Intrauterine Growth Retardation in Pregnant Women with Long QT Syndrome Treated with Beta-Receptor Blockers

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Abstract
Pregnant women with inherited long QT syndrome (iLQTS) are at an increased risk for preterm delivery and intrauterine growth retardation (IUGR) due to their underlying disease. Additionally, they are at a risk of arrhythmogenic events, particularly during the postpartum period because of physiological changes and increased emotional/physical stress. \(\beta\)-receptor blockers can effectively prevent life-threatening Torsades de Pointes ventricular tachycardia and they are the treatment of choice in iLQTS. Use of \(\beta\)-receptor blockers in pregnancy is recommended, although IUGR is commonly reported for prenatally exposed infants. IUGR, particularly in preterm infants, can result in adverse neonatal outcomes. This review was performed to support clinicians in their selection of \(\beta\)-receptor blocker treatment for their pregnant iLQTS women by (i) summarizing the available literature addressing the impact of different \(\beta\)-receptor blockers on IUGR and (ii) reporting additional aspects which might influence the \(\beta\)-receptor blocker selection. In general, experts recommend to use nonselective \(\beta\)-receptor blockers, such as nadolol and propranolol, for iLQTS management as these drugs seem to be superior in effectiveness. However, \(\beta\)-1-selective receptor blockers, such as bisoprolol or metoprolol, seem to affect less likely uterine contraction, peripheral vasodilation, and are associated with lower IUGR rates and fetal hypoglycemia. They are therefore recommended, except atenolol, as first-line therapy for pregnant women. Additionally, maternal factors such as iLQTS genotype, other underlying comorbidities (e.g., diabetes mellitus type 1, asthma bronchiale), and uteroplacental dysfunction or fetal factors have to be taken into account. Therefore, each woman with iLQTS who wants to become pregnant should be well-advised for a personalized \(\beta\)-receptor blocker therapy according to the individual risk-benefit evaluation by a multidisciplinary team of cardiologists, gynecologists, pediatric cardiologists, neonatologists, and clinical pharmacologists. During pregnancy, a close monitoring of IUGR and, after birth, monitoring of bradycardia, hypoglycemia, and respiratory depression in the neonate is mandatory. This review summarizes available data on \(\beta\)-receptor blocker-related risk for IUGR in prenatally.
exposed infants and illustrates which factors might influence β-receptor blocker selection with the aim to support clinicians in their pharmacological management of their pregnant iLQTS patients.

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Introduction

Nowadays, an increasing number of women with heart disease, including congenital arrhythmias such as the inherited long QT syndrome (iLQTS), become pregnant [1]. The iLQTS is caused by gene variants coding for ion channels, ion subunits, or regulatory proteins, representing a genetically diverse group of disorders of ventricular repolarization associated with an increased risk of life-threatening Torsades de Pointes ventricular tachycardias. The 3 major forms of iLQTS are LQTS type 1–3. The diagnosis of iLQTS relies on prolonged QT interval in the electrocardiogram (ECG) or prolongation of the heart beat corrected QT interval (QTc), respectively, clinical and family history, and/or genetic testing [2]. In the absence of secondary causes for QT prolongation, a QTc of 480 ms or longer in repeated ECGs, or in presence with unexplained syncope, a QTc of 460 ms or longer indicates iLQTS, or a Schwartz score 1.5–3 shows an intermediate probability of iLQTS [3, 4]. It is known that pregnancy in women with heart disease is a medical challenge as the underlying disease is a major risk factor for maternal morbidity and mortality [5], but can also cause fetal complications. Pregnant iLQTS women seem to be at an increased risk of placental or myometrial dysfunction, which might result in stillbirth, miscarriage, preterm delivery, and intrauterine growth retardation (IUGR). Stillbirths are 8 times more frequently reported in iLQTS compared to the normal pregnant population (4 vs. approximately 0.5%) and miscarriages are twice more frequent (16 vs. 8%) [6]. One reason for intrauterine fetal death might be the risk of iLQTS and bradycardia in the fetus [7], as in 2/3 of newborns from iLQTS mothers/fathers, an iLQTS genotype can be detected [6]. Additionally, the birth weight for newborns of iLQTS mothers seems to be lower if adjusted for gestational age compared to those of iLQTS fathers [6]. It is important to highlight that pregnant iLQTS women themselves are at an increased risk for cardiac events (CE) postpartum, particularly in case of a LQTS type 2 (LQTS2) [8–11]. The mothers have an alternated sleep pattern, lack of sleep, and an increased physical/emotional stress, which might lead to an increased sympathetic activity, with an increased gene-specific CE risk postpartum [9, 11, 12]. The new auditory stimuli of the crying infant might be an additional trigger for CE in LQTS2 [11]. As estrogen seems to have an anti-arrhythmogenic effect, the physiological estrogen decrease during lactation might be an additional risk factor for CE [13, 14]. During the 9-month postpartum period, a 2.7-fold increased CE risk and a 4.1-fold increased risk for life-threatening events compared to the preconception period for iLQTS women are reported [10].

