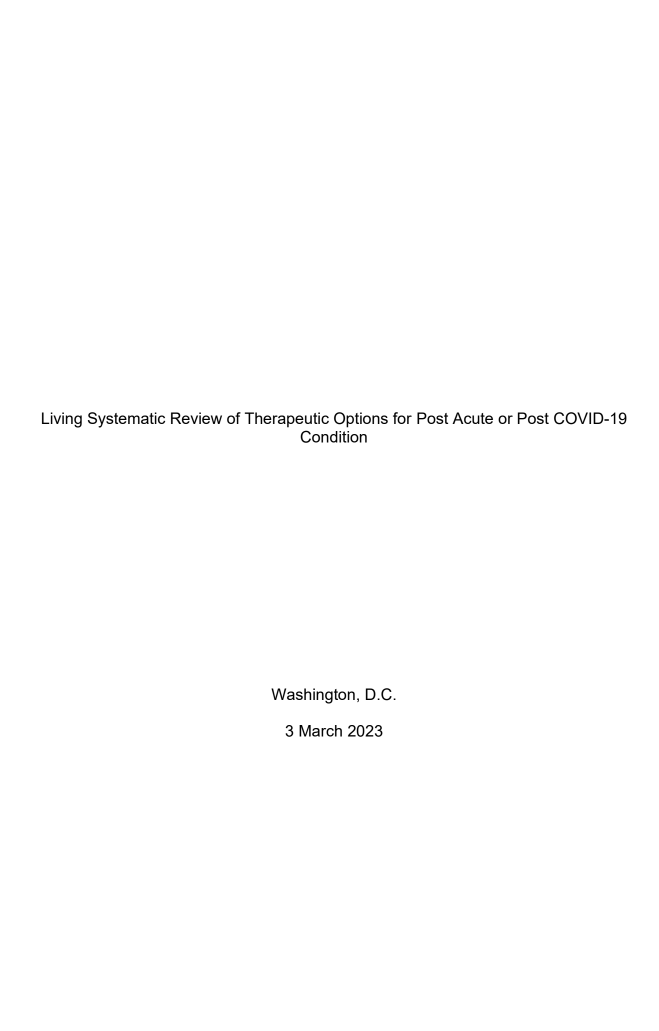


LIVING SYSTEMATIC REVIEW
OF THERAPEUTIC OPTIONS FOR
POST-ACUTE AND POST-COVID19
CONDITION

March 3th 2023



Living Systematic Review of Therapeutic Options for Post Acute or Post COVID-19 condition.

3 March 2023

PAHO/IMS/EIH/COVID-19/23-0006

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#### Disclaimer

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.

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# **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 states 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 the 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 ES13) 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 ES14) summarize the status of evidence for the 28 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 = 16)

Intervention		Overall number of studies including the intervention, n=16	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)
Fermented food supplements		2		2					
Physical training	NEW	2	0 9	10	1	1			
1_MNA		1			1	1			
Actovegin	NEW	1			1				
ADAPT_232 (adaptogens)		1			1				
Arginine_Vitamin C		1			1				
CQ10		1	3	1 1					
Cytoflavin		1			1				
Enzimes_Probiotics		1			1				
Hydrogen (nasal)		1			1	1			
Phytochemicals	NEW	1	1	1	1				
Leronlimab		1		1					
tDCS		1			1			1	
Telerehabilitation		1	1	1	1	4	1.		



**Table ES2.** Summary of findings on potential therapeutic options for P-ACC-related asthenia or fatigue (n = 14), as of 28 February 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	Arginine + Vitamin C	Uncertainty in potential benefits and harms. Further research is needed.
5	Coenzyme Q10	Uncertainty in potential benefits and harms. Further research is needed.
6	Cytoflavin	Cytoflavin may not improve fatigue. However, certainty of the evidence was low. Further research is needed.
7	Enzymes + probiotics	Enzymes + probiotics may improve fatigue. However, certainty of the evidence was low. Further research is needed.
8	Fermented food supplements	Uncertainty in potential benefits and harms. Further research is needed.
9	Hydrogen (nasal)	Uncertainty in potential benefits and harms. Further research is needed.
10	Leronlimab	Uncertainty in potential benefits and harms. Further research is needed.
11	Phytochemicals	Phytochemicals may improve fatigue and HRQL. However, certainty of the evidence was low. Further research is needed.
12	Physical training	Uncertainty in potential benefits and harms. Further research is needed.
13	Transcranial direct current stimulation (tDCS)	tDCS may not improve fatigue and may not increase adverse events. However, certainty of the evidence was low. Further research is needed.

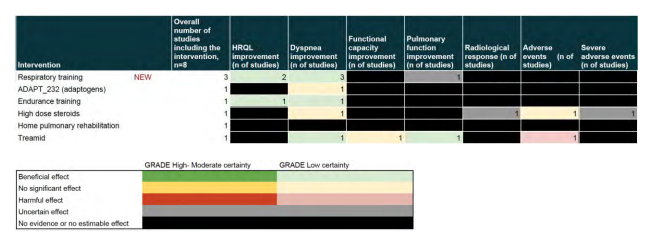
	Intervention	Summary of findings
14	Telerehabilitation	Uncertainty in potential benefits and harms. Further research is needed.

