‘Fit for surgery’: the relationship between cardiorespiratory fitness and postoperative outcomes

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Abstract
Surgery accounts for 7.7% of all deaths globally and the number of procedures is increasing annually. A patient’s ‘fitness for surgery’ describes the ability to tolerate a physiological insult, fundamental to risk assessment and care planning. We have evolved as obligate aerobes that rely on oxygen ($O_2$). Systemic $O_2$ consumption can be measured via cardiopulmonary exercise testing (CPET) providing objective metrics of cardiorespiratory fitness (CRF). Impaired CRF is an independent risk factor for mortality and morbidity. The perioperative period is associated with increased $O_2$ demand, which if not met leads to $O_2$ deficit, the magnitude and duration of which dictates organ failure and ultimately death. CRF is by far the greatest modifiable risk factor and optimal exercise interventions are currently under investigation in patient prehabilitation programmes. However, current practice demonstrates potential for up to 60% of patients, who undergo preoperative CPET, to have their fitness incorrectly stratified. To optimise this work we must improve the detection of CRF and reduce potential for interpretive error that may misinform risk classification and subsequent patient care, better quantify risk by expressing the power of CRF to predict mortality and morbidity compared to traditional cardiovascular risk factors, and improve patient interventions with the capacity to further enhance vascular adaptation. Thus, a better understanding of CRF, used to determine fitness for surgery, will enable both clinicians and exercise physiologists to further refine patient care and management to improve survival.
1 | INTRODUCTION

Surgery is amongst the leading risk factors for mortality and has been estimated to account for 7.7% of all deaths globally (Nepogodiev et al., 2019). By 2030, it is estimated that one-fifth of people aged 75 years and older in the United Kingdom alone will undergo surgery (Fowler et al., 2019). Therefore, to better understand and mitigate this risk, we need to consider not just the disease or surgical procedure, but also the phenotypical response and ability to cope with the physiological insult posed by major surgery. Furthermore, prophylactic intervention targeting modifiable risk factors prior to surgery, a process known as ‘rehabilitation’, requires investigation and optimisation. This review explores our relationship with oxygen (O₂), the elixir of life, and how its transport and use in the human body determines ‘fitness for surgery’.

2 | ORIGIN OF O₂ AND OUR DEPENDENCY ON OXIDATIVE METABOLISM

When the solar system emerged 4.6 billion years ago (Dickerson, 1978), Earth’s atmosphere was devoid of O₂, a vast difference compared with the modern day atmospheric inspired fraction of 20.93%. The emergence of life, likely originating in alkaline thermal vents at the bottom of the oceans, initially gave rise to the domains of archaea and bacteria (Miller & Bada, 1988). Approximately 1.5 billion years ago, cyanobacteria began to release O₂ into the atmosphere (Nisbet & Sleep, 2001). The organic compounds that emerged from the ‘primordial soup’ were photosynthetic, capturing solar radiation and creating the organic molecule glucose. In turn, the O₂ released into the atmosphere signalled a major evolutionary event, arguably described by two oxidation ‘pulses’, the Great Oxidation Event and the Neoproterozoic Event, or as a progressive evolution, the Great Oxidation Transition (Lyons et al., 2014). This gave rise to atmospheric O₂ and the evolution of O₂-dependent organisms, from primitive eukaryotic unicellular structures performing metabolism, locomotion and reproduction to present day Homo sapiens.

Figure 1 describes the production of paleo-atmospheric O₂ and the entire dependency of the respiring mammalian cell for the constancy of electron flow, with molecular O₂ serving as the terminal electron acceptor in mitochondrial oxidative phosphorylation. Homo sapiens, like all mammals, has a remarkable ability to harness O₂, allowing a rapid turnover of adenosine triphosphate (ATP) and affording cells, tissue and organs a coordinated stasis sustaining life. Mammalian evolution has thus produced a structural, functional and physiological organisation that efficiently coordinates the convective delivery and diffusive uptake of O₂, essential for successful life.

