

Original Research Article

A Retrospective Study On Clinical Features Of Early Neonatal Jaundice In Term Babies At Ratchaburi Hospital, Thailand

ABSTRACT

Aims-Background: Neonatal jaundice is the a common condition and sometimes it can lead to devastating neurological consequence like kernicterus.

Aim: This study was aimed to recognize clinical features and to know etiology of term neonatal jaundice.

Study design: Hospital-based retrospective study

Methodology: The study was conducted by reviewing 117 medical records of neonatal jaundice who were admitted at Ratchaburi Regional Hospital, Bangkok, Thailand from 1st October 2007 to 30th September 2008. Both the patient's and their mother's profiles, etiology and clinical features of jaundice were extracted.

Results: The results showed that female babies were more likely to ~~contract-develop~~ jaundice. The most common etiology was inconclusive jaundice (64.9%) followed by ABO incompatibility (17.9%) and breast feeding jaundice (10.2%). ~~In addition to these, o~~ther less common causes were G6PD deficiency, minor blood group incompatibility and cephalhematoma. The onset of the neonatal jaundice usually occurred on the 2nd to the 4th day of life and almost all newborns responded well to phototherapy. Most of interventions were started on the 2nd day of life. Moreover, exchange transfusion was needed in four cases. ~~Maximum microbilirubin and unconjugated bilirubin levels were two factors associated with double side phototherapy and/or exchange transfusion.~~

Conclusion: ~~As stated by our study, various etiologies of hyperbilirubinemia were found. From our study, the most common etiology was inconclusive jaundice which is followed by ABO incompatibility as a second cause but non-immune hemolysis and polycythemia were not discovered-encountered. in this study.~~ There was significant difference of haematocrit, NRC and reticulocytes between haemolytic and non-haemolytic groups. Detailed approach of history taking and physical examination, early investigations of jaundice work up and septic work up are recommended in eliciting various etiologies and preventing complications.

Comment [UP1]: This is not clear.

Keywords: Neonatal jaundice, etiology, clinical feature, retrospective

1. INTRODUCTION

The most common cause of readmission to hospital in healthy term infants is neonatal jaundice [1]. Paediatricians were aggressive in treating jaundice though several factors have changed the management of jaundice [2, 3]. Effective approach and evaluations for management are crucial in preventing bilirubin induced encephalopathy and long term neurodevelopment outcomes [4]. The possible factors exaggerating physiological jaundice ~~were-are~~ shown in table 1 [5].

25
26

Table 1: Possible factors exaggerating physiological jaundice

Factor	Clinical correlate
1. Bilirubin load to liver	<ul style="list-style-type: none">• Infants with polycythemia• Infants of diabetic mother• Collection of extravasated blood like cephalhematoma and intraventricular hemorrhage• Delayed cord clamping
2. Defective uptake from liver	<ul style="list-style-type: none">• Decreased Y protein due to caloric deprivation
3. Defective bilirubin conjugation	<ul style="list-style-type: none">• Due to decreased UDPG activity as seen in hypothyroidism and inhibitors in breast milk
4. Decreased hepatic excretion	<ul style="list-style-type: none">• Congenital infections
5. Inadequate hepatic perfusion	<ul style="list-style-type: none">• Hypoxia• Congenital heart diseases
6. Increased enterohepatic circulation	<ul style="list-style-type: none">• Unfed babies• Delayed passage of meconium

27

28 Studies in the 1980s and 1990s suggested that kernicterus from jaundice was rare and
29 that too many infants were being treated unnecessarily [6, 7]
30 Early detection, effective intervention and new approaches to prevention have been also
31 stimulated as it has a potential damage to developing brain [8]. Previous studies have
32 indicated a relationship between neonatal hyperbilirubinemia and diverse factors including
33 racial region, male gender, epidural anaesthesia, and instrumental delivery [9].
34

35

Risk factors for significant neonatal hyperbilirubinemia are:

36

1. Jaundice visible on the first day of life

37

2. A sibling of jaundice or anaemia

38

3. Unrecognized haemolysis (ABO, Rh and other blood incompatibility)

39

4. Nonoptimal feeding (Formula or breast feeding)

40

5. Deficiency of glucose 6 phosphate dehydrogenase

41

6. Infection, Infant of diabetic mother and immaturity

42

7. Cephalhematoma or bruise, Central haematocrit >65% (polycythemia)

43

8. East Asian, Mediterranean, Native American heritage [10].
44

45

Pathological Jaundice fulfils any of the following criteria:

