## Barnsley Hospital Lipid Optimisation Clinic

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## Why do we need a new lipid clinic?

- Only locality within SY ICB which doesn't have a secondary care lipid service provision.
- To improve access to newer and more specialised lipid lowering treatments.
- To reduce hospitalisation in patients at highest risk of CVD
- To reduce health inequalities in CVD
- To provide support to primary care teams


## The Clinic Team

- Dr A Q Negahban - Consultant Cardiologist, BHNFT
- Dr H Delaney, Consultant Lipidologist, STH
- Gillian Turrell, Lead Cardiology Pharmacist, BHNFT


## Routine treatment options

- Statins
- If issues with intolerance can try different statins (hydrophilic/lipophilic) and/or reduced frequency dosing schedules
- Ezetimibe
- As an adjunct to statins or an alternative if statin intolerant
- Fibrates no longer routinely recommended


Statins are the comerstone for prevention and treament of cardiovascular (CV) disease with a substantial evidence of reduction of morbidity and mortailty. Refer to Lipid Management Pathway and related NICE guidelines (CG181) CG71) for guidance on intiation, titration and monitoring of statin therapy. - In clinical trials, statins were found to be largely well tolerated (often with a similar adverse effect (AE) profile to placebo), however this is not reflected in clinical practice where up to $75 \%$ of people started on a statin will discontinue treatment within 2 years.

- Stopping statin therapy is associated with an increased risk of major CV events and there is growing concem that clinicians are labelling patients as associated with negative media coverage.
Definition of Statin Intolerance
- Intolerance to initial statin therapy is defined by NICE as the presence of clinically significant adverse effects that represent an unacceptable risk to the patient or that may reduce compliance with therapy.
Other definition: any adverse event (AEs) considered unacceptable by
the patient, and/or some laboratory abnormalities, both attributed to stati) the patient, and/or some laboratory abnorma
treatment and leading to its discontinuation.
Statin-associated muscle symptoms (SAMS)
- SAMS are one of the principal reasons for statin non-adherence and/or discontinuation. However, not all such symptoms should lead to a label of 'statin intolerance' as they may not be truly statin related muscle toxicity
(SRM) as demonstrated by resolution on de-chalenge and recurrence with re-challenge.
Non-Statin related musculoskeletal symptoms (Non SRM) - If patients report symptoms that are not typical of SRM (e.g. asymmetric
distribution, failure to rescolve with de-chalenge despite normal CK) conside distibuich, aliure to resodve wth de-chalenge despte normar Ck) consider other muscuioskeletal disorders, metabolic, degenerative or inflammatory e.g.
Vitamin D deficiency, polymyalgia reumatica. Check Bone profili, Vit D , CR .


## Considerations when starting a statin to reduce risk of SRM

 - Check baseline thyroid, liver and renal function, any potential drug interactions, Check bassine thyroid, iver and renal funcion, any potental drug interactand avoid the highest doses in at risk groups (See "Risk Factors' below).
Ask the person if they have had persistent generalised unexplained muscle pain, whether associated or not with previous lipid-lowering therapy. If
they have, measure CK. If CK levels are $>4 x$ ULN do not starts statininvestigation required.
Do not measure CK if person is asymptomatic.

- Warn patients about AEs, specifically muscle symptoms. Advise people
who are being treated with a statin to seek medical advice if they develop muscle symptoms (pain, tenderness or weakness). If this occurs, measure CK (see page 1).


## Risk factors for SRM and statin intolerance

Endogenous factors

- Female gender

Frailty (red 75 yrs)
Frally (reduced lean body mass)

- Impaired renal or hepatic function
- Persenal or family history of intele

Personal or family history of intolerance
to lipid-lowering therapies.

