**INTERACTION OF FACTORS DETERMINING CRITICAL POWER**

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**Running head:** Interaction of factors determining critical power

**ABSTRACT**

The physiological determinants of high-intensity exercise tolerance are important for both elite human performance and morbidity, mortality and disease in clinical settings. The asymptote of the hyperbolic relation between external power and time to task failure, critical power, represents the threshold intensity above which systemic and intramuscular metabolic homeostasis can no longer be maintained. After ~60 years of research into the phenomenon of critical power a clear understanding of its physiological determinants has emerged. The purpose of the present review is to critically examine this contemporary evidence in order to explain the physiological underpinnings of critical power. Evidence demonstrating that alterations in convective and diffusive oxygen delivery can impact upon critical power are first addressed. Subsequently, evidence is considered which shows that rates of muscle oxygen utilization, inferred via the kinetics of pulmonary oxygen consumption, can influence critical power. The data reveals a clear picture that alterations in the rates of flux along every step of the oxygen transport and utilization pathways influences critical power. It is also clear that critical power is influenced by motor unit recruitment patterns. On this basis it is proposed that convective and diffusive oxygen delivery act in concert with muscle oxygen utilization rates to determine the intracellular metabolic milieu and state of fatigue within the myocytes. This interacts with exercising muscle mass and motor unit recruitment patterns to ultimately determine critical power.

**KEY POINTS**

* Critical power represents the threshold intensity above which steady-state metabolism is no longer attainable, and within the last ~15 years, experimental data has emerged which illuminates its underpinning physiological determinants
* Here, we summarize this experimental data to demonstrate that critical power is a parameter of aerobic function that is affected by alterations in the capacities of each step in the oxygen transport and utilization pathways
* Convective/diffusive oxygen delivery and intracellular oxygen utilization rates interact with muscle fibre composition and motor unit recruitment profiles to determine the upper limit for steady-state exercise

1. **INTRODUCTION**

The determinants of exercise tolerance are of clear interest because of the strong relationships between exercise capacity and athletic performance [1,2], health in the general population, and clinical outcomes in disease populations [3,4]. Exercise intensity is, of course, a key factor that determines the tolerability of a given task. Moreover, for individuals or groups of individuals, partitioning the exercise intensity spectrum into domains where the physiological responses to a given task share common qualitative characteristics is an effective approach that can yield insight into the physiological determinants of exercise tolerance. Accordingly, the mechanisms of fatigue and determinants of exercise intolerance are not ubiquitous across the spectrum of exercise intensities [5]. However, above a particular, individual-specific power output, the consistent feature of exercise intolerance (and hence, impending task failure), is the inability for pulmonary oxygen uptake (O2) and [lactate] (L-) to attain a steady-state [6–9]. Thus, for each individual, there exists a range of intensities for which a steady-state in pulmonary O2 is attainable, and a range for which it is not [6,9–12], with the duration of sustainable exercise in the latter being significantly limited compared to the former. The threshold intensity that separates these two ranges of system behaviour, and its position relative to other landmarks of aerobic function (i.e. maximal O2 [O2max] and the lactate threshold [LT]), is therefore a fundamental determinant of the ability to sustain exercise [6,13–15].

This threshold intensity can be determined by undertaking 3-5 high-intensity constant-power output cycle ergometer tests to the point of task failure on separate days. The tests should be selected to last no less than 2 and no more than 15 minutes in duration [16–19], with the precise time to task failure and power output at which each test is conducted recorded. These durations are recommended for valid determination of this intensity, since it is essential that O2max is attained at the end of trial in order to meet the requirement for all prediction trials to be performed within the severe-intensity domain. When time to task failure is plotted against power output, the relationship is curvilinear, with the ability to sustain exercise falling away more rapidly at higher power outputs (Figure 1). This power-time relationship is well-described by a hyperbolic function [20], with an asymptote known as critical power (CP) and the curvature constant termed *W*' (i.e., W prime). This relationship is described by the following equation:

Where T is the tolerable duration and P the power output of a given exercise task [6,20,21]. When intensity is measured in units of speed, the asymptote is termed critical speed (CS) and the curvature constant *D*’ (i.e., with units of distance). This power-time relationship appears to be a universal feature of high-intensity exercise tolerance, being apparent in every species [22–26] and mode of exercise (with appropriate units of force, torque or velocity, [15,27–30]) in which it has been studied. This relationship can also be converted to its linear equivalents, either with work plotted against time:

Where W is work, CP the slope and *W*' the intercept of the equation, or with power plotted against the inverse of time:

or

Where CP is the intercept and W' is the slope of the equation.

Since the seminal work by Professor David Poole and colleagues in the late 1980’s, it has been repeatedly demonstrated that CP reflects the upper limit at which a metabolic steady state can be sustained. The basis for this has been the ubiquity of steady state behaviour of metabolic variables associated with aerobic function below, but not above CP. For example, O2 rises to O2max during exercise above, but not at or below, CP [6], accompanied by similarly inexorable trajectories of blood [lactate], [HCO3-] and pH [6,31]. Such findings were subsequently confirmed in different populations, including the elderly [32], chronic heart failure [33] and chronic obstructive pulmonary disease patients [34,35], and healthy children [36]. More recently, non-invasive (31P-magnetic resonance spectroscopy, near infrared spectroscopy) and invasive (i.e. muscle biopsy) studies have demonstrated the achievement of a steady state in the exercising muscle below, but not above CP, in muscle O2, [phosphocreatine] ([PCr]), [inorganic phosphate] [Pi], pH and muscle [lactate] [15,31,37]; for review see [8,11,12]. Critical speed (CS, an analogue of CP) has also been demonstrated to be a critical threshold for motor unit recruitment patterns, with Copp et al. demonstrating that exercise above CS was accompanied by disproportionate increases in blood flow to type IIb/d/x fibres in the rat hindlimb muscle [25].

Despite CP, and its analogues of external expression (i.e. critical speed, torque, force, etc.) being widely recognised as reflecting the threshold intensity above which a metabolic steady state cannot be sustained, its physiological antecedents have previously been obscure. Tables 1 & 2 detail interventional and observational approaches to understanding CP. Prior to the year 2010, intervention studies on CP were scant, and primarily confined to the effect of exercise training alongside additional measures of O2max and the gas exchange threshold/LT only, although one of the earliest studies on CP did show an independent effect of O2 availability on CP (albeit in just 2 participants; [21]). Nevertheless, such findings supported the notion of CP as being a parameter of aerobic function [20]. In contrast, since 2010 multiple experimental approaches have revealed those factors which, directly or indirectly, determine CP. The purpose of the present review is therefore to examine the physiological and biochemical underpinnings of this fundamental parameter of exercise tolerance. Particular attention will be paid to evidence generated over the last 10 - 12 years demonstrating that CP is a key parameter of aerobic function that can be affected by any step in the O2 transport and utilization pathway.

[INSERT TABLE 1 HERE]

[INSERT TABLE 2 HERE]

1. **INTERACTION OF FACTORS DETERMINING CRITICAL POWER**

That CP represents the threshold intensity above which exercise cannot be sustained in a steady state indicates that it is a parameter of aerobic function. Consequently, it follows that CP may be affected by any step in the O2 transport and utilization cascade, from atmospheric air down to the muscle mitochondria themselves. Specifically, these steps include: 1) transport of atmospheric O2 into the blood via pulmonary diffusion, 2) bulk transport of O2 to the muscle via convection (i.e., convective O2 delivery), 3) diffusion of O2 from capillary to muscle mitochondria (i.e., diffusive O2 delivery), and 4) the utilization of O2 by the muscle mitochondria (Figure 2). Whilst the respiratory system may constrain CP in chronic respiratory disease conditions such as COPD [34,35,38–42], in most young, healthy individuals, the respiratory system appears to be well-adapted to ensure a highly efficient and appropriate homeostatic response to high-intensity exercise [43]. Hence, the remainder of this review will focus on the impact of convective and diffusive O2 delivery and mitochondrial O2 utilization on CP, downstream of the respiratory system.

*2.1 Convective oxygen delivery*

Convective O2 delivery refers to that achieved via bulk movement of O2 within the circulation to the exercising muscles. Convective O2 delivery (O2, L.min-1) can thus be defined mathematically as the product of cardiac output (CO, L.min-1) and arterial O2 content (CaO2, mL O2 100 mL-1):

Where CaO2 is defined as:

Where 1.34 is Hüfner's constant describing the maximum O2 carrying capacity per gram of haemoglobin (ml O2 g−1 Hb), [Hb] is haemoglobin concentration (g dl-1), SaO2 is the arterial saturation of Hb, 0.03 is the solubility coefficient of O2 at body temperature (ml O2 100 ml−1 plasma kPa−1), and PaO2 is the arterial partial pressure of O2 (mmHg). This provides a measure of whole-body convective O2 delivery. However, the O2 flux to each portion of the exercising muscles is not uniform but varies according to regional metabolic demands, vascular control and fibre type [44–46].

