Review of early development of near-infrared spectroscopy and recent advancement of studies on muscle oxygenation and oxidative metabolism

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Abstract
Near-infrared spectroscopy (NIRS) has become an increasingly valuable tool to monitor tissue oxygenation (Toxy) in vivo. Observations of changes in the absorption of light with Toxy have been recognized as early as 1876, leading to a milestone NIRS paper by Jöbsis in 1977. Changes in the absorption and scattering of light in the 700–850-nm range has been successfully used to evaluate Toxy. The most practical devices use continuous-wave light providing relative values of Toxy. Phase-modulated or pulsed light can monitor both absorption and scattering providing more accurate signals. NIRS provides excellent time resolution (~ 10 Hz), and multiple source–detector pairs can be used to provide low-resolution imaging. NIRS has been applied to a wide range of populations. Continued development of NIRS devices in terms of lower cost, better detection of both absorption and scattering, and smaller size will lead to a promising future for NIRS studies.

Keywords Muscle · Oximetry · Tissue oxygenation · Oxidative metabolism · Exercise

Introduction
This article addresses the use of in vivo near-infrared spectroscopy (NIRS) for evaluating muscle oxygenation and oxidative metabolism, with a focus on the historical development of the method. There are a series of excellent review papers on the use of NIRS to study skeletal muscle [1–8]. There is also a recent article on principles, insights, and potential pitfalls of the noninvasive determination of muscle oxidative capacity by near-infrared spectroscopy [9] with corresponding commentaries [10]. NIRS has also been applied to study other tissues such as the brain [11, 12], but this topic will not be addressed in this review. NIRS has been used to study muscle oxygen levels in athletes, control subjects and patients with chronic illnesses or injuries [13, 14]. Given the increasing use of NIRS to study skeletal muscle, the purpose of this review is to guide future studies by providing: (1) descriptions of early studies that developed the instrumentation and methodologies that has led to current NIRS measurements, (2) The basic principles behind the NIRS measurements along with key limitations of the various NIRS based methodologies, and (3) a summary with selected examples of current the NIRS approaches used to understand the biochemistry and physiology of skeletal muscle oxidative metabolism.

Early studies with in vitro and in vivo oximetry leading to the use of NIR for the evaluation of skeletal muscle

In 1876, Karl von Vierordt observed by eye (visible light) spectral changes of hemoglobin (Hb) in trans-illuminated human fingers as well as in solutions containing Hb [15]. When circulation to the finger was occluded, the oxygenated Hb (oxy-Hb) bands disappeared and the deoxygenated Hb (deoxy-Hb) bands appeared, demonstrating the potential for qualitative oximetry, defined as a noninvasive method for monitoring tissue oxygenation. This was followed up a half century later with the building of reliable devices to perform in vitro spectrophotometry utilizing
visible light (400–650 nm) by Drabkin and Austin in 1932 [16], and Millikan in 1933 [17]. Millikan invented a colorimeter, which had been conventionally operated by the eye of the observer requiring subjective judgements and the potential for eye strain. The device objectively measured the degree of oxygenation of dilute Hb solutions using a differential copper copper-oxide photoelectric cell and two color filters. Later, it is known that Millikan developed the first portable ear oximeter using red and NIR light for monitoring black-outs of the pilots at extreme high altitude, which provided a prototype of a clinically useful oximeter [18]. In 1970s, Aoyagi at Nihon Kohden, a Japanese company, attempted to measure cardiac output using dye dilution methods along with a commercially available ear oximeter [18]. Initially, he found that light transmitted through the earlobe exhibited pulsatile variations that interfered with measurements of cardiac output. By balancing the red and infrared signals to cancel the pulsatile signal variations, he was able to accurately measure cardiac output using the kinetics of the dye washout. In addition, he successfully monitored the pulsating changes in the light transmission through the ear to measure arterial oxygen saturation. At the same time, a researcher with Minolta, another Japanese company, was conducting similar experiments and applied for a patent soon after. These experiments led to the marketing of a successful pulse oximeter by Minolta around 1978 [18].