46

1. Clinical jaundice appearing in the first 24 hour.

47

2. Increase on level of total bilirubin by more than 0.5 mg/dL/hr or 5mg/dL/24 hr.

48

3. Total serum bilirubin > 12 mg/dL in term infants and total serum bilirubin level 10- 14 mg/dL in preterm infants.

49

4. Direct reacting bilirubin > 2.0 mg/dL [11].
50

51

The primary concern with respect to exaggerated hyperbilirubinemia is the potential for neurotoxic effects, but general cellular injury also occurs [12]. Bilirubin can interfere with neuroexcitatory signals, impair nerve conduction (particularly auditory nerve) inhibit ion exchange and water transport in renal cells. **Blood brain barrier can be altered and entry to bilirubin to brain** can be affected by infections, acidosis, hypoxia, sepsis, prematurity and hyperosmolarity [13].
52
53
54
55
56

57

Risk factors of severe hyperbilirubinemia: [14]
58

59 *Major Risk Factors:*

- 60 1. Pre discharge microbilirubin level in high risk zone
- 61 2. Jaundice observed in 1st 24 hour
- 62 3. Blood group incompatibility with positive direct antiglobulin test, other known haemolytic
- 63 disease
- 64 4. Previous sibling received phototherapy
- 65 5. Cephalhematoma or significant bruising
- 66 6. Exclusive breast feeding if nursing is not going well and weight is excess
- 67 7. East Asian Race

68 *Minor Risk Factors:*

- 69 1. Pre discharge microbilirubinemia level in the high intermediate risk zone
- 70 2. Jaundice observed before discharge
- 71 3. Previous sibling with jaundice
- 72 4. Macrosomic infant of diabetic mother
- 73 5. Male Gender

74 *Decreased risk factors: Factors associated with decreased risk of neonatal jaundice*

- 75 1. Microbilirubin level in low risk zone
- 76 2. Exclusive bottle feeding
- 77 3. Black race
- 78 4. Discharge from hospital after 72 hour

79
80 ~~Although it is not common of hazardous hyperbilirubinemia~~ ($\geq 30\text{mg/dL}$)
81 ~~hyperbilirubinemia is uncommon~~, timely recognition, potent work up and compelling
82 management ~~are play~~ pivotal roles ~~to in the prevention of~~ chronic, bilirubin-induced
83 neurotoxicity [15]. This study was designed to ~~get a various document the~~ clinical profiles ~~and~~
84 ~~aetiology~~ of early neonatal jaundice. ~~Careful history taking and~~, physical examinations to
85 elicit information on risk factors were recommended for preventing development of
86 pathological jaundice. Moreover, early measurement of serum bilirubin and work up for
87 jaundice were necessary to know different etiologies.

Comment [UP2]: This is not necessary and should be deleted

88
89 **2. MATERIAL AND METHODS**

90
91 This study was a hospital based retrospective ~~record study review~~ of 117 medical records,
92 conducted at Ratchaburi regional hospital, Bangkok, Thailand from November 2008 to 2009
93 March. The clinical features of early neonatal jaundice ~~d~~ for newborns younger than 7 days
94 whose gestational age ranged ~~s~~ from 37 to 42 weeks were explored. ~~Babies with and~~ onset
95 of jaundice within 7 days requiring intervention ~~were included in the analysis. for jaundice~~
96 ~~was studied.~~ The exclusion criteria included preterm, low birth weights, having major
97 congenital anomalies and congenital infections, systemic infections before onset of jaundice
98 and serious illness including sepsis, meconium aspiration syndrome and severe birth
99 asphyxia.

