Increased variability of fetal heart rate during labour: a review of preclinical and clinical studies

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Abstract
Increased fetal heart rate variability (FHRV) in intrapartum cardiotocographic recording has been variably defined and poorly understood, limiting its clinical utility. Both preclinical (animal) and clinical (human) evidence support that increased FHRV is observed in the early stage of intrapartum fetal hypoxaemia but can also be observed in a subset of fetuses during the preterminal stage of repeated hypoxaemia. This review of available evidence provides data and expert opinion on the pathophysiology of increased FHRV, its clinical significance and a stepwise approach regarding the management of this pattern, and propose recommendations for standardisation of related terminology.

KEYWORDS
acidaemia, cardiotocography, electronic fetal monitoring, fetal heart rate, increased variability, marked variability, neonatal morbidity, pregnancy, saltatory pattern, ZigZag pattern

Tweetable abstract: Increased fetal heart rate variability is parasympathetic-mediated and is caused by acutely deteriorating placental function.

1 INTRODUCTION
Cardiotocographic (CTG) electronic fetal heart rate (FHR) monitoring is the gold standard for assessing fetal wellbeing during labour, although it has a very poor positive predictive value for fetal hypoxic–ischemic neural injury.\textsuperscript{1–5} In recent years, efforts have been made to improve the accuracy of fetal monitoring and the evaluation of intrapartum adaptation by emphasising that an adequate interpretation of a CTG tracing relies not only on recognition of FHR patterns but also on understanding the fetal physiology behind the patterns.\textsuperscript{6–11}

The evaluation of FHR patterns is based on baseline FHR, and the depth, duration, timing and frequency of FHR decelerations and associated changes in FHR variability (FHRV).\textsuperscript{12–16} Moderate levels of FHRV are associated with a well-oxygenated fetus, whereas reduced or absent FHRV is a warning sign of fetal compromise.\textsuperscript{17–20} Intriguingly, there is growing evidence suggesting that increased FHRV, characterised by high-amplitude oscillations of FHR, may be important.\textsuperscript{21,22} Experimental studies in fetal sheep have demonstrated that fetal compromise can be associated with transiently increased FHRV.\textsuperscript{23–25} Recently, on the basis of intrapartal visual evaluation, increased FHRV has been associated with increased risk of fetal acidaemia at birth and early neonatal complications in human labour.\textsuperscript{26–30} The definition and classification of increased FHRV vary in the literature and in CTG monitoring.
therefore believe that a simplified definition and classification of increased FHRV is uncertain.27,43,44 The aims of the present review are to delineate the pathophysiology of increased FHRV, clarify the related terminology, and elucidate its potential clinical utility. We further propose that broadly there exist two hypoxaemia-related patterns of increased FHRV during labour: a pattern which has variously been called the ‘ZigZag’ or ‘saltatory’ pattern and is more often observed earlier in labour and not associated with deep repetitive FHR decelerations; and secondly, a pattern of increased FHRV observed in association with deep FHR decelerations in late labour. Suppression of FHRV, particularly in the presence of deep decelerations, remains an ominous sign that requires clinical attention. Nonetheless, it is increasingly being understood that this is not a universal finding in fetuses at risk of intrapartum asphyxia and hypoxic–ischemic injury.45–48 We therefore believe that a simplified definition and classification of these two patterns of increased FHRV will help to increase awareness and to alert birth attendants.

