

# Computerized monitoring of COVID-19 trials, studies and registries in ClinicalTrials.gov registry

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Clinical trial registries can provide important information about relevant studies for a given condition to other researchers and the public. We developed a computerized informatics based approach to provide an overview and analysis of COVID-19 studies registered on ClinicalTrials.gov registry. Using the perspective of analyzing active or completed COVID-19 studies, we identified 401 interventional clinical trials, 287 observational studies and 64 registries. We analyzed features of each study type separately such as location, design, interventions and update history. Our results show that the United States had the most COVID-19 interventional trials, France had the most COVID-19 observational studies and France and the United States tied for the most COVID-19 registries on ClinicalTrials.gov. The majority of studies in all three study types had a single study site. For update history 'Study Status' is the most updated information and we found that studies located in Canada (2.70 updates per study) and the United States (1.76 updates per study) update their studies more often than studies in any other country. Using normalization and mapping techniques, we identified Hydroxychloroquine (92 studies) as the most common drug intervention, while convalescent plasma (20 studies) is the most common biological intervention. The primary purpose of most interventional trials is for treatment with 298 studies (74.3%). For COVID-19 registries we found the most common proposed follow-up time is one year (15 studies). Of specific importance and interest is COVID-19 vaccine trials, of which 12 were identified. Our informatics based approach allows for constant monitoring and updating as well as multiple applications to other conditions and interests.

1 **Computerized monitoring of COVID-19 Trials, Studies and**  
2 **Registries in ClinicalTrials.gov registry**

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## 42 **Abstract**

43

44 Clinical trial registries can provide important information about relevant studies for a given  
45 condition to other researchers and the public. We developed a computerized informatics based  
46 approach to provide an overview and analysis of COVID-19 studies registered on  
47 ClinicalTrials.gov registry. Using the perspective of analyzing active or completed COVID-19  
48 studies, we identified 401 interventional clinical trials, 287 observational studies and 64  
49 registries. We analyzed features of each study type separately such as location, design,  
50 interventions and update history. Our results show that the United States had the most COVID-  
51 19 interventional trials, France had the most COVID-19 observational studies and France and the  
52 United States tied for the most COVID-19 registries on ClinicalTrials.gov. The majority of  
53 studies in all three study types had a single study site. For update history ‘Study Status’ is the  
54 most updated information and we found that studies located in Canada (2.70 updates per study)  
55 and the United States (1.76 updates per study) update their studies more often than studies in any  
56 other country. Using normalization and mapping techniques, we identified Hydroxychloroquine  
57 (92 studies) as the most common drug intervention, while convalescent plasma (20 studies) is the  
58 most common biological intervention. The primary purpose of most interventional trials is for  
59 treatment with 298 studies (74.3%). For COVID-19 registries we found the most common  
60 proposed follow-up time is one year (15 studies). Of specific importance and interest is COVID-  
61 19 vaccine trials, of which 12 were identified. Our informatics based approach allows for  
62 constant monitoring and updating as well as multiple applications to other conditions and  
63 interests.

64

## 65 **1 Introduction**

66

67 The purpose of clinical trial registries, among others, is to inform the research community about  
68 currently ongoing studies. A registry can also be a sole source of study results for thousands of  
69 studies that would otherwise not publish a result article in a journal.(Zarin et al., 2019) For the  
70 current effort to address the COVID-19 epidemic, this function of registries is of great value. The  
71 fields of clinical informatics and clinical research informatics have an important role to play in  
72 fighting the epidemic.(Moore et al., 2020)

73

74 We focus on a single registry, ClinicalTrials.gov (CTG), and analyze COVID-19 registered  
75 studies. We chose to focus on CTG because it collects a rich set of metadata, supports record  
76 updates,(Fleminger & Goldacre, 2018) allows for basic summary results deposition and studies  
77 in CTG make up a large volume of all studies tracked by the World Health Organization  
78 registry.(Huser & Cimino, 2013) The goal of our study is to demonstrate how automated and  
79 clinical research informatics(Embi & Payne, 2009) methods can be used to analyze a set of  
80 closely related studies, as well as to use general statistical principles to analyze and understand  
81 key metrics for studies on a given condition or in a specific clinical domain. The computer code  
82 written in R language is open source and available at the project repository.(“regCOVID Project  
83 Repository,” 2020) Our project differs from past published analyses of COVID-19 clinical  
84 studies(Rosa & Santos, 2020; Fragkou et al., 2020; Checcucci et al., 2020) by not using manual  
85 review of study records and instead relying fully on study metadata recorded in the registry.

86 Unlike manual approaches, our approach of using automation to monitor COVID-19 studies  
87 allows for quick and efficient, continuous monitoring of the state of COVID-19 research.  
88 Automated queries can provide an instant overview of COVID-19 research. They also allow for  
89 computing and visualizing current COVID-19 research trends. Furthermore, the informatics  
90 approach we use improves the capability for effective modulation and allows individuals to  
91 specifically target the desired information wanted to inform their research decisions (such as on  
92 clinical guidelines, and the use of certain interventions). With the gradual publication of COVID-  
93 19 study results, this approach also allows for automated detection of registry deposition of basic  
94 summary results and the publication of a study result journal article that is clearly tied to a  
95 registered COVID-19 trial, study or registry.  
96

## 97 **2 Materials & Methods**

98  
99 Our study had two aspects. The first aspect was assuming a journalist perspective and the  
100 motivation was whether CTG registry can provide a useful overview of COVID-19 studies  
101 without any manual curation. We wanted to demonstrate on a COVID-19 use case whether  
102 existing study metadata currently collected by CTG registry are accurate and adequate for a  
103 journalist interested in a registry-based picture of COVID-19 research. The methods and results  
104 sections address mostly this first aspect.  
105

106 The second aspect was assuming an informatics or data science perspective and the motivation  
107 for this was to apply additional rules, data transformations and heuristics to CTG metadata that  
108 could characterize the quality of the registry data and possibly narrow the list of all CTG studies  
109 to a smaller set. The critical component of this second informatics aspect was a vision to  
110 generalize the fully computerized single disease report to all diseases. The discussion section of  
111 this article addresses this second aspect.  
112

### 113 **2.1 Set of analyzed studies**

114  
115 We used the Aggregate Analysis of ClinicalTrials.gov (AACT), which is a relational database  
116 version of CTG data that is created by parsing the XML (Extensible Markup Language)  
117 representation of each study.(AACT Team, 2020) It is published and maintained by Duke  
118 University. AACT data is typically four days behind CTG in terms of content or changes, which  
119 we deemed as acceptable. We performed separate analyses of COVID-19 studies based on their  
120 CTG study type of (1) interventional trials, (2) observational studies, and (3) registry-based  
121 studies (we use the term registries).(CTG Team, 2020a)  
122

123 We designed several inclusion criteria to focus only on COVID-19 studies in scope for our  
124 analysis. This consisted of first, creating a search strategy based on title or study keywords. We  
125 also looked at CTG study metadata to select studies with fields that we considered relevant based  
126 on the triggering of quality measures and the connection to the regulatory process.  
127

128 In terms of keyword and title search strategy, we evaluated three search approaches. The first  
129 method involved a search for the presence of keywords in the official title of the study. The  
130 keywords used were, 'covid', 'sars-cov', '2019-ncov', and 'coronavirus'. The second method  
131 searched for the same keywords in the free text condition field. The third method found studies

132 that had a Medical Subject Heading (MeSH) term for the study of ‘coronavirus infections’. We  
133 limited our search for each method to only include studies first submitted after 27 December  
134 2019 (the date of the official report from Wuhan hospital to the local center for disease control  
135 and prevention). For later analysis, we present data for the first search method (with results for  
136 all three methods being available on the study repository at [https://github.com/lhncbc/r-snippets-  
137 bmi/tree/master/regCOVID](https://github.com/lhncbc/r-snippets-bmi/tree/master/regCOVID)). The data presented below reflects the search performed on 11 May  
138 2020. The results of the search methods were validated as appropriate COVID-19 studies via the  
139 manual review of the titles of a subset of the search results.