A convenient means by which to experimentally alter CaO2, and hence, convective O2 delivery, is by varying the fraction of inspired O2 (FiO2). Although hypoxia-induced vasodilatation [47] and hyperoxia-induced vasoconstriction [48] often influence blood flow thereby helping to normalize muscle O2 delivery during exercise, many studies that have quantified skeletal muscle O2 delivery under these conditions have demonstrated that hyperoxia can enhance and hypoxia can impair skeletal muscle O2 delivery during exercise, respectively [49–54], thereby impacting upon intramyocyte *P*O2 (*P*O2im)[55]. Indeed, the early work of Moritani et al. [21] showed that, in a limited sample of two participants, inspiration of a hypoxic gas mixture (FiO2 0.09) resulted in a reduced CP compared to normoxia (i.e. FiO2 0.21; hypoxia: 106 ± 6 W, versus normoxia: 214 ± 4 W). Under conditions of more moderate hypoxia (FiO2 0.15) and in a larger sample of 11 subjects, Dekerle et al. [56] found that CP was reduced by 30 W in hypoxia compared to normoxia, consequent to a reduction in arterial O2 saturation of 12%. Notably, in this latter study, the percentage decrement in CP between hypoxia and normoxia was correlated with O2max in normoxia, suggesting that those with the greatest O2max values were better able to offset the reductions in convective O2 delivery brought about via hypoxia. It is not known if such a protective effect remains in highly trained athletes where pulmonary limitations to high-intensity exercise are more likely [57,58], causing reductions in arterial saturation and O2max even at modest simulated altitudes [59]. Similarly, however, Simpson et al. [60] reported a reduction in CP of 43 W using an FiO2 of 0.13, a finding which was consistent when CP was determined either via the conventional constant-load prediction trial method or via a 3-minute all-out test. Moreover, Valli et al. [61] demonstrated that at an altitude of 5050 m (equivalent FiO2 ~0.11) CP was reduced by 42 W. In all of these studies, SaO2 was reduced either at rest or during exercise in hypoxia, providing indirect evidence that hypoxia impaired convective O2 delivery which contributed to the reduced CP in each study. These findings were subsequently extended to arm cycle ergometry by La Monica et al. [62], who demonstrated that arm CP was reduced by 5 W in moderate (FiO2 0.14) normobaric hypoxia (~6% of normoxic CP).Whilst the magnitude of the effect of hypoxia on CP in these studies varied with the fitness of the participants (see for example Dekerle et al. [56]), Townsend et al. [63] demonstrated a progressive reduction in CP with decreasing FiO2. Hence, the extant literature is unanimously consistent with the notion that reductions in FiO2 (and by extension, convective O2 delivery) reduce CP.

The consistency of the effects of hyperoxia on CP are similar to those of hypoxia. This was first demonstrated by Vanhatalo et al. [37], who assessed the impact of an FiO2 of 0.7 on CP utilizing a single leg knee-extension exercise model. These authors showed that CP was increased in hyperoxia compared to normoxia, with a concomitant increase in muscle oxygenation (as determined via near-infrared spectroscopy, NIRS). The increase in CP was accompanied by a slower rate of change in muscle [PCr], [ADP], [Pi], and pH. Subsequently, these findings for small muscle mass exercise were confirmed for large muscle mass exercise by Goulding et al. [64,65]. Specifically, a hyperoxic inspirate (FiO2 of 0.5) resulted in increases in end-tidal *P*O2 (and, therefore, alveolar *P*O2) and muscle oxygenation determined via NIRS both at rest and during exercise [64,65]. As a result, CP was enhanced during cycle exercise in hyperoxia versus normoxia in both the supine [64] and upright [65] body positions, with the magnitude of improvement being ~10% in both studies. Hence, studies have consistently shown that CP is sensitive to both increased [37,64,65] and decreased [21,56,60–62] FiO2.

Another experimental intervention which has yielded insights into the dependency of CP on convective O2 delivery is via manipulations in the muscle contraction duty cycle. Muscle contraction, particularly during small muscle mass exercise where compressive forces can be high, increases intramuscular pressure, compresses blood vessels, increases impedance to flow and may cause temporary blood flow occlusion [66–69]. Hence, the muscular contraction cycle yields rhythmic alterations in intramuscular pressure, and hence blood flow, with the majority of flow occurring during the relaxation phase of contraction, [69–72]. Utilizing small muscle mass handgrip exercise, Broxterman et al. [73] directly tested the hypothesis that alterations in the duty cycle would cause concomitant alterations in convective O2 delivery, and hence CP, by measuring brachial artery blood flow via Doppler ultrasound during exercise with a 20% and 50% duty cycle (i.e. muscle contraction comprised 20 and 50%, respectively, of the total contraction-relaxation cycle). Brachial artery blood flow, and thus, convective O2 delivery, was greater in the 20% duty cycle when compared to the 50% duty cycle, with a concomitant increase in CP [73].

In extending the principle of altering convective O2 delivery to observe its effect on CP, Broxterman et al. [74,75] showed that during blood flow occlusion (which constrains O2 delivery to zero), CP was reduced to a negative value. Whilst a negative CP appears implausible, this finding demonstrates a reliance of CP on convective O2 delivery since there is no sustainable oxidative metabolism without blood flow. Resting (i.e. 0W) occlusion results in progressive depletion of [PCr] and muscle/capillary O2 stores [76,77], a feature consistent with non-steady -state conditions [15]. Accordingly, the magnitude of the negative CP during blood flow occlusion would be expected to be proportional to the resting metabolic rate, and as such is entirely plausible.

These findings were recently extended by Hammer et al. [78] where critical force (CF) was estimated during the final minute of repeated handgrip MVC efforts over a 5-minute duration. Under free-flowing conditions without occlusion, force progressively declined with time during the test until a plateau was reached in the final minute of the test, termed CF [78]. With muscle occlusion, however, force continuously declined with time, i.e., there was no plateau in force at end-exercise [78]. Following subsequent reperfusion, force was able to recover to a level not significantly different from CF determined under free-flowing conditions [78]. These authors also demonstrated that up to and including CF, end-exercise limb blood flow values were linearly related to the constant-force requirements of each task [79,80]. However, during exercise slightly above CF, end-exercise brachial artery blood flow demonstrated a plateau, being no different from the blood flow values obtained during exercise at CF [79]. These findings were subsequently extended to large muscle mass, whole-body exercise by the same authors [80]. Specifically, leg blood flow and limb vascular conductance were determined using Doppler ultrasound and calibrated finger plethysmography during exercise above and below CP [80]. Post-exercise increases in limb vascular conductance and leg blood flow post-exercise were observed following supra-CP but not sub-CP exercise [80]. The data of Hammer et al. [79,80] are in contrast to observations in the running rat [25] and from upright, incremental, large muscle mass exercise in humans [52,81] showing increases in limb blood flow up to O2max. Nevertheless, these findings raise the intriguing possibility that in certain contexts, CF/CP represents a threshold in relative muscular force that limits skeletal muscle perfusion during exercise. Moreover, the extant literature appears to be unanimously consistent with CP being determined, at least in part, by mechanisms related to convective O2 delivery.

*2.2 Diffusive oxygen transport*

Diffusive O2 transport refers to the diffusive movement of O2 from the capillaries to the muscle mitochondria where O2 serves as the final electron acceptor for the electron transport system. This process is described mathematically via Fick’s law of diffusion:

Where O2 corresponds to the rate of O2 flux, *D*O2 is the muscle diffusing capacity, and Δ*P*O2 is the partial pressure difference between the capillary and intramyocyte spaces (*P*O2cap and *P*O2im, respectively). This relationship dictates that elevations in O2 must be established via changes in either 1) changes in the driving force for O2 diffusion (i.e., Δ*P*O2 = *P*O2cap - *P*O2im), and/or 2) changes in effective diffusing capacity (i.e., *D*O2, determined primarily by the aggregate number of blood cells within capillaries adjacent to the myocyte at any given moment, [82,83]).

Fick’s Law of Diffusion predicts that alterations in FiO2 will bring about concomitant alterations in CP via altered O2 diffusion in addition to convection. For instance, hypoxia reduces and hyperoxia increases both estimated *P*O2cap [84]and *P*O2im [85], though to differing extents such that PO2 is reduced and increased, respectively. Hence, in the studies reviewed in *Convective Oxygen Delivery* wherein hypoxia reduced [21,56,60–62] and hyperoxia increased CP [37,64,65], it is also probable that alterations in the transcapillary driving force for O2 flux, and thus diffusive O2 delivery, also contributed to the alterations in CP observed therein, likely via the alterations this would be expected to have on *P*O2im [55].

Muscle capillarity is an important influence on *D*O2, and thus diffusive O2 delivery, since it determines the number of red blood cells adjacent to contracting fibres and thus the surface area available for O2 diffusion. Indeed, Mitchell et al. [86] recently demonstrated a striking relationship between CP and skeletal muscle capillary density (*r* = 0.50), capillary-to-fibre ratio (*r* = 0.88), and capillary contacts per type 1 fibre (*r* = 0.94) in a homogenous group of endurance-trained individuals (63.2 ± 4.1 mL.kg-1.min-1, range: 58.7–72.2 mL.kg-1.min-1). These findings indicate that enhancements to diffusive O2 flux enable a metabolic steady-state to be attained for a greater range of power outputs (i.e. extending the range upwards), thus increasing CP.

