Critical oxygen delivery, the microcirculation and cardiac surgery: What we know now and need to know!

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Introduction

Critical oxygen delivery ($DO_{2crit}$) is a key physiologic parameter for organ and organism survival. It defines physiologically the boundary of shock (1,2). The purpose of cardiopulmonary bypass is to support that physiologic variable ($DO_{2crit}$), keep the patient out of shock, while the heart and lungs are removed from the normal circulation during repair. An understanding of $DO_{2crit}$ is paramount to understanding shock and being able to treat it appropriately. Although many in cardiac surgery think they understand $O_2$ delivery, the process is complex and few understand the ramifications/limitations of $DO_{2crit}$. CPB has at times been compared to other shock states, but every effort is made to assure specific organ demands (brain, heart, kidney and intestine). The microcirculation has been under intense study in the last 10-15 years in highly specialized physiology laboratories. With new techniques we are learning a great deal about how $O_2$ fluxes through the complexity of arterioles, venules and capillaries that together constitute the microcirculation. Although wonderful animal models of haemorrhagic shock, and haemodilution have been studied, the data we have gained is dramatically limited. Almost all of the studies to date have been performed in rodents and most often in striated muscle. We know a good deal about endothelial cells, vaso-regulation, blood cell interactions with endothelial cells, the glycocalyx and these observations are intriguing. Unfortunately we make assumptions that what we see in striated muscle might well translate to brain, heart or kidney microcirculation. However that may not always be true. Some data and hypotheses generated from these experiments could well explain some of the phenomenon that we encounter in CPB.

$DO_{2crit}$

The physiologic definition of shock is a state in which there is an inadequacy of $O_2$ delivery, in relation to $O_2$ demand (1-3). $O_2$ delivery is maintained in a surplus state. All of us are aware of the $O_2$ content equation based upon haemoglobin ($Hgb$) level, saturation and the amount of dissolved $O_2$ in plasma (Table 1). That equation utilizes a constant (1.36) multiplied by the measured haemoglobin ($Hgb$) level. One should point out that not all $Hgb$ binds $O_2$ at the usual curve. This becomes important when a pharmaceutical company tries to design a haemoglobin.

Table 1

<table>
<thead>
<tr>
<th>Equation</th>
<th>Description</th>
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<tr>
<td>$CaO_2 \ (oxygen \ content) = (1.36 \times Hgb_{con} \times Hgb_{sat}) + (0.0031 \times O_2 \ PAO_2)$</td>
<td>Oxygen content equation based upon haemoglobin ($Hgb$) level and saturation.</td>
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<tr>
<td>$DO_2 \ (oxygen \ delivery) = CaO_2 \times CO$ (cardiac output or CPB flow)</td>
<td>Oxygen delivery equation.</td>
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<tr>
<td>$VO_2 = DO_2 \ (PcO_2 - P_{mitoO_2})$</td>
<td>The gradient of $O_2$ from erythrocyte ($Hgb$) to the mitochondria is driven by partial pressures within each sub-cellular region.</td>
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based oxygen carrier (HBOC) as a blood substitue or when Hgb is dramatically different (stored banked blood, sickle Hgb, foetal Hgb, etc.). The O₂ content equation also has a component for dissolved O₂ (0.0031 X PaO₂). In many clinical situations that dissolved O₂ is disregarded as an insignificant contributor to the total O₂ content. In severe anaemia, as well as in hyperbarics, the dissolved O₂ content may well be a major portion of the total O₂ content. It is dissolved O₂ that is the O₂ physiologically utilized for cellular metabolism.

O₂ content is important for O₂ delivery (Table 1). Cardiac output (cardiopulmonary bypass machine flow) is multiplied times the total O₂ content for an estimate of delivery. This is the classic teaching. What we have learned from the microcirculation in the last 10 years is that such calculated numbers may not reflect the actual delivery of O₂ to tissues. Calculated whole body numbers may not be real in terms of minute to minute biology. The microcirculation will auto-regulate its own O₂ delivery and extraction is based upon instantaneous tissue utilization, acid base levels and a number of other complex mechanisms. Hgb dissociation curves are dramatically manipulated through acid base equilibrium, chloride ion concentration, and 2,3 diphosphoglycerate (2,3 DPG) concentration. The production of 2,3 DPG is highly O₂ and energy dependent.