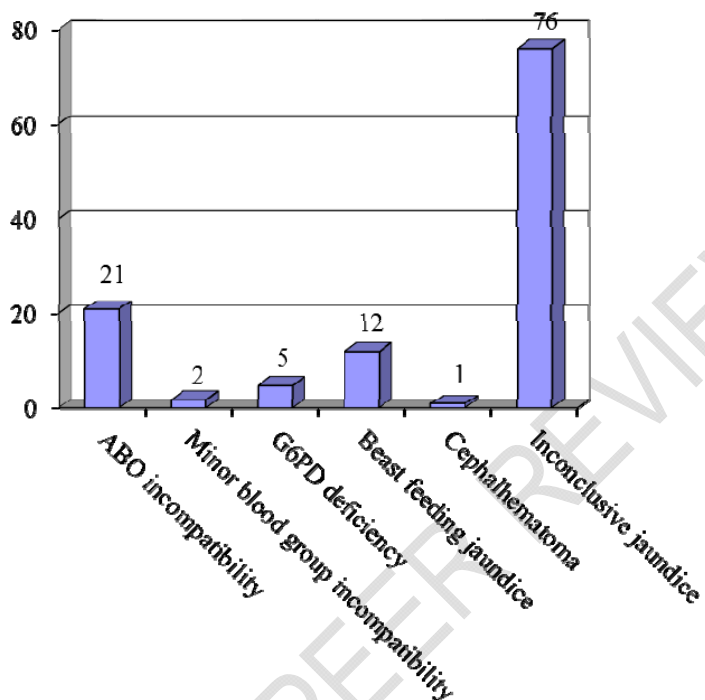
100 Required data from the selected records were collected and transferred into ~~a~~ case record
101 forms that ~~was constructed designed~~ based upon the variables from Ratchaburi Regional
102 Hospital and study objectives. The data obtained were analyzed with SPSS version 11.5. ~~for~~
103 ~~statistical analysis.~~ The research was funded by the Faculty of Tropical Medicine, Mahidol
104 University, Thailand in collaboration of SEAMEO TROPED, Thailand.

105
106 **3. RESULTS**

107
108 Figure 1 shows that ~~inconclusive jaundice~~ was the most common etiology in this study
109 (n=76, 64.9%), which was followed by ABO incompatibility (17.9%). Other diagnoses were
110 breast feeding jaundice (10.2%), G6PD deficiency (4.2%) and minor blood group

Comment [UP3]: What is inconclusive jaundice? This should be defined or explained.

111 incompatibility (1.7%). There was one case of cephalhematoma. There were no cases of
 112 non-immune haemolysis and polycythemia in this study.



113 Figure 1: Etiologies of Jaundice

114 Table 2: Characteristic maternal profile according to the etiology of jaundice

Maternal profile	No.	Blood group incompatibility		G6PD def. (n=5)	Breast feeding (n=12)	Cephal-hematoma (n=1)	Inconclusive (n=76)
		ABO (n=21)	Minor (n=2)				
Maternal age (n=113)							
15-20 years	26	5	0	0	1	0	20
21-35 years	76	15	2	5	7	0	47
36-45 years	11	1	0	0	4	0	6
Maternal complication (n=116)							
CPD	18	5	0	1	1	0	11
Hypertension	2	0	0	0	0	0	2
Diabetic	1	0	0	0	1	0	0
Hypertension and diabetic	1	0	0	0	0	0	1
No complication	94	16	2	3	10	1	62
Oxytocin using (n=115)							
Yes	3	0	0	0	0	0	3
No	112	21	2	5	11	1	72

Any medication during pregnancy (n=115)							
Yes	2	0	0	0	1	0	1
No	113	21	2	4	11	1	74
Previous neonatal jaundice history (n=67)							
Yes	5	0	0	0	1	0	4
No	62	13	0	4	5	1	39

117

118 | Table 2 shows that 47/76 cases (62%) were born from mothers aged between 21 to 35
 119 | years were diagnosed as inconclusive jaundice. In this study, oxytocin was used in 3
 120 | mothers and diagnoses of their infants were inconclusive jaundice. Among 5 cases that had
 121 | previous history of neonatal jaundice, 1 case was breast feeding jaundice and other 4 cases
 122 | were inconclusive jaundice. Other findings such as bruise/petechiae, hepatosplenomegaly
 123 | were not observed in this study.