The treatment of choice in patients with iLQTS is oral β-receptor blockers, as they are effective in reducing the CE risk [15, 16]. β-receptor blocker treatment seems to be safe and not teratogenic but is associated with IUGR, apnea, hypoglycemia, hyperbilirubinemia, and bradycardia in the prenatally exposed fetus and newborn infant [17–20]. However, bradycardia can be also related with clinical presentation of iLQTS in the newborn [21].

Risk factors for IUGR comprise a wide range and may be caused by fetal (e.g., congenital infection and genetic syndromes), placental (e.g., marginal or velamentous cord insertion, placenta praevia, abruptio placenta, and infarction), and maternal (e.g., smoking, advanced maternal age, very low or very high body mass index, underlying chronic diseases, anemia, and uterine malformation) factors. In pregnant iLQTS women, risk factors for IUGR might include the underlying disease and the β-receptor blocker treatment. IUGR, particularly in preterm infant can result in adverse neonatal outcome. To our knowledge, no overview about the impact of different β-receptor blocker on IUGR is available so far. The aim of this review is to summarize available data on β-receptor blocker-related risk for IUGR and to give an overview in which additional factors might influence β-blocker selection in pregnant iLQTS patients to support clinicians in their pharmacological management of their pregnant iLQTS patients.

Material and Methods

Based on a pharmacological clinical consultation, September 19, 2020, a literature review was performed in September 2020 to assess data addressing β-receptor blocker-related risk for IUGR in infants prenatally exposed to β-receptor blockers. Search strings in the PubMed database included β-blocker OR atenolol OR bisoprolol OR metoprolol OR carvedilol OR propranolol OR nadolol AND fetal/intrauterine growth retardation OR small for gestational age. The additional literature addressing efficacy data for β-receptor blockers in iLQTS that might influence the treatment selection in pregnancy were reviewed. Available full text publications (original articles, case reports, and reviews) were screened.
Additionally, relevant publications were identified manually from the bibliographies of references retrieved from the PubMed search. Data were extracted by a scientist in an electronic datasheet for IUGR. Additional information for the US Food and Drug Administration (FDA) pregnancy letter categories used before June 30, 2015 (Table 1) was retrieved and combined with data from the literature search.

### Results

We could identify 15 papers providing information on the topic IUGR in infants and prenatally exposure to β-receptor blocker. Of these, 8 cohort studies and 1 case report showed detailed data for IUGR according to prenatally β-receptor blocker exposure (Table 2). Main indication for the β-receptor blocker treatment during pregnancy was maternal hypertension. Five reviewed β-receptor blockers (metoprolol, bisoprolol, carvedilol, propranolol, and nadolol) were classified by the former FDA category (Table 1) as C and atenolol was classified as D. However, the most data available was for atenolol (n = 7). Only 1 case report could be identified for nadolol [22]. An important aspect influencing the β-receptor blocker selection for treatment is the efficacy of the β-receptor blocker, which seems to depend on the iLQTS genotype. Some studies comparing the efficacy of different β-blockers in patients with iLQTS are available but randomized controlled trials are lacking.

