

1 Heart morphology differences induced by intrauterine  
2 growth restriction and preterm birth measured on the  
3 ECG at preadolescent age

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22 **Abstract**

Intrauterine Growth Restriction (IUGR) and premature birth are associated with higher risk of cardiovascular diseases throughout adulthood. The aim of this study was to evaluate the influence of these factors in ventricular electrical remodeling in preadolescents. Electrocardiography was performed in a cohort of 33-IUGR, 32-preterm with appropriate weight and 60 controls. Depolarization and repolarization processes were studied by means of the surface ECG, including loops and angles corresponding to QRS and T-waves. The angles between the dominant vector of QRS and the frontal plane XY were different among the study groups: controls [20.03°(10.11°-28.64°)], preterm [25.48°(19.79°-33.56°)], and IUGR [27.77°(16.59°-33.23°)]. When compared to controls, IUGRs also presented wider angles between the difference of QRS and T-wave dominant vectors and the XY-plane (5.28°±12.15°

vs  $0.49^\circ \pm 14.15^\circ$ ,  $p < 0.05$ ) while preterms showed smaller frontal QRS-T angle ( $4.68^\circ (2.20^\circ - 12.89^\circ)$  vs  $6.57^\circ (2.72^\circ - 11.31^\circ)$ ,  $p < 0.05$ ). Thus, electrical remodeling is present in IUGR and preterm preadolescents, and might predispose them to cardiovascular diseases in adulthood. Follow-up studies are warranted.

23 *Keywords:* Premature birth, Intrauterine Growth Restriction, ventricular  
24 electrical remodelling

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## 25 **1. Introduction**

26 For many years, genetics and lifestyle have been considered as the main  
27 cardiovascular risk factors. However, by the 90s, Barker and colleagues  
28 demonstrated a strong association between cardiovascular diseases (CVD)  
29 and birthweight [1], and proposed that low birthweight may produce struc-  
30 tural and functional changes in key organs in postnatal life that predispose to  
31 CVD in adulthood, from which the concept of fetal programming emerged [1].  
32 In this paper, we studied two leading causes of low birthweight: intrauterine  
33 growth restriction (IUGR) and preterm delivery.

34 IUGR affects 7-10% of pregnancies and is defined as the failure of a fe-  
35 tus to achieve its growth potential [2]. It is usually associated with pla-  
36 cental insufficiency that determines fetal hypoxia, undernutrition and pres-  
37 sure/volume overload. Several studies have demonstrated that IUGR fetuses  
38 [3] and children with earlier IUGR [4] show significant changes in cardiac  
39 structure and function in the form of more spherical hearts with reduced  
40 longitudinal motion and impaired relaxation. On the other hand, prematu-  
41 rity represents 4-10% of deliveries and is defined by birth before 37 completed  
42 weeks of gestation. Preterm birth has multiple causes, including spontaneous  
43 preterm labor, intraamniotic inflammation and/or infection, preterm rupture  
44 of membranes, and labor induction due to IUGR or preeclampsia. Recently,  
45 increased cardiac mass together with short and small left ventricles have been  
46 described in adults born preterm [5].

47 In addition, both IUGR and prematurity have been associated with in-  
48 creased CVD in adulthood [6] as well as with arrhythmias i.e sudden death  
49 syndrome [7] and apnea-bradycardia episodes [8] in infants. However, little is  
50 known about the presence of electrical changes associated with these condi-  
51 tions that could predispose to long-term consequences. It has been proposed  
52 that electrical remodelling could occur primarily or secondary to the cardiac

53 structural remodelling described in these conditions [9].

54 Hence, the aim of the present study was to identify signs of ventricular  
55 electrical remodelling in preterm and growth restricted babies on reaching  
56 preadolescence by assessing changes in ventricular depolarization and re-  
57 polarization using the QRS complex and T-wave, respectively. This was  
58 assessed by analyzing the direction of the QRS and T vectors, as well as  
59 the angle between them, which is supposed to reflect possible deviations be-  
60 tween ventricular depolarization and repolarization. Finally, an analysis of  
61 QRS and T loop morphology was also performed in order to study in detail  
62 the presence of abnormalities in the ECG signal.

## 63 2. Materials and Methods

### 64 2.1. Study populations

65 The study population included 125 preadolescents, whose surface 12-  
66 lead ECG was recorded at a sampling rate of 1000 Hz in a tertiary cen-  
67 ter. 33 subjects had severe IUGR with medically-induced preterm delivery,  
68 32 were spontaneously born preterm with appropriate weight for gestational  
69 age (AGA) and 60 normally grown controls born at term. From now on,  
70 these three groups will be denoted by preterm-IUGR, preterm-AGA and  
71 controls. IUGR was defined by low birthweight below 10th centile and ab-  
72 normal umbilical artery Doppler (pulsatility index above the 95th centile),  
73 while adequate growth was considered if birthweight was above 10th centile  
74 for gestational age according to local standards [10]. Preterm birth was de-  
75 fined by delivery before 37 completed weeks of gestation. Gestational age was  
76 calculated by first-trimester crown-rump length measurement by ultrasound.  
77 Cases and controls were selected from a previously published cohort study  
78 that included 200 children at 2-6 years of age whose gestational age ranged  
79 from 25 to 41 weeks [4], which was conducted in a tertiary referral university  
80 hospital in Barcelona, Spain. We contacted all previous study participants  
81 to be included in this follow-up (6 years after the previous cardiovascular  
82 assessment) and 125 accepted to be recruited for the present study.

