





Surrey (East Surrey CCG, Guildford & Waverley CCG, North West Surrey CCG, Surrey Downs CCG & Surrey Heath) North East Hampshire & Farnham CCG and Crawley, Horsham & Mid-Sussex CCG

### Information sheet

# **IVABRADINE**

# **Treatment of Angina in Chronic Stable Angina**

# NHS Surrey's Medicines Management Committee classification: Amber\*

Amber\*: Drugs that require initiation by a specialist in secondary / tertiary care but due to more widespread experience in primary care GPs are generally happy to prescribe on specialist advice without the need for a formal shared care protocol. This information sheet is available on the internet (<a href="www.surreyhealth.nhs.uk">www.surreyhealth.nhs.uk</a>) forming part of Surrey PCT's traffic light document giving GPs appropriate advice / guidance and is not required to be sent to the GP with the clinic letter. A minimum of one month supply of medication will be provided by the initiating consultant.

### **RESPONSIBILITIES and ROLES**

### Specialist responsibilities

- 1. Assess the patient to determine suitability for prescribing of ivabradine; ensuring other treatment options have been fully explored.
- 2. Initiate treatment with Ivabradine (initial 1 month supply from Acute Trust Pharmacy).
- 3. Provision of advice to patient and GP on further management of treatment for optimum control of symptoms and related issues such as drug interactions.
- 4. Agree need for and frequency of review by secondary care in consultation with GP. Assess continued appropriateness of ivabradine and advise GP of any changes and additions to prescribed therapy if necessary.
- 5. Evaluating any concerns arising from treatment, including serious adverse events and drug interactions advising on appropriate action. Responding to any queries from the patient or GP relating to treatment of ivabradine.

#### **General Practitioner responsibilities**

- 1. Subsequent prescribing of ivabradine at the dose recommended.
- 2. Monitoring the patient's overall health and well being.
- 3. Observing patient for evidence of ADRs/abnormalities induced by ivabradine and raising with secondary care clinician if necessary.
- 4. Reviewing the patient on a regular basis to monitor heart rate and general condition in line with their usual appointments as deemed appropriate by the CHD register. If the resting heart rate falls below 50 beats per minute then consider reducing dose and seek opinion of referring consultant if necessary. If there is evidence of atrial fibrillation seek advice from referring consultant.
- 5. Ensuring advice is sought from the secondary care clinician if there is any significant change in the patient's physical health status.
- 6. Report any serious adverse events to the Consultant.

## Patient's / Carer's roles

- 1 Ask the specialist or GP for information, if he or she does not have a clear understanding of the treatment.
- 2 Share any concerns in relation to treatment with Ivabradine
- 3 Tell the specialist or GP of any other medication being taken, including over-the-counter products.
- 4 Read the patient information leaflet included with your medication and report any side effects or concerns you have to the specialist or GP

### **BACK-UP ADVICE AND SUPPORT**

| Contact Details | Specialist            | Telephone No. | Email address                               |
|-----------------|-----------------------|---------------|---|
| Specialist:     | Dr. Nandu Gandhi      | 01737 768511  | Nandkumar.Gandhi@sash.nhs.uk                |
|                 |                       | Ext. 6746     |   |
| Specialist:     | Dr. Shrilla Banerjee  | 01737 768511  | Shrilla.Banerjee@sash.nhs.uk                |
|                 |                       | Ext. 2028     |   |
| Specialist:     | Dr Rashid Iqbal       | 01737 768511  | Rashid.lqbal@sash.nhs.uk                    |
|                 |                       | Ext. 6975     |   |
| Specialist:     | Dr. James Sneddon     | 01737 768511  | James.Sneddon@sash.nhs.uk                   |
|                 |                       | Ext. 2026     |   |
| Specialist:     | Dr. Richard Allen     | 01737 768511  | Richard.Allen@sash.nhs.uk                   |
|                 |                       | Ext. 6945     |   |
| Specialist:     | Dr. Ansuman Saha      | 01737 768511  | Ansuman.Saha@sash.nhs.uk                    |
|                 |                       | Ext. 6746     |   |
| Hospital        | Surrey and Sussex     | 01737 768511  | MedicinesInformationPharmacists@sash.nhs.uk |
| Pharmacy:       | Medicines Information | Ext 6246      |   |
| Out of hours    | On-Call switchboard   | 01737 768511  |   |
| contact:        |                       |               |   |