124

125

126

Table 3: Maximum bilirubin level according to the etiology of jaundice

Maximum bilirubin level	No.	Blood group incompatibility		G6PD def. (n=5)	Breast feeding (n=12)	Cephal-hematoma (n=1)	Inconclusive (n=76)
		ABO (n=21)	Minor (n=2)				
Maximum microbilirubin level (mg/dL)							
<12	5	0	0	0	1	0	4
12-25	106	21	2	5	11	1	66
>25	6	0	0	0	0	0	6
Maximum unconjugated bilirubin level (mg/dL)							
<12	10	1	0	0	1	0	8
12-25	103	20	2	5	11	1	64
>25	4	0	0	0	0	0	4

127

128

129

130

131

Most of the cases had the maximum microbilirubin level in range 12-25 mg/dL which a correlates to with the result of maximum unconjugated bilirubin level significantly with Pearson coefficient level at 0.699. All cases that had bilirubin level above 25 were in inconclusive etiology. (Table 3)

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

Table 4 shows that the mean values of CBC profile and bilirubin levels according to etiology of hyperbilirubinaemia were demonstrated. Among mean value of maximum mean hematocrit (58.3%) was highest seen in babies with breast feeding jaundice. In all diagnosis, mean value of WBC and platelet count were within normal limit. NRC was found in only ABO incompatibility and inconclusive jaundice and higher mean of NRC in ABO incompatibility (131.5%). Mean of reticulocyte count percentage was 13.6 in minor blood group incompatibility which was the highest in all etiologies. Mean value of conjugated bilirubin was highest (0.7 mg/dL) in ABO incompatibility while as unconjugated bilirubin level was highest in minor blood group incompatibility (17.6 mg/dL). The diagnosis of G6PD deficiency had the highest maximum value of microbilirubin (20.5 mg/dL) among other causes of neonatal jaundice. Only one blood film had hypochromic picture in which and this was in the inconclusive diagnosis group. Almost all patients had normal size of red blood cells. Anisocytosis accounted for 11.9% and microcytes were 3.4%. On the other hand, only 11% of newborns had normal shape of red blood cells. The rest 89% of them had abnormal shape such as poikilocytosis, burr cells, target cells, spherocyte, ovalocyte, schistocyte and polychromasia.

148 As shown in Table 5, ~~account for all diagnosis~~, onset of jaundice was most commonly
 149 mostly started on the 2nd day after delivery (n=49; 41.9%). It was followed by the 3rd day after
 150 birth (n=35; 29.9%) and the 4th day (n=12; 10.3%). Early onsets of jaundice (within 24 hours
 151 after birth) were seen in 10 cases (8.5%) and most of them (70%) were inconclusive
 152 jaundice. One case of breast feeding jaundice and 5 cases of inconclusive jaundices
 153 presented with late onset of jaundice on the 6th and 7th day of birth. According to onset of
 154 jaundice, treatment was started mostly on 2nd day after birth. Duration of phototherapy was
 155 ranged from 1 to 4 days. There were four cases that needed exchange transfusion therapy
 156 for rescue. One case was ABO incompatibility with high onset of jaundice on the second day
 157 of life, microbilirubin level was 16.4 mg/dL and 14.9% reticulocyte count was 14.9%
 158 observed. Three cases were inconclusive jaundice with high onset of jaundice on the 2nd, the
 159 4th and the 5th day and the microbilirubin levels were 38.5, 32.9 and 31.1 mg/dL respectively,
 160 without evidence of haemolysis. ~~According to ABO blood group examination of the mothers~~
 161 ~~and the babies who were diagnosed as~~ For ABO incompatibility (n=11), all had blood group
 162 O mothers and blood group A and group B in newborn were 10 (47.6%) and 11 (52.4%)
 163 respectively.

164 Table 6 shows that the mean value of maximum and minimum haematocrit was
 165 significantly lower in haemolysis group than in non-haemolysis group significantly (P= 0.013
 166 and P<0.001). NRC count and percent of reticulocyte counts were higher in haemolysis
 167 group than in non-haemolysis group and the difference was with statistically significant.
 168 There were four cases that needed exchange transfusion therapy for rescue. One case was
 169 ABO incompatibility with high onset of jaundice on the second day of life, and microbilirubin
 170 level as 16.4 mg/dL on the 1st day after delivery that compatible with pathological jaundice
 171 and reticulocyte count was also increased (14.9% even on 1st day). Three cases were
 172 inconclusive jaundice with high onset of jaundice on the 2nd, the 4th and the 5th day and the
 173 microbilirubin level are 38.5, 32.9 and 31.1 mg/dL respectively, without evidence of
 174 haemolysis.