2 | FETAL ADAPTATION DURING LABOUR

During childbirth, uterine contractions result in repeated, brief reductions in uteroplacental perfusion, causing intermittent relative fetal and placental relative hypoxaemia. This reduction is associated with a transient fall in blood pH, base excess (BE) and oxygen tension, and a rise in carbon dioxide and base deficit (BD), even in normal, uncomplicated labour.49–52 The fetus compensates for moderate to severe intrapartum stress by activating the peripheral chemoreflex, leading first to FHR decelerations, presumptively to reduce myocardial oxygen demand, and second to trigger peripheral vasoconstriction preferentially to support blood flow to the heart, brain and adrenal glands.11,55–57 A healthy term fetus with a normally developed and functioning placenta is able to adapt to the typical frequency and intensity of uterine contractions without adverse consequences.58 However, if the interval between the contractions is too short, or placental function is compromised, prolonged impairment of oxygen delivery may lead to tissue hypoxia, metabolic acidemia, and persistent reduction in fetal cerebral oxygenation.59–64 If these episodes of hypoxaemia continue, fetal cardiac output is progressively compromised, leading to fetal hypotension and hypoxygenation, potentially resulting in hypoxic–ischemic brain injury.65–71 The progressive worsening intrapartum fetal hypoxaemia can be observed as changes in baseline FHR and deeper FHR decelerations,6,7,23,73 but once deeper decelerations are established there is typically little further change in FHR.74 Increased FHRV typically develops in the early stage of fetal hypoxia23,29,36,43 but can be seen also in FHR tracings of fetuses during the terminal stage of repeated asphyxia (Appendix S1).21,35,70

3 | PATHOPHYSIOLOGY OF INCREASED FHRV: INSIGHT FROM PRECLINICAL ANIMAL STUDIES

Increased FHRV patterns are seen rarely in antenatal FHR tracings, occurring almost exclusively during the active stage of labour.75,76 This suggests that labour-induced fetal stress, i.e. intermittent gas exchange disruption and consequent fetal hypoxaemia caused by intense uterine contractions, contributes to the intrapartum increased FHR pattern.29,77 Many studies in chronically instrumented fetal animals have used simulated intrapartum stress to improve clinical understanding of compensation mechanisms in the human fetus during birth, as well as the accompanying changes in FHR and FHRV.47,78–83 Animal studies can be broadly separated into those that study sustained periods of hypoxaemia and those that study intermittent periods of repeated hypoxaemia. The latter is more characteristic of the repetitive nature of hypoxaemia during intrapartum uterine contractions. Sustained periods of hypoxaemia can occur for example during sentinel events (e.g. placental abruption, cord prolapse, uterine rupture) but here we propose that sustained periods of mild hypoxaemia may have an underappreciated role in some instances of increased FHRV.

3.1 | Sustained hypoxaemia

In 1977, Dalton et al.78 reported increased FHRV during sustained moderate hypoxaemia in fetal sheep achieved by maternal inhalation of decreased oxygen, an observation that has been replicated multiple times.43,84,85 This is typically observed in the presence of bradycardia; for example in the study by Parer et al.43 FHR fell from 170 ± 22 to 139 ± 21 bpm after 5 minutes of hypoxaemia with a fall in mean pO2 from 20.7 to 11.3 mmHg.43 Likewise, more severe hypoxaemia (mean pO2 from 22.4 to 5.8 mmHg) induced by complete umbilical cord occlusion (UCO) results in increased FHR during the early minutes of UCO in association with marked bradycardia.47,86 Of particular interest, mild hypoxaemia in fetal sheep is associated with increased FHRV without a marked fall in FHR.43 Similarly, mild hypoxaemia in fetal monkeys was associated with an average fall in FHR from 199 to 178 bpm.87,88 However, it is notable that individual fetuses that show a less pronounced fall in pO2 displayed increased FHRV without a fall in FHR. In those that had a fall in FHR, this was often preceded by increased FHRV.88

3.2 | Repeated brief hypoxaemia

Repeated partial or complete UCOs have been used in fetal sheep to simulate the repetitive nature of hypoxaemia induced by uterine contractions. Each UCO is associated with a FHR deceleration, with more severe UCOs associated with deeper decelerations.4,89,90 These studies have shown that the early stages of fetal adaptation to repetitive brief hypoxaemia
are associated with increased FHRV between FHR decelerations. When UCO continues, the initial increase in FHRV diminishes and FHRV returns to baseline values. The terminal phase of UCO resulting in cardiovascular compromise and hypotension is associated with variable FHRV patterns. In the study by Westgate et al. in fetal sheep, two-thirds developed mild suppression of FHRV, with the remaining third showing a marked increase in FHRV. The mechanism underlying the differing FHRV patterns remains unknown.

