Influence of hypoxia in inflammatory synovitis

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Hypothesis
Persistent synovitis results from a hypoxia driven progressive transition to glycolytic metabolism that promotes transcriptional changes permissive to unresolved inflammation as the disease progresses. Erosion of cartilage and bone may involve gene expression characteristic to the “anoxic response system” of wound repair.

HYPOXIA AND MODULATION OF IMMUNE RESPONSES
The pathogenesis of rheumatoid arthritis (RA) is hypothesised to involve inappropriate triggering of the MHC controlled immune surveillance, resulting in altered T and B cell profiles. Recently, the cytokine profile of T helper lymphocytes has been associated with the disease. The cytokine repertoire of inflamed synovia is categorised as that of a Th1 response. The characteristic Th1 response displays increased expression of select cytokines such as TGFβ, IFNγ, TNFα, IL1, and IL2. In contrast, Th2 responses exhibit increased IL4, IL5, IL10, and IL13 (predominantly anti-inflammatory cytokines). While the levels of TNFα and IL1 are high within RA synovium,1 the level of IL2 is paradoxically low.2 This is atypical of a Th1 response, but may be accounted for by the overriding hypoxic condition of the RA synovium.

Hypoxia transcriptionally upregulates TNF and IL1 but downregulates IL2.3 The predominance of a given cytokine in the microenvironment of the responding Th cell influences Th1/Th2 differentiation.4 This phenomenon is believed to be regulated by phosphorylation of the relevant signal transducers and activators of transcription (STATs), after cytokine binding to their associated receptors. Based on the knowledge that hypoxia can modulate both cytokine expression and phosphorylation events, we propose that within inflamed synovial tissues, hypoxia underlies the functional polarisation of the Th1/Th2 lymphocytes and the apparent Th2 to Th1 switch in RA. The T cell cytokine profile of inflamed synovia may account for the development of an “unorganised” or abnormal inflammatory response. However, it does not convincingly explain the persistence of the existing inflammatory reaction, or more distant events such as the development of pannus tissue and the erosion of cartilage and bone. Evidence of an ancillary mechanism to cytokine induced inflammatory changes comes from a study of RA patients with HIV. In this situation, even when terminal loss of T cells occurs and all immune and cytokine activity ceases (patients in remission) destructive rheumatoid pathology still continues.5

The apparent RA synovial T cell hyporesponsiveness (or anergy) may also be related to hypoxia. It is known that transient carbonyl-aminoo condensation, or Schiff base formation, regulates T cell and antigen presenting cell (APC) interactions.6 Inhibition of T cell responses can thus occur, depending on the species and concentration of Schiff base forming aldehydes. Such low molecular weight species are generated during hypoxic metabolism. We have demonstrated the presence of several low molecular mass carbonyl compounds, notably acetone and acetoacetate, in RA synovial fluid (SF)7 that can potentially lead to the observed synovial T cell anergy.

Other studies indicate that the cytokine and growth factor profile of the Type A (macrophage) and Type B (fibroblastic) synoviocytes in culture, match more closely with that displayed by rheumatoid synovial tissues.8 These observations reflect a dominant and persistent cytokine and growth factor influence by resident synovial cells rather than the T lymphocytes; in particular, macrophage or Type A derived IL1, IL6, TNFα and GM-CSF. The production of macrophage (Type A synoviocyte) derived cytokines is also influenced by hypoxia.9

We therefore propose that chronic hypoxia and anoxia are the most significant factors mediating persistent synovitis and bone erosions and should be considered as a direct target for therapeutic manipulation.

RHEUMATOID SYNOVIIUM IS CHRONICALLY HYPOXIC
The hypoxic nature of rheumatoid synovium was originally suggested on the basis of measurements of oxygen tension within the inflamed cavity.10 Other studies reporting raised carbon dioxide tension, lactate, lowered glucose and acidosis also supported these observations.11 More indirect, but supportive evidence was also provided by studies of the synovial oxidative metabolism, revealing increased metabolic demand and shift to glycolytic metabolism.12 These observations were confirmed with our studies of the synovial membrane and fluid from rheumatoid joints,
using nuclear magnetic resonance (NMR) spectroscopy. The results of this sensitive technique displayed a profile of low molecular weight metabolites consistent with hypoxic metabolism. Decreased synovial pH, raised lactate, 3-D-hydroxybuturate, acetate levels, and ketone body formation confirmed a progressive shift to glycolytic metabolism, in accordance with chronic hypoxia. Further evidence came from studies utilising a polarographic needle electrode, which directly measured Po2 levels in diseased synovium and verified the hypoxic nature of inflamed synovia. The most hypoxic regions were found to be the innermost synovium in contact with the synovial fluid, irrespective of effusions. Our morphometric analysis of inflamed synovial tissues also supported these observations and revealed structural features that explain chronic hypoxia.