### Data on β-Receptor Blockers and IUGR in General

IUGR is reported as a known side effect in the fetus/infant which has been prenatally exposed to β-receptor blockers, and might be explained by an increase of blood flow resistance in both myometrium and placenta with a decrease in the uteroplacental blood flow [23–26]. Cuneo et al. [6] studied retrospectively 148 pregnancies from 80 LQTS mothers and 23 LQTS fathers, resulting in 118 live births. Of these pregnant females, 71% (n = 57) received β-receptor blockers. The birth weights of infants from mothers who used β-receptor blockers (2.8 ± 0.7 kg) were significantly different (p < 0.001) from the birth weights of infants of LQTS fathers (3.5 ± 0.5 kg) and if the birth weights were adjusted for gestational age, infants of LQTS mothers who used β-receptor blockers had lower birth weights as compared to infants from untreated mothers [6]. It seems that IUGR risk is more pronounced if β-receptor blocker therapy is started early in pregnancy and continued throughout the entire pregnancy [27]. However, IUGR is also observed if β-receptor blockers were only used after the first trimester, but perhaps less severe [28]. As most women with iLQTS will be treated with β-receptor blockers already before conception the fetus will be exposed during the entire pregnancy. Additionally, the risk for neonatal adverse events seems to increase with high β-receptor blocker dosages [29].

