

1 **European Resuscitation Council Guidelines 2025: Newborn resuscitation and support of transition of**  
2 **infants at birth**

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51 **[h1] Abstract**

52 This European Resuscitation Council (ERC) Guideline 2025 on Newborn Life Support is based on the  
53 International Liaison Committee on Resuscitation (ILCOR) Consensus on Science and Treatment  
54 Recommendations (CoSTRs) for Neonatal Life Support. The guideline presents a logical approach to  
55 resuscitation and support of transition to extra-uterine life, for both preterm and term newborn  
56 infant. The guideline includes factors before birth, training and education, thermal control,  
57 management of the umbilical cord after birth, initial assessment, airway, breathing and circulation  
58 assessment and interventions, emergency vascular access, low resource and out of hospital settings,  
59 communication with parents, and considerations on withholding and discontinuing life sustaining  
60 treatments. Life support guidelines for older infants and children are covered in the ERC Guidelines  
61 2025 on Paediatric Life Support.

62 **Abbreviations:**

63	CPR	Cardiopulmonary resuscitation
64	C:V	Compression to ventilation (ratio)
65	CI	Confidence intervals
66	CoSTR	Consensus on Science and Treatment Recommendations
67	CPAP	Continuous positive airway pressure
68	DCC	Delayed cord clamping
69	ECG	Electrocardiography
70	FRC	Functional residual capacity
71	GA	Gestational age
72	HCP	Health care professional
73	HR	Heart rate
74	HIE	Hypoxic ischemic encephalopathy
75	ICC	Immediate cord clamping
76	IO	Intraosseous
77	IV	Intravenous or intravascular
78	NICU	Neonatal Intensive Care Unit
79	NLS	Newborn Life Support
80	FiO <sub>2</sub>	Fractional inspired oxygen concentration
81	OR	Odds ratio
82	PLS	Paediatric Life Support
83	SpO <sub>2</sub>	Peripheral oxygen saturation
84	PEEP	Positive end-expired pressure
85	PPV	Positive pressure ventilation
86	RCT	Randomized controlled trial
87	ROSC	Return of spontaneous circulation
88	s	Second(s)
89	SGA	Supraglottic airway device
90	UVC	Umbilical venous catheter
91		

## 92 [h1] Introduction and scope

93 The ERC Guidelines 2025 on Newborn Life Support (NLS) includes both resuscitation at birth and  
94 support of transition from fetus to newborn infant across all gestational ages (GA). Newborn  
95 resuscitation differs fundamentally from resuscitation in every other age group due to the unique  
96 physiological transition from intrauterine to extrauterine life. The adaptation at birth requires a  
97 complex interplay of respiratory, cardiovascular, and metabolic changes, making timely and  
98 appropriate intervention crucial. It primarily focuses on supporting postnatal transition, with  
99 establishment of lung aeration, effective breathing and optimising pulmonary blood flow.

100 The evidence to support newborn resuscitation remains limited, with many recommendations  
101 extrapolated from animal studies, observational data, or expert consensus. The ERC NLS Writing Group  
102 recognises these challenges but has aimed to develop clear, evidence-informed guidelines that balance  
103 scientific rigor with practical applicability. By emphasising consistency, simplicity, and effective  
104 training, they serve as a foundation for improving newborn resuscitation practices across diverse  
105 healthcare environments.

106 The ERC Guidelines 2025 NLS are based on the International Liaison Committee on Resuscitation  
107 (ILCOR) Consensus on Science and Treatment Recommendations (CoSTRs) for Newborn Life Support  
108 (ILCOR 2025). For the purposes of this Guideline, the ILCOR recommendations were supplemented by  
109 focused literature reviews undertaken by the ERC NLS Writing Group for topics not reviewed by LCOR  
110 CoSTRs. When required, the guidelines were informed by expert consensus of the ERC NLS Writing  
111 Group. The ERC Guidelines 2025 NLS were drafted and agreed by the ERC NLS Writing Group members  
112 and the ERC Guidelines 2025 Steering Committee. This guideline was posted for public comment in  
113 May 2025. A total of [INSERT NUMBER] individuals from [INSERT COUNTRIES] submitted [INSERT  
114 NUMBER] comments, leading to [INSERT CHANGES] in the final version. Subsequently, the feedback  
115 was reviewed by the NLS writing group, and the guideline was thereafter updated where relevant. The  
116 ERC Guidelines 2025 NLS was presented to and approved by the ERC Board and the ERC General  
117 Assembly on xy June 2025. The methodology used for guideline development is presented in the  
118 Executive summary. (REF)

119 For consistency, the ERC Guidelines 2025 NLS describes a baby at birth as a 'newborn infant' and a  
120 baby or a neonate as an 'infant' throughout this guideline. The term 'parents' is used to describe the  
121 mother and partner/caregivers.

## 123 [h2] Newborn Life Support or Paediatric Life Support?

124 In agreement with the ERC Guidelines 2025 Paediatric Life Support (PLS) Writing Group, the ERC  
125 recommends the following:

- 126 • Use the ERC Guidelines 2025 NLS immediately after birth irrespective of birth location (i.e.  
127 hospital or home birth)
- 128 • The ERC Guidelines 2025 NLS can also be used during Neonatal Intensive Care Unit (NICU) stay,  
129 especially in preterm infants or term infants with primary respiratory problems.
- 130 • Use the ERC Guidelines 2025 PLS after first hospital discharge.
- 131 • Using the ERC Guidelines 2025 PLS during first hospital stay after birth is also reasonable in the  
132 following circumstances:
- 133 ○ after cardiac surgery,  
134 ○ in known cardiac arrhythmia, and  
135 ○ In other non-respiratory cardiac arrests
- 136 • Develop local policies defining which guideline to use for which infants, applicable to the  
137 healthcare setting. Factors to take into consideration include:
- 138 ○ individual NICU case-mix,  
139 ○ algorithm familiarity and training, and  
140 ○ human and organisational factors
- 141 • Teams may initiate resuscitation using the guideline they are most familiar with (NLS or PLS) and  
142 summon appropriate help and switch guideline if needed, in a timely and coordinated manner.

143

144 **[h2] Preterm infants at the threshold of viability**

145 The ERC Guidelines 2025 NLS applies predominantly to management of infants with gestational age (GA)  
146  $\geq 25$  weeks. Until more evidence from trials including the most preterm infants is available, the ERC NLS  
147 Writing Group recommends caution when applying the recommendations in this guideline to them. (1)

148

149 **[h2] Standardised gestational age cut-off across the guideline**

150 To ensure consistency and practical applicability, the ERC NLS Writing Group standardised the  
151 gestational age (GA) cut-off across all subtopics. Although many ILCOR reviews on preterm infants focus  
152 on infants  $< 34$  weeks, most studies include predominantly infants  $< 32$  weeks, therefore 32 weeks was  
153 adopted as a pragmatic cut-off. This aligns with the ERC Guidelines 2021 NLS and common clinical  
154 thresholds for determining the appropriate level of perinatal care.

155

156 **[h2] Key messages and key changes**

157 Key messages are presented in figure 1. Summary of key changes is presented in table 1.

158 <<INSERT Figure 1 Key messages infographic >>

- 159 << INSERT Table 1 Key changes >>
- 160 << INSERT Figure 2 NLS algorithm >>

This is a DRAFT-version

161 **[h1] Concise guideline for clinical practice**

162 **[h2] Factors before birth**

163 << INSERT Figure 3 Common factors associated with an increased risk of a need for stabilisation, or  
164 resuscitation at birth >>

165 **[h3] Staff attending births in hospitals**

166 Any infant may develop problems during birth. Local guidelines should indicate who should attend  
167 births taking into consideration identified risk factors (figure 3).

168 As a guide:

- 169 • An interprofessional team with appropriate experience and training in NLS proportionate to  
170 the expected risk should attend the birth.
- 171 • Neonatal staffing levels should acknowledge the potential need to deliver unexpected support  
172 in the delivery room.
- 173 • A process should be in place for rapidly mobilising extra team members with adequate  
174 resuscitation skills for any birth.

175 **[h3] Telemedicine**

- 176 • Consider the use of collaboration through telemedicine, as it facilitates providing remote advice.

177 **[h3] Equipment and environment**

- 178 • Regularly check all equipment to ensure it is ready for use.
- 179 • Ensure that equipment is easily accessible and organised in a standardised way.
- 180 • Consider human factor elements when organising equipment to maximise efficiency and  
181 minimise time delays-
- 182 • Resuscitation should take place in a warm, well-illuminated, draught-free area with a flat  
183 resuscitation surface and a radiant heater (see thermal control).

184 **[h3] Briefing**

- 185 • Team briefing is important and should be performed before birth.
- 186 • The purpose of briefing is to:
  - 187 ○ Review available clinical information
  - 188 ○ Assign roles and tasks
  - 189 ○ Check equipment and presence of personnel
  - 190 ○ Prepare the family
  - 191 ○ Use a checklist and/or cognitive aid to facilitate all of the above, reduce mental load, and  
192 enhance safety

193

**194 [h2] Education**

- 195 • Institutions or clinical areas where births may occur should provide sufficient opportunities and  
196 resources for healthcare professionals involved in neonatal resuscitation to receive regular  
197 training, maintaining up-to-date knowledge as well as technical and non-technical skills.
- 198 • The content and organisation of such training programmes may vary according to the needs of  
199 the providers and the local organisation.
- 200 • Undertake training at least once per year to prevent skill decay, preferably supplemented with  
201 more frequent short-duration booster sessions (e.g. every 3–6 months). For more information  
202 on training see ERC Guidelines 2025 Education for Resuscitation.

203

**204 [h2] Thermal Control****205 [h3] Standards**

- 206 • Maintain the temperature of newborn infants between 36.5°C and 37.5°C.
- 207 • Monitor the infant's temperature regularly or continuously after birth.
- 208 • Record the admission temperature as a prognostic and quality indicator.
- 209 • Rewarm infants who are hypothermic after birth; avoid hyperthermia.
- 210 • In appropriate circumstances, therapeutic hypothermia may be considered after resuscitation  
211 (see post-resuscitation care).

**212 [h3] Environment**

- 213 • Protect the infant from draughts. Ensure windows are closed and air-conditioning  
214 appropriately programmed.
- 215 • In infants >28 weeks, keep the delivery area at 23-25°C.
- 216 • In infants ≤28 weeks, keep the delivery area at >25°C.

**217 [h3] Newborn infants ≥32 weeks**

- 218 • Dry the infant immediately after birth and remove wet towels.
- 219 • Cover the head with a hat, and the body with dry towels.
- 220 • If no intervention is required, place the infant skin-to-skin with mother and cover both with  
221 towels.
- 222 • Ongoing careful observation of mother and infant is required, especially in more preterm and  
223 growth restricted infants to ensure they both remain normothermic.
- 224 • Consider the use of a plastic bag/wrap if skin-to-skin care is not possible.
- 225 • Place the infant on a warm surface using a preheated radiant warmer, if support of transition  
226 or resuscitation is required.

227 **[h3] Newborn infants <32 weeks**

- 228 • Dry the infant's head and cover with a hat.
- 229 • Put the infant's body in a plastic (polyethylene) bag or wrap without drying.
- 230 • Use a preheated radiant warmer.
- 231 • Consider the use of additional measures during delayed cord clamping to ensure thermal
- 232 stability (e.g., increasing room temperature, warm blankets, and thermal mattress).
- 233 • Be careful to prevent hypothermia during skin-to-skin care during assisted transition,
- 234 especially in the more preterm and/or growth restricted infants.
- 235 • Consider the use of heated humidified respiratory gases for infants receiving respiratory
- 236 support.
- 237 • Be aware of the risk of hyperthermia when multiple heat-preservation interventions are used
- 238 simultaneously, especially while using a thermal mattress.

239

240 **[h2] Management of the umbilical cord**

- 241 • Ideally, delayed cord clamping is performed in all births, after inflation of the lungs and before
- 242 uterotonics are given.

243 **[h3] Cord clamping**

- 244 • Discuss the options for managing cord clamping and the rationale with parents and
- 245 obstetricians before birth.
- 246 • Perform thermal management, tactile stimulation and initial assessment during delayed cord
- 247 clamping.
- 248 • Newborn infants without need for support: facilitate at least 60s of delayed cord clamping.
- 249 • Newborn infants in need of resuscitation: clamp the cord <30s to minimise any delay to
- 250 necessary interventions.
- 251 • If stabilisation with intact cord can be safely undertaken, longer delayed cord clamping is
- 252 preferred, especially in infants <34 weeks.

253 **[h3] Cord milking**

- 254 • Do **not** milk the cord in preterm infants <28 weeks.
- 255 • Consider intact cord milking as an alternative in infants ≥28 weeks, but only if delayed cord
- 256 clamping cannot be performed.

257

258 **[h2] Initial assessment**

- 259 • Perform initial assessment as soon as possible after birth, ideally during delayed cord  
260 clamping, drying and wrapping to:
- 261 ○ Identify the need for support and/or resuscitation
- 262 ○ Aid decisions relating to the appropriateness and duration of delayed cord clamping.
- 263 • Assess:
- 264 ○ Breathing
- 265 ○ Heart rate (HR)
- 266 ○ Muscle tone
- 267 • Provide thermal management and tactile stimulation during delayed cord clamping and  
268 assessment.
- 269 • Reassess breathing and HR frequently to assess any response and determine if further  
270 interventions are required.
- 271 **[h3] Breathing**
- 272 • Note the rate, depth, symmetry and work of breathing.
- 273 **[h3] Heart rate**
- 274 • Initial HR assessment can be performed with a stethoscope.
- 275 • Continuous HR assessment methods (pulse oximetry, Electrocardiography (ECG)) are preferred  
276 when interventions are indicated or during stabilisation of preterm newborns.
- 277 << INSERT Table 2 Assessment of breathing and heart rate >>
- 278 **[h3] Response to tactile stimulation**
- 279 • Gently stimulate the newborn infant by drying them, rubbing the soles of the feet or their  
280 back.
- 281 • Avoid more vigorous methods of stimulation, especially in preterm infants.
- 282
- 283 **[h3] Muscle tone & Colour**
- 284 • A very floppy infant is likely to need breathing support.
- 285 • Hypotonia is common in preterm infants.
- 286 • Do not use colour to assess oxygenation.
- 287 • Interpret pallor within clinical context, as it may have several causes such as blood loss,  
288 chronic anaemia, or asphyxia.
- 289 **[h3] Classification according to initial assessment**
- 290 • Based on the initial assessment, further actions can be implemented guided by the NLS  
291 algorithm (Figure 2). These are summarised in Figure 4.

292 << INSERT Figure 4 Schematic illustration of initial assessment and interventions >>

293

## 294 [h2] Newborn life support

- 295 • Ensure the **airway is open** and the **lungs are inflated**.
- 296 • Do not undertake subsequent interventions before the airway is open and the lungs have been  
297 inflated.
- 298 • Following initial assessment, start respiratory support if the infant is not breathing regularly or  
299 the HR is  $<100 \text{ min}^{-1}$ .

300

## 301 [h2] Airway

- 302 • Assess the effect of each airway technique by observing for chest movement and assessing HR.

### 303 [h3] Position

- 304 • Place the newborn infant on their back with the head supported in a neutral position (figure  
305 5).

306 << INSERT Figure 5 Head in a neutral position >>

- 307 • Gently pull the jaw forwards (jaw thrust) to open the airway (figure 6).

308 << INSERT Figure 6 Jaw thrust, lifting the lower jaw forwards enlarges the pharyngeal space >>

### 309 [h3] Two-person method

- 310 • Use the two-person method of airway support (jaw thrust) as this approach is more effective  
311 than single person jaw thrust.

### 312 [h3] Suction

- 313 • Do not routinely suction meconium from infant's' airways because it delays initiating  
314 ventilation.
- 315 • Consider physical airway obstruction if lung inflation is unsuccessful despite alternative airway  
316 opening techniques.
- 317 • Perform suction under direct vision.
- 318 • Rarely, with no response to inflations and no chest wall movement, an infant may require  
319 tracheal suctioning to relieve an airway obstruction below the vocal cords.

### 320 [h3] Airway devices

- 321 • Use airway devices only if competent personnel are available and trained in the appropriate  
322 equipment; if not continue with mask ventilation and call for help.

### 323 [h4] Supraglottic airway devices

324 Consider using an appropriate size supraglottic airway device (SGA) (see manufacturers guidelines):

- 325 • When face mask ventilation is ineffective
- 326 • When a more definitive airway is required as an alternative to tracheal intubation.
- 327 • Where tracheal intubation is not possible or deemed unsafe because of congenital
- 328 abnormality, a lack of equipment, or a lack of skill.
- 329 • When chest compressions are performed.
- 330 **[h4] Nasopharyngeal and Oropharyngeal airways**
- 331 • Consider nasopharyngeal or oropharyngeal airways, especially when facemask ventilation may
- 332 be difficult (e.g. micrognathia).
- 333 • Use oropharyngeal airways with caution in infants <34 weeks. They might contribute to airway
- 334 obstruction.
- 335 **[h4] Tracheal tube** Consider tracheal tube placement:
- 336 • When equipment and skills permit
- 337 • When facemask or SGA ventilation are ineffective
- 338 • With prolonged ventilation
- 339 • When suctioning the lower airways (removal of presumed tracheal blockage)
- 340 • When chest compressions are performed
- 341 When performing tracheal intubation:
- 342 • Have a range of different sized tubes available
- 343 • Use video laryngoscopy or, if not available, direct laryngoscopy
- 344 • Use exhaled CO<sub>2</sub> detection and clinical assessment to confirm tracheal intubation
- 345 ○ Be aware that exhaled CO<sub>2</sub> detection may be false negative in low or no cardiac output
- 346 states at birth
- 347 • Use appropriate imaging to confirm correct tube position
- 348 • If available use respiratory function monitoring to confirm tube position within the airway and
- 349 adequate ventilation (expired tidal volume 4 to 8 mL kg<sup>-1</sup> with minimal leak).
- 350
- 351 **[h2] Breathing**
- 352 • Inflate the lungs when the newborn infant is not breathing using a facemask or nasal prongs.
- 353 **[h3] Assisted ventilation**
- 354 **[h4] Lung inflation**
- 355 • If apnoeic, gasping or not breathing effectively, start positive pressure ventilation (PPV) as
- 356 soon as possible to inflate the lungs – ideally within 60s.

- 357 • Apply appropriately fitting nasal prongs or a facemask connected to a device for providing  
358 positive pressure ventilation.
- 359 • Give inflations with an inflation time of 1-3s:
- 360 ○ If a 1s inflation time is used, give 10 inflations
- 361 ○ If a 2-3s inflation time is used, give 5 inflations
- 362 • Infants <32 weeks: starting inflation pressure 25 cm H<sub>2</sub>O.
- 363 • Infants ≥32 weeks: starting inflation pressure 30 cm H<sub>2</sub>O.
- 364 • Use pulse oximetry and ECG.

