

1 **Gender disparities in COVID-19 clinical trial leadership**

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31 **Abstract**

32 **Objectives:** We aimed to compare the gender distribution of clinical trial leadership in COVID-19 clinical
33 trials.

34 **Methods:** We searched <https://clinicaltrials.gov/> and retrieved all clinical trials on COVID-19 from
35 January 1, 2020 to June 26, 2020. As a comparator group, we have chosen two fields that are not
36 related to emerging infections and infectious diseases: and considered not directly affected by the
37 pandemic : breast cancer and type 2 diabetes mellitus (T2DM) and included studies within the
38 aforementioned study period as well as those registered in the preceding year (pre-study period:
39 January 1, 2019 and December 31, 2019). Gender of the investigator was predicted using the
40 genderize.io API (application programming interface). The repository of the datasets used to collect and
41 analyse the data available at <https://osf.io/k2r57/>.

42 **Results:** Only 27.8% (430/1548) of principal investigators (PIs) among COVID-19-related studies were
43 women, which is significantly different compared to 54.9% (156/284) and 42.1% (56/133) for breast
44 cancer ($p<0.005$) and T2DM ($p<0.005$) trials over the same period, respectively. During this “pre-study”
45 period, the proportion of PIs who were predicted to be women were 49.7% (245/493) and 44.4%
46 (148/333) for breast cancer and T2DM trials, respectively and the difference was not statistically
47 significant when compared to results from the study period ($p>0.05$).

48 **Conclusion:** We demonstrate that less than one-third of COVID-19-related clinical trials are led by
49 women PIs, half the proportion observed in non-COVID-19 trials over the same period which remained
50 similar to the pre-study period. These gender disparities during the pandemic may indicate not only a
51 lack of women's leadership in international clinical trials and involvement in new projects but also may
52 reveal imbalances in women's access to research activities and funding during health emergencies.

53

54 **Key words:** COVID-19, coronavirus, pandemic, SARS-CoV-2, novel coronavirus, gender

55

56 **Introduction**

57

58 In addition to the human and financial loss associated with the novel Coronavirus Disease 2019 (COVID-
59 19) pandemic, COVID-19 has also had a significant impact on both the personal and professional life of
60 the global workforce, including that of the scientific research community [1-3]. Before COVID-19,
61 women occupied fewer leadership positions, led a fewer funded studies, and applied for and received
62 less grant funding than men when they did apply [4-7]. The employment gap that occurs when women
63 take parental leave impacts the rate of academic advancement and in turn the receipt of institutional
64 support to apply for and secure funding [6, 7]. These imbalances contribute to systemic inequalities that
65 hamper women's access to and progress in science [2, 7, 8]. A review of the gender distribution of 24
66 COVID-19 national task forces suggests that many committees are comprised of less than a quarter
67 women, indicating that women's voices and expertise have been excluded from decision making during
68 this unprecedented public health emergency [9].

69

70 For example, emerging data suggest that across all disciplines, despite an increased number of peer-
71 reviewed articles submitted to journals during the pandemic, women have published fewer papers than
72 men thus far this year [10]. This may indicate a similarly reduced involvement of women in research
73 leadership positions and an imbalanced distribution of grants and funding -- important indicators of
74 advancement in a scientist's academic career [4-7, 10, 11]. Being principal investigator (PI) on a clinical
75 trial is strongly associated with advancement to full professor among women academics in infectious
76 diseases [8].

77

78 The COVID-19 pandemic offers numerous opportunities in clinical research. These include trials to assess
79 the safety and efficacy of medical interventions, with protocols in various stages of implementation.
80 Here, we compare the gender distribution of clinical trial leadership in COVID-19 clinical trials.

81

82 **Materials and Methods**

83 We systematically searched <https://clinicaltrials.gov/> and retrieved all clinical trials on COVID-19
84 registered from January 1, 2020 to June 26, 2020 using “COVID” as a keyword. As a comparator group,
85 we have chosen two fields that are not related to emerging infections and infectious diseases, and
86 considered not directly affected by the pandemic: breast cancer and type 2 diabetes mellitus (T2DM).
87 We retrieved all clinical trials related to these comparator conditions registered at
88 <https://clinicaltrials.gov/> within the aforementioned study period as well as those registered in the
89 preceding year (pre-study period: January 1, 2019 and December 31, 2019). We retrieved the names of
90 investigators listed; study director, principal investigator (PI) (the person who is responsible for the
91 scientific and technical direction of the entire clinical study) and study chair (whose role involve toxicity
92 and accrual monitoring). Gender of the investigator was predicted using the genderize.io API
93 (application programming interface). This tool has been used to predict the gender of first names in
94 studies regarding gender bias [12, 13] and achieves a minimum accuracy of 82%, with an F1 score
95 (weighted average of precision and recall) of 90% for women and 86% for men [14]. Clinical trials were
96 excluded if i) investigator information was not provided; ii) the genderize.io API could not predict any of
97 the investigators’ gender from their first name; or iii) organization or company names were provided as
98 the investigator. The number of studies that were excluded for the above reasons are reported in the
99 supplementary flow diagram. An exploratory temporal analysis was conducted with the available data.
100 Categorical variables were summarized by frequencies and percentages. We compared groups using Chi-
101 square testing for equality of proportions with continuity correction [15]. The analysis was performed

