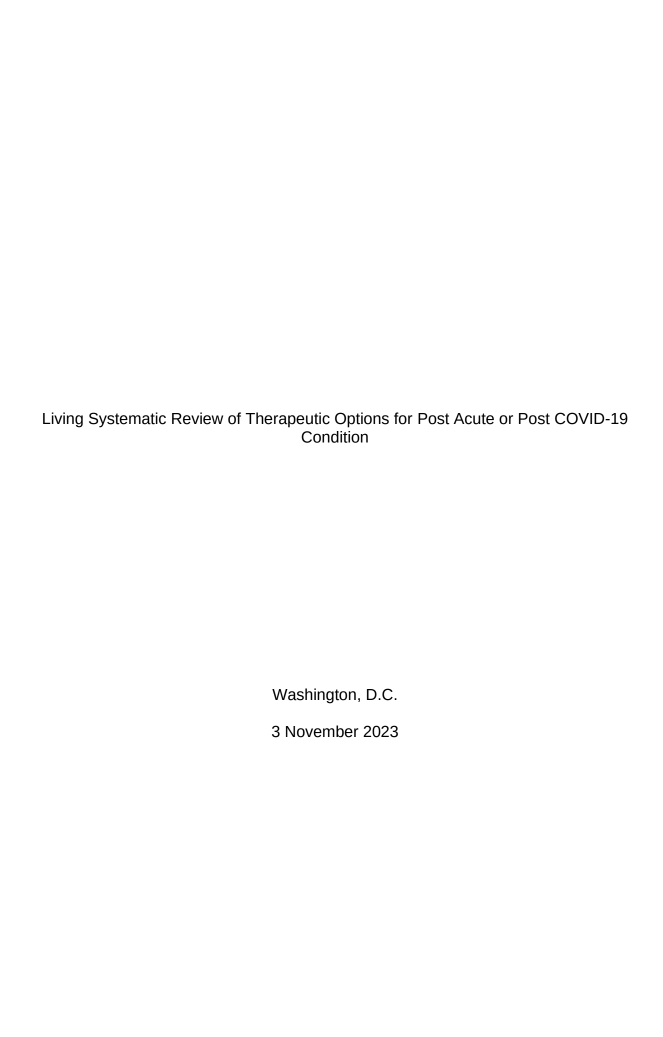


OF THERAPEUTIC OPTIONS FOR POST-ACUTE AND POST-COVID19 CONDITION

November 3 2023



Living Systematic Review of Therapeutic Options for Post Acute or Post COVID-19 Condition, 3 November 2023

PAHO/IMS/EIH/COVID-19/23-0026

© Pan American Health Organization, 2023

Some rights reserved. This work is available under the Creative Commons Attribution-NonCommercial-ShareAlike 3.0 IGO license (CC BY-NC-SA 3.0 IGO).

Under the terms of this license, this work may be copied, redistributed, and adapted for non-commercial purposes, provided the new work is issued using the same or equivalent Creative Commons license and it is appropriately cited. In any use of this work, there should be no suggestion that the Pan American Health Organization (PAHO) endorses any specific organization, product, or service. Use of the PAHO logo is not permitted.

All reasonable precautions have been taken by PAHO to verify the information contained in this publication. However, the published material is being distributed without warranty of any kind, either expressed or implied. The responsibility for the interpretation and use of the material lies with the reader. In no event shall PAHO be liable for damages arising from its use.

This document includes the results of a rapid systematic review of current available literature. The information included in this review reflects the evidence as of the date posted in the document. In recognition of the fact that there are numerous ongoing clinical studies, PAHO will periodically update this review and corresponding recommendations as new evidence becomes available.

Contents

| Acknowledgments | 6 |
|---|----|
| Funding | 6 |
| Executive summary | 7 |
| Summary of evidence | 7 |
| P-ACC-related asthenia or fatigue | 8 |
| P-ACC-related dyspnea | 12 |
| P-ACC-related neurocognitive symptoms or sleep disturbances | 14 |
| P-ACC-related olfactory and/or gustatory dysfunction | 17 |
| P-ACC-related cardiovascular system symptoms | 20 |
| P-ACC-related psychological distress | 21 |
| P-ACC-related thromboembolic risk | 22 |
| Pediatric inflammatory multisystem syndrome associated with SARS-CoTS) | |
| P-ACC prophylaxis | 24 |
| Changes since previous edition | 26 |
| Concluding remarks | 26 |
| Systematic review of therapeutic options for post acute or post COVID-19 ACC) | |
| Background | 28 |
| Methods | 29 |
| Search strategy | 29 |
| Study selection | 29 |
| Inclusion criteria | 30 |
| Living evidence synthesis | 30 |
| Results | 33 |
| Studies identified and included | 33 |
| Risk of bias | 34 |
| Main findings | 36 |
| P-ACC-related asthenia or fatigue | 36 |
| P-ACC-related dyspnea | 42 |
| P-ACC-related neurocognitive symptoms or sleep disturbances | 45 |
| P-ACC-related olfactory and/or gustatory dysfunction | 47 |

| P-ACC-related cardiovascular system symptoms | 50 |
|--|-----|
| P-ACC-related psychological distress | 51 |
| P-ACC-related thromboembolic risk | 52 |
| Full description of included studies | 56 |
| References | 121 |
| Annex 1. Summary of findings tables | 132 |
| | |

Acknowledgments

This document was developed by Ariel Izcovich, Martin Ragusa, Fernando Tortosa, Sasha Peiris, and Ludovic Reveiz from the knowledge Translation Program, Department of Evidence and Intelligence for Action in Health, and the Incident Management System for the response to COVID-19, Pan American Health Organization (PAHO). It was strengthened with the valuable contributions of: Pedro Ordunez, Antony Duttine and Carmen Martinez from the Noncommunicable Diseases and Mental Health department, PAHO, and Sylvain Aldighieri, Deputy Director of the Health Emergencies and Incident Management System for the response to COVID-19.

Funding

This work was partially funded by the Government of the United States of America.

Executive summary

Background

Post COVID-19 condition (PCC), also known as long COVID or post-acute sequelae of SARS-CoV-2 infection (PASC), is the continuation or development of new symptoms in the period after acute infection with SARS-CoV-2. The World Health Organization (WHO) definition of PCC states that these symptoms should be present after three months of the initial SARS-CoV-2 infection and last for at least two months with no other explanation. While PASC definitions state that persistent or new symptoms need to be present 30 days after a documented SARS-COV-2 infection or the onset of COVID-19 symptoms, post COVID-19 condition or post-acute sequelae of SARS-CoV-2 infection (P-ACC) can affect anyone exposed to SARS-CoV-2, regardless of age or severity of acute infection. Many of the reported symptoms are debilitating and have a strong negative impact on mental health and quality of life. While most patients recover, some may experience multiple outcomes, with multiple organ systems affected simultaneously, including cardiovascular, mental, metabolic, renal, and others.

This review compiles the following evidence on potential therapeutic options for P-ACC. It includes all the identified clinical forms, symptoms, and manifestations of P-ACC for which an intervention was assessed in at least one randomized controlled trial (RCT). It is hoped this information will support investigators, policymakers, and prescribers navigate the flood of relevant data to ensure that management of P-ACC, at both the individual and population levels, is based on the best available knowledge. This resource will be continually updated as more research is released into the public space.

Summary of evidence

All odd numbered tables (Table ES1 to ES15) present RCTs according to the reported P-ACC related organ/system affected and indicate the primary outcome measures used for each investigation and the level of certainty. The even numbered tables (Table ES2 to

ES16) summarize the status of evidence for the 51 potential therapeutic options for P-ACC for which studies were identified through this systematic review.

P-ACC-related asthenia or fatigue

Table ES1. List of RCTs on interventions for P-ACC-related asthenia or fatigue with primary outcome measures and certainty (n=36)

| Intervention | | Overall number of studies including the intervention, n=36 | HRQL improvement (n of studies) | Overall symptom improvement (n of studies) | Fatigue improvement (n of studies) | Functional capacity improvement (n of studies) | Strength improvement (n of studies) | Adverse events (n of studies) | Severe adverse events (n of studies) |
|--------------------------------------|-----|---|---------------------------------------|---|--|---|---|-------------------------------------|--|
| Physical training | NEW | 8 | 3 | | 4 | 5 | 2 | | |
| Telerehabilitation | | 5 | 4 | | 3 | 4 | | | |
| Fermented food supplements | | 2 | 2 | 2 | | | | | |
| tDCS | | 2 | 1 | | 2 | | | 1 | |
| 1_MNA | | 1 | | | 1 | 1 | | | |
| Actovegin | | 1 | | | 1 | | | | |
| ADAPT_232 (adaptogens) | | 1 | | | 1 | | | | |
| Amygdala and Insula Retraining (AIR) | | 1 | 1 | | 1 | | | | |
| Arginine_Vitamin C | | 1 | | | 1 | | 1 | | |
| Aromatherapy | | 1 | 1 | | 1 | | | | |
| AXA1125 | | 1 | | | 1 | 1 | | 1 | |
| CCSA | NEW | 1 | | | 1 | | | | |
| Cognitive behavioral therapy | | 1 | | | 1 | 1 | | | |
| CQ10 | | 1 | 1 | 1 | | | | | |
| Cytoflavin | | 1 | | | 1 | | | | |
| Echinochrome A | | 1 | | | 1 | | | | |
| Enzimes_Probiotics | | 1 | | | 1 | | | | |
| Hydrogen (nasal) | | 1 | | | 1 | 1 | | | |
| Immunodaat | | 1 | 1 | | 1 | | | | |
| Mindfulness | | 1 | | | 1 | | | | |
| Phytochemicals | | 1 | 1 | | 1 | | | | |
| Probiotics | | 1 | | | 1 | | | | |
| Leronlimab | | 1 | | 1 | | | | | |



Table ES2. Summary of findings on potential therapeutic options for P-ACC-related asthenia or fatigue (n=23), as of 3 November 2023

| | Intervention | Summary of findings |
|----|---|--|
| | | |
| 1 | 1-MNA | Uncertainty in potential benefits and harms. Further research is needed. |
| 2 | Actovegin | Actovegin may improve fatigue. However, certainty of the evidence was low. Further research is needed. |
| 3 | ADAPT-232 (adaptogens) | ADAPT-232 may not improve fatigue. However, certainty of the evidence was low. Further research is needed. |
| 4 | Amygdala and Insula Retraining (AIR) | Uncertainty in potential benefits and harms. Further research is needed. |
| 5 | Arginine + Vitamin C | Uncertainty in potential benefits and harms. Further research is needed. |
| 6 | Aromatherapy | Uncertainty in potential benefits and harms. Further research is needed. |
| 7 | AXA1125 | AXA1125 may increase fatigue improvement but may not increase functional capacity improvement. However, certainty of the evidence was low. Further research is needed. |
| 8 | CCSA | Uncertainty in potential benefits and harms. Further research is needed. |
| 9 | Coenzyme Q10 | Uncertainty in potential benefits and harms. Further research is needed. |
| 10 | Cognitive behavioral therapy | Cognitive behavioral therapy may increase fatigue improvement and functional capacity improvement. Further research is needed. |
| 11 | Cytoflavin | Cytoflavin may not improve fatigue. However, certainty of the evidence was low. Further research is needed. |
| 12 | Echinochrome A | Uncertainty in potential benefits and harms. Further research is needed. |
| 13 | Enzymes + probiotics | Enzymes + probiotics may improve fatigue. However, certainty of the evidence was low. Further research is needed. |

| | Intervention | Summary of findings |
|----|--|--|
| | | |
| 14 | Fermented food supplements | Uncertainty in potential benefits and harms. Further research is needed. |
| 15 | Hydrogen (nasal) | Uncertainty in potential benefits and harms. Further research is needed. |
| 16 | Immunodaat | Uncertainty in potential benefits and harms. Further research is needed. |
| 17 | Leronlimab | Uncertainty in potential benefits and harms. Further research is needed. |
| 18 | Mindfulness | Uncertainty in potential benefits and harms. Further research is needed. |
| 19 | Phytochemicals | Phytochemicals may improve fatigue and HRQL. However, certainty of the evidence was low. Further research is needed. |
| 20 | Physical training | Physical training may improve HRQL, strength and functional capacity. However, certainty of the evidence was low. Further research is needed. |
| 21 | Probiotics | Uncertainty in potential benefits and harms. Further research is needed. |
| 22 | Transcranial direct current stimulation (tDCS) | tDCS may improve fatigue and HRQL and may not increase adverse events. However, certainty of the evidence was low. Further research is needed. |
| 23 | Telerehabilitation | Telerehabilitation may improve fatigue and functional capacity. However, certainty of the evidence was low. Further research is needed. |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined 22 therapeutic options for P-ACC-related asthenia or fatigue.
- **Actovegin:** The results of one RCT suggest that actovegin may improve fatigue. However, certainty of the evidence was low because of imprecision. Further research is needed.

- ADAPT-232 (adaptogens): The results of one RCT suggest that ADAPT-232 may not improve fatigue. However, certainty of the evidence was low because of imprecision. Further research is needed.
- AXA1125 (amino acids + N-acetylcysteine): AXA1125 may increase fatigue improvement but may not increase functional capacity improvement. However, certainty of the evidence was low because of imprecision. Further research is needed.
- Cytoflavin: The results of one RCT suggest that cytoflavin may not improve fatigue. However, certainty of the evidence was low because of imprecision and risk of bias. Further research is needed.
- Enzymes + probiotics: The results of one RCT suggest that enzymes + probiotics may not improve fatigue. However, certainty of the evidence was low because of imprecision and risk of bias. Further research is needed.
- Physical training: The results of eight RCTs suggest that physical training may improve HRQL, strength, and functional capacity. However, certainty of the evidence was low because of imprecision and risk of bias. Two studies compared different training modalities but provided very low certainty evidence. Further research is needed.
- Transcranial direct current stimulation (tDCS): The results of two RCTs suggest that tDCS may improve fatigue and HRQL and may not increase adverse events. However, certainty of the evidence was low because of imprecision. Further research is needed.
- **Telerehabilitation:** The results of five RCTs suggest that telerehabilitation may improve fatigue and functional capacity. However, certainty of the evidence was low because of imprecision and risk of bias. Further research is needed.

P-ACC-related dyspnea

Table ES3. List of RCTs of interventions for P-ACC-related dyspnea with primary outcome measures and certainty (n=20)

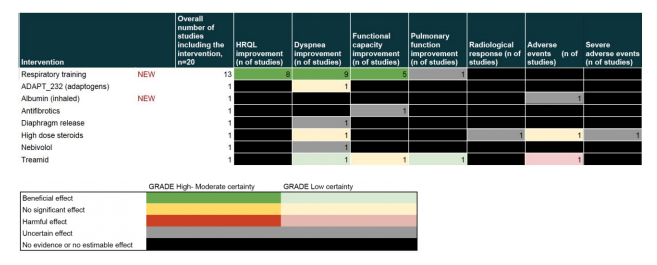


Table ES4. Summary of findings on potential therapeutic options for P-ACC-related dyspnea (n=8), as of 3 November 2023

| | Intervention | Summary of findings |
|---|------------------------|--|
| | | |
| 1 | ADAPT-232 (adaptogens) | ADAPT-232 may not improve dyspnea. However, certainty of the evidence was low. Further research is needed. |
| 2 | Albumin (inhaled) | Uncertainty in potential benefits and harms. Further research is needed. |
| 3 | Antifibrotics | Uncertainty in potential benefits and harms. Further research is needed. |
| 4 | Diaphragm release | Uncertainty in potential benefits and harms. Further research is needed. |
| 5 | High dose steroids | High dose steroids, compared to standard dose steroids, may not improve dyspnea and may not increase adverse events. However, certainty of the evidence was low. Further research is needed. |
| 6 | Nebivolol | Uncertainty in potential benefits and harms. Further research is needed. |

| | Intervention | Summary of findings |
|---|--|--|
| | | |
| 7 | Respiratory training/rehabilitation | Respiratory training/rehabilitation probably improves HRQL, dyspnea, and functional capacity |
| 8 | Treamid | Treamid may improve dyspnea and pulmonary function but may not improve functional capacity. Treamid may increase adverse events. However, certainty of the evidence was low. Further research is needed. |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined seven therapeutic options for P-ACC-related dyspnea.
- ADAPT-232 (adaptogens): The results of one RCT suggest that ADAPT-232 may not improve dyspnea. However, certainty of the evidence was low because of imprecision. Further research is needed.
- **High dose steroids**: The results of one RCT suggest that high dose steroids (prednisone 40 mg a day) may not improve dyspnea compared to standard dose steroids (prednisone 10 mg a day). However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
- Respiratory training/rehabilitation: The results of ten RCTs suggest that respiratory training probably improves HRQL, dyspnea and functional capacity.
- **Treamid:** The results of one RCT suggest that treamid may improve dyspnea and pulmonary function but may not improve functional capacity. However, certainty of the evidence was low because of imprecision. Further research is needed.

P-ACC-related neurocognitive symptoms or sleep disturbances

Table ES5. List of RCTs of interventions for P-ACC-related neurocognitive symptoms or sleep disturbances with primary outcome measures and certainty (n=10)

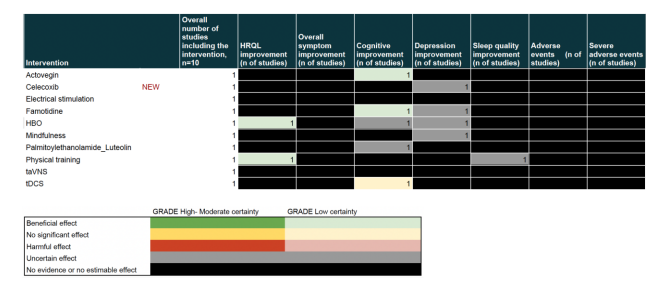


Table ES6. Summary of findings on potential therapeutic options for P-ACC-related neurocognitive symptoms or sleep disturbances (n=10), as of 3 November 2023

| | Intervention | Summary of findings |
|---|-------------------------|--|
| | | |
| 1 | Actovegin | Actovegin may improve cognition. However, certainty of the evidence was low. Further research is needed. |
| 2 | Celecoxib | Uncertainty in potential benefits and harms. Further research is needed. |
| 3 | Electric stimulation | Uncertainty in potential benefits and harms. Further research is needed. |
| 4 | Famotidine | Famotidine may improve cognition. However, certainty of the evidence was low. Further research is needed |
| 5 | Hyperbaric oxygen (HBO) | HBO may improve HRQL. However, certainty of the evidence was low. Further research is needed. |

| | Intervention | Summary of findings |
|----|--|---|
| | | |
| 6 | Mindfulness | Uncertainty in potential benefits and harms. Further research is needed. |
| 7 | Palmitoylethanolamide + Luteolin | Uncertainty in potential benefits and harms. Further research is needed. |
| 8 | Physical training | Uncertainty in potential benefits and harms. Further research is needed. |
| 9 | Transcutaneous auricular vagus nerve stimulation (taVNS) | Uncertainty in potential benefits and harms. Further research is needed. |
| 10 | Transcranial direct current stimulation (tDCS) | tCDS may not improve cognition. However, certainty of the evidence was low. Further research is needed. |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined nine therapeutic options for PCC neurocognitive symptoms or sleep disturbances.
- Actovegin: The results of one RCT suggest that actovegin may improve cognition. However, certainty of the evidence was low because of risk of bias. Further research is needed.
- Famotidine: The results of one RCT suggest that famotidine may improve cognition. However, certainty of the evidence was low because of imprecision. Further research is needed.
- **Hyperbaric oxygen (HBO):** The results of one RCT suggest that HBO may improve HRQL. However, certainty of the evidence was low because of imprecision. Further research is needed.

• Transcranial direct current stimulation (tDCS): The results of one RCT suggest that tDCS may not improve cognition. However, certainty of the evidence was low because of imprecision. Further research is needed.

P-ACC-related olfactory and/or gustatory dysfunction

Table ES7. List of RCTs of interventions for P-ACC-related olfactory and/or gustatory dysfunction with primary outcome measures and certainty (n=18)



Table ES8. Summary of findings on potential therapeutic options for P-ACC-related olfactory and/or gustatory dysfunction (n=12), as of 3 November 2023

| | Intervention | Summary of findings |
|---|------------------------|---|
| | | |
| 1 | ADAPT-232 (adaptogens) | ADAPT-232 may not improve olfactory symptoms. However, certainty of the evidence was low. Further research is needed. |
| 2 | Diode laser | Uncertainty in potential benefits and harms. Further research is needed. |
| 3 | EDTA | Uncertainty in potential benefits and harms. Further research is needed. |
| 4 | Gabapentin | Uncertainty in potential benefits and harms. Further research is needed. |

| | Intervention | Summary of findings |
|----|--|---|
| | | |
| 5 | Olfactory training | Olfactory training may improve olfactory symptoms. Further research is needed. |
| 6 | Omega-3 fatty acids | Uncertainty in potential benefits and harms. Further research is needed. |
| 7 | Palmitoylethanolamide + Luteolin | Palmitoylethanolamide + Luteolin may improve olfactory symptoms. However, certainty of the evidence was low. Further research is needed. |
| 8 | Pentasodium diethylenetriamine pentaacetate (DTPA) | Uncertainty in potential benefits and harms. Further research is needed. |
| 9 | Steroids (nasal) | Uncertainty in potential benefits and harms. Further research is needed. |
| 10 | Steroids | Steroids may not improve olfactory nor gustatory symptoms. Further research is needed. |
| 11 | Theophylline (nasal) | Uncertainty in potential benefits and harms. Further research is needed. |
| 12 | Vitamin A | Uncertainty in potential benefits and harms. Further research is needed. |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined nine therapeutic options for PCC olfactory and/or gustatory dysfunction.
- ADAPT-232 (adaptogens): The results of one RCT suggest that ADAPT-232 may improve olfactory symptoms. However, certainty of the evidence was low because of imprecision. Further research is needed.
- Olfactory training: The results of four RCTs suggest that olfactory training may improve olfactory symptoms. However, certainty of the evidence was low because of risk of bias and inconsistency. Further research is needed.

- Palmitoylethanolamide + Luteolin: The results of one RCT suggest that Palmitoylethanolamide + Luteolin may improve olfactory symptoms. However, certainty of the evidence was low because of imprecision. Further research is needed.
- **Steroids:** The results of two RCTs suggest that steroids may not improve olfactory nor gustatory symptoms. However, certainty of the evidence was low because of imprecision. Further research is needed.

P-ACC-related cardiovascular system symptoms

Table ES9. List of RCTs of interventions for P-ACC-related cardiovascular system symptoms with primary outcome measures and certainty (n=2)

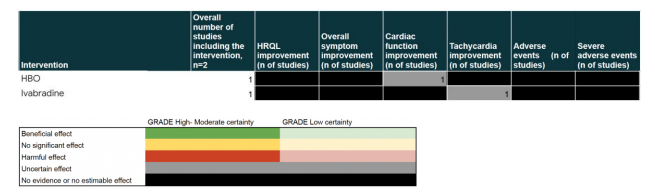


Table ES10. Summary of findings on potential therapeutic options for P-ACC-related cardiovascular system symptoms (n=2), as of 3 November 2023

| | Intervention | Summary of findings |
|---|-------------------|--|
| | | |
| 1 | Hyperbaric oxygen | Uncertainty in potential benefits and harms. Further research is needed. |
| 2 | Ivabradine | Uncertainty in potential benefits and harms. Further research is needed. |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined two therapeutic options for P-ACC- related cardiovascular system symptoms.
- The effects of assessed interventions are uncertain.

P-ACC-related psychological distress

Table ES11. List of RCTs of interventions for P-ACC-related psychological distress with primary outcome measures and certainty (n=2)

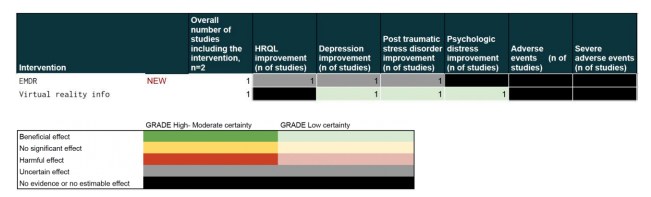


Table ES12. Summary of findings on potential therapeutic options for PCC psychological distress (n=2), as of 3 November 2023

| | Intervention | Summary of findings | | | | |
|---|--|---|--|--|--|--|
| 1 | Eye-movement desensitization and reprocessing (EMDR) | Uncertainty in potential benefits and harms. Further research is needed. | | | | |
| 2 | Virtual reality informational video | Virtual reality informational video may improve depression, post- traumatic stress, and psychological distress. However, certainty of the evidence was low. Further research is needed. | | | | |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined one therapeutic option for PCC psychological distress.
- Virtual reality informational video: The results of one RCT suggest that virtual reality informational video may improve depression, post-traumatic stress, and psychological distress. However, certainty of the evidence was low because of imprecision. Further research is needed.

P-ACC-related thromboembolic risk

Table ES13. List of RCTs of interventions for P-ACC-related thromboembolic risk with primary outcome measures and certainty (n=2)

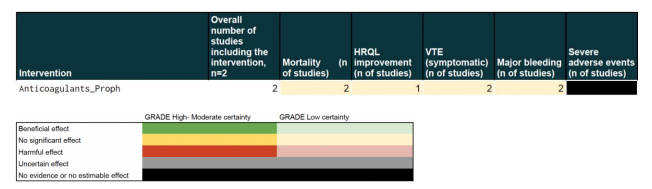


Table ES14. Summary of findings on potential therapeutic options for PCC thromboembolic risk (n=1), as of 3 November 2023

| | Intervention | Summary of findings | | | | |
|---|------------------------------------|--|--|--|--|--|
| 1 | Anticoagulants (prophylactic dose) | Anticoagulants may not have an important effect on mortality, VTE, major bleeding and HRQL. However, certainty of the evidence was low because of imprecision. Further research is needed. | | | | |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined one therapeutic option for PCC olfactory and/or gustatory dysfunction.
- Anticoagulants: The results of two RCTs suggest that anticoagulants (rivaroxaban and apixaban) may not have an important effect on mortality, HRQL, VTE or major bleeding. However, certainty of the evidence was low because of risk of imprecision. Further research is needed.