Further insight into the role of diffusive factors in determining CP/CF was provided by a series of experiments from Ansdell et al. that compared the power-duration relationship between the sexes during small- [87] and large-muscle mass exercise [88]. It was demonstrated that CF occurred at a greater relative percentage of the maximum voluntary contraction (MVC) in females compared to males during small-muscle mass, intermittent isometric single-leg knee extension exercise [87]. Conversely, there were no differences observed in the relative percentage of MVC at which CP occurred between males and females during large-muscle mass dynamic cycle exercise [88]. Females have previously been demonstrated to possess a greater degree of capillarity in skeletal muscle and greater proportion of type I fibres when compared to males [89–91], suggesting a greater capacity for diffusive O2 transport. Moreover, during small-muscle mass knee extension exercise, far greater mass-specific rates of blood flow are achieved when compared to cycle exercise, and hence, diffusive rather than convective factors constrain O2 transport to muscle mitochondria [52,81,92–97]. These authors [87,88] consequently interpreted their findings to indicate that during single-limb exercise where convective factors are not limiting, the sex difference in CF arises due to a greater skeletal muscle diffusive capacity of females [87,88]. Conversely, during dynamic cycle exercise where muscle O2 delivery is constrained by the central nervous system to prevent a dangerous fall in mean arterial pressure [98], convective O2 delivery may be relatively more important in determining CP than muscle diffusive capacity, leading to the lack of a sex difference in this mode of exercise [87,88].

Utilizing measurements of brachial artery blood flow via Doppler ultrasound and NIRS to determine muscle O2 extraction, Broxterman et al. [73] were able to estimate muscle O2 and thereby estimate the contributions of enhanced convective and diffusive O2 delivery to the changes in CP they observed between 20% and 50% duty cycles (discussed in *Convective Oxygen Delivery*). These authors demonstrated that the increase in *D*O2 in the 20% versus the 50% duty cycle was approximately double the increase in convective O2 delivery that occurred between the same trials (i.e., +69% versus +34%, respectively), implicating changes in diffusive, rather than convective, O2 delivery as being a more important determinant of CP in this situation. These authors suggested that the shorter duty cycle would have facilitated higher red blood cell velocity and therefore increased the surface area of the capillary involved in gas exchange (i.e. longitudinal capillary recruitment, [99]), thereby enhancing *D*O2 and contributing to the increased CP. Interestingly, this observation is also consistent with the suppositions of Ansdell et al. [87,88] noted above, namely that diffusive factors may be more important for constraining CP during small versus large-muscle mass exercise. That *D*O2 is an independent determinant of CP was recently confirmed by Colburn et al. [100]. Specifically, the vascular ATP-sensitive K+ channel inhibitor glibenclamide decreased CS in rats, and this was accompanied by a 25% decrease in *D*O2 determined from measurements of skeletal muscle blood flow, arterial O2 content, and interstitial and microvascular O2 pressures [100]. Collectively, therefore, there is now a growing body of evidence to indicate that CP can be influenced by factors dictating the rate of diffusion of O2 from capillary to mitochondria.

*2.3 Oxygen utilization*

A sentinel parameter defining the skeletal muscle bioenergetics system is the time constant of the fundamental phase of muscle V̇O2 kinetics (i.e., τV̇O2), which is reflective of the time taken to attain 63% of the V̇O2 amplitude in response to a change in metabolic demand [101–104], and is closely reflected by the pulmonary τV̇O2 [103]. Pulmonary τV̇O2 is therefore a highly convenient assay of the time course of changes in oxidative phosphorylation that occur at the onset of exercise or during changes in metabolic rate. At the onset of exercise, therefore, the delayed response of pulmonary and muscle V̇O2 kinetics that is encapsulated by the parameter τV̇O2 necessitates an energy deficit that must be met via a reduction in O2 stores and an increased rate of substrate-level phosphorylation [103,105,106]. This “O2 deficit” is a function of τV̇O2 and the steady-state increment V̇O2 [105], at least for work rates where a steady-state is rapidly attained. The magnitude of this O2 deficit at exercise onset is critical, since it determines 1) the degree of reliance on nonoxidative sources of energy provision (i.e., depletion of [PCr] and [glycogen] and consequent accumulation of [L-] and [H+]), 2) the magnitude of metabolic perturbation incurred during the rest-to-work transition (i.e., Δ[PCr], Δ[ADP], Δ[Pi], extracellular [K+] accumulation, loss of sarcoplasmic Ca2+ release and sensitivity), 3) the extent of fatigue induction sustained, and 4) the loss of skeletal muscle efficiency induced during the rest-to exercise transition [8,10,14,101,102,104,107–110]. V̇O2 kinetics would therefore appear to be central in setting the tolerability of exercise. Indeed, very low τV̇O2 values (i.e., fast V̇O2 kinetics) are observed in endurance athletes [111] and trained individuals [112], whereas very large τV̇O2 values (i.e., slow V̇O2 kinetics) are observed in the elderly [113] and chronically ill [102]. However, until relatively recently, an independent role for τV̇O2 in determining CP had not been considered.

Murgatroyd et al. [14] characterized relationships between τV̇O2 and CP by normalizing exercise intensity across individuals such that the tolerable duration of exercise was uniform (6 minutes). They demonstrated a strong, inverse correlation between τV̇O2 and CP ( r = 0.95), consistent with the notion that τV̇O2 hasan independent role in determining CP. Moreover, when this analysis was extended across human populations spanning the extremes of aerobic function (i.e., healthy young trained individuals, young inactive individuals, healthy elderly, and chronic obstructive pulmonary disease patients), the relationship between τV̇O2 and CP was strong, inverse, and linear [104]. These authors interpreted this relationship causally: by minimizing the reliance on substrate level phosphorylation, and hence the accumulation of fatigue-related metabolites during the transition, a lower τV̇O2 (i.e.faster V̇O2 kinetics) allows a higher power production to be achieved for a given magnitude of O2 deficit accumulation. CP represents the upper limit of the metabolic steady state, and by extension also signifies the upper limit of an O2 deficit below which muscle fatigue, reduction in work efficiency and the O2 deficit itself will stabilise. All else being equal, therefore, faster V̇O2 kinetics will result in a higher CP. However, despite the strong rationale and cross-sectional evidence supporting a mechanistic link between τV̇O2 and CP, until recently, this hypothesis had not received direct experimental scrutiny.

In the first of a series of studies examining the purported determining effect of τV̇O2 on CP, Goulding et al. [114] examined the influence of prior heavy (“priming”) exercise on pulmonary V̇O2 kinetics and CP during supine and upright cycling. A prior bout of priming exercise does not speed V̇O2 kinetics (i.e. reduce τV̇O2) during upright cycle exercise in young, healthy individuals. However, during exercise in the supine position, muscle perfusion pressure is impaired and τV̇O2becomes O2 delivery-dependent [114–120]. Hence in a young healthy population, prior heavy exercise (which enhances muscle O2 delivery, [115,121,122]) would be expected to reduce τV̇O2during supine but not upright cycling. Accordingly, should τV̇O2 exert a determining effect on CP, an increase in CP during supine, but not upright, exercise would be observed following priming exercise as compared to control conditions. It was demonstrated that when priming exercise was conducted in the supine position, τV̇O2 was indeed reduced and CP concomitantly increased, whereas during upright exercise, both τV̇O2 and CP were unaffected [114]. These findings therefore provided the first experimental evidence that τV̇O2 is mechanistically related to CP.

Because of the nature of the priming intervention utilized in this first study [114], however, it was not possible to separate any independent effect of a reduced τV̇O2 (i.e. slowed V̇O2 kinetics) on CP from that of an improved O2 availability as a consequence of the priming exercise. Indeed, the strong correlation observed between τV̇O2 and CP for upright exercise was absent for supine exercise [114]. Hence it remained plausible that, at least in supine exercise, other physiological factors, such as muscle O2 availability, and its distribution relative to V̇O2, determine CP, with the concomitant improvements in τV̇O2 and CP being an artefact of shared physiological determinants, without any dependence of CP on τV̇O2 *per se*. Hence, confirmation or refutation of the hypothesis that τV̇O2 is an independent determinant of CP required an intervention that could alter τV̇O2 without any concomitant alterations in muscle O2 delivery, such that the independent effect of τV̇O2 on CP could be observed. When exercise is initiated from an elevated baseline work rate, τV̇O2 is greater than when compared to work initiated from a baseline of unloaded cycling [123–126]. Importantly, this slowing of the O2 kinetics appears to occur independently of any alterations to O2 availability [127–129]. Hence, we conducted two further studies that assessed the influence of exercise initiated from an elevated baseline work rate on τV̇O2 and CP in the upright [130] and supine [117] positions. In both of these studies, τV̇O2 was greater (i.e., V̇O2 kinetics was slower) and CP was correspondingly reducedduring work-to-work exercise compared with when exercise was initiated from a baseline of unloaded cycling [117,130]. Crucially, indicators of O2 availability determined via NIRS were either improved [130] or unchanged [117] during work initiated from an elevated baseline, suggesting that the slowing of O2*p*kinetics brought about by this intervention was wholly independent of changes in microvascular O2 availability. Taken together, these findings therefore demonstrate an independent effect of τV̇O2 on CP [130], and that this effect persisted even in situations where O2 delivery is substantially impaired [117].