365 << INSERT Table 3 Inflation time and number of inflations >>

366 **[h4] Assessment**

- 367 • During lung inflations: look for chest movement.
- 368 ○ Visible chest movement during inflations indicates a patent airway and delivered volume.
- 369 ○ Failure of the chest to move may indicate that the airway is not open, or that insufficient  
370 inflation pressure/volume is delivered
- 371 • After lung inflations: check HR
- 372 ○ An increase in HR within 30s of positive pressure ventilation, or a stable HR >100 min<sup>-1</sup>,  
373 usually confirms adequate ventilation/oxygenation
- 374 ○ HR <100 min<sup>-1</sup> or decreasing usually suggests continued hypoxia and almost always  
375 indicates inadequate ventilation

376 **[h4] If there is a HR response**

- 377 • Continue uninterrupted positive pressure ventilation until the infant begins to breathe  
378 adequately and the HR >100 min<sup>-1</sup>.
- 379 • Aim for a positive pressure ventilation rate of 30 ventilations min<sup>-1</sup> with an inflation time of  
380 approximately 1s.
- 381 • Adapt inflation pressure based on clinical observation (chest movement and HR) and/or  
382 respiratory function monitor data.
- 383 • Reassess breathing and HR every 30s, until the newborn infant is deemed stabilised.
- 384 • Consider inserting SGA or tracheal tube if apnoea continues or if mask ventilation is not  
385 effective.

386 **[h4] If there is no HR response**

387 If there is **no** HR response **and** the chest is not moving with inflations:

- 388 • Call for help.
- 389 • Recheck equipment.

- 390 • Perform airway opening technique of choice.
- 391 • If the airway opening techniques are ineffective, increase inflation pressure.
- 392 • Repeat inflations after every airway opening technique or after increasing inflation pressure.
- 393 • Re-assess chest movement and HR until visible chest movement **or** HR response.
- 394 • Reduce inflation pressure when chest movement is seen and clinical improvement.
- 395 • If being used, check with a respiratory function monitor that expired tidal volume is within
- 396 target range (4 to 8 mL kg<sup>-1</sup>, depending on gestational age).
- 397 Without adequate lung inflation, chest compressions will be ineffective:
- 398 • **Confirm effective ventilation** through observed chest movement or other measures of
- 399 respiratory function.
- 400 • **Then** progress to chest compressions.
- 401 **[h3] Continuous positive airway pressure (CPAP) and Positive end-expired pressure (PEEP)**
- 402 • Use either nasal prongs or a facemask as device-patient interface
- 403 • Start with CPAP at 6 cm H<sub>2</sub>O as initial breathing support in:
  - 404 ○ Spontaneously breathing infants <32 weeks with respiratory distress
  - 405 ○ Spontaneously breathing infants ≥32 weeks with respiratory distress requiring
  - 406 supplemental O<sub>2</sub>
- 407 • In infants needing positive pressure ventilation, start with PEEP at 6 cm H<sub>2</sub>O.
- 408 **[h3] Ventilation devices**
- 409 • Use appropriately sized nasal prongs or facemask.
- 410 • Ensure effective seal with minimal force on the facemask.
- 411 • Where possible, use a T-piece resuscitator capable of providing either CPAP or positive
- 412 pressure ventilation +PEEP when giving ventilatory support, especially in the preterm infant.
- 413 • Self-inflating bags should be available as backup:
  - 414 ○ Take care not to deliver excessive volumes
  - 415 ○ Be aware that CPAP might not be effectively delivered even when a PEEP valve is used
- 416 **[h3] Oxygen**
- 417 • Use pulse oximetry and O<sub>2</sub>-blenders during resuscitation or stabilisation in the delivery area.
- 418 • Check O<sub>2</sub> and saturations every 30s.
- 419 • Titrate inspired O<sub>2</sub> to achieve target SpO<sub>2</sub> between the 25<sup>th</sup> -75<sup>th</sup> percentile (Table 4).
- 420 • Infants ≥32 weeks needing respiratory support:
  - 421 ○ Start at 21% O<sub>2</sub>
- 422 • Infants <32 weeks:

- 423 ○ Start at  $\geq 30\%$  O<sub>2</sub>
- 424 ○ Avoid SpO<sub>2</sub> <80% and/or bradycardia at 5 minutes of age
- 425 << INSERT Table 4 Target oxygen saturation ranges >>
- 426 < INSERT Table 5 summary of positive pressure ventilation and oxygen >>

427

## 428 [h2] Circulation

### 429 [h3] Chest compressions

- 430 • Start chest compressions if the HR remains <60 min<sup>-1</sup> after at least 30s of effective ventilation.
- 431 • When starting chest compressions:
  - 432 ○ Increase O<sub>2</sub> to 100%
  - 433 ○ Call for experienced help if not already summoned
  - 434 ○ Anticipate the need to secure the airway and establish vascular access for drugs
- 435 • Use a 3:1 compression-to-ventilation ratio (C:V), aiming for 90 compressions and 30
- 436 ventilations (120 events) per minute.
- 437 • Use the two-thumb-hands-encircling-technique with overlapping or adjacent thumbs to
- 438 deliver chest compressions (figure 7).
- 439 • Compress to a depth of one-third of the anterior-posterior chest diameter.
- 440 • Allow full chest recoil between compressions.
- 441 • Re-assess HR every 30s.
- 442 • If HR <60 min<sup>-1</sup>, secure the airway with an SGA or tracheal tube (if competent and not already
- 443 done) with minimal interruptions to ongoing chest compressions.
- 444 • After SGA placement or tracheal intubation continue with the 3:1 C:V ratio.
- 445 • Titrate O<sub>2</sub> against the oxygen saturation once a reliable value is achieved (table 4)
- 446 • Discontinue chest compressions if the HR is >60 min<sup>-1</sup>; check for output (e.g. auscultation,
- 447 pulse check, pulse oximetry, signs of life)

448 << INSERT Figure 7 Demonstration of the two-thumbs-hand-encircling-technique >>

449

### 450 [h3] Vascular access

#### 451 [h4] Umbilical Venous Access

- 452 • Use the umbilical vein for rapid emergency vascular access during resuscitation at birth.
- 453 • Perform emergency umbilical venous catheter (UVC) placement under clean rather than
- 454 sterile conditions to ensure timely vascular access is secured.
- 455 • Consider the use of emergency umbilical venous catheter until some days after birth as it

456 may still be achievable.

457 **[h4] Intraosseous Access**

- 458 • Use intraosseous (IO) access as an alternative method of emergency vascular access for
- 459 drugs/fluids.
- 460 • Consider device-specific weight limitations for IO related equipment.
- 461 • Ensure there is no extravasation when administering drugs/fluids.
- 462 • Do not aspirate blood; even when correctly positioned, it is often not possible.

463 **[h4] Support of transition / post-resuscitation care**

- 464 • If venous access is required following resuscitation, peripheral access may be adequate
- 465 unless multiple infusions and/or vasopressors are required in which case central access may
- 466 be preferred.

467

468 **[h2] Drugs during resuscitation at birth**

469 Resuscitation drugs may be considered where, despite adequate control of the airway, effective  
470 ventilation, and chest compression for at least 30s, HR remains  $<60 \text{ min}^{-1}$  and is not increasing.

471 **[h3] Adrenaline**

- 472 • Umbilical venous catheter or IO is the preferred route.
  - 473 o Give 20 micrograms  $\text{kg}^{-1}$  (0.2  $\text{mLkg}^{-1}$  of 1:10,000 adrenaline [0.1 mg/mL])
  - 474 o Give subsequent doses every 4 minutes if HR remains  $<60 \text{ min}^{-1}$
- 475 • If no umbilical venous catheter/IO access but intubated:
  - 476 o Give intra-tracheal adrenaline at dose of 100 micrograms  $\text{kg}^{-1}$  (1  $\text{mLkg}^{-1}$  of 1:10,000
  - 477 adrenaline [0.1 mg/mL])
  - 478 o As soon as umbilical venous catheter/ IO access is obtained, immediately give a dose via
  - 479 this route, irrespective of when the intra-tracheal dose was given

480 **[h3] Glucose**

- 481 • If possible, check the blood glucose value during resuscitation.
- 482 • If blood glucose is low: give glucose 250  $\text{kg}^{-1}$  (2.5 mL/kg of 10% glucose).

483 **[h3] Intravascular volume replacement**

- 484 • Give 10 mL/kg of group O Rh-negative blood or isotonic crystalloid solution if suspected blood
- 485 loss or in a newborn infant unresponsive to other resuscitative measures.

486

487 **[h2] Absence of an adequate response** despite appropriate resuscitation measures

- 488       • Consider other factors which may be impacting the response to resuscitation, and which  
489           require addressing such as the presence of pneumothorax, hypovolaemia, congenital  
490           abnormalities, equipment failure.

491

492 **[h2] Low resource or remote settings**

493 Births outside the hospital may be considered birth in a remote or lower resource setting, and not all  
494 hospitals have the same resources.

495 **[h3] Planned home births**

- 496       • Ideally, two trained HCPs should be present at all home births.  
497       • Have at least one HCP competent in providing inflations, PPV and CC to the newborn infant.  
498       • Have a minimum set of equipment of an appropriate size for the newborn infant available.  
499       • Have a clear plan of who will attend, what equipment will be available, and how transfer will  
500           be arranged if newborn support is required and agreed this with parents when formulating the  
501           home birth plan.  
502       • HCPs attending home births should have pre-defined plans for unexpected or difficult  
503           situations, including knowing how to communicate with receiving healthcare facilities for the  
504           mother and newborn infant.

505 **[h3] Unexpected births outside the hospital**

- 506       • Emergency services should be prepared and trained for such events and carry appropriate  
507           equipment, especially related to thermal care and support of airway and breathing.  
508       • Equipment to support thermal care and oxygenation should be available.

509 **[h3] Temperature control out of hospital**

- 510       • Involved HCPs should have a heightened awareness of the increased risk of hypothermia in  
511           infants born (unexpectedly) out of hospital.  
512       • They should perform regular temperature checks and intervene if the temperature is too low.  
513       • Most interventions for infants born in hospital (see temperature management) can also be  
514           applied outside the hospital.  
515       • Place compromised, preterm (<37 weeks), and/or growth restricted infants in a preheated  
516           incubator for thermal control and transport.

517

518 **[h2] Post-resuscitation care**

- 519 • Once effective ventilation and circulation are established, the infant should be cared for in or  
520 transferred to an environment in which close monitoring and anticipatory care can be  
521 provided.

522 **[h3] Glucose management**

- 523 • Measure blood glucose values early and regularly until they have stabilised in the normal  
524 range; especially in newborns resuscitated at birth, those at risk of hypoxic-ischaemic  
525 encephalopathy (HIE), and/or receiving intravenous glucose.
- 526 • Avoid hypoglycaemia, hyperglycaemia, and large swings in blood glucose value.

527 **[h3] Thermal Care**

- 528 • Monitor the infant's temperature frequently or continuously after resuscitation.
- 529 • Maintain temperature between 36.5°C and 37.5°C and rewarm if the temperature is below  
530 this.

531 **[h3] Therapeutic Hypothermia**

- 532 • Consider inducing therapeutic hypothermia (33-34°C) *after* completion of resuscitation and  
533 detailed assessment of the infant in (near) term infants with clinical, biochemical, and (if  
534 available) neurophysiological evidence of HIE.
- 535 • Use appropriate eligibility criteria and strictly defined protocols to guide the cooling process;  
536 inappropriate application of therapeutic hypothermia may be harmful.
- 537 • Arrange safe transfer to an appropriately equipped facility where monitoring and treatment  
538 can be continued.
- 539 • Monitor (rectal) temperature during transport and, if available, apply active cooling with a  
540 servo-controlled device while transferring the infant.

541 **[h3] Oxygenation & Ventilation**

- 542 • Consider additional monitoring of post-ductal oxygen saturation to identify pulmonary  
543 hypertension.
- 544 • Avoid hypoxia and hyperoxia.
- 545 • Avoid inadvertent hypocapnia during mechanical ventilation.

546 **[h3] Documentation & Prognostication**

- 547 • Keep an accurate time-based record of the infant's clinical state, interventions and responses  
548 during resuscitation to facilitate retrospective review.
- 549 • Record APGAR scores.

550 **[h3] Clinical team debriefing**

- 551 • Use performance-focused, interdisciplinary/interprofessional team debriefings following  
552 resuscitation or other non-routine situations to optimise individual and team performance as  
553 well as systems issues (e.g., emergency supplies, equipment).

554

555 **[h2] Communication with parents**

556 **[h3] Where intervention is anticipated**

- 557 • The decision to attempt resuscitation of an extremely preterm or clinically complex infant  
558 should be taken in close consultation with the parents and senior paediatricians, midwives,  
559 and obstetricians.
- 560 • Discuss the options, including the potential need and magnitude of resuscitation and the likely  
561 prognosis before birth, so that an individualised management plan can be agreed.
- 562 • Ensure concise and factual documentation of discussions is recorded in mother's notes before  
563 birth and in the infant's notes after birth.

564 **[h3] For every birth**

- 565 • If parents want and resources allow, enable parents to be present during the stabilisation or  
566 resuscitation.
- 567 • Consider the views of the resuscitation team, parents and circumstances.
- 568 • Ensure that parents are fully informed about the progress of the care provided to their infant.
- 569 • Identify a member of healthcare staff to support parents and be aware that witnessing the  
570 resuscitation of their infant will be distressing for them.
- 571 • Encourage parents to hold or touch their infant as soon as possible after resuscitation; this  
572 should be facilitated especially when the resuscitation was unsuccessful.
- 573 • Ensure an accurate record is kept of the resuscitation and of any subsequent parental  
574 communication.
- 575 • Provide an explanation of any procedures and why they were required.
- 576 • Facilitate further discussions later to enable parents to reflect and to aid their understanding  
577 of events.
- 578 • Provide additional support for parents following resuscitation at birth.

579

580 **[h2] Discontinuing or withholding resuscitation**

- 581 • Use national or regional outcomes and guidelines to interpret these recommendations

- 582 • When discontinuing, withdrawing or withholding resuscitation, care should be focused on the  
583 comfort and dignity of the infant and family and should ideally involve senior  
584 paediatric/neonatal staff.

### 585 [h3] Discontinuing resuscitation

- 586 • If the HR remains absent despite resuscitation, review clinical factors (for example  
587 presence/absence of dysmorphic features, gestation of the infant), effectiveness of  
588 resuscitation, and the views of other members of the clinical team about continuing  
589 resuscitation.
- 590 • If the HR of a newborn infant remains absent for more than 20 minutes after birth despite the  
591 provision of all recommended steps and exclusion of reversible causes, consider stopping  
592 resuscitation.
- 593 • For preterm infants (particularly extremely preterm), it may be appropriate to discontinue  
594 resuscitation earlier than 20 minutes. The decision should be individualised.
- 595 • Where there is partial or incomplete HR improvement despite apparently adequate  
596 resuscitative efforts, the choice is much less clear. It may be appropriate to take the infant to  
597 the intensive care unit and later consider withdrawing life sustaining treatment.
- 598 • Where life-sustaining treatment is withheld or withdrawn, infants should be provided with  
599 appropriate palliative (comfort focused) care.

### 600 [h3] Withholding resuscitation

- 601 • Decisions to withhold life sustaining treatment should be made in advance of birth together  
602 with parents in the light of regional/national evidence on outcome if resuscitation and active  
603 (survival focused) treatment is attempted.
- 604 • In situations where there is extremely high (e.g. >90%) predicted neonatal mortality and  
605 unacceptably high morbidity in surviving infants, attempted resuscitation and active (survival  
606 focused) management is usually not appropriate.
- 607 • Resuscitation is nearly always indicated in conditions associated with lower (e.g. <50%)  
608 neonatal mortality and what is deemed to be acceptable morbidity. This will include most  
609 infants with congenital malformations and most infants >24 weeks or above in high resource  
610 settings with access to neonatal intensive care.
- 611 • Resuscitation should usually be commenced in situations where there is uncertainty about the  
612 outcome and there has been no chance to have prior discussions with parents.
- 613 • In situations where there is high mortality (e.g. >50%) and/or a high rate of morbidity, and  
614 where the anticipated burden of medical treatment for the child is high, parental wishes

615 regarding resuscitation are usually supported. It may be appropriate to provide full  
616 resuscitation, to provide some measures (but withhold other interventions) or to provide  
617 comfort focused care. Provision of antenatal palliative care support can be beneficial to  
618 parents in the face of certain or uncertain poor outcomes.

This is a DRAFT-version

## 619 [h1] Evidence informing the guidelines

620

## 621 [h2] Newborn Life Support or Paediatric Life Support?

622 The ERC Guidelines 2025 NLS applies mainly to newborn infants at birth and in the immediate  
623 postnatal phase, i.e. during perinatal transition. There is no clear definition of when transition ends.  
624 Thus, evidence-based recommendations on when to convert from NLS to Paediatric life support (PLS)  
625 guidelines are challenging to produce.

## 626 [h3] Epidemiology

627 Neonatal intensive care units (NICU) often set distinct age thresholds for admitting and keeping  
628 infants. Some transfer infants to paediatric units at 44 weeks' postmenstrual age (2), whereas other  
629 NICUs transfer infants as late as 24 months. (3) Moreover, some NICUs operate separately from  
630 birthing hospitals, affecting the patient case-mix. Thus, the incidence of NICU resuscitation with chest  
631 compressions and/or adrenaline varies between 0.25% to 1-2% of infants (4-7), and a significant  
632 proportion of CPR events in paediatric intensive care units (PICUs) is in infants <1 year of age. (8) Most  
633 NICU arrests are respiratory in origin (6, 7), with more respiratory-related instances including tracheal  
634 tube and airway complications in NICU resuscitation events compared with PICU/cardiac intensive care  
635 unit resuscitation events. (9) Pulseless electrical activity or asystole occurs in 13% of NICU resuscitation  
636 events, while ventricular tachycardia or fibrillation incidents are rare in NICUs.(10)

## 637 [h3] Differences between NLS and PLS guidelines

638 Neonatal resuscitation guidelines prioritise ventilation to stabilise bradycardic or asystolic newborns.  
639 (11) Paediatric resuscitation guidelines emphasise chest compressions while managing ventilation  
640 carefully to prevent overventilation by the HCP. (12) The two guidelines also diverge in areas such as  
641 thermal care for different maturity of newborns, synchronisation of breaths with compressions after  
642 tracheal intubation, and use of medications and adjunctive methods.(13) Unlike paediatric guidelines,  
643 neonatal guidelines omit management strategies for (septic) shock and arrhythmias other than  
644 bradycardia/asystole, leaving out rhythm evaluation and defibrillation.

## 645 [h3] Evidence informing the transition from NLS to PLS

646 Several approaches on the use of NLS or PLS have been suggested or are used, such as location-based,  
647 age-based, patient-based or provider-based approaches. (13) A location-based approach would be to  
648 take education and training implications into consideration by choosing guidelines based on location  
649 (e.g., NICU or PICU). Observational studies that found differences in outcomes after PICU versus NICU  
650 CPR events did not account for prematurity and low birth weight in infants in the NICU. (14) The  
651 presence of fluid-filled lungs only during immediate perinatal transition may provide a rationale for

652 changing from the neonatal to paediatric resuscitation guideline using a time-based approach. For  
653 example, after the first 24 hours of life, (13, 15) or at 44 weeks postmenstrual age as a cut-off point. A  
654 patient-oriented approach may be to focus on the pathophysiology of the bradycardia or arrest, as in  
655 cases of congenital and acquired heart disease. (16) A provider-based approach may have many  
656 similarities with a location-based approach, but if HCP are trained in both NLS and PLS they can apply  
657 both.

