World Journal of **Pediatric Surgery**

Congenital diaphragmatic hernia survival in an English regional ECMO center

Elizabeth O'Connor.¹ Rvo Tamura.¹ Therese Hannon.² Sundeep Harigopal ⁰.³ Bruce Jaffrav 💿 1

ABSTRACT

To cite: O'Connor E, Tamura R, Hannon T, et al. Congenital diaphragmatic hernia survival in an English regional ECMO center. World Jnl Ped Surgerv 2023;6:e000506. doi:10.1136/ wjps-2022-000506

Received 5 October 2022 Accepted 22 March 2023

Check for updates

C Author(s) (or their employer(s)) 2023. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

¹Paediatric surgery, The Great North Children's Hospital, Newcastle upon Tyne, Tyne & Wear, UK ²Fetal medicine and obstetrics, Royal Victoria Infirmary, Newcastle upon Tyne, Tyne & Wear, UK ³Neonatal medicine, Royal Victoria Infirmary, Newcastle upon Tyne, Tyne & Wear, UK

Correspondence to

Dr Bruce Jaffray; bruce.jaffray@ nhs.net

Introduction Congenital diaphragmatic hernia (CDH) remains a cause of neonatal death. Our aims are to describe contemporary rates of survival and the variables associated with this outcome, contrasting these with our study of two decades earlier and recent reports.

Materials and methods A retrospective review of all infants diagnosed in a regional center between January 2000 and December 2020 was performed. The outcome of interest was survival. Possible explanatory variables included side of defect, use of complex ventilatory or hemodynamic strategies (inhaled nitric oxide (iNO), highfrequency oscillatory ventilation (HFOV), extracorporeal membrane oxygenation (ECMO), and Prostin), presence of antenatal diagnosis, associated anomalies, birth weight, and gestation. Temporal changes were studied by measuring outcomes in each of four consecutive 63-month periods.

Results A total of 225 cases were diagnosed. Survival was 60% (134 of 225). Postnatal survival was 68% (134 of 198 liveborn), and postrepair survival was 84% (134 of 159 who survived to repair). Diagnosis was made antenatally in 66% of cases. Variables associated with mortality were the need for complex ventilatory strategies (iNO, HFOV, Prostin, and ECMO), antenatal diagnosis, rightsided defects, use of patch repair, associated anomalies, birth weight, and gestation. Survival has improved from our report of a prior decade and did not vary during the study period. Postnatal survival has improved despite fewer terminations. On multivariate analysis, the need for complex ventilation was the strongest predictor of death (OR=50, 95% CI 13 to 224, p<0.0001), and associated anomalies ceased to be predictive.

Conclusions Survival has improved from our earlier report, despite reduced numbers of terminations. This may be related to increased use of complex ventilatory strategies.

INTRODUCTION

In 2003, we published our institutional diaphragmatic hernia (congenital diaphragmatic hernia (CDH)) survival study.¹ In contrast to most studies of that era, we described no improvement in survival over the preceding 10 years, perhaps because we were able to identify deaths occurring in peripheral hospitals and terminations.

We reported total survival of 32% and liveborn survival of 55%. However, we did

WHAT IS ALREADY KNOWN ON THIS TOPIC

 \Rightarrow Survival for babies is dependent on severity of pulmonary hypoplasia.

WHAT THIS STUDY ADDS

 \Rightarrow Survival rates are improving despite fewer terminations. Survival without fetoscopic tracheal occlusion (FETO) appears similar to FETO centers. Extracorporeal membrane oxygenation does not significantly affect outcomes.

HOW THIS STUDY MIGHT AFFECT RESEARCH. PRACTICS OR POLICY

- \Rightarrow Survival appears to improve as a result of small incremental improvements.
- Outcomes of FETO should be compared to this benchmark.

not analyze the effect of additional cardiac anomalies nor did we attempt to distinguish differing postnatal times of death.

The intervening decades have seen several important innovations in the treatment of CDH. The prognostic utility of the liver position and the fetal lung:head ratio measured on antenatal scan was recognized² and subsequently refined.³ Grading of the risk of mortality has allowed the deployment of novel interventions, such as fetoscopic tracheal occlusion (FETO), with reported improvements in outcomes.⁴ However, the utility of such measurements remains dependent on antenatal detection, which may still be relatively low.⁵ Extracorporeal membrane oxygenation (ECMO) seems to offer promise,⁶ but its precise role remains controversial,⁷ and concerns about neurological morbidity persist.⁸ Other therapies that have been deployed in the treatment of CDH since our original study include inhaled nitric oxide (iNO)⁹ and high-frequency oscillatory ventilation (HFOV).¹⁰

In the context of these evolving therapies, we were interested in comparing survival in our institution in the last 20 years, contrasting these with our earlier report

BMJ

and contemporary descriptions of outcomes for this condition.

METHODS

Patients and data collection

We recorded all CDH cases between January 2000 and December 2020. Cases were identified from institutional records, including the regional antenatal service and a dedicated database of CDH repairs. The outcome of interest was survival. We categorized deaths as (1) elective terminations, (2) intrauterine death, (3) postnatal, (4) death without surgical repair, and (5) postnatal death following surgical repair.