The determining effect of τV̇O2 on CP observed in healthy populations [64,114,117,130] was later confirmed in a study that assessed the impact of priming exercise on V̇O2 kinetics and CP in a population of individuals with type 1 diabetes [131]. In this population, priming exercise speeded V̇O2 kinetics and increased CP during subsequent severe-intensity cycle exercise. Notably, these effects were accompanied by a concomitant speeding of muscle deoxygenation kinetics determined via NIRS [131]. Since the muscle deoxygenation signal derived via NIRS represents the relative balance between O2 delivery and utilization within the interrogated region, a relative speeding of muscle deoxygenation kinetics suggests that the effects of priming exercise on τV̇O2 were predominantly due to an upregulation of otherwise impaired intracellular mechanisms of mitochondrial O2 utilization, rather than O2 delivery [131]. Taken together, therefore, substantial recent evidence has accumulated to demonstrate that rates of intracellular O2 utilization at the onset of exercise, encapsulated by τV̇O2, can influence CP independently of factors related to mitochondrial O2 provision.

1. **INTERACTION OF FACTORS DETERMINING CP**

The studies of Goulding et al. [8,64,65,114,117,130,131] provide convincing evidence that τV̇O2 is an independent determinant of CP. As reviewed above, there is also evidence for an independent determining role of convective and diffusive O2 delivery in influencing CP. That each of τV̇O2, convective and diffusive O2 delivery has an independent role in determining CP is evinced by the fact that each one can alter CP without a concomitant change in the other.

The proportion of CP explained by τV̇O2 has been reported to be as high as 90% in a homogenous participant group where relative exercise intensity was precisely controlled (i.e. a tolerable duration of 6 minutes across subjects) [14]. Our own data has demonstrated *R*2 values of 0.64 to 0.90 for the relationship between CP and τV̇O2 during upright exercise [60,111, ,127,128]. Collation of this data across differing exercise intensity domains and populations, including hyperoxia conditions, yields an *R*2 = 0.60 (Figure 3A; data from ref. [131] previously unpublished), with a slope of ~0.03 W.kg-1.s-1. However, this includes data from diseased populations (type 1 diabetes [131]) and hyperoxia [65], both of which might be expected to confound the analysis since the latter may distort the relationship between pulmonary and muscle τV̇O2 and the former has a slope (0.01 W.kg-1.s-1) significantly different to the healthy populations. Exclusion of diseased and hyperoxic data blunts the strength of the relationship between CP and τV̇O2 (*R*2 = 0.43 (Figure 3B). However, the strength of this relationship increases markedly when only moderate intensity exercise in healthy participants is considered (*R*2 = 0.79; Figure 3C). The slope of the relationship between τV̇O2 and CP was preserved across this latter analysis, and taken together, CP appears to be well predicted from τV̇O2 when the latter is precisely determined for a given relative exercise intensity, varying by ~0.03 W.kg-1 per second change in τV̇O2. However, and perhaps exemplified by the data from type 1 diabetes [131] and hyperoxia [65], when this relationship is expanded to cover the range of values for τV̇O2 encountered across the animal kingdom (Figure 3D), the relationship with CP appears curvilinear, but nevertheless preserved, suggesting a fundamental linkage of CP with muscular bioenergetics across species. Moreover, when the human-only data are considered and the speed of oxygen uptake kinetics expressed as a rate constant (i.e. 1/τV̇O2), the relationship with CP is linear (Figure 3E). Accordingly, when the scope of human aerobic fitness is considered, the relationship between CP and τV̇O2 can be considered to be hyperbolic, with previously published linear relationships [14,104] being an artefact of participant homogeneity. By contrast, only one previous study has titrated the effect of oxygen delivery on CP [63]. Here, the reduction in CP with increasing altitude as a proxy for oxygen delivery was established, simulated by changes to FiO2. A nonlinear (3rd order polynomial) relationship was established with increases in altitude producing progressively larger reductions in CP. CP was reduced by 74W with a 4000m increase in altitude, though any such relationship will inevitably be impacted by the effect of reductions in FiO2 increasing τV̇O2 (i.e. slowing O2 kinetics).

Given the evidence reviewed herein, we therefore propose that each of mitochondrial O2 utilization (encapsulated by the τV̇O2 parameter), convective and diffusive O2 delivery exert independent effects on CP such that intracellular O2 utilization and O2 transport interact to determine CP (Figure 4). Exceptions to this include where pulmonary limitations (e.g. [34]) are dominant factors in limiting exercise tolerance to the extent that they dictate the shape of the power – duration relationship.

The precise mechanisms underpinning such an interaction have not been fully elucidated, however a starting point is to consider the inexorable loss of intracellular homeostasis, and thus unsustainable rise in O2 deficit, during exercise above, but not below, CP. This is accompanied by a mirror-like association between peripheral fatigue [107,132] and the loss of exercise efficiency [109,133,134] that occurs during exercise above CP [135]. Of the factors which accumulate as a result of the O2 deficit, [Pi] is a prime candidate for the common denominator between fatigue and efficiency due to its central role in muscle fatigue and task failure [136]. A recent *in silico* study by Korzeniewski & Rossiter [10] tested the hypothesis that accumulation of [Pi] during the transition from rest-to-work could explain both the loss of intracellular homeostasis during supra-CP exercise and the fatigue-related termination of exercise. Using a validated model of the human bioenergetic system, Korzeniewski & Rossiter [10] defined a “critical” (i.e. threshold) [Pi] above which further [Pi] accumulation drove an increase in the requirements for ATP turnover (i.e., an increased ATP cost of muscle contraction) and a “peak” (i.e. limiting) [Pi] at which exercise would cease. The additional ATP turnover driven by [Pi] accumulation resulted in a self-propagating positive feedback loop where additional ATP turnover resulted in increased [Pi], which caused fatigue and additional ATP turnover, and so on until the pre-defined peak [Pi] (and accompanying muscle V̇O2max) was achieved. By contrast, when [Pi] accumulated below or only marginally above critical [Pi] this positive feedback loop stabilized such that [Pi] did not attain peak values and muscle oxygen uptake attained a steady state. Based on these findings we therefore recently proposed a model whereby muscle O2 consumption kinetics determine CP by dictating the magnitude of O2 deficit (and thus [Pi], amongst other factors) accumulated during a given exercise transition [8]. Slow V̇O2 kinetics begets large intracellular perturbations whereas fast V̇O2 kinetics engenders smaller intracellular perturbations for a given metabolic rate at exercise onset [102,104,110,137]. Accordingly, more rapid V̇O2 kinetics will enable a higher exercise intensity before a critical value of [Pi] is breached, thereby increasing critical power, all else being equal. Importantly, simulating alterations in *P*O2im within the computer model of Korzeniewski & Rossiter [10] resulted in the changes in τV̇O2 and CP predicted by the evidence reviewed in each of the previous sections [10].

Alongside other O2 deficit-related factors, that breaching a critical [Pi] results in an inexorable cascade of increasing [Pi], fatigue and ATP turnover is also consistent with the evidence reviewed herein whereby convective and diffusive O2 delivery has a determining effect on CP. O2 delivery is known to regulate the concentrations of phosphate metabolites at a given metabolic rate, such that when intracellular *P*O2 is higher, the intracellular perturbations incurred for [Pi], [PCr] and [ADP] are reduced, whereas the reverse is true when intracellular *P*O2 is lower [49,50,54,138]. From these observations it follows that the aforementioned effects of convective and diffusive O2 delivery and intracellular O2 utilization on CP stem from their impact upon the intracellular metabolic state, or more specifically, the rate of ATP turnover at which a critical threshold for [Pi] (which is itself a proxy for a collection of intracellular metabolites reflecting the intracellular state of fatigue) is attained. Hence, faster V̇O2 kinetics, as well as increased O2 delivery, exert their effects on CP via reducing the intracellular metabolic perturbations required to sustain a given rate of ATP turnover, thus enabling a higher power output to be achieved before CP is reached.

1. **INTEGRATION OF MECHANISMS: WHOLE-BODY**

This model of CP being an emergent property of the metabolic derangements established at the onset of exercise may provide an explanation for the metabolic bases of CP at the level of a single fibre, however, it does not pretend to be a complete explanation of CP at the integrative whole-body level. This is despite the *in silico* approach of Korzeniewski & Rossiter [10] being “chimeric”, in that it is built using the data of whole-body and –muscle responses of V̇O2, [PCr], [Pi] and pH and reflecting a variety of muscle fibre types, averaged into a single response. In practice, the exercise transition is undertaken by muscle fibres across the spectrum of function, with differing underlying oxidative phosphorylation activities, each-step activation intensities, convective & diffusive O2 supply and fatigue characteristics [139–141]. Additionally, the location of a given fibre with respect to the skin surface has implications for the relative O2 delivery [45,118–120,142]. Nevertheless, findings at the whole muscle level are congruent with the notion that metabolic inertia at the onset of exercise determines CP via its effect on the accumulation of Pi and other O2 deficit-related metabolites that are implicated in the fatigue process. Type I fibres possess faster V̇O2 kinetics, better metabolic control, and maintain greater values for capillary and interstitial *P*O2 at rest and during contractions [139,140,143–150]. Hence, as detailed earlier, in human biopsy studies the proportion of type I fibres and indices of muscle fibre capillarization have been shown to be closely associated with CP [31,86].

