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Exercise effect on symptom severity, morbidity and mortality in viral infections: a systematic review and meta-analysis.

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20 **Short running head:** Exercise effects during virus infections.

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25 Abstract

26 There is a knowledge gap regarding the consequences of exercise during acute 27 infections in humans and contradictory findings in animal studies, compromising public 28 health advice on the potential benefits of physical activity for immunity. Here, we carried out a meta-analysis of studies of the effects of moderate exercise (ME) and 29 30 exercise until fatigue (EF) on symptom severity, morbidity and mortality during viral infection in animal models. The systematic review on PubMed, Scopus, Embase, Web 31 32 of Science, Cochrane and EBSCOhost (CINAHL and SPORT Discus) identified 8 33 controlled studies, with 15 subgroups within them. The studies exposed the animals (mice [7 studies] and monkeys [1 study]) to exercise immediately before or after viral 34 inoculation (HSV-1, H1N1 influenza and B.K. virus) with follow-up for 21 days. ME 35 36 significantly reduced morbidity (OR 0.43 [0.19; 0.98], P = 0.04) with no change for symptom severity (SMD -3.37 [-9.01; 2.28], P = 0.24) or mortality (OR 0.48 [0.08; 3.03], 37 P = 0.43). In contrast, EF gave a trend towards increased symptom severity (SMD) 38 39 0.96 [-0.06; 1.98], P = 0.07) and mortality (OR 1.47 [0.96;2.28], P = 0.08) with no change in morbidity (OR 1.22 [0.60;2.5], P = 0.58). We conclude that in animals 40 41 moderate exercise during infection is advantageous, whilst exercise until fatigue should be avoided. Further research is required to determine if moderate exercise may 42 43 also be beneficial in humans during infection.

44

45 **Keywords:** Immunity, Virus Infection, Physical Activity, Exercise, Survival.

46

48 Introduction

49 There is a considerable literature supporting the benefits of exercise for human health in general (10) and for the immune system in particular (7, 16, 21). The positive 50 51 effects of exercise include an enhanced response to vaccination (39), improved immune surveillance mediated by redistribution of immune cells to tissues following 52 exercise (30, 52), increased apoptosis of senescent T cells potentially rejuvenating the 53 immune system (33, 51), and with maintained physical activity in to old age there is 54 evidence that the negative effects of age upon immune phenotype and immune 55 56 responses can be reduced (17, 46). Regards effect of exercise on occurrence, severity and duration of acute respiratory infections, a comprehensive meta-analysis showed 57 exercise reduced the severity of symptoms and the number of symptom days (22). 58 59 However, the benefits of exercise during an acute infection have received less attention and some of the data concerning the immune response to acute exercise, 60 such as temporary lymphopenia (45, 49, 50) and reduced salivary immunoglobulin A 61 levels (36), have been interpreted as indicating immune suppression (27). Whilst 62 alternative interpretations of the exercise immunology literature have been made (7), 63 recommendations for conservative exercise protocols, and even exercise restriction 64 at times of infection persist (20, 26). Crucially, as there is also good evidence that 65 exercising skeletal muscle is a major positive regulator of immune function (3, 16), it 66 67 is also possible that exercise could enhance the immune response against viruses 68 and bacteria and reduce the burden of latent viral infections (1, 22, 54). In the absence of infection, exercising skeletal muscle is the major producer of a range of cytokines 69 70 including IL-6 which in this context has anti-inflammatory actions (3), for example induction of IL-10, and IL-1RA production by macrophages (40). Muscle also produces 71

cytokines such as IL-7 and IL-15 which support the function of the thymus and
enhance the survival and function of immune cells (24, 43).

74 Opinions, reviews and practical guidelines discussing the risk of acute exercise 75 during infection in humans, including the recent COVID-19 pandemic, are based largely on indirect evidence (18, 20, 57, 59) and only two controlled trials in humans 76 77 have been reported that directly tested exercise effects on symptom severity during 78 infections (55, 56). In the first, rhinovirus 16 was inoculated into moderately fit young 79 adults who then underwent 40 minutes of aerobic exercise at 70% of their reserve 80 heart rate for the following 7 days (56). There was no difference in the scores for the 81 number of symptoms or the symptom severity measure between the exercise and 82 control groups during the 10 day follow-up (56). However in this study symptom 83 severity was assessed only by weighing the mucus (nasal secretion) instead of a more robust and/or sensitive method to confirm the duration of infection (56). A second study 84 85 by the same group assigned non-physically active young adults with a naturally 86 acquired upper respiratory tract infection to either 30 minutes aerobic exercise at 70% of target heart rate for five days, or control period with no exercise (55). The study 87 88 found no difference between mean symptom score and mean number of days with symptoms (55). Although these studies do not make a case for the benefits of exercise 89 90 during infection, in healthy young individuals, they also do not support the school of 91 thought that exercise is detrimental at times of infection.

In contrast to the lack of human studies, studies in animals, including mice and primates, have considered the influence of exercise on infection outcomes (11, 31). These studies have shown negative effects of exercise carried out before virus inoculation, with increased symptom severity, morbidity and mortality in mice and monkeys compared to controls (11, 31). These apparent differences in outcome

97 between humans and animals could be caused by the differences in volume and 98 intensity of exercise performed in animals, as some studies have used exercise with 99 exhaustive protocols (11). In fact, after a marathon race, humans undergo some 100 reduction in delayed-type hypersensitivity response, salivary IgA, T cell function, NK 101 cell activity, macrophage function, granulocyte oxidative burst together with increase 102 in neutrophil/lymphocyte ratio, cytokines and stress hormones that would lead to 103 transient immune dysfunction (37).