Professor Frans F. Jöbsis of Duke University is regarded as the pioneer of medical applications of NIRS [19]. Before his 1977 milestone article, reflectance spectrophotometry and surface fluorescence using visible light were primarily used for investigating large solid organs such as the brain. Jöbsis extended NIRS research to other tissues, including skeletal muscle. He learned about optical monitoring techniques as a postdoctoral fellow in Britton Chance’s laboratory at the University of Pennsylvania. Using ultraviolet and visible regions, he attempted to study the redox state of the cytochrome c oxidase (cyt c) to understand the behavior of the mitochondrial respiration [20]. As the photons in the ultraviolet and visible regions do not travel deep into the tissue, surgical exposure of tissue was needed. In open-skull animal setups, he had attempted to study the heme a component, the absorption peak of which appears in the orange-colored region at about 605 nm and the heme a3, in the violet region at about 445 nm. Jöbsis was puzzled by the observation that cyt c in skeletal muscle was less reduced than other tissues, and less reduced than cyt c obtained from isolated mitochondria. The disparity in the oxidation/reduction (redox) states of the respiratory chain between skeletal muscle and other tissues motivated him to study the in vivo behavior of cyt c in skeletal muscle. Using NIRS on skeletal muscle, he found that the copper atom associated with heme a3 did not respond to anoxia, and therefore may be reduced under normoxic conditions; whereas, the heme-a copper was at least partially reducible [19].

Unfortunately, the observation that Hb absorbs more intensely than cytochromes in the violet region made these measurements more difficult in intact organs under normal circulation conditions. In this sense, Jöbsis noticed that the NIR region was more appropriate to study the relative absorption strength of Hb and cyt c. He has presented his rationale for deciding to adopt NIR light for tissue monitoring as follows [21]. Briefly, he enjoyed a dinner with his family, the menu of which featured a grilled chuck roast on December 28, 1976. The very American cut of beef still contains part of the shoulder blade of the steer; a flat piece of bone perhaps 3- or 4-mm thick, about the same as the human skull. He held the pink object up against the light and noticed that the shadow of a finger could easily be noted in the diffuse red light coming through the bone. If the red light could, then certainly NIR light at the longer wavelengths would penetrate the human skull and provide access to the brain. In addition, it was possible that other tissues could also be monitored in a minimally invasive way [21].

In 1977, in Science [19], Jöbsis reported Hb oxygenation, total-Hb, and cyt c under hypoxic conditions in the exposed heart and in the brain without surgical intervention. He demonstrated using animal preparations that oxygen sufficiency for cyt c can be recorded using NIR effectively. He also extended the experiment to the human head and successfully monitored the decrease in total-Hb in the brain by the hypocapnea induced by voluntary hyperventilation. Later, simultaneous measurements were made of both tissue oxygen tension (PtO2) and the redox ratio of cyt c from rat cerebral cortex, in situ. These studies showed that decreased PtO2 was accompanied by cyt c reduction [22]. Hoshi et al. developed a new approach for measuring the redox state of cyt c in the brain under normal blood-circulation conditions in rats [23]. When fractional inspired oxygen was decreased in a stepwise manner from 100 to < 10%, at which point the concentration of oxy-Hb [oxy-Hb] decreased by approximately 60%, cyt c started to reduce. Increases in arterial PO2 (PaO2) under hypoxic conditions caused an increase in [oxy-Hb], whereas further oxidation of cyt c was not observed. The dissociation of the responses of Hb and cyt c was also clearly observed after the injection of epinephrine under severely hypoxic conditions; that is, cyt c was re-oxidized with increasing blood pressure, whereas Hb oxygenation was not changed. Using newly developed time-resolved NIR (NIR_TRS). Chance reported that the intensity profile of photon migration in tissues permits determination of the path length and thus concentration changes in oxy-Hb/myoglobin (Mb) in the resting and ischemic muscle model [24]. NIR_TRS emits short light pulses (100 ps) and counts photons, which are scattered and absorbed in the tissue. The distribution of the photons migrated follows the
photon-diffusion equation and the mean light path length and tissue optical properties can be determined by solving the equation [24].