83 Digital standard 12-lead surface ECGs were recorded using a Gem Heart  
84 One recorder (Gem-Med SL, Spain), at an equivalent paper speed of 50mm/s  
85 and a gain of 10 mm/mV. This recorder provided digital recordings in SCP  
86 format. The acquisition process was performed by a trained nurse.

87 *2.2. Signal Preprocessing*

88 ECG was delineated so that QRS complexes and T waves, as well as their  
 89 onset and end, were detected. This was performed by a wavelet transform  
 90 delineation technique [11]. Then, baseline wander was reduced by means of  
 91 cubic splines [12]. The vectorcardiogram was synthesized using the inverse  
 92 Dower matrix [13].

93 *2.3. Loop alignment and averaging*

94 In order to attenuate noise, respiratory influence and muscular activity,  
 95 depolarization and repolarization loops were first aligned with respect to a  
 96 reference loop and then averaged. The reference loop is selected as the first  
 97 visually checked normal loop,  $\mathbf{Z}_R$ , consisting of a  $3 \times (k + 2\Delta)$  matrix con-  
 98 taining at each row the leads X, Y and Z, respectively. Spatial and temporal  
 99 alignment was performed in terms of scaling, rotation and time synchroniza-  
 100 tion of the loops, as presented in [14].

Thus, three transformations were considered to perform the alignment.  
 This process can be described as:

$$\mathbf{Z} = \alpha \mathbf{Q} \mathbf{Z}_R \mathbf{J}_\tau \quad (1)$$

101 where  $\mathbf{Z}$  and  $\mathbf{Z}_R$  denote the  $3 \times K$  and  $3 \times (K + 2\Delta)$  matrices that contain  
 102 in each row  $K$  or  $K + 2\Delta$  samples corresponding to leads X,Y,Z of the loop  
 103 to be aligned and the reference loop, respectively. Scaling was controlled by  
 104 the positive parameter  $\alpha$ , that allows loop expansion or contraction, whereas  
 105 rotational changes of the heart were introduced by the  $3 \times 3$  rotation matrix  
 106  $\mathbf{Q}$ . Finally, time synchronization was described by the integer time shift  $\tau$   
 107 in the shift matrix  $\mathbf{J}_\tau$ , which was defined as

$$\mathbf{J}_\tau = \begin{bmatrix} 0_{\Delta+\tau} \\ I \\ 0_{\Delta-\tau} \end{bmatrix}$$

108 where  $\tau = -\Delta, \dots, \Delta$ , the identity matrix  $I$  is  $K \times K$ , and the top and bottom  
 109 zero matrices are  $(\Delta + \tau) \times K$  and  $(\Delta - \tau) \times K$ , respectively. Parameter  $\Delta$   
 110 corresponds to the  $2\Delta$  additional samples that the reference loop  $\mathbf{Z}_R$  has in  
 111 order to allow for time synchronization of observations of different subsets of  
 112  $K$  samples.

The optimal estimates for  $\alpha$ ,  $\tau$  and  $\mathbf{Q}$  were determined by solving the minimization problem

$$\epsilon_{min}^2 = \min_{\alpha, \mathbf{Q}, \tau} \|\mathbf{Z} - \alpha \mathbf{Q} \mathbf{Z}_R \mathbf{J}_\tau\|_F^2 \quad (2)$$

113 where  $\epsilon^2$  represents the error, calculated as the Frobenius norm of the differ-  
 114 ence between the actual and the reference loop subjected to the transforma-  
 115 tions.  $\epsilon^2$  is minimized by first finding the estimates for  $\alpha$  and  $\mathbf{Q}$  for every  
 116 value of  $\tau$  and then selecting the value of  $\tau$  that minimizes the error. For a  
 117 fully detailed explanation of the alignment process see [14].

118 Depolarization loops are aligned and the estimated transformations for  
 119 each heartbeat are applied to align the repolarization loops. Then, depolar-  
 120 ization and repolarization loops are averaged in order to obtain a clean and  
 121 robust mean QRS and T loop. Then, the dominant vectors of the average  
 122 loops, denoted by  $\mathbf{v}_{QRS}$  and  $\mathbf{v}_T$ , are obtained by averaging the whole set of  
 123 vectors that form each loop. These set of vectors describe the dominant di-  
 124 rection of the electrical wavefront along the depolarization ( $\mathbf{v}_{QRS}$ ) and the  
 125 repolarization ( $\mathbf{v}_T$ ) processes.

