

Pregnancy outcomes in women with cardiovascular disease: evolving trends over 10 years in the ESC Registry Of Pregnancy And Cardiac disease (ROPAC)

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Aims

Reducing maternal mortality is a World Health Organization (WHO) global health goal. Although maternal deaths due to haemorrhage and infection are declining, those related to heart disease are increasing and are now the most important cause in western countries. The aim is to define contemporary diagnosis-specific outcomes in pregnant women with heart disease.

Methods and results

From 2007 to 2018, pregnant women with heart disease were prospectively enrolled in the Registry Of Pregnancy And Cardiac disease (ROPAC). Primary outcome was maternal mortality or heart failure, secondary outcomes were other cardiac, obstetric, and foetal complications. We enrolled 5739 pregnancies; the mean age was 29.5. Prevalent diagnoses were congenital (57%) and valvular heart disease (29%). Mortality (overall 0.6%) was highest in the pulmonary arterial hypertension (PAH) group (9%). Heart failure occurred in 11%, arrhythmias in 2%. Delivery was by Caesarean section in 44%. Obstetric and foetal complications occurred in 17% and 21%, respectively. The number of high-risk pregnancies (mWHO Class IV) increased from 0.7% in 2007–2010 to 10.9% in 2015–2018. Determinants for maternal complications were pre-pregnancy heart failure or New York Heart Association >II, systemic ejection fraction <40%, mWHO Class 4, and anticoagulants use. After an increase from 2007 to 2009, complication rates fell from 13.2% in 2010 to 9.3% in 2017.

Conclusion

Rates of maternal mortality or heart failure were high in women with heart disease. However, from 2010, these rates declined despite the inclusion of more high-risk pregnancies. Highest complication rates occurred in women with PAH.

Keywords

Pregnancy • Maternal mortality • Congenital Heart Disease • Cardiomyopathies • Pregnancy complications

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Introduction

Maternal heart disease complicates between 1% and 4% of pregnancies, and accounts for up to 15% of maternal deaths.^{1,2} According to the most recent statistics from the World Health Organization (WHO), maternal mortality in developed economies is around 12 per 100 000 live births (0.012%) and 239 per 100 000 live births (0.2%) in emerging economies, with large disparities both between and within countries.³ While leading causes such as haemorrhage and infection are declining, mortality due to maternal heart disease is increasing^{2,4,5} and in developed countries, heart disease is now the leading cause. The burden of maternal heart disease is expected to increase due to improved survival of women with congenital heart disease (CONHD) and the trend to delay motherhood, with the associated rise in comorbidities such as metabolic syndrome. Thus, heart disease may become an even greater contributor to maternal deaths worldwide.²

Research is difficult to perform in the field of heart disease and pregnancy and observational studies such as CARPREG and ZAHARA have been the primary source of evidence.^{6–9} Large prospective registries were needed to improve the quality and level of knowledge relating to this growing population. Therefore, in 2007, the EURObservational Research Programme (EORP) of the European Society of Cardiology (ESC) established the Registry Of Pregnancy And Cardiac disease (ROPAC). This important data source assembles granular clinical data on large numbers of pregnant patients with heart disease from real-life daily practice, informing our understanding of the outcomes in this population with the ultimate goal of improving patient outcomes. An interim analysis of the first 1321 patients was reported in 2013¹⁰ and an overview of the full registry is presented here.

The aims of the current study are to provide contemporary information on the impact of a wide range of cardiovascular diseases on maternal and foetal mortality and morbidity and identify key areas where current management can be improved.

Methods

Study design

The Registry Of Pregnancy And Cardiac disease (ROPAC) is an international, prospective, observational registry of pregnant women with CONHD, valvular heart disease (VHD), cardiomyopathy (CMP), or ischaemic heart disease (IHD). Pregnancies were included prospectively from January 2007 until January 2018. From 2013, patients with pulmonary arterial hypertension (PAH) or aorta pathology (AOP) were also enrolled. An interim analysis of the first 1321 pregnancies (PREG1) was published in 2013. Comparisons were made between the PREG1 and the data collected from 2011 onwards (PREG2). We included data from all pregnancies up to 1 week post-partum. Women with a primary diagnosis of PAH or AOP were excluded from the analysis of trends over time because they have not been enrolled in ROPAC from the start of the registry.