- 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 12 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.
- 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.
- Transcranial direct current stimulation (tDCS): The results of one RCT suggest that tDCS may not improve fatigue and may not increase adverse events. However, certainty of the evidence was low because of imprecision. 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 = 8)



**Table ES4.** Summary of findings on potential therapeutic options for P-ACC-related dyspnea (n = 6), as of 28 February 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	Endurance training	Endurance training may improve health-related quality of life (HRQL) and dyspnea. However, certainty of the evidence was low. Further research is needed.
3	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.
4	Home pulmonary rehabilitation	Uncertainty in potential benefits and harms. Further research is needed.
5	Respiratory training	Respiratory training may improve HRQL and dyspnea. However, certainty of the evidence was low. Further research is needed.

	Intervention	Summary of findings
6	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.

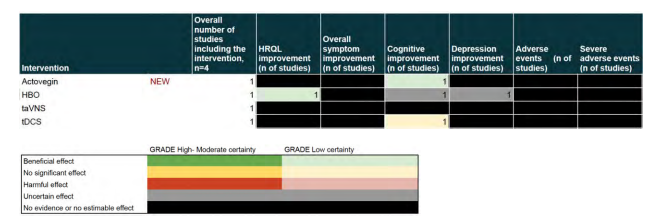
- Therapeutic options: According to the WHO International Clinical Trials Registry Platform (ICTRP), multiple potential interventions are being assessed in hundreds of clinical trials and observational studies. In this review, we identified and examined five therapeutic options for P-ACC-related dyspnea.
- 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.
- Endurance training: The results of one RCT suggest that endurance training may improve dyspnea and HRQL compared to standard physiotherapy. However, certainty of the evidence was low because of imprecision and risk of bias. 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:** The results of three RCTs suggest that respiratory training may improve dyspnea and HRQL. However, certainty of the evidence was low because of imprecision, inconsistency and risk of bias. Further research is needed.



• **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

**Table ES5.** List of RCTs of interventions for P-ACC-related neurocognitive symptoms with primary outcome measures and certainty (n = 4)



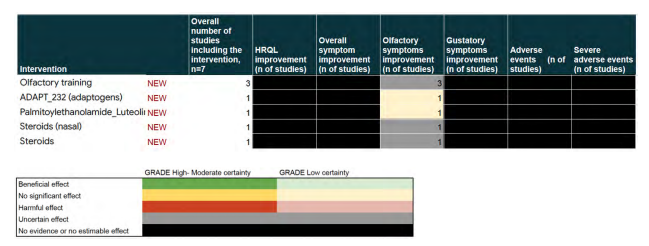
**Table ES6.** Summary of findings on potential therapeutic options for P-ACC-related neurocognitive symptoms (n = 4), as of 28 February 2023

	Intervention	Summary of findings
1	Actovegin	Actovegin may improve cognition. However, certainty of the evidence was low. Further research is needed.
2	Hyperbaric oxygen (HBO)	HBO may improve HRQL. However, certainty of the evidence was low. Further research is needed.
3	Transcutaneous auricular vagus nerve stimulation (taVNS)	Uncertainty in potential benefits and harms. Further research is needed.
4	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 three therapeutic options for PCC neurocognitive symptoms.
- **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.
- **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 = 7)



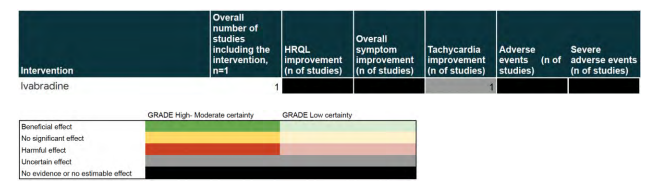
**Table ES8.** Summary of findings on potential therapeutic options for P-ACC-related olfactory and/or gustatory dysfunction (n = 5), as of 28 February 2023

	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	Olfactory training	Uncertainty in potential benefits and harms. Further research is needed.
3	Palmitoylethanolamide + Luteolin	Palmitoylethanolamide + Luteolin may not improve olfactory symptoms. However, certainty of the evidence was low. Further research is needed.
4	Steroids (nasal)	Uncertainty in potential benefits and harms. Further research is needed.
5	Steroids	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 five 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.
- Palmitoylethanolamide + Luteolin: The results of one RCT suggest that Palmitoylethanolamide + Luteolin may not improve olfactory 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 = 1)



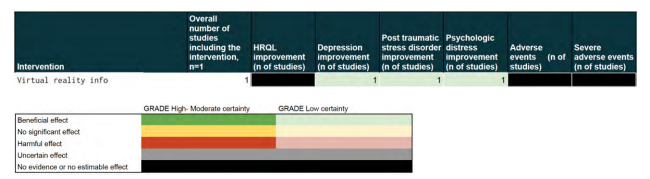
**Table ES10.** Summary of findings on potential therapeutic options for P-ACC-related cardiovascular system symptoms (n = 1), as of 28 February 2023

	Intervention	Summary of findings
1	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 one therapeutic option 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 = 1)



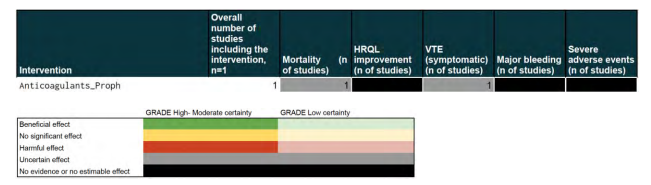
**Table ES12.** Summary of findings on potential therapeutic options for PCC psychological distress (n = 1), as of 28 February 2023

	Intervention	Summary of findings
1	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 = 1)



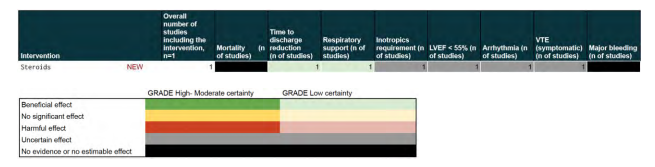
**Table ES14.** Summary of findings on potential therapeutic options for PCC thromboembolic risk (n = 1), as of 28 February 2023

	Intervention	Summary of findings
1	Anticoagulants (prophylactic dose)	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 one therapeutic option for PCC olfactory and/or gustatory dysfunction.
- The effects of assessed interventions are uncertain.

