Methemoglobinemia during the Use of Glyceryl Trinitrate Patches in Neonates: Two Case Reports

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Abstract

Methemoglobinemia can result in severe hypoxia. It has been frequently reported during the use of inhaled nitric oxide, but can occur where nitrate containing medications are used. Glyceryl trinitrate (GTN) patches have been used in the treatment of digital and limb ischemia in prematurely born infants. Little is known about the pharmacokinetics of GTN when incorporated into patches. Studies of other topical forms of nitroglycerine have shown a wide range of absorption. It is likely that the increased permeability of the prematurely born infant’s skin would facilitate absorption. We describe the use of GTN patches in two very prematurely born infants used to treat limb/digit ischemia. This resulted in methemoglobinemia and resultant increase in their supplementary oxygen requirements. Removal of the patches was associated with a reduction in their methemoglobin levels and the supplementary oxygen requirements back to baseline levels. In conclusion, routine monitoring of methemoglobin levels should be undertaken when GTN patches are used in very prematurely born infants.

Keywords
► methemoglobinemia
► preterm
► glyceryl trinitrate
► ischemia

Methemoglobinemia is formed as a result of the oxidization of iron from its ferrous (Fe²⁺) to ferric (Fe³⁺) state and can occur following exposure not only to inhaled nitric oxide (iNO) but also nitrite containing medications. Methemoglobinemia has been frequently reported in infants treated with iNO.¹ The levels of methemoglobinemia relate to the iNO levels used.¹ Methemoglobin has a higher oxygen-binding capacity than hemoglobin and thus affected infants can suffer severe hypoxia. Ischemia can be caused by a combination of vascular damage and concomitant vasoconstriction.² Glyceryl trinitrate (GTN) patches are used in the treatment of ischemic digits and limbs in neonates²,³ and in adults.⁴ GTN is converted to NO in the vascular smooth muscle, and this activates guanylate cyclase and increases the levels of cyclic guanosine monophosphate (cGMP);³ cGMP causes relaxation of the vascular smooth muscle in the affected artery and
dilation of the collateral circulation. GTN patches, however, remain unlicensed for use in neonates and little is known about the pharmacokinetics of GTN when used in patch formation in prematurely born neonates. Methemoglobinemia is not listed as a side effect of the use of GTN patches. The absorption of nitroglycerin from topical ointments has been shown to be variable in infants. GTN patches vary in size and dose. A 10-cm² patch typically contains 40 mg of nitroglycerin and is designed to release this at 0.2 mg/h; however, in one study of seven infants, an application of 1 mg in an ointment resulted in blood levels of between 0.03 and 3.36 ng/mL. The skin of prematurely born infants has increased permeability compared with term born infants which might lead to greater absorption of topical treatments. We present two cases of methemoglobinemia which occurred during the use of GTN patches to treat ischemic digits and limbs of very prematurely born infants. In both cases, 9 cm² patches were used which contained 18.7 mg GTN.

**Case 1**

An infant with a birth weight of 650 g was born at 24 weeks of gestational age following the onset of spontaneous preterm labor. During initial resuscitation, it was noted that the lower limbs of the infant were dusky and this did not resolve despite improving the blood pressure levels by administering volume replacement and inotropes. One GTN patch was applied to each lower limb over the area of the posterior tibial arteries. These were left in place as the infant was stabilized and transferred to the local tertiary unit. A radial arterial line was inserted the following day, day 2 after birth. The thumb and forefinger distal to the radial arterial line became poorly perfused shortly after insertion and a further GTN patch was applied to the hand. This was in addition to the two patches in situ on the feet. The oxygen requirement was noted to rise from an inspired oxygen fraction (FiO₂) of 0.21 to 0.4 within 6 hours of application of the third patch. The infant’s inspired oxygen concentration was altered as necessary to keep the oxygen saturation levels between 92 and 95%. Serial capillary blood gases showed an increase in methemoglobinemia from 1.1% to 9.5%, with a corresponding reduction in the oxygenation index (mean airway pressure × FiO₂ × 100 divided by PaO₂, kPa), from 24 to 8. The tips of the toes that were necrotic sloughed off after several weeks, but no amputation was required.

**Discussion**

We have demonstrated GTN patch use in very prematurely born infants can result in methemoglobinemia. In both infants, this was associated with an increase in their inspired oxygen requirement. In the first case, the ischemia completely resolved, but in the second resolution was incomplete and as a consequence, the toes became necrotic. In both cases due to the development of the methemoglobinemia, the patches were removed prematurely, which may have reduced their efficacy. Previous case reports have suggested a treatment duration of 28 days.

It is not clear whether the methemoglobinemia was the cumulative effect of all the patches applied or after a certain number of patches a threshold was reached. It is known, however, that prematurely born infants have lower levels of methemoglobin reductase which makes them more susceptible to methemoglobinemia. Once the patches were removed, the methemoglobin levels decreased and the supplementary oxygen requirement returned to the level prior to treatment with the patches. Thus, we suggest the use of the GTN patches resulted in the methemoglobinemia. As a consequence, we recommend if GTN patches are used in a very prematurely born infant, frequent assessment of methemoglobin levels should be undertaken.

Conflict of Interest

None.

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