140

141 In terms of study inclusion criteria based on structured CTG’s study metadata fields we used  
142 study status. Study status reflects the progress of the study from ‘not yet recruiting’ to  
143 ‘recruiting’ to ‘completed’ (or other statuses such as ‘terminated’ or ‘suspended’).(CTG Team,  
144 2020a) We elected to limit the scope of our analysis to reflect currently ongoing or completed  
145 COVID-19 studies. Given this assumption, we excluded studies with ‘not yet recruiting’ study  
146 status. Under existing quality assurance rules used by CTG, studies in status ‘not yet recruiting’  
147 do not have to provide the study location (in terms of country) and we considered study country  
148 to be essential study metadata. Given our chosen analytical perspective of currently ongoing or  
149 completed COVID-19 studies, we excluded studies with an *unusual completion* status of  
150 ‘terminated’, ‘suspended’, or ‘withdrawn’. However, we provide some results for unusually  
151 completed COVID-19 studies because when combined with the ‘Reason for termination’ field,  
152 such studies may provide important insights.

153

## 154 **2.2 Analysis**

155

156 For all study types, we analyzed a set of study metadata described below. Study metadata  
157 specific to a given study type (e.g., those collected only for observational studies or registries)  
158 are described in subsequent sections.

159

### 160 **2.2.1 Number of studies over time**

161

162 Using the date when the study was first registered on CTG, we counted the number of total  
163 studies for each study type on a given date. We created a plot showing the temporal trend over  
164 time.

165

### 166 **2.2.2 Study sites and country**

167

168 Despite the existence of national registries, CTG registry contains studies from many countries.  
169 For example, as of 2 June 2020, 61% of recruiting studies were located solely outside the US  
170 (according to CTG’s overview page(“Trends, Charts, and Maps - ClinicalTrials.gov”)).

171 Additional incentive for registration on CTG are FDA rules that require studies submitted in  
172 support of new drug applications to FDA to be registered at CTG. Similar requirements of CTG  
173 registration also exist from many study funders such as NIH, and journals such as International  
174 Committee of Medical Journal Editors (ICMJE) member journals. We analyzed the geographic  
175 data for each study by reviewing the location fields in the CTG study record and counted the  
176 number of sites and identified the country of their locations. Each study could consequently  
177 include one or multiple countries.

178

### 179 **2.2.3 Study update activity**

180

181 The clinical trial registry allows principal investigators to update the public about study  
182 completion, the final number of enrolled participants and basic summary results. High public  
183 interest in updates about COVID-19 studies was the main motivation for measuring update  
184 activity and study record recency.

185

186 We quantified the level of update activity for a study by looking at the number of updates and  
187 what fields are updated for a given study after its initial registration. We attempted to classify  
188 type of updates into technical updates (e.g., study sites changes) and updates of significant public  
189 interest (actual primary completion date or deposition of study results). We also evaluated the  
190 recency of the CTG study record by evaluating the number of days between the last update and  
191 the current date. Study update data is not available via AACT or the CTG Application Protocol  
192 Interface (API).(CTG Team, 2020b) However, information about study updates is available via  
193 the CTG website. To obtain this information we wrote an R script to scrape the data into a  
194 computable form.

195

196 We also evaluated the level of update activity for studies based in each country and found studies  
197 from which countries were more active in updating CTG study records.

198

### 199 **2.2.4 Study design**

200

201 We analyzed CTG metadata pertaining to study design to classify studies. One feature analyzed  
202 for all study types was study enrollment (number of participants). CTG allows the reporting of  
203 estimated and actual enrollment into the trial. Study record managers can use this mechanism to  
204 publicly post updates about the number of enrolled participants.

205

## 206 **2.3 Study type specific analysis**

207

### 208 **2.3.1 Interventional trials**

209

#### 210 *2.3.1.1 Interventional trial specific features*

211

212 We analyzed certain features which are specific to interventional trials alone. This includes  
213 phase, primary purpose, and number and type of arms.

214

#### 215 *2.3.1.2 Intervention type*

216

217 For interventional trials specifically we analyzed the intervention types for the collection of  
218 COVID-19 studies. To do this we used CTG's metadata field of intervention type. CTG  
219 classifies each intervention as drug, device, biological, procedure, radiation, genetic, dietary  
220 supplement, behavioral, combination product, diagnostic test, and other. We counted the number  
221 of studies associated with each intervention type. Each study could include one or multiple

222 intervention types. If multiple intervention types were included, we counted each study based on  
223 the combination of intervention types associated with the study. For example, NCT04334512, a  
224 ‘Study of Quintuple Therapy to Treat COVID-19 Infection’, included two interventions of type  
225 ‘Drug’ (Hydroxychloroquine and Azithromycin) and three interventions of type ‘Dietary  
226 supplement’ (Vitamin C, Vitamin D and Zinc). This study was counted exactly once under a  
227 composite intervention type that consisted of the alphabetically sorted combination of two types:  
228 ‘dietary supplement | drug’, with the ‘|’ representing the term ‘and’.

229

230 Our analysis of intervention types and names revealed that placebo as an intervention name is  
231 often used and captured under type ‘Drug’ or ‘Biological’. CTG type classification does not  
232 include placebo as a separate intervention type, however, we decided to experimentally create it  
233 and assign it based on a rule that looked for the term placebo in the intervention name.

234

### 235 2.3.1.3 Interventions

236

237 Researchers and the public are most interested to see which drugs (or other interventions) are  
238 being tested in relation to COVID-19. CTG allows study record administrators to specify  
239 intervention using free text and further assign interventions to study arms. Because of the vital  
240 importance of interventions and the correct counting of studies using the same intervention, we  
241 did implement a limited computerized method of processing free text interventions to achieve  
242 some semantic harmonization. After free text string transformations into harmonized  
243 intervention terms, we counted the number of studies that included a given intervention. We also  
244 evaluated the temporal change in the amount of studies for the most common interventions by  
245 showing the number of new studies on a weekly scale (as seen in a figure in the Results section).