# 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 28 February 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.				

# Key findings

• 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.

#### Changes since previous edition

- Actovegin for P-ACC-related neurocognitive symptoms: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Home pulmonary rehabilitation for P-ACC-related dyspnea: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- **Steroids for PIMS-TS:** New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Phytochemicals for P-ACC-related asthenia or fatigue: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Physical training for P-ACC-related asthenia or fatigue: New evidence included affecting results interpretation and/or certainty of the evidence judgments.
- Respiratory training for P-ACC-related dyspnea: New evidence included without significant changes.

# 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 evidence-based 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 post acute 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 states 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 the 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 **OVerview** Evidence (L·OVE: available used the Living 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: <a href="https://app.iloveevidence.com/loves/5e6fdb9669c00e4ac072701d?question\_domain=un\_defined&section=methods">https://app.iloveevidence.com/loves/5e6fdb9669c00e4ac072701d?question\_domain=un\_defined&section=methods</a>). 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 28 February 2023. 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%.

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: <a href="https://app.magicapp.org/">https://app.magicapp.org/</a>) to generate the tables summarizing our findings, which are included in Annex 1.

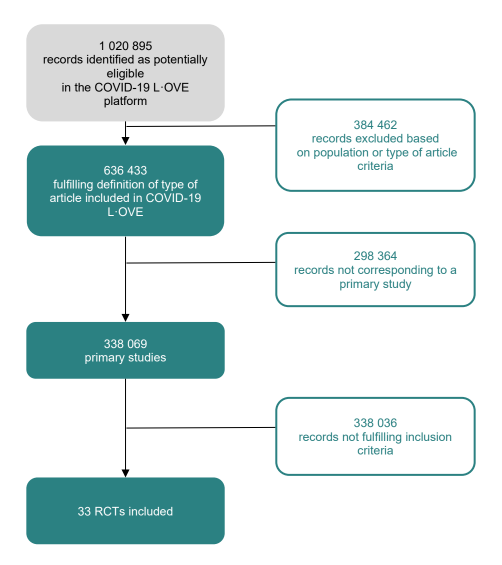


#### Results

#### Studies identified and included

The study identification and selection process is shown in Figure 1. A total of 28 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 randomization process	Risk-of-bias due to deviations from the intended interventions	Risk-of-bias due to missing outcome data	Risk-of-bias in measurement of the outcome	Risk-of-bias in selection of the reported result	Overall Risk-of-bias judgement	
						Mortality	HRQL, symptom specific outcomes
Vaira 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	Low	Low	Low	Low	Low	Low	Low
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
UK Phyto-V	High	Some Concerns	Low	Some Concerns	Low	High	High
Rodriguez-Blanco	High	Some Concerns	Low	Some Concerns	Low	High	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 ⊕⊕○○



#### **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 one RCT including 47 patients in which tDCS was compared against standard of care. Our results showed:

tDCS may not improve fatigue, RR 0.95 (95% CI 0.5 to 1.79); RD −2.4% (95% CI −22.8% to 36.4%); 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 ⊕⊕○○

#### Endurance training

#### See Summary of findings Table A8 in Annex 1

We identified one RCT including 60 patients in which endurance training was compared against standard physiotherapy. Our results showed:

- Endurance training may improve HRQL, RR 1.48 (95% CI 0.92 to 2.37); RD 21% (95% CI −3.4% to 60%); Low certainty ⊕⊕○○
- Endurance training may improve dyspnea, RR 2.03 (95% CI 0.98 to 4.21);
   RD 24% (95% CI −0.4% to 76%); 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

#### See Summary of findings Table A10, Annex 1

We identified three RCTs including 271 patients in which different modalities of respiratory training were compared with standard of care. Our results showed:

- Respiratory training may improve HRQL, RR 1.93 (95% CI 1.3 to 2.86); RD 24.1% (95% CI 7.8% to 48.1%); Low certainty ⊕⊕○○ (see Figure 2)
- Respiratory training may improve dyspnea, RR 1.86 (95% CI 1.38 to 2.49); RD 22.9% (95% CI 10.1% to 39.7%); Low certainty ⊕⊕○○

**Figure 2.** HRQL in RCTs comparing respiratory training with standard of care for treatment of patients with P-ACC-related dyspnea

Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Rob = High ENO Breathe McNarry Rodriguez-Blanco Fixed effect model Random effects model Heterogeneity: $I^2 = 95\%$ ,	-		1.45 24.56 [1.86	[ 0.63; 1.91] [ 0.99; 2.12] 10.28; 58.67] [ 1.38; 2.49] 0.76; 13.71]	28.1% 60.4% 11.5% 100.0%	33.8% 34.7% 31.5% 
Fixed effect model Random effects mode Heterogeneity: $I^2 = 95\%$ , Residual heterogeneity: $I^2$	$\tau^2 = 1.5383, p < 0.01$	0.1 0.5 1 2 10		[ 1.38; 2.49] 0.76; 13.71]	100.0% 	100.0%

#### 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

#### 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 ⊕⊕○○

### 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 ⊕⊕⊖○

#### Palmitoylethanolamide + Luteolin

#### See Summary of findings Table A16, Annex 1

We identified one RCT including 126 patients in which palmitoylethanolamide + luteolin was compared with standard of care. Our results showed:

Palmitoylethanolamide + luteolin may not improve olfactory symptoms, RR 1.11 (95% CI 0.68 to 1.81); RD 4.1% (95% CI −11.7% to 29.7%); Low certainty ⊕⊕⊖○

## 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 A17, 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

The effects of the assessed interventions are uncertain.



