

| 1 | [Title] European Resuscitation Council Guidelines 2025: Adult Advanced Life Support |
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| 2 | |
| 3 | Jasmeet Soar* |
| 4 | Southmead Hospital, North Bristol NHS Trust, Bristol, UK |
| 5 | |
| 6 | Bernd W. Böttiger |
| 7 | Medical Faculty, University of Cologne, Köln, Germany |
| 8 | German Red Cross, Berlin, Germany |
| 9 | Institute for Emergency Care, Health Pedagogy and Nursing Science, Faculty of Health Sciences, |
| 10 | University of Pécs, Pécs, Hungary |
| 11 | |
| 12 | Pierre Carli |
| 13 | Université Paris Cité, Hôpital Universitaire Necker, Paris Centre, Assistance Publique Hôpitaux de |
| 14 | Paris, France |
| 15 | |
| 16 | Francesc Carmona Jiménez |
| 17 | Sistema d'Emergències Mèdiques, Barcelona, Spain |
| 18 | Campus Docent Sant Joan de Déu-UVic (SJD-RCP), Sant Boi de Llobregat, Barcelona, Spain |
| 19 | |
| 20 | Diana Cimpoesu |
| 21 | "Grigore T Popa" University of Medicine and Pharmacy, Iasi, Romania; |
| 22 | Emergency Department, Emergency County Hospital "Sf.Spiridon", Iasi, Romania |
| 23 | |
| 24 | Gareth Cole |
| 25 | Sudden Cardiac Arrest UK |
| 26 | |
| 27 | Keith Couper |
| 28 | Critical Care Unit, University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK; |
| 29 | Warwick Medical School, University of Warwick, Coventry, UK |
| 30 | |
| 31 | Sonia D'Arrigo |
| 32 | Department of Intensive Care, Emergency Medicine and Anaesthesiology, Fondazione Policlinico |
| 33 | Universitario A. Gemelli-IRCCS – Università Cattolica del Sacro Cuore, Rome, Italy |
| 34 | |
| | |



- 35 Charles D. Deakin
- 36 University Hospital Southampton NHS Foundation Trust, Southampton, UK; South Central Ambulance
- 37 Service NHS Foundation Trust, UK
- 38
- 39 Jacqueline Eleonora Ek
- 40 Department of Medicine, Mater Dei Hospital, Malta.
- 41
- 42 Mathias J. Holmberg
- 43 Department of Anaesthesiology and Intensive Care, Aarhus University Hospital, Aarhus, Denmark
- 44
- 45 Aurora Magliocca
- 46 Department of Pathophysiology and Transplantation Milan University, Italy
- 47
- 48 Nikolaos Nikolaou
- 49 Cardiology Department, Konstantopouleio General Hospital, Athens, Greece
- 50
- 51 Peter Paal
- 52 Department of Anaesthesiology and Intensive Care Medicine, St. John of God Hospital, Paracelsus
- 53 Medical University, Salzburg, Austria
- 54
- 55 Helen Pocock
- 56 South Central Ambulance NHS Foundation Trust, Bicester, UK
- 57 Warwick Clinical Trials Unit, Coventry, UK
- 58
- 59 Claudio Sandroni
- 60 Department of Intensive Care, Emergency Medicine and Anaesthesiology, Fondazione Policlinico
- 61 Universitario A. Gemelli-IRCCS Università Cattolica del Sacro Cuore, Rome, Italy
- 62
- 63 Tommaso Scquizzato
- 64 Department of Anaesthesia and Intensive Care, IRCCS San Raffaele Scientific Institute, Milan, Italy 65
- 66 Markus B. Skrifvars
- 67 Department of Anaesthesia and Intensive Care Medicine, University of Helsinki and Helsinki
- 68 University Hospital, Helsinki, Finland



69

- 70 Francesca Verginella
- 71 Regional Emergency Dispatch Centre of Friuli Venezia Giulia, SORES FVG, Italy
- 72
- 73 Joyce Yeung
- 74 Warwick Clinical Trials Unit, Warwick Medical School, University of Warwick, Warwick, UK
- 75 University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK
- 76
- 77 Jerry P. Nolan
- 78 University of Warwick, Warwick Medical School, Warwick, UK
- 79 Royal United Hospital, Bath, UK
- 80
- 81 *Corresponding author
- 82 jasmeet.soar@nbt.nhs.uk



83 [h1]Abstract

- 84 These European Resuscitation Council (ERC) Guidelines 2025 Adult Advanced Life Support (ALS)
- 85 guidelines are based on the International Liaison Committee on Resuscitation (ILCOR) Consensus on
- 86 Cardiopulmonary Resuscitation Science with Treatment Recommendations (CoSTR). The evidence
- 87 informing the ALS guidelines is also included. When ILCOR has not addressed a specific topic, the ERC
- 88 ALS Writing Group has provided its own guidance and the evidence supporting it. This section
- 89 provides guidelines for ALS treatments for adults with in- or out-of-hospital cardiac arrest. The ERC
- 90 Guidelines 2025 ALS emphasise providing early and effective ALS interventions to improve survival
- 91 from cardiac arrest in adults.
- 92

93 [h1] Key words

- 94 Cardiac arrest
- 95 Cardiopulmonary resuscitation
- 96 Advanced Life Support



97 [h1] Common abbreviations used in text 98 AED automated external defibrillator 99 AF atrial fibrillation 100 ALS advanced life support 101 BLS basic life support 102 ΒP blood pressure 103 CHD coronary heart disease 104 CoSTR Consensus on Science and Treatment Recommendation 105 CPR cardiopulmonary resuscitation 106 DC direct current 107 DNACPR do not attempt cardiopulmonary resuscitation 108 DSD dual (double) sequential defibrillation 109 ECG electrocardiogram 110 ECPR extra-corporeal cardiopulmonary resuscitation 111 EMS emergency medical service 112 ERC **European Resuscitation Council** 113 ETCO₂ End-tidal carbon dioxide 114 ICD implantable cardioverter defibrillator in-hospital cardiac arrest 115 IHCA 116 ILCOR International Liaison Committee on Resuscitation 117 10 intraosseous 118 IV intravenous 119 OHCA out-of-hospital cardiac arrest 120 PEA pulseless electrical activity 121 POCUS point of care ultrasound 122 ROSC return of spontaneous circulation 123 SBAR situation, background, assessment, recommendation. 124 SCA sudden cardiac arrest 125 SGA supraglottic airway 126 STEMI ST-elevation myocardial Infarction 127 TOR termination of resuscitation 128 VT ventricular tachycardia 129 VF ventricular fibrillation 130 VF/pVT ventricular fibrillation/pulseless VT



131 [h1]Introduction

- 132 This European Resuscitation Council (ERC) Guidelines 2025 Adult Advanced Life Support (ALS)
- 133 includes the advanced interventions that can be used in addition to basic life support (BLS) and
- automated external defibrillator (AED) by health care professionals. These ALS treatments are helpful
- 135 when started quickly and early during cardiac arrest. ALS includes the prevention and treatment of
- 136 both in-hospital cardiac arrest (IHCA) and out-of-hospital cardiac arrest (OHCA), the ALS algorithm,
- 137 manual defibrillation, airway management during cardiopulmonary resuscitation (CPR), drugs and
- 138 their delivery, and the treatment of peri-arrest arrhythmias. These ERC Guidelines 2025 are based on
- 139 the International Liaison Committee on Resuscitation (ILCOR) Consensus on Science and Treatment
- 140 Recommendations (CoSTR) for ALS.¹ For these ERC Guidelines, the ILCOR recommendations were
- supplemented by focused literature reviews undertaken by the ERC ALS Writing Group for those
- 142 topics not reviewed by ILCOR. When required, the guidelines were informed by the expert consensus
- 143 of the writing group membership. For the first time we have had a patient public representative (GC)
- 144 on the ALS Writing Group.
- 145 The scope of the 2025 ALS Guidelines was posted for feedback from National Resuscitation Councils
- 146 and public comment and several new topics were added based on the scoping process. The ALS
- 147 guidelines were drafted and agreed by the ALS Writing Group members and the ERC Guidelines 2025
- 148 Steering Committee. This Guideline was posted for public comment in May/June 2025. A total of
- 149 [INSERT NUMBER] individuals from [INSERT COUNTRIES] submitted [INSERT NUMBER] comments,
- 150 which resulted in [INSERT CHANGES] in the final version. Review of these comments led to XX
- 151 changes. The Guideline was presented to and approved by the ERC General Assembly on XXXXXX.
- 152 The methodology used for guideline development is presented in the Executive summary.²



- 153 [h1] Summary of key changes
- 154
- 155 The key messages are presented in Figure 1. Table 1 summarises the major changes that have been
- 156 made.
- 157

158 **Figure 1. Advanced Life Support** – key messages

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Figure 1

Advanced Life Support – key messages

- Start ALS early every second counts!
- Ensure *effective* ventilation breaths with high-quality chest compressions
- Defibrillate *early* for shockable rhythms use *correct* apical (lateral) defibrillator pad placement – switch to anterior-posterior pads if 3 ineffective shocks
- Give IV adrenaline early for non-shockable arrests
- Identify and treat reversible causes without delay
- · Consider ECPR early if conventional CPR is failing

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161 **Table 1. What's new in Guidelines 2025 for Advanced Life Support?**

| | Guidelines 2021 | Guidelines 2025 |
|------------|---|--|
| Headlines | Emphasis on: | Greater emphasis on: |
| | High quality chest | Starting ALS as early as |
| | compression | possible to help save more |
| | Premonitory signs to | lives. |
| | prevent cardiac arrest | Effective oxygenation and |
| | Using basic and advanced | ventilation breaths with high |
| | airway techniques in a step | quality chest compressions. |
| | wise fashion – only rescuers | Correct apical (lateral) pad |
| | with a high success rate | position for defibrillation. |
| | should use tracheal | Use of waveform |
| | intubation | capnography to confirm |
| | Early adrenaline for non- | correct tracheal tube |
| | shockable rhythms | placement |
| | | • What's out? |
| | | Calcium and sodium |
| | | bicarbonate have no role |
| | | during CPR except for very |
| | | specific indications. |
| | | • Precordial thump is no longer |
| | | included in these guidelines. |
| ALS in low | Not mentioned in guidelines | ALS guidelines may have to be |
| resource | 2021 | adapted according to resources |
| settings | | and there may need to be a |
| | | greater focus on prevention, early |
| | | first aid, and basic life support |
| | | measures. |
| | | Rescuers should be aware that |
| | | even in high-income settings, ALS |
| | | may be constrained by limited |
| | | resources. |



| | | • | A two-tiered approach |
|---------------|-----------------------------|---|---------------------------------------|
| | | | incorporating basic and advanced |
| | | | interventions may be the safest |
| | | | and most effective. |
| CPR-induced | Not mentioned in guidelines | • | Rescuers may consider using |
| consciousness | 2021 | | sedative or analgesic drugs (or |
| | | | both) in small doses to prevent |
| | | | pain and distress to patients who |
| | | | are conscious during CPR. |
| | | • | Neuromuscular blocking drugs |
| | | | alone should not be given to |
| | | | conscious patients. |
| | | • | Drug regimens may be based on |
| | | | those used in critically ill patients |
| | | | and according to local protocols |
| | | | such as small doses of opioids, |
| | | | ketamine and/or midazolam. |



| Defibrillation: | • | Not clarified in 2021 | ٠ | Manual defibrillators should only |
|-----------------|---|-----------------------------------|---|-------------------------------------|
| AED versus | | | | be used by rescuers who can |
| manual | | | | quickly and accurately identify a |
| defibrillation | | | | cardiac arrest rhythm (within 5 |
| during ALS | | | | seconds) and, if needed, deliver a |
| | | | | safe shock with minimal |
| | | | | interruption (less than 5 seconds) |
| | | | | to chest compressions. |
| | | | ٠ | ALS providers must be proficient |
| | | | | in using both an AED and a |
| | | | | manual defibrillator. |
| | | | • | If an AED is already in use when |
| | | | | ALS providers arrive, they should |
| | | | | follow its shock prompts. When |
| | | | | possible, they should transition to |
| | | | | a manual defibrillator during a 2- |
| | | | | minute CPR cycle. |
| Manual | • | Mentioned in the supporting | • | Immediate defibrillation of |
| defibrillation | | text: The 2015 ERC ALS | | (ventricular fibrillation)VF of any |
| strategy | | Guideline stated that if there is | | amplitude (even fine VF) should |
| | | doubt about whether the | | be attempted. |
| | | rhythm is asystole or extremely | | |
| | | fine VF, do not attempt | | |
| | | defibrillation; instead, continue | | |
| | | chest compressions and | | |
| | | ventilation. We wish to clarify | | |
| | | that when the rhythm is clearly | | |
| | | judged to be VF a shock should | | |
| | | be given. | | |



| | I | |
|--------------|-------------------------------------|-------------------------------------|
| Refractory | • For refractory VF, consider using | • For refractory VF, defined as |
| ventricular | an alternative defibrillation pad | continuous VF after three |
| fibrillation | position (e.g. anterior- | consecutive shocks, and having |
| | posterior) | ensured correct antero-lateral |
| | • Do not use dual (double) | pad positioning, consider using a |
| | sequential defibrillation for | defibrillation vector change by |
| | refractory VF outside of a | using an alternative defibrillation |
| | research setting. | pad position (e.g. antero- |
| | | posterior). |
| | | Dual (double) sequential |
| | | defibrillation (DSD), involves |
| | | using a combination of antero- |
| | | lateral and antero-posterior pad |
| | | positioning, discharged in close |
| | | succession. Given the practical |
| | | challenges of using two |
| | | defibrillators to deliver DSD and |
| | | the limited evidence for its |
| | | efficacy the ERC does not |
| | | recommend its routine use. |
| Bag-mask | Not emphasised in 2021 | Deliver effective bag-mask |
| ventilation | | ventilation breaths by optimising |
| | | mask seal and airway patency |
| | 2 | and if necessary and feasible, use |
| | | a two-person technique for bag- |
| 5 | | mask ventilation. |
| Choice of | • This was not specified in 2021 | When using a supraglottic airway |
| supraglottic | | (SGA), an i-gel is preferred to a |
| airway | | laryngeal tube. |
| | | |

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| Conformation | Use waveform capnography to | • A sustained ETCO ₂ trace on |
|-----------------|----------------------------------|--|
| of correct | confirm tracheal tube position. | waveform capnography must be |
| tracheal tube | | used to confirm correct |
| placement | | placement of the tracheal tube. |
| Mechanical | • This was not specified in 2021 | Use a volume-controlled mode |
| ventilator | | during chest compressions set |
| settings during | | the ventilator to a tidal volume of |
| chest | | 6-8 mL kg ⁻¹ (predicted body |
| compressions | | weight), or to achieve a visible |
| | | chest rise, the maximum inspired |
| | | oxygen, a respiratory rate of 10 |
| | | min ⁻¹ , an inspiratory time of 1–2 |
| | | seconds, a positive end expiratory |
| | | pressure (PEEP) 0-5 cm H_2O , the |
| | | peak pressure alarm at 60-70 cm |
| | | H ₂ O, and the flow trigger off. |
| Vascular access | Attempt intravenous (IV) access | • Attempt intravenous (IV) rather |
| | first to enable drug delivery in | than intraosseous (IO) access |
| | adults in cardiac arrest. | first, to enable drug delivery in |
| | Consider intraosseous (IO) | adults in cardiac arrest. |
| | access if attempts at IV access | • If IV access cannot be rapidly |
| | are unsuccessful or IV access is | achieved within two attempts, it |
| | not feasible | is reasonable to consider IO |
| | 7 | access as an alternative route for |
| | | vascular access during adult |
| 5 | | cardiac arrest. |



| Use of calcium, | • This was not made explicit in | n • Do not routinely give calcium, |
|-----------------|---------------------------------|--|
| sodium | 2021 | sodium bicarbonate or |
| bicarbonate | | corticosteroids during cardiac |
| and | | arrest. |
| corticosteroids | | |
| | | |
| | | |
| | | |
| | | |
| ALS in highly- | Not mentioned in guidelines | s • A sudden decrease in ETCO ₂ may |
| monitored | 2021 | indicate a cardiac arrest or very |
| cardiac arrest, | | low cardiac output state. |
| and physiology- | | Consider starting chest |
| guided CPR | | compressions if the systolic blood |
| | | pressure decreases and remains |
| | | < 50 mmHg despite interventions. |
| | | In adults with continuous intra- |
| | | arterial blood pressure |
| | | monitoring, we suggest that |
| | | adrenaline is initially given in |
| | | small increments (e.g., 50–100 μg |
| | | IV) rather than a 1 mg bolus. |
| | | A pragmatic approach during |
| | 7 | physiology-guided CPR is to aim |
| | | for a diastolic blood pressure of \geq |
| 5 | | 30 mmHg (when using intra- |
| | | arterial blood pressure |
| | | monitoring) and an $ETCO_2 \ge 25$ |
| | | mmHg (3.3 kPa). |
| 1 | | |



| Peri-arrest | • | • | The 2025 has a greater emphasis |
|-------------|---|---|------------------------------------|
| arrhythmias | | | on those arrhythmias that require |
| | | | immediate treatment before or |
| | | | after cardiac arrest. |
| | | • | The tachycardias section has |
| | | | been renamed tachyarrhythmias. |
| | | • | There is greater emphasis on use |
| | | | of electrical cardioversion with a |
| | | | synchronised shock for patients |
| | | | immediately after ROSC, or who |
| | | | are unstable. |

162 163



164 [h1]Concise guidelines for clinical practice 165 166 [h2]Prevention of in-hospital cardiac arrest 167 The ERC recommends: 168 Shared decision making and advanced care planning which integrates resuscitation decisions 169 with emergency care treatment plans to increase clarity of treatment goals and also prevent 170 inadvertent deprivation of other indicated treatments, besides CPR. These plans should be 171 recorded in a consistent manner (See ERC Guidelines 2025 Ethics in Resuscitation).³ 172 Hospitals use a track and trigger early warning score system for the early identification of 173 patients who are critically ill or at risk of clinical deterioration. 174 Hospitals train staff in the recognition, monitoring and immediate care of the acutely ill patient. 175 Hospitals empower all staff to call for help when they identify a patient at risk of physiological 176 deterioration. This includes calls based on clinical concern, rather than solely on vital signs. 177 Hospitals have a clear policy for the clinical response to abnormal vital signs and critical illness. 178 This may include a critical care outreach service and, or emergency team (e.g. medical 179 emergency team, rapid response team). 180 Hospital staff use structured communication tools to ensure effective handover of information. 181 Patients receive care in a clinical area that has the appropriate staffing, skills, and facilities for 182 their severity of illness. 183 Hospitals should review cardiac arrest events to identify opportunities for system improvement 184 and share key learning points with hospital staff. 185 Participation in national cardiac arrest audit as a benchmark for local performance. 186 187 [h2] Prevention of out-of-hospital cardiac arrest 188 Coronary heart disease (CHD) is the leading cause of sudden cardiac arrest (SCA), responsible for 189 80% of cases, particularly in older patients. Non-ischaemic cardiomyopathies contribute to 10-190 15% of SCA cases. In younger individuals, the main causes of SCA include inherited heart 191 diseases, congenital heart defects, myocarditis, and substance abuse. In these patient groups, 192 risk stratification is possible, and preventive treatments may be effective. 193 Predicting SCA is challenging because most cases happen in individuals with undiagnosed heart 194 disease. As a result, detecting early warning signs, implementing an efficient emergency medical 195 services (EMS) system, and focusing on the prevention of cardiovascular disease (CVD) risk 196 factors are crucial in the general population.



197 Symptoms such as chest pain, syncope (especially during exercise, while sitting or supine), 198 palpitations, dizziness or sudden shortness of breath that are consistent with cardiac ischaemia 199 or an arrhythmia should be investigated. 200 Overtly healthy young adults who have SCA can also have preceding signs and symptoms (e.g. 201 syncope/pre-syncope, chest pain and palpitations) that should alert healthcare professionals to 202 seek expert help to prevent cardiac arrest. 203 Young adults presenting with characteristic symptoms of arrhythmic syncope should have a 204 specialist cardiology assessment, which should include an electrocardiogram (ECG) and in most 205 cases echocardiography, 24-hour ECG monitoring and an exercise test. 206 Systematic evaluation in a clinic specialising in the care of those at risk for SCA is recommended 207 in family members of young victims of SCA or those with a known cardiac disorder resulting in an 208 increased risk of SCA. 209 Identification of individuals with inherited conditions and screening of family members can help 210 prevent deaths in young people with inherited heart disorders. 211 Follow current European Society of Cardiology (ESC) guidelines for the diagnosis and 212 management of syncope and arrhythmias. 213 214 [h2]Treatment of in-hospital cardiac arrest 215 Start ALS as early as possible. 216 Hospital systems should aim to recognise cardiac arrest, start CPR immediately, defibrillate 217 rapidly (<3 minutes) for shockable rhythms, give adrenaline rapidly for non-shockable rhythms, 218 and identify and treat reversible causes. 219 All hospital staff should be able to recognise cardiac arrest rapidly, call for help, start CPR and 220 defibrillate (attach an AED and follow the AED prompts, or use a manual defibrillator). 221 Hospitals should adopt a standard 'Cardiac Arrest Call' telephone number (2222). 222 Hospitals should have a resuscitation team that immediately responds to IHCAs. 223 The hospital resuscitation team should include team members who have completed an 224 accredited adult ALS course that incorporates teamwork and leadership training. 225 Resuscitation team members should have the key skills and knowledge to manage a cardiac 226 arrest including manual defibrillation, advanced airway management, intravenous access, intra-227 osseous access, and identification and treatment of reversible causes. 228 The resuscitation team should meet at the beginning of each shift for introductions and 229 allocation of team roles.



230 Hospitals should standardise resuscitation equipment. 231 Termination of resuscitation rules (TOR) should not be used as a sole strategy for terminating an 232 in-hospital resuscitation attempt. 233 234 [h2] Treatment of out-of-hospital cardiac arrest 235 Start ALS as early as possible – EMS systems should be organised to provide a rapid ALS response 236 with sufficient qualified personnel. This may include a prehospital critical care team. 237 Adults with non-traumatic OHCA should be considered for transport to a cardiac arrest centre 238 according to local protocols. 239 Emergency medical systems should consider implementing criteria for the withholding and 240 termination of resuscitation (TOR) taking into consideration specific local legal, organisational 241 and cultural context (See ERC Guidelines 2025 Ethics in Resuscitation³). 242 Emergency medical systems should monitor staff exposure to resuscitation and low exposure 243 should be addressed to increase EMS team experience in resuscitation. 244 245 [h2]Debriefing 246 Use data-driven, performance-focused debriefing of rescuers to improve CPR quality and patient 247 outcomes (See Education guidelines⁴). 248 249 [h2] ALS in low-resource settings 250 Advanced Life Support guidelines may have to be adapted according to resources and there may 251 need to be a greater focus on prevention, early first aid, and basic life support measures (See the 252 System Saving Lives⁵ and First Aid guidelines⁶). 253 Rescuers should be aware that even in high-income settings, ALS may be constrained by limited 254 resources. 255 A two-tiered approach incorporating basic and advanced interventions may be the safest and 256 most effective. 257 258 [h2] CPR-induced consciousness 259 Cardiopulmonary resuscitation induced consciousness is uncommon but increasingly reported. 260 Rescuers may consider using sedative or analgesic drugs (or both) in small doses to prevent pain 261 and distress to patients who are conscious during CPR. 262 Neuromuscular blocking drugs alone should not be given to conscious patients.



- The optimal drug regimen for sedation and analgesia during CPR is uncertain. Regimens may be
 based on those used in critically ill patients and according to local protocols such as small doses
 of fentanyl, ketamine and/or midazolam.
- 266

267 [h2] Defibrillation

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269 [h3] Automated External Defibrillation (AED) versus manual defibrillation during ALS

- Manual defibrillators should only be used by rescuers who can quickly and accurately identify a
 cardiac arrest rhythm (within 5 seconds) and, if needed, deliver a safe shock with minimal
- interruption (less than 5 seconds) to chest compressions.
- Advanced Life Support providers must be proficient in using both an AED and a manual
 defibrillator.
- If an AED is already in use when ALS providers arrive, they should follow its shock prompts.
 When possible, they should transition to a manual defibrillator during a 2-minute CPR cycle.
- 277 [h3] Defibrillation strategy
- Continue CPR while a defibrillator is retrieved and pads applied. High quality CPR improves the
 chances of successful defibrillation.
- Give a shock as early as possible when appropriate.
- Deliver shocks with minimal interruption to chest compressions and minimise the pre-shock and
 post-shock pause. This is achieved by continuing chest compressions during defibrillator
 charging, delivering defibrillation with an interruption in chest compressions of less than 5
 seconds and then immediately resuming chest compressions.
- Immediate defibrillation of (ventricular fibrillation)VF of any amplitude (even fine VF) should be
 attempted.
- Immediately resume chest compressions after shock delivery. If there is a combination of clinical
 and physiological signs of return of spontaneous circulation (ROSC) such as return of
- 289 consciousness, purposeful movement, arterial waveform or a sharp rise in ETCO₂, consider
- 290 stopping chest compressions for rhythm analysis, and if appropriate, a pulse check.
- In defibrillators that display an ECG with motional artefact removed (from chest compressions), a
- rhythm compatible with ROSC may guide the decision to perform a pulse check every two
- 293 minutes. Asystole requires no interruption to chest compressions.
- 294
- 295 [h3] Safe and effective defibrillation



