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1 **Ivermectin Prophylaxis Used for COVID-19 Reduces COVID-19 Infection and**
2 **Mortality Rates: A City-Wide, Prospective Observational Study of 223,128**
3 **Subjects Using Propensity Score Matching.**

4
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29 **Key-words:** COVID-19, SARS-CoV-2, ivermectin, prophylaxis, prevention,
30 coronavirus

31
32 **Acromyums:** COPD = Chronic Obstructive Pulmonary Disease; CVD = cardiovascular
33 disease; MI = Myocardial infarction; T2D = Type 2 Diabetes

41 **Abstract**

42

43 **Background:** Ivermectin has demonstrated different mechanisms of action that
44 potentially protect from both COVID-19 infection and COVID-19-related comorbidities.
45 Based on the studies suggesting efficacy in prophylaxis combined with the known safety
46 profile of ivermectin, a citywide prevention program using ivermectin for COVID-19 was
47 implemented in Itajaí, a Southern city in Brazil in the state of Santa Catarina. The
48 objective of this study was to evaluate the impact of regular ivermectin use on subsequent
49 COVID-19 infection and mortality rates.

50 **Materials and methods:** We analyzed data from a prospective, observational study of
51 the citywide COVID-19 prevention with ivermectin program which occurred between
52 July 2020 to December of 2020 in Itajaí, Brazil. Study design, institutional review board
53 approval, and analysis of registry data occurred after completion of the program. The
54 program consisted of inviting the entire population of Itajaí to a medical visit in order to
55 enroll in the program and to compile baseline, personal, demographic and medical
56 information. In the absence of contraindications, ivermectin was offered as an optional
57 treatment to be taken 2 consecutive days every 15 days at a dose of 0.2mg/kg/day. In
58 cases where a participating citizen of Itajaí became ill with COVID-19, they were
59 recommended to not use ivermectin or any other medication in early outpatient treatment.
60 Clinical outcomes of infection, hospitalization, and death were automatically reported
61 and entered into the registry in real time. Study analysis consisted of comparing
62 ivermectin users with non-users using cohorts of infected patients propensity score
63 matched (PSM) by age, sex, and comorbidities. COVID-19 infection and mortality rates
64 were analyzed with and without use of propensity score matching.

65 **Results: Of the 223,128 citizens of Itajaí considered for the study, a total of 159,561**
66 **subjects were included in the analysis; 113,845 (71.3%) regular ivermectin users and**
67 **45,716 (23.3%) non-users. Of these, 4,311 ivermectin users were infected, among**
68 **which 4,197 from the city of Itajaí (3.7% infection rate) and 3,034 non-users (from**
69 **Itajaí) were infected (6.6% infection rate), a 44% reduction in COVID-19 infection**
70 **rate (Risk ratio (RR), 0.56; 95% confidence interval (95%CI), 0.53 – 0.58; p <**
71 **0.0001). Using PSM, two cohorts of 3,034 subjects suffering COVID-19 infection were**
72 **compared. The regular use of ivermectin led to a 68% reduction in COVID-19 mortality**
73 **[25 (0.8%) versus 79 (2.6%) among ivermectin non-users; risk ratio (RR), 0.32; 95%**
74 **confidence interval (CI), 0.20 – 0.49; p < 0.0001]. When adjusted for residual variables,**

75 reduction in mortality rate was 70% (RR, 0.30; 95%CI 0.19 – 0.46; p < 0.0001). There
76 was a 56% reduction in hospitalization rate (44 versus 99 hospitalizations among
77 ivermectin users and non-users, respectively; RR, 0.44; 95%CI, 0.31 – 0.63; p < 0.0001).
78 After adjustment for residual variables, reduction in hospitalization rate was 67% (RR,
79 0.33; 95%CI 0.23 – 0.66; p < 0.0001).

80 **Conclusion:** In this large, propensity score matched study, regular use of ivermectin as a
81 prophylactic agent was associated with significantly reduced COVID-19 infection,
82 hospitalization, and mortality rates.

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109 **Introduction**

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111 Ivermectin has been demonstrated to have not only extensive anti-parasitic actions^{1,2}, but
112 also anti-viral, anti-bacterial, and anti-protozoan properties. Ivermectin has been long
113 proposed for use as a repurposed antiviral agent⁴⁻⁶. Indeed, antiviral effects of ivermectin
114 have been reported against both RNA and DNA types of viruses, including HIV-1,
115 Yellow fever (YFV), Japanese encephalitis, tick-borne encephalitis, West Nile, Zika
116 (ZKV), Dengue fever, Chikungunya (CHIKV), Venezuelan equine encephalitis and the
117 Pseudorabies virus^{3,5,7}, as well as functioning in regulation of proteins involved in
118 antiviral responses⁸.

119

120 Additional actions of ivermectin described include agonism activity to the X-LBD
121 binding receptor (FXR), with multiple potential metabolic benefits^{9,10}; neuronal
122 regeneration^{11,12}, prevention of muscle hypoxia¹³, anti-inflammatory activity to
123 Interferon (INF)¹⁴, nuclear factor- κ B (NF- κ B), lipopolysaccharide (LPS)¹⁵ and JAK-
124 STAT pathway, PAI-1^{16,17}; generation of P21 activated Kinase 1 (PAK-1)^{18,19}; reduction
125 of Interleukin-6 (IL-6) levels¹⁵; allosteric modulation of P2X4 receptor²⁰; inhibition of
126 high mobility group box 1 (HMGB1)^{21,22}; suppression of mucus hypersecretion,
127 diminished recruitment of immune cells and production of cytokines in the lung²³.
128 Ivermectin is also described to induce Th1-type immune response against protozoans²⁴,
129 and anti-coagulant action through binding to the S protein of some viruses²⁵.

130

131 The hypothesis that ivermectin could be protective against COVID-19 is
132 substantiated by its multi-pathway, anti-inflammatory effects^{15,26} and multi-antiviral
133 mechanisms. COVID-19 pathogenesis is largely understood as an inflammation-mediated
134 hemagglutinating infection disrupting pulmonary, vascular and endothelial systems,
135 leading to a multi-systemic disease. *In vitro* and *in-silico*, ivermectin has demonstrated
136 anti-SARS-CoV-2 activity through more than 20 direct and indirect mechanisms^{2,27,28}.

137

138 Ivermectin has demonstrated preliminary protective effects against SARS-CoV-2
139 infection in terms of reducing times to clinical recovery, rates of disease progression and
140 mortality^{2,29,30}. However, more robust studies with larger sample sizes are still

141 recommended to confirm the possible beneficial effects ivermectin confers in COVID-
142 19.

