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DOI: 10.1002/pd.4912 PRENATAL **DIAGNOSIS** 

# **ORIGINAL ARTICLE**

# An exploration of the potential utility of fetal cardiovascular MRI as an adjunct to fetal echocardiography

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# **ABSTRACT**

Objectives Fetal cardiovascular magnetic resonance imaging (MRI) offers a potential alternative to echocardiography, although in practice, its use has been limited. We sought to explore the need for additional imaging in a tertiary fetal cardiology unit and the usefulness of standard MRI sequences.

Methods Cases where the diagnosis was not fully resolved using echocardiography were referred for MRI. Following a three-plane localiser, fetal movement was assessed with a balanced steady-state free precession (bSSFP) cine. Single-shot fast spin echo and bSSFP sequences were used for diagnostic imaging.

Results Twenty-two fetal cardiac MRIs were performed over 12 months, at mean gestation of 32 weeks (26–38 weeks). The majority of referrals were for suspected vascular abnormalities (17/22), particularly involving the aortic arch (n=10) and pulmonary vessels (n=4). Single-shot fast spin echo sequences produced 'black-blood' images, useful for examining the extracardiac vasculature in these cases. BSSFP sequences were more useful for intracardiac structures. Real-time SSFP allowed for dynamic assessment of structures such as cardiac masses, with enhancement patterns also allowing for tissue characterisation in these cases.

Conclusions Fetal vascular abnormalities such as coarctation can be difficult to diagnose by using ultrasound. Fetal MRI may have an adjunctive role in the evaluation of the extracardiac vascular anatomy and tissue characterisation. © 2016 The Authors. *Prenatal Diagnosis* published by John Wiley & Sons, Ltd.

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Conflict of interest: None

# INTRODUCTION

Conventional fetal echocardiography is the mainstay for prenatal diagnosis of congenital cardiac lesions, based on its ease of use, availability and high diagnostic accuracy.<sup>1–3</sup> However, individual fetal and maternal factors can have a deleterious effect on the quality of ultrasound imaging,<sup>4,5</sup> and there remain inherent difficulties in diagnosing specific forms of congenital heart disease in the fetus.<sup>3,6–8</sup> Despite this, due to technical and safety considerations, alternatives to ultrasonographic techniques have traditionally been extremely limited.<sup>9</sup>

Magnetic resonance imaging (MRI) has been an established adjunct for postnatal assessment of the cardiovascular system since the 1980s, 10 with routine studies able to deliver three-dimensional angiography, detailed real-time imaging, cardiac volumes and vascular flow measurements. 11 Cardiovascular MR imaging in the fetus, however, presents

several challenges not present after birth. The fetal heart is extremely small, with a total length of between 2 and 4 cm depending on gestational age, <sup>12</sup> and beats with high heart rates of up to 160 beats per minute. <sup>13</sup> Electrocardiographic 'gating', a means of improving spatiotemporal resolution, is not yet practicable in the fetus, although several non-invasive methods are under investigation. <sup>14,15</sup> Unpredictable three-dimensional fetal motion, maternal bulk motion and fetal and maternal breathing movements present additional difficulties, <sup>16</sup> with no additional benefit from attempted fetal sedation. <sup>5</sup>

The first report of attempted fetal cardiac MRI was in 2005, using real-time sequences with an attempt to quantify ventricular volumes.<sup>17</sup> There followed a number of small case series of fetuses with normal hearts<sup>18,19</sup> and congenital heart disease.<sup>20</sup> Whilst several subsequent reviews have attempted to develop defined protocols,<sup>21,22</sup> few have focused on the

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clinical role and utility of fetal cardiac MRI,<sup>5</sup> and there is currently no consensus on routine fetal indications or technical protocols for cardiovascular MRI in clinical practice.

The increasing use of MRI for prenatal evaluation of the fetal brain<sup>23</sup> and other organs<sup>24</sup> offers the potential for more widespread access to the expertise and infrastructure required to deliver fetal MRI programmes. We sought to explore both the need for adjunctive fetal cardiac imaging in a tertiary fetal cardiology unit and the usefulness of standard MRI sequences in delivering additional diagnostic information.

# **AIMS**

Over a one-year period, fetal MRI was requested on clinical grounds by the fetal cardiology service, reserved for any difficult fetal cases where the diagnosis was in doubt or could benefit from additional imaging. We present an analysis of the pattern of referrals we subsequently observed and the diagnostic utility of standard MRI sequences in this context.