**Basic principles for and operation of in vivo NIRS**

The NIR (700–2000 nm) shows much less scattering, and thus better penetration into biological tissue than visible light (360–700 nm). However, light absorption by water progressively increases at wavelengths above 900 nm, which limits the light penetration into tissue above these wavelengths in larger water- and lipid-containing biological tissue (a peak at around 976 nm and 927 nm, respectively). Thus, NIRS measurements usually adopt wavelengths in the range of 700–850 nm [19]. The major absorbing compounds of this wavelength region are intravascular Hb, intramuscular Mb, and mitochondrial cyt c. NIRS measurements (for example, 760 nm and 850 nm in this case) rely on O₂-dependent absorption changes that occur in the heme in the brain, muscle, liver, etc. (Fig. 1). There are several types of NIRS devices: NIR single-distance continuous wave spectroscopy (NIRSDCWS), multi-distance continuous wave spectroscopy (NIRMDCWS), NIRTRS, phase modulation spectroscopy (NIRPMS) (Fig. 2), diffuse correlation spectroscopy (NIRDNS), and diffuse reflectance spectroscopy (NIRDNS) [25]. NIRDCS uses coherent NIR light to penetrate deep tissues and measures speckle fluctuations of the diffuse light, which are sensitive to the motions of red blood cells in tissues [8]. NIRDNS, a new rapidly developing technique, can continuously measure blood flow in the superficial muscles. NIRDNS methodology uses the unique approach of monitoring muscle blood flow (mBF) by measuring the optical phase shift caused by moving blood cells. While NIRTRS and NIRPMS are more accurate due to their ability to monitor changes in both absorption and scattering, they are limited in practicality by higher costs and complexity. This leaves the continuous wavelength devices (NIRSDCWS and NIRMDCWS) as affordable alternatives for measuring changes in oxygen levels in tissues, even if these devices cannot account for changes in scattering.

To measure wider areas of the limb and image regional differences in skeletal muscle oxygenation, continuous wavelength multi-channel or imaging devices were built and applied to exercise physiology [5, 26–30]. NIRDNS would be suitable to low-cost, user-friendly wearable/wireless 2D imagers in different pathophysiological conditions and in sports sciences [13]. NIRDNS can be used to reconstruct images of the internal distribution of optical absorption and scattering coefficients. However, NIRDNS requires physically accurate model, which is parameterized by the spatial distribution of scattering and absorption properties in the media [25]. The properties of the model need to be adjusted iteratively until the predicted measurements from

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![Fig. 1](image1.png)

*Fig. 1* Changes in absorbance (optical density) in accordance to a wavelength. The optical density (absorption) increases at 760 nm when the oxygenated hemoglobin (O₂Hb) is deoxygenated (HHb). When blood volume increases, the line shifts to the upper (increase in absorbance) Copyright © 2017 Willingham and McCully from Ref. [14]

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![Fig. 2](image2.png)

*Fig. 2* Schematic illustration of main types of near-infrared (NIR) spectroscopy instrumentation. **A** NIR continuous wave spectroscopy with single-distance (NIRSDCWS) or multi-distance (NIRMDCWS). **B** NIR time-resolved spectroscopy (NIRTRS). **C** NIR phase modulation spectroscopy (NIRPMS).
the forward model match the physical measurements from the device [25].

To continuously monitor muscle oxygenation during untethered human locomotion in the field or a clinical setup, small portable NIRS devices have been built [31]. For example, a wireless NIRMDCWS system which is the size of a cell phone was developed by Artinis Medical Systems (the Netherlands) [32], which is one of the first commercially available wireless and portable devices using 2-channel and 6-laser diodes as a light source. A portable NIRMDCWS device has been developed by Astem Inc. (Japan), which consists of an LED light source, two photodiodes, microprocessors, and a Bluetooth networking module (less than 100 g) [33]. An American company has developed a small, inexpensive NIRS device (Moxy) for sports applications [34], which adopts Monte Carlo modeling and specifically measures Hb oxygen saturation (SO2) with 1-channel and 4-LEDs as a light source. Another American company has developed an even less expensive NIRS device for the athletic market using portable NIRMDCWS [35].

