

1 **SARS-CoV-2, COVID-19 and inherited arrhythmia syndromes**

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38 **Abstract**

39 Ever since the first case was reported at the end of 2019, the SARS-COV-2 virus and associated
40 lung disease COVID-19 has spread throughout the world and has become a pandemic. In
41 particular, the high transmission rate of the virus has made it a threat to public health globally.
42 Currently, there is no proven effective therapy against the virus, and the impact on other
43 diseases is also uncertain, especially inherited arrhythmia syndrome.

44 Arrhythmogenic effect of COVID-19 can be expected, potentially contributing to disease
45 outcome. This may be of importance for patients with an increased risk for cardiac arrhythmias,
46 either secondary to acquired conditions or co-morbidities or consequent to inherited syndromes.

47 Management of patients with inherited arrhythmia syndromes such as Long QT syndrome,
48 Brugada syndrome, Short QT syndrome and Catecholaminergic Polymorphic Ventricular
49 Tachycardia in the setting of the COVID-19 pandemic may prove particularly challenging.

50 Depending on the inherited defect involved, these patients may be susceptible to pro-
51 arrhythmic effects of COVID-19-related issues such as fever, stress, electrolyte disturbances
52 and use of antiviral drugs. We here describe the potential COVID-19 associated risks and
53 therapeutic considerations for patients with distinct inherited arrhythmia syndromes and
54 provide recommendations, pending local possibilities, for their monitoring and management
55 during this pandemic.

56

57 **Introduction**

58 Ever since the first case was reported at the end of 2019, the SARS-COV-2 virus and
59 associated lung disease COVID-19 has spread throughout the world and has become a
60 pandemic. In particular, the high transmission rate of the virus has made it a threat to public
61 health globally.^{1,2} Currently, there is no proven effective therapy against the virus, and the
62 impact on other diseases is also uncertain.

63 SARS-CoV-2 is an RNA virus, a member of coronavirus family of viruses, similar to
64 SARS-CoV.³ Like SARS-CoV, SARS-CoV-2 infects humans by binding to the angiotensin-
65 converting enzyme 2 (ACE2) receptor on the surface of the cell through its spike domain.³
66 Infected patients present with a variety of manifestations. The most common clinical symptom
67 is fever (88.7%). Other symptoms include cough (67.8%), shortness of breath (18.7%), myalgia
68 or arthralgia (14.9%), headache (13.6%), diarrhea (3.8%), sore throat (13.9%), and sputum
69 production (33.7%) and fatigue (38.1%).⁴ Studies have shown that while the vast majority of
70 patients have minor symptoms, it is also possible for infected cases to become critically ill,
71 especially older individuals (above 60 years old) or patients with comorbidities.^{1,2} Severely
72 affected patients may have acute respiratory distress (15.6%) which requires invasive
73 mechanical ventilation (14.5%) and extracorporeal membrane oxygenation (2.9%).⁴

74 **Possible cardiac effects of SARS-COV-2 corona virus**

75 A registry of 1099 cases with COVID-19 reported a higher prevalence of hypertension
76 (23.7% vs. 13.4%) and coronary artery disease (5.8% vs. 1.8%) in severely affected versus
77 non-severely affected patients.⁴ Another study, of 138 hospitalized COVID-19
78 patients compared patients admitted to the intensive care unit (ICU) and non-ICU patients.
79 Higher rates of hypertension (58.3% vs. 21.6%, $p < 0.001$) and cardiovascular disease (25.0%
80 vs. 10.8%, $p = 0.04$) were observed in ICU patients.¹ This indicates that patients with pre-
81 existing cardiovascular disease may have a worse prognosis than others although age could be
82 one of the confounders. Furthermore, it is also essential to understand that although most
83 clinical presentations relate to the respiratory system, the disease may also impact on the
84 cardiovascular system.⁵ Besides the respiratory system, ACE2 is expressed in the human
85 cardiovascular system including the heart⁶ and a number of mechanisms have been put forward
86 whereby SARS-CoV-2 may cause myocardial injury. These include mechanisms involving
87 derangement of ACE2 signal pathways (animal studies have shown that cellular ACE2 levels
88 decrease upon SARS-CoV infection),⁶ cytokine storm and myocarditis.^{7,8} Occurrence of
89 myocardial involvement and severity thereof varies among affected individuals. While
90 myocardial damage evidenced by high cardiac markers such as hs-TnI has been
91 recognized⁹ and fulminant myocarditis has been reported,⁸ whether cardiovascular
92 complications include malignant arrhythmias is not yet known. In the afore-mentioned study
93 of 138 hospitalized COVID-19 patients, arrhythmia (not further specified) was reported in 17%