Pediatric inflammatory multisystem syndrome associated with SARS-CoV-2 (PIMS-TS)

Table ES13. List of RCTs of interventions for PIMS-TS with primary outcome measures and certainty (n=1)

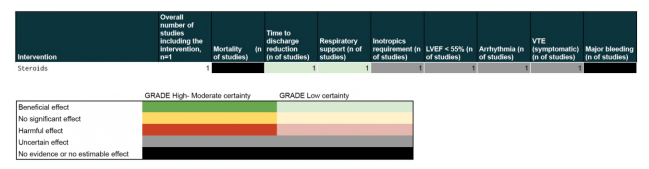


Table ES14. Summary of findings on potential therapeutic options for PCC thromboembolic risk (n=1), as of 3 November 2023

| | Intervention | Summary of findings | | | | |
|---|--------------|--|--|--|--|--|
| 1 | Steroids | Steroids may reduce time to discharge and respiratory support requirements. However, certainty of the evidence was low for risk of bias and imprecision. Further research is needed. | | | | |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined one therapeutic option for PCC olfactory and/or gustatory dysfunction.
- **Steroids:** The results of one RCT suggest that steroids may reduce time to discharge and respiratory support requirements. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.

P-ACC prophylaxis

Table ES15. List of RCTs of interventions for P-ACC prophylaxis with primary outcome measures and certainty (n=5)

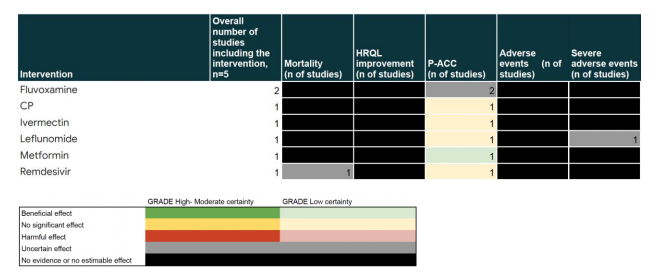


Table ES16. Summary of findings on potential therapeutic options for P-ACC prophylaxis (n=6), as of 3 November 2023

| | Intervention | Summary of findings | | | | |
|---|---------------------|---|--|--|--|--|
| 1 | Convalescent plasma | Convalescent plasma may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |
| 2 | Fluvoxamine | Fluvoxamine may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |
| 3 | Ivermectine | Ivermectin may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |
| 4 | Leflunomide | Leflunomide may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |
| 5 | Metformin | Metformin may reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |

| | Intervention | Summary of findings | | | | |
|---|--------------|--|--|--|--|--|
| 6 | Remdesivir | Remdesivir may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | |

- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined six therapeutic options for PCC olfactory and/or gustatory dysfunction.
- **Metformin:** The results of one RCT suggest that metformin may reduce P-ACC. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
- Ivermectin: The results of one RCT suggest that ivermectin may reduce P-ACC. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
- Convalescent plasma: The results of one RCT suggest that convalescent plasma may not reduce P-ACC. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
- Remdesivir: The results of one RCT suggest that remdesivir may not reduce P-ACC. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
- **Leflunomide:** The results of one RCT suggest that leflunomide may not reduce P-ACC. However, certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.

• Fluvoxamine: The results of two RCTs suggest that fluvoxamine may not reduce P-ACC. However, certainty of the evidence was low because of imprecision. Further research is needed.

Changes since previous edition

- Celecoxib for P-ACC related neurocognitive symptoms: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Budesonide for P-ACC related olfactory and/or gustatory dysfunction: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Albumin (inhaled) for P-ACC related dyspnea: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Gabapentin for P-ACC related olfactory and/or gustatory dysfunction: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Respiratory training for P-ACC related dyspnea: New evidence included not affecting results interpretation and/or certainty of the evidence judgments.
- Physical training for P-ACC related asthenia and/or fatigue: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Eye-movement desensitization and reprocessing (EMDR) for P-ACC related psychological distress: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- EDTA for P-ACC related olfactory and/or gustatory dysfunction: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- CCSA for P-ACC related asthenia and/or fatigue: New evidence included not affecting results interpretation and/or certainty of the evidence judgments.

Concluding remarks

- The Pan American Health Organization (PAHO) is continually monitoring ongoing research on any possible therapeutic options. As evidence emerges, PAHO will immediately assess and update its position, particularly as it applies to any special population subgroups such as children, expectant mothers, and those with immune conditions.
- PAHO is also mindful of the emerging differential impact of PCC on ethnic and minority groups and is continuously seeking data that could help in mitigating excess risk of severe illness or death in minority subgroups. These groups are plagued by social and structural inequities that bring to bear a disproportionate burden of COVID-19 illness.
- The safety of the patient suffering from COVID-19 is a key priority to improve the quality of care in the provision of health services.
- Adequately designed and reported clinical trials are crucial for the practice of evidencebased medicine. Most of the research to date on PCC has very poor methodology that is hidden and very difficult to validate. Greater transparency and better designed studies are urgently needed.

Systematic review of therapeutic options for postacute or post COVID-19 condition (P-ACC)

Background

Post COVID-19 condition (PCC), also known as long COVID or post-acute sequelae of SARS-CoV-2 infection (PASC), is the continuation or development of new symptoms in the period after acute infection with SARS-CoV-2 (1–4). The World Health Organization (WHO) definition of PCC states that these symptoms should be present after three months of the initial SARS-CoV-2 infection and last for at least two months with no other explanation (1, 2). While PASC definitions state persistent or new symptoms need to be present 30 days after a documented SARS-COV-2 infection or the onset of COVID-19 symptoms (3, 4). Post COVID-19 condition or post-acute sequelae of SARS-CoV-2 infection (P-ACC) can affect anyone exposed to SARS-CoV-2, regardless of age or severity of acute infection. Many of the reported symptoms are debilitating and have a strong negative impact on mental health and quality of life (5). While most patients recover, some may experience multiple outcomes, with multiple organ systems affected simultaneously, including cardiovascular, mental, metabolic, renal, and others (3, 6). Recommendations for the management of patients with PCC are continuously being developed and need to evolve as evidence of interventions effects becomes available (7).

In this review, we compiled the following evidence on potential therapeutic options for P-ACC. We included all the identified clinical forms, symptoms, and manifestations of P-ACC for which an intervention was assessed in at least one randomized controlled trial (RCT). We hope this information will support investigators, policymakers, and prescribers navigate the flood of relevant data to ensure that management of P-ACC, at both the individual and population levels, is based on the best available knowledge. We will endeavor to continually update this resource as more research is released into the public space.

Methods

We used Living OVerview Evidence (L-OVE: available the of from: https://iloveevidence.com) platform to identify studies for inclusion in this review. This platform is a system that maps PICO (Patient-Intervention-Comparison-Outcome) questions to a repository developed by Epistemonikos Foundation. This repository is continuously updated through searches in electronic databases, preprint servers, trial registries, and other resources relevant to COVID-19. The latest version of the methods, the total number of sources screened, and a living flow diagram and report of the project is updated regularly on the L-OVE website (8).

Search strategy

We systematically searched in L·OVE for COVID-19. The search terms and databases covered are described on the L·OVE search strategy methods page (available from: https://app.iloveevidence.com/loves/5e6fdb9669c00e4ac072701d?question_domain=un_defined§ion=methods). The repository is continuously updated, and the information is transmitted in real time to the L·OVE platform. It was last checked for this review on 3 November2023. The searches covered the period from the inception date of each database, and no study design, publication status, or language restriction was applied.

Study selection

The results of the searches in the individual sources were de-duplicated by an algorithm that compares unique identifiers (database identification number, digital object identifier [DOI], trial registry identification number), and citation details (i.e., author names, journal, year of publication, volume, number, pages, article title, and article abstract). Then, the information matching the search strategy was sent in real time to the L-OVE platform, where at least two authors independently screened the titles and abstracts yielded against the inclusion criteria. We obtained the full reports for all titles that appeared to meet the inclusion criteria or required further analysis and then decided about their inclusion.

Inclusion criteria

We aimed to find all available RCTs for potential therapeutic interventions for P-ACC with study designs that included head-to-head comparisons, or control groups with no intervention or a placebo. Target patient populations included both adults and children with persistent, or new, symptoms or clinical manifestations after acute COVID-19. We used the term Post Acute or Post COVID-19 condition (P-ACC) to refer to the population included in our review (studies reporting on patients with persistent or new symptoms after acute COVID-19 independently of the time of onset of those symptoms) (1–4). We focused on comparative effectiveness studies that provide evidence on outcomes of crucial importance to patients (mortality, health-related quality of life [HRQL], and disease-specific symptoms).

Living evidence synthesis

An artificial intelligence algorithm deployed in the Coronavirus/COVID-19 topic of the L-OVE platform provides instant notification of articles with a high likelihood of being eligible. The authors review them, decide upon inclusion, and update the living web version of the review accordingly. If meta-analytical pooling is possible from retrieved evidence, we will do this to derive more precise estimates of effect and derive additional statistical power. No electronic database search restrictions were imposed.

For any meta-analytical pooling, if and when data allow, we pool all studies and present the combined analysis with relative and absolute effect sizes. To assess interventions' absolute effects, we applied relative effects to baseline risks (risks with no intervention). For baseline risks we used the mean risk in the control groups from included RCTs. For continuous outcomes, when possible, we calculated relative and absolute effects by estimating the proportion of patients with important improvement or deterioration following published guidance (9).

For result interpretations and imprecision assessment we used a minimally contextualized approach that considers whether the 95% confidence interval (CI) includes the null effect, or, when the point estimate is close to the null effect, whether the 95% CI lies within the

boundaries of small but important benefit and harm that corresponds to every outcome assessed (10, 11).

We used the following absolute effects thresholds to define important benefits and harms: Mortality, +/-1%; HRQL improvement, +/-2%; Overall symptom improvement, +/-5%; Functional capacity improvement, +/-5%; Strength improvement, +/-5%; Fatigue improvement, +/-5%; Pulmonary function improvement, +/-10%; Radiological response, +/-10%; Cognitive improvement, +/-5%; Depression improvement, +/-5%; Olfactory symptoms improvement, +/-5%; Gustatory symptoms improvement, +/-5%; Tachycardia improvement, +/-5%; Venous thromboembolism (VTE) (symptomatic), +/-3%; Post-traumatic stress disorder improvement, +/-5%; Psychological distress improvement, +/-5%; Major bleeding, +/-3%; Severe adverse events, +/-3%; Adverse events, +/-5%; Time to discharge reduction, +/-4%; Respiratory support requirement +/-2%; Inotropic requirement +/-2%; Left ventricular ejection fraction deterioration (LVEF <55%) +/-5%; Arrhythmia +/-5%; P-ACC, +/-3%.

For some interventions when we found significant heterogeneity, we performed subgroup analysis considering: 1) risk of bias (high/moderate vs low risk of bias); and 2) intervention characteristics (e.g., different doses or administration schemes). When we observed significant differences between subgroups, we presented individual subgroups' estimates of effect and certainty of the evidence assessment.

A risk of bias assessment was applied to RCTs focusing on randomization, allocation concealment, blinding, attrition, or other biases relevant to the estimates of effect (Table 1) (12). The GRADE approach was used to assess the certainty of the body of evidence for every comparison on an outcome basis (13).

Study selection, data extraction, and risk of bias assessment were performed, independently and in parallel, by two reviewers. Discrepancies were resolved by discussion.

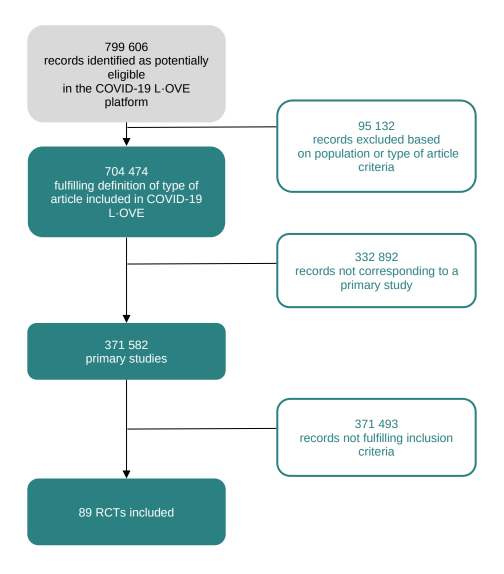
We used MAGIC authoring and publication platform (available from: https://app.magicapp.org/) to generate the tables summarizing our findings, which are included in Annex 1.

Results

Studies identified and included

The study identification and selection process is shown in Figure 1. A total of 89 RCTs were selected for inclusion. A list of excluded studies is available upon request.

Figure 1. Study identification and selection process



Risk of bias

Overall, our risk of bias assessment for the limited reported RCTs found high risk of bias due to suboptimal randomization, allocation concealment, and blinding (as well as other methodological and reporting concerns). Most RCTs were also very small in size and had small event numbers. The methods were very poor overall, and the reporting was suboptimal. In general, follow-up was short. The risk of bias assessment of each RCT is presented in Table 1.

Table 1. Risk of bias of included RCTs

| Study | Risk-of-bias arising from | Risk-of-bias due to deviations | Risk-of-bias due to | Risk-of-bias in | Risk-of-bias in selection | Overall Risk-of-bias judgement | |
|----------------------------------|---------------------------|---------------------------------|----------------------|--------------------|---------------------------|--------------------------------|-------------------|
| | randomization process | from the intended interventions | missing outcome data | measurement of the | of the reported result | Mortality | HRQL, symptom |
| | | | | outcome | | | specific outcomes |
| √aira LA et al | High | Some Concerns | Low | Some Concerns | Low | High | High |
| RC 4-7-2020 (Abdelalim AA et al) | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Di Stadio | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Chudzik M et al | High | Some Concerns | Low | Some Concerns | Low | High | High |
| CITADEL | High | Some Concerns | Low | Some Concerns | Low | High | High |
| MICHELLE | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Zilberman-Itskovich | Low | Low | Low | Low | Low | Low | Low |
| Botek M et al | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Jadhav KP et al | High | Some Concerns | Low | Some Concerns | Low | High | High |
| COLDSTER | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Oliver-Mas | Low | Low | Low | Low | Low | Low | Low |
| Nambi | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Di Stadio 2 | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Hansen | Low | Low | Low | Low | Low | Low | Low |
| Tosato | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Rathi | | | | | | | |
| | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Bazdyrev | Low | Low | Low | Low | Low | Low | Low |
| King | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| ICU-VR | High | Some Concerns | Low | Some Concerns | Low | High | High |
| ENO Breathe | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Pires | High | Some Concerns | Low | Some Concerns | Low | High | High |
| McNarry | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Srinivasan | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Kharaeva_Moderate | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Kharaeva_Severe | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Gaylis | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Karosanidze | Low | Low | Low | Low | Low | Low | Low |
| Badran | Low | Low | Low | Low | Low | Low | Low |
| COVANOS | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| RECOVER | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Kutashov | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Vallier | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Swissped RECOVERY | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| JK Phyto-V | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Rodriguez-Blanco | High | Some Concerns | Low | Some Concerns | Low | High | High |
| COVID-OUT - Metformin | Low | Low | High | Low | Low | High | High |
| COVID-OUT - Ivermectin | | | | Low | | | |
| | Low | Low | High | | Low | High | High |
| COVID-OUT - Fluvoxamine | Low | Low | High | Low | Low | High | High |
| Santana | Low | Low | Low | Low | Low | Low | Low |
| CSSC-004 | Low | Low | Some Concerns | Low | Low | Low | Some Concerns |
| Deshpande | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Oal Negro | High | Some Concerns | Low | Some Concerns | Low | High | High |
| nsCOVID | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Rutkowski | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| SOLIDARITY - Finland | Low | Low | Low | Low | Low | Low | Low |
| Hawkins | Low | Low | Low | Low | Low | Low | Low |
| Schepens | Low | Low | Low | Low | Low | Low | Low |
| Kusumawardani | High | Some Concerns | Low | Some Concerns | Low | High | High |
| SCENT2 | Low | Low | Low | Low | Low | Low | Low |
| Hausswirth | High | Some Concerns | Low | Some Concerns | Low | High | High |
| /ersace | High | Low | Low | Low | Low | High | High |
| ACTIV-4C | Low | Low | Low | Low | Low | Low | Low |
| Simpson | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Stavrou | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Zulbaran-Rojas | High | | | | | High | High |
| • | | Some Concerns | Low | Some Concerns | Low | | - |
| Kalayeh | High | Some Concerns | Low | Some Concerns | Low | High | High |
| DEFEAT-COVID | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Kerget | High | Some Concerns | Low | Some Concerns | Low | High | High |
| arahani | Low | Low | Low | Low | Low | Low | Low |
| Finnigan | Low | Low | Low | Low | Low | Low | Low |
| Longobardi | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| | | | | | | | |

| Kuut | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
|---------------------|------|---------------|------|---------------|------|------|------|
| Shabaan | Low | Low | Low | Low | Low | Low | Low |
| RECOVE | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Del Corral | Low | Low | Low | Low | Low | Low | Low |
| Imam | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Nagy | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Ibrahim | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Samper-Pardo | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Momtazmanesh | Low | Low | Low | Low | Low | Low | Low |
| Leitman | Low | Low | Low | Some Concerns | High | High | High |
| Brichetti | Low | Low | Low | Low | Low | Low | Low |
| Lerner | Low | Low | Low | Low | Low | Low | Low |
| Rasmussen | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Caprioli | Low | Low | Low | Low | Low | Low | Low |
| Nambi_2 | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Chung | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Toussaint | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Oliveira | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Da Silva | Low | Low | Low | Low | Low | Low | Low |
| Tanhan | High | Some Concerns | Low | Some Concerns | Low | High | High |
| TERECO | Low | Low | Low | Low | Low | Low | Low |
| Ansari | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Hautefort | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Ampio | High | Some Concerns | Low | Some Concerns | Low | High | High |
| GRACE | Low | Low | High | Low | Low | High | High |
| Mirenayat | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Espinoza-Bravo | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Senén | High | Some Concerns | Low | Some Concerns | Low | High | High |
| COVEMERALD | High | Some Concerns | Low | Some Concerns | Low | High | High |
| Abdelazim | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| Tanashyan | Low | Low | Low | Low | Low | Low | Low |
| Ogonowska-Slodownik | Low | Some Concerns | Low | Some Concerns | Low | Low | High |
| | | | | | | | |

Main findings

P-ACC-related asthenia or fatigue

Actovegin

See Summary of findings Table A1, Annex 1

We identified one RCT including 444 participants in which Actovegin was compared against standard of care. Our results showed:

Actovegin may improve fatigue, relative risk (RR) 1.84 (95% CI 1.59 to 2.14); risk difference (RD) 39.7% (95% CI 27.7% to 56.3%); Low certainty ⊕⊕○○

ADAPT-232 (adaptogens)

See Summary of findings Table A2, Annex 1

We identified one RCT including 99 participants in which ADAPT-232 was compared against standard of care. Our results showed:

ADAPT-232 may not improve fatigue, relative risk (RR) 1.02 (95% CI 0.84 to 1.24);
 risk difference (RD) 1.6% (95% CI −12.6% to 18.9%); Low certainty ⊕⊕○○

Cytoflavin

See Summary of findings Table A3, Annex 1

We identified one RCT including 200 patients in which cytoflavin was compared against standard of care. Our results showed:

Cytoflavin may not improve fatigue, RR 1.02 (95% CI 0.98 to 1.06); RD 2.1% (95% CI −1.9% to 6.2%); Low certainty ⊕⊕○○

Enzymes + probiotics

See Summary of findings Table A4, Annex 1

We identified one RCT including 200 patients in which enzymes + probiotics were compared against standard of care. Our results showed:

Enzymes + probiotics may improve fatigue, RR 6.07 (95% CI 3.79 to 9.71);
 RD 76% (95% CI 41.8% to 85%); Low certainty ⊕⊕○○

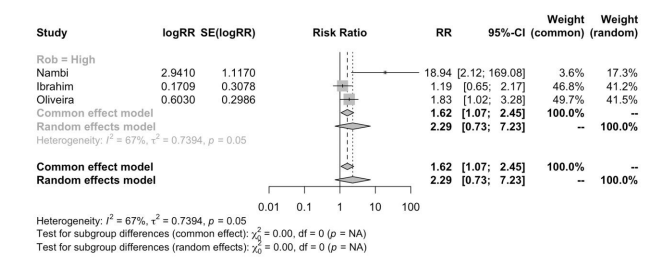
Physical training

See Summary of findings Table A30, Annex 1

We identified six RCTs including 304 patients in which physical training was compared against standard of care. Our results showed:

- Physical training may improve HRQL, 1.62 (95% CI 1.07 to 2.45); RD 21.1% (95% CI 2.4% to 49.2%); Low certainty ⊕⊕○○
- Physical training may improve fatigue, RR 4.66 (95% CI 1.96 to 11.09); RD 44.4% (95% CI 11.7% to 87.9%); Low certainty ⊕⊕○○
- Physical training may improve functional capacity, RR 1.81 (95% CI 1.23 to 2.65); RD 17.4% (95% CI 4.9% to 35.4%); Low certainty ⊕⊕○○
- Physical training may improve strenght, RR 3.13 (95% CI 1.02 to 9.55); RD
 11.3% (95% CI 0.1% to 45.4%); Low certainty ⊕⊕○○

Figure 1. HRQL improvement in RCTs comparing physical training with standard of care for treatment of patients with P-ACC-related asthenia/fatigue



Phytochemicals

See Summary of findings Table A5, Annex 1

We identified one RCT including 147 patients in which phytochemicals were compared against standard of care. Our results showed:

- Phytochemicals may improve HRQL, RR 1.33 (95% CI 1.03 to 1.71); RD 18% (95% CI 1.8% to 39%); Low certainty ⊕⊕○○
- Phytochemicals may improve fatigue, RR 1.24 (95% CI 0.95 to 1.62); RD 13.1% (95% CI -2.5% to 33.5%); Low certainty ⊕⊕○○

Transcranial direct current stimulation (tDCS)

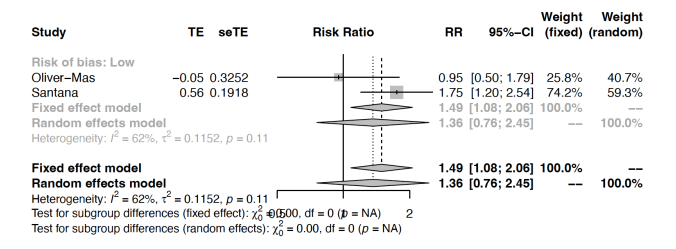
See Summary of findings Table A6, Annex 1

We identified two RCTs including 117 patients in which tDCS was compared against standard of care. Our results showed:

tDCS may improve fatigue, RR 1.36 (95% CI 0.76 to 2.45); RD −16.9% (95% CI − 11.2% to 53%); Low certainty ⊕⊕⊖⊖ (see figure 2.)