A very key and little recognized fact, is that in striated muscle the haematocrit (Hct) of blood in the capillary network is approximately 15% (4,5). Even if the aorta and large arterioles carry a haematocrit of 40% the precapillary sphincter cells along with a complex set of physics (micro-tubular rheology) allows that red cells cannot be stacked tighter in the capillaries than the 15% Hct (Figure1). We

**Figure 1: Erythrocytes traversing a capillary.** Note that they “stack” and appear to be touching. However, the cells are all viewed in an obtuse angle and they all flex/fold to fit through microcirculation. One can see the plasma gaps and appreciate the distance form the red cell membrane outer limit to the edge of the capillary. In the lower picture, at a slightly greater magnification one should appreciate the lining of the capillary. This area is made up of the glycocalyx and represents a highly complex glycosaminoglycan surface which holds a number of key molecules that have anti-coagulation and anti-inflammation function. The thickness of the glycocalyx can be measure using these microvascular techniques. Photomicrograph care of Ivo Torres-Filho, MD, PhD, VCURES microcirculation laboratory.
do not know if in other key tissues such as heart, kidney or brain whether this 15% Hct limit exist as well.

If cardiac output (CPB flow) drops then total calculated O₂ delivery will drop as well. The compensatory event that occurs in the capillaries will be that O₂ extraction rises to meet tissue O₂ demands. Eventually if either systemic or local flow drops enough, or if anaemia is so bad (less than 15-20% Hct) then a level of critical O₂ is encountered. For the vast majority of our lives all of our tissues exist with a luxury O₂ delivery and this is known as flow independent O₂ delivery (Figure 2). At the point at which O₂ extraction has hit its limits or if anaemia is so severe (<15% Hct) then flow dependent O₂ extraction occurs (Figure 2). As one approaches and exceeds that interesting physiologic point a number of key events happen.

This inflection point is known as the point of critical O₂ delivery or DO₂crit (1-3). When an animal or a tissue is to the left, on the curve, of DO₂crit it is by definition in anaerobic metabolism. The length of time spent to the left of DO₂crit has a direct correlation to survival. With anaerobic metabolism cells do not die immediately. They begin to create metabolic acids, lactate being the most important. The mitochondria shift their ATP production to anaerobic biochemistry but the other cellular organelles involved with DNA/RNA transcription cease to work and protein synthesis drops off dramatically. As this imbalance continues, cellular ion pumps, particularly calcium flows change eventually leading to other side effects such as cellular, mitochondrial oedema and eventually apoptosis. But it all begins to be dysfunctional with the point of DO₂crit. So it is that the study of DO₂crit ought

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**Figure 2: Supply independent and supply (flow dependent) oxygen delivery.** Note that as there is a drop in oxygen delivery there is an increase in oxygen extraction The point of DO₂crit is the definition of when shock occurs. The more time spent to the left of DO₂crit affects tissue, organ and whole body survival. Left of DO₂crit can be thought of as “the killing zone”.

VO₂

[Oxygen delivery dependence]

[Lactate]

Accumulative O₂ debt

O₂ extraction ratio

Critical DO₂

Venous O₂ sat

Oxygen delivery DO₂
To further understand DO$_{2 crit}$ by way of illustration, if a mountain climber ascends the Himalayas he will increase his/her Hct leading to increased O$_2$ carrying capacity. This is in direct response to a lowered partial pressure of O$_2$ increasing erythropoietin form the kidney. The cardiac output will increase also but factors such as viscosity and fluid losses will lead to a point of diminishing enhanced cardiac output. Eventually viscosity and cardiac output increases reach their maximum. Once the climber enters a level above 24,000 feet the PaO$_2$ is so reduced that all human physiology will be forced to the left of DO$_{2 crit}$. This is known in the climbing world as “killing zone”. The entire body is constantly anaerobic in an oxygen debt. Of interest total O$_2$ carrying capacity has dramatically increased but the dissolved O$_2$ content has dropped so low that all the increased carrying capacity cannot make up for the reduced dissolved O$_2$ and diffusivity of O$_2$ combined with viscosity overtake the compensatory mechanisms.

**O$_2$ debt**

Oxygen debt is a concept not often understood (nor even studied) in cardiac surgery. O$_2$ debt is the total amount of O$_2$ (quantity of O$_2$ per cc tissue volume X time) not delivered to an organism or tissue (Table 1) (6,7). In trauma and haemorrhagic shock it is well understood that the amount of O$_2$ debt again correlates with survival or death (1-3, 8-10). The amount of O$_2$ debt is directly related to the amount of left shift beyond DO$_{2 crit}$ and the length of time spent in that physiologic (shock) state. A rough way to estimate this is to look at lactate levels and even rising potassium levels (a result of tissue ion leakage).