Data, definitions, study endpoints, and statistical analysis

The study protocol, methods, and the outcomes of the first 1321 pregnancies included were published in 2013.¹⁰ The primary endpoint was maternal mortality and/or heart failure. Information on data, definitions,

endpoints, and statistical methods can be found in the [Supplementary material](#) online, [Appendix](#).

Results

A total of 5739 pregnancies were recruited from 138 centres in 53 countries. The list of participating centres is given in the [Supplementary material](#) online, [Appendix](#). Sixty percent of women were from developed countries. The mean maternal age was 29.5 years, and 45% were nulliparous. There were 96 twin pregnancies (1.7%). The baseline characteristics for PREG1, PREG2, and the total cohort are presented in [Table 1](#). Details of diagnosis are shown in [Supplementary material](#) online, [Figure S1](#). There were missing data in 1.7% of baseline variables.

[Table 2](#) summarizes the cardiovascular, obstetric, and foetal outcomes, for PREG1, PREG2, and the total cohort. In [Supplementary material](#) online, [Tables S1–S3](#), the baseline characteristics and outcomes are summarized per diagnostic group and divided in emerging and developed countries and in [Supplementary material](#) online, [Table S4](#), the differences in characteristics and outcome are compared between women who have undergone pre-conceptional counselling and those who have not.

The PREG1 and the PREG2 cohorts differed significantly with respect to the percentage nulliparous women (49.2% vs. 43.5%, $P < 0.001$), women from emerging countries (21.3% vs. 45.3%, $P < 0.001$), women with a diagnosis of CONHD (62.0% vs. 56.0%, $P < 0.001$), prior interventions (61.3% vs. 53.2%, $P < 0.001$), and pre-pregnancy cardiac medication use (20.4% vs. 40.8%, $P < 0.001$). Also, the distribution of women in the different mWHO classes differed significantly, the most remarkable difference being the percentage of women in mWHO IV.

Maternal cardiovascular outcome and clinical determinants of study endpoint

Maternal mortality and/or heart failure occurred in 629 pregnancies (11%), with maternal death in 34 women (0.6%). In four cases, the cause of death was unknown and five patients died from non-cardiac causes, four of these being septic shock. Refractory heart failure was the cause of death in 15 patients, five of which were related to severe PAH. Four patients died from complications of mechanical valve thrombosis, two from endocarditis, two patients from primary cardiac arrest with ventricular fibrillation, and two other patients from haemodynamic complications following abortion and pulmonary embolism, respectively.

Heart failure complicated 611 pregnancies (11%), of which 44 (7%) occurred solely postpartum. In women with a mechanical valve, 22 (7%) pregnancies were complicated by valve thrombosis, which was lethal in four (18%). Of the six women with known valvular replacement surgery none died, while of the two women with known thrombolytic therapy one died due to intracerebral haemorrhage. [Supplementary material](#) online, [Table S5](#) shows the women with valve thrombosis and their outcome. In the AOP group, aortic dissection occurred in four (1.8%), at a mean of 34 weeks gestation. In [Figure 1](#), the occurrence of cardiovascular events is summarized for the different diagnosis groups and in [Supplementary material](#) online, [Figure S2](#), it is subdivided for the additional sub-groups. Within PREG1 were