- tDCS may improve HRQL, RR 1.37 (95% CI 1.09 to 1.71); RD −26% (95% CI − 6.7% to 30%); Low certainty ⊕⊕○○
- tDCS may not increase adverse events, RR 0.83 (95% CI 0.26 to 2.73); RD −3.4% (95% CI −15.5% to 36%); Low certainty ⊕⊕○○

Figure 2. Fatigue in RCTs comparing tDCS with standard of care for treatment of patients with P-ACC-related asthenia/fatigue



AXA1125

See Summary of findings Table A27, Annex 1

We identified one RCT including 41 participants in which AXA1125 was compared against standard of care. Our results showed:

- AXA1125 may improve fatigue, RR 1.07 (95% CI 0.79 to 1.44); RD 5.1% (95% CI −16.6% to 34.5%); Low certainty ⊕⊕○○
- AXA1125 may not improve functional capacity, RR 0.87 (95% CI 0.51 to 1.48); RD
 -8.1% (95% CI –30% to 29.3%); Low certainty ⊕⊕○○

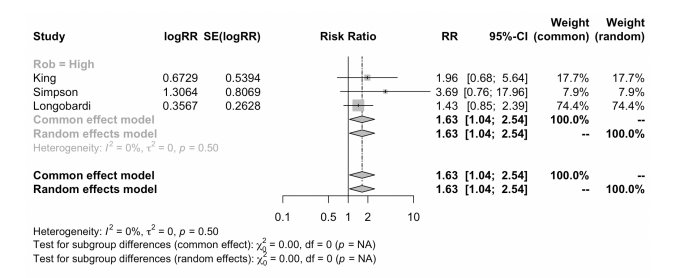
Telerehabilitation

See Summary of findings Table A28, Annex 1

We identified five RCTs including 246 participants in which telerehabilitation was compared against standard of care. In addition, we identified one study in which synchronous and asynchronous telerehabilitation strategies were compared. Our results showed:

- Telerehabilitation may improve fatigue, RR 1.63 (95% CI 1.04 to 2.54); RD 19.1% (95% CI 1.2% to 46.8%); Low certainty ⊕⊕○○ (see figure 3.)
- Telerehabilitation may improve functional capacity, RR 1.37 (95% CI 1.11 to 1.67);
 RD 13.8% (95% CI 2% to 29.8%); Low certainty ⊕⊕○○

Figure 3. Fatigue in RCTs comparing telerehabilitation with standard of care for treatment of patients with P-ACC-related asthenia/fatigue



Cognitive behavioral therapy

See Summary of findings Table A29, Annex 1

We identified one RCT including 114 participants in which cognitive behavioral therapy was compared against standard of care. Our results showed:

- Behavioral therapy may improve fatigue, RR 2.2 (95% CI 1.35 to 3.58); RD 31.6% (95% CI 9.2% to 68%); Low certainty ⊕⊕○○
- Behavioral therapy may improve functional capacity, RR 1.37 (95% CI 1.08 to 1.73); RD 22.4% (95% CI 4.8% to 44.7%); Low certainty ⊕⊕○○

P-ACC-related dyspnea

ADAPT-232 (adaptogens)

See summary of findings Table A7 in Annex 1

We identified one RCT including 99 patients in which ADAPT-232 was compared against standard of care. Our results showed:

ADAPT-232 may not improve dyspnea, RR 1 (95% CI 0.94 to 1.06); RD 0% (95% CI −5.4% to 5.7%); Low certainty ⊕⊕○○

High dose steroids

See Summary of findings Table A9, Annex 1

We identified one RCT including 130 patients in which high dose steroids (prednisone 40 mg a day) was compared against standard dose steroids (prednisone 10 mg a day). Our results showed:

- High dose steroids may not improve dyspnea, RR 1 (95% CI 0.87 to 1.15); RD 0% (95% CI −11% to 13%); Low certainty ⊕⊕○○
- High dose steroids may not increase adverse events, RR 0.92 (95% CI 0.75 to 1.13); RD −6.2% (95% CI −19.3% to 10%); Low certainty ⊕⊕○○

Respiratory training/rehabilitation

See Summary of findings Table A10, Annex 1

We identified ten RCTs including 616 patients in which different modalities of respiratory training/rehabilitation were compared with standard of care. In addition, we identified two studies that compared home based respiratory training vs. inpatient respiratory training, two studies comparing VR respiratory training vs. conventional respiratory training and one study that compares incentive spirometry vs. conventional respiratory training. Our results showed:

- Respiratory training/rehabilitation probably improves RR 1.41 (95% CI 1.18 to 1.67); RD 22.3% (95% CI 9.8% to 36.5%); Moderate certainty ⊕⊕⊕○ (see Figure 4)
- Respiratory training/rehabilitation probably improves dyspnea, RR 2.11 (95% CI 1.17 to 3.8); RD 34.2% (95% CI 5.2% to 86.4%); Moderate certainty ⊕⊕⊕○
- Respiratory training/rehabilitation may improve functional capacity, RR 1.41 (95%) CI 1.05 to 1.9); RD 20.5% (95% CI 2.5% to 44.9%); Moderate certainty ⊕⊕⊕○

Figure 4. HRQL in RCTs comparing respiratory training/rehabilitation with standard of care for treatment of patients with P-ACC-related dyspnea.

| Study | logRR S | E(logRR) | Risk Ratio | RR | 95%-CI | Weight (common) | - |
|--|------------------|--------------------------------------|------------|-------------------------------------|--|---|---|
| Rob = High ENO Breathe McNarry RECOVER Nambi_2 Common effect model Random effects model Heterogeneity: $I^2 = 0\%$, τ^2 | | 0.3090 0.2639 0.2410 0.1050 | | 1.68 1.48 1.56 1.61 | [1.28; 4.30] [1.00; 2.82] [0.92; 2.37] [1.27; 1.92] [1.36; 1.91] | 4.5% 6.2% 7.5% 39.3% 57.5% | 7.0% 9.0% 10.4% 27.4% 53.8% |
| Rob = Low Del Corral Del Corral_Insp TERECO Common effect model Random effects model Heterogeneity: $l^2 = 0\%$, τ^2 | | 0.2229 0.2154 0.1332 | | 1.00 1.13 1.16 | [0.94; 2.26] [0.66; 1.53] [0.87; 1.47] [0.95; 1.41] | | 11.7% 12.3% 22.2% 46.2% |
| Common effect model Random effects model Heterogeneity: $I^2 = 36\%$, 1 | $x^2 = 0.0178$, | | 0.5 1 2 | | [1.23; 1.60] [1.18; 1.67] | 100.0% | 100.0% |

Test for subgroup differences (common effect): χ_1^2 = 6.18, df = 1 (ρ = 0.01) Test for subgroup differences (random effects): χ_1^2 = 6.18, df = 1 (ρ = 0.01)

Treamid

See Summary of findings Table A11, Annex 1

We identified one RCT including 59 patients in which treamid was compared with standard of care. Our results showed:

- Treamid may improve dyspnea, RR 1.96 (95% CI 0.9 to 4.25); RD 21.7% (95% CI -2.3% to 73.7%); Low certainty ⊕⊕○○
- Treamid may improve functional capacity, RR 1.1 (95% CI 0.64 to 1.9); RD 0.4% (95% CI 16.2% to 39.8%); Low certainty ⊕⊕○○
- Treamid may increase adverse events, RR 1.19 (95% CI 0.56 to 2.5); RD 5.5% (95% CI −12.7% to 43.6%); Low certainty ⊕⊕○○

P-ACC-related neurocognitive symptoms or sleep disturbances

Actovegin

See Summary of findings Table A12, Annex 1

We identified one RCT including 44 patients in which actovegin was compared with standard of care. Our results showed:

Actovegin may improve cognition, RR 1.19 (95% CI 1.06 to 1.33); RD 12.7% (95% CI 4.2% to 22.3%); Low certainty ⊕⊕○○

Famotidine

See Summary of findings Table A31, Annex 1

We identified one RCT including 50 patients in which famotidine was compared with standard of care. Our results showed:

- Famotidine may improve cognition, RR 1.33 (95% CI 1.05 to 1.69); RD 24.6% (95% CI 3.7% to 51.1%); Low certainty ⊕⊕○○
- It is uncertain if famotidine improves depresion, RR 3.71 (95% CI 0.83 to 16.6);
 RD 20.6% (95% CI -1.3% to 92.4%); Very low certainty ⊕○○○

Hyperbaric oxygen (HBO)

See Summary of findings Table A13, Annex 1

We identified one RCT including 73 patients in which HBO was compared with standard of care. Our results showed:

HBO may improve HRQL, RR 1.3 (95% CI 0.84 to 2); RD 13.9% (95% CI −7.4% to 46.9%); Low certainty ⊕⊕○○

Transcranial direct current stimulation (tDCS)

See Summary of findings Table A14, Annex 1

We identified one RCT including 47 patients in which tDCS was compared with standard of care. Our results showed:

tDCS may not improve HRQL, RR 0.59 (95% CI 0.33 to 1.05); RD −27.5% (95% CI −44.8% to 3.4%); Low certainty ⊕⊕○○

P-ACC-related olfactory and/or gustatory dysfunction

ADAPT-232 (adaptogens)

See Summary of findings Table A15, Annex 1

We identified one RCT including 99 patients in which ADAPT-232 was compared with standard of care. Our results showed:

ADAPT-232 may not improve olfactory symptoms, RR 0.89 (95% CI 0.79 to 1.01);
 RD −10.3% (95% CI −20.5% to 1.4%); Low certainty ⊕⊕○○

Olfactory training

See Summary of findings Table A32, Annex 1

We identified four RCTs including 308 patients in which olfactory training was compared with standard of care. Our results showed:

Olfactory training may improve olfactory symptoms, RR 1.49 (95% CI 0.96 to 2.31); RD 20% (95% CI −1.6% to 53.6%); Low certainty ⊕⊕○○

Figure 5. Olfactory symptoms in RCTs comparing olfactory training with standard of care for treatment of patients with P-ACC-related olfactory and/or gustatory dysfunction.

| Study | logRR S | E(logRR) | Risk Ratio | RR | 95%-CI | Weight (common) | Weight (random) |
|---|---|--------------------------------------|------------|---------------------------------------|--|--|----------------------------------|
| Rob = High Di Stadio_2 Pires COVANOS Chung_2 Common effect mod Random effects mod Heterogeneity: $l^2 = 63^\circ$ | del | 0.1340 0.4158 0.2606 0.4530 | | 0.87 1.16 - 1.65 1.79 | [1.68; 2.84] [0.39; 1.97] [0.70; 1.93] [0.68; 4.01] [1.44; 2.23] [0.96; 2.31] | 68.7% 7.1% 18.2% 6.0% 100.0% | 38.6% 17.6% 27.9% 15.8% |
| Common effect mod Random effects mod Heterogeneity: $I^2 = 63^\circ$ Test for subgroup differest for subgroup differest for subgroup differences. | del del $\%$, $\tau^2 = 0.1124$, μ rences (commor | $p = 0.05$ n effect): $\chi_Q^2 = 0$ | | | [1.44; 2.23] [0.96; 2.31] | 100.0% | 100.0% |

Palmitoylethanolamide + Luteolin

See Summary of findings Table A16, Annex 1

We identified one RCT including 130 patients in which palmitoylethanolamide + luteolin was compared with standard of care. Our results showed:

Palmitoylethanolamide + luteolin may improve olfactory symptoms, RR 3.11 (95% CI 1.47 to 6.66); RD 35.5% (95% CI 7.8% to 83.3%); Low certainty ⊕⊕○○

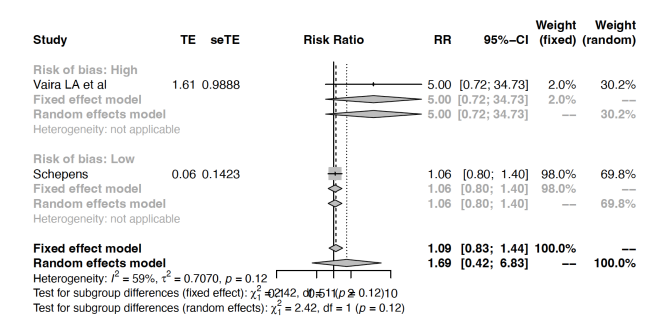
Steroids

See Summary of findings Table A17, Annex 1

We identified two RCTs including 131 patients in which steroids were compared with standard of care. Our results showed:

- Steroids may not improve olfactory symptoms, RR 1.09 (95% CI 0.83 to 1.44); RD
 3.3% (95% CI −6.2% to 16.1%); Low certainty ⊕⊕○○ (figure 5)
- Steroids may not improve gustatory symptoms, RR 1.01 (95% CI 0.67 to 1.53); RD 0.5% (95% CI −14.6% to 23.3%); Low certainty ⊕⊕○○

Figure 6. Olfactory symptoms in RCTs comparing steroids with standard of care for treatment of patients with P-ACC-related olfactory and/or gustatory dysfunction.



P-ACC-related cardiovascular system symptoms

The effects of the assessed interventions are uncertain.

P-ACC-related psychological distress

Virtual reality (VR) informational video

See Summary of findings Table A18, Annex 1

We identified one RCT including 89 patients in which a virtual reality-based (VR) intervention was compared with standard of care. Our results showed:

- VR informational video may improve depression, RR 1.21 (95% CI 0.95 to 1.54);
 RD 14% (95% CI −3.7% to 36.7%); Low certainty ⊕⊕○○
- VR informational video may improve post-traumatic stress, RR 1.18 (95% CI 0.98 to 1.42); RD 13.8% (95% CI −1.5% to 32.3%); Low certainty ⊕⊕○○
- VR informational video may improve psychological distress, RR 1.49 (95% CI 1.08 to 2.05); RD 25.5% (95% CI 4.1% to 55.1%); Low certainty ⊕⊕○○

P-ACC-related thromboembolic risk

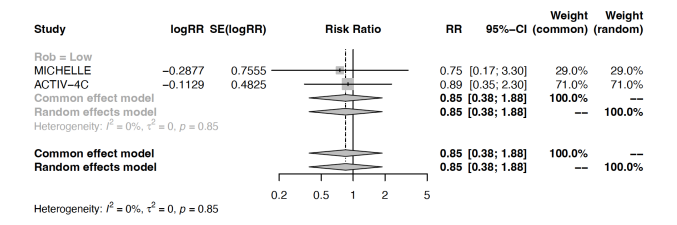
Anticoagulants

See Summary of findings Table A19, Annex 1

We identified two RCTs including 1535 patients with COVID-19 after hospitalization in which anticoagulants in prophylactic dose were compared with standard of care. Our results showed:

- Anticoagulants may not have an important effect on mortality, RR 0.85 (95% CI 0.38 to 0.88); RD -0.3% (95% CI −1.2% to 1.8%); Low certainty ⊕⊕⊖⊝ (Figure 6)
- Anticoagulants may not have an important effect on RR 0.99 (95% CI 0.78 to 1.24); Low certainty ⊕⊕○○
- Anticoagulants may not have an important effect on VTE, RR 1 (95% CI 0.29 to 3.45); RD 0% (95% CI −2.3% to 7.9%); Low certainty ⊕⊕○○ (based on low RoB studies)
- Anticoagulants may not have an important effect on VTE, RR 2.01 (95% CI 0.18 to 22.1); RD 0.1% (95% CI −0.1% to 1.2%); Low certainty ⊕⊕○○

Figure 7. Mortality in RCTs comparing anticoagulants with standard of care for treatment of patients with COVID-19 after hospitalization.



Pediatric inflammatory multisystem syndrome associated with SARS-CoV-2 (PIMS-TS)

Steroids

See Summary of findings Table A20, Annex 1

We identified one RCT including 75 patients in which systemic steroids were compared with intravenous immunoglobulins (IVIG). Our results showed:

- Steroids may reduce time to discharge, RR 1.09 (95% CI 0.88 to 1.39); RD 4.5% (95% CI -6% to 19.5%); Low certainty ⊕⊕○○
- Steroids may reduce respiratory support requirements, RR 0.49 (95% CI 0.27 to 0.89); RD -28.2% (95% CI -40.5% to -5.9%); Low certainty ⊕⊕○○

P-ACC prophylaxis

Metformin

See Summary of findings Table A21, Annex 1

We identified one RCT including 1125 patients in which metformin was compared with standard of care. Our results showed:

Metformin may reduce P-ACC, RR 0.59 (95% CI 0.39 to 0.88); RD -4.3% (95% CI -6.4% to -1.2%); Low certainty ⊕⊕○○

Ivermectin

See Summary of findings Table A22, Annex 1

We identified one RCT including 739 patients in which metformin was compared with standard of care. Our results showed:

Metformin may reduce P-ACC, RR 0.99 (95% CI 0.61 to 1.62); RD 0% (95% CI - 1.7% to 2.6%); Low certainty ⊕⊕○○

Convalescent plasma

See Summary of findings Table A23, Annex 1

We identified one RCT including 882 patients in which metformin was compared with standard of care. Our results showed:

Convalescent plasma may not reduce P-ACC, RR 0.93 (95% CI 0.77 to 1.12); RD
 -2.4% (95% CI -7.9% to -4.2%); Low certainty ⊕⊕○○

Remdesivir

See Summary of findings Table A24, Annex 1

We identified one RCT including 181 patients in which metformin was compared with standard of care. Our results showed:

Remdesivir may not reduce P-ACC, RR 1.06 (95% CI 0.53 to 2.13); RD 0.8% (95% CI -6.9% to -16.4%); Low certainty ⊕⊕○○

Leflunomide

See Summary of findings Table A25, Annex 1

We identified one RCT including 172 patients in which leflunomide was compared with standard of care. Our results showed:

Leflunomide may not reduce P-ACC, RR 1.28 (95% CI 0.92 to 1.77); RD 11.2% (95% CI -3.2% to 31.1%); Low certainty ⊕⊕○○

Fluvoxamine

See Summary of findings Table A26, Annex 1

We identified two RCTs including 680 patients in which fluvoxamine was compared with standard of care. Our results showed:

Fluvoxamine may not reduce P-ACC, RR 0.99 (95% CI 0.81 to 1.21); RD -0.4% (95% CI -8.4% to 9.3%); Low certainty ⊕⊕○○

Full description of included studies

Tables 2 to 10 list all the identified studies that were included in this systematic review by intervention and P-ACC-related organ system affected. The treatments are arranged in alphabetical order. Study or author names, publication status, patient populations, interventions, sources of bias, outcomes, effect sizes, and certainty are listed for each study.

Table 2. Description of included studies and interventions effects for P-ACC-related asthenia or fatigue

| | 1-MNA Uncertainty in potential benefits and harms. Further research is needed. | | | | | | |
|--|--|--|--------------------------|--|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | ı | RCT | | | | |
| Chudzik et al. (14); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 30 days of acute COVID-19). 25 assigned to 1-MNA 58 mg a day and 25 assigned to standard of care. | Median age 49.5, male 32%, hypertension 14%, diabetes 2% | Not reported (NR) | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: Very low certainty ⊕○○○ Strength improvement: No information Adverse events: No information Severe adverse events: No information | | |

| Actov | Actovegin Actovegin may improve fatigue. However, certainty of the evidence was low. Further research is needed. | | | | | | |
|---|--|----------------------------|--------------------------|--|---|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| Kutashov et al. (15); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after NR days of acute COVID-19). 222 assigned to Actovegin 1200 mg a day for 60 days and 222 assigned to standard of care. | Mean age 67.6, male 31.98% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: RR 1.84 (95% CI 1.59 to 2.14); RD 39.7% (95% CI 27.7.6% to 53.6%); Low certainty ⊕⊕⊖⊖ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information | | |

ADAPT-232 (adaptogens) ADAPT-232 may not improve fatigue. However, certainty of the evidence was low. Further research is needed. Study; Patients and Comorbidities Additional Risk of bias and study Interventions publication status interventions interventions limitations effects vs standard of care (SOC) and analyzed **GRADE** certainty of the evidence **RCT** Karosanidze et Patients with P-Mean age 48.9, male NR Low risk of bias **HRQL** al. (16); Peer ACC (asthenia or improvement: No reviewed; 2022 fatigue after 30 information days of acute COVID-19). 49 Overall symptom improvement: No assigned to ADAPT-232 information (adaptogens) 60 mL a day for 14 **Fatigue** days and 50 improvement: assigned to RR 1.02 (95% CI standard of care. 0.84 to 1.24); RD 1.6% (95% CI -12.6% to 18.9%); Low certainty $\Theta\ThetaOO$ **Functional** capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information

Amygdala and Insula Retraining (AIR)

Uncertainty in potential benefits and harms. Further research is needed.

| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
|---|--|------------------------------------|--------------------------------------|--|--|
| | | | RCT | | |
| Toussaint et al (17); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 50 assigned to Amygdala and Insula Retraining (AIR) 40 to 60 minutes a day and 50 assigned to standard of care. | Mean age 43.6 | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information |
| | Uncertainty | Arginine v in potential benefits a | + Vitamin C nd harms. Further res | earch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | | RCT | | |

| Tosato et al. (18); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 28 days of acute COVID-19). 23 assigned to Arginine + Vitamin C 1.66 g/500 mg for 28 days and 23 assigned to standard of care. | Mean age 50.5 ± 14, male 34.8%, interval between COVID-19 and enrolment 254 days, hospitalization during COVID-19 56.5% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Strength improvement: Very low certainty ⊕○○○ Adverse events: No information Severe adverse events: No information |
|---|---|---|------------------------------------|--|--|
| | Uncertainty | Aroma | atherapy nd harms. Further rese | earch is needed. | |
| Study | | • | | | Interventions |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | СТ | | |

| Hawkins et al (19); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 150 days of acute COVID-19). 20 assigned to Aromatherapy Twice a day for 14 days and 20 assigned to standard of care. | Male 0% | NR | Low risk of bias | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information |
|--|--|----------------------------|--------------------------|------------------------------------|--|
| AXA1125 may in | | | | ity improvement. Howe | ver, certainty of the |
| | | evidence was low. Fu | Titler research is need | ea. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Finnigan et al (20); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 21 assigned to AXA1125 33.9 gr twice a day for 4 weeks and 20 assigned to standard of care. | Mean age 43.6, male 31.7%, | NR | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: RR 1.07 (95% CI 0.79 to 1.44); |

| | | | | | RD 5.1% (95% CI |
|---|---|--|-------------------------------------|--|--|
| | | | | | -16.6% to 34.5%); Low certainty ⊕⊕○○ |
| | | | | | Functional capacity improvement: RR 0.87 (95% CI 0.51 to 1.48); RD - 8.1% (95% CI - 30% to 29.3%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ |
| | | | | | Strength improvement: No information |
| | | | | | Adverse events: No information |
| | | | | | Severe adverse events: No information |
| CCSA | (ethylmethylhy | vdroxypyridine | trimethylhyd | rosinium propio | nate and |
| | | | acid anion) | | |
| Study; publication status | Uncertainty Patients and | succinate | acid anion) | | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| Study; | Uncertainty Patients and interventions | Succinate y in potential benefits a Comorbidities | e acid anion) nd harms. Further res | earch is needed. | Interventions effects vs standard of care (SOC) and GRADE certainty of |
| Study; | Patients and interventions analyzed Patients with post COVID-19 condition (asthenia | Succinate y in potential benefits a Comorbidities | Additional interventions | earch is needed. | Interventions effects vs standard of care (SOC) and GRADE certainty of |
| Study; publication status Tanashyan et al (21); Peer | Patients and interventions analyzed Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 15 assigned to CCSA | Succinate in potential benefits a Comorbidities Mean age 35, male | Additional interventions | earch is needed. Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence HRQL improvement: No |
| Study; publication status Tanashyan et al (21); Peer | Patients and interventions analyzed Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 15 | Succinate in potential benefits a Comorbidities Mean age 35, male | Additional interventions | earch is needed. Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence HRQL improvement: No information Overall symptom improvement: No |

| | _ Uncertaint\ | Coenzyme o in potential benefits a | e Q10 (CQ10) nd harms. Further reso | earch is needed. | information Strength improvement: No information Adverse events: No information Severe adverse events: No information |
|---|--|--|--|---------------------------------------|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Hansen et al. (22); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 84 days of acute COVID-19). 59 assigned to coenzyme Q10 500 mg a day for 6 weeks and 60 assigned to standard of care. | Median age 49, male 25.2%, obesity 33.6%, interval between COVID-19 and enrolment 288.55 days, hospitalization during COVID-19 15.1% | | Low risk of bias | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: Very low certainty ⊕○○○ Fatigue improvement: No information Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information |

| Cognitive beha | avioral therapy may in | crease fatigue improve | navioral therap ment and functional c eeded. | D y apacity improvement. F | urther research is |
|--|--|--|--|---|--|
| publication status | interventions analyzed | | interventions | limitations | effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | · | КСТ | | |
| Kuut et al. (23) Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 57 assigned to Cognitive behavioral therapy for 17 weeks and 57 assigned to standard of care. | Mean age 46, male 27%, interval between COVID-19 and enrolment 188 days, hospitalization during COVID-19 11.5% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: RR 2.2 (95% CI 1.35 to 3.58); RD 31.6% (95% CI 9.2% to 68%); Low certainty ⊕⊕○○ Functional capacity improvement: RR 1.37 (95% CI 1.08 to 1.73); RD 22.4% (95% CI 1.08 to 44.7%); Low certainty ⊕⊕○○ Strength improvement: No information Adverse events: No information Severe adverse |

| Cytofla | vin may not improve fa | | oflavin nty of the evidence wa | s low. Further research | events: No information |
|--|---|---|-----------------------------------|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| CITADEL trial (24), Putilina et al.; Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 30 to 90 days of acute COVID-19). 50 assigned to cytoflavin 2 tablets a day for 25 days and 50 assigned to standard of care. | Mean age 40.4 ± 12, male 57%, hypertension 38%, diabetes 4% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: RR 1.02 (95% CI 0.98 to 1.06); RD 2.1% (95% CI -1.9% to 6.2%); Low certainty + Compared to the compared to |

| | Uncertainty | Echino in potential benefits a | chrome A nd harms. Further reso | earch is needed. | |
|--|---|-----------------------------------|------------------------------------|------------------------------------|---|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Brichetti et al (25) Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 24 assigned to Echinochrome A and 22 assigned to standard of care. | Age between 18 and 60, male 40%, | NR | Low risk of bias Notes: | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information |

Enzymes + probiotics Enzymes + probiotics may improve fatigue. However, certainty of the evidence was low. Further research is needed. Patients and Comorbidities Additional Risk of bias and study Interventions Study; publication status interventions interventions limitations effects vs standard analyzed of care (SOC) and GRADE certainty of the evidence **RCT** Patients with P-Mean age 41.2 ± 13, NR High risk of bias **HRQL** Rathi et al. (26); ACC (asthenia or male 63.5%, interval improvement: No Peer reviewed; fatigue after acute between COVID-19 Notes: Concealment information 2022 and enrolment 19.5 COVID-19). 100 of allocation and assigned to days, one blinding probably Overall symptom enzymes + comorbidity 14.5% inappropriate. improvement: No probiotics information **ImmunoSEB** (500 mg/capsule) + **Fatigue** ProbioSEB CSC3 improvement: (5 billion CFUs RR 6.07 (95% CI /capsule) and 100 3.79 to 9.71); assigned to RD 76% (95% CI 41.8% to 85%); standard of care. Low certainty $\Theta\Theta$ **Functional** capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information

Fermented food supplements

Uncertainty in potential benefits and harms. Further research is needed.

| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | |
|---|--|---|-----------------------------|--|--|--|
| RCT | | | | | | |
| Kharaeva et al. (27); Peer reviewed; 2022 | Patients with P-ACC after moderate infection (asthenia or fatigue after acute COVID-19). 68 assigned to fermented food supplements 14 g twice a day for 20 days and 29 assigned to standard of care. | Age 38–69, male 51.5%, hypertension 36.1%, diabetes 15.5%, chronic lung disease 14.4%, obesity 19.6%, hospitalization during COVID-19 46.4% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: Very low certainty ⊕○○○ Fatigue improvement: No information | |
| Kharaeva et al. (27); Peer reviewed; 2022 | Patients with P-ACC after severe infection (asthenia or fatigue after 0 days of acute COVID-19). 64 assigned to fermented food supplements 14 g twice a day for 20 days and 27 assigned to standard of care. | Age 36–65, male 47.2%, diabetes 28.6%, chronic lung disease 20.9%, asthma 3.3%, chronic heart disease 37.5%, obesity 40.6%, hospitalization during COVID-19 41.8% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information | |
| Hydrogen (nasal) Uncertainty in potential benefits and harms. Further research is needed. | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | |

| RCT | | | | | | |
|---|--|--|--------------------------|--|--|--|
| Botek et al. (28); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 21 to 35 days of acute COVID-19). 26 assigned to hydrogen (nasal) 300 mL/min for 14 days and 24 assigned to standard of care. | Mean age 40, male 52%, interval between COVID-19 and enrolment 25 days | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: Very low certainty ⊕○○○ Strength improvement: No information Adverse events: No information Severe adverse events: No information | |
| Immunodaat Uncertainty in potential benefits and harms. Further research is needed. | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | |
| RCT | | | | | | |

| Deshpande trial (29); Preprint; 2022 | Patients with post COVID-19 condition. 26 assigned to Immunodaat 500 mg a day for 30 days and 28 assigned to standard of care. | Mean age 38.9, male 59.4% Lero | nlimab | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information | |
|---|---|---------------------------------|--------------------------|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | |
| RCT | | | | | | |
| Gaylis et al. (30); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 90 days of acute COVID-19). 27 assigned to Leronlimab 700 mg a week for 8 weeks and 26 assigned to standard of care. | NR | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: Very low certainty ⊕○○○ Fatigue improvement: No information | |

| | | | | | Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information | |
|--|---|---------------------------|--------------------------|---|--|--|
| | Mindfulness training Uncertainty in potential benefits and harms. Further research is needed. | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | |
| | | ı | RCT | | | |
| Hausswirth et al (31); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 90 days of acute COVID-19). 17 assigned to a mindfulness based | Mean age 47.9, male 26.5% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Fatigue | |

| Phy | sical training may im | | al training and functional capacit | y. Further research is n | improvement: No information Adverse events: No information Severe adverse events: No information |
|--|---|--------------------------------|---------------------------------------|--|--|
| publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Nambi et al. (32); Peer reviewed; 2022 | Patients with P-ACC (sarcopenia after acute COVID-19). 36 assigned to aerobic training (high intensity) and 37 assigned to aerobic training (standard intensity). | Mean age 63.5, male 100% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: RR 1.62 (95% CI 1.07 to 2.45); RD 21.1% (95% CI 2.4% to 49.2%); Low certainty ⊕⊕○○ |
| Blanco et al. (33) Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 40 days of acute COVID-19). 24 assigned to endurance training rehabilitation (ETR) (10 breathing and strength-based exercises) for 14 days, and 24 assigned to standard of care. | Mean age 40.7, male 22.91% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕ ○ ○ Functional capacity improvement: RR 1.81 (95% CI 1.23 to 2.65); RD 17.4% (95% CI 4.09 to 25.49 ()) |
| (34) Jimeno- Almazán et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 43 assigned to physical training 3 days a week for 8 | Mean age 45.3 ± 8, male 31% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | 4.9% to 35.4%); Low certainty ⊕⊕⊖⊖ Strength improvement: RR 3.13 (95% CI 1.02 to 9.55); RD 11.3% (95% CI |

| Ibrahim et al. (35) Peer reviewed; 2022 | weeks and 20 assigned to standard of care. Patients with post COVID-19 condition (asthenia or fatigue after acute COVID-19). 24 assigned to physical training 4 times a week for 10 weeks and 24 assigned to standard of care. | Mean age 63, male 39.6%, diabetes 81.2%, chronic lung disease 66.6%, CHD 83.3%, cancer 12.5%, | NR | High risk of bias Notes: Concealment of allocation and blinding probably inappropriate. | 0.1% to 45.4%); Low certainty ⊕⊕○○ Adverse events: No information Severe adverse events: No information |
|---|---|--|----|---|---|
| Rasmussen et al (36) Peer reviewed; 2022 | Patients with post COVID-19 condition. 14 assigned to Aerobic training (high intensity) three time a week for 12 weeks and 14 assigned to standard of care. | days, hospitalization | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | |
| Oliveira et al;(37) Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after acute COVID-19). 31 assigned to physical training 60-minutes, twiceweekly for 12 weeks and 28 assigned to standard of care. | Mean age 52.3 ± 11.9, male 42.4%, hypertension 44.1%, diabetes 10.6%, CHD 13.6% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | |
| Ogonowska- Slodownik et al (38); Peer reviewed; 2022 | Pediatric patients with post COVID-19 condition (asthenia or fatigue after acute COVID-19). 48 assigned to Physical training twice a week, 45 minutes for 8 weeks and 26 assigned to standard of care. | Mean age 10.8, male 41.9% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | |

| Espinoza-Bravo et al (39); Preprint; 2022 Phytochemica Study; publication status | Patients and | | hemicals | High risk of bias Notes: Non-blinded study which might have introduced bias. nce was low. Further res Risk of bias and study limitations | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty ⊕○○○ Functional capacity improvement: Very low certainty ⊕○○○ Strength improvement: No information Adverse events: No information Severe adverse events: No information Severe adverse events: No information Severe adverse events: No information |
|--|---|---|----------|---|--|
| | | F | RCT | | |
| UK Phyto-V trial (40) Thomas et al; Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after acute COVID-19). 74 assigned to phytochemicals one capsule a day | Mean age 53, male 56%, obesity 35%, interval between COVID-19 and enrolment 108 days, hospitalization during COVID-19 63% | NR | High risk of bias Notes: Concealment of allocation and blinding probably inappropriate. | HRQL improvement: RR 1.33 (95% CI 1.03 to 1.71); RD 18% (95% CI 1.8% to 39%); Low certainty $\oplus \oplus \bigcirc$ |

| | <u> </u> | <u> </u> | | | |
|---------------------------------------|---|--|--|------------------------------------|---|
| | and 73 assigned to standard of care. | | | | Overall symptom improvement: No information Fatigue improvement: RR 1.24 (95% CI 0.95 to 1.62); RD 13.1% (95% CI -2.5% to 33.5%); Low certainty ⊕⊕○○ Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No |
| | | | | | information |
| | | D | | | |
| | Uncertainty | Prol in potential benefits a | DIOTICS nd harms. Further rese | earch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Caprioli et al (41) Preprint; 2022 | COVID-19 condition (asthenia or fatigue after 28 days of acute | Mean age 54.1 ± 7.9, male 52.6%, interval between COVID-19 and enrolment 756 days, hospitalization during COVID-19 57.9% | NR | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Fatigue improvement: Very low certainty |

| tDCS may improv | | | | ion (tDCS) er, certainty of the evide | Functional capacity improvement: No information Strength improvement: No information Adverse events: No information Severe adverse events: No information on information |
|---|---|---|--------------------------|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Oliver-Mas et al. (42); Preprint; 2022 | Patients with P-ACC (asthenia or fatigue after 180 days of acute COVID-19). 23 assigned to transcranial direct current stimulation (tDCS) 1 session a week for 8 weeks and 24 assigned to standard of care. | Mean age 45.6, male 21.3%, hypertension 12.8%, diabetes 4.3%, interval between COVID-19 and enrolment 620 days, hospitalization during COVID-19 14.9% | NR | Low risk of bias | HRQL improvement: RR 1.37 (95% CI 1.09 to 1.71); RD – 26% (95% CI – 6.7% to 30%); Low certainty ⊕⊕⊖⊖ Overall symptom improvement: No information |
| Santana et al (43); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 90 days of acute COVID-19). 35 assigned to transcranial direct current stimulation (tDCS) 10 sessions and 35 assigned to standard of care. | Mean age 53, male 35.7%, hypertension 17.1%, diabetes 14.3%, chronic lung disease 5.7%, CHD 7.1%, , hospitalization during COVID-19 25.7% | NR | Low risk of bias | Fatigue improvement: RR 1.36 (95% CI 0.76 to 2.45); RD − 16.9% (95% CI − 11.2% to 53%); Low certainty ⊕⊕○○ Functional capacity improvement: No information Strength |

| Telerehabilitation | n may improve fatigue | and functional capacity | abilitation y. However, certainty deeded. | of the evidence was low | improvement: No information Adverse events: RR 0.83 (95% CI 0.26 to 2.73); RD − 3.4% (95% CI − 15.5% to 36%); Low certainty ⊕⊕○○) Severe adverse events: No information |
|---|---|---|--|--|---|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | КСТ | | |
| King et al. (44); Preprint; 2022 | Patients with P-ACC (asthenia or fatigue after 110 days of acute COVID-19). 11 assigned to telerehabilitation twice weekly for 10 weeks and 10 assigned to standard of care. | Mean age 48.5 ± 13, male 47.6%, interval between COVID-19 and enrolment 366 days, hospitalization during COVID-19 19% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: Very low certainty Overall symptom improvement: No information Fatigue improvement: |
| Simpson et al (45); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 14 days of acute COVID-19). 15 assigned to telerehabilitation 45 to 60 min sessions, twice a week for 4 weeks and 12 assigned to standard of care. | Mean age 58 ± 12, male 58%, interval between COVID-19 and enrolment 14 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | RR 1.63 (95% CI 1.04 to 2.54); RD 19.1% (95% CI 1.2% to 46.8%); Low certainty ⊕⊕○○ Functional capacity improvement: RR 1.37 (95% CI 1.11 to 1.67); RD 13.8% (95% CI 2% to 29.8%); Low |
| Longobardi et al. (46); Peer | Patients with post COVID-19 | Mean age 61, male 50%, hypertension | NR | High risk of bias | certainty ⊕⊕○○ |

| reviewed; 2022 | condition (asthenia or fatigue after 84 days of acute COVID-19). 21 assigned to home physical training 3 times a week for 16 weeks and 20 assigned to standard of care. | 56%, diabetes 36%, chronic lung disease 16%, CHD 20%, obesity 28%, interval between COVID-19 and enrolment 159 days, hospitalization during COVID-19 100% | | Notes: Non-blinded study which might have introduced bias. | Strength improvement: No information Adverse events: No information Severe adverse events: No information |
|---|---|--|---|--|---|
| Samper-Pardo et al. (47); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 84 days of acute COVID-19). 52 assigned to Telerehabilitation through specificaly designed app for 12 weeks and 48 assigned to standard of care. | Mean age 48.3 ± 9.26, male 20%, hypertension %, diabetes %, chronic lung disease %, asthma %, CHD %, CKD %, cancer %, obesity %, interval between COVID-19 and enrolment days, hospitalization during COVID-19 % | %, lopinavir- ritonavir %, tocilizumab %, azithromycin %, convalescent plasma %; Vaccinated % | High risk of bias Notes: Non-blinded study which might have introduced bias. | |
| Da Silva et al. (48); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after acute COVID-19). 28 assigned to Telerehabilitation three time a week for 8 weeks and 29 assigned to standard of care. | Mean age 55 ± 11, male 56%, diabetes 42%, chronic lung disease 16%, CHD 65%, obesity 60% | NR | Low risk of bias | |
| Tanhan et al. (49); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 0 days of acute COVID-19). 16 assigned to synchronous telerehabilitation three day a week and 16 assigned to asynchronous telerehabilitation | Mean age 54, male 46.9%, interval between COVID-19 and enrolment 49 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Fatigue improvement: No information Functional capacity improvement: Very low certainty ⊕○○○ |

| | | Strength improvement: No information |
|--|--|--|
| | | Adverse events: No information |
| | | Severe adverse events: No information |

Table 3. Description of included studies and interventions effects for P-ACC-related dyspnea

| ADAPT-232 (adaptogens) ADAPT-232 may not improve fatigue. However, certainty of the evidence was low. Further research is needed. | | | | | | | | |
|--|---|-------------------------|--------------------------|---------------------------------------|--|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | F | RCT | | | | | |
| Karosanidze et al. (16); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 30 days of acute COVID-19). 49 assigned to ADAPT-232 (adaptogens) 60 mL a day for 14 days and 50 assigned to standard of care. | Mean age 48.9, male 14% | NR | Low risk of bias | HRQL improvement: No information Dyspnea improvement: RR 1. (95% CI 0.94 to 1.06); RD 0% (95% CI - 5.4% to 5.6%); Low certainty Tunctional capacity improvement: No information Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse events: No information | | | |

Albumin (inhaled)
Uncertainty in potential benefits and harms. Further research is needed.

| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
|--|---|--------------------------------|--------------------------|---|---|--|--|
| | | F | RCT | | | | |
| Ampio trial; Other; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 28 days of acute COVID-19). 15 assigned to albumin (inhaled) and 16 assigned to standard of care. | Mean age 52.1 ± 13, male 43.8% | NR | High risk of bias Notes: Concealment of allocation and blinding probably inapropriate. | HRQL improvement: No information Dyspnea improvement: No information Functional capacity improvement: No information Pulmonary function improvement: No information Radiological response: No information Adverse events: Very low certainty ⊕○○○ Severe adverse events: No information | | |
| Antifibrotics Uncertainty in potential benefits and harms. Further research is needed. | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | ı | RCT | | | | |

| Kerget et al. (50); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 84 days of acute COVID-19). 15 assigned to pirfenidone 600 to 1800 mg a day for 3 months and 15 assigned to nintendanib 300 mg a day for 3 months | Mean age 65.6, male 40% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Dyspnea improvement: No information Functional capacity improvement: Very low certainty ⊕○○○ Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse events: No |
|---|---|---|-------------------------------------|--|---|
| | Uncertainty | Diaphra in potential benefits a | gm release nd harms. Further res | earch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Nagy et al; (51); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities acute COVID-19). 26 assigned to diaphragm release three sessions a week for 6 weeks and 26 assigned to standard of care. | Mean age 40 | NR | High risk of bias Notes: Concealment of allocation and blinding probably inapropriate. | HRQL improvement: No information Dyspnea improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information |

| | | | | | Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse events: No information |
|---|--|--------------------------------|---|--|--|
| | | | oivolol | | |
| Study; publication status | Patients and | c in potential benefits a | nd harms. Further reso Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Dal Negro et al (52); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 84 days of acute COVID-19). 8 assigned to Nebivolol 2.5 mg a day and 8 assigned to standard of care. | Mean age 50.5 ± 17.2, male 63% | NR | High risk of bias Notes: Concealment of allocation and blinding probably inappropriate. | HRQL improvement: No information Dyspnea improvement: Very low certainty ⊕○○○ Functional capacity improvement: No information Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse |

| | | | | | events: No | | | | | |
|---|--|---|--------------------------|---|---|--|--|--|--|--|
| | Respiratory training/rehabilitation Respiratory training probably improves HRQL, dyspnea and functional capacity | | | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | | | |
| | | F | RCT | | | | | | | |
| ENO Breathe trial (53), Philip et al.; Peer reviewed; 2022 | Patients with P-ACC (dyspnea and/or lung radiological abnormalities after 30 days of acute COVID-19). 58 assigned to ENO Breathe 6-week program and 71 | Mean age 49.5 ± 12, male 17.3%, interval between COVID-19 and enrolment 320 days, hospitalization during COVID-19 17.3% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: RR 1.41 (95% CI 1.18 to 1.67); RD 22.3% (95% CI 9.8% to 36.5%); Moderate certainty ⊕⊕⊕⊖ | | | | | |
| | assigned to standard of care. | | | | improvement: RR 2.11 (95% CI 1.17 | | | | | |
| McNarry et al. (54); Peer reviewed; 2022 | Patients with P-ACC (dyspnea and/or lung radiological abnormalities after acute COVID-19). 37 assigned to inspiratory muscle training 3 sessions a week for 8 weeks and 37 assigned to standard of care. | Mean age 46.6 ± 12, male 12.8%, interval between COVID-19 and enrolment 270 days | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. Intention-to-treat (ITT) analysis for primary outcome not available. | to 3.8); RD 34.2% (95% CI 5.2% to 86.4%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Functional capacity improvement: RR 1.41 (95% CI 1.05 to 1.9); RD 20.5% (95% CI 2.5% to 44.9%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ | | | | | |
| Srinivasan et al. (55); Peer reviewed; 2022 | Patients with P-ACC (dyspnea and/or lung radiological abnormalities after acute COVID-19). 24 assigned to respiratory training 3 times a day for 6 weeks and 24 assigned to standard of care. | NR | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Pulmonary function improvement: Very low certainty ⊕○○○ Radiological response: No information Adverse events: | | | | | |

| Rodriguez- | Patients with P- | Mean age 40.7, male | NR | High risk of bias | No information |
|---|--|---|----|--|------------------------------|
| Blanco et al; (33) Peer reviewed; 2022 | ACC (asthenia or fatigue after 40 days of acute COVID-19). 24 assigned to respiratory training (10 breathing and strength-based exercises) for 14 days, and 24 assigned to standard of care. | 22.91% | | Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Severe adverse events: No |
| InsCOVID trial (56); Palau et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 90 days of acute COVID-19). 13 assigned to inspiratory muscle training twice a day for 12 weeks and 13 assigned to standard of care. | Mean age 50.4 ± 12.2, male 58%, hypertension 12%, interval between COVID-19 and enrolment 362 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | |
| RECOVER trial. (57), Romanet et al.; Preprint; 2022 | Patients with P-ACC (dyspnea and/or lung radiological abnormalities after 90 days of acute COVID-19). 27 assigned to endurance training rehabilitation (ETR) two (1 h) sessions per week for 10 weeks and 33 assigned to standard of care. | Mean age 58.2, male 61.6%, diabetes 36.7%, chronic lung disease 8.3%, chronic heart disease 5%, cancer 5%, interval between COVID-19 and enrolment 173 days, hospitalization during COVID-19 100% | | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | |
| RECOVE trial (34); Jimeno- Almazán et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 84 days of acute COVID-19). 40 assigned to respiratory training | Mean age 45.3 ± 8, male 31% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | |

| | twice a day for 8 weeks and 20 assigned to standard of care. | | | |
|---|--|--|-------------------------|---|
| Del Corral et al (58) Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 84 days of acute COVID-19). 44 assigned to Respiratory training home-based, 40 minutes a day, 6 days a week for 8 weeks and 44 assigned to standard of care. | Mean age 46.4, male 28.5%, interval between COVID-19 and enrolment 350.7 days, hospitalization during COVID-19 31.8% | Corticosteroids 5.7% | Low risk of bias |
| Nambi et al (59) Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 28 days of acute COVID-19). 68 assigned to Home pulmonary rehabilitation four time a week for 8 weeks and 68 assigned to standard of care. | Mean age 48, male 47.8%, hypertension 30.1%, chronic lung disease 5.9%, CHD 8%, obesity 15.4%, | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. |
| TERECO trial (60); Li et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 0 days of acute COVID-19). 55 assigned to home pulmonary rehabilitation 3 to 4 sessions a week for | Mean age 50.6 ± 10.98, male 44.5%, hypertension 21.9%, diabetes 13.6%, chronic lung disease 5.9%, CHD 7.6%, obesity 15.3%, interval between COVID-19 and enrolment 71 days, hospitalization during COVID-19 100% | NR | Low risk of bias Notes: |

| | 6 weeks and 50 assigned to standard of care. | | | | |
|---|--|---|----|--|---|
| Mirenayat et al (61); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 7 days of acute COVID-19). 26 assigned to Respiratory training Twice a day for 4 weeks and 26 assigned to standard of care. | Mean age 50.02, male 55.7%, | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | |
| Senén trial (62); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 84 days of acute COVID-19). 18 assigned to Respiratory training 2 daily sessions, 6 days a week for 8 weeks and 19 assigned to standard of care. | Mean age 47 ± 7.1, male 27%, hypertension 14%, diabetes 0%, chronic lung disease 8%, CKD 0%, interval between COVID-19 and enrolment 359 days | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | |
| Vallier et al; (63) Peer reviewed; 2022 | Patients with P-ACC (dyspnea and/or lung radiological abnormalities after acute COVID-19). 8 assigned to home pulmonary rehabilitation four times a week for 4 weeks and 9 assigned to inpatient rehabilitation four | Mean age 54.8 ± 16, male 70.6%, interval between COVID-19 and enrolment 141 days, hospitalization during COVID-19 76.5% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Dyspnea improvement: Very low certainty ⊕○○○ Functional capacity improvement: |

| Simpson et al (45); Peer reviewed; 2022 | times a week for 4 weeks Patients with post COVID-19 condition (asthenia or fatigue after 14 days of acute COVID-19). 15 assigned to telerehabilitation 45 to 60 min sessions, twice a week for 4 weeks and 12 assigned to standard of care. | Mean age 58 ± 12, male 58%, interval between COVID-19 and enrolment 14 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Very low certainty October 19 Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse events: No information |
|---|---|---|----|--|---|
| Rutkowski et al (64); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after of acute COVID-19). 18 assigned to VR respiratory training five sessions a week for 3 weeks and 14 assigned to conventional respiratory training. | Mean age 57.8 ± 4.9, male 37.5% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: No information Dyspnea improvement: Very low certainty ⊕○○○ Functional capacity improvement: Very low certainty ⊕○○○ |
| Stavrou et al (65); Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung radiological abnormalities after 60 days of acute COVID-19). 10 assigned to VR respiratory training and 10 assigned to standard of care. | Mean age 53.9, male 80%, interval between COVID-19 and enrolment 60 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | Pulmonary function improvement: No information Radiological response: No information Adverse events: No information Severe adverse events: No information |

| Kusumawardani | Patients with post | Mean age 46, male | NR | High risk of bias | HRQL |
|---|---|---|--|---|---|
| et al (66); Peer reviewed; 2022 COVID-19 condition (dyspnea and/or lung | obesity 55%, interval | | Notes: Non-blinded study. Concealment | improvement: No information | |
| | radiological abnormalities after acute COVID-19). | between COVID-19 and enrolment 22.5 days, hospitalization | | of allocation probably inappropriate. | Dyspnea improvement: No information |
| | 10 assigned to incentive spirometry 5 times a day for four weeks and 10 | during COVID-19 100% | | | Functional capacity improvement: No information |
| | assigned to conventional respiratory training. | | | | Pulmonary function improvement: Very low certainty ⊕○○○ |
| | | | | | Radiological response: No information |
| | | | | | Adverse events: No information |
| | | | | | Severe adverse events: No information |
| High dose steroid | ls may not improve dy | spnea and may not inc | (high dose) rease adverse events. earch is needed. | . However, certainty of tl | he evidence was low. |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | ı | RCT | | |
| COLDSTER trial (67); Dhooria et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (dyspnea and/or lung | Mean age 57, male 68%, one commorbiditie 73% | NR | High risk of bias Notes: Non-blinded study which might | HRQL improvement: No information |
| | radiological abnormalities after 21 to 49 days of acute COVID-19). 65 assigned to High dose steroids Prednisone 40 mg a day descending | | | have introduced bias. | Dyspnea improvement: RR 1 (95% CI 0.87 to 1.15); RD 0% (95% CI −11.1% to 12.7%); Low certainty ⊕⊕⊖⊖ |
| | progressively to 10 | | | | Functional |

| | mg a day for 6 weeks and 65 assigned to standard of care. | | | | capacity improvement: No information Pulmonary function |
|---|--|---------------------------------|-----------------------------|---|--|
| | | | | | improvement: No information |
| | | | | | Radiological response: Very low certainty ⊕○○○ |
| | | | | | Adverse events: RR 0.92 (95% CI 0.75 to 1.13); RD − 6.2% (95% CI − 19.3% to 10%); Low certainty ⊕⊕○○ |
| | | | | | Severe adverse events: Very low certainty ⊕○○○ |
| Treamid may imp | | | | onal capacity. Treamid n research is needed. | nay increase adverse |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Bazdyrev et al. (68); Peer reviewed; 2022 | Patients with P- ACC (dyspnea and/or lung radiological | Mean age 55 ± 11, male 44.1% | NR | Low risk of bias | HRQL improvement: No information |
| | abnormalities after acute COVID-19). 29 assigned to treamid 50 mg a day for 28 days and 30 assigned to standard of care. | | | | Dyspnea improvement: RR 1.96 (95% CI 0.9 to 4.25); RD 21.7% (95% CI −2.3% to 73.7%); Low certainty ⊕⊕○○ |
| | | | | | Functional capacity improvement: |

| | | | RR 1.10 (95% CI 0.64 to 1.90); RD 4.3% (95% CI −16.2% to 39.8%); Low certainty ⊕⊕⊖⊖ |
|--|--|--|---|
| | | | Pulmonary function improvement: RR 2.48 (95% CI 1 to 6.17); RD 24.7% (95% CI 0% to 86.1%); Low certainty $\oplus \oplus \bigcirc$ |
| | | | Radiological response: Very low certainty ⊕○○○ |
| | | | Adverse events: RR 1.19 (95% CI 0.56 to 2.50); RD − 5.5% (95% CI − 12.7% to 43.6%); Low certainty ⊕⊕⊖⊖ |
| | | | Severe adverse events: No information |