102 using R (Version 4.0.2). The repository of the datasets used to collect and analyse the data available at
103 <https://osf.io/k2r57/>.

104

105 **Results**

106 We identified 2 345 COVID-19-related clinical trials. Of those, 1 448 had at least one investigator listed
107 (i.e., principal investigator, study director, or study chair) whose gender could be predicted. In the
108 comparator group, we identified 449 trials on breast cancer and 272 on T2DM that were registered. Of
109 those, 274 breast cancer studies and 139 T2DM studies had at least one investigator whose gender
110 could be predicted.

111

112 Overall 27.8% (430/1548) of PIs among COVID-19-related studies were predicted to be women, which is
113 significantly different compared to 54.9% (156/284) and 42.1% (56/133) for breast cancer ($p<0.005$) and
114 T2DM ($p<0.005$) trials over the same period, respectively (Table 1). While there has been a small
115 increase in the proportion of PIs who were predicted to be women in May 2020, clinical research
116 leadership for COVID-19 among this group was below 25% for the remainder of the study period
117 (Supplementary Material). While 31.4% (76/242) of study chairs were predicted to be women in COVID-
118 19-related studies, 32.1% (9/28) ($p=0.7$) and 63.6% (7/11) ($p<0.01$) were predicted to be women in
119 breast cancer and T2DM trials, respectively. Proportion of study chairs were not significantly different
120 across the three fields.

121

122 We also reviewed comparator group studies registered before January 1, 2020 to determine whether
123 the pandemic might have affected gender distribution of trial leadership. We identified 839 clinical trials
124 related to breast cancer and 533 on T2DM over a 12-month period prior to January 1, 2020. Of those,
125 573 breast cancer studies and 359 T2DM studies yielded at least one investigator whose gender could

126 be predicted. During this “pre-study” period, the proportion of PIs who were predicted to be women
127 were 49.7% (245/493) and 44.4% (148/333) for breast cancer and T2DM trials, respectively and the
128 difference was not statistically significant when compared to results from the study period ($p>0.05$).
129

130 **Discussion**

131 In this study, we demonstrate that less than one-third of COVID-19-related clinical trials are led by
132 women PIs, half the proportion observed in non-COVID-19 (breast cancer and T2DM) trials over the
133 same period. The proportion of PIs in breast cancer and T2DM studies also remained similar to the pre-
134 study period. These gender disparities during the pandemic may indicate not only a lack of women’s
135 leadership in international clinical trials and involvement in new projects, but also may reveal
136 imbalances in women's access to research activities and funding during health emergencies [2, 16].
137

138 The COVID-19 pandemic offers numerous opportunities for research and leadership that could equalize
139 opportunity in a new field, but our results suggest the opposite. The pandemic has reinforced the
140 prevailing gender norms in which men continue to be allocated a disproportionate share of the funding,
141 as well as leadership and authorship roles [9, 10, 16]. One potential contributor for this discrepancy is
142 the speed demanded by the research agenda during the pandemic. The sense of urgency in starting
143 clinical trials may lead to an abandonment of any checks and balances around equality and inclusion
144 that would have otherwise encouraged the involvement of women scientists. Many women scientists
145 have already raised concerns about institutional funding distribution lacking gender balance or being left
146 out of research activities despite their expertise [2, 16]. During COVID-19 pandemic, a UK study showed
147 that women were more than twice as likely to take on childcare and schooling responsibilities of
148 children than men, while male academic counterparts leverage professional relationships and networks
149 more effectively [1, 2, 16].

150

151 As a community, we must recognise that there is a tendency to “turn to men” in times of crisis both for
152 leadership and scientific expertise [2, 3, 16, 17], highlighting the need to challenge this culture. Research
153 and academia are already competitive; being in the central decision-making group is often challenging
154 due to gender norms, along with roles and rules on how these groups are established and maintained;
155 during health emergencies, these same authoritative circles become more difficult for women scientists
156 to join [2, 16]. Our findings suggest that there is a need for transparency in opportunities and funding
157 that requires actively identifying and addressing the structurally implicit and unconscious biases that
158 favour men. For example, in recent years, the campaign against MANELs (Male-only Panels) has already
159 met considerable support in the scientific community and several influential journals have published
160 policies and editorials in support of women in science and medicine.