To address the discordance between studies and attempt to come to some consensus regarding exercise and the response to viral infections, we aimed to carry out a meta-analysis of the effects of moderate exercise (ME) and exercise until fatigue (EF) on symptom severity, morbidity and mortality during viral infection in animal studies.

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110 Methods

111 All details of the review protocol can be seen on PROSPERO 112 (CRD42021277401). We searched on PubMed (MEDLINE), adapted to Embase 113 Scopus, Web of Science, Cochrane, and EBSCOhost, April 19, 2021, for controlled 114 studies testing the effect of any type of exercise in animals during infection. They could 115 be purposely or naturally exposed to infection by any virus, and they needed to report 116 the impact on morbidity, symptom severity, and mortality in the exercise and control 117 groups. Duplicates were automatically removed using the Mendeley reference 118 manager system and the selection of studies was done by two independent reviewers 119 on the Rayyan-Systematic Reviews system (38).

120 Morbidity was assessed as the percentage of sick animals on the last day of 121 follow up which was at day 21; symptom severity was assessed by scales that would

122 consider different symptoms such as ruffled fur, inactivity, hunched back, and redness 123 around eyes, nose, or mouth; mortality was assessed by percentage of deaths during 124 the study period. Thus, morbidity and mortality were assessed as odds ratio (OR and 125 95% CI), according to the following equation OR = (n events in EXERCISE / n total in 126 EXERCISE) / (n events in CONTROL/ n total in CONTROL). The symptom severity 127 was calculated as the standardized mean difference (SMD) and 95% CI between 128 exercise and control means at a given day.

129 The three meta-analyses were performed using Comprehensive Meta-Analysis software, version 3.3.070. When there was statistical significance for heterogeneity. 130 131 randomized effect models were selected and when there was no significant heterogeneity, fixed effects were applied. The inconsistency between studies was 132 133 reported as a percentage (I²), based on difference between expected heterogeneity 134 (df) and true heterogeneity (Q-value). The subgroups within studies were clustered according to exercise protocols performed until fatigue (EF) or protocols of moderate 135 136 intensity (ME). Although, one of the studies described its exercise group as prolonged exercise, it was analyzed as EF (32). Q tests were applied to group comparisons, 137 138 considering 95% confidence. Egger's tests were performed to check the risk of publication bias in each meta-analysis. 139

140 **Results**

141 Supplementary figure 1 details the flowchart of selection of studies that led to 142 inclusion of 8 controlled studies, with 15 subgroups within them. The characteristics of 143 the studies are summarized on table 1.

144

145 **Please insert Table 1 here*

146

147 The meta-analysis of morbidity included 249 animals in the EXERCISE and 393 animals in the CONTROL. Since some studies had more than one group of 148 intervention and control (e.g.: males and females), each controlled intervention was 149 150 included as a separated study for analysis (4, 6, 31, 34). The overall hypothesis test showed the meta-analysis was not significant (OR 0.90 [0.46; 1.77], P = 0.77), with 151 significant heterogeneity and inconsistency across studies (P < 0.001; $I^2 = 67.94\%$), 152 and significant risk of bias (Egger test, P = 0.02). Figure 1a shows EF did not alter 153 154 morbidity compared to CONTROL (OR 1.22 [0.60;2.5], P = 0.58), while ME 155 significantly reduced morbidity in comparison to CONTROL (OR 0.43 [0.19; 0.98], P = 0.04). 156

The meta-analysis of symptom severity included 178 animals in the EXERCISE 157 158 and 182 animals in the CONTROL. The days that each study reported the severity 159 peak of symptoms were included for analysis, except for one study that reported the 160 first day of symptoms rather than its severity peak (6). The overall hypothesis test 161 showed there was no significant difference between the EXERCISE and CONTROL groups (SMD 0.05 [-1.04; 1.14], P = 0.93), with significant heterogeneity and 162 inconsistency across studies (P < 0.001; $I^2 = 94.66\%$), and non-significant risk of bias 163 (Egger test, P = 0.75). Figure 1b shows EF trended towards a higher severity of 164 165 symptoms compared to CONTROL (SMD 0.96 [-0.06; 1.98], P = 0.07), with no 166 difference between ME and CONTROL (SMD -3.37 [-9.01; 2.28], P = 0.24).

The meta-analysis of mortality included 371 animals in the EXERCISE and 370 animals in the CONTROL. The overall hypothesis test showed the meta-analysis was not significant (OR 1.07 [0.51; 2.21], P = 0.17), with significant heterogeneity and inconsistency across studies (P < 0.001; l^2 = 76.31%), and non-significant risk of bias (Egger test, P = 0.94). Figure 1c shows EF trended towards higher mortality than

172 CONTROL (OR 1.47 [0.96; 2.28], P = 0.08), while ME was no different from CONTROL
173 (OR 0.48 [0.08; 3.03], P = 0.43).

174

175 **Please insert figure 1 a, b and c here**

176

177 SYRCLES's risk of bias tool (25) showed low quality within the primary studies, 178 in which the large majority of them did not report whether group allocation was 179 adequately concealed, whether caregivers and outcome assessors were blinded; 180 whether the animals were selected at random for outcome assessment, and 181 incomplete outcome were not reported (Supplementary Table 1). At last, there was 182 low quality of evidence (score 2) for the severity of symptoms and Mortality meta-183 analyses, whilst there was very low quality of evidence (score 1) for the morbidity 184 meta-analysis assessed by the GRADE approach (23). In summary, the three meta-185 analyses lost two points due to its considerable inconsistency and low quality in their 186 primary studies (score between 4 and 5 on SYRCLES); only the morbidity meta-187 analysis lost one more point due to its significant risk of publication bias; and all three 188 led to precise results by direct evidence.