94 of total patients and in 16 of 36 patients admitted to the ICU.¹ Therefore, an arrhythmogenic
95 effect of COVID-19 could be expected, potentially contributing to disease outcome. This may
96 be of importance for patients with an increased risk for cardiac arrhythmias, either secondary
97 to acquired conditions, co-morbidities, or consequent to inherited syndromes. Management of
98 patients with inherited arrhythmia syndromes such as Long QT syndrome, Brugada syndrome,
99 Short QT syndrome and Catecholaminergic Polymorphic Ventricular Tachycardia in the
100 setting of the COVID-19 pandemic may prove particularly challenging. Depending on the
101 inherited defect involved, these patients may be susceptible to pro-arrhythmic effects of
102 COVID-19-related issues such as fever, stress, electrolyte disturbances and use of antiviral
103 drugs. Hence, additional precautions and preventive measures are recommended, including
104 ECG monitoring, aggressive antipyretic treatment, and more stringent social distancing to
105 prevent infection.¹⁰ We here describe the potential COVID-19 associated risks and therapeutic
106 considerations for patients with distinct inherited arrhythmia syndromes and provide
107 recommendations for their monitoring and management during this pandemic.

108 **Long QT syndrome**

109 The Long QT syndrome (LQTS) is characterised by abnormally prolonged ventricular
110 repolarization and an increased risk of the malignant arrhythmia *Torsades de Pointes* and
111 ventricular fibrillation that may lead to sudden death. LQTS is an inheritable condition caused
112 by pathogenic variants in genes encoding ion channels (primarily *KCNQ1*, *KCNH2*, *SCN5A*).

113 An often-faced clinical situation, however, is acquired QT-interval prolongation, that occurs
114 for instance during myocardial ischemia, hypothermia, as a result of treatment with a wide
115 range of drugs, hypokalaemia or sepsis. Severe QTc-prolongation due to these conditions might
116 similarly result in malignant arrhythmias. Rather commonly, patients who have severe forms
117 of acquired QT-prolongation also have a genetic predisposition for QTc-prolongation,^{11,12} but
118 without such extreme provocation these patients generally have normal QT-intervals. In fact,
119 many LQTS patients may also have QT-intervals within normal limits in resting conditions,¹³
120 although this still puts them at higher risk for malignant arrhythmias,¹⁴ especially during
121 provocations such as the use of QTc-prolonging drugs.¹⁵ Whereas severe forms of inherited
122 LQTS often surface during (early) childhood (from infants to adolescents),^{14,16} acquired QT-
123 prolongation generally occurs in older patients because these critical provocative events more
124 often occur in older patients.

125 *Long QT syndrome and COVID-19*

126 There are several issues that require attention when discussing COVID-19 in relation to
127 inheritable or acquired QT-prolongation.

128 The most important determinant of risk for malignant arrhythmias in patients with LQTS
129 or in acquired QT-prolongation, is the use of one or more QTc prolonging drugs in the setting
130 of severe manifestations of COVID-19. Many drugs (either with cardiac or non-cardiac
131 indications) have the ability to block cardiac potassium currents, impairing ventricular

132 repolarisation with subsequent prolongation of the QT-interval and an increased risk for
133 malignant arrhythmias.¹⁵ In addition, many drugs may alter drug metabolism, e.g. due to
134 inhibition of CYP3A4, which may further increase plasma levels of QT-prolonging drugs and
135 further increase risk. Of special interest in COVID-19 is that there are indications that
136 chloroquine and hydroxychloroquine might be of value.¹⁷