246

247 From prior studies, there is an obvious need to harmonize semantically different interventions  
248 expressed as free text across different studies. (Cepeda, Lobanov & Berlin, 2013) For example,  
249 the intervention term ‘ruxolitinib’ can semantically harmonize entries of ‘Ruxolitinib Oral  
250 Tablet’ (in study NCT04334044), ‘Ruxolitinib 10 MG’ (NCT04338958) and ‘Ruxolitinib’  
251 (NCT04331665). Initial normalization involved the removal of extra white space and the  
252 conversion of each term to lower case. Representing drug dose form was out of scope so further  
253 normalization removed commonly occurring dose form terms, such as, ‘tablet’, ‘injection’, and  
254 ‘pill’.

255

256 Studies with multiple interventions were counted multiple times under each individual  
257 intervention. In some cases, the free text string for a single intervention (in the CTG data entry  
258 field) specified a combination of several interventions. Our transformation approach in such  
259 cases kept the combination as well as expanded the single entry into multiple separate  
260 interventions and counted each intervention separately. For example, NCT04334928 has an  
261 intervention that includes a combination of Emtricitabine and Tenofovir Disoproxil. In this case  
262 the study is counted once for Emtricitabine, once for Tenofovir, and once for the combination of  
263 Emtricitabine and Tenofovir. In some cases, the manual mapping reduced the term granularity  
264 and used a higher-level term.

265

#### 266 2.3.1.4 COVID-19 vaccine trials

267

268 A segment of COVID-19 interventional trials with high importance and significant public  
269 interest are vaccine trials. CTG maintains a hierarchy of intervention types but vaccine as an  
270 intervention does not have a designated intervention type and is subsumed under the intervention  
271 type 'Biological'. Because there is no special vaccine intervention type, our method for finding  
272 vaccine trials was based on a string search for the term 'vaccine' in the official title of the study.  
273 Once we created a COVID-19 vaccine trials subset, we applied on this set the same series of  
274 analyses and metrics mentioned above.

275

#### 276 2.3.2 Observational studies and registries

277

278 Observational studies and registries have metadata features that are not recorded for  
279 interventional trials. Such analyzed features were: time perspective and observational model for  
280 the set of observational studies and registries. In addition, one feature recorded and analyzed for  
281 only registries was follow up time.

282

### 283 3 Results

284

285 We developed a methodology to search and extract metadata on COVID-19 clinical studies. The  
286 database is a subset of the AACT database of ClinicalTrials.gov data. The database and result  
287 files can be found in our github repository at [https://github.com/lhncbc/r-snippets-](https://github.com/lhncbc/r-snippets-bmi/tree/master/regCOVID)  
288 [bmi/tree/master/regCOVID](https://github.com/lhncbc/r-snippets-bmi/tree/master/regCOVID). The repository includes the R code ([https://github.com/lhncbc/r-](https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid_code_for_analysis.R)  
289 [snippets-bmi/blob/master/regCOVID/regCovid\\_code\\_for\\_analysis.R](https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid_code_for_analysis.R)), with comments  
290 explaining how it works, to obtain and analyze the data, as well as all comma separated value  
291 (CSV) data files used during the analysis. The repository also includes additional result data files  
292 not included in this paper but described in the repository documentation. The repository also  
293 includes a list of descriptions for each data file ([https://lhncbc.github.io/r-snippets-](https://lhncbc.github.io/r-snippets-bmi/regCOVID/regCOVID_data_file_descript.html)  
294 [bmi/regCOVID/regCOVID\\_data\\_file\\_descript.html](https://lhncbc.github.io/r-snippets-bmi/regCOVID/regCOVID_data_file_descript.html)) for easy use. For example, the files,  
295 `regCovid_all_studies-a.csv`, `regCovid_int-a.csv`, `regCovid_obs-a.csv`, and `regCovid_registry-`  
296 `a.csv` are the lists of all studies, interventional trials, observational studies, and registries  
297 generated from search method A respectively. These files include all 64 columns from the  
298 AACT studies table, such as NCT ID, official title, start date, primary completion data, and  
299 enrollment. The description file has more than 80 entries and provides guidance and descriptions  
300 for each included file in the analysis. Also included in the repository is an example of part of the  
301 code used in the analysis ([https://github.com/lhncbc/r-snippets-](https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid_example.R)  
302 [bmi/blob/master/regCOVID/regCovid\\_example.R](https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid_example.R)) and a quick-start tutorial  
303 ([https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid\\_Tutorial.md](https://github.com/lhncbc/r-snippets-bmi/blob/master/regCOVID/regCovid_Tutorial.md)) that  
304 shows users how to easily access and use our code and load the data files into R to review our  
305 results and perform their own analysis.

306

307 While this paper includes results from the main analysis done on 11 May 2020, the repository  
308 report is updated weekly and offers up to date results.

309

#### 310 3.1 Set of analyzed studies

311

### 312 **3.1.1 Search strategy**

313

314 We found that the first search method, using the official title of the study, was the most  
315 comprehensive and included the most COVID-19 studies. The numbers listed below reflect only  
316 the search strategy and not applying the criteria based on study status. As of 11 May 2020, the  
317 first search method returned a total of 1302 studies. The second search method, based on the free  
318 text condition field, found fewer records (1165 studies). The third method based on the MeSH  
319 term, returned 328 studies. The significant difference in the studies captured in the third search  
320 strategy is likely due to the fact, that there is no specific MeSH term for COVID-19 at this point  
321 and the MeSH condition field is not required and is left blank for many studies (38.2% of studies  
322 captured in the first search method left MeSH condition term blank).

323

324 We then applied metadata inclusion criteria (studies that are active, recruiting or completed and  
325 are not expanded access). This reduced the set for the first search method to 752 studies, the set  
326 from the second search method to 680 studies, and the set from the third search method to 210  
327 studies.

328

329 This led us to select the set of COVID-19 studies generated from the first, most comprehensive  
330 search method, based on study title.

331

### 332 **3.1.2 Final study set**

333

334 In terms of completion and presence of results, 48 studies in the final set were completed at the  
335 time of this analysis. None have provided summary results to this point. It is important to note  
336 that studies are typically required to submit results within one year after the primary completion  
337 date. (“FDAAA 801 and the Final Rule - ClinicalTrials.gov”) Also, at the time of the analysis,  
338 106 studies have past their primary completion date (12 studies when using primary completion  
339 day + 30 days) declared in the latest study record and have a status that indicates the study is still  
340 ongoing. This indicates that the record is possibly not kept current. Administrators do typically  
341 have 30 days after a status change to update the record (see 42 Code of Federal Regulation  
342 [CFR] 11.64(a)(1)(ii)). (“Frequently Asked Questions - ClinicalTrials.gov”) In an extreme case,  
343 20 studies of those 106 studies have a status of ‘not yet recruiting’ and are past their primary  
344 completion date.