189

190 **Discussion**

Eichner (18) first questioned why someone should exercise during an infection if the workout intensity will be suboptimal to increase performance or skills. However, the loss of strength, muscle mass, and cardiorespiratory capacity are remarkable after a few days of de-training, such as during bed rest with or without an infection (2, 13, 42). An argument therefore could be made for maintaining exercise routines during an infection to avoid deconditioning. This may be even more important in older individuals

who are already at increased risk of sarcopenia and frailty (48, 58). Older adults also
have compromised immune systems which increase their risk of infections and of
succumbing to more severe symptoms, as demonstrated in the COVID-19 pandemic
(19).

201 Here we showed that moderate exercise could be a tool to boost immune 202 responses as we found a significant reduction in morbidity in animal studies of viral infection using such exercise programmes. Many physiological mechanisms could be 203 204 mediating such benefits. Acute exercise sessions repeated over several weeks 205 increase antibody production and cell-mediated responses during vaccination (39) and transiently enhance immune system features such as reducing the number of 206 207 senescent lymphocytes in the circulation (29, 44), increasing in blood counts for 208 neutrophils, lymphocytes, monocytes, and natural killer cells (37). Through the 209 increase in cortisol and adrenaline, and possibly also increased blood and lymph 210 circulation, exercise stimulates leukocyte circulation, release of cytokines, 211 chemokines, in turn facilitating antigen recognition, processing, and presentation, as 212 well as cell migration to lymph nodes and cell differentiation (39, 41).

213 In contrast, we found that exercise to fatigue trended towards an increase in the 214 severity of symptoms and mortality. The exact mechanism that explains the 215 differences between types of exercise are unknown. However, the exercise to fatigue 216 could affect different pathways that contribute to reduced immune responses. For 217 example, the generation of Damage Associated Molecular Patterns (DAMPs) from 218 damaged muscle which is then recognized by TLR receptors and could lead to 219 immune paresis (8, 28). Production of immune suppressive stress hormones such as 220 cortisol would also impact on immunity and reduction in energy availability with these 221 longer duration exercise protocols could compromise lymphocyte proliferation which

is highly energy dependent (15, 37, 47). It is worth noting that animals who are forced
to perform exercise would be more stressed than during voluntary exercise, which
would trigger a negative immune response (9, 14, 53).

225 Considering that the studies in the meta-analysis were performed in previously 226 healthy, young animals, the effect of exercise during infections in a high-risk population 227 such as older animals or humans remains to be determined. The only two studies 228 testing exercise effects during infection in humans were in healthy young adults but 229 showed that moderate exercise did not alter symptom severity (55, 56). As these 230 adults would have highly functional immune systems, the benefits of exercise may be 231 more marked in an older population with compromised immunity (16).

The main limitation of this study was the high inconsistency and low quality of evidence in each analysis suggesting that more studies will be necessary to identify the potential causes of heterogeneity between studies. Also, since most of the analyses were heterogeneous, we believe the difference between studies might be caused by a variety of factors such as: type and dose of pathogen; the mode, duration and intensity of exercise; and timing of virus administration in relation to exercise treatment.

Another potential limitation was the inclusion of two exercise interventions to fatigue in monkeys (31) in the meta-analysis assessing morbidity. However, we ran a separate analysis without these interventions and confirmed the same results as the analysis with all studies included (OR 1.056 [0.448; 2.489], P = 0.901) In conclusion, while exercise to fatigue trended to increase symptom severity and mortality during infections in animals, moderate exercise did not and significantly

²⁴⁵ reduced mortality. Future studies should test the effect of moderate intensity exercise

- ²⁴⁶ during infections in humans as a potential therapy to reduce symptom burden and
 ²⁴⁷ accelerate recovery.
- 248

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256

257 Statement of Ethics

258 An ethics statement is not applicable as this study is based exclusively on 259 published literature.

260

- 261 **Conflict of Interest Statement**
- 262 The authors have no conflicts of interest to declare.

263

264 **References**

1. Agha NH, Mehta SK, Rooney B, Laughlin MS, Markofski MM, Pierson DL,

266 Katsanis E, Crucian BE, Simpson RJ. Exercise as a countermeasure for

267 latent viral reactivation during long duration space flight. FASEB J 34: 2869–

268 2881, 2020. doi: 10.1096/fj.201902327R.

- 269 2. Alibegovic AC, Højbjerre L, Sonne MP, Van Hall G, Stallknecht B, Dela F,
- 270 **Vaag A**. Impact of 9 days of bed rest on hepatic and peripheral insulin action,

- insulin secretion, and whole-body lipolysis in healthy young male offspring of
- 272 patients with type 2 diabetes. *Diabetes* 58: 2749–2756, 2009. doi:
- 273 **10.2337/db09-0369**.
- 274 3. Bay ML, Pedersen BK. Muscle-Organ Crosstalk: Focus on
- Immunometabolism. *Front Physiol* 11: 1–8, 2020. doi:
- 276 10.3389/fphys.2020.567881.
- 277 4. Brown AS, Davis JM, Murphy EA, Carmichael MD, Carson JA, Ghaffar A,
- 278 **Mayer EP**. Susceptibility to HSV-1 infection and exercise stress in female
- 279 mice: Role of estrogen. *J Appl Physiol* 103: 1592–1597, 2007. doi:
- 280 10.1152/japplphysiol.00677.2007.
- 281 5. Brown AS, Davis JM, Murphy EA, Carmichael MD, Carson JA, Ghaffar A,
- 282 Mayer EP. Susceptibility to HSV-1 infection and exercise stress in female
- 283 mice: Role of estrogen. *J Appl Physiol* 103: 1592–1597, 2007. doi:
- 284 10.1152/japplphysiol.00677.2007.
- 285 6. Brown AS, Davis MM, Murphy EA, Carmichael MD, Ghaffar A, Mayer EP.
- 286 Gender differences in viral infection after repeated exercise stress. *Med Sci*
- 287 Sports Exerc 36: 1290–1295, 2004. doi:
- 288 10.1249/01.MSS.0000135798.72735.B3.
- 289 7. **Campbell JP**, **Turner JE**. Debunking the myth of exercise-induced immune
- 290 suppression: Redefining the impact of exercise on immunological health
- 291 across the lifespan. *Front Immunol* 9: 1–21, 2018. doi:
- 292 10.3389/fimmu.2018.00648.
- 293 8. Cavalcante PAM, Gregnani MF, Henrique JS, Ornellas FH, Araújo RC.
- 294 Aerobic but not Resistance Exercise Can Induce Inflammatory Pathways via
- Toll-Like 2 and 4: a Systematic Review. *Sport Med open* 3: 42, 2017. doi:

296 10.1186/s40798-017-0111-2.

