

Proxalutamide reduces COVID-19 risk: real-time meta analysis of 4 studies

@CovidAnalysis, July 2025, Version 11
<https://c19early.org/xmeta.html>

Abstract

Significantly lower risk is seen for mortality, ventilation, hospitalization, recovery, and viral clearance. 4 studies from 2 independent teams in 2 countries show significant benefit.

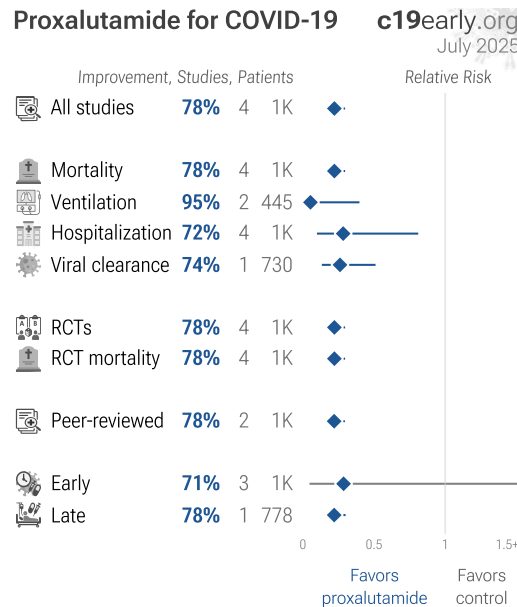
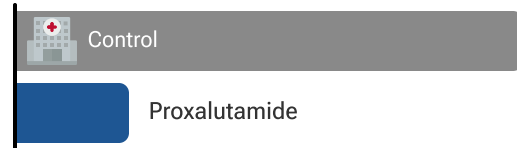
Meta analysis using the most serious outcome reported shows 78% [70-83%] lower risk. Results are similar for peer-reviewed studies. Currently all studies are RCTs.

Studies to date are from only 2 different groups.

No treatment is 100% effective. Protocols combine safe and effective options with individual risk/benefit analysis and monitoring. All data and sources to reproduce this analysis are in the appendix.

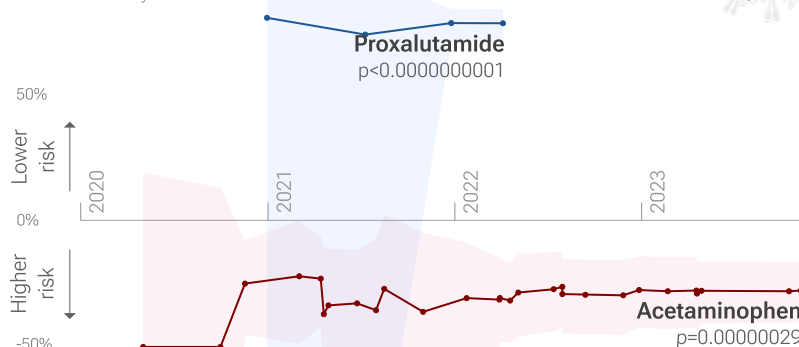
Zheng *et al.* present another meta analysis for proxalutamide, showing significant improvements for mortality, mechanical ventilation, hospitalization, and clinical improvement.

Serious Outcome Risk



Evolution of COVID-19 clinical evidence

Meta analysis results over time



PROXALUTAMIDE FOR COVID-19 – HIGHLIGHTS

Proxalutamide reduces risk with very high confidence for mortality and in pooled analysis, high confidence for hospitalization, and low confidence for ventilation, recovery, and viral clearance.

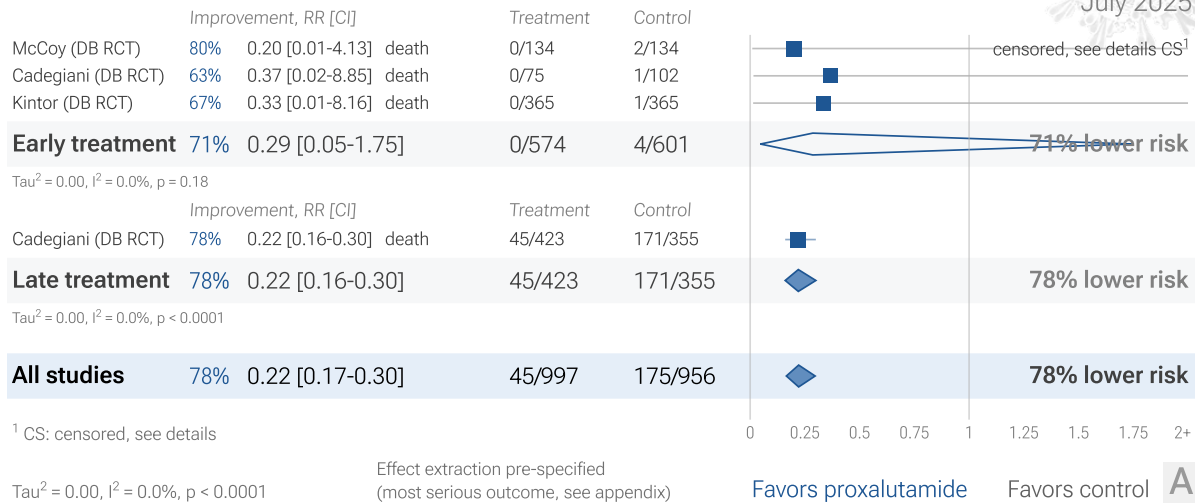
34th treatment shown effective in December 2021, now with $p < 0.0000000001$ from 4 studies, recognized in 3 countries.

Real-time updates and corrections with a consistent protocol for 172 treatments. Outcome specific analysis and combined evidence from all studies including treatment delay, a primary confounding factor.

4 proxalutamide COVID-19 studies

c19early.org

July 2025



Timeline of COVID-19 proxalutamide studies (pooled effects)

c19early.org

July 2025

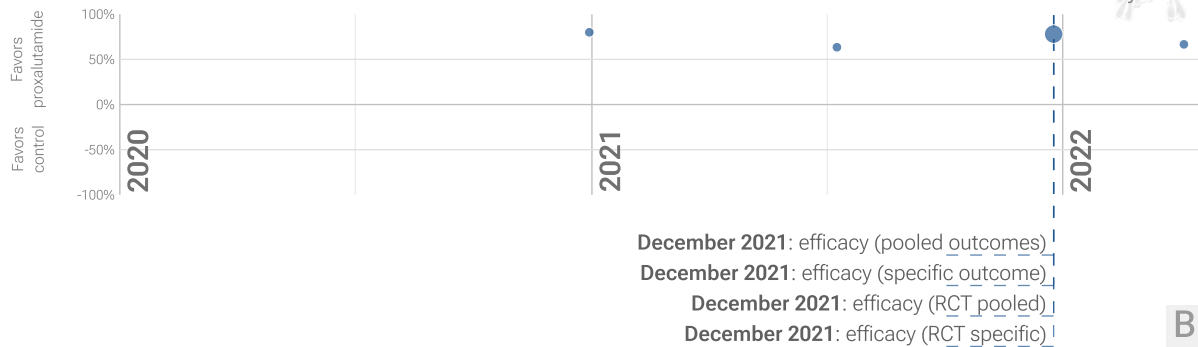


Figure 1. A. Random effects meta-analysis. This plot shows pooled effects, see the specific outcome analyses for individual outcomes. Analysis validating pooled outcomes for COVID-19 can be found below. Effect extraction is pre-specified, using the most serious outcome reported. For details see the appendix. **B. Timeline of results in proxalutamide studies.** The marked dates indicate the time when efficacy was known with a statistically significant improvement of $\geq 10\%$ from ≥ 3 studies for pooled outcomes, one or more specific outcome, pooled outcomes in RCTs, and one or more specific outcome in RCTs.

Introduction

Immediate treatment recommended

SARS-CoV-2 infection primarily begins in the upper respiratory tract and may progress to the lower respiratory tract, other tissues, and the nervous and cardiovascular systems, which may lead to cytokine storm, pneumonia, ARDS, neurological injury³⁻¹⁵ and cognitive deficits^{6,11}, cardiovascular complications¹⁶⁻²⁰, organ failure, and death. Even mild untreated infections may result in persistent cognitive deficits²¹ — the spike protein binds to fibrin leading to fibrinolysis-resistant blood clots, thromboinflammation, and neuropathology. Minimizing replication as early as possible is recommended.

Many treatments are expected to modulate infection

SARS-CoV-2 infection and replication involves the complex interplay of 100+ host and viral proteins and other factors^{A,22-29}, providing many therapeutic targets for which many existing compounds have known activity. Scientists have predicted that over 9,000 compounds may reduce COVID-19 risk³⁰, either by directly minimizing infection or replication, by supporting immune system function, or by minimizing secondary complications.

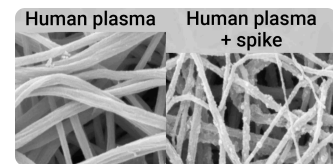


Figure 2. SARS-CoV-2 spike protein fibrin binding leads to thromboinflammation and neuropathology, from².

Analysis

We analyze all significant controlled studies of proxalutamide for COVID-19. Search methods, inclusion criteria, effect extraction criteria (more serious outcomes have priority), all individual study data, PRISMA answers, and statistical methods are detailed in Appendix 1. We present random effects meta-analysis results for all studies, studies within each treatment stage, individual outcomes, peer-reviewed studies, and Randomized Controlled Trials (RCTs).

Treatment timing

Figure 3 shows stages of possible treatment for COVID-19. Prophylaxis refers to regularly taking medication before becoming sick, in order to prevent or minimize infection. Early Treatment refers to treatment immediately or soon after symptoms appear, while Late Treatment refers to more delayed treatment.

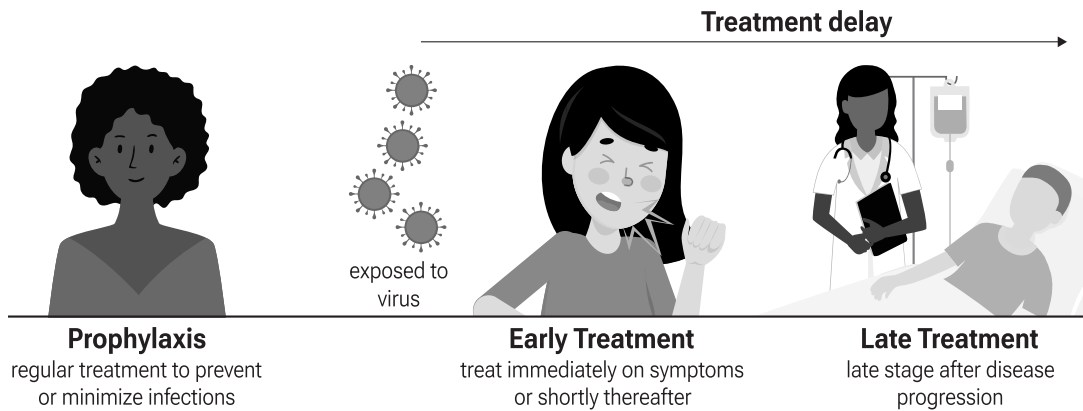


Figure 3. Treatment stages.

Preclinical Research

2 *In Vitro* studies support the efficacy of proxalutamide^{31,32}.

Preclinical research is an important part of the development of treatments, however results may be very different in clinical trials. Preclinical results are not used in this paper.