- Hypothyroidism

Classification of statin related muscle toxicity (SRM)

| SRM | Phenotype | Incidence | Deffinition |
| :---: | :---: | :---: | :---: |
| SRM 0 | CX elevation <4x ULN | 1.5-26\% | No muscle symptoms |
| SRM 1 | Myalgia, tolerable | 190/100,000 <br> Patient-years: $0.3-33 \%$ | Muscle symptoms without CK elevation |
| SRM 2 | Myalgia, intolerable | 0.2-211,000 | Muscle symptoms, CK $<4 x$ ULN, complete resolution on dechallenge |
| SRM 3 | Myopathy | $\begin{aligned} & 5 / 100,000 \\ & \text { Patientyears } \end{aligned}$ | CK elevation $>4 \times$ ULN $<10 \mathrm{x}$ ULN $\pm$ muscle symptoms, complete resolution on dechallenge |
| SRM 4 | Severe myopathy | 0.11\% | CK elevation > $>10 x$ ULN $<50 x$ ULN, muscle symptoms, complete resolution on dechallenge |
| SRM 5 | Rhabdomyolysis | 0.1-8.4/100.000 | CK elevation $>10 x$ ULN with evidence of renal impairment + muscle symptoms or CK $>50 \mathrm{x}$ ULN |
| SRM 6 | Autoimmune-mediated necrofzing myosis (SINAM) | $\underset{\substack{-2 \text { milion per } \\ \text { year }}}{ }$ | Detection of HMGCR antibodies, HMGCR expression in muscle biopsy showing autoimmune myosifis, incomplete resolution on dechalenge |

HMGCR $=3$-hydrony 3 -metrygutary coennyme A reductase ULN $=$ upper imt of nomal

- SRM is a spectrum from myalgia to severe myopathy
- SRM 0 - does not preclude statin therapy, consider reducing starting dose - SRM 1-3 manage according to pathway
- When SRM4 is suspected, without evidence of impaired renal function, discontinue statin therapy immediately and refer for outpatient assessment. Assess and treat possible contributory factors and re-assess the need for a station. Intensify lifestyle modifications and consider alternative lipid lowering regimens. - If inabdomyolysis (SRM5) is suspected, immediately stop statins, urgently refer to inpatient assessment and management including intravenous rehydration as required to preserve renal function. Do not wait for measurement of urinary myoglobin. Post recovery, manage as for SRM4.
- Statin induced necrotizing autoimmune myositis (SINAM) (SRMG) should be
suspected in patients with progressive muscle weakness and ongoing CK suspected in patients with progressive muscle weakness and ongoing CK and avoidance of re-exposure to statins. Re-assess the need for lipid lowering therapy - may be eligible for treatment with PCSK9 inhibitor (NICE TA 393, 394).


## nitial Consultation

- Be aware of "nocebo effect" and

Reinforce heathy lifestyle habits
(e.g. exercise, reducing weight)

- Listen to the concems of each patient.

Explain LDL-C targets and strategles
to lower LDL-C/non-HDL-C

- Discuss options to reduce LDL-C/
non-HDL-C with pros and cons
- Explain the benefits of statins
- Evaluate and identify any risk factors
and address (e.g. drug interactions)

Work with patients io identify and Work with patients to identify and
agree best options and next steps
follow up
Follow up on agreed plan and address any issues/concern.

- Advise patients to contact you if they experience muscle symptoms Ongoing patient education and regular review helps addressing concerns around medicine safety and underline the importance of adherence.
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 are press ribed a placebo
Statin celuctance is an altilusnal state of
aversion to taiking statins (chen without proo exposurie).