Those animals that are successfully resuscitated after haemorrhage will regain a normal blood pressure, cardiac output, and oxygen carrying capacity. But, if they had an existing O$_2$ debt it may well be that all the vital signs appear normal (BP, hear rate, etc) yet repayment of O$_2$ debt is not complete. O$_2$ debt cannot be measured in our usual operating rooms or ICUs. To measure O$_2$ debt a metabolic measurement must be precisely made of O$_2$ uptake and CO$_2$ production. When O$_2$ uptake does not meet the demands set by CO$_2$ production O$_2$ debt is occurring. We as yet do not know the sub-cellular mechanisms of repair that go on with repayment of O$_2$ debt but it probably is at least the restoration of ATP stores as well as the recreation/repair of dysfunctional or destroyed cellular protein machinery. In O$_2$ debt it may well be that a trauma victim could be 1-3 litres behind in O$_2$ delivery (1-3). Generally, more than 3 litres behind in O$_2$ delivery will mean certain death. To replete that amount of debt may take hours, depending upon O$_2$ delivery. Those patients who can repay their O$_2$ debt within 60-90 minutes often survive whereas those that cannot repay O$_2$ debt by 4 hours or more will almost certainly die. No one has ever investigated cardiac surgery patients in terms of DO$_{2 crit}$ and O$_2$ debt.

**O$_2$ flux**

The microcirculation is where the “rubber meets the road” in terms of tissue O$_2$ delivery. The microcirculation is a complex, highly dynamic, redundant network of arterioles, capillaries and venules. Flow is not constant through all vascular channels at all times. Erythrocyte flow stops and starts depending upon tissue demands. Many channels cannot be seen with routine trans-illumination microscopy if there are no red cells within the lumens. We do know that at some times plasma flows through channels either devoid of erythrocytes or at different flow rates than the erythrocytes are moving. O$_2$ flows from all vascular channels out to the tissues (Figure 3).

That fact cannot be overemphasized. It is not just capillaries that interact in the delivery of O$_2$ to cells. Arterioles and venules contribute to O$_2$ delivery but generally there is a net-
work dependent upon one or more feeder arterioles. Any cell cannot survive if it is further than 40-50 microns from a vascular O₂ source. If arteries and venules are in juxtaposition they actually transfer O₂ between them and venules can be very active in the delivery of O₂ to tissues. At any given time, in most tissue only about 30% of capillaries are open and flowing at one time. This fact allows for increased O₂ demand to be supplied by a regulated mechanism of delivery. Unfortunately most of what we know regarding the microcirculation is from striated muscle with assumptions made to other tissue. Again, we know relatively little about flow in the microcirculation during CPB.

O₂ moves from haemoglobin in red cells into the surrounding plasma and from that plasma out to the tissues. Although such a process sounds easy, the route of an O₂ molecule leaving haemoglobin and entering a mitochondria or onto myoglobin is difficult (11-13). O₂ is poorly soluble in water. Plasma is essentially water with some proteins, hormones and of course cellular elements. Each red cell has approximately 300,000,000 molecules of Hgb and on each Hgb there are 4 O₂ molecules. Per cc of blood there are 4-5,000,000 red blood cells. It would therefore seem that the amount of available O₂, no matter what the demand would be massively in excess. However Hgb binds O₂ very tightly. We now understand that the movement of O₂ from Hgb to target sites is dependent upon the erythrocyte acting as a localized super charger of dissolved O₂. Remember it is the dissolved O₂ that is available for metabolic function. The larger the plasma gap from the surface of an erythrocyte the larger is the resistance to movement of O₂ (Figure 4). The erythrocyte functions with a corona of O₂ surrounding it and as one moves by Angstroms away from the cell membrane the partial pressure of O₂ drops. The way in which Hgb recharges the surrounding plasma is through the biochemistry of changed O₂ binding. The way the microcirculation auto-regulates O₂ supply to local tissue demand is by increasing red cell transit time and by increasing O₂ extraction ratio. Remember the system is limited by the 15% Hct physics of stacking red cells in capillaries. Of interest

**Figure 3:** The flux of O₂ from arterioles, venules and capillaries. One should realize that O₂ freely flows from levels of high partial pressure to areas of lower partial pressure. Hgb exerts a pull and push based upon relative saturation. Mitochondria pull O₂ based upon their needs but the movement of O₂ is based upon its solubility in water based (plasma) and lipids (membrane) as well as its binding and release from Hgb.
there is a fascinating network interaction that leads to countercurrent movement of \( O_2 \) from venules to capillaries and from arterioles to both other vessel types. So \( O_2 \) is in constant flux diffusing down gradients, but convectively carried by plasma and red cell movements.