Table 4. Description of included studies and interventions effects for PCC neurocognitive symptoms or sleep disturbances

| Actovegir | Actovegin Actovegin may improve cognition. However, certainty of the evidence was low. Further research is needed. | | | | | | | |
|---|--|----------------------------|---------------------------------|--|---|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | R | СТ | | | | | |
| Kutashov et al. (15); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after NR days of acute COVID-19). 222 assigned to Actovegin 1200 mg a day for 60 days and 222 assigned to standard of care. | Mean age 67.6, male 31.98% | NR | High risk of bias Notes: Non- blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: RR 1.19 (95% CI 1.06 to 1.33); RD 12.7% (95% CI 4.2% to 22.3%); Low certainty ⊕⊕⊖⊖ Depression improvement: No information Adverse events: No information Severe adverse events: No information | | | |
| | Uncertainty in | Celeo | COXID I harms. Further resea | arch is needed. | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | R | CT | | | | | |

| Ansari trial (69); Peer reviewed; 2022 | Patients with post COVID-19 condition (depression after 20 days of acute COVID-19). 31 assigned to celecoxib 200 mg a day for 6 weeks and 31 assigned to standard of care. | Mean age 27.8, male 61%, hypertension 11.3%, interval between COVID-19 and enrolment 87.5 days, | Corticosteroids 25.8%, remdesivir 11.3%, | High risk of bias Notes: Concealment of allocation and blinding probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: No information Depression improvement: Very low certainty OAdverse events: No information Severe adverse events: No information |
|--|--|--|--|---|--|
| | Uncertainty in | Electric s | timulation d harms. Further resea | arch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | R | СТ | | |
| Zulbaran-Rojas et al (70); Peer reviewed; 2022 | Patients with post COVID-19 condition (neurocognitive after acute COVID-19). 10 assigned to Electrical stimulation and 8 assigned to standard of care. | Mean age 51.7, male 27.8%, hypertension 44.4%, diabetes 33.3%, interval between COVID- 19 and enrolment 299 days, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non- blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: No information Depression improvement: No information Adverse events: |

| | | | | | Severe adverse events: No information | | | | |
|--|--|---|---|---------------------------------------|---|--|--|--|--|
| Famotidin | e may improve cognit | | otidine by of the evidence was | s low. Further research | is needed. | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | | |
| | | R | СТ | | | | | | |
| Momtazmanesh et al (71); Peer reviewed; 2022 | Patients with post COVID-19 condition (neurocognitive after 20 days of acute COVID-19). 25 assigned to famotidine 80 mg a day and 25 assigned to standard of care. | Mean age 36.3, male 54%, hypertension 12%, diabetes 4%, CHD 2%, cancer 8%, obesity 22%, interval between COVID-19 and enrolment 29 days, hospitalization during COVID-19 100% | Corticosteroids 16%, remdesivir 84% | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: RR 1.33 (95% CI 1.05 to 1.69); RD 24.6% (95% CI 3.7% to 51.1%); Low certainty ⊕⊕⊖⊖ Depression improvement: No information Adverse events: No information Severe adverse events: No information | | | | |
| | may improve UDOL | | oxygen (HBO) | Further receased in the | odod | | | | |
| НВО | HBO may improve HRQL. However, certainty of the evidence was low. Further research is needed. | | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | | |
| | | R | СТ | | | | | | |

| Zilberman- Itskovich et al. (72); Peer reviewed; 2022 | Patients with P-ACC (neurocognitive symptoms after 90 days of acute COVID-19). 37 assigned to HBO 1 session a day for 40 days and 36 assigned to standard of care. | Mean age 48, male 39.7%, hypertension 8.2%, diabetes 2.7%, chronic lung disease 0%, asthma 4.1%, cancer 0%, obesity 27.4%, interval between COVID-19 and enrolment 165 days, hospitalization during COVID-19 16.4% | NR | Low risk of bias | HRQL improvement: RR 1.30 (95% CI 0.84 to 2); RD 13.9% (95% CI −7.4% to 46.9%); Low certainty ⊕⊕⊖⊖ Overall symptom improvement: No information Cognitive improvement: Very low certainty ⊕⊖⊖⊖ Depression improvement: Very low certainty ⊕⊖⊖⊖ Adverse events: No information Severe adverse events: No information |
|--|--|--|---------------------------------------|---|---|
| | Uncertainty | Mindfuln in potential benefits a | ess training nd harms. Further res | earch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Hausswirth et al (31); Peer reviewed; 2022 | Patients with post COVID-19 condition (asthenia or fatigue after 90 days of acute COVID-19). 17 assigned to a mindfulness based intervention (Rebalance®) 2 to | Mean age 47.9, male 26.5% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: No |

| | 3 sessions (30 min) a week for 4 weeks and 17 assigned to standard of care. | | | | information Depression improvement: Very low certainty ⊕○○○ Adverse events: No information Severe adverse events: No information |
|---|---|--|--------------------------|--|--|
| | | Palmitoylethan in potential benefits a | | | |
| Study; publication status | Patients and | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | , | i | RCT | | |
| Versace et al (73); Peer reviewed; 2022 | Patients with post COVID-19 condition (neurocognitive after acute COVID-19). 17 assigned to Palmitoylethanolam ide + Luteolin 1400/400mg a day for 8 weeks and 17 assigned to standard of care. | Mean age 50.8, male 35.3% | NR | High risk of bias Notes: pseudo- randomized | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: Very low certainty ⊕○○○ Depression improvement: No information Adverse events: No information Severe adverse events: No information |

| | Physical training Uncertainty in potential benefits and harms. Further research is needed. | | | | | | | |
|--|--|---|--------------------------|--|--|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | | RCT | | | | | |
| Kalayeh et al (74); Preprint; 2022 | Patients with post COVID-19 condition (sleep disturbances after 84 days of acute COVID-19). 17 assigned to endurance training rehabilitation (ETR) Three times a week for eight weeks and 15 assigned to standard of care. | Mean age 25, male 100%, interval between COVID-19 and enrolment 165 days, | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Cognitive improvement: No information Depression improvement: No information Sleep quality improvement: Very low certainty ⊕○○○ Adverse events: No information Severe adverse events: No information | | | |
| | | Ous auricular value of the potential benefits a | | mulation (taVNS earch is needed. | 5) | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | | RCT | | | | | |

| Badran et al. (75); Preprint; 2022 | | | | ion (tDCS) ertainty of the evidence | improvement: No information Overall symptom improvement: No information Cognitive improvement: No information Depression improvement: No information Adverse events: No information Severe adverse events: No information was low. Further |
|--|---|---|--------------------------|---------------------------------------|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | I | RCT | | |
| Oliver-Mas et al. (42); Preprint; 2022 | Patients with P-ACC (asthenia or fatigue after 180 days of acute COVID-19). 23 assigned to transcranial direct current stimulation (tDCS) 1 session a week for 8 weeks and 24 assigned to standard of care. | Mean age 45.6, male 21.3%, hypertension 12.8%, diabetes 4.3%, interval between COVID-19 and enrolment 620 days, hospitalization during COVID-19 14.9% | NR | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Cognitive improvement: RR 0.59 (95% CI 0.33 to 1.05); RD -27.5% (95% CI -44.8% to 3.4%); Low certainty $\oplus \oplus \bigcirc$ Depression |

| | | information |
|--|--|---------------------------------------|
| | | Adverse events: No information |
| | | Severe adverse events: No information |

Table 5. Description of included studies and interventions effects for PCC olfactory and/or gustatory dysfunction

| ADAPT-232 ma | ADAPT-232 (adaptogens) ADAPT-232 may not improve olfactory symptoms. However, certainty of the evidence was low. Further research is needed. | | | | | | |
|--|---|---------------------------|-----------------------------|---------------------------------------|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| Karosanidze et al. (16); Peer reviewed; 2022 | Patients with P-ACC (asthenia or fatigue after 30 days of acute COVID-19). 49 assigned to ADAPT-232 (adaptogens) 60 mL a day for 14 days and 50 assigned to standard of care. | Mean age 48.9, male 14% | NR | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: RR 0.89 (95% CI 0.79 to 1.01); RD − 10.3% (95% CI − 20.5% to 1.4%); Low certainty ⊕⊕○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information | | |
| | | Diod | e laser | | | | |
| | Uncertainty | / in potential benefits a | nd harms. Further reso | earch is needed. | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |

| Shabaan et al;(76); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 28 days of acute COVID-19). 18 assigned to Diode laser 6-minute session and 18 assigned to standard of care. | Mean age 41.5, male 22.2%, hospitalization during COVID-19 22.2% | NR | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty Outline Companies Gustatory symptoms improvement: No information Adverse events: No information Severe adverse |
|---|--|---|--------------------------|---|--|
| | | | | | events: No information |
| | | (Ethylene Diar gabapentin fatty acids a | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Abdelazim et al (77); Peer reviewed; 2022 | Patients with post COVID-19 condition (olfactory and/or gustatory dysfunction after 168 days of acute COVID-19). 25 assigned to EDTA 0.1 ml for 3 months and 25 assigned to standard of care. | Mean age 40.5, male 38%, hypertension 16%, diabetes 22%, asthma 6%, enrolment 180 days, hospitalization during COVID-19 % | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty Gustatory symptoms |

| Study; publication status | Patients and | Gabagabapentin fatty acids a | apentin are uncertain. Further Additional interventions | research is needed. Risk of bias and study limitations | information Adverse events: No information Severe adverse events: No information Interventions effects vs standard of care (SOC) and |
|--|---|-----------------------------------|--|---|---|
| | | | | | GRADE certainty of the evidence |
| | | F | RCT | | |
| Mahadev et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (olfactory and/or gustatory dysfunction after 84 days of acute COVID-19). 18 assigned to gabapentin 900 to 3600 mg a day for 4 weeks and 26 assigned to standard of care. | Mean age 43, male 25% | NR | High risk of bias Notes: Significant lost to follow-up | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty ⊕○○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information |
| | Olfactory traini | Olfacto ng may improve olfacto | ry training ory symptoms. Further | research is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |

| | RCT | | | | | | |
|---|---|---|---------------------------|--|---|--|--|
| Di Stadio et al. (79); Peer reviewed; 2022 | Patients with P-ACC (olfactory and/or gustatory dysfunction after 180 days of acute COVID-19). 76 assigned to olfactory training and 88 assigned to standard of care. | Mean age 40.7, male 27.6%, hypertension 1.7%, diabetes 0%, chronic heart disease 5.2% | | High risk of bias Notes: Non-blinded study which might have introduced bias. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms | | |
| Pires et al. (80); Preprint; 2022 | Patients with P-ACC (olfactory and/or gustatory dysfunction after 30 days of acute COVID-19). 26 assigned to advanced olfactory training with 8 essential oils: rose, eucalyptus, clove and lemon, citronella, mint, vanilla and cedarwood and 54 assigned to standard of care. | Mean age 37.6, male 35% | Steroids (nasal) 23.8% | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | improvement: Very low certainty ⊕○○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information | | |
| COVANOS trial (81), Lechner et al; Peer reviewed; 2022 | Patients with P-ACC (olfactory and/or gustatory dysfunction after 30 days of acute COVID-19). 25 assigned to olfactory training for 12 weeks and 26 assigned to standard of care. | Mean age 44, male 13.8%, hypertension 8.9%, diabetes 1.1%, chronic lung disease 0%, asthma 12.6%, chronic heart disease 0%, cancer 2.1% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | | | |
| Chung et al. (82); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfatory or gustatory disfunction after 84days of acute COVID-19). 8 assigned to olfactory training thrice daily for 4 weeks and 5 assigned to | Mean age 52.5, male 30.7%, cancer 15.4%, interval between COVID-19 and enrolment 154 days, | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | | | |

| Study; publication status | Patients and | Omega-3 fomega 3 fatty acids an Comorbidities | Fatty Acids re uncertain. Further re | esearch is needed. Risk of bias and study limitations | effects vs standard of care (SOC) and GRADE certainty of |
|---|--|---|--------------------------------------|--|---|
| | | | | | the evidence |
| | | F | RCT | | |
| Lerner et al (83) Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after acute COVID-19). 57 assigned to Omega 3 2000 mg a day and 60 assigned to standard of care. | Mean age 41, male 21.4% | Corticosteroids 6.6% | Low risk of bias | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty ⊕○○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information |
| Palmitoylethanolar | | Palmitoylethand | | eolin nty of the evidence was | low. Further research |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | ı | СТ | | |

| Di Stadio et al; (79) Peer reviewed; 2023 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 180 days of acute COVID-19). 94 assigned to Palmitoylethanolam ide + Luteolin 700/70 mg a day and 36 assigned to standard of care. | Mean age 40.4 ± , male 46.2%, hypertension 2.3%, diabetes 0% | NR | High risk of bias Notes: Concealment of allocation and blinding probably inapropriate. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: RR 3.11 (95% CI 1.47 to 6.66); RD 35.5% (95% CI 7.8% to 83.3%); Low certainty ⊕⊕○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information |
|---|--|--|--------------------------|--|--|
| | | um diethylenetr fects of DTPA are unce | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Imam et al;(84); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 0 days of acute COVID-19). 33 assigned to DTPA 2% nasal spray, three times a day | Mean age 39.4, male 42.4%, hypertension 22.7%, diabetes 22.7%, asthma 7.6%, interval between COVID-19 and enrolment 97 days, | | High risk of bias Notes: Concealment of allocation and blinding probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms |

| | for one month and 33 assigned to standard of care. | | | | improvement: Very low certainty ⊕○○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information |
|--|---|--|--------------------------|--|---|
| | Uncortointe | Steroic | ds (nasal) | oorah is paadad | |
| Study; publication status | Patients and | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | КСТ | | |
| RC 4-7-2020 trial (85), Abdelalim et al.; Peer reviewed; 2022 | Patients with P-ACC (olfactory and/or gustatory dysfunction after acute COVID-19). 50 assigned to Mometasone 2 puffs (100 µg) once daily in each nostril for 3 weeks and 50 assigned to standard of care. | Mean age 29, male 46%, hypertension 14%, diabetes 16%, hospitalization during COVID-19 31% | Steroids 13% | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty |
| Hautefort et al (86); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 30 days of acute COVID-19). 62 assigned to budesonide 2 mg a day for 30 days and 61 assigned to standard of care. | Mean age 40.3, male 32.5%, hypertension %, diabetes 1.6% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: Very low certainty ⊕○○○ |

| | Steroids Steroids may not improve olfactory nor gustatory symptoms. Further research is needed. | | | | | | | |
|--|--|--|--------------------------|--|---|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | | F | RCT | | | | | |
| Vaira et al. (87); Peer reviewed; 2022 | Patients with P-ACC (olfactory and/or gustatory dysfunction after acute COVID-19). 9 assigned to prednisone 1 mg/kg a day and 9 assigned to standard of care. | Mean age 42.1, male 38.8% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Olfactory symptoms | | | |
| Schepens et al (88); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 28 days of acute COVID-19). 57 assigned to Prednisone 40 mg a day for 10 days and 56 assigned to standard of care. | Median age 49, male 36.5%, interval between COVID-19 and enrolment 56 days | Vaccinated 79.1% | Low risk of bias | improvement: RR 1.09 (95% CI 0.83 to 1.44); RD 3.3% (95% CI −6.2% to 16.1%); Low certainty ⊕⊕○○ Gustatory symptoms improvement: RR 1.01 (95% CI 0.67 to 1.53); RD 0.5% (95% CI −14.6% to 23.3%); Low certainty ⊕⊕○○ Adverse events: No information Severe adverse events: No information | | | |

Theophylline (nasal)
Uncertainty in potential benefits and harms. Further research is needed.

| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
|--|---|---|----------------------------------|--|---|
| | | F | RCT | | |
| SCENT2 trial (89); Gupta et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 90 days of acute COVID-19). 26 assigned to Theophylline (nasal) 400 mg twice a day for 6 weeks and 25 assigned to standard of care. | Mean age 44.7, male 29.4%, interval between COVID-19 and enrolment 387 days | NR | Low risk of bias | HRQL improvement: Very low certainty ⊕○○○ Overall symptom improvement: No information Olfactory symptoms improvement: Very low certainty ⊕○○○ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: Very low certainty ⊕○○○ |
| | Uncertainty | Vita vin potential benefits a | nmin A nd harms. Further reso | earch is needed. | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence |
| | | F | RCT | | |
| Chung et al (82); Peer reviewed; 2022 | Patients with post COVID-19 condition (Olfactory and/or gustatory dysfunction after 84 days of acute COVID-19). 9 | male %, hypertension 5.9%, diabetes 0%, cancer | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information |

| assigned to Vitamin A 25000 IU for 14 days and 8 assigned to standard of care. | | Olfactory symptoms improvement: Very low certainty |
|--|--|---|
| | | Gustatory symptoms improvement: No information |
| | | Adverse events: No information |
| | | Severe adverse events: No information |

Table 6. Description of included studies and interventions effects for PCC cardiovascular system symptoms

| | Hyperbaric oxygen Uncertainty in potential benefits and harms. Further research is needed. | | | | | | |
|---|---|---|---------------------------------|---|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| Leitman et al. (90) Peer reviewed; 2022 | Patients with post COVID-19 condition (cardiological symptoms after 60 days of acute COVID-19). 16 assigned to HBO five sessions per week for two months and 13 assigned to standard of care. | Mean age 47, male 41.7%, hypertension 10%, chronic lung disease 5%, CHD 1.7%, interval between COVID-19 and enrolment 158 days, hospitalization during COVID-19 13.3% | NR | High risk of bias Notes: Post-Hoc analysis based on a surrogate marker of cardiac function | HRQL improvement: No information Overall symptom improvement: No information Cardiac function improvement: Very low certainty ⊕○○○ Tachycardia improvement: No information Adverse events: No information Severe adverse events: No information | | |
| | Uncertainty | lvab in potential benefits a | radine nd harms. Further res | earch is needed. | | | |
| | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | i | RCT | | | | |

| Jadhav et al. (91); Peer reviewed; 2022 | Patients with P-ACC (cardiovascular symptoms after 0 to 14 days of acute COVID-19). 25 assigned to Ivabradine 5 to 10 mg and 25 assigned to standard of care. | Mean age 48.8 ± 7.66 | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Overall symptom improvement: No information Tachycardia improvement: Very low certainty ⊕○○○ Adverse events: No information Severe adverse events: No information |
|---|---|----------------------|----|--|--|
|---|---|----------------------|----|--|--|

Table 7. Description of included studies and interventions effects for PCC psychological distress

| Eye-movement desensitisation and reprocessing (EMDR) Uncertainty in potential benefits and harms. Further research is needed. | | | | | | | |
|--|--|--|-----------------------------|--|---|--|--|
| publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| trial (92); Bates et al; Peer reviewed; 2022 | Patients with post COVID-19 condition (psychological distress after 84 days of acute COVID-19). 11 assigned to EMDR up to eight 60 to 90 minutes sessions and 12 assigned to standard of care. | Mean age 58 ± 15.3, male 61.5%, CHD 15.4%, CKD 3.8%, cancer 3.8%, hospitalization during COVID-19 100% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: Very low certainty ⊕○○○ Depression improvement: Very low certainty ⊕○○○ Post-traumatic stress improvement: Very low certainty ⊕○○○ Psychological distress improvement: No information Adverse events: No information Severe adverse events: No information | | |
| Virtual reality infor | mational video may ir | /irtual reality in mprove depression, pos f the evidence was low. | st-traumatic stress, an | nd psychological distres | s. However, certainty | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |

| ICU-VR trial (93), Vlake et al.; Peer reviewed; 2022 | Patients with P-ACC (psychological distress after 90 days of acute COVID-19). 45 assigned to virtual reality 14-minute informational video session once and 44 assigned to standard of care. | Mean age 60, male 36% | NR | High risk of bias Notes: Non-blinded study. Concealment of allocation probably inappropriate. | HRQL improvement: No information Depression improvement: RR 1.21 (95% CI 0.95 to 1.54); RD 14% (95% CI − 3.7% to 36.7%); Low certainty ⊕⊕○○ |
|--|--|-----------------------|----|--|--|
| | | | | | Post-traumatic stress improvement: RR 1.18 (95% CI 0.98 to 1.42); RD 13.8% (95% CI −1.5% to 32.3%); Low certainty ⊕⊕○○ |
| | | | | | Psychological distress improvement: RR 1.49 (95% CI 1.08 to 2.05); RD 25.5% (95% CI 4.1% to 55.1%); Low certainty ⊕⊕○○ |
| | | | | | Adverse events: No information Severe adverse events: No information |

Table 8. Description of included studies and interventions effects for P-ACC-related thromboembolic risk

| Anticoagulants ma | Anticoagulants (prophylactic dose) Anticoagulants may not have an important effect on mortality, VTE, major bleeding and HRQL. However, certainty of the evidence was low because of imprecision. Further research is needed. | | | | | | | |
|------------------------------|--|---------------|--------------------------|---------------------------------------|---|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and | | | |

| | | | | | GRADE certainty of the evidence |
|---|---|---|-----|--|---|
| | | F | RCT | | |
| (94), Ramacciotti et al.; Peer reviewed; 2022 | Patients with P-ACC (at increased risk of VTE after acute COVID-19). 159 assigned to rivaroxaban 10 mg a day for 35 days and 159 assigned to standard of care. | Mean age 57.1, male 60%, interval between COVID-19 and enrolment 8 days, hospitalization during COVID-19 100% | | High risk of bias Notes: Non-blinded study which might have introduced bias to symptoms, VTE and adverse events outcomes. | Mortality: RR 0.85 (95% CI 0.38 to 0.88); RD -0.3% (95% CI -1.2% to 1.8%); Low certainty ⊕⊕⊖⊖ HRQL improvement: RR 0.99 (95% CI 0.78 to 1.24); Low certainty ⊕⊕⊖⊖ VTE (symptomatic): RR 1 (95% CI 0.29 to 3.45); RD 0% (95% CI -2.3% to |
| Peer reviewed; 2022 | Patients with post COVID-19 condition (thromboembolic events after 0 days of acute COVID-19). 607 assigned to Apixaban 5 mg a day for 30 days and 610 assigned to standard of care. | | NR | Low risk of bias | Major bleeding: RR 2.01 (95% CI 0.18 to 22.1); RD 0.1% (95% CI −0.1% to 1.2%); Low certainty ⊕⊕○○ Severe adverse events: No information |

 Table 9. Description of included studies and interventions effects for PIMS-TS

| Steroids Steroids may reduce time to discharge and respiratory support requirements. However, certainty of the evidence was low. Further research is needed. | | | | | | | |
|--|---|--|--------------------------|---|--|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| Swissped RECOVERY trial (96); Welzel et al; Peer reviewed; 2022 | Patients with PIMS-TS. 37 assigned to methylprednisolone 10 mg/kg a day for 3 days and 38 assigned to IVIG 2 gr/kg once | Mean age 9.1, male 75%, underlying chronic disease 11% | NR | High risk of bias Notes: Non-blinded study which might have introduced bias. | Mortality: No information Time to discharge reduction: RR 1.09 (95% CI 0.88 to 1.39); RD 4.5% (95% CI -6% to 19.5%); Low certainty ⊕⊕○○ Respiratory support: RR 0.49 (95% CI 0.27 to 0.89); RD -28.2% (95% CI -40.5% to -5.9%); Low certainty ⊕⊕○○ Inotropic requirements: Very low certainty ⊕○○○ LVEF <55%: Very low certainty ⊕○○○ Arrhythmia: Very low certainty ⊕○○○ VTE: Very low certainty ⊕○○○ VTE: Very low certainty ⊕○○○ Major bleeding: No information | | |

Table 10. Description of included studies and interventions effects for P-ACC prophylaxis

| Convale | Convalescent plasma Convalescent may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | | | |
|---|--|--|---|---|---|--|--|
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| CSSC-004 trial (97); Kelly et al; Preprint; 2022 | Patients with mild to moderate COVID-19. 445 assigned to convalescent plasma 250 ml once and 437 assigned to standard of care. | Median age 43, male 42.6%, hypertension 23.5%, diabetes 8.2%, obesity 16%, | Vaccinated 22% | High risk of bias Notes: Significant loss to follow-up | Mortality: No information HRQL improvement: No information P-ACC: RR 0.93 (95% CI 0.77 to 1.12); RD -2.4% (95% CI -7.9% to 4.2%); Low certainty ⊕⊕⊖⊖ Adverse events: No information Severe adverse events: No information | | |
| Fluvoxam | ine may not reduce P- | | examine tainty of the evidence | was low. Further resear | ch is needed. | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | |
| | | F | RCT | | | | |
| COVID-OUT trial (98); Bramante et al; Preprint; 2022 | Patients with mild to moderate COVID-19. 298 assigned to Fluvoxamine 50 mg once followed by 100 mg a day for 14 days and 297 assigned to | Median age 44.5, male 45.8%, hypertension 26.9%, diabetes 1.1%, obesity 47.2%, | Corticosteroids 1.5%, monoclonal antibodies 4.2%; Vaccinated 56.4% | High risk of bias Notes: Significant loss to follow-up | Mortality: No information HRQL improvement: No information P-ACC: RR 0.99 (95% CI 0.81 to | | |

| | standard of care. | | | | 1.21); RD -0.4% | | | | |
|---|--|---|--|---|---|--|--|--|--|
| Farahani et al. (99): Peer reviewed; 2022 | Patients with post COVID-19 condition (P-ACC prophylaxis after 0 days of acute COVID-19). 42 assigned to Fluvoxamine 100 mg a day for 10 days and 43 assigned to standard of care. | Mean age 38.5, male 51.2%, hypertension 15.3%, diabetes 6.5%, CHD 11.6%, | Vaccinated 100% | Low risk of bias | (95% CI -8.4% to 9.3%); Low certainty ⊕⊕⊖⊖ Adverse events: No information Severe adverse events: No information | | | | |
| lverme | ctin may not reduce P | | mectin nty of the evidence wa | s low. Further research | is needed. | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | | | | | |
| | | F | RCT | | | | | | |
| COVID-OUT - Ivermectin trial (98); Bramante et al; Preprint; 2022 | Patients with mild to moderate COVID-19. 377 assigned to Ivermectin 390-470 mcg/kg per day for 3 days and 361 assigned to standard of care. | Median age 45.5, male 44%, hypertension 26.7%, diabetes 2%, obesity 48.8% | Corticosteroids 1.5%, monoclonal antibodies 4.2%; Vaccinated 52.2% | High risk of bias Notes: Significant loss to follow-up | Mortality: No information HRQL improvement: No information P-ACC: RR 0.99 (95% CI 0.61 to 1.62); RD 0% (95% CI -1.7% to -2.6%); Low certainty ⊕⊕⊖⊖ Adverse events: No information Severe adverse events: No information | | | | |
| Lefluno | Leflunomide Leflunomide may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | | |

| | RCT | | | | | | | |
|--|--|---|--|---|---|--|--|--|
| DEFEAT-COVID trial (100); Kralj-Hans et al; Peer reviewed; 2022 | Patients with moderate to severe COVID-19). 81 assigned to leflunomide 100 mg/day for 3 days followed by 20 mg/day for 7 days and 91 assigned to standard of care. | Mean age 55.8, male 67%, diabetes 22%, chronic lung disease 12%, CHD 39%, cancer 3%, obesity 4%, interval between COVID-19 and enrolment 0 days, hospitalization during COVID-19 100% | Corticosteroids 95%, remdesivir %, hydroxychloroquine 47%, tocilizumab 2.3%, | High risk of bias Notes: Non-blinded study which might have introduced bias. | Mortality: No information HRQL improvement: No information P-ACC: RR 1.28 (95% CI 0.92 to 1.77); RD 11.2% (95% CI -3.2% to 31.1%); Low certainty ⊕⊕⊖⊖ Adverse events: No information Severe adverse events: Very low certainty ⊕⊖⊖⊖ | | | |
| Metfe | Metformin Metformin may reduce P-ACC. However, certainty of the evidence was low. Further research is needed. | | | | | | | |
| Study; publication status | Patients and interventions analyzed | Comorbidities | Additional interventions | Risk of bias and study limitations | Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence | | | |
| | , | F | RCT | | | | | |
| COVID-OUT - Metformin trial (98); Bramante et al; Preprint; 2022 | Patients with mild to moderate COVID-19. 564 assigned to metformin 1500 mg a day for 14 days and 561 assigned to standard of care. | Median age 45.5, male 45.3%, hypertension 22.8%, diabetes 1.6%, obesity 47.4% | Steroids 1.5%, remdesivir %, monoclonal antibodies 4.2%; Vaccinated 55.6% | High risk of bias Notes: Significant loss to follow-up | Mortality: No information HRQL improvement: No information P-ACC: RR 0.59 (95% CI 0.39 to 0.88); RD -4.3% (95% CI -6.4% to -1.2%); Low certainty ⊕⊕○○ Adverse events: No information Severe adverse events: No information | | | |

| Study; | Remdesivir Remdesivir may not reduce P-ACC. However, certainty of the evidence was low. Further research is needed. tudy; Patients and comorbidities Additional interventions limitations Risk of bias and study limitations effects vs standard | | | | | | | | | |
|--|---|---|--------------------------|------------------|--|--|--|--|--|--|
| | analyzed | | | | of care (SOC) and GRADE certainty of the evidence | | | | | |
| | , | F | RCT | | | | | | | |
| SOLIDARITY - Finland trial (101); Nevalainen et al; Peer reviewed; 2022 | COVID-19 condition (P-ACC | Mean age 58.4, male 60.2%, diabetes 22.1%, hospitalization during COVID-19 100% | Corticosteroids 71.8% | Low risk of bias | Mortality: Very low certainty ⊕○○○ HRQL improvement: No information P-ACC: RR 1.06 (95% CI 0.53 to 2.13); RD 0.8% (95% CI -6.9% to 16.4%); Low certainty ⊕⊕○○ Adverse events: No information Severe adverse events: No information | | | | | |