Moreover, given that type I fibres maintain greater values for capillary and interstitial *P*O2 at rest and during contractions (presumably due to enhanced capillarization) [139,140,143–150], these data are consistent with the present proposal that τV̇O2, convective and diffusive O2 delivery each exert independent, interactive determining effects on CP. However, the eventual external outcome of interest from all of these processes, i.e., CP, will also be a function of factors such as (relative) exercising muscle mass, the local musculoskeletal lever system dynamics and co-ordination, the extent of localized fatigue within working muscle groups and motor-unit recruitment. Indeed, that our data demonstrates a significant relationship between τV̇O2 and CP expressed in W.kg-1, but not W (data not shown) speaks to the role of exercising muscle mass in the eventual determination of CP. The individual muscles of the quadriceps muscle group have been shown to produce divergent patterns of [PCr] depletion and [Pi] accumulation within distinct muscle regions (72 cm3 voxels) during fatiguing (incremental) exercise [151]. Exercising muscle mass may therefore also play a role in the extent of such muscle metabolite heterogeneity, and thus the degree of metabolic perturbation within distinct muscle regions, which in turn contributes towards setting CP.

Morgan and colleagues [152,153] provided insight into how muscle recruitment patterns may act to determine CP. During repeated intermittent isometric contractions, the ingestion of acetaminophen led to a smaller reduction in torque across 60 MVCs when compared to a placebo [153]. This was associated with a greater preservation of muscle activation with acetaminophen as assessed via electromyography (EMG). Subsequently it was shown that during upright cycle ergometry, acute acetaminophen ingestion increased CP and preserved muscle activity throughout the duration of exercise when compared to a placebo [152]. These findings suggest blunting neuromuscular fatigue development and preserving muscle activation enhances CP, and thus demonstrates the importance of motor unit recruitment profiles.

The interaction between muscle recruitment patterns and muscle O2 delivery in determining CP is perhaps most strikingly illustrated by the recent study of Hammer et al. [78], discussed previously (see *Convective O2 delivery*). These authors showed that, following muscle reperfusion, both muscular activity and force production returned to levels not different from that observed under free-flowing conditions [78]. Hence, muscle occlusion constrained muscular recruitment and thus critical force, however, once muscle perfusion was restored to pre-occlusion conditions, both muscle recruitment and force-generating capacity were restored. These findings illustrate that CP represents an intricate balance between muscle O2 supply, muscle recruitment patterns and peripheral fatigue development.

To summarize, CP is sensitive to muscle fibre type composition because it is a parameter of aerobic function. Hence, the oxidative characteristics inherent within type I fibres, such as rapid V̇O2 kinetics, greater rates of blood flow and higher capillary and interstitial *P*O2 values, allow the attainment of high rates of ATP utilization with minimal derangement of the intracellular metabolic milieu. Therefore, all else being equal, individuals with a relatively greater proportion of type I skeletal muscle fibres will tend to possess greater CP values when compared to individuals of equivalent training status with a greater proportion of type II fibres. Moreover, animal data indicate that CP appears to be a critical threshold for the recruitment of high-order motor units containing a high fraction of type II fibres [25]. Hence, individuals with more type I fibres will attain a relatively greater fraction of their V̇O2 max before reaching the threshold for progressive recruitment of type II fibres, i.e., CP (as seen in highly trained humans, [111,154]). Interventions that increase motor unit recruitment are also conducive to high CP values, as a greater number of motor units/muscle fibres performing a given task will lessen the metabolic strain on each individual fibre. Hence, when muscular recruitment is increased, each fibre is able to maintain intramuscular metabolite accumulation below its critical threshold for a wider range of ATP utilization rates, thus enabling a greater CP, as suggested for the effects of priming by Burnley et al. [155]. Hence, although there is clearly a role for convective and diffusive O2 delivery and intracellular O2 utilization in determining CP, understanding the physiology of CP at the level of integrative physiology is only possible via consideration of how these factors interact with muscle fibre type composition and recruitment patterns.

1. **CONCLUSIONS**

CP separates the heavy and severe exercise intensity domains wherein qualitatively divergent physiological responses are observed, such that CP represents the threshold intensity above which a metabolic steady-state cannot be attained during exercise. Hence, CP is fundamental to the understanding of human endurance performance and the causes of exercise limitation in populations where exercise tolerance is impaired. Over the past 15 years or so, evidence has emerged that CP also represents a key threshold for a variety of aspects of physiological system behaviour, such as muscle fibre recruitment, blood flow and vascular control, as well as muscle fatigue. Accordingly, a wide range of evidence has emerged, spanning each step of the oxygen transport pathway, that CP is a fundamental parameter of aerobic function. It has been demonstrated that alterations in delivery of O2 to the exercising muscles, via both convection and diffusion, impact upon CP. The rates of O2 utilization during exercise, particularly during the transition from rest-to-work, also plays a key role in determining CP by governing the degree of matching between the rates of ATP utilization and production. These factors each interact with one another, and via this interaction determine the degree of intracellular metabolic disturbance required to sustain a given power output. How each of these factors interacts to determine CP at the whole-body level will be dependent upon the muscle fibre type composition and their recruitment patterns during exercise.

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**CONFLICTS OF INTEREST**

Richie Goulding and Simon Marwood declare that they have no conflicts of interest relevant to the content of this review.

**AUTHOR CONTRIBUTIONS**

RPG conceived the work and wrote the first draft. Both authors conducted the literature search, revised the manuscript critically for important intellectual content and approved the final version to be submitted. Both authors agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

**ETHICAL APPROVAL**

Ethical approval was not required for this review paper.

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**Figure 1.** The hyperbolic power-duration curve that defines the sustainable duration of exercise in the severe-intensity domain. This hyperbolic relationship is defined by two parameters: the power asymptote, known as the critical power (CP), and the curvature constant W' (denoted by the rectangular dashed blue lines above CP and expressed in kilojoules). CP defines the boundary between the heavy and severe exercise intensity domains and represents the highest power output for which a metabolic steady state may be attained. The W' comprises a fixed and finite volume of work that is expendable above CP. During severe-intensity exercise, task failure occurs when W' = 0. GET: gas exchange threshold.

**Figure 2.** This schematic illustrates an adaptation of Wasserman’s classic “Gears” diagram. It demonstrates Wasserman’s conception of how the respiratory, cardiovascular and neuromuscular systems conflate to enable exercise to be sustained. O2 flows from the atmosphere through the lungs, pulmonary and peripheral circulation to the muscle mitochondria where it is ultimately consumed. CO2 produced by the contracting muscle flows along the same pathway in reverse. Muscle work leads to increased cardiac output and redistribution of blood flow, and increased ventilation in response to both the increased metabolism and evolution of CO2 from the blood as the result of lactic acid buffering. The efficacy of these processes determines the ability to sustain muscular exercise. These concepts are reconsidered in this review within the context of critical power. This figure was created with BioRender.com and was exported under a paid subscription. O2O2 matching, matching of oxygen delivery to local oxygen consumption; O2, cellular oxygen consumption; CO2,cellular carbon dioxide production; matching, matching of ventilation to perfusion; *A*, alveolar ventilation; *D*, dead space ventilation, *E*, minute ventilation, O2, pulmonary oxygen consumption; CO2, pulmonary carbon dioxide production. Adapted from Wasserman et al. [156], with permission.