#### 126 *2.4. Angles estimation based on dominant vectors of depolarization and re-* 127 *polarization loops*

128 We estimated the angle between the dominant vectors of depolarization  
 129 and repolarization phases in the three dimensional space ( $\theta_{RT}$ , the so-called  
 130 spatial QRS-T angle). It is defined as the angle measured in the plane that  
 131 formed by the maximum vectors of the QRS complex and the T-wave. It  
 132 usually differs from the angle of the projections of the QRS and T axes in  
 133 the frontal plane XY, denoted by  $\theta_{RT-XY}$ , and which was also calculated in  
 134 our study.

135 Next, we estimated the absolute angles between vectors  $\mathbf{v}_{QRS}$  and  $\mathbf{v}_T$  and  
 136 the three orthogonal planes,  $\Phi_{R-\mathcal{P}}$  and  $\Phi_{T-\mathcal{P}}$ , where  $\mathcal{P} \in \{XZ, XY, YZ\}$   
 137 denotes each orthogonal plane formed by the two directions referred to  $\mathcal{P}$ .  
 138 Finally, we also estimated the difference between them with respect to each  
 139 orthogonal plane.

#### 140 *2.5. Loop morphology*

141 Loop morphology was assessed by means of planarity and roundness mea-  
 142 surements. *Planarity* is a measure of how well the VCG can be approximated  
 143 by a plane or whether it is not possible, since the VCG is so curved that it is

144 really distributed along the 3D spatial loop. *Roundness* measurements were  
145 referred to 2D planar projections of the VCG, taking values in the range [0-1]  
146 (where 1 means that the projection is closer to a circle than to a straight line,  
147 whereas roundness close to 0 reflects the opposite and intermediate situations  
148 - an ellipsoidal shape of the VCG-) with different axes ratios as function of  
149 roundness.

If we denote the eigenvalues of the average loop matrix as  $\lambda_i, i = 1, 2, 3$ , sorted in descending order, the planarity of the loops is defined as

$$\rho_L = \frac{\lambda_3}{\lambda_1} \quad (3)$$

where  $\rho$  can take values between zero (entirely planar loop) and one (loop that equally extends into the three dimensions). Features will be obtained for both the depolarization and repolarization loops, being indicated by the subscript  $L$  values  $R$  or  $T$ , respectively. Planarity can also be analysed by features  $\sigma_L$  and  $\delta_L$ , which can take values in the range [0-1] (reflecting more planar loops when  $\sigma_L$  is close to 1 and  $\delta$  is close to 0):

$$\sigma_L = \frac{\lambda_1 + \lambda_2}{\lambda_1 + \lambda_2 + \lambda_3} \quad (4)$$

$$\delta_L = \frac{\lambda_3}{\lambda_1 + \lambda_2} \quad (5)$$

QRS and T wave complexity was defined as the ratio between the second to the first eigenvalues, denoted by  $C$ :

$$C_L = \frac{\lambda_2}{\lambda_1} \quad (6)$$

The geometrical interpretation of  $C$  refers to the roundness of the loop [15]. This feature was also obtained for each plane  $\mathcal{P} \in \{XZ, XY, YZ\}$ , corresponding to the loop projection onto each orthogonal plane. Eigenvalues obtained on the projection were  $\nu_1, \nu_2$ , sorted in descending order. Thus, the roundness of the loops projected on each plane was defined as

$$\varrho_{L-\mathcal{P}} = \frac{\nu_2}{\nu_1} \quad (7)$$

150 *2.6. Statistical analysis*

151 A descriptive analysis including angles, planarity and characteristics was  
 152 performed using mean  $\pm$  standard deviation or median (interquartile range)  
 153 to assess the differences between each group (Preterm-AGA and Preterm-  
 154 IUGR) and controls. The results were compared by t-test or Wilcoxon-  
 155 Mann-Whitney to compare means and standard deviations or medians and  
 156 interquartile ranges, respectively, and  $\chi^2$  to compare percentages. Multivari-  
 157 ate analysis by linear or quantile regression was used to adjust for age, height,  
 158 gender and heart rate.

159 **3. Results**

160 Estimated angles and morphology measurements were obtained on the  
 161 cohort of 125 subjects described in Section 2.1, who form the three groups  
 162 under study: 60 controls, 32 preterm-AGA, 33 preterm-IUGR. The median  
 163 and interquartile ranges of the explored parameters are shown in Tables 2  
 164 and 3, whereas perinatal and anthropometric characteristics of the study  
 165 populations are shown in Table 1.

Table 1: Baseline and perinatal characteristics of the study populations. AGA (appropriate growth for gestational age), IUGR (intrauterine growth restriction). Data are mean  $\pm$  standard deviation or median (interquartile range). \*p<0.05 by T-test, Wilcoxon-Mann-Whitney or  $\chi^2$  as compared to controls.