References

- 1. A clinical case definition of post COVID-19 condition by a Delphi consensus. 6 October, 2021. Available from: https://apps.who.int/iris/rest/bitstreams/1376291/retrieve
- A clinical case definition for post COVID-19 condition in children and adolescents by expert consensus (16 February 2023). Available from: https://www.who.int/publications/i/item/WHO-2019-nCoV-Post-COVID-19-condition-CA-Clinical-case-definition-2023-1
- 3. Zhang H, Zang C, Xu Z, Zhang Y, Xu J, Bian J, et al. Data-driven identification of post-acute SARS-CoV-2 infection subphenotypes. Nat Med [Internet]. 2022 Dec 1 [cited 2023 Jan 6]; Available from: https://www.nature.com/articles/s41591-022-02116-3
- 4. Nalbandian A, Sehgal K, Gupta A, Madhavan MV, McGroder C, Stevens JS, et al. Post-acute COVID-19 syndrome. Nat Med. 2021 Apr;27(4):601–15.
- 5. Malik P, Patel K, Pinto C, Jaiswal R, Tirupathi R, Pillai S, et al. Post-acute COVID-19 syndrome (PCS) and health-related quality of life (HRQoL)—A systematic review and meta-analysis. Journal of Medical Virology. 2022 Jan;94(1):253–62.
- 6. Han Q, Zheng B, Daines L, Sheikh A. Long-Term Sequelae of COVID-19: A Systematic Review and Meta-Analysis of One-Year Follow-Up Studies on Post-COVID Symptoms. Pathogens. 2022 Feb 19;11(2):269.
- 7. World Health Organization. Clinical management of COVID-19 Living guideline. Geneva: WHO; 15 September 2022. Available from: https://reliefweb.int/report/world/clinical-management-covid-19-living-guideline-15-september-2022
- 8. The L·OVE Platform. Methods for the special L·OVE of coronavirus infection [Internet] Santiago: Epistemonikos Foundation; 2020 [cited 7 December 2020]. Available from: https://app.iloveevidence.com/covid-19
- 9. Guyatt GH, Thorlund K, Oxman AD, Walter SD, Patrick D, Furukawa TA, et al. GRADE guidelines: 13. Preparing Summary of Findings tables and evidence profiles—continuous outcomes. Journal of Clinical Epidemiology. 2013 Feb;66(2):173–83.
- 10. Hultcrantz M, Rind D, Akl EA, et al. The GRADE Working Group clarifies the construct of certainty of evidence. J Clin Epidemiol 2017; 87: 4–13.
- 11. Zeng L, Brignardello-Petersen R, Hultcrantz M, et al. GRADE guidelines 32: GRADE offers guidance on choosing targets of GRADE certainty of evidence ratings. Journal of Clinical Epidemiology 2021; 137: 163–75.

- 12. Sterne JAC, Savović J, Page MJ, Elbers RG, Blencowe NS, Boutron I, et al. RoB 2: A revised tool for assessing risk of bias in randomised trials. BMJ 2019;366:14898. Available from: https://doi.org/10.1136/bmj.14898.
- 13. Guyatt GH, Oxman AD, Vist GE, et al. GRADE: an emerging consensus on rating quality of evidence and strength of recommendations. BMJ 2008; 336: 924–26.
- 14. Chudzik M, Burzyńska M, Kapusta J. Use of 1-MNA to Improve Exercise Tolerance and Fatigue in Patients after COVID-19. Nutrients. 2022 Jul 22;14(15):3004.
- 15. Kutashov VA. Actovegin use in patients with cognitive impairment after coronavirus infection (COVID-19). RJTAO. 2021 Apr 25;13(2):65–72.
- 16. Karosanidze I, Kiladze U, Kirtadze N, Giorgadze M, Amashukeli N, Parulava N, et al. Efficacy of Adaptogens in Patients with Long COVID-19: A Randomized, Quadruple-Blind, Placebo-Controlled Trial. Pharmaceuticals. 2022 Mar 11;15(3):345.
- 17. Toussaint LL, Bratty AJ. Amygdala and Insula Retraining (AIR) Significantly Reduces Fatigue and Increases Energy in People with Long COVID. Zahiruddin S, editor. Evidence-Based Complementary and Alternative Medicine. 2023 Jul 17;2023:1–8.
- 18. Tosato M, Calvani R, Picca A, Ciciarello F, Galluzzo V, Coelho-Júnior HJ, et al. Effects of l-Arginine Plus Vitamin C Supplementation on Physical Performance, Endothelial Function, and Persistent Fatigue in Adults with Long COVID: A Single-Blind Randomized Controlled Trial. Nutrients. 2022 Nov 23;14(23):4984.
- 19. Hawkins J, Hires C, Keenan L, Dunne E. Aromatherapy blend of thyme, orange, clove bud, and frankincense boosts energy levels in post-COVID-19 female patients: A randomized, double-blinded, placebo controlled clinical trial. Complementary Therapies in Medicine. 2022 Aug;67:102823.
- 20. Finnigan LEM, Cassar MP, Koziel MJ, Pradines J, Lamlum H, Azer K, et al. Efficacy and tolerability of an endogenous metabolic modulator (AXA1125) in fatigue-predominant long COVID: a single-centre, double-blind, randomised controlled phase 2a pilot study. eClinicalMedicine. 2023 Apr;101946.
- 21. Tanashyan M, Morozova S, Raskurazhev A, Kuznetsova P. A prospective randomized, double-blind placebo-controlled study to evaluate the effectiveness of neuroprotective therapy using functional brain MRI in patients with post-covid chronic fatigue syndrome. Biomedicine & Pharmacotherapy. 2023 Dec;168:115723.
- 22. Hansen KS, Mogensen TH, Agergaard J, Schiøttz-Christensen B, Østergaard L, Vibholm LK, et al. High-dose coenzyme Q10 therapy versus placebo in patients with post COVID-19 condition: A randomized, phase 2, crossover trial. The Lancet Regional Health Europe. 2022 Nov;100539.

- 23. Kuut TA, Müller F, Csorba I, Braamse A, Aldenkamp A, Appelman B, et al. Efficacy of cognitive behavioral therapy targeting severe fatigue following COVID-19: results of a randomized controlled trial. Clinical Infectious Diseases. 2023 May 8;ciad257.
- 24. Putilina MV, Teplova NV, Bairova KI, Petrikeeva AE, Shabalina NI. The result of prospective randomized study CITADEL the efficacy and safety of drug cytoflavin in postcovid rehabilitation. Z nevrol psikhiatr im SS Korsakova. 2021;121(10):45.
- 25. Brichetti V, Rubilar T, Tejada J, Montecino P, Crespi-Abril A, Barbieri E, et al. EuroQol-5D-3L in Long Covid patients After Supplementation with EchA Marine [®], a Sea Urchin Eggs Extract: a double-blinded, multicentrical study [Internet]. Pain Medicine; 2023 Jun [cited 2023 Jul 19]. Available from: http://medrxiv.org/lookup/doi/10.1101/2023.05.31.23290798
- 26. Rathi A, Jadhav SB, Shah N. A Randomized Controlled Trial of the Efficacy of Systemic Enzymes and Probiotics in the Resolution of Post-COVID Fatigue. Medicines. 2021 Aug 30;8(9):47.
- 27. Kharaeva Z, Shokarova A, Shomakhova Z, Ibragimova G, Trakhtman P, Trakhtman I, et al. Fermented Carica papaya and Morinda citrifolia as Perspective Food Supplements for the Treatment of Post-COVID Symptoms: Randomized Placebo-Controlled Clinical Laboratory Study. Nutrients. 2022 May 25;14(11):2203.
- 28. Botek M, Krejčí J, Valenta M, McKune A, Sládečková B, Konečný P, et al. Molecular Hydrogen Positively Affects Physical and Respiratory Function in Acute Post-COVID-19 Patients: A New Perspective in Rehabilitation. IJERPH. 2022 Feb 10;19(4):1992.
- 29. Deshpande S, Mundhe N, Deshpande V, Tamoli S, Mahadik S, Pawar V. Potential use of Immunodaat® (Botanical extract of Elderberry -Sambucus Nigra L.) in the management of Post Covid-19 symptoms- a comparative, multi-centric, randomized, clinical study [Internet]. Allergy and Immunology; 2022 Oct [cited 2023 Mar 23]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.10.04.22280680
- 30. Gaylis NB, Ritter A, Kelly SA, Pourhassan NZ, Tiwary M, Sacha JB, et al. Reduced Cell Surface Levels of C-C Chemokine Receptor 5 and Immunosuppression in Long Coronavirus Disease 2019 Syndrome. Clinical Infectious Diseases. 2022 Sep 30;75(7):1232–4.
- 31. Hausswirth C, Schmit C, Rougier Y, Coste A. Positive Impacts of a Four-Week Neuro-Meditation Program on Cognitive Function in Post-Acute Sequelae of COVID-19 Patients: A Randomized Controlled Trial. IJERPH. 2023 Jan 11;20(2):1361.
- 32. Nambi G, Abdelbasset WK, Alrawaili SM, Elsayed SH, Verma A, Vellaiyan A, et al. Comparative effectiveness study of low versus high-intensity aerobic training with

- resistance training in community-dwelling older men with post-COVID 19 sarcopenia: A randomized controlled trial. Clin Rehabil. 2022 Jan;36(1):59–68.
- 33. Rodriguez-Blanco C, Bernal-Utrera C, Anarte-Lazo E, Gonzalez-Gerez JJ, Saavedra-Hernandez M. A 14-Day Therapeutic Exercise Telerehabilitation Protocol of Physiotherapy Is Effective in Non-Hospitalized Post-COVID-19 Conditions: A Randomized Controlled Trial. JCM. 2023 Jan 18;12(3):776.
- 34. Jimeno-Almazán A, Buendía-Romero Á, Martínez-Cava A, Franco-López F, Sánchez-Alcaraz BJ, Courel-Ibáñez J, et al. Effects of a concurrent training, respiratory muscle exercise, and self-management recommendations on recovery from post-COVID-19 conditions: the RECOVE trial. Journal of Applied Physiology. 2023 Jan 1;134(1):95–104
- 35. Ibrahim AA, Hussein HM, Ali MS, Kanwal R, Acar T, Shaik DH, et al. A randomized controlled trial examining the impact of low vs. moderate-intensity aerobic training in post-discharge COVID-19 older subjects. European Review for Medical and Pharmacological Sciences. 2023 May;27(9):4280–91.
- 36. Rasmussen IE, Løk M, Durrer CG, Foged F, Schelde VG, Budde JB, et al. Impact of high-intensity interval training on cardiac structure and function after COVID-19: an investigator-blinded randomized controlled trial. Journal of Applied Physiology. 2023 Jun 30;japplphysiol.00078.2023.
- 37. Oliveira KCVD, Ferreira APDL, Silva DDA, Monteiro JDS, Silva KV, Lucena LCD, et al. The impact of post-COVID multicomponent rehabilitation. Fisioter mov. 2023;36:e36112.
- 38. Ogonowska-Slodownik A, Labecka MK, Maciejewska-Skrendo A, McNamara RJ, Kaczmarczyk K, Starczewski M, et al. Effect of Water-Based vs. Land-Based Exercise Intervention (postCOVIDkids) on Exercise Capacity, Fatigue, and Quality of Life in Children with Post COVID-19 Condition: A Randomized Controlled Trial. JCM. 2023 Sep 28;12(19):6244.
- 39. Espinoza-Bravo C, Arnal-Gómez A, Martínez-Arnau FM, Núñez-Cortés R, Hernández-Guillén D, Flor-Rufino C, et al. Effectiveness of Functional or Aerobic Exercise Combined with Breathing Techniques in Telerehabilitation for Patients with Long COVID: A Randomized Controlled Trial. Physical Therapy. 2023 Sep 2;pzad118.
- 40. Thomas R, Williams M, Aldous J, Yanagisawa Y, Kumar R, Forsyth R, et al. A Randomised, Double-Blind, Placebo-Controlled Trial Evaluating Concentrated Phytochemical-Rich Nutritional Capsule in Addition to a Probiotic Capsule on Clinical Outcomes among Individuals with COVID-19—The UK Phyto-V Study. COVID. 2022 Mar 22;2(4):433–49.

- 41. Marinoni B, Rimondi A, Bottaro F, Ciafardini C, Amoroso C, Muià M, et al. The Role of VSL#3® in the Treatment of Fatigue and Other Symptoms in Long Covid-19 Syndrome: a Randomized, Double-blind, Placebo-controlled Pilot Study (DELong#3) [Internet]. Psychiatry and Clinical Psychology; 2023 Jun [cited 2023 Jul 23]. Available from: http://medrxiv.org/lookup/doi/10.1101/2023.06.28.23291986
- 42. Oliver-Mas S, Delgado-Alonso C, Delgado-Álvarez A, Díez-Cirarda M, Cuevas C, Fernández-Romero L, et al. Transcranial Direct Current Stimulation (tDCS) for Post-COVID Fatigue: A Randomized, Double-Blind, Controlled Pilot Study. SSRN Journal [Internet]. 2022 [cited 2022 Dec 21]; Available from: https://www.ssrn.com/abstract=4216601
- 43. Santana K, França E, Sato J, Silva A, Queiroz M, de Farias J, et al. Non-invasive brain stimulation for fatigue in post-acute sequelae of SARS-CoV-2 (PASC). Brain Stimulation. 2023 Jan;16(1):100–7.
- 44. King M, Byrne A, Denehy L, Graham P, Douglas B, de Toni P, et al. Feasibility of a Group-Based Telerehabilitation Intervention for Long COVID Management. [Internet]. In Review; 2022 Mar [cited 2022 Dec 27]. Available from: https://www.researchsquare.com/article/rs-1452186/v1
- 45. Simpson AJ, Green A, Nettleton M, Hyde L, Shepherdson J, Killingback C, et al. Group-based pulmonary telerehabilitation is feasible, safe, beneficial and well-received in patients who have been hospitalised with COVID-19. ERJ Open Res. 2023 Mar;9(2):00373–2022.
- 46. Longobardi I, Goessler K, de Oliveira Júnior GN, Prado DML do, Santos JVP, Meletti MM, et al. Effects of a 16-week home-based exercise training programme on health-related quality of life, functional capacity, and persistent symptoms in survivors of severe/critical COVID-19: a randomised controlled trial. Br J Sports Med. 2023 May 10;bjsports-2022-106681.
- 47. Samper-Pardo M, León-Herrera S, Oliván-Blázquez B, Méndez-López F, Domínguez-García M, Sánchez-Recio R. Effectiveness of a telerehabilitation intervention using ReCOVery APP of long COVID patients: a randomized, 3-month follow-up clinical trial. Sci Rep. 2023 May 16;13(1):7943.
- 48. Da Silva MMC, Viana DR, Colucci MG, Gonzaga LA, Arcuri JF, Frade MCM, et al. Effects of a cardiopulmonary telerehabilitation using functional exercises in individuals after COVID-19 hospital discharge: A randomized controlled trial. J Telemed Telecare. 2023 Aug 9;1357633X231188394.

- 49. Tanhan A, Ozer AY, Timurtas E, Batirel A, Polat MG. Is asynchronous telerehabilitation equal to synchronous telerehabilitation in COVID-19 survivors with classes 4–6? J Telemed Telecare. 2023 Aug 7;1357633X231189761.
- 50. Kerget B, Çil G, Araz Ö, Alper F, Akgün M. Comparison of two antifibrotic treatments for lung fibrosis in post-COVID-19 syndrome: A randomized, prospective study. Medicina Clínica. 2023 Mar;S0025775323000817.
- 51. Nagy EN, Elimy DA, Ali AY, Ezzelregal HG, Elsayed MM. Influence of Manual Diaphragm Release Technique Combined with Inspiratory Muscle Training on Selected Persistent Symptoms in Men with Post-Covid-19 Syndrome: A Randomized Controlled Trial. JRM. 2022 Oct 20;54:jrm00330.
- 52. Dal Negro RW, Turco P, Povero M. Nebivolol: an effective option against long-lasting dyspnoea following COVID-19 pneumonia a pivotal double-blind, cross-over controlled study. Multidis Res Med [Internet]. 2022 Dec 23 [cited 2023 Mar 23];17. Available from: https://mrmjournal.org/mrm/article/view/886
- 53. Philip KEJ, Owles H, McVey S, Pagnuco T, Bruce K, Brunjes H, et al. An online breathing and wellbeing programme (ENO Breathe) for people with persistent symptoms following COVID-19: a parallel-group, single-blind, randomised controlled trial. The Lancet Respiratory Medicine. 2022 Sep;10(9):851–62.
- 54. McNarry MA, Berg RMG, Shelley J, Hudson J, Saynor ZL, Duckers J, et al. Inspiratory muscle training enhances recovery post-COVID-19: a randomised controlled trial. Eur Respir J. 2022 Oct;60(4):2103101.
- 55. Srinivasan V., Kandakurti P.K., Alagesan J., Suganthirababu P., Jenifer Augustina S., Anitha A., et al. Efficacy of pursed lip breathing with bhastrika pranayama vs incentive spirometry in rehabilitating post Covid 19 follow up-a randomized control study. Turkish Journal of Physiotherapy and Rehabilitation. 2021;32(3):402–7.
- 56. Palau P, Domínguez E, Gonzalez C, Bondía E, Albiach C, Sastre C, et al. Effect of a home-based inspiratory muscle training programme on functional capacity in postdischarged patients with long COVID: the InsCOVID trial. BMJ Open Resp Res. 2022 Dec:9(1):e001439.
- 57. Romanet C, Wormser J, Fels A, Lucas P, Prudat C, Sacco E, et al. Effectiveness of endurance training rehabilitation after hospitalisation in intensive care for COVID-19-related acute respiratory distress syndrome on dyspnoea (RECOVER): a randomised controlled, open-label multicentre trial [Internet]. Respiratory Medicine; 2022 Aug [cited 2022 Dec 30]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2022.08.29.22279327

- 58. Del Corral T, Fabero-Garrido R, Plaza-Manzano G, Fernández-de-las-Peñas C, Navarro-Santana M, López-de-Uralde-Villanueva I. Home-based respiratory muscle training on quality of life and exercise tolerance in long-term post-COVID-19: Randomized controlled trial. Annals of Physical and Rehabilitation Medicine. 2023 Feb;66(1):101709.
- 59. Nambi G, Alghadier M, Vellaiyan A, Ebrahim EE, Aldhafian OR, Mohamed SHP, et al. Role of Tele-Physical Therapy Training on Glycemic Control, Pulmonary Function, Physical Fitness, and Health-Related Quality of Life in Patients with Type 2 Diabetes Mellitus (T2DM) Following COVID-19 Infection—A Randomized Controlled Trial. Healthcare. 2023 Jun 17;11(12):1791.
- 60. Li J, Xia W, Zhan C, Liu S, Yin Z, Wang J, et al. A telerehabilitation programme in post-discharge COVID-19 patients (TERECO): a randomised controlled trial. Thorax. 2022 Jul;77(7):697–706.
- 61. Mirenayat MS, Moradkhani A, Abedi M, Abedini A, Zahiri R, Karimzadeh S, et al. Role of Inspiratory Muscle Training on Pulmonary Rehabilitation in Patients with COVID-19: A Pilot Study. Tanaffos. 2022 Apr;21(4):466–71.
- 62. Senén AB, Fernández AG, López JG, Rodríguez JB, Brejano MG, Guillén PC, et al. Functional rehabilitation based on therapeutic exercise training in patients with postacute COVID syndrome (RECOVER). Revista Española de Cardiología (English Edition). 2023 Oct;S188558572300261X.
- 63. Vallier JM, Simon C, Bronstein A, Dumont M, Jobic A, Paleiron N, et al. Randomized controlled trial of home-based vs. hospital-based pulmonary rehabilitation in post COVID-19 patients. Eur J Phys Rehabil Med [Internet]. 2023 Jan [cited 2023 Feb 9]; Available from: https://www.minervamedica.it/index2.php?show=R33Y9999N00A23012602
- 64. Rutkowski S, Bogacz K, Rutkowska A, Szczegielniak J, Casaburi R. Inpatient post-COVID-19 rehabilitation program featuring virtual reality—Preliminary results of randomized controlled trial. Front Public Health. 2023 Feb 6;11:1121554.
- 65. Stavrou VT, Vavougios GD, Kalogiannis P, Tachoulas K, Touloudi E, Astara K, et al. Breathlessness and exercise with virtual reality system in long-post-coronavirus disease 2019 patients. Front Public Health. 2023 Feb 23;11:1115393.
- 66. Kusumawardani RI, Tinduh D, Poerwandari D, Marhana IA, Melaniani S. The effectiveness of incentive spirometry exercise on pulmonary function in COVID-19 survivors: a randomized controlled trial study. Bali Medical Journal. 2023;12(1):539–44.
- 67. Dhooria S, Chaudhary S, Sehgal IS, Agarwal R, Arora S, Garg M, et al. High-dose versus low-dose prednisolone in symptomatic patients with post-COVID-19 diffuse

- parenchymal lung abnormalities: an open-label, randomised trial (the COLDSTER trial). Eur Respir J. 2022 Feb;59(2):2102930.
- 68. Bazdyrev E, Panova M, Brachs M, Smolyarchuk E, Tsygankova D, Gofman L, et al. Efficacy and safety of Treamid in the rehabilitation of patients after COVID-19 pneumonia: a phase 2, randomized, double-blind, placebo-controlled trial [Internet]. In Review; 2022 Jul [cited 2022 Dec 27]. Available from: https://www.researchsquare.com/article/rs-1845321/v1
- 69. Ansari S, Sanjari Moghaddam H, Basti FA, Salehi M, Akhondzadeh S. Efficacy and safety of celecoxib monotherapy for treatment of moderate depressive symptoms following COVID-19 infection: A randomized, double-blind, placebo-controlled trial. Journal of Psychosomatic Research. 2023 Nov;174:111471.
- 70. Zulbaran-Rojas A, Lee M, Bara RO, Flores-Camargo A, Spitz G, Finco MG, et al. Electrical stimulation to regain LOWER EXTREMITY muscle perfusion and endurance in patients with POST-ACUTE sequelae of SARS CoV -2: A randomized controlled trial. Physiological Reports [Internet]. 2023 Mar [cited 2023 May 5];11(5). Available from: https://onlinelibrary.wiley.com/doi/10.14814/phy2.15636
- 71. Momtazmanesh S, Ansari S, Izadi Z, Shobeiri P, Vatankhah V, Seifi A, et al. Effect of famotidine on cognitive and behavioral dysfunctions induced in post-COVID-19 infection: A randomized, double-blind, and placebo-controlled study. Journal of Psychosomatic Research. 2023 Sep;172:111389.
- 72. Zilberman-Itskovich S, Catalogna M, Sasson E, Elman-Shina K, Hadanny A, Lang E, et al. Hyperbaric oxygen therapy improves neurocognitive functions and symptoms of post-COVID condition: randomized controlled trial. Sci Rep. 2022 Jul 12;12(1):11252.
- 73. Versace V, Ortelli P, Dezi S, Ferrazzoli D, Alibardi A, Bonini I, et al. Co-ultramicronized palmitoylethanolamide/luteolin normalizes GABAB-ergic activity and cortical plasticity in long COVID-19 syndrome. Clinical Neurophysiology. 2023 Jan;145:81–8.
- 74. Kalayeh MB, Gaeini AA, Kordi MR, Kalayeh MB. Effects of Eight-Week Resistance Training on the Quality of Life and Sleep Quality of Untrained Men with a History of COVID-19 [Internet]. In Review; 2023 Apr [cited 2023 May 5]. Available from: https://www.researchsquare.com/article/rs-2818393/v1
- 75. Badran BW, Huffman SM, Dancy M, Austelle CW, Bikson M, Kautz SA, et al. A pilot randomized controlled trial of supervised, at-home, self-administered transcutaneous auricular vagus nerve stimulation (taVNS) to manage long COVID symptoms [Internet]. In Review; 2022 Jun [cited 2022 Dec 30]. Available from: https://www.researchsquare.com/article/rs-1716096/v1