143

144 Since the onset of the COVID-19 pandemic, the use of inexpensive options based
145 on a consistently beneficial signal of efficacy, a well-established safety profile,
146 favourable cost-effectiveness, ivermectin is a highly attractive intervention for the patient
147 centred medicine practiced by frontline clinicians, with use aligning strongly with the
148 bioethical principles for medical practice outlined in Article 36 of the Helsinki
149 declaration³¹.

150

151 However, despite this favorable risk/benefit profile and absence of therapeutic
152 alternatives, ivermectin has yet to be approved for prophylaxis and treatment of COVID-
153 19 by agencies throughout the world, including FDA (Food & Drug Administration;
154 United States of America), EMA (European Medicines Agency; Europe) and ANVISA
155 (Agência Nacional de Vigilância Sanitária – Brazilian Health Regulatory Agency;
156 Brazil).

157

158 The ability to prescribe ivermectin or any other off-label drug for COVID-19 has
159 long been at the discretion of frontline physicians once all risks, uncertainties, potential
160 benefits, and patients' rights are exposed, and informed consent has been obtained. Of
161 particular note, in Brazil, this follows the medical autonomy to determine the best
162 therapeutic strategies for individuals, as per the Medical Code of Ethics of the Brazilian
163 Board of Medical Doctors; the Federal Council of Medicine – Conselho Federal de
164 Medicina (CFM), that determines the obligations and rights of medical doctors in
165 Brazil³².

166

167 **Since vaccines for COVID-19 were not available in Brazil until 2,021, and the**
168 **lack of prophylactic alternatives in the absence of vaccines**, Itajai, a city in the
169 Southern Brazilian state of Santa Catarina, initiated a population wide government
170 program for COVID-19 prophylaxis. The medical-focused decision parameters
171 established are based on the distribution of ivermectin to whole populations in different
172 countries. To ensure the safety of the population, a well-controlled computer program
173 was developed to compile and maintain all relevant demographic and clinical data. The

174 use of ivermectin was optional and based on patients' preferences given its benefits as a
175 preventative agent was unproven.

176

177 This study's objective is to assess the impact on important clinical outcomes when
178 ivermectin is used as prophylaxis for COVID-19. The prophylaxis program occurred in
179 addition to the standard non-pharmacological strategies of masking and social distancing,
180 as part of a citywide program conducted in outpatient settings.

181

182

183 **Material and Methods**

184

185 *Study population*

186

187 This was a prospective, observational study. Although study design, IRB approval, and
188 data analysis occurred after completion of the voluntary prophylaxis program, all data
189 were collected prospectively in real-time with mandated reporting to the registry of all
190 events as they occurred during the citywide governmental COVID-19 prevention with
191 ivermectin program, from July 2020 to December 2020, developed in the city of Itajaí, in
192 the state of Santa Catarina, Brazil. Demographic and clinical data was reported from
193 medical records of patients followed in a large outpatient setting; a provisional outpatient
194 clinic set in the Convention Center of Itajaí, and several secondary outpatient settings, as
195 part of the Universal Health System (SUS).

196

197 The objective was to determine the number of patients affected by COVID-19
198 (positivity rate of rtPCR-SARS-CoV-2), risk of death due to COVID-19 (whether
199 infected or not), and COVID-19 mortality rate (risk of death from COVID-19) of those
200 who used and did not use ivermectin prophylactically for COVID-19. This data was
201 analyzed stratified by age, sex, presence of comorbidities, and correlated demographic
202 characteristics.

203

204 The present retrospective analysis of the prospectively collected data was
205 approved by the CONEP - National Research Ethics Council (CONEP) under the number
206 4.821.082 with the project number CAAE: 47124221.2.0000.5485. Although study
207 design, IRB approval, and data analysis occurred after completion of the voluntary

208 prophylaxis program, all data were collected prospectively in real-time with mandated
209 reporting to the registry of all events as they occurred during the citywide governmental
210 COVID-19 prevention with ivermectin program, from **July 7, 2020 to December 2,**
211 **2020**, developed in the city of Itajaí, in the state of Santa Catarina, Brazil.

212

213

214 *Study procedures and data collection*

215

216 Optional, voluntary prophylactic use of ivermectin was offered to patients during regular
217 medical visits between July 7, 2020 and December 31, 2020 in 35 different sites,
218 including 34 local SUS health centres and a large temporary patient setting. Doctors
219 working in these sites were free to prescribe ivermectin prophylactically. Subjects that
220 did not use ivermectin either refused or their primary care physicians opted not to offer
221 ivermectin.

222

223 The program was conducted in all 35 sites, 24/7, with the initial enrollment in the
224 program occurring during a two-week time frame, due to the large number of subjects to
225 evaluate in the entire population of Itajaí. In order to avoid underreported data, strict
226 procedure sequencing was followed: 1. Registration and recording of patient data,
227 documented by assistants; 2. Weighing subjects (Subject weight was essential to calculate
228 the appropriate dose of ivermectin); 3. Brief medical evaluation of past medical history,
229 comorbidities, use of medications and contraindications to drugs; 4. Medical prescription
230 of prophylactic doses of ivermectin, according to medical judgment and following a
231 subject's informed consent related to potential benefits, risks, and side effects. **Regarding**
232 **drug interactions with ivermectin, use of warfarin was a contra-indication for**
233 **prophylaxis with ivermectin due to drug interactions. Subjects under chronic use of**
234 **glucocorticoids, protease inhibitors and anti-epileptics were recommended to**
235 **schedule regular medical visits every six to eight weeks. Subjects were recommended**
236 **to inform medical doctors about the use of ivermectin, in case one or more of the**
237 **following medications were prescribed: warfarin, azithromycin, dexamethasone,**
238 **prednisone or prednisolone (Hydrocortisone or cortisone are not commercially**
239 **available in regular pharmacies in Brazil).**

240

241 All details of this citywide program and campaign had been previously agreed
242 upon between the city local department of the National Healthcare System (SUS), city
243 mayor, and local public prosecutors.

244

245 The following data were analyzed, adjusted as confounding factors, and used as
246 variables for balancing and matching groups for the employment of propensity score
247 matching (PSM) in the present study: age, sex, past medical history, previous diseases;
248 myocardial infarction (MI), stroke: existing comorbidities; type 2 diabetes (T2D), asthma,
249 chronic obstructive pulmonary disease (COPD), hypertension, dyslipidemia,
250 cardiovascular diseases (CVD), cancer (any type), and other pulmonary diseases: habits
251 (past or current smoking). Additional data analyzed included self-reported comorbidities
252 and medications used.