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

#### See Summary of findings Table A18, 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 ⊕⊕⊖⊝

# Full description of included studies

Tables 2 to 8 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

	Uncertainty	1 in potential benefits a	-MNA and harms. Further res	search 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		
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; 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  Fatigue improvement: RR 1.02 (95% CI 0.84 to 1.24); RD 1.6% (95% CI −12.6% to 18.9%); Low certainty ⊕⊕○○  Functional capacity improvement: No information  Strength improvement: No information  Adverse events: No information  Severe adverse events: No information			

	Uncertainty	Arginine y 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
		ı	RCT		
Tosato et al. (17); 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

	Coenzyme Q10 (CQ10) 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						
Hansen et al. (18); 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%	NR	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				

Cytofla	Cytoflavin Cytoflavin 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				
			 RCT						
CITADEL trial (19), 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 ⊕⊕○○  Functional capacity improvement: No information  Strength improvement: No information  Adverse events: No information  Severe adverse events: No information				

Enzymes +	probiotics may impro		+ probiotics	e was low. Further resea	ırch 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		
Rathi et al. (20); Peer reviewed; 2022	Patients with P-ACC (asthenia or fatigue after acute COVID-19). 100 assigned to enzymes + probiotics ImmunoSEB (500 mg/capsule) + ProbioSEB CSC3 (5 billion CFUs /capsule) and 100 assigned to standard of care.	Mean age 41.2 ± 13, male 63.5%, interval between COVID-19 and enrolment 19.5 days, one comorbidity 14.5%	NR	High risk of bias  Notes: Concealment of allocation and blinding probably inappropriate.	HRQL improvement: No information  Overall symptom improvement: No information  Fatigue improvement: RR 6.07 (95% CI 3.79 to 9.71); RD 76% (95% CI 41.8% to 85%); Low certainty ⊕⊕○○  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				
		F	RCT						
Kharaeva et al. (21); 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. (21); 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				



	Uncertainty	Hydrog v in potential benefits a	en (nasal) 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		
Botek et al. (22); 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

	Leronlimab 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	СТ						
Gaylis et al. (23); 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 ① ① ○ Fatigue improvement: No information  Functional capacity improvement: No information  Strength 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		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. (24); 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: Very low certainty ⊕○○○  Overall symptom improvement: No information				
Rodriguez- Blanco et al; (25) 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.	Fatigue improvement: Very low certainty ⊕○○○  Functional capacity improvement: Very low certainty ⊕○○○  Strength improvement: Very low certainty ⊕○○○  Adverse events: No information  Severe adverse events: No information				



Phytochemicals Phytochemicals may improve fatigue and 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		
		F	RCT				
UK Phyto-V trial; (26) 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 and 73 assigned to standard of care.	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 ⊕⊕⊖⊖  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		

tDCS may not	Transcranial direct current stimulation (tDCS) tDCS may not improve fatigue and may not increase adverse events. 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						
Oliver-Mas et al. (27); 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  Fatigue improvement: RR 0.95 (95% CI 0.5 to 1.79); RD − 2.4% (95% CI − 22.8% to 36.4%); Low certainty ⊕⊕○○  Functional capacity improvement: No information  Strength 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				

	Telerehabilitation 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						
King et al. (28); 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: Very low certainty ⊕○○○  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 ⊕⊕○○  Functional capacity improvement: No information  Pulmonary function improvement: No information  Radiological response: No information  Adverse events: No information  Severe adverse events: No information			

Endurance training  Endurance training may improve HRQL and dyspnea. 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				
RECOVER trial. (29), 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%	NR	High risk of bias  Notes: Non-blinded study. Concealment of allocation probably inappropriate.	HRQL improvement: RR 1.48 (95% CI 0.92 to 2.37); RD 21.2% (95% CI -3.4% to 60.6%); Low certainty  ①①  Dyspnea improvement: RR 2.03 (95% CI 0.98 to 4.21); RD 24.4% (95% CI -0.4% to 75.9%); Low certainty  ①①  Functional capacity improvement: No information  Pulmonary function improvement: No information  Radiological response: No information  Adverse events: No information  Severe adverse events: No information		

## High dose steroids

High dose steroid	High dose steroids may not improve dyspnea and may not increase adverse events. 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					
COLDSTER trial (30), Dhooria et al.; Peer reviewed; 2022	Patients with P-ACC (dyspnea and/or lung radiological abnormalities after 21 to 49 days of acute COVID-19). 65 assigned to prednisone 40 mg a day descending progressively to 10 mg a day for 6 weeks and 65 assigned to prednisone 10 mg a day for 6 weeks	Mean age 57, male 68%, one comorbidity 73%	NR	High risk of bias  Notes: Non-blinded study which might have introduced bias.	HRQL improvement: No information  Dyspnea improvement: RR 1 (95% CI 0.87 to 1.15); RD 0% (95% CI -11.1% to 12.7%); Low certainty ⊕⊕⊖⊖  Functional 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 ⊕⊕⊖⊖			