658 In the absence of evidence, an approach to education and training tailored towards the decision to use  
659 one or both guidelines based on individual unit case-mix and epidemiology of cardiac arrests may be  
660 reasonable. The ERC recommends creation of local policies applicable to the healthcare setting (good  
661 practice statement).

662

## 663 [h2] The most preterm infants at the limits of viability

664 Survival and outcomes of preterm infants continue to improve, especially for those born at extremely  
665 low gestational age (GA). (17) Following recently changed guidance concerning the initial stabilisation  
666 of the most preterm infants, i.e. those born below 25 weeks (18, 19), these infants are now  
667 increasingly being offered survival-oriented care. (20-22) However historically, trials in neonatal  
668 medicine have, almost without exception, excluded the most preterm infants. (23)

669 Consequently, the ERC NLS Writing Group cautions that recommendations given in the ERC Guidelines  
670 2025 NLS are based on evidence from studies of higher GA infants and any extrapolation of such  
671 evidence to the most preterm infants will not fully take into account their distinct physiology and  
672 response to treatment. (24)

673

## 674 [h2] Factors before birth

### 675 [h3] Perinatal transition

676 Survival at birth involves major physiological changes during perinatal transition from fetal to newborn  
677 life. First, lung liquid-clearance and aeration need to occur after which pulmonary gas exchange can be  
678 established. (25) Most newborn infants transition smoothly, but some have problems with transition  
679 and without timely and adequate support might need resuscitation. (26-29) Approximately 11% of all  
680 newborns receive interventions with large variation between hospitals (1.4–38.1%). (30) Newborn  
681 infants born via caesarean section receive an intervention (19.6%) more often than vaginally born  
682 infants (5.9%), with most common interventions being CPAP (7%), O<sub>2</sub> supplementation (8%),  
683 suctioning (6%) and non-invasive ventilation (4%). (30) Less common interventions include tracheal  
684 intubation (1%), cardiac compressions (0.1%) adrenaline administration (0.1%), intraosseous access  
685 (0.01%) and SGA insertion (0.01%). (30) Intervention frequencies varied considerably between

686 hospitals and countries. (30) In preterm infants, the need for respiratory support is higher, with  
687 almost all infants born with a GA <30 weeks receiving CPAP and/or positive pressure ventilation (PPV).  
688 (30)

### 689 [h3] Risk factors

690 Several maternal and fetal prenatal and intrapartum factors increase the risk for compromised  
691 transition and the need for resuscitation. A recent multicentre survey and an ILCOR evidence update  
692 confirm previously identified risk factors for needing assistance after birth. (30, 31) There is no  
693 universally applicable model to predict risk for resuscitation or need of support during transition, and  
694 the list of risk factors in the guidelines is not exhaustive. Elective caesarean delivery at term, in the  
695 absence of other risk factors, does not increase the risk of needing newborn resuscitation. (32, 33)  
696 In accordance with the unchanged ILCOR recommendation, the ERC recommends that when an infant  
697 is delivered at term by caesarean section under regional anaesthesia an HCP capable of performing an  
698 initial assessment and assisted ventilation should be present at the birth. It is not necessary for a  
699 provider skilled in neonatal intubation to be present at that delivery. (31)

### 700 [h3] Staff attending births in hospitals

701 It is not always possible to predict the need for stabilisation or resuscitation before an infant is born.  
702 Therefore, the ERC recommends that those in attendance at birth need to be able to undertake initial  
703 resuscitation steps effectively. The experience of the team and their ability to respond in a timely  
704 manner can improve outcomes of term (34) and preterm infants. (35-37) It is essential that  
705 resuscitation teams can respond rapidly if not present from the beginning. In a simulation-based study  
706 on term neonatal resuscitation a significant increase in workload was demonstrated in 2-person teams  
707 compared to 3-person teams. (38) The ERC advises a process should be in place for rapidly mobilising  
708 extra HCPs with appropriate resuscitation skills.

### 709 [h3] Telemedicine

710 In hospitals with low birth rates, it can be difficult for staff to maintain neonatal resuscitation skills.  
711 (39-41) Video telemedicine may help address these challenges by providing immediate access to  
712 neonatal specialists, allowing a neonatologist to virtually assist with neonatal resuscitation at remote  
713 locations, which ultimately might improve patient outcomes. (42-45) Limited observational data  
714 suggests that video telemedicine may improve the quality of neonatal resuscitation and reduce the  
715 need for neonatal transfers and can be introduced without significant adverse effects. (42, 46-50)  
716 The ERC recommends that where the technology is available and/or access to a neonatologist is not  
717 immediately available, telemedicine use is considered.

### 718 [h3] Equipment and environment

719 Suggestions have been made on standardising an optimal layout of a resuscitation area, (51) but no  
720 published evidence has demonstrated improvement in patient outcome due to specific arrangements.  
721 However, some studies suggest a reduced retrieval time for emergency materials when organized  
722 according to specific frameworks such as the ABC protocol (airway, breathing, circulation) (52), task-  
723 based package approach (53), or with a focus on emergency supplies or airway management. (37)

### 724 [h3] Briefing, Debriefing & Checklists

725 Briefing with role allocation and the use of checklists improve team functioning and communication  
726 and are suggested. (54, 55) Evidence on the isolated impact of briefing on patient outcomes is  
727 challenging, as it is typically implemented within quality improvement bundles. However, an ILCOR  
728 scoping review (2021) on the effect of briefing and debriefing on the outcome of neonatal  
729 resuscitation concluded that ‘improvements in the process of care (...), short term clinical outcomes  
730 and a reduction in communication problems’ were associated with briefing and debriefing. (56, 57)  
731 The use of checklists during briefings and debriefings may help improve team communication and  
732 process, but there is little evidence of effect on clinical outcomes. (58, 59) The ERC recommends  
733 (de)briefing the team present at birth and suggests the use of cognitive aids.

734

### 735 [h2] Education

736 For an in-depth discussion on resuscitation education principles, see ERC 2025 Guidelines Education  
737 for Resuscitation (ref). Research on educational methods in neonatal resuscitation is evolving, but due  
738 to study heterogeneity with non-standardised outcome measures, there is still little evidence on the  
739 effect of different training modalities on clinical outcome. (60-62) Nevertheless, available studies on  
740 the clinical impact of neonatal resuscitation training are summarised in table 6.

### 741 [h3] Frequency of training

742 Infrequent neonatal resuscitation training and rare clinical exposure lead to skills decay. Two  
743 observational studies using video analysis found that annual training may be insufficient (63), as skills  
744 deteriorate within 3–6 months, highlighting the benefits of high-frequency, short-duration sessions.  
745 (28, 64) After NLS courses significant knowledge and skill decay was found within three months, with  
746 technical skills declining faster than knowledge. (65) Another study focusing on neonatal ventilation  
747 skills, found that airway patency requires training every 4.5 months, and mask seal every 1.5 months.  
748 (63) The ERC recommends training at a minimum interval of 12 months, preferably supplemented with  
749 more frequent short booster sessions every 3-6 months.

### 750 [h3] Technical skills, behavioural skills and self-efficacy

751 Optimal newborn life support requires neonatal providers to possess not only technical expertise but  
752 also behavioural skills, including team collaboration competences, crisis resource management, and

753 personal resilience. (66, 67) A 2021 ILCOR systematic review about team competences and training for  
754 resuscitation found improved skills performance during clinical resuscitation and suggests its inclusion  
755 in basic and advanced life support courses (68, 69). Providers need strong confidence to perform NLS  
756 optimally, initiate and persist in resuscitation, and stay resilient under pressure. (70-72) Confidence  
757 can be achieved, among others, through practice and reflection, and observational learning, where  
758 participants are motivated to attain a similar level of performance as they observed in their peers. (70,  
759 73, 74) The ERC recommends incorporating team collaboration competencies in newborn life support  
760 training.

### 761 [h3] System training

762 *In situ* neonatal simulation training is highly effective not only for training in human factors and  
763 teamwork, it also enables adaptation of team composition, the environment and equipment to create  
764 ideal circumstances for newborn resuscitation performance. (75-78) Simulation training may also be  
765 used to rigorously test new neonatal environments or procedures. The ERC recommends that  
766 simulation training forms part of resuscitation training.

767

### 768 [h2] Thermal Control

769 The World Health Organization recommends keeping newborn temperatures between 36.5°C and  
770 37.5°C.(79) Exposed, wet newborn infants cannot maintain their body temperature in a room that  
771 feels comfortably warm for adults. The mechanisms (convection, conduction, radiation, evaporation)  
772 and effects of hypothermia and how to avoid these have been reviewed elsewhere. (80, 81)  
773 Hypothermia may impair respiratory function, lower the arterial oxygen tension, cause elevated  
774 pulmonary vascular resistance, and increase the risk of metabolic acidosis, hypoglycaemia, and  
775 bradycardia. (80) Two recent systematic reviews showed associations between admission hypothermia  
776 and various morbidities (intraventricular haemorrhage, bronchopulmonary dysplasia, sepsis,  
777 retinopathy of prematurity) and mortality in very low birthweight infants (<1500g) and very preterm  
778 infants, respectively. (82, 83)

779 As the admission temperature of non-asphyxiated infants is associated with morbidity and mortality at  
780 all gestations and in all settings (69, 84, 85), the ERC recommends to record temperature as both a  
781 predictor of outcome and a quality indicator. (11) A recent systematic review and network meta-  
782 analysis showed that (the combination of) plastic bags/wraps, plastic caps, thermal mattress, and  
783 heated humidified gases in the delivery room reduced major brain injury and mortality in preterm  
784 infants. (86) The ERC recommends that as a minimum, plastic bags/wraps and hats are used in preterm  
785 infants at birth, and where available heated humidified gases are also used at the earliest opportunity  
786 in preterm infants.

### 787 [h3] Temperature monitoring

788 Temperature monitoring is key to avoiding hypothermia. However, there is very little evidence to  
789 guide the optimal placement of temperature monitoring probes on the infant. In a study of 122  
790 preterm infants (28-36 weeks) randomised to different sites for temperature monitoring, dorsal,  
791 thoracic, and axillary sited probes all had comparable temperature measurements. (87) There are no  
792 published studies comparing the use of rectal temperature probes. The ILCOR NLS Task Force does not  
793 specify the site where the temperature should be determined. (31, 88, 89)

794 In infants <1500g immediately after birth, servo-controlled thermoregulation did not improve  
795 admission normothermia compared with using a radiant warmer in manual mode. (90) ILCOR stated  
796 that there is insufficient published human evidence to suggest for or against the use of a radiant  
797 warmer in servo-controlled mode compared with manual mode in infants <34 weeks directly after  
798 birth.(91) In newborns who are unintentionally hypothermic after birth, ILCOR concluded that there is  
799 insufficient evidence to recommend either a rapid ( $\geq 0.5^{\circ}\text{C}/\text{hour}$ ) or slow ( $< 0.5^{\circ}\text{C}/\text{hour}$ ) rewarming rate.  
800 (92)

801 The ERC recommends that all newborn infants undergoing resuscitation and all preterm infants  
802 undergoing support of transition have their temperature monitored frequently or continuously during  
803 resuscitation until stabilisation.

### 804 [h3] Hyperthermia

805 Hyperthermia ( $\geq 38.0^{\circ}\text{C}$ ) should be avoided, because it is associated with adverse effects. (11) Infants  
806 born to febrile mothers have a higher incidence of perinatal respiratory compromise, neonatal  
807 seizures, early mortality, and cerebral palsy. (93-95) Animal studies indicate that hyperthermia during  
808 or following ischaemia is associated with a progression of cerebral injury. (95)

### 809 [h3] Term and near-term infants $\geq 34$ weeks

810 ILCOR treatment recommendation (89, 96) suggests a room temperature of  $23-25^{\circ}\text{C}$  in infants  $\geq 34$   
811 weeks. (89, 96) If support of transition or resuscitation is not required, immediate skin-to-skin care is  
812 good practice to maintain normothermia. A Cochrane review involving 46 trials and 3850 dyads of  
813 mothers with their (predominantly healthy term and some late preterm) newborns concluded that  
814 skin-to-skin contact may be effective in maintaining thermal stability and improve maternal bonding  
815 and breast-feeding rates. (97) Aligning with ILCOR, the ERC recommends the use of plastic bags/wraps  
816 where skin-to-skin contact is not possible, and resuscitation is not required.

### 817 [h3] Preterm infants <34 weeks

818 For infants <34 weeks a room temperature of  $23-25^{\circ}\text{C}$  is suggested. (88, 91) For infants <28 weeks,  
819 room temperature should ideally be  $>25^{\circ}\text{C}$ . (80, 81, 98) The use of plastic bags or wraps (without  
820 drying) is advocated in infants <34 weeks. Further thermal control while using radiant warmers in the

821 delivery area can be achieved with a combination of warmed blankets, cap, thermal mattress, heated  
822 humidified respiratory gases, and skin-to-skin care. With these interventions, both hypothermia and  
823 hyperthermia are possible and require attention. (91) Quality improvement programs, including the  
824 use of checklists, continuous feedback, and debriefing have shown to significantly reduce the  
825 incidence of hypothermia at admission in very preterm infants. (98, 99)

### 826 [h3] Delivery or operating room cuddles

827 Following stabilisation after birth it may be possible to offer physical contact between the parents and  
828 their baby in the form of supervised skin-to-skin contact or a cuddle. Studies have considered the  
829 feasibility of a delivery room cuddle in relation to physiological variables (HR, temperature). (100, 101)  
830 The effect of delivery room cuddles on thermoregulation was conflicting; with some studies reported  
831 no difference (100-103) and others reported more hypothermia in infants who received skin to skin  
832 care after birth. (102-106) There is emerging evidence of positive effect on maternal bonding (100,  
833 107, 108) and that delivery room cuddles may promote breast feeding in near term and term infants.  
834 (108) However, there is also evidence of potential risks including accidental extubation,  
835 disconnections, or apnoea. (95, 98)

836 Current evidence is insufficient to provide a specific recommendation and there is no ILCOR evidence  
837 review on this topic. Discussing the possibility of a delivery room cuddle on an individual basis is  
838 reasonable, if the clinical team feels confident to support this. However, the practicality of offering this  
839 will not be clear until after their baby has been born. If a delivery room cuddle is impractical,  
840 encourage brief physical contact, e.g. touching their baby's hand as an alternative. Where  
841 resuscitation measures are required, this takes priority.

842

### 843 [h2] Umbilical cord clamping

844 There is no universally accepted definition of 'delayed' or 'deferred' cord clamping (DCC), only that it  
845 does not occur immediately after birth. Early or immediate cord clamping (ICC) has been defined as  
846 less than 30sec after birth, later or delayed cord clamping as >30sec after birth or when cord pulsation  
847 has ceased. (109, 110) Physiological based cord clamping (PBCC) is not based on time, but on  
848 physiological parameters (i.e. when breathing has been initiated). (111, 112) When possible,  
849 interventions for stabilising the infant may take place close to the mother with intact cord. (113)  
850 The ERC recommends facilitating at least 60s of DCC for newborn infants without need for support;  
851 and to clamp the cord <30s to minimise delay in interventions in those in need for resuscitation. If  
852 intact cord stabilisation can be safely performed, longer DCC is preferred, especially in newborn  
853 infants <34 weeks.

### 854 [h3] Rationale: experimental and observational studies

855 Although ICC was introduced to reduce postpartum haemorrhage (114), its impact was minimal and  
856 primarily associated with reduced birth weight. (115, 116) Clamping the cord before lung inflation and  
857 the increase in pulmonary circulation has occurred results in reduced ventricular preload and  
858 increased left ventricular afterload. (117), impairing the circulation and causing hypoxia. (111, 117) A  
859 second rationale for DCC is placental transfusion – blood redistribution from placenta to newborn  
860 which can account for up to 25% of placental volume. (118, 119) Gravity and uterine contractions do  
861 not drive this transfusion (120, 121), but spontaneous breathing of the infant might. Therefore,  
862 clamping should ideally be delayed until breathing has been established. (122)

### 863 [h3] Infants $\geq$ 34 weeks

864 A 2019 Cochrane review found that DCC compared to ICC increased birth weight, neonatal  
865 haemoglobin, and reduced iron deficiency at 3-6 months, without increasing polycythaemia. (123) A  
866 2021 ILCOR meta-analysis of 33 trials in newborns  $\geq$ 34 weeks confirmed these findings, showing no  
867 effect on mortality or need for resuscitation. (110) DCC improved early ( $\leq$ 24 hours) and later (7 days)  
868 haematological and circulatory parameters, but had no impact on longer term anaemia,  
869 neurodevelopment, or phototherapy. (110)

870 Evidence on DCC in (near) term newborns needing resuscitation is limited. One study found no HR  
871 difference between cord intact resuscitation and ICC, (124) while two RCTs reported better vital  
872 parameters, higher Apgar scores, and reduced need for ventilation and/or chest compression. (125,  
873 126) Only one trial reported mortality with no difference. (126) Admission temperatures were similar  
874 across all three trials. (124-126)

### 875 [h3] Infants $<$ 34 weeks

876 Multiple trials have compared DCC with ICC in preterm infants. Most used DCC for 30-60s, excluding  
877 infants needing immediate resuscitation. Studies using intact cord resuscitation applied longer  
878 clamping times. A 2021 ILCOR meta-analysis (infants  $<$ 34 weeks) DCC ( $\geq$ 30s) may slightly improve  
879 survival (109), with better cardiovascular stability, less inotropic support, improved haematological  
880 indices, and fewer transfusions – without effects on prematurity complications (or adverse maternal  
881 outcomes). Subgroup analysis showed a possible positive link between survival and DCC duration (109,  
882 127, 128). A separate systematic review and individual participant data meta-analysis confirmed  
883 reduced mortality with DCC vs ICC, but no difference in morbidity or transfusion rates.(127)

884 A network meta-analysis comparing short (15-45s), medium (45-120s), and long ( $>$ 120s) cord clamping  
885 times with ICC and cord milking found the strongest survival benefit with longer delays (mortality OR  
886 0.31, 95% CI 0.11-0.80). (128) They concluded that for newborns requiring resuscitation/stabilisation,  
887 longer DCC is only feasible with intact umbilical cord. (128)

888 Three multi-centre RCTs on intact cord resuscitation have been completed. Two used fixed clamping  
889 times and one used physiological criteria. The VentFirst trial (<29 weeks) found no difference in  
890 intraventricular haemorrhage or mortality between 120s DCC with intact cord ventilation vs DCC 30-  
891 60s and ventilation afterwards. (129) No difference was reported in the composite outcome of death,  
892 severe intraventricular haemorrhage, and bronchopulmonary dysplasia between 3-minute intact cord  
893 resuscitation and cord milking. (130) In the ABC3 trial physiologically based cord clamping vs 30-60s  
894 DCC showed no overall difference in intact survival, but improved outcomes in male infants and with  
895 increased intact cord resuscitation experience. (131)

### 896 [h3] Umbilical cord milking

897 Umbilical cord milking has been considered an alternative to DCC when DCC is not feasible. (132) In  
898 'intact cord milking', the cord is milked 3–5 times before clamping, promoting faster blood transfer. In  
899 'cut cord milking', a ~25 cm cord segment is milked after clamping, usually during resuscitation.  
900 (132) Experimental studies show intact cord milking causes significant fluctuations in cerebral blood  
901 flow. (133, 134) A large clinical trial in preterm infants was stopped early due to increased risk of  
902 severe intraventricular haemorrhage in the <28 weeks GA subgroup who were randomised to  
903 umbilical cord milking. (135) Meta-analyses in preterm infants showed no differences in mortality or  
904 morbidity. (127, 128) Umbilical cord milking reduced transfusion need compared to ICC, but to DCC. A  
905 recent cluster RCT in 1730 non-vigorous infants ≥35 weeks found no difference in mortality or NICU  
906 admission between intact cord milking and ICC. (136) The reported reduction in moderate-to-severe  
907 hypoxic ischemic encephalopathy (HIE) (RR 0.49, 95% CI: 0.25–0.97) was based on unadjusted data and  
908 may reflect later clamping. The ERC recommends that for all infants the focus should be on DCC  
909 instead of umbilical cord milking. The ERC recommends avoiding cord milking in infants <28 weeks,  
910 acknowledging that intact cord milking is an alternative to DCC in infants ≥28 weeks, only if DCC  
911 cannot be performed.