296 Minimise the risk of fire by taking off any oxygen mask or nasal cannulae and by placing them at 297 least 1 m away from the patient's chest. Oxygen exhaust from ventilation circuits should be 298 directed away from the chest. Ventilator circuits should remain attached. 299 Charging the defibrillator in anticipation of each rhythm check may minimise hands-off time 300 prior to shock delivery and is an acceptable alternative strategy if delivered without causing peri-301 shock pauses. 302 A shock with a manual defibrillator can be safely delivered without interrupting mechanical chest 303 compression. 304 Do not defibrillate during manual chest compressions (even when wearing clinical gloves), as 305 that practice is not safe to the rescuer. 306 307 [h3] Defibrillation pads and paddles 308 There is insufficient evidence to recommend a specific pad or paddle size for optimal external 309 defibrillation in adults. When available, defibrillation pads are preferable to paddles as they offer practical benefits for 310 311 routine monitoring and defibrillation. Pads enable the operator to stand clear during 312 defibrillation and minimise pre- and post-shock pauses to chest compressions by enabling hands-313 free operation. Better contact with the chest wall may also reduce the risk of arcing and 314 subsequent fires. 315 When using defibrillation paddles, apply firm force to both defibrillation paddles to optimise skin 316 contact, minimise transthoracic impedance and reduce the risk of electrical arcing. 317 Antero-lateral pad position is the position of choice for initial pad/paddle placement. In 318 particular, ensure that the apical (lateral) pad is positioned correctly (i.e. below the armpit in the 319 mid-axillary line). 320 Antero-posterior pad position may be used for vector change defibrillation following three failed 321 shocks in cases of refractory shockable rhythms. The anterior pad is placed to the left of the 322 sternum, avoiding as much breast tissue as possible. The posterior pad is placed at the same 323 height, centred just medial to the left scapula. 324 In patients with an implantable pacemaker/defibrillator (ICD), place the pad greater than 8 cm 325 away from the device, or use an alternative pad position. Consider an alternative pad position 326 when the patient is in the prone position (bi-axillary), or in a refractory shockable rhythm (see 327 below). 328 329 [h3] Energy levels and number of shocks



| 330 | • | Use single shocks followed by a 2-minute cycle of chest compressions. |
|-----|-----|--|
| 331 | • | The use of up to three stacked shocks may be considered only if initial ventricular |
| 332 | | fibrillation/pulseless ventricular tachycardia (VF/pVT) occurs during a witnessed, monitored |
| 333 | | cardiac arrest with a defibrillator immediately available, e.g. during cardiac catheterisation or in a |
| 334 | | high dependency area. (For the purposes of adrenaline administration after three failed shocks, |
| 335 | | the initial three stacked shocks should be counted as the initial shock). |
| 336 | • | Energy levels: |
| 337 | | • For biphasic waveforms (rectilinear biphasic or truncated exponential biphasic, but not |
| 338 | | pulsed biphasic), defibrillation shock energy levels for the first shock is at least 150 J. |
| 339 | | For pulsed biphasic waveforms, deliver the first shock at 130-150 J. |
| 340 | • | If the first shock is not successful and the defibrillator is capable of delivering shocks of higher |
| 341 | | energy, it is reasonable to increase the energy for subsequent shocks. |
| 342 | • | If the rescuer is unaware of the recommended energy settings of the defibrillator, for an adult |
| 343 | | use the highest energy setting for all shocks. |
| 344 | • | Use standard energy levels in obese patients. |
| 345 | | |
| 346 | [h3 |] Refractory ventricular fibrillation |
| 347 | • | Consider escalating the shock energy, after a failed shock. |
| 348 | • | For refractory VF, defined as continuous VF after three consecutive shocks, and having ensured |
| 349 | | correct antero-lateral pad positioning, consider using a defibrillation vector change by using an |
| 350 | | alternative defibrillation pad position (e.g. antero-posterior). After a failed third shock, prepare |
| 351 | | to place a fresh set of pads, at the time of the following rhythm check. Optimise transthoracic |
| 352 | | impedance by shaving the anticipated area of placement for the anterior pad (if necessary). |
| 353 | | Dual (double) sequential defibrillation (DSD), involves using a combination of antero-lateral and |
| 354 | | antero-posterior pad positioning, discharged in close succession and has been advocated for use |
| 355 | | in refractory shockable rhythms. Given the practical challenges of using two defibrillators to |
| 356 | | deliver DSD and the limited evidence for its efficacy the ERC does not recommend its routine |
| 357 | | use. |
| 358 | | |
| 359 | [h3 |] Ventricular fibrillation waveform analysis for optimising shock success |
| 360 | • | Rescuers should give defibrillation shocks according to AED prompts or use a manual defibrillator |
| 361 | | for ventricular fibrillation/pulseless ventricular tachycardia (VF/pVT) according to the ALS |
| 362 | | algorithm – there is currently no role for VF waveform analysis (e.g., based on amplitude) for |
| 363 | | identifying the optimal time for defibrillation. |
| | | |



| 364 | | |
|-----|-----|--|
| 365 | [h3 | 3] Patients with actively discharging implantable cardioverter defibrillators |
| 366 | • | Rescuers may sense a significant shock across their arms if a shock is delivered by an ICD while |
| 367 | | they are performing external chest compressions, even when wearing clinical gloves. |
| 368 | • | If an ICD fails to terminate a shockable rhythm, conventional external shocks should be |
| 369 | | delivered, placing any defibrillation pad/paddle more than 8 cm from the defibrillator box (as |
| 370 | | above). |
| 371 | • | If the ICD is incorrectly detecting arrhythmias and shocking inappropriately, a magnet placed |
| 372 | | over the ICD can temporarily stop shocks but will not disable pacing (if programmed). |
| 373 | | |
| 374 | [h2 | 2]Airway and ventilation |
| 375 | • | During CPR, start with basic airway techniques and progress stepwise according to the skills of |
| 376 | | the rescuer until effective ventilation is achieved. |
| 377 | • | Give the highest feasible inspired oxygen during CPR. |
| 378 | • | Start effective ventilation breaths as soon as possible ensuring the rate and tidal volume are |
| 379 | | appropriate to prevent both inadequate ventilation (hypoventilation) and excessive ventilation |
| 380 | | (hyperventilation). |
| 381 | • | Deliver effective bag-mask ventilation breaths by optimising mask seal and airway patency and if |
| 382 | | necessary and feasible, use a two-person technique for bag-mask ventilation. |
| 383 | • | Give each breath over 1 second to achieve a visible chest rise. |
| 384 | • | When using a supraglottic airway (SGA), an i-gel is preferred to a laryngeal tube. |
| 385 | • | Tracheal intubation should only be attempted by rescuers with a high success rate and with the |
| 386 | | use of continuous waveform capnography. The expert consensus is that a high tracheal |
| 387 | | intubation success rate is over 95% within two attempts at intubation. |
| 388 | • | Aim for less than a 5-second interruption in chest compression for tracheal intubation. |
| 389 | • | Use direct or video laryngoscopy for tracheal intubation according to local protocols and rescuer |
| 390 | | experience. |
| 391 | • | A sustained ETCO ₂ trace on waveform capnography must be used to confirm correct placement |
| 392 | | of the tracheal tube. |
| 393 | • | Once a tracheal tube or a SGA has been inserted, ventilate the lungs at a rate of 10 min ⁻¹ and |
| 394 | | continue chest compressions without pausing during ventilations. With a SGA, if gas leakage |
| 395 | | results in inadequate ventilation, pause compressions for ventilation using a compression- |
| 396 | | ventilation ratio of 30:2. |



| 397 | When using mechanical ventilation in volume-controlled mode during chest compressions set |
|-----|---|
| 398 | the ventilator to a tidal volume of 6-8 mL kg ⁻¹ (predicted body weight), or to achieve a visible |
| 399 | chest rise, the maximum inspired oxygen, a respiratory rate of 10 min ⁻¹ , an inspiratory time of 1– |
| 400 | 2 seconds, a positive end expiratory pressure (PEEP) 0-5 cm H_2O , the peak pressure alarm at 60- |
| 401 | 70 cm H_2O , and the flow trigger off. |
| 402 | If standard airway management strategies (oropharyngeal airway and bag-mask/supraglottic |
| 403 | airway/ tracheal tube) fail during cardiac arrest, appropriately trained rescuers should attempt |
| 404 | surgical cricothyroidotomy to enable oxygenation and ventilation. |
| 405 | |
| 406 | [h2] Drugs and fluids |
| 407 | [h3] Vascular access |
| 408 | • Attempt intravenous (IV) rather than intraosseous (IO) access first, to enable drug delivery in |
| 409 | adults in cardiac arrest. |
| 410 | • If IV access cannot be rapidly achieved within two attempts, it is reasonable to consider IO access |
| 411 | as an alternative route for vascular access during adult cardiac arrest. |
| 412 | |
| 413 | [h3] Vasopressor drugs |
| 414 | • Give adrenaline 1 mg as soon as possible for adult patients in cardiac arrest with a non-shockable |
| 415 | rhythm. |
| 416 | • Give adrenaline 1 mg after the third shock for adult patients in cardiac arrest with a shockable |
| 417 | rhythm. |
| 418 | • Repeat adrenaline 1 mg every 3-5 minutes whilst ALS continues. |
| 419 | |
| 420 | [h3] Antiarrhythmic drugs |
| 421 | • Give amiodarone 300 mg IV for adult patients in cardiac arrest who are in VF/pVT after three |
| 422 | shocks have been administered. |
| 423 | • Give a further dose of amiodarone 150 mg IV for adult patients in cardiac arrest who are in |
| 424 | VF/pVT after five shocks have been administered. |
| 425 | • Give the first dose of amiodarone after three shocks, and the second dose after five shocks, |
| 426 | irrespective of whether the shockable rhythms are sequential (refractory) or intermittent |
| 427 | (recurrent). |
| 428 | • Lidocaine 100 mg IV may be used as an alternative if amiodarone is not available or a local |
| 429 | decision has been made to use lidocaine instead of amiodarone. An additional bolus of lidocaine |
| 430 | 50 mg can also be given after five defibrillation attempts. |



| 431 | |
|-----|---|
| 432 | [h3] Thrombolytic drugs |
| 433 | Consider immediate thrombolytic drug therapy when pulmonary embolism is the suspected or |
| 434 | confirmed cause of cardiac arrest. |
| 435 | In select patients with suspected pulmonary embolism, consider CPR for 60-90 minutes after |
| 436 | administration of thrombolytic drugs. |
| 437 | |
| 438 | [h3] Fluids |
| 439 | Give fluids during CPR only if cardiac arrest is caused by hypovolaemia. |
| 440 | Use either isotonic saline or balanced crystalloids for fluid infusion during CPR. |
| 441 | |
| 442 | [h3] Other drugs |
| 443 | • Do not routinely give calcium, sodium bicarbonate or corticosteroids during cardiac arrest. |
| 444 | |
| 445 | [h2] ALS in highly-monitored cardiac arrest, and physiology-guided CPR |
| 446 | • A sudden decrease in ETCO ₂ may indicate a cardiac arrest or very low cardiac output state. |
| 447 | • Consider starting chest compressions if the systolic blood pressure decreases and remains < 50 |
| 448 | mmHg despite interventions. |
| 449 | In adults undergoing continuous intra-arterial blood pressure monitoring, we suggest that |
| 450 | adrenaline is initially given in small increments (e.g., 50–100 μg IV) rather than a 1 mg bolus. If a |
| 451 | total of 1 mg has been given with no response, ensure that there is no extravasation and |
| 452 | consider giving further IV adrenaline doses of 1 mg every 3-5 minutes. |
| 453 | A pragmatic approach during physiology-guided CPR is to aim for a diastolic blood pressure of ≥ |
| 454 | 30 mmHg (when using intra-arterial blood pressure monitoring) and an ETCO ₂ \geq 25 mmHg (3.3 |
| 455 | kPa). |
| 456 | |
| 457 | [h2] Waveform capnography during advanced life support |
| 458 | • Use waveform capnography to confirm correct tracheal tube placement during CPR. |
| 459 | Use waveform capnography to monitor the quality of CPR. |
| 460 | • An increase in ETCO ₂ during CPR may indicate that ROSC has occurred. However, chest |
| 461 | compression should not be interrupted based on this sign alone. Use a combination of clinical |
| 462 | and physiological signs of ROSC (e.g., consciousness, purposeful movement, arterial waveform, |
| | |



| 463 | rise in ETCO ₂) before stopping chest compressions for rhythm analysis, and if appropriate, a pulse |
|-----|---|
| 464 | check. |
| 465 | • Do not use a low ETCO ₂ value alone to decide if a resuscitation attempt should be stopped. |
| 466 | |
| 467 | [h2] Use of ultrasound imaging during advanced life support |
| 468 | Only skilled operators should use intra-arrest point-of-care ultrasound (POCUS). |
| 469 | POCUS must not cause additional or prolonged interruptions in chest compressions. |
| 470 | POCUS may be useful to identify treatable causes of cardiac arrest such as cardiac tamponade |
| 471 | and tension pneumothorax. |
| 472 | Right ventricular dilation in isolation during cardiac arrest should not be used to diagnose |
| 473 | pulmonary embolism. |
| 474 | Do not use POCUS for assessing contractility of the myocardium as a sole indicator for |
| 475 | terminating CPR. |
| 476 | |
| 477 | [h2] Devices |
| 478 | [h3] Mechanical chest compression devices |
| 479 | Consider mechanical chest compressions only if high-quality manual chest compression is not |
| 480 | practical or compromises provider safety. |
| 481 | When a mechanical chest compression device is used, minimise interruptions to chest |
| 482 | compression during device application by using only trained teams familiar with the device. |
| 483 | |
| 484 | [h3] Resuscitative endovascular balloon occlusion of the aorta (REBOA) |
| 485 | • Do not use REBOA during cardiac arrest unless being evaluated in a clinical trial. |
| 486 | |
| 487 | [h3] Intra-arrest cooling |
| 488 | • We do not recommend intra-arrest cooling during advanced life support (unless there is severe |
| 489 | hyperthermia). |
| 490 | |
| 491 | [h2] Extracorporeal CPR |
| 492 | ECPR may be considered as a rescue therapy for selected adults with IHCA and OHCA when |
| 493 | conventional CPR is failing to restore spontaneous circulation, in settings in which this can be |
| 494 | implemented. |
| 495 | |
| | |



| 496 | [h2 | 2] Pe | eri-arrest arrhythmias | | | |
|-----|--|--|---|--|--|--|
| 497 | • | Th | e 2025 ALS guidelines and algorithms focus on those arrhythmias that require immediate | | | |
| 498 | | tre | atment before or after cardiac arrest. | | | |
| 499 | • | Rescuers should seek expert advice if the arrhythmia and/or life-threatening features persist. | | | | |
| 500 | • | The assessment and treatment of all arrhythmias address the condition of the patient (stable | | | | |
| 501 | | vei | rsus unstable) and the nature of the arrhythmia. Persistent arrhythmias require careful | | | |
| 502 | | evaluation, as they are often linked to underlying structural heart disease and may indicate | | | | |
| 503 | | un | resolved issues such as myocardial ischaemia. In addition to an arrhythmia occurring | | | |
| 504 | | im | mediately after ROSC, life-threatening features in an unstable patient include: | | | |
| 505 | | 0 | Shock – recognised by hypotension (e.g., systolic blood pressure < 90 mmHg) along with | | | |
| 506 | | | signs of compensatory mechanisms, such as increased sympathetic activity, and evidence of | | | |
| 507 | | | inadequate organ perfusion | | | |
| 508 | | 0 | Syncope – as a consequence of reduced cerebral blood flow. | | | |
| 509 | | 0 | Heart failure – manifested by pulmonary oedema (failure of the left ventricle) and/or raised | | | |
| 510 | | | jugular venous pressure (failure of the right ventricle). | | | |
| 511 | | 0 | Myocardial ischaemia – may present with chest pain (angina) or may occur without pain as | | | |
| 512 | | | an isolated finding on the 12-lead ECG (silent ischaemia). | | | |
| 513 | | | | | | |
| 514 | [h3 | B]Ta | chyarrhythmias | | | |
| 515 | • | Ele | ectrical cardioversion is the preferred treatment for tachyarrhythmia in the unstable patient | | | |
| 516 | | dis | playing potentially life-threatening adverse signs or immediately after ROSC. | | | |
| 517 | Electrical cardioversion is recommended for stable patients with monomorphic VT who have | | | | | |
| 518 | | str | uctural heart disease or when it is unclear whether there is underlying heart muscle damage. | | | |
| 519 | • | Со | nscious patients require careful anaesthesia or sedation before attempting synchronised | | | |
| 520 | | car | dioversion – be aware of the risk of haemodynamic deterioration with anaesthesia/sedation. | | | |
| 521 | • | Wł | nen cardioverting atrial or ventricular tachyarrhythmias, the shock must be synchronised to | | | |
| 522 | | 00 | cur with the R wave of the ECG. | | | |
| 523 | • | Fo | r atrial fibrillation: | | | |
| 524 | | 0 | An initial synchronised shock at maximum defibrillator output, rather than an escalating | | | |
| 525 | | | approach, is a reasonable strategy based on current data. | | | |
| 526 | • | Fo | r atrial flutter and paroxysmal supraventricular tachycardia: | | | |
| 527 | | 0 | Give an initial shock of 70-120 J. | | | |
| 528 | | 0 | Give subsequent shocks using stepwise increases in energy. | | | |
| 529 | • | Fo | r ventricular tachycardia with a pulse: | | | |

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530 Use energy levels of 120-150 J for the initial shock. 0 531 • Consider stepwise increases in energy if the first shock fails to achieve sinus rhythm. 532 If cardioversion fails to restore sinus rhythm and the patient remains unstable, give amiodarone 533 300 mg intravenously over 10–20 min (or procainamide 10–15 mg/kg over 20 min) and re-534 attempt electrical cardioversion. The loading dose of amiodarone can be followed by an infusion 535 of 900 mg over 24 h. 536 Pharmacological treatment may be considered in haemodynamically stable patients with 537 monomorphic ventricular tachycardia if there is an increased risk with sedation or anaesthesia. 538 Consider amiodarone for acute heart rate control in patients with AF and haemodynamic 539 instability and severely reduced left ventricular ejection fraction (LVEF). For stable patients with 540 LVEF < 40% consider the smallest dose of beta-blocker to achieve a heart rate less than 110 min⁻ 541 ¹. Add digoxin if necessary. 542 543 [h3]Bradycardia 544 If bradycardia is accompanied by adverse signs, give atropine 500 µg IV (IO) and, if necessary, 545 repeat every 3–5 min to a total of 3 mg. 546 If treatment with atropine is ineffective, consider second-line drugs. These include isoprenaline 547 (5 μ g min⁻¹ starting dose), and adrenaline (2–10 μ g min⁻¹). 548 For bradycardia in patients with cardiac transplant or spinal cord injury, consider giving 549 aminophylline (100–200 mg slow intravenous injection). 550 Do not give atropine to patients with cardiac transplants – it can cause a high-degree AV block or 551 even sinus arrest – use aminophylline. 552 Consider giving glucagon if beta-blockers or calcium channel blockers are a potential cause of the 553 bradycardia. 554 Do not give atropine to patients with high-degree atrioventricular block and wide QRS. It is 555 ineffective and may worsen the block. 556 Consider pacing in patients who are unstable, with symptomatic bradycardia refractory to drug 557 therapies. 558 Establish early transvenous pacing in unstable patients with symptomatic bradycardia. 0 559 Consider transthoracic (transcutaneous) pacing as a bridge to transvenous pacing or when 560 transvenous pacing is not readily available. 561 Whenever a diagnosis of asystole is made, check the ECG carefully for the presence of P waves 562 because, unlike true asystole, this is more likely to respond to cardiac pacing.



- If atropine is ineffective and transvenous/transcutaneous pacing is not immediately available, fist
 pacing can be attempted while waiting for pacing equipment.
- 565

566 [h2]Uncontrolled organ donation after circulatory death

- When there is no ROSC, consider uncontrolled organ donation after circulatory death in settings
- 568 where there is an established programme, and in accordance with local protocols and legislation.
- 569
- 570



| 571 | [h1] Evidence informing the guidelines |
|-----|--|
| 572 | |
| 573 | [h2] Prevention of in-hospital cardiac arrest |
| 574 | IHCA occurs in about 1.5 patients per 1000 admitted to hospital. ⁷⁻¹¹ There are two main strategies to |
| 575 | prevent cardiac arrest and the need for attempted CPR: |
| 576 | Patient-focused decision-making to determine if CPR is appropriate. |
| 577 | Identifying and treating physiological deterioration early to prevent cardiac arrest. |
| 578 | |
| 579 | [h3] Emergency care treatment and CPR decisions |
| 580 | Most patients who die in hospital do not have a resuscitation attempt. ¹²⁻¹⁵ The ERC Guidelines on |
| 581 | Ethics guidelines promote shared decision-making and advanced care planning which integrates |
| 582 | resuscitation decisions with emergency care treatment plans to increase clarity of treatment goals |
| 583 | and also prevent inadvertent deprivation of other indicated treatments, besides CPR. Further |
| 584 | information is provided in the ERC Guidelines 2025 Ethics in Resuscitation. |
| 585 | |
| 586 | [h3] Physiological deterioration |
| 587 | In-hospital cardiac arrest is often preceded by physiological deterioration. ^{16,17} This provides an |
| 588 | opportunity to recognise deterioration and prevent cardiac arrest. The 5 key steps have been |
| 589 | conceptualised as the in-hospital chain of survival: 'staff education', 'monitoring', 'recognition', the |
| 590 | 'call for help' and the 'response'. ¹⁸ This ERC guidance is based on an ILCOR COSTR and systematic |
| 591 | review of adult rapid response systems, and UK guidance for early warning scores and recognising |
| 592 | and responding to deterioration of acutely ill adults in hospital, the ILCOR's 'Ten Steps Toward |
| 593 | Improving In-Hospital Cardiac Arrest Quality of Care and Outcomes' and the Society of Critical Care |
| 594 | Medicine's guidelines on "Recognizing and Responding to Clinical Deterioration Outside the ICU". ^{9,19-} |
| 595 | 22 |
| 596 | |
| 597 | [h4]Staff education |
| 598 | Education should include knowing the importance of timely and serial measurement of vital signs for |
| 599 | the early prediction of patient deterioration, a structured ABCDE-type approach that includes |
| 600 | assessment and initial treatment interventions, use of structured communication tools such as |
| 601 | Situation-Background-Assessment-Recommendation (SBAR), and how to call for help and escalate |
| 602 | care. ²⁰ Staff should also be aware of treatment escalation plans, protocols for critical care admission, |

603 implementation of local policies regarding do-not-attempt CPR (DNACPR) decisions and the



management of end-of-life care. Timely treatment escalation and DNACPR decisions will avoid
 ineffective treatments or treatments patients may not wish to have(see Ethics in Resuscitation).³

606

607 [h4]Monitoring

608 Most patients with IHCA have an initial non-shockable rhythm, and preceding signs of respiratory 609 depression or shock are common.^{7,8,23} Although the supporting evidence is low to very low certainty, 610 there is consensus that to help detect deterioration and critical illness early, all patients should have 611 a documented plan for vital sign monitoring that includes which physiological measurements should 612 be recorded and how frequently.^{24,25} This can be addressed by using a standardised early warning score (EWS) system for all patients.²⁶ The choice of system depends on local circumstances and 613 614 should align with national guidelines. For example in the UK, the National Early Warning Score 615 (NEWS) is endorsed by the National Institute for Health and Care Excellence (NICE) guidelines.^{19,20} 616 Higher trained and nurse staffing levels are associated with lower rates of failure to respond to 617 abnormal vital signs, and improved patient outcomes.^{27,28} There is a lack of randomised controlled 618 trials (RCTs) or consensus on which patients should undergo continuous ECG or other continuous 619 vital sign monitoring.¹¹ In a registry-based study, settings where patients are closely monitored were 620 associated with improved survival irrespective of initial rhythm.²⁹ The use of artificial intelligence to 621 predict patient deterioration has gained interest during recent years but currently the evidence does 622 not support its wider adoption without further studies on effectiveness and impact on clinical 623 management.³⁰ Implementation of an automated predictive model to identify high-risk patients in 19 624 United States hospitals was associated with decreased mortality.³¹

625

626 [h4]Recognition

In non-ICU patients, strategies to simplify and standardise tracking of a patient's condition, and recognising acute illness or deterioration, and triggering a response include early warning score systems. These scoring systems have a predefined graded and escalating response according to the patient's early warning score. The early warning score is used to identify ward patients needing escalation of care, increasing vital sign monitoring, and may improve identification of deterioration, and reduce time to emergency team activation.³² Clinical concern from nurses and other members of the multidisciplinary team can also indicate patient deterioration.^{33,34}

....

635 [h4]The call for help

All staff should be empowered to call for help and also trained to use structured communication
 tools such as SBAR to ensure effective communication.³⁵⁻³⁷ The response to patients who are critically



ill or who are at risk of becoming critically ill is often provided by a rapid response system (which
includes medical emergency team (MET), rapid response team (RRT), or critical care outreach team
(CCOT)). Any member of the healthcare team can initiate a call to such a team according to explicit
activation criteria. In some hospitals, the patient, and their family and friends, are also encouraged
to activate the team.³⁸⁻⁴⁰ Such a patient safety initiative is being implemented throughout English

- 643 hospitals.⁴¹
- 644

645 [h4]Response

- 646 The response to patients who are or at risk of being critically ill is often provided by a
- 647 MET/RRT/CCOT. These teams usually comprise critical care medical and nursing staff who respond to
- 648 specific calling criteria. They replace or coexist with traditional cardiac arrest teams, which typically
- only respond to patients already in cardiac arrest. These teams regardless of composition should
- 650 function 24/7. Systematic reviews, meta-analyses and multicentre studies suggest that
- 651 RRT/MET/CCOT systems reduce the rate of IHCA and hospital mortality.^{42,43} These data led ILCOR to
- 652 suggest that hospitals consider the introduction of rapid response systems to reduce the incidence of
- 653 IHCA and in-hospital mortality (weak recommendation, low-certainty evidence).²¹ Team
- interventions often involve simple tasks such as starting oxygen therapy and IV fluids, as well as more
- 655 complex decision-making such as transferring the patient to the intensive care unit (ICU) or initiating
- discussions regarding DNACPR, treatment escalation or end-of-life care plans (See Ethics guidelines³).
- An important part of the response is to place a patient at risk of deterioration, or an already
- 658 deteriorating patient, in an appropriate setting. Patients should be treated in a clinical area that is
- equipped and staffed to meet the patient's needs. A quality improvement process should be
- 660 implemented for RRT/MET/CCOT systems.⁹
- 661

662 [h2] Prevention of out-of-hospital cardiac arrest

In high-income settings, sudden cardiac arrest (SCA) is the third leading cause of death. Survival
 following OHCA is generally no more than 10% or less,⁴⁴⁻⁴⁶ and a positive survival trend has been
 observed in only half of the countries studied recently which underscores the importance of OHCA
 prevention.^{47,48}

- 667
- 668 Even seemingly healthy young adults who experience SCA may present with prodromal signs and
- 669 symptoms—such as syncope, pre-syncope, chest pain, or palpitations—that should prompt
- 670 healthcare professionals to seek expert evaluation to prevent cardiac arrest.⁴⁹⁻⁵⁸
- 671



- 672 There is no ILCOR systematic review on this topic and existing guidelines of the European Society of
- 673 Cardiology (ESC), the American Heart Association (AHA) and ERC⁵⁹ were also considered.
- 674

675 [h3] Epidemiology and pathophysiology of sudden cardiac arrest

- 676 Coronary heart disease (CHD) is the underlying cause of SCA in 80% of cases, especially in older
- 677 patients, and non-ischaemic cardiomyopathies account for another 10-15%.⁶⁰ In the young, inherited
- diseases, congenital heart disease, myocarditis and substance abuse are predominant causes.
- 679 Knowledge of the causes of SCA assists in early treatment and the prevention of OHCA (Table 2).
- 680
- 681 **Table 2. Causes of sudden cardiac arrest (SCA)** Adapted from Kandala⁶⁰ and Winkel.⁶¹

| Coronary heart disease | | |
|--|--|--|
| ST-segment elevation | | |
| Other myocardial infarction | | |
| Unstable angina | | |
| Silent ischaemia | | |
| Electrical heart disease, often associated with SCA in the young | | |
| Long QT-syndrome (LQTS) | | |
| Short QT syndrome | | |
| Brugada syndrome | | |
| Catecholaminergic Polymorphic Ventricular Tachycardia (CPVT) | | |
| Triadin knock-out syndrome (TKOS) | | |
| Arrhythmogenic bi-leaflet mitral valve prolapse | | |
| Drug or medication induced | | |
| Non-atherosclerotic coronary artery anomalies | | |
| Congenital heart disease | | |
| Hypertrophic cardiomyopathy (HCM) | | |
| Dilated cardiomyopathy (DCM) | | |
| Valvular heart disease | | |

682

683 [h3] Coronary heart disease (CHD)

- 684 Arrhythmias triggered by acute myocardial infarction (AMI) or subsequent myocardial scarring can
- 685 result in SCA.⁶² About two-thirds of SCDs occur as the first CHD event or in individuals considered to
- 686 be at low risk.⁶⁰During the last 50 years primary prevention and secondary revascularisation have



- 687 reduced CHD age-adjusted mortality.⁶⁰The percentage of SCDs associated with CHD remains 688 unchanged, suggesting that there are interactions between CHD and triggering events such as 689 autonomic nervous system dysfunction, electrolyte disturbances, drug toxicity and individual genetic 690 profiles.⁶⁰Cardiac electrophysiology studies can identify patients with CHD at high versus low risk of 691 SCA.⁶³ Additional factors such as heart failure (HF) and left ventricular hypertrophy (LVH) predispose 692 to ventricular arrhythmias (polymorphic VT and VF). How to effectively identify patients at high risk 693 of SCA with HF and LVH is uncertain.⁶⁴ Changes in left ventricular geometry affect the likelihood of 694 developing VT and VF. The only indicator that has been identified to be consistently associated with 695 an increased risk of SCA in the setting of CHD and left ventricular (LV) dysfunction is LV ejection 696 fraction (LVEF).⁶² LVEF is used to indicate the need for an ICD for the primary and secondary 697 prevention of SCA.65 698 Cardiac MRI has been proposed as a tool for detecting scar burden and assessing risk of SCA. 699 Recently, artificial intelligence (AI) and deep learning analysis of the scars have been used to assess 700 patient-specific prognosis.⁶⁶ Despite considerable progress, the ability to recognise the risk of SCA
- 701 before the event remains very limited.⁶².
- 702