Table 1 Baseline characteristics of PREG1 and PREG2 cohorts and the total cohort

	Total cohort (n = 5739)	PREG1 cohort (n = 1321)	PREG2 cohort (n = 4418)	P-value
Demographics				
Age, mean (SD)	29.5 (5.6)	29.2 (5.7)	29.6 (5.6)	0.01
Nulliparity	2573 (44.8%)	652 (49.2%)	1921 (43.5%)	<0.001
Emerging country	2281 (39.7%)	282 (21.3%)	1999 (45.3%)	<0.001
Diagnosis				
Congenital heart disease	3295 (57.4%)	822 (62.0%)	2473 (56.0%)	<0.001
Valvular heart disease	1648 (28.7%)	341 (25.7%)	1307 (29.6%)	0.01
Cardiomyopathy	438 (7.6%)	91 (6.9%)	347 (7.9%)	0.23
Pregnancy-induced during current pregnancy	23 (5.3%) ^a	0 (0%)	23 (6.6%)	<0.001
Ischaemic heart disease	95 (1.6%)	26 (2.0%)	69 (1.6%)	0.32
Pre-pregnancy clinical characteristics				
Current smoking	228 (4%)	45 (3.3%)	183 (4.1%)	0.34
Hypertension	380 (7%)	88 (6.7%)	292 (6.6%)	0.89
History of diabetes mellitus	90 (2%)	21 (1.6%)	69 (1.6%)	1
History of atrial fibrillation	106 (2%)	24 (1.8%)	82 (1.9%)	0.91
Signs of heart failure	596 (11%)	136 (10.3%)	460 (10.4%)	0.88
Prior intervention	3160 (55%)	812 (61.3%)	2348 (53.2%)	0.08
NYHA class				0.35
I	4207 (73.3%)	937 (70.7%)	3270 (74.1%)	
II	1191 (20.8%)	323 (24.5%)	868 (19.7%)	
III	176 (3.1%)	40 (3.0%)	136 (3.1%)	
IV	28 (0.5%)	4 (0.3%)	24 (0.5%)	
Cardiac medication	2069 (36.1%)	270 (20.4%)	1799 (40.8%)	<0.001
mWHO class				0.05
I	1185 (20.6%)	284 (21.4%)	901 (20.4%)	
II	828 (14.4%)	226 (17.1%)	602 (13.6%)	
II–III	2698 (47.0%)	720 (54.3%)	1978 (44.8%)	
III	593 (10.3%)	88 (6.6%)	505 (11.4%)	
IV	407 (7.1%)	7 (0.5%)	400 (9.1%)	

P-values were calculated between the PREG1 and PREG2 cohorts (independent samples *t*-test and χ^2 tests where appropriate).

^aPercentage of cardiomyopathies.

significantly less thrombotic events compared with PREG2 (0.5% vs. 1.8%, $P = 0.006$).

In *Figure 2*, the results from the regression analyses are shown for the primary outcome for the total cohort. Pre-pregnancy New York Heart Association (NYHA) >II, systemic ventricular ejection fraction (EF) of <40%, signs of heart failure, mWHO IV, and anticoagulation use were independent predictors of the occurrence of death/heart failure.

Study endpoints over time

The incidence of death/heart failure changed over time (*Figure 3*). Assuming a linear trend since 2007, the annual incidence decreased by 0.8%/year ($P < 0.001$). However, our data suggest that there was an early increase until 2010, with a subsequent annual decrease of 1.3%/year ($P < 0.001$) thereafter. Noteworthy, this decline was reached, while the percentage of mWHO IV patients increased from around 1% in the first years up to 10% in the last years (*Supplementary material online, Figure S3*). Time trends were similar in the three main diagnostic groups, but most pronounced in CMP.

Supplementary material online, Figure S5 shows trend analyses for emerging and developed countries.

Obstetric and foetal outcome

A total of 992 pregnancies (17%) were complicated by obstetric events during pregnancy. Emergency caesarean section was performed in 537 pregnancies (9%), 84 (16%) were for cardiac reasons. Preeclampsia occurred in 3% and was seen more commonly in patients with CMP and PAH (7% and 11%, respectively). In women with peripartum CMP during current pregnancy ($n = 23$), six contemporaneously suffered from preeclampsia (26.1%).

In 1186 pregnancies (21%), one or more foetal complications occurred. In particular prematurity was prevalent (16%). These rates were highest in patients with CMP (26%), PAH (49%), and AOP (21%). In *Supplementary material online, Figure S6*, the occurrence of obstetric and foetal events is summarized per diagnostic group. Cardiovascular, and all obstetric and foetal outcomes varied according to mWHO categories (*Supplementary material online, Figure S3*).