Results

Table 1 summarizes the results for all stages combined, for Randomized Controlled Trials, for peer-reviewed studies, and for specific outcomes. Table 2 shows results by treatment stage. Figure 4 plots individual results by treatment stage. Figure 5, 6, 7, 8, 9, 10, and 11 show forest plots for random effects meta-analysis of all studies with pooled effects, mortality results, ventilation, hospitalization, recovery, viral clearance, and peer reviewed studies.

	Relative Risk	Studies	Patients
All studies	0.22 [0.17-0.30] ****	4	1,953
Peer-reviewed	0.22 [0.16-0.30] ****	2	1,046
RCTs	0.22 [0.17-0.30] ****	4	1,953
Mortality	0.22 [0.17-0.30] ****	4	1,953
Ventilation	0.05 [0.01-0.40] **	2	445
Hospitalization	0.28 [0.10-0.81] *	4	1,953
RCT mortality	0.22 [0.17-0.30] ****	4	1,953
RCT hospitalization	0.28 [0.10-0.81] *	4	1,953

Table 1. Random effects meta-analysis for all stages combined, for Randomized Controlled Trials, for peer-reviewed studies, and for specific outcomes. Results show the relative risk with treatment and the 95% confidence interval. ** $p < 0.01$ *** $p < 0.001$ **** $p < 0.0001$.

	Early treatment	Late treatment
All studies	0.29 [0.05-1.75]	0.22 [0.16-0.30] ****
Peer-reviewed	0.20 [0.01-4.13]	0.22 [0.16-0.30] ****
RCTs	0.29 [0.05-1.75]	0.22 [0.16-0.30] ****
Mortality	0.29 [0.05-1.75]	0.22 [0.16-0.30] ****
Ventilation	0.05 [0.01-0.40] **	
Hospitalization	0.19 [0.07-0.54] **	0.67 [0.54-0.82] ***
RCT mortality	0.29 [0.05-1.75]	0.22 [0.16-0.30] ****
RCT hospitalization	0.19 [0.07-0.54] **	0.67 [0.54-0.82] ***

Table 2. Random effects meta-analysis results by treatment stage. Results show the relative risk with treatment and the 95% confidence interval. ** $p < 0.01$ *** $p < 0.001$ **** $p < 0.0001$.

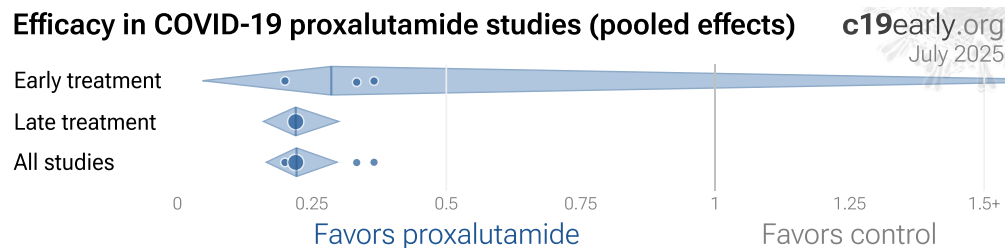


Figure 4. Scatter plot showing the most serious outcome in all studies, and for studies within each stage. Diamonds shows the results of random effects meta-analysis.

4 proxalutamide COVID-19 studies

c19early.org

July 2025

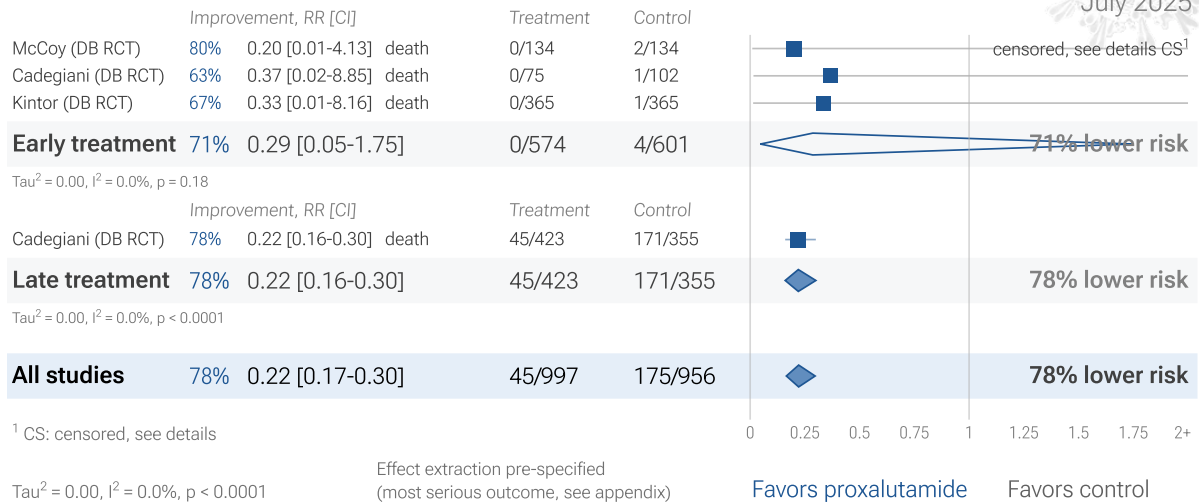


Figure 5. Random effects meta-analysis for all studies. This plot shows pooled effects, see the specific outcome analyses for individual outcomes. Analysis validating pooled outcomes for COVID-19 can be found below. Effect extraction is pre-specified, using the most serious outcome reported. For details see the appendix.

4 proxalutamide COVID-19 mortality results

c19early.org

July 2025

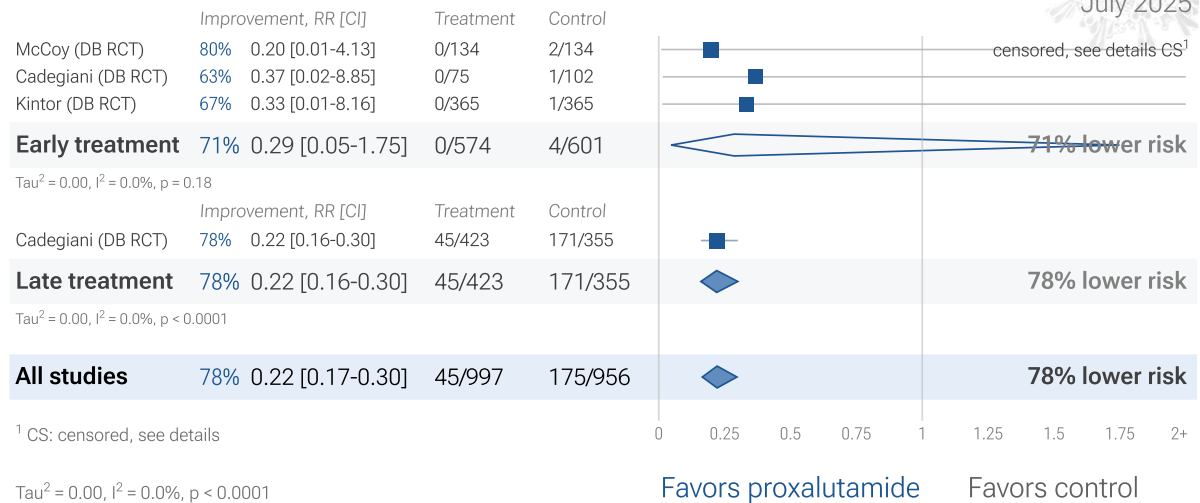


Figure 6. Random effects meta-analysis for mortality results.

2 proxalutamide COVID-19 mechanical ventilation results

c19early.org

July 2025

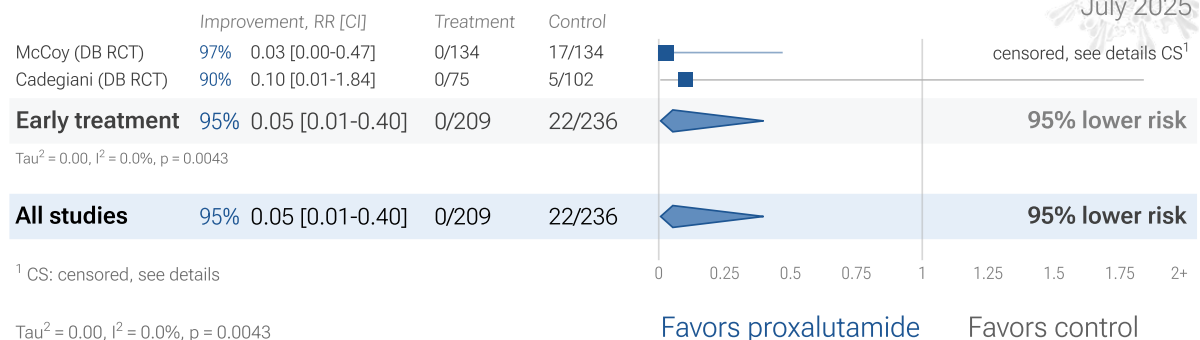


Figure 7. Random effects meta-analysis for ventilation.

4 proxalutamide COVID-19 hospitalization results

c19early.org

July 2025

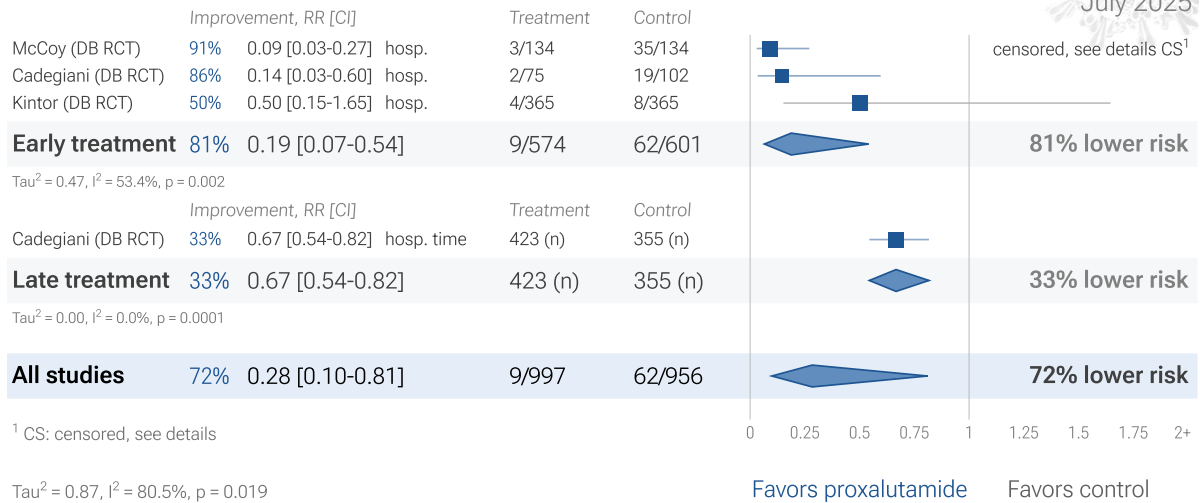


Figure 8. Random effects meta-analysis for hospitalization.