137 Chloroquine is one of the most widely used anti-malarial drugs world-wide, but it has also
138 been investigated as a potential broad-spectrum anti-viral drug.¹⁸ Amongst its mechanisms,
139 chloroquine appears to interfere with the terminal glycosylation of ACE2 and may thus
140 negatively influence virus-receptor binding and abrogate infection.¹⁹⁻²¹ However, chloroquine
141 is closely related to quinidine, and while the latter is used as an anti-arrhythmic drug in Brugada
142 syndrome and idiopathic forms of ventricular fibrillation, it is also well known for its QT-
143 prolonging effects and has been associated with QT related malignant arrhythmias. Luckily,
144 the QT-prolonging effect of chloroquine is very modest, and in general it does not result in
145 clinically significant QT-prolongation in patients without LQTS.²² Hydroxychloroquine sulfate,
146 a less toxic derivative of chloroquine, is widely used in the chronic treatment of autoimmune
147 diseases without significant effects on ECG parameters,²³ and was recently shown to also
148 efficiently inhibit SARS-CoV-2 infection *in vitro*.²⁴ However, both chloroquine and
149 hydroxychloroquine are metabolised by CYP3A4, and COVID-19 treatment with
150 (hydroxy)chloroquine can be combined with additional anti-viral treatments such as ritonavir

151 plus lopinavir (both potent CYP3A4 inhibiting drugs; their combination is associated with QT-
152 prolongation), azithromycin (besides a macrolide antibiotic also investigated for its antiviral
153 properties, with also (weak) CYP3A4 inhibition and associated with QT-prolongation)^{25,26}, or
154 remdesivir (an investigational drug for which metabolism and possible QT prolonging effects
155 are not yet resolved). Combining (hydroxy)chloroquine with these drugs might thus result in
156 higher plasma levels and significant QT-prolongation. Hence, we advise monitoring QT-
157 intervals and cardiac rhythm if starting these drugs given the increased risk for malignant
158 arrhythmias (Figure 1). In addition, physicians should be aware of the alpha-blocking effects
159 of (hydroxy)chloroquine, which might result in hypotension.

160 Another issue is fever. The effect of fever is, in contrast to patients with for example BrS
161 (see below), much less evident in patients with LQTS. A possible exception are patients, with
162 specific LQTS 2 mutations, presenting with fever-triggered arrhythmias which are based on
163 temperature sensitive mutant channels (i.e. less current with higher temperature).²⁷ As most
164 patients hospitalised for COVID-19 have fever,⁴ patients with known LQTS will thus generally
165 not be at increased risk. The separate contribution of fever in acquired QT-prolongation is not
166 well known, but sepsis is a denominator of risk of acquired QT-prolongation²⁸, and septic shock
167 is one of the clinical scenarios in COVID-19.⁴

168 Finally, interpretation of the QT-interval is not easy,²⁹ but guidance is available.¹³ While
169 COVID-19 patients admitted to Intensive Care Units will often have continuous ECG

170 monitoring available, ECG monitoring of inpatients who are being treated in an airborne
171 isolation room can be challenging. Nevertheless, if possible, we advise (Figure 1) to
172 monitor QT-intervals at baseline and at 4h after administration of (hydroxy)chloroquine and/or
173 anti-viral therapy in patients with congenital or acquired LQTS, patients already taking other
174 QT-prolonging drugs, and patients with structural heart disease or bradycardia. A second ECG
175 is recommended after 1-3 days. In all other patients, QTc-interval monitoring should be
176 performed 24h after start of therapy. During the course of (hydroxy)chloroquine and/or anti-
177 viral therapy, QTc-interval monitoring is furthermore indicated in case of worsening
178 kidney/liver function and electrolyte disorders (in particular K^+ , Ca^{2+} and Mg^{2+}), especially in
179 LQTS patients or patients with abnormal QT-intervals at baseline. Of particular concern is the
180 COVID-19 associated diarrhea which may lead to hypokalemia with adverse effects on the
181 QTc interval. In addition, beta-blocker treatment should be considered if the patient is not yet
182 treated. Cardiologists throughout Europe, Canada and the US have initiated a QT-interval
183 registry for COVID-19 patients treated with chloroquine, hydroxychloroquine and/or anti-viral
184 drugs and contribution is open to all.