Possible explanatory variables were sex, gestation, birth weight, antenatal diagnosis, side of defect, and congenital anomalies. Cardiac anomalies were categorized as (1) atrial septal defect (ASD) / ventricular septal defect (VSD), (2) patent ductus arteriosus (PDA), or (3) complex. Surgical repair was (1) sutured or (2) prosthetic patch. We did not document defect size during this study, but the use of patches will be a surrogate marker for larger defects. The timing of repair was early (<2 months of age) or late (>2 months of age). We chose this cutoff, recognizing that a proportion of CDH cases manifest no respiratory distress at birth and present later, typically with bowel-related symptoms. We wished to analyze this cohort separately from the cohort presenting with immediate respiratory symptoms.

Complex ventilatory/hemodynamic strategies were (1) iNO, (2) HFOV, (3) ECMO, and (4) prostaglandin E (Prostin, Pfizer).

Statistical analysis

We examined temporal patterns by dividing the period of 21 years into four consecutive 63-month intervals, performing a χ^2 goodness-of-fit test for each outcome. We analyzed the association between survival and explanatory variables using a χ^2 test.

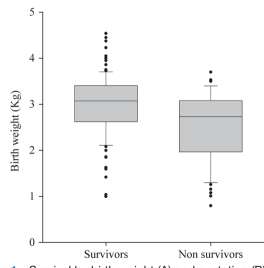


Figure 1 Survival by birth weight (A) and gestation (B).

We obtained the OR for survival with predictors using logistic regression. Variables that were significantly associated with survival were then combined in a multivariate model using a forward entry method. Because of the strong collinearity of iNO, HFOV, and Prostin use, we only used iNO in the multivariate model. We hypothesized that there may be differing effects on infants who died prior to repair and therefore performed the analysis twice, with a separate analysis for those who survived to undergo repair.

All analyses were performed using SPSS V.24. Graphs were constructed with SigmaPlot V.13 (Systat, California, USA).

RESULTS

Of 225 cases, 148 (66%) were identified antenatally. After 26 elective terminations and 1 intrauterine death, 198 were liveborn. Birth weight and gestation were greater among survivors (2.9 kg (0.6) vs 2.5 kg (0.8), p<0.0001; 38.0 vs 37.5 weeks, p=0.001) (figure 1). Of the 198 cases, 127 (64%) were male and 154 (78%) had a left-sided defect.

Thirty-nine of 198 (20%) died due to respiratory failure before repair. One child underwent FETO but was delivered less than 48 hours later and survived with no complex ventilation.

Survival

The overall survival was 134 of 225 (60%). Liveborn survival was 134 of 198 (68%), with no sex difference, 85 of 127 (67%) for male cases vs 49 of 71 (66%) for female cases. Of 91 deaths, 27 (30%) occurred antenatally; 39 (43%) occurred postnatally; and 25 (27%) occurred after repair.

Surgical intervention group

Of 159 CDH cases who were repaired, 139 (87%) underwent early repair; 123 (77%) had sutured repair; 36

44 42-40-38-36-34-32-30-28-26-Survivors Non survivors

	Time period/observed frequency						
Variable	1	2	3	4	Expected frequency	χ ²	P value
Number of cases	52	57	66	50	56	2.7	0.4
Antenatally diagnosed	29	38	46	35	37	4	0.3
Terminations	2	4	15	5	6	15.5	<0.0001
Live birth	52	55	53	48	52	0.5	0.9
Surgical repair	37	42	41	39	40	0.4	0.9
Alive	30	40	35	29	34	2	0.5
HFOV	15	19	26	19	22	3	0.4
iNO	21	19	29	29	22	3	0.4
Prostin	2	15	23	18	14	16	0.001
ECMO	1	0	5	5	5	4	0.2
Significant cardiac anomaly	10	12	14	14	13	1.1	0.7
Isolated non-cardiac anomaly	4	5	4	2	4	1.5	0.6

 Table 1
 Temporal patterns and frequency of outcomes and explanatory variables over four consecutive time intervals, each of 63 months

ECMO, extracorporeal membrane oxygenation; HFOV, high-frequency oscillatory ventilation; iNO, inhaled nitric oxide.

(23%) had prosthetic patches; and 134 (84%) survived. The mean age of cases with early repair was 7 days (SD 8 days) vs 1073 days (SD 1628 days) for cases with late repair. The proportion of survival cases with early repair was 118 of 139 (85%) vs 16 of 20 (80%) for the late repair group (χ^2 =1.1, p=0.3) with a mean follow-up of 6 years. Besides, 108 of 123 (88%) cases with sutured repairs survived, while 26 of 36 (72%) cases with patches survived (χ^2 =5, p=0.02). We only used one latissimus dorsi patch in a girl whose prosthetic patch had become infected.