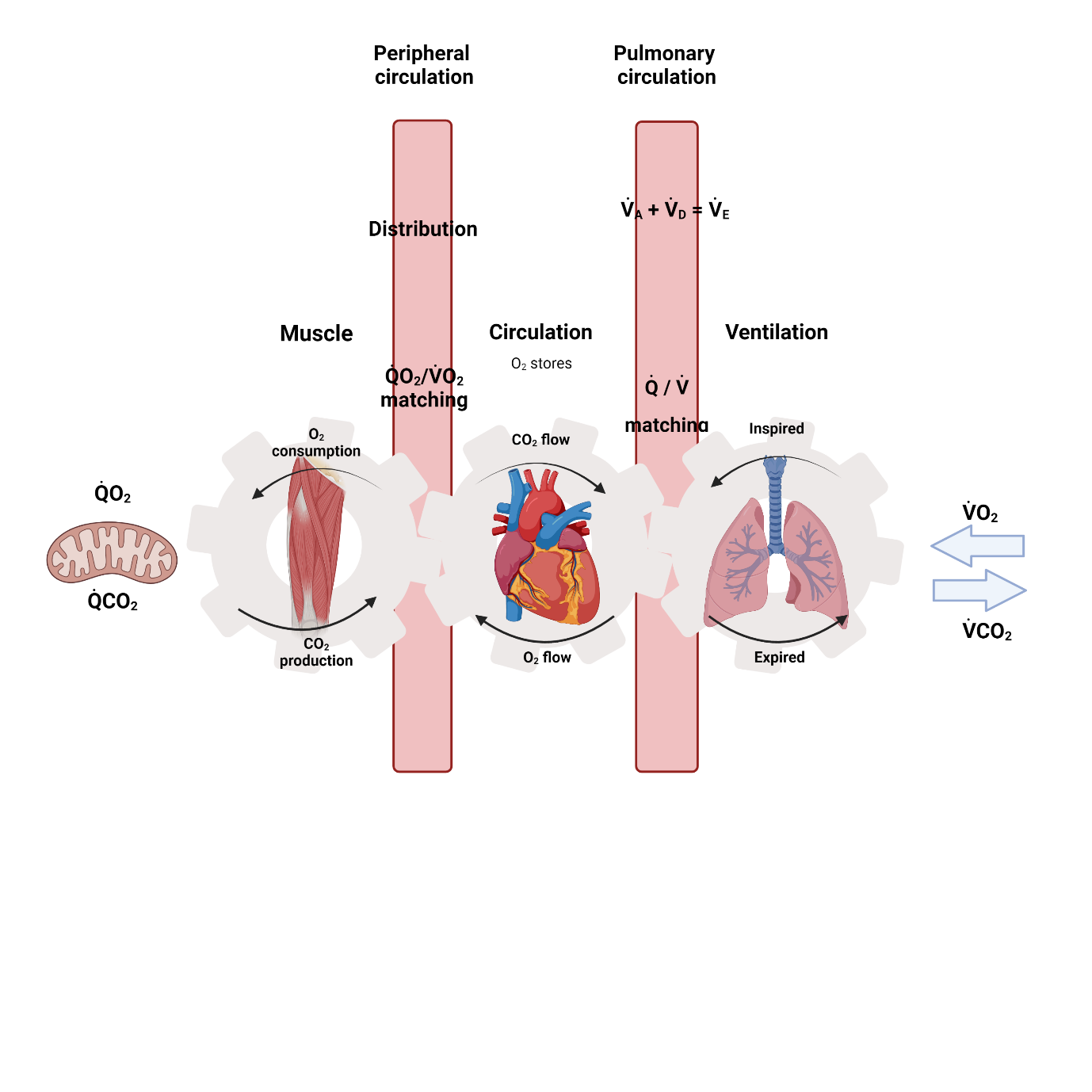
**Figure 3.** Panels A-C show the relationship between the fundamental phase time constant of pulmonary oxygen uptake kinetics (τV̇O2) and critical power normalized by body mass across a series of 4 experiments performed by Goulding et al. [65,114,130,131]. Panel A displays all conditions from these studies in which τV̇O2 was characterized with a high degree of confidence, including both moderate and heavy intensity exercise, normoxia and hyperoxia (fraction of inspired O2 = 0.5), and in patients with type 1 diabetes. Panel B displays the same relationship with removal of data points where τV̇O2 was characterized in hyperoxic conditions and in type 1 diabetes (see *Interaction of factors determining CP”* for discussion). Panel C displays the relationship when only normoxic, moderate intensity exercise transitions in healthy participants are utilized. Note the increase in the *R2* value as the conditions become more uniform with respect to exercise intensity, population and fraction of inspired O2. Panel D shows the relationship between τV̇O2 and critical O2 across various human populations; elite athletes [157], young trained, active young, healthy elderly, and patients with chronic obstructive pulmonary disease) and other species where measurements of τV̇O2 and critical power have both been conducted (i.e. the thoroughbred racehorse, rat, ghost crab and lungless salamander). The figure is derived from values reported in the literature of 28 papers published between 1982 and 2010; human populations were originally reported by Rossiter [104], with groups which were approximately matched for age, *O2* max, and health status. Electronic Supplementary Material Table S1 should be consulted for details regarding derivation of critical O2 in different species. Panel E shows human-only data from panel D of CP (ml.kg-1.min-1) plotted as a function of 1/ τV̇O2 (i.e. the rate constant, *k*). There is a notable linear relationship across what can be regarded as the complete range of human fitness, indicating that the relationship between τV̇O2 and CP is hyperbolic, with previously published linear relationships likely being a function of participant homogeneity, and thus reflecting only a truncated portion of the hyperbolic relationship. COPD, chronic obstructive pulmonary disease.

**Figure 4.** This schematic illustrates each of the factors that has been demonstrated to impact upon critical power. Convective and diffusive O2 delivery act in concert with muscle O2 utilization to determine the degree of intracellular metabolic perturbation and fatigue induction incurred during the rest-to-exercise transition. The extent of such metabolic perturbations, in turn, determine whether or not an exercise bout can be met in a metabolic steady-state within a given myocyte. Within a given individual, whether or not an extant power output is met in a whole-body steady-state will depend on the muscle fibre type composition of the individual, the muscle recruitment patterns employed during the task, and the extent of metabolic derangement and fatigue induction incurred in the recruited fibres during the rest-to-exercise transition. This figure was created with BioRender.com and was exported under a paid subscription. , cardiac output; CaO2, arterial oxygen content, PO2cap, capillary O2 pressure; *P*O2im, intramyocyte O2 pressure; DO2, muscle diffusive capacity.