345

346 To understand how our metadata study inclusion criteria affects the final set, we briefly analyzed  
347 the set of studies excluded due to our study metadata criteria. The studies removed due to  
348 metadata included 516 studies that were not yet recruiting, 10 that were withdrawn, 5 that were  
349 suspended and 2 that were terminated. The reasons for termination of the 2 studies were ‘We  
350 cannot meet number of subjects as recently published similar studies’ for NCT04357535 and  
351 ‘The epidemic of COVID-19 has been controlled well in China, no eligible patients can be  
352 enrolled at present’ for NCT04257656. The interventions of the terminated studies were ACE-I  
353 (angiotensin-converting enzyme inhibitors) and ARB (angiotensin receptor blocker) for the  
354 former and Remdesivir for the latter. Our study type criteria also excluded 17 studies with a  
355 study type of ‘Expanded access’.

356

## 357 **3.2 Studies over time**

358

359 Figure 1 shows the number of registered studies over time by study type. Interventional trials are  
360 most numerous. An important regulatory consideration is that, in the US, applicable  
361 interventional clinical trials are required to register,<sup>13</sup> while registration of observational studies  
362 and registries is optional. When considering the submission date, the first interventional trial, and  
363 the first study overall, was submitted to CTG on 23 January 2020, while the first observational  
364 study was submitted on 26 January 2020 and the first registry was not submitted until 12 March  
365 2020.

366  
367 **Figure 1.**

### 369 **3.3 Analysis by study type**

#### 370 **3.3.1 Interventional trials**

371  
372 We identified a total of 401 COVID-19 interventional trials from CTG. These 401 studies  
373 included a total of 1666 interventions.  
374

##### 375 *3.3.1.1 Study sites and country*

376  
377 The majority of interventional trials had just one study site (259 studies, 64.6%%). 41 studies  
378 had two sites and 18 studies had three sites, the second and third highest study counts. The study  
379 with the most sites was NCT04292730 ('Study to Evaluate the Safety and Antiviral Activity of  
380 Remdesivir in Participants With Moderate Coronavirus Disease (COVID-19) Compared to  
381 Standard of Care Treatment') with 183 sites.

382  
383 As for country of operation, Table 1 shows the count of interventional trials by the country or  
384 countries that have at least one site that is part of the study.

385  
386 **Table 1.**

387  
388 The vast majority of studies 385 (96.0%) only included sites in a single country. Table 1 results  
389 indicate that the most common country for interventional trials was the United States with 121  
390 studies (30.2%) followed by China with 49 studies (12.2%).  
391

##### 392 *3.3.1.2 Update activity*

393  
394 We evaluated the amount of interventional trials that had updates after the study was first  
395 submitted to CTG (full update data are available in a report and as Comma Separated Value  
396 [CSV] files in the study github repository). ("regCOVID," 2020) At the time of the analysis,  
397 71.1% (285 studies) of the 401 interventional trials show the presence of at least one update  
398 since first being submitted to CTG. The study with the most updates was NCT04280705,  
399 'Adaptive COVID-19 Treatment Trial (ACTT)' with 18 updates. The most common public  
400 interest and overall feature updated for COVID-19 interventional trials was 'Study Status',  
401 which was updated 643 times including at least once by each of the 285 studies that had at least  
402 one update. Other commonly updated public interest fields include 'Recruitment Status' (212

403 updates from 199 studies) and ‘Outcome Measures’ (137 updates from 108 studies). The second  
404 most commonly updated field overall, and most common technical field, was  
405 ‘Contacts/Locations’, which was updated 393 times by 223 studies. Using 11 May 2020 as the  
406 current date, we also looked at the amount of days since last update to evaluate how current the  
407 existing CTG record is and found that the average amount of days since the last update is 20.6  
408 days for all COVID-19 interventional trials.

409

410 Table 2 shows the amount of updates by studies in each country and the ratio of the number of  
411 updates compared to the number of studies in a given country. The table is limited to countries  
412 with at least eight studies. The country with the highest update rate is Canada with 2.70 updates  
413 per study, followed by the United States with 1.76 updates per study.

414

415 **Table 2.**

416

417

### 418 3.3.1.3 *Study design and interventional trial specific features*

419

420 **Study phase and size:** Considering study phase and study size (or enrollment; number of  
421 participants), Table 3 shows the counts of studies and percentage by study phase, as well as  
422 study size indicators: 1<sup>st</sup> quartile, median, and 3<sup>rd</sup> quartile for the participants enrolled (either  
423 actual or anticipated) for the set of studies of each phase.

424

425 Table 3 shows that the phase with the most studies is N/A with 111 studies (27.7%), which  
426 represents studies of intervention type device or behavioral. The second most common phase is  
427 Phase 2 with 108 studies (26.9%). Unsurprisingly the phase with the highest enrollment is Phase  
428 3 with a median of 500 participants, while the lowest enrollment is Early Phase 1 with a median  
429 enrollment of 10 participants.

430

431

432 **Table 3.**

433

434 **Arms:** Considering number of study arms, most interventional trials have two arms (245 studies,  
435 61.1%), while 73 studies (18.2%) have just one arm.

436

437 **Primary Purpose:** Considering study primary purpose, Table 4 presents the breakdown into 8  
438 purpose categories. In 298 (74.3%) of the analyzed COVID-19 interventional trials, the primary  
439 purpose was treatment. For 41 (10.2%) the primary purpose was prevention.

440

441 **Table 4.**

442

443 **Arm Type:** CTG allows study managers to specify the type of each study arm. Each study arm is  
444 named and is classified as a specified arm type. Each study could have one or multiple arms of  
445 the same type. For example, NCT04321993, ‘Treatment of Moderate to Severe Coronavirus  
446 Disease in Hospitalized Patients’, has three arms of type ‘Experimental’ and one arm of type ‘No  
447 Intervention’. One arm in this study has patients receiving an intervention of  
448 Lopinavir/Ritonavir, the second has patients receiving an intervention of Hydroxychloroquine,

449 and the third has patients receiving an intervention of Baricitinib. This study also has a fourth arm  
450 of patients receiving no intervention.

451

452 Considering types of all COVID-19 interventional trials, we found that the most common arm  
453 type is 'Experimental', which appears 489 times. Table 5 shows the complete data for arm type  
454 in the set of 401 interventional trials.

455

456 **Table 5.**

457

458 Different diseases at different maturity of clinical research may be employing a different design,  
459 such as the inclusion of a placebo or active comparator. We calculated the placebo index, which  
460 is the percentage of interventional trials that have a placebo or sham comparator arm. Each study  
461 can have one or multiple arms that are assigned a placebo comparator. For our set of COVID-19  
462 studies, the placebo index was 28.7% (115 of 401 total trials). We also calculated the active  
463 comparator index, which is the percentage of trials with at least one active comparator arm, and  
464 found that 28.9% (116 trials) have at least one active comparator arm.

465

#### 466 3.3.1.4 *Intervention type*

467

468 Table 6 shows the count of studies by intervention type. Intervention type 'Drug' is the most  
469 common (137 studies [34.2%]). The combination of drug and placebo intervention type was the  
470 second most prevalent with 75 studies (18.7%). Biological was the next most prevalent type with  
471 32 studies (8.0%). Based on our methodology for classifying intervention types, each study can  
472 be counted only under one composite intervention type.