The pathlength of the light varies due to optical characteristics of tissue such as muscle, adipose tissue, blood volume, and motion artifact created by muscle contractions. NIRSDCWS can only provide the relative values of tissue oxygenation due to undeterminable optical path and its length, but is commercially available because of inexpensive and portable instrumentation. To calculate the relative changes in oxy-Hb/Mb, deoxy-Hb/Mb, and total-Hb/Mb, the equation of a 2-, or multiple-wavelength method can be applied according to the modified Beer–Lambert law [36]. Because of the difficulty in quantifying NIRSDCWS, muscle oxy-Hb/ Mb, deoxy-Hb/Mb, and total-Hb/Mb are usually expressed in the following way:

\[
\text{Arbitrary units [OD],} \\
\mu M \times \text{cm, or} \\
\mu M \times (\text{DPF} \times \text{source-detector spacing}),
\]

where OD, optical density; DPF, differential pathlength factor defined by pathlength in terms of fold of the light source to detector distance.

It has been reported in a few studies that changes in pathlength were less than 10% during and after the end of arterial occlusion and during exercise [37–39]. DPF is determined in the thigh muscle and decreased slightly (around 5%), but significantly from baseline to peak cycle exercise [38]. There is still not an enough data available on whether and how much optical pathlength changes during varying interventions, such as arterial occlusion, muscular contractions, and recovery from hyperemia. The absolute optical path length can be measured by NIRTRS and NIRPMS [40]. NIRDCS and NIRDRS have been developed for measuring changes in muscle oxygenation and mBF, and are able to compute muscle oxygen consumption (mVO2) [41]. The similar absorption spectra of Hb and Mb make it difficult to differentiate the two by optical properties alone. What we know of the differences between the two molecules is based on 1proton-magnetic resonance spectroscopy (1H-MRS) using the water suppression to separate their deoxy forms [42]. Studies using 1H-MRS have suggested <10% Hb and 90% Mb contribution to the overall NIR signals [43]. Computer modeling has quantified the Mb contribution to the NIRS signal in human muscle to be 85–95% of the total signal [44]. Attempts have been made to use wavelength shift analysis of the NIRS signal from muscle tissue to determine relative concentrations of Hb and Mb [45]. Wavelength shift analysis found that Hb accounts for only 20% of the overall signal in human muscle, and the position of the NIRS deoxy-heme peak at 760 nm is linear with Hb/Mb concentration ratio. A recent study has examined the question of the NIRS signal origin by measuring simultaneously the 1H-MRS, 31phosphorus (31P)-MRS, and NIRS signals in finger flexor muscles during the transition from rest to contraction, recovery, ischemia, and reperfusion. The experiment results support a predominant Mb contribution to the NIRS signal from muscle [46]. Most experimental analysis often referred a study by Seiyama et al. [47], which reported on the comparative NIRS observation with and without a blood substitute. With a blood substitute, the NIRS signal decreases precipitously, which leads to the supposition that Hb contributes predominantly to the NIRS signal. Yet, recent studies with buffer perfused hindlimb and with a proper physiologically monitored conditions show that NIRS can still detect a robust signal [48, 49]. Additional studies will be needed to clarify not only the contribution of Mb/Hb to the NIRS signal, but also the relative kinetics of Mb and Hb deoxygenation during a variety of interventions under different conditions.

When using reflected light, that is the path of light from source and detector pair which is sent orthogonally into the tissue, the pattern of the continuous light path follows a “banana shaped” curve. The penetration depth into the tissue is approximately equal to half the distance between the light source and the detector [50]. In a study using NIRTRS [51], the mean depth of the light penetration was found to be greater (approximately 20 mm at a 30-mm optode separation) than half of the emitter–detector separation reported in many previous studies [48], mainly due to the time gating for photon counting designed for selectively collecting photons in a deeper tissue. The volume of tissue being evaluated using continuous light and 3-cm separation distance was estimated to be ~4 cm³ in the milk model by Chance et al. [52], when it is assumed that the volume of interest be a rotation body of the ellipse revolved across 180° or a hemisphere (Fig. 3a). If a banana-like pattern is assumed rather than a hemisphere [53], this volume might be overestimated.
The reason why NIR_CWS cannot determine the absolute value of oxygenation using reflected light is the unknown path length of NIR light in the tissue, which is generally assumed to be a banana shape as discussed above. The presence of skin and adipose tissue (subcutaneous adipose tissue thickness: SATT) above the muscle will increase penetration depth, as a result of the reduced absorption of light due to the reduced concentrations of heme in the SATT. Conversely, increased blood volume in the skeletal muscle will increase light absorption and reduce penetration depth. Melanin in the skin also absorbs light in the NIR region, such that darker skin can reduce reflected light intensity and penetration depth [54].