175
 176 Table 4: Laboratory profile according to the etiology of jaundice

Laboratory profile (Mean; Range; SD)	Blood group incompatibility		G6PD def. (n=5)	Breast feeding (n=12)	Inconclusive (n=76)
	ABO (n=21)	Minor (n=2)			
CBC profile					
Maximum of Hct (%)	53.1	54.5	58.2	58.3	56.4
	38-68	48-61	52-70	50-66	32-70
	(7.3)	(9.2)	(7.2)	(5.9)	(6.7)
Minimum of Hct (%)	43.9	47.5	51.6	51.1	49.9
	31-57	38-57	47-63	38-60	30-62
	(7.4)	(13.4)	(6.5)	(6.6)	(7.1)
WBC count (X10 ³ /mm ³)	14.4	14.8	11.6	13.3	15.9
	6.9-24.5	13.9-15.7	7.8-20.5	9.1-20.5	6.0-17.3
	(4.7)	(1.3)	(5.1)	(3.6)	(23.3)
Platelet count (X10 ³ /mm ³)	291.0	345.5	234.8	236.3	275.8
	145-494	314-377	165-357	66-398	89-599
	(93.6)	(44.5)	(89.9)	(98.7)	(84.2)
NRC (/mm ³)	131.5	0	0	0	9.9
	0-1470				0-232
	(321.4)				(40.5)
% Reticulocyte count	8.6	13.6	4.6	7.1	6.0
	3-14.9	3.5-23.6	2.8-8.0	0.7-13.4	0-15.1
	(3.8)	(14.2)	(2.4)	(4.1)	(3.4)

Comment [UP4]: How possible is this? If there were 11 mothers, the total number of babies with A and B blood groups should also be 11.

Comment [UP5]: Write in full before the abbreviation.

Comment [UP6]: This is repeating lines 152-155 above.

Bilirubin					
Conjugated (mg/dL)	0.7 0.2-6.0 (1.2)	0.3 0.3-0.4 (0.0)	0.4 0.2-0.4 (0.1)	0.4 0.2-0.7 (0.2)	0.5 0.0-10.6 (1.2)
Unconjugated (mg/dL)	16.0 11.9-21.4 (2.6)	17.6 16.4-18.8 (1.7)	16.6 12.4-21.1 (3.7)	15.6 9.4-19.1 (2.8)	16.8 7.3-37.8 (4.9)
Maximum of MB	17.4 12.4-24.1 (2.8)	18.1 17.5-18.7 (0.8)	20.5 17.7-22.2 (1.9)	17.1 10.3-21.5 (3.0)	18.3 8.9-38.5 (4.9)

177
178
179

Table 5: Onset and treatment of neonatal jaundice

Characteristic of jaundice	No.	Blood group incompatibility		G6PD def. (n=5)	Breast feeding (n=12)	Cephal-hematoma (n=1)	Inconclusive (n=76)
		ABO (n=21)	Minor (n=2)				
ONSET OF JAUNDICE							
1 st day of birth	10	2	0	0	1	0	7
2 nd day of birth	49	14	1	4	6	0	24
3 rd day of birth	35	5	0	0	3	1	26
4 th day of birth	12	0	1	1	1	0	9
5 th day of birth	5	0	0	0	0	0	5
6 th day of birth	1	0	0	0	0	0	1
7 th day of birth	5	0	0	0	1	0	4
TREATMENT							
Day of start therapy							
1 st day of birth	10	2	0	0	1	0	7
2 nd day of birth	51	15	2	4	6	0	24
3 rd day of birth	32	3	0	0	3	0	26
4 th day of birth	11	0	0	1	1	0	9
5 th day of birth	4	0	0	0	0	0	4
6 th day of birth	3	1	0	0	0	0	2
7 th day of birth	4	0	0	0	1	0	3
8 th day of birth	2	0	0	0	0	1	1
Phototherapy duration (Days) (Mean; Range; SD)							
Single phototherapy	1.4	1		1.2	1.3	1	1.1
	0-2			1-2	0-3		0-4
	(0.6)			(0.4)	(0.9)		(0.7)
Double phototherapy	0.5	1		0.8	0.3	0	0.5
	0-2	0-2		0-1	0-1		0-2
	(0.7)	(1.4)		(0.4)	(0.5)		(0.7)
All type	1.9	2.0		2.0	1.7	1	1.6
	1-4	1-3		1-3	1-4		1-6