- 297 9. Chen C, Nakagawa S, An Y, Ito K, Kitaichi Y, Kusumi I. The exercise-
- 298 glucocorticoid paradox: How exercise is beneficial to cognition, mood, and the
- brain while increasing glucocorticoid levels. *Front Neuroendocrinol* 44: 83–102,
- 300 2017. doi: 10.1016/J.YFRNE.2016.12.001.
- 301 10. Chodzko-Zajko WJ, Proctor DN, Fiatarone Singh MA, Minson CT, Nigg
- 302 CR, Salem GJ, Skinner JS. American College of Sports Medicine position
- 303 stand. Exercise and physical activity for older adults. *Med Sci Sports Exerc*
- 304 41: 1510–1530, 2009. doi: 10.1249/MSS.0b013e3181a0c95c.
- 305 11. Davis JM, Kohut ML, Colbert LH, Jackson DA, Ghaffar A, Mayer EP.
- 306 Exercise, alveolar macrophage function, and susceptibility to respiratory
- 307 infection. J Appl Physiol 83: 1461–1466, 1997. doi:
- 308 10.1152/jappl.1997.83.5.1461.
- 309 12. Davis JM, Murphy EA, Brown AS, Carmichael MD, Ghaffar A, Mayer EP.
- 310 Effects of moderate exercise and oat β -glucan on innate immune function and
- 311 susceptibility to respiratory infection. *Am J Physiol Regul Integr Comp Physiol*
- 312 286: 366–372, 2004. doi: 10.1152/ajpregu.00304.2003.
- 13. Demangel R, Treffel L, Py G, Brioche T, Pagano AF, Bareille M-P, Beck A,
- 314 **Pessemesse L, Candau R, Gharib C, Chopard A, Millet C**. Early structural
- 315 and functional signature of 3-day human skeletal muscle disuse using the dry
- immersion model. *J Physiol* 595: 4301–4315, 2017. doi: 10.1113/JP273895.
- 317 14. Dhabhar FS. Effects of stress on immune function: the good, the bad, and the
 318 beautiful. *Immunol Res* 58: 193–210, 2014. doi: 10.1007/s12026-014-8517-0.
- 319 15. Dorneles GP, da Silva IM, Santos MA, Elsner VR, Fonseca SG, Peres A,
- 320 **Romão PRT**. Immunoregulation induced by autologous serum collected after

321 acute exercise in obese men: a randomized cross-over trial. *Sci Rep* 10: 1–16,

322 2020. doi: 10.1038/s41598-020-78750-z.

- 323 16. Duggal NA, Niemiro G, Harridge SDR, Simpson RJ, Lord JM. Can physical
 324 activity ameliorate immunosenescence and thereby reduce age-related multi-
- 325 morbidity? Nat Rev Immunol 19: 563–572, 2019. doi: 10.1038/s41577-019-
- 326 **0177-9**.
- 17. Duggal NA, Pollock RD, Lazarus NR, Harridge S, Lord JM. Major features
- 328 of immunesenescence, including reduced thymic output, are ameliorated by
- high levels of physical activity in adulthood. *Aging Cell* 17, 2018. doi:
- 330 10.1111/acel.12750.
- 18. Eichner ER. Infection, Immunity, and Exercise. *Phys Sportsmed* 21: 125–135,
 1993. doi: 10.1080/00913847.1993.11710319.
- 333 19. Gerdes EOW, Vanichkachorn G, Verdoorn BP, Hanson GJ, Joshi AY,
- 334 Murad MH, Rizza SA, Hurt RT, Tchkonia T, Kirkland JL. Role of
- 335 senescence in the chronic health consequences of COVID-19. .
- 336 20. Gleeson M, Bishop N WN. Exercise immunology. 1st Editio. London: August
 337 14, 2013, 2013.
- 338 21. Gleeson M, Bishop NC, Stensel DJ, Lindley MR, Mastana SS, Nimmo MA.
- 339 The anti-inflammatory effects of exercise: mechanisms and implications for the
- 340 prevention and treatment of disease. *Nat Rev Immunol* 11: 607–615, 2011.
- doi: 10.1038/nri3041.
- 342 22. Grande AJ, Keogh J, Silva V, Scott AM, AJ G, Keogh J. Exercise versus no
- 343 exercise for the occurrence, severity, and duration of acute respiratory
- 344 infections (Review). Cochrane Database Syst Rev, 2020. doi:
- 345 10.1002/14651858.CD010596.pub3.www.cochranelibrary.com.