#### **METHODS**

Referrals were based on the judgement of the attending fetal cardiologists (JS, OM, VZ and GS). All examinations took place between June 2014 and May 2015. Pre-existing ethical approval was in place for a long-standing project examining brain growth in fetuses with CHD, which also permitted the use of cardiac sequences (REC: 07/H0707/105). All participants provided written consent and received concomitant fetal brain

imaging. The timing of the MRI was determined by the clinical teams, taking into account the gestational age at diagnosis, the potential impact of new information on fetal counselling and management, and the timeframe for optimal imaging quality (generally later gestation, when the fetus is larger with fewer major translational movements). A paediatric cardiologist with expertise in cardiac MR was present at every study (RR or KP). Each mother's tympanic temperature was taken before and after the MRI or if they became uncomfortable during the study. Conservative limits of a rise in temperature of  $\geq 1\,^{\circ}\text{C}$  and/or an absolute estimated temperature in the fetus of greater than 38 °C (i.e. maternal temperature +0.5 °C) <sup>17</sup> were set; breach of either would lead to the study being discontinued.

A large field of view (covering the whole fetus) bSSFP cine (real-time) scan with low spatial resolution but higher temporal resolution (303 ms) was used following the initial localiser sequence to demonstrate the degree of gross fetal movement.<sup>25</sup> Diagnostic sequences were generally limited to half-Fourier acquisition single-shot fast spin-echo ('SSFSE') and balanced steady-state free precession ('bSSFP') gradient echo sequences, as these sequences were felt to offer reasonable signal contrast and spatiotemporal resolution within established limits of radiofrequency power deposition<sup>26</sup> and acoustic noise<sup>27</sup>. These were either multi-slice imaging to provide whole heart coverage using parallel overlapping slices or single-slice imaging in targeted scan planes where dynamic

Table 1 Imaging parameters

Sequence	Multi-slice SSFSE	Multi-slice bSSFP	Single-slice bSSFP
TR/TE (ms)	15 000/100	4.3/2.1	3.9/1.9
Flip angle (degrees)	90	90	60
Field of view (mm)	350 × 350	350 × 350	400 × 300
In plane resolution (mm)	$1.4 \times 1.4$	$1.5 \times 2.3$	1.8 × 2.3
Slice thickness (mm)	2.5	5	5
Order	16 interleaved/TR	Linear slices	Dynamic single slice
SENSE factor	2	1	2
Partial-Fourier factor	5/8	1	5/8
Single-slice duration (ms)	468	647	159

SSFSE, single shot fast spin echo; bSSFP, balanced steady-state free precession; TR, repetition time; TE, echo time; SENSE, sensitivity encoding

Table 2 Referral indications

Indication	N	MRI successful	No cardiac data obtained	No postnatal data
Possible coarctation	6	5	la	_
Aortic arch anatomy	4	3	l a	_
Pulmonary vasculature	4	3	1 <sub>p</sub>	_
Other vascular	3	3	_	1 (TOP 35w)
Cardiac mass	3	3	_	1 (TOP 32w)
Cardiac diverticulum	2	2	_	_
TOTAL (%)	22 (100%)	19 (86%)	3 (14%)	2 (9%)

TOP, termination of pregnancy.

<sup>&</sup>lt;sup>a</sup>Excessive fetal movement.

<sup>&</sup>lt;sup>b</sup>Unable to visualise pulmonary veins due to lung compression from large hernia.