### Data on Different β-Receptor Blockers and IUGR

Tanaka et al. [23] analyzed maternal and neonatal outcomes in 158 pregnancies of women with cardiovascular
| β-receptor blocker | Pharmacokinetic properties | Data | Former FDA category | Placenta permeable | Additional notes |
|-------------------|----------------------------|------|---------------------|------------------|-----------------|
| **β-1-receptor selective blockers** |
| Atenolol | Protein binding: 6–16% [53]  
Metabolism: limited hepatic [53]  
Elimination half-life: in utero exposed neonates: 16 h (mean) [54], adults: 6–7 h (longer in case of renal impairment) [53] | 33% newborns had IUGR (n = 2/6); atenolol dose: 25–50 mg/day [23]  
18% of <2,500 g newborns (n = 115/638) [30]  
3.8% of <1,500 g newborns (n = 24/638) [30]  
48.7% of newborns (n = 38/78) [27]  
Newborns (n = 15) had a lower birthweight compared to nonexposed newborns (2,620 vs. 3,530 g; 440–1,380 g; atenolol dose: 50–200 mg/day, [32])  
Exposed newborns were significantly lighter compared to infants exposed to other β-receptor blockers [33]  
75% newborns had IUGR (n = 3/4) [29]  
70% newborns exposed <15 weeks gestation showed growth ≤10th centile (n = 26/50) [55]  
40% newborns exposed <15 weeks gestation showed growth <3rd centile (n = 15/50) [55]  
51% newborns exposed 15–30 weeks gestation showed growth ≤10th centile (n = 15/29) [55]  
17% newborns exposed 15–30 weeks gestation showed growth ≤3rd centile (n = 5/29) [55] | D | Yes [17, 53] | Not recommended in pregnancy  
The highest risk for IUGR of all analyzed β-receptor blockers |
| Bisoprolol | Protein binding: ~30% [56]  
Metabolism: extensively hepatic [56]  
Elimination half-life: 9–12 h (normal renal function) [56] | No IUGR was observed (n = 0/5), bisoprolol dose: 5–10 mg/day [23] | C | Yes [17] | Effective QTc shortening in LQTS1/LQTS2 [40]  
Good long-term tolerance [40]  
Seems similar effective to nadolol/propranolol in preventing major CE [39] |
| Metoprolol | Protein binding: ~10–12% [57]  
Metabolism: extensively hepatic via CYP2D6 [57]  
Elimination half-life: neonates: 5–10 h, adults: 3–4 h (7–9 h in poor CYP2D6 metabolizers or hepatic impairment) [57] | 17% of newborns had IUGR (n = 2/12), metoprolol dose: 20–120 mg/day [23]  
13.3% of <2,500 g newborns (n = 43/324) [30]  
3.1% of <1,500 g newborns (n = 10/324) [30] | C | Yes [17, 58] | PK changes during pregnancy with need of aggressive dosage adaptation [59]  
Might be effective in patient with mutation W305L [60]  
Risk of breakthrough CE in LQTS1/LQTS2 [35] |
| Carvedilol | α and β-receptor blocking agent [61]  
Protein binding: >98%, primarily to albumin [61]  
Metabolism: extensively hepatic [61]  
Elimination half-life: adults: ~7–10 h [61] | 7% of IUGR was observed (N = 1/13), carvedilol dose: 2.5–20 mg/day [23] | C | Yes (data from rats) [17] | Class III antiarrhythmic effect might be beneficial [62] |
| Propranolol | β-1-receptor and β-2-receptor blockade [63]  
Protein binding: newborns: 68%; adults: ~90% [63]  
Metabolism: extensively hepatic [63]  
Elimination half-life: adults: Immediate release formulation: 3–6 h; extended-release formulations: 8–10 h [63] | 36% of IUGR was observed (n = 8/22), propranolol dose: 15–60 mg/day [23]  
7.6% of <2,500 g newborns (n = 37/489) [30]  
1.6% of <1,500 g newborns (n = 8/489) [30]  
29% newborns had IUGR (n = 2/7) [29] | C | Yes [17] | Risk for breakthrough CE in high-risk patients [41]  
Efficacy seems comparable with nadolol (first line) [35, 42, 43] |