346	23.	Guyatt GH, Oxman AD, Vist GE, Kunz R, Falck-Ytter Y, Alonso-Coello P,
347		Schünemann HJ. GRADE: an emerging consensus on rating quality of
348		evidence and strength of recommendations. BMJ 336: 924–926, 2008. doi:
349		10.1136/bmj.39489.470347.AD.
350	24.	Haugen F, Norheim F, Lian H, Wensaas AJ, Dueland S, Berg O, Funderud
351		A, Skålhegg BS, Raastad T, Drevon CA. IL-7 is expressed and secreted by
352		human skeletal muscle cells. Am J Physiol - Cell Physiol 298, 2010. doi:
353		10.1152/ajpcell.00094.2009.
354	25.	Hooijmans CR, Rovers MM, De Vries RBM, Leenaars M, Ritskes-Hoitinga
355		M, Langendam MW. SYRCLE's risk of bias tool for animal studies. BMC Med
356		Res Methodol 14: 1–9, 2014. doi: 10.1186/1471-2288-14-43.
357	26.	Jaworski CA, Rygiel V. Acute Illness in the Athlete. Clin Sports Med 38: 577-
358		595, 2019. doi: 10.1016/j.csm.2019.05.001.
359	27.	Kakanis MW, Peake J, Brenu EW, Simmonds M, Gray B, Hooper SL,
360		Marshall-Gradisnik SM. The open window of susceptibility to infection after
361		acute exercise in healthy young male elite athletes. Exerc Immunol Rev 16:
362		119–137, 2010.
363	28.	Kawai T, Akira S. Toll-like Receptors and Their Crosstalk with Other Innate
364		Receptors in Infection and Immunity. Immunity 34: 637–650, 2011. doi:
365		10.1016/j.immuni.2011.05.006.
366	29.	Krüger K, Alack K, Ringseis R, Mink L, Pfeifer E, Schinle M, Gindler K,
367		Kimmelmann L, Walscheid R, Muders K, Frech T, Eder K, Mooren F-C.
368		Apoptosis of T-Cell Subsets after Acute High-Intensity Interval Exercise. Med
369		Sci Sports Exerc 48: 2021–2029, 2016. doi:
370		10.1249/MSS.000000000000979.

371	30.	Krüger K, Lechtermann A, Fobker M, Völker K, Mooren FC. Exercise-
372		induced redistribution of T lymphocytes is regulated by adrenergic
373		mechanisms. Brain Behav Immun 22: 324–338, 2008. doi:
374		10.1016/j.bbi.2007.08.008.
375	31.	Levinson SO, Milzer A, Lewin P. Effect of fatigue, chilling and mechanical
376		trauma on resistance to experimental poliomyelitis. Am J Epidemiol 42: 204–
377		213, 1945. doi: 10.1093/oxfordjournals.aje.a119037.
378	32.	Lowder T, Padgett DA, Woods JA. Moderate exercise protects mice from
379		death due to influenza virus. Brain Behav Immun 19: 377–380, 2005. doi:
380		10.1016/j.bbi.2005.04.002.
381	33.	Mooren FC, Krüger K. Apoptotic lymphocytes induce progenitor cell
382		mobilization after exercise. J Appl Physiol 119: 135–139, 2015. doi:
383		10.1152/japplphysiol.00287.2015.
384	34.	Murphy EA, Davis JM, Brown AS, Carmichael MD, Van Rooijen N, Ghaffar
385		A, Mayer EP. Role of lung macrophages on susceptibility to respiratory
386		infection following short-term moderate exercise training. Am J Physiol - Regul
387		Integr Comp Physiol 287: 1354–1358, 2004. doi: 10.1152/ajpregu.00274.2004.
388	35.	Murphy EA, Davis JM, Carmichael MD, Gangemi JD, Ghaffar A, Mayer EP.
389		Exercise stress increases susceptibility to influenza infection. Brain Behav
390		Immun 22: 1152–1155, 2008. doi: 10.1016/j.bbi.2008.06.004.
391	36.	Nieman DC, Henson DA, Fagoaga OR, Utter AC, Vinci M. Change in
392		salivary IgA following a competitive marathon race. Int J Sports Med 23: 69–
393		75, 2002. doi: 10.1055/s-2002-19375.

37. Nieman DC, Wentz LM. The compelling link between physical activity and the
body's defense system. *J Sport Heal Sci* 8: 201–217, 2019. doi:

- 396 10.1016/j.jshs.2018.09.009.
- 397 38. Ouzzani M, Hammady H, Fedorowicz Z, Elmagarmid A. Rayyan-a web and
 398 mobile app for systematic reviews. *Syst Rev* 5: 1–10, 2016. doi:
- 399 10.1186/s13643-016-0384-4.
- 400 39. Pascoe AR, Fiatarone Singh MA, Edwards KM. The effects of exercise on
- 401 vaccination responses: a review of chronic and acute exercise interventions in
- 402 humans. *Brain Behav Immun* 39: 33–41, 2014. doi: 10.1016/j.bbi.2013.10.003.
- 403 40. Pedersen BK. Anti-inflammatory effects of exercise: role in diabetes and
- 404 cardiovascular disease. *Eur J Clin Invest* 47: 600–611, 2017. doi:
- 405 **10.1111/eci.12781**.
- 406 41. Pedersen L, Idorn M, Olofsson GH, Lauenborg B, Nookaew I, Hansen RH,
- 407 Johannesen HH, Becker JC, Pedersen KS, Dethlefsen C, Nielsen J, Gehl
- 408 **J**, **Pedersen BK**, **Thor Straten P**, **Hojman P**. Voluntary Running Suppresses
- 409Tumor Growth through Epinephrine- and IL-6-Dependent NK Cell Mobilization
- 410 and Redistribution. *Cell Metab* 23: 554–562, 2016. doi:
- 411 10.1016/j.cmet.2016.01.011.
- 412 42. **Ried-Larsen M**, **Aarts HM**, **Joyner MJ**. Effects of strict prolonged bed rest on 413 cardiorespiratory fitness: systematic review and meta-analysis. *J Appl Physiol*
- 414 123: 790–799, 2017. doi: 10.1152/japplphysiol.00415.2017.
- 415 43. Rinnov A, Yfanti C, Nielsen S, Akerström TCA, Peijs L, Zankari A, Fischer
- 416 **CP**, **Pedersen BK**. Endurance training enhances skeletal muscle interleukin-
- 417 **15** in human male subjects. *Endocrine* **45**: **271–278**, **2014**. doi:
- 418 10.1007/s12020-013-9969-z.
- 419 44. Sardeli AV, Mori MA, Lord JM. Effect of exercise on acute senescent
- 420 *lymphocyte counts: a systematic review and meta-analysis.* 2021.