473

474 **Table 6.**

475

476

#### 477 3.3.1.5 *Interventions*

478

479 There were a total of 449 distinct interventions listed prior to the implementation of our  
480 normalization and mapping process. Once the interventions were mapped the amount of  
481 normalized interventions was reduced to 403. The full mapping is available at the study  
482 repository (file: `intervention_map2.xlsx`). ("regCOVID Project Repository," 2020) Table 7 shows  
483 the most common interventions used in COVID-19 interventional trials. Given our counting  
484 methodology for interventions, each study can be counted multiple times in Table 7 because  
485 combined interventions are expanded into their components as well as kept as a combination.  
486 The most common drug intervention was Hydroxychloroquine with 92 studies, followed by  
487 Azithromycin with 24 studies. The two (Hydroxychloroquine and azithromycin) appeared  
488 together four times. The most common combination intervention was Lopinavir/Ritonavir with  
489 16 studies. We also found the presence of interventions most likely listed as a comparator or a  
490 non-intervention group, rather than a specific intervention. This is seen as 99 studies have  
491 placebo listed as an intervention while another 40 studies have standard care listed.

492

493 **Table 7.**

494

495 As for non-drug interventions, the most common biological is convalescent plasma with 20  
496 studies. Other leading interventions for different types (not shown in Table 7) include oxygen  
497 supplying equipment for device with six studies and Vitamin C for dietary supplements with four  
498 studies. We also found that the same intervention can be listed as different intervention types.  
499 For example, convalescent plasma was listed for 14 studies as the intervention type biological,  
500 three times as other and 3 times as drug. We combined each intervention to count as the most  
501 commonly used intervention type when counting the intervention. For this case of convalescent  
502 plasma, that would count as 20 studies and categorize convalescent plasma as having the type  
503 biological.

504

505 **Interventions over time:** We also evaluated the temporal change for the most common  
506 interventions by analyzing the amount of new studies weekly for the most common interventions  
507 as seen in Figure 2.

508

509

510 **Figure 2.**

511

512 The plot shows how most interventions, including the most common intervention of  
513 hydroxychloroquine, peak in new weekly studies in early April. The plot also shows the later  
514 emergence of other interventions, such as convalescent plasma (shown in green).

515

### 516 3.3.1.6 *COVID-19 vaccine interventional trials*

517

518 Our search method for vaccine trial intervention type studies identified 12 COVID-19 vaccine  
519 trials, that also met our inclusion criteria of being active, recruiting or completed. Due to their  
520 high significance and increased public interest, it is interesting to consider how frequently such  
521 trials are updated. A total of 9 trials (75.0% of the 12 vaccine trials) have at least one update and  
522 the median amount of updates is two. Considering the study country, six different countries have  
523 at least one vaccine trial, with China (5 vaccine trials) having the most, followed by the US with  
524 3 trials. Five of the trials were Phase 1, six were Phase 1/Phase 2 and one was Phase 2. Of note is  
525 the fact that Phase 1 trials are not “applicable clinical trials” (as defined in US regulations) and  
526 such trials have no mandatory registration. (“FDAAA 801 and the Final Rule -  
527 ClinicalTrials.gov”) Exactly half of vaccine interventional trials (6 trials) had more than one site.  
528 As for design, the average number of arms was 5.4 with a median overall trial enrollment of  
529 119.5 participants. The 12 vaccine interventional trials also included 52 experimental arms and  
530 seven placebo comparator arms. The full overview of all metadata parameters for vaccine trials  
531 (as well as for observational studies and registries described in subsequent sections) is available  
532 at the study repository. (“regCOVID,” 2020)

533

### 534 3.3.2 **Observational studies**

535

536 We found a total of 287 observational studies. Similarly, to interventional trials, the vast majority  
537 of observational studies had just one site (238 studies, 82.9%). The country with the most  
538 observational studies was France with 75 (26.1%), followed by the United States with 47

539 (16.4%). Observational studies are updated less frequently than interventional trials as only  
540 52.6% (151 studies) of the COVID-19 observational studies have been updated since first being  
541 submitted to CTG (compared to the 71.1% of interventional trials that have been updated at least  
542 once). The observational study with the most updates was NCT04334954 ‘SARS-COV2  
543 Pandemic Serosurvey and Blood Sampling’ with 25 updates since registration on 6 April 2020.  
544 The most commonly updated public interest feature for observational studies was the ‘Study  
545 Status’ which was updated 270 times by 151 studies and the most common technical feature  
546 updated was ‘Contacts/Locations’ with 99 updates from 83 studies.

547

548 The median enrollment for observational studies was 353 participants. One feature of  
549 observational study design is the time perspective. A majority of the observational studies  
550 analyzed were prospective (180 studies, 62.7%), as opposed to 58 studies (20.2%) which were  
551 retrospective. For observational model, 167 of the observational studies (58.2%) use a cohort  
552 model. The second most commonly used model for the analyzed observational studies was case  
553 (45 studies, 15.7%).

554

555 Contrary to our expectation, we found observational studies that included interventions in their  
556 CTG record. Of the 287 observational studies, 179 (62.4%) listed something in the free-text  
557 intervention field. However, this number is misleading as in many cases the listed intervention  
558 was something that stated that there was no intervention (such as ‘no intervention’,  
559 ‘observation’, ‘non-interventional’, etc.). Of the listed interventions most are listed as  
560 intervention type ‘Other’ (86 studies, 30.0%) or ‘Diagnostic Test’ (34, 11.9%).

561

### 562 **3.3.3 Registries**

563

564 We analyzed a total of 64 COVID-19 registries (shorter term for registry-based studies). Of these  
565 registries 52 (81.3%) were limited to one site. The largest number of sites was 53. The countries  
566 with the most COVID-19 registries were France and the United States with 9 studies each.  
567 Similar to observational studies, just over half of the analyzed registries, 51.6% (33 registries),  
568 have been updated at least once since their first registration. Also similar to observational  
569 studies, the most common public interest update for registries is to the study status, which has  
570 been updated 56 times by all 33 registries with an update, and the most common technical update  
571 is to the contacts and locations with 28 updates from 18 studies.

572

573 The median enrollment for the set of registries was 388 participants. Registries have many  
574 specific design features that differentiate them from other study types. One is the presence of a  
575 targeted follow-up time. The most common follow-up time for the analyzed registries was one  
576 year for 15 studies (23.4%), which was listed as either ‘1 year’ or ‘12 months’ and was combined  
577 to get the accurate value. The shortest follow-up time was one day for NCT04331171,  
578 ‘Epidemiological Observation From a Smartphone Self-monitoring Application for Suspected  
579 COVID-19 Patients’ Triage’, while the longest targeted follow-up duration for a registry was 20  
580 years, for NCT04359602, ‘COVID-19 Recovered Volunteer Research Participant Pool Registry’.  
581 For registries, CTG collects their observational model (similar to observational studies). The  
582 majority of registries, 48 (75.0%), use a cohort model. Also similar to observational studies,  
583 registries can include a time perspective. However, unlike observational studies, no registries are

584 retrospective. Instead the time perspective is usually either prospective (50 studies, 78.1%), or  
585 cross-sectional (6 studies, 9.4%). A cross-sectional perspective means that the observation or  
586 intervention is made at a single point in time rather than on a continuous or recurring basis.(CTG  
587 Team, 2020a)

588

589 Like observational studies, more than half (53.1%, 34 of 64 registries) included an intervention  
590 in the free text field. These interventions also include many that are not representative of an  
591 actual intervention and rather state the absence of an intervention just like with the previously  
592 mentioned observational studies. This is also shown in the intervention type as 19 of the 34  
593 registries (55.9%) have an intervention type of ‘other’.

