

Knowledge, understanding and learning to improve young lives

National Child Mortality Database Professional Advisory Group

Terms of Reference

Document Version: 1.8

VERSION HISTORY

Date	Document Version	Document Revision History	Document Author/Reviser
22 nd May 2018	1.0	Initial draft	Vicky Sleap
9 th July 2018	1.1	Revised draft following comments for collaborators	Vicky Sleap
31st January 2019	1.2	Further revised draft prior to publication on website	Sylvia Stoianova / Vicky Sleap
25 th June 2019	1.3	Further revision and final for publication on the NCMD website	Sylvia Stoianova
3 rd August 2020	1.4	Additional member added	Sylvia Stoianova
29 th July 2022	1.5	Members/organisations added and members/organisations removed	Ghazala Jones
4 th October 2023	1.6	Members/organisations added and members/organisations removed	Gaja Wright
29 th February 2024	1.7	Members/organisations added and members/organisations removed	Gaja Wright
25 rd April	1.8	Additional members added	Gaja Wright

Chair: Dr James Fraser

Members:

Organisation	Name
Association for Child & Adolescent Mental	Barbara Maughan
Health ¹	
Association of Child Death Review Professionals	Joanna Garstang
(ACDRP)	
Association of Paediatric Palliative Medicine	Emily Harrop
(APPM)	Anna-Karenia Anderson
British Association for Community Child Health ¹	Pending confirmation of representative
Royal College of Emergency Medicine	Sian Thomas
British Association of Paediatric Surgeons	Pending confirmation of representative
British Association of Perinatal Medicine ¹	Stephen Wardle
British Association of Social Workers	Luke Geoghegan
Children and Young People's Cancer Clinical	Rachael Hough
Reference Group	
Children's Cancer & Leukaemia Group 1	Jesicca Bate
Children and Young People's Cancer Clinical	Julia Chisholm
Reference Group	
Children's Commissioner for England	Harriet Waldegrave
College of Paramedics	Will Broughton
Coroners' Society of England and Wales	Pending confirmation of representative
National Director, Children and Young People's	Prathiba Chitsabesan
Mental Health Services (CYPMHS) Clinical	
Reference Group (CRG)	
Children and Young People's Mental Health	Tina Irani
Services (CYPMHS) Clinical Reference Group	
(CRG)	
Department for Education	Melanie Cawthorn
English Ambulance Services via Association of	Nicola Albutt, Lucy Gascoigne
Ambulance Chief Executives (AACE)	
Institute of Criminal Justice Studies, University of	John Fox
Portsmouth	
Institute of Health Visiting	Philippa Bishop
UK Preterm Clinical Network and NIHR Research	Nigel Simpson
Delivery Network (Reproductive Health and	
Childbirth national specialty group)	
National Association of Head Teachers	Pending confirmation of representative
National Genetic Risk Expert Group	Nasaim Khan
National Medical Examiner for England and	Alan Fletcher Pending confirmation
Wales	
National Network of Designated Healthcare	Peter Green
Professionals	
National Police Chiefs Council	Eamonn Bridger
NHS England	Kay Pope

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¹ Key Clinical groups representing the major causes and places of death

Office of the Chief Coroner	Cathy Yallop	
Office for Health Improvement and Disparities	Helen Duncan and Marilena Korkodilos	
Office for National Statistics	Pending confirmation of representative	
Paediatric Critical Care Society	Miriam Fine Goulden	
PICANet	Richard Feltbower	
	Sarah Seaton	
Police National Child Death Working Group	Eamonn Bridger (see above)	
Health & Social Care, Northern Ireland (HSNI)	Sinead Magill	
Quality, Patient Safety and Maternity,	Karen Todd	
Department of Health of Social Care		
Royal College of Anaesthetists	John Pappachan	
Royal College of Emergency Medicine	Damian Roland	
Royal College of GPs	Janice Allister and	
	Bryony Kendall	
Royal College of Midwives	Mervi Jokinen	
Royal College of Nursing (England)	Pending confirmation of representative	
Royal College of Obstetrics & Gynaecology	Pending confirmation of representative	
Royal College of Paediatrics and Child Health	Ingrid Wolfe	
Royal College of Pathologists	Gauri Batra and Prof Marta Cohen	
Scottish Child Death Review Steering Group	Nanisa Feilden	
Shooting Stars Children's Hospice	Jo Cohen	
Trauma Audit & Research Network ¹	Antoinette Edwards	
University of the West of England, Health and	Dawn Odd	
Social Sciences Department		
Welsh Child Death Review Programme	Rosalind Reilly	
GP/ Public Health Medical Associate	Nikesh Parekh	

Background

The National Child Mortality Database (NCMD) programme has been commissioned to collect data, via the child death review process, on all child deaths in England. The NCMD programme team quality assures and analyses all data to provide timely, accurate and informative outputs to relevant stakeholders in order to facilitate learning and actions to reduce modifiable child mortality.