Temporal patterns

During the four equal time periods, there was no difference in the observed and expected frequencies of cases diagnosed, antenatal diagnosis, live birth, infants undergoing repair and survival. More terminations occurred in the third period (χ^2 =15.5, p<0.0001) (table 1 and figure 2). There was no change in the frequencies of use of HFOV, iNO or ECMO, while the frequency of prostin changed significantly (χ^2 =16, p=0.001) (table 1).

Antenatal diagnosis

Of all cases, 148 (66%) had antenatal diagnosis and 77 (34%) had normal antenatal scan and postnatal diagnosis. Of the 198 live births, 68 of 121 (56%) antenatal diagnosed cases survived, and 66 of 77 (86%) postnatal diagnosed cases survived (χ^2 =18, p<0.0001) (figure 3).

Following repair (159 cases), antenatal diagnosis remained a predictor of death, with survival in 68 of 87 cases diagnosed antenatally (78%) vs 66 of 72 cases diagnosed postnatally (92%) (χ^2 =5, p=0.02).

Antenatal diagnosis was strongly associated with HFOV (χ^2 =30, p<0.001), iNO (χ^2 =17, p<0.001), ECMO (χ^2 =4, p=0.03), and Prostin use (χ^2 =21, p<0.0001) (figure 3).

Left-sided defects were significantly more likely to be antenatally diagnosed (118 in 170 (69%) vs 24 in 49 (49%); χ^2 =10.2, p=0.016). Two cases of bilateral defects were also diagnosed antenatally.

The adverse survival of the right side worsened if diagnosed antenatally. Only 5 out of 18 (28%) live births with a antenatal diagnosis of right-sided defect survived vs 63 out of 103 (61%) live births with an antenatal diagnosis of left-sided defect (χ^2 =7, p=0.008).

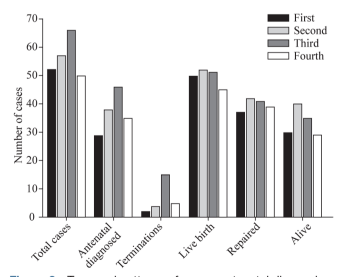


Figure 2 Temporal patterns of cases, antenatal diagnosis, terminations, live birth, repairs and survival during four consecutive periods of 63 months for the duration of the study.

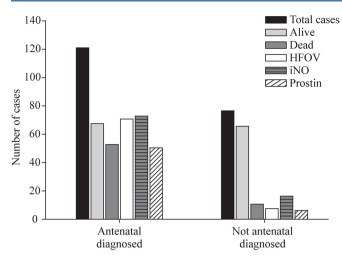


Figure 3 Incidence and association of antenatal diagnosis with outcomes. HFOV, high-frequency oscillatory ventilation; iNO, inhaled nitric oxide.

Terminations

We found only a weak correlation between the annual termination rate and the annual postnatal survival rate (R^2 =0.23, p=0.025).

Associated anomalies

We also discovered that of the 225 cases, 21 (9%) had isolated non-cardiac anomaly, and 52 (23%) had an isolated cardiac anomaly.

Of the 198 live births, 115 (58%) cases had echocardiogram; 43 (22%) had either an ASD or a VSD; 37 (19%) had an isolated PDA; and 7 (4%) had a more complex cardiac anomaly. Of the 83 cases that did not have echocardiogram, 62 (75%) were alive and could be assumed to have no significant cardiac anomaly. Of the remaining 21 cases, 16 died before repair, with 12 dying either on the day of birth or on the following day. Five of the 198 cases (3%) died without cardiac assessment after repair.

We assessed the influence of cardiac anomalies on survival by combining the ASD/VSD group with the more complex anomalies to give a group of 50 with a significant cardiac anomaly. Postnatal survival of cases with either no cardiac anomaly, an ASD/VSD, a more complex cardiac anomaly, or a PDA was the same $(\chi^2=1.9, p=0.5)$.

Twenty-one babies had an additional non-cardiac anomaly, of whom 6 cases (29%) survived, while 15 (71%) died (χ^2 =17, p<0.001). Of the 21 babies with an additional anomaly, a total of 17 (81%) were diagnosed antenatally, compared with 139 of 214 babies (65%) with no additional anomaly (χ^2 =2.1, p=0.1). Children were not more likely to be diagnosed antenatal if there was an associated anomaly.

The adverse effect of additional anomalies was specific to non-cardiac anomalies. Thirty (60%) of 50 infants with an isolated cardiac anomaly survived vs 104 (70%) of 148 infants without an isolated cardiac anomaly (χ^2 =1.8, p=0.2).

High-frequency ventilation or nitric oxide or Prostin administration

Of the 198 live births, 76 cases (38%) underwent HFOV, and 90 (45%) received iNO (table 2). The two strategies were used concurrently in most cases (n=73). Seventeen received iNO without HFOV, and 6 underwent HFOV without iNO. Both were strongly associated with death. We found that 35 (39%) of 90 cases who received iNO survived (χ^2 =62, p<0.0001), while 27/79 (19%) of those who underwent HFOV survived (χ^2 =67, p<0.0001) (table 2).