1 proxalutamide COVID-19 recovery result

c19early.org

July 2025

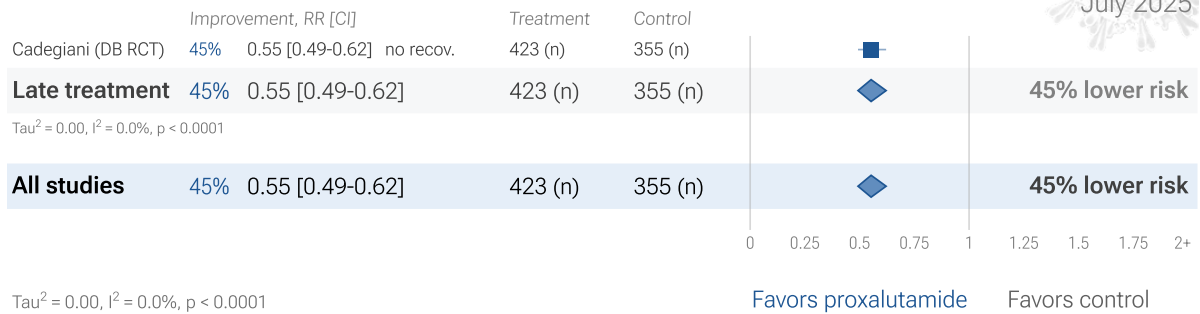


Figure 9. Random effects meta-analysis for recovery.

1 proxalutamide COVID-19 viral clearance result

c19early.org

July 2025

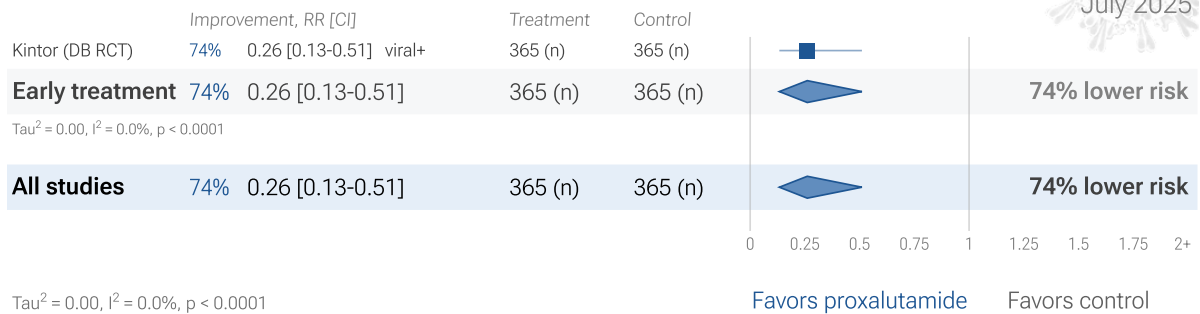


Figure 10. Random effects meta-analysis for viral clearance.

2 proxalutamide COVID-19 peer reviewed studies

c19early.org

July 2025

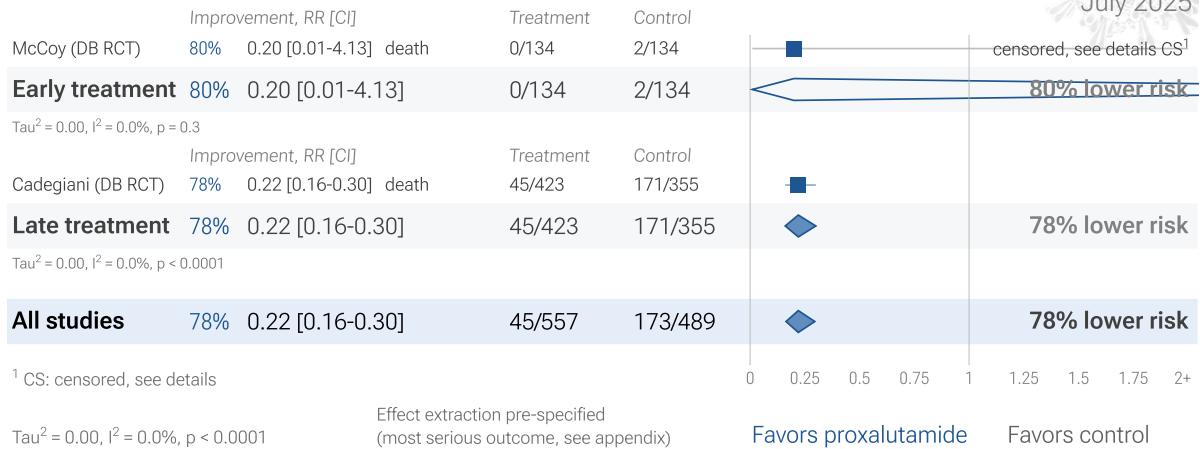


Figure 11. Random effects meta-analysis for peer reviewed studies. Effect extraction is pre-specified, using the most serious outcome reported, see the [appendix](#) for details. Analysis validating pooled outcomes for COVID-19 can be found below. Zeraatkar *et al.* analyze 356 COVID-19 trials, finding no significant evidence that preprint results are inconsistent with peer-reviewed studies. They also show extremely long peer-review delays, with a median of 6 months to journal publication. A six month delay was equivalent to around 1.5 million deaths during the first two years of the pandemic. Authors recommend using preprint evidence, with appropriate checks for potential falsified data, which provides higher certainty much earlier.

Davidson *et al.* also showed no important difference between meta analysis results of preprints and peer-reviewed publications for COVID-19, based on 37 meta analyses including 114 trials.

Randomized Controlled Trials (RCTs)

Currently all studies are RCTs.

Heterogeneity

Heterogeneity in COVID-19 studies arises from many factors including:

Treatment delay

The time between infection or the onset of symptoms and treatment may critically affect how well a treatment works. For example an antiviral may be very effective when used early but may not be effective in late stage disease, and may even be harmful. Oseltamivir, for example, is generally only considered effective for influenza when used within 0-36 or 0-48 hours^{35,36}. Baloxavir marboxil studies for influenza also show that treatment delay is critical — Ikematsu *et al.* report an 86% reduction in cases for post-exposure prophylaxis, Hayden *et al.* show a 33 hour reduction in the time to alleviation of symptoms for treatment within 24 hours and a reduction of 13 hours for treatment within 24-48 hours, and Kumar *et al.* report only 2.5 hours improvement for inpatient treatment.

Treatment delay	Result
Post-exposure prophylaxis	86% fewer cases ³⁷
<24 hours	-33 hours symptoms ³⁸
24-48 hours	-13 hours symptoms ³⁸
Inpatients	-2.5 hours to improvement ³⁹

Table 3. Studies of baloxavir marboxil for influenza show that early treatment is more effective.

Figure 12 shows a mixed-effects meta-regression for efficacy as a function of treatment delay in COVID-19 studies from 172 treatments, showing that efficacy declines rapidly with treatment delay. Early treatment is critical for COVID-19.

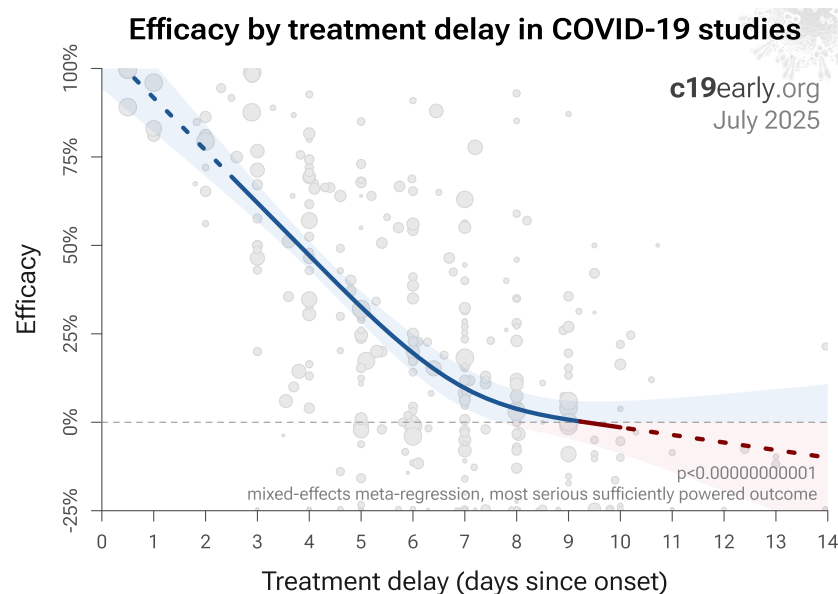


Figure 12. Early treatment is more effective. Meta-regression showing efficacy as a function of treatment delay in COVID-19 studies from 172 treatments.

Patient demographics

Details of the patient population including age and comorbidities may critically affect how well a treatment works. For example, many COVID-19 studies with relatively young low-comorbidity patients show all patients recovering quickly with or without treatment. In such cases, there is little room for an effective treatment to improve results, for example as in *López-Medina et al.*

SARS-CoV-2 variants

Efficacy may depend critically on the distribution of SARS-CoV-2 variants encountered by patients. Risk varies significantly across variants⁴¹, for example the Gamma variant shows significantly different characteristics⁴²⁻⁴⁵. Different mechanisms of action may be more or less effective depending on variants, for example the degree to which TMPRSS2 contributes to viral entry can differ across variants^{46,47}.

Treatment regimen

Effectiveness may depend strongly on the dosage and treatment regimen.

Medication quality

The quality of medications may vary significantly between manufacturers and production batches, which may significantly affect efficacy and safety. *Williams et al.* analyze ivermectin from 11 different sources, showing highly variable antiparasitic efficacy across different manufacturers. *Xu et al.* analyze a treatment from two different manufacturers, showing 9 different impurities, with significantly different concentrations for each manufacturer.

Other treatments

The use of other treatments may significantly affect outcomes, including supplements, other medications, or other interventions such as prone positioning. Treatments may be synergistic⁵⁰⁻⁶⁶, therefore efficacy may depend strongly on combined treatments.

Effect measured

Across all studies there is a strong association between different outcomes, for example improved recovery is strongly associated with lower mortality. However, efficacy may differ depending on the effect measured, for example a treatment may be more effective against secondary complications and have minimal effect on viral clearance.

Meta analysis

The distribution of studies will alter the outcome of a meta analysis. Consider a simplified example where everything is equal except for the treatment delay, and effectiveness decreases to zero or below with increasing delay. If there are many studies using very late treatment, the outcome may be negative, even though early treatment is very effective. All meta analyses combine heterogeneous studies, varying in population, variants, and potentially all factors above, and therefore may obscure efficacy by including studies where treatment is less effective. Generally, we expect the estimated effect size from meta analysis to be less than that for the optimal case. Looking at all studies is valuable for providing an overview of all research, important to avoid cherry-picking, and informative when a positive result is found despite combining less-optimal situations. However, the resulting estimate does not apply to specific cases such as early treatment in high-risk populations. While we present results for all studies, we also present treatment time and individual outcome analyses, which may be more informative for specific use cases.

Pooled Effects

Pooled effects are no longer required to show efficacy as of December 2021

This section validates the use of pooled effects for COVID-19, which enables earlier detection of efficacy, however pooled effects are no longer required for proxalutamide as of December 2021. Efficacy is now known based on specific outcomes for all studies and when restricted to RCTs.