The NCMD programme is funded by NHS England and commissioned by the Healthcare Quality Improvement Partnership (HQIP) on behalf of NHS England. It is delivered by the University of Bristol in collaboration with UCL Partners, the Anna Freud Centre and the NCMD system software provider company QES Ltd (Quality Education Solutions). Parent, patient and public involvement (PPPI) in this programme is enabled through the Lullaby Trust, Sands (Stillbirth and neonatal death charity) and Child Bereavement UK (CBUK). The programme is currently funded until July 2026 with a possibility for an extended funding for a further 2 years until July 2028.

The overalls aim of the programme are as follows:

- Capture, analyse and disseminate appropriate data and learning from 100% of child death reviews
- Study and analyse the demographics, patterns, causes and associated risk factors of child mortality in England

- Provide information in a timely fashion that can be used by relevant NHS, health and social care organisations to make policy decisions and interventions with the aim of reducing the number of preventable child deaths
- Produce routine aggregate data tables and benchmarked child mortality data at the local level to support local response by NHS regions and Integrated Care Boards (ICBs)
- Provide information in a timely fashion that can be used by organisations in wider society to make policy decisions and interventions with the aim of reducing the number of preventable child deaths
- Support cross-system work to reduce the number of preventable child deaths, particularly in relation to groups with health inequalities, e.g. those who experience factors which impact on health determinants such as disability, socio-economic status, deprivation and poor housing
- Identify and issue relevant alerts for:
 - modifiable factors in child deaths, both related to the NHS and in wider society
 - clusters of deaths
 - newly emerging causes and risk factors for death
- Produce reports, briefings, present data, share case studies and make recommendations to relevant agencies for implementation
- Drive improvements in the quality of child death reviews, e.g. through training and support for the professionals involved

The Professional Advisory Group provides a formal interface between NCMD and the professional community. Its meetings provide a forum in which organisations can:

- a) raise practical issues with regards to the operation of NCMD
- b) contribute to discussions around themed reports and other data arising from analysis of its database
- c) inform the long term development of its program.

Its membership includes organisations that represent the cohort of children who die and who on a national level can affect policy change.

Purpose / Role of the Group

The members of the group are asked to:

- 1. Provide a line of communication between the NCMD programme team and their organisation and its members (and vice versa).
- 2. Advise on how best to promote NCMD activities and outputs arising from its programmes of work.
- 3. Advise on how best to engage the members of their organisation in the work of the NCMD programme.
 - Advise on prioritisation and content of NCMD reports.
- 4. Provide specialist support/advice as required.
- 5. Attend all meetings (or send a deputy) and be familiar with all circulated content for meetings.

Meetings and Quoracy

1. Meetings will be held 2 times a year.

- 2. Attendance may be in person or by teleconference facility (e.g. Microsoft Teams).
- 3. Communication between the NCMD programme team and the membership of the Professional Advisory Group will be via email.
- 4. All papers / agenda items will be forwarded at least 5 working days in advance of the meeting.
- 5. All meetings will be minuted. Travel expenses (standard class travel) will be paid.
- 6. A minimum number of members, i.e. an appropriate quorum and organisations' representatives will be required for decision-making purposes. The minimum number required to be quorate will be 15 members.

Appendix A NCMD Governance Structure

NCMD National Child Mortality Database **Management and Governance** Structure for the NCMD **Programme NHS England UOB Dean of UOB Pro UOB Deputy CYP Policy Team Health Sciences** Vice-Chancellor Vice-Chancellor **UOB Head of HQIP Bristol Medical** School NCMD **NCMD Project** (UOB) (Karen Luyt: Responsible Officer) Monitoring **Steering Group** Group **NCMD PPPI NCMD Project** Professional Stakeholder Group Operational Advisory Group (PAG) (Includes Tier 1, 2 Group and 3 Membership)