	Home pulmonary rehabilitation 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					
Vallier et al; (31) 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 times a week for 4 weeks	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: Very low certainty ⊕○○○  Pulmonary function improvement: No information  Radiological response: No information  Adverse events: No information  Severe adverse events: No			
Respiratory training	Respiratory training Respiratory training may improve HRQL and dyspnea. 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							
ENO Breathe trial (32), 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 assigned to standard of care.	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 1.93 (95% CI 1.30 to 2.86); RD 24.1% (95% CI −7.8% to 48.1%); Low certainty ⊕⊕○○  Dyspnea improvement: RR 1.86 (95% CI			
McNarry et al. (33); 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.	1.38 to 2.49); RD 22.9% (95% CI 10.1% to 39.7%); Low certainty ⊕⊕⊖⊖  Functional capacity improvement: No information  Pulmonary function			
Srinivasan et al. (34); 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.	improvement: Very low certainty  Comparison  Radiological response: No information  Adverse events: No information  Severe adverse events: No			
Rodriguez- Blanco et al; (25) Peer reviewed; 2022	Patients with P-ACC (asthenia or fatigue after 40 days of acute COVID-19). 24 assigned to respiratory training (10 breathing and strength-based exercises) for 14	Mean age 40.7, male 22.91%	NR	High risk of bias  Notes: Non-blinded study. Concealment of allocation probably inappropriate.				



High dose steroid	days, and 24 assigned to standard of care.	spnea and may not inc	se steroids rease adverse events.	However, certainty of th	ne evidence was low.
Study; publication status	Patients and interventions analyzed	Further reso	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence
		į į	RCT		
Bazdyrev et al. (35); Preprint; 2022	Patients with P-ACC (dyspnea and/or lung radiological abnormalities after acute COVID-19). 29 assigned to treamid 50 mg a day for 28 days and 30 assigned to standard of care.	Mean age 55 ± 11, male 44.1%	NR	Low risk of bias	HRQL improvement: No information  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 ⊕⊕⊖⊖
		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

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			

Hyperbaric oxygen (HBO)
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			
		F	RCT					
Itskovich et al. (36); 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			
	Transcutaneous auricular vagus nerve stimulation (taVNS)  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					



Badran et al. (37); Preprint; 2022  tDCS may not Study; publication status				on (tDCS) ertainty of the evidence Risk of bias and study limitations	HRQL 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  Interventions effects vs standard of care (SOC) and GRADE certainty of
			ЭСТ		the evidence
			RCT		
Oliver-Mas et al. (27); 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	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





			⊕⊕○○
			Depression improvement: No 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 (adaptogens) ADAPT-232 may not improve olfactory symptoms. 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 analyzed of care (SOC) and **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 14% reviewed; 2022 fatigue after 30 information days of acute COVID-19). 49 Overall symptom assigned to improvement: No ADAPT-232 information (adaptogens) 60 mL a day for 14 Olfactory days and 50 symptoms assigned to improvement: standard of care. RR 0.89 (95% CI 0.79 to 1.01); RD -10.3% (95% CI -20.5% to 1.4%); Low certainty $\Theta\ThetaOO$ Gustatory symptoms improvement: No information Adverse events: No information Severe adverse events: No information

Olfactory 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							
Di Stadio et al. (38); 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%	Steroids 44%, vitamins 20.7%, alpha lipoic/nicetile 26.7%	High risk of bias  Notes: Non-blinded study which might have introduced bias.	improvement: No information  Olfactory symptoms improvement: Very low certainty ⊕○○○  Gustatory		
Pires et al. (39); 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.			
COVANOS trial (40), 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	disease 0%, asthma 12.6%, chronic heart disease 0%, cancer	NR	High risk of bias  Notes: Non-blinded study which might have introduced bias.			



Palmitoylethand Study; publication status	Patients and			eolin certainty of the evidence Risk of bias and study limitations	
		F	RCT		
Di Stadio et al. (38); Peer reviewed; 2022	Patients with P-ACC (olfactory and/or gustatory dysfunction after 180 days of acute COVID-19). 88 assigned to palmitoylethanolam ide + luteolin 700/70 mg a day and 38 assigned to standard of care.	Mean age 42.1, male 24.6%, hypertension	Steroids 32.5%,	Low risk of bias	HRQL improvement: No information  Overall symptom improvement: No information  Olfactory symptoms improvement: RR 1.11 (95% CI 0.68 to 1.81); RD 4.1% (95% CI -11.7% to 29.7%); Low certainty  ① Gustatory symptoms improvement: No information  Adverse events: No information  Severe adverse events: No information

	Uncertainty	Steroic	ds (nasal) 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		
RC 4-7-2020 trial (41), 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 ⊕○○○  Gustatory symptoms improvement: No information  Adverse events: No information  Severe adverse events: No information
	Uncertainty	Ste in potential benefits a	eroids 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		



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

	Ivabradine 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			
		I	RCT					
Jadhav et al. (43); 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

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.							
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				
	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	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 1.08 to 55.1%); Low certainty ⊕⊕○○  Adverse events: No information		

				Severe adverse events: No information
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**Table 8.** Description of included studies and interventions effects for P-ACC-related thromboembolic risk

	Anticoagulants (prophylactic dose) 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					
	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%	NR	High risk of bias  Notes: Non-blinded study which might have introduced bias to symptoms, VTE and adverse events outcomes.	Mortality: Very low certainty ⊕○○○ HRQL improvement: No information  VTE (symptomatic): Very low certainty ⊕○○○ Major bleeding: No information  Severe adverse events: No information			

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

Steroids may redu	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 (46); 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 ⊕○○○			

			<b>Major bleeding:</b> No information

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# Annex 1. Summary of findings tables

#### **Summary of findings Table A1.**

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

Intervention: Actovegin

Comparator: Standard of care (SOC)

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

Comparator: Standard of care (SOC)

<b>Outcome</b> Timeframe	Study results and measurements	Absolute effect estimates		Certainty of the evidence	Plain language summary
		SOC	ADAPT-232	(Quality of evidence)	riaiii ianguage summary
Fatigue improvement	Relative risk 1.02 (95% CI 0.84 to 1.24) Based on data from 99 participants in 1 study Follow-up 21 days		816 per 1000 more per 1000 wer to 192 more)	<b>Low</b> Due to very serious imprecision <sup>a</sup>	Adapt-232 may have little or no difference on fatigue improvement

a. **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

Comparator: Standard of care (SOC)

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

a. Decrease in 12 units of the MFI score.