Combining studies is required

For COVID-19, delay in clinical results translates into additional death and morbidity, as well as additional economic and societal damage. Combining the results of studies reporting different outcomes is required. There may be no mortality in a trial with low-risk patients, however a reduction in severity or improved viral clearance may translate into lower mortality in a high-risk population. Different studies may report lower severity, improved recovery, and lower mortality, and the significance may be very high when combining the results. "*The studies reported different outcomes*" is not a good reason for disregarding results. Pooling the results of studies reporting different outcomes allows us to use more of the available information. Logically we should, and do, use additional information when evaluating treatments—for example dose-response and treatment delay-response relationships provide additional evidence of efficacy that is considered when reviewing the evidence for a treatment.

Specific outcome and pooled analyses

We present both specific outcome and pooled analyses. In order to combine the results of studies reporting different outcomes we use the most serious outcome reported in each study, based on the thesis that improvement in the most serious outcome provides comparable measures of efficacy for a treatment. A critical advantage of this

approach is simplicity and transparency. There are many other ways to combine evidence for different outcomes, along with additional evidence such as dose-response relationships, however these increase complexity.

Ethical and practical issues limit high-risk trials

Trials with high-risk patients may be restricted due to ethics for treatments that are known or expected to be effective, and they increase difficulty for recruiting. Using less severe outcomes as a proxy for more serious outcomes allows faster and safer collection of evidence.

Validating pooled outcome analysis for COVID-19

For many COVID-19 treatments, a reduction in mortality logically follows from a reduction in hospitalization, which follows from a reduction in symptomatic cases, which follows from a reduction in PCR positivity. We can directly test this for COVID-19.

Analysis of the the association between different outcomes across studies from all 172 treatments we cover confirms the validity of pooled outcome analysis for COVID-19. Figure 13 shows that lower hospitalization is very strongly associated with lower mortality ($p < 0.000000000001$). Similarly, Figure 14 shows that improved recovery is very strongly associated with lower mortality ($p < 0.000000000001$). Considering the extremes, *Singh et al.* show an association between viral clearance and hospitalization or death, with $p = 0.003$ after excluding one large outlier from a mutagenic treatment, and based on 44 RCTs including 52,384 patients. Figure 15 shows that improved viral clearance is strongly associated with fewer serious outcomes. The association is very similar to *Singh et al.*, with higher confidence due to the larger number of studies. As with *Singh et al.*, the confidence increases when excluding the outlier treatment, from $p = 0.000000082$ to $p = 0.0000000033$.

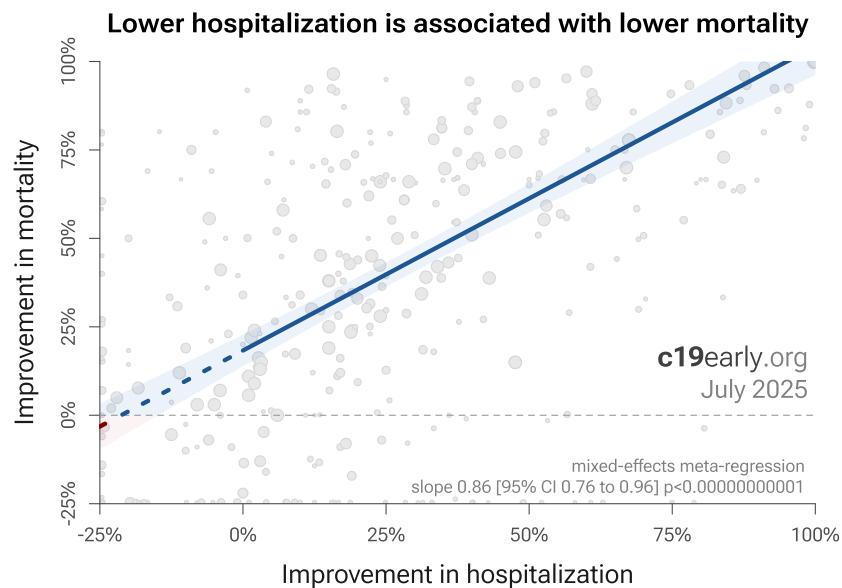


Figure 13. Lower hospitalization is associated with lower mortality, supporting pooled outcome analysis.

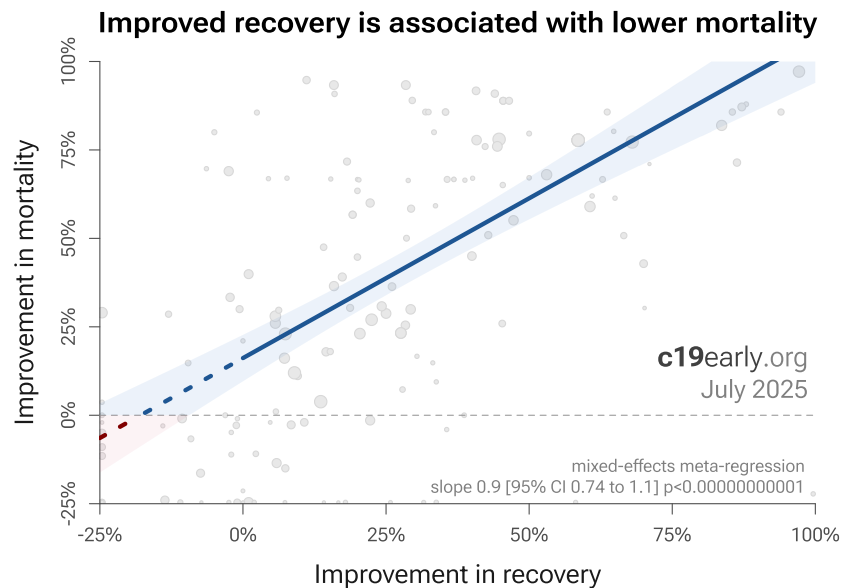


Figure 14. Improved recovery is associated with lower mortality, supporting pooled outcome analysis.

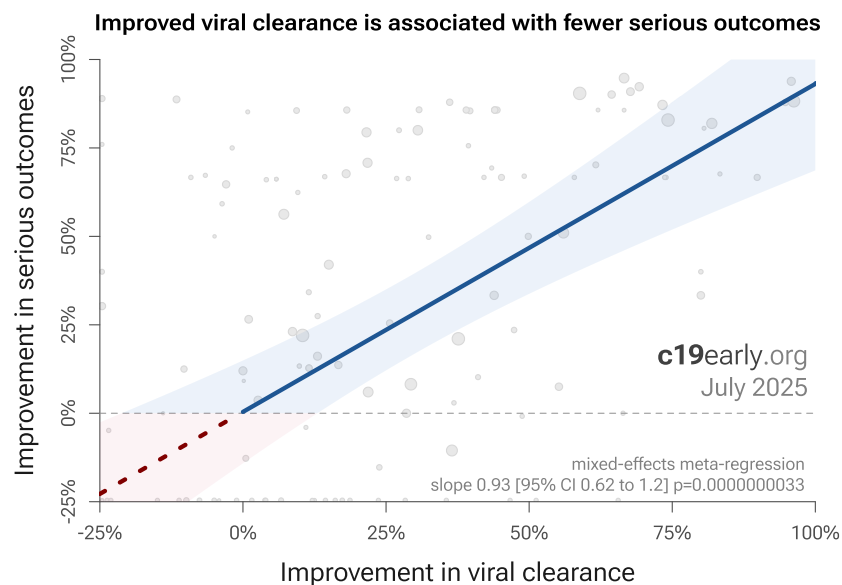


Figure 13. Improved viral clearance is associated with fewer serious outcomes, supporting pooled outcome analysis.

Pooled outcomes identify efficacy 5 months faster (7 months for RCTs)

Currently, 55 of the treatments we analyze show statistically significant efficacy or harm, defined as $\geq 10\%$ decreased risk or $> 0\%$ increased risk from ≥ 3 studies. 88% of these have been confirmed with one or more specific outcomes, with a mean delay of 4.9 months. When restricting to RCTs only, 57% of treatments showing statistically significant efficacy/harm with pooled effects have been confirmed with one or more specific outcomes, with a mean delay of 7.3 months. Figure 16 shows when treatments were found effective during the pandemic. Pooled outcomes often resulted in earlier detection of efficacy.

Time when COVID-19 studies showed efficacy

c19early.org
July 2025

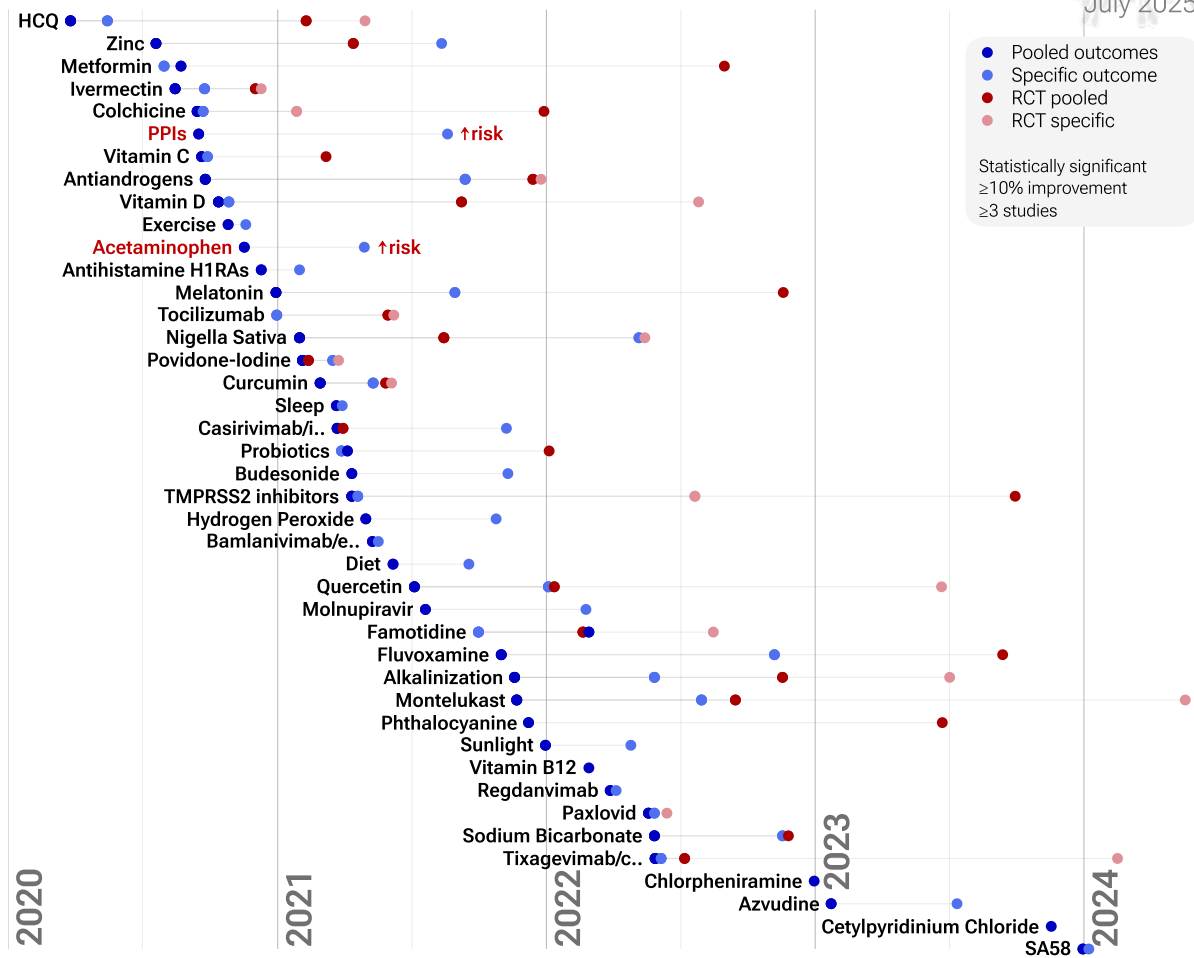


Figure 16. The time when studies showed that treatments were effective, defined as statistically significant improvement of $\geq 10\%$ from ≥ 3 studies. Pooled results typically show efficacy earlier than specific outcome results. Results from all studies often shows efficacy much earlier than when restricting to RCTs. Results reflect conditions as used in trials to date, these depend on the population treated, treatment delay, and treatment regimen.