- 76. Shabaan AA, Kassem I, Mahrous AI, Aboulmagd I, Badrah M, Attalla M, et al. Diode laser in management of loss of taste sensation in patients with post-COVID syndrome: a randomized clinical trial. BMC Oral Health. 2023 May 6;23(1):263.
- 77. Abdelazim MH, Mandour Z, Abdelazim AH, Ismaiel WF, Gamal M, Abourehab MAS, et al. Intra Nasal Use of Ethylene Diamine Tetra Acetic Acid for Improving Olfactory Dysfunction Post COVID-19. Am J Rhinol Allergy. 2023 Nov;37(6):630–7.
- 78. Mahadev A, Hentati F, Miller B, Bao J, Perrin A, Kallogjeri D, et al. Efficacy of Gabapentin For Post–COVID-19 Olfactory Dysfunction: The GRACE Randomized Clinical Trial. JAMA Otolaryngol Head Neck Surg [Internet]. 2023 Sep 21 [cited 2023 Oct 5]; Available from: https://jamanetwork.com/journals/jamaotolaryngology/fullarticle/2809346
- 79. Di Stadio A, Cantone E, De Luca P, Di Nola C, Massimilla EA, Motta G, et al. Parosmia COVID-19 Related Treated by a Combination of Olfactory Training and Ultramicronized PEA-LUT: A Prospective Randomized Controlled Trial. Biomedicines. 2023 Apr 6;11(4):1109.
- 80. Pires ÍAT, Steffens ST, Mocelin AG, Shibukawa DE, Leahy L, Saito FL, et al. Intensive Olfactory Training in Post-COVID Patients: A Randomized Multicenter Clinical Trial [Internet]. 2022 Jan [cited 2022 Dec 29]. Available from: https://preprints.scielo.org/index.php/scielo/preprint/view/3301/version/3492
- 81. Lechner M, Liu J, Counsell N, Gillespie D, Chandrasekharan D, Ta NH, et al. The COVANOS trial insight into post-COVID olfactory dysfunction and the role of smell training. Rhin. 2022 Jun 1;60(3):188–99.
- 82. Chung TWH, Zhang H, Wong FKC, Sridhar S, Lee TMC, Leung GKK, et al. A Pilot Study of Short-Course Oral Vitamin A and Aerosolised Diffuser Olfactory Training for the Treatment of Smell Loss in Long COVID. Brain Sciences. 2023 Jun 30;13(7):1014.
- 83. Lerner DK, Garvey KL, Arrighi-Allisan A, Kominsky E, Filimonov A, Al-Awady A, et al. Omega-3 Fatty Acid Supplementation for the Treatment of Persistent COVID-Related Olfactory Dysfunction. Am J Rhinol Allergy. 2023 Jun 1;194589242311747.
- 84. Imam MS, Abdelazim MH, Abdelazim AH, Ismaiel WF, Gamal M, Abourehab MAS, et al. Efficacy of pentasodium diethylenetriamine pentaacetate in ameliorating anosmia post COVID-19. American Journal of Otolaryngology. 2023 Jul;44(4):103871.
- 85. Abdelalim AA, Mohamady AA, Elsayed RA, Elawady MA, Ghallab AF. Corticosteroid nasal spray for recovery of smell sensation in COVID-19 patients: A randomized controlled trial. American Journal of Otolaryngology. 2021 Mar;42(2):102884.

- 86. Hautefort C, Corré A, Poillon G, Jourdaine C, Housset J, Eliezer M, et al. Local budesonide therapy in the management of persistent hyposmia in suspected non-severe COVID-19 patients: Results of a randomized controlled trial. International Journal of Infectious Diseases. 2023 Nov:136:70–6.
- 87. Vaira LA, Hopkins C, Petrocelli M, Lechien JR, Cutrupi S, Salzano G, et al. Efficacy of corticosteroid therapy in the treatment of long- lasting olfactory disorders in COVID-19 patients. Rhin. 2020 Dec 1;0(0):0–0.
- 88. Schepens EJA, Blijleven EE, Boek WM, Boesveldt S, Stokroos RJ, Stegeman I, et al. Prednisolone does not improve olfactory function after COVID-19: a randomized, double-blind, placebo-controlled trial. BMC Med. 2022 Nov 16;20(1):445.
- 89. Gupta S, Lee JJ, Perrin A, Khan A, Smith HJ, Farrell N, et al. Efficacy and Safety of Saline Nasal Irrigation Plus Theophylline for Treatment of COVID-19–Related Olfactory Dysfunction: The SCENT2 Phase 2 Randomized Clinical Trial. JAMA Otolaryngol Head Neck Surg. 2022 Sep 1;148(9):830.
- 90. Leitman M, Fuchs S, Tyomkin V, Hadanny A, Zilberman-Itskovich S, Efrati S. The effect of hyperbaric oxygen therapy on myocardial function in post-COVID-19 syndrome patients: a randomized controlled trial. Sci Rep. 2023 Jun 10;13(1):9473.
- 91. Kartik Pandurang J, Pankaj V J. Ivabradine versus carvedilol in the management of palpitation with sinus tachycardia among recovered COVID-19 patients. J Cardiol Cardiovasc Me. 2020 Dec 23;5(3):176–80.
- 92. Bates A, Golding H, Rushbrook S, Shapiro E, Pattison N, Baldwin DS, et al. A randomised pilot feasibility study of eye movement desensitisation and reprocessing recent traumatic episode protocol, to improve psychological recovery following intensive care admission for COVID-19. Journal of the Intensive Care Society. 2023 Aug;24(3):309–19.
- 93. Vlake JH, van Bommel J, Wils EJ, Bienvenu J, Hellemons ME, Korevaar TI, et al. Intensive Care Unit–Specific Virtual Reality for Critically Ill Patients With COVID-19: Multicenter Randomized Controlled Trial. J Med Internet Res. 2022 Jan 31;24(1):e32368.
- 94. Ramacciotti E, Barile Agati L, Calderaro D, Aguiar VCR, Spyropoulos AC, de Oliveira CCC, et al. Rivaroxaban versus no anticoagulation for post-discharge thromboprophylaxis after hospitalisation for COVID-19 (MICHELLE): an open-label, multicentre, randomised, controlled trial. The Lancet. 2022 Jan;399(10319):50–9.

- 95. Wang TY, Wahed AS, Morris A, Kreuziger LB, Quigley JG, Lamas GA, et al. Effect of Thromboprophylaxis on Clinical Outcomes After COVID-19 Hospitalization. Ann Intern Med. 2023 Apr;176(4):515–23.
- 96. Welzel T, Atkinson A, Schöbi N, Andre MC, Bailey DGN, Blanchard-Rohner G, et al. Methylprednisolone versus intravenous immunoglobulins in children with paediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2 (PIMS-TS): an open-label, multicentre, randomised trial. The Lancet Child & Adolescent Health. 2023 Feb;S2352464223000202.
- 97. Gebo KA, Heath SL, Fukuta Y, Zhu X, Baksh S, Abraham AG, et al. Early Treatment, Inflammation and Post-COVID Conditions [Internet]. Infectious Diseases (except HIV/AIDS); 2023 Feb [cited 2023 Mar 17]. Available from: http://medrxiv.org/lookup/doi/10.1101/2023.02.13.23285855
- 98. Bramante C, Buse JB, Liebovitz D, Nicklas J, Puskarich M, Cohen KR, et al. Outpatient Treatment of COVID-19 and the Development of Long COVID Over 10 Months: A Multi-Center, Quadruple-Blind, Parallel Group Randomized Phase 3 Trial [Internet]. SSRN; 2023 [cited 2023 Mar 14]. Available from: https://www.ssrn.com/abstract=4375620
- 99. Farahani RH, Ajam A, Naeini AR. Effect of fluvoxamine on preventing neuropsychiatric symptoms of post COVID syndrome in mild to moderate patients, a randomized placebocontrolled double-blind clinical trial. BMC Infect Dis. 2023 Mar 31;23(1):197.
- 100. Kralj-Hans I, Li K, Wesek A, Lamorgese A, Omar F, Ranasinghe K, et al. Leflunomide treatment for patients hospitalised with COVID-19: DEFEAT-COVID randomised controlled trial. BMJ Open. 2023 Apr;13(4):e068179.
- 101. Nevalainen OPO, Horstia S, Laakkonen S, Rutanen J, Mustonen JMJ, Kalliala IEJ, et al. Effect of remdesivir post hospitalization for COVID-19 infection from the randomized SOLIDARITY Finland trial. Nat Commun. 2022 Oct 18;13(1):6152.

Annex 1. Summary of findings tables

Summary of findings Table A1.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Actovegin

| Outcome Timeframe | Study results and | Absolute effect estimates | | Certainty of the | Diain language augument |
|---|---|---|---------------------|--|-------------------------|
| | measurements | SOC | Actovegin | Evidence (Quality of evidence) | Plain language summary |
| Fatigue improvement (CI 95% 1.59 - 2.* Based on data from participants in 1 st | Relative risk: 1.54 (Cl 95% 1.59 - 2.14) Based on data from 444 | 471 per 1000 | 725 per 1000 | Low Due to very serious risk of | Actovegin may improve |
| | participants in 1 study Follow up 90 days | Difference: 254 more per 1000 (CI 95% 278 more - 537 more) | | bias ¹ | fatigue |

Risk of Bias: very serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate sequence generation/ generation of comparable groups, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: no serious. 95% CI include important benefits and harms.

Summary of findings Table A2.

Population: Patients with P-ACC-related asthenia or fatigue Intervention: ADAPT-232

| Outcome Timeframe | Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language summary |
|-----------------------|--|--|------------------------|--------------------------------|---|
| | measurements | SOC | ADAPT-232 | (Quality of evidence) | riain language summary |
| Fatigue improvement B | Relative risk 1.02 (95% CI 0.84 to 1.24) Based on data from 99 | 800 per 1000 | 816 per 1000 | Low Due to very serious | Adapt-232 may have little or no difference on fatigue |
| | participants in 1 study Follow-up 21 days | Difference: 16 more per 1000 (95% CI 128 fewer to 192 more) | | imprecision ¹ | improvement |

^{1.} Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A3.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Cytoflavin

| Outcome Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language | |
|---------------------------|--|--|---------------------------|---|---------------------|
| Timeframe | measurements SOC Cytoflavin | Cytoflavin | (Quality of evidence) | summary | |
| Fatigue | Relative risk 1.02 (95% CI 0.98 to 1.06) improvementa Based on data from 200 participants in 1 study Follow-up 25 days | 979 per 1000 | 999 per 1000 | Low Due to serious risk of bias, | Cytoflavin may have |
| inprovement | | Difference: 20 more per 1000 (95% CI 20 fewer to 21 more) | | Due to serious imprecision ^b | fatigue improvement |

a. Decrease in 12 units of the MFI score.

Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: serious. Low number of patients.

Summary of findings Table A4.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Enzymes + probiotics Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language |
|--|---|---------------------------|--------------------------------------|---|---|
| | measurements | SOC | Enzymes + probiotics | (Quality of evidence) | summary |
| Fatigue improvement (95% CI 3.71 to Based on data from participants in 1 | Relative risk 6.07 (95% CI 3.71 to 9.71) Based on data from 200 | 150 per 1000 | 911 per 1000 | Low Due to serious risk of bias, | Enzymes + probiotics may increase fatique |
| | participants in 1 study Follow-up 25 days | | 1 more per 1000 nore to 850 more) | Due to serious imprecision ^a | improvement |

a. Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: serious. Low number of patients.

Summary of findings Table A5.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Phytochemicals

| Outcome | Study results and | Absolute effect estimates | | Certainty of the | Plain language | |
|---------------------|---|---|------------------------|--|-------------------------------------|--|
| Timeframe | measurements | soc | Phytochemicals | Evidence (Quality of evidence) | summary | |
| HRQL improvement | Relative risk: 1.33 (Cl 95% 1.03 - 1.71) Based on data from 147 | 543 per 1000 | 722 per 1000 | Low Due to serious risk of | Phytochemicals may increase HRQL | |
| | participants in 1 study Follow up 30 days | Difference: 179 more per 1000 (CI 95% 16 more - 386 more) | | bias, Due to serious imprecision ¹ | improvement | |
| Fatigue improvement | Relative risk: 1.24 (CI 95% 0.95 - 1.62) Based on data from 147 | 539 per 1000 | 668 per 1000 | Low Due to serious risk of | Phytochemicals may increase fatigue | |
| | participants in 1 study Follow up 30 days | Difference: 129 more per 1000 (CI 95% 27 fewer - 334 more) | | bias, Due to serious imprecision ² | improvement | |

- Risk of Bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: serious. Low number of patients.
- 2. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.

Summary of findings Table A6.

Population: Patients with P-ACC-related asthenia or fatigue Intervention: Transcranial direct current stimulation (tDCS)

| Outcome | Charles manufactured | Absolute et | ffect estimates | Certainty of the | | |
|----------------------|--|--|--|--|---|--|
| Outcome Timeframe | Study results and measurements | SOC | Transcranial direct current stimulation (tDCS) | Evidence (Quality of evidence) | Plain language summary | |
| Fatigue improvement | Relative risk: 1.36 (CI 95% 0.76 - 2.45) Based on data from 117 | 468 per 1000 | 636 per 1000 | Low Due to very serious | Transcranial direct current stimulation (tdcs) may have | |
| | participants in 2 studies Follow up 32.5 days | Difference: 168 more per 1000 (CI 95% 112 fewer - 672 more) | | imprecision ¹ | little or no difference on fatigue improvement | |
| HRQL improvement | Relative risk: 1.37 (CI 95% 1.09 - 1.71) Based on data from 70 participants in 1 study Follow up 35 days | 705 per 1000 | 966 per 1000 | Low Due to very serious | Transcranial direct current stimulation (tdcs) may | |
| | | Difference: 261 more per 1000 (CI 95% 63 more - 295 more) | | imprecision ² | improve HRQL | |
| Adverse events | Relative risk: 0.83 (Cl 95% 0.26 - 2.73) Based on data from 47 participants in 1 study Follow up 30 days | 208 per 1000 | 173 per 1000 | Low Due to very serious | Transcranial direct current stimulation (tdcs) may have | |
| | | Difference: 35 fewer per 1000 (Cl 95% 154 fewer - 360 more) | | Due to very serious imprecision ³ | little or no difference on adverse events | |

- 1. Imprecision: very serious. 95% CI include important benefits and harms;
- 2. **Imprecision: very serious.** 95% CI include important benefits and harms;
- 3. **Imprecision: very serious.** 95% CI include important benefits and harms;

Summary of findings Table A7.

Population: Patients with P-ACC-related dyspnea Intervention: ADAPT-232

| Outcome | Study results and | Absolute effect estimates | | Certainty of the | Plain language |
|------------------------|---|---------------------------|------------------------------------|-----------------------------------|---|
| Timeframe | | soc | ADAPT-232 | evidence (Quality of evidence) | summary |
| Dyspnea improvement | Relative risk 1.0 (95% CI 0.94 to 1.06) Based on data from 99 participants in 1 study Follow-up 21 days | 980 per 1000 | 980 per 1000 | Low Due to very serious | ADAPT-232 may have little or no difference on |
| pioromon | | | fewer per 1000 ewer to 20 more) | imprecision ^a | dyspnea improvement |

a. Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A8.

Population: Patients with P-ACC-related dyspnea

Intervention: Endurance training Comparator: Standard of care (SOC)

| Outcome | Study results and measurements | Absolute eff | ect estimates | Certainty of the | Plain language summary |
|--|--|--|----------------------------|---|------------------------|
| Timeframe | | soc | Endurance training | evidence (Quality of evidence) | |
| Relative risk 1.48 (95% CI 0.92 to 2.37) Based on data from 60 | 441 per 1000 | 980 per 1000 | Low Due to serious risk of | Endurance training may increase HRQL | |
| | participants in 1 study Follow-up 21 days | Difference: 0 fewer per 1000 (95% CI 59 fewer to 20 more) | | bias, Due to serious imprecision ^b | improvement |
| Dyspnea Relative risk 2.03 (95% CI 0.98 to 4.21) Based on data from 60 participants in 1 study Follow-up 21 days | 236 per 1000 | 980 per 1000 | Low Due to serious risk of | Endurance training may increase dyspnea | |
| | | Difference: 0 fewer per 1000 (95% CI 59 fewer to 20 more) | | bias, Due to serious imprecision ^d | improvement |

a. Increment of 7 units in the SF-12 scale.

b. **Risk of bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. **Imprecision: serious.** Low number of patients.

c. Increment of 7 units in the SF-12 scale.

d. Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: serious. Low number of patients.

Summary of findings Table A9.

Population: Patients with P-ACC-related dyspnea

Intervention: High dose steroids (i.e., prednisone 40 mg a day) Comparator: Standard dose steroids (i.e., prednisone 10 mg a day)

| Outcome | Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language |
|------------------------|---|--|---------------------|--|---|
| Timeframe | measurements | Standard dose steroids | High dose steroids | (Quality of evidence) | summary |
| Dyspnea improvement | Relative risk 1.0 (95% Cl 0.87 to 1.15) Based on data from 130 participants in 1 study | 862 per 1000 | 862 per 1000 | Low Due to serious risk of bias, Due to serious | have little or no difference |
| | Follow-up 42 days | Difference: 0 fe (95% CI 112 few | | imprecision ^a | on dyspnea improvement |
| Radiological response | Relative risk 1.33 (95% CI 0.69 to 2.59) Based on data from 60 | 185 per 1000 | 246 per 1000 | Very low Due to serious risk of bias, | We are uncertain whether high dose steroids |
| | participants in 1 study Follow-up 21 days | Difference: 61 r (95% CI 57 fewe | | Due to very serious imprecision ^b | increases or decreases radiological response |
| Adverse events | Relative risk 0.92 (95% CI 0.75 to 1.13) Based on data from 60 | 769 per 1000 | 707 per 1000 | Low Due to serious risk of bias, | High dose steroids may have little or no difference |
| | participants in 1 study Follow-up 21 days | Difference: 62 f e (95% CI 192 few | | Due to serious imprecision ^c | on adverse events |
| Severe adverse | (| 15 per 1000 | 45 per 1000 | Very low Due to serious risk of bias, | We are uncertain whether high dose steroids |
| events | Based on data from 60 participants in 1 study Follow-up 21 days | Difference: 30 r (95% CI 10 fewe | | Due to very serious imprecision ^d | increases or decreases severe adverse events |

- a. Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: serious. Low number of patients.
- b. **Risk of bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. **Imprecision: very serious.** 95% CI includes important benefits and harms.
- c. Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: serious. Low number of patients.
- d. Risk of bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias. Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A10.

Population: Patients with P-ACC-related dyspnea Intervention: Respiratory training/rehabilitation

| Outcome | Study results and | Absolute ef | Absolute effect estimates Certainty of the | | | |
|--|--|---|--|--|--|--|
| Timeframe | measurements | SOC | Respiratory training | Evidence (Quality of evidence) | Summary | |
| HRQL improvement | Relative risk: 1.41 (CI 95% 1.18 - 1.67) Based on data from 644 participants in 8 studies | 493 per 1000 Difference: 20 | 695 per 1000 2 more per 1000 | Moderate Due to serious risk of bias ¹ | Respiratory training probably increases HRQL improvement | |
| | Follow up 90 days | (CI 95% 89 m | ore - 330 more) | | | |
| Functional capacity improvement | Relative risk: 1.41 (CI 95% 1.05 - 1.9) Based on data from 381 | 467 per 1000 | 658 per 1000 | Moderate Due to serious risk of | Respiratory probably increases functional | |
| | participants in 5 studies Follow up 138 days | Difference: 191 more per 1000 (CI 95% 23 more - 420 more) | | bias ² | capacity improvement | |
| Dyspnea improvement | Relative risk: 2.11 (CI 95% 1.17 - 3.8) Based on data from 579 | 309 per 1000 | 652 per 1000 | Moderate Due to serious risk of | Respiratory training probably increases | |
| prevenien | participants in 9 studies Follow up 84 days | | 3 more per 1000 ore - 865 more) | bias ³ | dyspnea improvement | |
| Pulmonary function improvement Base partic | Relative risk: 12.8 (CI 95% 1.65 - 99.5) Based on data from 26 | 66 per 1000 | 845 per 1000 | Very low Due to serious risk of bias, | We are uncertain whether respiratory training increases or decreases | |
| | participants in 1 studies Follow up 90 days | Difference: 779 more per 1000 (CI 95% 43 more - 6501 more) | | Due to very serious imprecision ⁴ | pulmonary function improvement | |

- 1. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias;
- 2. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;
- 3. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias;
- 4. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** 95% CI include important benefits and harms;

Summary of findings Table A11.

Population: Patients with P-ACC-related dyspnea

Intervention: Treamid

| Outcome | Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language | |
|---------------------------------|---|--|------------------------------------|--|---|--|
| Timeframe | measurements | SOC | Treamid | (Quality of evidence) | summary | |
| Functional capacity improvement | Relative risk 1.1 (95% CI 0.64 to 1.9) Based on data from 59 | 445 per 1000 | 490 per 1000 | Low Due to very serious | Treamid may have little or no difference on | |
| improvement | participants in 1 study Follow-up 28 days | | more per 1000 wer to 401 more) | imprecision ^a | functional capacity improvement | |
| Dyspnea | Relative risk 1.96 (95% CI 0.9 to 4.25) | 227 per 1000 | 445 per 1000 | Low | Treamid may increase | |
| improvement | Based on data from 59 participants in 1 study Follow-up 28 days | Difference: 218 more per 1000 (95% CI 23 fewer to 738 more) | | Due to very serious imprecision ^b | dyspnea improvement | |
| Pulmonary function improvement | Relative risk 2.48 (95% CI 1.0 to 6.17) Based on data from 59 | 167 per 1000 | 414 per 1000 | Low Due to very serious | Treamid may increase pulmonary function | |
| шрюченен | participants in 1 study Follow-up 28 days | | ' more per 1000 er to 863 more) | imprecision ^c | improvement | |
| Adverse events | Relative risk 1.19 (95% CI 0.56 to 2.5) | 290 per 1000 | 345 per 1000 | Low | Treamid may increase | |
| Adverse events | Based on data from 59 participants in 1 study Follow-up 28 days | | more per 1000 wer to 435 more) | Due to very serious imprecision ^d | adverse events | |

Imprecision: very serious. 95% CI includes important benefits and harms. Imprecision: very serious. 95% CI includes important benefits and harms. Imprecision: very serious. 95% CI includes important benefits and harms.

Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A12.

Population: Patients with P-ACC-related neurocognitive symptoms or sleep disturbances

Intervention: Actovegin

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | Plain language |
|-----------------------------|---|--|---------------------|-----------------------------------|-----------------------|
| | | SOC | Actovegin | Evidence (Quality of evidence) | summary |
| Cognitive improvement | Odds ratio: 1.19 (CI 95% 1.06 - 1.33) Based on data from 444 participants in 1 study | 673 per 1000 | 710 per 1000 | Low Due to very serious risk of | Actovegin may improve |
| improvement | | Difference: 37 more per 1000 (Cl 95% 13 more - 384 fewer) | | bias ¹ | cognition |

^{1.} **Risk of Bias: very serious.** Inadequate sequence generation/generation of comparable groups, resulting in potential for selection bias, Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Indirectness: serious.** Non appropriately established MID; **Imprecision: no serious.** 95% CI include important benefits and harms.

Summary of findings Table A13.

Population: Patients with P-ACC-related neurocognitive symptoms or sleep disturbances

Intervention: Hyperbaric oxygen (HBO) Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | Plain language |
|---------------------------|--|---|------------------------|---|---|
| | | soc | НВО | (Quality of evidence) | summary |
| HRQL improvement | Relative risk 1.3 (95% CI 0.84 to 2.0) Based on data from 73 participants in 1 study | 469 per 1000 | 610 per 1000 | Low Due to very serious | HBO may increase HRQF |
| | | Difference: 141 more per 1000 (95% CI 75 fewer to 469 more) | | imprecision ^a | improvement |
| Cognitive improvement | Odds ratio 2.84 (95% Cl 1.09 to 7.37) Based on data from 73 participants in 1 study | 667 per 1000 | 850 per 1000 | Very low Due to extremely | We are uncertain whether HBO increases or |
| | | Difference: 183 more per 1000 (95% CI 19 more to 22 more) | | serious imprecision, Due to serious indirectness ^b | decreases cognitive improvement |
| Depression improvement | Odds ratio 35.9 (95% CI 2.72 to 474.6) Based on data from 73 participants in 1 study Follow-up 28 days | 681 per 1000 | 987 per 1000 | Very low Due to extremely | We are uncertain whether HBO increases or |
| | | Difference: 306 more per 1000 (95% CI 172 more to 312 more) | | serious imprecision, Due to serious indirectness ^c | decreases depression improvement |

a. Imprecision: very serious. 95% CI includes important benefits and harms.

Indirectness: serious. Non appropriately established minimal important difference (MID). Imprecision: extremely serious.
 95% CI includes important benefits and harms.

c. Indirectness: serious. Non appropriately established MID. Imprecision: extremely serious. 95% CI includes important benefits and harms.