Table 2 Outcomes of PREG1 and PREG2 cohorts and the total cohort

	Total cohort (n = 5739)	95% CI	PREG1 cohort (n = 1321)	95% CI	PREG2 cohort (n = 4418)	95% CI	P-value
Cardiovascular outcome							
Maternal mortality and/or heart failure	629 (11.0%)	10.2–11.8%	161 (12.2%)	10.5–14.1%	468 (10.6%)	9.7–11.5%	0.11
Maternal mortality	34 (0.6%)	0.4–0.8%	9 (0.7%)	0.4–0.13%	25 (0.6%)	0.4–0.8%	0.63
Heart failure event	611 (10.6%)	9.9–11.5%	153 (11.5%)	10.0–13.4%	458 (10.3%)	9.5–11.3%	0.29
Supraventricular arrhythmia	95 (1.7%)	1.4–2.0%	11 (0.9%)	0.5–0.15%	84 (1.9%)	1.5–2.3%	0.06
Ventricular arrhythmia	90 (1.6%)	1.3–1.9%	27 (2.0%)	1.4–3.0%	63 (1.4%)	1.1–1.8%	0.30
Thrombotic event	87 (1.5%)	1.2–1.9%	6 (0.5%)	0.2–1.0%	81 (1.8%)	1.5–2.3%	0.006
Aortic dissection	5 (0.1%)	0.0–0.2%	0 (0%)	0.0–0.3%	5 (0.1%)	0.0–0.3%	0.22
Obstetric outcome							
Pregnancy-induced hypertension	150 (2.6%)	2.2–3.1%	25 (2.4%)	1.3–2.8%	125 (2.8%)	2.4–3.4%	0.57
(Pre-)eclampsia	159 (2.8%)	2.4–3.2%	44 (3.3%)	2.5–4.4%	115 (2.6%)	2.2–3.1%	0.35
Post-partum haemorrhage	170 (3.0%)	2.6–3.4%	38 (2.9%)	2.1–3.9%	132 (3.0%)	2.5–3.5%	0.89
Caesarean section	2681 (46.7%)	45.4–48.0%	538 (40.6%)	38.1–43.3%	2143 (48.6%)	47.0–50.0%	<0.001
Emergency caesarean section	766 (13.3%)	12.5–14.3%	204 (15.4%)	13.6–17.5%	562 (12.7%)	11.8–13.7%	0.01
Foetal outcome							
Foetal mortality	72 (1.3%)	1.0–1.6%	20 (1.5%)	1.0–2.3%	52 (1.2%)	0.9–1.5%	0.56
Neonatal mortality	33 (0.6%)	0.4–0.8%	8 (0.6%)	0.3–1.2%	25 (0.6%)	0.4–0.8%	0.88
Premature birth	905 (15.8%)	14.8–16.7%	200 (15.1%)	13.3–17.2%	705 (16.0%)	14.9–17.1%	0.58
Low apgar scores	397 (6.9%)	6.3–7.6%	95 (7.2%)	5.9–8.7%	282 (6.4%)	5.7–7.1%	0.48
IUGR	254 (4.4%)	3.9–5.0%	70 (5.3%)	4.2–6.6%	184 (4.2%)	3.6–4.8%	0.25
Low birth weight	673 (11.7%)	10.9–12.6%	12 (0.9%)	0.5–1.6%	661 (15.0%)	13.9–16.0%	<0.001

P-values were calculated between the PREG1 and PREG2 cohort (χ^2 tests). All events were monitored between enrolment date (start of pregnancy or first trimester) and 1 week after delivery.