253

254 Patients who presented signs or the diagnosis of COVID-19 before July 7, 2020,
255 were excluded from the sample. Other exclusion criteria were contraindications to
256 ivermectin and subjects below 18 years of age. The dose and frequency of ivermectin
257 treatment was 0.2mg/kg/day; *i.e.*, giving one 6mg-tablet for every 30kg. for 2 consecutive
258 days every 15 days.

259

260 During the study, subjects who became infected with COVID-19 were diagnosed
261 with a positive rtPCR-SARS-CoV-2 and then underwent a specific medical visit to assess
262 COVID-19 clinical manifestations and severity. All subjects were recommended not to
263 use ivermectin, nitazoxanide, hydroxychloroquine, spironolactone or any other drug
264 claimed to be effective against COVID-19. The city did not provide or support any
265 specific pharmacological outpatient treatment for subjects infected with COVID-19.

266

267 They were questioned for the presence of common COVID-19 symptoms. These
268 included chills, high-grade fever, cough, myalgia, fatigue, anosmia, ageusia, sore throat,
269 headache, nasal congestion, sneeze, runny nose, hemoptysis, nauseas, vomiting,
270 abdominal pain, diarrhea, cutaneous rash, arthralgia, chest pain, eye pain and pinkeye,
271 and presence of alert signs, including shortness of breath, signs of hypoxia, signs of
272 coagulation abnormalities and an altered level of consciousness. Systolic and diastolic
273 blood pressure, heart rate, respiratory rate, oxygen saturation, and axillar temperature
274 were measured. The same signs and symptoms, and vital signs were collected at each

275 following medical visit during COVID-19. Individual data was compiled and reviewed
276 by the researchers.

277

278 Registry data of all patient records from the city of Itajaí between July 7, 2020 and
279 December 2, 2020, including those who used ivermectin and did not use ivermectin were
280 reviewed. Subjects who tested positive for COVID-19 during the study were considered
281 for this analysis, whether they used ivermectin or not. Of the infected subjects, two groups
282 were considered: subjects who used ivermectin prophylactically (treated group) and
283 subjects who did not use ivermectin prophylactically (untreated group). Any missing data
284 from patients were actively searched by the investigators, via phone or in person. Since
285 this is a citywide program, all recorded data must have matched the exact number of
286 COVID-19 cases and deaths of the city. This strict interval avoids differences in terms of
287 periods of exposure.

288

289 Due to the uncertainty of reinfection with COVID-19, subjects with a history of
290 previous COVID-19 did not participate in the program although they were still permitted
291 to use ivermectin prophylactically. Limiting parameters of the government system
292 allowed the recording of a first episode of COVID-19 infection only. **Subjects below 18**
293 **years old and subjects with diagnosis of COVID-19 before July 7, 2020 were**
294 **excluded from all datasets and analysis.**

295

296 **From the registry of the city population (223,128 inhabitants), subjects below**
297 **18 years old (61,583 subjects) were removed. Of the 161,545 subjects above 18 y/o**
298 **from the city of Itajaí, we removed the 1,984 COVID-19 cases that occurred before**
299 **July 7, 2020, among subjects above 18 y/o, remaining 159,561 subjects. Subjects**
300 **above 18 y/o were considered those who were born before June 30, 2002.**

301

302 **A total of 147,223 subjects participated in the program of ivermectin**
303 **prophylaxis used for COVID-19. Of these, 24,304 subjects were below 18 y/o. Of the**
304 **122,919 ivermectin users above 18 y/o, 8,346 were from other cities, and 728 had**
305 **COVID-19 before July 7, 2020, although they used ivermectin afterwards. In total,**
306 **113,845 subjects that participated in the program remained in the dataset. The**
307 **45,716 non-participants, remaining subjects among the 159,561 subjects were**
308 **considered as the ivermectin non-users.**

309

310 Finally, city-wide COVID-19 hospitalization and mortality rates of Itajaí were
311 compared between the period before the program (before July 7, 2020) and during the
312 program between July 7, 2020 and December 2, 2020) aiming to evaluate whether a
313 program of prophylaxis with ivermectin for COVID-19 would cause a positive impact in
314 the overall numbers of the city, despite only partial adoption. Chances of dying from
315 COVID-19 in the overall population, according to use or non-use of ivermectin
316 (irrespective of COVID-19 infection) were only calculated prior to matching. Conversely,
317 mortality rate, i.e., among those who were infected by the SARS-CoV-2, was calculated
318 for both pre and post-matched cohorts. Analysis of hospitalization and mortality rates
319 before matching, mortality rate in subpopulations among ivermectin users, among
320 ivermectin non-users, and mortality rate ratios between iveremctin users and non-users in
321 subpopulations, before and after propensity score matching, and STROBE checklist are
322 presented in the **Supplement Appendix 1**.

323

324

325 *Statistical analysis*

326

327 **The full underlying data for the present analysis was analyzed by two independent**
328 **statisticians, and discrepancies evaluated by a third statistics expert.** In this outpatient
329 study of those who tested positive for SARS-CoV-2, mortality rate was evaluated
330 according to each parameter, that adjusted against other variables (for multivariate
331 regression analysis) and used for balancing and matching groups, including age intervals,
332 sex, history of smoking, prophylactic ivermectin use, T2D, asthma, COPD,
333 cardiovascular diseases and other pulmonary diseases, hypertension, current cancer (any
334 type), history of stroke and/or MI. Groups, baseline characteristics, and mortality rates
335 were presented before matching and after matching.

336

337 Before matching, a generalized linear mixed model was employed, assuming the
338 binomial distribution for the residues and including the fixed classificatory effects of each
339 of these parameters. Age intervals were adjusted for the evaluation of ivermectin
340 prophylactic use as an independent predictor of death from COVID-19. Unadjusted and
341 multivariate Poisson- adjusted probabilities to survive from COVID-19 (p-value),
342 according to each parameter were provided.

343

344 PSM was performed for mortality risk between ivermectin and non-ivermectin
345 users. COVID-19 infection rate and risk of dying were also calculated matching for
346 variables. After PSM, a second adjustment (‘double adjustment’) with multivariate linear
347 regression was performed for residual variables^{33,34}.

348

349 The statistical approach for missing data depended on the percentage of missing
350 data for each parameter. However, due to the registry system design mandating that all
351 data variables be filled to be formally included in the registry, only erroneously entered
352 (illogical) data were found. In such instances, medical record review was performed to
353 obtain the accurate data.

354

355 The program used for the analysis was the Statistical Analysis Software
356 (SAS/STAT) (SAS Institute Inc., Cary, North Carolina, USA).