Table 3 Summary table of all fetuses referred for fetal cardiac MRI

	Age	5 days	3 days	3 days	ı	4 days	I	1	ı	2 days
	Outcome	Hybrid procedure	Coarctation repair	Hybrid procedure	Outpatient review	Hybrid procedure	Outpatient review	Sugical resection	Outpatient review	Surgical repair
	Postnatal diagnosis	Bonderline left heart Coarcation of the aorta (echocardiography and surgery)	Coarctation of the aota (echocardiography and surgery)	As per prenatal imaging (postnatal MRI and surgery)	Structurally normal heart (echocardiography)	As per prenatal imaging (echocardiography and surgery)	Structurally normal heart (echocardiography)	Leff atrial rhabdomyoma (histology)	As per prenatal imaging (echocardiography)	As per prenatal imaging (echocardiography and surgery)
	Additional MRI benefits	Concomitant MRI brain	I	1	I	No pulmonary lymphangedasia	I	Tissue characterisation	Tissue characterisation Concomitant MRI brain	1
Q.A.	MRI findings	Abandoned: excessive Fetal movement	Hypoplasia of the aortic isthmus 'Posterior shelf' (Figure 1) Ventricular asymmetry IV < RV High risk of coarctation	Large, tortuous, solitary pulmonary veins bilaterally (Figure 3)	Normal cardiac connections No aneurysm seen	Dilated pulmonary veins with nomal anatomy	Mild ventricular asymmetry Normal arch and isthmus Low suspicion of coarctation	Leff atrial mass; atrial myxoma, haemangioma or habdomyoma (Figure 7)	Solitary RV rhabdomyoma (Figure 8)	Type B interruption Subclavian in continuity with descending Ao
	Co-morbidity	Cerebral ventriculomegaly	None	None	None	None	None	Possible tuberous sclerosis	None	22q11 microdeletion (postnatal)
	MRI indication	Evidence of CoA?	Evidence of CoA?	Pulmonary venous anatomy?	Ventricular aneurysm/ diverticulum?	Pulmonary venous anatomy?	Evidence of CoA?	Nature of cardiac mass?	Nature of cardiac mass?	Aortic arch anatomy?
	Fetal diagnosis (USA)	AVSD Ventricular asymmetry	Possible coardation	Hypoplastic left heart syndrome	Right ventricular aneurysm Small pericardial effusion	Hypoplastic left heart syndrome Restrictive atrial septum	Muscular VSD Possible coarctation	Left atrial mass	RV apex tumour	Mataligned VSD Interrupted aortic arch
	۵	0	0	m	m	0	0	7	-	5
	>	59	32	35	88	£ 8	34	33	38	33
	u	-	~	m	4	40	0	$\sim$	ω	0

Table 3 (Continued)

	Age	9 days	32/40	29 days	1	I	6 days	I	1	29/40	1
	Outcome	Surgical repair	Twin 1 well, twin 2: selective feticide	No intervention to date	Outpatient review	Outpatient review	Surgical repair	Outpatient review	Outpatient review	Medical termination	Outpatient review
	Postnatal diagnosis	As per prenatal imaging (postnatal MR and surgery)	Selective feticide: no postmortem Matemal germline mosaicism for tuberous sclerosis	Mild coardation of the aorta only (Figure 5) (postnatal MRI)	As per prenatal imaging (postnatal CT)	Right-sided aortic arch ALSA (echocardiography)	Coarctation of the aosta (echocardiography and surgery)	As per prenatal imaging (postnatal MRI)	As per antenatal imaging (echocardiography)	Normal pulmonary veins Absent diaphragm bilaterally (postmortem)	As per prenatal imaging (echocardiography)
	Additional MRI benefits	ı	Tissue characterisation MRI brain: multiple rhabdomyomas Twin 1: normal heart	ı	ı	1	I	No airway compression	l	Large diaphragmatic hemia with pulmonary hypoplasia	I
GA	MRI findings	Common arterial trunk with arterial duct No interruption	Twin 1: normal Twin 2: three intramyocardial tumours suggestive of habdomyomas on 12 weighting	Hypoplasia of the aortic isthmus Long gap LCCA-LSA Highrisk neonatal coarctation (Figure 4)	Confluent right and left PAs	Inadequate views (fetal motion)	Moderate transverse arch hypoplasia Highrisk coarctation	Twin 1:R arch, L dua, ALSA Twin 2: normal	large RV, PA and Ao Normal connections No AVMs	Not able to visualise pulmonary veins due to lung compression	Widemouthed contractile diverticulum at base of right ventricle, 10 × 12 mm (Figure 6)
	Co-morbidity	22q11 microdeletion Left hydronephrosis	DCDA twins	None	None	None	None	None	None	Diaphragmatic hemia	SCO
ı	MRI indication	Aortic arch anatomy?	Nature of cardiac masses?	Evidence of CoA?	Pulmonary artery anatomy?	Aortic arch anatomy?	Evidence of CoA?	Aortic arch anatomy?	Other vascular: exclude large AVMs	Pulmonary venous anatomy?	Ventricular aneurysm/ diverticulum?
	Fetal diagnosis (USA)	Common arterial trunk ?Interruption of aortic arch	Twin 1: normal Twin 2: multiple cardiac masses	&Coardation of the aorta Bilateral SVCs	Dysplastic pulmonary valve Severe pulmonary regurgitation Maternal abesity	Right-sided aortic arch Left-sided arterial duct ALSA	Small aortic arch ?Coardation of the aorta	Twin 1: R arch, L duct, ALSA Twin 2: normal	Cardiomegaly, RV>LV Dilated great vessels Normal aortic isthmus	Diaphragmatic hemia	Possible RV diverticulum
	۵	9	7	-	•	4	5	5	-	4	м
	>	33	28	32	29	33	33	36	33	26	30
	u	2	Ξ	12	<u>~</u>	4	15	9		<u>~</u>	6