Variable	Frequency	Death in the presence of variable (%) or mean (SD) among non-survivors for continuous data, n (%)	Death in absence of variable (%) or mean (SD) among survivors for continuous data, n (%)	OR of death (95% CI)	P value
Antenatal diagnosis	121/198	53/121 (44)	11/77 (14)	4.6 (2.2 to 9.7)	<0.0001
Right-sided defect	43/198	19/43 (44)	45/155 (29)	1.9 (0.9 to 3.8)	0.06
Isolated additional anomaly	15/198	9/15 (60)	55/183 (30)	3.4 (1.1 to 10.2)	0.02
Isolated cardiac anomaly	50/198	20/50 (40)	44/148 (30)	1.5 (0.8 to 3.0)	0.2
PDA	61/198	18/61 (29)	46/137 (34)	0.8 (0.4 to 1.5)	0.5
High-frequency ventilation	79/198	52/79 (66)	12/119 (10)	17 (8 to 36)	< 0.0001
Nitric oxide	90/198	55/90 (61)	9/108 (8)	17 (8 to 38)	< 0.0001
ECMO	11/198	7/11 (64)	57/187 (31)	4 (1.1 to 14)	0.03
Prostin	58/198	30/50 (52)	34/140 (24)	3 (1.7 to 6)	< 0.0001
Birth weight (kg)		2.49 (0.8)	2.99 (0.6)	0.4 (0.2 to 0.6)	< 0.001
Gestation (weeks)		35 (5)	37 (2)	0.82 (0.7 to 0.9)	< 0.001

Table 3 Univariate analysis of variables related to survival of infants who survived to undergo repair					
Variable	Frequency	Death in the presence of variable or mean (SD) among non-survivors for continuous data, n (%)	Death in the absence of variable or mean (SD) among survivors for continuous data, n (%)	OR of death (95% CI)	P VALUE
Antenatal diagnosis	87/159	19/87 (22)	6/78 (8)	3 (1 to 8)	0.02
Right-sided defect	34/159	10/34 (29)	15/125 (12)	3 (1.2 to 7.0)	0.02
Defect patched	36/159	10/36 (28)	15/123 (12)	3 (1.1 to 6.8)	0.03
Isolated additional anomaly	9/159	3/9 (33)	22/150 (15)	3 (0.6 to 12.0)	0.1
Isolated cardiac anomaly	39/159	9/39 (23)	16/120 (13)	1.9 (0.7 to 4.8)	0.1
PDA	49/159	6/49 (12)	19/110 (17)	0.6 (0.2 to 1.7)	0.4
HFOV	43/159	16/43 (37)	9/116 (8)	7 (2.8 to 17.0)	< 0.0001
iNO	54/159	19/54 (35)	6/105 (6)	9 (3.3 to 24.0)	<0.0001
Prostin	41/159	13/41 (32)	12/118 (10)	4 (1.6 to 10.0)	0.002
ECMO	9/159	5/9 (56)	20/150 (13)	8 (2 to 32)	0.003
Birth weight		2.3 (0.8)	2.9 (0.7)	0.3 (0.17 to 0.6)	<0.0001
Gestation		35 (3)	37 (2)	0.7 (0.6 to 0.8)	<0.0001

ECMO, extracorporeal membrane oxygenation; HFOV, high-frequency oscillatory ventilation; iNO, inhaled nitric oxide; PDA, patent ductus arteriosus.

Fifty-eight of 198 live births (29%) received Prostin, which was also associated with death (30 of 58 (52%) vs 34 of 140 (24%); χ^2 =14, p<0.0001) (table 2).

Extracorporeal membrane oxygenation

Eleven of 198 (6%) live births underwent ECMO, with 4 of 11 (36%) surviving vs 130 of 187 (69%) of those who did not undergo ECMO (χ^2 =5.2, p=0.02) (table 2). Of the four survivors, one had a neurological deficit, compared with 5 of 130 (3%) of those who did not undergo ECMO (exact test p=0.1).

Six of 11 cases (55%) required emergency laparotomy or thoracotomy due to complications (usually bleeding), with 4 dying, vs 5 of 187 (3%) of non-ECMO infants (χ^2 =40, p<0.0001).

Seven repairs were performed while on ECMO: four cases died and two of the three survivors were neurologically impaired. Three had repair prior to ECMO, and one required ECMO after repair.

Only 1 in 11 children (9%) were alive after ECMO, did not require emergency surgery, and had no neurological deficits.

Logistic regression analysis of explanatory variables

Univariate analysis showed the following variables to be associated with mortality: antenatal diagnosis; rightsided defects; the use of patch, HFOV, iNO, Prostin and ECMO; birth weight; and gestation (tables 2 and 3). On multivariate analysis, only the use of iNO (p<0.0001) and birth weight (p=0.008) were significantly associated with postnatal survival (table 4). For infants who survived to undergo repair, only iNO (p=0.002) (and therefore HFOV and Prostin), antenatal diagnosis (p=0.047), and the side of the defect (p=0.04) were significant (table 5).