185 In summary, we advise (Figure 1):

- 186 • QTc-interval monitoring when using (hydroxy)chloroquine in COVID-19 patients
- 187 • QTc-interval monitoring when using or combining anti-viral drugs in COVID-19
188 patients

- 189 • QTc-interval monitoring in patients with known LQTS, acquired QT-prolongation or
190 conditions associated with acquired QT-prolongation (e.g. use of other QT-prolonging
191 drugs, structural heart disease, bradycardia <50/min, liver and renal disease)
- 192 • When QTc is above 500msec, we advise consultation with a cardiologist (“QT-
193 specialist”) for guidance (which might, e.g., result in intensified monitoring, raising
194 potassium levels, and/or discontinuation of one or more QT-prolonging drugs)
- 195 • Patients with acquired LQTS or patients using a combination of QT-prolonging drugs
196 should have a high serum potassium level. Avoiding hypokalemia is not enough and
197 the adagium should be "a serum potassium of 5 is better than 4."³⁰

198 **Brugada syndrome**

199 Brugada syndrome (BrS) is a familial arrhythmia syndrome disorder characterized by the
200 type 1 Brugada ECG pattern in the right precordial leads of the ECG (coved type ST-elevation
201 and T wave inversion in lead V1 and/or V2) and an increased risk for ventricular fibrillation
202 and sudden cardiac death. Up to 30% of patients with BrS carry a loss-of-function pathogenic
203 variant (mutation) in *SCN5A*, the gene that encodes the cardiac sodium channel, as the
204 pathophysiological substrate of their disease.³¹ The most frequently used drugs for SARS-CoV-
205 2 and COVID-19 patients are not on the list of drugs to be avoided by BrS patients.³² However,
206 attention to BrS patient management is relevant in the setting of the SARS-CoV-2 outbreak
207 since ECG manifestations of the disorder may be uncovered during fever, and since fever has

208 been unequivocally associated with life-threatening arrhythmic events (LTE) in patients with
209 the disorder.³³

210 The importance of fever in BrS patients is now well-established.³³⁻³⁵ In 24 patients with
211 BrS, 3 of whom had a fever-triggered cardiac arrest, the increase in body temperature reduced
212 the PR interval in control individuals, but increased PR interval, QRS width, and the maximum
213 J-point in BrS patients.³⁴ Another study showed that fever-associated BrS seems to be
214 associated with a higher future risk of LTE's compared to drug-induced type 1 pattern.³⁵ Finally,
215 fever seems to be particularly relevant in children.³³ Indeed, in a registry with symptomatic
216 BrS patients (the SABRUS registry) approximately 6% of LTE's were associated with fever
217 and the highest rate of fever-triggered LTE's was observed in the very young (65%, age \leq 5
218 years). In the age range 16 to 70 years, only 4% of the LTE's was related to fever. In the elderly
219 (>70 years) this percentage increased to 25%.³³

220 In the setting of fever, the presence of a pathogenic variant in *SCN5A* may be particularly
221 relevant. In a single center series of 111 patients with BrS, 22 presented with a cardiac arrest,
222 4 of which were fever related. Three of these 4 patients harbored a pathogenic variant in
223 *SCN5A*.³⁴ In the SABRUS registry, the percentage of *SCN5A* pathogenic variants was 77% in
224 children and 27% in adults with a LTE.³³ The authors also performed an analysis of all
225 published cases (up to 2018) with fever-triggered LTE's (40 patients in 22 reports) revealed
226 the presence of a putatively pathogenic variant in *SCN5A* was found in 13 (68%) of 19 patients

