Phototherapy: A Simple and Safe Treatment for Neonatal Jaundice

Abstract

Phototherapy: A simple and safe treatment for Neonatal Jaundice. Jaundice is a common finding in premature and full term newborns. Phototherapy has been widely used in the management of Neonatal unconjugated Hyperbilirubinemia for over five decades. Phototherapy devices include Fluorescent, Halogen, Fiberoptic or Light-Emitting Diode light sources. Safety issues and possible complications of phototherapy are discussed. It is a non invasive, safe and easily available therapy worldwide. This review will focus on the use of phototherapy in newborn nurseries.

Keywords: Phototherapy; Hyperbilirubinemia; Light treatment; Newborn

Introduction

In 1958, Cremer et al. [1] demonstrated that serum bilirubin concentrations fall faster in premature infants exposed to sunlight or blue fluorescent bulbs. However, phototherapy wasn’t commonly used until 1968, when Lucey et al. [2] conducted the first large clinical trial using daylight fluorescent tubes for the treatment of jaundice. Today, it is commonly applied in most nurseries throughout the world in order to prevent kernicterus. Risk factors for the development of severe jaundice and kernicterus include asphyxia, sepsis, acidosis, hypoalbuminemia and hemolysis [3]. The goal of phototherapy is to avoid an exchange transfusion. Any baby placed under phototherapy requires a history and physical examination, as well as laboratory screening for pathologic jaundice. Bilirubin is one of the few substances in the body that absorbs light. Under lights, normal bilirubin (4Z, 15Z-bilirubin) undergoes photoisomerization through the skin to form photobilirubin and lumirubin. These photo products are water-soluble, bypassing the liver’s conjugating system, and are rapidly excreted in both urine and bile. Maintenance of adequate hydration and good urine output enhances the efficacy of phototherapy.

Neonatal Jaundice

Jaundice is the visible yellow appearance of the skin that occurs with chemical hyperbilirubinemia. It happens in adults with serum bilirubin greater than 2mg/dl, and in neonates with serum bilirubin greater than 5 mg/dl. Bilirubin is formed by the breakdown of heme. Newborns have differences in bilirubin production and elimination leading to a rise in bilirubin levels referred to as physiological hyperbilirubinemia that is present in as many as 60% of all normal neonates in the first several days after birth. Pathological newborn hyperbilirubinemia can be caused by increased bilirubin production such as hemolysis or decreased bilirubin clearance such as prematurity. Effective treatments for high bilirubin concentration in infants include intravenous gamma-globulin [3] phototherapy and exchange transfusion. The exact total serum bilirubin level, at which phototherapy should be started depends on several risk factors including prematurity, age of the infant and other risk factors [4].

Indications

To help pediatricians in their decisions on when to start phototherapy, the American Academy of Pediatrics, published in 2011 guidelines on the management of hyperbilirubinemia in newborns 35 or more weeks gestation [5]. Maisels et al. [6] in 2012 provided an approach (Table 1) to use phototherapy and exchange transfusion in preterm infants less than 35 weeks of gestation. The AAP committees on fetus and newborn guidelines were evidence-based but Maisels et al. [6] recommendations are consensus-based. Infants with congenital erythropoietic porphyria are contraindications to the use of phototherapy.

Table 1: Suggested use of phototherapy and exchange transfusion in preterm infants <35 weeks gestation age Maisels et al. [7].

| Gestational Age (week) | Phototherapy | Exchange Transfusion |
|------------------------|--------------|----------------------|
| Total Serum Bilirubin (mg dl-1) | Initiate Phototherapy | Total Serum Bilirubin (mg dl-1) |
| <28 0/7 | 5-6 | 11-14 |
| 280/7 – 296/7 | 6-8 | 12-14 |
| 300/7 – 316/7 | 8-10 | 13-16 |
| 320/7 – 336/7 | 10-12 | 15-18 |
| 340/7 – 346/7 | 12-14 | 17-19 |

Types of Phototherapy

The traditional phototherapy units using fluorescent tubes contain standard blue (Westinghouse F20T 12B), daylight (F20 T12D) and cool white (F20 T12CW) lamps. The most effective lights are those with a high energy output near the maximum absorption peak of bilirubin (450 to 460 nm) [7]. Special blue lights (Phillips TL 52/20W, Westinghouse 20 watt F20 T12BB) are the most efficient for neonatal phototherapy because they have more than twice the energy output at 450 mm than the standard blue bulbs. Investigators using these special blue bulbs report they have achieved a more rapid reduction of serum bilirubin than with daylight or standard blue bulbs [8]. However,
the special blue bulbs have been found to cause nausea and dizziness among the nursery staff. A combination of four special blue lamps placed in the center of the phototherapy unit with two day light lamps on either side has been found to provide excellent irradiance without producing significant discomfort to staff members. Non-fluorescent halogen lamps (spotlights) produce a more intense light over a smaller surface but are more expensive than fluorescents. If they are placed closer than 50 cm, halogen lamps unlike fluorescent bulbs incur the risk of burns to the infant.