357

358 **For transparency reasons, two datasets will be made public upon peer-**
359 **reviewed publication, of the 7,345 COVID-19 cases and of the 113,845 participating**
360 **subjects considered for the present analysis.**

361

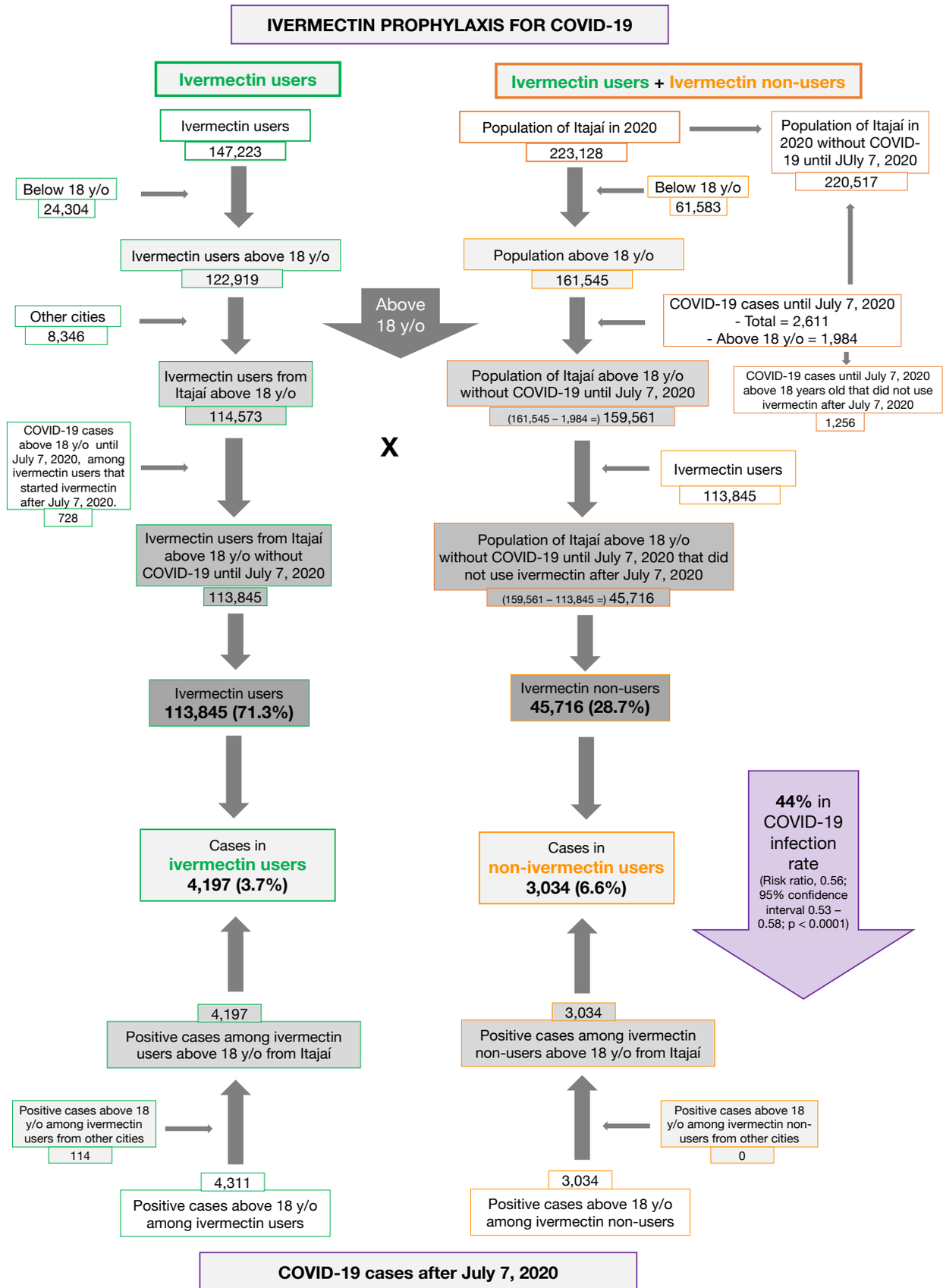
362 **Results**

363

364 **Figure 1.** Underlying data for the study on ivermectin prophylaxis used for COVID-19.

365

366



369

370 **A detailed description of the data considered for the present analysis is**
371 **illustrated in Figure 1. Of the 220,517 citizens of Itajaí without COVID-19 until July**
372 **7, 2020, 159,561 were above 18 years old. Of the 159,561 citizens above 18 y/o without**
373 **COVID-19 until July 7, 2020, 113,845 (71.3% of the population above 18 years old)**
374 **received ivermectin before being infected by COVID-19. A total of 45,716 citizens**
375 **(28.7%) did not receive or did not want to receive ivermectin during the program,**
376 **including as a prophylactic or as treatment after having COVID-19.**

377

378 **Of the 113,845 prophylaxed subjects from the city of Itajaí, 4,197 had a positive**
379 **rtPCR-SARS-CoV-2 (3.7% infection rate), while 3,034 of the 37,027 untreated subjects**
380 **had positive rtPCR-SARS-CoV-2 (6.6% infection rate), a 44% reduction in COVID-19**
381 **infection rate (Risk ratio (RR), 0.56; 95% confidence interval (95%CI), 0.53 - 0.58;**
382 **p < 0.0001). An addition of 114 subjects that used ivermectin and were infected were**
383 **originally from other cities but were registered as part of the program, in a total of**
384 **4,311 positive cases among ivermectin users. For the present analysis, the 4,311**
385 **positive cases among subjects that used ivermectin and 3,034 cases among subjects**
386 **that did not use ivermectin were considered. After PSM, two cohorts of 3,034 subjects**
387 **were created.**

388

389 **Baseline characteristics of the 7,345 subjects included prior to PSM and the**
390 **baseline characteristics of the 6,068 subjects in the matched groups are shown in Table**
391 **1. Prior to PSM, ivermectin users had a higher percentage of subjects over 50 years old**
392 **(p < 0.0001), higher prevalence of T2D (p = 0.0004), hypertension (p < 0.0001), CVD (p**
393 **= 0.03), and a higher percentage of caucasians (p = 0.004), than non-users. After PSM,**
394 **all baseline parameters were similar between groups. Figure 2 summarizes the main**
395 **findings of this study.**

396

397 **Table 1.** Baseline characteristics of subjects enrolled in study before matching and after
398 propensity score matched.