# **Supporting Information**

Ivabradine is an novel antianginal therapy licensed for use in patients with chronic stable angina in sinus rhythm, who have a contra-indication or intolerance to beta-blockers. It is a pure heart rate lowering agent and has been shown to be as effective as beta-blockers in anti-anginal and anti-ischaemic activity, but as yet is not supported by cardiovascular outcomes data.

Ivabradine should only be initiated by a consultant cardiologist or a primary care angina specialist, who should issue the first prescription. Baseline blood pressure and pulse must be provided to GP to aid monitoring.

| Contra-indications   | Cautions  |
|--|---|
| <ul> <li>Sick sinus syndrome</li> </ul>                                    | <ul> <li>Pre-existing cardiac arrhythmias</li> </ul>              |
| <ul> <li>Bradycardia (resting heart rate &lt;60bpm)</li> </ul>             | <ul> <li>Concurrent HR lowering agents</li> </ul>                 |
| <ul> <li>Cardiogenic shock and acute MI</li> </ul>                         | <ul> <li>Mild heart failure (NYHA class I-II)</li> </ul>          |
| <ul> <li>Within 4 weeks of CVA</li> </ul>                                  | - Post-CVA  |
| <ul> <li>Sino-atrial block &amp; 3<sup>rd</sup> degree AV-block</li> </ul> | <ul> <li>Retinis pigmentosa</li> </ul>                            |
| <ul> <li>Moderate to severe heart failure (NYHA class III-IV)</li> </ul>   | <ul> <li>Hypotension (avoid if BP &lt; 90/50mmHg)</li> </ul>      |
| <ul> <li>Congenital QT syndrome</li> </ul>                                 | <ul> <li>Hepatic insufficiency (avoid if severe)</li> </ul>       |
| <ul> <li>Pacemaker dependent patients</li> </ul>                           | <ul> <li>Severe renal insufficiency (CrCl&lt;15ml/min)</li> </ul> |
| <ul> <li>Unstable angina</li> </ul>  |   |
| <ul> <li>Pregnancy and lactation</li> </ul>                                |   |

## Commonly Used Interacting Drugs (See BNF for a full list of drug interactions)

- Amiodarone or disopyramide increased risk of ventricular arrhythmias
- Macrolide antibiotics, particularly clarithromycin and erythromycin avoid concomitant use
- Imidazole anti-fungals, particularly ketoconazole and itraconzole avoid concomitant use
- Nelfinavir and ritonavir avoid concomitant use
- Sotalol increased risk of ventricular arrhythmias
- · Calcium channel blockers, specifically diltiazem and verapamil avoid concomitant use
- Mefloquine avoid concomitant use

## Initiation: Ivabradine is usually initiated at a dose of 5mg twice daily.

After 3-4 weeks dose may be increased to 7.5mg twice daily if required for greater symptom control on advice of hospital consultant. If the patient is elderly or 5mg twice daily is not tolerated, the dose can be reduced to 2.5mg twice daily.

### Monitoring

- Obtain baseline BP, pulse before initiation and after each change in dose
- In the absence of adverse effects, review within 4 weeks and consider increasing the dose if required for better symptor
- Visual symptoms were the most common adverse effects reported. Luminous phenomena were reported in 15% of patient and therefore new patients should be warned about this potential transient side effect.

  Other potential side effects include bradycardia, AV 1<sup>st</sup> degree block, ventricular extrasystoles, headache and dizziness.

Ivabradine is still a black triangle drug and therefore all adverse effects should be reported to the CSM using the yellow car system, even if well documented.

This information sheet does not replace the SPC, which should be read in conjunction with this guidance. Prescribers should also refer to the appropriate paragraph in the current edition of the BNF.

The GP has the right to refuse to agree to shared care, in such an event the total clinical responsibility will remain with the consultant.