	Controls n=60	Preterm AGA n=32	Preterm IUGR n=33
Birthweight (g)	3444 $\pm$ 379.4	1872 $\pm$ 614.7*	1162 $\pm$ 351.9*
Gestational age at delivery (weeks)	39.5 $\pm$ 1.3	31.6 $\pm$ 3.1*	32.0 $\pm$ 2.4*
Birthweight centile	56(37-81)	48(30-85)	0(0-0)*
Caucasian ethnicity (%)	95.1	93.9	80.0*
Male gender (%)	53.2	39.4	37.1
Age (years)	12.1(10.5-12.4)	10.4(8.5-11.3)*	9.41(8.3-11.4)*
Heart rate (bpm)	77.66 $\pm$ 11.83	88.30 $\pm$ 10.94*	81.4 $\pm$ 9.26*
Height (cm)	148.5(137.8-154.3)	135.2(129.7-145.1)*	135(129.2-147.5)*
Weight (kg)	43.3(35.9-48.0)	32.8(28.2-41.1)*	34.4 (28.5-41.7)*
Body mass index (kg/ m <sup>2</sup> )	19.0(17.1-20.6)	17.24(15.9-20.1)	18.1(16.4-19.6)

166 As expected, both preterm-IUGR and preterm-AGA cases presented lower  
167 birthweight and gestational age at delivery as compared with controls. Preterm-  
168 IUGR cases also showed a lower birthweight centile as compared with controls  
169 and preterm-AGA. The proportion of males was similar among groups, while  
170 the preterm-IUGR subgroup presented a lower prevalence of Caucasian eth-  
171 nicity. Heart rate was significantly higher for preterm-AGA and preterm-  
172 IUGR, since these subjects had lower stroke volume which was compensated  
173 by increasing the heart rate [4]. Both preterm-IUGR and preterm-AGA cases  
174 showed significantly lower age, height and weight (with preserved body mass  
175 index) at evaluation as compared with controls. All subjects were asymp-  
176 tomatic, and none of them received medication for any cardiac condition.

177 Tables 2 and 3 also highlight obtained p-values lower than 0.05 by apply-  
178 ing the Wilcoxon-Mann-Witney or the Student's t-test for the comparisons  
179 between the three different groups of subjects. We studied differences be-  
180 tween the control group and the preterm-AGA and preterm-IUGR groups.

181 Figure 1 depicts the graphical comparison between the three groups (one  
182 example of each group under study, in pairs). In this Figure, a comparison  
183 of the preterm-AGA and the preterm-IUGR groups with respect to the con-  
184 trol group was represented, whereas Figure 2 shows the comparison between  
185 the dominant vectors of the depolarization and repolarization loops for the  
186 analysis of one subject from the preterm-IUGR and another subject from the  
187 preterm-AGA. Furthermore, for the sake of clarity in graphical representa-  
188 tion, the projections of the loops on the plane with clearer differences were  
189 represented (instead of representing them in the three-dimensional space).

190 In general, preterm-AGA and preterm-IUGR subjects presented larger  
191 values for angle measurement  $\Phi_{R-XY}$  than controls, being also statistically  
192 significant when adjusting the data by age, height, gender and heart rate. In  
193 the particular case of preterm-IUGR, the angle difference  $\Phi_{R-XY} - \Phi_{T-XY}$  was  
194 also significant, being even larger than for the preterm-AGA group.

195 Roundness measurements revealed significant differences between preterm-  
196 AGA and control groups, presenting the latter ones lower roundness measure-  
197 ments when analyzing the depolarization loops in the planes XZ and YZ.