3 | FROM MOUTH TO MITOCHONDRIA: CONVECTIVE AND DIFFUSIVE DETERMINANTS OF O₂ TRANSPORT

Early measurements describing O₂ uptake (V̇O₂) in humans at the onset of intense movement were conducted by Hill and Lupton (1923) and demonstrated a rapid and exponential response, as skeletal muscle has the capacity to increase rate of metabolism by an astounding 50- to 100-fold above its resting requirements. This challenges a rapid delivery of O₂ to the mitochondrial inner membrane for use as the terminal electron acceptor, whereby oxidative phosphorylation generates ATP. O₂ is transported by convection, which describes its movement within the airways and circulation-driven aero- and hydrostatic pressure gradients, and by diffusion, the passive movement down a concentration gradient such as between the alveolar compartment and pulmonary capillary bed and between the systemic microcirculation and tissue.

Figure 2 illustrates the major organs and processes, both convective and diffusive, that describe the ‘O₂ cascade’. Following inspiration of air into the lungs, O₂ diffuses down a concentration gradient at the alveolar–capillary membrane, minimally dissolves in plasma and predominantly binds with haemoglobin (Hb), an allosteric protein with affinity for four molecules of O₂. Deoxygenated venous blood is therefore saturated with O₂ in the pulmonary capillaries, the concentration of which is proportional to the concentration of Hb, its P₅₀ and the partial pressure exerted by O₂ on the plasma at a given temperature (Henry’s law). Oxygenated blood then travels the vascular system driven by the heart. This convective component is referred to as ‘O₂ delivery’ (QO₂), the product of cardiac output (Q) and arterial O₂ content (Q × CaO₂), and is complete when O₂ diffuses across the microcirculatory capillary beds and reaches the mitochondrial matrix where it is used as the terminal electron carrier. V̇O₂, as described by Fick’s principle is equal to the product of Q and the difference between arterial and mixed venous oxygen content (CaO₂ − Cvo₂). Notably, in health, the principal ‘rate limiting’ steps for maximal O₂ uptake (V̇O₂max) are attributed to the perfusive (QO₂) and diffusive components of the cascade (Wagner, 2000).

4 | METRICS AND MEANING: ASSESSMENT OF CARDIORESPIRATORY FITNESS

The advent of breath-by-breath measurement technology has allowed us to measure the capacity of the O₂ transport system and determine metrics describing the magnitude of cardiorespiratory fitness (CRF), which not only describes an individual’s ability to perform physical activity, but is linked to cardiovascular health (Ross et al., 2016) and
**New Findings**

- **What is the topic of this review?**
  The relationships and physiological mechanisms underlying the clinical benefits of cardiorespiratory fitness (CRF) in patients undergoing major intra-abdominal surgery.

- **What advances does it highlight?**
  Elevated CRF reduces postoperative morbidity/mortality, thus highlighting the importance of CRF as an independent risk factor. The vascular protection afforded by exercise prehabilitation can further improve surgical risk stratification and postoperative outcomes.

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...longevity (Blair et al., 1989). Cardiopulmonary exercise testing (CPET) is used to objectively measure the ability of a patient to uptake O₂ and typically involves an incremental exercise test to symptom-limited exhaustion. CPET can also identify underlying pathology and evaluate the impact of chronic comorbidities on O₂ uptake. Recently, the use of CPET has been widely adopted in patients prior to major surgery and approximately 30,000 tests are conducted annually in the UK alone (Reeves et al., 2018). These data are used to support patient care decisions, plan appropriate postoperative critical care, and direct prehabilitation programs aimed at improving CRF (Levett et al., 2018).

Three primary metrics describing CRF are typically reported when conducting CPET:

1. Peak oxygen consumption ($\dot{V}_{O_2}^{\text{peak}}$), defined as the $\dot{V}_{O_2}$ attained during an incremental test to exhaustion, expressed in absolute terms (ml min$^{-1}$) or relative to body mass (ml kg$^{-1}$ min$^{-1}$), which can be subject to allometric scaling, and measured as the highest value recorded, often occurring during the final 20 s of a test. Whilst $\dot{V}_{O_2}^{\text{peak}}$ is reflective of a patient’s ‘best effort’, it may not necessarily reflect a true highest value, defined as $\dot{V}_{O_2}^{\text{max}}$ with an observed plateau present in the $O_2$ uptake work-rate slope of Hill & Lupton (1923) demonstrated in Figure 3. Controversy exists here, and evidence suggests only a minority of continuous tests, even in young healthy people, yield a measurable plateau (Day et al., 2003; Poole & Jones, 2017). Nevertheless, an exercise test to exhaustion is important since it allows for the site of transport limitation across the O₂ cascade to be identified (Wagner, 2000).