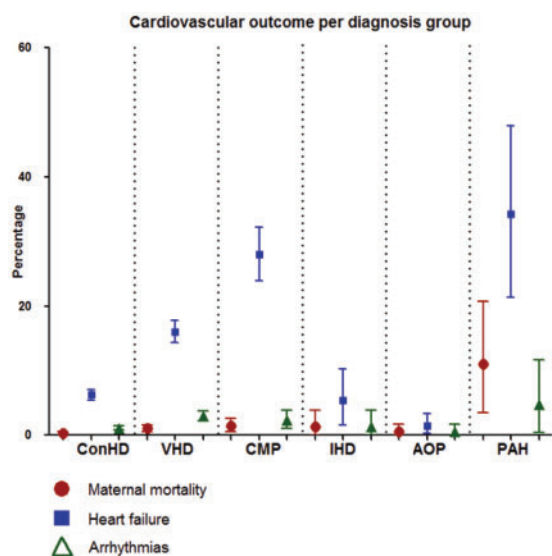
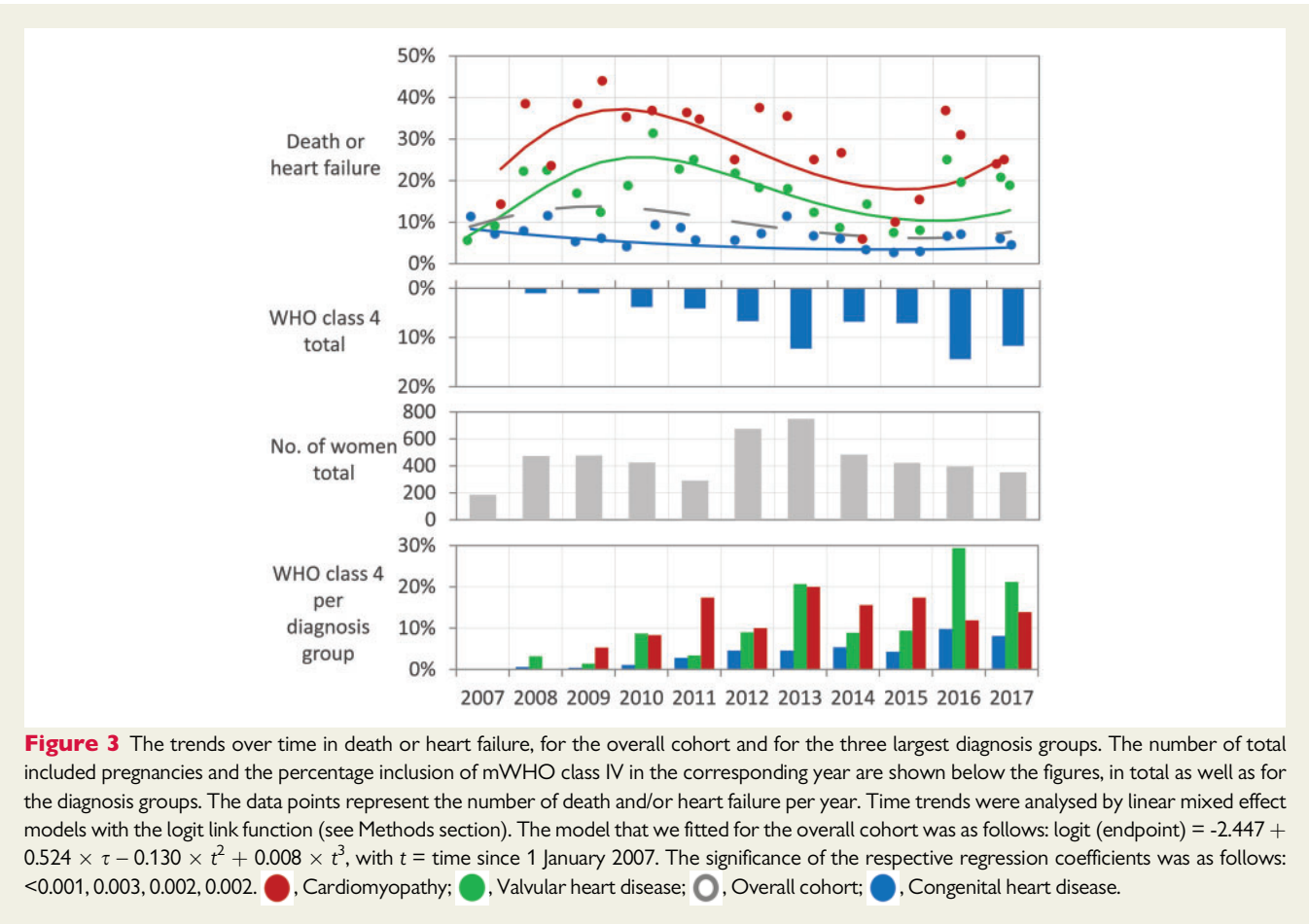
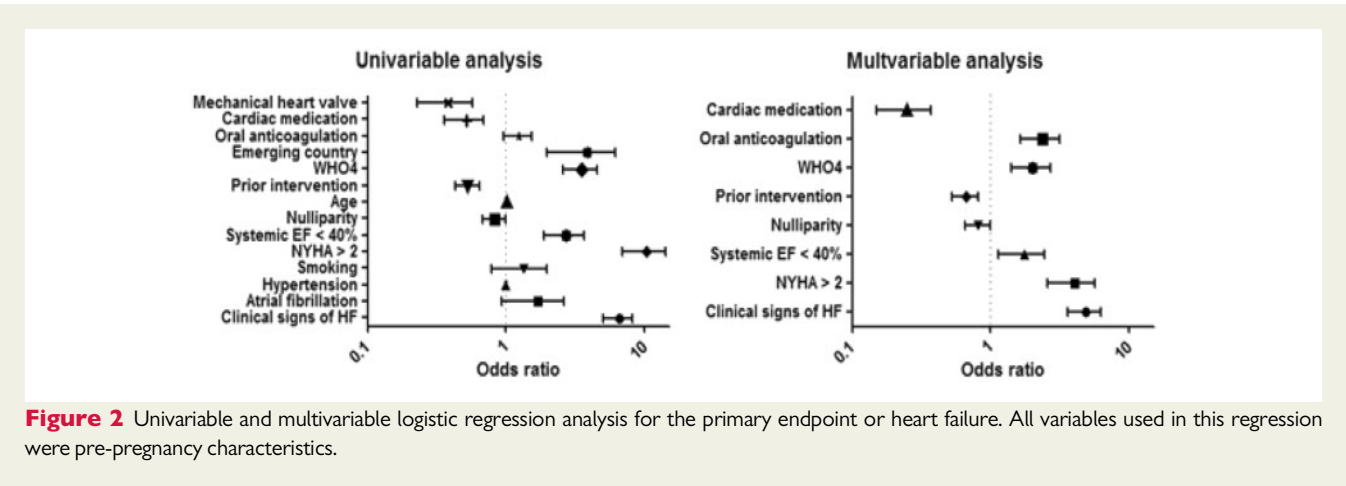