### **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  Relative risk 6.07 (95% Cl 3.71 to 9.71) Based on data from 200 participants in 1 study Follow-up 25 days	<b>150</b> per 1000	<b>911</b> per 1000	Low  Due to serious risk of bias,  Due to serious imprecision <sup>a</sup>	Enzymes + probiotics may increase fatigue improvement	
	participants in 1 study	Difference: <b>761 more per 1000</b> (95% CI 407 more to 850 more)			

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: serious.** Low number of patients.

### **Summary of findings Table A5.**

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

Intervention: Phytochemicals

Outcome Stu	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language	
Timeframe	measurements	SOC	Phytochemicals	(Quality of evidence)	summary	
HRQL improvement	Relative risk: 1.33 (CI 95% 1.03 - 1.71) Based on data from 147	<b>543</b> per 1000	<b>722</b> per 1000	<b>Low</b> Due to serious risk of	Phytochemicals may increase HRQL	
	participants in 1 study Follow up 30 days	Difference: <b>179 more per 1000</b> (CI 95% 16 more - 386 more)	bias, Due to serious imprecision <sup>1</sup>	improvement		
Fatigue improvement	Relative risk: 1.24 (CI 95% 0.95 - 1.62) Based on data from 147	<b>539</b> per 1000	<b>668</b> per 1000	<b>Low</b> Due to serious risk of	Phytochemicals may increase fatigue	
	participants in 1 study Follow up 30 days	Difference: <b>129 more per 1000</b> (CI 95% 27 fewer - 334 more)		bias, Due to serious imprecision²	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;
- 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)

		Absolute e	ffect estimates	Certainty of the	
Outcome Timeframe		evidence (Quality of evidence)	Plain language summary		
Fatigue improvement	Relative risk 0.95 (95% CI 0.5 to 1.79) Based on data from 47	<b>458</b> per 1000	<b>435</b> per 1000	<b>Low</b> Due to very serious	Transcranial direct current stimulation (tDCS) may have little or no
	participants in 1 study Follow-up 25 days	Difference: 23 fewer per 1000 (95% CI 229 fewer to 362 more)	imprecision <sup>a</sup>	difference on fatigue improvement	
Adverse events	Based on data from 47 participants in 1 study Difference: 351		<b>173</b> per 1000	<b>Low</b> Due to very serious	Transcranial direct current stimulation (tDCS) may have little or no
		5 fewer per 1000 ewer to 360 more)	imprecision <sup>b</sup>	difference on adverse events	

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

b. **Imprecision: very serious.** 95% CI includes 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	measurements	soc	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	<b>980</b> per 1000	<b>980</b> per 1000	<b>Low</b> Due to very serious	ADAPT-232 may have little or no difference on
	participants in 1 study Follow-up 21 days	Difference: <b>0 fewer per 1000</b> (95% CI 59 fewer to 20 more)		imprecision <sup>a</sup>	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	Absolute eff	fect estimates	Certainty of the	
Timeframe	rame measurements	evidence (Quality of evidence)	Plain language summary		
Relative risk 1.48 (95% Cl 0.92 to 2.37) Based on data from 60	<b>441</b> per 1000	<b>980</b> per 1000	<b>Low</b> Due to serious risk of	Endurance training may increase HRQL	
	participants in 1 study Follow-up 21 days		Difference: <b>0 fewer per 1000</b> (95% CI 59 fewer to 20 more)	bias, Due to serious imprecision <sup>b</sup>	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	<b>236</b> per 1000	<b>980</b> per 1000	Low Due to serious risk of	Endurance training may increase dyspnea	
		Difference: <b>0 fewer per 1000</b> (95% CI 59 fewer to 20 more)		bias, Due to serious imprecision <sup>d</sup>	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)

<b>Outcome</b> Timeframe	Study results and measurements	Absolute effe Standard dose steroids	ct estimates  High dose steroids	Certainty of the evidence (Quality of evidence)	Plain language summary
Dyspnea improvement	Relative risk 1.0 (95% CI 0.87 to 1.15) Based on data from 130 participants in 1 study Follow-up 42 days	862 per 1000 Difference: 0 fe (95% CI 112 few		Low  Due to serious risk of bias,  Due to serious  imprecision <sup>a</sup>	High dose steroids may have little or no difference on dyspnea improvement
Radiological response	Relative risk 1.33 (95% CI 0.69 to 2.59) Based on data from 60 participants in 1 study Follow-up 21 days	185 per 1000 Difference: 61 r (95% CI 57 few		Very low  Due to serious risk of bias,  Due to very serious  imprecision <sup>b</sup>	We are uncertain whether high dose steroids increases or decreases radiological response
Adverse events	Relative risk 0.92 (95% CI 0.75 to 1.13) Based on data from 60 participants in 1 study Follow-up 21 days	769 per 1000 Difference: 62 f (95% CI 192 few		Low Due to serious risk of bias, Due to serious imprecision <sup>c</sup>	High dose steroids may have little or no difference on adverse events
Severe adverse events	Relative risk 3.0 (95% CI 0.32 to 28.09) Based on data from 60 participants in 1 study Follow-up 21 days	15 per 1000 Difference: 30 r (95% CI 10 fewer		Very low  Due to serious risk of bias,  Due to very serious  imprecisiond	We are uncertain whether high dose steroids 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 Comparator: Standard of care (SOC)