DISCUSSION

In the contemporary era, in an English regional center, we describe overall survival for the anomaly of CDH of 60%, postnatal survival of 68%, and postrepair survival of 84%. The following variables are individually associated with significantly poorer survival: lower birth weight, antenatal diagnosis, right-sided defects, non-cardiac-associated anomalies, requirement for patch repair, and use of complex ventilatory/hemodynamic strategies. Taken in combination, only lower birth weight and the need for complex ventilatory/hemodynamic strategies remained significant predictors of death. Our outcomes reflect our management practices, which may differ from those of other institutions.

Table 4	Multivariate analysis of variables related to survival
of live bi	rths.

Variable	OR of death (95% CI)	P value
Antenatal diagnosis	3 (0.9 to 10)	0.06
iNO	53 (13 to 224)	<0.0001
Isolated additional anomaly	3 (0.5 to 21)	0.2
Prostin	0.6 (0.2 to 1.6)	0.3
ECMO	3 (0.6 to 12.0)	0.1
Birth weight	0.3 (0.1 to 0.7)	0.008
Gestation	0.5 (0.7 to 1.1)	0.5
ECMO extracorporeal m	ambrane oxygenation: iNO ir	haled nitric

ECMO, extracorporeal membrane oxygenation; iNO, inhaled nitric oxide.

 Table 5
 Multivariate analysis of variables related to survival of infants who survived to undergo repair

OR of death (95% CI)	P value					
9 (1-83)	0.047					
5 (1-26)	0.04					
0.4 (0.1 to 1.926)	0.3					
1.4 (0.3 to 5)	0.6					
23 (3 to 180)	0.002					
2.2 (0.4 to 10)	0.4					
3 (0.6 to 18)	0.1					
0.3 (0.7 to 1.5)	0.1					
0.8 (0.5 to 1.1)	0.2					
	OR of death (95% Cl) 9 (1-83) 5 (1-26) 0.4 (0.1 to 1.926) 1.4 (0.3 to 5) 23 (3 to 180) 2.2 (0.4 to 10) 3 (0.6 to 18) 0.3 (0.7 to 1.5)					

ECMO, extracorporeal membrane oxygenation; iNO, inhaled nitric oxide.

Defect side and size, associated anomalies and effect of cardiac anatomy

Right-sided defects had poorer survival (54%) than leftsided defects (71%). The poorer survival of patched repairs corroborates studies relating the size of defects to survival.¹¹ Others report poorer survival with patches.¹² There are claims of improved outcome with high rates of patch,¹³ but survival was identical to our 84% with a conservative approach to patch. The use of patches is a surrogate marker for larger defects in our series. This will not be true for centers that repair most defects by patch. We report elsewhere the association between patch use and recurrence.¹⁴

An additional non-cardiac anomaly decreases the probability of survival but cardiac anomalies do not. This corroborates the findings of our earlier report.¹

Survival correlates with levels of beta natriuretic peptide, indicative of heart strain.¹⁵ Theoretically, therefore, cardiac defects that decrease pulmonary hypertension might offer survival advantages, but we found that neither ASD or VSD nor PDA improved survival. Others have noted that even with suprasystemic pulmonary pressures, atrial level shunting is left to right.¹⁶ We lack depth of information about pressures and flow directions. Prostin was associated with a poorer outlook, as it was used selectively in the presence of significant pulmonary hypertension or signs of right heart strain.

Antenatal diagnosis

Among live births, those antenatally diagnosed had worse survival (56% vs 86%). Antenatal diagnosis also predicted the need for HFOV, iNO, and Prostin, all strongly associated with mortality. While it might be thought that antenatal diagnosis would be more likely in the context of multiple anomalies; in fact, the excess mortality among antenatal cases was not because of additional anomalies, which is a separate, independent predictor of mortality.

Comparison with previous studies

There was a weak relationship between the termination rate and postnatal survival. We could not reproduce the association between the termination rate and postnatal survival in our earlier report.¹ There was no pattern of change with time during the study period.

In the 1990s, we reported an overall survival of 71 of 185 cases (38% vs 60% in the current study), postnatal survival of 71 of 129 live births (55% vs 68%), and postrepair survival of 71 of 111 cases (63% vs 84%). All three indices improved from our earlier report.¹

We report postrepair survival of 144/169 (84%) . In a study of late mortality, ventilation beyond 30 days and ECMO predicted death,¹⁷ agreeing with our multivariate analysis of postrepair mortality, where complex ventilation remained the strongest predictor of death.

Compared with our earlier report, improved rates of survival are accompanied by increasing rates of antenatal diagnosis (52%-66%) and decreased terminations (24%-12%). This may be explained first by the strong association between antenatal diagnosis and complex ventilatory strategies and second by our increased use of these strategies. In our earlier study,¹ 12 of 64 cases (19%) received iNO vs 90 of 198 cases (45%) in the current study, while 13 of 66 cases (20%) underwent HFOV previously vs 76 of 198 cases (40%).