Figure 1 The occurrence of maternal cardiovascular events for the different diagnosis groups. AOP, aortic pathology; CMP, cardiomyopathy; CONHD, congenital heart disease; IHD, ischemic heart disease; PAH, pulmonary arterial hypertension; VHD, valvular heart disease.

Discussion

ROPAC is a prospective registry containing 5739 pregnancies in women with structural and IHD, PAH, and AOP. This is by far the largest registry of its kind ever undertaken. The overall maternal mortality rate in our study was 0.6%, which is much higher than that observed in the normal pregnant population.¹¹ The most important complication found in our registry is heart failure, which complicated 11% of pregnancies, of which 7% occurred for the first time in the first week postpartum. These results highlight that intensive management and monitoring of women at risk of heart failure is essential throughout pregnancy and also after delivery.

One of the WHO Millennium Development Goals was to reduce maternal deaths by 75% between 1990 and 2015; this was partially successful, as the rate fell by 44%.³ The new Sustainable Development Goals have targeted a reduction in worldwide maternal mortality to <70 deaths per 100 000 live births by 2030.³ To achieve this target, the rate of decline in maternal mortality will have to increase to 7.5% per year. This is unlikely to be achieved unless the increasing contribution to mortality made by heart disease is reversed. Data from ROPAC can help by determining which patients are at high risk and identifying areas in which the greatest improvements can be achieved.

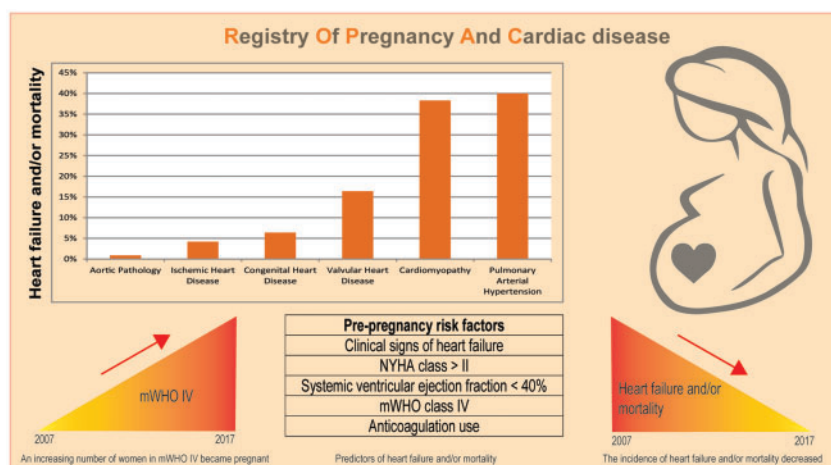


Diagnostic groups: low-risk and high-risk patients

The highest mortality/heart failure rates were found in women with PAH, CMP, and VHD as also described by previous series.^{12–20} The trends over time showed that especially in VHD and CMP there was a decrease in maternal mortality or heart failure, while in CONHD the rate was and remained low. Our study

emphasizes that patients with PAH are still, despite advances in management, a high-risk group and that they should be advised against pregnancy.

