Sunderland CCG does not support the prescribing of nefopam 30mg tablets in primary care

- **Nefopam** is a centrally acting non-opioid analgesic with associated antimuscarinic and antihistaminergic effects recommended for persistent pain unresponsive to other non-opioid analgesics.¹
- The **BNF** indicates nefopam may have a place in the relief of persistent pain unresponsive to other non-opioid analgesics, but prescribers need to consider carefully whether the anticipated benefits outweigh the risks of adverse effects, especially in high risk groups including the elderly.
- Most of the **studies** assessing the efficacy of nefopam are either single dose or short term based; the majority of these involve parenteral administration which is not supported by the UK marketing authorisation. The evidence base for the efficacy of nefopam is weak, conflicting or absent² ³ ⁴ ⁵ in reducing pain in patients with RA or postoperative period.
- **Adverse effects are common** and include nausea, sweating, dizziness, vomiting, hallucinations, confusion, urinary retention, headache, insomnia, tachycardia, palpitations convulsions and anaphylaxis.
- Nefopam scores 2 on the anticholinergic burden scale (ACB).⁶ Each definite anticholinergic may increase the risk of cognitive impairment by 46% over 6 years. For each point increase in the ACB total score, a decline in MMSE score of 0.33 points over 2 years has been suggested. Additionally, each one point increase in the ACB total score has been correlated with a 26% increase in the risk of death.
- Nefopam is toxic in overdose with observed clinical manifestations including seizures, first degree heart block, right bundle branch block, ventricular tachycardia, acute renal failure, cerebral oedema and pulseless electrical activity. Four deaths following intentional nefopam overdose have been reported. The fatal dose, known in one case only, was 1.8g.
- Nefopam has abuse potential primarily through its psychostimulant-like effects, which are probably linked to its dopamine reuptake inhibition properties. There are reports it is being identified on drug screening results. Where nefopam has been used for its abuse potential withdrawal may lead to depression,⁷ no information is available about the necessity to taper dosing following long term use at usual or standard doses.
- NB: Locally Newcastle Gateshead CCG and North Tyneside CCG are stopping the use of nefopam.

**Bottom line** what does this mean in practice?
- don’t initiate nefopam for acute or chronic pain except under advice from specialist pain service
- do not continue nefopam post discharge following secondary care acute initiation
- only continue nefopam in line with the recommendations of the specialist pain service

¹ SAFER MEDICATION USE Nefopam No 14 January 2015 RDTC