Absolute values from NIRS can be measured using NIRTRS [24, 25, 39] and NIRPMS [55–58]. NIRTRS emits short light pulses (100 ps) and counts photons in a several cm apart from the light emission on the skin surface. The emitted photons are scattered and absorbed in the tissue and arrive at the detectors with a varying timing, which depends on the path of each photon travels. The photons that travel shallower arrive earlier, while those that travel deeper arrive later. The distribution of the photons migrated follows the photon-diffusion equation and the light path length, absorption coefficient, and reduced scattering coefficient can be computed by solving the equation. NIRPMS, a frequency-domain method, usually uses intensity-modulated light at a radio frequency from 50 MHz to 1 GHz and monitors the migrated light intensity (DC), amplitude (AC), and phase shift, which comprises the time of the light travelled in the tissue. Intensity-modulated light propagates through tissue with a coherent front, forming photon-density wave. The detected photon-density wave is delayed because of the phase velocity of the wave being altered by the optical characteristics of the tissue. In a study, it is suggested that the absolute scattering and absorption coefficients can be accurately determined by the combination of the phase shift and the DC with relationships provided by diffusion theory [58].

There are several studies which used NIRTRS for monitoring muscle deoxygenation during resting arterial occlusion and exercise [38, 59–61]. Although NIRTRS can quantitatively measure muscle oxygenation, the values are still influenced by the SATT and tissue heterogeneity. The reason is that the measured oxygenation/deoxygenation values would be diluted by the interference of the lower basal Hb/Mb concentration and/or lower metabolic rates in the SATT [1] (Fig. 4a). The relationship between the slope ($S$) of change in [deoxy(Hb/Mb)] and SATT (mm) was examined and attempted to be expressed the following equation: $S = 12.73 - 1.49 \cdot \text{SATT}$, which might be case sensitive for each experiment (Fig. 4b) [62]. However, this technique does not always overcome the accurate quantification of tissue oxygenation [62]. Thus, a further research needs to be warranted to quantify tissue oxygenation using NIRTRS.
metabolism, including peripheral and cardiorespiratory measurements. Peripheral measurements include tissue $O_2$ microelectrodes, Mb $O_2$ saturation by spectrophotometric analysis [65], and NADH analysis from exposed tissue surfaces [66]. In an alternative approach to evaluating muscle oxidative metabolism, MRS has been developed to measure in vivo active forms of high-energy phosphorus metabolites and intramuscular pH [67, 68]. Since these studies, MRS has evolved into the “gold standard” for noninvasive detection of skeletal muscle bioenergetics. The limitation to $^{31}$P-MRS is that it is relatively expensive option to standard $^1$H-MRS systems and is not readily available. In addition, changes in high-energy phosphorus metabolites are influenced by both the delivery of oxygen as well as oxidative capacity, making the interpretation of the results somewhat imprecise [69]. In comparison to MRS methodology, the strength of NIRS for measuring skeletal muscle oxidative metabolism is that it is relatively inexpensive and portable, making it far more assessable. NIRS devices can make biochemical measurements at frequent intervals on even frail or vulnerable populations [70], and can be employed in both laboratory- and field-based studies. The ability to collect data during human locomotion is a major reason NIRS lends itself to the study of exercise and athletic performance.

Acceptable approaches for the in vivo muscle NIRS evaluation

A number of studies have reported the validity of NIRS-measured oxy-Hb/Mb and deoxy-Hb/Mb signals in animals and humans under steady-state conditions by comparing
these signals with venous blood [31, 39, 71–73]. Thus, it is generally accepted that NIRS-oxygenation/deoxygenation signal has a considerable agreement with the changes in venous saturation under varying oxygenation status of the human muscles. However, there have been a few studies that have failed to validate NIRS measurements especially under non-steady-state conditions [74, 75]. A possible explanation for the discrepancies found in these studies is that the NIRS signal contains information from arterioles, capillaries, venules, and intracellular Mb. The O$_2$ gradient from an arteriole to a venule is large in normoxic conditions such that variations in blood volume from arteriole to venule could alter the NIRS signal without change in venous oxygen signals [1]. The lower oxygen levels during hypoxic conditions would reduce this effect and produce a good agreement with values determined by NIRS and blood samples [1]. However, as it is suggested $< 10\%$ Hb and $90\%$ Mb contribution to the overall NIR signals [44, 46], we have to carefully consider the above-mentioned interpretation.