703 [h3] Sudden cardiac arrest in the young

704 Sudden cardiac arrest in the young (SCDY, 5-35 years of age) accounts for 7% of all SCDs;⁶¹ with an 705 incidence of 1-8/100 000 fatalities per year.⁶⁷ In adolescent SCA, 50% of patients had misinterpreted 706 symptoms before death.⁵⁷ CHD is the most frequent cause of explained SCDY, but 25-31% of the 707 cases remain unexplained after post mortem examination (Sudden Arrhythmic Death Syndrome-708 SADS).⁶⁸ The majority of inherited cardiac diseases can be treated if diagnosed, yet most young SCA 709 victims are not diagnosed.⁵⁵ Premonitory signs of SCDY were present in only 29% in one study, and thus less common than in older patients.⁶⁹ QT-prolonging and psychotropic drugs, alone or in 710 711 combination, increase the risk of SCA.⁶¹ Investigation of cardiac arrest survivors or post mortem 712 examination is crucial to identify inherited cardiac disease in unexplained cases of SCA; this should 713 result in a cardiac investigation of first-degree relatives. In one study, this screening resulted in a 714 diagnosis of an inherited cardiac disease in over half of the families.⁷⁰ In a large retrospective SCDY 715 study, a cause was identified in 113/180 patients (62.8%), the rest were classified as idiopathic VF.⁷¹ 716 With improvements in diagnosis (e.g. provocation drug testing for cardiac channelopathies and 717 coronary vasospasm, genetic testing), the number of unexplained SCDs should decrease.⁷¹ (See Epidemiology).47 718

- 719
- 720 [h3] Non-atherosclerotic coronary artery anomalies



| Coronary artery embolism, coronary arteritis (e.g. Kawasaki disease, polyarteritis nodosa), spasm |
|---|
| and myocardial bridging have all been described with SCA. |
| |
| [h3] Congenital heart disease |
| Congenital coronary anomalies are present in 1% of all patients. SCA because of congenital coronary |
| anomalies is exercise-related and accounts for 17% of SCA in young athletes. ^{60,69} |
| |
| [h3] Hypertrophic cardiomyopathy (HCM) |
| Hypertrophic cardiomyopathy is the most common genetic disorder of the heart, and the most |
| common cause of SCDY. ⁷² It often remains clinically silent until SCA presents as the first cardiac event. |
| The incidence of SCA in families with HCM is 2-4% a year and 4-6% in children and adolescents. ⁶⁰ |
| |
| The 2022 ESC guideline on the management of ventricular arrhythmias and prevention of sudden |
| cardiac arrest proposed 10 new key aspects that may improve the management of SCD. ⁷³ (Table 3). |
| |
| Table 3. Key points from the European Society of Cardiology guidelines for the treatment of |
| ventricular arrhythmias and sudden cardiac death (Adapted from Könemann, 202373) |
| 1 Development of public BLS and access to automatic defibrillators |
| 2 Focus on the management of electrical storm |
| 3 Increased relevance of cardiac MRI |
| 4 Increased relevance of catheter ablation |
| 5 Implementation of SCA risk scores and calculators |
| 6 New algorithms for diagnostic evaluation |
| 7 Upgrade of genetic counselling and testing |
| 8 Algorithm for antiarrhythmic drug therapy |
| |

9 Individualized risk stratification

10 Change regarding primary electrical diseases

738

739 The prediction of SCA presents an epidemiological paradox: although high-risk patients have a

740 greater individual risk, the absolute number of OHCA is higher in the much larger low-risk general

population. Preventing and predicting SCA is challenging, as most events occur in individuals from

the general population without known heart disease. In response, *The Lancet Commission on SCA*

recently issued a call for multidisciplinary action to reduce the burden of SCA, addressing all aspects



- 744 of prevention and treatment.⁷⁴ The development of high-quality, population-based registries of
- 745 OHCA is important for improving our understanding and prediction of SCA.
- 746 However, there are currently no established strategies or guidelines for preventing OHCA in the
- 747 general population.
- 748
- 749 SCA may be associated with a range of factors—some related to cardiovascular disease, others linked
- 750 to the broader socioeconomic environment (e.g., obesity, climate, pollution, lifestyle).
- 751
- 752 Many drugs prescribed commonly (antibiotics, antidepressants) impacting cardiac electrophysiology
- 753 mainly by prolonging the QT interval, may also increase the risk of OHCA.⁷⁵ Recently, proton pump
- 754 inhibitors, were also identified to be associated with this risk even in patients without cardiovascular
- 755 disease.76
- 756
- 757 Al and machine learning models offer new possibilities by linking cardiac arrest patient's files and the
- 758 health records of the general population. It may identify new factors driving risk of SCA and lead to
- 759 improved targeted screening at the patient level.⁷⁷ AI may also help to predict SCA with pulseless
- 760 electrical activity (PEA) and the understanding of mechanisms and warning symptoms, in this
- 761 population with a poorer prognosis of survival.⁷⁸
- 762

763 [h2] Premonitory signs

- 764 Approximately 50% of cardiac arrests occur in individuals with undiagnosed CHD. ^{62,79} Many SCA 765 victims have a history of cardiac disease and warning signs before cardiac arrest, most commonly 766 chest or upper abdominal pain or dyspnoea that has not been acted on by the patient or health care 767 professionals.^{80,81} Approximately one third of elderly patients will have symptoms in the days or 768 hours before cardiac arrest; primarily chest pain, dyspnoea, syncope, and/or cold sweats.^{81,82} In 1960 769 OHCA patients, 9.4% had been assessed by an ambulance crew within the preceding 48 h.⁸³ 770 Emergency care in patients with symptoms is associated with improved survival.⁸⁰ Early recognition 771 of acute coronary syndrome (ACS) by EMS teams with 12-lead ECG capabilities and reduction of time 772 to reperfusion may prevent SCA.84 773 The most effective approach to prevent SCA in the general population remains the quantification of
- 774 the individual risk of developing CHD, followed by control of risk factors.⁸⁵ Syncope can be an
- 775 important premonitory sign of SCA.
- 776
- 777 [h3] Syncope



- 5778 Syncope occurring during strenuous exercise, while sitting or in the supine position should always
- raise the suspicion of a cardiac cause; in other situations it is more likely to be vasovagal syncope or
- postural hypotension.⁸⁴ In patients with known cardiac disease, syncope (with or without prodrome,
- particularly recent or recurrent) is an independent risk factor for increased risk of death.^{65,72,86-96}
- High-risk (suggesting a serious condition) and low-risk features (suggesting a benign condition) of
- 783 patients with syncope at initial evaluation in the emergency department have been published by the
- 784 ESC (Table 4).⁶⁵ Early EMS acquisition of a 12 lead-ECG may be helpful.
- 785
- 786 Table 4. High risk features suggesting a serious condition in patients with syncope at initial

787 evaluation in the emergency department

- 788 Adapted from Brignole 2018.⁶⁵ ECG electrocardiogram, ICD implantable cardioverter defibrillator,
- 789 LVEF left ventricular ejection fraction, SCA sudden cardiac arrest, VT ventricular tachycardia.

Syncopal event features

Major

New onset of chest discomfort, breathlessness, abdominal pain or headache⁹⁷⁻⁹⁹

Syncope during exertion or when supine¹⁰⁰

Sudden onset palpitation immediately followed by syncope¹⁰⁰

<u>Minor</u>

No warning symptoms or short (<10 sec) prodrome ¹⁰⁰⁻¹⁰³

Family history of SCA at young age¹⁰⁴

Syncope in the sitting position¹⁰⁵

Past medical history

<u>Major</u>

Severe structural or coronary artery disease (heart failure, low LVEF or previous myocardial infarction)^{97,99}

Physical examination

Major

Unexplained systolic blood pressure <90 mmHg^{97,99}

Persistent bradycardia (<40 min⁻¹) in an awake state, in absence of physical training

Undiagnosed systolic murmur

ECG

<u>Major</u>

ECG changes consistent with acute ischaemia



Mobitz II second- and third-degree atrioventricular (AV) block Slow atrial fibrillation (AF) (<40 min⁻¹) Persistent sinus bradycardia (<40 min⁻¹) or repetitive sinoatrial block or sinus pauses >3 second in an awake state, in absence of physical training Bundle branch block, intraventricular conduction disturbance, ventricular hypertrophy or Q waves consistent with ischaemic heart disease or cardiomyopathy98,103 Sustained and non-sustained VT Dysfunction of an implantable cardiac device (pacemaker or ICD) ST-segment elevation with type 1 morphology in leads V1-V3 (Brugada pattern) QTc >460 ms in repeated 12-lead ECGs indicating long QT syndrome (LQTS)⁸⁴ Minor (high-risk only if history consistent with arrhythmic syncope) Mobitz I second-degree AV block and 1st degree AV block with markedly prolonged PR interval Asymptomatic inappropriate mild sinus bradycardia (40-50 bpm.)¹⁰³ Paroxysmal supraventricular (SVT) or atrial fibrillation¹⁰⁶ Pre-excited QRS complex Short QTc interval (<=340 ms)⁸⁴ Atypical Brugada patterns⁸⁵ Negative T waves in right precordial leads, epsilon waves suggestive of arrhythmogenic right

ventricular cardiomyopathy (ARVC)⁸⁵

790

Screening programs for athletes may be helpful but vary between countries.¹⁰⁷⁻¹⁰⁹ In one study from 791 792 the United Kingdom between 1996 and 2016, 11,168 athletes received cardiovascular screening and 793 diseases associated with SCA were identified in 0.38% (n=42).¹¹⁰ The incidence of SCA in competitive 794 athletes is higher than in non-athletes.¹¹⁰ Sub populations that have been identified as at increased 795 risk include male, black ethnicity, basketball or football players.¹¹¹ Screening commonly includes 796 physical examination and ECG. The ECG has a false positive risk because of some ECG features that 797 are particular to athletes. Despite more specialised screening, the risk decreases globally but SCA 798 may still occur. Consequently awareness, CPR training, and availability of AEDs during sport remain 799 important for the protection of athletes.^{112,113}

800

801 [h3] Preventive measures against sudden cardiac arrest

802 Prevention of SCA focuses on identifying and managing medical conditions that may contribute to or

803 exacerbate arrhythmias, assessing the risk posed by the arrhythmia itself, and evaluating the risk-


804 benefit ratio of potential therapies. Interventions may include anti-arrhythmic medications, ICDs,

- 805 and catheter ablation or surgery.^{65,114} The effective management of non-cardiovascular diseases
- 806 associated with an increased risk of cardiac arrest has also been proposed as a strategy for
- 807 preventing SCA.¹¹⁵ For instance, a large registry study found that treatment of sleep apnoea with
- 808 continuous positive airway pressure (CPAP) was associated with a lower risk of OHCA compared with
- 809 patients who did not receive treatment.¹¹⁶
- 810
- 811 Noninvasive telemetry or implantable devices transmitting the ECG are currently used in selected
- 812 groups of patients to detect high risk arrhythmias and prevent SCA. More recently, connected
- 813 devices with arrhythmia detection capabilities (smartwatch, smartphone applications) have been
- 814 introduced and may be helpful in detecting asymptomatic AF, however their potential role in the
- 815 general population to detect SCA arrhythmias is unknown.^{117,118} A recent review on smartwatches
- 816 identified 57 publications including 24 cohort studies mostly focused on AF, and often detected by
- 817 Apple Watch^{TM, 119} Automated cardiac arrest diagnosis, using smart devices such as wearables and
- 818 phones, remains an innovative field of research. It promotes the possibility of transforming
- 819 unwitnessed SCA to witnessed events. However, most of the studies published have been aimed at
- 820 feasibility and concern a small population.¹²⁰ They need to be validated in diverse populations and
- then integrated into EMS protocols. They carry the risk of false positive alarms which may be
- 822 responsible for patient anxiety and stress and inappropriate activation of the EMS response,
- 823 overwhelming the available resources.¹²¹
- Educating the public to report on symptoms before SCA and to help a person in cardiac arrest is
 important.⁸⁰ An awareness campaign on chest pain was associated with an increase in EMS calls and
 a reduction in OHCA incidence and may serve in part, as an effective primary prevention strategy for
 OHCA. The campaign period was associated with an 8.8% (Incident rate ratio [IRR] 1.09, 95% CI: 1.07,
 1.11) increase in the incidence of EMS attendances for chest pain and a 5.6% (IRR 0.94, 95% CI: 0.92,
 0.97) reduction in OHCA attendances.¹²²
- 830

831 [h2] Treatment of in-hospital cardiac arrest

Cardiac arrest treatment principles, such as early defibrillation, early delivery of adrenaline and
delivery of high-quality CPR, are consistent across both the IHCA and OHCA settings. In the hospital
setting, the immediate availability of trained clinical staff and equipment provides an opportunity for
the rapid identification of cardiac arrest and initiation of treatment. An IHCA can be defined as any
cardiac arrest that occurs on the hospital premises. This may include a cardiac arrest in patients,



- 837 hospital visitors or staff, in a variety of hospital settings. For IHCA, BLS and ALS interventions can
- often start and take place at the same time (Figure 2).
- 839
- 840 **Figure 2. In-hospital resuscitation algorithm** ABCDE airway breathing circulation disability exposure;
- 841 AED automated external defibrillator; ALS advanced life support; BP blood pressure; CPR
- 842 cardiopulmonary resuscitation; ETCO₂ end-tidal carbon dioxide; IV intravenous; SBAR situation,
- background, assessment, recommendation; SpO₂ oxygen saturation measured with pulse oximetry.
- 844



845 846

847 [h3] Initial responders

The clinical skill of the initial responder may range from a non-clinical member of staff trained in BLS to an ALS provider. Irrespective of skill level, the initial action of the initial responder is to recognise cardiac arrest, immediately start CPR, call for help and facilitate rapid defibrillation. Delays in starting treatment reduce the likelihood of a successful outcome.^{123,124}

852

The process for calling for help may differ between hospitals or locations within a hospital. If the responder is alone, they may need to leave the patient to call for help. Where a telephone system is used to activate the emergency team, the standard European number (2222) should be used.¹²⁵ To date, hospital uptake of the standard number across European countries has been variable.¹²⁶⁻¹²⁸



- 858 Following the completion of initial actions and provided sufficient staff are available, staff should
- 859 collect ALS equipment and prepare to hand over to the resuscitation team using a standardised
- 860 communication system, such as SBAR (Situation, Background, Assessment, Recommendation) or
- 861 RSVP (Reason, Story, Vital Signs, Plan).^{35,129,130} Each clinical area in a hospital should consider patient
- acuity, risk of cardiac arrest, and geographical location (e.g. distance for the resuscitation team to
- travel) in determining the specific training needs of staff.
- 864

865 [h3] Resuscitation team

- 866 The resuscitation team may take the form of a traditional cardiac arrest team that responds only to
- 867 cardiac arrest events or a medical emergency team/ rapid response team (MET/RRT) that responds
- 868 to both cardiac arrests and critically unwell patients. ILCOR recommends accredited ALS-level
- training for healthcare staff (strong recommendation based on very low certainty evidence) as this
- 870 type of training is associated with improved patient outcomes.^{21,131} ILCOR also recommends that ALS
- 871 training incorporates team and leadership training (weak recommendation based on very low
- 872 certainty evidence) because it is associated with improved patient and process outcomes.²¹
- 873
- Resuscitation teams often form on an ad hoc basis depending on hospital work rosters and may
 include individuals from a range of specialities (e.g. emergency medicine, acute medicine, cardiology,
 critical care, anaesthesia). Lack of knowledge of team member roles, including who is acting as team
 leader can lead to errors during ALS for IHCA.^{132,133} A team meeting at the beginning of each shift for
 introductions and allocation of roles is straightforward to implement and may support effective
 team-working during resuscitation, although its effect on patient outcomes is uncertain.¹³⁴
- 880

881 [h3] Equipment

- Hospitals should ensure that clinical areas have immediate access to resuscitation equipment and
 drugs to facilitate rapid resuscitation of the patient in cardiac arrest. Missing or malfunctioning
 equipment contributes to treatment delays.^{132,135} Equipment should be standardised throughout the
 hospital and regularly checked to ensure proper functioning. In contrast to OHCA, most patients who
 have an IHCA will have vascular access at the time of cardiac arrest, facilitating rapid administration
 of time-critical drugs, such as adrenaline.¹³⁶⁻¹³⁸
- 889 [h3] Termination of CPR rules for in-hospital cardiac arrest
- 890 Research on termination of resuscitation (TOR) rules for IHCA remains limited. An ILCOR systematic



review published in 2021 identified only very low-certainty evidence for a single clinical decision rule,

the UN10 rule, which included three variables: unwitnessed, initial non-shockable rhythm, and

resuscitation duration of more than 10 minutes.¹³⁹⁻¹⁴¹ While the rule had a false-positive rate of 0% in

the derivation cohort, later validation revealed a false-positive rate above the 1% threshold deemed

acceptable for clinical use.³ Consequently, ILCOR concluded that no identified tool was reliable in

896 predicting death after IHCA and specifically recommended against using the UN10 as a sole strategy

- 897 for termination of resuscitation in IHCA (strong recommendation, very low–certainty evidence).¹⁴²
- 898 Following the 2021 ILCOR systematic review, a Scandinavian study enrolling IHCA patients from
- Denmark, Sweden, and Norway developed and validated five TOR rules for IHCA.¹⁴³ The best-
- 900 performing rule included four variables (unwitnessed, unmonitored, initial rhythm of asystole, and
- 901 resuscitation duration of 10 minutes or more). This rule incorrectly predicted 30-day mortality in 6
- 902 per 1000 cases, and proposed termination in 110 per 1000 cardiac arrests, potentially reducing futile
- 903 resuscitation attempts. Notably, a large observational study demonstrated that survival to hospital
- discharge is possible even after prolonged IHCA (>1 hour), although survival rates were less than 1%
- 905 after 40 minutes of resuscitation.¹⁴⁴ An Austrian study found that machine learning models
- 906 effectively predicted failure to achieve ROSC and poor functional outcomes while CPR was ongoing;
- 907 however, the positive predictive value was insufficient to justify early termination of resuscitation
- 908 efforts.¹⁴⁵

909 In alignment with ILCOR, the ERC does not recommend using TOR rules as the sole basis for TOR in

910 IHCA. While the results of the Scandinavian study are promising, further external validation is

911 needed before the rule can be considered for use in clinical practice.¹⁴³ TOR rules should be

912 validated in local, regional, or national cohorts before implementation and revisited as survival rates

- 913 change. Decisions to terminate resuscitation should also consider the local legal, organisational, and
- 914 cultural context. The ERC Guidelines 2025 on Ethics in Resuscitation provide additional guidance on
- 915 the use of termination of resuscitation rules.³
- 916

917 [h2]Treatment of out-of-hospital cardiac arrest

918 This section provides an overview of specific ALS issues related to resuscitation for OHCA. Further

919 information is available in the ERC Guidelines 2025 Basic Life Support, Cardiac Arrest in Special

- 920 Circumstances, Systems Saving Lives, Epidemiology, Post-resuscitation Care, and Ethics in
- 921 Resuscitation.^{3,5,113,128,146,147}
- 922



| 923 | [h3] Transfer of patients with OHCA |
|-----|--|
| 924 | The aim of ALS for OHCA is to provide the same interventions as available in hospital as early as |
| 925 | possible, and to rapidly transfer the patient to hospital for those interventions that are not feasible |
| 926 | out-of-hospital. A recent systematic review addressed the benefit of rapid transport from the scene |
| 927 | to definitive in-hospital care versus extended on-scene resuscitation in OHCA. ¹⁴⁸ Nine studies (8 |
| 928 | cohort studies, one RCT) were included. In pooled analysis, expedited (or earlier) transfer was not |
| 929 | predictive of survival to discharge (odds ratio [OR] 1.16, 95% confidence interval [CI] 0.53 to 2.53, I^2 = |
| 930 | 99%, p = 0. 65) or favourable neurological outcome (OR 1.06, 95% CI 0.48 to 2.37, I ² = 99%, p = 0.85). |
| 931 | The certainty of evidence was assessed as very low with a moderate risk of bias. Significant |
| 932 | heterogeneity was observed mostly related to the region of the EMS studied. In a large North |
| 933 | American registry a propensity-matched cohort, which included 27,705 patients, survival to hospital |
| 934 | discharge occurred in 4.0% of patients who underwent intra-arrest transport compared with 8.5% |
| 935 | who received on-scene resuscitation (risk difference, 4.6% [95% Cl, 4.0%- 5.1%]). ¹⁴⁹ |
| 936 | |
| 937 | [h3] Care at cardiac arrest centres |
| 938 | An ILCOR systematic review assessed the benefits of care at a dedicated cardiac arrest centre. The |
| 939 | resulting ILCOR treatment recommendations include ¹⁵⁰ : |
| 940 | Adult patients with non-traumatic OHCA should be considered for transport to a cardiac |
| 941 | arrest centre, according to local protocols. |
| 942 | Adult patients with non-traumatic OHCA should be cared for at a cardiac arrest centre |
| 943 | whenever possible. |
| 944 | Health care networks should establish local protocols to develop and maintain a cardiac |
| 945 | arrest network. |
| 946 | The ERC has adopted these recommendations on cardiac arrest centres and further details can be |
| 947 | found in ERC Guidelines 2025 for Systems Saving Lives and Post Resuscitation Care. |
| 948 | |
| 949 | [h3] Initial Treatment of OHCA |
| 950 | Several patient and CPR factors affect outcome from OHCA (Table 5). Community programmes of lay |
| 951 | bystander CPR and AED use improve outcome from OHCA. ¹⁵¹ Chest compressions and early |
| 952 | defibrillation are the cornerstones of CPR in OHCA. The only definitive treatment for VF remains |
| 953 | prompt defibrillation. ¹⁵² |
| | |

954 **Table 5. Patient and resuscitation factors affecting outcome from OHCA**. Adapted from Kandala

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955 2017.⁶⁰ AED denotes automated external defibrillator, CPR cardiopulmonary resuscitation.

| Patient | |
|---|--|
| Age | |
| Sex | |
| Comorbidities (cardiac, pulmonary, renal, trauma) | |
| Special circumstances | |
| Cardiopulmonary resuscitation | |
| Location (private vs. public) | |
| Witnessed vs. unwitnessed cardiac arrest | |
| Bystander CPR | |
| Type of bystander CPR (compression only vs. standard) | |
| First cardiac arrest rhythm | |
| Use of AED by bystander | |
| Time to return of spontaneous circulation | |

956

957 [h3] EMS personnel and interventions

958 ILCOR conducted a systematic review examining how EMS exposure to and experience with OHCA 959 impacts outcomes.¹⁵³ The largest study in this review linked exposure of paramedics to OHCA, 960 defined as the number of times a paramedic had attended an OHCA, to patient survival to hospital 961 discharge.¹⁵⁴ Increasing exposure in the preceding three years was associated with increased survival 962 to discharge: ≤6 exposure (control group), >6-11 exposures (adjusted odds ratio [aOR] 1.26, 95% CI 963 1.04-1.54), 11-17 exposures (aOR 1.29, 95%Cl 1.04-1.59), >17 exposures (aOR 1.50, 95%Cl 1.22-964 1.86).¹⁵⁴ Another large observational study reported that increased exposure of the treating paramedic was associated with increased ROSC (<15 exposures [control group] vs. ≥15 exposures, 965 966 aOR 1.22, 95%CI 1.11-1.36).¹⁵⁵ The ILCOR CoSTR concluded that EMS should monitor exposure of 967 their clinical personnel to resuscitation and implement strategies to address low exposure or ensure 968 that treating teams have members with recent exposure (weak recommendation, very-low certainty 969 of evidence).²²



970 There is no recommendation on the optimal number of members in prehospital ALS teams. A recent 971 review of 22 articles published between 2005 and 2023 assessed the effectiveness of ALS CPR 972 performed by two-member teams and found insufficient evidence to support adapting ALS protocols 973 to such settings. ¹⁵⁶ A recent review in 2023 identified four non-RCT studies and concluded that 974 prehospital ALS care with a ratio of on-scene ALS-trained personnel >50% could improve survival-todischarge with a certainty of evidence rated as very low.¹⁵⁷ A large national cohort analysis 975 976 conducted in a system with prehospital ALS and the initial dispatch of more than one EMS crew 977 found both a higher number of EMS crew members (three or more) and a higher proportion of ALS 978 providers in the first-contact EMS crew. This was associated with improved neurological recovery in 979 adults with non-traumatic OHCA. Specifically, good neurological recovery was associated with an 980 adjusted odds ratio (95% confidence interval) of 1.23 (1.06–1.43) for three-member crews, and 1.28 981 (1.17–1.40) for crews with a higher proportion of ALS-trained providers.¹⁵⁸ 982

983 A 2024 ILCOR review compared prehospital critical care for OHCA with standard prehospital ALS.^{22,159} 984 Prehospital critical care was defined as care involving enhanced clinical competencies beyond 985 standard ALS, delivered by dedicated EMS teams dispatched to critically ill patients. These teams 986 were staffed by physicians (specialised in emergency medicine, anaesthesia, critical care, or intensive 987 care) or specially trained critical care paramedics. This was compared with standard prehospital ALS. 988 The review included 15 articles. Prehospital critical care was associated with improved outcomes for 989 several measures: survival to hospital admission, ROSC (OR 1.95, 95% CI 1.35–2.82), survival to 990 hospital discharge (OR 1.34, 95% CI 1.10–1.63), 30-day survival (OR 1.56, 95% CI 1.38–1.75), and 991 favourable neurological outcome at 30 days (OR 1.56, 95% CI 1.38–1.75). ILCOR recommends that 992 adults with non-traumatic OHCA receive care from prehospital critical care teams in EMS systems 993 with sufficient resource infrastructure (weak recommendation, low certainty of evidence). However, 994 the included studies did not report on resource costs, cost-effectiveness, impact on health equity, or 995 implementation feasibility; therefore, these aspects were not analysed. 996 Prehospital critical care teams also provide the opportunity to complement ALS with more advanced 997 and invasive resuscitation techniques, such as ECPR, balloon occlusion, and emergency 998 thoracotomy.¹⁶⁰ These techniques, their targets populations, and their potential impacts on outcome

999 1000

1001 [h3] Termination of CPR rules for out-of-hospital cardiac arrest

are discussed in other parts of the guidelines.