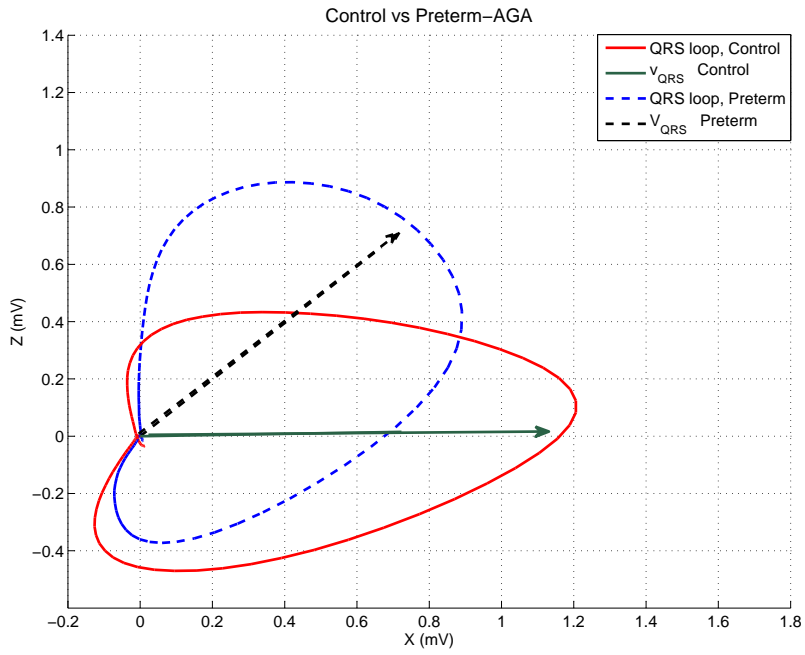


Table 2: Vectocardiographic angle features of the study population.  $\theta_{RT}$  and  $\theta_{RT-XY}$  denote the so-called spatial QRS-T and frontal QRS-T angles, respectively. Absolute angles between vectors  $\mathbf{v}_{QRS}$  or  $\mathbf{v}_T$  and the three orthogonal planes of the space are denoted by  $\Phi_{R-\mathcal{P}}$  and  $\Phi_{T-\mathcal{P}}$ , where  $\mathcal{P} \in \{XZ, XY, YZ\}$  refers to each orthogonal plane. AGA (appropriate growth for gestational age), IUGR (intrauterine growth restriction). Subscripts <sub>R</sub> and <sub>T</sub> refer to the depolarization and repolarization loops, respectively. Data are mean  $\pm$  standard deviation or median (interquartile range). \* p-value  $< 0.05$  by Wilcoxon-Mann-Whitney or t-test as compared to controls. † p-value  $< 0.05$  adjusted by age, height, gender and heart rate, controls as reference category.

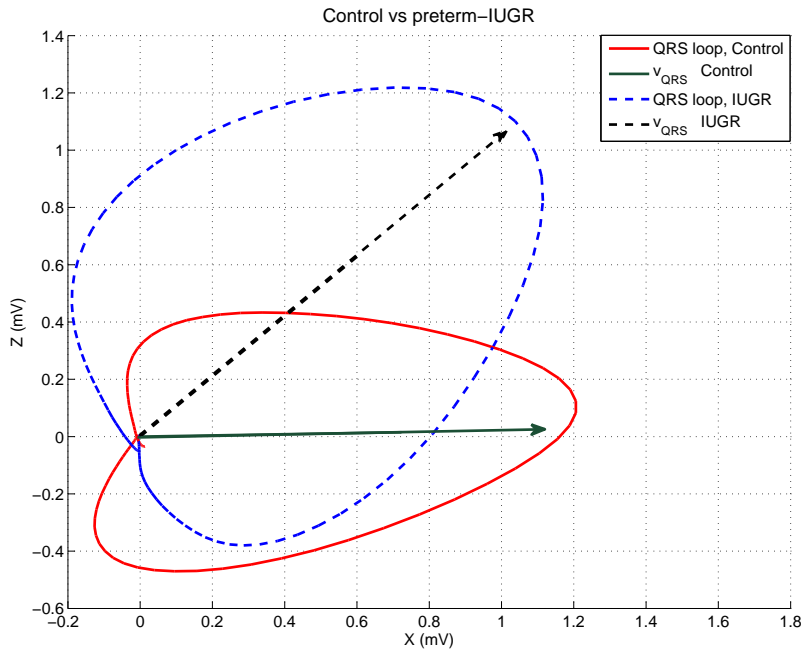
	Controls n=60	Preterm AGA n=32	Preterm IUGR n=33
$\theta_{RT}$	15.28(8.390-24.10)	10.63(5.539-20.76)	16.58(6.890-22.73)
$\theta_{RT-XY}$	6.57(2.72-11.31)	4.68(2.20-12.89)†	6.52(3.88-15.82)
$\Phi_{R-XZ}$	37.60(29.91-40.82)	32.64(28.90-36.78)*	34.24(28.58-39.91)
$\Phi_{R-XY}$	20.03(10.11-28.64)	27.77(16.59-33.23)*	25.48(19.79-33.56)*†
$\Phi_{R-YZ}$	45.95 $\pm$ 9.31	45.85 $\pm$ 9.02	43.06 $\pm$ 8.92
$\Phi_{T-XZ}$	36.42(32.05-38.83)	31.89(29.07-38.14)	32.01(28.58-38.63)†
$\Phi_{T-XY}$	29.39(13.71-27.26)	23.59(17.63-26.80)	21.52(12.77-25.89)
$\Phi_{T-YZ}$	47.24(38.75-52.86)	46.57(41.63-53.30)	47.86(41.16-56.55)
$\Phi_{R-XZ} - \Phi_{T-XZ}$	1.70(-3.97-5.43)	-0.52(-6.01-4.69)	0.88(2.33-6.10)
$\Phi_{R-XY} - \Phi_{T-XY}$	-0.491 $\pm$ 14.15	3.001 $\pm$ 12.99	5.279 $\pm$ 12.15*†
$\Phi_{R-YZ} - \Phi_{T-YZ}$	-0.662 $\pm$ 12.50	-1.752 $\pm$ 10.67	-4.854 $\pm$ 12.74

