Sacubitril Valsartan ▼ for Heart Failure

Sacubitril Valsartan (Entresto® ▼) – Amber, specialist initiated including dose tritration to a stable dose

Criteria for use

• NICE TA388 recommends sacubitril valsartan as an option for treating symptomatic chronic heart failure with reduced ejection fraction, only in people:
  • with New York Heart Association (NYHA) class II to IV symptoms  AND
  • with a left ventricular ejection fraction (LVEF) of 35% or less  AND
  • who are already taking a stable and optimised dose of angiotensin-converting enzyme (ACE) inhibitors or angiotensin II receptor-blockers (ARBs)

Treatment with sacubitril valsartan should be started by a heart failure specialist with access to a multidisciplinary heart failure team. ALWAYS PRESCRIBE GENERICALLY

Local Wiltshire Pathway for access

• NICE recommend this drug is best suited to be initiated and titrated in a community heart failure clinic, where they can select the most suitable patients and monitor them closely for side-effects and titrate the dose accordingly
• Once stable, the patient can then be sent out to the GP for them to continue to prescribe
• Wiltshire CCG does not currently have such a service
• As a result of the successful pilot, a limited community heart failure clinic service is due to start in South Wiltshire at the beginning of June 2016
• Discussions are on-going with respect to developing a service across all of Wiltshire
• At the current time, any suitable patients will have to be referred to secondary care cardiology clinics for initiation and titration until a community service is commissioned
• GPs should not initiate this drug themselves (unless they have a specialist interest in heart failure)

Important Drug Information

Sacubitril/valsartan must not be used concurrently with ACE inhibitors due to the risk of life threatening angioedema. It should also not be used with ARBs due to its angiotensin II receptor blocking activity

• Sacubitril/valsartan is a fixed dose combination preparation containing a first-in-class: Angiotensin Receptor Antagonist Neprilysin inhibitor (ARNi). It is a complex of valsartan and sacubitril

Dose Information (PRESCRIBE GENERICALLY)

• For full drug information please refer to the Summary of Product Characteristics https://www.medicines.org.uk/emc/medicine/31244
• STOP the ACE-I that the patient was on for at least 36 hours before starting sacubitril/valsartan. (In practice, a more pragmatic 2 days washout will be used, but the SPC stipulates at least 36 hours). A washout period is not needed if switching from a ARB to Sacubitril/valsartan
• The recommended starting dose is 100mg BD with or without food
• The dose should be doubled every 2-4 weeks to the target dose of 200mg BD as tolerated by the patient
• U&Es should be checked prior to initiation and before each dose titration
• Patients should have been titrated up to maximum tolerated doses of ACE-I s or ARBS before trying Sacubitril/Valsartan
• If systolic blood pressure between 100-110mmHg or moderate renal impairment (eGFR 30-60 ml/min/1.73m2) consider starting with lowest dose (24/26mg)
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Further information on use in renal and hepatic impairment, adverse effects and drug interactions

- **Drug interactions of note (not exhaustive):** (see SPC for full details)
  - Statins, PDE5 inhibitors, Rifampicin, ciclosporin and anti-virals, Metformin, Lithium, NSAIDS, Cox-2 inhibitors

- **Contraindications (not exhaustive):**
  - Do not use with Aliskiren
  - Patients with a known history of angioedema related to previous ACE-I or ARB therapy
  - SBP is less than 100mmHg
  - Renal and Hepatic impairment - no dose adjustment required in mild or moderate (eGFR 30-90mls/min/1.73m²) renal impairment or mild hepatic impairment.
  - Caution is recommended when using sacubitril/valsartan in severe renal impairment or end stage renal disease or moderate hepatic impairment; use a starting dose of 50mg BD
  - Overall the adverse effect profile including the incidence of hypotension and associated adverse effects, particularly the propensity for sacubitril/valsartan to precipitate renal impairment (due to decreased renal perfusion) is unclear from the available trial data
  - Side effects listed as very common (≥ 1/10) in the SPC are hyperkalaemia, hypotension and renal impairment
  - Careful upward titration may be needed to ensure patient tolerability and safety which in turn may lead to a decrease in overall efficacy and mortality. A lower strength is licensed to facilitate a lower initial dose and upward titration.
  - Sacubitril/Valsartan is a black triangle drug, please monitor for and report any side effects via the yellow card reporting system: www.mhra.gov.uk/yellowcard

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  - Caution is recommended when using sacubitril/valsartan in severe renal impairment or end stage renal disease or moderate hepatic impairment; use a starting dose of 50mg BD

- **Cost information**
  - Sacubitril 24mg/valsartan 26mg 56 tabs: £91.56 Dose: 1 bd
  - Sacubitril 49mg/valsartan 51mg 56 tabs: £91.56 Dose: 1 bd
  - Sacubitril 97mg/valsartan 103mg 56 tabs: £91.56 Dose: 1 bd
  - Due to the flat pricing structure, it is important that prescribers ensure dose optimisation to minimise cost (e.g. 1 x 100mg NOT 2 x 50mg)
  - The cost of treating one patient with sacubitril/valsartan for one year = £1,098 vs £13-£41 for enalapril 10-20 mg bd

- **Tolerability**
  - If patients experience tolerability issues such as those listed below, dose adjustment of concomitant medications, temporary down-titration or discontinuation of sacubitril/valsartan may be required. This should be dealt with by the specialist team.
  - Systolic blood pressure ≤95mmHg
  - Symptomatic hypotension
  - Hyperkalaemia (If above 5.4mmol/l consider discontinuation)
  - Renal dysfunction
  - Symptomatic hypotension and decreased renal function is more likely to occur if the patient has been volume depleted, e.g. by diuretic therapy, concomitant NSAID use, dietary salt restriction, diarrhoea or vomiting.
  - If a patient has tolerability problems after previously tolerating it then seek further advice on management.

- **Other information**
  - The number needed to treat, to prevent one primary event and one death was 21 and 32 respectively (from Paradigm HF trial)
  - There is limited data beyond 27 months. The long term efficacy and safety profile of sacubitril/valsartan has not been defined. There is also limited data in patients aged less than 18 years and patients over 75 years of age
  - There is limited comparative data with alternative treatments for heart failure
  - Improvement in quality of life measures have not been adequately assessed to date
  - There is limited clinical experience with use of sacubitril/valsartan in NYHA class IV patients. Hence this should be used with caution in such patients.

- **Contact details of local services**
  - RUH: david.skirrow@nhs.net (Cardiology Pharmacist) Dr Jacob Easaw (Consultant cardiologist) secretary: 01225 825442
  - SFT: Helen Moule, secretary to Dr Anthony Jones (Consultant Cardiologist) 01722 429223
  - GWH: GWH.Heartfunctionteam@nhs.net 01793 604273
  - For patient warning cards and information see: www.entresto.co.uk

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