912

## 913 [h2] Initial assessment

### 914 [h3] Breathing

915 Not crying may be due to apnoea and can function as a marker of inadequate breathing needing  
916 support. (137) In an observational study of almost 20,000 infants (>22 weeks GA) just after birth in a  
917 rural hospital setting, 11% were not crying, around half of whom were assessed as apnoeic. About 10%  
918 of those assessed as breathing at birth became apnoeic. Breathing without crying compared to  
919 breathing and crying was associated with a 12-fold increase in morbidity. (137) The presence or  
920 adequacy of breathing effort in preterm infants can be difficult to assess as breathing can be very  
921 subtle and is often missed. (138, 139) When breathing was perceived as inadequate infants were more

922 likely to receive interventions. (140, 141) The ERC recommends assessing rate, depth, symmetry and  
923 work of breathing.

### 924 [h3] Heart rate

925 HR is the most sensitive indicator of a successful response to resuscitation interventions. (124, 142,  
926 143) There is no published evidence clearly defining the thresholds for intervention during newborn  
927 resuscitation. Historically, heart rates of  $>100 \text{ min}^{-1}$  were pragmatically selected as reassuring and  $<60$   
928  $\text{min}^{-1}$  as prompting interventions. (144) A 2023 ILCOR review found no new evidence on alternative HR  
929 thresholds. (145) In uncompromised breathing term infants undergoing DCC, the HR is usually above  
930  $100 \text{ min}^{-1}$ . (143) In an observational study in resuscitated term/near-term infants, initial HRs at birth  
931 were distributed showing bimodal peaks around 60 and  $165 \text{ min}^{-1}$ . (146) In preterm infants  $<30$  weeks  
932 the HR did not stabilise until it reaches approximately  $120 \text{ min}^{-1}$  and, in some, stability was only  
933 achieved once the HR was  $>150 \text{ min}^{-1}$ . (147) A recent study in extremely or very preterm neonates  
934 with favourable outcome established that the 10<sup>th</sup> percentile of HR at 2, 5, 10, and 15 minutes after  
935 birth were 70, 109, 126, and  $134 \text{ min}^{-1}$  respectively, indicating varying expected HR values during the  
936 immediate postnatal transition period. (148)

### 937 [h4] Heart rate assessment

938 The main methods of HR assessment are auscultation, pulse oximeter, and ECG. The advantages and  
939 disadvantages of these are summarised in Table 7. Most studies excluded newborns who were  
940 bradycardic at birth, required resuscitation or were very preterm infants, which limits applicability of  
941 study results. (69, 149, 150) Auscultation by stethoscope is simple and enables rapid assessment of HR  
942 in any setting, including low resource settings (ERC practice statement). The 2024 ILCOR review  
943 suggests that if resources allow, ECG to continuously assess HR is reasonable, with pulse oximeter and  
944 auscultation as alternatives. (149, 150) It is currently unclear if speed/precision of HR assessment at  
945 birth is associated with clinically important differences in interventions, performance or outcomes.  
946 (149, 150) There is insufficient evidence to recommend the use of digital stethoscopes, Doppler  
947 ultrasound, dry electrode technology or other techniques to assess HR at birth. (149, 150) The ERC  
948 recommendations align with ILCOR. Initial HR assessment can be done by auscultation; continuous HR  
949 assessment through ECG or pulse oximetry are recommended with ongoing resuscitation.

950 << INSERT Table 7: Methods of HR assessment and evidence >>

### 951 [h3] Tactile Stimulation

952 ILCOR systematic reviews on both cord management and tactile stimulation suggest tactile stimulation  
953 immediately after birth in infants with inadequate breathing efforts, regardless of method of umbilical  
954 cord management. (140, 151, 152) Tactile stimulation should not delay the initiation of breathing  
955 support if required. The optimal type and length of tactile stimulation as well as differences in different

956 gestational ages is unknown. (152) An RCT in a preterm population reported that repetitive stimulation  
957 improved oxygen saturations and reduced need for supplemental inspired oxygen. (153) Data from an  
958 observational study shows that tactile stimulation at birth is associated with more spontaneous  
959 breathing, especially if the cord was still intact. (137) The ERC recommends performing tactile  
960 stimulation on all newborn infants at birth, especially if breathing is inadequate, but it must not delay  
961 initiation of breathing support if required.

### 962 [h3] Tone and Colour

963 Healthy infants are cyanosed at birth, reflecting lower *in utero* saturations, but this improves within  
964 approximately 30s of the onset of effective breathing. (139) Peripheral cyanosis is common and does  
965 not, by itself, indicate hypoxia. Persistent pallor despite ventilation may indicate significant acidosis,  
966 or, more rarely, hypovolaemia with intense cutaneous vascular vasoconstriction. Colour is an  
967 unreliable marker of oxygenation, and it should not be used to judge oxygenation. (154) The ERC  
968 recommends using pulse oximetry to measure oxygen saturations in preference to using colour as a  
969 proxy for oxygenation.

970

### 971 [h2] Airway

972 Airway obstruction is most commonly caused by suboptimal airway positioning, lack of pharyngeal  
973 tone, and adducted vocal cords especially in preterm newborns. (155, 156) There is no evidence that  
974 normal lung fluid and secretions cause obstruction. (157) In line with ILCOR, the ERC recommendation  
975 is not to routinely suction clear fluid from the oropharynx.

### 976 [h3] Position

977 With flexion and extension of the neck, the newborn airway can easily become occluded. (158)  
978 Evidence on the mechanisms of airway occlusion in the newborn is limited. A retrospective analysis of  
979 images of the airway of 53 sedated infants between 0-4 months undergoing cranial MRI indicates how,  
980 in extension, obstruction might occur through anterior displacement of the posterior airway at the  
981 level of the tongue. (159) Video review of airway position and airway obstruction also found  
982 hyperextension of the neck is associated with airway obstruction. (160) Therefore, the ERC  
983 recommends a neutral head position to ensure optimal airway patency in newborn infants.

### 984 [h3] Jaw thrust and two-person method

985 Studies in children demonstrate that anterior displacement of the mandible enlarges the pharyngeal  
986 space through lifting the epiglottis away from the posterior pharyngeal wall, reversing the narrowing  
987 of the laryngeal inlet. (161) Two-person manual ventilation techniques are superior to single handed  
988 airway support: it reduces mask leak and is more effective (158, 162-164), which is recommended by  
989 the ERC.

**990 [h3] Preterm newborns**

991 Vocal cord adduction is a cause of airway obstruction at birth in preterm infants <30 weeks. (156) In an  
992 observational study of 56 preterm infants <32 weeks significant mask leak (>75%) and/or obstruction  
993 to inspiratory flow (75%) were identified using respiratory function monitoring in 73% of interventions  
994 during the first 2 minutes of PPV. (165) In an animal model of premature birth, phase contrast X-rays  
995 demonstrated that the larynx and epiglottis were predominantly closed (adducted) in those with un-  
996 aerated lungs and unstable breathing patterns, making intermittent PPV ineffective unless there was  
997 an inspiratory breath, and only opening once the lungs were inflated. (155) This may be an explanation  
998 for the challenges in inflating preterm infant lungs, but a solution for overcoming these phenomena is  
999 not yet known.

**1000 [h3] Suction**

1001 Routine oropharyngeal and nasopharyngeal suction in newborn infants has not been shown to  
1002 improve respiratory function, may delay other necessary manoeuvres and the onset of spontaneous  
1003 breathing and is associated with adverse events. (166-170)

1004 The ERC, following ILCOR, does not recommend routine intrapartum oropharyngeal and  
1005 nasopharyngeal suction for newborn infants with clear or meconium-stained amniotic fluid. (157) If  
1006 suctioning is attempted it should be undertaken under direct vision, ideally using a laryngoscope and a  
1007 wide-bore catheter or Yankauer sucker. Bulb suction can be useful if no vacuum source is available. A  
1008 meconium aspirator, attached to a tracheal tube, can clear thick material from the trachea, applied  
1009 suction should not exceed 150 mmHg (20 kPa). (171, 172)

**1010 [h3] Meconium**

1011 Lightly meconium-stained liquor is common and usually does not cause difficulty with transition. Non-  
1012 vigorous newborns delivered through meconium-stained amniotic fluid are at significant risk for  
1013 requiring advanced resuscitation and a neonatal team competent in advanced resuscitation may be  
1014 required. Routine suctioning of non-vigorous infants can delay initiating ventilation and there is no  
1015 evidence to support intrapartum suctioning nor routine tracheal intubation and suctioning of vigorous  
1016 infants born through meconium-stained liquor. (173-175) Evidence from retrospective registry-based  
1017 studies (176, 177), meta-analyses (178-180), a post policy change impact analysis (181), and ILCOR  
1018 2025 (ILCOR 2025) all support omitting suctioning in favour of immediate ventilation.

1019 The ERC recommends against routine suction of either pharynx or trachea in newborn infants born  
1020 through meconium -stained liquor and recommend providing standard NLS. If there is evidence of  
1021 airway obstruction, the ERC recommends suction under direct vision in the first instance. Rarely,  
1022 airway obstruction may occur below the level of the larynx, and this may require tracheal suction.

**1023 [h3] Airway devices**

**1024 [h4] Supraglottic airway devices (SGA)**

1025 SGAs are effective in newborns, particularly if facemask ventilation or tracheal intubation is  
1026 unsuccessful or not feasible. (69) A systematic review showed that SGAs were more effective than bag-  
1027 mask ventilation in terms of shorter resuscitation and shorter duration of ventilation with less need for  
1028 tracheal intubation. (182) Bag-mask ventilation was effective in more than 80% of enrolled infants.  
1029 Efficacy of an SGA was comparable to tracheal intubation.

1030 Aligned with ILCOR, ERC recommends using an SGA as a valid alternate airway device, particularly if  
1031 tracheal intubation is unsuccessful or intubation skills are unavailable. (69)(ILCOR 2025)  
1032 Studies generally included infants with birth weight >1500 g or GA≥34 weeks, so evidence supporting  
1033 SGAs in more premature infants is limited. (182, 183) A 2024 Cochrane update found no or little  
1034 difference in neonatal morbidities and mortality when giving surfactant via an SGA compared with a  
1035 tracheal tube. (184) The SGA has not been evaluated in the setting of meconium-stained fluid, during  
1036 chest compressions, or for the administration of emergency intra-tracheal medications. ILCOR  
1037 considers it reasonable to use an SGA during chest compressions if tracheal intubation is not  
1038 possible/unsuccessful (good practice statement) (ILCOR 2025) and the ERC aligns with this.

**1039 [h4] Oropharyngeal airway**

1040 Although the oropharyngeal airway is effective in children (185), there is no published evidence  
1041 demonstrating effectiveness in maintaining airway patency at birth. In an RCT of 137 preterm  
1042 newborns where gas flow through a mask was measured, obstructed inflations were more common in  
1043 the oropharyngeal compared to control group.(186) However, by helping lift the tongue and  
1044 preventing it occluding the laryngeal opening, an oropharyngeal airway may facilitate airway support  
1045 where difficulty is experienced and where other airway opening techniques, like jaw thrust, fail to  
1046 improve ventilation.

**1047 [h4] Nasopharyngeal airway**

1048 A nasopharyngeal airway may help establish an airway where there is congenital upper airway  
1049 abnormality (187) and has been used successfully in preterm infants at birth. (138, 185-187)

**1050 [h4] Tracheal tube**

1051 Safe tracheal intubation is facilitated by well-trained clinicians with appropriate equipment, and the  
1052 use of an intubation checklist. (188) Once insertion, confirmation of tracheal tube position by clinical  
1053 assessment and the use of exhaled CO<sub>2</sub> detection is required. (189) Contingency plans should be made  
1054 for an unexpectedly difficult airway. Declining intubation skills mean unsuccessful intubation is more  
1055 common and safe airway management around intubation attempts is vital. (190, 191)

1056 << INSERT Table 8. Approximate tracheal tube size by gestation and approximate lengths for oral and  
1057 nasal intubation (11, 192) >>

**1058 [h4] Video Laryngoscopy**

1059 A 2024 ILCOR systematic review of video laryngoscopy versus direct laryngoscopy (193, 194) found  
1060 higher overall intubation success rates and higher first attempt success rates using a video  
1061 laryngoscopy compared to a direct laryngoscope. These findings are confirmed by a 2025 systematic  
1062 review. (195)

1063 Where resources and training allow, the ERC recommends using a VL to intubate newborn infants,  
1064 especially in settings where less experienced staff are intubating. Direct laryngoscopy remains a  
1065 reasonable option, and such a laryngoscope should be available as a backup device.

**1066 [h3] Exhaled CO<sub>2</sub>**

1067 Detection of exhaled CO<sub>2</sub> alongside clinical assessment is used to confirm tube placement in the  
1068 trachea in newborn infants, even from >400g. (11, 196-199) Failure to detect exhaled CO<sub>2</sub> strongly  
1069 suggests tube misplacement. (197, 200) Studies relating to exhaled CO<sub>2</sub> have mostly excluded infants  
1070 in need of extensive resuscitation. False negative CO<sub>2</sub> detection can occur in cases of poor or absent  
1071 pulmonary blood flow, tracheal obstruction, low or absent cardiac output in resuscitation at birth and  
1072 in birthweight <1500g. (198, 200) Where there is no CO<sub>2</sub> detection after tracheal intubation, tracheal  
1073 tube position should be rechecked by VL or direct laryngoscopy to avoid unnecessary tracheal tube  
1074 removal.

1075 Like ILCOR (199), the ERC recommends using exhaled CO<sub>2</sub> detection combined with clinical  
1076 assessment, to confirm tracheal tube placement.

1077 Both qualitative (colorimetric) and quantitative (waveform) CO<sub>2</sub> detection methods have been  
1078 successfully used in intubated newborn infants (201). Colorimetric detection offers a simple, easy to  
1079 use, cheap device where a colour change indicates exhaled CO<sub>2</sub>. Waveform detection provides a  
1080 continuous graphical and numerical representation of exhaled CO<sub>2</sub> throughout the respiratory cycle,  
1081 allowing for continuous monitoring, but requires specialised equipment and power sources may not be  
1082 readily available in all delivery area settings. Colorimetric detection failed to detect correct tube  
1083 placement in one-third of delivery area intubation in very preterm infants in one study. (202).

1084 Although waveform capnography is more sensitive in adults, limited newborn data advise caution,  
1085 especially if being used during resuscitation. (203-205) The ERC can not recommend one method over  
1086 the other.

1087 Exhaled CO<sub>2</sub> can be used in non-intubated patients (206-209) Exhaled CO<sub>2</sub> detector use with interfaces  
1088 such as SGAs are standard in adult patients, but as newborn physiology differs markedly from that of  
1089 older children and adults, practices of proven benefit for older patients may not apply to neonatal  
1090 patients, especially during perinatal transition.

1091 The ERC currently can not recommend the routine use of exhaled CO<sub>2</sub> detection in non-intubated  
1092 newborn infants in the delivery area.

### 1093 [h3] Respiratory flow monitoring

1094 Flow monitoring via a respiratory function monitor was reported in an RCT to confirm tracheal tube  
1095 position faster and more reliably than end-tidal CO<sub>2</sub> detection. (210) An ILCOR systematic review (211)  
1096 and a 2025 evidence update (ILCOR 2025) found insufficient evidence to recommend for or against the  
1097 use of respiratory function monitoring at birth, and ERC recommendations align with ILCOR.

1098

## 1099 [h2] Breathing

### 1100 [h3] Initial inflations and assisted ventilation

1101 Lung inflation must begin without delay in apnoeic or inadequately breathing newborn infants. An  
1102 observational study in low-resource settings found a 16% increase in morbidity/mortality for every 30s  
1103 delay in starting ventilation. (212) Optimal inflation pressure, inspiratory and expiratory times, and  
1104 duration of PPV remain uncertain.

### 1105 [h3] Facemask

1106 Facemask ventilation is limited by leaks, often caused by poor mask fit or suboptimal technique, both  
1107 of which contribute to mask leak which can be improved after training. (163, 213) A clinical study  
1108 demonstrated obstruction and/or leak (>75%) during initial ventilations in preterm infants. (165) An  
1109 observational study in preterm infants <32 weeks suggested that the application of a facemask to  
1110 support breathing might induce apnoea by triggering the trigeminocardiac reflex in spontaneously  
1111 breathing infants. However, the significance of this is currently unclear (214).

### 1112 [h3] Nasal prongs

1113 While facemasks are most commonly used, nasal interfaces (prongs or masks) have been found to be  
1114 as effective as facemasks. (215) Emerging studies suggest nasal prongs also reduce delivery room  
1115 intubation and PPV use in infants <28 weeks. (216, 217) (218). The ERC recommends performing PPV  
1116 using either a facemask or nasal prongs.

### 1117 [h3] Inflation duration

1118 Initial inflations or spontaneous breathing establish functional residual capacity (FRC). (219, 220) There  
1119 is ongoing debate about the optimal inflation duration. (221-229) This should not be confused with  
1120 sustained inflations (i.e. inflations ≥5s), which are not recommended by ILCOR or the ERC. (11, 31)  
1121 Previous ERC guidance recommended 2-3s inflations (11), while other NLS guidelines around the world  
1122 support shorter durations of inflations (~1s). (230, 231) Limited evidence shows no clear advantage of  
1123 longer (2-3s) over shorter (~1s) inflations (229, 230, 232, 233), so the ERC permits either approach.

1124 As a HR response may not be seen until at least 20s of PPV in bradycardic infants, (234, 235) the  
1125 number of inflations recommended varies depending on the length of inflation. The ERC recommends  
1126 either 5 inflations of 2-3s, or 10 inflations of ~1s, based on pragmatic consensus.