594

## 595 **4 Discussion**

596

597 Based on our two perspectives, we discuss separately COVID-19 studies results (journalist  
598 perspective) and data science implications (informatics perspective).

599

### 600 **4.1 COVID-19 studies**

601

602 Our study developed a computerized approach of retrieving COVID-19 studies from CTG  
603 registry for analysis. CTG’s study metadata facilitates the useful classification of studies into  
604 many relevant subgroups (e.g., by study design, size, phase, recruitment status or intervention).  
605 Availability of this data in a structured form (either via CTG’s API or via structured XML or  
606 relational data files) provides analytical views that would be difficult or impossible to achieve  
607 without a registry. As of 11 May 2020 ( the date of primary analysis), no study had deposited  
608 basic summary results.

609

610 The results presented above were summarized as of 11 May 2020. Refreshed and more current  
611 data (released weekly) can be obtained at the project repository. (“regCOVID Project  
612 Repository,” 2020; “regCOVID,” 2020). Weekly updated reports allow researchers, journalists  
613 or the general public to quickly obtain a snapshot of the ongoing COVID-19 research. For  
614 example, a weekly report intervention section (similar to Table 7) can reveal to many research  
615 teams concentrated on COVID-19 what interventions are being studied with what intensity. This  
616 analytical view would require tens of manual queries using the generic CTG web interface.

617

#### 618 **4.1.1 Study limitations**

619

620 Our study has several limitations. First, we only used CTG registry to look for COVID-19  
621 studies. Within this registry, we evaluated three search strategies, but some relevant COVID-19  
622 studies may possibly be missed. Without a benchmark gold standard of all COVID-19 studies,  
623 the recall of our search strategy cannot be evaluated. It was out of scope of this study to establish  
624 the precision of our search. Second, our semantic harmonization of interventions is based on  
625 manual mapping by a single expert. Third, there are significant limitations of the informatics-  
626 based approach compared to manual review.

627

#### 628 **4.1.2 Related studies**

629

630 For example, COVID-19 Evidence Service from Centre for Evidence-Based Medicine at  
631 University of Oxford offers more comprehensive reviews. It was out of scope of our project to  
632 offer results comparable to human review. Fragkou et al. used a search and manual review  
633 methodology to compile and analyze a set of COVID-19 interventional trials and their  
634 interventions.(Fragkou et al., 2020) Checcucci et al. did a literature and clinical trial registries  
635 search based on built-in search criteria to review COVID-19 vaccine trials.(Checcucci et al.,  
636 2020) Rosa et al. did a manual search of CTG to analyze COVID-19 trials using repurposed  
637 interventions. Considering the existing published studies, we conclude that our study is the first  
638 study to rely solely on computerized data science methods to compile and analyze a set of  
639 COVID-19 interventional trials, observational studies and registries(Rosa & Santos, 2020). Our  
640 approach of using computerized data science methods allows for the continuous monitoring of  
641 the current state of COVID-19 research with minimal additional effort compared to a resource  
642 intensive manual review methodology. During a continuously changing public health emergency,  
643 this ability for any researcher to quickly and efficiently monitor changes and trends in clinical  
644 research is invaluable in informing the direction of their research efforts.

645

## 646 **4.2 Data science perspective**

647

648 During the creation of a fully computerized, disease-focused report about ongoing or completed  
649 clinical studies, we observed several informatics themes described below. Before we describe  
650 individual lessons learned, we want to re-emphasize how computable representation of clinical  
651 study metadata is a crucial enabler for creating disease-based research snapshots. Moreover,  
652 several features of ClinicalTrials.gov registry proved to be highly valuable for our project. Such  
653 features are: structured representation of study metadata (XML and relational database format),  
654 registry support for result deposition and record updates, and legal and funding source policy  
655 requirements to maintain accurate registry records. In our analysis, we were able to build on  
656 prior clinical informatics research projects. Our project also shows value in further developing  
657 clinical informatics methods for data and metadata representation, semantic harmonization  
658 through terminologies and standards. The following informatics lessons were learned:

659

660 **Updates:** Our study is the first to analyze the frequency of updates to a study in CTG. We  
661 believe that adding the ability to access study updates to the CTG's API would be a useful  
662 addition. Our results indicate that analyzing study update activity is helpful in distinguishing  
663 studies with possibly outdated metadata (e.g., studies in status 'not yet recruiting' but are past  
664 their anticipated completion date with some grace period allowed for record updating). Our study  
665 is also the first to analyze update activity by country of study.

666

667 **Intervention (free text):** CTG collects intervention as free text and for some studies, provides a  
668 corresponding concept in Medical Subject Headings (MeSH) terminology. This intervention  
669 harmonization as MeSH concept is done post hoc rather than during study metadata entry by the  
670 study record manager. We found that the MeSH intervention concept is present in less than half  
671 (47%) of COVID-19 analyzed studies. This analysis prompted us to develop the denormalization  
672 and mapping method that we used.

673

674 Another intervention-related observation is the difference in how intervention combinations are  
675 listed in the free text field. In some cases, the combination intervention (e.g., 2 drugs given to

676 some study group in combination) is recorded as two separate entries and the group or arm free-  
677 text description provides a way to clarify the combined usage. In other case the same  
678 intervention combination is recorded together as a single entry. This dual way of recording  
679 combined interventions formed our methodology for the most comprehensive approach of  
680 counting interventions (count them as both combinations and as separate interventions). We did  
681 not analyze arm description and so we did not combine separated interventions, which may have  
682 been assigned to the same arm and used in combination. This may possibly lead to the  
683 undercounting of certain intervention combinations.

684

685 **Registries:** We find valuable that CTG currently allows registration of observational studies and  
686 registries. Designing a user interface for registration and study representation format that can  
687 accommodate various designs and studies is a challenging task. Due to specific characteristics of  
688 certain study types, further customization of user interface or additional data quality checks may  
689 further improve the registry value to many stakeholders. For example, registries do not typically  
690 post one-time study results and may not have the same concept of primary completion date.  
691 Instead, annual or other regular interval updates about number of participants and summary  
692 results for participant flow may be more applicable. Clarifications in the user interface for  
693 entering interventions for registries (and for observational studies) may prevent entries which  
694 declare a formally drug typed intervention with the title ‘no intervention’.

695

#### 696 **4.2.1 Generalizing report to other diseases**

697

698 Our emphasis on fully computerized analysis of a COVID-19 set of studies was motivated by our  
699 larger vision to apply the R scripted report for all MeSH encoded diseases found within the CTG  
700 registry. We refer to this result as the regCTG project and report repository. regCTG allows  
701 analysis of research by MeSH keyword for all clinical domains. We generated reports for all  
702 MeSH terms with at least 100 registered studies. A collection of nearly 1000 disease-based  
703 reports is available at <https://github.com/lhncbc/CRI/tree/master/regCTG>.