1002 Termination of resuscitation (TOR) rules guide EMS in deciding whether to continue resuscitation or 1003 transport a patient with ongoing CPR. An ILCOR review found that current TOR rules may result in



1004 some missed survivors but do also prevent premature termination of resuscitation.^{1,22,161} ILCOR

1005 makes a conditional recommendation based on very-low certainty evidence that EMS systems may

1006 use TOR rules to guide decisions on stopping resuscitation or transporting with ongoing CPR – the

1007 rules should only be implemented after local validation, ensuring acceptable specificity and

- alignment with local culture, values, and context. Further ethical guidance is available in the ERC
- 1009 Guideline 2025 Ethics in Resuscitation.³
- 1010

1011 [h2]Debriefing

1012 In 2020, ILCOR conducted a systematic review of debriefing following cardiac arrest²¹, including four 1013 observational studies.¹⁶²⁻¹⁶⁵ At that time, debriefing was associated with improved hospital survival, 1014 ROSC, and CPR quality. In 2024, ILCOR conducted a new systematic review, incorporating ten nonrandomised studies—six involving adult patients^{134,162,164-167} one paediatric¹⁶³, and two neonatal 1015 1016 cardiac arrest.^{168,169} This updated review revealed that, despite the very low certainty of evidence because of significant risks of bias and inconsistency, post-event debriefing was either associated 1017 1018 with no effect or with improvements in ROSC, survival to hospital discharge, favourable neurological 1019 outcomes, and enhanced CPR quality. The ILCOR review did not identify any negative consequences, 1020 such as emotional trauma to the debriefed team or significant resource demands (including costs), 1021 associated with debriefing after cardiac arrest in the studies examined. Based on these findings, 1022 ILCOR suggests, and the ERC recommends, implementing post-event debriefing following adult 1023 cardiac arrest (weak recommendation, very low certainty of evidence). This recommendation stems 1024 from the review's conclusion that debriefing has a neutral to positive impact on critical and 1025 important outcomes, which likely outweighs any potential undesirable effects. The ERC Guidelines 1026 2025 Education for Resuscitation include further details on these issues.⁴

1027

1028 [h2] ALS in low-resource settings

1029 This review was informed by a narrative ILCOR review. ¹⁷⁰ Low-resource settings in CPR are most 1030 often associated with low-income countries. High-income countries may also have low-resource 1031 settings (e.g., mass-casualty incidents, at night or bad weather, natural disasters, pandemics or war) 1032 or in space (e.g. site difficult to access - mountains and sea - or remote, for example aircraft, oil 1033 platforms, ships). Implementation of resuscitation guidelines from well-resourced settings may not 1034 be applicable in low-resource settings because of lack of logistics, personnel and infrastructure. 1035 Several organisations from the global south and the World Health Organization (WHO) have 1036 developed and are further developing resuscitation guidelines tailored to low-resource settings.¹⁷⁰⁻¹⁹⁰

1037 These guidelines often focus on emergency prevention, first aid and basic life support. The ERC



- 1038 special circumstances guidelines have addressed some settings which can also have low resources, such as cardiac arrest inflight, on cruise ships and mass casualty incidents.^{113,185} An adapted approach 1039 1040 towards the chain of survival has been proposed to address the different needs and opportunities in 1041 low-resource settings¹⁷⁰ and is described in more detail in the ERC guidelines for Systems Saving 1042 Lives.⁵ 1043 1044 [h2] CPR-induced recovery of consciousness 1045 This guidance is based on an ILCOR summary statement 2024¹⁹¹, which is an update of the previous 1046 2021 summary statement¹⁹² and a scoping review from 2022, which included eight observational 1047 studies, 26 case studies, and three reviews.¹⁹³ A 2025 scoping review identified two additional observational studies,^{194,195} a case series,¹⁹⁶ and one review on prehospital guidelines to treat CPR-1048 1049 induced consciousness.¹⁹⁷ ILCOR has made the following good practice statements: 1050 In settings in which it is feasible, rescuers may consider using sedative or analgesic drugs (or 1051 both) in very small doses to prevent pain and distress to patients who are conscious during CPR. 1052 Neuromuscular-blocking drugs alone should not be given to conscious patients. 1053 The optimal drug regimen for sedation and analgesia during CPR is uncertain. Regimens can be 1054 based on those used in critically ill patients and according to local protocols). 1055 1056 CPR-induced consciousness can be defined as 'a demonstration of consciousness whilst undergoing CPR with no measurable spontaneous cardiac output displayed'.¹⁹³ The incidence of CPR-induced 1057 1058 consciousness is 0.23%-0.90% in observational studies of patients¹⁹⁸ and 48-57% of experienced healthcare professionals have reported observing patients with CPR-induced consciousness.¹⁹⁹ CPR-1059 1060 induced consciousness is associated with witnessed and shockable cardiac arrests, younger age, and 1061 better outcomes.¹⁹⁶ Longer CPR attempts in patients with CPR-induced consciousness may be reasonable.196 1062 1063 1064 Patients with CPR-induced consciousness may interfere with and prevent effective CPR^{193 196} and 1065 sedation may be required. Consider using sedatives or analgesics (or both) in very small doses to 1066 prevent pain and distress in patients who are conscious or where consciousness cannot be ruled out
 - during CPR. The optimal drug regimen for sedation and analgesia during CPR is uncertain, and local
 and national protocols exist. ¹⁹⁷ Use drug regimens based on those used in critically ill patients such
 as small doses of fentanyl, ketamine and/or midazolam. Do not give neuromuscular blocking drugs
 alone to conscious patients. Consciousness during CPR or awareness without visible consciousness



- 1071 may lead to post-traumatic stress disorder for clinicians, bystanders, and cardiac arrest
- 1072 survivors.^{195,200}
- 1073

1074 [h2] ALS treatment algorithm

1075 Cardiac arrest is associated with either shockable rhythms (VF/pVT) or non-shockable rhythms 1076 (asystole and PEA). The main difference in the treatment of shockable rhythms is the need for 1077 attempted defibrillation. Other interventions, including high-quality chest compressions with 1078 minimal interruption, airway management and effective ventilation, venous access, administration of 1079 adrenaline and the identification and treatment of reversible causes, are common for all arrests. The 1080 ALS algorithm (Figure 3) provides an overview of these key interventions. These are based on the 1081 expert consensus of the ERC ALS Writing Group. The ALS cardiac arrest algorithm is applicable to all 1082 adult cardiac arrests. Additional interventions may be indicated for cardiac arrest caused by special 1083 circumstances.^{113,201} To improve understanding of timing of shocks and drugs we have provided some 1084 example flow charts - these examples do not cover every possible scenario or transition between 1085 cardiac arrest rhythms or ROSC (Figure 4).

1086

Figure 3. Advanced Life Support algorithm ABCDE airway, breathing, circulation, disability, exposure
 CPR cardiopulmonary resuscitation; ECG electrocardiogram; EMS emergency medical system; io

1089 intraosseous; IV intravenous; PEA pulseless electrical activity; PaCO₂ arterial partial pressure of

- 1090 carbon dioxide; ROSC return of spontaneous circulation; SpO₂ oxygen saturation measured with
- 1091 pulse oximetry; VF ventricular fibrillation; VT ventricular tachycardia.







1093

1094 **Figure 4. Advanced Life Support** – example scenarios for timing of shocks and drugs [not every

1095 **possible scenario is included]** PEA pulseless electrical activity; VF ventricular fibrillation; VT

1096 ventricular tachycardia.



1102



1103 [h3] Automated External Defibrillation versus manual defibrillation during ALS

1104 This guidance is based on an ILCOR Evidence update from 2020²⁰² and a scoping review from January

- 1105 2020 to January 2025 which identified three systematic reviews, ²⁰³⁻²⁰⁵ and two observational
- 1106 studies.^{206,207}
- 1107

1108 ALS providers require frequent training and advanced ECG recognition skills for manual defibrillation. 1109 ²⁰² They should preferentially use manual defibrillation but also be skilled in AED use. Despite the 1110 faster time to first shock with AEDs, studies suggest that this does not translate into improved 1111 survival when used instead of manual defibrillators. For instance, a study in paramedics reported 1112 that while AEDs improved the time to first shock within 2 minutes (aOR 1.72; 95% Cl 1.32–2.26; 1113 P<0.001), they were associated with a reduction in survival to hospital discharge (aOR 0.71; 95% CI 0.55–0.92; P=0.009), event survival (aOR 0.74; 95% CI 0.62–0.88; P=0.001), and prehospital ROSC 1114 1115 (aOR 0.81; 95% CI, 0.68–0.96; P=0.01) compared with manual defibrillation.²⁰⁸ Manual defibrillation 1116 has been associated with shorter pauses in chest compressions, which is critical for maintaining coronary and cerebral perfusion during resuscitation.²⁰⁹ Software that filters chest compression 1117 1118 artefacts and enables rhythm analysis during CPR and performing chest compressions while charging 1119 for defibrillation also provides an advantage for manual defibrillation over use of AEDs. 1120 1121 In two-tier systems, ALS providers may arrive after an AED has already been attached to the patient 1122 and is in use. If an AED is already in use when ALS providers arrive, they should follow its shock 1123 prompts. When possible, they should transition to a manual defibrillator during a 2-minute CPR 1124 cycle. There is insufficient evidence to recommend the placement of an additional manual 1125 defibrillator in a vector-changing position when an AED is already in place.²¹⁰

1126

1127Compared with following AED prompts, manual defibrillation may result in more inappropriate1128defibrillation shocks (because a shock would not have been advised by an AED), and more missed1129shocks (because a shock would have been advised by an AED).²¹¹ A shock should be given if the ALS1130provider is in doubt whether fine VF or asystole are displayed on the monitor. In recommending this1131the ERC balanced the risks of not shocking VF versus shocking a patient in asystole. If rescuers are1132not confident in making shockable versus non-shockable rhythm decisions rapidly (within 5 seconds)1133during a resuscitation attempt they should use the defibrillator in an AED mode.1134

1135 [h3] Manual defibrillation



1136 Attempted defibrillation is a vital component of CPR as it has the potential to terminate VF/pVT and 1137 achieve ROSC. Although defibrillation is indicated in approximately 20% of cardiac arrests, more than 1138 80% of those who survive present in a shockable rhythm.²¹² As defibrillation effectiveness decreases 1139 with both time and VF duration, defibrillation attempts must be timely, whilst remaining efficient 1140 and safe.²¹³ Knowledge of how to use a defibrillator (manual or AED) is key for rescuers performing 1141 ALS. Rescuers who use a manual defibrillator should aim to take less than 5 seconds to recognise a 1142 shockable cardiac arrest rhythm and make the decision to give a shock to minimise interruption to 1143 chest compressions. If the delay exceeds 5 seconds, consider resuming chest compressions, switch 1144 the device to AED mode and then allow the device to analyse immediately. 1145 Since 2015, ERC defibrillation guidelines have referred solely to biphasic energy waveforms and

- defibrillation pads. In these 2025 guidelines, we reintroduce recommendations on the use of
- defibrillation paddles which in some countries, remain in clinical use.²¹⁴ The evidence for this section
- 1148 is based on ILCOR 2020 CoSTRs, the ERC 2015 ALS Guidelines, and expert consensus.^{151,202,214}
- 1149

1150 [h4] Strategies for minimising the peri-shock pause

1151 The delay between stopping chest compressions and shock delivery (the pre-shock pause) must be 1152 kept to an absolute minimum; even a 5-10 second delay will reduce the chances of the shock being 1153 successful.²¹⁵⁻²²⁰ The pre-shock pause can be reduced to less than 5 seconds by continuing 1154 compressions during charging of the defibrillator and by having an efficient team coordinated by a 1155 leader who communicates effectively.^{221,222} The safety check to avoid rescuer contact with the 1156 patient at the moment of defibrillation should be undertaken rapidly but efficiently. The delay 1157 between shock delivery and recommencing chest compressions (the post-shock pause) is minimised 1158 by immediately resuming chest compressions after shock delivery.²²³ If there are both clinical and 1159 physiological signs of ROSC (e.g., return of consciousness, movement, arterial waveform, increase in

- 1160 ETCO₂), chest compressions can be paused briefly for rhythm analysis. The entire process of manual
- 1161 defibrillation should be achievable with less than a 5 second interruption to chest compressions.
- 1162

1163 [h4] CPR versus defibrillation as the initial treatment

1164 A 2020 ILCOR systematic review addressed whether a specified period (typically 1.5-3 minutes) of 1165 chest compressions before shock delivery compared with a short period of chest compressions (to

- 1166 enable defibrillator start-up) before shock delivery affected resuscitation outcomes. Outcomes were
- 1167 no different when CPR was provided for up to 180 seconds before attempted defibrillation,
- 1168 compared with rhythm analysis and attempted defibrillation first.¹⁵¹ Therefore, the routine delivery
- of a pre-specified period of CPR (e.g., 2-3 minutes) before rhythm analysis and shock delivery is not



- 1170 recommended. ILCOR made a weak recommend based on low certainty evidence for rescuers
- providing a short period of CPR until the defibrillator is ready for rhythm analysis in unmonitored
- 1172 cardiac arrest. Defibrillation should then be delivered as indicated, without delay. Immediate
- 1173 defibrillation of VF of any amplitude should be attempted at the end of each 2-minute cycle.
- 1174

1175 [h4] Anticipatory defibrillator charging

- 1176 Using this method, the defibrillator is charged as the end of a compression cycle is approached, but 1177 before the rhythm is checked. When compressions are paused briefly to check the rhythm, a shock 1178 can be delivered immediately (if indicated) from a defibrillator that is already charged, thereby 1179 avoiding a period of further chest compressions while the defibrillator is being charged. This method 1180 was reviewed by ILCOR in 2020 as the technique is already in use in some countries as an alternative to the conventional sequence.²²⁴ Manikin studies show anticipatory charging is feasible, and can 1181 1182 reduce the overall number of interruptions to chest compression, but can increase pre-, post, and 1183 peri-shock pause duration. This technique may be a reasonable alternative for use by well-drilled
- 1184 teams that can minimise pre- post, and peri-shock pause duration. Clinical studies are required to
- 1185 determine the best technique for manual defibrillation.
- 1186

1187 [h3] Safe use of oxygen during defibrillation

In an oxygen-enriched atmosphere, sparking from poorly applied defibrillator paddles can cause a fire and significant burns to the patient.²²⁵⁻²³⁰ Although defibrillation pads may be safer than paddles with regards to arcing and spark generation, recommendations for the safe use of oxygen during defibrillation remain unchanged in these guidelines. The risk of fire during attempted defibrillation

- 1192 can be minimised by taking the following precautions:
- Take off any oxygen mask or nasal cannulae and place them at least 1 m away from the patient's
 chest.
- Leave the ventilation bag or ventilation circuit connected to the tracheal tube or supraglottic
 airway, any oxygen exhaust is directed away from the chest.
- If the patient is connected to a ventilator, for example in the operating room or critical care unit,
 leave the ventilator tubing (breathing circuit) connected to the tracheal tube.
- 1199

1200 [h3] Pad contact with the chest and anatomical position

- 1201 A 2024 ILCOR systematic review found no RCTs since the 2021 guidelines regarding optimal
- 1202 defibrillation pad position and evidence from two observational studies in OHCA patients was rated
- 1203 as very low certainty.¹⁹¹



- 1204 The techniques described below aim to place external defibrillation pads (self-adhesive pads) in an 1205 optimal position to maximise transmyocardial current density and minimise transthoracic 1206 impedance. No human studies have evaluated the pad position as a determinant of ROSC or survival 1207 from VF/pVT.¹⁵¹ Transmyocardial current during defibrillation is likely to be maximal when pads are 1208 placed so that the area of the heart that is fibrillating lies directly between them (i.e. ventricles in 1209 VF/pVT, atria in AF). Therefore, the optimal pad position may not be the same for ventricular and 1210 atrial arrhythmias. The antero-lateral pad position is preferred as the initial pad position for VF/VT 1211 because it is easier to place and avoids interruptions to CPR while the posterior pad is positioned.
- 1212

1213 [h4] Pad placement for ventricular arrhythmias and cardiac arrest

- 1214 In adults, place defibrillator pads or paddles in the anterior-lateral position to optimise placement
- 1215 speed and minimise interruptions to chest compressions. One pad/paddle should be positioned
- below the patient's right clavicle, just to the right of the upper sternal border. The other pad/paddle
- 1217 should be placed on the patient's left mid-axillary line, below the armpit (Figure 5).
- 1218

1219 Figure 5. Correct antero-lateral pad placement for defibrillation



- 1220
- 1221

1222 In adults, if the initial antero-lateral position is not feasible, consider using the anterior-posterior pad 1223 position if trained (Figure 6). Place the anterior pad on the left side of the chest, between the midline 1224 and the nipple. For female patients, place the anterior pad to the left of the lower sternum, ensuring 1225 it avoids breast tissue as much as possible.²³¹ The posterior pad should be placed on the left side of 1226 the patient's spine, just below the scapula.

1227



1228 Figure 6. Antero-posterior pad placement





1249 Atrial fibrillation is usually maintained by functional re-entry circuits in the left atrium. As the left 1250 atrium is located posteriorly in the thorax, pad positions that result in a more posterior current 1251 pathway may theoretically be more effective for atrial arrhythmias. Although some studies have 1252 shown that antero-posterior pad placement is more effective than the traditional antero-apical 1253 position in elective cardioversion of atrial fibrillation^{234,235}, the majority have failed to show any clear advantage of any specific pad position.²³⁶⁻²³⁹ Efficacy of cardioversion may be less dependent on pad 1254 1255 position when using biphasic impedance-compensated waveforms.²³⁸⁻²⁴⁰ The following pad positions 1256 are safe and effective for cardioversion of atrial arrhythmias: 1257 Traditional sternal-apical position. 1258 Anterior-posterior position (one pad anteriorly, over the left precordium, and the other pad 1259 posteriorly to the heart, just below the left scapula). 1260 1261 [h4] Pad placement to avoid implantable medical devices 1262 More patients are presenting with implantable medical devices (e.g. permanent pacemaker,

1263 implantable cardioverter defibrillator (ICD)). Medic Alert bracelets are recommended for these

1264 patients. Implantable medical devices may be damaged during defibrillation if current is discharged

1265 through pads placed directly over the device.^{241,242} Place the pad away from the device (at least 8 cm)

1266 or use an alternative pad position (anterior-lateral, anterior-posterior).^{241,243,244}

1267

1268 [h3] Hands-on defibrillation

1269 By allowing continuous chest compressions during the delivery of the defibrillation shock, hands-on 1270 defibrillation would minimise peri-shock pause and allow continuation of chest compressions during 1271 defibrillation. However, the benefits of this approach are unproven, and further studies are required 1272 to assess the safety and efficacy of this technique. A post-hoc analysis of a multi-centre trial did not 1273 observe any benefit when shocks were delivered without pausing manual or mechanical chest 1274 compressions.²⁴⁴ Only Class 1 electrical safety gloves, but not standard clinical gloves (or bare hands), 1275 provide a safe level of electrical insulation for hands-on defibrillation.²⁴⁵ There have been no new 1276 studies since the 2021 guidelines and the ERC recommendation 2025 therefore remains unchanged.59,214 1277

1278

1279 [h3] Respiratory phase

1280 Positive end expiratory pressure (PEEP) increases transthoracic impedance and should be minimised

1281 where possible during defibrillation. Auto-PEEP (gas trapping) may be particularly high in patients

1282 with asthma and may necessitate higher than usual energy values for defibrillation.²⁴⁶



1283

1284 [h3] One shock versus three stacked shock sequence

1285 In 2010, it was recommended that when defibrillation was required, a single shock should be 1286 provided with immediate resumption of chest compressions after the shock.²⁴⁷ This recommendation 1287 was made for two reasons. Firstly, to minimise peri-shock interruptions to chest compressions and 1288 secondly, given the greater efficacy of biphasic shocks, if a biphasic shock failed to defibrillate, a 1289 further period of chest compressions could be beneficial. Studies have not shown that any specific 1290 shock strategy is of benefit for any survival end-point.^{248,249} There is no conclusive evidence that a 1291 single-shock strategy is of benefit for ROSC or recurrence of VF compared with three stacked shocks, 1292 but given the evidence suggesting that outcome is improved by minimising interruptions to chest 1293 compressions, the ERC continued in 2020 to recommend single shocks for most situations. 1294 1295 When defibrillation is warranted, give a single shock and resume chest compressions immediately 1296 following the shock.¹⁵¹ Do not delay CPR for rhythm reanalysis or a pulse check immediately after a 1297 shock. Continue CPR for 2 min until rhythm reanalysis is undertaken and another shock given (if 1298 indicated). Even if the defibrillation attempt is successful, it takes time until the post shock 1299 circulation is established and it is very rare for a pulse to be palpable immediately after 1300 defibrillation.^{250,251} Patients can remain pulseless for over 2 min and the duration of asystole before ROSC can be longer than 2 min in as many as 25% of successful shocks.²⁵² In patients where 1301 1302 defibrillation achieves a perfusing rhythm, the effect of chest compressions on re-inducing VF is not clear.253 1303 1304 1305 [h4] Monitored and witnessed shockable cardiac arrest

If a patient has a monitored and witnessed cardiac arrest (e.g. in the catheter laboratory, coronary
care unit, or other monitored critical care setting in or out-of-hospital) and a manual defibrillator is
rapidly available:

- Confirm cardiac arrest and shout for help.
- If the initial rhythm is VF/pVT, give up to three quick successive (stacked) shocks.
- Rapidly check for a rhythm change and, if appropriate, ROSC after each defibrillation
 attempt.
- Start chest compressions and continue CPR for 2 min if the third shock is unsuccessful.
- 1314 This three-shock strategy may also be considered for an initial, witnessed VF/pVT cardiac arrest if the
- patient is already connected to a manual defibrillator. Although there are no data supporting a three-
- 1316 shock strategy in any of these circumstances, it is unlikely that chest compressions will improve the



- already very high chance of ROSC when defibrillation occurs early in the electrical phase²⁵⁴,
- 1318 immediately after onset of VF/pVT. For giving drugs after three stacked shocks:
- Give the first dose of amiodarone (or lidocaine) after the third shock if VF/pVT persists.
- Consider the stacked shocks as the first shock in the ALS algorithm for the purposes of
 adrenaline dosing.
- 1322

1323 [h3] Waveforms

- 1324 Biphasic waveforms are now well established as a safe and effective waveform for defibrillation.
- 1325 Biphasic defibrillators compensate for the wide variations in transthoracic impedance by
- electronically adjusting the waveform magnitude and duration to ensure optimal current delivery to
- 1327 the myocardium, irrespective of the patient's size (impedance compensation). There are two main
- 1328 types of biphasic waveform: the biphasic truncated exponential (BTE) and rectilinear biphasic (RLB).
- 1329 A pulsed biphasic waveform is also in clinical use, in which the current rapidly oscillates between
- 1330 baseline and a positive value before inverting in a negative pattern.³²
- 1331

1332 [h3] Energy levels

Defibrillation requires the delivery of sufficient electrical energy to defibrillate a critical mass of myocardium, abolish the wavefronts of VF and enable restoration of spontaneous synchronised electrical activity in the form of an organised rhythm. The optimal energy for defibrillation is that which achieves defibrillation whilst causing the minimum of myocardial damage.²⁵⁵ Selection of an appropriate energy level also reduces the number of repetitive shocks, which in turn limits

- 1338 myocardial damage.²⁵⁶
- 1339

1340 Optimal energy levels for defibrillation are unknown. The recommendations for energy levels are 1341 based on a consensus following careful review of the current literature. Although delivered energy 1342 levels are selected for defibrillation, it is the transmyocardial current that achieves defibrillation; the electrical current correlates well with successful defibrillation and cardioversion.²⁵⁷ A recent large 1343 1344 database study revealed no significant association between initial energy level and outcomes, such 1345 as ROSC or survival to hospital discharge.²⁵⁸ Defibrillation shock energy levels for the common BTE 1346 and RLB waveforms are unchanged from the 2021 guidelines, with pulsed biphasic energy levels increased to a minimum to 130 J to reflect the clinical evidence.^{59,259} Studies of defibrillation energy 1347 1348 levels in obese patients have generally found that defibrillation efficacy for any given energy level is 1349 unchanged.²⁶⁰



1350

1351 **[h4]** First shock

1352 Few studies have been published with which to refine the current defibrillation energy levels set in 1353 the 2010 guidelines.²⁶¹ There is no evidence that one biphasic waveform or device is more effective 1354 than another. First shock efficacy of the BTE waveform using 150-200 J has been reported as 86-1355 100%.²⁶²⁻²⁶⁷ First shock efficacy of the RLB waveform using 120 J has been reported as 85%.²⁶⁸ Four 1356 studies have suggested equivalence with lower and higher starting energy biphasic defibrillation (BTE waveform)²⁶⁹⁻²⁷² although one has suggested that initial low energy (150 J) defibrillation is associated 1357 1358 with better survival.²⁷³ Although human studies have not shown harm (raised biomarkers, ECG changes, ejection fraction) from any biphasic waveform up to 360 J^{269,274}, several animal studies have 1359 1360 suggested the potential for harm with higher energy levels.²⁷⁵⁻²⁷⁸ 1361 1362 The initial biphasic shock should be no lower than 150 J for RLB and BTE waveforms. For pulsed 1363 biphasic waveforms, begin at 130-150 J. Ideally, the initial biphasic shock energy should be at least

1364 150 J for all biphasic waveforms to simplify energy levels across all defibrillators, particularly because 1365 the type of waveform delivered by a defibrillator is not marked. Manufacturers should display the 1366 effective waveform dose range on the face of the biphasic defibrillator. If the rescuer is unaware of 1367 the recommended energy settings of the defibrillator, for an adult, use the highest energy setting for

1368 all shocks.

1369

1370 [h4] Second and subsequent shocks

1371 The 2010 guidelines recommended either a fixed or escalating energy strategy for defibrillation. 1372 Several studies of BTE waveform show that although an escalating strategy reduces the number of 1373 shocks required to restore an organised rhythm compared with low fixed-dose biphasic defibrillation, 1374 and may be needed for successful defibrillation, 279,280 rates of ROSC or survival to hospital discharge are not significantly different between strategies.²⁶⁹⁻²⁷¹ Additionally, a rectilinear biphasic protocol 1375 1376 using a fixed low energy level showed high cardioversion rates (>90%) but a significantly lower ROSC 1377 rate for recurrent VF could not be excluded.²⁸¹ Several in-hospital studies using an escalating shock 1378 energy strategy have shown improvement in cardioversion rates (compared with fixed dose 1379 protocols) in non-arrest rhythms.²⁸²⁻²⁸⁷ 1380 In 2025, there remains no evidence to support either a fixed or escalating energy protocol. Both

1381 strategies are acceptable; however, if the first shock is not successful and the defibrillator is capable

- 1382 of delivering shocks of higher energy, it is reasonable to increase the energy for subsequent shocks.
- 1383



| 1384 | [h3] Recurrent ventricular fibrillation (refibrillation) |
|------|---|
| 1385 | Recurrence of fibrillation is usually defined as 'recurrence of VF during a documented cardiac arrest |
| 1386 | episode, occurring after initial termination of VF while the patient remains under the care of the |
| 1387 | same providers (usually out-of-hospital).' Refibrillation is common and occurs in >50% of patients |
| 1388 | following initial first-shock termination of VF. ²⁷⁹ Two studies showed termination rates of subsequent |
| 1389 | refibrillation were unchanged when using fixed 120 J or 150 J shock protocols respectively ^{281,288} , but |
| 1390 | a larger study showed termination rates of refibrillation declined when using repeated 200 J shocks, |
| 1391 | unless an increased energy level (360J) was selected. ²⁷⁹ In a retrospective analysis, conversion of VF |
| 1392 | to an organised rhythm was higher if the VF had first appeared after a perfusing rhythm, than after |
| 1393 | PEA or asystole. ²²² |
| 1394 | |
| 1395 | In view of the larger study suggesting benefit from higher subsequent energy levels for |
| 1396 | refibrillation ²⁷⁹ , the ERC recommends that if a shockable rhythm recurs after successful defibrillation, |
| 1397 | and the defibrillator can deliver shocks of a higher energy, it is reasonable to increase the energy for |
| 1398 | subsequent shocks. |
| 1399 | |
| 1400 | [h3] Refractory ventricular fibrillation |
| 1401 | Refractory VF is defined as fibrillation that persists after three or more shocks. It was initially |
| 1402 | reported as occurring in approximately 20% of patients who present in VF ²⁷⁹ , but more recent studies |
| 1403 | have suggested that the true incidence may be a low as 5%. The majority of patients cardiovert |
| 1404 | successfully, but refibrillate during the two-minute period of chest compression following the |
| 1405 | shock. ²⁸⁹ . These cases of refibrillation may therefore make up a large proportion of cases of |
| 1406 | refractory VF. Duration of VF correlates negatively with good outcome. Actively search for and |
| 1407 | correct any reversible causes (Figure 3. ALS algorithm). Ensure that the defibrillation energy output is |
| 1408 | on the maximum setting. Check that the defibrillation pads are placed correctly (particularly the |
| 1409 | apical [lateral] pad, when using the antero-lateral pad position). Consider using an alternative |
| 1410 | defibrillation pad position (e.g., antero-posterior) – this is unchanged from previous guidance but has |
| 1411 | been given increased prominence. |
| 1412 | |
| 1413 | [h4] Dual/double sequential defibrillation (DSD) |
| 1414 | Patients in refractory VF have significantly lower rates of survival than patients who respond to |
| 1415 | standard resuscitation treatments. Double sequential defibrillation (DSD) is the use of two |
| 1416 | defibrillators to deliver two overlapping shocks or two rapid sequential shocks, one with standard |

1417 pad placement and the other with either anterior-posterior or additional antero-lateral pad



1418 placement. The technique has been suggested as a possible means of increasing VF termination rates. With numerous case reports and some observational studies²⁹⁰⁻²⁹⁷, in 2020 ILCOR reviewed the 1419 1420 efficacy of this technique and based on very low certainty evidence made a weak recommendation 1421 against the routine use of a DSD strategy in comparison with standard defibrillation strategy for 1422 cardiac arrest with a refractory shockable rhythm.^{83,223} More recently, ILCOR again reviewed the evidence¹ following the publication of a RCT of OHCA incorporating DSD. ²¹⁰ The updated ILCOR 1423 1424 recommendation suggested that a DSD (weak recommendation, low certainty of evidence) or a 1425 vector change defibrillation strategy (weak recommendation, very low certainty of evidence) may be 1426 considered for adults with cardiac arrest who remain in VF/pVT after three or more consecutive 1427 shocks. However, in view of the practical challenges of delivering DSD using two defibrillators and the 1428 limited evidence for its efficacy, the ERC guidelines do not recommend its introduction into routine

1429 practice.

1430

1431 [h4] Manual pressure augmentation

1432 The application of manual pressure on defibrillation pads to reduce impedance and thereby increase

1433 defibrillation shock success has been studied. There is evidence that this technique may be helpful

1434 for treating resistant cardioversion of atrial fibrillation.²⁹⁸ However, the safety and efficacy of manual

1435 pressure augmentation has not been studied in the context of shockable OHCA.