The mortality rate for the CMP group was 1.1% and heart failure occurred in 28%. Current guidelines advise against pregnancy if the systemic ventricular EF is $<30\%$.¹² Our data suggest that all patients with dilated CMP, even with relatively preserved EF, are high risk and should be counselled accordingly.^{21–23}



Take home figure From 2007 to 2018, pregnant women with heart disease were included in the Registry Of Pregnancy And Cardiac disease (ROPAC). During that time, the incidence of 'maternal death or heart failure' dropped, despite the inclusion of more seriously ill women (mWHO IV). Especially women with cardiomyopathies or pulmonary arterial hypertension were at risk of 'maternal death or heart failure'. Pre-pregnancy risk factors were signs of heart failure prior to pregnancy, NYHA class > II, systemic ventricular ejection fraction of below 40%, mWHO class IV conditions and anticoagulation use.

Patients with CONHD constitute the largest subgroup within this registry and have a relatively favourable pregnancy outcome,¹⁰ with a mortality rate of 0.2%. However, due to the heterogeneous nature of this group, the type and complexity of CONHD need to be taken into account. Indeed, the heart failure rate in the patients with complex CONHD was 13%, while it was 5% and 6% for the simple and moderate defects, respectively. The vast majority of patients had their condition diagnosed and treated at a young age, allowing ample opportunity for pre-pregnancy counselling and optimization prior to pregnancy, perhaps accounting for the relatively good outcomes.

In the AOP group, the rate of dissection was 1.8% and we found a strikingly high rate of delivery by caesarean section (52%). There are limited data on the effect of labour and active pushing on the risk of aortic dissection, and current guidelines are solely based on expert opinion.^{24–26} Many physicians feel safe when using caesarean section, however, using epidural anaesthesia with the avoidance of pushing might be as good or even better for mother and baby,²⁷ but more research is clearly required to investigate the optimal mode and timing of delivery in these women. A dedicated ROPAC registry for patients with AOP just started within the EORP, to further analyse the risks and outcomes during pregnancy.

A large proportion of patients with VHD had rheumatic heart disease (56%). These women show a tendency to present late in pregnancy, meaning that those caring for them often only become aware of the condition when complications arise.²⁸ This may explain the high rates of maternal mortality (1%) and heart failure (17%) in this group. Especially severe mitral stenosis is associated with high complication rates.²⁸ Seven percent of patients with a mechanical valve suffered from valve thrombosis, of which 18% died. There is still no consensus with respect to the best anticoagulation strategy,^{29–32} another area where research is urgently required. Women with a mechanical prosthesis are currently classified as mWHO III. However, given the high rates of severe and potentially lethal

complications and miscarriages, perhaps Class IV is more appropriate and a particular cautious approach is warranted to counselling these women and close observation with dedicated supervision is recommended when pregnancy occurs. In the recently published ESC guidelines on cardiovascular disease and pregnancy, women with mechanical prostheses are now advised to be referred to a tertiary care centre, because of these high complication rates.¹ A dedicated ROPAC registry for patients with valvular prostheses also started in 2018, to specifically investigate which anticoagulation regimen is associated with the lowest risks and to further analyse the outcomes during pregnancy.

The women with IHD were typically older, were more often multipara and had surprisingly good maternal outcome, with no mortality and 4% heart failure. Evidence in this field is still very limited and these patients are often being approached with much caution and pregnancy being discouraged. Clearly larger series are required to facilitate better counselling.

The results from our study are in line with recommendations from the 2018 ESC guidelines for the management of cardiovascular diseases during pregnancy.¹ Indeed all women with heart disease should be managed in a hospital. Women in mWHO Class I and II can be cared for in a local hospital and women in Class II–III in a referral hospital, while women in Class III and IV should be managed in specialized, tertiary centres by a specialized, multidisciplinary team. Pre-conceptional counselling is advised for women in all mWHO classes as it is associated with better outcome.