704 We consider this generalization from a COVID-19 research report to a research report for nearly  
705 1000 diseases an important result of our project.

706

707

708 In another follow-up research project for this COVID-19 case study, we have also built a  
709 disease-intervention snapshot knowledge base (called D-SHOT) that lists all interventions  
710 appearing in interventional trials for a given condition. (“Project Repository for Disease  
711 Snapshot”) This knowledge base of disease-intervention pairs has many parameters for each  
712 intervention, such as date when first introduced, count of regularly completed studies or count of  
713 unusually completed studies (‘terminated’, ‘suspended’, or ‘withdrawn’) studying that  
714 intervention. Experience from semantic harmonization of CTG’s free text field into terminology  
715 concepts gained during this COVID-19 project was crucial in these two follow-up projects by  
716 our team. A related, non-open source project called Sherlock, proprietary to Johnson and  
717 Johnson is similarly parsing CTG’s terms into formal concepts.(Cepeda, Lobanov & Berlin,  
718 2013)

719

### 720 **4.3 Weekly results updates**

721 While, the main analysis presented above was done on 11 May 2020 (main analysis date), thanks  
722 to the computed nature of the analysis, we have been producing weekly updated reports  
723 (available at the github repository). We have also been improving and adding to the automated  
724 report since the main analysis date based on the deposition of the first study results and the  
725 appearance of study results publications. As of the main analysis date, there were zero studies  
726 with results deposited on CTG. Because of data evolution during the article review and revision  
727 preparation, the latest weekly report on our github repository (as of 13 August 2020; update  
728 analysis date) now shows 3 interventional trials and one registry with results posted. Analysis of  
729 linked PubMed publications for completed interventional trials, found that of the 83 completed  
730 interventional trials at the point of secondary analysis, 9 had linked PubMed publications  
731 (10.8%).

732

733 For the weekly reports and data in our github repository, we welcome change requests submitted  
734 by interested researchers. For researchers re-using our code and interested in making  
735 modifications, a free registration to access the AACT database is required (obtainable within  
736 hours).

737

## 738 **5 Conclusions**

739

740 We developed a computerized, data science driven approach to monitoring COVID-19 interventional  
741 trials, observational studies and registries. We report on several metrics for the 401 interventional trials,  
742 287 observational studies and 64 registries as of our analysis date on 11 May 2020. More current and  
743 weekly refreshed data is available at our github repository. We also demonstrated that our COVID-19  
744 disease focused report can be generalized to all diseases represented within a clinical trial registry.

745

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757 metadata fields not required by US regulations).

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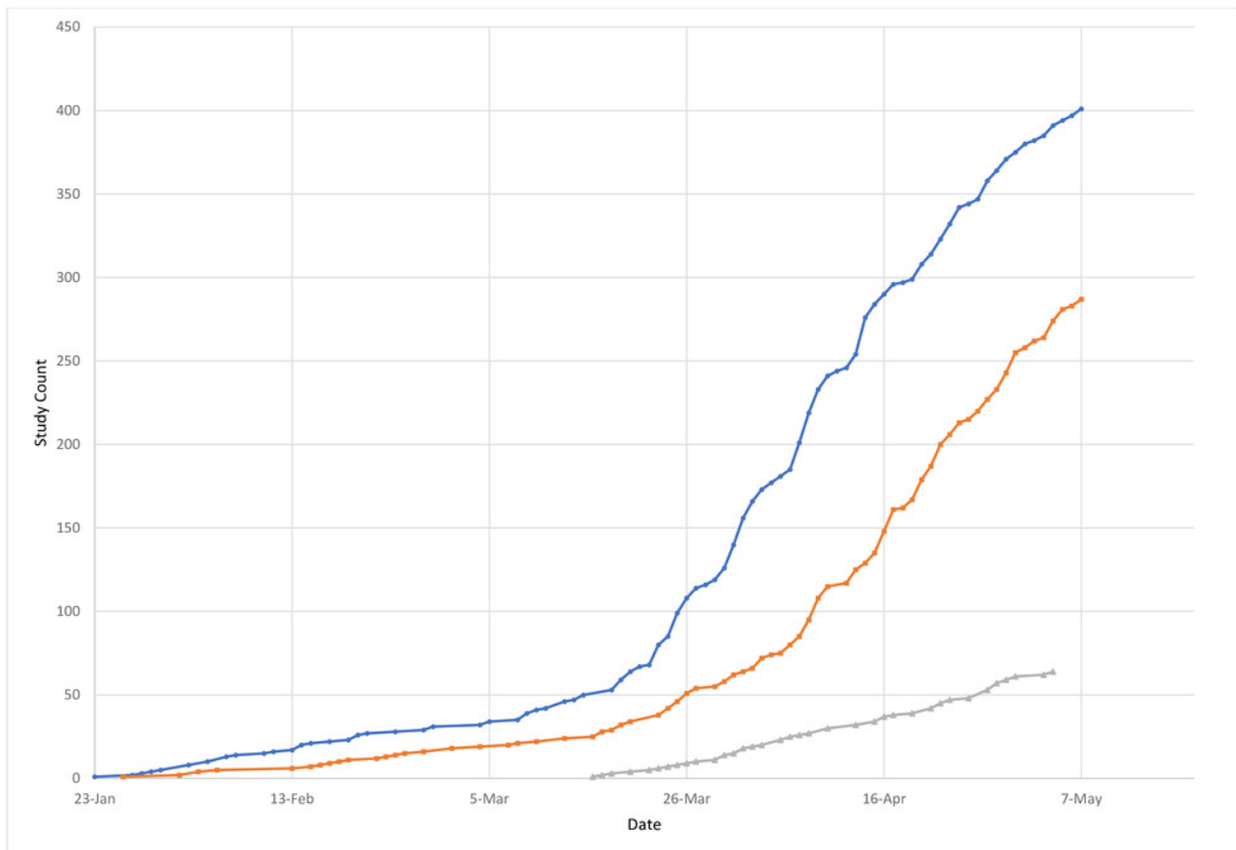
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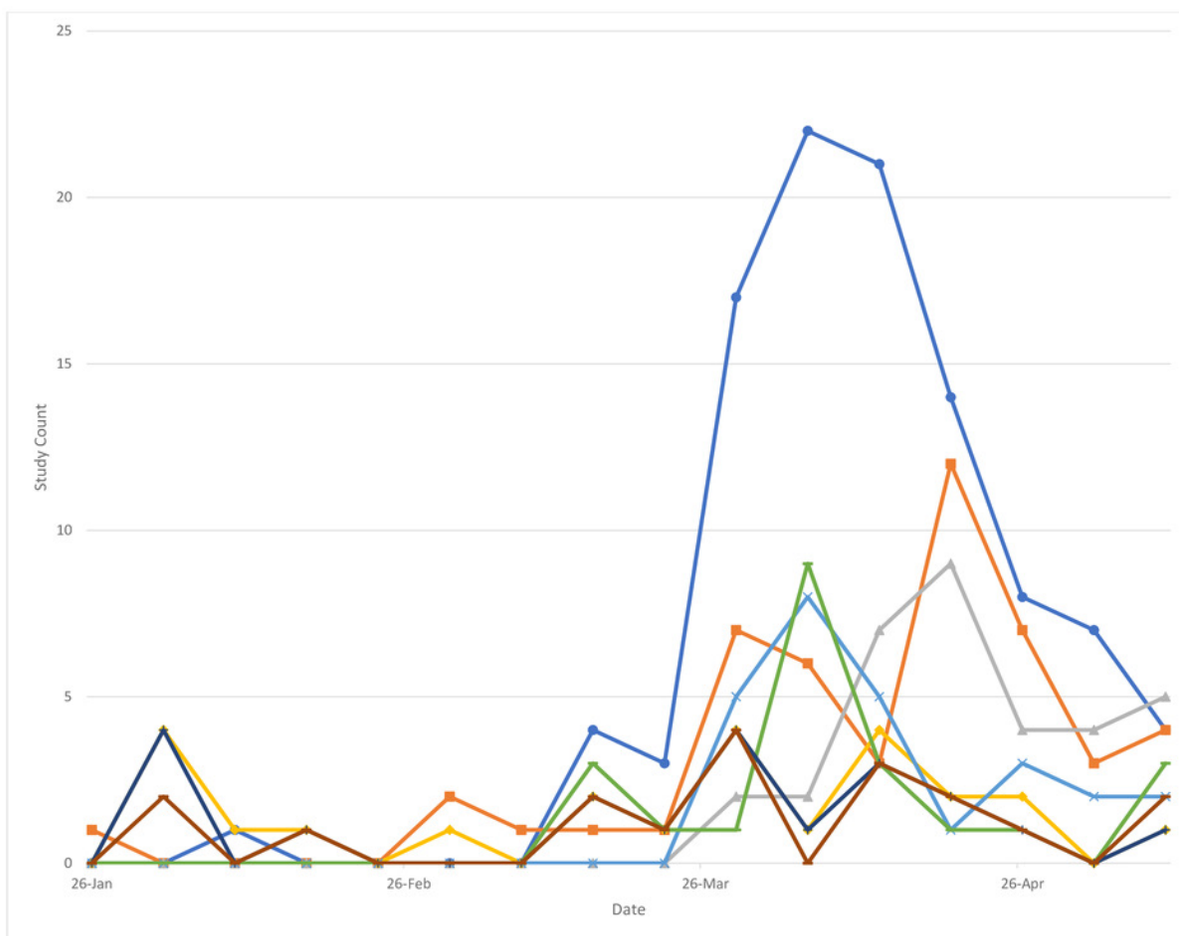
# Figure 1