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162 The evidence while sparse indicates that teams that are diverse in terms of gender, ethnicity, and social
163 background produce better health science, are more highly cited, generate a broader range of ideas and
164 innovations, and better represent society [2, 16, 18, 19]. Not only can these women drive discovery and
165 innovation, but they can act to address health disparities and provide role models for the next
166 generation of women scientists [2, 16, 18, 19]. Ensuring gender representation would also reflect the
167 commitment of the global community to promoting gender equality in academic medicine and research:
168 inclusion, diversity, representation, progression, and success for all. Therefore, the disadvantage not
169 only affect women themselves and their research career but has much more profound implications for
170 the wider society especially given the disproportionate burden of such outbreaks for communities who
171 are marginalized due to their gender, sexuality, class, ethnicity, and ability [20-22].

172

173 Our analysis has some limitations. We could include only ~50-75% of trials for which an investigator's
174 gender could be algorithmically predicted because the majority of studies had no investigator
175 information, or the investigator names were not distinguishable (supplementary material). Furthermore,
176 while such algorithms allow for the rapid analysis of gender disparities such as those conducted here,
177 they can also be exclusionary to gender non-conforming, non-binary, and trans individuals. Beyond
178 these limitations, although there were several observational studies in our dataset, clinicaltrials.gov may
179 be biased towards randomised control trial registration and women may be more likely to be involved in
180 observational studies, which still demonstrates gender disparities in types of trials women lead. Also, we
181 did not consider studies that received private funding, which may not have been registered on
182 clinicaltrials.gov; however, it is worth noting that clinicaltrials.gov is an international database with
183 widespread international representation. Finally, while we attempted to provide a comparison with two
184 other fields, a potential for bias could arise from the difference of gender distributions of researchers
185 working in the fields of infectious diseases, breast cancer and diabetes.

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187 In summary, while the COVID-19 pandemic has thus far provided many new opportunities for research,
188 with numerous clinical trials initiated worldwide, a disproportionate proportion of PIs leading COVID-19
189 related studies are predicted to be men, despite women accounting for 70% of the global health
190 workforce [16]. Our demonstration of gender differences in trial leadership argue for revised policies
191 and strategies that encourage the participation and leadership of women in pandemic research. This
192 may include setting up review committees that are gender balanced, available funding to be provided to
193 equal number of PIs, or funding gender balanced trial teams, and overall ensuring that funding agencies
194 are aware of the lack of women leadership in clinical trials.

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196 **Authors contributions**

197 MC: conceptualisation, methodology, investigation, literature review, data curation, writing – original
198 draft. SH and MM: investigation, data curation, formal analysis, writing – review and editing; JS, KK, PS:
199 methodology, writing – review and editing, supervision. CO: conceptualisation, methodology,
200 investigation, literature review, writing – original draft, supervision.

201

202 **Financial support and sponsorship**

203 None

204

205 **Conflict of interests**

206 MC, SH, JS, MM have none to disclose. CO has received honoraria, fees for lectures, and advisory boards
207 from Gilead, MSD, Viiv, and Janssen. She has also received research grants to her institution from the
208 above-mentioned companies. PES has received honoraria, fees for lectures, and advisory boards from
209 Gilead, Merck, Janssen, and ViiV; he has also received research grants to his institution from Gilead and
210 ViiV. KK has received personal fees from GSK, outside the submitted work.

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	Jan 1, 2020 - June 26, 2020					before Jan 1, 2020		
	COVID-19	Breast Cancer	p value	T2DM	p value	Breast Cancer	T2DM	p value
PI	27.8% (430/1548)	54.9% (156/284)	<0.01	42.1% (56/133)	<0.01	49.7% (245/493)	44.4% (148/333)	0.15
Study Director	28.7% (72/251)	48.9% (23/47)	<0.01	22.2% (4/18)	0.75	30.5% (29/95)	47.6% (40/84)	0.02
Study Chair	31.4% (76/242)	32.1% (9/28)	1	63.6% (7/11)	0.98	33.3% (26/78)	40.4% (19/47)	0.54

Table 1: Proportion of women leadership in clinical trials between January 1, 2020 and June 26, 2020 and before January 1, 2020

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226 **Supplementary material:**

227 Flow diagram of process of selection

228 Gender distribution over time (months)

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