2. Anaerobic threshold (AT), a submaximal index of CRF defined as the $\dot{V}_{O_2}$ above which anaerobic metabolism supplements oxidative phosphorylation with additional carbon dioxide (CO₂) production, creating a deflection point on a plot of pulmonary CO₂ output versus $O_2$ uptake (Figure 4). The AT is also commonly reported as a percentage of $\dot{V}_{O_2}^{\text{peak}}$ or $\dot{V}_{O_2}^{\text{max}}$. Whilst the AT signifies a transition where increased glycolysis raises the muscles lactate efflux into the blood above its removal rate with associated metabolic acidosis, a multitude of definitions and controversies exist (Poole et al., 2021). Thus in the context of preoperative CPET, AT refers to the gas exchange threshold (GET, sometimes also referred to as the ventilatory threshold), typically measured using the ‘gold standard’ V-slope (Beaver et al., 1986) method of determination. GET is expressed in ml kg$^{-1}$ min$^{-1}$ or ml min$^{-1}$.

3. The ventilatory equivalent for carbon dioxide ($V_{eq CO_2}$), defined as a ratio of minute ventilation to CO₂ production and usually reported at the GET. $V_{eq CO_2}$ reflects the composite efficiency

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**FIGURE 1** The production of oxygen (O₂), dependency of mitochondrial oxidative phosphorylation upon O₂, and the evolution of Homo sapiens to support O₂ delivery. Adapted from Bailey (2019)
Ventilation
Alveolar-capillary diffusion
Circulatory perfusion
Microvascular diffusion

**FIGURE 2** The oxygen (O₂) transport system characterised by pulmonary ventilation, alveolar-capillary diffusion, circulatory perfusion driven by the cardiovascular system and diffusion across the microcirculatory capillary beds. The volume of O₂ transport, described by Fick’s principle, is determined by the product of convective (cardiac output) and diffusive O₂ transport terms (and is the product of cardiac output and the difference between the O₂ content of arterial and venous blood).

**FIGURE 3** Schematic representation of O₂ consumption at the limit of exercise tolerance during CPET. (a) \( \dot{V}O_2^{\text{peak}} \) reported as the highest value recorded. (b) \( \dot{V}O_2^{\text{max}} \) idealised as a true highest value with observed plateau present. \( \dot{V}O_2^{\text{peak}} \), peak oxygen consumption; \( \dot{V}O_2^{\text{max}} \), maximal oxygen uptake

of the ventilatory response, including the breathing pattern and adaptive changes in pulmonary gas exchange in response to exercise. Elevated values for \( V_{eqCO2} \) occur in heart failure, respiratory disease and pulmonary hypertension (ATS/ACCP, 2003; Snowden et al., 2010; Sun et al., 2001) consequent to diminished perfusion, ventilation-perfusion mismatching, or diffusion limitation and changes in breathing pattern, which increase dead space ventilation.

5 | CRF AND SURGERY: LINK TO SURVIVAL

Mortality following major surgery is a significant risk despite progress being made in surgical technologies, anaesthesia and peri-operative care. In colorectal surgery, mortality is reported at 3.2% within 90 days (NBOCA, 2017) with complication rates above 30% (Lucas & Pawlik, 2014). Similarly, in-hospital mortality for elective abdominal aortic aneurysm (AAA) repair is 2.9% for open repair and 0.4% for endovascular repair (VSQI, 2017). Furthermore, the insult of major colorectal surgery has been shown to reduce CRF by ~40%, with hospital stays of 7–9 days, and only 50% of patients regaining pre-operative CRF levels after 6 months (Jensen et al., 2011).