Sardeli AV, Mori MA, Lord JM. Effect of Exercise on Acute Senescent 421 45. 422 Lymphocyte Counts: A Systematic Review and Meta-Analysis. . 423 Sardeli AV, Tomeleri CM, Cyrino ES, Fernhall B, Cavaglieri CR, Chacon-46. 424 **Mikahil MPT**. Effect of resistance training on inflammatory markers of older adults: A meta-analysis. *Exp Gerontol* 111: 188–196, 2018. doi: 425 426 10.1016/j.exger.2018.07.021. 427 47. Sarin H, Gudelj I, Honkanen J, Ihalainen JK, Vuorela A, Lee JH, Jin Z, Terwilliger JD, Isola V, Ahtiainen JP, Häkkinen K, Jurić J, Lauc G, 428 429 Kristiansson K, Hulmi JJ, Perola M. Molecular pathways mediating 430 immunosuppression in response to prolonged intensive physical training, lowenergy availability, and intensive weight loss. Front Immunol 10, 2019. doi: 431 432 10.3389/fimmu.2019.00907. 433 48. Shamliyan T, Talley KMC, Ramakrishnan R, Kane RL. Association of frailty with survival: a systematic literature review. Ageing Res Rev 12: 719–736, 434 435 2013. doi: 10.1016/j.arr.2012.03.001. 49. Shek P, Sabiston B, Buguet A, Radomski M. Strenuous Exercise and 436 437 Immunological Changes. Int J Sports Med 16: 466-474, 1995. doi: 10.1055/s-2007-973039. 438 439 50. Shephard RJ. Adhesion molecules, catecholamines and leucocyte 440 redistribution during and following exercise. Sport Med 33: 261–284, 2003. doi: 10.2165/00007256-200333040-00002. 441 Shephard RJ. Aging, Persistent Viral Infections, and Immunosenescence: 442 51. Can Exercise "Make Space"? Yearb Sport Med 2011: 144–146, 2011. doi: 443 10.1016/j.yspm.2011.03.027. 444 445 Simpson RJ, Boßlau TK, Weyh C, Niemiro GM, Batatinha H, Smith KA, 52.

446		Krüger K. Exercise and adrenergic regulation of immunity. Brain Behav
447		Immun 97: 303–318, 2021. doi: 10.1016/j.bbi.2021.07.010.
448	53.	Simpson RJ, Campbell JP, Gleeson M, Krüger K, Nieman DC, Pyne DB,
449		Turner JE, Walsh NP. Can exercise affect immune function to increase
450		susceptibility to infection? Exerc Immunol Rev 26: 8–22, 2020.
451	54.	Simpson RJ, Hussain M, Baker F, Bigley AB, Peek MK, Stowe RP.
452		Cardiorespiratory fitness is associated with better control of latent herpesvirus
453		infections in a large ethnically diverse community sample: Evidence from the
454		Texas City Stress and Health Study. Brain Behav Immun 66: e35, 2017. doi:
455		10.1016/J.BBI.2017.07.128.
456	55.	Weidner T, Schurr T. Effect of exercise on upper respiratory tract infection in
457		sedentary subjects. Br J Sports Med 37: 304–306, 2003. doi:
458		10.1136/bjsm.37.4.304.
459	56.	Weidner TG, Cranston T, Schurr T, Kaminsky LA. The effect of exercise
460		training on the severity and duration of a viral upper respiratory illness. Med
461		Sci Sports Exerc 30: 1578–1583, 1998. doi: 10.1097/00005768-199811000-
462		00004.
463	57.	Wilson MG, Hull JH, Rogers J, Pollock N, Dodd M, Haines J, Harris S,
464		Loosemore M, Malhotra A, Pieles G, Shah A, Taylor L, Vyas A, Haddad
465		FS, Sharma S. Cardiorespiratory considerations for return-to-play in elite
466		athletes after COVID-19 infection: A practical guide for sport and exercise
467		medicine physicians. Br J Sports Med 54: 1157–1161, 2020. doi:
468		10.1136/bjsports-2020-102710.
469	58.	Xu J, Wan CS, Ktoris K, Reijnierse EM, Maier AB. Sarcopenia Is Associated
470		with Mortality in Adults: A Systematic Review and Meta-Analysis.

- 471 *Gerontology*: 1–16, 2021.
- 472 59. **Zhu W**. Should, and how can, exercise be done during a coronavirus
- 473 outbreak? An interview with Dr. Jeffrey A. Woods. J. Sport Heal. Sci. 9: 105-
- 474 107, 2020.
- 475

Figures and Tables

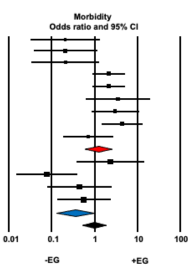
Table 1. Characteristics of the studies included.