### 3.3 Autonomic origin

During the prepartum period, in a healthy normoxic fetus, FHRV is complexly and constantly regulated by both the sympathetic and parasympathetic nervous systems, which are integrated at the sinoatrial node in concurrence with its own inborn rhythm. Over the past decades, the pathophysiology of increased FHRV during labour has been explained by the hypothesis that during rapid hypoxaemia the fetus has insufficient time to release catecholamines, leading to impaired central organ perfusion, and a magnified autonomic response caused by instability of sympathetic and parasympathetic nervous systems.

In contrast, more recent studies in fetal sheep have employed multiple forms of autonomic blockade during repeated UCOs to illustrate that FHRV during labour (once repetitive decelerations are apparent) is solely mediated by the parasympathetic nervous system, as recently reviewed. For example, neither complete β-adrenergic blockade with propranolol nor chemical sympathectomy with 6-hydroxydopamine neurotoxin reduced FHRV during repeated UCOs. In contrast, FHRV was abolished with parasympathetic blockade with either atropine sulphate or bilateral vagotomy. Likewise, during sustained periods of moderate fetal hypoxaemia induced by maternal hypoxaemia, atropine but not propranolol prevented the increase in FHRV. The mechanisms underlying the shift from dual sympathetic and parasympathetic control of FHRV during normoxia to parasympathetic dominance during both repetitive and sustained hypoxaemia are unknown but may involve feedback inhibition from high circulating catecholamine concentrations. Increased FHRV during both sustained and intermittent hypoxaemia is therefore likely mediated by increased parasympathetic activity, although the upstream mechanisms driving increased parasympathetic activity are likely distinct in each scenario.

### 4 HUMAN STUDIES

#### 4.1 Definitions and incidences

Periods of increased or high-amplitude FHRV that are occasionally observed in routine intrapartum FHR recordings have been referred to by multiple terms over the years. Initially, these patterns were referred to as 'marked irregularity' by Hon and Lee, 'rapid baseline fluctuations' by Caldeyro-Barcia et al. and 'high-amplitude oscillations' by Hammacher et al. The current literature includes descriptions of 'marked variability', the saltatory pattern and the ZigZag pattern. Table 1 gives a summary of the terminology, definitions and incidences of increased FHRV used by current human studies and clinical guidelines.

Although the saltatory pattern is well known, it is notable that in a recent study of a large obstetric cohort, only six (1.0%) of the 582 CTG recordings showed increased FHRV; the duration of a single increased FHRV episode lasted between 15 and 25 minutes, and in one (0.2%) case was >25 minutes (28 minutes). Furthermore, not a single increased FHRV pattern with a duration of >30 minutes was found in the cohort of 5150 childbirths. These findings are in agreement with suggestion that the saltatory pattern, as defined by FIGO and NICE, is almost nonexistent.

### 4.2 Association with fetal acidaemia and compromise

A recent study including 8580 births by Polnaszek et al. showed that marked variability patterns occurred in 6.7% of the 149 cases with cord blood acidaemia (UA pH <7.10). Marked variability was associated with an increased risk of elevated cord blood lactate and an increased risk of respiratory distress, although no association with composite neonatal morbidity was found. In their study, episodes of marked variability were most common during the final 10 minutes prior to birth, becoming progressively less common in the 2 hours studied prior to birth. The authors concluded that marked FHRV in isolation does not predict neonatal acidaemia. This is in agreement with the study by O’Brien-Abel and Benedetti, who concluded that a pattern of increased FHRV can be considered benign when observed in the absence of other abnormal periodic FHR changes, and in the presence of normal FHR variability before and after the high-amplitude oscillations of FHR.