Accurate prediction of surgical risk is required to facilitate shared decision making, improve patient outcomes and plan peri-operative care. Traditionally, subjective clinical acumen alone was used; however, objective scoring systems are available including the Portsmouth Physiological and Operative Severity Score for the Enumeration of Mortality and Morbidity (P-POSSUM; Whiteley et al., 1996), American Society of Anaesthesiologists (ASA) physical status, Charlson Comorbidity Index, and measures of cardiac function (Moyes et al., 2013). These systems are generally weak, and complementary ‘biomarkers’ are needed. CRF, a modifiable risk factor, has long been (inversely) associated with all-cause mortality (Kokkinos et al., 2010; Mandsager et al., 2018), and evidence also suggests that impaired CRF
MECHANISTIC LINK BETWEEN CRF AND POSTOPERATIVE OUTCOME

The model presented (Figure 5) presumes the existence of an obligatory baseline level of CRF (such as the threshold values for $\dot{V}O_2\text{peak}$, GET or $V_{eqCO_2}$ found in Table 1) to survive an increased demand for $O_2$ during the perioperative period. If the patient is unable to meet this presumed $O_2$ demand, chronic hypoxaemia and limited $\dot{Q}_O_2$ may be responsible for increased morbidity and mortality for any severity of disease. Whilst a detailed mechanistic understanding explaining why impaired CRF is associated with poor postoperative outcome remains to be elucidated, the presence of an $O_2$ deficit during the perioperative period is fundamental to this model.

The surgical stress response is characterised by an increased $O_2$ demand as demonstrated by Ciaffoni et al. (2016), measured directly (via in-airway sensors) beginning in the intraoperative period (Figure 6). The underlying mechanisms responsible for the perioperative elevation in $\dot{V}O_2$ can be explained by complex changes in metabolic demand. These comprise hormonal, haematological and immunological changes, manifest by increased $\dot{Q}$ and $O_2$ consumption as the delivery of nutrient and $O_2$-rich blood supports energy processes, tissue repair and protein synthesis (Gillis & Wischmeyer, 2019). Shoemaker et al. (1988) also describe a substantial increase in $O_2$ demand, from an average of $110 \text{ ml min}^{-1} \text{ m}^{-2}$ at rest to...
170 ml m⁻¹ m⁻² following major surgery, consequent to the strong systemic inflammatory response. Thus, a patient with greater $V_{O2}$ reserve may help mitigate this cardiovascular burden.

Surgery, is also known to result in oxidative stress with consequent increases in free radical formation (Arsalani-Zadeh et al., 2011; Bailey et al., 2006, 2007). This is particularly prominent during abdominal surgery given the potential for ischemia–reperfusion, leukocyte activation, mitochondrial dysfunction and concurrent depletion of antioxidants in the postoperative period due to increased consumption due to increased consumption of oxygen species (ROS) in the perioperative period is also important. Ciaffoni et al. (2016) also demonstrated concurrent elevation of CO₂ production during the intraoperative period, which may be equally important in terms of 'clearance' for the maintenance of normal acid–base balance (Bailey et al., 2017) and requires further mechanistic investigation.

The contribution of anaesthesia to the production of reactive oxygen species (ROS) in the perioperative period is also important. O₂ is one of the most used drugs in anaesthetic practice. Reducing cellular hypoxia is a clinical priority, and thus the sickest patients who are likeliest to suffer the adverse consequences of hyperoxia and ROS formation are the likeliest to receive supplemental O₂. There is a good case for accepting lower levels of arterial oxygenation to minimise ROS damage in the perioperative period. However, because of genetic variability in susceptibility to damage by ROS it is impossible to