227 tested.³³ Moreover, in a multicenter pediatric population of 106 patients, 10 patients had a LTE
228 during follow-up, which was triggered by fever in 27%; all of the latter patients were positive
229 for a pathogenic *SCN5A* variant. Finally, preliminary data in a pediatric cohort indicated that
230 mainly children with a *SCN5A* mutation developed a type 1 ECG during fever (43.8% of
231 children who developed a type 1 ECG during fever had a *SCN5A* mutation vs 4.2% of children
232 without a type 1 during fever) and had events during follow-up (7/21 vs 0/47).³⁶ These studies
233 collectively indicate that sodium channel function is sensitive to temperature. This sensitivity
234 may be due to altered temperature-sensitive kinetics, in particular accelerated inactivation,³⁷
235 and/or decreased sodium channel expression at higher temperatures.³⁸ Also in other sodium
236 channel mediated diseases, increased temperature sensitizes patients to disease-related
237 symptoms.^{39,40}

238 Based on the above we feel that the following recommendations are pertinent:

- 239 1. All patients with Brugada syndrome should self-treat with
240 paracetamol/acetaminophen immediately if they develop signs of fever and self-isolate.
- 241 2. Patients without an ICD who are at higher risk due to fever include:
 - 242 a. sodium channel disease with or without a type 1 ECG pattern,
 - 243 b. children and young adults (under 26 years old) and the elderly (over 70 years)
 - 244 with Brugada syndrome; and

245 c. all patients with a spontaneous type 1 Brugada pattern and/or cardiac syncope.

246 3. If these higher risk patients develop a high fever (>38.5C) despite paracetamol
247 treatment, they will need to attend the emergency department*. The emergency
248 department must be forewarned to allow assessment by staff with suitable protective
249 equipment. Assessment should include an ECG** and monitoring for arrhythmia. If an
250 ECG shows the type 1 Brugada ECG pattern, then the patient will need to be observed
251 until fever and/or the ECG pattern resolves. If all ECGs show no sign of the type 1 ECG
252 pattern, then they can go home to self-isolate.

253 4. Patients who are not part of the higher risk group and have a drug-induced type 1 ECG
254 pattern, no symptoms of syncope and no sign of a spontaneous type 1 pattern at any
255 other time are at lowest risk and can afford to self-isolate at home. The risk of visiting
256 the emergency department and contracting COVID-19 is likely to outweigh the risk of
257 a LTE. Attendance at hospital should then be dictated by other clinical features, such
258 as palpitations or (pre-)syncope etc. The same advice holds for patients with an ICD.

259 * attendance at the emergency department may require regulation according to the capacity
260 of service and risk of COVID-19 infection.

261 ** ideally three different ECGs with V1 and V2 in the 4th, 3rd and 2nd intercostal spaces

262 Management in the hospital should include monitoring of ECG abnormalities and
263 arrhythmia, as well as efforts to reduce the body temperature (with antipyretic drugs, preferably
264 paracetamol/acetaminophen, or eventually ibuprofen). More generally, BrS patients, in
265 particular those with a pathogenic or likely pathogenic variant in *SCN5A*, are advised to self-
266 isolate in their private environment.

267 **Short QT syndrome**

268 Short QT syndrome (SQTS) is a familial arrhythmia syndrome characterized by short QT
269 intervals on the ECG and a significant rate of ventricular arrhythmias.⁴¹ It is a heterogeneous
270 disease caused by pathogenic variants in at least three different potassium channel genes
271 (*KCNH2*, *KCNQ1* and *KCNJ2*) and the cardiac chloride-bicarbonate exchanger gene
272 (*SLC4A3*).⁴² It is an extremely rare disease; in a recent systematic literature review only 110
273 cases were described.⁴³ No specific triggers for LTE, including fever, have been described.
274 Hence, based on current knowledge, SQTS patients do not seem to be at particular risk when
275 they are affected by COVID-19.

276 Potential drugs for COVID-19 patients, like chloroquine, might actually be beneficial for
277 SQTS patients due to lengthening of their QT-interval, as has been suggested by modelling
278 data for SQTS type 1 (*KCNH2*-related⁴⁴) and type 3 (*KCNJ2* related^{44,45}). There are no clinical
279 data as far as we are aware.