Limitations

Pooled analysis could hide efficacy, for example a treatment that is beneficial for late stage patients but has no effect on viral clearance may show no efficacy if most studies only examine viral clearance. In practice, it is rare for a non-antiviral treatment to report viral clearance and to not report clinical outcomes; and in practice other sources of heterogeneity such as difference in treatment delay is more likely to hide efficacy.

Summary

Analysis validates the use of pooled effects and shows significantly faster detection of efficacy on average. However, as with all meta analyses, it is important to review the different studies included. We also present individual outcome analyses, which may be more informative for specific use cases.

Discussion

Publication bias

Publishing is often biased towards positive results. Trials with patented drugs may have a financial conflict of interest that results in positive studies being more likely to be published, or bias towards more positive results. For example with molnupiravir, trials with negative results remain unpublished to date (CTRI/2021/05/033864 and

CTRI/2021/08/0354242). For proxalutamide, there is currently not enough data to evaluate publication bias with high confidence.

Limitations

Summary statistics from meta analysis necessarily lose information. As with all meta analyses, studies are heterogeneous, with differences in treatment delay, treatment regimen, patient demographics, variants, conflicts of interest, standard of care, and other factors. We provide analyses for specific outcomes and by treatment delay, and we aim to identify key characteristics in the forest plots and summaries. Results should be viewed in the context of study characteristics.

Some analyses classify treatment based on early or late administration, as done here, while others distinguish between mild, moderate, and severe cases. Viral load does not indicate degree of symptoms — for example patients may have a high viral load while being asymptomatic. With regard to treatments that have antiviral properties, timing of treatment is critical — late administration may be less helpful regardless of severity.

Details of treatment delay per patient is often not available. For example, a study may treat 90% of patients relatively early, but the events driving the outcome may come from 10% of patients treated very late. Our 5 day cutoff for early treatment may be too conservative, 5 days may be too late in many cases.

Comparison across treatments is confounded by differences in the studies performed, for example dose, variants, and conflicts of interest. Trials with conflicts of interest may use designs better suited to the preferred outcome.

In some cases, the most serious outcome has very few events, resulting in lower confidence results being used in pooled analysis, however the method is simpler and more transparent. This is less critical as the number of studies increases. Restriction to outcomes with sufficient power may be beneficial in pooled analysis and improve accuracy when there are few studies, however we maintain our pre-specified method to avoid any retrospective changes.

Studies show that combinations of treatments can be highly synergistic and may result in many times greater efficacy than individual treatments alone⁵⁰⁻⁶⁶. Therefore standard of care may be critical and benefits may diminish or disappear if standard of care does not include certain treatments.

This real-time analysis is constantly updated based on submissions. Accuracy benefits from widespread review and submission of updates and corrections from reviewers. Less popular treatments may receive fewer reviews.

No treatment or intervention is 100% available and effective for all current and future variants. Efficacy may vary significantly with different variants and within different populations. All treatments have potential side effects. Propensity to experience side effects may be predicted in advance by qualified physicians. We do not provide medical advice. Before taking any medication, consult a qualified physician who can compare all options, provide personalized advice, and provide details of risks and benefits based on individual medical history and situations.

Notes

Zheng *et al.* present another meta analysis for proxalutamide, showing significant improvements for mortality, mechanical ventilation, hospitalization, and clinical improvement.

Other studies

Additional preclinical or review papers suggesting potential benefits of proxalutamide for COVID-19 include⁷³⁻⁷⁸. We have not reviewed these studies in detail.

Perspective

Results compared with other treatments

SARS-CoV-2 infection and replication involves a complex interplay of 100+ host and viral proteins and other factors²²⁻²⁹, providing many therapeutic targets. Over 9,000 compounds have been predicted to reduce COVID-19 risk³⁰, either by directly minimizing infection or replication, by supporting immune system function, or by minimizing secondary complications. Figure 17 shows an overview of the results for proxalutamide in the context of multiple COVID-19 treatments, and Figure 18 shows a plot of efficacy vs. cost for COVID-19 treatments.

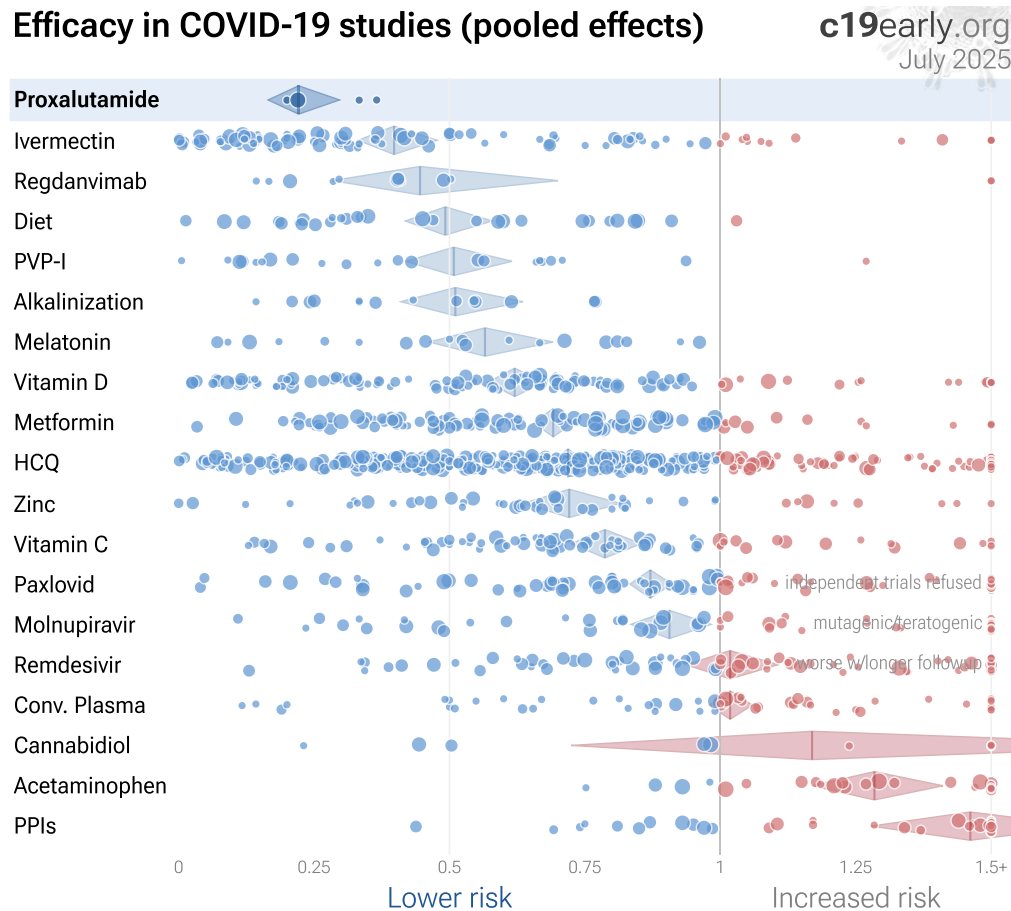


Figure 17. Scatter plot showing results within the context of multiple COVID-19 treatments. Diamonds shows the results of random effects meta-analysis. 0.6% of 9,000+ proposed treatments show efficacy⁷⁹.

Efficacy vs. cost for COVID-19 treatments

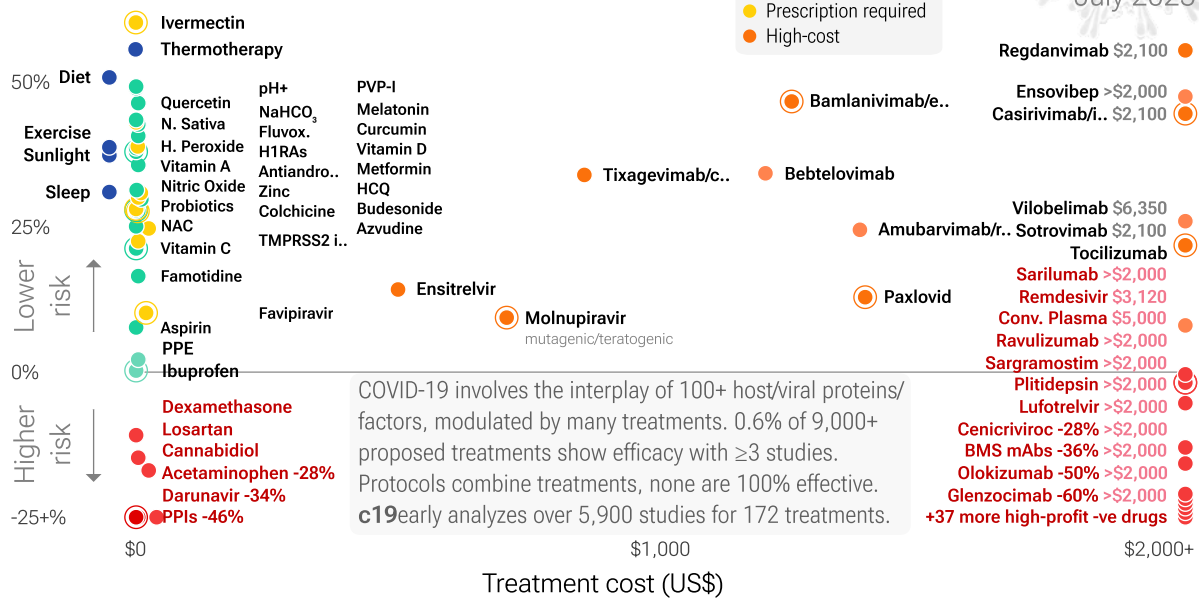


Figure 18. Efficacy vs. cost for COVID-19 treatments.

Conclusion

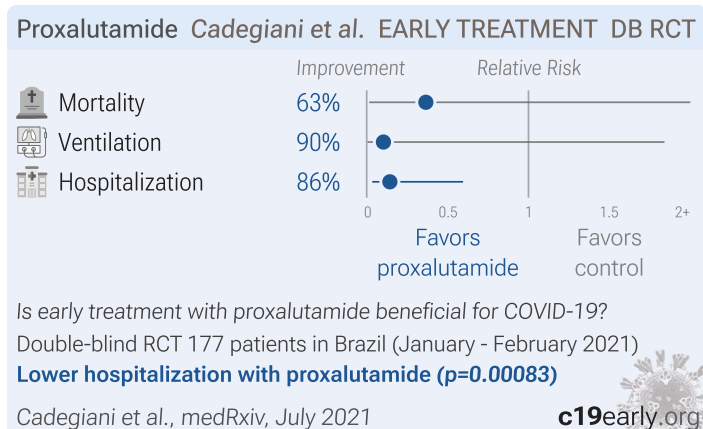
Studies to date show that proxalutamide is an effective treatment for COVID-19. Significantly lower risk is seen for mortality, ventilation, hospitalization, recovery, and viral clearance. 4 studies from 2 independent teams in 2 countries show significant benefit. Meta analysis using the most serious outcome reported shows 78% [70-83%] lower risk. Results are similar for peer-reviewed studies. Currently all studies are RCTs.