### Table 1

| Author                   | Patients (n) | $V_{O2,peak}$ risk threshold (ml kg⁻¹ min⁻¹) | GET risk threshold (ml O₂ kg⁻¹ min⁻¹) | $V_{eqCO2}$ risk threshold | Risk thresholds defined/adopted | Postoperative outcome |
|--------------------------|--------------|---------------------------------------------|-------------------------------------|---------------------------|--------------------------------|----------------------|
| **Intra-abdominal surgery** |              |                                             |                                     |                           |                                |                      |
| Older et al. (1993)      | 187          | Not measured                                | Yes < 11.0                          | Not measured              | Adopted                        | Hospital mortality   |
| Older et al. (1999)      | 548          | Not measured                                | Yes < 11.0                          | No                        | Adopted                        | Hospital mortality   |
| Wilson et al. (2010)     | 847          | Not measured                                | Yes < 10.9                          | Yes > 34                  | Adopted                        | Mortality 90 days    |
| Snowden et al. (2010)    | 116          | No                                          | Yes < 10.1                          | No                        | Defined                        | Morbidity: comp      |
| **Vascular AAA surgery** |              |                                             |                                     |                           |                                |                      |
| Carlisle and Swart (2007)| 130          | Yes                                         | Yes                                 | Yes > 42                  | Defined                        | Mortality: 2 years   |
| Hartley et al. (2012)    | 415          | Yes < 15.0                                  | Yes < 10.2                          | Yes > 42                  | Adopted                        | Mortality: 30 days, 90 days |
| Prentis et al. (2012)    | 185          | No                                          | Yes < 10.0                          | No                        | Defined                        | Morbidity: LoS       |
| Goodyear et al. (2013)   | 188          | Not measured                                | Yes < 11.0                          | Not measured              | Adopted                        | Mortality: 30 days, LoS |
| Grant et al. (2015)      | 506          | Yes < 15.0                                  | Yes < 10.2                          | Yes > 42                  | Adopted                        | Mortality: 3 years   |
| Rose et al. (2018a)      | 124          | Yes < 13.1                                  | No                                  | Yes ≥34                   | Defined                        | Mortality: 2 years   |
| **Colorectal surgery**   |              |                                             |                                     |                           |                                |                      |
| Lai et al. (2013)        | 269          | Not measured                                | Yes < 11.0                          | Not measured              | Adopted                        | Mortality: 2 years, LoS |
| West et al. (2014b)      | 136          | Yes < 16.7                                  | Yes < 10.1                          | Yes > 32                  | Defined                        | Morbidity: comp      |
| West et al. (2014a)      | 105          | Yes < 18.6                                  | Yes < 10.6                          | No                        | Defined                        | Morbidity: comp      |
| Wilson et al. (2019)     | 1375         | Not measured                                | No                                  | Yes > 39                  | Defined                        | Mortality: 90 days   |
| **Upper gastrointestinal surgery** |          |                                             |                                     |                           |                                |                      |
| McCullough et al. (2006) | 109          | Yes < 15.8                                  | No                                  | No                        | Defined                        | Morbidity: comp      |
| Nagamatsu et al. (2001)  | 91           | Yes < 800 ml                                | Yes                                 | Not measured              | Defined                        | Morbidity: comp      |
| Moyes et al. (2013)      | 108          | No                                          | Yes < 9.0                           | No                        | Defined                        | Morbidity: comp      |
| Patel et al. (2019)      | 120          | Yes < 17.0                                  | No                                  | No                        | Defined                        | Morbidity: comp      |

Risk thresholds relate to a level of CRF below which an inferior postoperative outcome has been observed and are either defined from the respective study data or have been adopted from other studies and applied to the study data. Abbreviations: AAA, (open) abdominal aortic aneurysm; comp, complications; GET, gas exchange threshold; LoS, hospital length of stay; $V_{eqCO2}$, ventilatory equivalent for carbon dioxide; $V_{O2,peak}$, peak oxygen consumption.
Potential mechanisms that enhance survival

Figure 6  Pulmonary O2 uptake during the intraoperative period of a patient undergoing abdominal aortic aneurysm repair, taken from Ciaffoni et al. (2016). Events are represented by knife to skin (A); reduction in ventilator driving pressure (B); aortic clamp applied (C); fall in blood pressure (D); metaraminol (fast-acting α-agonist) bolus; infusion rate increased from 2 to 5 ml h−1 (E); sequential removal of iliac artery clamps (F, G); increase in ventilator driving pressure (H); and removal of superior retractor restricting rib cage movement (I).

predict which patients would be most vulnerable and when. Furthermore, some anaesthetics (e.g., ketamine) have been shown to interfere with mitochondrial function, promote dismutase activity and affect ROS handling, albeit in animal studies (Venâncio et al., 2015).

