

## **Sudden Cardiac Death where death occurs in ICU** **Operating Guideline**

### **Introduction**

Some deaths resulting from out of hospital cardiac arrest (OOHCA) are due to inherited cardiac causes. The 2022 ESC Guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death<sup>1</sup> contain some applicable information but are not prescriptive in follow up requirements. However, based on these international guidelines, local pathways, and local protocols the following NHS Lothian guideline has been developed in conjunction with colleagues in Clinical genetics, Cardiology and ICU specifically for patients admitted to ICU with a sudden cardiac death syndrome where there is a strong suspicion of an inherited cardiac condition (ICC). This guideline may be used by other ICU's in South East Scotland within the regional ICC network (NHS Borders, Forth Valley and Fife)

Patients discharged to cardiology from ICU will undergo appropriate screening for underlying cardiac disease including a possible inherited cardiac condition.

This guideline applies to patients who do not survive beyond ICU.

### **Which patients does this guideline apply to?**

OOHCA Patients meeting the following criteria:

- Deaths in general intensive care unit (ICU)
- Patients under 60
- Unlikely to be due to an acute coronary syndrome
- Non-cardiac precipitant excluded
- Sufficient information to complete the death certificate (MCCD)\*
- Cardiology review/advice confirms suspicion of ICC

\*Please be as specific as possible for the underlying cause of death on the MCCD. A discussion with the cardiology team is advised. Some examples of underlying causes of death on the MCCD that might trigger the application of the guideline may include dilated cardiomyopathy, hypertrophic cardiomyopathy or arrhythmogenic right ventricular cardiomyopathy, the channelopathies (Brugada syndrome, Long QT syndrome, Polymorphic catecholamine induced VT), and less commonly familial aortopathy (acute dissection) which typically end up on cardiac ICU. In many cases this level of specificity may not be possible without further post-mortem investigation. In these cases, less specific terms, provided they appear in ICD11, are permitted.

### **Which patients does this guideline not apply to?**

- Deaths following OOHCA where there is clear rationale to refer to the Procurator Fiscal, typical examples being deaths associated with suspected illicit drug use, deaths following trauma, suicide
- Deaths following OOHCA where there is insufficient information to complete the MCCD
- Both of these cases should be referred to the Procurator Fiscal.
  - Please ring the duty biochemist to store the closest available samples to the time of admission and send sample: Urine, drugs of abuse (all sites).
- For the full list of reasons to refer to Procurator Fiscal please see this [link](#).
- For patients with OOHCA due to coronary heart disease under 60 years, who die in ICU, first degree relatives should be advised to attend their GP for cardiovascular risk assessment.

Family contact: The ICU Team should identify and record contact details of at least one family member who agrees to act as a liaison between the family and the Clinical Genetics team for ongoing communications regarding screening for an inherited cardiac condition. This family member needs to consent for their contact details to be passed to Clinical Genetics. These details should be included in the email sent to Clinical Genetics referring the patient (index case) to the Cardiac Genetics MDT.

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## For included patients

If patient meets criteria set out in *NHSL Critical Care Sudden Cardiac Death Where Death Occurs on ICU*. Store DNA sample by sending EDTA blood to Molecular Genetics Lab Western General Hospital marking the sample for storage using this [link](#). See appendix for completion instructions. Obtain "Urine drugs of abuse- all sites" if not already done.

**Proceeding to organ donation?** If suspicion of sudden cardiac death do not donate cardiac tissue

No

Yes

Approach relatives for hospital PM. Use forms available from this intranet [link](#). MCCD required\*.

Approach relatives for limited hospital PM of remaining tissue. Use forms available from this intranet [link](#). MCCD required\*.

**Hospital PM accepted and performed.**  
Non cardiac cause of death identified.

**Hospital PM accepted and performed.**  
Is negative or suggests cardiomyopathy.

**Hospital PM declined.**  
Offer consideration of genetics review for limited investigations.

**MCCD may be amended by pathologist. Inform family. Further action as appropriate.**

Refer to Cardiac Genetics MDT by secure email  
[wgh.clinicalgenetics@nhslothian.scot.nhs.uk](mailto:wgh.clinicalgenetics@nhslothian.scot.nhs.uk) include

1. Patient's CHI
2. Date of death
3. Copy of discharge letter.
4. Details of next of kin/most appropriate contact relative:

Name  
CHI or DOB  
Address  
Telephone

Cardiac MDT meets every month. Genetic testing takes at least 8 weeks to be completed.

\* When completing the PM request form, please endeavour to tick section 2A & 2B: Heart. It is unlikely the pathologist will retain the heart beyond a day or two before returning it to the body but this provides flexibility

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## **INHERITED CARDIAC CONDITIONS/GENETICS MDT:**

This is a regional meeting for South - East Scotland held at 0900h on the last Friday of each month in the Cardiology department at RIE and can be attended on Teams.

The responsible ICU consultant/doctor will be invited to attend the ICC MDT meeting when the patient is scheduled for discussion. Attendance is not mandatory but would be extremely valuable in providing clinical context. The MDT will review and discuss clinical details and will decide if gene testing is appropriate or not. The MDT will then decide which gene panel analysis should be used. The Clinical Genetics team will contact the family, via the liaison contact, explore family history and consent for testing if appropriate.