Clinical determinants of study endpoint

Pre-pregnancy predictors of heart failure and/or mortality were signs of heart failure or NYHA >II, systemic ventricular EF of <40%, mWHO IV, and anticoagulants use. This highlights the importance of pre-conception counselling and optimization of cardiac function before pregnancy. Some patients can benefit from adjusting or

introducing medication to improve their cardiovascular status before pregnancy. The use of anticoagulation as a predictor for adverse outcome is likely to be explained only partly by the effect of the medication itself. The underlying reason for the medication, such as mechanical valves, arrhythmias or dilated CMP probably have an effect.

Study endpoints over time

A remarkable finding was that over the period of the study, an increasing number of pregnancies in women with mWHO IV disease were included. This is surprising since conventional advice to women in this high-risk group is to avoid pregnancy. This is not due to the later inclusion of patients with PAH and AOP since these groups were excluded from this analysis. Of course, this increase might be (partly) explained by selection of patients due to centre and country inclusion, but clustering within country and site was controlled for. While recommendations and guidelines are now available and often discussed with the patients, we cannot prevent women from choosing to become pregnant and we not infrequently encounter high-risk women who have become pregnant despite our advice. A further explanation of this change in risk profile is that physicians have become more comfortable with managing women with heart disease, feel better capable of dealing with their problems and have therefore modified their advice. Alternatively, it may simply reflect the existence of increasing numbers of high-risk patients in the population. Indeed, the first wave of patients with surgically corrected complex congenital heart defects such as women with a Fontan circulation are now reaching adulthood and have the wish to become pregnant. This increasing number of high-risk pregnancies is a challenge and requires careful management. Centralization of care for these patients may optimise outcome. It is also notable, that despite this increase in risk profile, there was not an increase in mortality with time. [Supplementary material online, Figure S4](#) shows that there were around 11% of women in mWHO IV in the years 2015–2017, yet still the mortality was lower than in the beginning of the study period, when the proportion mWHO IV was only 1%. In fact, from 2010 to 2011 onwards, the incidence of death or heart failure showed a significant decrease. This might indicate greater awareness of the specific problems and an improvement in the management of pregnant women with heart disease, which may be related to the publication in 2011 of the first ESC guideline on the management of pregnancy in women with cardiovascular disease. Although this may be a coincidence, it is possible that the well-publicized availability of this central resource has led to an increased awareness of the improvement in management. Of course, the improvement in outcome could also be associated with political, socioeconomic or access-to-healthcare factors. Our findings are mirrored by those of the United Kingdom report on maternal mortality which also shows a fall in mortality in recent years.² Not surprisingly, we saw the greatest improvement in outcome in emerging countries, probably because this is where the mortality was highest and consequently, where the greatest improvement could be achieved.

Limitations

While ROPAC is prospective and the largest registry including all groups of heart disease, which describes real-life clinical practice, it is limited by several factors:

- Because we report only mortality up to one week our estimates are highly likely to have underestimated this component of the primary endpoint compared with other studies and this should be considered when making comparison.
- Many pregnant women with heart disease in our cohort were managed in specialized, tertiary centres with an interest in this field. The outcomes of other, non-tertiary centres may be different and caution is required in generalizing our results to the whole population.
- Although the major advantage of a prospective registry is the way in which it can document clinical reality, there might still be some form of selection bias present, as we cannot guarantee (although it was requested and stressed) that consecutive patients from all centres were included.
- Some countries and centres have included more patients than others. Furthermore, the number and constitution of participating centres varied over time, creating a heterogeneous cohort in which interpreting results is difficult. However, we adjusted for this in our analyses.

Conclusions

While pregnancy in women with heart disease in general is associated with increased risks of maternal mortality (0.6%) and cardiovascular, obstetric, and foetal complications, many patients tolerate pregnancy well. Patients with PAH and CMP are at particularly high risk, while women with simple CONHD have relatively good outcomes. Pregnancy in women with low-risk conditions should not be discouraged, while clear and timely advice not to become pregnant should be provided to very high-risk women. Pre-pregnancy predictors for mortality and/or heart failure are NYHA class of >II, mWHO IV, systemic ventricular EF of <40%, signs of heart failure, and the use of anticoagulants.

Until 2010, maternal mortality and/or heart failure increased. Thereafter, these rates declined, especially in emerging countries, despite increasing numbers of high-risk patients. Further studies are required to assess the optimal management of these patients, with particular emphasis on cardiac medication use, including anticoagulation regimes and mode of delivery.

Supplementary material

[Supplementary material](#) is available at *European Heart Journal* online.

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