Studies to date are from only 2 different groups.

Zheng *et al.* present another meta analysis for proxalutamide, showing significant improvements for mortality, mechanical ventilation, hospitalization, and clinical improvement.

Study Notes

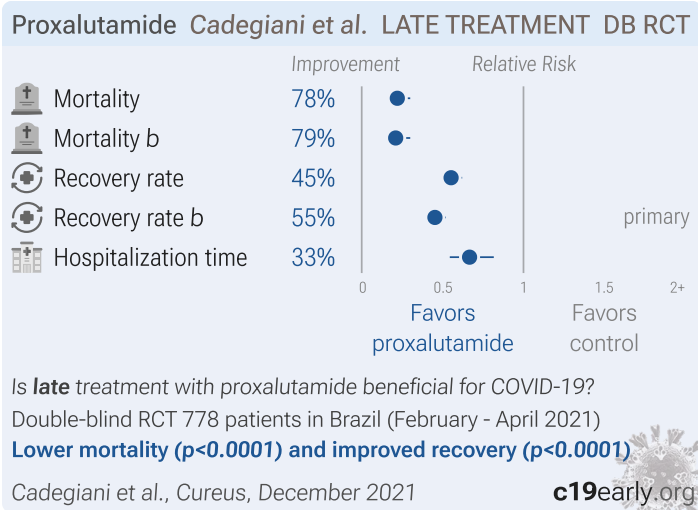
Cadegiani



SEE ALSO

RCT 177 women in Brazil, 75 treated with proxalutamide, showing significantly lower hospitalization with treatment.

Cadegiani

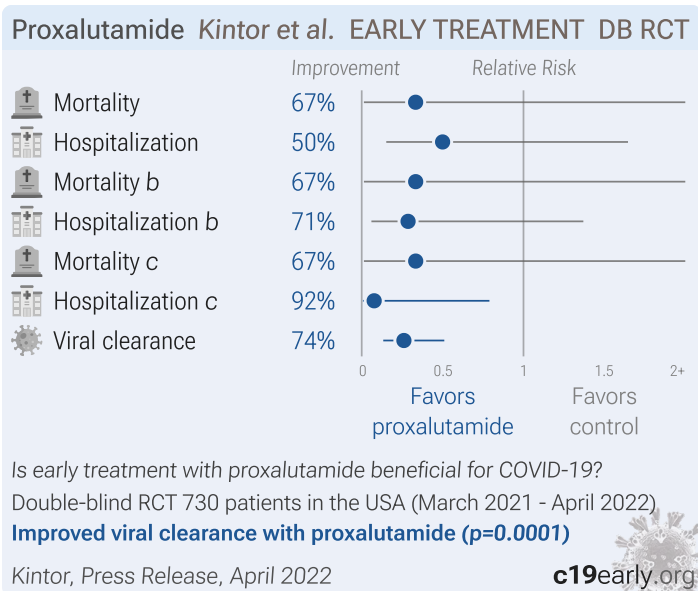


SEE ALSO

The High-Impact Medical Journal Editors Harassment Of The World's Leading Clinical Researcher of Repurposed Dru...

RCT 778 hospitalized patients in Brazil, 423 treated with proxalutamide, showing significantly lower mortality and improved recovery with treatment. NCT04728802 and NCT05126628. Authors note that cases in this trial were predominantly the P.1 Gamma variant, for which proxalutamide may be more effective compared to other variants.

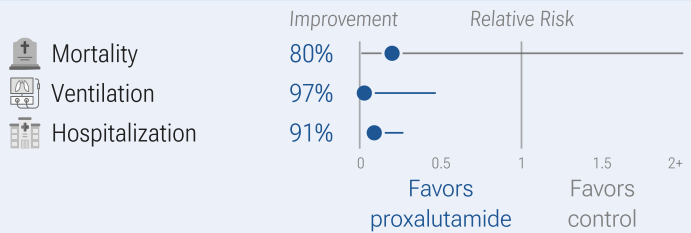
Kintor



RCT 733 outpatients, 99% in the USA, showing lower hospitalization/death, and significantly reduced viral load with proxalutamide treatment. The viral clearance result is from Ma et al.

McCoy

Proxalutamide McCoy et al. EARLY TREATMENT DB RCT



Is early treatment with proxalutamide beneficial for COVID-19?

Double-blind RCT 268 patients in Brazil (June - July 2020)

Lower ventilation ($p < 0.0001$) and hospitalization ($p < 0.0001$)

McCoy et al., *Frontiers in Medicine*, Dec 2020

c19early.org

SEE ALSO

The High-Impact Medical Journal Editors Harassment Of The World's Leading Clinical Researcher of Repurposed Dru...

RCT 268 male patients in Brazil, 134 treated with proxalutamide, showing significantly lower hospitalization and mechanical ventilation.

This paper was retracted, however no specific reason is provided, the editors have ignored the authors, and the "external expert" was reportedly funded by Pfizer. For details see⁸⁰.

The retraction notice states: "The investigation found that the claims made in the conclusions were not adequately supported by the methodology of the study. In particular, as confirmed by an external expert, the process of allocation to treatment and control was not sufficiently random."

The lack of any detail on what conclusion is not supported and why, or details of any issues in randomization, suggests the paper was censored rather than retracted.

Appendix 1. Methods and Data

We perform ongoing searches of PubMed, medRxiv, Europe PMC, ClinicalTrials.gov, The Cochrane Library, Google Scholar, Research Square, ScienceDirect, Oxford University Press, the reference lists of other studies and meta-analyses, and submissions to the site c19early.org. Search terms are proxalutamide and COVID-19 or SARS-CoV-2. Automated searches are performed twice daily, with all matches reviewed for inclusion. All studies regarding the use of proxalutamide for COVID-19 that report a comparison with a control group are included in the main analysis. Studies with major unexplained data issues, for example major outcome data that is impossible to be correct with no response from the authors, are excluded. This is a living analysis and is updated regularly.

We extracted effect sizes and associated data from all studies. If studies report multiple kinds of effects then the most serious outcome is used in pooled analysis, while other outcomes are included in the outcome specific analyses. For example, if effects for mortality and cases are reported then they are both used in specific outcome analyses, while mortality is used for pooled analysis. If symptomatic results are reported at multiple times, we use the latest time, for example if mortality results are provided at 14 days and 28 days, the results at 28 days have preference. Mortality alone is preferred over combined outcomes. Outcomes with zero events in both arms are not used, the next most serious outcome with one or more events is used. For example, in low-risk populations with no mortality, a reduction in mortality with treatment is not possible, however a reduction in hospitalization, for example, is still valuable. Clinical outcomes are considered more important than viral outcomes. When basically all patients recover in both treatment and control groups, preference for viral clearance and recovery is given to results mid-recovery where available. After most or all patients have recovered there is little or no room for an effective treatment to do better, however faster recovery is valuable. An IPD meta-analysis confirms that intermediate viral load reduction is more

closely associated with hospitalization/death than later viral load reduction⁸¹. If only individual symptom data is available, the most serious symptom has priority, for example difficulty breathing or low SpO₂ is more important than cough. When results provide an odds ratio, we compute the relative risk when possible, or convert to a relative risk according to *Zhang et al.* Reported confidence intervals and *p*-values are used when available, and adjusted values are used when provided. If multiple types of adjustments are reported propensity score matching and multivariable regression has preference over propensity score matching or weighting, which has preference over multivariable regression. Adjusted results have preference over unadjusted results for a more serious outcome when the adjustments significantly alter results. When needed, conversion between reported *p*-values and confidence intervals followed *Altman, Altman (B)*, and Fisher's exact test was used to calculate *p*-values for event data. If continuity correction for zero values is required, we use the reciprocal of the opposite arm with the sum of the correction factors equal to 1⁸⁵. Results are expressed with $RR < 1.0$ favoring treatment, and using the risk of a negative outcome when applicable (for example, the risk of death rather than the risk of survival). If studies only report relative continuous values such as relative times, the ratio of the time for the treatment group versus the time for the control group is used. Calculations are done in Python (3.13.5) with *scipy* (1.16.0), *pythonmeta* (1.26), *numpy* (2.3.1), *statsmodels* (0.14.4), and *plotly* (6.2.0).

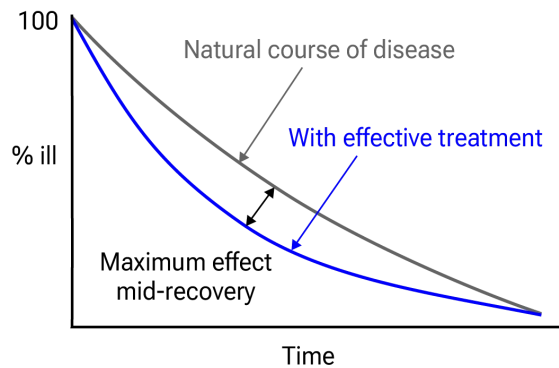


Figure 19. Mid-recovery results can more accurately reflect efficacy when almost all patients recover. *Mateja et al.* confirm that intermediate viral load results more accurately reflect hospitalization/death.

Forest plots are computed using *PythonMeta*⁸⁶ with the DerSimonian and Laird random effects model (the fixed effect assumption is not plausible in this case) and inverse variance weighting. Results are presented with 95% confidence intervals. Heterogeneity among studies was assessed using the *I*² statistic. Mixed-effects meta-regression results are computed with R (4.4.0) using the *metafor* (4.6-0) and *rms* (6.8-0) packages, and using the most serious sufficiently powered outcome. For all statistical tests, a *p*-value less than 0.05 was considered statistically significant. *Grobid* 0.8.2 is used to parse PDF documents.

We have classified studies as early treatment if most patients are not already at a severe stage at the time of treatment (for example based on oxygen status or lung involvement), and treatment started within 5 days of the onset of symptoms. If studies contain a mix of early treatment and late treatment patients, we consider the treatment time of patients contributing most to the events (for example, consider a study where most patients are treated early but late treatment patients are included, and all mortality events were observed with late treatment patients). We note that a shorter time may be preferable. Antivirals are typically only considered effective when used within a shorter timeframe, for example 0-36 or 0-48 hours for oseltamivir, with longer delays not being effective^{35,36}.

We received no funding, this research is done in our spare time. We have no affiliations with any pharmaceutical companies or political parties.

A summary of study results is below. Please submit updates and corrections at <https://c19early.org/xmeta.html>.

Early treatment

Effect extraction follows pre-specified rules as detailed above and gives priority to more serious outcomes. For pooled analyses, the first (most serious) outcome is used, which may differ from the effect a paper focuses on. Other outcomes are used in outcome specific analyses.