Table 3: Vectocardiographic loops morphology of the study population.  $\rho_L$ ,  $\sigma_L$ , and  $\delta_L$  denote measures of planarity of loop  $L$ .  $C_L$  and  $\varrho_{L-\mathcal{P}}$  refer to roundness measurements.  $\mathcal{P} \in \{XZ, XY, YZ\}$  refers to the projection of the loop in each orthogonal plane  $\mathcal{P}$ , whereas the subscript  $L$  will be replaced by  $R$  or  $T$ , depending on whether it refers to the depolarization or repolarization loops. AGA (appropriate growth for gestational age), IUGR (intrauterine growth restriction). Subscripts  $R$  and  $T$  refer to the depolarization and repolarization loops, respectively. Data are median (interquartile range). \* p-value < 0.05 by Wilcoxon-Mann-Whitney or t-test as compared to controls. † p-value < 0.05 adjusted by age, height, gender and heart rate, controls as reference category.

	Controls n=60	Preterm AGA n=32	Preterm IUGR n=33
$\rho_R$	0.044(0.029-0.067)	0.036(0.022-0.058)	0.037(0.022-0.051)
$\rho_T$	0.031(0.021-0.054)	0.037(0.028-0.061)	0.034(0.021-0.048)
$\sigma_R$	0.967(0.953-0.979)	0.975(0.956-0.982)	0.971(0.961-0.982)
$\sigma_T$	0.973(0.955-0.981)	0.968(0.946-0.976)	0.972(0.955-0.981)
$\delta_R$	0.033(0.022-0.045)	0.024(0.017-0.046)	0.028(0.017-0.040)
$\delta_T$	0.028(0.019-0.044)	0.031(0.024-0.054)	0.027(0.019-0.044)
$C_R$	0.316(0.222-0.423)	0.264(0.171-0.331)	0.291(0.202-0.420)
$C_T$	0.135(0.103-0.189)	0.149(0.123-0.192)	0.134(0.096-0.189)
$\varrho_{R-XZ}$	0.384(0.257-0.489)	0.291(0.194-0.396)*†	0.335(0.247-0.516)
$\varrho_{R-XY}$	0.113(0.066-0.162)	0.086(0.049-0.155)†	0.117(0.074-0.187)
$\varrho_{R-YZ}$	0.330(0.211-0.452)	0.227(0.164-0.331)*†	0.276(0.212-0.346)
$\varrho_{T-XZ}$	0.157(0.124-0.198)	0.166(0.113-0.204)	0.141(0.102-0.235)
$\varrho_{T-XY}$	0.054(0.037-0.067)	0.072(0.042-0.110)*	0.054(0.032-0.095)
$\varrho_{T-YZ}$	0.201(0.151-0.272)	0.155(0.111-0.235)*	0.171(0.134-0.281)

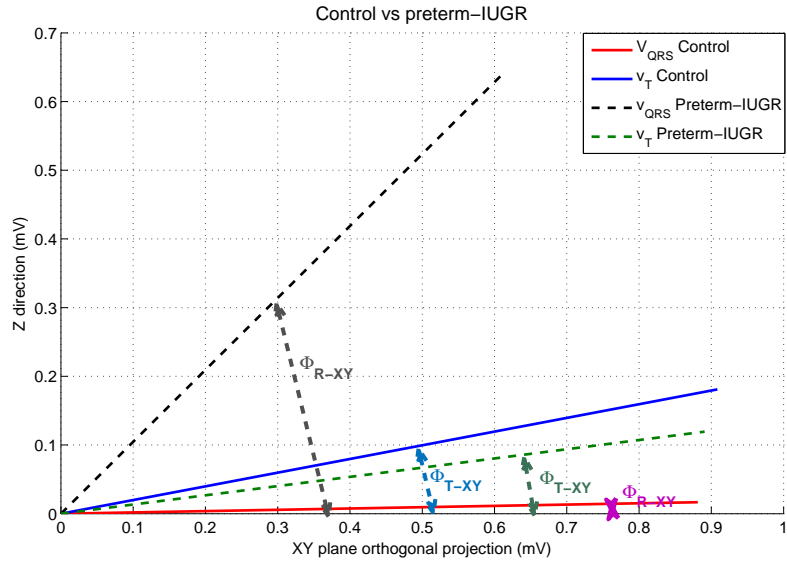


(a)

