



## Leflunomide (Arava) GP Information Sheet

### Leflunomide

Leflunomide is an immunomodulatory agent which arrests activated lymphocytes thought to be involved in inflammatory arthritis pathogenesis. The active metabolite has a long half-life usually 1-4 weeks.

### Pre Treatment Assessment

FBC, LFT, U&E and blood pressure

### Administration:

Oral - the tablets should be swallowed whole with plenty of water. Absorption is not affected by the food.

### A typical dose regime is:

10mg or 20mg daily depending on disease severity and patient tolerance.

### Precautions and Contraindications

Leflunomide may inhibit the metabolism of **warfarin** and **phenytoin**. It has an extremely long elimination half-life and interaction with these drugs and with other DMARDs may occur even after Leflunomide has been discontinued. **Therefore patients on warfarin and Leflunomide will require careful monitoring of INR levels.**

If a severe and undesirable side-effect occurs a washout procedure is available that will rapidly remove its active metabolite. This involves administration of cholestyramine 8g three times a day or activated charcoal 50g five times a day.

**Female patients should not procreate within two years of discontinuing Leflunomide.** Effective contraception is essential in women during treatment and 2 years post treatment. Blood concentrations of its active metabolite should be measured 2 years after discontinuation before pregnancy occurs. The 2 years waiting period can be reduced to 3 months with the wash out procedure.

**Leflunomide is contraindicated in patients with severe liver impairment (ALT>2 x ULN), moderate to severe renal failure, severe immunodeficiency states for example AIDS, severe hypoproteinemia, and nephrotic syndrome.**

### Time to Response

Begins after 6-8 weeks, but improvements may continue for 4-6 months.

### Monitoring

FBC, LFT and BP check monthly for 3 months then every 3 months thereafter

### Actions to be taken

Neutrophils $<2.0 \times 10^9/l$	Hold drug and repeat in 2 weeks
Platelets $<150 \times 10^9/l$	
$>2$ but $<3$ fold rise in ALT or AST (from upper limit of reference range)	Reduce dose and monitor
$>3$ fold rise	Hold Leflunomide and repeat LFT within 2-4 weeks

**Please note that in addition to absolute values for haematological indices, a rapid fall or a consistent downward trend in any value should prompt caution and extra vigilance.**

### Side Effects

Eczema, dry skin, itching, urticaria, oral ulceration and alopecia, (Diffuse hair loss may occur in about 10% of patients, usually reversible on dose reduction or discontinuation)

**NB:** In case of ulceration stomatitis, stop treatment. If Stevens-Johnson syndrome or toxic epidermal necrolysis occur treatment should be stopped. A complete washout is essential in such cases.

**Haematological:** Leucopenia, anaemia mild thrombocytopenia, eosinophilia and rarely agranulocytosis.

**Gastrointestinal:** Nausea, vomiting, anorexia, abdominal pain, taste disturbance and diarrhoea (usually self-limiting)

**Hepatic:** Hepatotoxicity, pancreatitis, severe liver dysfunction rare but small LFT elevation more common. **Patients should be advised that alcohol consumption should be avoided, or kept to a minimum.**

**Nervous System:** Headaches, dizziness, asthenia, paraesthesia, insomnia, migraine, vertigo and anxiety.

**Musculo-Skeletal System:** Tenosynovitis, tendon rupture.

**Cardiovascular:** Hypertension may occur in about 10% of patients. Pre-existing hypertension predisposes.

**Allergic Reactions:** Mild allergic reactions may occur (rash, pruritis urticaria). Anaphylaxis rare.

**Infection:** Severe infection may necessitate stopping the drug and administering a washout. Patients with previous TB need careful monitoring as there is increased risk of reactivation.

**Vaccinations:** No clinical data is available on the efficacy and safety of vaccinations and Leflunomide treatment. Vaccination with live vaccine is therefore not recommended. The prolonged half-life of Leflunomide should be considered when contemplating live vaccine after stopping the drug.

**Please consult up to date relevant literature (data sheets) or BNF when prescribing this agent. Please contact the Rheumatology team if you have any other queries regarding the prescribed medication.**

### Contact Details

Davida Hehir, Clinical Nurse Specialist in Rheumatology: (021) 4922645/087 6011746  
Arun Augustine, Clinical Nurse Specialist in Rheumatology: (021) 4234517/087 4518911