Cadegiani, 7/10/2021, Double Blind Randomized Controlled Trial, Brazil, preprint, 7 authors, study period 4 January, 2021 - 28 February, 2021.

risk of death, 63.4% lower, $RR\ 0.37$, $p = 1.00$, treatment 0 of 75 (0.0%), control 1 of 102 (1.0%), NNT 102, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm).

	risk of mechanical ventilation, 89.7% lower, RR 0.10, $p = 0.07$, treatment 0 of 75 (0.0%), control 5 of 102 (4.9%), NNT 20, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm).
	risk of hospitalization, 85.7% lower, RR 0.14, $p < 0.001$, treatment 2 of 75 (2.7%), control 19 of 102 (18.6%), NNT 6.3.
Kintor, 4/5/2022, Double Blind Randomized Controlled Trial, placebo-controlled, USA, preprint, 1 author, study period 5 March, 2021 - 1 April, 2022, trial NCT04870606 (history).	risk of death, 66.7% lower, RR 0.33, $p = 1.00$, treatment 0 of 365 (0.0%), control 1 of 365 (0.3%), NNT 365, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm), 1+ days of treatment, group sizes approximated.
	risk of hospitalization, 50.0% lower, RR 0.50, $p = 0.38$, treatment 4 of 365 (1.1%), control 8 of 365 (2.2%), NNT 91, 1+ days of treatment, group sizes approximated.
	risk of death, 66.6% lower, RR 0.33, $p = 1.00$, treatment 0 of 360 (0.0%), control 1 of 361 (0.3%), NNT 361, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm), >1 day of treatment, group sizes approximated.
	risk of hospitalization, 71.3% lower, RR 0.29, $p = 0.18$, treatment 2 of 360 (0.6%), control 7 of 361 (1.9%), NNT 72, >1 day of treatment, group sizes approximated.
	risk of death, 66.6% lower, RR 0.33, $p = 1.00$, treatment 0 of 346 (0.0%), control 1 of 347 (0.3%), NNT 347, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm), >7 days of treatment, group sizes approximated.
	risk of hospitalization, 92.3% lower, RR 0.08, $p = 0.03$, treatment 0 of 346 (0.0%), control 6 of 347 (1.7%), NNT 58, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm), >7 days of treatment, group sizes approximated.
	risk of no viral clearance, 73.9% lower, RR 0.26, $p < 0.001$, treatment 365, control 365, group sizes approximated, day 7.
McCoy, 12/30/2020, Double Blind Randomized Controlled Trial, Brazil, peer-reviewed, 15 authors, study period 15 June, 2020 - 28 July, 2020, censored, see details, trial NCT04446429 (history).	risk of death, 80.0% lower, RR 0.20, $p = 0.50$, treatment 0 of 134 (0.0%), control 2 of 134 (1.5%), NNT 67, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm).
	risk of mechanical ventilation, 97.1% lower, RR 0.03, $p < 0.001$, treatment 0 of 134 (0.0%), control 17 of 134 (12.7%), NNT 7.9, relative risk is not 0 because of continuity correction due to zero events (with reciprocal of the contrasting arm).
	risk of hospitalization, 91.0% lower, RR 0.09, $p < 0.001$, treatment 3 of 134 (2.2%), control 35 of 134 (26.1%), NNT 4.2.

Late treatment

Effect extraction follows pre-specified rules as detailed above and gives priority to more serious outcomes. For pooled analyses, the first (most serious) outcome is used, which may differ from the effect a paper focuses on. Other outcomes are used in outcome specific analyses.

Cadegiani (C), 12/25/2021, Double Blind Randomized Controlled Trial, Brazil, peer-reviewed, 15 authors, study period 1 February, 2021 - 15 April, 2021, trial NCT04728802 (history).	risk of death, 78.0% lower, RR 0.22, $p < 0.001$, treatment 45 of 423 (10.6%), control 171 of 355 (48.2%), NNT 2.7, adjusted per study, 28 days, Cox proportional hazards.
	risk of death, 79.0% lower, RR 0.21, $p < 0.001$, treatment 34 of 423 (8.0%), control 138 of 355 (38.9%), NNT 3.2, adjusted per study, 14 days, Cox proportional hazards.
	recovery rate, RR 0.55, $p < 0.001$, treatment 423, control 355, adjusted per study, inverted to make $RR < 1$ favor treatment, 28 days, Cox proportional hazards.
	recovery rate, RR 0.45, $p < 0.001$, treatment 423, control 355, adjusted per study, inverted to make $RR < 1$ favor treatment, 14 days, Cox proportional hazards, primary outcome.
	hospitalization time, 33.3% lower, relative time 0.67, $p < 0.001$, treatment 423, control 355.

Supplementary Data

Supplementary Data

Footnotes

- a. Viral infection and replication involves attachment, entry, uncoating and release, genome replication and transcription, translation and protein processing, assembly and budding, and release. Each step can be disrupted by therapeutics.

References

- Zheng** et al., *Small-molecule antivirals treatment for COVID-19: A systematic review and network meta-analysis*, International Journal of Antimicrobial Agents, doi:10.1016/j.ijantimicag.2024.107096.
- Ryu** et al., *Fibrin drives thromboinflammation and neuropathology in COVID-19*, Nature, doi:10.1038/s41586-024-07873-4.
- Rong** et al., *Persistence of spike protein at the skull-meninges-brain axis may contribute to the neurological sequelae of COVID-19*, Cell Host & Microbe, doi:10.1016/j.chom.2024.11.007.
- Yang** et al., *SARS-CoV-2 infection causes dopaminergic neuron senescence*, Cell Stem Cell, doi:10.1016/j.stem.2023.12.012.
- Scardua-Silva** et al., *Microstructural brain abnormalities, fatigue, and cognitive dysfunction after mild COVID-19*, Scientific Reports, doi:10.1038/s41598-024-52005-7.
- Hampshire** et al., *Cognition and Memory after Covid-19 in a Large Community Sample*, New England Journal of Medicine, doi:10.1056/NEJMoa2311330.
- Duloquin** et al., *Is COVID-19 Infection a Multiorganic Disease? Focus on Extrapulmonary Involvement of SARS-CoV-2*, Journal of Clinical Medicine, doi:10.3390/jcm13051397.
- Sodagar** et al., *Pathological Features and Neuroinflammatory Mechanisms of SARS-CoV-2 in the Brain and Potential Therapeutic Approaches*, Biomolecules, doi:10.3390/biom12070971.
- Sagar** et al., *COVID-19-associated cerebral microbleeds in the general population*, Brain Communications, doi:10.1093/braincomms/fcae127.
- Verma** et al., *Persistent Neurological Deficits in Mouse PASC Reveal Antiviral Drug Limitations*, bioRxiv, doi:10.1101/2024.06.02.596989.
- Panagea** et al., *Neurocognitive Impairment in Long COVID: A Systematic Review*, Archives of Clinical Neuropsychology, doi:10.1093/arclin/aca042.
- Ariza** et al., *COVID-19: Unveiling the Neuropsychiatric Maze —From Acute to Long-Term Manifestations*, Biomedicines, doi:10.3390/biomedicines12061147.
- Vashisht** et al., *Neurological Complications of COVID-19: Unraveling the Pathophysiological Underpinnings and Therapeutic Implications*, Viruses, doi:10.3390/v16081183.
- Ahmad** et al., *Neurological Complications and Outcomes in Critically Ill Patients With COVID-19: Results From International Neurological Study Group From the COVID-19*



- Critical Care Consortium, The Neurohospitalist, doi:10.1177/19418744241292487.
15. **Wang** et al., SARS-CoV-2 membrane protein induces neurodegeneration via affecting Golgi-mitochondria interaction, *Translational Neurodegeneration*, doi:10.1186/s40035-024-00458-1.
 16. **Eberhardt** et al., SARS-CoV-2 infection triggers pro-atherogenic inflammatory responses in human coronary vessels, *Nature Cardiovascular Research*, doi:10.1038/s44161-023-00336-5.
 17. **Van Tin** et al., Spike Protein of SARS-CoV-2 Activates Cardiac Fibrogenesis through NLRP3 Inflammasomes and NF- κ B Signaling, *Cells*, doi:10.3390/cells13161331.
 18. **Borka Balas** et al., COVID-19 and Cardiac Implications—Still a Mystery in Clinical Practice, *Reviews in Cardiovascular Medicine*, doi:10.31083/j.rcm2405125.
 19. **Altaweel** et al., An In-Depth Insight into Clinical, Cellular and Molecular Factors in COVID19-Associated Cardiovascular Ailments for Identifying Novel Disease Biomarkers, Drug Targets and Clinical Management Strategies, *Archives of Microbiology & Immunology*, doi:10.26502/ami.936500177.
 20. **Saha** et al., COVID-19 beyond the lungs: Unraveling its vascular impact and cardiovascular complications—mechanisms and therapeutic implications, *Science Progress*, doi:10.1177/00368504251322069.
 21. **Trender** et al., Changes in memory and cognition during the SARS-CoV-2 human challenge study, *eClinicalMedicine*, doi:10.1016/j.eclinm.2024.102842.
 22. **Dugied** et al., Multimodal SARS-CoV-2 interactome sketches the virus-host spatial organization, *Communications Biology*, doi:10.1038/s42003-025-07933-z.
 23. **Malone** et al., Structures and functions of coronavirus replication-transcription complexes and their relevance for SARS-CoV-2 drug design, *Nature Reviews Molecular Cell Biology*, doi:10.1038/s41580-021-00432-z.
 24. **Murigneux** et al., Proteomic analysis of SARS-CoV-2 particles unveils a key role of G3BP proteins in viral assembly, *Nature Communications*, doi:10.1038/s41467-024-44958-0.
 25. **Lv** et al., Host proviral and antiviral factors for SARS-CoV-2, *Virus Genes*, doi:10.1007/s11262-021-01869-2.
 26. **Lui** et al., Nsp1 facilitates SARS-CoV-2 replication through calcineurin-NFAT signaling, *Virology*, doi:10.1128/mbio.00392-24.
 27. **Niarakis** et al., Drug-target identification in COVID-19 disease mechanisms using computational systems biology approaches, *Frontiers in Immunology*, doi:10.3389/fimmu.2023.1282859.
 28. **Katiyar** et al., SARS-CoV-2 Assembly: Gaining Infectivity and Beyond, *Viruses*, doi:10.3390/v16111648.
 29. **Wu** et al., Decoding the genome of SARS-CoV-2: a pathway to drug development through translation inhibition, *RNA Biology*, doi:10.1080/15476286.2024.2433830.
 30. **c19early.org**, c19early.org/treatments.html.
 31. **Qiao** et al., Proxalutamide reduces SARS-CoV-2 infection and associated inflammatory response, *PNAS*, doi:10.1073/pnas.222180912.
 32. **Ma** et al., Mechanisms of action (MOA) for proxalutamide, an androgen receptor (AR) antagonist, for the treatment of mild, moderate and severe COVID-19 patients, *AACR Annual Meeting 2022*, www.abstractsonline.com/pp8/#/1/10517/presentation/18533.
 33. **Zeraatkar** et al., Consistency of covid-19 trial preprints with published reports and impact for decision making: retrospective review, *BMJ Medicine*, doi:10.1136/bmjmed-2022-0003091.
 34. **Davidson** et al., No evidence of important difference in summary treatment effects between COVID-19 preprints and peer-reviewed publications: a meta-epidemiological study, *Journal of Clinical Epidemiology*, doi:10.1016/j.jclinepi.2023.08.011.
 35. **Treanor** et al., Efficacy and Safety of the Oral Neuraminidase Inhibitor Oseltamivir in Treating Acute Influenza: A Randomized Controlled Trial, *JAMA*, 2000, 283:8, 1016-1024, doi:10.1001/jama.283.8.1016.
 36. **McLean** et al., Impact of Late Oseltamivir Treatment on Influenza Symptoms in the Outpatient Setting: Results of a Randomized Trial, *Open Forum Infect. Dis.* September 2015, 2:3, doi:10.1093/ofid/ofv100.
 37. **Ikematsu** et al., Baloxavir Marboxil for Prophylaxis against Influenza in Household Contacts, *New England Journal of Medicine*, doi:10.1056/NEJMoa1915341.
 38. **Hayden** et al., Baloxavir Marboxil for Uncomplicated Influenza in Adults and Adolescents, *New England Journal of Medicine*, doi:10.1056/NEJMoa1716197.
 39. **Kumar** et al., Combining baloxavir marboxil with standard-of-care neuraminidase inhibitor in patients hospitalised with severe influenza (FLAGSTONE): a randomised, parallel-group, double-blind, placebo-controlled, superiority trial, *The Lancet Infectious Diseases*, doi:10.1016/S1473-3099(21)00469-2.
 40. **López-Medina** et al., Effect of Ivermectin on Time to Resolution of Symptoms Among Adults With Mild COVID-19: A Randomized Clinical Trial, *JAMA*, doi:10.1001/jama.2021.3071.
 41. **Korves** et al., SARS-CoV-2 Genetic Variants and Patient Factors Associated with Hospitalization Risk, *medRxiv*, doi:10.1101/2024.03.08.24303818.
 42. **Faria** et al., Genomics and epidemiology of the P.1 SARS-CoV-2 lineage in Manaus, Brazil, *Science*, doi:10.1126/science.abh2644.
 43. **Nonaka** et al., SARS-CoV-2 variant of concern P.1 (Gamma) infection in young and middle-aged patients admitted to the intensive care units of a single hospital in Salvador, Northeast Brazil, February 2021, *International Journal of Infectious Diseases*, doi:10.1016/j.ijid.2021.08.003.
 44. **Karita** et al., Trajectory of viral load in a prospective population-based cohort with incident SARS-CoV-2 G614 infection, *medRxiv*, doi:10.1101/2021.08.27.21262754.
 45. **Zavascki** et al., Advanced ventilatory support and mortality in hospitalized patients with COVID-19 caused by Gamma (P.1) variant of concern compared to other lineages: cohort study at a reference center in Brazil, *Research Square*, doi:10.21203/rs.3.rs-910467/v1.

