

PENTHROX[®] (methoxyflurane)

Patient Alert Card

IMPORTANT SAFETY INFORMATION

This medicine can cause **LIVER** or **KIDNEY** problems

Please READ and KEEP this Patient Alert Card

Date of Administration: _____

Patient Name: _____

Dear Patient and Caregiver,

This medicine can cause LIVER and KIDNEY PROBLEMS which can occur in the days or weeks after use.

If this does occur, it can be life-threatening so **seek medical attention IMMEDIATELY and show the healthcare professional this card if you experience any of the symptoms listed opposite:**

1. Loss of appetite

2. Nausea

3. Vomiting

4. Dark urine

5. Reduced or excessive
urination

6. Pale stools

7. Pain/ache or sensitivity to
touch in your right stomach
area (below your ribs)

8. Swelling of feet or lower legs

9. Jaundice (yellowing of the skin
and/or eyes)

Please read the Patient Information Leaflet, which should have been provided to you when you were treated, for full details about this medicine.

Tell your doctor, pharmacist or nurse about any side effects you get, **including ones not listed in this card.**

You can also report side effects directly via HPRA Pharmacovigilance. Website: www.hpra.ie

Any suspected adverse reactions should also be reported to Galen Pharma Ireland Limited on 048 3833 4974 and select the customer services option, or e-mail customer.services@galen-pharma.com.

By reporting side effects you can help provide more information on the safety of this medicine.

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