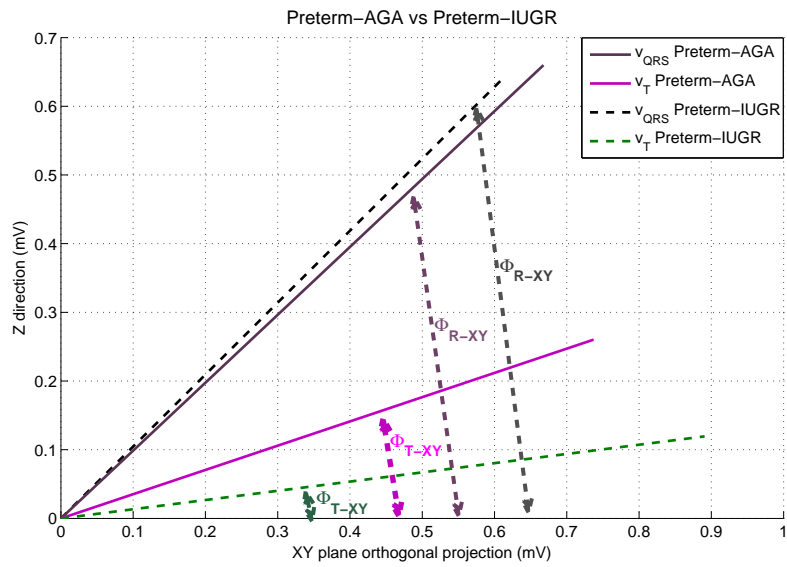


(b)

Figure 1: (Average of all depolarization loops for a patient and their corresponding dominant vectors ( $v_{QRS}$ ) projected onto the XZ-plane. (a) Subject of the control group vs. a subject from the preterm group. (b) Same subject of the control group vs. a subject from the IUGR group.



(a)



(b)

Figure 2: (a) Projection of dominant vectors of the average depolarization and repolarization loops on XY plane (and the respective absolute angles) for a subject of the control group and a subject from the preterm-IUGR group. (b) Projection of dominant vectors of the average depolarization and repolarization loops on XY plane (and the respective absolute angles) for a subject of the preterm-AGA group and a subject from the preterm-IUGR group.

#### 198 4. Discussion of results

199 This paper reports for the first time significant changes in depolarization-  
200 repolarization angles and VCG loop morphology in IUGR and preterm indi-  
201 viduals, respectively. These findings support the notion of electrophysiolog-  
202 ical remodeling in both IUGR and prematurity, and could partially explain  
203 the higher cardiovascular morbidity previously reported in these populations.

204 This study has included a group of preadolescents who suffered severe  
205 IUGR in uterus that lead to medically induced preterm delivery. This  
206 preterm-IUGR group was characterized by a wider QRS-T angle ( $\theta_{RT}$ ) in  
207 relation to body plane, with no significant differences in roundness and pla-  
208 narity. While a non-significant trend to higher values of spatial ( $\theta_{RT}$ ) and  
209 frontal ( $\theta_{RT-XY}$ ) QRS-T angles was observed, significantly higher angles in  
210 the frontal plane (XY) could be demonstrated with respect to controls (Ta-  
211 ble 2). Despite being wider than in controls, their angle values were within  
212 the normal limits [16]. Usually, a wider QRS-T angle is derived by changes  
213 in T-wave axis, which has been reported to be a risk factor for cardiac events  
214 [17, 18]. However, in our study wider QRS-T angle in the XY plane was  
215 mainly due to changes in QRS axis (although there were also changes in the  
216 T-wave -significant for XZ plane, but not significant for XY frontal plane-),  
217 suggesting more significant depolarization rather than repolarization abnor-  
218 malities. Wider spatial QRS-T angle has been proven to be a powerful pre-  
219 dictor of cardiovascular mortality in general population [19, 20, 17] and of  
220 sudden arrhythmic death in survivors of acute myocardial infarction [21]. It  
221 has been also described in smokers, myocardial hypertrophy, diabetes, and  
222 high blood pressure [22]. Regarding electrical remodeling, our results are  
223 in line with previous reports suggesting lower acceleration and deceleration  
224 capacity in very preterm-IUGR fetuses [23, 24] and higher QT and JT dis-  
225 persion [25] in IUGR fetuses and newborns. The electrical findings are also  
226 consistent with previous echocardiographic studies in IUGR demonstrating  
227 significant changes in the cardiac structure and function of these individu-  
228 als in uterus but also in postnatal age [4]. Additional results included in  
229 the Supplementary data section showed significant correlations between left  
230 ventricular mass and depolarization-repolarization angles for preterm IUGR  
231 subjects. This can be related to the fact that severe IUGR cases typically  
232 show more spherical hearts with impaired relaxation and longitudinal mo-  
233 tion together with a decrease in stroke volume which is compensated by an  
234 increase in heart rate in order to maintain cardiac output [26, 27]. These car-

235 diac changes are thought to be a response to the placental insufficiency which  
236 is usually associated with fetal hypoxia, undernutrition and pressure/volume  
237 overload [28]. We speculate that the link between our electrical results and  
238 the previously reported structural changes could be explained by the loca-  
239 tion of the Purkinje system inside the endocardium. The pressure overload  
240 and hypoxia -usually associated with IUGR- is known to mostly affect the  
241 endocardial fibers leading to a decreased longitudinal motion and could also  
242 predominantly affect the endocardial Purkinje system and change the vec-  
243 tor gradient of depolarization and repolarization. It is also shown that this  
244 pressure overload mainly affects the basal part of the ventricle which might  
245 contribute to the heterogeneity in the distribution of the endocardial pres-  
246 sure, explaining wider angles as the T- vectors might not be in the same  
247 direction as the QRS ones [22].