1436

1437 [h4] Percutaneous stellate ganglion block in refractory electrical storm

Percutaneous stellate ganglion block (PSGB) is an emerging, minimally invasive strategy for electrical
 storm treatment, with growing evidence supporting its efficacy and safety.²⁹⁹ A large, prospective,
 multicentre study provides evidence in favour of the effectiveness and safety of PSGB for treating

refractory electrical storm.³⁰⁰ A secondary analysis of this study, including 14 patients who were

treated with PSGB during IHCA due to refractory or recurrent VF, found no major complications; 11

1443 (78%) patients survived for at least 24 hours, and 7 (50%) were discharged with favourable

1444 neurological outcome.³⁰⁰ However, data on its use in patients experiencing refractory cardiac arrest is

1445 very limited, and larger studies are needed before its introduction into routine practice.

1446

1447 **[h3]** Analysis of rhythm during chest compression

1448 New software technology in some defibrillators enables the removal of ECG motion artefact

generated during chest compressions to show the real-time underlying waveform during CPR. An

- 1450 ILCOR systematic review found no studies in humans evaluating this technology, leading to a weak
- recommendation based on very low certainty evidence to suggest against the routine use of artefact-



- 1452 filtering algorithms for analysis of electrocardiographic rhythm during CPR.¹⁵¹ In making its
- recommendation ILCOR placed a priority on avoiding the costs of a new technology where
- 1454 effectiveness remains to be determined. The ILCOR task force acknowledged that some EMS already
- 1455 use artefact-filtering algorithms for rhythm analysis during chest compressions, and strongly
- 1456 encouraged EMS to report their experience to build the evidence base regarding these technologies
- 1457 in clinical practice.
- 1458

1459 [h4] Implantable cardioverter defibrillators

- 1460 Implantable cardioverter defibrillators (ICDs) are becoming increasingly common as they are
- 1461 implanted more frequently in an aging population. They are implanted because a patient is at risk
- 1462 from, or has had, a life-threatening shockable arrhythmia. They are usually embedded under the
- 1463 pectoral muscle below the left clavicle (in a similar position to pacemakers, from which they cannot
- 1464 be immediately distinguished). More recently, extravascular devices can be implanted
- subcutaneously in the left chest wall, with a lead running parallel to the left of the sternum.³⁰¹ In a
- 1466 recent RCT, the subcutaneous ICD (S-ICD) was non-inferior to the transvenous ICD with respect to
- 1467 device-related complications and inappropriate shocks.³⁰²
- 1468

On sensing a shockable rhythm, an ICD will discharge approximately 40 J (approximately 80J for
subcutaneous devices) through an internal pacing wire embedded in the right ventricle. On detecting
VF/pVT, ICD devices will discharge no more than eight times but may reset if they detect a new
period of VF/pVT. Patients with fractured ICD leads may suffer repeated internal defibrillation as the
electrical noise is mistaken for a shockable rhythm. In these circumstances, the patient is likely to be
conscious, with the ECG showing a relatively normal rate. A magnet placed over the ICD will turn off
the defibrillation function.²⁴³

1476

Discharge of an ICD may cause pectoral muscle contraction in the patient, and shocks to the rescuer
have been documented.³⁰³ Given the low energy values discharged by conventional ICDs, it is unlikely
that any harm will come to the rescuer, but minimising contact with the patient whilst the device is
discharging is prudent. Surface current from subcutaneous ICDs is significant and may cause a
perceptible shock to the rescuer.^{304,305} Cardioverter and pacing function should always be reevaluated following external defibrillation, both to check the device itself and to check
pacing/defibrillation thresholds of the device leads.

1484



- 1485 Pacemaker spikes generated by devices programmed to unipolar pacing may confuse AED software
- 1486 and emergency personnel and may prevent the detection of VF.³⁰⁶ The diagnostic algorithms of
- 1487 modern AEDs can be insensitive to such spikes.
- 1488

1489 [h4] Ultraportable automated external defibrillators

- 1490 A recent ILCOR scoping review found no evidence of ultraportable AED device performance, clinical
- 1491 or safety outcomes, particularly in relation to their novel low-energy waveforms.³⁰⁷ ILCOR
- 1492 recommends that the safety and efficacy of these devices are established prior to clinical
- 1493 introduction and this recommendation is supported by the ERC.
- 1494

1495 [h3]Patients with actively discharging implantable cardioverter defibrillators

- 1496 Patients considered at risk of cardiovascular collapse from malignant arrhythmias may have a pre-
- 1497 emptive ICD placed. There are two main types:
- a) Transvenous (standard) ICD: Conventional placement beneath the left pectoral muscle, with a lead
- 1499 inserted into the right ventricle (Figure 7).
- 1500
- 1501 Figure 7. Position of a transvenous (standard) Implantable Cardioverter Defibrillator (ICD)
- 1502



- 1503
- 1504
- 1505 b) Subcutaneous ICD: Placed beneath the skin in the left lateral chest wall with a lead placed
- 1506 subcutaneously along the left sternal border (Figure 8). Typically delivers 40J on discharge.
- 1507
- 1508 Figure 8. Position of a Subcutaneous Implantable Cardioverter Defibrillator





1509 1510

1511 If either device senses a shockable rhythm, the ICD will deliver a defibrillation shock between the ICD

and the distal section of the wire. Externally, this is often not apparent with a standard ICD, but a S-

1513 ICD depolarises more chest muscle to make the shock visible externally.

- 1514 Most ICDs deliver up to 6–8 shocks per episode before stopping. During this time, the shock intervals
- are approximately 5-20 seconds (depending on the internal programming). A lead fracture may cause
- 1516 intermittent erroneous sensing, resulting in resetting of the discharge programme and near-
- 1517 continuous internal defibrillation. Because the S-ICD does not have intracardiac leads, it requires a
- 1518 higher energy output to ensure effective defibrillation. Rescuers may sense a significant shock across
- 1519 their arms while performing external chest compressions, even when wearing clinical gloves. If an
- 1520 ICD/S-ICD fails to terminate a shockable rhythm, conventional external shocks should be delivered,
- 1521 placing any defibrillation pad/paddle >8 cm from the defibrillator box (as above).
- 1522

1523 [h3] Waveform analysis during CPR

1524 It is possible to predict, with varying reliability, the success of defibrillation from the fibrillation waveform.^{219,308-329} If optimal defibrillation waveforms and the optimal timing of shock delivery can 1525 1526 be determined in prospective studies, it should be possible to prevent the delivery of unsuccessful high energy shocks and minimise myocardial injury. Since the 2021 guidelines, ⁵⁹ one small RCT, 10 1527 observational studies, one systematic review and one narrative review were identified.^{204,330-341} Most 1528 1529 of the studies identified were retrospective and observational, assessing the ability of VF waveform 1530 analysis to predict defibrillation success or ROSC. The only RCT showed the prospective real-time use 1531 of VF waveform analysis using Amplitude Spectrum Area (AMSA) analysis during CPR.³³⁰ Amplitude 1532 Spectrum Area is the most studied parameter (9/10 studies) and showed the highest accuracy to predict defibrillation success^{300,331-333,336,337}, although its predictivity was inferior to a machine 1533



1534 learning algorithm during chest compression.³³⁷ The small RCT that compared AMSA-guided CPR 1535 with standard CPR showed no evidence of improvement in termination of VF (primary outcome), 1536 ROSC or in long-term survival.³³⁰ Recent developments include the use of convolutional neural 1537 networks to calculate AMSA during continuous CPR.³³⁹ Additionally, information other than the 1538 outcome of an immediate defibrillation can be obtained from the waveform analysis. For instance, 1539 two retrospective observational studies assessed the ability of AMSA to detect coronary occlusion, 1540 showing lower AMSA values in case of acute myocardial infarction.³⁴⁰ A recent ILCOR systematic 1541 review update³⁴² found two observational studies focusing on software-based cardiac rhythm 1542 analysis during CPR in OHCA patients.^{343,344} Both studies were observational and used historical 1543 controls, showing improved CPR quality metrics with rhythm analysis during compressions. None of 1544 the studies evaluated survival rates or neurological outcomes, leaving it uncertain whether these 1545 technologies enhance patient outcomes. ILCOR found insufficient evidence to make a treatment 1546 recommendation, given the absence of RCTs or well-controlled observational studies. Consistent 1547 with previous recommendations, there remains insufficient evidence to support routine use of VF 1548 waveform analysis to guide the optimal timing for a shock attempt.

1549

1550 [h2]Airway and ventilation

1551 Since 2015 the ERC has recommended a stepwise approach to airway management during CPR.²¹⁴ Three large RCTs of airway management for OHCA have been published since 2015 that have 1552 supported this guidance.³⁴⁵⁻³⁴⁷ An ILCOR systematic review addressed whether a specific advanced 1553 1554 airway management strategy (use of supraglottic airway devices (SGA) or tracheal intubation) 1555 improved outcome from cardiac arrest in comparison with an alternative airway management 1556 strategy.^{348,349} Seventy-eight observational studies were included; nine of these addressed the timing 1557 of advanced airway management. Eleven controlled trials were included but only three of these were 1558 RCTs.³⁴⁹ The first of these RCTs compared early tracheal intubation with bag-mask ventilation 1559 (tracheal intubation delayed until after ROSC) in a physician-staffed EMS system.³⁴⁶ The result of this 1560 non-inferiority trial that recruited over 2,000 patients was inconclusive – 4.3% versus 4.2% for 28-day 1561 survival with favourable functional outcome (Cerebral Performance Category [CPC] 1-2), no 1562 significant difference. Notably, the tracheal intubation success rate was 98% and 146 patients in the 1563 bag-mask ventilation group underwent 'rescue intubation' (i.e., crossed over); 100 of these were 1564 because of regurgitation. In a comparison of initial laryngeal tube insertion with tracheal intubation 1565 in 3,000 OHCAs by paramedics in the United States, 72-h survival (primary outcome) was higher in 1566 the laryngeal tube group (18.2% versus 15.3%; p =0.04).³⁴⁷ However, the overall tracheal intubation 1567 success rate was just 51% making it possible that the lower survival rate in the tracheal intubation



1568group was a reflection of the poor tracheal intubation success rate. The third of these RCTs was a1569comparison of the initial insertion of an i-gel supraglottic airway (SGA) with tracheal intubation in1570OHCA treated by paramedics in the United Kingdom (UK).345 Among the more than 9,000 patients1571enrolled, there was no difference in the primary outcome of favourable functional survival (modified

- 1572 Rankin Scale [mRS] ≤ 3; 6.4% versus 6.8%; P =0.33).
- 1573

A cluster RCT in Taiwan compared insertion of i-gel with tracheal intubation in 936 OHCA patients and found no difference in the rate of sustained ROSC, the primary outcome.³⁵⁰ Three observational studies have compared the use of the laryngeal tube with i-gel in OHCA. Rates of successful airway placement and survival to hospital discharge were higher with the i-gel than the laryngeal tube.³⁵¹⁻³⁵³

1579 A large observational cohort study of IHCA from the American Heart Association (AHA) Get with the 1580 Guidelines-Resuscitation (GWTG-R) registry, compared outcomes in patients intubated at any given 1581 minute within the first 15 min after cardiac arrest onset against matched patients still receiving CPR at risk of being intubated within the same minute.³⁵⁴ The matching was based on time-dependent 1582 1583 propensity scores, pairing 43,314 intubated patients and non-intubated patients with similar 1584 propensity for intubation. Compared with not intubating, tracheal intubation was associated with a 1585 lower rate of ROSC (risk ratio [RR] 0.97; 95% CI 0.96-0.99; p < 0.001), lower survival to hospital 1586 discharge (RR 0.84; 95% CI 0.81-0.87; p < 0.001), and worse neurological outcome (RR 0.78; 95% CI 1587 0.75-0.81; p < 0.001). Two ongoing RCTs are comparing insertion of an i-gel with tracheal intubation in IHCA.355,356 1588

1589

After reviewing the evidence for airway management during cardiac arrest, the ILCOR ALS Task Force made the following treatment recommendations³⁴⁹, which the ERC has adopted, and these remain unchanged following an evidence update in 2025¹:

We suggest using bag-mask ventilation or an advanced airway strategy during CPR for adult
 cardiac arrest in any setting (weak recommendation, low to moderate certainty of evidence).

- 1595 If an advanced airway is used, we suggest a SGA for adults with OHCA in settings with a low
 1596 tracheal intubation success rate (weak recommendation, low certainty of evidence).
- If an advanced airway is used, we suggest an SGA or tracheal intubation for adults with OHCA in
 settings with a high tracheal intubation success rate (weak recommendation, very low certainty
 of evidence).
- If an advanced airway is used, we suggest an SGA or tracheal intubation for adults with IHCA
 (weak recommendation, very low certainty of evidence).



1602

- 1603 Patients often have more than one type of airway intervention, typically starting with basic and 1604 advancing to more complex techniques that are inevitably applied later during cardiac arrest - the 1605 stepwise approach.^{345,357} The best airway, or combination of airway techniques will vary according to 1606 patient factors, the phase of the resuscitation attempt (during CPR, after ROSC), and the skills of 1607 rescuers. If basic airway techniques enable effective ventilation, there may be no need to progress to 1608 advanced techniques until after ROSC. One potential advantage of inserting an advanced airway is 1609 that it enables chest compressions to be delivered continuously without pausing during ventilation. 1610 Most patients with ROSC remain comatose and will need tracheal intubation and mechanical 1611 ventilation (See Post-resuscitation Care).¹⁴⁶ 1612
- 1613 **[h3]** Airway obstruction
- 1614 Patients requiring CPR often have an obstructed airway, usually secondary to loss of consciousness,
- 1615 but occasionally it may be the primary cause of cardiorespiratory arrest.
- 1616

1617 [h4] Basic airway management and adjuncts

- 1618 Three manoeuvres may improve the patency of an airway obstructed by the tongue or other upper 1619 airway structures: head tilt, chin lift, and jaw thrust. Despite a total lack of published data on the use 1620 of nasopharyngeal and oropharyngeal airways during CPR, they are often helpful, and sometimes
- 1621 essential, to maintain an open airway, particularly when CPR is prolonged.
- 1622

1623 [h3] Oxygen during CPR

- 1624 During cardiac arrest, the blood flow and oxygen reaching the brain are low, even with effective CPR.
- 1625 Based on the physiological rationale and expert opinion, ILCOR and ERC recommends giving the
- 1626 highest feasible inspired oxygen concentration during cardiac arrest to maximise oxygen delivery to
- 1627 the brain thereby minimising hypoxic-ischaemic injury.²⁰² Observational studies have shown that a
- $1628 \qquad \text{higher PaO}_2 \, \text{during CPR is associated with a higher likelihood of ROSC and patient survival.}^{358,359} \, \text{After}$
- $1629 \qquad \text{ROSC, as soon as SpO}_2 \text{ can be measured reliably or arterial blood gas values are obtained, titrate the}$
- 1630 inspired oxygen to achieve an arterial oxygen saturation of 94-98% or arterial partial pressure of
- 1631 oxygen (PaO₂) of 10–13 kPa (75–100 mmHg). Rescuers should be aware that pulse oximetry can
- 1632 overestimate the true oxygen saturation in people with darker skin tones.^{360,361}
- 1633
- 1634 **[h3]** Choking



1635 The initial management of foreign body airway obstruction (choking) is addressed in the ERC 1636 Guidelines 2025 First Aid.⁶ This topic has been recently assessed by an ILCOR evidence update³⁶² and 1637 a previous systematic review.³⁶³ In an unconscious patient with suspected foreign body airway 1638 obstruction if initial basic measures are unsuccessful use laryngoscopy and forceps to remove the 1639 foreign body under direct vision. To do this effectively requires training. Some patients with foreign 1640 body airway obstruction may require emergency front of neck access and this is addressed further 1641 below.

1642

1643 [h3] Ventilation

1644 [h4] Ventilation breath volume

1645 Recent studies suggest that effective ventilation and oxygenation is often done poorly during ALS. 1646 Advanced life support providers should give artificial ventilation as soon as possible for any patient in 1647 whom spontaneous ventilation is inadequate or absent. This is usually achieved with a self-inflating 1648 bag attached to a facemask or an advanced airway. Deliver each breath over approximately 1 second, 1649 giving a volume that corresponds to normal chest movement. The chest should visibly rise; this 1650 represents a compromise between giving an adequate volume, minimizing the risk of gastric 1651 inflation, and allowing adequate time for chest compressions.

1652

1653 [h4] 30:2 versus asynchronous ventilation

1654 Although the delivery of continuous chest compressions during face-mask ventilation was previously 1655 thought to increase the risk of regurgitation, a trial of continuous versus interrupted chest 1656 compressions during CPR (CCC Trial) that enrolled more than 23,000 patients showed no statistically 1657 significant difference in survival to discharge.³⁶⁴ ILCOR has subsequently recommended that when 1658 using bag mask, EMS providers perform CPR either using a 30:2 compression-ventilation ratio 1659 (pausing chest compressions for ventilation) or continuous chest compressions without pausing 1660 while delivering positive pressure ventilation (strong recommendation, high-quality evidence).³⁶⁵ In 1661 Europe, the most common approach during CPR with an unprotected airway is to give two 1662 ventilations after each sequence of 30 chest compressions. A secondary analysis of the CCC trial 1663 examined the frequency of effective ventilations delivered with bag-mask and measured via thoracic 1664 bioimpedance during pauses in chest compressions among 1976 OHCA patients of the 30:2 arm of 1665 the trial.³⁶⁶ The study revealed that the quality of ventilation was notably poor: in 60% of the 1666 patients, lung inflation was observed in less than half of the chest compression pauses, with a 1667 median time to the first ventilation greater than 4 minutes. Patients who experienced lung inflations 1668 in at least 50% of pauses showed higher rates of ROSC (RR 1.3 [95% CIs 1.2-1.5]; p< 0.0001), survival



1669at hospital discharge (RR 2.2 [95% CIs 1.6-3.0]; p< 0.0001) and survival with favourable neurological</th>1670outcomes (RR 2.8 [95% CIs 1.8-4.3]; p< 0.0001) (1). The ERC recommends, that advanced life support</td>1671providers should ensure effective ventilation when using a bag-mask. If ventilation is inadequate,1672efforts should be made to optimise bag-mask ventilation by improving the mask seal, maintaining1673airway patency, and, when possible, employing a two-person technique.367,368

1674

1675 Once a tracheal tube or an SGA has been inserted, ventilate the lungs at a rate of 10 min⁻¹ and continue chest compressions without pausing during ventilations.³⁶⁹ The laryngeal seal achieved with 1676 1677 an SGA may not be good enough to prevent at least some gas leaking when inspiration coincides 1678 with chest compressions. Moderate gas leakage is acceptable (unless there is a significant risk of infection),^{370,371} particularly as most of this gas will pass up through the patient's mouth. If excessive 1679 1680 gas leakage results in inadequate ventilation of the patient's lungs, chest compressions will have to 1681 be interrupted to enable ventilation, using a compression - ventilation ratio of 30:2. One 1682 observational study of OHCA patients who had received prolonged CPR with a mechanical device has 1683 documented worse blood gases among those whose airway was managed with an SGA compared 1684 with tracheal intubation.³⁷²

1685

1686 [h4] Ventilation rate

1687 Since the 2021 guidelines review, one small RCT, a secondary analysis of an RCT, and three 1688 observational studies assessing ventilation rate during CPR have been identified through a systematic search strategy.³⁷³⁻³⁷⁷ A small RCT randomised 46 patients after tracheal intubation to receive either 1689 1690 10 or 20 breaths/min delivered with mechanical ventilation in volume-controlled mode with a tidal 1691 volume of 6 mL kg⁻¹.³⁷³ The study was terminated early, failing to achieve the planned sample size, 1692 and showed a higher minute ventilation in the group randomised to 20 breaths min⁻¹ (primary 1693 outcome) with no evidence of improvement in hypercapnia, hypoxia, and ROSC rates. A secondary 1694 analysis of the Pragmatic Airway Resuscitation Trial (PART) compared laryngeal tubes with tracheal 1695 intubation evaluated the association between ventilation rates and outcomes.³⁷⁴ A median 1696 ventilation rate of 8 min⁻¹ was observed in both groups, and the duration of hypoventilation (defined 1697 as $<6 \text{ min}^{-1}$) was associated with decreased rates of ROSC and hospital survival. The duration of mild 1698 hyperventilation (>12 to 16 min⁻¹) was associated with improved ROSC, survival to hospital discharge 1699 and survival with favourable neurological outcome. A prospective observational study evaluating 1700 prehospital respiratory mechanics after advanced airway placement showed a respiratory rate of 8 1701 min⁻¹ during CPR with a lower tidal volume (6 mL kg⁻¹) delivered compared with the post ROSC phase.³⁷⁵ Two retrospective observational studies failed to demonstrate an association between 1702



1703 ventilation rates and patient outcomes.^{376,377} Although there is increasing evidence in ALS settings

- 1704 showing that respiratory rate is frequently below 10 min⁻¹ during CPR and that the duration of
- 1705 hypoventilation is associated with worse outcomes, there remains insufficient evidence to establish
- 1706 an optimal ventilation rate during CPR.
- 1707
- 1708

1709 [h4] Mechanical ventilation during CPR

- 1710 Two RCTs and two observational studies comparing mechanical ventilation and bag ventilation during
- 1711 CPR were identified.³⁷⁸⁻³⁸¹ The RCTs were feasibility studies comparing mechanical ventilation
- delivered in volume-controlled mode (tidal volume 6-7 mL kg⁻¹, RR 10 min⁻¹, FiO₂ 1.0) with bag
- 1713 ventilation after advanced airway placement. Each study included 60 patients and no sample size
- 1714 calculations were performed. They showed that the use of mechanical ventilation during both
- 1715 mechanical³⁷⁸ and manual³⁷⁹ chest compressions was feasible. Neither of these trials found
- 1716 differences in oxygenation, ROSC, or survival. A prospective observational study indicated that
- 1717 mechanical ventilation is associated with reduced PaCO₂ values; however, it did not affect the rates
- 1718 of ROSC, survival, or survival with favourable neurological outcomes.³⁸⁰ The retrospective study
- 1719 compared mechanical ventilation delivered in cardiopulmonary ventilation mode (298 patients) with
- bag ventilation (2268 patients), showing that mechanical ventilation may increase the rate of ROSC
- 1721 with no effect on survival with favourable neurological outcomes.³⁸¹
- 1722 Although studies show that the use of mechanical ventilation during chest compressions is feasible,
- 1723 there is insufficient evidence to support the use of mechanical ventilation over manual bag
- ventilation or to recommend a specific ventilation mode. We suggest that when using mechanical
- ventilation in volume-controlled mode during CPR, the following settings are used: tidal volume of 6-
- 1726 8 mL kg⁻¹ (predicted body weight) or sufficient to cause a visible chest rise, maximum inspired
- 1727 oxygen, respiratory rate of 10 min⁻¹, an inspiratory time of \geq 1 second, PEEP 0-5 cm H₂O, alarm for
- 1728 peak pressure set at 60-70 cm H_2O , and the flow trigger off.
- 1729

1730 [h4] Ventilation during mechanical chest compressions

- Ventilation during mechanical chest compressions may be particularly challenging because of the
 mechanical forces applied to the chest, causing lung volume to decrease below functional residual
 capacity (FRC).^{382,383} One RCT and one retrospective observational study evaluated ventilation during
- 1734 mechanical chest compressions.^{384,385} The pilot RCT randomised 30 OHCA patients undergoing
- 1735 mechanical chest compressions after tracheal intubation into one of the following three ventilation
- 1736 strategies: 1) biphasic positive airway pressure (BIPAP) with assisted spontaneous breathing; 2)



1737 continuous positive airway pressure (CPAP) and 3) volume-controlled ventilation. This study showed 1738 a higher tidal volume delivered in BIPAP mode than in CPAP mode, while no differences in the rate of 1739 ROSC were detected. The retrospective observational study evaluated the frequency of bag-1740 ventilation during pauses in mechanical chest compressions delivered at a 30:2 ratio and showed 1741 inadequate ventilation during 3-second pauses in mechanical chest compressions – two inflations 1742 were successfully provided in 45% of compression pauses, and no ventilation was delivered in 19% of 1743 compression pauses.³⁸⁵ A particularly low likelihood of delivering two successful insufflations was 1744 detected in the first four minutes after mechanical chest compression device placement. The best 1745 method for ventilation during mechanical chest compression is uncertain. 1746

1747 [h4] Passive oxygen delivery

1748 In the presence of a patent airway, chest compressions alone may result in some ventilation of the lungs.³⁸⁶ Oxygen can be delivered passively, either via an adapted tracheal tube (Boussignac 1749 1750 tube)^{387,388}, or with the combination of an oropharyngeal airway and standard oxygen mask with a non-rebreather reservoir.³⁸⁹ In theory, a SGA can also be used to deliver oxygen passively but this has 1751 1752 yet to be studied. One study has shown higher neurologically favourable survival with passive oxygen 1753 delivery (oral airway and oxygen mask) compared with bag-mask ventilation after shockable OHCA, 1754 but this was a retrospective analysis and is subject to numerous confounders.³⁸⁹ A trial of continuous 1755 or interrupted chest compressions during CPR (CCC Trial) included a subgroup of patients who were 1756 treated with passive oxygenation but until further data are available, passive oxygen delivery without 1757 ventilation is not recommended for routine use during CPR.³⁶⁴

- 1758
- 1759 **[h3] Choice of airway devices**

1760 Disadvantages of tracheal intubation over bag-mask ventilation include:

The risk of an unrecognised misplaced tracheal tube – in patients with OHCA the reliably
 documented incidence ranges from 0.5% to 17%: emergency physicians – 0.5%³⁹⁰; paramedics –
 2.4%³⁹¹, 6%^{392,393}, 9% ³⁹⁴, and 17%³⁹⁵

- A prolonged period without chest compressions while tracheal intubation is attempted. In a
 study of prehospital tracheal intubation by paramedics during 100 cardiac arrests the median
 duration of the interruptions in CPR associated with tracheal intubation attempts was 110 s
 (interquartile range 54 198 s; range 13 446 s).³⁹⁶ Tracheal intubation attempts accounted for
 almost 25% of all CPR interruptions.
- A comparatively high failure rate. Intubation success rates correlate with the tracheal intubation
 experience attained by individual paramedics.³⁹⁷ The high failure rate of 51% documented in the



- PART trial³⁴⁷ is similar to those documented in some prehospital systems more than 20 years
 ago.^{398,399}
- Tracheal intubation is a difficult skill to acquire and maintain. In one study, anaesthesia residents
 required about 125 intubations with direct laryngoscopy in the operating room setting before
 they were able to achieve a tracheal intubation success rate of 95% under such optimal
 conditions.⁴⁰⁰
- Healthcare personnel who undertake prehospital tracheal intubation should do so only within a
 structured, monitored program, which should include comprehensive competency-based training
 and regular opportunities to refresh skills.
- 1780

The ILCOR recommendation is that only systems that achieve high tracheal intubation success rates should use this technique.³⁴⁹ ILCOR did not recommend a particular success rate but suggested it should be similar to that achieved in the RCT comparing early tracheal intubation with bag-mask ventilation (tracheal intubation delayed until after ROSC) in a physician-staffed EMS system.³⁴⁶ The tracheal intubation success rate in this study was 98%. The expert consensus of the ERC 2021 ALS Guidelines writing group was that a high success rate is greater than 95% with up to two intubation attempts.⁵⁹

1788

1789 Rescuers must weigh the risks and benefits of tracheal intubation against the need to provide 1790 effective chest compressions. To avoid any interruptions in chest compressions, unless alternative 1791 airway management techniques are ineffective, it is reasonable to defer tracheal intubation until 1792 after ROSC. In settings with personnel skilled in advanced airway management, laryngoscopy should 1793 be undertaken without stopping chest compressions; a brief pause in chest compressions will be 1794 required only as the tube is passed through the vocal cords. The tracheal intubation attempt should 1795 interrupt chest compressions for less than 5 seconds; if intubation is not achievable within these 1796 constraints, recommence bag-mask ventilation. After tracheal intubation, tube placement must be 1797 confirmed immediately (see below) and the tube must be secured adequately.