## Statin-based approaches to manage muscle symptoms

## - Adopt person-centred approach as described above.

- Apply a repetitive "De-Challenge". . "Re-Challenge" approach to establish if
symptoms are caused by a statin(s) and the best statin regimen for each patient. - Switch to a different statin or re-challenge with the same statin using a lower dose or frequency (intermittent dosages)
Patients who do not tolerate statins on a daily basis, alternate day or twice-weekly dosing is a good option.
- Rosuvastatin and atorvastatin have longer half-lives, permitting their use on a non-daily regime.
Adding ezetimibe to a lower dose statin may be better tolerated with robust of LDL-C / non-HDL-C.
Once a new regime is tolerated, dose / frequency can be up-titrated slowly to
goals with minimal or no muscle complaints.
It is important to note that cardiovascular benefits have not been proven for all the
above appraaches but any reduction of LOL-C/ non-HDL-C is beneficial
LDL-C lowering options for patients with genuine statin intolerance
- Refer to the AAC Lipid Management Algorithm. (dllick here)

Consider ezetmibe, (NICE TA 385) therapy as per algorith

- Consider ezetimbe combined with bempedoic acid (NICE TA 694) as per algorithm
- Consider inclisiran if elligible for treatment according to NICE TA 733
- Consider PCSK91 if eligible for treatment according to NICE TA 393, 394


## Non-muscle related statin side effects

May vary between different statins. In clinical trials some side effects often associated with statins are not statistically different from placebo.
Most commonly reported: gastrointestinal disturbance and asymptomatic increases in hepatic transaminases (ALT or AST). May affect up to 1 in 10 statin users. Rarer side effects include: Hepatotoxicity, new onset Type 2 Diabetes (benefits outweigh risk, do not stop statin). Renal insufficiency, proteinuria, Neurocognitive and neuroiogical impairments (no apparent link from RCTs), Intracranial haemorthage (conflicting evidence, benefit outweigh possible harm), Intersititial lung disease, Pancreatitis, Skin disorders including alopecia, Lupus-like reaction, Sleep disturbance, headache, dizziness, fatigue, depression, sexual dysfunction. Management: If symptoms appear statin related, consider de-challenge and re-challenge or change to a different statin (e.g. hydrophilic instead of lipophilic). Liver enzyme abnormalities - minor increases in liver enzymes ( $<2 \mathrm{x}$ ULN) may be seen within the first three months of statin therapy, temporary discontinuation and further assessment is warranted filevels exceed $3 x$ ULN. Several studies have conffrmed that the cardiovascular benefits of statin treatment in high-risk populations outweigh the rare adverse effects, such as mabdomyolysis.

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## Specialist treatment options

- Bempedoic acid plus ezetimibe
- Statins contraindicated or not tolerated
- Ezetimibe alone ineffective in optimising lipid management
- Inclisiran as per NICE TA733
- History of cardiovascular disease
- LDL-C persistently above $2.6 \mathrm{mmol} / \mathrm{I}$ despite maximum tolerated statin therapy +/- other lipid lowering therapies
- Initiated or recommended by specialist clinic within Barnsley


## Specialist Treatment options cont.....

- PCSK9 inhibitors as per NICE TA393 (Alirocumab) and TA394 (Evolocumab)
- Patients at high risk of CVD
- Persistently high LDL-C levels
- Icosapent Ethyl as per NICE TA805
- Established CVD
- Fasting triglycerides $\geq 1,7 \mathrm{mmol} / \mathrm{l}$ and taking statins
- LDL-C >1.04mmol/l but $\leq 2.6 \mathrm{mmol} / \mathrm{l}$

Theoretical \% LDL-C Reduction Obtained by Different Lipid-Lowering Combination Therapies


Figure created using data from Masana L, et al. Rev Esp Cardiol (Engl Ed). 2016;69:342-343
Masana L, et al. Curr Cardiol Rep. 2020;320:122-128
Wichalyo S, ot al. Clin Drug Investig. 2021;41:843-851

## Which Patients?

- Not familial hypercholesterolaemia patients requiring genetic testing $\rightarrow$ STH
- Mixed dyslipidaemia patients
- Working with Dr Dowling and Khawer Ashfaq to define initial referral criteria:
- Highest risk of CVD
- Routine treatment options already optimised or not tolerated
- Social deprivation scores


## Questions...?