248 The present study also included a group of normally grown cases who were  
249 spontaneously born preterm. This preterm-AGA group was characterized  
250 by a reduced frontal QRS-T angle and roundness with preserved planarity.  
251 Similar results in the QRS-T angle were reported in healthy middle-aged  
252 subjects with a genetic polymorphism of the KCNH2(HERG), a gene with  
253 functional electrical properties. Smaller values in the QRS-T angle could  
254 reflect a desynchronization between repolarization and depolarization that  
255 could be associated with increased cardiovascular mortality(7). However, it  
256 has been suggested that frontal QRS-T angle turned out to be a bad sub-  
257 stitute for spatial QRS-T angle [29, 16]. Therefore, this result needs to be  
258 carefully interpreted. The less round projections of the VCG observed in  
259 preterm-AGA could also be associated with higher cardiovascular risk. The  
260 electrical changes observed in preterm-AGA are consistent with a previous  
261 study suggesting longer QT interval, higher QT dispersion and shorter PR  
262 interval in young adults born preterm [30]. We speculate that these elec-  
263 trical findings could be related to the cardiac structural changes previously  
264 described in individuals born prematurely. Adults born preterm show smaller  
265 and shorter hearts with increased left ventricular mass and impaired longi-  
266 tudinal motion [5]. This cardiovascular remodelling seems to be an adaptive  
267 response to the relative pressure overload that the immature cardiovascular  
268 system suffers during the neonatal period after a premature delivery [31],  
269 though inflammation, infection or other pathophysiological mechanisms po-  
270 tentially associated with spontaneous preterm delivery could also explain the  
271 cardiac response observed in these individuals. The different pathophysiological  
272 mechanisms observed in IUGR and spontaneous preterm delivery could

273 determine the distinct pattern of electrical remodelling observed in preterm-  
274 AGA and preterm-IUGR in our study.

275 Strengths and limitations of the present deserve further comments. Our  
276 work is pioneer in establishing the electrophysiological consequences of pre-  
277 maturity and IUGR by VCG. The study populations were well-defined and  
278 phenotyped from fetal life, permitting a clear distinction between sponta-  
279 neous and medically-induced preterm delivery, which are usually mixed up  
280 in the literature. IUGR cases were also carefully selected including umbilical  
281 artery Doppler in the definition, which is an excellent surrogate of severe pla-  
282 cental insufficiency. In addition, the VCG measurements used here are less  
283 susceptible to noise and definition problems as compared to the conventional  
284 ECG analysis including QT dispersion. Despite these strengths, we acknowl-  
285 edge potential limitations. Firstly, the relatively limited sample size might  
286 have prevented demonstrating significant differences in some angular mea-  
287 surements and planarity. Secondly, we acknowledge that differences in age  
288 at evaluation between cases and controls might have influenced the results.  
289 Thirdly, potential confounding variables such as smoking, mental stress or  
290 family history were not included in the analysis. Fourthly, the study was  
291 performed in preadolescent age which warrants future studies to evaluate the  
292 persistence of these findings in older ages and its potential association with  
293 cardiovascular adverse events. Additionally, while VCG analysis based on  
294 the ECG is a non-invasive low-cost approach that has been shown to be use-  
295 ful for risk assessment, it is still not widely used in clinical practice. Finally,  
296 future studies are warranted to better determine the mechanisms underlying  
297 the electrical changes observed in these populations.

## 298 5. Conclusions

299 Our study demonstrates for the first time particular patterns of electrical  
300 remodeling associated with both prematurity and IUGR. Significantly wider  
301 angles between the depolarization dominant vector and the frontal XY body  
302 plane were observed for preterm-IUGR subjects, resulting in significantly  
303 wider angles between depolarization and repolarization vectors. Significantly  
304 lower angles were observed for the repolarization vector and the XZ plane in  
305 preterm-IUGR subjects with respect to controls. The classical frontal QRS-  
306 T angle was significantly narrower in preadolescents who were spontaneously  
307 born preterm with respect to controls. Moreover, significantly less roundness

308 measurements for the depolarization loop projections in all three orthogonal  
309 planes were also observed.

310 These findings might be related to the previously described cardiac struc-  
311 tural changes and increased cardiovascular risk in these populations. Future  
312 studies are warranted to confirm these results and further describe the elec-  
313 trical characteristics of these individuals and its potential long-term conse-  
314 quences.

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