	Pre-Matching				Propensity Score Matched		
	Overall (n = 7,345)	Ivermectin users (n = 4,311)	Non- ivermectin users	<i>p-value</i>	Overall (n = 6,068)	Ivermectin users (n = 3,034)	Non- ivermectin users

			(n = 3,034)				(n = 3,034)
Age							
Mean ± SD	42.0 ± 14.7	43.5 ± 14.9	39.8 ± 14.2	< 0.0001	39.7 ± 14.0	39.67 ± 13.8	39.8 ± 14.2
< 30 y/o	1730 (23.6%)	886 (20.5%)	844 (27.8%)		1,691 (27.9%)	844 (27.9%)	847 (27.8%)
30-50 y/o	3703 (50.4%)	2121 (49.2%)	1582 (52.2%)		3,155 (52.0%)	1,573 (51.9%)	1,582 (52.1%)
> 50 y/o	1912 (26.0%)	1304 (30.3%)	608 (20.0%)		1,222 (20.1%)	614 (20.2%)	608 (20.1%)
Sex				<i>0.31</i>			
Female	3983 (54.2%)	2359 (54.7%)	1624 (53.5%)		3,231 (53.2%)	1,607 (53.0%)	1,624 (53.5%)
Male	3362 (45.8%)	1952 (45.3%)	1410 (46.5%)		2,837 (46.8%)	1,427 (47.0%)	1,410 (46.5%)
Race							
Caucasians	5437 (74.0%)	3245 (75.3%)	2192 (72.2%)	0.004	4,398 (72.5%)	2,206 (72.7%)	2,192 (72.3%)
Afro-Brazilians	209 (2.8%)	109 (2.5%)	100 (3.3%)	0.052	193 (3.2%)	93 (3.1%)	100 (3.3%)
Mixed	1583 (22.6%)	901 (20.9%)	682 (22.5%)	<i>0.10</i>	1,364 (22.5%)	93 (3.1%)	100 (3.3%)
Asian-Brazilians	116 (1.6%)	56 (1.3%)	60 (2.0%)	0.023	113 (1.9%)	53 (1.8%)	60 (2.0%)
Type 2 diabetes				0.0004			
Yes	214 (2.9%)	151 (3.5%)	63 (2.1%)		141 (2.3%)	78 (2.6%)	63 (2.1%)
No	7131 (97.1%)	4160 (96.5%)	2971 (97.9%)		5,927 (97.7%)	2,956 (97.4%)	2,971 (97.9%)
Asthma				0.067			
Yes	26 (0.3%)	20 (0.5%)	6 (0.2%)		21 (0.3%)	15 (0.5%)	6 (0.2%)
No	7319 (99.7%)	4291 (99.5%)	3028 (99.8%)		6,047 (99.7%)	3,019 (99.5%)	3,028 (99.8%)
COPD				<i>0.72</i>			
Yes	13 (0.2%)	7 (0.2%)	6 (0.2%)		12 (0.2%)	6 (0.2%)	6 (0.2%)
No	7332 (99.8%)	4304 (99.8%)	3028 (99.8%)		6,056 (99.8%)	3,028 (99.8%)	3,028 (99.8%)
Hypertension				< 0.0001			
Yes	528 (7.2%)	362 (8.4%)	166 (5.5%)		343 (5.6%)	177 (5.8%)	166 (5.5%)
No	6817 (92.8%)	3949 (91.6%)	2868 (94.5%)		5,725 (94.4%)	2,857 (94.2%)	2,868 (94.5%)
CVD				0.03			
Yes	56 (0.8%)	41 (1.0%)	15 (0.5%)		32 (0.5%)	17 (0.6%)	15 (0.5%)
No	7289 (99.2%)	4270 (99.0%)	3019 (99.5%)		6,036 (99.5%)	3,017 (99.4%)	3,019 (99.5%)
Other pulmonary diseases				<i>0.53</i>			
Yes	15 (0.2%)	10 (0.2%)	5 (0.2%)		9 (0.1%)	4 (0.1%)	5 (0.1%)
No	7330 (99.8%)	4301 (99.8%)	3029 (99.8%)		6,059 (99.9%)	3,030 (99.9%)	3,029 (99.9%)
Cancer (any type)				<i>0.66</i>			

Yes	32 (0.4%)	20 (0.5%)	12 (0.4%)		22 (0.4%)	10 (0.3%)	12 (0.4%)
No	7313 (99.6%)	4291 (99.5%)	3023 (99.6%)		6,046 (99.6%)	3,024 (99.7%)	3,022 (99.6%)
Current smoking				0.76			
Yes	110 (1.5%)	63 (1.5%)	47 (1.5%)		95 (1.6%)	48 (1.6%)	47 (1.6%)
No	7235 (98.5%)	4248 (98.5%)	2987 (98.5%)		5,973 (98.4%)	2,986 (98.4%)	2,987 (98.4%)
History of MI				0.26			
Yes	15 (0.2%)	11 (0.3%)	4 (0.1%)		8 (0.1%)	4 (0.1%)	4 (0.1%)
No	7330 (99.8%)	4300 (99.7%)	3030 (99.9%)		6,060 (99.9%)	3,030 (99.9%)	3,030 (99.9%)
History of stroke				0.56			
Yes	21 (0.3%)	11 (0.3%)	10 (0.3%)		21 (0.4%)	11 (0.4%)	10 (0.3%)
No	7324 (99.7%)	4300 (99.7%)	3024 (99.7%)		6,047 (99.6%)	3,023 (99.6%)	3,024 (99.7%)