diseases, such as (i) congenital heart disease and pulmonary hypertension; (ii) aortic disease; (iii) valvular heart disease; (iv) coronary artery disease and acute coronary syndrome; (v) cardiomyopathy and heart failure; and (vi) arrhythmia, who were exposed to carvedilol \((n=13,\) dose: 2.5–20 mg/day), propranolol \((n=22,\) dose: 15–60 mg/day), metoprolol \((n=12,\) dose: 20–120 mg/day), atenolol \((n=6,\) dose: 25–50 mg/day), and bisoprolol \((n=5,\) dose: 5–10 mg/day). IUGR occurred in 1 patient (7%) with carvedilol [23]. Duan et al. [30] analyzed data from 4,847 pregnant women treated with β-receptor blockers and compared them with data of 374,391 pregnant women without β-receptor blocker treatment. Diagnoses of hypertension, hyperlipidemia, diabetes, heart failure, and a history of arrhythmia were more common among patients exposed to β-receptor blockers. The 4 most commonly prescribed β-receptor blockers were labetalol \((n=3,357),\) atenolol \((n=638),\) propranolol \((n=489),\) and metoprolol \((n=324).\) The risk of fetal growth in infants exposed to metoprolol or bisoprolol was not significantly different from the nonexposed group (metoprolol: adjusted OR 1.3, 95% CI 0.9–1.9; bisoprolol: adjusted OR 1.3, 95% CI 0.9–1.9), whereas atenolol was associated with an increased risk of IUGR (adjusted OR 2.4, 95% CI 1.7–3.3) [30]. The mean birth weight and percentage low birth weight \(<2,500\) g were 3,058 ± 748 g and 18.0% for atenolol, 3,163 ± 702 g and 13.3% for metoprolol, 3,286 ± 651 g and 7.6% for propranolol, and 3,353 ± 554 g and 5.2% for nonexposed controls [30]. Kayser et al. [31] analyzed 294 neonates of hypertensive mothers treated with metoprolol or bisoprolol during the second and/or third trimester and compared these infants with 225 methyldopa-exposed infants and 588 infants of nonhypertensive mothers. The rate of IUGR was significantly higher in infants who had long-term β-receptor blocker exposure (24.1%) as compared to those exposed to methyldopa (10.2%, OR 2.5, 95% CI 1.2–5.2) and the nonhypertensive ones (9.9%, OR 4.3, 95% CI 2.6–7.1) [31]. Lydakis et al. [27] performed a study in 312 pregnancies of 223 women. Of these pregnancies, 35.9% were complicated by gestational hypertension, 6.1% by pre-eclampsia in previously normotensive women, in 57.4% of pregnancies there was a positive history of chronic hypertension, and finally 10.9% of pregnancies had pre-eclampsia superimposed on chronic hypertension. In 91 of these pregnancies (29.2%) no treatment was administered, whereas 78 women were treated with atenolol monotherapy (25%), 53 with different drugs as antihypertensive monotherapy (17%), and 90 with multiple drug combinations (28.8%) [27]. Atenolol was found to be associated with a lower birth weight, a trend towards a higher prevalence of prematurity (<37 weeks) and IUGR as compared to other antihypertensive monotherapies, or to no treatment [27]. Butters et al. [32] treated 15 pregnant women with essential hypertension with atenolol, whereas the other 14 women received placebo. All infants in the atenolol group had a significantly lower birth weight as compared to those in the placebo group (2,620 vs. 3,530 g; 910 [440–1,380] g). They concluded that atenolol given from the end of the first trimester in pregnant women with mild hypertension is associated with IUGR [32]. These observations are in line with Lip et al.
[33] who showed, by reviewing records of 398 pregnant women, that infants born to women taking atenolol (n = 76) had a significantly lower birth weight (p < 0.001) than infants born to women taking other β-blockers, other antihypertensive drugs, or no therapy. Details are shown in Table 2.