One of the challenges of measuring muscle oxygenation levels during exercise concerns comparing oxygen levels between individuals with different SATT values [76]. People with higher SATT values will have lower oxygen levels and lower metabolic rates. Appropriate and acceptable approaches for accounting for the effect of SATT values on muscle oxygenation are described as the following: (1) directly measure the absolute value in the absorption and scattering coefficient computing the optical path length and absolute Hb/Mb parameters using NIR$_{TRS}$ and NIR$_{PMS}$, (2) normalize signal using a physiological calibration by determining muscle O$_2$ store (0–100% level of oxygenation) and oxidative metabolic rate during arterial occlusion, (3) adjust oxygen levels using a calibration equation that includes measurements of SATT, and (4) perform kinetics measurements at the onset of or following muscle contractions or muscle contractions either voluntarily or by electrical stimulation. The kinetic measurements are acceptable because only the muscle tissue responds with increased metabolic activity and the kinetic measurements are reported with rate constants in units of time (or 1/time).

**Physiological calibration and oxygen consumption measurements by arterial occlusion**

In vivo NIR$_{SDCWS}$ provides relative changes in oxygen levels due to the inability to correct for the effects of scattering related to varying amounts of SATT and unknown path lengths of light [1]. A simple and common method for calibrating NIR$_{SDCWS}$ signals is “physiological calibration” to use the range of muscle deoxygenation caused by arterial occlusion followed by reactive hyperemia [50]. The arterial occlusion method is based on the assumptions that a period of ischemia will result in the complete disappearance of oxy-Hb/Mb in a measurement area, and that the reactive hyperemia after occlusion will almost completely eliminate deoxy-Hb/Mb. While oxy-Hb/Mb and deoxy-Hb/Mb in arbitrary units may vary between measurement sites and individuals, the arterial occlusion calibration adjusts for these changes making inter-individual or inter-occasion comparisons possible. The units of measurement for the NIRS device now become %Hb/MbO$_2$, and the values vary from 0 to 100%. This is in contrast to the units provided by the NIR$_{SDCWS}$ devices provided by Artinis, LTDs that report units in tissue oxygen index (TSI) values. TSI values are also presented in units of %, but because the TSI is a percentage of ratios of absorbance, the values typically have a maximal range of ~35 to ~75% as absorption at the two separate wavelengths can never reach zero even if the concentration of oxygenated or deoxygenated hemoglobin is zero. An important consideration when performing a physiological calibration is the duration of the ischemic period. Starting from resting conditions, 5–6 min of ischemia is sufficient to reach functionally zero oxygen levels in most humans [50, 77] because phosphocreatine (PCr) begins to decline after this amount of time. The use of a short duration of exercise prior to inflation of the cuff increases metabolic rate, and reduces the time to reach zero oxygen levels to 3–4 min [78]. Leaving the ischemic period long enough to confirm zero oxygen levels will also result in some depletion of PCr. The result of this is that there will be some post ischemia oxygen consumption. This oxygen consumption will slow the rate of return of oxygen levels compared to reactive hyperemia without an oxygen debt. If the investigator wants to use an ischemic period to induce and measure reactive hyperemia, then a balance must be considered between the need to confirm zero oxygen levels to obtain the physiological calibration, and the need to avoid phosphocreatine depletion and post ischemic oxygen consumption delaying the recovery of oxygen levels. Quantitative calibration of NIR$_{SDCWS}$ signal is possible in a combination with MRS measurement by applying a 15-min period of ischemia to the muscles (Fig. 5) [77]. The rate of decline of muscle oxy-Hb/Mb during ischemia can be compared with that of muscle phosphocreatine (PCr) in mM per second or a conversion to mM O$_2$ per second. Interestingly, muscle metabolic rate appears to remain constant in the presence or absence of oxygen. As a result, this method provides quantitative values of both muscle oxygen stores and mVO$_2$.