First Author, Year	Species	Age	sex		Exercise time- point	Intensity category/ exactly	Туре	Volume/Duration	Morbidity	Mortality	Symptoms Severity
Levinson, 1945 (31)	Monkeys (Macaca mulatta)	NR	M/F	BKV (intracerebrally)	Post inoculation	EF/ Fatigue	Swimming	2-3 hours/ 4 d	Yes (days 11-14)	-	Yes *
Davis,1997	Mice 4		М	HSV-1	Before	EF/ Fatigue	Running (treadmill)	2.5–3.5 hours/ 3 d	Yes (day	Yes (day	
(11)	MICE	wk	IVI	(Intranasal)	inoculation	ME/ NR	Running (treadmill)	30 minutes/ 3 d	21);	21)	-
Brown, 2004 (6)	Mice	~60 d	M/F	HSV-1 (Intranasal)	Before inoculation	EF/ 70– 80% VO2 max.	Running (treadmill)	135 ± 5 min/ 3 d	Yes (day 21)	Yes (day 21)	Yes (1° day of symptom)
Davis, 2004 (12)	Mice	4 wk	М	HSV-1 (Intranasal)	Post inoculation	ME/ 68- 78% VO2 max.	Running (treadmill)	1 hour/ 6 d	Yes (day 21);	Yes (day 21)	
Murphy, 2004 (34)	Mice	4 wk	М	HSV-1 (Intranasal)	Post inoculation	ME/ 75- 90% VO2 max.	Running (treadmill)	1 hour/ 6 d	Yes (day 21);	Yes (day 21)	Yes (day 7)
Lowder, 2005 (32)	Mice	20- 24 wk	Μ	H3N2 (Intranasal)	Post inoculation	EF/ 65- 70% VO2 max.	Running (treadmill)	2.5 hours/ 3 d;	Yes (day 21);	Yes** (day 21)	

						ME/ 65- 70% VO2 max.	Running (treadmill)	30 min/ 3 d			
Brown, 2007 (5)	Mice	7 wk	F	HSV-1 (Intranasal)	Before inoculation	EF/ 70– 80% VO2 max.	Running (treadmill)	20 min/ 3 d,	Yes (day 21);	Yes (day 21)	Yes (days 12, 16-21)
Murphy, 2008 (35)	Mice	4 wk	М	H1N1 (Intranasal)	Post inoculation	EF/ 70– 80% VO2 max.	Running (treadmill)	20 min/ 3 d	Yes (day 21);	Yes (day 21)	Yes (day 7)

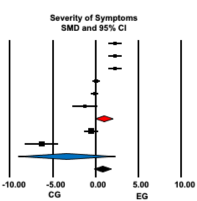
Legend: BKV: BK virus; d: days; EF: Exercise-fatigue; ME: moderate exercise; F: Female; H1N1: Influenza A virus subtype H1N1; HSV-1: herpes simplex virus 1; M: Male; Min: Minutes; NR: Not report; $\dot{V}O_{2 max}$ refers to the maximum amount of oxygen you can utilize during exercise; wk: weeks;* Assessed by incidence of paralysis and degree of involvement (not included in the meta-analysis); **The animals were followed up for 30 days, but the 21th day was meta-analysed in order to maintain consistency between studies.

First author,	OR	LL	UL	p-value	Events	/ Total	Relative	
year					EG	CG	weight	
Brown, 2007a	0.206	0.032	1.339	0.098	10 / 15	16 / 18	8.23	
Brown, 2007b	0.206	0.038	1.125	0.068	13/20	19/21	9.12	
Brown, 2007c	0.206	0.033	1.274	0.089	14 / 21	15/17	8.47	
Brown, 2004a	2.103	0.890	4.967	0.090	29/44	21/44	14.64	
Brown, 2004b	2.103	0.881	5.017	0.094	28/43	21/43	14.57	
Murphy, 2008	3.500	0.630	19.441	0.152	19/21	15/21	9.02	
Davis, 1997	3.000	0.837	10.753	0.092	11/22	6/22	11.69	
Levinson, 1945a	4.333	1.439	13.053	0.009	32 / 40	12/25	12.90	
Levinson, 1945b	0.706	0.187	2.659	0.607	32 / 40	22/26	11.36	
EF Overall (R)	1.222	0.595	2.510	0.585	188 / 266	147 / 237	100	
Murphy, 2004a	2.348	0.378	14.586	0.360	32/34	36/41	19.88	
Murphy, 2004b	0.076	0.015	0.391	0.002	24 / 41	33/35	24.58	
Davis, 1997	0.448	0.081	2.482	0.358	2/17	6/22	22.64	
Davis, 2004	0.562	0.136	2.326	0.427	3/21	12/58	32.89	
EM Overall (F)	0.434	0.192	0.979	0.044	61 / 113	87 / 156	100	
Summarized effect (R)	0.971	0.516	1.826	0.926	249 / 379	234 / 393	100	



Tau²= 0.99; Q= 37,43; df= 12; p< 0.001; l²= 67.94%; test for overall effect z: -0.30 (p= 0.77).

First author,	OR	LL	UL	p-value	Samp	Relative	
year					EG	CG	weight
Brown, 2007a	2.250	1.477	3.023	0.000	21	21	16.82
Brown, 2007b	2.250	1.477	3.023	0.000	21	21	16.82
Brown, 2007c	2.250	1.477	3.023	0.000	21	21	16.82
Brown, 2004a	0.067	-0.351	0.485	0.754	44	44	18.04
Brown, 2004b	-0.095	-0.518	0.328	0.661	43	43	18.03
Murphy, 2008	-1.262	-2.723	0.199	0.091	3	7	13.49
EF Overall (R)	0.960	-0.064	1.983	0.066	153	157	100
Murphy, 2004a	-0.555	-1.355	0.245	0.174	12	13	51.19
Murphy, 2004b	-6.318	-8.237	-4.399	0.000	13	12	48.81
EM Overall (R)	-3.368	-9.014	2.278	0.242	25	25	100
Summarized effect (R)	0.822	-0.185	1.829	0.110	178	182	100



Tau²= 2.21; Q=131.20; df= 7; p<0.001; l²= 94.66%; test for overall effect z:0.09 (p= 0.93)