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Table 3 (Continued)

	Age		35/40	33/40
	Outcome	Outpatient review	Medical termination	Premature Died 1 day of age (atrial arrhythmia)
	Postnatal diagnosis	Structurally namal heart (echocardiography)	Medical termination Genetics: Jacobsen syndrome (no post-morten)	Coarctation of the aorta (echocardiography)
	Additional MRI benefits	1	1	Deficient cerebellar vermis diagnosed
ďγ	MRI findings	Leftward rotation of cardiac axis with tortwars arterial duct Normal vascular anatomy	Rightward rotation of cardiac axis Bildreral SVCs Normal vascular anatomy	Dominant RV, bilateral SVCs, hypoplastic acrtic arch, high probability of coarctation
	Co-morbidity	None	Jacobsen syndrome	Deficient cerebellar vermis
	MRI indication	Other vascular: arterial duct	Other vascular	Evidence of CoA?
	Fetal diagnosis (USA)	Abnormal arterial duct	Bilateral SVCs, apex to right ?Other vascular abnormality	Bilateral SVCs ¿Coarctation of the aorta
	Q	0	4	0
	>	28	22	22
	u	20	21	22

ventricular septal defect; ICCA, left common carotid artery; ISA, left subclavian artery; PA, pulmonary artery; ALSA, aberrant left subclavian arteriovenous malformation; Ao, aorta; CHAOS, congenital high airway obstruction syndrome; RV, right ventricle; VSD, left ventricle, left pulmonary artery; DCSA, doubly committed sub-arterial; AVM, AVSD, atrioventricular septal defect; CoA, coarctation of the aorta; LV, artery;

information was of interest (real-time; Table 1). If needed, further shortened localiser sequences were also used to account for fetal movement. Sequences were repeated as needed to a maximum study time of 60 min.

#### **RESULTS**

Twenty-two fetal cardiac MRI examinations were performed over the referral period. The average gestational age at diagnosis  $23^{+4}$  weeks (median  $23^{+3}$  weeks, range  $15^{+4}$ – $34^{+0}$  weeks) and the average age at MRI was  $31^{+3}$  weeks (median  $33^{+2}$  weeks, range  $22^{+2}$ – $38^{+1}$  weeks). Two twin pregnancies were included, each with one healthy and one affected twin.

The most common indications were for possible coarctation of the aorta in six patients, other arch abnormalities in four patients (including aortic interruption in two patients, right aortic arch and aberrant left subclavian artery) and assessment of pulmonary vessels in four patients (Table 2). In total, referrals for clarification of the extracardiac cardiovascular anatomy comprised 77% of all referrals (17/22 cases). No examinations were abandoned early for maternal claustrophobia. One patient experienced a rise in temperature to 38.3 °C towards the end of the study, which was then discontinued.

Three examinations failed to provide any useful cardiovascular information. In two cases, this was due to excessive fetal movement (at 29 and 33 weeks). In one case, assessment of the pulmonary veins in a 26-week fetus was impossible due to severe lung compression from a large diaphragmatic hernia; however, this important prognostic information was used to help counsel the parents prior to the demise of the fetus at 29 weeks. No postnatal data were available in two cases (one intrauterine death at 32 weeks and one medical termination at 35 weeks). Autopsy was not performed in either case. In all remaining patients, the MRI diagnosis was confirmed postnatally. A comprehensive summary of all studies performed is shown in Table 3.