### β-Receptor Blockers and Data on Efficacy and iLQTS Type

The efficacy of β-receptor blockers seems to be influenced by the genotype [34]. Chockalingam et al. [35] studied ECGs and clinical parameters in 382 LQTS1/LQTS2 patients treated with propranolol (n = 134; median dose 1.8 mg/kg/day), metoprolol (n = 147; median dose 0.9 mg/kg/day), and nadolol (n = 101, median dose 0.9 mg/kg/day). The baseline heart rate (76 ± 16 bpm) was comparable among the different β-receptor blocker groups, whereas baseline QTc was different (472 ± 46 ms). The on-therapy heart rate and QTc were different compared to the baseline heart rate and QTc within each β-blocker group, respectively (p < 0.001 for all paired comparisons). However, while the change in heart rate with β-receptor blocker initiation was comparable among the groups (p = 0.9), the change in QTc with propranolol was greater than that with metoprolol (p = 0.003) and nadolol (p = 0.004). Additionally, the cohort was subdivided in a normal baseline QTc (≤450 ms), a borderline (451–480 ms), and a prolonged baseline QTc (>480 ms). The QTc shortening with propranolol, metoprolol, and nadolol in the normal QTc subset and in the borderline QTc subset was comparable, but was significantly different in the prolonged QTc subset (49 ± 43, 30 ± 40, 27 ± 29 ms, p = 0.01). They concluded that propranolol had a significantly better QTc shortening effect compared to metoprolol and nadolol, particularly in patients with prolonged QTc [35]. As Kaplan-Meier analysis showed a significantly lower event-free survival for symptomatic patients on metoprolol compared to nadolol/propranolol, they stated that metoprolol should not be used in symptomatic LQTS1/LQTS2 patients, whereas propranolol and nadolol seem equally effective [35]. However, the QTc shortening effect of propranolol seems to be dose dependent, as particularly high concentrations result in a significant shortening of the QTc interval [36, 37]. Chattrath at al. [38] studied 28 patients with iLQTS treated with atenolol (n = 12), propranolol (n = 10), metoprolol (n = 4), and nadolol (n = 2) for breakthrough CE. During a median follow-up of 46 months, 7 patients (25%) developed 15 breakthrough CE. Of these, 10 occurred in atenolol and 3 in propranolol compliant patients; 2 were related with noncompliance. They concluded that atenolol treatment might be an important factor for breakthrough CE [38]. Fazio et al. [39] evaluated 34 LQTS patients with an average follow-up time of 93 months for (i) no treatment (31 months), (ii) treatment with nadolol or propranolol (31 months), and (iii) treatment with bisoprolol (31 months). They observed in the follow-up period 2 major (arrhythmic syncope resuscitated by electrical defibrillation) and 12 minor (non-arrhythmic syncope ± loss of consciousness) CEs. Both major events occurred without β-receptor blocker treatment, whereas the minor events were distributed as follows: 3 without treatment, 2 with nadolol, 5 with propranolol, and 2 with bisoprolol [39]. Steinberg et al. [40] compared retrospectively ECG parameters and CE of 114 LQTS1/LQTS2-patients treated with bisoprolol (n = 59), nadolol (n = 16), or atenolol (n = 39). The baseline heart rate and QTc interval were similar between the groups. QTc shortening was observed in the individuals on bisoprolol (ΔQTc −5 ± 31 ms; p = 0.049) and nadolol (ΔQTc −13 ± 16 ms; p = 0.02), but not on atenolol (ΔQTc +9 ± 24 ms; p = 0.16) [40]. A median follow-up was 3 years for bisoprolol and nadolol and 6 years for atenolol [40]. During follow-up, one CE was observed in the bisoprolol group (1.7%) and 2 for the atenolol group (5.1%; p = 0.45), none occurred in nadolol treated patients. Based on that, they pointed out, that bisoprolol is effective in QTc shortening for LQTS1 and LQTS2 and is well tolerated during long-term administration [40]. Abu-Zeitone compared in 1,530 patients from the Rochester, New York-based LQTS registry, the efficacy of nadolol (n = 259; dose: <18 and ≥18 years: 1.0 ± 0.8 mg/kg/day), propranolol (n = 679; dose: <18 years: 2.3 ± 1.5 mg/kg/day, ≥18 years: 2.1 ± 2.3 mg/kg/day), atenolol (n = 441; dose: <18 years: 1.0 ± 0.7 mg/kg/day, ≥18 years: 0.7 ± 0.3 mg/kg/day), and metoprolol (n = 151; dose: <18 years: 1.4 ± 1.0 mg/kg/day, ≥18 years: 1.2 ± 0.9 mg/kg/day) in LQTS and in genotype positive patients with LQTS1/LQTS2 [41]. In LQTS1 (n = 379), the risk reduction for any β-receptor blocker was 57% (p < 0.01) with insufficient evidence of different efficacy for any specific drug. In LQTS2 (n = 406) there was a significant variability in efficacy by drug, with nadolol being the only β-receptor blocker showing a significant risk reduction in CE (hazard ratio 0.40; p < 0.05). The authors concluded that nadolol is the only β-receptor blocker with a significant risk reduction in LQTS2 patients [41]. Recurrent CE occurred less frequently in patients with initially prescribed metoprolol, nadolol, and atenolol compared with propranolol (p = 0.002) [41]. Schwartz recommends that initial treatment of iLQTS should al-
ways involve β-receptor blockers, with propranolol and nadolol being the 2 most effective ones, whereas bisoprolol, metoprolol, atenolol, and carvedilol being less effective [42]. Similarly, Ackermann et al. [43], highlighted recently that the protective effect of the nonselective β-receptor blockers nadolol and propranolol in management of iLQTS is superior than that observed with β-1-selective receptor blockers. They report that there is substantial consensus among experts to prefer nadolol as a first-choice treatment in patients with iLQTS [43]. In line with these suggestions is the observation from Mazzanti et al. [44], showing the superiority of nadolol in risk reduction of CEs for iLQTS patients.