Evaluation of muscle energy metabolism using NIRS is difficult because the measured oxygenation levels do not specifically reflect mVO$_2$, rather they reflect the balance between muscle DO$_2$ and mVO$_2$. To distinguish mVO$_2$ from DO$_2$ using NIRS, two approaches have been used; the transient arterial occlusion method and the venous occlusion method. The transient occlusion uses 10–30 s of arterial occlusion provided by a pneumatic tourniquet. With no
blood entering or leaving the tissue of interest, changes in NIRS-measured oxygen levels now reflect $\text{mVO}_2$ [76, 77, 79–82]. Resting $\text{mVO}_2$ of young health males determined by NIRSDCW was found to be a small amount of variability (23.0 ± 1.2%/min) [70], and to be consistent between studies by different investigators [83]. NIRTRS has also been used to measure resting $\text{mVO}_2$, providing results in absolute units (0.82 $\mu$M s$^{-1}$) [39]. In the former study, a significant correlation was found between NIRSDCW measured $\text{mVO}_2$ and MRS measured PCr ($r^2 = 0.99$, $p < 0.01$), and ADP ($r^2 = 0.98$, $p < 0.01$) concentrations. The linear relationships between the NIRS and MRS measured indicators supports both the thermodynamic [84, 85] and the kinetic [86] regulation models of muscle mitochondrial respiration. There is an intimate relationship between NIRS-measured $\text{mVO}_2$ using the transient arterial occlusion method and the rate of PCr recovery, a biochemical process of ATP resynthesis via oxidative phosphorylation after muscle contractions [82]. These studies suggest that the initial rate of muscle deoxygenation during transient arterial occlusion is a direct measure of $\text{mVO}_2$. Venous occlusion has also been used with NIRS [87]. The change in NIRS signals with patent arteries and venous occlusion is similar to venous plethysmography [88, 89]. In this approach, venous occlusion can provide measurements of muscle blood flow.

Repeated transient arterial occlusions after exercise can provide successive $\text{mVO}_2$ values, and the recovery of $\text{mVO}_2$ values after exercise provides information that is basically same as that determined from the kinetics of PCr levels after exercise [90, 91]. The recovery kinetics of PCr has been developed as a method of measuring mitochondrial capacity [69, 84, 92–94]. Thus, time constant (Tc) or rate constant $k$ (1/Tc) for $\text{mVO}_2$ recovery is an indicator for evaluating muscle oxidative capacity (Fig. 6) [14, 95]. The advantages of the NIRS method of monitoring the kinetics of $\text{mVO}_2$ compared to the $^{31}$P-MRS measuring the kinetics of PCr are that in addition to the lower cost, greater availability, and greater ease of use, recovery of $\text{mVO}_2$ can be made independent of blood flow limitations while recovery of PCr is not. The measurements of $\text{mVO}_2$ are made during transient ischemia and thus do not depend on oxygen delivery. However, caution must be taken to make sure that oxygen levels are high enough in the tissue to assure that $\text{mVO}_2$ is not oxygen limited [96].

**Sensitivity adjustment by measuring SATT**

Muscle NIR signal intensity is greatly influenced by overlying SATT [1, 97, 98]. SATT values can vary from 3 mm over the forearm muscles of lean subjects, to greater than 20 mm over the vastus lateralis muscles of obese subjects. In addition to increasing penetration depth, increasing SATT also reduces the signal intensity coming from the deeper tissues, assumed to be skeletal muscle. For example, a SATT thickness of 5 mm reduces the signal intensity by approximately 20% with a light source–detector separation being 30–40 mm. The use of shorter separation distances, 15 or 20 mm with a 5-mm fat thickness, attenuates the signal intensity by 30 and 60%, respectively [26]. Perhaps of more importance, SATT values greater than 10 mm attenuate the signal from the deeper muscle by over 90% [98, 99], making the study of people with obesity very difficult. A correction equation based on optical properties has been used to adjust values of absolute StO2 based on SATT values [100]. SATT independently confounds NIRSDCW-derived StO2 by overestimating actual skeletal muscle oxygenation and by decreasing the magnitude of exercise-induced changes in oxygen levels. Several available NIRS units use the multiple source–detector pairs to separate out signals primarily from skeletal muscle from signals coming from SATT [39, 101, 102].