280 We therefore do not believe that there is a particular concern when SQTS patients are
281 infected with SARS-CoV-2.

282 **Catecholaminergic Polymorphic Ventricular Tachycardia**

283 Catecholaminergic Polymorphic Ventricular Tachycardia (CPVT) is a familial arrhythmia
284 syndrome characterized by adrenergic-related ventricular arrhythmias (i.e. during exercise, or
285 stress).⁴¹ It is a heterogeneous disease with pathogenic variants in *RYR2* encoding the human
286 Ryanodine receptor 2 as the most important contributor.⁴⁶ First line treatment comprises
287 intensive beta blocker therapy. In insufficiently responsive cases flecainide should be added or
288 left sympathetic denervation should be conducted.^{41,46} ICD therapy should be avoided.⁴⁷

289 As mentioned above, exercise and emotional circumstances constitute specific triggers for
290 LTE. An increased heart rate alone (pacing-induced), as an important symptom of fever, does
291 not appear to be sufficient for the induction of ventricular arrhythmias.⁴⁸ Fever, as a specific
292 trigger has not been described. Whether or not the stressful circumstances that COVID-19
293 patients find themselves in will lead to an increased burden of arrhythmias can only be
294 speculated upon.

295 The antiviral therapy proposed for COVID-19 is not expected to lead to increased risk.
296 The only potential deleterious pharmacological interaction in these patients are drugs with
297 alpha or beta adrenoceptor mimetic activity, which may be used in cases in need of
298 hemodynamic support. Intravenous epinephrine has been used to unmask ventricular

299 arrhythmias and initial data suggested that epinephrine was more effective than exercise testing
300 in unmasking ventricular arrhythmias.⁴⁹ Later studies revealed, however, a low sensitivity and
301 high specificity (with the exercise test as the gold standard⁵⁰). Nevertheless, based on their
302 pathophysiological mechanism of action, epinephrine, isoproterenol and dobutamine, all alpha
303 and/or B1 receptor agonists, should probably be avoided. Milrinone, the most widely used
304 phosphodiesterase 3 inhibitor, acts by decreasing the degradation of cyclic adenosine
305 monophosphate (cAMP). This may potentially stimulate the RyR2 receptor and must thus be
306 used with caution. However, with continuation of the beta blockers (as we recommend, see
307 below) this may not be that relevant because betablockers suppress milrinone-induced
308 increased Ca-leak.⁵¹ CPVT patients, in particular those who were symptomatic prior to
309 diagnosis, should stay on their beta blocker treatment with or without flecainide as long as is
310 tolerated hemodynamically. Flecainide does have interactions with Ritonavir/Lopinavir and
311 chloroquine, yet we believe that it is an important enough therapy not to stop in these
312 particularly stressful circumstances.

313 Based on the above we also suggest avoidance of epinephrine in the setting of a VT/VF
314 arrest if possible. This is probably the only resuscitation setting where epinephrine is
315 contraindicated.⁵²

316 **Conclusion**

317 Patients with inherited arrhythmia syndromes may be at an increased pro-arrhythmic risk
318 in the setting of COVID-19 infection, necessitating additional precautions and specialized
319 management. Preventive measures should include stringent social distancing to prevent
320 infection, aggressive antipyretic treatment to reduce fever in Brugada syndrome patients, and
321 ECG monitoring in Long QT syndrome patients treated with antiviral drugs.