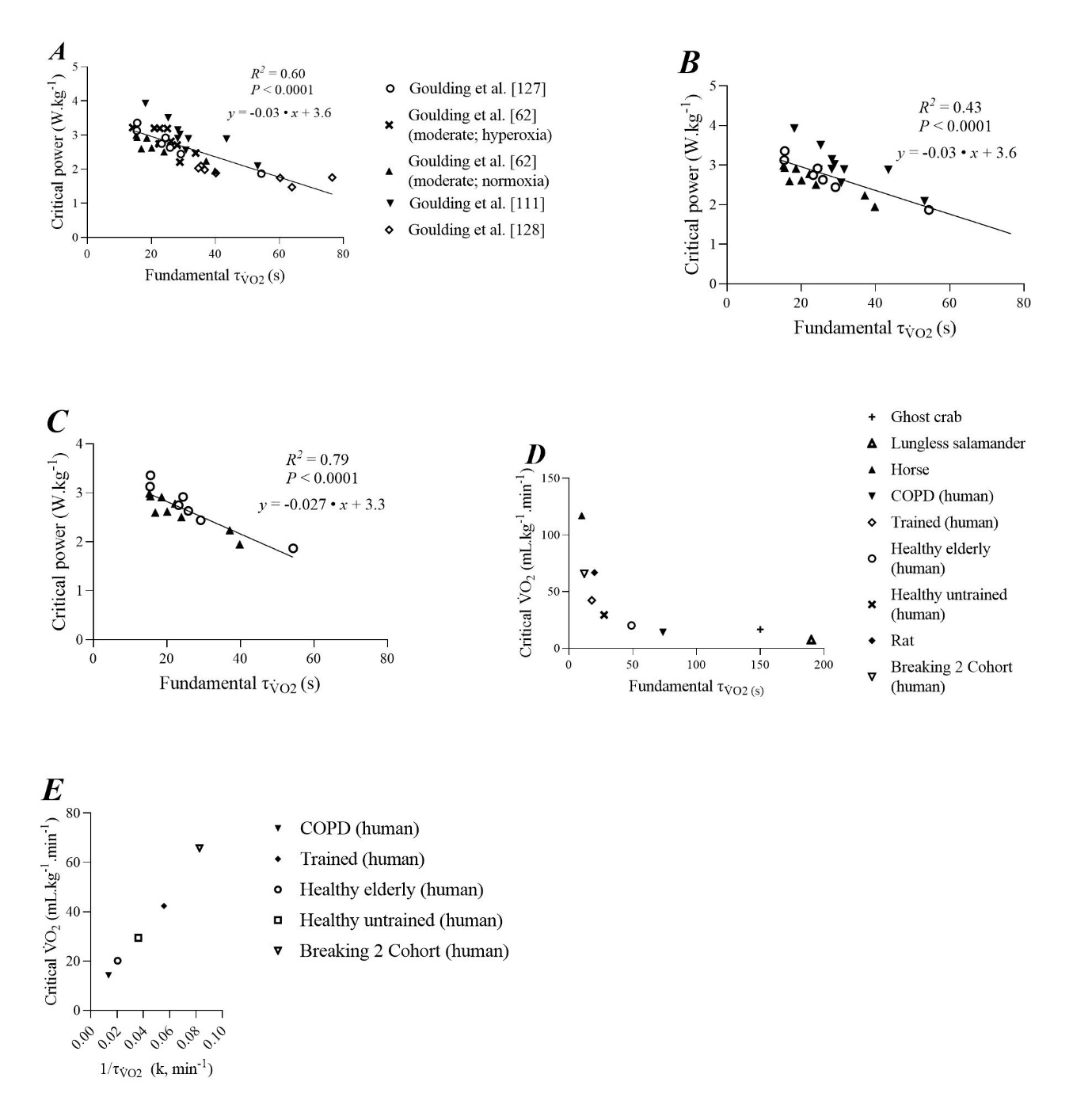
**Figure 1.**



**Figure 2.**



**Figure 3.**



**Figure 4.**

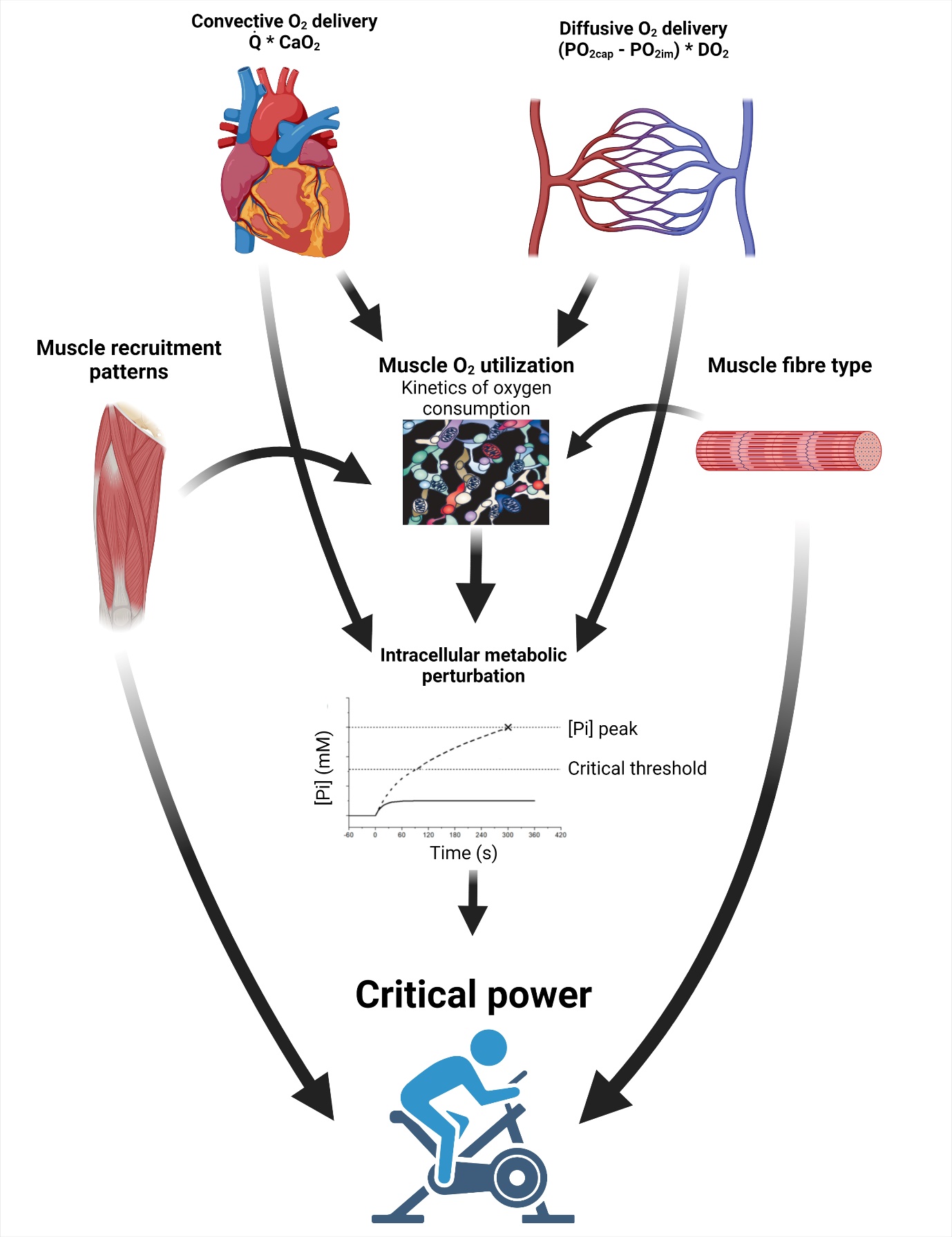


Table 1. Summary of studies that have altered CP via chronic or acute interventions

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Author / date** | **Population** | **Mode** | **Intervention** | **Effect on CP** | **Physiological effects of intervention** | **CP determination method** |
| Moritani et al. [21] | H (2) | Upright cycling | Hypoxia (FiO2 = 0.09) | ↓ CP  (106 vs 214 W) |  | 4CWR |
| Gaesser & Wilson [158] | ETM (2) HM (3) | Upright cycling | Endurance training  (6 weeks) | ↑ CP  (228 vs 201 W) | ↔V̇O2peak | 4CWR |
| Gaesser & Wilson [158] | ETM (3) HM (3) | Upright cycling | HIIT  (6 weeks) | ↑ CP  (254 vs 220W) | ↑V̇O2peak | 4CWR |
| Poole et al. [159] | HM (8) | Upright cycling | HIIT  (7 weeks) | ↑ CP  (288 vs 325 W) | ↑V̇O2peak, ↑LT | 5CWR |
| Jenkins & Quigley [160] | HM (12) | Upright cycling | Endurance training  (8 weeks) | ↑ CP  (255 vs 196 W) | ↑V̇O2peak | 3CWR |
| Hill [161] | HM (13) HF (11) | Upright cycling | Cadence  (100 rpm vs 60rpm) | ↓ CP  (195 vs 207 W) |  | 4CWR |
| Serres et al. [162] | COPD (8) | Upright single leg knee extension | Endurance training  (3 weeks) | ↑ CP  (1.8 vs 1.3 kg.s-1) | ↑V̇O2peak, ↑MVC | 3CWR |
| Puente-Maestu et al. [38] | COPDM (27) | Upright cycling | Endurance training  (6 weeks) | ↑ CP  (65 vs 58 W) | ↑V̇O2peak, ↓peak blood [La], ↓V̇epeak | 3CWR |
| Barker et al. [163]\* | ETM (5) ATM (6) | Upright cycling | Cadence  (100 rpm vs 60rpm) | ↓ CP  (189 vs 297 W) |  | 4CWR |
| Vanhatalo et al. [164] | HM (8)  HF (1) | Upright cycling | HIIT  (4 weeks) | ↑ CP  (255 vs 230 W) | ↑V̇O2peak, ↑GET | 3MT |
| Miura et al. [165] | HM (6)  HF (2) | Upright cycling | Heavy priming exercise | ↑ CP  (177 vs 169 W) |  | 4CWR |
| Vanhatalo et al. [37] | HM (7) | Prone knee extension | Hyperoxia  (FiO2 = 0.7) | ↑ CP  (18 vs 16 W) | ↓rate of change: muscle [ADP], [PCr], [Pi], pH; ↑PCr, ↑Δ[HbO2], ↓Δ[HHb], ↑TOI , ↑TD[HHb], ↔ HHb] 1 | 4CWR |
| Corn & Barstow [166] | HM (7) | Upright cycling | N-acetylcysteine  (acute oral supplementation) | ↑ CP  (232 vs 226 W) | ↑GSH, ↑EMGMPF (RF), ↓EMGRMS (VL) | 4CWR |
| Dekerle et al. [56] | HM (5)  HF (6) | Upright cycling | Hypoxia  (FiO2 = 0.15) | ↓ CP  (190 vs 220 W) | ↓SaO2 | 3-4CWR |
| Valli et al. [61] | HM (4)  HF (2) | Upright cycling | Hypoxia  (altitude = 5050m) | ↑ CP  (123 vs 81 W) | ↓V̇O2peak, ↓blood [lactate], ↓SaO2, ↓O2 pulse | 3 CWR |
| Broxterman et al. [73] | HM (8) | Handgrip | Duty cycle  (50% vs 20%) | ↓ CP  (3.9 vs 5.1 W) | ↓Q̇BA, ↑iEMG, ↓EMGMPF, ↓mV̇O2, ↔ [THb], ↓end-exercise [HHb] 2 | 3-4CWR |
| Mueller et al. [167] | ETM (11) | Upright cycling | Resistance + vibration training  (8 weeks) | ↑ CP  (296 vs 286 W) | ↑capillary:fibre, ↑thigh LBM, ↑MyHC1 & ↑MyHC2 CSA,  ↔ SDH | 4CWR |
| Broxterman et al. [168]\* | ETM (5) ATM (5) | Upright cycling | Cadence  (100 rpm vs 60rpm) | ↓ CP  (196 vs 214 W) |  | 4 CWR |
| Black et al. [169] | HM (10) | Upright cycling | Pacing  (self vs. constant load) | ↑ CP  (265 vs 250 W) | ↓MRTV̇O2, ↑VO2 in first 60 s | 3-4TT / 3-4CWR |
| Broxterman et al. [75] | HM (6) | Handgrip | Blood flow occlusion | ↓ CP  (-0.7 vs 4.1 W) | ↓EMGRMS, ↑[HHb], ↓[HbO2], ↓[THb] 2 | 4CWR |
| Parker-Simpson et al. [60] | HF (13) | Upright cycling | Hypoxia  (FiO2 = 0.13) | ↓ CP  (132 vs 175 W) ↓ EP  (134 vs 172 W) | ↓V̇O2max | 5CWR & 3MT |
| Deb et al. [170] | ETM (11) | Upright cycling | Hypoxia  (FiO2 = 0.145) +/- sodium bicarbonate | ↓ CP  (265 vs 263 vs 301 W) | ↓SaO2 | 3MT |
| Goulding et al. [114] | HM (10) | Supine cycling | Heavy priming exercise | ↑ CP  (185 vs 177 W) | ↓V̇O2, ↔V̇O2max, ↑[HbO2], ↑[HHb] 1 | 4CWR |
| Townsend et al. [63] | ETM (9) | Upright cycling | Hypoxia  (FiO2 = 0.18, 0.159, 0.14, 0.123) | ↓ CP  (257, 235, 218, 196 vs 270 W) |  | 3TT |
| Clark et al. [171] | ETM (6) | Upright cycling | 2 hours heavy exercise | ↓ CP  (282 vs 306 W) |  | 3MT |
| Goulding et al. [117] | HM (8) | Supine cycling | Exercise transition from elevated baseline | ↓ CP  (132 vs 146 W) | ↑V̇O2, ↔V̇O2max, ↔ [HbO2], ↑[HHb], ↓Δ[HHb]/ΔV̇O2 3 | 4CWR |
| Goulding et al. [130] | HM (7) | Upright cycling | Exercise transition from elevated baseline | ↓ CP  (203 vs 213 W) | ↑ V̇O2, ↑ [HbO2], ↑[HHb], ↓Δ[HHb]/ΔV̇O2 1 | 4CWR |
| La Monica et al. [62] | HM (21) | Upright arm cycling | Hypoxia  (FiO2 = 0.14) | ↓ CP  (85 vs 90 W) | ↓V̇O2peak | 4CWR |
| Mitchell et al. [86] | ETM (21) | Upright cycling | SIT, SIT + blood flow restriction  (4 weeks) | ↑ CP  (302, 302 vs 292 W) | ↑V̇O2peak, ↔capillarity, ↔mitochondrial protein content | 3-5CWR |
| Clark et al. [172] | HM (14) | Upright cycling | 2 hours heavy exercise | ↓ CP  (CWR: 256, EP: 256 vs EP: 287 W) | ↓muscle [glycogen], ↔V̇O2peak | 4CWR & 3MT |
| Clark et al. [173] | ETM (16) | Upright cycling | 2 hours heavy exercise | ↓ CP  (236 vs 260 W) | ↓muscle [glycogen], ↔V̇O2peak | 3MT |
| Goulding et al. [64] | HM (8) | Supine cycling | Hyperoxia  (FiO2 = 0.5) | ↑ CP  (148 vs 134 W) | ↑V̇O2max, ↓V̇O2, ↑[HbO2], ↔[HHb] 3 | 4CWR |
| Morgan et al. [152] | HM (16) | Upright cycling | Acetaminophen  (acute oral supplementation) | ↑ CP  (297 vs 288 W) | ↑EMGRMS, ↔V̇O2peak | 3MT |
| Waldron et al. [174] | HM (12) | Upright cycling | Taurine  (acute oral supplementation) | ↑ CP  (212 vs 197 W) | ↑post-exercise blood [lactate] | 3MT |
| Goulding et al. [65] | HM (9) | Upright cycling | Hyperoxia  (FiO2 = 0.5) | ↑ CP  (216 vs 197 W) | ↑V̇O2max, ↔ V̇O2 ↑PetO2, ↑[HbO2], ↓[HHb], ↔[HHb] 3 | 4CWR |
| Goulding et al. [131] | T1DM (7) | Upright cycling | Heavy priming exercise | ↑ CP  (161 vs 149 W) | ↓V̇O2, ↔V̇O2max, ↔[HbO2], ↓[HHb] 1 | 4CWR |
| Karabiyik et al. [175] | TM (32) | Upright cycling | SIT  (4 weeks)  +/- hypoxia (FiO2 = 0.135) | ↑ CP  (200 vs 170 W)# | ↑post-ramp blood [lactate], ↔V̇O2peak | 3MT |
| Collins et al. [176] | HM (5)  HF (6) | Upright cycling | Endurance training  (8 weeks) | ↑ CP  (161 vs 140 W) | ↑V̇O2max | 3-6CWR |
| Collins et al. [176] | HM (6)  HF (5) | Upright cycling | HIIT (8 weeks) | ↑ CP  (176 vs 140 W) | ↑V̇O2max | 3-6CWR |