All mammalian species (we do not know about reptiles and fish) have the same level of \( \text{DO}_2_{\text{crit}} \) in terms of Hct. That one observation should be contemplated for a bit, as it has profound implications. Whether you are a mouse, a rat, a pig, goat, chimp or human at or near 15% Hct flow independent oxygen delivery is maxed out, \( O_2 \) extraction ratio has hit its limit and lactate production begins (14-16). This means that at 3.5-4 gm/dl Hgb, no matter what else is done, shock will occur. Blood pressure may be preserved (although likely it will be depressed) and cardiac output is maximized but the cells somewhere in the organism will revert to anaerobic glycolysis and metabolic acid production will begin.

Even if everything else is done correctly with a level below 3.5-4 gm/dl Hgb \( O_2 \) debt is occurring. Therefore 3.5-4 gm/dl becomes a floor below which we cannot electively accept going especially at normothermia. No one knows the \( \text{DO}_2_{\text{crit}} \) at different levels of hypothermia, although \( O_2 \) usage drops about 4% per degree. Therefore during CPB, understanding \( \text{DO}_2_{\text{crit}} \), Hgb and microcirculation \( O_2 \) fluxes, a wide range of basic physiology could/should be studied. Not only does temperature change \( O_2 \) demand but it changes extraction ratio, oxy-Hgb curves, acid base etc.

In terms of transfusion, historically when first conceived in the early years of the 20\textsuperscript{th} century and blood banking was not yet viable, the trigger for transfusion was a level of Hgb between 3-5 gm/dl. This was the point at which cardiac failure and unacceptable deaths increased. Of note, in Jehovah’s Witnesses it is not until the levels of Hgb drop to around 5 gm/dl or below that death rates rise in data bases following outcomes both in cardiac surgery and other surgeries (17,18). Both of these facts seem to relate to the limit of the microcirculation to function at or near \( \text{DO}_2_{\text{crit}} \). In CPB we have long had the debate about what is the “best” Hgb or Hct to transfuse. Both measurements are surrogates for potential \( O_2 \) delivery. One day perhaps we can understand and talk in terms of \( \text{DO}_2_{\text{crit}} \) and study \( O_2 \) debt in CPB rather than such gross measurements as Hgb and Hct.

Measurements in experimental microcirculation work

Today the use of microvascular/microcirculation research techniques is moving from the highly instrumented animal research laboratory to the operating room. In the research laboratory, the standard has been trans-illumination intra-vital and confocal microscopy. These techniques use one of several standard
animal preps to view a representative piece of tissue left intact to its native circulation. Hamster cremaster muscle, Hamster cheek pouch, rat and other animal mesentery and rat spinotrapezius muscle preparations have all been utilized. Recently some exciting work using an imbedded plastic “window” in the rat skull has made it possible for surface microscopy investigations of brain blood flow (19-21). Work is underway to adapt such techniques to intact spinal cord blood flow as well (22).

From these preparations capillary density, vessel flow rates and vessel sizes can be measured off-line. Usually videos are captured of the vessels to be interrogated and then computer programmes are adapted for automated or semi-automated calculations of parameters. Cell types, erythrocytes, platelets and white cells can be distinguished. White cell rolling, sticking and diapedesis can be followed at a site of capillary or vessel interest. Work with laser injury has been able to create distinct lesions of endothelial cells to assess platelet adhesion, clot formation and anticoagulation pharmaceuticals. The lining of the endothelial cells with the glycosaminoglycans is available for study as well. It’s size can be measured using overlaid digital subtraction photomicroscopy. Using a number of molecular markers with immune-fluorescence, the presence, clearance and production of key endothelial cell products such as nitric oxide, hydrogen peroxide, endothelin etc. can be directly visually assessed. Furthermore, again with immune-fluorescent techniques individual endothelial cells can be seen to be healthy or undergoing apoptosis.