Outcome	Outcome Study results and	Absolute eff	ect estimates	Certainty of the	Blata I.	
Timeframe	measurements	soc	Respiratory training	Evidence (Quality of evidence)	Plain language summary	
HRQL improvement	Relative risk: 1.93 (CI 95% 1.3 - 2.86) Based on data from 203 participants in 2 studies Follow up 118 days	<b>259</b> per 1000	<b>500</b> per 1000	Low  Due to serious imprecision, Due to	Respiratory training may increase hrql improvement	
		Difference: <b>241 more per 1000</b> (Cl 95% 78 more - 482 more)		serious risk of bias <sup>1</sup>	morease man improvement	
Dyspnea improvement	Relative risk: 1.86 (CI 95% 1.38 - 2.49) Based on data from 271	<b>266</b> per 1000	<b>495</b> per 1000	<b>Low</b> Due to serious risk of bias,	Respiratory training may increase dyspnea	
	participants in 3 studies Follow up 83 days		<b>9 more per 1000</b> nore - 396 more)	Due to serious inconsistency <sup>2</sup>	improvement	
Pulmonary function improvement Relative risk: 1.17 (CI 95% 0.66 - 2.07) Based on data from 48 participants in 1 study Follow up 42 days	<b>459</b> per 1000	<b>537</b> per 1000	Very low Due to serious risk of bias,	We are uncertain whether respiratory training increases or decreases		
	participants in 1 study		more per 1000 ewer - 491 more)	Due to very serious imprecision <sup>3</sup>	pulmonary function 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.
- 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. **Incosistency: serious.** CI not overlapping.
- 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: very serious.** 95% CI includes important benefits and harms.

### **Summary of findings Table A11.**

Population: Patients with P-ACC-related dyspnea

Intervention: Treamid

Outcome	Outcome Study results and	Absolute eff	ect estimates	Certainty of the evidence	Plain language	
Timeframe	measurements	soc	Treamid	(Quality of evidence)	summary	
Functional capacity	Relative risk 1.1 (95% CI 0.64 to 1.9) Based on data from 59	<b>445</b> per 1000	<b>490</b> per 1000	<b>Low</b> Due to very serious	Treamid may have little or no difference on	
шрюченен	participants in 1 study Follow-up 28 days		more per 1000 wer to 401 more)	imprecision <sup>a</sup>	functional capacity improvement	
Dyspnea		<b>227</b> per 1000	<b>445</b> per 1000	Low	Treamid may increase	
improvement	Based on data from 59 participants in 1 study Follow-up 28 days	Difference: <b>218 more per 1000</b> (95% CI 23 fewer to 738 more)		Due to very serious imprecision <sup>b</sup>	dyspnea improvement	
Pulmonary function improvement	Relative risk 2.48 (95% CI 1.0 to 6.17) Based on data from 59	<b>167</b> per 1000	<b>414</b> per 1000	<b>Low</b> 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 <sup>c</sup>	improvement	
Relative risk 1.19 (95% CI 0.56 to 2.5) Based on data from 59 participants in 1 study Follow-up 28 days	<b>290</b> per 1000	<b>345</b> per 1000	Low	Treamid may increase		
	participants in 1 study	Difference: <b>55 more per 1000</b> (95% CI 128 fewer to 435 more)		Due to very serious imprecision <sup>d</sup>	adverse events	

- a. **Imprecision: very serious.** 95% CI includes important benefits and harms.
- b. **Imprecision: very serious.** 95% CI includes important benefits and harms.
- c. Imprecision: very serious. 95% CI includes important benefits and harms.
- d. **Imprecision: very serious.** 95% CI includes important benefits and harms.

### **Summary of findings Table A12.**

Population: Patients with P-ACC-related neurocognitive symptoms

Intervention: Actovegin

Comparator: Standard of care (SOC)

Outcome	Study results and	Absolute effect estimates		Certainty of the	Plain language
Timeframe	measurements	soc	Actovegin	Evidence (Quality of evidence)	summary
Cognitive improvement	, ,	<b>673</b> per 1000	<b>710</b> per 1000	Low Due to very serious risk of	Actovegin may improve
·	participants in 1 study		more per 1000 ore - 384 fewer)	bias <sup>1</sup>	cognition

3. **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

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

Outcome	Study results and	Absolute eff	fect estimates	Certainty of the	Plain language	
Timeframe	measurements	soc	НВО	(Quality of evidence)	summary	
HRQL improvement	Relative risk 1.3 (95% CI 0.84 to 2.0) Based on data from 73	<b>469</b> per 1000	<b>610</b> per 1000	<b>Low</b> Due to very serious	HBO may increase HRQF improvement	
	participants in 1 study Difference		I more per 1000 ver to 469 more)	imprecision <sup>a</sup>		
Cognitive improvement	Odds ratio 2.84 (95% Cl 1.09 to 7.37) Based on data from 73	<b>667</b> per 1000	<b>850</b> per 1000	Very low Due to extremely serious imprecision,	We are uncertain whether HBO increases or	
improvement	participants in 1 study		3 more per 1000 ore to 22 more)	Due to serious indirectness <sup>b</sup>	decreases cognitive improvement	
Depression (95% CI 2.72	Odds ratio 35.9 (95% CI 2.72 to 474.6)	<b>681</b> per 1000	<b>987</b> per 1000	Very low Due to extremely	We are uncertain whether HBO increases or	
improvement	Based on data from 73 participants in 1 study Follow-up 28 days		6 more per 1000 ore to 312 more)	serious imprecision, Due to serious indirectness <sup>c</sup>	decreases depression improvement	