		(0.7)	(1.4)	(0.7)	(1.0)		(0.9)
Exchange transfusion	4	1	0	0	0	0	3

Table 6: Laboratory profile according to etiology of jaundice

Laboratory	Haemolysis (n=24)	Non haemolysis (n=93)	P value
CBC profile			
Maximum of Hct (%)	53.0 38-68 (7.2)	56.8 32-70 (6.5)	0.013
Minimum of Hct (%)	44.2 31-57 (7.7)	50.1 30-63 (6.9)	<0.001
WBC count (X10 ³ /mm ³)	14.5 6.9-24.5 (4.4)	15.3 6.0-17.3 (21.2)	>0.05
Platelet count (X10 ³ /mm ³)	304.1 145-497 (98.1)	265.4 66-599 (83.4)	>0.05
NRC count (/mm ³)	115.0 0-1470 (303.0)	8.1 0-232 (36.8)	0.001
% Reticulocyte count	8.7 3.0-23.6 (4.9)	6.0 0-15.1 (3.5)	0.003
4. Bilirubin			
Conjugated (m/dL)	0.7 0.16-6.0 (1.2)	0.5 0-10.6 (1.1)	>0.05
Unconjugated (m/dL)	16.4 11.9-21.4 (2.7)	16.6 7.3-37.8 (4.6)	>0.05
Maximum of MB	17.6 12.4-24.1 (2.7)	18.2 8.9-38.5 (4.6)	>0.05

DISCUSSION

This study was conducted to describe the various clinical profiles of early neonatal jaundice. Careful history taking, physical examinations, early measurement of serum bilirubin and work up for jaundice to elicit information on risk factors and different etiologies. According to exclusion criteria of current study, ~~this study fails to illustrate the preterm babies neonatal with neonatal jaundice in preterm as the previous study which showed jaundice occurred in 60% of term infants and 80% of preterm infants were not studied.~~

Comment [UP7]: Not a complete sentence so the meaning is not clear

206 It is crucial to assess characteristic maternal and neonatal profile during hospital stay and
207 on discharge for risk of neonatal jaundice [16].

208 In our study, there was no difference between haemolysis and non-haemolysis group
209 according to spontaneous delivery which was similar to the previous study finding [17].
210 Increased frequency of jaundice is obviously associated with maternal usage of epidural
211 anaesthesia. We found that the minimum haematocrit was significantly lower and
212 reticulocyte count is significantly higher in haemolytic group compared to non-haemolytic
213 group which were compatible with pathogenesis of hyperbilirubinemia. [18].

214 According to etiologies and maternal profile, nearly 50% of ABO incompatibility are
215 primigravida in this study which is similar to the previous study establishing that
216 approximately 50% of the ABO haemolytic jaundice cases occur in first born infants and
217 there is no predictable pattern of recurrence in subsequent infants [19]. Rh haemolytic
218 disease and sepsis are etiologies that have increased risk of bilirubin encephalopathy than
219 ABO incompatibility [20]. Previous research showed inconclusive jaundice was the most
220 common etiology as demonstrated in our study.

221 Theoretically, G6PD deficiency usually occurs in males although heterozygous females
222 may manifest the mild features of disease [21]. In our study, newborns who diagnosed as
223 G6PD deficiency were all males (4.2%). ~~which female with G6PD deficiency could not~~
224 ~~demonstrate.~~

225 Infection ~~was~~ one of the risk factors of hyperbilirubinemia [22]. Unexplained
226 unconjugated hyperbilirubinemia may be a first ~~sign (is it sing or sign)~~ of neonatal sepsis as
227 bacterial sepsis can contribute to neonatal jaundice [23]. Our study ~~could did not~~
228 demonstrate ~~the a~~ higher WBC count in ~~the~~ non-haemolytic group compared to ~~the~~
229 haemolytic group. One of the possible explanations was ~~not only infections but also that~~
230 unidentified non-infectious etiologies ~~stands as grounds may also play a significant role in~~
231 ~~the aetiology for of~~ non-haemolytic ~~jaundice group.~~

232 Measuring serum bilirubin concentrations exactly 24 hours after delivery or at a later time
233 just before discharge may improve the sensitivity of this predictor for hyperbilirubinemia [24].
234 Predictive bilirubin nomogram is an effective predictor for significant hyperbilirubinemia
235 according to their risks: high, intermediate and low [25].