### 1127 [h3] Inflation pressure

1128 Inflation pressures of 30 cm H<sub>2</sub>O are usually sufficient to inflate the fluid-filled lungs of apnoeic term  
1129 infants, based on historical cohort studies. (220, 236, 237) However, a prospective cohort study of 821  
1130 term and near-term infants resuscitated using bag-mask ventilation found median peak pressures of  
1131 36 cmH<sub>2</sub>O required for successful stabilisation. (238) For preterm infants, an initial inflation pressure of  
1132 25 cmH<sub>2</sub>O is considered reasonable (224, 239-241), though higher pressures may be needed due to  
1133 greater airway resistance. If no chest movement is observed, the ERC recommends increasing inflation  
1134 pressures, regardless of GA, to achieve lung inflation.

### 1135 [h3] Ventilation rate

1136 Evidence on the optimal ventilation rate for newborn resuscitation is limited. In an observational study  
1137 of 434 late preterm and term infants, ventilation at 30 breaths min<sup>-1</sup> achieved adequate tidal volumes  
1138 without hypocarbia, with the best CO<sub>2</sub> clearance at 10-14 mL/kg. (242) An observational study suggest  
1139 that PPV rates >60 min<sup>-1</sup> compared to rates < 60 min<sup>-1</sup> fail to achieve adequate tidal volumes. (243).  
1140 Other studies suggest optimum rates for PPV are 30-40 min<sup>-1</sup>. (220, 236). The ERC recommends PPV  
1141 rate of 30-40 min<sup>-1</sup> once lungs have been inflated.

### 1142 [h3] Effectiveness of inflations

1143 The primary sign of adequate lung inflation is a rapid HR increase, usually within 20-30s of onset of  
1144 effective ventilations. (142, 244, 245) Chest wall movement usually indicates lung inflation, although  
1145 this may be less obvious in preterm infants. (246) Excessive chest movements may indicate excessive  
1146 tidal volumes, which should be avoided. If HR improves but breathing remains inadequate, PPV must  
1147 continue.

1148 Failure of the HR to increase is most often due to suboptimal airway control or inadequate ventilation.  
1149 (163, 213, 247) Adjustments to head/airway position (158), choosing alternative airway opening  
1150 techniques, or increased inflation pressures may be needed. (238) In preterm infants, mask pressure,  
1151 glottal closure, or triggering the trigeminocardiac reflex may impair PPV. (155, 165, 248, 249) Although  
1152 exhaled CO<sub>2</sub> monitoring can sometimes detect such obstructions and mask leaks, current evidence is  
1153 insufficient to recommend its routine use to assess quality of PPV. (209)

### 1154 [h3] Sustained inflations >5 seconds

1155 Animal studies suggested longer inflations may have physiological benefits (250, 251), but clinical  
1156 benefits in human infants have not been demonstrated. In preterm infants, there is evidence of  
1157 possible harm from sustained inflation >5s. (252) A Cochrane systematic review found that sustained

1158 inflations (15-20s) were not better than intermittent ventilation ( $\leq 1s$ ) for reducing mortality,  
1159 intubation, need for respiratory support or bronchopulmonary dysplasia.(253) An ILCOR review advises  
1160 against routine use of sustained inflations  $>5s$  in preterm infants receiving PPV at birth, due to a  
1161 possible increase in mortality in infants  $<28$  weeks. ((227, 229) ILCOR did not recommend a specific  
1162 inflation duration for late preterm and term infants due to low-confidence in the estimates of the  
1163 effect. The ERC guidance aligns with ILCOR and recommends against sustained inflations  $>5s$  in  
1164 preterm infants.

### 1165 [h3] CPAP and PEEP

1166 Successful respiratory transition at birth relies on alveolar aeration, lung inflation, and formation of  
1167 FRC. (254) Most premature infants can breathe at birth but often struggle to obtain and maintain FRC.  
1168 (255, 256)(251) The need for respiratory support at birth is inversely correlated to GA. (257, 258)  
1169 Animal studies show that a few early inflations at high tidal volumes can cause lung injury and  
1170 inactivate surfactant. (259, 260) Preclinical studies demonstrated that applying CPAP or PEEP  
1171 immediately after birth assists lung inflation. (261) Unlike CPAP, PEEP is only present during exhalation  
1172 and is applied during manual or mechanical ventilation. (262) While other non-invasive respiratory  
1173 supports are under investigation, CPAP remains the gold-standard for newborn infants  $<32$  weeks.  
1174 (263)

### 1175 [h4] CPAP for infants $<32$ weeks

1176 Large RCTs show that starting CPAP at birth, compared with intubation and ventilation, significantly  
1177 reduces death and bronchopulmonary dysplasia. (264-268) An ILCOR systematic review recommends  
1178 starting CPAP promptly in spontaneously breathing preterm infants with respiratory distress instead of  
1179 TI and PPV. (31, 269)(ILCOR 2025). The ERC guidelines align with this recommendation. Whilst some  
1180 RCTs used CPAP levels up to 8 cmH<sub>2</sub>O (264, 265), an observational study shows that levels of 5 - 6  
1181 cmH<sub>2</sub>O are most commonly used in practice. (270) Comparative studies on optimal CPAP levels remain  
1182 limited.(271, 272) A 2021 Cochrane Review concluded that it was not possible to recommend a specific  
1183 starting level. (272) Animal data suggest that CPAP at 15 cmH<sub>2</sub>O (with O<sub>2</sub> 60%) improves lung inflation  
1184 compared to 4 – 8 cmH<sub>2</sub>O.(273) An ongoing trial is studying dynamic CPAP (8-12 cm H<sub>2</sub>O) vs static  
1185 CPAP (6 cm H<sub>2</sub>O) at birth. (clinicaltrials.gov NCT04372953)  
1186 Until higher-quality evidence is available, and based on indirect measures showing better lung inflation  
1187 at 6 cm H<sub>2</sub>O (263), aligning with the European Consensus Guidelines on the Management of  
1188 Respiratory Distress Syndrome (274), the ERC recommends starting CPAP at 6 cm H<sub>2</sub>O in  
1189 spontaneously breathing preterm infants  $<32$  weeks.

### 1190 [h4] CPAP for infants $\geq 32$ weeks

1191 The 2022 ILCOR CoSTR, states that there is insufficient evidence for or against routine CPAP in late  
1192 preterm and term infants. (275) However, late preterm infants and term infants with conditions such as  
1193 transient tachypnoea of the newborn, or those requiring supplemental O<sub>2</sub>, may benefit from CPAP (good  
1194 practice statement).(276) The ERC considers it reasonable to start CPAP at 6 cm H<sub>2</sub>O in newborn infants  
1195 ≥32 weeks with respiratory distress needing supplemental O<sub>2</sub>.

#### 1196 [h4] PEEP during PPV

1197 Self-inflating bags can be equipped with PEEP valves to deliver defined PEEP during PPV, but cannot  
1198 provide CPAP, even with attached bias gas flow. (277) ILCOR recommends using PEEP during initial PPV  
1199 of preterm newborn infants at birth (31). Accordingly, the ERC recommends starting with a PEEP of 6  
1200 cmH<sub>2</sub>O in preterm newborn infants receiving PPV.

#### 1201 [h3] Assisted ventilation devices

1202 Recent reviews summarize the principles of interfaces, devices and settings for delivering CPAP, PEEP  
1203 and PPV during fetal-to-neonatal transition.(277) (278) T-piece resuscitators deliver more consistent  
1204 CPAP/PEEP compared to self-inflating bags.(277) ILCOR (2021) concluded T-piece resuscitators may  
1205 slightly improve outcomes like survival, intraventricular haemorrhage and bronchopulmonary  
1206 dysplasia over self-inflating bags.(279, 280) Thus, the ERC recommends using T-piece resuscitators for  
1207 PPV at birth, but self-inflating bags should be available as a backup if gas supply fails.

1208

#### 1209 [h2] Oxygen

#### 1210 [h3] Term infants and late preterm infants ≥ 32 weeks

1211 Lower inspired O<sub>2</sub> concentrations may result in suboptimal oxygenation where there is significant lung  
1212 disease (281), while higher O<sub>2</sub> may delay spontaneous breathing in term infants. (282) ILCOR  
1213 recommended starting with 21% O<sub>2</sub> in infants ≥35 weeks receiving respiratory support at birth, and  
1214 advises against 100% O<sub>2</sub>.(31) An updated systematic review (2164 patients) demonstrated 27% lower  
1215 short-term mortality with room air vs 100% O<sub>2</sub>, without differences in neurodevelopment or hypoxic  
1216 ischaemic encephalopathy (HIE).(283) For neonates born at 32-34<sup>+6</sup> weeks, ILCOR found insufficient  
1217 evidence for specific O<sub>2</sub> recommendations. (ILCOR 2025) The ERC recommends starting with 21% O<sub>2</sub> in  
1218 infants ≥32 weeks and titrating O<sub>2</sub> to achieve target saturations.

#### 1219 [h3] Preterm infants <32 weeks

1220 In preterm infants, higher supplemental O<sub>2</sub> improves breathing effort and oxygenation and results in  
1221 shorter mask ventilation and higher minute volumes. (248, 284)

1222 The NetMotion individual patient data network meta-analysis (1055 infants from 12 studies)  
1223 suggested high O<sub>2</sub>>90% may reduce all-cause mortality compared to lower O<sub>2</sub> (<30% and 50-  
1224 65%). (285). An updated ILCOR study level meta-analysis found insufficient evidence to definitively

1225 recommend high (>50%) vs low ( $\leq$ 50%) O<sub>2</sub> (1804 infants from 16 studies + NetMotion, infants <35  
1226 weeks) (285, 286) [ILCOR 2025]. For infants <32 weeks, the ERC recommends starting resuscitation  
1227 with  $\geq$ 30% O<sub>2</sub> and adjusting O<sub>2</sub> to achieve and maintain target saturations.

### 1228 [h3] Target oxygen saturations

1229 In 2010 target oxygen saturation curves were published, however, these data predated DCC and the  
1230 majority of included infants were  $\geq$ 37 weeks. (287) In 2024, different saturation and HR references  
1231 were published for infants <32 weeks stabilised according to more current guidelines. All included  
1232 infants had favourable outcomes, defined as survival without cerebral injury. (148) Most (92%) of the  
1233 cohort received O<sub>2</sub> and CPAP (91%) (148). Table 9 provides an overview of target saturation ranges  
1234 that are also used in the Newborn Resuscitation Programme (NRP).(230) (148, 287)

1235 A systematic review showed that failing to reach SpO<sub>2</sub>  $\geq$ 80% at 5 minutes doubled the risk of death  
1236 and severe intraventricular haemorrhage in very preterm infants. (288)

1237 Nearly all infants <32 weeks require supplemental O<sub>2</sub> after birth (148, 287, 288), but achieving target  
1238 saturations in the minutes after birth can be challenging; only 12% reached 80% SpO<sub>2</sub> at 5 minutes of  
1239 life. (288) SpO<sub>2</sub> readings <60% are considered inaccurate (289). Dark skin tones may be associated with  
1240 oxygen saturation discrepancies, with a higher incidence of occult hypoxaemia (290) , although limited  
1241 data suggests that the discrepancy may be less pronounced in neonates. (291, 292)

1242 There is no direct evidence for the optimal oxygen saturations to strive for after birth. The ERC  
1243 provides a consensus-based recommendation on uniform oxygen saturation targets across all GAs,  
1244 balancing out the perceived detrimental effects of hypoxia that may be worse than those for hyperoxia  
1245 (table 4).

1246 <<INSERT Table 4. Target saturations>>

1247 << INSERT Table 9. Overview of oxygen saturation targets >>

1248 << INSERT figure 8: Oxygen saturations in healthy infants at birth without medical intervention (3<sup>rd</sup>, 10<sup>th</sup>,  
1249 25<sup>th</sup>, 50<sup>th</sup>, 75<sup>th</sup>, 90<sup>th</sup>, 97<sup>th</sup> centiles. Reproduced with permission - from Dawson 2010 e1340 (287). >>

1250 << INSERT figure 9: Oxygen saturations during the first 15 minutes after birth of infants<32 weeks with  
1251 favorable outcome (10th, 50th, and 90th centile (bold lines) of A, SpO<sub>2</sub> (%); (10th, 50th, and 90th centile  
1252 (bold lines), (gray lines: 5th, 15th, 20th, 25th, 30th, 35th, 40th, 45th, 55th, 60th, 65th, 70th, 75th, 80th,  
1253 85th, 95th centile). From Wolfsberger et al 2024 (148).

### 1254 [h3] Titration of oxygen

1255 Timely adjustment in delivery of O<sub>2</sub> is critical to avoid hypoxia, hyperoxia and bradycardia. The ERC  
1256 recommends reviewing O<sub>2</sub> every 30s (293) and adjusting O<sub>2</sub> to achieve target SpO<sub>2</sub>. There can be a  
1257 delay between titration of intended O<sub>2</sub> and delivery of O<sub>2</sub> to the baby. One study suggests a T-piece

1258 resuscitator takes 19s (IQR 0-57) to achieve the desired O<sub>2</sub> at the distal end, (294) and another that  
1259 nasal prongs may reduce this delay.(295)

### 1260 [h3] Cerebral tissue oxygenation monitoring

1261 Using the same population of preterm infants, the application of different statistical methodology has  
1262 resulted in differing conclusions. (296-298) The ERC, following the 2024 ILCOR recommendation,  
1263 recommends that near infrared spectroscopy monitoring of cerebral oxygenation in the delivery  
1264 room should only be considered where resources permit, preferably within structured research trials  
1265 to help close knowledge gaps. (297)(ILCOR 2025)

1266

### 1267 [h2] Circulatory support

1268 Circulatory support with chest compression is effective only after successful lung inflation and  
1269 subsequent oxygen delivery to the heart. Ventilation may be compromised during chest compressions,  
1270 so ensuring effective ventilation before starting chest compression is critical. (299)

### 1271 [h3] Threshold for initiating and discontinuing chest compressions

1272 The HR threshold to initiate chest compression at birth (<60 min<sup>-1</sup>) was based on expert opinion  
1273 and limited animal data. (145, 300) No human studies have compared different HR thresholds for  
1274 initiating chest compression in human newborns (145), and current practice remains to start chest  
1275 compression if HR is <60 min<sup>-1</sup> after successful lung inflation.

1276 In asystolic piglets, starting chest compression after 30 or 60s of PPV resulted in comparable  
1277 outcomes, however, delaying chest compression beyond 90s worsened outcomes. (144). Two  
1278 narrative reviews reconsidered the HR threshold for starting chest compression at birth (300) (301),  
1279 suggesting chest compression could potentially be delayed an additional 30s of PPV, if HR is rising after  
1280 30s of ventilation, but still <60 min<sup>-1</sup>, though more research is needed.

1281 The ERC recommends considering an additional 30s of PPV when HR is still <60 min<sup>-1</sup> but increasing.

1282 The ERC also recommends checking the HR every 30s unless using continuous monitoring (pulse  
1283 oximetry, ECG). Whilst chest compression may be discontinued when HR is >60 min<sup>-1</sup>, a continued  
1284 increase in rate and confirmation of cardiac output, e.g. auscultation, pulse check, pulse oximetry,  
1285 signs of life, are necessary to truly demonstrate improvement. Stability often occurs only when HR  
1286 exceeds 120 min<sup>-1</sup>.(146, 147)

### 1287 [h3] Compression technique

1288 ERC recommendations align with ILCOR, whose 2023 systematic review reaffirmed that for infants at  
1289 birth, the two-thumb-hands-encircling-the-chest method should be used to deliver CC, because it  
1290 results in improved compression depth, less fatigue, and better digit placement than the two-finger  
1291 technique. (145)(ILCOR 2025) Two overlapping or adjacent thumbs should be placed on the lower

1292 third of the sternum from either the lateral or over-the-head position. (302, 303) The over-the-head  
1293 position may facilitate umbilical catheterisation. Alternative techniques were also considered but were  
1294 not superior.(145)

### 1295 [h3] Compression depth

1296 In a post-transitional piglet model, compressions of 25-40% depth achieved ROSC, while 12.5%  
1297 compression depth did not.(304) Evidence in human newborn infants is lacking(145), although deeper  
1298 compressions improved blood pressures in post-surgical infants.(305). Full recoil between  
1299 compressions is important. (306-310) The ERC recommends compressing the sternum one-third of the  
1300 anterior-posterior chest diameter (good practice statement), allowing full recoil between  
1301 compressions.

### 1302 [h3] Compression-to-ventilation ratio

1303 ILCOR (2023) found insufficient evidence to change the recommended 3:1 C:V ratio, aiming for 90  
1304 compressions and 30 ventilations per minute. (31, 145) However, the quality of compressions and  
1305 ventilations is probably more important than the rate. (311) Animal studies suggested chest  
1306 compression with sustained inflations improved outcomes over 3:1 C:V, but human trials remain  
1307 inconclusive. (311-314). The ERC continues to recommend a 3:1 C:V ratio for resuscitation at birth,  
1308 even after securing the airway.

### 1309 [h3] Supplemental oxygen during chest compressions

1310 Available evidence remains insufficient to alter the recommendation of increasing O<sub>2</sub> to 100% when  
1311 starting chest compression (good practice statement). (11, 145) No human studies have compared  
1312 21% with 100% O<sub>2</sub> (or any other O<sub>2</sub> concentration) during chest compression (315), and animal  
1313 studies reported no major differences in time to ROSC, mortality, inflammation, or oxidative  
1314 stress between concentrations. (145, 315) Both hypoxia and hyperoxia can be detrimental (145,  
1315 315). In a transitional term ovine model of asphyxia-induced cardiac arrest, 21% O<sub>2</sub> was  
1316 associated with lower cerebral oxygen levels and higher brain lactic acid after ROSC compared to  
1317 were reported with the 100% O<sub>2</sub>.(316) Rapid weaning of inspired O<sub>2</sub> after ROSC may prevent  
1318 hyperoxia and thereby possibly mitigate oxidate stress and organ damage. Thus, ERC recommends  
1319 that, once HR recovers, O<sub>2</sub> should be actively reduced, guided by pulse oximetry (good practice  
1320 statement).

### 1321 [h3] Prompts and feedback devices

1322 Earlier studies suggested exhaled CO<sub>2</sub> monitoring and pulse oximetry may be useful in detecting ROSC.  
1323 (317-320) ILCOR reviewed 16 studies examining chest compression in relation to (audio)visual  
1324 feedback devices, auditory feed forward devices, audiovisual prompts provided by a decision  
1325 support tool, capnography, and blood pressure monitoring, but findings were difficult to compare

1326 due to heterogeneity. (145) Currently, ERC cannot recommend the clinical use of prompts or  
1327 feedback devices to assess chest compression during neonatal resuscitation

### 1328 [h3] Automated chest compression devices

1329 Mechanical chest compression devices are used in adults but not yet in newborns. (321) In a neonatal  
1330 asphyxiated piglet model, machine-delivered chest compression improved stroke volume and left  
1331 ventricular contractility compared to manual chest compression. (322) More research is needed  
1332 before clinical use in human newborn infants can be recommended.