1798

1799 [h4] Video laryngoscopy

Video laryngoscopy (VL) is being used increasingly in anaesthetic and critical care practice.^{401,402}
 Preliminary studies indicate that compared with direct laryngoscopy (DL), VL during CPR improves
 laryngeal view and tracheal intubation success rates^{403,404}, reduces the risk of oesophageal
 intubation⁴⁰⁵ and reduces interruptions to chest compressions.⁴⁰⁶ One systematic review concluded
 that in the prehospital setting, VL decreased the first-attempt tracheal intubation success rate (RR,



1805 0.57; P < 0.01; high-quality evidence) and overall success rate (RR, 0.58; 95% CI, 0.48-0.69; moderatequality evidence) by experienced operators.⁴⁰⁷ A later meta-analysis of OHCAs and IHCAs that 1806 1807 included six observational studies and one RCT documented higher first pass success rate and better 1808 grade of view with VL compared with DL.⁴⁰⁸ A secondary analysis of an RCT also documented a higher 1809 rate of successful tracheal intubation on the first attempt with VL compared with DL.⁴⁰⁹ Several 1810 different video laryngoscopy systems are available and they do not all perform in the same way. The 1811 expert consensus of the writing group and recommendation of the ERC is that the rescuer's choice of 1812 direct laryngoscopy or video laryngoscopy should be guided by local protocols and rescuer 1813 experience.

1814

1815 **[h3] Confirmation of correct placement of the tracheal tube**

1816 Unrecognised oesophageal intubation is the most serious complication of attempted tracheal 1817 intubation in patients with and without cardiac arrest. One large RCT comparing bag-mask 1818 ventilation and tracheal intubation during CPR by skilled prehospital physicians reported accidental but recognised oesophageal intubation in 10% of attempts.³⁴⁶ Another study found the rate of 1819 1820 unrecognised oesophageal intubation in OHCA by paramedics was 5%.³⁹³ Therefore, accurate means 1821 to verify correct tracheal tube placement in CPR are of greatest importance. The previous evidence supporting this guideline is summarised in longstanding ILCOR recommendations.^{32,223,410} The 2022 1822 1823 PUMA Guidelines for the prevention of unrecognised oesophageal intubation state clearly that the 1824 only accurate method to confirm tracheal placement of the tracheal tube is the presence of a 1825 sustained $ETCO_2$ trace for at least seven breaths on waveform capnography in patients with and 1826 without cardiac arrest.⁴¹¹ Clinical assessment such as chest and abdominal auscultation, observation 1827 of chest expansion, and fogging of the tube cannot be used to confirm tracheal placement if 1828 waveform capnography suggests otherwise. The 'No Trace = Wrong Place' campaign by the UK Royal 1829 College of Anaesthetists emphasises that immediately after tracheal intubation (even during CA) the absence of exhaled CO₂ strongly suggests oesophageal intubation.⁴¹² If waveform capnography does 1830 1831 not confirm correct placement of the tracheal tube in patients without cardiac arrest the PUMA 1832 Guidelines suggest using repeat laryngoscopy to view the passage of the tube, use of a flexible 1833 bronchoscope or ultrasound of the neck. Portable monitors make capnographic initial confirmation 1834 and continuous monitoring of tracheal tube position feasible in both in- and out-of-hospital settings 1835 where tracheal intubation is performed. Ultrasonography of the neck or visualisation with a flexible 1836 fibreoptic scope by skilled operators can also be used to confirm the presence of a tracheal tube in 1837 the trachea.413

1838



1839 **[h3] Cricoid pressure**

- 1840 The use of cricoid pressure in cardiac arrest is not recommended (expert consensus). Cricoid
- 1841 pressure can impair ventilation, laryngoscopy, tracheal tube and SGA insertion, and may even cause
- 1842 complete airway obstruction.414
- 1843

1844 [h3] Securing the tracheal tube and supraglottic airway

- 1845 Accidental dislodgement of a tracheal tube can occur at any time but may be more likely during CPR
- 1846 and during transport. An SGA is more prone to being dislodged than a tracheal tube.³⁴⁵ The most
- 1847 effective method for securing the tracheal tube or SGA has yet to be determined. Use either
- 1848 conventional tapes or ties, or purpose-made holders.
- 1849

1850 [h3] Emergency Front of Neck Access (eFONA)

1851 In rare cases, it may be impossible to oxygenate a patient in cardiac arrest using bag-mask 1852 ventilation, or to insert a tracheal tube or supraglottic airway. This may occur in patients with

- 1852 ventilation, or to insert a tracheal tube or supraglottic airway. This may occur in patients with
- 1853 extensive facial trauma or laryngeal obstruction caused by oedema, tumour or a foreign body. In
- 1854 2023, ILCOR undertook a scoping review which summarised evidence from 69 studies across four
- 1855 domains: incidence of eFONA; success rates of eFONA attempts; clinical outcomes in patients with an
- 1856 eFONA attempt; and complications associated with eFONA attempts.⁴¹⁵ The review found that none
- 1857 of the identified studies focussed specifically on cardiac arrest, and noted that the available evidence
- 1858 was highly heterogeneous. Consistent with a systematic review on pre-hospital eFONA success rates,
- 1859 the scoping review observed that, when attempted, eFONA success rates were typically high.⁴¹⁶
- 1860 Based on the available evidence, ILCOR identified that it was not possible to undertake a systematic
- 1861 review, but made a good practice statement supporting the use of cricothyroidotomy in patients
- 1862 where standard airway management and ventilation strategies have been unsuccessful.¹ An
- 1863 Australian observational study, published in 2024, reported on 80 cricothyroidotomy attempts of
- 1864 which 56 occurred in OHCA patients.⁴¹⁷ The reported incidence was 1.1 cricothyroidotomies per
- 1865 1,000 attempted resuscitations with a success rate in cardiac arrest of 58.9%. The use of a surgical
- 1866 approach was associated with a higher success rate than a needle approach (88.2 % v. 54.6 %,
- 1867 p = 0.003). Consistent with Difficult Airway Society recommendations, the ERC recommends, where
- 1868 feasible, the use of a scalpel-bougie cricothyroidotomy.⁴¹⁸
- 1869
- 1870 [h2] Drugs and fluids
- 1871
- 1872 **[h3] Vascular access**



1873The effectiveness of drugs in cardiac arrest is time-dependent.419-421The IO route has been proposed1874as an alternative strategy for initial vascular access in adult cardiac arrest, based on the perceived1875ease of insertion and an RCT showing that the time to drug administration was fastest with a tibial IO1876strategy, compared to both a peripheral IV or humeral IO strategy.422,423Observational studies show

- 1877 that the use of the IO route in clinical practice has increased over recent years.^{424,425}
- 1878

In 2024, three large RCTs compared an IO-first with an IV-first strategy in adult cardiac arrest.⁴²⁶⁻⁴²⁸ In 1879 1880 contrast to earlier research, none of these trials found that the IO route facilitated more rapid drug 1881 administration. These three trials, which comprised 9,332 patients, formed the basis of a systematic 1882 review and meta-analysis led by the ILCOR ALS Task Force.⁴²⁹ The meta-analysis showed that an IO-1883 first strategy, compared with an IV-first strategy, did not improve the rate of survival to 30-days or 1884 discharge with favourable neurological outcome (OR 1.07; 95% CI 0.88–1.30; low-certainty evidence) 1885 and 30-day survival (OR 0.99: 95% CI 0.84–1.17; moderate-certainty evidence), but may reduce the 1886 rate of sustained ROSC (OR 0.89, 95% CI 0.80–0.99; moderate-certainty evidence). There was no 1887 evidence that the treatment effect was influenced by baseline patient characteristics or the 1888 anatomical site of the IO cannula. In formulating its treatment recommendation, ILCOR noted the 1889 routine availability of the peripheral IV route across international systems, the high cost of IO 1890 cannulae, the additional training requirement associated with intraosseous use, and the evidence 1891 from the meta-analysis suggesting that an IO-first strategy reduces the odds of achieving ROSC.¹ 1892 ILCOR noted that no RCTs had been undertaken in an IHCA but observed that the IO route is 1893 infrequently required in such cases.

1894

1895 Consistent with ILCOR, the ERC recommends that the IV route should be used as the primary route 1896 for vascular access in adult cardiac arrest. We recognise that the IO route may be a reasonable 1897 vascular access strategy where IV access cannot be rapidly achieved. Consistent with the two 1898 European RCTs, we recommend that two IV vascular access attempts are made before considering 1899 attempting IO access.^{427,428}

1900

1901There has been recent interest in the use of intramuscular (IM) adrenaline for cardiac arrest based1902on animal studies and observational data.1903study, a single dose of 5 mg IM adrenaline in 420 patients with OHCA was associated with improved1904survival to hospital admission, survival to hospital discharge, and functional survival compared with1905standard adrenaline dosing.433The role of the IM route for adrenaline in cardiac arrest needs to be1906studied with RCTs before considering its inclusion in guidelines.


1907

1908 [h3] Vasopressors

1909 ILCOR recently updated their review of vasopressors in cardiac arrest.¹ The systematic reviews and 1910 meta-analyses evaluated standard dose adrenaline (1 mg) versus placebo, high dose (5-10 mg) versus 1911 standard dose (1 mg) adrenaline, adrenaline versus vasopressin, and adrenaline and vasopressin versus adrenaline alone.^{191,434-436} The evidence showed that adrenaline (1 mg) improved ROSC, 1912 1913 survival to hospital admission, survival to hospital discharge, and long-term survival (up to 12) 1914 months), although the effect on favourable neurological outcome remains uncertain. By contrast, the 1915 use of high-dose adrenaline or vasopressin (with or without adrenaline) did not improve long-term 1916 survival or favourable neurological outcome. 1917 1918 Based on this evidence, ILCOR makes a strong recommendation in favour of adrenaline during CPR (strong recommendation, low certainty of evidence).¹⁹¹ The justification and evidence to decision 1919 1920 framework highlights that the Task Force placed a very high value on the apparent life-preserving 1921 benefit of adrenaline, even if the absolute effect size is likely to be small and the effect on survival 1922 with favourable neurological outcome is uncertain. 1923 1924 The evidence supporting adrenaline use comes primarily from two placebo-controlled trials.^{437,438}

1925 The PARAMEDIC2 trial followed the ERC ALS 2015 Guidelines, which recommended adrenaline 1926 administration as soon as vascular access was obtained for non-shockable rhythms, and after three unsuccessful defibrillation attempts for shockable rhythms.^{214,438} Analysis of this trial continues to 1927 1928 provide new evidence, with long-term follow-up data showing sustained survival benefits at 12 1929 months.⁴³⁹ Meta-analyses with the earlier PACA trial found greater effects of adrenaline on ROSC and 1930 survival to hospital discharge in initial non-shockable rhythms compared to shockable rhythms, 1931 although this difference was less pronounced for longer-term survival and favourable neurological 1932 outcomes.²¹² A secondary analysis of time to drug administration found that while relative treatment 1933 effects of adrenaline remained constant, survival and favourable neurological outcome rates 1934 decreased over time, suggesting potential benefit from early intervention.419 1935

1936 Consequently, ILCOR recommends that adrenaline is administered as soon as feasible for non-

shockable rhythms (PEA/asystole) (strong recommendation, very low certainty of evidence). For

1938 shockable rhythms (VF/pVT), ILCOR suggests administration of adrenaline after initial defibrillation

1939 attempts are unsuccessful during CPR (weak recommendation, very low certainty of evidence).

1940



1941 Consistent with the ILCOR Treatment Recommendations, the ERC recommends that adrenaline 1 mg 1942 is administered as soon as possible for adult patients in cardiac arrest with a non-shockable rhythm. 1943 For patients with a shockable rhythm persisting after three initial shocks, administer adrenaline 1 1944 mg. Repeat adrenaline 1 mg every 3-5 minutes whilst ALS continues. While practical challenges may 1945 affect the timing of adrenaline in the prehospital setting, early administration remains a priority, 1946 particularly for non-shockable rhythms where alternative interventions are limited. 1947 1948 The optimal dose of adrenaline remains unclear. Pharmacokinetic and observational data suggest 1949 further investigation is needed regarding dosing strategies.^{440,441} This includes the standard 1 mg 1950 dose, the cumulative effects of repeated doses, and potential alternative approaches such as titrated 1951 dosing in closely monitored settings. Alternative routes such as intracoronary administration, have 1952 been studied in cardiac catheterisation laboratories, but evidence is insufficient to support these approaches.442,443 1953 1954 If three stacked shocks have been given for a witnessed and monitored shockable cardiac arrest 1955 1956 where immediate defibrillation is possible, these initial three stacked shocks should be considered as 1957 the first shock with regards to timing of the first dose of adrenaline. After these stacked shocks, 1958 providers should continue resuscitation attempts and adrenaline dosing according to the standard 1959 ALS algorithm. 1960 1961 A recent cost-effectiveness analysis of PARAMEDIC2 incorporating both direct survival benefits and 1962 increased organ donation rates supports the use of adrenaline during cardiac arrest, although costs 1963 may vary between healthcare systems.444 1964 1965 Consistent with the ILCOR treatment recommendation, the ERC does not support the use of 1966 vasopressin during cardiac arrest. 1967 1968 [h3] Antiarrhythmic drugs ILCOR updated the CoSTR for antiarrhythmic drugs in 2018.⁴⁴⁵ This was followed by ILCOR evidence 1969 1970 updates in July 2023 and October 2024.^{1,191} The 2018 CoSTR made the following recommendations: 1971 We suggest the use of amiodarone or lidocaine in adults with shock refractory VF/pVT (weak 1972 recommendation, low-quality evidence).



| 1973 | • We suggest against the routine use of magnesium in adults with shock-refractory VF/pVT (weak |
|------|---|
| 1974 | recommendation, very low-quality evidence). |
| 1975 | The confidence in effect estimates is currently too low to support an ALS Task Force |
| 1976 | recommendation about the use of bretylium, nifekalant, or sotalol in the treatment of adults in |
| 1977 | cardiac arrest with shock-refractory VF/pVT. |
| 1978 | The confidence in effect estimates is currently too low to support an ALS Task Force |
| 1979 | recommendation about the use of prophylactic antiarrhythmic drugs immediately after ROSC in |
| 1980 | adults with VF/pVT cardiac arrest. |
| 1981 | |
| 1982 | Despite many studies, the ILCOR evidence updates in 2023 did not identify any compelling new data |
| 1983 | that would justify a further systematic review or changes to the treatment recommendations. ¹⁹¹ The |
| 1984 | 2024 evidence update ¹ identified one additional small pilot RCT of the beta-blocker landiolol |
| 1985 | including 36 patients. ⁴⁴⁶ Given the interest in beta-blockers for cardiac arrest and in particular |
| 1986 | esmolol and landiolol, and the fact that sotalol is not considered a primary beta blocker, the ILCOR |
| 1987 | ALS Task Force clarified the third bullet point of its 2018 treatment recommendation to state: |
| 1988 | The confidence in effect estimates is currently too low to support an ALS Task Force |
| 1989 | recommendation about the use of beta-blocker drugs, bretylium, or nifekalant, or sotalol in the |
| 1990 | treatment of adults in cardiac arrest with shock-refractory VF/pVT. |
| 1991 | |
| 1992 | This 2025 guidance is therefore primarily based on the 2018 ILCOR systematic review that identified |
| 1993 | evidence from 14 RCTs and 17 observational studies that included lidocaine, amiodarone, |
| 1994 | magnesium, bretylium, nifekalant, procainamide, sotalol and beta-blockers.447 Meta-analysis of |
| 1995 | randomised trials in adults, found that none of the antiarrhythmic drugs improved survival or |
| 1996 | favourable neurological outcome compared to placebo. |
| 1997 | |
| 1998 | The largest randomised trial compared amiodarone, lidocaine or placebo in patients with VF/pVT |
| 1999 | refractory after at least one defibrillation attempt. Compared with placebo, amiodarone and |
| 2000 | lidocaine increased survival to hospital admission. However, there was no difference in survival to |
| 2001 | discharge or favourable neurological survival at discharge between groups. ⁴⁴⁸ In the pre-defined sub- |
| 2002 | group of bystander-witnessed cardiac arrests, amiodarone and lidocaine increased survival to |
| 2003 | hospital discharge compared with placebo. Survival was also higher with amiodarone than with |
| 2004 | placebo after EMS-witnessed arrest. |

2005



These data led ILCOR to suggest that amiodarone or lidocaine could be used in adults with shock refractory VF/pVT (weak recommendation, low quality evidence).⁴⁴⁵ The values and preferences analysis indicates that the Task Force prioritised the pre-defined and reported sub-group analysis from the ALPS study, which showed greater survival with amiodarone and lidocaine in patients with a witnessed cardiac arrest. ILCOR did not support the use of magnesium, bretylium, nifekalant or procainamide.

2012

2013 The ERC updated its guidelines in 2018 to recommend that amiodarone should be given after three 2014 defibrillation attempts, irrespective of whether they are consecutive shocks, or interrupted by CPR, or for recurrent VF/pVT during cardiac arrest.⁴⁴⁹ The initial recommended dose is amiodarone 2015 2016 300 mg; a further dose of 150 mg may be given after five defibrillation attempts. The 2017 recommendation in favour of amiodarone was based on 21 of 24 National Resuscitation Councils of Europe reporting that amiodarone was the main drug used during CPR.⁴⁴⁹ Lidocaine 100 mg may be 2018 2019 used as an alternative if amiodarone is not available, or a local decision has been made to use 2020 lidocaine instead of amiodarone. An additional bolus of lidocaine 50 mg can also be given after five 2021 defibrillation attempts.449

2022

2023 [h3] Thrombolytic therapy

2024 ILCOR evidence updates in 2022 and 2024 for the treatment of pulmonary embolism did not identify 2025 sufficient new evidence to justify an update of the previous 2020 ILCOR systematic review.^{191,192} We 2026 also considered a recent systematic review that identified 13 studies with 804 patients.⁴⁵⁰ The 2020 ILCOR CoSTR pooled evidence from a sub-group analysis of the TROICA trial⁴⁵¹ and four observational 2027 2028 studies which examined the use of thrombolytic drugs in cardiac arrest caused by suspected or 2029 confirmed pulmonary embolism (PE).⁴⁵²⁻⁴⁵⁵ The studies did not find evidence that thrombolytic drugs 2030 improved neurological outcome.^{451,454} By contrast, in one study, 30-day survival was higher in the 2031 intervention group $(16\% \text{ vs } 6\%; \text{P} = 0.005)^{455}$ but not in three other studies which examined survival to discharge.⁴⁵²⁻⁴⁵⁴ ROSC also improved in one study ⁴⁵³but not in two others. ^{452,454} In making a weak 2032 2033 recommendation for the use of thrombolytic drugs for suspected or confirmed PE and cardiac arrest 2034 based on very low certainty evidence, the ILCOR Task Force considered that the potential benefits 2035 outweighed the potential harm from bleeding.²²³

2036

The ERC endorses the recommendation from ILCOR, which aligns with the ERC guidelines in 2015
 and 2021.^{59,214} The ERC does not support the routine use of thrombolytic drugs in cardiac arrest,
 unless the cause is suspected or confirmed PE. When thrombolytic drugs have been administered,



2040 based on evidence from case series consider continuing CPR attempts for at least 60-90 minutes

2041 before termination of resuscitation attempts.⁴⁵⁶⁻⁴⁵⁸ As for all cardiac arrests the duration of the CPR

attempt should take into account the possibility of reversing the underlying cause, and other factors

2043 such as patient comorbidity and frailty.

2044

2045 [h3] Fluids and blood components

2046 ILCOR has not recently addressed the use of fluids during cardiac arrest. The ERC ALS Writing Group 2047 performed its own search up to February 2025. No RCTs have evaluated the routine administration of 2048 fluids versus no fluids as a treatment strategy for cardiac arrest. Two large RCTs provide indirect 2049 evidence from treatment strategies designed to induce hypothermia, which included the administration of up to 2 L of ice-cold IV fluids during OHCA⁴⁵⁹ or immediately after ROSC.⁴⁶⁰ The 2050 2051 studies found no improvement in short^{459,460} or long-term outcomes.⁴⁶¹ The studies reported evidence of reduced ROSC in patients with VF,⁴⁵⁹ increased rate of re-arrest,⁴⁶⁰ and higher rates of 2052 pulmonary oedema.^{459,460} It is not possible to determine from these studies whether the harmful 2053 2054 effects were related to fluid volume, the rate of infusion, or the temperature of the infused fluids.⁴⁶² 2055 The routine rapid infusion of large-volume fluids should be avoided unless there is evidence or 2056 suspicion of a hypovolaemic cause of the cardiac arrest. A clinical assessment, the history of events 2057 before cardiac arrest, and when the skills are available, POCUS can help identify hypovolaemia during 2058 resuscitation.

2059

A recent systematic review on fluid therapy during and after CPR for non-traumatic CA confirmed the limited evidence for the use of either isotonic saline or balanced crystalloids, and the potential benefit of hypertonic fluids, and mentioned the potential role of optimised fluid resuscitation with accurate clinical assessment of volumes status during CPR, including considering medical history, exam findings and POCUS.⁴⁶³

2065

Either isotonic saline or balanced crystalloids can be considered during CPR. A multicentre RCT (n=432) compared the infusion of two units each of packed red blood cells (PRBC) and fresh frozen plasma versus 1 L of isotonic saline in patients with traumatic haemorrhagic shock.⁴⁶⁴ The study was stopped earlier than the planned sample size of 490 patients due to the COVID-19 pandemic. No difference was observed in the primary outcome parameter (composite of mortality and lactate clearance failure) and secondary outcomes: i) mortality and ii) lactate clearance failure); serious adverse events were comparable. The role of intra- and peri-arrest administration of hypertonic



- fluids⁴⁶⁵, blood, and blood products for non-traumatic cardiac arrest remains uncertain,⁴⁶⁴ and these
 products should only be used in clinical trials.
- 2075

2076 [h3] Other drugs

2077 [h4] Calcium

2078 ILCOR updated the CoSTR for calcium administration during cardiac arrest based on a systematic 2079 review that included three randomized trials and eight observational studies in adult cardiac 2080 arrest.⁴⁶⁶ The largest and most recent trial of 391 patients comparing calcium chloride with placebo 2081 during OHCA was terminated early because an interim analysis suggested potential harm.^{467,468} While 2082 no statistically significant differences were observed in ROSC, neurological outcomes at 30 days, or 2083 survival up to 1 year, patients receiving calcium had lower rates of favourable neurological outcome 2084 at 90 days and 1 year, with a signal of harm across multiple outcomes. Two older, smaller trials of 2085 cardiac arrest patients with refractory asystole or pulseless electrical activity, along with most observational studies, also failed to show any survival benefit.^{466,469,470} These data led ILCOR to 2086 2087 recommend against routine calcium administration in both OHCA (strong recommendation, 2088 moderate certainty evidence) and IHCA (weak recommendation, low certainty evidence).¹ Consistent 2089 with ILCOR, the ERC recommends not routinely giving calcium during cardiac arrest. As noted in the 2090 ERC Guidelines 2025 Special Circumstances, for cardiac arrest caused by suspected hyperkalaemia,

- 2091 ILCOR found insufficient evidence to recommend for or against calcium administration.¹¹³
- 2092

2093 [h4] Sodium bicarbonate

2094 ILCOR updated the CoSTR for sodium bicarbonate administration during cardiac arrest based on a 2095 systematic review including three RCTs and three observational studies in OHCA.¹ Meta-analyses 2096 found no benefit of sodium bicarbonate administration compared to standard care for short-term 2097 survival, survival to hospital discharge, and survival with favourable neurological outcome at one 2098 month.⁴⁷¹⁻⁴⁷³ Supporting observational data showed similar results.⁴⁷⁴⁻⁴⁷⁹ Although the evidence is 2099 inconclusive, sodium bicarbonate is commonly given during cardiac arrest.⁴⁷⁹⁻⁴⁸¹ ILCOR suggests 2100 against routine sodium bicarbonate administration in both OHCA (weak recommendation, low 2101 certainty evidence) and IHCA (weak recommendation, very low certainty evidence).¹ The ERC does 2102 not support the routine administration of sodium bicarbonate during cardiac arrest, unless there is a 2103 specific indication.

2104

2105 [h4] Corticosteroids



2106 The guidance for administering corticosteroids during cardiac arrest, whether given alone or in 2107 combination with vasopressin, is based on an ILCOR evidence update and systematic review of 2108 individual participants data.^{1,482,483} In adult IHCA, three RCTs comparing vasopressin and 2109 methylprednisolone to placebo found higher rates of ROSC and favourable neurological outcome at 2110 hospital discharge, but no significant improvement in survival, longer-term neurological outcome, or health-related quality of life up to 90 days.^{136,484,485} Secondary analyses of the largest trial also found 2111 2112 no difference in haemodynamic or long-term outcomes for vasopressin and 2113 methylprednisolone.^{486,487} The two earlier smaller trials that administered corticosteroids both 2114 during and after ROSC reported improved outcomes overall.^{484,485} Given that no consistent survival 2115 benefit was identified in meta-analyses, ILCOR suggests against using vasopressin and corticosteroids 2116 in addition to usual care for adult cardiac arrest.¹ For corticosteroids as a standalone intervention 2117 during cardiac arrest, early small trials have found no meaningful benefit, including three trials in OHCA and one in IHCA.^{488,489} Based on these data, and considering the practical challenges of 2118 2119 incorporating additional drugs into resuscitation protocols, the ERC recommends against the routine 2120 use of corticosteroids either alone or in combination with vasopressin during cardiac arrest except 2121 when this is done as part of a clinical trial.

2122

2123 [h2] ALS for cardiac arrest in highly-monitored settings

2124 Patients in highly monitored settings such as critical care areas, operating or recovery rooms, and 2125 cardiac catheterisation laboratories are closely and continuously monitored, and causes of 2126 unexpected cardiac arrest may be promptly reversible, particularly when detected immediately (e.g., 2127 a sudden arrhythmia in the ICU or a relative overdose of induction drug in the operating room). If 2128 blood pressure is monitored continuously using an indwelling arterial cannula, sudden changes in 2129 blood pressure can be detected almost immediately and a pulsatile arterial waveform may be seen 2130 even if peripheral and central pulses are no longer palpable. A sudden decrease in cardiac output, 2131 including cardiac arrest, may be detected very quickly by a sudden decrease in ETCO₂ if this is being 2132 monitored continuously. These are just some of the features of cardiac arrest in a highly monitored 2133 setting that may warrant adjustment of the doses of resuscitation drugs, changes in defibrillation 2134 strategy, and different indications for starting chest compressions.

- 2135
- 2136 A recent Royal College of Anaesthetists National Audit in the United Kingdom (National Audit Project
- 2137 7) estimated the incidence of intra-operative cardiac arrest to be 3 in 10,000.⁴⁹⁰ Of the 548 patients
- with intraoperative cardiac arrest during the one-year period of study, half had invasive blood
- 2139 pressure monitoring. Sustained ROSC was achieved in 78% of patients and 62% were alive at the



time of reporting. Only 70 patients (13%) had a shockable rhythm. Among the patients with an initial
rhythm of PEA or bradycardia, the three most reported triggers for starting CPR were an impalpable
pulse (57%), severe hypotension (47%), and a reduction in ETCO₂ (25%).