Another recent study of 1070 fetuses who had fetal scalp blood sampled during labour showed that increased fetal scalp blood lactate level was associated with increased short-term FHRV. The association was observed in all four 30-minute epochs during the last 2 hours prior to birth. These findings support the concept that the early stages of intrapartum fetal hypoxaemia is associated with increased FHRV.

Recently, Tarvonen et al. investigated the episodes of increased FHRV ≥2 minutes in duration (the ZigZag pattern) in a retrospective study of 194 CTG tracings of fetuses with low Apgar scores and their 51 healthy controls. The ZigZag pattern was associated with both cord blood acidaemia and high concentrations of cord blood erythropoietin (EPO) at birth. Fetal hypoxaemia strongly stimulates EPO synthesis and hence high plasma EPO concentration is a marker of the severity of fetal hypoxaemia.
Further work by Tarvonen et al. has highlighted the association between the ZigZag pattern and FHR decelerations. The presence of increased FHRV or late decelerations, or both, in the CTG recordings during the last 2 hours of labour has been shown to increase by three-fold the likelihood of severe hypoxaemia-related complications (i.e. UA pH <7.10 and/or BE < -12.0 mEq/l and/or 5-minute Apgar score <4 and/or intubation for resuscitation and/or grade II/III neonatal encephalopathy) in newborn infants. A CTG recording with both ZigZag pattern and late decelerations occurred in 76.9% (123/160) of cases with severe neonatal complications but in only 5.6% (201/3620) of cases with no complications.

Strikingly, in the vast majority of cases, a rapid transition was observed from an initially 'normal' or 'reassuring' FHR trace without decelerations, to the pattern of increased FHRV and the subsequent appearance of late decelerations. The median time between the end of the first ZigZag episode and the onset of late decelerations was 9 minutes.

Two previous case reports, and one study with a population of high-risk patients, further support the concept that the concurrent occurrence of increased FHRV and late decelerations indicates an increased risk of severe hypoxaemia.
4.3 | Risk factors

Observations in human fetuses suggest that maternal and fetal background factors may play a role in the origin and development of intrapartum increased FHRV. Among a cohort of 5150 childbirths, the ZigZag pattern only occurred in term and post-term pregnancies, with an increasing incidence with advancing gestation. This finding confirms a previous observation that the presence of increased FHRV is rare in preterm fetuses. These observations are in agreement with a study in which umbilical cord plasma EPO levels increased significantly after 40 weeks of gestation in pregnancies with spontaneous onset of labour, suggesting that placental insufficiency after 40 weeks of gestation may contribute to the occurrence of the hypoxaemia-related increased FHRV.

Recently, male sex of the fetus, nulliparous pregnancy and post-term pregnancy of ≥42 weeks were independently associated with the ZigZag pattern. Consequently, the presence of any of these three independent risk factors, or a combination of them, increased the likelihood of the occurrence of increased FHRV up to 16-fold. Another recent study showed that fetuses of women with gestational diabetes mellitus (GDM) were more likely to have the ZigZag pattern than in pregnancies of women with no GDM. Moreover, fetuses of GDM mothers with two abnormal oral glucose tolerance test values had the strongest association with the intrapartal ZigZag pattern. These findings are in agreement with previous studies in GDM pregnancies, in which impaired glucose metabolism diagnosed in early pregnancy, as a result of more severe form of GDM, is associated with both functional and structural placental changes.