**Author/date:** \*latter publication uses a sub-set of data taken from the former publication

**Population**: M, healthy male; F, healthy female; ET, endurance trained; COPD, chronic obstructive pulmonary disease; AT, anaerobically trained; T1D, type 1 diabetes; (n), number of participants

**Intervention:** FiO2, fraction of inspired O2; HIIT, high intensity interval training; SIT, sprint interval training; rpm, revolutions per minute, +/-, with and without

**Effect on CP**: CP, critical power; ↑, increased; ↓, decreased; #, values for CP estimated from visual inspection of figures

**Physiological effects of intervention (all factors considered for chronic interventions, only those factors measured during the determination of CP considered for acute interventions):** ↔, unchanged; V̇O2, rate of oxygen uptake; VO2, total oxygen consumed;V̇O2peak, highest V̇O2 recorded but not verified with additional tests >CP; V̇O2max, maximal V̇O2 recorded following verification from additional trials >CP; MRTV̇O2, mean response time of V̇O2; V̇O2, time constant of V̇O2 kinetics; LT, lactate threshold; MVC, maximal voluntary contraction; La, lactate, V̇epeak, highest ventilation measured; GET, gas exchange threshold; ADP, adenosine diphosphate; PCr, phosphocreatine; Pi, inorganic phosphate; HbO2, oxygenated haemoglobin; HHb, deoxygenated haemoglobin; THb, total haemoglobin; [HHb], time constant of [HHb] kinetics;SaO2, arterial oxygen saturation; Q̇BA, brachial artery blood flow; mV̇O2: muscle V̇O2 estimated via combined near infrared spectroscopy and doppler ultrasound; iEMG, integrated electromyography; EMGMPF, electromyography median power frequency; EMGRMS, electromyography root mean squared; LBM, lean body mass; MyHC1, myosin heavy chain 1; MyHC2, myosin heavy chain 2; CSA, cross-sectional area; SDH, succinate dehydrogenase; PetO2, end-tidal pressure of O2; RF, rectus femoris; VL, vastus lateralis; 1[HHb], [THb], [HbO2] determined via near infrared spectroscopy on the VL; 2[HHb], [THb], [HbO2] determined via near infrared spectroscopy on the flexor digitorum superficialis; 3[HHb], [THb], [HbO2] determined via near infrared spectroscopy on the VL & RF

**CP determination method:** nCWR, number of constant work-rate trials; 3MT, 3-minute all-out test; nTT, number of time-trials

Table 2: Summary of studies demonstrating physiological or performance factors that correlate with CP

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Study** | **Population** | **Mode** | **Correlation details** | **CP determination method** |
| Neder et al. [34] | MH (10) & MCOPD (8) | Upright cycling | CP (W) correlated with MRTV̇O2 (s)during severe intensity exercise in MH  (r = -0.65) but not MCOPD | 4CWR |
| Murgatroyd et al. [14] | HM (14) | Upright cycling | CP (W) correlated with V̇O2 (s)during severe intensity exercise  (r = -0.95) | 4CWR |
| Black et al. [13] | ETM (10) | Upright cycling | CP (W) correlated with 10mile TT performance (min) (r = - 0.83) | 3MT |
| Vanhatalo et al. [31] | HM (4) HF (4) | Upright cycling | CP (W) correlated with %type 1 fibre (r = 0.67) | 3MT |
| Goulding et al. [114] | HM (10) | Upright cycling | CP (W.kg-1) correlated with V̇O2 (s)during heavy intensity exercise  (r = -0.80) | 4CWR |
| Byrd et al. [177] | ATM (15) | Upright cycling | CP (W) correlated with LBM (kg) (r = 0.59) | 3MT |
| Goulding et al. [130] | HM (7) | Upright cycling | CP (W.kg-1) correlated with V̇O2 (s)during moderate intensity exercise  (r = -0.95) | 4CWR |
| Mitchell et al. [86] | ETM (14) | Upright cycling | CP (W) correlated with: %type 1 fibre (r = 0.79), no. of capillary contacts in type 1 (r = 0.94) & type 2 (r = 0.68) fibres, capillary:fibre (r = 0.88) | 3-4CWR |
| Goulding et al. [131] | HM (9) | Upright cycling | CP (W.kg-1) correlated with V̇O2 (s)during moderate intensity exercise  (r = -0.92) | 4CWR |
| Smyth & Muniz-Pamares [28] | HM & HF (31,190) | Running | CS (m.s-1) correlated with marathon time (min) (r = -0.83) | 3TT |
| Collins et al. [176] | HM (11) HF (11) | Upright cycling | CP (W) correlated with: V̇O2max (ml.min-1) (r=0.96), Ppeak (W) (r=0.97), Pmax (W) (r=0.84), Leg LBM(kg)) (r=0.81)  CP (W.kg-1) correlated with: V̇O2max (ml.min-1.kg-1) (r=0.85), Ppeak (W.kg-1) (r=0.89), Pmax (W.kg-1) (r=0.52) | 3-6CWR |

**Population**: M, healthy male; F, healthy female; ET, endurance trained; COPD, chronic obstructive pulmonary disease; (n), number of participants

**Correlation details:** MRTV̇O2, mean response time of V̇O2; V̇O2, time constant of V̇O2 kinetics; TT, time-trial; LBM, lean body mass; V̇O2max, maximal V̇O2 recorded following verification from additional trials >CP; Ppeak, highest power attained during an incremental exercise test; Pmax, highest power attained during a 30-s all-out exercise test

**CP determination method:** nCWR, number of constant work-rate trials; 3MT, 3-minute all-out test; nTT, number of time-trials