236 Among four cases ~~preceeded for whom~~ exchange transfusion ~~was done~~, one case was
237 ABO incompatibility and the rest three cases were inconclusive jaundice. Three cases of
238 inconclusive jaundice ~~denied had no~~ evidence of haemolysis ~~in which and~~ onset of jaundice
239 ~~are were~~ on 2nd, 4th and 5th day ~~and while~~ the microbilirubin levels ~~are were~~ 38.5, 32.9 and
240 31.1 mg/dL respectively. All neonatal jaundice with high bilirubin levels ~~is~~ would ~~be treated~~
241 ~~with require~~ aggressive treatment (such as exchange transfusion and double ~~side~~
242 phototherapy) to prevent complications of hyperbilirubinemia [26]. In newborn with bilirubin
243 level above 20 mg/dL, there is noticeable association with kernicterus which has 70% long
244 term consequences [27].

245

246 5. CONCLUSION

247

248 ~~As stated by our study, various etiologies of hyperbilirubinemia were found. Our study has~~
249 ~~shown that~~ the most common etiology was inconclusive jaundice ~~closely followed by~~
250 ~~because it was only rely on medical record. As we know before, the second common~~
251 ~~etiology was abo~~ ABO incompatibility. Neonatal jaundice caused by non-immune hemolysis
252 and polycythemia were not found in this study. Supporting neonatal sepsis and infections are
253 main concern for neonatal jaundice, wbc count was highest in the inconclusive jaundice. In
254 all diagnosis, mean values of platelet count were within normal limit. Mean value of
255 reticulocyte count was highest in the diagnosis of minor blood group incompatibility. There
256 was significant difference of haematocrit, NRC and reticulocytes between haemolytic and
257 non-haemolytic groups. As the study was the hospital based retrospective record study, the
258 subjective data such as clinical features was not so gained as expected. To retrieve the

259 comprehensive data, the prospective study should be suggested. Nevertheless, the study
260 highlighted the etiologies of neonatal jaundice of study group, demographic data, maternal
261 data and also laboratory profiles, onset and duration of treatments. **(Please the authors**
262 **should give deductive and logical conclusion that has contribution to knowledge)**
263
264

265 **COMPETING INTERESTS**

266
267 Authors have declared that no competing interests exist.
268

269 **CONSENT**

270
271 NA
272

273 **ETHICAL APPROVAL**

274
275 This research was approved_ and funded by the Faculty of Tropical Medicine, Mahidol
276 University, Thailand.
277

278 **REFERENCES**

- 279
280 1. Mitra S, Rennie J. Neonatal jaundice: aetiology, diagnosis and treatment. *Br J Hosp*
281 *Med (Lond)*. 2017; 2; 78(12):699-704. doi: 10.12968/hmed.2017.78.12.699.
- 282 2. Brown AK. Bilirubin metabolism with special reference to neonatal jaundice. *Adv*
283 *Pediatr* 1962;-12:-121-87.
- 284 3. Watchko JF, Oksi FA. Bilirubin 20 mg/dl=vagintiphobia. *Pediatr*. 1983; 71: 660-3.
- 285 4. Olusanya BO, Teeple S, Kassebaum NJ. The Contribution of Neonatal Jaundice to
286 Global Child Mortality: Findings from the GBD 2016 Study. *Paediatrics*. 2018;
287 141(2). pii: e20171471. doi: 10.1542/peds.2017-1471
- 288 5. Lalitha KG. Neonatal jaundice. In: Ghai OP, Gupta P, Paul VK, editors. *Ghai*
289 *Essential Paediatrics*. 5th ed. New Delhi: Interprint; 1993.
- 290 6. Newman AJ, Gross S. Hyperbilirubinemia in breast fed infants. *Pediatr* 1983;
291 32:995-1000.
- 292 7. Newman TB, Klebanoff MA. Neonatal hyperbilirubinemia and long term outcome.
293 *Pediatr*. 1993; 92(5):651-7.
- 294 8. Phyllis AD, Daniel SS, David KS. Neonatal hyperbilirubinemia. *N Eng J Med*. 2001;
295 344(8): 211.
- 296 9. Campbell N, Harvey D, Norman AP. Increased frequency of neonatal jaundice in a
297 maternity hospital. *Br Med J*. 1976; 1:548-52.
- 298 10. CDC-MMWR (Morbidity and Mortality Weekly Report). Kernicterus in full term
299 infants- United States, 1994- 1998. 2001;-50(23):-491-4.
- 300 11. Anthony JP, Barbara JS. Jaundice and hyperbilirubinemia in newborn. In: Robert
301 MK, Richard EB, Hal BJ, editors. *Nelson Textbook of Pediatrics*.18th ed. London:
302 WB Saunders; 2007.
- 303 12. Chuniaud L, Dessante M, Chantoux F, Blondeau JP, Francon J, Trivin F.
304 Cytotoxicity of bilirubin for human fibroblasts and rat astrocytes in culture. *Clin Chim*
305 *Acta*. 2006; 256(2):103-14.
- 306 13. Bratlid D. How bilirubin gets into the brain. *Clin Perinatol*. 1990;-17(2):449-65.
- 307 14. Pamela GL. Jaundice in the Newborn. In: Ronald MP, James DS, Dale AN, editors.
308 *Paediatric hospital medicine, Textbook of inpatient management*. Philadelphia:
309 Lippincott Williams & Wilkins; 2003.