1333

## 1334 [h2] Vascular access

### 1335 [h3] Umbilical venous catheter (UVC) and intraosseous (IO) access

1336 No new evidence was identified comparing umbilical venous catheter (UVC) route or use of  
1337 intravenous (IV) cannulas against the intraosseous (IO) route in the newborn for drug administration in  
1338 any setting in an ILCOR systematic review. (31) A systematic review on the use of IO in neonates in any  
1339 situation identified one case series and 12 case reports of IO device insertion into 41 neonates  
1340 delivering several drugs including adrenaline and fluid/blood. (323) First attempt success rates for IO  
1341 varied from 50-86%. Both UVC and IO access have complications associated and adverse events have  
1342 been described. (31, 324-327).

1343 The actual route and method used may depend on local availability of equipment, training and  
1344 experience. (31) There is limited evidence on the effectiveness of IO devices immediately after birth,  
1345 or the optimal site or type of device (328, 329), although simulation studies undertaken in a delivery  
1346 room setting suggest that the IO route can be faster to insert and use than UVC. (330, 331) Proximal  
1347 tibia is the anatomical site usually used in newborn infants. (332) IO access might be possible in  
1348 preterm infants. However, device-specific weight limitations must be considered. ERC recommends, in  
1349 alignment with ILCOR, to use UVC as the primary method of vascular access at birth, and if UVC is not  
1350 feasible, or birth occurs in another setting, the IO route is a reasonable alternative.

### 1351 [h3] Peripheral Access

1352 No studies were identified reviewing the use of peripheral IV cannulation in infants requiring  
1353 resuscitation at birth. A retrospective analysis of 61/70 stable newborn preterm infants in a single  
1354 centre showed that peripheral IV cannulation is feasible and successful in most cases at first attempt.  
1355 (333)

1356

## 1357 [h2] Drugs

1358 Drugs are rarely indicated in resuscitation of the newborn infant. (30, 334) (335) Bradycardia is usually  
1359 caused by profound hypoxia and the key to resuscitation is inflating the fluid filled lungs and

1360 establishing adequate ventilation. However, if the HR remains less than 60 min<sup>-1</sup> despite effective  
1361 ventilation and chest compressions, it is reasonable to consider the use of drugs. Knowledge of the  
1362 efficacy of drugs in newborn resuscitation is largely limited to retrospective studies, as well as  
1363 extrapolation from animals and adult humans. (336)

### 1364 [h3] Adrenaline

1365 A systematic review identified two observational studies involving 97 newborns comparing doses and  
1366 routes of administration of adrenaline. (337) There were no differences between IV and endotracheal  
1367 adrenaline for the primary outcome of death at hospital discharge, for failure to achieve return of  
1368 spontaneous circulation, time to return of spontaneous circulation or proportion receiving additional  
1369 epinephrine. There were no differences in outcomes between 2 endotracheal doses. No human  
1370 newborn studies were found addressing IV dose or dosing interval (very low certainty evidence).  
1371 Recent animal data show no differences in response to doses between 0.2, 0.4, or 0.8 IU/kg  
1372 vasopressin, or 0.02 mg/kg adrenaline and support intravenous administration as the most effective  
1373 route for adrenaline. (338) Despite the lack of newborn human data it is reasonable to use adrenaline  
1374 when effective ventilation and chest compressions have failed to increase HR above 60 beats min<sup>-1</sup>.  
1375 ILCOR suggests that if adrenaline is used, an initial dose of 10-30 micrograms kg<sup>-1</sup> (0.1–0.3 mL kg<sup>-1</sup> of  
1376 1:10,000 adrenaline [1mg in 10 mL]) should be administered intravenously [ILCOR 2025]. If  
1377 intravascular access is not yet available, endotracheal adrenaline at a larger dose of 50 - 100  
1378 micrograms kg<sup>-1</sup> (0.5 – 1.0 mL kg<sup>-1</sup> of 1:10,000 adrenaline [1 mg in 10 mL]) is suggested but should not  
1379 delay attempts at establishing vascular access. (339) If HR remains <60 min<sup>-1</sup> further doses - preferably  
1380 intravascularly - every 3-5 min are suggested. If the response to tracheal adrenaline is inadequate  
1381 ILCOR suggests an IV dose is given as soon as vascular access is established regardless of the interval  
1382 between doses (31, 69, 340)(ILCOR 2025) For pragmatic reasons, ERC recommends using iv/io route  
1383 preferably at a dose of 20 mcg kg<sup>-1</sup> or an endotracheal dose of 100 micrograms kg<sup>-1</sup>; and to repeat  
1384 further doses of adrenaline every 4 minutes if required.

### 1385 [h3] Glucose

1386 Dysglycaemia (hyper or hypoglycaemia) is common during neonatal resuscitation and may be  
1387 associated with poorer resuscitation outcomes. Hypoglycaemia is an important additional risk factor  
1388 for perinatal brain injury. (341) The definition of hypoglycaemia in the context of resuscitation is  
1389 unknown. Hyperglycaemia is a stress response and does not need to be corrected during resuscitation  
1390 but may need to be addressed during post resuscitation care. The ERC recommendation is to check  
1391 blood glucose during a prolonged resuscitation and if low, IV or IO glucose should be given as a 250 mg  
1392 kg<sup>-1</sup> bolus (2.5 mLkg<sup>-1</sup> of 10% glucose). After successful resuscitation formal steps to prevent both  
1393 hypoglycaemia and hyperglycaemia should be instituted.

**1394 [h3] Intravascular volume replacement**

1395 Early intravascular volume replacement is indicated for newborns with blood loss who are not  
1396 responding to resuscitation (31). Therefore, if there has been suspected blood loss or the newborn  
1397 appears to be hypovolaemic and has not responded adequately to other resuscitative measures then  
1398 consider giving volume replacement with crystalloid or red cells. Blood loss causing acute  
1399 hypovolaemia in the newborn is a rare event. There is little to support the use of volume replacement  
1400 in the absence of blood loss when the newborn is unresponsive to ventilation, chest compressions and  
1401 adrenaline. However, because blood loss may be occult and distinguishing normovolaemic newborns  
1402 with shock due to asphyxia from those who are hypovolaemic can be problematic, a trial of fluid  
1403 administration may be considered. (11, 31)

1404 The ERC recommends in the absence of suitable blood (i.e. group O Rh-negative blood), isotonic  
1405 crystalloid rather than albumin is the solution of choice for restoring intravascular volume and to give  
1406 a bolus of 10 mLkg<sup>-1</sup> initially. If successful it may need to be repeated to maintain an improvement.  
1407 When resuscitating preterm newborns, fluid is rarely needed and has been associated with  
1408 intraventricular and pulmonary haemorrhages when large volumes are infused rapidly. (342)

**1409 [h3] Sodium bicarbonate**

1410 ILCOR concluded that a 2005 treatment recommendation on the use of sodium bicarbonate during  
1411 prolonged resuscitation was not supported by a systematic review using contemporary ILCOR methods  
1412 of evidence appraisal; consequently, the recommendation for the routine use of sodium bicarbonate  
1413 has been withdrawn from the 2025 CoSTR (ILCOR 2025). Indeed, there may be harm associated with  
1414 its use, as it is hyperosmolar and generates CO<sub>2</sub> which may impair myocardial and cerebral function.  
1415 (343). Given insufficient data to recommend routine use of bicarbonate in resuscitation of the  
1416 newborn the ERC has followed ILCOR's recommendation and removed it from the guideline.

**1417 [h3] Naloxone**

1418 Naloxone is very seldomly used during newborn resuscitation (writing group experience). There is no  
1419 high-certainty evidence for the use of naloxone during resuscitation.(344) Consequently, the ERC  
1420 cannot recommend the use of Naloxone in that setting.

1421

**1422 [h2] Low resource or remote settings**

1423 Infants born unplanned out of hospital are often in a remote area with lower resources and at higher  
1424 risk of needing resuscitation. Resuscitation then needs to be provided by out of hospital practitioners,  
1425 possibly with less experience of neonatal resuscitation. Stabilisation is followed by additional  
1426 challenges of safe transfer to an appropriate healthcare facility. Hypoxia and hypothermia are

1427 common and should be anticipated and proactively managed. (84, 345-347) Not all hospital settings  
1428 have the same resources, and remote locations may benefit from use of telemedicine.

### 1429 **[h3] Planned home births**

1430 A systematic review of eight studies involving 14637 low risk planned home births compared with  
1431 30177 low risk planned hospital births concluded that the risks of neonatal morbidity and mortality  
1432 were similar.(51) However, unplanned births are more at risk of needing resuscitation and despite risk  
1433 stratification, infants born at home may still require resuscitation. (348) Those attending home births  
1434 must have appropriate skills to manage this. Thermal care with a focus on prevention of hypothermia  
1435 is essential irrespective of birth location. (347) This can be supported, by increasing room temperature  
1436 in the birth location (e.g., turn heating up, close windows), use of warming mattresses or skin-to-skin  
1437 contact. Plastic bags can be used for preterm babies as a useful thermal care adjunct alongside a heat  
1438 source.

1439

## 1440 **[h2] Post-resuscitation care**

### 1441 **[h3] Glucose management**

1442 Hypoglycaemia may occur after perinatal asphyxia because of rapid glucose consumption during  
1443 anaerobic metabolism, stress-induced hyperinsulinism, impaired gluconeogenesis, and concomitant  
1444 risk factors. (349, 350) Conversely, hyperglycaemia may result from endogenous stress hormone  
1445 release, adrenaline administration, and reduced insulin sensitivity. Both hypoglycaemia and  
1446 hyperglycaemia occur frequently after resuscitation: approximately 1 in 7 and 1 in 4 newborns in the  
1447 first 6 hours, increasing to 1 in 5 and 1 in 2 newborns at  $\geq 24$  hours after birth, respectively. (349)  
1448 Infants with hypoxic-ischaemic encephalopathy and severe acidosis are particularly at risk.  
1449 Animal studies suggest hypoxic cerebral injury is worsened by both hypoglycaemia and  
1450 hyperglycaemia. (351, 352) (353) Research in human infants with hypoxic-ischaemic encephalopathy  
1451 has revealed that initial hypoglycaemia and glycaemic lability are associated with more brain injury on  
1452 MRI, lower cognitive scores, and poorer neurological outcome. (354-357) Hyperglycaemia and  
1453 glycaemic lability were also associated with amplitude-integrated electroencephalographic evidence of  
1454 worse global brain function and seizures. (358)  
1455 Hypoglycaemia and hyperglycaemia are associated with higher mortality rates, and (early)  
1456 hypoglycaemia ( $\leq 12$  hours after birth) also causes more neurodevelopmental impairment in newborns  
1457 treated with therapeutic hypothermia for moderate-to-severe hypoxic-ischaemic encephalopathy.  
1458 (359, 360) A systematic review and meta-analysis confirmed the association of hypoglycaemia and  
1459 hyperglycaemia with death and worse neurodevelopmental outcome in babies with neonatal  
1460 encephalopathy. (361) Early hypoglycaemia and hyperglycaemia were independently associated with

1461 death and/or severe neurodevelopmental impairment at 18 months in infants with moderate-to-  
1462 severe hypoxic-ischaemic encephalopathy, irrespective of cooling. (362) Fluctuating glucose levels in  
1463 neonates with hypoxic-ischaemic encephalopathy also correlate with unfavourable outcomes. (363,  
1464 364)

1465 ILCOR concluded that evidence on glucose management is scarce. (349) Only two good practice  
1466 statements could be issued and these are the ERC recommendations: 1) measure blood glucose  
1467 concentration early and regularly after resuscitation until normoglycaemia is achieved; 2) titrate  
1468 infusion of intravenous glucose against the infant's blood glucose values to avoid hypoglycaemia and  
1469 iatrogenic hyperglycaemia. Although the optimal blood glucose target range for newborns with HIE is  
1470 uncertain (365, 366), it seems appropriate to maintain blood glucose  $\geq 2.6$  mmol/l (47 mg/dL) (good  
1471 practice statement). (350, 360)

### 1472 [h3] Thermal Care

1473 If therapeutic hypothermia is not indicated, hypothermia after birth should be corrected because of its  
1474 association with poor outcomes. (96) Infants should be maintained within the normal temperature  
1475 range (36.5°C - 37.5°C). (96) (79)

1476 Hyperthermia ( $\geq 38^\circ\text{C}$ ) after cardiopulmonary resuscitation is also associated with unfavourable  
1477 outcomes (death, moderate or severe disability) in neonates, children, and adults. (367-371) A  
1478 secondary analysis of an RCT comparing whole-body cooling with standard care in term infants with  
1479 hypoxic-ischaemic encephalopathy demonstrated that the risk of death or moderate-to-severe  
1480 disability was increased 3.6-5.9-fold for every 1°C increase in temperature (372). Hyperthermia should  
1481 thus be avoided. (373) The ERC recommends monitoring temperature and aiming for normothermia.

### 1482 [h3] Therapeutic Hypothermia

1483 A Cochrane review including 11 RCTs comprising 1505 term and late preterm infants calculated that  
1484 therapeutic hypothermia resulted in a statistically significant and clinically important reduction in the  
1485 combined outcome of mortality or major neurodevelopmental disability to 18 months of age and  
1486 concluded that newborn infants at term or near-term with evolving moderate-to-severe hypoxic-  
1487 ischaemic encephalopathy (HIE) should be offered therapeutic hypothermia. (372) A more recent  
1488 systematic review and meta-analysis including 29 RCTs with 2926 infants  $\geq 35$  weeks of gestation with  
1489 HIE showed that therapeutic hypothermia diminishes the risk of neurological disability and cerebral  
1490 palsy. (373) The overall effect of therapeutic hypothermia on mortality was uncertain.

1491 Cooling should be performed in NICUs with the capabilities for multidisciplinary care, using clearly  
1492 defined protocols. During transfer to a NICU, servo-controlled active cooling is the preferred method  
1493 to maintain hypothermia in the desired range. (374) Treatment should commence within 6 hours of  
1494 birth, target a temperature between 33°C and 34°C, continue for 72 hours, with rewarming over at

1495 least four hours. (375) A clinical trial of 364 infants randomised to receive longer (120 hours) or deeper  
1496 (32°C) cooling found no evidence of benefit of longer cooling or lower temperatures. (376) Animal data  
1497 strongly suggest that the effectiveness of cooling is related to early intervention. Hypothermia  
1498 initiated at 6-24 hours after birth may have benefit, but there is uncertainty in its effectiveness. (377)  
1499 Such therapy can be considered on an individual basis. Current evidence is insufficient to recommend  
1500 routine therapeutic hypothermia for infants with mild HIE. (378) The ERC recommends applying  
1501 therapeutic hypothermia in term newborns ( $\geq 37$  weeks) with evolving moderate-to-severe HIE in low-  
1502 and middle-income countries as long as appropriate supportive neonatal care can be provided. There  
1503 is insufficient evidence to offer a recommendation on therapeutic hypothermia in low- and middle-  
1504 income countries for late preterm infants (34 to 37 weeks).

### 1505 [h3] Oxygenation & Ventilation

1506 Evidence on oxygen targets in infants with perinatal asphyxia is lacking. It seems prudent to  
1507 continuously monitor oxygen saturations and regularly draw arterial blood gases. (379) Considering  
1508 the increased risk of pulmonary hypertension in infants with hypoxic-ischaemic encephalopathy,  
1509 sometimes aggravated by therapeutic hypothermia, measurement of pre and post-ductal saturations  
1510 is sensible. (380-384) Both hypoxaemia and hyperoxaemia can be detrimental. (385) The ERC  
1511 recommends titrating O<sub>2</sub> to avoid hypoxaemia and hyperoxaemia, and to aim for normocapnia.  
1512 A review of nine retrospective studies reported that hypocapnia in newborns with HIE is associated  
1513 with adverse short- and long-term outcomes. (386) A retrospective cohort study including 188 infants  
1514 managed with therapeutic hypothermia for HIE showed that hypocapnia was associated with more  
1515 severe brain injury on MRI in a dose-dependent fashion. (387) Targeting normocapnia appears sensible  
1516 after neonatal resuscitation. (385)

### 1517 [h3] Prognostication

1518 The Apgar score was designed to focus attention on the newborn and to identify infants needing  
1519 interventions. (388) Individual components of the score (e.g. breathing, HR) reflect the physiological  
1520 relationships during postnatal transition. Lower scores at one minute are associated with more  
1521 interventions at 5 and 10 minutes (30). Although the overall Apgar score is widely recorded in clinical  
1522 practice and for research purposes, its applicability has been questioned because of large inter- and  
1523 intra-observer variations and racial bias. (389) (139, 390, 391) A retrospective study involving 42  
1524 infants (23-40 weeks) found a significant discrepancy (average 2.4 points) between retrospective  
1525 video-based Apgar scores and scores applied by those attending the birth. (389) Individual  
1526 components of the Apgar score are used to guide resuscitation, but the overall Apgar scores are not.  
1527 Apgar scores are calculated after resuscitation and are often required by institutions and national  
1528 registries.

1529 Several studies have looked at the prognostic ability of clinical parameters, biochemical results,  
1530 medication use, neuroimaging, and neurophysiological studies to predict neurodevelopmental  
1531 outcomes of newborn infants (treated) with (hypothermia for) hypoxic-ischaemic encephalopathy.  
1532 (392-399) However, a recent systematic review concluded that all clinical prediction models proposed  
1533 so far have methodological limitations hampering their routine use in clinical practice. (400)  
1534 The ERC cannot recommend a specific clinical prediction model.

### 1535 [h3] Clinical team debriefing

1536 Debriefing after a resuscitation may help improve team performance in subsequent resuscitation  
1537 events. (401) A meta-analysis revealed that team debriefings after simulated events outperformed  
1538 non-debriefing teams by approximately 25%. (402) Another meta-analysis of 61 studies evaluated the  
1539 effectiveness of After-Action Reviews following training and clinical events, indicating an average  
1540 improvement effect size of 0.79 (Cohen's d) in task performance, cognitive skills, attitudes toward  
1541 training/learning. (403)

1542 An ILCOR review on the effect of debriefing on clinical outcome (resuscitation skills and knowledge)  
1543 and patient outcome (good neurological outcome, survival at discharge, survival to hospital) found  
1544 studies with no effect, but also improved favourable neurological outcome, survival to discharge,  
1545 ROSC, chest compression depth, rate and fraction and adherence to guidelines. No undesirable effects  
1546 from debriefing could be demonstrated. The ERC recommends post-event debriefing after neonatal  
1547 cardiac arrest in settings that have adequate resources (55).