It is further worth appreciating that both maternal and fetal infections have been previously linked with increased FHRV. In preterm fetal sheep, acute exposure to high-dose lipopolysaccharide (a bacterial cell wall component that induces a systemic inflammatory response) triggered an increase in FHRV between 2 and 4 hours after exposure. This increase in FHRV, however, is only observed after an acute inflammatory stimulus that triggers a rapid inflammatory and cardiovascular response, whereas a stable inflammation response has little effect on FHRV even if prolonged. In two recent studies, 4% (nine of 224) and 33% (four of 12) of human fetuses of COVID-19 parturients showed the ZigZag pattern in CTG tracing. Similarly, chorioamnionitis and funisitis have been associated with the ZigZag pattern during labour. It remains unclear whether these human findings are due to infection/inflammation exacerbating hypoxaemia during labour or are an independent effect of infection/inflammation increasing FHRV. Unfortunately, the effect of infection/inflammation on FHRV during labour-like hypoxaemia has not been studied in animals.

4.4 | Nomenclature

Increased FHRV is defined as baseline amplitude changes of ≥25 bpm. Based on recent findings, even short episodes of ≥1 minute of increased FHRV are associated with unfavourable fetal and neonatal outcomes. Nonetheless, confusion about terminology used to describe increased FHRV patterns has been a longstanding problem. It is well known that standardised terminology to describe intrapartum CTG may avoid miscommunication among clinicians caring for parturients and can improve the safety of childbirth. Furthermore, unified terms when evaluating whether FHR patterns are reassuring or nonreassuring (i.e. normal, suspicious or pathological) help providers to decide when to intervene. Currently, according to FIGO, RCOG and ACOG CTG guidelines and the wider literature, a number of different terms of increased FHRV patterns are used that actually describe the same FHRV phenomenon. We therefore propose that terms such as ‘saltatory pattern’, ‘ZigZag pattern’ and ‘marked variability’ should be abandoned, and the common term ‘increased variability’ should be used in clinical guidelines in the same way the term ‘reduced variability’ already reflects decreased levels of FHRV.

5 | LINKING THE PRECLINICAL AND CLINICAL FINDINGS

Collectively the evidence from human labour suggests that increased FHRV broadly occurs in two situations: (1) in association with repetitive FHR decelerations, manifesting as brief periods of increased FHRV mainly between decelerations, and (2) increased FHRV without repetitive FHR decelerations. This second pattern is more typically observed earlier during labour and manifests as a relatively longer period of increased FHRV. Figure 1 shows typical examples of these two patterns in intrapartum CTG recordings. In this section we attempt to parallel human observations with insight from preclinical animal studies in order to explain the pathophysiological origins of FHRV.

5.1 | Increased FHRV with concurrent decelerations

The occurrence of increased FHRV associated with FHR decelerations appears to match the scenario described above in the ‘repeated brief hypoxaemia’ section. These animal studies illustrate that FHRV increases firstly during the early phase of adaptation to repetitive brief hypoxaemia, and therefore increased FHRV can be expected to occur early after the first appearance of frequent FHR decelerations during labour. During prolonged exposure to intense repetitive brief hypoxaemia, fetal cardiovascular adaptation can progressively fail in association with worsening acidemia. Animal evidence suggests that this late phase can be associated with either increased or reduced FHRV. Increased FHRV can sometimes be observed during the nadir of repetitive brief decelerations, but there is little understanding about whether this has any different implications than increased FHRV between decelerations.

The human findings that increased FHRV, particularly towards the end of labour, is associated with increased risk of fetal
Increased variability of fetal heart rate

Increased FHRV without concurrent decelerations

The pattern of increased FHRV during labour observed without FHR decelerations (including the patterns previously called saltatory or ZigZag patterns) has remained poorly understood. This pattern is more commonly observed in early labour. In considering the potential pathophysiological mechanisms, the following observations are important:

- The absence of FHR decelerations suggests the absence of significant repetitive brief hypoxaemia episodes.
- The pattern is observed in the presence of either a stable baseline FHR or on top of a modest fall in baseline FHR (although this may be obscured by increased FHRV).
- The duration of these patterns is longer than a typical uterine contraction.
- The majority of risk factors associated with the pattern are equally risk factors for impaired placental function, including severe form of GDM, post-term pregnancy, elevated fetal cord blood EPO concentration, high placental weight relative to birthweight, and the occurrence of late decelerations.