First author,	OR	OR LL UL p-v		p-value	o-value Events / Total				
year					EG	CG	weight		
Brown, 2007a	3.696	0.744	18.355	0.110	7/21	2/21	7.85		
Brown, 2007b	2.442	0.705	8.452	0.159	12/21	8/21	12.41		
Brown, 2007c	1.542	0.452	5.260	0.489	13/21	11/21	12.67		
Brown, 2004a	1.000	0.320	3.126	1.000	7/44	7/44	14.39		
Brown, 2004b	1.000	0.390	2.564	1.000	12/43	12/43	19.73		
Murphy, 2008	2.919	0.651	13.082	0.162	18/21	14/21	8.86		
Davis, 1997	3.648	0.880	15.118	0.074	9/22	4/22	9.77		
Lowder, 2005	0.568	0.181	1.782	0.332	8 / 26	11/26	14.32		
EF Overall (R)	1.505	0.942	2.404	0.087	86 / 219	69 / 219	100		
Murphy, 2004a	1.463	0.530	4.035	0.462	32/41	30/41	21.34		
Murphy, 2004b	0.045	0.013	0.154	0.000	12/42	37/41	20.75		
Davis, 1997	0.519	0.081	3.310	0.488	2/22	4/22	18.55		
Davis, 2004	0.102	0.017	0.614	0.013	2/21	10/21	18.77		
Lowder, 2005	6.039	1.702	21.426	0.005	21/26	11/26	20.60		
EM Overall (R)	0.477	0.075	3.030	0.433	70/152	91 / 151	100		
Summarized effect (R)	1.404	0.892	2.211	0.143	156 / 371	159 / 370	100		

Tau²= 1.35; Q= 50.66; df= 12; p< 0.001; l²= 76.31%; test for overall effect z: 0.17 (p= 0.87).

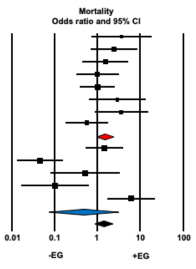
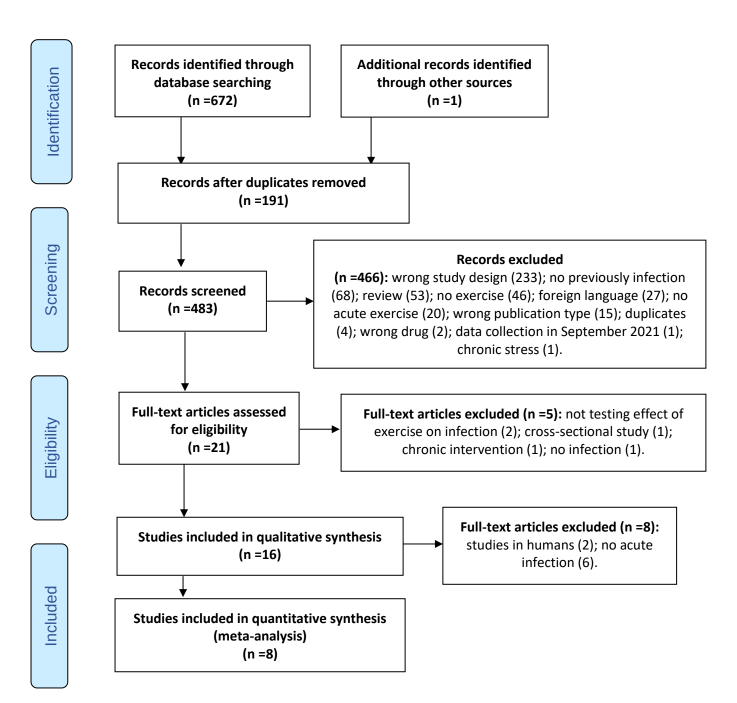


Figure 1. Forest plots of the effect of acute exercise on symptom severity (a), morbidity (b) and mortality (c) during acute virus infections. CG: control group; CI: confidence interval; df: degrees of freedom; EG: exercise group; F: fixed effect; I²: percentage of inconsistency between studies; LL: Lower limit; OR: Odds ratio; PBS: PBS liposomes; Q: true heterogeneity; R: random effect; SMD: standardized mean difference; UL: Upper limit; Brown, 2007a: intact (Sham) group; Brown, 2007b: ovariectomized group; Brown, 2007c: ovariectomized and estrogen-supplemented group; Brown, 2004a: female group; Brown, 2004b: male group; Levinson, 1945a: Cage control group; Levinson, 1945b: Water control group; Murphy, 2004a: clodronate encapsulated liposomes intranasally administered group; Murphy, 2004b: PBS liposomes intranasally administered group.



Supplementary Figure 1. The flowchart of selection of studies. Of note the final analysis did not include the 2 human studies and the focus was on the 8 animal studies.

Supplementary Table 1. SYRCLE Risk of Bias in the studies included.

First author,

year	1	2	3	4	5	6	7	8	9	10	Total
Levinson, 1945	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Davis,1997	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Brown, 2004	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Davis, 2004	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Murphy, 2004	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Lowder, 2005	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Brown, 2007	Yes	Yes	NR	Yes	No	NR	NR	NR	Yes	No	4
Murphy, 2008	Yes	Yes	NR	Yes	No	NR	NR	Yes	Yes	No	5

Legend: 1: allocation sequence adequately generated and applied; 2: similar groups at baseline or adjusted for confounders in the analysis; 3: group allocation adequately concealed; 4: animals randomly housed during the experiment; 5: caregivers blinded; 6: animals selected at random for outcome assessment; 7:outcome assessor blinded; 8: incomplete outcome data adequately addressed; 9: Reports of the study free of selective outcome reporting; 10: apparently free of other risk of bias; NR: Not Reported.