Vascular O$_2$ content, as well as tissue O$_2$ content can be directly measured in real time during microcirculation research. With the use of specific laser wavelengths of light a technique of phosphorescence quenching has been perfected. This technique uses a known amount of phosphorous attached to albumin. With the right laser light it gives a decay curve directly and inversely related to the partial pressure of O$_2$. Such techniques allow for assessments of vascular and tissue O$_2$ delivery in real time under any desired Hgb, Hct or shock (low BP, haemorrhage etc.) to be investigated. Phosphorescence quenching cannot be done in humans and neither can routine intra-vital microscopy. However about 10 years ago a new technique was commercially created, orthogonol polarization video microscopy (23-25). This technique allows for using polarized light at 550nm which is the wavelength reflected by Hgb. By using this technique and shining the device the orthogonol (90° reflected light) forms a picture or red blood cells flowing through the microcirculation. Video images can then be made of nail beds, oral mucosa and even rectal mucosa in humans during

Figure 5: Before and after retinal angiograms of a human after cardiopulmonary bypass. Note the changes in the microcirculation and the loss of capillary network. These are transient and are thought to be due to micro-air emboli. The retinal microcirculation is a direct reflection of the brain microcirculation.
any number of adverse physiologic conditions. Measurements of red cell velocity, red cell concentration, vascular diameter etc. can all be made from off line analysis. Work from our centre has created a technique using Raman spectroscopy (light scattering) at the right wavelength such that microvascular oxygen content and Hgb O₂ saturation can be read without touching the organ or organism. This means that in the future we should be able to get readings of tissue or even cellular O₂ amounts in humans without using phos- phorescence quenching.

Studies in CPB and microcirculation

The use of orthogonal polarization video analysis has led to some recent literature regarding the changes of the microcirculation during CPB (23-25). In a small study from Belgium 9 patients undergoing cardiac surgery were compared to 6 patients undergoing cardiac surgery without CPB and 7 patients undergoing thyroidectomy (complete controls) (25). At baseline, prior to surgery the percentage of perfused vessels was the same in all groups. When anaesthesia was induced the levels of vessel perfusion dropped to about 70% perfused. Of interest during CPB the levels of perfused vessels dropped to 53% and when patients concluded their surgery (on entrance to ICU) the perfusion had begun to come back. Those that underwent CPB had a lower perfusion than either thyroid patients who had normalized or non CPB cardiac patients (64%). Those who under went CPB still had only about 60% of vessels perfused even with normalized haemodynamics. The severity of obstructed vessels correlated with systemic lactate measurement. Others have confirmed the same thing that once CPB is begun there is a measurable decrease in microvascular flow index (26-28). In our research we have found that micro-air embolism is a universal event during CPB (29,30). Furthermore air embolism causes destruction of the glycocalyxx, up regulates white cell sticking and has effects upon hydrogen peroxide reperfusion injury of endothelial cells. Whether these mechanisms are important in routine CPB microcirculation events or are more rare situations we simply do not know.

In animal work with trans-illumination video microscopy the effects of some vasoconstrictors have been examined as well as basic mechanisms of CPB. In a study of small bowel microcirculation it was shown in rats on CPB that even if the haemodynamics were maintained in a normal range (stable and normal mean blood pressure) there was a decrease in functional capillary density, arterio-lar vasoconstriction, and blood velocity reduction. Increased leukocyte accumulation occurred with more sticking and rolling of leukocytes during CPB as well as an extravasation of albumin. These observations signal that endothelial cells and the glycocalyx are dysfunctional, but they have not been directly studied to date. The use of phenylephrine, vasopressin and other vasoconstrictors to enhance or normalize blood pressure appear to be particularly bad on the maintenance of microvascular perfusion, capillary density and O₂ delivery. Large blood vessel flow went up whereas small vessels (where O₂ is transferred) dropped. The endothelium is responsible for vasoconstriction/dilation, local blood flow, inflammatory mediation, coagulation mediation, vascular permeability and vascular growth/repair. Think about how many of these events we manipulate in cardiac surgery and how few of them we truly understand (31). The microcirculation is where blood and endothelium interact.

The future

The initial foray into microcirculation biology research with CPB is disturbing. Observations that flow in the key units for O₂ flux are decreased dramatically suggests that even though we do our best to support haemodynamics, the complexity of the microcirculation and the endothelial biology leads to a
disregulation of DO$_{2\text{crit}}$. Mechanisms for this can easily be suggested. They include the near universal micro emboli that occur with CPB, inflammatory events, changes in hormones, nitric oxide synthesized and perhaps many more. The fact that we use CPB in an attempt to maintain homeostasis and preserve organs during repair of the heart and lungs suggests that at best we are far from performing anything normal. This is not new news. However the widespread efforts by anaesthesiologists and perfusionists to maintain blood pressure in a normal range using infused vasoconstrictors again suggests that we are sailing in waters we know little about. The use of understanding DO$_{2\text{crit}}$ and O$_2$ debt coupled with advanced physiologic measurements of the microcirculation, endothelial blood interface will surely yield exciting results in the future. With these studies will come new models for testing pharmacologic interventions, new CPB techniques and strategies that should make CPB safer and improve outcomes.

References

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