# MRI sequences

Single-shot fast spin echo sequences produced T2-weighted images with black-blood-like contrast with excellent contrast between the vasculature and higher signal intensity of the thymus, lung and myocardium. These sequences were the most useful for assessment of extra-cardiac vascular anatomy as these structures are less mobile throughout the cardiac cycle (Figures 1–4). They were, however, still prone to artefact from gross fetal movement. It is important to note that three-dimensional interrogation of multi-slice sequences was not possible, as even small fetal movements produced non-contiguous anatomy between slices (see the *Future directions* section). An example of a standard postnatal 'black-blood' sequence is shown in Figure 5.

Balanced SSFP images produced bright blood images with good contrast between the blood pool and surrounding tissue, particularly intracardiac structures (Figures 6–8); however, they were more susceptible to motion and/or flow artefacts. Real-time bSSFP allowed for dynamic imaging of the beating fetal heart and was the most reliable sequence for assessment of moving structures, i.e. aneurysms and diverticulums

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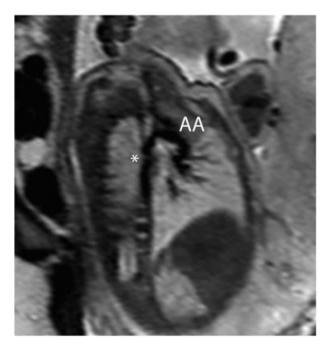


Figure 1 Single-shot fast spin-echo (SSFSE) black-blood image of the aorta in a 32-week fetus with coarctation of the aorta, confirmed postnatally. A characteristic indentation in the region of the aortic isthmus (a 'posterior shelf') is clearly visualised (\*). AA, aortic arch

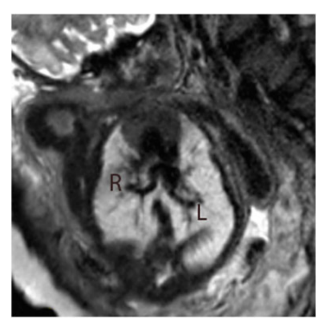


Figure 3 Single-shot fast spin-echo (SSFSE) black-blood image in a 35-week fetus with hypoplastic left heart syndrome with highly unusual pulmonary venous drainage. Tortuous, solitary pulmonary veins are seen bilaterally draining to confluence connected directly to the left atrium. R, right pulmonary vein; L, left pulmonary vein



Figure 2 Single-shot fast spin-echo (SSFSE) black-blood image in a high transverse orientation in a normal fetus. This corresponds to the standard 'three-vessel view' in fetal echocardiography showing the V-shaped connection between arterial duct (+) and the aorta (\*) adjacent to the superior caval vein (S). The thymus gland is seen anteriorly (T)

(Figure 6). Differences in enhancement patterns between sequences allowed for tissue characterisation of intracardiac masses by using established MRI techniques.<sup>28</sup>

# **DISCUSSION**

We instituted an open referral process for fetal MRI in a tertiary UK fetal cardiac centre. Whilst previous explorations of the clinical utility of fetal cardiac MRI have focused predominantly on intracardiac anatomy,5 the most common indication in our series was clarification of the extracardiac vascular anatomy, in keeping with established evidence that abnormalities involving these vessels, such as the coarctation of the aorta<sup>8,29,30</sup> and anomalies of the pulmonary veins,<sup>6,7</sup> are amongst the most challenging to define using ultrasound alone. Novel 3D and 4D echocardiographic techniques, such as spatiotemporal image correlation, have shown some potential for adjunctive imaging and can be visualised in three dimensions, but are still subject to many of the same limitations as conventional ultrasound. 12 Promisingly, however, and in keeping with animal models,31 our initial data suggest a promising strength of fetal MRI in vascular imaging, particularly using SSFSE sequences. To date, the only other report examining the clinical utility of fetal cardiac MRI used bSSFP sequences only, at a similar median gestational age to our series, interpreted blindly in 40 fetuses with CHD. Whilst this showed reasonable sensitivity for detection of abnormal intracardiac anatomy, the authors concede that MRI added no new diagnostic information when compared to ultrasound. In the same study, however, 15 fetuses were classified postnatally as having an abnormal aortic arch, of which five had been missed on ultrasound, six were missed on MRI and four were detected on both.5 Unlike in our study, no SSFSE sequences were used for vascular imaging.