1548

### 1549 [h2] Communication with the parents

1550 The principles governing the need for good communication with parents are derived from clinical  
1551 consensus and enshrined in published European guidance. (404, 405) Mortality and morbidity for  
1552 newborns varies according to region, ethnicity and to availability of resources. (406-408) Social science  
1553 studies indicate that parents wish to be involved in decisions to resuscitate or to discontinue life  
1554 support in severely compromised infants. (409, 410) Local survival and outcome data are important in  
1555 appropriate counselling of parents. The institutional approach to management (for example at the  
1556 border of viability) affects the subsequent results in surviving infants. (411)

1557 The ERC is supportive of family presence during cardiopulmonary resuscitation. (412) Healthcare  
1558 professionals are increasingly offering family members the opportunity to remain present during  
1559 resuscitation, and this is more likely if the resuscitation takes place within the delivery room. Parents'  
1560 wishes to be present during resuscitation should be supported where possible. (31, 413, 414)  
1561 There is insufficient evidence to indicate an interventional effect from parental presence on patient or  
1562 family outcome. Being present during the resuscitation of their baby seems to be a positive experience

1563 for some parents but there are concerns among professionals and family members that it may impair  
1564 performance. (31, 414)

1565 In a single centre review of management of birth at the bedside, parents who were interviewed were  
1566 supportive, but some found witnessing resuscitation difficult. (415) Clinicians involved felt the close  
1567 proximity improved communication, but interviews suggested support and training in dealing with  
1568 such situations might be required for staff. (416) In a retrospective survey of clinicians' workload  
1569 during resuscitation the presence of parents appeared to be beneficial in reducing perceived  
1570 workload. (417)

1571 Qualitative evidence emphasizes the need for support during and after any resuscitation, without which  
1572 the birth may be a negative experience with post traumatic consequences. (418, 419) There should be  
1573 an opportunity for the parents to reflect, ask questions about details of the resuscitation and be  
1574 informed about the support services available (413). It may be helpful to offer any parental witness of  
1575 a resuscitation the opportunity to discuss what they have seen at a later date. (418, 419)

1576 Decisions to discontinue or withhold resuscitation should ideally involve senior paediatric staff.

1577 The ERC recommends that where practically possible and parental inclination allows, parents should  
1578 be supported and empowered to be present during the resuscitation of their newborn infant with  
1579 appropriate support from staff. Decisions to discontinue or withhold resuscitation should involve  
1580 senior paediatric staff.

1581

1582 **[h2] Discontinuing or withholding treatment.**

1583 **[h3] Discontinuing resuscitation**

1584 Failure to achieve return of spontaneous circulation in newborn infants after 10-20 minutes of  
1585 intensive resuscitation is associated with a high risk of mortality and a high risk of severe  
1586 neurodevelopmental impairment among survivors. There is no evidence that any specific duration of  
1587 resuscitation universally predicts mortality or severe neurodevelopmental impairment.

1588 The outcomes of infants whose heart rate has been absent for longer than 10 minutes are not  
1589 universally poor. (420) (421) (422) An ILCOR systematic review identified 13 studies involving 271  
1590 infants with at least 10 minutes of asystole, bradycardia or pulseless electrical activity. Of these  
1591 infants, 70% died, 18% survived with moderate/severe neurodevelopmental impairment, and 11%  
1592 survived without moderate/severe impairment. (423) Another review identified 820 infants with  
1593 absent heart rate >10 minutes after birth: 40% survived; 21% survived with moderate to severe  
1594 neurodevelopmental impairment and 19% without moderate or severe neurodevelopmental  
1595 impairment. (424) A secondary analysis of the Optimising Cooling Trial, found that a 10-minute Apgar  
1596 score of 0 alone did not predict well death or moderate or severe disability. (425) It can be helpful to

1597 consider clinical factors, effectiveness of resuscitation and the views of other members of the clinical  
1598 team about continuing resuscitation. (426)

1599 The ERC recommends discontinuing resuscitation after prolonged cardiopulmonary resuscitation if all  
1600 recommended interventions have been applied and potentially reversible causes excluded. A  
1601 reasonable time to consider this is around 20 minutes after birth.

1602 In extremely preterm infants, prolonged resuscitation is associated with lower survival rates and  
1603 higher morbidity; it may be appropriate to discontinue resuscitation sooner. (423, 427) The decision  
1604 should be individualized. The decision to cease resuscitation is a medical decision, but it is important,  
1605 where possible, to give the family updates during the resuscitation and advance warning that there is a  
1606 high chance the baby will not survive.

### 1607 **[h3] Withholding resuscitation**

1608 In situations where there is extremely high predicted mortality and severe morbidity in surviving  
1609 infants, withholding resuscitation may be reasonable, particularly when there has been the  
1610 opportunity for prior discussion with parents. (17-19, 428, 429) Examples from the published literature  
1611 include extreme prematurity (GA <22 weeks and/or birth weight less than 350 g) (429), and anomalies  
1612 such as anencephaly and bilateral renal agenesis. Withholding resuscitation and discontinuation of life-  
1613 sustaining treatment during or following resuscitation are considered by many to be ethically  
1614 equivalent and clinicians should not be hesitant to withdraw treatment when it would not be in the  
1615 best interests of the infant. (430) The ERC recommends a consistent and coordinated approach to  
1616 individual cases by the obstetric and neonatal teams which actively involves the parents. In conditions  
1617 where there is low survival and a relatively high rate of morbidity, and where the anticipated burden  
1618 to the child is high, parental wishes regarding resuscitation should be sought and supported. (405)

1619 **[h1] Conflicts of interest Statement**

1620 MH, VM, JF, MR, ALS, DT, MW and JM are members of ILCOR NLS taskforce. MH is the science chair of  
 1621 the ERC NLS science and education committee. VM is a member of the RC NLS UK. JF is Chair of RC NLS  
 1622 UK. CCR is content expert for the ILCOR NLS-TF. He has received grants from the UK Government for  
 1623 clinical trials and studies. He has received speakers' honoraria and consultancy fees from Chiesi  
 1624 Pharmaceuticals, Italy. MR has received grants from German Government to introduce telemedicine in  
 1625 East Saxony and has received speakers' honorarium from the following companies: Chiesi,  
 1626 AstraZeneca, Pfizer. ES is CEO of SIMCharacters Training GmbH, a company specialising in simulation-  
 1627 based training for neonatal and paediatric emergency care. TS is the president of Polish Neonatal  
 1628 Society. He has received speakers' honoraria from the following companies: Masimo, Medtronic,  
 1629 AstraZeneca, Sanofi. AtP is content expert for ILCOR NLS. He is consultant Research and Development  
 1630 for Fisher&Paykel Healthcare, inventor of the Concord Resuscitation table and received unrestricted  
 1631 grants from SLE. DT is emeritus member of the ILCOR NLS taskforce. He has received speaker  
 1632 honorarium from Chiesi Farmaceutici. MW has received speakers' honoraria from the following  
 1633 companies: Chiesi, Monivent, Schuelke. JM is the education cochair of the ERC NLS science and  
 1634 education committee and a member of the RCUK NLS committee

1635

1636 **[h1] Acknowledgements**

1637 Sylvia Obermann, representative for the Dutch organisation Care4Neo.

1638

1639 **[h1] List of figures**

1640 Figure 1 Key messages

1641 Figure 2 NLS algorithm 2025

1642 Figure 3 Common factors associated with an increased risk of a need for stabilization, or  
 1643 resuscitation at birth

1644 Figure 4 Schematic representation of initial assessment & actions

1645 Figure 5 Head in a neutral position. Face is horizontal (middle picture), neither flexed (left) or  
 1646 extended (right)

1647 Figure 6 Jaw thrust, lifting the lower jaw forwards enlarges the pharyngeal space

1648 Figure 7 Two thumbs encircling technique for CC

1649 Figure 8 Oxygen saturations in healthy infants at birth without medical intervention

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1652

1653 **[h1] List of Tables**

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1662	Table 9	Overview target oxygen saturation ranges

This is a DRAFT-version

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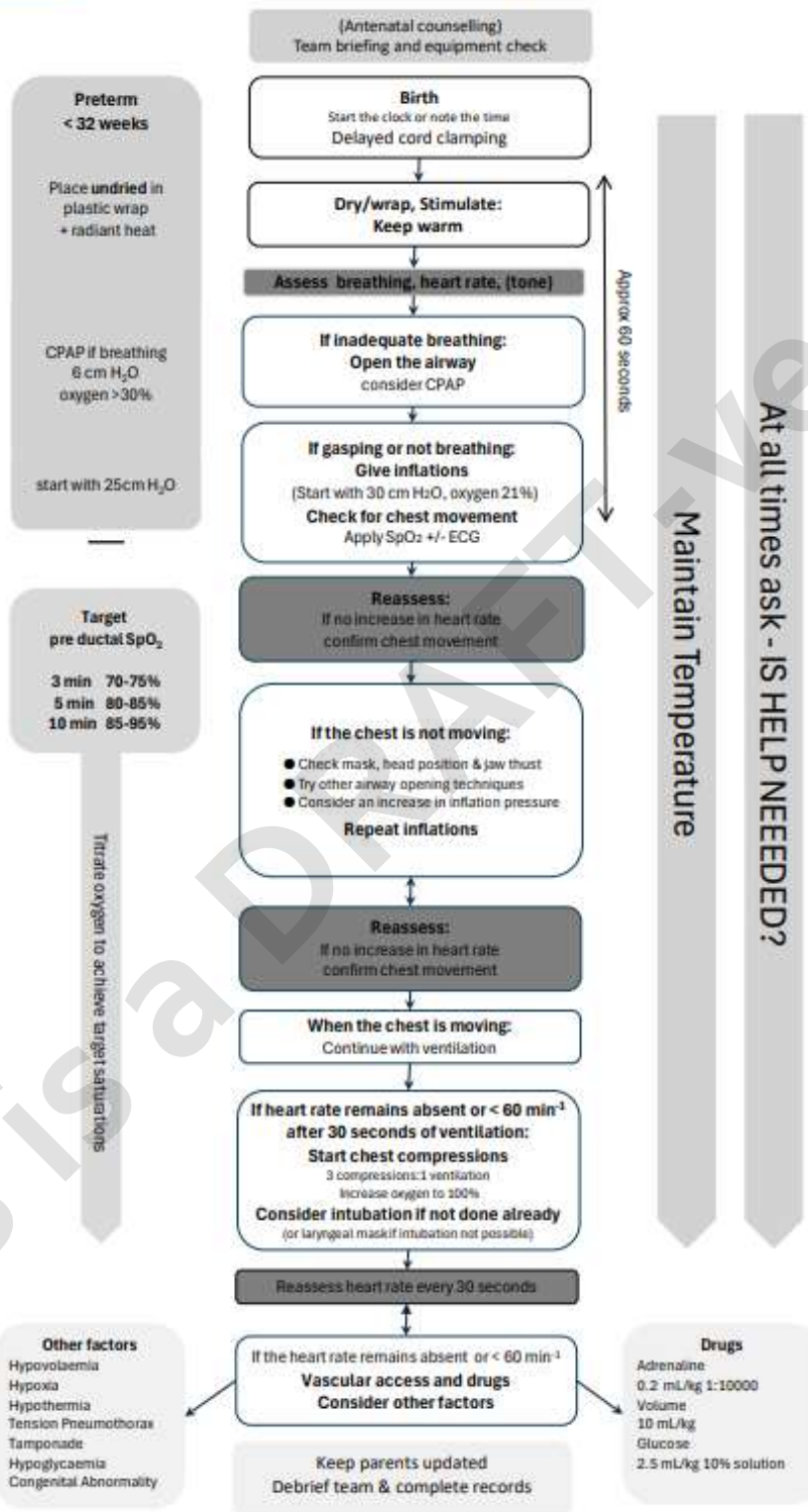
3015 **Figure 1** **Key Messages ERC Guidelines 2025 Neonatal Life Support**

- 3016
- 3017 1. **Delay cord clamping for at least 60 seconds;**
- 3018 longer may be beneficial.
- 3019 No cord milking <28w.
- 3020 2. **Dry, wrap, stimulate & keep warm:**
- 3021 most infants do not need advanced help at birth
- 3022 3. **Assess breathing & heart rate;**
- 3023 this will guide the next steps
- 3024 4. **Open the airway & inflate the lungs;**
- 3025 these are fundamental steps in supporting transition and newborn resuscitation
- 3026 5. **Chest Compressions and drugs are rarely needed;**
- 3027 when needed preferred vascular access is UVC
- 3028 6. **Infants <32 weeks require an altered approach**
- 3029 7. **Unexpected out of hospital births require an altered approach**
- 3030 8. **Involve Parents;**
- 3031 communication before, surrounding and after birth is key

3032 Figure 2 NLS algorithm 2025



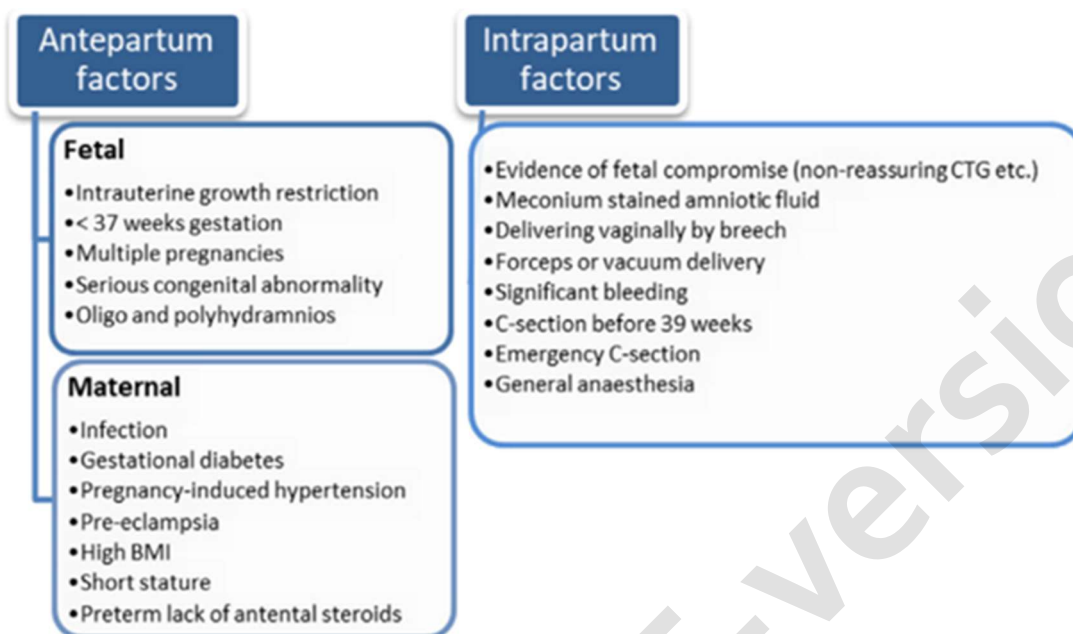
### Newborn Life Support



Draft ERC NLS Algorithm V4b - April 2025

3033

3034 **Figure 3** Common factors associated with the need for stabilisation, or resuscitation at birth

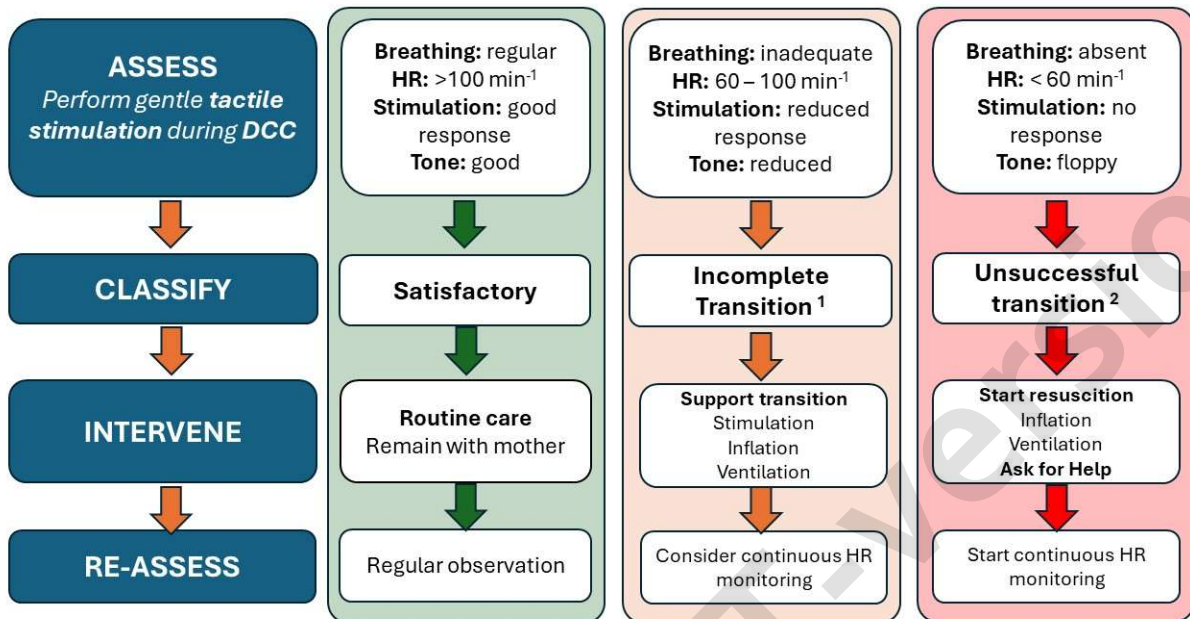


3035

This is a DRAFT-version

3036

**Figure 4: Schematic Illustration of Initial Assessment & Interventions**



3037

3038 <sup>1</sup> Slow HR may indicate hypoxia, so airway and breathing require support. Ventilatory support will  
 3039 likely be adequate for a higher HR and adequate transition.