Light-Emitting Diode (LED) lights are now commercially available for use in the United States [9]. The Neo Blue LED systems incorporate optimal blue LED technology and are manufactured by Natus Medical Inc., San Carlos, CA, USA. Neo Blue LED’s emit blue light in the 450-470 nm spectrum. They are the safest phototherapy devices available because they do not emit light in the ultraviolet and the infrared radiation range. The virtual absence of heat when delivering overhead Neo Blue phototherapy should be less likely to cause insensible water loss [10]. Fiberoptic phototherapy systems first appeared on the market in 1989. They are widely considered to be equally as effective as and more convenient than overhead lights. Light is delivered from a halogen bulb (bulb life = 450 hours) through a fiber optic cable and is emitted filtered from the sides and ends of fibers inside a plastic blanket which is protected by a disposable cover. Infants lie on the blanket or are held with the blanket wrapped around them, and the need for eye patches - otherwise required in neonatal phototherapy - is eliminated.

Technology currently provides the clinician with three different modes of phototherapy delivery: fiber optic, low intensity, and high intensity phototherapy. For low intensity phototherapy, overhead lamps are typically set at a distance of 50 cm from the patient. High intensity phototherapy has been defined by the American Academy of Pediatrics as a spectral irradiance of at least 30 MW per square meter per nanometer. High-intensity phototherapy is achieved by using a unit with eight special blue lamps or Neo Blue LED systems 25 cm above the naked infant who is on a fiber optic phototherapy blanket in a bassinet, while wearing a tie-on surgeon’s mask as a diaper. This method allows maximum skin exposure and achieves an irradiance as high as 50uw/cm²/nm.

However, as lamps are lowered close to the infant, there is an increase in the heterogeneity of irradiation, with a much greater increase at the center than at the periphery. Lining the bassinet with a white cloth produces greater homogeneity or irradiance. Infants lying in the bassinet should be less likely to cause insensible water loss [10]. Fiber optic phototherapy units first appeared on the market in 1989. They are widely considered to be equally as effective as and more convenient than overhead lights. Light is delivered from a halogen bulb (bulb life = 450 hours) through a fiber optic cable and is emitted filtered from the sides and ends of fibers inside a plastic blanket which is protected by a disposable cover. Infants lie on the blanket or are held with the blanket wrapped around them, and the need for eye patches - otherwise required in neonatal phototherapy - is eliminated.

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However, as lamps are lowered close to the infant, there is an increase in the heterogeneity of irradiation, with a much greater increase at the center than at the periphery. Lining the bassinet with a white cloth produces greater homogeneity or irradiance and an increase in the amount of indirect reflected irradiance. Home phototherapy [11] should be limited to otherwise healthy term infants who are older than 48 hours with bilirubin levels between 15 and 20 mg/dl and no hemolysis. Parents have to be able to monitor the baby’s temperature and hydration status. Home visits by a nurse experienced in evaluating newborns are performed, with results of bilirubin levels available on a timely basis.

Monitoring Light Intensity

The energy output in the 425 to 475 mm range delivered to the patient by any source of light must be monitored every day. A different sensor should be used with fiber optic phototherapy, and measurements are performed directly at the surface of the pad. Caution must be exercised in interpreting measurements from photometers or spectroradiometers, as different instruments provide readouts in varying units. According to current knowledge, there is a dose-response relationship between bilirubin degradation and phototherapy intensity up to a reading of approximately 40 uw/cm²/nm.

Fluorescent lamps decay with use and should not be left on until they burn out. If these lamps are kept adequately cool, their irradiance drops about 10 percent in the first hundred hours of use and 20 percent after 3,600 hours. In contrast halogen lamps used in fiber optic devices or overhead spotlights, have a lifetime irradiation drop of less than a few percent. Therefore, measuring their output is unnecessary as these bulbs can burn out before needing replacement. It should also be noted that skin pigmentation does not reduce effectiveness of phototherapy.

Safety Issues and Possible Complications

Current literature on phototherapy use provides a variety of safety considerations for the clinician. A Plexiglas shield should always be in position under the fluorescent bulbs. This shield acts as a filter for erythemogenic ultraviolet radiation (below 320 nm) and protects in the event of lamp breakage. All phototherapy devices available for use in the United States [9] are widely considered to be equally as effective as and more convenient than overhead lights. Light is delivered from a halogen bulb (bulb life = 450 hours) through a fiber optic cable and is emitted filtered from the sides and ends of fibers inside a plastic blanket which is protected by a disposable cover. Infants lie on the blanket or are held with the blanket wrapped around them, and the need for eye patches - otherwise required in neonatal phototherapy - is eliminated.

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[16], the percentage of fluid intake should be increased in an inverse ratio to the size of the infant (20 to 60 percent above maintenance). Studies have shown there is no difference in fluid loss between fiber optic and conventional phototherapy. During non-LED lights phototherapy, there is an increase in skin and muscle blood flow, insensible water loss, skin temperature, heart rate, and respiratory rate. These effects can be partially decreased by vigilant monitoring of vital signs every two to three hours and by covering the servo probe with a heat reflecting device. Phototherapy may produce a transient rash, transient loose green stools, lethargy, or abdominal distention. Recent studies have suggested that phototherapy is associated with childhood bronchial asthma [17] and is a risk factor for insulin-dependent diabetes [18].

**Conclusion**

Phototherapy is safe [19,20] inexpensive, efficient, and easy to use. With high intensity phototherapy, even severe hemolytic hyperbilirubinemia can be adequately controlled [21].

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