46. **Willett** et al., The hyper-transmissible SARS-CoV-2 Omicron variant exhibits significant antigenic change, vaccine escape and a switch in cell entry mechanism, medRxiv, doi:10.1101/2022.01.03.21268111.
47. **Peacock** et al., The SARS-CoV-2 variant, Omicron, shows rapid replication in human primary nasal epithelial cultures and efficiently uses the endosomal route of entry, bioRxiv, doi:10.1101/2021.12.31.474653.
48. **Williams**, T., Not All Ivermectin Is Created Equal: Comparing The Quality of 11 Different Ivermectin Sources, Do Your Own Research, doyourownresearch.substack.com/p/not-all-ivermectin-is-created-equal.
49. **Xu** et al., A study of impurities in the repurposed COVID-19 drug hydroxychloroquine sulfate by UHPLC-Q/TOF-MS and LC-SPE-NMR, Rapid Communications in Mass Spectrometry, doi:10.1002/rcm.9358.
50. **Jitobaom** et al., Favipiravir and Ivermectin Showed in Vitro Synergistic Antiviral Activity against SARS-CoV-2, Research Square, doi:10.21203/rs.3.rs-941811/v1.
51. **Jitobaom (B)** et al., Synergistic anti-SARS-CoV-2 activity of repurposed anti-parasitic drug combinations, BMC Pharmacology and Toxicology, doi:10.1186/s40360-022-00580-8.
52. **Jeffreys** et al., Remdesivir-ivermectin combination displays synergistic interaction with improved in vitro activity against SARS-CoV-2, International Journal of Antimicrobial Agents, doi:10.1016/j.ijantimicag.2022.106542.
53. **Ostrov** et al., Highly Specific Sigma Receptor Ligands Exhibit Anti-Viral Properties in SARS-CoV-2 Infected Cells, Pathogens, doi:10.3390/pathogens10111514.
54. **Alsaïdi** et al., Griffithsin and Carrageenan Combination Results in Antiviral Synergy against SARS-CoV-1 and 2 in a Pseudoviral Model, Marine Drugs, doi:10.3390/md19080418.
55. **Andreani** et al., In vitro testing of combined hydroxychloroquine and azithromycin on SARS-CoV-2 shows synergistic effect, Microbial Pathogenesis, doi:10.1016/j.micpath.2020.104228.
56. **De Forni** et al., Synergistic drug combinations designed to fully suppress SARS-CoV-2 in the lung of COVID-19 patients, PLoS ONE, doi:10.1371/journal.pone.0276751.
57. **Wan** et al., Synergistic inhibition effects of andrographolide and baicalin on coronavirus mechanisms by downregulation of ACE2 protein level, Scientific Reports, doi:10.1038/s41598-024-54722-5.
58. **Said** et al., The effect of *Nigella sativa* and vitamin D3 supplementation on the clinical outcome in COVID-19 patients: A randomized controlled clinical trial, Frontiers in Pharmacology, doi:10.3389/fphar.2022.1011522.
59. **Fiaschi** et al., In Vitro Combinatorial Activity of Direct Acting Antivirals and Monoclonal Antibodies against the Ancestral B.1 and BQ.1.1 SARS-CoV-2 Viral Variants, Viruses, doi:10.3390/v16020168.
60. **Xing** et al., Published anti-SARS-CoV-2 in vitro hits share common mechanisms of action that synergize with antivirals, Briefings in Bioinformatics, doi:10.1093/bib/bbab249.
61. **Chen** et al., Synergistic Inhibition of SARS-CoV-2 Replication Using Disulfiram/Ebselen and Remdesivir, ACS Pharmacology & Translational Science, doi:10.1021/acscptsci.1c00022.
62. **Hempel** et al., Synergistic inhibition of SARS-CoV-2 cell entry by otamixaban and covalent protease inhibitors: pre-clinical assessment of pharmacological and molecular properties, Chemical Science, doi:10.1039/D1SC01494C.
63. **Schultz** et al., Pyrimidine inhibitors synergize with nucleoside analogues to block SARS-CoV-2, Nature, doi:10.1038/s41586-022-04482-x.
64. **Ohashi** et al., Potential anti-COVID-19 agents, cepharanthine and nelfinavir, and their usage for combination treatment, iScience, doi:10.1016/j.isci.2021.102367.
65. **Al Krad** et al., The protease inhibitor Nirmatrelvir synergizes with inhibitors of GRP78 to suppress SARS-CoV-2 replication, bioRxiv, doi:10.1101/2025.03.09.642200.
66. **Thairu** et al., A Comparison of Ivermectin and Non Ivermectin Based Regimen for COVID-19 in Abuja: Effects on Virus Clearance, Days-to-discharge and Mortality, Journal of Pharmaceutical Research International, doi:10.9734/jpri/2022/v34i44A36328.
67. **Singh** et al., The relationship between viral clearance rates and disease progression in early symptomatic COVID-19: a systematic review and meta-regression analysis, Journal of Antimicrobial Chemotherapy, doi:10.1093/jac/dkac045.
68. **Kintor**, Kintor Pharma's Proxalutamide Demonstrated Reduction in Hospitalization/Mortality for Patients with Mild to Moderate COVID-19 in Phase III MRCT Study, Press Release, www.prnewswire.com/news-releases/kintor-pharmas-proxalutamide-demonstrated-reduction-in-hospitalization-mortality-for-patients-with-mild-to-moderate-covid-19-in-phase-iii-mrct-study-301518525.html.
69. **Cadegiani** et al., Proxalutamide (GT0918) Reduces the Rate of Hospitalization in mild-to-moderate COVID-19 Female Patients: A Randomized Double-Blinded Placebo-Controlled Two-Arm Parallel Trial, medRxiv, doi:10.1101/2021.07.06.21260086.
70. **Cadegiani (B)** et al., Proxalutamide Significantly Accelerates Viral Clearance and Reduces Time to Clinical Remission in Patients with Mild to Moderate COVID-19: Results from a Randomized, Double-Blinded, Placebo-Controlled Trial, Cureus, doi:10.7759/cureus.13492.
71. **McCoy** et al., Proxalutamide (GT0918) Reduces the Rate of Hospitalization for COVID-19 Male Outpatients: A Randomized Double-Blinded Placebo-Controlled Trial, Frontiers in Medicine, doi:10.3389/fmed.2021.668698.
72. **Cadegiani (C)** et al., Final Results of a Randomized, Placebo-Controlled, Two-Arm, Parallel Clinical Trial of Proxalutamide for Hospitalized COVID-19 Patients: A Multiregional, Joint Analysis of the Proxa-Rescue AndroCoV Trial, Cureus, doi:10.7759/cureus.20691.
73. **Baby** et al., Exploring TMPRSS2 Drug Target to Combat Influenza and Coronavirus Infection, Scientifica, doi:10.1155/sci5/3687892.

74. **Ponnampalli** et al., COVID-19: Vaccines and therapeutics, Bioorganic & Medicinal Chemistry Letters, doi:10.1016/j.bmcl.2022.128987.
75. **Guo** et al., Multi-omics in COVID-19: Driving development of therapeutics and vaccines, National Science Review, doi:10.1093/nsr/nwad161.
76. **Oliver** et al., Different drug approaches to COVID-19 treatment worldwide: an update of new drugs and drugs repositioning to fight against the novel coronavirus, Therapeutic Advances in Vaccines and Immunotherapy, doi:10.1177/25151355221144845.
77. **Yang (B)** et al., Bench-to-bedside: Innovation of small molecule anti-SARS-CoV-2 drugs in China, European Journal of Medicinal Chemistry, doi:10.1016/j.ejmech.2023.115503.
78. **Zhong** et al., Recent advances in small-molecular therapeutics for COVID-19, Precision Clinical Medicine, doi:10.1093/pcmedi/pbac024.
79. **c19early.org (B)**, c19early.org/timeline.html.
80. **twitter.com**, twitter.com/FlavioCadegiani/status/1534716400073326592.
81. **Mateja** et al., The choice of viral load endpoint in early phase trials of COVID-19 treatments aiming to reduce 28-day hospitalization and/or death, The Journal of Infectious Diseases, doi:10.1093/infdis/jiaf282.
82. **Zhang** et al., What's the relative risk? A method of correcting the odds ratio in cohort studies of common outcomes, JAMA, 80:19, 1690, doi:10.1001/jama.280.19.1690.
83. **Altman, D.**, How to obtain the P value from a confidence interval, BMJ, doi:10.1136/bmj.d2304.
84. **Altman (B)** et al., How to obtain the confidence interval from a P value, BMJ, doi:10.1136/bmj.d2090.
85. **Sweeting** et al., What to add to nothing? Use and avoidance of continuity corrections in meta-analysis of sparse data, Statistics in Medicine, doi:10.1002/sim.1761.
86. **Deng, H.**, PyMeta, Python module for meta-analysis, www.pymeta.com/.