Discussion

β-receptor blockers are effective in reducing life-threatening CE in iLQTS and therefore should be continued during pregnancy and postpartum as pregnant iLQTS women are at a risk to develop CE [8–10, 17, 45]. β-receptor blockers are not teratogenic, but they are categorized by former FDA category as “C,” whereas atenolol is categorized as “D.” They can cause IUGR commonly and additionally bradycardia, apnea, hypoglycemia, and hyperbilirubinemia have been reported in the newborn infant [17–20]. Pregnant iLQTS women are not only at a risk for IUGR due to β-receptor blocker treatment, but also due to their underlying disease. IUGR can result in adverse neonatal outcomes. Fetuses with IUGR are at a risk for preterm delivery and they have a higher risk of mortality and perinatal complications [46]. Long-term outcome can be affected by an increased risk for neurodevelopmental impairment and growth delay [46]. Treatment recommendations for pregnant iLQTS women and data on IUGR in infants exposed prenatally to β-receptor blockers are scarce. LQTS management in pregnant women is challenging, as several factors addressing the fetus, such as IUGR and the mother, such as efficacy of the β-receptor blocker treatment, have to be taken into account.

Data on the efficacy of β-receptor blockers are heterogeneous but several studies and experts state that nonselective β-receptor blockers, such as nadolol and propranolol, might be superior in iLQTS management, particularly in patients with LQTS2 [35, 41–43]. Nadolol is approved by the FDA. Unfortunately, nadolol is not available in all European countries. Study data on prenatal nadolol exposure and IUGR are scarce. Nadolol has a 30% protein binding and a long half-life with a risk of causing adverse events in the newborn still several hours after birth [22]. The longer duration of action and its protein binding seems to make it less desirable for use in pregnancy [47]. Propranolol and nadolol, both nonselective β-receptor blockers, seem not to be the recommended first choice in general in pregnant women. However, if a woman with iLQTS was treated with nadolol effectively before becoming pregnant or if a LQTS2 genotype is known, continuation of the treatment with nadolol or start of nadolol should be considered. During our literature search we could identify most data related to IUGR for the β-1-selective receptor blocker atenolol. All available studies highlighted a negative impact on fetal growth, and therefore atenolol should be avoided in pregnancy. The other β-1-selective receptor blockers seem to affect less likely the uterine contraction and peripheral vasodilation and have therefore a lower rate of IUGR [17]. Based on the very limited data with respect to the current recommendation to favor in general β-1-selective receptor blockers in pregnancy there might be an indication that bisoprolol is the one with a relatively low risk for IUGR.

We have to emphasize that only scarce data are available for bisoprolol and IUGR (n = 5) and that data on the efficacy of bisoprolol in LQTS1/LQTS2 are limited, although, available studies show good and comparable efficacy to nadolol/propranolol in preventing major CE and well tolerated long-term tolerance [23, 39, 40]. Based on the fact that there are so few data available, we are not able to recommend the optimal choice of a β-receptor blocker for use in pregnant iLQTS women. For a personalized β-receptor blocker treatment, it is important, that each woman diagnosed with iLQTS who wants to become pregnant should be consulted by a multidisciplinary team consisting of cardiologists, gynecologists, pediatric cardiologists, neonatologists, and clinical pharmacologists. This team should evaluate the current medication for individual risks in respect to mother and fetus, underlying comorbidities of the mother, the iLQTS genotype, the pharmacologic β-receptor blocker properties, and risk-benefit assessment in case of a β-receptor blocker switch. Desirable drug properties in pregnancy due to reduced maternal-fetal exchange are, for example, a low lipophilicity, an intrinsic sympathomimetic activity, a high protein binding, and a large molecular weight [23, 48]. If the multidisciplinary team considers a β-receptor blocker switch (i) a wearable cardioverter defibrillator or (ii) an inpatient monitoring might be beneficial during the period of up-titration, which might be associated with a higher risk for CE. Irrespective of β-receptor blocker selection, each pregnant iLQTS women treated with a...
β-receptor blocker should be monitored regularly for fetal growth. It seems that pregnancy associated plasma protein, regular uterine artery Doppler, serial fundal height measurements, and middle cerebral artery Doppler are useful to early detect IUGR [49, 50]. Additionally, it is recommended to determine the time of delivery guided by umbilical artery and ductus venosus blood flow patterns [17]. After delivery, each infant who was prenatally exposed to a β-receptor blocker should be monitored for bradycardia, hypoglycemia, and respiratory depression, preferentially if possible in a neonatal care unit. Assuming linear pharmacokinetics for these β-receptor blockers, 95% will be eliminated after 5 times of their half-lives (Table 2) resulting in the following elimination time periods: (i) metoprolol after 25–50 h, (ii) propranolol after 15–50 h, (iii) carvedilol after 35–50 h, (iv) bisoprolol after 45–60 h, and for (v) nadolol in 100–120 h. Embryotox (www.embryotox.de) stated for bisoprolol, metoprolol, propranolol, and carvedilol that beta-receptor blocker-related side effects should improve in the infant 48 h after delivery [51].

There are several limitations of this review. Despite the fact that we used a comprehensive search strategy to identify relevant publications there is the risk of reporting bias as we might have missed some articles by our defined search strings and the fact that we have not performed a systematic review. Several studies were retrospective cohort studies, and both the underlying disease and the β-receptor blocker might have had a combined effect on IUGR. In addition, the majority of data on IUGR exposed infants is collected from pregnant women with hypertension and not from iLQTS patients. Furthermore, we cannot recommend a first-line β-receptor blocker treatment for pregnant iLQTS patients because in addition to IUGR other maternal and fetal factors have to be taken into account. We have summarized these factors to assist treating physicians in making the optimal choice for each individual pregnant woman with iLQTS.

**Conflict of Interest Statement**

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**Author Contributions**

Conceptualization, T.W., B.D., and J.N.v.d.A.; data curation, T.W.; writing – original draft preparation, T.W.; writing – review and editing, B.D. and J.N.v.d.A.; supervision, B.D. and J.N.v.d.A.

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