The Congenital Diaphragmatic Hernia Study Group (CDHSG) suggests that survival has improved over the same period.¹⁸ Improved survival after ECMO was also reported.¹⁹ Our liveborn survival of 68% resembles the 70% of a multi-institutional German study²⁰ and a Californian 72%.²¹ A Danish study of population size identical to ours reported 80% survival with no ECMO use.¹² Our 56% survival among antenatal diagnosed cases, and 86% survival with no antenatal diagnosis resembles the CDHSG 65% and 83%.²² A UK audit described 1-year survival of 69%, with no change in disease incidence or survival.²³ While this equates to our postrepair survival of 84%, we are skeptical that neonates who died before repair are captured by hospital episode statistics, as the authors concede. Our 30-day postoperative mortality of 3% is similar to the UK figure of 4%.

The Antenatal CDH registry group (ACDHRG) described right-sided defects and lower gestation as predictors of death, as we did.³ We did not subcategorize left-sided defects by liver position and did not measure the fetal lung:head circumference ratio (LHR). Others comment on variable use, difficulties in standardization, and reproducibility of these measurements.^{24 25} Although the fetal LHR is proposed as being prognostic, it should be noted that the area under the curve of the receiver operator characteristic is only 0.76, indicative of only a 'fair' test.³ There remains a considerable overlap between survivors and non-survivors when using the fetal LHR. We note other groups proposing further refinements to the use of antenatal measurements,²⁶ suggesting that we may yet have to arrive at a definitive antenatal prognostic test.

There are conflicting reports on the significance of liver position for left-sided defects, with the same group suggesting that it is²⁷ and is not³ prognostic. The overall survival for the ACDHRG infants with left-sided defects was 126 of 339 (62%), the same as our live-birth survival of 63 of 103 (61%). The same group reported 11 of 25 (44%) survival cases in those with right-sided defects, while our study showed 5 of 18 (28%) survival cases. However, the ACDHRG selected fetuses with isolated defects, while we reported all cases.

Fetoscopic tracheal occlusion

In a randomized trial, FETO was shown to reduce the risk of death among high-risk fetuses with isolated leftsided defects, with survival of 40% compared with 15% in a control group.⁴ However, the same authors reported a population cohort when all patients with CDH were included and FETO was applied, with survival of 62%, which was lower than our 68% with no FETO.²⁸ A large proportion of the babies in the population-based study underwent FETO, 77 of 162 (47%), with 36 surviving. Another contrast is that 78% of patients received patch compared with our 23%. It is difficult to reconcile these survival figures. Clearly, there has been no attempt to distinguish high-risk fetuses in our series, and these babies must be present. However, survival is better than where FETO is liberally applied.

Applying FETO to right-sided defects, results showed that 44 of 116 (38%) survived, but the controls had particularly poor survival of 3 of 26 (12%).²⁹ We reported survival of 4 of 15 (27%) patients with no FETO.

Ventilatory strategies

The association between iNO, HFOV, Prostin, and death indicates that these therapies were applied to the sickest babies. We suggest the failure of others to demonstrate this indicates the therapies being applied to babies who could have survived without their use.^{30 31} Some reports use iNO in 70% of cases,³² which seems difficult to reconcile with our figure of 45% of cases requiring iNO.

Extracorporeal membrane oxygenation

ECMO survival (52%) is identical to the Extracorporeal Life Support Organization registry³³ and higher than the 36% survival where high risk was defined by the lowest achievable paCO₂.³⁴ Among babies who underwent ECMO in our series, emergency surgery for complications was 38%, and neurological deficits were 60%, indicating the complexity and risks of this therapy. We are one of five UK centers offering ECMO, yet only 6% of inborn babies receive ECMO, with a further 10 referred from the UK and Eire. A UK audit reported that 4% of patients were offered ECMO between 2003 and 2016.23 An Austrian study reported -1/3 one third of patients receiving ECMO, but survival was identical to ours.³⁵ ECMO was deployed in 28% of patients in California, again with survival similar to ours.²¹ In contrast, an American survey reported 13% of patients undergoing

ECMO, with postrepair 30-day mortality of 9%.³⁶ Wide variations in ECMO use and survival have been documented,³⁷ implying ECMO use among infants who could survive without ECMO. Given the incidence of neurological damage, this would be a significant error.^{38 39} There are reports of high survival of patients with no ECMO.¹² We do not believe that ECMO will significantly reduce mortality in the UK.

Volume: outcome

We repaired an average of eight cases per annum. A systematic review showed that while there is no accepted definition of high volume, by most standards, we are a high-volume center.⁴⁰ A UK study indicated that more than six repairs would be high volume.²³ The median number of repairs in Californian centers was seven,²¹ with a survival advantage toward higher volume. Because of our numbers of patients and being an ECMO center, it seems unlikely that our findings are explained by lack of experience.

Limitations of this study

Our study suffers from the following limitations. Treatment was in all cases pragmatic and therefore subject to random variation. It is entirely possible that babies might have been allocated to different treatment modalities had differing clinicians been responsible. As we state in our introduction, therapy has changed in the study time period, and willingness to adopt therapies will vary during the study period. We describe contemporary outcomes in an English regional center and accept that these may not be generalizable to other healthcare settings.