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

Intervention: Transcranial direct current stimulation (tDCS)

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

a. 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

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

Outcome Study results and	Study results and	Absolute effect estimates		Certainty of the	Plain language summary
Timeframe	Timeframe measurements Pa	Palmitoylethanola mide + Luteolin	evidence (Quality of evidence)		
Olfactory symptoms	Relative risk 1.11 (95% CI 0.68 to 1.81) Based on data from 126 participants in 1 study Follow-up 90 days	<b>368</b> per 1000	<b>408</b> per 1000	<b>Low</b> Due to very serious	Palmitoylethanolamide + luteolin may have little or no
		Difference: <b>40 more per 1000</b> (95% CI 118 fewer to 298 more)		imprecision <sup>a</sup>	difference on olfactory symptoms improvement

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

### **Summary of findings Table A17.**

Population: Patients with P-ACC-related psychological distress

Intervention: Virtual reality informational video

<b>Outcome</b> Timeframe	Study results and measurements	Absolute effect estimates		Certainty of the	
		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 participants in 1 study Follow-up 90 days	<b>682</b> per 1000	<b>825</b> per 1000	<b>Low</b> Due to serious risk of bias,	Virtual reality informational video may
		Difference: <b>143 more per 1000</b> (95% CI 34 fewer to 368 more)		Due to serious imprecision <sup>a</sup>	increase depression improvement
Post-traumatic stress disorder improvement	Relative risk 1.18 (95% CI 0.98 to 1.42) Based on data from 89 participants in 1 study Follow-up 90 days	<b>773</b> per 1000	<b>912</b> per 1000	Low Due to serious risk of bias,	Virtual reality informational video may increase post-traumatic
		Difference: <b>139 more per 1000</b> (95% CI 15 fewer to 227 more)		Due to serious imprecision <sup>b</sup>	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	<b>523</b> per 1000	<b>779</b> per 1000	<b>Low</b> Due to serious risk of bias,	Virtual reality informational video may
		Difference: <b>256 more per 1000</b> (95% CI 42 more to 549 more)		Due to serious imprecision <sup>c</sup>	increase psychological distress improvement

- a. Risk of bias: serious. Imprecision: serious. Low number of patients.
- b. **Risk of bias: serious. Imprecision: serious.** Low number of patients.
- c. Risk of bias: serious. Imprecision: serious. Low number of patients.

## **Summary of findings Table A18.**

Population: Patients with PIMS-TS

Intervention: Steroids Comparator: IVIG

<b>Outcome</b> 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 <sup>1</sup>	Relative risk: 1.09 (Cl 95% 0.88 - 1.39) Based on data from 75 participants in 1 study Follow up 28	500 545 per 1000 per 1000 Difference: 45 more per 1000 (CI 95% 60 fewer - 195 more)	Low Due to serious risk of bias, Due to serious imprecision <sup>2</sup>	Steroids may decrease time to discharge
Respiratory support	Relative risk: 0.49 (CI 95% 0.27 - 0.89) Based on data from 75 participants in 1 study Follow up 28	553 271 per 1000 per 1000  Difference: 282 fewer per 1000 (CI 95% 404 fewer - 61 fewer)	Low Due to serious risk of bias, Due to serious imprecision <sup>3</sup>	Steroids may decrease respiratory support 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 Difference: 126 fewer per 1000 (CI 95% 257 fewer - 126 more)	Very low Due to serious risk of bias, Due to serious imprecision, Due to very serious imprecision <sup>4</sup>	We are uncertain whether steroids increases or decreases inotropic requirements
Left ventricular fraction deterioration	Relative risk: 0.57 (CI 95% 0.21 - 1.54) Based on data from 75 participants in 1 study Follow up 28	237 135 per 1000 per 1000 Difference: 102 fewer per 1000 (CI 95% 187 fewer - 128 more)	Very low  Due to serious risk of bias,  Due to serious  imprecision, Due to very  serious imprecision <sup>5</sup>	We are uncertain whether steroids increases or decreases LVEF deterioration
Arrhythmia	Relative risk: 2.05 (CI 95% 0.19 - 21.7) Based on data from 75 participants in 1 study Follow up 28	26 53 per 1000 per 1000 Difference: 27 more per 1000 (Cl 95% 21 fewer - 538 more)	Very low  Due to serious risk of bias,  Due to serious  imprecision, Due to very  serious imprecision <sup>6</sup>	We are uncertain whether steroids increases or decreases Arrhythmias
Venous thromboembolic events	Relative risk: 0.34 (CI 95% 0.01 - 8.14) Based on data from 75 participants in 1 study Follow up 28	39 13 per 1000 per 1000 Difference: 26 fewer per 1000 (CI 95% 39 fewer - 278 more)	Very low  Due to serious risk of bias,  Due to serious  imprecision, Due to very  serious imprecision <sup>7</sup>	We are uncertain whether steroids increases or decreases VTE

- 1. Proportion of patients discharged on day 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: 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;