The additional demand for O2 is not solely constrained to the intraoperative period. Shoemaker et al. (1992) measured VO2 in 253 high-risk patients (defined by criteria with a >30% surgical mortality rate) before, during and immediately after major surgery. These values were compared with the estimated VO2 requirements of the patients (using resting preoperative control values) to calculate the magnitude of O2 deficit. Patients who died (n = 64) had organ failure and a mean O2 deficit of 33.2 l m−2, compared with 21.6 l m−2 for survivors with organ failure (n = 31), and 9.2 l m−2 for survivors without organ failure or major complications (n = 158). These findings highlight the clinical significance of the cumulative O2 deficit across the perioperative period and corresponding implications for development of organ failure and ultimately death (Figure 7). Furthermore, the authors also investigated the time course and types of emerging complications up to 10 days following surgery as illustrated in Figure 8. Interestingly, the recovery ‘slopes’ of the O2 deficit in Figure 8 are much the same between survivors (with organ failure) and non-survivors, and just the intraoperative and early postoperative magnitude is greater, which may suggest this to be the more critical component.

7  |  Potential mechanisms that enhance survival

Whilst mechanistic bases explaining the link between (elevated) CRF and postoperative outcome require further elucidation, evidence demonstrates that patients with low CRF are associated with poor postoperative outcome, likely explained by the prevailing magnitude of perioperative O2 deficit. Importantly, CRF is a modifiable risk factor and a primary component of prehabilitation strategies (Macmillan, 2019; Tew et al., 2018). Prehabilitation represents an opportunity to improve patient preparation for surgery and is multi-modal in nature comprising exercise training and improving nutritional and psychological status (Scheede-Bergdahl et al., 2019). Prehabilitation aims to improve patient CRF to better tolerate the surgical stress response, leading to a reduced risk of perioperative complications and improved postoperative outcome (Tew et al., 2018). The theoretical potential for this strategy is illustrated in Figure 9.

Few studies have investigated the potential to improve CRF prior to surgery using exercise interventions and those that have mainly comprise small sample sizes demonstrating proof of principle (Rose et al., 2020; Simonsen et al., 2020). West et al. (2015) deployed an exercise intervention in patients following neoadjuvant chemoradiotherapy prior to surgery. The intervention group comprised 22 patients with 17 controls who followed a high intensity interval training (HIIT) protocol, three times per week for 6 weeks. Following neoadjuvant chemoradiotherapy, VO2 at GET was significantly reduced by a mean of 1.9 ml kg−1 min−1. Conversely, 6 weeks of subsequent HIIT sessions increased O2 uptake at GET by 2.1 ml kg−1 min−1, whereas it did not change in the controls. In a systematic review, Loughney et al. (2016) concluded that preoperative exercise interventions are safe and feasible, yet there are insufficient controlled trials to draw reliable conclusions about their efficacy and feasibility. Recently, clinical guidelines and recommendations for preoperative exercise training in patients awaiting major non-cardiac surgery have been published (Tew et al., 2018). However, it is again acknowledged that further research is needed to identify the optimal exercise prescription in different clinical scenarios, particularly in the short preoperative time frame encountered in urgent cancer surgery.

While interest lies in preoperative exercise training, clear translational evidence to improved postoperative outcomes is yet to be established, with studies by West et al. (2015) underpowered for this endpoint. The most current systematic review (of 22 studies) with meta-analysis claimed that whilst prehabilitation improved preoperative functional capacity (measured by 6-min walk distance, albeit unlike West et al. (2015) objective measures of CRF including VO2peak and GET were not improved) and substantially reduced hospital stay, it did not reduce postoperative complications, 30-day hospital readmissions or postoperative mortality (Waterland et al., 2021). These findings need to be considered cautiously given the small sample sizes, heterogeneity of exercise interventions, limited reporting of objective measures of CRF, and lack of consensus on standardised endpoints of included studies.