**Kinetics parameter measurements**

There has been a lot of research interest in the study of onset and offset kinetics of muscle oxidative metabolism during exercise [103, 104]. Muscle oxygenation determined by NIRS has been used to provide relevant information on onset and offset kinetics [46]. Early studies measured changes deoxy-Hb/Mb signals during ramp cycling exercise

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**Fig. 5** Changes in phosphocreatine (PCr) and oxygenation in human forearms during 15 min of arterial occlusion measured by near-infrared and $^{31}$P-magnetic resonance spectroscopy. No significant changes were found in pH or ATP throughout arterial occlusion. Copyright © The American Physiological Society. Reproduced by permission of the publisher. Adopted from Ref. [77]
to differentiate trained cyclists from physically inactive subjects [105]. The rate of deoxygenation at the onset of exercise [72, 106] recovery time of muscle reoxygenation after submaximal to maximal exercise [41, 107–112] and the rate of reoxygenation after brief high-intensity exercise [112] are among indicators for evaluating muscle oxidative capacity. These studies have reported a good agreement between faster PCr recovery kinetics and faster oxygenation kinetics measured with NIRS [113]. A different outcome was obtained after maximal short-duration isometric exercise, where higher oxidative capacity muscle (faster PCr kinetics) was inversely related to the rate of muscle reoxygenation after the exercise. [114]. The result of this study was attributed to the hypothesis that muscle reoxygenation rate after this type of short high-intensity exercise may be influenced more by VO₂ than by DO₂, when O₂ demand is still high and O₂ supply is not fully activated. In a study, a new method was proposed to noninvasively approximate muscle capillary blood flow kinetics from the kinetics of the primary component of pulmonary O₂ uptake and deoxy-Hb/Mb in humans during exercise [101].

Muscle blood flow has been measured using the kinetic changes in oxygen levels after reactive hyperemia [115]. As mentioned earlier, if muscle oxygen levels are reduced to near zero by 5–6 min of ischemia, without significantly lowering PCr levels, then the return of oxygen levels with reactive hyperemia reflects the indicator wash-in methods used to measure muscle blood flow. Several studies have used these approaches by measuring the rate of recovery of oxygen saturation with NIRS to evaluate blood flow in the calf muscles of people with peripheral arterial disease [114, 115].
Conclusions and perspectives

Observations of changes in the absorption of light with oxygen levels have been recognized as early as 1876. Gradual advances lead to a milestone paper by Jöbsis in 1977 illuminating the potential of NIRS to study tissue oxygen levels. Subsequent studies have shown NIRS to be useful for the in vivo evaluation of changes in muscle oxygenation and oxidative metabolism during these 40 years. The most practical and inexpensive devices use continuous wave light, which requires assumptions related to changes in scattering to work successfully. More expensive devices using phase-modulated or pulsed light can monitor both absorption and scattering, and can provide more accurate signals under a wider range of conditions. Compared to other imaging methods, NIRS provides excellent time resolution, and multiple source–detector pairs can be used to provide low-resolution imaging. To image whole body activity, we might invent NIRS device embedded in exercise clothing, which can be also used outside the laboratory with the help of energy harvesting technologies (solar batteries, light source with sunlight and band pass filters). NIRS device will be used in combination with magnetic resonance imaging (MRI), MRS, electromyogram (EMG), respiratory gas analysis, etc. However, along with applied clinical studies, basic research is still needed on topics such as the origin of the NIRS signals, the NIR penetration depth or measurement area in tissue including the effect of non-muscular tissue, changes in optical properties during wide range of oxygenation status, varying subjects, and exercise modality. Continued development of NIRS devices in terms of lower cost, better detection of both absorption and scattering, and smaller size will lead to a promising future for NIRS studies.

Author contributions TH wrote about early development of near-infrared spectroscopy and methodological section and organized throughout the manuscript. KKMC wrote about application of near-infrared spectroscopy to sports and clinical science.

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Compliance with ethical standards

Conflict of interest Takafulmi Hamaoka declares that he has no conflict of interest. Kevin K. McCully is the President of Infrared Rx, Inc, and NIRS software company.

Ethical approval All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants included in the study.

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