2143

In adults during general anaesthesia, it has been suggested that chest compressions should be
started if the systolic blood pressure decreases and remains < 50 mmHg despite interventions.^{491,492}
This approach is likely also applicable to any patient undergoing invasive arterial monitoring (e.g., in
an critical care setting). A systolic blood pressure < 50 mmHg is likely to be associated with
impalpable pulses⁴⁹³ and the benefit from the additional cerebral and coronary blood flow produced
by chest compressions will likely outweigh the risk of harm from chest compressions.⁴⁹¹

2150

2151 Although the standard ERC ALS guidelines recommend a dose of 1 mg adrenaline, the optimal dose 2152 of adrenaline to treat cardiac arrest is unknown.⁴⁹⁴ Smaller doses are likely appropriate when 2153 adrenaline is first given IV for profound hypotension, when there is a high probability of a low-flow 2154 state during PEA or severe bradycardia, or when there is a very short time between the onset of 2155 cardiac arrest and injection of adrenaline. With continuous monitoring, cardiac arrest can be 2156 diagnosed immediately and treated rapidly with the likelihood of restoring spontaneous circulation 2157 very quickly. Under these circumstances, giving a 1 mg bolus of adrenaline may lead to severe 2158 hypertension and tachyarrhythmias. Thus, we suggest that adrenaline is given initially in increments 2159 (e.g., $50-100 \ \mu g \ IV$) rather than a 1 mg bolus, but if 1 mg in total has been given with no response, 2160 further IV adrenaline doses of 1 mg are given at the usual 3-5 minute intervals. Australasian 2161 guidelines for resuscitation after cardiac surgery include similar recommendations.⁴⁹⁵ When 2162 continuously monitoring arterial blood pressure it is reasonable to aim to achieve a diastolic blood 2163 pressure higher than 30 mmHg by combining high-quality chest compressions and titrated 2164 adrenaline (by giving 50–100 µg boluses or an infusion) (see physiology-guided CPR). 496,497 2165

The International Liaison Committee on Resuscitation does not recommend the precordial thump for
established cardiac arrest, citing its low success rate documented in a systematic review – although
already de-emphasised in previous ERC guidelines, we have removed it completely from this
guideline.⁴⁹⁸ In two studies of patients undergoing electrophysiological testing, it terminated
malignant ventricular arrhythmias (VT or VF) in only 1 of 80 and 2 of 155 cases, respectively—all
cases were VT, and none of the 49 VF cases responded.^{499,500} In an OHCA study, a precordial thump



- 2172 restored circulation in 5 of 103 patients (3 VF, 2 pVT) but worsened rhythms in 10 cases.⁵⁰¹ Another 2173 Italian OHCA study found it effective in only 3 of 144 patients, all initially in asystole.⁵⁰² 2174 2175 [h2] Physiology-guided CPR 2176 The quality of CPR is associated with survival and the ability to monitor the quality of CPR is fundamental to its improvement.⁵⁰³ The quality of CPR can be measured by monitoring the 2177 2178 performance of the rescuer to ensure adherence to guidelines; this might include compression rate, 2179 depth and recoil, chest compression fraction and ventilation rate and volume. Many of the current 2180 defibrillator-monitors are capable of monitoring and providing these metrics in real time. Another 2181 way of monitoring CPR performance is to assess its effect on the patient's physiology using intra-2182 arterial blood pressure, capnography or cerebral oximetry. 2183 2184 During low flow conditions, ETCO₂ values are more strongly associated with cardiac output and pulmonary blood flow than with ventilation.⁵⁰⁴ The use of ETCO₂ to guide the quality of chest 2185 2186 compressions was first described almost 40 years ago.⁵⁰⁵ Two observational studies have 2187 subsequently shown that chest compression depth but not rate is associated with ETCO₂ values, ^{506,507} 2188 although a third observational study did show a small association between compression rate and 2189 ETCO₂.⁵⁰⁸ In a pig model of neonatal asphyxial cardiac arrest, chest compressions aimed at maximising ETCO₂ increased the rate of ROSC compared with a control group of standard CPR.⁵⁰⁹ 2190 2191 2192 A systematic review of haemodynamic-directed feedback during CPR identified six animal studies.⁵¹⁰ 2193 Four of these studies examined the effect of haemodynamic-directed CPR on survival in swine. The 2194 hemodynamic-directed CPR groups had chest compression depth titrated to a systolic blood pressure 2195 (SBP) of 100 mmHg and vasopressors titrated to maintain a coronary perfusion pressure (CPP) >20 2196 mmHg. Pooled results showed that 35/37 (94.6%) of the animals survived in the haemodynamic-2197 directed CPR groups and 12/35 (34.3%) survived in the control groups (p < 0.001). 2198 2199 Most of the clinical studies of physiology-directed CPR have involved children. In a stepped-wedge 2200 cluster randomised trial in 18 paediatric intensive care units, a bundled intervention comprising 2201 physiologically focused CPR training at the point of care and structured clinical event debriefings did not improve survival to hospital discharge with favourable functional outcome.⁵¹¹ In this study, when 2202 2203 an arterial line or ETCO₂ was in place, clinicians were educated to aim for a systolic blood pressure
- $2204 \qquad \text{above 100 mmHg in older children (i.e. not neonates or infants) and an ETCO_2 above 25 mmHg.}$
- 2205 Observational data from this study demonstrate that achieving a mean diastolic BP (DBP) during CPR



| 2206 | greater than 30 mmHg for older children is associated with higher rates of survival to hospital |
|------|--|
| 2207 | discharge. ⁴⁹⁶ |
| 2208 | |
| 2209 | In one of the first clinical studies in adults on this topic, invasive blood pressure was measured during |
| 2210 | CPR in 104 patients. ⁵¹² A chest compression rate of 100–120 min ⁻¹ and compression depth \ge 6 cm |
| 2211 | was associated with a DBP ≥30 mmHg in both femoral and radial recordings. However, although |
| 2212 | there was a weak upward trend in blood pressure as compression depth increased in individual |
| 2213 | subjects, deeper compression depth did not result in a higher blood pressure in all patients. In a |
| 2214 | more recent study, 80 OHCA adults had invasive blood pressure monitoring during CPR delivered by |
| 2215 | helicopter emergency medicine service (HEMS) clinicians. ⁴⁹⁷ The maximum, average and delta-DBP |
| 2216 | (difference between the initial and maximum values); and maximum and average mean arterial |
| 2217 | pressure (MAP) were positively associated with ROSC. Maximum DBP had an optimal threshold value |
| 2218 | of 35 mmHg (sensitivity 94.1 %; specificity 58.7 %) for predicting ROSC. The odds ratio for ROSC was |
| 2219 | 1.05 (95 % CI 1.03–1.08) for every 1 mmHg increase in maximum DBP. |
| 2220 | |
| 2221 | A consensus statement from the American Heart Association suggested several physiological targets |
| 2222 | during CPR, all of which were based on expert consensus:503 |
| 2223 | • If both an arterial line and a central venous catheter are in situ during cardiac arrest, aim to |
| 2224 | achieve a coronary perfusion pressure (CPP) > 20 mmHg. |
| 2225 | • If only an arterial line is in situ during cardiac arrest, aim to maintain a DBP > 25 mmHg. |
| 2226 | • If only capnography is available during cardiac arrest, aim to achieve an ETCO ₂ > 20 mmHg (3.3 |
| 2227 | КРа). |
| 2228 | |
| 2229 | Given the recent observational data, ^{496 512 497} the ERC suggests aiming for a diastolic blood pressure |
| 2230 | of ≥ 30 mmHg when using intra-arterial blood pressure monitoring for physiology-guided CPR. |
| 2231 | |
| 2232 | Other potential physiological targets during CPR include cerebral oximetry and EEG. Higher cerebral |
| 2233 | oxygenation saturation during CPR is associated with ROSC ⁵¹³ , but although theoretically cerebral |
| 2234 | oxygenation could be used as a physiological target during CPR, this has yet to be investigated (see |
| 2235 | below). |
| 2236 | |
| 2237 | [h2] Waveform capnography during advanced life support |



2238 End-tidal CO_2 (ETCO₂) is the partial pressure of carbon dioxide (PCO₂) measured at the end of 2239 expiration. It reflects cardiac output, tissue perfusion, pulmonary blood flow, and minute ventilation. 2240 Carbon dioxide is produced in perfused tissues by aerobic metabolism, transported by the venous 2241 system to the right side of the heart and pumped to the lungs by the right ventricle, where it is 2242 removed by alveolar ventilation. 2243 2244 Waveform capnography enables a continuous, non-invasive measurement of PCO₂ in the exhaled air 2245 during CPR. In the typical capnogram, the ETCO₂ recorded at the end of the expiratory plateau best 2246 reflects the alveolar PCO2.⁵¹⁴ ETCO₂ is most reliable when the patient's trachea is intubated, but it 2247 can also be detected with a supraglottic airway device or bag mask.⁵¹⁵ 2248 2249 The aims of monitoring waveform capnography during CPR include: 2250 Confirming tracheal tube placement (this has been addressed above). 2251 Monitoring the quality of CPR. ETCO₂ monitoring during CPR enables the measurement of the ventilation rate helping rescuers avoid hypo- or hyperventilation.^{377,515} Observational studies on 2252 2253 adults with in-hospital or OHCA have shown that ETCO₂ values are proportional to the chest 2254 compression depth measured using transthoracic impedance. In contrast, variations in the chest compression rate do not affect ETCO₂.^{506,507} 2255 2256 • **Detecting ROSC during CPR**. When ROSC occurs, ETCO₂ may rise three times above the 2257 values during CPR. Capnography during CPR might, therefore, help avoid unnecessary chest 2258 compression or potentially harmful administration of adrenaline in a patient with ROSC. However, no 2259 consistent amount of ETCO₂ rise has been identified as a criterion for diagnosing that ROSC has 2260 occurred. In addition, the rise in ETCO₂ associated with ROSC is not immediate and can start 10 minutes or more before a palpable pulse is detected. 516-518 2261

2262 Prognostication during CPR. Lower ETCO₂ values during CPR indicate a lower likelihood of • 2263 ROSC. Previous evidence suggested that failure to achieve an ETCO₂ value >10 mmHg (1.33 kPa) was associated with a poor outcome.^{519,520} This threshold has also been suggested as a criterion for 2264 2265 withholding e-CPR in refractory cardiac arrest.⁵²¹ However, recent evidence has shown that patients 2266 may survive with ETCO2 values below 10 mmHg during CPR. In a study on 617 adults with refractory 2267 OHCA from non-shockable rhythm, of whom 615 (99.3%) died before hospital discharge, ETCO₂ at 30 2268 minutes was still above 10 mmHg in 88% of non-survivors. In one of the two survivors, 30-min ETCO₂ was below 10 mmHg.⁵²² In a study on 14,122 adult non-traumatic OHCA from all rhythms, 4.2% of 2269 2270 the 9,226 patients who subsequently achieved ROSC and 3.3% of those who survived to hospital 2271 discharge had ETCO₂ <10 mmHg at 20 minutes.⁵²³ In that study, the adjusted odds of mortality in



patients with maximum prehospital ETCO₂ values above 50 mmHg (6.67 Kpa) were 50% higher than
 in patients with ETCO₂ values of 30-40 mmHg (4.0 - 5.33 KPa), probably reflecting hypoventilation.
 Other factors affecting ETCO₂ during cardiac arrest include the presence of airway closure⁵²⁴, and the

2275 cause of arrest.⁵²⁵⁻⁵²⁷

- 2276 Besides single values of ETCO₂, the changes in ETCO₂ during CPR have also been investigated to predict ROSC.^{528,529} In a recent secondary analysis of 1113 patients enrolled in the multicentre cluster 2277 2278 randomised PART trial, in which patients were ventilated with supraglottic airways or endotracheal 2279 tube, the median ETCO₂ increased from 30.5 at 10 minutes to 43.0 mmHg (p for trend < .001) five 2280 minutes before the end of resuscitation in ROSC patients, while it declined from 30.8 to 22.5 mmHg (p for trend < .001) in non-ROSC patients, respectively.⁵³⁰ After adjusting for major confounders, the 2281 2282 slope of the ETCO₂ during CPR remained associated with both ROSC and 72-h survival (OR 1.45 [1.31-2283 1.61] and 1.33 [1.20-1.45] respectively). ETCO₂ trends are probably more appropriate than point 2284 values for predicting ROSC during CPR.⁵²⁰ In the study on refractory non-shockable OHCA mentioned 2285 above, while only one of the two survivors to discharge had an ETCO₂ above 10 mmHg at 30 minutes, 2286 both had an ETCO₂ increase from the initial value to 30 minutes (specificity 13% vs 33%).⁵²² In a 2287 multicentre study on 668 OHCA patients of any aetiology and rhythm, both an ETCO₂ value at 2288 intubation greater than 20 mmHg (2.67KPa) and its increase 10 minutes later were independent 2289 predictors of survival to hospital admission and survival at hospital discharge. However, after 2290 adjustment for bystander and CPR status, presenting rhythm and EMS arrival time, only the ETCO₂ 2291 change at 10 minutes remained an independent predictor of outcome.⁵³¹
- 2292 Evidence regarding the prognostic value of ETCO₂ is based on observational, unblinded studies,
- which may have caused a self-fulfilling prophecy. Although ETCO₂ values below 10 mmHg after
- prolonged CPR are strongly associated with unfavourable outcome (no ROSC or death before hospital
- discharge), the ERC recommends that they should not be used alone as a mortality predictor or for
- 2296 deciding to stop a resuscitation attempt
- 2297

2298 [h2] Near infra-red spectroscopy (NIRS), and EEG monitoring during CPR

Although NIRS and EEG monitoring during CPR were within the scope of the 2025 ALS guidelines update, the ERC has not made any recommendations for their use and the reasons are set out below. Near infrared spectroscopy is a non-invasive method to monitor the oxygenation of the combination of arterial and venous blood in the brain. NIRS monitoring can be conducted with external electrodes placed on the forehead of the patient. With the emission of infrared light, the monitor can provide the regional oxygen saturation proportion in tissue and blood. When applied to the forehead of a patient this will provide an estimation of the amount of oxygen saturation in brain tissue. Given the



2306 risk of hypoxic-ischaemic brain injury in patients undergoing CPR this has raised much interest. 2307 According to a systematic review conducted in 2021, NIRS monitoring may provide information on 2308 the likelihood of ROSC.⁵³² The systematic review including 17 studies showed that higher initial 2309 cerebral regional oxygen saturation (rSO₂) values were associated with the likelihood of ROSC. Thus 2310 far only one study has compared ETCO₂ and NIRS and this study suggested superiority for NIRS compared with ETCO₂.⁵³³ This finding should however be confirmed in future studies. On the other 2311 2312 hand, studies have failed to show any association of NIRS values with the quality of chest 2313 compressions, the amount of oxygen in the blood or the administration of adrenaline.^{359,534} Thus, it is 2314 unclear if NIRS can be used to modify the delivery of resuscitation measures in any way. No study to 2315 date has assessed the cost effectiveness of NIRS monitoring. Therefore, there is not enough evidence 2316 to recommend NIRS as a monitoring tool for patients undergoing CPR.

2317

EEG monitors the electric activity of the brain noninvasively. Changes in EEG correlate with changes in cerebral blood flow. Some studies have assessed the feasibility of using EEG during resuscitation to monitor the efficacy and outcome of CPR.⁵³⁵ The EEG methods used include traditional multichannel EEG as well as the raw EEG signal from monitors of the depth of anaesthesia, including the bispectral index (BIS®). The existing evidence is inconclusive and comes mainly from smaller observational studies and case-reports. At present, the use of EEG is not recommended by the ERC either for monitoring the performance of CPR or estimating the outcome.

2325

2326 [h2]Use of ultrasound imaging during advanced life support

Point-of-care ultrasound (POCUS) imaging is already commonly used in emergency care settings. Its
 use during CPR is also increasing. Previous and current guidance emphasises the need for skilled
 POCUS operators and minimising interruption in chest compression to acquire images.^{59,214}

2330 2331 An ILCOR systematic review assessed the role of POCUS during cardiac arrest as a prognostic tool in 2332 which assessment of cardiac motion informs the likelihood of achieving ROSC and clinical decisions 2333 to terminate resuscitation.⁵³⁶ The review identified several limitations such as inconsistent definitions 2334 and terminology around sonographic evidence of cardiac motion, low inter-rater reliability of 2335 findings, low sensitivity and specificity for outcomes, confounding from self-fulfilling prophecy when 2336 terminating resuscitation in unblinded settings as well as unspecified timing of POCUS. The review 2337 concluded that no sonographic finding had sufficiently diagnostic accuracy to support its use as a 2338 sole criterion to terminate resuscitation. A 2025 ILCOR Evidence Update¹ identified additional studies 2339 with small sample sizes, and heterogeneous POCUS findings and clinical outcomes.⁵³⁷⁻⁵⁴⁰ All studies



were potentially biased by lack of blinding of the resuscitating team to POCUS findings. There was a
lack of agreement in the interpretation of acquired views in most of studies, which highlighted the
inherent real-time clinical challenge of time-limited image acquisition and the need for skilled POCUS
operators.

2344

2345 A further ILCOR systematic review assessed the role of POCUS to diagnose treatable causes of cardiac 2346 arrest such as cardiac tamponade, pneumothorax, pulmonary embolism, myocardial infarction, 2347 aortic dissection, and hypovolaemia.⁵⁴¹ There was a high degree of clinical heterogeneity and a 2348 critical risk of bias, which precluded a meta-analysis. The certainty of evidence was very low, and 2349 individual studies were difficult to interpret. However, the review stressed the issue of 2350 misinterpreting POCUS findings as the cause of cardiac arrest as opposed to an incidental finding. For 2351 example, unilaterally absent lung sliding could indicate a small pneumothorax or mainstem bronchial 2352 intubation, rather than a tension pneumothorax. Likewise, visualised peritoneal fluid could be 2353 ascites, rather than an acute haemorrhage; a pericardial effusion could be present without cardiac 2354 tamponade; and right heart dilation can occur during CPR without a massive pulmonary embolism. 2355 Right ventricular dilation a few minutes after the onset of cardiac arrest as blood moves from the 2356 systemic circulation to the right heart along its pressure gradient ⁵⁴² was consistently observed in a 2357 porcine model of cardiac arrest caused by hypovolaemia, hyperkalaemia, and primary arrhythmia. ⁵⁴³ 2358 This is a common trans-oesophageal echocardiography (TOE) observation in patients with OHCA 2359 regardless of the cause. 544

2360

2361 Since 2015, the ERC ALS guidelines recommend a transthoracic sub-xiphoid view with the probe 2362 placed just before chest compressions are paused for a planned rhythm assessment²¹⁴, minimising additional interruptions in chest compressions, which may delay or impede other therapies. 545,546 2363 2364 One proposed strategy to deal with this is to record brief sonographic video clips during 2365 pulse/rhythm checks (less than 10 seconds) and then view/interpret them after the resumption of 2366 chest compressions. Alternatively, sonographers may pre-localise the approximate acoustic window 2367 during chest compressions with subsequent fine-tuning during pauses in CPR.⁵⁴⁷ Another possibility 2368 for reducing hands-off time during cardiac arrest is the use of TOE. In 2021, a systematic review of 2369 TOE in cardiac arrest concluded that because of the heterogeneity of studies, small sample size and 2370 inconsistent reference standard, the evidence for TOE in cardiac arrest resuscitation is of low certainty and is affected by a high risk of bias.⁵⁴⁸ Further studies are needed to better understand the 2371 2372 true diagnostic accuracy of TOE in identifying reversible causes of arrest and cardiac contractility. 2373 Since then, additional case series and observational studies have been published, which showed that



- in experienced hands, TOE provides useful diagnostic and therapeutic information and that the rate
 of adequate cardiac visualisation can be excellent with a low complication rate.⁵⁴⁹ During CPR, TOE
 may help to improve hand positioning for chest compression and improve left ventricular
 compression.⁵⁵⁰ In an observational study, improving left ventricular compression was associated
 with increased ROSC.⁵⁴⁰ The use of TOE during cardiac arrest requires additional equipment and
- 2379 expertise.
- 2380

2381 [h2] Devices

2382 [h3] Mechanical chest compression devices

2383 Since the 2021 guidelines⁵⁹, ILCOR has published updated recommendations for the use of 2384 mechanical chest compression. The ILCOR 2024 systematic review identified 14 reports of 11 RCTs 2385 conducted post-2000.¹ Trials before this date were not included because of the significant changes to 2386 CPR and cardiac arrest treatment that have occurred since 2000. Three new RCTs were identified 2387 since the 2021 ERC guidelines, each providing very low certainty evidence.⁵⁵¹⁻⁵⁵³ One study, enrolling 2388 1191 patients, assessed the use of the LUCAS device following OHCA. No difference in ROSC (RR 0.90 2389 [95% CI 0.62 to 1.32]) or survival to 30 days (RR 0.89 [95% CI 0.41 to 1.92]) was found when 2390 comparing LUCAS with manual CPR.⁵⁵¹ Two studies compared LUCAS with manual CPR following 2391 IHCA.^{552,553} One study enrolled patients with IHCA (but not in the emergency department) with nonshockable rhythms (n=127).⁵⁵² No benefit in survival with favourable neurological outcome was 2392 2393 found with LUCAS compared with manual CPR (RR 1.13 [95% CI 0.13 to 9.72]). A trial enrolling 2394 patients sustaining witnessed cardiac arrest in the emergency department found no difference in ROSC when using a LUCAS device compared with manual CPR (RR 0.80 [95% CI 0.55 to 1.173]).⁵⁵³ No 2395 2396 new RCTs reporting safety outcomes were identified in the ILCOR systematic review. Consistent with 2397 the ILCOR treatment recommendations, the ERC recommends considering mechanical chest 2398 compression only if high-quality manual chest compressions are not practical (e.g., during 2399 percutaneous coronary intervention or ECMO cannulation) or compromise provider safety (e.g. 2400 during transport). When a mechanical device is used, delays in the initial defibrillation attempt 2401 should be avoided and pauses to chest compression during device deployment minimised. 2402 Mechanical chest compression should only be used by trained teams familiar with the device. 2403 2404 [h3] Resuscitative endovascular balloon occlusion of the aorta (REBOA) 2405 Resuscitative endovascular balloon occlusion of the aorta (REBOA) is a technique in which blood flow

through the aorta is occluded by inflating an intra-aortic balloon. The use of REBOA has not yet been

addressed by ILCOR but was included in the scope of the ERC guidelines. REBOA has been used in the



2408 management of haemorrhagic shock and traumatic cardiac arrest.^{554,555} However, in a recent RCT the

in-hospital use of REBOA in patients with exsanguinating traumatic haemorrhage did not improve 90-

- 2410 day survival and may even have increased mortality compared with standard care alone.⁵⁵⁶ It has
- recently been proposed as an adjunctive treatment for patients with non-traumatic cardiac arrest
- because of its potential to redistribute blood flow to organs proximal to the occlusion, thereby
- 2413 increasing cerebral and coronary perfusion.⁵⁵⁷
- 2414
- 2415 In animal models of non-traumatic cardiac arrest, inflation of REBOA increases central diastolic
- 2416 arterial pressure, which is a surrogate marker of coronary artery pressure and cerebral
- 2417 perfusion.^{558,559} However, human evidence is currently limited to case reports⁵⁶⁰⁻⁵⁶⁴ and small case
- series with a total of 78 patients.⁵⁶⁵⁻⁵⁷³ These studies aimed to assess the feasibility of positioning
- 2419 REBOA during resuscitation. REBOA positioning was successful in 72 cases (92%) and ROSC was
- achieved in 35 cases (45%) but only 16 (20.5%) had sustained ROSC. Approximately 80% of patients
- with sustained ROSC died (13/16); of these, nearly half died from brain injury^{561,562,568,573} and less
- 2422 frequently from withdrawal of life-sustaining treatment or re-arrest after balloon deflation.⁵⁶⁷
- 2423 Currently, there are no data to support the use of REBOA in non-traumatic OHCA. There are two (one
- 2424 pre-hospital; one emergency department) ongoing efficacy RCTs which aim to evaluate the ability of
- REBOA to increase ROSC in adult, witnessed, non-traumatic cardiac arrest and its potential effects on
 survival and neurological outcome.^{574,575}
- ----

2427

2428 [h3] Intra-arrest cooling

2429 Cooling of patients during CPR may potentially alleviate reperfusion injury and prevent or reduce 2430 hypoxic-ischaemic brain injury. The most recent ILCOR systematic review on temperature control in 2431 cardiac arrest patients found no evidence of improved outcome in patients treated with pre-hospital 2432 cooling in patients with OHCA.⁵⁷⁶⁻⁵⁷⁸ These reviews did not separate cooling conducted during or 2433 after ROSC. This may be important with respect of both the effects and side effects of the 2434 intervention. A systematic review focusing on cooling during CPR was published in 2021.⁵⁷⁹ This 2435 review identified four RCTs (two of high certainty and two of moderate certainty of evidence) including 2305 patients that compared intra-arrest cooling with cooling started in the hospital. 459,580-2436 2437 ⁵⁸² Two studies used evaporative intranasal cooling: one used cooling with cold IV fluids (up to 2000 2438 mL) and one used a combination of cold fluids (up 2000 mL) and external cooling with gel pads. In 2439 comparison with the control patients, the use of cooling during CPR was not associated with 2440 improved favourable neurological outcome (OR 0.96 [95% Cls 0.68-1.37]; p= 0.84), any change in 2441 ROSC (OR 1.11 [95% CIs 0.83-1.49]; p= 0.46) or survival to hospital discharge (OR 0.91 [95% CIs 0.73-



- 2442 1.14]; p=0.43). A further analysis of the method of intra-arrest cooling did not appear to influence
- 2443 the results. Therefore, the use of intra arrest cooling is not currently recommended by the ERC. The
- 2444 ongoing PRINCESS 2 trial will assess the effect of intranasal evaporative cooling in patients
- 2445 resuscitated from a shockable initial rhythm.⁵⁸³
- 2446

2447 [h2] Extracorporeal CPR

- 2448 Extracorporeal CPR (ECPR) is the rapid deployment of veno-arterial extracorporeal membrane
- 2449 oxygenation (VA-ECMO) during ongoing CPR to restore and maintain organ perfusion in patients in
- whom conventional CPR is unsuccessful in achieving a sustained ROSC.⁵⁸⁴ The use of ECPR has
- 2451 continued to increase for both IHCA and OHCA in recent years⁵⁸⁵ despite uncertainty in
- 2452 evidence.586,587
- 2453 Since 2020, three RCTs of ECPR have been published. The Advanced Reperfusion Strategies for 2454 Patients with OHCA and Refractory Ventricular Fibrillation Trial (ARREST) demonstrated higher 2455 survival to hospital discharge with ECPR compared to conventional CPR in OHCA patients with an 2456 initial shockable rhythm.⁵⁸⁸ The Prague OHCA study found improved survival with a favourable 2457 neurological outcome at 30 days but not at six months (primary outcome) among 264 patients with 2458 all-rhythms refractory OHCA.⁵⁸⁹ Of note, these were both single-centre trials performed in well-2459 established ECPR systems. A third, multi-centre trial conducted in 10 cardiosurgical centres, Early 2460 Initiation of Extracorporeal Life Support in Refractory OHCA Trial (INCEPTION) showed no difference 2461 between ECPR and conventional CPR in OHCA patients with an initial shockable rhythm.⁵⁹⁰ 2462 Additionally, a small pilot RCT, the Extracorporeal Cardiopulmonary Resuscitation for Refractory Out-2463 of-Hospital Cardiac Arrest (EROCA) trial, failed to meet the feasibility goal of transporting patients to an ECPR-capable emergency department within 30 minutes from OHCA.⁵⁹¹ A meta-analysis of these 2464 2465 four trials demonstrated higher survival with favourable neurological outcome with ECPR compared to conventional CPR.⁵⁹² A Bayesian meta-analysis found a posterior probability of a clinically relevant 2466 2467 ECPR-based treatment effect on 6-month neurologically favourable survival of 71% in patients with 2468 all rhythms and 76% in shockable rhythms.⁵⁹³ For IHCA, RCTs have been conducted, and a meta-2469 analysis of observational data found that ECPR was effective in improving survival and favourable neurological outcomes.594 2470
- 2471

The use of ECPR during cardiac arrest was addressed by an ILCOR systematic review in 2022, which was updated in 2024.^{191,587} The ILCOR ALS Task suggested, and the ERC recommends that ECPR may be considered as a rescue therapy for selected adults with IHCA and OHCA when conventional CPR is failing to restore spontaneous circulation in settings where this can be implemented (weak



recommendation). The overall certainty of evidence was rated as low for OHCA and as very low for
IHCA (downgraded further because all evidence was in OHCA).¹

2478

2479 ECPR is not intended for the entire population of cardiac arrest patients, with only approximately 10% of OHCA cases being eligible candidates.⁵⁸⁹ However, there are no universally agreed selection 2480 2481 criteria for ECPR, and practices vary widely among centres, particularly regarding the inclusion of 2482 older patients, those with initial non-shockable rhythms (pulseless electrical activity or asystole), and 2483 those with longer low-flow times. A recent systematic review and meta-analysis found that younger 2484 age, initial shockable rhythm, witnessed arrest, immediate CPR, pre-cannulation ROSC at any time, 2485 and shorter low-flow times are associated with increased likelihood of survival with favourable 2486 neurological outcome.⁵⁹⁵ Adopting more liberal selection criteria impacts survival and survival with favourable neurological outcome⁵⁹⁶⁻⁵⁹⁸ but also influences organ donation rates.⁵⁹⁹ Conversely, more 2487 2488 restrictive criteria may exclude potential survivors and organ doners.