Clearly, there is a requirement for a higher quality of evidence from large, randomised control trials, and clinical trials are ongoing with results awaited. Examples include: Van Rooijen et al. (2019), an international multicentre multimodal prehabilitation intervention including exercise, nutrition and psychological coping strategies.
within an enhanced recovery after surgery (ERAS) protocol (Trial ID NTR5947); a comparison of hospital-based supervised exercise, supported home-based exercise versus usual care to investigate patient recovery after bowel cancer surgery (PREPARE-ABC, 2020; Trial ID ISRCTN82233115); and Wessex Fit-4 Cancer Surgery (Southampton University, 2020) investigating the effectiveness of a community-based structured responsive exercise training programme with or without psychological support (Trial ID NCT03509428).

From a mechanistic perspective, similarities exist between the physiological insult of surgery and the acute response to an exercise stimulus. Primarily, an increased cellular demand for \( \dot{V}_O_2 \), consequent to oxidative phosphorylation required to regenerate ATP, is required to enable continued physical activity. As a chronic adaptive response to exercise, an improved ability to increase \( \dot{V}_O_2 \) is associated with elevated mRNA of peroxisome proliferator-activated receptor \( \gamma \) coactivator 1-\( \alpha \) (Gibala et al., 2009), a moderator of skeletal muscle mitochondrial biogenesis. An increase in citrate synthase (a marker of muscle oxidative capacity) has also been reported (Burgomaster et al., 2005), and an increase in oxidative stress (Bailey et al., 2010, 2018; Davies et al., 1982; Radák et al., 1999), which is attenuated following exercise training (Fatouros et al., 2004).

The mechanisms of this exercise-induced response have been linked to improvements in total antioxidant capacity (Fatouros et al., 2004; Radák et al., 1999), which is considered a marker of the body’s defence system to neutralise excessive and deleterious free radical and associated ROS formation (Ghiselli et al., 2000). Total antioxidant capacity has been enhanced following exercise training in both animal (Liu et al., 2000) and human (Fatouros et al., 2004) models. However, whether the long-term exercise-induced increase in total antioxidant capacity, and thus reduction in oxidative stress, is a key factor in improving postoperative outcomes remains to be elucidated. Exercise training been associated not only with a reduction in oxidative stress, but also with improved vascular function and consequent \( O_2 \) transport (Wray et al., 2011). Systemic and cerebrovascular function has
FIGURE 9  The fundamental principle underlying exercise prehabilitation whereby CRF is improved prior to surgery, thus reducing the risk of postoperative complications, and enhancing recovery as indicted by the green plot. Adapted from Clegg et al. (2013). The dashed line represents the cut-off between independent (ward-based care) and dependent (high dependency unit, intensive care unit) patient recovery.

been shown to improve following HIIT (Calverley et al., 2020; Molmen-Hansen et al., 2012), the potential consequence of an ‘optimised’ blood flow-shear phenotype, triggering calcium influx into the hyperpolarised endothelial cells (Cooke et al., 1991) upregulating endothelial nitric oxide synthase (Bolduc et al., 2013).

8  |  OPTIMISING RISK QUANTIFICATION AND PATIENT MANAGEMENT

The evidence reviewed suggests that impaired CRF is both an independent and a modifiable risk factor associated with postoperative outcome. Yet the strength of this relationship, used to predict postoperative outcome, is not effectively compared against traditional cardiovascular risk factors such as ischaemic heart disease, lung disease, or diabetes and obesity. This comparison has been addressed epidemiologically for all-cause deaths (outside of the surgical setting) within the Aerobics Centre Longitudinal Study, in which low CRF was found to be a greater risk factor than hypertension, smoking, high cholesterol, diabetes and obesity (Blair, 2009).

Attributable fractions describe the percentage of deaths that would not occur if a risk factor were removed from a population and account for both the risk of mortality associated with that condition and its prevalence in the population, as illustrated in Figure 10. This approach could be conducted in the surgical setting to help optimise risk quantification and further highlight the clinical importance of CRF relative to traditional risk factors.