Studies over time by study type



## Figure 2

Plot of new studies weekly for selected frequent COVID-19 interventions over time



**Table 1** (on next page)

Table 1. List of countries where COVID-19 interventional trials are conducted.

1 **Table 1.** List of countries where COVID-19 interventional trials are conducted.

Country	Study Count	Percentage
United States	121	30.2%
China	49	12.2%
France	42	10.5%
Spain	23	5.7%
Italy	19	4.7%
Brazil	10	2.5%
Canada	10	2.5%
Iran, Islamic Republic of	10	2.5%
Germany	8	2.0%
Mexico	8	2.00%

2

3

**Table 2** (on next page)

Number of updates per study by country (for countries with at least 8 COVID-19 interventional trials)

1 **Table 2.** Number of updates per study by country (for countries with at least 8 COVID-19  
2 interventional trials)

Country	Total Updates	Study Count	Changes Per Study
Canada	27	10	2.70
United States	213	121	1.76
Germany	14	8	1.75
Brazil	17	10	1.70
Spain	38	23	1.65
China	65	49	1.33
France	53	42	1.26
Iran	12	10	1.20
Mexico	7	8	0.88
Italy	13	19	0.68

3

**Table 3** (on next page)

Overview of studies by study phase and number of participants (study size)

1 **Table 3.** Overview of studies by study phase and number of participants (study size)

Phase	Study Count	Percentage	1st Qu.	# of participants: median (IQR)*	3rd Qu.
N/A	111	27.7%	49.5	120	330
Early Phase 1	7	1.7%	10	10	40
Phase 1	17	4.2%	20	40	54
Phase 1/Phase 2	23	5.7%	20	72	190
Phase 2	108	26.9%	60	145	273.75
Phase 2/Phase 3	34	8.5%	108	269.5	433.5
Phase 3	74	18.5%	245	500	1215
Phase 4	27	6.7%	83	200	450

2 \* IQR is interquartile range (1<sup>st</sup> quartile [25<sup>th</sup> percentile] and 3<sup>rd</sup> quartile [75<sup>th</sup> percentile])

3

**Table 4** (on next page)

Primary purpose of COVID-19 interventional trials

1 **Table 4.** Primary purpose of COVID-19 interventional trials

Primary Purpose	Study Count	Percentage
Treatment	298	74.3%
Prevention	41	10.2%
Other	19	4.7%
Supportive Care	17	4.2%
Diagnostic	15	3.7%
Health Services Research	7	1.7%
Basic Science	2	0.5%
Screening	2	0.5%

2

**Table 5** (on next page)

Count of the number of arms by arm type

1 **Table 5.** Count of the number of arms by arm type

Arm Type	Arm Count
Experimental	489
Active Comparator	160
Placebo Comparator	118
No Intervention	87
Other	43
Sham Comparator	3

2

**Table 6** (on next page)

Count of intervention types included in interventional trials

1 **Table 6.** Count of intervention types included in interventional trials

Composite Intervention Type	Study Count	Percentage
Drug	137	34.2%
Drug Placebo	75	18.7%
Biological	32	8.0%
Other	31	7.7%
Device	22	5.5%
Drug Other	22	5.5%
Behavioral	12	3.0%
Biological Placebo	12	3.0%
Diagnostic Test	10	2.5%
Procedure	8	2.0%
All Other Types*	40	10.0%

2 \*This row combines rare Composite Intervention Types, such as 'Drug|Biological' , 'Dietary  
 3 Supplement', or 'Device|Procedure' (see repository report for full table of intervention types)(“r-  
 4 snippets-bmi/regCOVID at master · lhncbc/r-snippets-bmi”)  
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**Table 7** (on next page)

Most frequent interventions by study count (with a minimum study count of 13)

1 **Table 7.** Most frequent interventions by study count (with a minimum study count of 13)

Intervention Type	Intervention	Study Count	Date when first appeared
Placebo	Placebo	99	20-Feb-2020
Drug	Hydroxychloroquine	92	06-Feb-2020
Other	Standard care	40	23-Jan-2020
Drug	Azithromycin	24	23-Mar-2020
Drug	Ritonavir	24	28-Jan-2020
Drug	Tocilizumab	21	09-Mar-2020
Biological	Convalescent plasma	20	23-Mar-2020
Drug	Lopinavir	20	28-Jan-2020
Drug	Lopinavir/Ritonavir	16	30-Jan-2020
Drug	Chloroquine	13	19-Mar-2020

2