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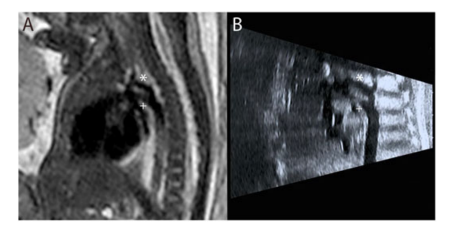


Figure 4 Single-shot fast spin-echo (SSFSE) black-blood image in a 32-week fetus with suspected coarctation of the aorta. The hypoplastic aortic arch (\*) is visualised superior to the dominant ductal arch (+) on fetal MRI (A), confirming prenatal ultrasound findings (B)



Figure 5 Postnatal MRI of the same patient at 5 months of age. No postnatal interventions have been performed. Despite prenatal findings, there is only mild coarctation, with isthmal hypoplasia and a posterior shelf at the point of insertion of the ductal ligament (+) (a vestige of the fetal arterial duct). The proximal aortic arch is within normal limits (\*)

The single most frequent single referral indication in our series was for suspected coarctation of the aorta, a notoriously challenging condition to predict prenatally due to both the technical difficulty of imaging this region and the obligatory patency of the arterial duct in fetal life, the postnatal closure of which can only then reveal life-threatening aortic constriction. Alongside extracardiac and genetic associations, a number of anatomical fetal risk factors for the condition have been suggested, such as the absolute and relative diameter of aortic isthmus, the presence of a posterior shelf, ventricular/great artery asymmetry and the persistence of a left-sided superior caval vein. In fetal MRI, imaging quality is unaffected by artefact from the fetal spine and

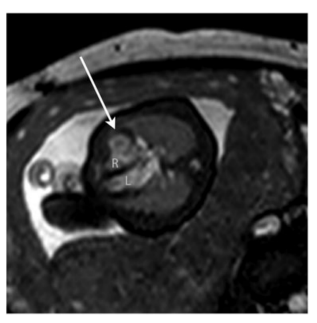


Figure 6 Still from a bSSFP cine (real-time) sequence in a 30-week fetus with a large right ventricular diverticulum (arrowed). R, right ventricle; L, left ventricle

ribcage, offering the potential for more accurate visualisation of these subtle anatomical factors. It is well established, however, that even a detailed anatomical assessment is not sufficient to accurately predict postnatal coarctation; for example, one of our cases with clearly visualised hypoplasia of the aortic arch before birth has gone on to develop only a mild form of coarctation, which has yet to require intervention (Figures 4 and 5). Indeed, it is likely that coarctation has a particularly complex aetiology, not yet fully understood, related to intrinsic myocardial abnormalities and/or a reduction in flow through the left heart. This has only recently started to be explored by using functional ultrasound techniques, rather than relying on anatomical findings alone.<sup>32</sup> The addition of more advanced MRI techniques in the future may help to define a more robust method of stratifying fetuses

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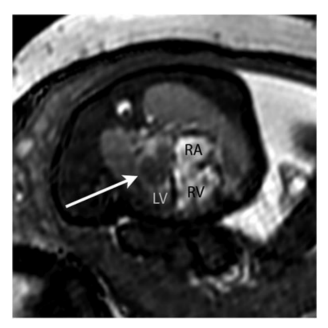


Figure 7 Still from a bSSFP cine (real-time) sequence in a 33-week fetus with a large left atrial mass (arrowed). The MRI differential diagnosis included myxoma, haemangioma or rhabdomyoma. The latter was confirmed postnatally. RA, right atrium; RA, right ventricle; LV, left ventricle

with suspected coarctation, providing a more detailed physiological assessment of affected fetuses (see the *Future directions* section).

In addition to vascular imaging, in three cases the combined use of SSFSE sequences provided the unique ability to characterise cardiac tumours before birth, a potential use of prenatal MRI with a high prognostic impact.<sup>33</sup> Multi-slice and real-time bSSFP sequences were most useful for intracardiac imaging, particularly for demonstrating the anatomy of gross lesions such as cardiac masses and diverticulums.