The outcome of the MDT and results of any investigations will be reported back to the family, and the patient's GP by the clinical genetics team. Further necessary genotypic or phenotypic screening of family members will be coordinated through the ICC MDT.

## **References**

1. 2022 ESC Guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death | European Heart Journal | Oxford Academic.

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<p><b>PATIENT DETAILS</b> (<i>printed label preferred</i>)</p> <p style="text-align: right;">Sex M / F</p>  <p><i>Required: Name, date of birth, CHI or 1<sup>st</sup> line home address and postcode</i></p> <hr/> <p>I have discussed genomic/ genetic testing with my health professional and I understand that:</p> <ul style="list-style-type: none"> <li>1) <b>Family implications</b> The results of my test may have implications for other members of my family. I acknowledge that my results may sometimes be used to inform the appropriate health care of others. This could be done in discussion with me, or in such a way that I am not personally identified in this process.</li> <li>2) <b>Uncertainty</b> The results of my test <i>may</i> reveal genetic variation whose significance is not yet known. Deciding whether such variation is significant may require sharing of information about me including (inter)national comparisons with variation in others. I acknowledge that interpretation of my results may change over time as such evidence is gathered.</li> <li>3) <b>Unexpected information</b> The results of my test <i>may</i> reveal a chance of a disease in the future, and nothing to do with why I am having this test. These may be found by chance, whilst focussing on the reason for my test, and I may then need further tests to understand their significance. If these additional findings are to be looked for, I will be given more information about this.</li> <li>4) <b>DNA storage</b> Normal laboratory practice is to store the DNA extracted from my sample even after the current testing is complete. My sample might be used as a 'quality control' for other testing, for example, that of family members.</li> <li>5) <b>Data storage</b> Data from my genetic test will be stored to allow for possible future interpretations.</li> <li>6) <b>Health records</b> Results from my genomic test and my test report will be part of my Patient Health Record.</li> </ul> <p>Note of other specific issues discussed (e.g. referral to particular research programmes, insurance):</p> <hr/> <p>*I agree to genetic or genomic investigations for</p>	<p><b>REFERRER DETAILS</b></p> <p>Name: _____</p> <p>Report to: <div style="border: 1px solid black; padding: 2px; width: fit-content; margin-top: 5px;"><b>ICU consultant details</b></div></p> <p><input type="checkbox"/> Email report to: _____ (Lab preference)</p> <p>Other contact details: _____</p> <p><b>DISEASE / CONDITION</b></p> <div style="border: 1px solid black; padding: 5px; text-align: center; margin-top: 10px;"><b>Sudden cardiac death</b></div> <p><b>TESTS REQUESTED</b> (SPECIFY GENE / VARIANT IF KNOWN)</p> <div style="border: 1px solid black; padding: 5px; text-align: center; margin-top: 10px;"><b>Storage, for discussion at ICC MDT</b></div> <p><b>CLINICAL DETAILS</b></p> <div style="border: 1px solid black; padding: 5px; text-align: center; margin-top: 10px;"><b>See email referral to ICC MDT</b></div> <p>Discussed with Clinical Genetics? Y / N <div style="border: 1px solid black; padding: 2px; width: fit-content; float: right;"><b>Debbie Mackin/Philip Greene</b></div></p> <p>If yes, name Clinical Genetics contact: _____</p> <p><b>SAMPLE DETAILS</b></p> <p>Taken by: Name (print) _____</p> <p>Date taken: ____/____/____ Signature _____</p> <p>High risk (see over) Y / N <input type="checkbox"/> URGENT (<i>phone lab to discuss</i>)</p> <p><input checked="" type="checkbox"/> Blood in Potassium EDTA (KE) <i>All DNA tests including microarray and QF-PCR</i></p> <p><input type="checkbox"/> Blood in Lithium Heparin <i>Cell culture-based tests – G-banded karyotyping</i></p> <p><input type="checkbox"/> DNA, mouthwash, tissue etc. (<i>Please state</i>): _____</p> <p>Arrange for immediate transport to the laboratory by van service or first class post. If sending later blood specimens should be refrigerated. <b>(DO NOT FREEZE)</b></p> <p><b>LAB USE ONLY</b></p>
<p>.....</p> <p><b>Patient/Parent Signature</b> <div style="border: 1px solid black; padding: 2px; width: fit-content; margin-top: 5px;"><b>Relative's signature</b></div> DATE ____/____/____</p> <p>Discussion undertaken by .....</p> <p>(Clinician's name, designation and signature) .....</p> <p><small>*Insert e.g. to investigate the cause of my child's developmental delay/ family history of cancer/ heart disease etc.</small></p>	<p><b>Incomplete or illegible forms, or use of incorrect blood tubes, will cause delay or rejection of samples.</b></p> <p>Authority for issue: Austin Diamond      GENE-WM275      v8 (Issue date 21.03.2019)</p>

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