399 COPD = chronic obstructive pulmonary disease; CVD = cardiovascular disease; MI = myocardial infarction; SD = standard deviation
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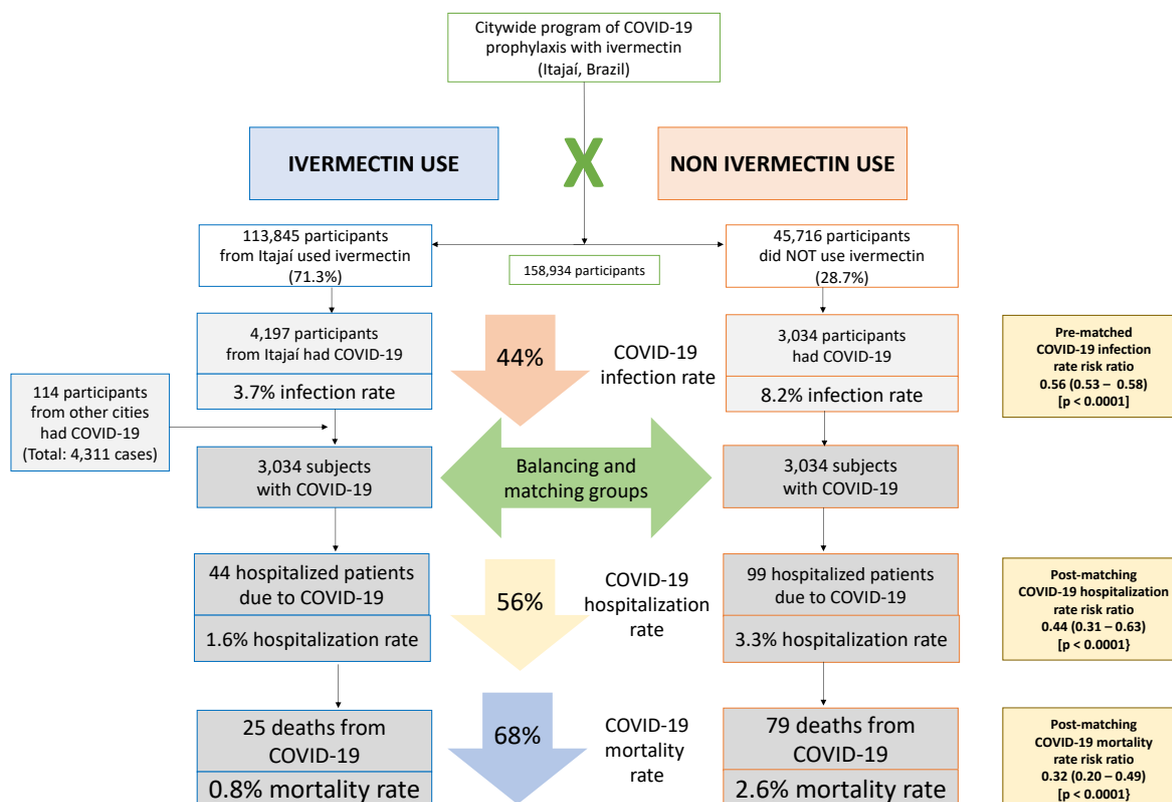
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Figure 2. Summary of the findings.

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412 *Hospitalization and mortality rates in ivermectin users and ivermectin non-users in*
413 *propensity score matched analysis*

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415 As described in **Table 2**, after employing PSM, of the 6,068 subjects (3,034 in each
416 group), there were 44 hospitalizations among ivermectin users (1.6% hospitalization rate)
417 and 99 hospitalizations (3.3% hospitalization rate) among ivermectin non-users, a 56%
418 reduction in hospitalization rate (RR, 0.44; 95%CI, 0.31 – 0.63). When adjustment for
419 variables was employed, reduction in hospitalization rate was 67% (RR, 0.33; 95%CI 0.23
420 – 0.66; p < 0.0001).

421

422 There were 25 deaths among ivermectin users (0.8% mortality rate) and 79 deaths
423 among non-ivermectin users (2.6% mortality rate), a 68% reduction in mortality rate (RR,
424 0.32; 95%CI 0.20 – 0.49). When PSM was adjusted, reduction in mortality rate was 70%
425 (RR, 0.30; 95%CI 0.19 – 0.46; p < 0.0001).

426

427 **Table 2.** Propensity score matched hospitalization and mortality rate among ivermectin users and
428 non-users.

		Overall	IVM users	Non-IVM users	PSM mortality risk ratio (95%CI) and p-value [p]	Adjusted PSM mortality risk ratio (95%CI) and p-value [p]
COVID-19 infection	Infected population (n)	6,068	3,034	3,034	-	-
COVID-19 hospitalization	Hospitalization due to COVID-19	143	44	99	-	-
	Hospitalization rate* (in case of COVID-19) (%)	2.3%	1.6%	3.3%	0.44 (0.31 – 0.63) [< 0.0001]	0.33 (0.23 – 0.46) [< 0.0001]
COVID-19 death	COVID-19 deaths (n)**	104	25	79	-	-
	Mortality rate (among infected subjects) (%)	1.7%	0.8%	2.6%	0.32 (0.20 – 0.49) [< 0.0001]	0.30 (0.19 – 0.46) [< 0.0001]

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IVM = ivermectin; PSM = propensity score matching; CI = confidence interval; *Only subjects hospitalized in public hospitals; **All deaths, including from public and private hospitals, and in-home.

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Determinants of COVID-19 mortality through propensity score matched analysis

Table 3 describes resulting risk factors for COVID-19 death amongst the overall population through PSM analysis. Risk factors for mortality in COVID-19 included aging ($p < 0.0001$), male sex ($p = 0.015$), T2D ($p < 0.0001$), hypertension ($p < 0.0001$), asthma ($p = 0.011$), COPD ($p < 0.0001$), other pulmonary diseases ($p = 0.048$), history of MI ($p = 0.034$) and history of stroke ($p < 0.0001$). To detect independent risk factors, post-PSM adjustment for variables showed that ivermectin ($p < 0.0001$; 70% reduction in mortality risk) and female sex ($p = 0.022$; 38% reduction in mortality risk) were protectors, whereas T2D ($p = 0.041$; 79% increase in mortality risk), hypertension ($p = 0.008$; 98% increase in mortality risk), and, marginally, other pulmonary diseases ($p = 0.061$; 468% increase in mortality risk) and history of stroke ($p = 0.054$; 97% increase in mortality risk) were identified as independent risk factors.

Table 3. Propensity score matched COVID-19 mortality rate according to each characteristic, in overall population, ivermectin users, and non-users.

Propensity Score Matched Groups				
Variable	Overall (n = 6,068)	Death (%)	Unadjusted COVID-19 mortality risk ratio and p-value [p]	Multivariate adjusted COVID-19 mortality risk ratio and p-value [p]
Ivermectin use - n (%)			0.32 (0.20 – 0.49) [< 0.0001]	0.30 (0.19 – 0.46) [< 0.0001]
Yes	3,034	25 (0.8%)		
No	3,034	79 (2.6%)		
Age - n (%)			[< 0.0001]	[< 0.0001]
< 30 y/o	1,691	1 (0.1%)		
30-50 y/o	3,155	12 (0.4%)		
> 50 y/o	1,222	91 (7.4%)		
Sex- n (%)			0.62	0.64