We propose that this pattern represents a comparatively mild but more prolonged hypoxaemia than the pattern observed in association with repetitive variable FHR decelerations, consistent with animal experiments that have modelled ‘sustained hypoxaemia’. Indeed, there is evidence in both fetal sheep and monkeys that mild hypoxaemia can trigger a parasympathetic-mediated increase in FHRV either in the absence of a fall in FHR, or with only a modest fall in FHR. The fact that these patterns are frequently not synchronised with uterine contractions, suggests that they reflect an acute deterioration of placental function leading to a relatively prolonged period of mild fetal hypoxaemia. Supporting this, in the recent studies by Tarvonen et al. a close temporal association was found between an initially ‘normal’ or ‘reassuring’ FHR trace without decelerations, the appearance of the pattern of increased FHRV and the subsequent appearance of late decelerations. This rapid appearance of FHR decelerations suggests a deterioration of placental reserve or uteroplacental gas exchange.
Alternatively, and considering the association of the increased FHRV with GDM, the fetus is an obligate user of glucose and thus increases oxygen consumption during acute hyperglycaemia which can lead to hypoxaemia and acidemia.\textsuperscript{137–133} An acute maternal hyperglycaemia during labour may therefore exacerbate labour-induced hypoxaemia and contribute to the pattern of increased FHRV. Consistent with this speculation, GDM is associated with an increased risk of fetal hypoxaemia and acidemia in labour.\textsuperscript{30,134–136}

It is further notable that greater duration of a single increased FHRV episode was associated with higher risk of fetal compromise: a mean duration of 4.8 minutes was associated with no neonatal complications, 6.5 minutes was associated with moderate complications, and 10.7 minutes was associated with severe complications.\textsuperscript{29} It is important to note that this pattern of increased FHRV predominantly occurs during early labour but is associated with later outcomes at birth. Based on these observations, we propose that the pattern of increased FHRV represents mild hypoxaemia, which is not a significant threat to the fetal wellbeing but identifies a fetus with impaired uteroplacental function that is at increased risk of failure to adapt to the challenge of repetitive hypoxaemia during labour. Supporting this concept, Table 2 shows that among a cohort of term fetuses,\textsuperscript{29} a high placental weight to birthweight ratio was associated with higher rates of increased FHRV during labour. It has been previously reported that placental enlargement may be an indicator of chronic fetoplacental hypoxaemia and is associated with increased risk of fetal compromise.\textsuperscript{112,113} The association of higher risk of complications with longer durations of increased FHRV likely reflects more prolonged hypoxaemia and may reflect more significant impairment of placental function.

### 6 | CLINICAL CONSIDERATIONS

The need for research of increased FHRV in conjunction with other FHR patterns has been called for in previous papers.\textsuperscript{27,137,138} Based on the cohort of 4988 term deliveries including 160 cases with hypoxaemia-related fetal and neonatal complications,\textsuperscript{29} we estimated the number of caesarean deliveries that need to be performed (NNT) to prevent one case of cord blood acidemia or neonatal hypoxaemia-related morbidity (Table 3).\textsuperscript{139} In the setting of the combined

| TABLE 2 | Odds ratios (ORs) with 95% confidence interval (CI) for occurrence of increased FHRV pattern in CTG recording according to quartiles of placental weight to birthweight ratio in 4988 term deliveries |
| Quartiles of placental weight to birthweight ratio | Increased FHRV | Increased FHRV | Crude OR | 95% CI | Adjusted\textsuperscript{a} OR | 95% CI |
| Present (n = 582) | Absent (n = 4406) | |
| 1st | 110 | 1102 | Reference | Reference |
| 2nd | 121 | 1101 | 1.10 | 0.84–1.44 | 1.05 | 0.80–1.41 |
| 3rd | 155 | 1103 | 1.41 | 1.09–1.82 | 1.40 | 1.07–1.80 |
| 4th | 196 | 1100 | 1.79 | 1.39–2.29 | 1.77 | 1.37–2.27 |

Note: The ORs and 95% CIs for the increased FHRV pattern were estimated by fitting logistic regression models. The logistic regression analysis was performed using R version 3.6.0.