Conclusions

We have found that despite fewer terminations, survival has increased. Survival is predicted by birth weight, antenatal diagnosis, need for complex ventilatory/hemodynamic therapies, and non-cardiac anomalies but not cardiac anomalies. Right side and repair by patch are deleterious among those repaired. Overall survival is comparable to centers offering FETO, and ECMO has not significantly reduced mortality.

Acknowledgements The authors are grateful to Ms Victoria Lane, FRCS, for her helpful review of the manuscript.

Contributors EO'C and RT extracted data, suggested further variables of interest, and read and approved the manuscript. SH and TH agreed on the study design, contributed ideas regarding possible explanatory variables, and read and reviewed manuscript. BJ designed the database, performed the analysis, and wrote the manuscript. All authors approved the final manuscript as submitted and agreed to be accountable for all aspects of the work. Bruce Jaffray is the guarantor.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Not applicable.

Ethics approval The authors determined that this study did not require ethical approval by use of the NHS online tool.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. Data are available from the author as an Excel file upon request.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs

Sundeep Harigopal http://orcid.org/0000-0002-5329-5864 Bruce Jaffray http://orcid.org/0000-0002-8906-5016

REFERENCES

- 1 Stege G, Fenton A, Jaffray B. Nihilism in the 1990s: the true mortality of congenital diaphragmatic hernia. *Pediatrics* 2003;112(3 Pt 1):532–5.
- 2 Metkus AP, Filly RA, Stringer MD, et al. Sonographic predictors of survival in fetal diaphragmatic hernia. J Pediatr Surg 1996;31:148–51.
- 3 Jani J, Nicolaides KH, Keller RL, et al. Observed to expected lung area to head circumference ratio in the prediction of survival in fetuses with isolated diaphragmatic hernia. Ultrasound Obstet Gynecol 2007;30:67–71.
- 4 Deprest JA, Nicolaides KH, Benachi A, et al. Randomized trial of fetal surgery for severe left diaphragmatic hernia. N Engl J Med 2021;385:107–18.
- 5 Gallot D, Boda C, Ughetto S, *et al*. Prenatal detection and outcome of congenital diaphragmatic hernia: a French registry-based study. *Ultrasound Obstet Gynecol* 2007;29:276–83.
- 6 UK collaborative randomised trial of neonatal extracorporeal membrane oxygenation. UK collaborative ecmo trail group. *Lancet* 1996;348:75–82.
- 7 Morini F, Goldman A, Pierro A. Extracorporeal membrane oxygenation in infants with congenital diaphragmatic hernia: a systematic review of the evidence. *Eur J Pediatr Surg* 2006;16:385–91.
- 8 Hanekamp MN, Mazer P, van der Cammen-van Zijp MHM, *et al.* Follow-Up of newborns treated with extracorporeal membrane oxygenation: a nationwide evaluation at 5 years of age. *Crit Care* 2006;10:R127.
- 9 Campbell BT, Herbst KW, Briden KE, et al. Inhaled nitric oxide use in neonates with congenital diaphragmatic hernia. *Pediatrics* 2014;134:e420–6.
- 10 Snoek KG, Capolupo I, van Rosmalen J, et al. Conventional mechanical ventilation versus high-frequency oscillatory ventilation for congenital diaphragmatic hernia: a randomized clinical trial (the VICI-trial). Ann Surg 2016;263:867–74.
- 11 Congenital Diaphragmatic Hernia Study Group, Lally KP, Lally PA, et al. Defect size determines survival in infants with congenital diaphragmatic hernia. *Pediatrics* 2007;120:e651–7.
- 12 Larsen UL, Jepsen S, Strøm T, et al. Congenital diaphragmatic hernia presenting with symptoms within the first day of life; outcomes from a non-ECMO centre in Denmark. *BMC Pediatr* 2020;20:196.
- 13 Suply E, Rees C, Cross K, et al. Patch repair of congenital diaphragmatic hernia is not at risk of poor outcomes. J Pediatr Surg 2020;55:1522–7.
- 14 Tamura R, O'Connor E, Jaffray B. Surgeon level variation in outcome of repair of congenital diaphragmatic hernia with particular reference to the management of recurrence. J Pediatr Surg 2021;56:2207–14.
- 15 Tang Y, Ji Y. Do B-type natriuretic peptide levels accurately predict outcome in infants with congenital diaphragmatic hernia? *J Pediatr* 2021;229:311–2.
- 16 Wehrmann M, Patel SS, Haxel C, et al. Implications of atriallevel shunting by echocardiography in newborns with congenital diaphragmatic hernia. J Pediatr 2020;219:43–7.
- 17 Kim AG, Mon R, Karmakar M, et al. Predicting lethal pulmonary hypoplasia in congenital diaphragmatic hernia (CDH): institutional experience combined with CDH registry outcomes. J Pediatr Surg 2020;55:2618–24.
- 18 Gupta VS, Harting MT, Lally KP, et al. Has survival improved for congenital diaphragmatic hernia? A 25-year review of more than

5,000 patients from the congenital diaphragmatic hernia Study Group. *J Am Coll Surg* 2020;231:S199–200.

