during pregnancy. Trace amounts of the drug appear in breast milk and it should not be used during breast feeding unless unavoidable.

**STORAGE**
Store at cool and dry place, protect from light and moisture. Keep out of the children’s reach.

**HOW SUPPLIED**
- Kop® 50 tablet : Box containing 5x10 tablets in blister packs.
- Kop® 100 tablet : Box containing 5x10 tablets in blister packs.
- Kop® IM injection : Box containing 2x5 ampoules injection.
- Kop® 2.5% Gel : Each tube contains 20 gm gel.
- Kop® 100 SR Capsule : Box containing 5x10 capsules in blister packs.
- Kop® 200 SR Capsule : Box containing 3x10 capsules in blister packs.