Summary of findings Table A14.

Population: Patients with P-ACC-related neurocognitive symptoms or sleep disturbances Intervention: Transcranial direct current stimulation (tDCS)

| | | Absolute e | effect estimates | Certainty of the evidence (Quality of evidence) | Plain language summary |
|----------------------|--|---|--|---|--|
| Outcome Timeframe | Outcome Study results and measurements | SOC | Transcranial direct current stimulation (tDCS) | | |
| Cognitive | Relative risk 0.59 Cognitive (95% CI 0.33 to 1.05) Based on data from 47 participants in 1 study Follow-up 30 days | 667 per 1000 | 394 per 1000 | Low Due to very serious | tDCS may have little or no difference on cognitive |
| improvement | | Difference: 273 fewer per 1000 (95% CI 447 fewer to 33 more) | | imprecision ^a | improvement |

Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A15.

Population: Patients with P-ACC-related olfactory and/or gustatory dysfunction Intervention: ADAPT-232 Comparator: Standard of care (SOC)

| | Study results and | Absolute effect estimates | | Certainty of the evidence | Plain language |
|---|---|---------------------------|--------------------------------|---|----------------|
| | measurements | soc | ADAPT-232 | (Quality of evidence) | summary |
| Olfactory symptoms improvement Relative risk 0.89 (95% CI 0.79 to 1.01) Based on data from 99 participants in 1 study Follow-up 21 days | 960 per 1000 | 854 per 1000 | Low Due to very serious | ADAPT-232 may have little or no difference on | |
| | Difference: 106 fewer per 1000 (95% CI 202 fewer to 10 more) | | imprecision ^a | olfactory symptoms | |

a. Imprecision: very serious. 95% CI includes important benefits and harms.

Summary of findings Table A16.

Population: Patients with P-ACC-related olfactory and/or gustatory dysfunction Intervention: Palmitoylethanolamide + Luteolin Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | |
|---|--|---------------------------|--------------------------------------|---|------------------------|
| | | SOC | Palmitoylethanola mide + Luteolin | Evidence (Quality of evidence) | Plain language summary |
| Olfactory symptoms improvement Relative risk: 3.13 (CI 95% 1.47 - 666.0) Based on data from 130 participants in 1 study Follow up 90 days | 167 per 1000 | 523 per 1000 | Low Due to very serious | Palmitoylethanolamide + luteolin may increase | |
| | Difference: 356 more per 1000 (Cl 95% 78 more - 833 more) | | imprecision ¹ | olfactory symptoms improvement | |

^{1.} **Imprecision: very serious.** 95% CI include important benefits and harms;

Summary of findings Table A17.

Population: Patients with P-ACC-related olfactory and/or gustatory dysfunction

Intervention: Steroids

| | Study results and | Absolute eff | ect estimates | Certainty of the | Plain language summary | |
|--|---|--|------------------------|---|--|--|
| | measurements | soc | Steroids | Evidence (Quality of evidence) | | |
| Relative risk: 1.09 Olfactory symptoms (CI 95% 0.83 - 1.44) improvement Based on data from 131 | | 365 per 1000 | 398 per 1000 | Low Due to very serious | Steroids may have little or no difference on olfactory | |
| improvoment | participants in 2 studies Follow up 52 days | Difference: 33 more per 1000 (CI 95% 62 fewer - 161 more) | | imprecision ¹ | symptoms | |
| Gustatory symptoms improvement (CI 95% 0.67 Based on data participants in | Relative risk: 1.01 (Cl 95% 0.67 - 1.53) Based on data from 113 | 443 per 1000 | 447 per 1000 | Low Due to serious risk of bias, | Steroids may have little or no difference on | |
| | participants in 1 study Follow up 84 days | Difference: 4 more per 1000 (CI 95% 146 fewer - 235 more) | | Due to serious imprecision ² | gustatory symptoms | |

^{1.} **Imprecision: very serious.** Low number of patients, Wide confidence intervals.

^{2.} **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.

Summary of findings Table A18.

Population: Patients with P-ACC-related psychological distress Intervention: Virtual reality informational video

| _ | | Absolute ef | fect estimates | Certainty of the | |
|----------------------------------|--|--|-----------------------------------|--|---|
| Outcome Timeframe | Study results and measurements | SOC | Virtual informational video | Evidence (Quality of evidence) | Plain language summary |
| Depression improvement | Relative risk 1.21 (95% CI 0.95 to 1.54) Based on data from 89 | 682 per 1000 | 825 per 1000 | Low Due to serious risk of bias, | Virtual reality informational video may |
| Improvement | participants in 1 study Follow-up 90 days | Difference: 143 more per 1000 (95% CI 34 fewer to 368 more) | | Due to serious imprecision ^a | increase depression improvement |
| Post-traumatic stress disorder | Relative risk 1.18 (95% CI 0.98 to 1.42) Based on data from 89 participants in 1 study Follow-up 90 days | 773 per 1000 | 912 per 1000 | Low Due to serious risk of bias, | Virtual reality informational video may increase post-traumatic |
| improvement | | Difference: 139 more per 1000 (95% CI 15 fewer to 227 more) | | Due to serious imprecision ^b | stress disorder improvement |
| Psychologic distress improvement | Relative risk 1.49 (95% CI 1.08 to 2.05) Based on data from 89 participants in 1 study Follow-up 90 days | 523 per 1000 | 779 per 1000 | Low Due to serious risk of bias, inf | Virtual reality informational video may |
| | | Difference: 256 more per 1000 (95% CI 42 more to 549 more) | | Due to serious imprecision ^c | increase psychological distress improvement |

Risk of bias: serious. Imprecision: serious. Low number of patients.

b.

Risk of bias: serious. Imprecision: serious. Low number of patients. Risk of bias: serious. Imprecision: serious. Low number of patients.

Summary of findings Table A19.

Population: Patients with P-ACC-related thromboembolic risk

Intervention: Anticoagulants Comparator: Standard of care

| Outcome | Study results and | Absolute effect estimates | Certainty of the Evidence | Plain language | |
|------------------|--|---|---|---|--|
| Timeframe | measurements | SOC Anticoagulants | (Quality of evidence) | summary | |
| HRQL improvement | Relative risk: 0.99 (Cl 95% 0.78 - 1.24) Based on data from 1217 participants in 1 study Follow up 90 days | | Low Due to very serious imprecision ¹ | Anticoagulants may have little or no difference on HRQL | |
| VTE | Relative risk: 1.0 (Cl 95% 0.29 - 3.45) Based on data from 1535 participants in 2 studies Follow up 32.5 days | 32 per 1000 per 1000 Difference: 0 fewer per 1000 (Cl 95% 23 fewer - 78 more) | Low Due to very serious imprecision ² | Anticoagulants may have little or no difference on VTE | |
| Mortality | Relative risk: 0.85 (CI 95% 0.38 - 1.88) Based on data from 1535 participants in 2 studies Follow up 32.5 days | 20 17 per 1000 per 1000 Difference: 3 fewer per 1000 (Cl 95% 12 fewer - 18 more) | Low Due to very serious imprecision ³ | Anticoagulants may have little or no difference on mortality | |
| Major bleeding | Relative risk: 2.01 (CI 95% 0.18 - 22.1) Based on data from 1535 participants in 2 studies Follow up 32.5 days | 1 2 per 1000 per 1000 Difference: 1 more per 1000 (CI 95% 1 fewer - 21 more) | Low Due to very serious imprecision ⁴ | Anticoagulants may have little or no difference on major bleeding | |

- 1. **Imprecision: very serious.** Low number of patients, Wide confidence intervals.
- 2. **Imprecision: very serious.** Low number of patients, Wide confidence intervals.
- 3. **Imprecision: very serious.** Low number of patients, Wide confidence intervals.
- 4. **Imprecision: very serious.** Low number of patients, Wide confidence intervals.

Summary of findings Table A20.

Population: Patients with PIMS-TS

Intervention: Steroids Comparator: IVIG

| | | Ah as luta offert actionates | | | |
|---|---|---|---|--|--|
| Outcome Timeframe | Study results and measurements | Absolute effect estimates IVIG Steroids | Certainty of the Evidence (Quality of evidence) | Plain language summary | |
| Time to discharge time reduction ¹ | Relative risk: 1.09 (CI 95% 0.88 - 1.39) Based on data from 75 participants in 1 study | 500 545 per 1000 per 1000 Difference: 45 more per 1000 | Low Due to serious risk of bias, Due to serious | Steroids may decrease time to discharge | |
| | Follow up 28 | (Cl 95% 60 fewer - 195 more) | imprecision ² | | |
| Respiratory support | Relative risk: 0.49 (Cl 95% 0.27 - 0.89) Based on data from 75 | 553 271 per 1000 per 1000 | Low Due to serious risk of bias, | Steroids may decrease respiratory support | |
| | participants in 1 study Follow up 28 | Difference: 282 fewer per 1000 (CI 95% 404 fewer - 61 fewer) | Due to serious imprecision ³ | requirements | |
| Inotropic requirements | Relative risk: 0.68 (CI 95% 0.35 - 1.32) Based on data from 75 participants in 1 study Follow up 28 | 395 269 per 1000 per 1000 | Very low Due to serious risk of bias, Due to serious | We are uncertain whether steroids increases or | |
| · | | Difference: 126 fewer per 1000 (CI 95% 257 fewer - 126 more) | imprecision, Due to very serious imprecision ⁴ | decreases inotropic requirements | |
| Left ventricular fraction deterioration | Relative risk: 0.57 (Cl 95% 0.21 - 1.54) Based on data from 75 | 237 135 per 1000 per 1000 | Very low Due to serious risk of bias, Due to serious | We are uncertain whether steroids increases or | |
| | participants in 1 study Follow up 28 | Difference: 102 fewer per 1000 (CI 95% 187 fewer - 128 more) | imprecision, Due to very serious imprecision ⁵ | decreases LVEF deterioration | |
| Arrhythmia | Relative risk: 2.05 (Cl 95% 0.19 - 21.7) Based on data from 75 | 26 53 per 1000 per 1000 | Very low Due to serious risk of bias, Due to serious | We are uncertain whether steroids increases or | |
| | participants in 1 study Follow up 28 | Difference: 27 more per 1000 (Cl 95% 21 fewer - 538 more) | imprecision, Due to very serious imprecision ⁶ | decreases Arrhythmias | |
| Venous thromboembolic | Relative risk: 0.34 (CI 95% 0.01 - 8.14) Based on data from 75 | 39 13 per 1000 per 1000 | Very low Due to serious risk of bias, Due to serious | We are uncertain whether steroids increases or | |
| events | participants in 1 study Follow up 28 | Difference: 26 fewer per 1000 (Cl 95% 39 fewer - 278 more) | imprecision, Due to very serious imprecision ⁷ | decreases VTE | |

- 1. Proportion of patients discharged on day 6.
- 2. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Wide confidence intervals.
- 3. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Wide confidence intervals.
- 4. Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: very serious. Wide confidence intervals.
- 5. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: very serious.** Wide confidence intervals.

- 6. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: very serious.** Wide confidence intervals.
- 7. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: very serious.** Wide confidence intervals.

Summary of findings Table A21.

Population: Patients with COVID-19 Intervention: Metformin to prevent P-ACC

| Outcome Study results and | Absolute effect estimates | | Certainty of the | Plain language | |
|--|---|---------------------|--|-----------------------------------|----------------------|
| Timeframe | | SOC | Metfomin to prevent P-ACC | Evidence (Quality of evidence) | summary |
| P-ACC | Relative risk: 0.59 (CI 95% 0.39 - 0.88) P-ACC Record on data from 1135 | 105 per 1000 | 62 per 1000 | Low Due to serious risk of | Metformin may reduce |
| Based on data from 1125 participants in 1 study Follow up 300 days | Difference: 43 fewer per 1000 (Cl 95% 64 fewer - 13 fewer) | | bias, Due to serious imprecision ¹ | P-ACČ | |

^{1.} **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.

Summary of findings Table A22.

Population: Patients with COVID-19 Intervention: Ivermectin to prevent P-ACC

| Outcome | Study results and | Absolute effect estimates | | Certainty of the | Plain language |
|-----------|---|--|-----------------------------|---|--------------------|
| Timeframe | | soc | Ivermectin to prevent P-ACC | Evidence (Quality of evidence) | summary |
| P-ACC | P-ACC Relative risk: 0.99 (CI 95% 0.61 - 1.62) Based on data from 738 participants in 1 study Follow up 300 days | 105 per 1000 | 104 per 1000 | Low Due to serious risk of bias, | Ivermectin may not |
| | | Difference: 1 fewer per 1000 (CI 95% 41 fewer - 65 more) | | Due to serious imprecision¹ | reduce P-ACC |

^{1.} **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.

Summary of findings Table A23.

Population: Patients with COVID-19

Intervention: Convalescent plasma to prevent P-ACC

| Outcome Study results and | Absolute effect estimates | | Certainty of the | Plain language | |
|---|--|--|--|--|---------|
| Timeframe | measurements | SOC | CP to prevent P- ACC | Evidence (Quality of evidence) | summary |
| Relative risk: 0.93 (CI 95% 0.77 - 1.12) | 343 per 1000 | 319 per 1000 | Low Due to serious risk of bias, Due to serious imprecision ¹ | Convalescent plasma may not reduce P-ACC | |
| | Based on data from 882 participants in 1 study Follow up 90 days | Difference: 24 fewer per 1000 (CI 95% 79 fewer - 41 more) | | | |

^{1.} **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.

Summary of findings Table A24.

Population: Patients with COVID-19 Intervention: Remdesivir to prevent P-ACC

| Outcome | Study results and | Absolute effect estimates | | Certainty of the | Plain language | |
|-----------|--|--|--|---------------------------------------|--|--|
| Timeframe | measurements | soc | Remdesivir to prevent P-ACC | Evidence (Quality of evidence) | summary | |
| Mortality | Relative risk: 0.85 (CI 95% 0.25 - 2.83) Based on data from 181 participants in 1 study Follow up 365 days | 60 per 1000 | 51 per 1000 | Very low Due to serious risk of bias, | We are uncertain whether remdesivir to prevent p-acc | |
| | | Difference: 9 fewer per 1000 (CI 95% 45 fewer - 110 more) | | Due to very serious imprecision | increases or decreases mortality | |
| P-ACC | Relative risk: 1.06 (CI 95% 0.53 - 2.13) Based on data from 181 | 145 per 1000 | 154 per 1000 | Low Due to serious risk of bias. | Remdesivir may not | |
| | participants in 1 study Follow up 365 days | | 9 more per 1000 ewer - 164 more) | Due to serious imprecision | reduce P-ACC | |

- 1. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** Low number of patients, Wide confidence intervals.
- Risk of Bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: serious. Low number of patients.

Summary of findings Table A25.

Population: Patients with COVID-19

Intervention: Leflunomide to prevent P-ACC

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | Plain language |
|-----------------------------|---|---------------------------|---|--|--|
| | | soc | Leflunomide to prevent P-ACC | Evidence (Quality of evidence) | summary |
| P-ACC | Relative risk: 1.28 (CI 95% 0.92 - 1.77) Based on data from 172 participants in 1 study Follow up 90 days | | 521 per 1000 14 more per 1000 ewer - 313 more) | Low Due to serious risk of bias, Due to serious imprecision ¹ | Remdesivir may not reduce P-ACC |
| Severe adverse events | Relative risk: 1.76 (CI 95% 0.81 - 3.85) Based on data from 214 participants in 1 study Follow up 90 days | | 144 per 1000 2 more per 1000 ewer - 234 more) | Very low Due to serious risk of bias, Due to very serious imprecision ² | We are uncertain whether remdesivir to prevent pacc increases or decreases mortality |

- 1. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: serious.** Low number of patients.
- Risk of Bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: very serious. Low number of patients, Wide confidence intervals.

Summary of findings Table A26.

Population: Patients with COVID-19 Intervention: Fluvoxamine to prevent P-ACC Comparator: SOC

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | Plain language |
|---|--|---|------------------------------|--|---------------------|
| | | soc | Fluvoxamine to prevent P-ACC | Evidence (Quality of evidence) | summary |
| P-ACC (CI 95% 0.81 - 1.2 Based on data from participants in 2 stu | Relative risk: 0.99 (CI 95% 0.81 - 1.21) | 444 per 1000 | 440 per 1000 | Low | Fluvoxamine may not |
| | participants in 2 studies Follow up 192 days | Difference: 4 fewer per 1000 (Cl 95% 84 fewer - 93 more) | | Due to very serious imprecision ¹ | reduce P-ACC |

^{5.} **Imprecision: very serious.** Wide confidence intervals.

Summary of findings Table A27.

Population: Patients with P-ACC-related asthenia or fatigue Intervention: AXA1125

| Outcome | Study results and | Absolute effect estimates | | Certainty of the Evidence | Plain language |
|---------------------|--|---|------------------------------------|---|---|
| Timeframe | measurements | SOC | AXA1125 | (Quality of evidence) | summary |
| Fatigue improvement | Relative risk: 1.07 (Cl 95% 0.79 - 1.44) Based on data from 41 participants in 1 study Follow up 28 days | 780 per 1000 | 835 per 1000 | Low Due to very serious | Axa1125 may increase |
| | | Difference: 55 more per 1000 (CI 95% 164 fewer - 343 more) | | imprecision ¹ | fatigue improvement |
| Functional capacity | Relative risk: 0.87 (Cl 95% 0.51 - 1.48) Based on data from 41 participants in 1 study Follow up 28 days | 607 per 1000 | 528 per 1000 | Low | Axa1125 may not |
| improvement | | | fewer per 1000 ewer - 291 more) | Due to very serious imprecision ² | increase functional capacity improvement |
| Adverse events | Relative risk: 2.62 (Cl 95% 1.0 - 6.89) | 200 per 1000 | 524 per 1000 | , | We are uncertain whether |
| | Based on data from 41 participants in 1 study Follow up 28 days | | 4 more per 1000 ver - 800 more) | Due to extremely serious imprecision ³ | axa1125 improves or worsen adverse events |

- **Imprecision: very serious.** 95% CI include important benefits and harms.
- **Imprecision:** very serious. 95% CI include important benefits and harms.
- Imprecision: ~extreme_serious. 95% CI include important benefits and harms.

Summary of findings Table A28.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Telerehabilitation Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and | Absolute effect estimates | | Certainty of the | Plain language |
|---------------------------------|--|--|---|--|--|
| | measurements | SOC | Telerehabilitation | (Quality of evidence) | summary |
| HRQL improvement | Relative risk: 1.08 (Cl 95% 0.86 - 1.36) Based on data from 205 participants in 4 studies | 475 per 1000 | 513 per 1000 | Very low Due to very serious imprecision, Due to | We are uncertain whether telerehabilitation increases or decreases |
| | Follow up 66.5 days | Difference: 38 more per 1000 (Cl 95% 66 fewer - 171 more) | | serious risk of bias ¹ | hrql improvement |
| Fatigue improvement | Relative risk: 1.63 (CI 95% 1.04 - 2.54) Based on data from 89 | 304 per 1000 | 636 per 1000 | Low Due to serious risk of | Telerehabilitation may increase fatigue |
| | participants in 3 studies Follow up 79 days | | 68 more per 1000 fewer - 672 more) | bias, Due to serious imprecision ² | improvement |
| Functional capacity improvement | Relative risk: 1.37 (CI 95% 1.11 - 1.67) Based on data from 219 | 399 per 1000 | 547 per 1000 | Low Due to serious risk of | Telerehabilitation may |
| | participants in 4 studies Follow up 81 days | | 48 more per 1000 more - 267 more) | bias, Due to serious imprecision ³ | capacity improvement |

- 1. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: very serious.** 95% CI include important benefits and harms, Wide confidence intervals;
- Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: serious. Low number of patients;
- 3. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** 95% CI include important benefits and harms, Low number of patients;

Summary of findings Table A29.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Cognitive behavioral therapy Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | |
|---------------------------------|---|--|------------------------------------|--|---|
| | | soc | Cognitive behavioral therapy | Evidence (Quality of evidence) | Plain language summary |
| Fatigue improvemen | Relative risk: 2.2 (CI 95% 1.35 - 3.58) Based on data from 114 | 263 per 1000 | 636 per 1000 | Low Due to serious risk of bias, | Cognitive behavioral therapy may increase |
| | participants in 1 studies Follow up 119 days | Difference: 168 more per 1000 (CI 95% 112 fewer - 672 more) | | Due to serious imprecision ¹ | fatigue improvement |
| Functional capacity improvement | Relative risk: 1.37 (CI 95% 1.08 - 1.73) Based on data from 114 | 614 per 1000 | 841 per 1000 | Low Due to serious risk of bias, | Cognitive behavioral therapy may increase |
| | participants in 1 studies Follow up 119 days | Difference: 227 more per 1000 (CI 95% 49 more - 448 more) | | Due to serious imprecision ² | functional capacity improvement |

^{1.} **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Low number of patients;

^{2.} **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** 95% CI include important benefits and harms, Low number of patients;

Summary of findings Table A30.

Population: Patients with P-ACC-related asthenia or fatigue

Intervention: Physical training Comparator: Standard of care (SOC)

| Outcome | Study results and | Absolute effect estimates | | Certainty of the Evidence | Summary |
|--------------------------------------|--|--|--------------------------------------|---|--|
| Timeframe | measurements | soc | Physical training | (Quality of evidence) | Julillary |
| HRQL improvement ¹ | Relative risk: 1.62 (CI 95% 1.07 - 2.47) Based on data from 180 | 340 per 1000 | 551 per 1000 | Low Due to serious risk of bias, | Physical training may increase HRQL |
| | participants in 3 studies Follow up 70 days | Difference: 211 more per 1000 (Cl 95% 24 more - 500 more) | | Due to serious imprecision ² | improvement |
| Functional capacity | Relative risk: 1.81 (CI 95% 1.23 - 2.65) | 393 per 1000 | 711 per 1000 | Low Due to serious risk of bias, | Physical training may |
| improvement ³ | Based on data from 226 participants in 5 studies Follow up 56 days | Difference: 318 more per 1000 (CI 95% 90 more - 648 more) | | Due to serious imprecision ⁴ | increase functional capacity improvement |
| Fatigue | Relative risk: 2.1 (CI 95% 0.8 - 5.02) Based on data from 213 | 121 per 1000 | 254 per 1000 | Low Due to serious risk of bias, | Uncertainty about |
| improvement ⁵ | participants in 4 studies Follow up 38.5 days | | 33 more per 1000 ewer - 486 more) | Due to very serious imprecision ⁶ | interventions effects |
| Strength improvement ⁷ | Relative risk: 3.13 (Cl 95% 1.02 - 9.55) | 53 per 1000 | 166 per 1000 | Low Due to very serious | We are uncertain whether cognitive |
| | Based on data from 132 participants in 2 studies Follow up 70 days | | 13 more per 1000 nore - 453 more) | imprecision, Due to serious imprecision ⁸ | behavioral therapy increases or decreases strength improvement |

- 1. Decrease in 12 units of the MFI score
- Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: serious. Low number of patients;
- 3. Decrease in 12 units of the MFI score
- 4. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** 95% CI include important benefits and harms, Low number of patients;
- 5. Decrease in 12 units of the MFI score
- 6. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: very serious.** Low number of patients, Wide confidence intervals;
- 7. Decrease in 12 units of the MFI score
- 8. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Low number of patients;

Summary of findings Table A31.

Population: Patients with P-ACC-related neurocognitive symptoms

Intervention: Famotidine

| Outcome | Study results and | Absolute effect estimates | | Certainty of the | Plain language | |
|------------------------|--|--|---|------------------------------------|--|------------------------|
| Timeframe | measurements | soc | Famotidine | Evidence (Quality of evidence) | summary | |
| Cognitive improvement | Relative risk: 1.33 (CI 95% 1.05 - 1.69) Based on data from 50 | 739 per 1000 | 983 per 1000 | Low Due to very serious | Due to very serious Famotidine may impr | Famotidine may improve |
| | participants in 1 study | Difference: 244 more per 1000 (CI 95% 37 more - 510 more) | | imprecision ¹ | cognition | |
| Depression improvement | Relative risk: 3.71 (CI 95% 0.83 - 16.6) Based on data from 50 | 76 per 1000 | 282 per 1000 | Very low Due to extremely serious | We are uncertain whether famotidine increases or | |
| | participants in 1 study | | 96 more per 1000 ewer - 926 more) | imprecision ² | decreases depression improvement | |

^{1.} **Indirectness: serious.** Non appropriately established MID; **Imprecision: very serious.** Low number of patients;

^{2.} Indirectness: serious. Imprecision: ~extremely_serious. 95%CI includes important benefits and harms,

Summary of findings Table A32.

Population: Patients with P-ACC-related olfactory and/or gustatory dysfunction

Intervention: Olfactory training Comparator: Standard of care (SOC)

| Outcome Timeframe | Study results and measurements | Absolute effect estimates | | Certainty of the | Plain language |
|--------------------------------|---|---------------------------|--------------------------------------|--|--|
| | | soc | Olfactory training | Evidence (Quality of evidence) | summary |
| Olfactory symptoms improvement | Relative risk: 1.49 (Cl 95% 0.96 - 2.31) Based on data from 308 | 409 per 1000 | 609 per 1000 | Low Due to serious risk of bias, Due to serious inconsistency ¹ | Olfactory training may increase olfactory symptoms improvement |
| | participants in 4 studies Follow up 59.5 days | | 00 more per 1000 ewer - 536 more) | | |

^{1.} **Risk of Bias: serious. Inconsistency: serious.** The confidence interval of some of the studies do not overlap with those of most included studies/ the point estimate of some of the included studies.;

This review compiles the evidence on potential therapeutic options for post COVID-19 condition (PCC). Included are all the identified clinical forms, symptoms and manifestations of PCC for which an intervention was assessed in at least one randomized controlled trial.