- 19 Cuestas J, Lohmann P, Hagan JL, *et al.* Mortality trends in neonatal ECMO for pulmonary hypoplasia: a review of the extracorporeal life support organization database from 1981 to 2016. *J Pediatr Surg* 2021;56:788–94.
- 20 Wittekindt B, Doberschuetz N, Schmedding A, *et al.* Epidemiology and one-year follow-up of neonates with CDH-data from health insurance claims in Germany. *Children (Basel)* 2021;8:160.
- 21 Apfeld JC, Kastenberg ZJ, Gibbons AT, et al. Treating center volume and congenital diaphragmatic hernia outcomes in california. J Pediatr 2020;222:146–53.
- 22 Burgos CM, Frenckner B, Luco M, et al. Prenatally versus postnatally diagnosed congenital diaphragmatic hernia - side, stage, and outcome. J Pediatr Surg 2019;54:651–5.
- 23 Wang Y, Honeyford K, Aylin P, et al. One-Year outcomes for congenital diaphragmatic hernia. BJS Open 2019;3:305–13.
- 24 Perrone EE, Abbasi N, Cortes MS, et al. Prenatal assessment of congenital diaphragmatic hernia at North American fetal therapy network centers: a continued plea for standardization. Prenat Diagn 2021;41:200–6.
- 25 Kim ÅG, Norwitz G, Karmakar M, *et al.* Discordant prenatal ultrasound and fetal MRI in CDH: wherein lies the truth? *J Pediatr Surg* 2020;55:1879–84.
- 26 Niemiec SM, Louiselle AE, Phillips R, et al. Third-trimester percentage predicted lung volume and percentage liver herniation as prognostic indicators in congenital diaphragmatic hernia. *Pediatr Radiol* 2023;53:479–86.
- 27 Jani J, Keller RL, Benachi A, et al. Prenatal prediction of survival in isolated left-sided diaphragmatic hernia. Ultrasound Obstet Gynecol 2006;27:18–22.
- 28 Clohse K, Rayyan M, Deprest J, et al. Application of a postnatal prediction model of survival in CDH in the era of fetal therapy. J Matern Fetal Neonatal Med 2020;33:1818–23.
- 29 Russo FM, Cordier A-G, Basurto D, *et al.* Fetal endoscopic tracheal occlusion reverses the natural history of right-sided congenital diaphragmatic hernia: European multicenter experience. *Ultrasound Obstet Gynecol* 2021;57:378–85.
- Fuyuki M, Usui N, Taguchi T, *et al.* Prognosis of conventional vs. high-frequency ventilation for congenital diaphragmatic hernia: a retrospective cohort study. *J Perinatol* 2021;41:814–23.
 Derraugh G, Levesque M, Schantz D, *et al.* High-Frequency vs.
- 31 Derraugh G, Levesque M, Schantz D, et al. High-Frequency vs. conventional ventilation at the time of CDH repair is not associated with higher mortality and oxygen dependency: a retrospective cohort study. *Pediatr Surg Int* 2020;36:1275–80.
- 32 Herich K, Schaible T, Reinhard J, et al. Ino therapy in patients with congenital diaphragmatic hernia-discrepancy between widespread use and therapeutic effects. *Klin Padiatr* 2019;231:320–5.
- 33 Thiagarajan RR, Barbaro RP, Rycus PT, et al. Extracorporeal life support organization registry international report 2016. ASAIO J 2017;63:60–7.
- 34 Jancelewicz T, Langham MR Jr, Brindle ME, *et al.* Survival benefit associated with the use of extracorporeal life support for neonates with congenital diaphragmatic hernia. *Ann Surg* 2022;275:e256–63.
- 35 Brandt JB, Werther T, Groth E, et al. Risk factors for mortality in infants with congenital diaphragmatic hernia: a single center experience. Wien Klin Wochenschr 2021;133:674–9.
- 36 Zani-Ruttenstock E, Zani A, Eaton S, et al. First population-based report of infants with congenital diaphragmatic hernia: 30-day outcomes from the American College of surgeons national quality improvement program. *Eur J Pediatr Surg* 2019;29:62–7.
- 37 Lewit RA, Jancelewicz T. Sources of regional and center-level variability in survival and cost of care for congenital diaphragmatic hernia (CDH). *J Pediatr Surg* 2021;56:130–5.
- 38 Kim F, Bernbaum J, Connelly J, et al. Survival and developmental outcomes of neonates treated with extracorporeal membrane oxygenation: a 10-year single-center experience. J Pediatr 2021;229:134–40.
- 39 Montalva L, Raffler G, Riccio A, *et al*. Neurodevelopmental impairment in children with congenital diaphragmatic hernia: not an uncommon complication for survivors. *J Pediatr Surg* 2020;55:625–34.
- 40 Morche J, Mathes T, Jacobs A, et al. Relationship between volume and outcome for surgery on congenital diaphragmatic hernia: a systematic review. J Pediatr Surg 2020;55:2555–65.

8