Abbreviations: CI, confidence interval; CTG, cardiotocography; FHRV, fetal heart rate variability; OR, odds ratio.

\textsuperscript{a}Adjusted for parity, gestational age at delivery, maternal age ≥35 years, gestational diabetes mellitus, pre-eclampsia, maternal fever ≥38.0°C, smoking, fetal sex and birthweight z-score.

| TABLE 3 | Estimated number of caesarean deliveries needed to be performed for the CTG patterns predicting hypoxaemia-related fetal and neonatal complications at 120–90 minutes and at 120–0 minutes before birth in term fetuses (n = 4988) |
| FHR pattern | Number (n = 4988) | NNT | Number (n = 4988) | NNT |
| CTG with prolonged decelerations (with a duration of ≥3 min) and/or tachycardia episodes and/or reduced variability and/or uterine tachysystole but without increased FHRV pattern or late decelerations | 3320 (66.6) | 78.0 | 3851 (77.2) | 60.1 |
| CTG with late decelerations (increased FHRV pattern overlooked) | 253 (5.1) | 37.5 | 1934 (38.8) | 49.4 |
| CTG with increased FHRV pattern or late decelerations | 214 (4.3) | 23.3 | 1565 (31.4) | 28.7 |
| CTG with increased FHRV pattern or late decelerations or both | 311 (6.2) | 16.8 | 2096 (42.0) | 21.0 |
| CTG with increased FHRV pattern and late decelerations | 97 (1.9) | 3.9 | 531 (10.6) | 9.0 |

Note: Data are presented as number (%). Increased FHRV pattern: FHR baseline amplitude changes of >25 bpm with a duration of ≥2 minutes. Hypoxaemia-related fetal and neonatal complications: UA pH <7.10 and/or BE < −12.0 mEq/l and/or 5-minute Apgar score <4 and/or intubation for resuscitation and/or grade II/III neonatal encephalopathy.

Abbreviations: CTG, cardiotocography; FHR, fetal heart rate; NNT, number needed to treat.\textsuperscript{139}
occurrence of increased FHRV pattern followed by repetitive late decelerations at 120–90 minutes before birth, the NNT was suggested at four, which is relatively low (Table 3). However, when evaluated over the last 2 hours (120–0 minutes) of labour, the NNT was nine (Table 3). Similarly, the combined occurrence of increased FHRV and late decelerations at 120–90 minutes before birth had an adjusted odds ratio (aOR) of 33.0, and 120–0 minutes before birth an aOR of 5.4, for hypoxaemia-related morbidity when compared with cases without these FHR patterns. Hypothetically, these findings may reflect a longer exposure time to intrapartum hypoxaemia in fetuses showing the pattern of increased FHRV and late decelerations occurring at or earlier than 90 minutes before birth compared with fetuses showing the same pattern FHR patterns for the first time immediately before birth. These NNTs are comparable to those which Cahill et al. have reported concerning FHR deceleration area, and deceleration area combined with fetal tachycardia, as discriminatory for fetal acidemia and neonatal morbidity (NNTs five and six, respectively).

Previously, Downs and Zlomke have suggested that a clinician should consider intrauterine resuscitation methods (Appendix S1) to improve the fetal environment in utero when an increased FHRV pattern occurs in isolation in an intrapartum CTG tracing. Indeed, in a recent study, the majority of NICHD Category II FHR tracings (in which category the increased FHRV pattern is included) were improved to normal Category I within 60 minutes of intrauterine resuscitation interventions.