			(0.42 – 0.91) [0.015]	(0.44 – 0.93) [0.022]
Female	3,231	43 (1.3%)		
Male	2,837	61 (2.2%)		
Race - n (%)			[0.24]	[0.44]
Caucasians	4,398	79 (1.8%)		
Afro-Brazilians	193	6 (3.1%)		
Mixed	1,364	17 (1.3%)		
Asian-Brazilians	113	2 (1.9%)		
Type 2 diabetes - n (%)			10.0 (6.32-15.8) [< 0.0001]	1.79 (1.03 – 3.12) [0.041]
Yes	141	20 (14.2%)		
No	5,927	84 (1.4%)		
Hypertension - n (%)			8.83 (5.99 – 13.0) [< 0.0001]	1.98 (1.19 – 3.30) [0.008]
Yes	343	36 (10.5%)		
No	5,725	68 (1.2%)		
Asthma - n (%)			5.64 (1.49 – 21.4) [0.011]	1.74 (0.52 – 5.81) [0.36]
Yes	21	2 (9.5%)		
No	6,047	102 (1.7%)		
COPD - n (%)			15.0 (5.52 – 40.7) [< 0.0001]	1.71 (0.68 – 4.31) [0.25]
Yes	12	3 (25.0%)		
No	6,056	101 (1.7%)		
Cardiovascular diseases - n (%)			7.54 (2.96 – 19.3) [< 0.0001]	1.22 (0.44 – 3.37) [0.70]
Yes	32	4 (12.5%)		
No	6,036	100 (1.7%)		
Other pulmonary diseases - n (%)			6.54 (1.02 – 41.9) [0.048]	5.68 (0.92 – 35.0) [0.061]
Yes	9	1 (11.1%)		
No	6,059	103 (1.7%)		
Cancer (any type) - n (%)			2.67 (0.39 – 18.3) [0.32]	1.97 (0.30 – 12.9) [0.48]
Yes	22	1 (4.6%)		

No	6,046	103 (1.7%)		
Current smoking - n (%)			1.23 (0.31 – 4.92) [0.77]	0.36 (0.08 – 1.70) [0.20]
Yes	95	2 (2.1%)		
No	5,973	102 (1.7%)		
History of MI - n (%)			7.35 (1.16 – 46.5) [0.034]	1.91 (0.17 – 21.6) [0.60]
Yes	8	1 (12.5%)		
No	6,060	103 (1.7%)		
History of stroke - n (%)			17.6 (8.72 – 35.7) [< 0.0001]	1.97 (0.99 – 3.92) [0.054]
Yes	21	6 (28.6%)		
No	6,047	98 (1.6%)		

455 COPD = Chronic obstructive pulmonary disease; CVD = cardiovascular disease; MI = myocardial infarction

456

457 In a comparison of city-wide COVID-19 hospitalization rates prior to and during
 458 the program, COVID-19 mortality decreased from 6.8% before the program with
 459 prophylactic use of ivermectin, to 1.8% after its beginning (RR, 0.27; 95%CO, 0.21 –
 460 0.33; $p < 0.0001$), and in COVID-19 mortality rate, from 3.4% to 1.4% (RR, 0.41; 95%CI
 461 0.31 – 0.55; $p < 0.0001$). (Table 4).

462

463 **Table 4.** Hospitalization and mortality rates registered in the city of Itajaí, Brazil, before
 464 versus after the beginning of the citywide program with ivermectin use as prophylaxis for COVID-
 465 19, independent of the ivermectin use status.

	Overall	Until July 30th	After July 30th	Relative risk ratio (95%CI)	<i>p-value</i>
Infected COVID-19 population (n)	9956	2663	7293	-	-
Infected non-hospitalized COVID-19 population (n)	9641	2481	7160	-	-
Hospitalized COVID-19 population (n)	315	182	133	-	-
COVID-19 hospitalization rate COVID-19 (%)	3.2%	6.8%	1.8%	0.27 (0.21 – 0.33)	<0.0001
Overall number of COVID-19 deaths	192	90	102	-	-
Overall mortality rate (%)	1.9%	3.4%	1.4%	0.41 (0.31 – 0.55)	<0.0001

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495 **Discussion**

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This prospective, citywide COVID-19 ivermectin prophylaxis program resulted in significant reductions of COVID-19 infections, hospitalizations, and deaths. The ivermectin non-users were two times more likely to die from COVID-19 than ivermectin users in the overall population analysis. **Since groups were compared for the exposure during the same period, in a parallel manner, changes in transmission rates would affect ivermectin users and non-users equally.**

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505

The city of Itajai, in the state of Santa Catarina, Brazil, started a citywide program of prophylaxis with ivermectin in July 2020 as part of several initiatives to reduce the burden

506 of COVID-19. ivermectin was used, based on the existing literature at that time and on
507 the virtual absence of risks. The National Health System (Sistema Único de Saúde – SUS)
508 that functions as a full healthcare support to the entire population allowed the city to
509 establish a non-restricted population program. This program included a support structure
510 consisting of a large outpatient clinic located at the Convention Center of Itajaí. This
511 outpatient clinic became the main locale of assistance for COVID-19 patients, supported
512 by multiple public facilities where general practitioners regularly saw patients.

513

514 The use of ivermectin was optional unless contraindicated, and given upon
515 medical discretion. A structured medical-based program with a medical visit and
516 evaluation of basic demographic characteristics and comorbidities offered ivermectin as
517 an optional prophylaxis to those who agreed to participate in this preventive treatment
518 program. Health status was assessed and data was entered prospectively throughout the
519 period of the program, in a fully digitized system provided by the national health system
520 (SUS). Since the system existed prior to the pandemic, a significant number of the
521 population were already registered with their health information, including past and
522 current diseases, use of medications and other characteristics. The adaptations made to
523 the SUS for the pandemic preparedness, prior to the initiation of this ivermectin outpatient
524 program, allowed a structured, well-organized collection of the data that monitored any
525 missing values, reinforcing the reliability of the results.

526

527 An important conservative bias was present. Major risk factors for severe COVID-
528 19 and mortality due to COVID-19, including aging, diabetes, and hypertension, were
529 more present among ivermectin users, which may have underestimated the benefits
530 measured. Ivermectin was demonstrated to be particularly effective in subjects above 49
531 years old in terms of reduction of absolute risk, which corresponds to the group at the
532 highest risk for COVID-19. This allows the understanding that prophylactic use of
533 ivermectin can be particularly impactful in older subjects. In addition, ivermectin seemed
534 to reduce the exceeding risk of hypertension, T2D, and other diseases.

535

536 In accordance with the literature, subjects with higher age, diabetes and males
537 were less likely to survive ($p < 0.05$ for all), only aging remained as an independent risk
538 factor after PSM ($p < 0.0001$). However, prophylactic ivermectin use appears to mitigate

539 the additional risk of COVID-19 death due to T2D, hypertension, and cardiovascular
540 diseases.

541

542 The narrative that using preventive & early treatment therapies will have people
543 relax their caution of remaining socially & physically distanced to allow more COVID-
544 19 related infections is not supported here. This study data demonstrates that the use of
545 preventive ivermectin significantly lowers the infection rate, and benefits outweigh the
546 supposed increased risk of changes in social behaviours. Hence, we can speculate that the
547 prophylactic use of ivermectin could play an important role in the reduction of the
548 pandemic burden.