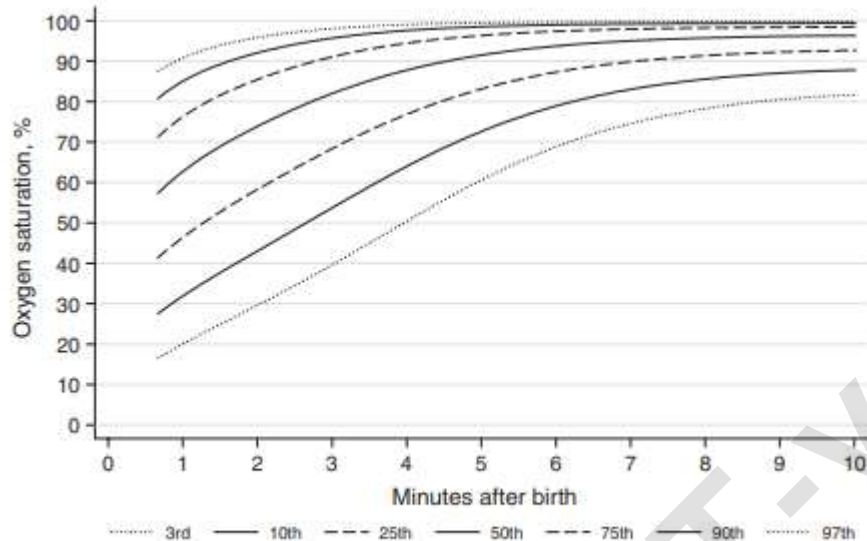
3040 <sup>2</sup> HR suggestive of significant hypoxia, so airway and breathing support required urgently

3041

- 3042 **Figure 5** **Head in a neutral position. Face is horizontal**
- 3043 **Figure 6** **Jaw thrust, lifting the lower jaw forwards enlarges the pharyngeal space**
- 3044 **Figure 7** **Two thumbs encircling technique for chest compressions**

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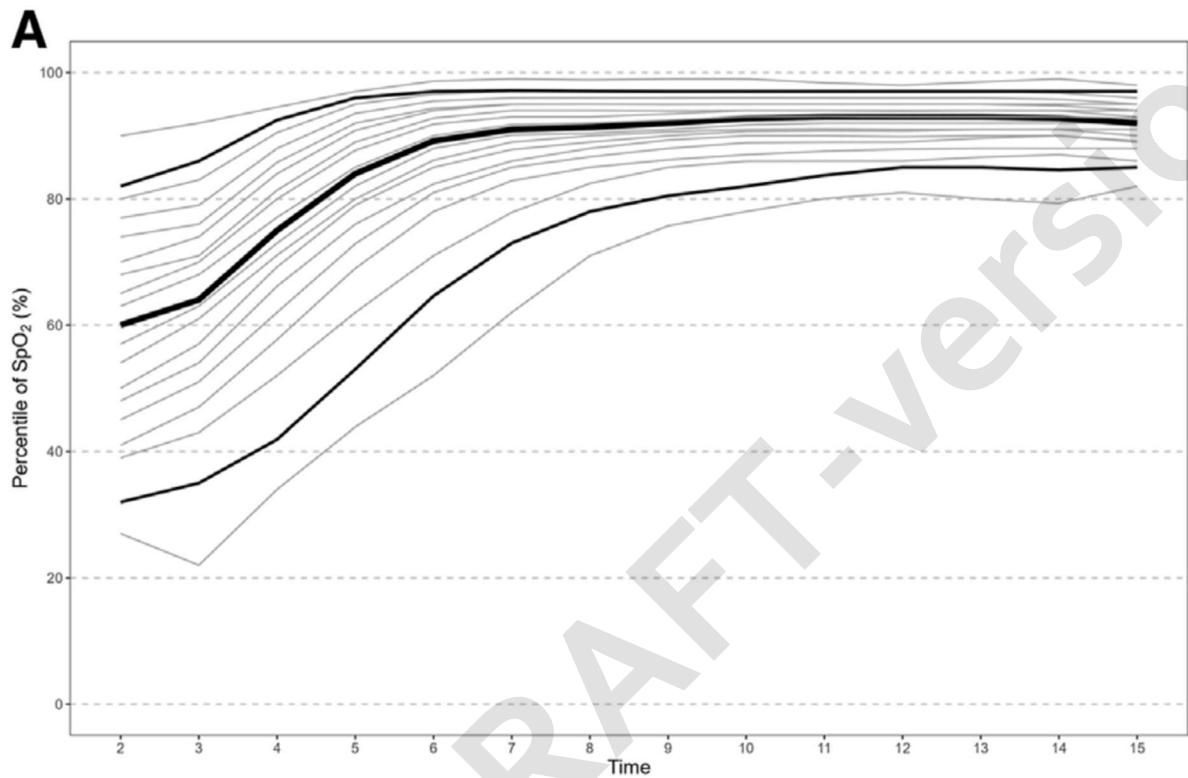
3045 **Figure 8** Oxygen saturations in healthy infants at birth without medical intervention (3<sup>rd</sup>, 10<sup>th</sup>,  
3046 25<sup>th</sup>, 50<sup>th</sup>, 75<sup>th</sup>, 90<sup>th</sup>, 97<sup>th</sup> centiles. Reproduced with permission - from Dawson et al 2010  
3047 (287).



**FIGURE 1** Third, 10th, 25th, 50th, 75th, 90th, and 97th SpO<sub>2</sub> percentiles for all infants with no medical intervention after birth.

3048  
3049

3050 **Figure 9** Oxygen saturations during the first 15 minutes after birth of infants <32 weeks with  
3051 favorable outcome of SpO<sub>2</sub> (%); (10th, 50th, and 90th centile (bold lines);  
3052 5th, 15th, 20th, 25th, 30th, 35th, 40th, 45th, 55th, 60th, 65th, 70th, 75th, 80th, 85th,  
3053 95th centile (gray lines)). From Wolfsberger et al 2024 (148).



3054

3055 **Table 1** Key changes in NLS 2025 Guidelines

3056

Topic	ERC 2021 NLS Guidelines	ERC 2025 NLS Guidelines
When to use Newborn Life Support (NLS) or Paediatric Life Support (PLS) algorithms	Not included	The NLS and PLS Writing Groups have included aligned statements relating to when it might be appropriate to use either resuscitation algorithm. Both writing groups consider it reasonable for teams to initiate resuscitation of an infant outside the delivery area using the guideline most familiar to them (either NLS or PLS) whilst summoning appropriate help and switching algorithm in a timely fashion if necessary.
Applicability of guideline to the most preterm infants at the limits of viability	Not included	The guideline acknowledges the paucity of resuscitation data available from extremely preterm infants especially <25 weeks and cautions that these guidelines are based upon evidence from predominantly older gestational ages, which limits applicability to extremely low gestational ages
Telemedicine	Not included	Telemedicine can provide remote advice and health systems should consider how this can be used.
Environment and equipment	All equipment must be regularly checked and ready for use. Where possible, the environment and equipment should be prepared in advance of the birth of the infant.	Equipment should be easily accessible and organised in a standardised way.  Consider human factor elements when organising equipment and training to maximise efficiency and to minimise time delays.
Delayed cord clamping (DCC)	Where immediate resuscitation or stabilisation is not required, aim for delayed cord clamping of at least 60 s. A longer period may be more beneficial.	Although recommendations about delayed cord clamping have not changed significantly, there is even more emphasis on the importance of delayed cord clamping for all newborn infants, especially preterm infants. In newborn infants needing resuscitation, clamp the cord <30s to minimise delay to necessary interventions.
Cord milking	Where delayed cord clamping is not possible consider cord milking in infants >28 weeks gestation	The guideline reinforces <b>not</b> milking the cord in preterm infants <28 weeks and focusses on trying to perform delayed cord clamping if possible. Cut cord milking is acknowledged as a reasonable alternative if >28 weeks and delayed cord clamping not possible.
Initial assessment - colour	As part of initial assessment, observe tone (& colour).	There is a reduced emphasis on skin colour during initial assessment. This reflects the

		subjective nature of detecting cyanosis or pallor especially in different skin tones.
Initial assessment - heart rate (HR)	Determine the heart rate with a stethoscope and a saturation monitor +/- electrocardiogram (ECG) for later continuous assessment.	The guideline recognises the increasing role for ECG as a continuous method of HR evaluation which is more precise than other methods. However, auscultation with a stethoscope remains a reasonable first option.
Airway management	If there is no heart rate response and the chest is not moving with inflations consider a 2-person mask support if single handed initially Securing the airway via tracheal intubation or insertion of a laryngeal mask.	Use the two-person method of airway support (jaw thrust) if sufficient providers are available as this approach is more effective than single person jaw thrust.  A supraglottic airway should be used if mask ventilation is ineffective
Airway – no chest wall movement – increasing pressures	If there is no heart rate response and the chest is not moving with inflations, consider a gradual increase in inflation pressure	If there is no HR response, the chest is not moving with inflations and airway opening techniques are ineffective, increase inflation pressure. Reduce inflation pressure when chest movement seen and clinical improvement.
Airway - video laryngoscopy	The use of video laryngoscopy may aid endotracheal tube placement.	Use video laryngoscopy if available. This reflects evidence of increased first attempt success in tracheal intubation when video laryngoscopy is used. A conventional direct laryngoscope should be available as an alternative.
Breathing - CPAP / PEEP	In spontaneously breathing preterm infants consider CPAP using either mask or nasal prongs. Use PEEP at minimum of 5-6 cm H <sub>2</sub> O when providing positive pressure ventilation (PPV) to these infants.	CPAP and PEEP is now recommended at a level of <b>6 cm H<sub>2</sub>O</b> . This guideline acknowledges that CPAP may be considered in infants >32 weeks GA with respiratory distress if they require supplemental O <sub>2</sub> .
Breathing - Inflations	Give five inflations maintaining the inflation pressure for up to 2-3 s using an appropriately fitting facemask.	The guideline acknowledges the lack of evidence of superiority or inferiority for initial longer (2-3s) versus shorter (1s) inflation times and now suggests initial inflation times between 1 and 3 seconds. <b>If 1s inflation time is used → give 10 initial inflations</b> <b>If 2 – 3s inflation time is used → give 5 initial inflations</b> Initial breathing support can be delivered via facemask or nasal prongs.

Breathing – Initial oxygen concentration	Infants $\geq 32$ weeks needing respiratory support: start with 21% O <sub>2</sub> . Infants $> 28$ weeks but $< 32$ weeks start with 21-30% O <sub>2</sub> . Infants $< 28$ weeks gestation start with 30% O <sub>2</sub>	Initial oxygen concentration according to gestation has been simplified: Infants $\geq 32$ weeks needing respiratory support: start with 21% O <sub>2</sub> Infants $< 32$ weeks: start with $\geq 30\%$ O <sub>2</sub>
Breathing – Oxygen target saturations	Aim to achieve target SpO <sub>2</sub> $> 25$ th percentile for healthy term infants. <b>Time after birth: target SpO<sub>2</sub></b> <ul style="list-style-type: none"> <li>• 2 min: 65%</li> <li>• 5 min: 85%</li> <li>• 10 min: 90%</li> </ul>	SpO <sub>2</sub> target ranges incorporating newer data from preterm infants in addition to the established data from mostly term infants before DCC was standard practice now result in a target range of acceptable SpO <sub>2</sub>  <b>Time after birth: target SpO<sub>2</sub></b> <ul style="list-style-type: none"> <li>• 3min: 70 - 75 %</li> <li>• 5 min: 80 - 85 %</li> <li>• 10 min: 85 – 95 %</li> </ul> Reduce O <sub>2</sub> if saturations exceed 95%.
Circulation	If chest compressions are required consider securing the airway, ideally with a tracheal tube.	When chest compressions are performed, a supraglottic airway or tracheal tube should be considered, depending on training and experience.
Drugs - Adrenaline	An intravenous dose of adrenaline of 10 - 30 micrograms kg <sup>-1</sup> (0.1-0.3 mLkg <sup>-1</sup> of 1:10,000 adrenaline [0.1 mg/mL]) every 3-5 minutes.	The dose and time intervals of intravenous or intraosseous adrenaline have been simplified: 20 micrograms kg <sup>-1</sup> (0.2 mLkg <sup>-1</sup> of 1:10,000 adrenaline [0.1 mg/mL]) every 4 minutes.
Drugs – Sodium Bicarbonate	Sodium bicarbonate may be considered in a prolonged unresponsive resuscitation with adequate ventilation to reverse intracardiac acidosis.	Removed from 2025 guideline
Drugs - Naloxone	A dose of naloxone may help in the few infants who, despite resuscitation, remain apnoeic with good cardiac output when the mother is known to have received opioids in labour.	Removed from 2025 guideline
Glucose	An intravenous dose of 250 mg kg <sup>-1</sup> (2.5 mL kg <sup>-1</sup> of 10% glucose) is suggested in a prolonged resuscitation to reduce the likelihood of hypoglycaemia	There is greater emphasis on checking blood glucose during resuscitation and treating only if it is low, rather than empirical treatment of presumed hypoglycaemia during resuscitation. The guideline acknowledges the potential for harm from both hypoglycaemia and hyperglycaemia.
Low resource and remote settings	Not included	The guideline considers out of hospital births as low resource settings, especially when unexpected and/or preterm birth.

		Includes a section on identifying and managing common problems of hypothermia, hypoxia and safe transfer to hospital.
Parent input into Guideline 2025	Not included	Guideline has been developed with input by a parent organisation in relevant sub-sections

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3059 **Table 2 Assessment of breathing and heart rate**

	Assessment	Intervention
<b>Breathing assessment</b>		
Regular	Satisfactory	None required
Slow, gasping or grunting	Inadequate	Assess – may require intervention
Not breathing	Absent	Intervention required
<b>HR assessment</b>		
>100 min <sup>-1</sup> (fast)	Satisfactory	None required
60 – 100 min <sup>-1</sup>	Inadequate	Assess – may require intervention
<60 min <sup>-1</sup> (very slow or absent)	emergency	Intervention required

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3061 **Table 3** Inflation time and number of inflations

Inflation time	Number of inflations
1 sec	10 inflations
3 sec	5 inflations

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3063 **Table 4** Target oxygen saturation ranges

3064 Derived from Dawson et al. 2010 and Wolfsberger et al. 2024, and consensus within the NLS

3065 WG (148, 230, 287)

Time after birth	SpO <sub>2</sub> [%]	
3 min	70 - 75	3066
5 min	80 - 85	3067
10 min	85 - 95	3068
		3069
		3070

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3071 **Table 5 Summary of positive pressure ventilation and oxygen**

GA	PIP	PEEP	O <sub>2</sub>
≥32 weeks	30 cm H <sub>2</sub> O	6 cm H <sub>2</sub> O	21%
<32 weeks	25 cm H <sub>2</sub> O	6 cm H <sub>2</sub> O	≥30%

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**Table 6 Summary of studies on effect of NLS training**

Reference	Study design	Setting	Intervention	Sample size	Results
Agudelo-Pérez (2022) (431)	Review 11 studies with NWKM level IV, all HBB studies; 8 pre-post intervention, 2 prospective cohort studies, 1 clinical trial;	LMIC	One-day trainings (HBB) in various intervals	n=412.741 newborns	↓ overall neonatal mortality ↓ intrapartum stillbirth ↓ 1d-mortality
Bayoumi (2022)(432)	Pre-post intervention; 1 level III unit (18000/a births)	HIC	5 in-situ simulation trainings and 27 workshops in post-era (2016-2021)	n = 799 in courses, n = 1.199 newborns, n = 326 intubations ev	↑ success rate LISA ↓ duration of intubation
Bhatia (2021) (433)	Pre-Post intervention; tertiary unit with 3 sites (9000/a births), multidisciplinary	HIC	10-12 in-situ simulation workshops per year (2012-2018)	n = 445 HCW, n > 40.000 newborns, n = 11.284 resuscitations	↓ perinatal mortality ↓ chest compressions, ↓ medication
Mayer (2022)(434)	Retrospective observation; 5 hospitals (2 district, 2 regional, 1 tertiary)	MIC	Annual one-day training (HBB, 2016- 2020)	n = 4795 HCW, n = 123.898 newborns	↓ neonatal mortality
Miledler (2024) (60)	Pre-Post intervention; 1 level IV unit (3500/a births), multidisciplinary	HIC	41 in-situ simulation trainings in 4 months	n = 48 HCW, n = 28 resuscitations	↑ 5-minute Apgar score
Lima (2023) (435)	Pre-post intervention; 5 hospitals (secondary healthcare regions)	MIC	n = 700 training sessions in 106 NRP courses	n = 431 HCW	↓ neonatal mortality in DR

Lindhard (2021) (436)	Review 2 Studies with NWKM level IV Lebanon: pre-post intervention Mexico: pair-matched study	LMIC	Lebanon: QI with 10 ex-situ simulation workshops (22 hospitals, 3 years); Mexico: 2 simulation trainings (12 hospitals)	Lebanon: n = 256 HCW, n = 8.4398 births; Mexico: n = 450 HCW;	↓ neonatal mortality, ↑ team performance
Patel (2017) (437)	Review 20 studies with NWKM level IV	LMIC	Variability in neonatal resuscitation training curricula, from basic to advanced life support	n = 1.653.805 newborns; variability in participants of the interventions	↓ neonatal mortality ↓ stillbirth mortality ↓ perinatal mortality
Schwindt (2022) (61)	Pre-Post intervention; 1 level II unit (2000/a births), multidisciplinary	HIC	11 in-situ simulation trainings in post-era (2015-2019)	n = 35 core and 200 additional HCW, n = 13.950 newborns, n = 826 resuscitations	↓ chest compressions
Vadla (2022) (438)	3-year prospective clinical observation study	LIC	High-frequency, self-guided skills training (simulator with automatic feedback)	n = 10.481	↓ time to first ventilation ↓ pauses in ventilation = neonatal mortality
Vadla (2024) (62)	Prospective observational study; 1 hospital site (3000/a births)	LIC	Annual one-day training (HBB 2nd, 2017-2021) + Low dose high frequency trainings	n = 12.983 newborns, n = 1.320 resuscitations	↓ neonatal mortality

Collected evidence from studies in neonatal settings (60-62, 431-439) on the impact of simulation-based training, focusing on Kirkpatrick Level IV

outcomes (clinical resuscitation outcomes) as defined by the New World Kirkpatrick Model (440).

Abbreviations: HBB: Helping babies breathe; HCW: healthcare workers; HIC: high income country; LISA: less invasive surfactant administration; LIC: low income country; MIC: middle income country; NRP; Neonatal Resuscitation Program NWKM: new world Kirkpatrick model; PPV: positive pressure ventilation; QI: Quality improvement

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**Table 7 Heart rate assessment**

Method of HR assessment	Continuous HR monitoring?	Advantages	Disadvantages	Recommendations
<b>Auscultation with stethoscope</b>	No	Rapid assessment Cheap Simple  Readily available in all settings	Intermittent HR monitoring Less reliable compared to other HR methods	Auscultation might be used for quick first assessment Auscultation reasonable alternative for HR assessment Auscultation (+/- pulse oximetry) should be used if ECG is unavailable, malfunctioning or PEA is suspected
<b>Pulse Oximetry</b>  <i>Ideally placed on the right hand or wrist.</i>	Yes	Continuous HR monitoring Provides a measure of oxygenation and perfusion	May underestimate HR as ECG in first 2-5 mins Interference in values caused by: <ul style="list-style-type: none"> <li>○ Signal dropout</li> <li>○ Movement</li> <li>○ Hypoperfusion</li> <li>○ Lighting</li> </ul> Potential cost implication	Unclear whether connecting sensor to infant first or to the pulse oximetry first confers advantage
<b>Electrocardiogram (ECG)</b>	Yes	Continuous HR monitoring Faster and more accurate than pulse oximetry	May indicate a HR in absence of cardiac output May adhere poorly to infants with vernix	Use of ECG is reasonable to assess HR after birth

			Potential cost implication	ERC recommends ECG should not replace pulse oximetry for further treatment, should be used in addition
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Sources for Table 7: (5, 149, 150, 441-451) (289). Abbreviations: HR: heart rate; ECG: electrocardiogram; PEA: pulseless electrical activity

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**Table 8 Approximate tracheal tube size by gestation and approximate lengths for oral and nasal intubation**

<b>Gestation (weeks)</b>	<b>Internal diameter (mm)</b>	<b>Length at lips (cm)</b>	<b>Nasal intubation length (cm)</b>
23-24	2.5	5.5	6.5
25-26	2.5	6.0	7.0
27-29	2.5	6.5	7.5
30-32	3.0	7.0	8.0
33-34	3.0	7.5	8.5
35-37	3.5	8.0	9.0
38-40	3.5	8.5	9.5
41-43	4.0	9.0	10.0

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**Table 9 Overview of oxygen saturation target ranges (148, 230, 287).**

	Dawson (287)		Wolfsberger (148)		Dawson (287)		NRP (230)
	<32 weeks, n=29		<32 weeks, n=207		>=37 weeks, n=308		
	P25	P75	P25	P75	P25	P75	
3 min	67	83	51	77	71	90	70-75
5 min	82	91	73	92	83	96	80-85
10 min	89	95	89	95	94	98	85-95

(148, 230, 287) NRP: Newborn Resuscitation Program.

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