Like all biomarkers, CRF is a dynamic metric subject to natural variation and thus needs to be interpreted with caution. Such variation encompasses both analytical and biological components which can be described using the concept of critical difference, indicative of the magnitude of variation around a true homeostatic point at any given time. Rose et al. (2018b) introduced the concept of critical difference to preoperative CPET and found differences of ±19%, 13% and 10% for \( \dot{V}_{O_2} \) -GET, \( \dot{V}_{O_2}{peak} \) and \( V_{eqCO2} \) -GET. The translational impact upon patient fitness stratification in their study demonstrated that up to 60% of patients were of indeterminate fitness, where for example, they could not be sure that a patient had a ‘true’ GET < 11 ml O\(_2\) kg\(^{-1}\) min\(^{-1}\) when variation was accounted for. A revised stratification model was formulated using zones along a spectrum of fitness; thus, clinicians are advised to look beyond a single cut-point and instead advocate a dynamic range of CPET values indicative of surgical risk (Wilson, 2018).

Furthermore, whilst inter-observer agreement, using intra-class correlation coefficient (ICC), for numerical values of GET (ICC 0.83 (0.75–0.90)) and \( \dot{V}_{O_2}{peak} \) (ICC 0.88 (0.84–0.92)) indicating good to excellent relative reliability (Abbott et al., 2018), inter-observer agreement regarding whether or not a reportable value existed was less consistent. This suggests that guidance for identification of reportable values could be improved.

Patient stratification should be optimised using the most effective metrics of CRF, with accompanying threshold values, which are indicative of risk specific to patient populations and surgical procedures. Table 1 highlights that many studies, including the seminal work of Older et al. (1993), have simply adopted threshold values developed by other studies sometimes using different patient populations and surgical procedures. Furthermore, CRF is commonly described using \( \dot{V}_{O_2}{peak} \), GET or \( V_{eqCO2} \) as discussed; however, alternative metrics may provide superior prognostic utility in some...
settings. For example, if a patient is unable or unwilling to exercise to exhaustion, a submaximal measure of CRF relating O2 consumption to workload achieved, such as the O2 uptake efficiency slope (OUES; Hollenberg & Tager, 2000; Bongers et al., 2017), may be more effective.

Female inclusion rate in peer-reviewed publications of perioperative CPET is reported at only 31% and may have a bearing on the interpretation of data (Thomas et al., 2020). Surprisingly, despite evidence that CRF is lower in females across the lifespan, given smaller body size, skeletal muscle mass, peak cardiac output and Hb concentration (Jackson et al., 2009; Fleg et al., 2005), sex is not considered during surgical risk stratification. If a simple dose-response relationship exists between low CRF and postoperative survival, we would expect females to be at increased risk given these congenital constraints. Furthermore, other risk factors such as cardiovascular disease (CVD), which may vary between the sexes, require investigation to appraise a potential compensatory effect for CRF and consequent changes in its prognostic potential on postoperative outcome.

9 | CONCLUSION

The current review has explored the intimate relationship between O2 transport and postoperative outcome, emphasising how preoperative CRF is an independent risk factor for postoperative mortality and morbidity, when patients undergo major intra-abdominal surgery. There is increased O2 demand during the perioperative period and patients must meet this demand to avoid tissue hypoxia, the presence and magnitude of which dictates postoperative morbidity and mortality. This relationship can be used to assess patient risk, plan perioperative care and optimise patient management using exercise as a modifiable intervention. However, there is a clear need to improve the physiological detection and interpretation of CRF, better quantify risk to specific populations, sex and surgical procedure, and better understand the optimal management of patients including the mode of exercise intervention and its timing. Collectively, a better understanding of CRF used to determine fitness for surgery will enable clinicians and physiologists alike to direct patient care more effectively and ultimately improve survival.

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COMPETING INTERESTS

D.M.B. is Chair of the Life Sciences Working Group and member of the Human Spaceflight and Exploration Science Advisory Committee to the European Space Agency and member of the Space Exploration Advisory Committee to the UK Space Agency.

AUTHOR CONTRIBUTIONS

All authors have read and approved the final version of this manuscript and 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. All persons designated as authors qualify for authorship, and all those who qualify for authorship are listed.

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