2489

The ERC, European Extracorporeal Life Support Organization (Euro-ELSO), European Society of 2490 2491 Intensive Care Medicine (ESICM), European Society of Emergency Medicine (EuSEM), European 2492 Society of Anaesthesia and Intensive Care (ESAIC), and the European Board of Cardiovascular 2493 Perfusion (EBCP) have produced consensus guidelines for ECPR in adults and the brief guidance 2494 below is based on these guidelines. Given the time-sensitive nature of ECPR, in systems where ECPR 2495 is available, it is crucial for clinicians to promptly recognise when a patient with ongoing ALS is in 2496 refractory cardiac arrest (i.e., three consecutive unsuccessful defibrillation attempts or 10 minutes of 2497 resuscitation in case of non-shockable rhythms) and might be a suitable candidate, enabling rapid 2498 activation of the ECPR team. 2499 Commonly used selection criteria for initiating ECPR include: 2500 Younger patients (50-75 years), no perceived frailty, and absence of major comorbidities. 2501 Witnessed cardiac arrest with immediate CPR (i.e., a no-flow duration of \leq 5 minutes) 2502 2503 In addition to the previous criteria, the following criteria are commonly applied:

- Initial shockable rhythm or PEA rhythms.
- Estimated time to establish ECPR from starting CPR (i.e., low-flow time) is ideally within 45 to
 60 minutes.
- Known or suspected treatable underlying cause of cardiac arrest.
- ROSC at any time prior to cannulation



| 2509 | Presence of signs of life during CPR |
|------|---|
| 2510 | • ETCO ₂ > 10 mmHg (1.3 kPa) |
| 2511 | Mechanical CPR during transport |
| 2512 | Whether initiating ECPR for OHCA in the pre-hospital setting is superior to in-hospital setting remains |
| 2513 | uncertain ^{18,19} , and ongoing clinical trials aim to address this question (ON-SCENE [ClinicalTrials.gov |
| 2514 | NCT04620070], RACE [NCT06789978], PACER [NCT06177730], PRECARE trial). |
| 2515 | |
| 2516 | [h2] Peri-arrest arrhythmias |
| 2517 | Prompt identification and treatment of life-threatening arrhythmias may prevent cardiac arrest or its |
| 2518 | recurrence. This section offers guidance and treatment algorithms for the non-specialist ALS |
| 2519 | provider. The scope is to focus on arrhythmias occurring pre-arrest or immediately post-ROSC and |
| 2520 | causing life-threatening instability. In case of persistent arrhythmia, the first goal should be to |
| 2521 | ascertain the patient's stability and seek advice from a specialist or more experienced physician. |
| 2522 | Electrical cardioversion is required in the peri-arrest patient with a clinically unstable |
| 2523 | tachyarrhythmia, while pacing is used in refractory, clinically important bradycardia. The key |
| 2524 | interventions are summarised in Figure 9. and Figure 10. |
| 2525 | |
| 2526 | Figure 9. Peri-arrest Tachyarrhythmia Algorithm ABCDE airway, breathing, circulation, disability, |
| 2527 | exposure; BP blood pressure; DC direct current; ECG electrocardiogram; EF ejection fraction, |
| 2528 | IV intravenous; ROSC return of spontaneous circulation; SpO $_2$ oxygen saturation measured with pulse |
| 2529 | oximetry; VT ventricular tachycardia. |
| 2530 | |





2531

- 2532
- 2533 **Figure 10. Bradycardia Algorithm** ABCDE airway, breathing, circulation, disability, exposure
- 2534 BP blood pressure; ECG electrocardiogram; IV intravenous; SpO₂ oxygen saturation measured with

2535 pulse oximetry.



2536

- 2537 These guidelines follow recommendations published by international cardiology societies, including
- the European Society of Cardiology (ESC), the European Heart Rhythm Association (EHRA), the
- 2539 European Society of Cardiothoracic Surgeons, the American Heart Association (AHA), the American



College of Cardiology (ACC) and the Heart Rhythm Society (HRS).^{85,114,600-603} Table 6 summarises the
 supporting evidence for vagal manoeuvres and some of the more commonly used drugs for the
 treatment of arrhythmias.

2543 Ventricular arrhythmias occurring peri-arrest usually originate from a complex interplay between 2544 underlying structural or electrical heart disease, external triggering factors and dominance of the 2545 sympathetic limb of the autonomic nervous system.⁶⁰⁴ Reversible triggering factors include acute 2546 myocardial ischaemia, electrolyte imbalance, fever or hypothermia, hormonal factors, sepsis, 2547 starvation and decompensated heart failure.^{605,606} For patients with wide QRS arrhythmias peri-2548 arrest, expert help should be pursued whenever possible to either help with the termination of the 2549 arrhythmia or to prevent recurrences.

2550 Treatment of ventricular arrhythmias depends on the haemodynamic consequences of the

arrhythmia, its morphology on ECG and the underlying myocardial substrate.⁶⁰⁵ For patients with

2552 monomorphic VT and structural heart disease or uncertain myocardial substrate, current European

2553 Society of Cardiology (ESC) guidelines (which are supported by the ERC) recommend synchronised

external cardioversion, even when the patient is haemodynamically stable, as haemodynamic

2555 deterioration may occur if monomorphic VT is not treated. Pharmacological treatment may be an

alternative if the risk of sedation/anaesthesia is high.⁶⁰⁰

2557

2558 For patients with polymorphic VT, precipitating factors should be sought and removed.⁶⁰⁷ In case

there is a prolongation of the QTc interval at baseline, IV Mg^{2+} and K^{+} infusion might help. Also

2560 increasing heart rate with isoproterenol or temporary pacing should be considered.⁶⁰⁸

In the case of recurrent ventricular arrhythmias, uncontrolled with non-invasive measures, consider
 reprogramming the ICD (if one is already in place), suppressing sympathetic activity (beta-blockers,
 sedation, autonomic modulation), starting mechanical circulatory support and referring the patient
 for catheter ablation.^{604,608}

2565

The ESC has published recent guidelines for the acute management of regular tachycardias without
an established diagnosis. The guidelines for treating regular narrow QRS (≤ 120 ms) and lifethreatening wide QRS (> 120 ms) tachycardias have been incorporated into the tachycardia
algorithm. The ESC Guidelines provide more detailed recommendations and evidence for treating
rhythms once a specific diagnosis of the rhythm has been made.¹¹⁴



- 2572 In a randomised trial involving haemodynamically stable patients with wide QRS-complex tachycardia 2573 of unknown aetiology, procainamide was associated with fewer major adverse cardiac events and a
- higher proportion of tachycardia termination within 40 min, compared with amiodarone.⁶⁰⁹ However,
- 2575 in many countries, procainamide is either unavailable or unlicensed.
- 2576

In critical low-perfusion conditions like hypovolaemic, cardiogenic, or distributive shock, sinus
tachycardia is a compensatory response to enhance blood flow to ischaemic tissues. In patients with
sinus tachycardia, the primary focus of treatment should be to address the underlying cause of sinus
tachycardia, such as through fluid administration. Rescuers should avoid trying to normalise the
heart rate through rate-control strategies, such as beta-blocker administration, as this may result in
patient deterioration or even haemodynamic collapse.

2583

In acute settings, rescuers should evaluate and manage underlying causes or triggering factors that initiate AF such as sepsis, fluid overload or cardiogenic shock.⁶¹⁰ The selection of rate versus rhythm control strategy and the appropriate drug will depend on the patient's characteristics, the presence of heart failure and the haemodynamic profile.⁶¹¹ In patients with AF and acute or worsening haemodynamic instability, electrical cardioversion is recommended.^{612,613}

2589 The rate of pharmacological cardioversion to sinus rhythm in patients with recent onset AF may be 2590 high, ranging from 50 to 95% in 10 min to 6 hours with the use of fast-acting drugs such as flecainide 2591 (200–300 mg orally once), propafenone (450–600 mg orally once), ibutilide (if patient < 60 kg, 0.6 mg 2592 kg⁻¹; if patient> 60 Kg, 1mg IV. The dose may be repeated after 10 min if the initial dose is ineffective) 2593 and vernakalant (3 mg kg⁻¹ IV over a 10-minute period. If AF persists 15 minutes after the completion 2594 of the initial infusion, a second dose of 2 mg kg⁻¹ may be administered over 10 minutes).⁶¹⁴ However, 2595 only amiodarone is indicated in patients with severe left ventricular hypertrophy, heart failure with 2596 reduced ejection fraction and coronary artery disease, which are common in patients presenting 2597 with cardiac arrest.^{615,616} The rate of successful restoration of sinus rhythm with amiodarone is 44% at 8-12 h to several days after IV administration.⁶¹⁵ In selected patients, rate control may be 2598 2599 sufficient. For this purpose, beta-blockers and diltiazem/verapamil are preferred over digoxin 2600 because of their rapid onset of action and effectiveness in patients with high sympathetic tone. For 2601 patients with left ventricular ejection fraction less than 40%, consider the smallest dose of beta-2602 blocker to achieve a heart rate of less than 110 min⁻¹ and add digoxin if necessary.⁶¹³



- 2603 Evidence for the treatment of patients with bradycardia was included in the American College of 2604 Cardiology (ACC)/American Heart Association (AHA)/Heart Rhythm Society (HRS) guidelines 2605 published in 2019, and these form the basis of the ERC bradycardia algorithm (Figure 10. Bradycardia 2606 algorithm).⁶⁰¹ If bradycardia is accompanied by adverse haemodynamic signs, atropine remains the 2607 first-choice drug.²¹⁴ Conversely, atropine is ineffective and may worsen the block in patients with high-degree atrioventricular block and wide QRS complexes.⁶⁰¹ When atropine is ineffective, second-2608 2609 line drugs include isoprenaline (5 mcg.min⁻¹ starting dose) and adrenaline (2 to 10 microgram.min⁻¹). 2610 For bradycardia in cases of sympathetic denervation, such as heart transplant or spinal cord injury, 2611 consider giving aminophylline (100-200 mg, slowly IV).⁶⁰¹ Atropine can cause a high-degree AV block or even sinus arrest in heart transplant patients.⁶¹⁷ 2612
- 2613

2614 Consider giving IV glucagon (one or more initial IV 3 to 10 mg IV push doses over 1–2 minutes. If

2615 effective, start a continuous infusion at 2-5 mg.h⁻¹) if beta-blockers or calcium channel blockers are

themselves a potential cause of the bradycardia.⁶¹⁸ Consider pacing in unstable patients, with

2617 symptomatic bradycardia refractory to drug therapy (see below).

2618

2619 [h3]Cardioversion

2620 Electrical cardioversion is the preferred treatment for tachycardia in the unstable patient displaying 2621 potentially life-threatening adverse haemodynamic signs (Figure 9. ERC Peri-arrest Tachyarrhythmia 2622 algorithm).^{600,602,604} The shock must be synchronised to occur with the R wave of the ECG: VF can be 2623 induced if a shock is delivered during the T wave which is a relative refractory portion of the cardiac 2624 cycle.⁶¹⁹ Synchronisation can be difficult in VT because of the wide-complex and variable forms of 2625 ventricular arrhythmia. Inspect the synchronisation marker carefully for consistent recognition of the 2626 R wave. If needed, choose another lead and/or adjust the amplitude. If synchronisation fails, give 2627 unsynchronised shocks to the unstable patient in VT to avoid prolonged delay in restoring sinus 2628 rhythm. Ventricular fibrillation (VF) or pulseless VT (pVT) require unsynchronised shocks. Conscious 2629 patients require anaesthesia or sedation before attempting synchronised cardioversion.

2630

2631 [h4] Cardioversion for atrial fibrillation

There is no single optimal position for external defibrillation electrodes. A meta-analysis of 10 RCTs comparing anterior-posterior with antero-lateral electrode positioning in AF showed no difference in sinus rhythm restoration.⁶²⁰ Applying active compression to the anterior defibrillation pad with anterior-posterior pads is associated with lower defibrillation thresholds, lower total energy delivery, fewer shocks for successful electric cardioversion, and higher success rates.²⁹⁸ More data are needed



2637 before specific recommendations can be made for optimal biphasic energy levels and different 2638 biphasic waveforms. Biphasic rectilinear and biphasic truncated exponential (BTE) waveforms show 2639 similar high efficacy in the elective cardioversion of atrial fibrillation.⁶²¹ An RCT showed that 2640 maximum fixed energy electrical cardioversion (360 J BTE) was more effective in achieving sinus rhythm one minute after cardioversion than an energy-escalating strategy.⁶²² There was no increase 2641 2642 in adverse events. An initial synchronised shock at maximum defibrillator output rather than an 2643 escalating approach is a reasonable strategy based on current data. In stable patients, follow 2644 appropriate guidelines on the need for anticoagulation before cardioversion to minimise stroke 2645 risk.613 2646 2647 [h4] Cardioversion for atrial flutter and paroxysmal supraventricular tachycardia 2648 Atrial flutter and paroxysmal supraventricular tachycardia (SVT) generally require less energy than atrial fibrillation for cardioversion.⁶²³ The ERC recommends giving an initial shock of 70-120 J. Give 2649 2650 subsequent shocks using stepwise increases in energy, as required.

2651

2652 [h4] Cardioversion for ventricular tachycardia with a pulse

The energy required for cardioversion of VT with pulse depends on the morphological characteristics and rate of the arrhythmia. Ventricular tachycardia with a pulse responds well to initial shocks with energy levels of 120-150 J. Consider stepwise increases if the first shock fails to achieve sinus rhythm.²⁵⁵

2657

2658 [h4] Pacing

2659 Consider pacing in unstable patients with symptomatic bradycardia refractory to drug therapy. 2660 Immediate pacing is indicated especially when the block is at or below the His-Purkinje level. Transvenous pacing should be established as soon as possible.^{601,603} Consider transthoracic 2661 2662 (transcutaneous) pacing as a bridge to transvenous pacing or when transvenous pacing is not readily 2663 available. Whenever a diagnosis of asystole is made, check the ECG carefully for the presence of P 2664 waves because this will likely respond to cardiac pacing. The use of epicardial wires to pace the 2665 myocardium following cardiac surgery is effective. Do not attempt pacing for asystole unless P waves 2666 are present; it does not increase short or long-term survival in-hospital or out-of- hospital.^{624,625} For 2667 haemodynamically unstable, conscious patients with bradyarrhythmia, for whom transthoracic 2668 pacing is not readily available, percussion pacing may be attempted as a bridge to electrical pacing,



- although its effectiveness has not been established.^{93,151,626} Give serial rhythmic blows with the
- 2670 closed fist over the left lower edge of the sternum to pace the heart at a physiological rate of 50-70
- 2671 min⁻¹. Transthoracic and percussion pacing can cause discomfort, so consider giving analgesic or
- 2672 sedative drugs to conscious patients.

| Drug | Indication | Timing | Dose/delivery | Notes |
|----------------|-------------|-------------------|-------------------|--------------------------------|
| /procedure | | | | |
| Vagal | Narrow QRS | | Blow into a 10 mL | Preferably in the supine |
| Manoeuvre | tachycardia | | syringe with | position with leg |
| | | | sufficient force | elevation ⁶²⁷ |
| | Wide QRS | | to move the | |
| | tachycardia | | plunger | |
| Adenosine | Narrow QRS | Recommended | Incremental, | If no evidence of pre- |
| | tachycardia | if vagal | starting at 6 mg, | excitation on resting |
| | | manoeuvres fail | followed by 12 | ECG ^{114,628} |
| | Wide QRS | | mg IV. | |
| | tachycardia | | An 18 mg dose | When using an 18 mg |
| | | | should then be | dose, consider the |
| | | | considered | tolerability/side effects in |
| | | | | the individual patient. |
| Verapamil or | Narrow QRS | Consider if vagal | Verapamil (0.075 | Should be avoided in |
| diltiazem | tachycardia | manoeuvres | – 0.15 mg/kg IV | patients with |
| | | and adenosine | [average 5–10 | haemodynamic |
| | | fail | mg] over 2 min) | instability, HF with |
| | | | Diltiazem [0.25 | reduced LV ejection |
| +.5 | | | mg/kg IV(average | fraction (<40)% ⁶²⁸ |
| | | | 20 mg) over 2 | |
| | | | min]. | |
| Beta-blockers | Narrow QRS | Consider if vagal | Esmolol (0.5 | More effective in |
| (IV esmolol or | tachycardia | manoeuvres | mg/kg IV bolus or | reducing the heart rate |
| metoprolol) | | and | 0.05–0.3 | than in terminating |
| | | adenosine fail | mg/kg/min | tachycardia |
| | | | infusion) | |



| | | | Metoprolol (2.5– | |
|--------------|--------------|-------------------|-------------------|-------------------------------|
| | | | 15 mg given IV in | |
| | | | 2.5 mg boluses), | |
| Procainamide | Wide QRS | Consider if vagal | 10-15 mg/kg IV | 114,609 |
| | tachycardia | manoeuvres | over 20 min | |
| | | and | | |
| | | adenosine fail | | |
| Amiodarone | Narrow and | Consider if vagal | 300 mg IV over | 114,600,629 |
| | wide QRS | manoeuvres | 10-60 min | |
| | tachycardia | and | according to | |
| | | adenosine fail | circumstances – | |
| | | | followed by | |
| | | | infusion of 900 | |
| | | | mg in 24h | |
| Magnesium | Polymorphic | ~ | 2 g IV over 10 | Magnesium can suppress |
| | wide QRS | | minutes. Can be | episodes of TdP without |
| | tachycardia | | repeated once if | necessarily shortening |
| | (torsades de | | necessary. | QT, even when serum |
| | pointes - | | | magnesium |
| | TdP) | | | concentration is |
| | | | | normal ^{600,604,630} |
| | | | | |

2673 **Table 6. Recommendations for the acute management of narrow and wide QRS tachycardia** Drugs

- 2674 may be administered via peripheral IV in an emergency. HF heart failure; LV left ventricular.
- 2675 [h2] Uncontrolled organ donation after circulatory death
- 2676 Less than half of patients achieve ROSC after resuscitation from cardiac arrest.^{46,631} When standard
- 2677 ALS fails to achieve ROSC, there are three broad treatment strategies:⁶³²
- Stop resuscitation and declare death.
- In selected patients, consider ECPR.
- In settings with an uncontrolled organ donation after circulatory death (uDCD) programme,
- 2681 continue CPR to preserve organ perfusion and transport the patient to a hospital with a
- 2682 designated uDCD pathway, following local protocols and legal requirements.



2683

This guideline focuses on uDCDs. These are defined as donors after unsuccessful resuscitation from unwitnessed (Maastricht category I) or witnessed (Maastricht category II) cardiac arrest, either inhospital or out-of-hospital.⁶³³ The ERC Post resuscitation care guidelines include guidance for organ donation pathways following brain death (donation after brain death: DBD) or controlled donation after circulatory death (cDCD, Maastricht category III donors) in patients who achieve ROSC or are treated with ECPR.¹⁴⁶ We acknowledge the ethical, cultural, and legislative issues that lead to variations in the use of uDCD.

2691

2692 There is a mismatch between the availability and demand across the world. Uncontrolled donation 2693 after circulatory death (uDCD) provides an opportunity for cardiac arrest victims in whom ROSC 2694 cannot be achieved to become organ donors. In Europe, uDCD is currently undertaken in Austria, Belgium, Israel, Italy, Lithuania, Portugal, Russia, and Spain.^{633,634} Organs that can be recovered 2695 2696 include kidneys, liver, pancreas and lungs. In 2025, an ILCOR systematic review showed that the rates 2697 of early dysfunction (primary non-function or delayed graft function) of kidneys and long-term 2698 function of livers recovered from uDCD were higher than in DBD or cDCD.¹ However, in most studies, 2699 long-term graft function was comparable.⁶³⁴⁻⁶³⁸ This difference may be partly due to the longer warm 2700 ischaemia time of uDCD compared with other organ recovery approaches.

2701

2702 There is no universal consensus on selection criteria for uDCD, and identifying a potential donor 2703 currently follows regional/national protocols. Criteria generally include age above those of consent 2704 (variable by nation, but usually above18 years) and not over 55-65 years, a no-flow time (the interval 2705 from emergency call or witnessed arrest to CPR start) within 15 minutes, and a total warm ischaemia 2706 time (the interval between cardiac arrest and the start of organ preservation) not longer than 150 2707 minutes. Exclusion criteria generally include trauma, homicide, or suicide as a cause of arrest and 2708 comorbidities such as cancer, sepsis, and, according to local programmes and the targeted organ to 2709 transplant, kidney and liver disease.⁶³⁹ In a 2024 study conducted on the Parisian registry, of 19,976 2710 adult patients who were resuscitated from 2011 to 2020, 12,890 (65%) had no ROSC, 9461 (47%) met 2711 termination of resuscitation (TOR) criteria, and 6720 (52%) could be considered for uDCD kidney 2712 donors.631

2713

Uncontrolled donation after circulatory death is a time-critical, resource-intensive, complex and
 ethically challenging process.⁶⁴⁰ Following the termination of resuscitation efforts, a 'no-touch'
 period is observed to rule out the possibility of autoresuscitation, i.e., ROSC after CPR has been



- 2717 stopped or life-sustaining measures have been withdrawn in the intensive care unit. In most
- 2718 countries where uDCD is practised, the prescribed duration of the no-touch period is five minutes⁶⁴¹,
- but some require 20 minutes.⁶⁴² An updated systematic review on autoresuscitation identified seven
- 2720 observational studies, of which one investigated OHCA.⁶⁴³ Among 840 patients whose resuscitation
- 2721 measures were terminated on site, the study reported five cases of ROSC occurring 3 to 8 minutes
- after cessation of CPR. Three of these five patients died on scene, while two died in hospital, one
- within 1.5h and the other within 26h.
- 2724 After the no-touch period, organ preservation procedures are started and continued until consent for
- 2725 organ recovery is ascertained.⁶⁴⁴⁻⁶⁴⁶ Obtaining consent from a surrogate decision maker (e.g., a family
- 2726 member) is particularly challenging for uDCD, given the unexpected nature of the arrest, the
- 2727 considerable time pressure, and the difficult environment of an emergency department. Establishing
- 2728 clear local protocols and legislative and societal acceptance is crucial for this process.⁶⁴⁷ Previous
- 2729 consent registered on a donor card or a public registry is invaluable and must be rapidly retrieved.
- 2730 Adopting an opt-out system with implied consent to donation is an effective strategy to improve the
- 2731 rates of organ donation if the legal and cultural context allows.⁶⁴⁸ The ERC Guidelines 2025: Ethics in
- 2732 Resuscitation includes further details on these issues.³
- 2733 For abdominal organs, organ preservation typically uses extracorporeal circulation with membrane
- 2734 oxygenation via a femoro-femoral bypass.⁶⁴⁷ Catheters with balloons are used to limit circulation to

the abdominal cavity.⁶⁴⁹

- 2736
- 2737



2738 [h1] Conflicts of interest

| 2739 | |
|------|--|
| 2740 | JS declares payment from Elsevier for his role as Editor of Resuscitation. He reports research |
| 2741 | funding from the UK National Institute for Health Research in relation to the AIRWAYS3 trial. |
| 2742 | |
| 2743 | BWB declares that he is an Editor of the Journal Notfall + Rettungsmedizin, Co-Editor of the Brazilian |
| 2744 | Journal of Anesthesiology, and fees for lectures/advice from Forum für medizinische Fortbildung |
| 2745 | (FomF), BD Deutschland GmbH, Boehringer Ingelheim Deutschland GmbH, Doccla Deutschland |
| 2746 | GmbH. He is an Editorial Board member of Resuscitation. |
| 2747 | |
| 2748 | KC reports funding from the National Institute of Health and Care Research and Resuscitation Council |
| 2749 | UK (paid to institution) for cardiac arrest research, and role as an Editorial Board member of |
| 2750 | Resuscitation. |
| 2751 | |
| 2752 | AM declares payment from Elsevier for her role as Young ERC Editor of Resuscitation Plus. |
| 2753 | |
| 2754 | JPN declares payment from Elsevier for his role as Editor-in-Chief of Resuscitation. He reports |
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| 2760 | Pharmaceuticals GmbH Member of the AOP Health Group |
| 2761 | |
| 2762 | TS declares payment from Elsevier for his role as Social Media Editor of Resuscitation and |
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| 2764 | |
| 2765 | PC, CDD, CS, MBS declare that they are Editorial Board members of Resuscitation. |
| 2766 | |
| 2767 | JY reports funding from the National Institute of Health and Care Research and Resuscitation Council |
| 2768 | UK (paid to institution) for cardiac arrest research, and role as an editorial board member of |
| 2769 | Resuscitation Plus and Perioperative Medicine. |
| 2770 | |
| 2771 | FCJ, DC, GC, SD'A, JEE, MJH, PP, HP, FV declare no conflicts of interest. |
| | |

European Resuscitation Council Science Park I Galileilaan 11 ISALA – 3.12b 2845 Niel, Belgium www.erc.edu 

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| 2787 | [h1] Figure legends |
|------|--|
| 2788 | |
| 2789 | Figure 1. Advanced Life Support – key messages |
| 2790 | |
| 2791 | Figure 2. In-hospital resuscitation algorithm ABCDE airway breathing circulation disability exposure; |
| 2792 | AED automated external defibrillator; ALS advanced life support; BP blood pressure; CPR |
| 2793 | cardiopulmonary resuscitation; ETCO ₂ end-tidal carbon dioxide; IV intravenous; SBAR situation, |
| 2794 | background, assessment, recommendation; SpO $_2$ oxygen saturation measured with pulse oximetry. |
| 2795 | |
| 2796 | Figure 3. Advanced Life Support algorithm ABCDE airway, breathing, circulation, disability, exposure |
| 2797 | CPR cardiopulmonary resuscitation; ECG electrocardiogram; EMS emergency medical system; io |
| 2798 | intraosseous; IV intravenous; PEA pulseless electrical activity; PaCO $_2$ arterial partial pressure of |
| 2799 | carbon dioxide; ROSC return of spontaneous circulation; SpO_2 oxygen saturation measured with |
| 2800 | pulse oximetry; VF ventricular fibrillation; VT ventricular tachycardia. |
| 2801 | |
| 2802 | Figure 4. Advanced Life Support – example scenarios for timing of shocks and drugs [not every |
| 2803 | possible scenario is included] PEA pulseless electrical activity; VF ventricular fibrillation; VT |
| 2804 | ventricular tachycardia. |
| 2805 | |
| 2806 | Figure 5. Correct antero-lateral pad placement for defibrillation |
| 2807 | |
| 2808 | Figure 6. Antero-posterior pad placement |
| 2809 | |
| 2810 | Figure 7. Position of a transvenous (standard) Implantable Cardioverter Defibrillator (ICD) |
| 2811 | |
| 2812 | Figure 8. Position of a Subcutaneous Implantable Cardioverter Defibrillator |
| 2813 | |
| 2814 | Figure 9. Peri-arrest Tachyarrhythmia Algorithm ABCDE airway, breathing, circulation, disability, |
| 2815 | exposure; BP blood pressure; DC direct current; ECG electrocardiogram; EF ejection fraction, |
| 2816 | IV intravenous; ROSC return of spontaneous circulation; SpO ₂ oxygen saturation measured with pulse |
| 2817 | oximetry; VT ventricular tachycardia. |
| 2818 | |
| 2819 | Figure 10. Bradycardia Algorithm ABCDE airway, breathing, circulation, disability, exposure |



| 2820 | BP blood pressure; ECG electrocardiogram; IV intravenous; SpO $_2$ oxygen saturation measured with |
|------|---|
| 2821 | pulse oximetry. |
| 2822 | |
| 2823 | [h1] Table legends |
| 2824 | |
| 2825 | Table 1. What's new in Guidelines 2025 for Advanced Life Support? |
| 2826 | |
| 2827 | Table 2. Causes of sudden cardiac arrest (SCA) Adapted from Kandala ⁶⁰ and Winkel. ⁶¹ |
| 2828 | |
| 2829 | Table 3. Key points from the European Society of Cardiology guidelines for the treatment of |
| 2830 | ventricular arrhythmias and sudden cardiac death (Adapted from Könemann, 202373) |
| 2831 | |
| 2832 | Table 4. High risk features suggesting a serious condition in patients with syncope at initial |
| 2833 | evaluation in the emergency department Adapted from Brignole 2018.65 ECG electrocardiogram; |
| 2834 | ICD implantable cardioverter defibrillator; LVEF left ventricular ejection fraction; SCA sudden cardiac |
| 2835 | arrest; VT ventricular tachycardia. |
| 2836 | |
| 2837 | Table 5. Patient and resuscitation factors affecting outcome from OHCA. Adapted from Kandala |
| 2838 | 2017. ⁶⁰ AED automated external defibrillator; CPR cardiopulmonary resuscitation. |
| 2839 | Table 6. Recommendations for the acute management of narrow and wide QRS tachycardia Drugs |
| 2840 | may be administered via peripheral IV in an emergency. HF heart failure; LV left ventricular. |
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2844 [h1] References

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