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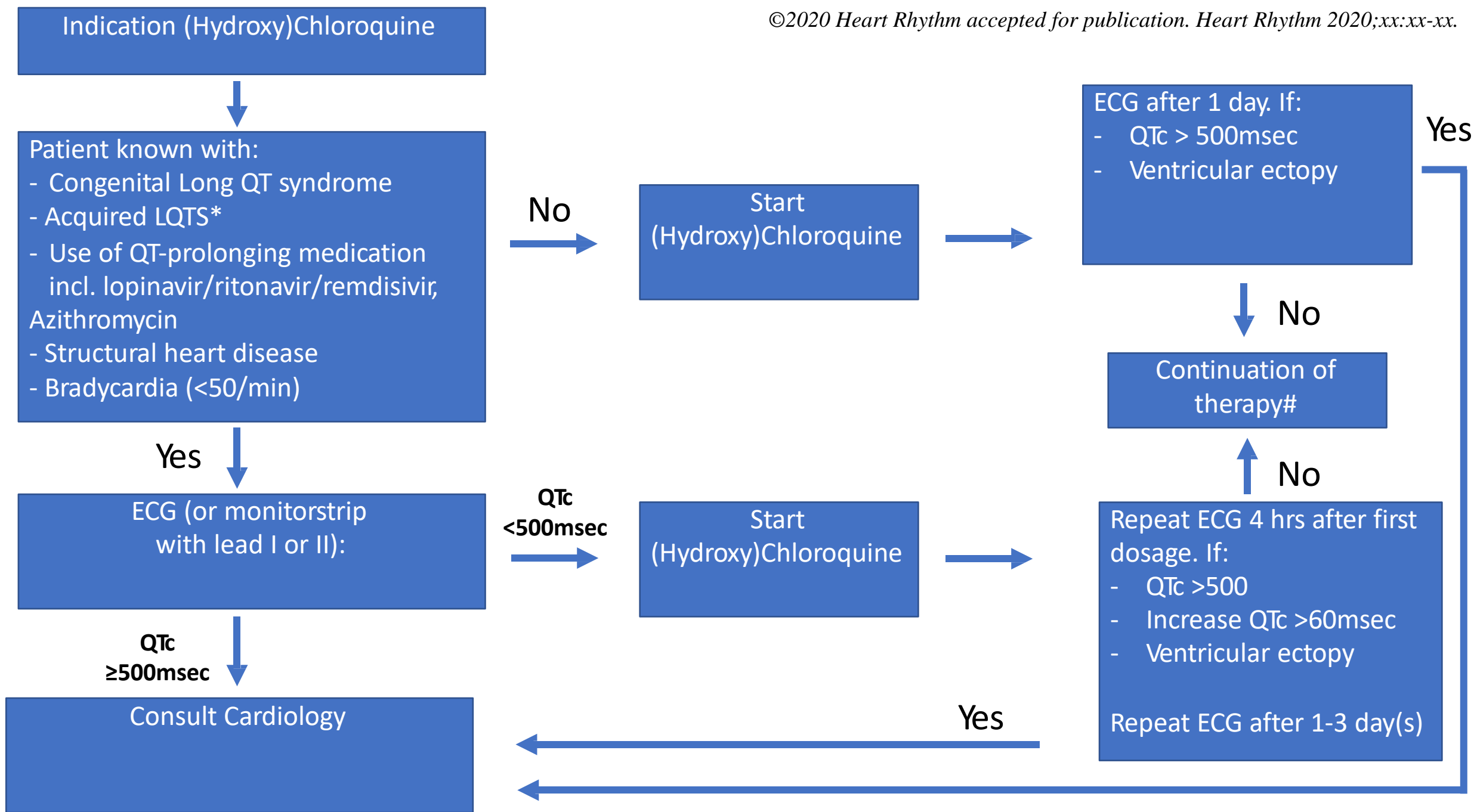
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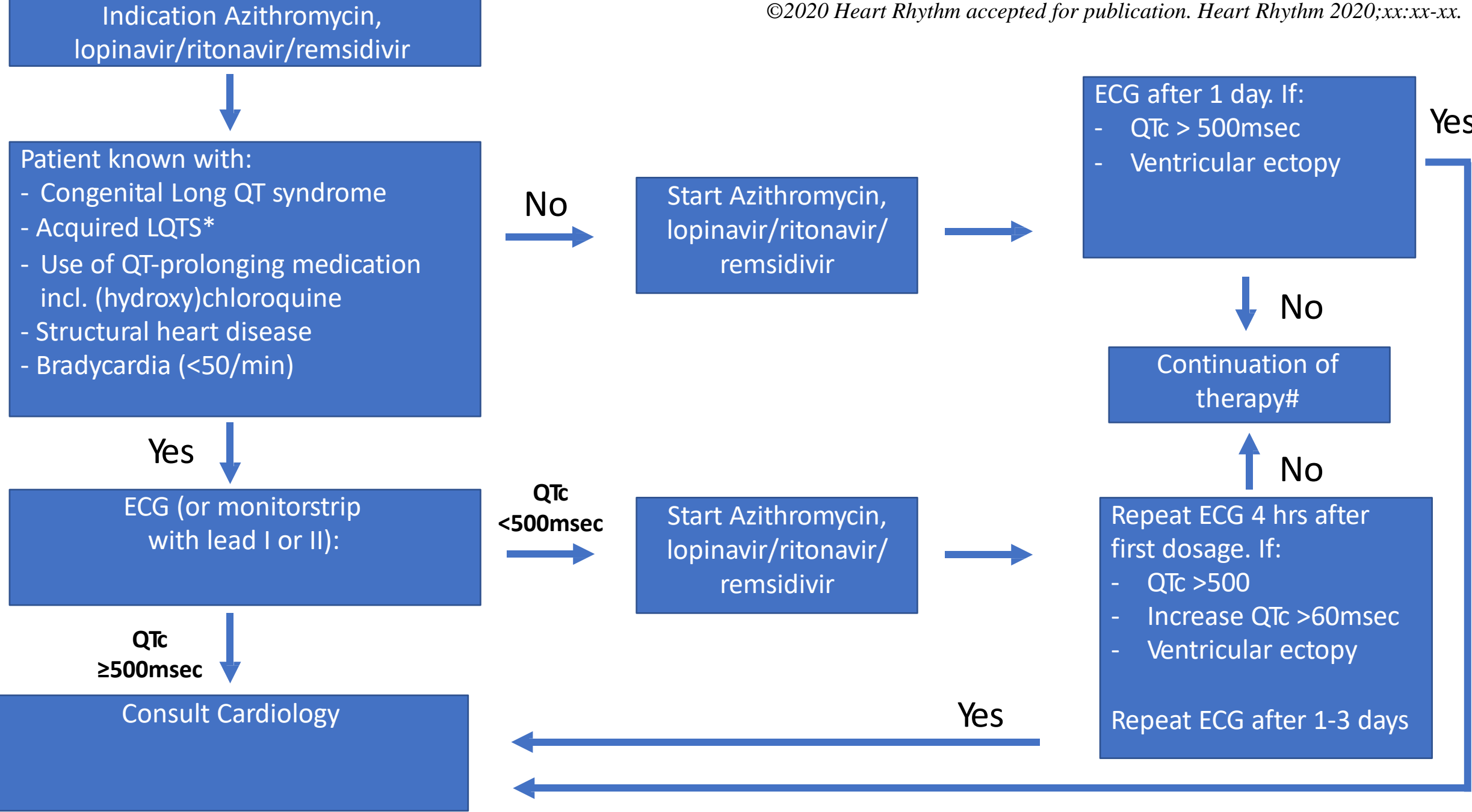
454 **Figure legends**

455 Figure 1: Flowchart of proposed guidance of QTc monitoring in patients receiving
456 (hydroxy-)chloroquine and/or antiviral drugs and /or azathromycin. It should be noted that not
457 every LQTS patient has the same risk. The length of the QTc interval is of importance (as is
458 implicit in the flowchart) but also gender, age and the genotype are important. LQT2 patients
459 may be at higher risk than LQT1 patients for example. The consulted cardiologist should have
460 sufficient experience with QT-related arrhythmic problems.

461



*: earlier QTc prolongation with medication; #: if ventricular ectopy, arrhythmia, dizziness or loss of consciousness: consult cardiology



*: earlier QTc prolongation with medication; #: if ventricular ectopy, arrhythmia, dizziness or loss of consciousness: consult cardiology