### 7 | CONCLUSIONS

In recent studies with relatively large obstetric cohorts, the occurrence of increased FHRV in intrapartum CTG tracing has been associated with fetal acidemia and a greater risk of neonatal complications. These studies have further associated increased FHRV with severe form of GDM, post-term pregnancy, elevated fetal cord blood EPO concentration, high placental weight relative to birthweight, as well as the occurrence of late decelerations of FHR, all of which are associated with fetal hypoxaemia. Although caution is needed when extrapolating from animal studies, multiple parallels can be observed across species, suggesting a conserved FHRV response to hypoxaemia. In particular, we here present a new hypothesis to explain the pattern of increased FHRV that spans multiple contractions early in labour, and suggest that this mild hypoxaemia may represent acute deterioration of placental function that identifies the fetus at risk of failing to adapt to labour. Indeed, the risks factors for this pattern all impair placental function; however, further work is needed to understand the potential mechanisms. Although this pattern is not synchronised with contractions, it is rare antenatally, suggesting that uterine contractions, or changes in uterine tone, during labour are part of the mechanism. Hence, computerised spectral analysis of FHRV is an important but still developing area of FHR monitoring. Future work should seek to examine whether patterns of increased FHRV occurring early and late during the process of fetal compromise can be distinguished by spectral analysis and other modern signal analysis techniques.

We believe there is already sufficient evidence to illustrate that increased FHRV observed in early labour can represent an early warning sign of impaired placental function, and additionally that increased FHRV in the setting of deep repetitive decelerations can indicate impending fetal compromise. An important corollary, and a significant departure from standard teaching, is that although suppression of FHRV is undoubtedly an ominous sign, the absence of suppressed FHRV cannot be relied upon to exclude fetal compromise. The specific features of increased FHRV, alone and combined with late decelerations, should be incorporated into clinical interpretation, electronic fetal monitoring guidelines and computerised algorithms to improve the performance of fetal surveillance during labour. The authors believe that the evidence presented in this review may improve fetal surveillance, enable timely conservative intrauterine resuscitation measures and recognition of loss of fetal compensatory reserve, and may improve the clinical decision making on intrapartum CTG recordings with episodes of increased FHRV. Standardisation of the terminology surrounding increased FHRV will likely improve the uptake of this knowledge.

### AUTHOR CONTRIBUTIONS

MJT conceived this review. MJT and CAL undertook the publication search. MJT prepared the first draft of the manuscript. MJT, CAL, SA, AJG and KAT contributed to interpreting the findings, editing and critically revising the manuscript, approved the final version of the manuscript and have agreed to be accountable for all aspects of the work. All persons who qualify for authorship are listed and all persons designated as authors, qualify for authorship.

### ACKNOWLEDGEMENTS

None.

### CONFLICT OF INTERESTS

Open access funding provided by University of Helsinki including Helsinki University Central Hospital. Mikko Tarvonen has received support from the Foundation for Paediatric Research, Finska Läkaresällskapet and Olga & Vilho Linnamo Foundation. Sture Andersson has received grants from a Special Governmental Subsidy for Clinical Research, Finska Läkaresällskapet, and the Society for Paediatric Research in Finland. Christopher Lear and Alistair Gunn received funding for this review from the Health Research Council of New Zealand. The sponsors had no role in the study design; collection, analysis, or interpretation of data, writing of the report, or in the decision to submit the report for publication. The authors have no interests to declare. Completed disclosure of interest forms are available to view online as supporting information.
DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

ETHICAL APPROVAL
No human or animal subjects were involved in this manuscript.

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How to cite this article: Tarvonen MJ, Lear CA, Andersson S, Gunn AJ, Teramo KA. Increased variability of fetal heart rate during labour: a review of preclinical and clinical studies. BJOG. 2022;129:2070–2081. https://doi.org/10.1111/1471-0528.17234