549

550 Even after adjustments to measure the most relevant variables that could influence
551 COVID-19 related outcomes, including age, sex, comorbidities, and habits, aiming to
552 avoid overestimation of the effects of ivermectin and to resemble a randomized clinical
553 trial, prophylactic ivermectin proved to be protective for the overall population, with a
554 reduction of 46% in death rate and $p < 0.0001$ after employment of PSM.

555

556 The protection provided by ivermectin when used prophylactically for COVID-
557 19 may have reflected in the reduction in COVID-19 hospitalization and mortality rates
558 observed in a populational level. Compared to before the beginning of the program,
559 COVID-19 hospitalization and mortality rates were reduced by 73% and 59%,
560 respectively ($p < 0.0001$ for both). These reductions were obtained when overall
561 population of the city of Itajaí, as well as overall number of COVID-19 cases,
562 hospitalizations, and deaths, were considered, irrespective of the percentage of patients
563 using ivermectin prophylactically. **There were no changes in SARS-CoV-2 variants,
564 infectivity and pathogenicity between before and during the program.**

565

566 When compared to all other major cities in the State of Santa Catarina, where
567 Itajaí is located, differences in COVID-19 mortality rate between before July 7, 2020 and
568 between July 7, 2020 and December 21, 2020, Itajaí is ranked number one, and far from
569 the second place³⁵. These results indicate that medical-based optional prescription,
570 citywide covered ivermectin can have a positive impact in the healthcare system.
571 **However, the present results do not provide sufficient support for the hypothesis
572 that ivermectin could be an alternative to COVID-19 vaccines.**

573

574 Due to the large number of participants, this citywide program was unable to
575 supervise whether ivermectin users were using ivermectin regularly, **although the**
576 **accumulated number of ivermectin tablets was strictly controlled.** This occurred to
577 be a potential another conservative bias, since the effects of ivermectin on prophylaxis
578 could be underestimated due to adherence to the recommended frequency of ivermectin
579 use.

580

581 While ivermectin is a multi-target drug³⁶, its maximum benefits occur when it's
582 present at minimum concentration in a wide range of sites to inhibit multiple metabolic
583 and inflammatory pathways. However, although the dose of ivermectin employed in the
584 program was smaller than the minimum to reach the concentration required to act in these
585 multiple sites, the reduction in infection, mortality, and death rates in the infected group
586 that used ivermectin prophylactically was surprisingly lower. Long-term or accumulated
587 ivermectin could also play a critical role for its long-term protection against COVID-19.

588

589 *Limitations*

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591 Being a prospective observational study which allowed subjects to self select
592 between treatment vs. non-treatment instead of relying on randomization, important
593 confounders may have been differentially present which could otherwise explain the
594 differences observed. Given that the benefits measured occurred despite negative risk
595 factors being more present in the treatment group, this suggests the benefits are likely
596 accurate and unbiased. Further, studies relying on PSM techniques have been shown
597 to consistently agree with those employing randomization^{37,38}, again supporting the
598 likelihood the benefits measured are accurate, The prevailing type of SARS-CoV-2 in the
599 city was unknown due to the lack of genotyping surveillance during the period of the
600 program. Whether the prophylaxis proposed in this program would be as effective in other
601 SARS-CoV-2 variants is unclear. Also, there was not a strict control of whether infected
602 subjects used any specific drug in case of COVID-19 infection, this allows the possibility
603 that the differences may be explained by differences in the use of ivermectin or other
604 medications as treatment.

605

606 *Final discussion*

607

608 In this city-wide ivermectin prophylaxis program, a large, statistically significant
609 decrease in mortality rate was observed after the program began among the entire
610 population of city residents. When comparing subjects that used ivermectin regularly,
611 non-users were two times more likely to die from COVID-19 while ivermectin users were
612 7% less likely to be infected with SARS-CoV-2 ($p = 0.003$).

613

614 Although this study is not a randomized, double-blind, placebo-controlled clinical
615 trial, the data was prospectively collected and resulted in a massive study sample that
616 allowed adjustment for numerous confounding factors, thus strengthening the findings of
617 the present study.

618

619 Due to the well-established, long-term safety profile of ivermectin, with rare
620 adverse effects, the absence of proven therapeutic options to prevent death caused by
621 COVID-19, and while effectiveness of vaccines in real-life all-cause mortality analyses
622 lacks, we recommend that ivermectin could be considered as a preventive strategy, in
623 particular for those at higher risk of complications from COVID-19 or at higher risk of
624 contracting the illness, **not as a substitute for COVID-19 vaccines, but as an additional
625 tool, particularly during periods of high transmission rates.**

626

627

628 **Conclusion**

629

630 In a city-wide ivermectin program with prophylactic, optional ivermectin use for COVID-
631 19, ivermectin was associated with significantly reduced COVID-19 infection,
632 hospitalization, and death rates from COVID-19.

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638 **Statements**

639

640 *Conflict of Interest*

641

642 The authors declare no conflict of interest regarding the drug, ivermectin, and potential
643 commercial benefits of the expansion of its use for COVID-19, or any other related gains.
644 Dr Lucy Kerr received funding from Vitamedic, that manufactures ivermectin, unrelated
645 to this study. Dr. Flavio A. Cadegiani was contracted by Vitamedic for consulting services
646 unrelated to this study, and donated the full budget for COVID-19 patient care and
647 research. Other authors have no conflicts of interest.

648

649 *Data availability statement*

650

651 Dataset is available under reasonable request by institutions and organizations.

652

653 *Author contributions*

654

655 Lucy Kerr designed the study. Washington Luiz Olivato Assagra and Fernando Carlos
656 Proença developed the computer program, compiled and ran the data. Raysildo Barbosa
657 Lôbo, Fernando Baldi, Flavio A. Cadegiani and Juan J. Chamie designed and performed
658 the statistical analyses. Lucy Kerr, Flavio A. Cadegiani, Fernando Baldi and Pierre Kory
659 performed the analyses and interpretation of clinical and demographic data generated by
660 the statistical analysis. Fernando Carlos Proença was responsible for the medical
661 surveillance, subjects follow-up and other aspects related to the program administration
662 of the present analysis. Raysildo Barbosa Lôbo and Lucy Kerr were responsible for
663 resources, supervision and project administration related to the analyses. Pierre Kory,
664 Juan J Chamie and Jennifer Hibberd reviewed the data and the manuscript. All authors
665 contributed to the writing of the original draft and final reviewed manuscript. All authors
666 have read and approved the manuscript.

667

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669

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671 sites where the citywide programs were conducted. No other funding sources were
672 obtained.

673

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675

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682 led to the present analysis.

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822 **Table list**

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835 **Figure list**

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837 Figure 1. Underlying data for the study on ivermectin prophylaxis used for COVID-19.

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