- 310 15. Kuzniewicz MW, Wickremasinghe AC, Wu YW, McCulloch CE, Walsh EM, Wi S, et
311 al. Incidence, Etiology, and Outcomes of Hazardous Hyperbilirubinemia in
312 Newborns. *Paediatrics*. 2014; 134 (3). doi:10.1542/peds.2014-0987
313 16. Maisels MJ, Kring E. Length of stay, jaundice and hospital readmission. *Pediatric*.
314 1998; 101(6): 995-8.
315 17. Sarici SU, Yurdakok M, Serdar MA, Oran O, Erdem G, Tekinalp G, et al. An early
316 (sixth hour) serum bilirubin measurement is useful in predicting the development of
317 significant hyperbilirubinemia and severe ABO haemolytic disease in a selective
318 high risk population of newborns with ABO incompatibility. *Paediatric*. 2002; 109(4);
319 53.
320 18. David E. Neonatal Jaundice. *BMJ Clin Evid*. 2007; 12:319 -28.
321 19. Hinkes MT, Cloharty JP. Neonatal hyperbilirubinemia. In: Cloharty JP, Stork AR,
322 editors. *Manual of neonatal care*. 5th ed. Philadelphia: Lippincott Williams & Wilkins;
323 1998:175-211.
324 20. Gamaleldin R, Iskander I, Seoud I, Aboraya H, Aravkin A, Sampson PD, et al. Risk
325 Factors for Neurotoxicity in Newborns With Severe Neonatal Hyperbilirubinemia.
326 *Paediatrics*. 2011; 128(4): e925–e931. doi: [10.1542/peds.2011-0206]
327 21. Malcolm IL, David IT, Sunil S. Jaundice. In: Malcolm IL, David IT, Sunil S. editors,
328 *Essential Neonatal Medicine* 4th ed. Oxford: Blackwell Publishing; 2008: 130-41.
329 22. Rennie JM, Robertson NRC. Physiological jaundice. In: Rennie JM, Robertson NRC,
330 editors. *A manual of neonatal intensive care*. 4th ed. London, 2002: 419.
331 23. Lindar N, Yatsiv I, Tsur M, Matoth I. Unexplained neonatal jaundice as an early
332 diagnostic sign of septicemia in the newborn. *Journal of Perinatology*. 8(4):325-7.
333 24. Seidman DS, Shaltiel ZE, Paz I, Gale R. Predicting the Risk of Jaundice in full term
334 Healthy Newborns: A Prospective Population-Based Study. *Journal of Perinatology*.
335 1999; 19(8): 564-7.
336 25. Bhutani VK, Johnson L, Sivieri EM. Predictive ability of a pre-discharge hour-specific
337 serum bilirubin for subsequent significant hyperbilirubinemia in healthy term and
338 near-term newborns. *Paediatrics*. 1999; 103(1):6-14.
339 26. Tan KL. Neonatal Jaundice. In: Robinson MJ and Lee EL editors. *Paediatric*
340 *Problems in Tropical Countries*. 2nd ed. London. Dr. K C Chaudhuri Foundation;
341 1983: 91-8.
342 27. Ip S, Chung M, Kulig J, O'Brien R, Sege R, Glicken S, et al. An evidence-based
343 review of important issues concerning neonatal hyperbilirubinemia. *Paediatrics*.
344 2004; 114(1):e130-53.PMID: 15231986.

345

346 **DEFINITIONS, ACRONYMS, ABBREVIATIONS**

347 Here is the Definitions section. This is an optional section.

348 **Term:** Definition for the term