Finally, it should also be noted that several of the MRI studies reported in our series concomitantly provided important

complementary imaging that could directly or indirectly impact the cardiac prognosis, for example describing the devastating extent of pulmonary compression from a very large diaphragmatic hernia in a fetus that did not survive to term. Fetal MRI can also be used to assess lung parenchymal lesions and airway compromise in CHD<sup>34</sup> and has been used to demonstrate the presence of pulmonary lymphangectasia in a fetus with hypoplastic left heart syndrome, an important prognostic finding.<sup>35</sup> MRI of the fetal brain may provide crucial information to clinicians and parents regarding medium-term and long-term prognoses, which may have a profound effect on both prenatal and postnatal management independent of the underlying cardiac diagnosis.<sup>23,36,37</sup>

#### **Future directions**

As we observed, even small fetal movements can disrupt the spatial relationship of multi-slice sequences, making post-hoc analysis extremely challenging. The application of novel motion-corrected slice-volume registration algorithms to multiple MRI image stacks has been used to generate accurate 3D volumes of the fetal thorax. These exciting techniques have the potential to further enhance the capability of fetal MRI for vascular imaging, both by improving image quality and reducing the gestation age at which useful MR data can be obtained, and are currently under investigation by our unit.

Other advanced MRI techniques can offer novel means of evaluation of physiological parameters of the fetal heart and circulation. The application of retrospective 'metric-optimised' gating to phase contrast sequences can be used to quantify vascular flows and is currently being explored in a wide variety of settings. Alelated technology allows for the retrospective gating of single slice real-time sequences to dramatically improve the spatiotemporal resolution for cardiac imaging. The potential extension of these real-time sequences to whole-heart coverage allows for the possibility of performing comprehensive volumetric measurements throughout the cardiac cycle, providing estimates of chamber volumes and ventricular function.

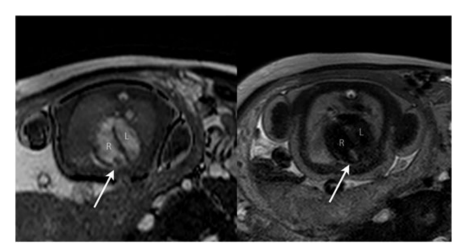


Figure 8 Balanced SSFP bright blood image (A) and single-shot fast spin-echo (SSFSE) black-blood image (B) in a 38-week fetus with an RV mass (arrowed). Based on the size, position and tissue signal on multiple sequences, a solitary ventricular rhabdomyoma was suspected; no other masses were seen including on concomitant fetal brain MRI. All findings were confirmed postnatally. R, right ventricle; L, left ventricle

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These innovations will not only help to bring prenatal cardiovascular MRI closer to its postnatal counterpart but could also offer the potential to enhance our fundamental understanding of the nature and aetiology of congenital heart defects – including challenging prenatal diagnoses such as coarctation of the aorta – and their interactions with other organ systems such as the placenta<sup>45</sup> and the developing fetal brain.<sup>37</sup> The incorporation of all of the above techniques is the subject of ongoing research within our departments.

# CONCLUSION

Abnormalities of the fetal vasculature such as coarctation of the aorta can be challenging to diagnose with ultrasound, even in experienced hands. Standard sequence MRI may have an adjunctive role in the assessment of the extracardiac vascular anatomy and tissue characterisation. Developments such as motion-corrected slice-volume registration and metric-optimised gated phase contrast flow quantification and real-time techniques, alongside increasingly accessible infrastructure, may further develop the role of MRI as an adjunct to echocardiography in the investigation and diagnosis of congenital heart disease in the fetus.

#### LIMITATIONS

We have described 22 cases that were selected for additional imaging from a single institution. Due to the increased likelihood

of image corruption from uncontrolled fetal movement, we tended to perform additional imaging in the third trimester. Whilst our median gestational age of 33 weeks is comparable to other published series, this does represent a current limitation of the modality. As imaging was performed for clinical evaluation, the acquisition and interpretation of MRI images was not blinded to previous assessments. In two fetuses that did not survive to term, no autopsy information was available due to lack of parental consent.

# WHAT'S ALREADY KNOWN ABOUT THIS TOPIC?

 Fetal cardiac MRI offers the potential to be a safe adjunct to echocardiography; however, there is no consensus on routine fetal indications or technical protocols for cardiovascular MRI in clinical practice. Previous studies have focused mainly on its utility for intracardiac imaging.

# WHAT DOES THIS STUDY ADD?

 Abnormalities involving the extracardiac vasculature were the most common group of referrals for additional imaging in clinical practice. Fast spin echo MRI sequences offer the potential for better visualisation of these structures than ultrasound alone. Additional MRI benefits such as tissue characterisation can add further value in selected cases.

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