Although synovitis was long considered to be an angiogenesis driven pathology, in our analyses, the capillary density was calculated to be a third of that in normal synovium. In addition, the average distance of the capillaries from the joint cavity was found to increase in rheumatoid synovia. These measurements indicate inadequate perfusion because of failure of angiogenesis to vascularise innermost synovium. We interpret this as the failure of angiogenesis to keep pace with synovial thickening and have discussed the implications in the context of inflammatory synovitis. Indeed, the prevalence of such events in inflamed rheumatoid synovium is exemplified by the presence of an avascular and predominantly hypoxic "pannus" tissue. We further argue that the loss of highly organised vascular structure causes relatively less uniform perfusion through the tissue. Subsequent poor blood flow aggravates hypoxia by diminishing the oxygen gradient out of the vessels and reduces the capillary release rate in small, remote capillaries. With sustained hypoxia, the capillaries become "paralysed" and lose their normal vasoactive responses. In addition to this hypoperfusion, mobility of the inflamed joint causes increased intra-articular pressure, further restricting synovial blood flow. Other hypoxia induced factors prevail. The presence of saturable, high affinity vascular endothelin (ET-1) binding sites within RA synovial sections and the localisation of ET-1 like immunoreactivity to synovial microvascular endothelial cells, we believe, is a further indication of a hypoxia mediated contribution to the reduction in local synovial perfusion.

Vascular insufficiency, hypoxia, and inflammatory cell infiltration leads to synovial fibroblastic hyperplasia with a resultant net increase in metabolic demand. The high metabolic demand predicts a predominantly glycolytic metabolism, a situation seen during wound repair in an essentially anoxic environment. Increased activities of enzymes such as glyceraldehyde-3-phosphate dehydrogenase, glucose-6-phosphate dehydrogenase, lactate dehydrogenase, and mitochondrial cytochrome oxidase within inflamed synovia demonstrate increased glycolytic activity. Increased intracellular but decreased extracellular sulphydryl content within the inflamed synovium is a strong indication of the disrupted redox balance. We suggest, this apparent redox imbalance in rheumatoid synovium is a manifestation of chronic hypoxia and movement induced, intermittent hypoxia/reperfusion.

CELLULAR RESPONSES TO HYPOXIA
A number of biochemical changes precede cell injury induced by hypoxia (reviewed by Khan and O’Brien). These changes are consequences of the inactivation of oxidative phosphorylation in the electron transport chain. A significant drop in the ATP levels follow increases in cytosolic calcium and sodium, phospholipase A2 activation, and membrane phospholipid degradation. Increased lactate/pyruvate ratios indicate a concomitant increase in the free cytoplasmic NADH/NAD+ ratios. Increased blood ketone body formation (acetoacetic/β-hydroxybutyric acid ratio) is also an indicator of mitochondrial NADH/NAD+ ratio and redox potential. Increased NADH levels thus infer a reducing environment in which cells are hypoxia sensitive. This sensitivity often represents a switch to genetic programmes that control appropriate responses to prolonged changes in intracellular oxygen tension. At the cellular level, the initial impact of hypoxia is on the regulation of intermediary metabolism, affecting genes encoding enzymes responsible for glucose transport, glycolysis, and gluconeogenesis, regulating the biochemical responses discussed above.

CELLULAR RESPONSES TO ANOXIA
Anderson and Stoler have coined the term "anoxic response system" to describe events that occur at extremely low oxygen tensions observed in early wound sites. This system has been devised during evolution for the purpose of wound debridement and entails a switch to glycolytic metabolism, release of cathepsins and endonucleases to remove damaged and dead tissue.

TISSUE RESPONSES TO HYPOXIA
At the tissue level, hypoxia induces coordinated activation of a network of genes that regulate processes leading to tissue reorganisation and regeneration. The resident cell types characterising the tissue, respond to hypoxia by a tightly controlled sequential process. The sequence of events begins with the "sensing" of the environmental change at the cellular level via signalling mechanisms involving the extracellular matrix and cell surface "sensor" molecules. The sensor then transduces the signal to the intracellular environment through second messenger systems that subsequently activate transcription factors (such as hypoxia inducible factor-1 (HIF-1)) and initiate specific transcriptional events. The nature and mechanism of activation of this apparently universal "hypoxia sensor" has been reviewed by Bunn and Poyton. It has also long been established that the sensor is a haemoflavoprotein.
HYPOXIA AND XANTHINE OXIDOREDUCTASE

Xanthine oxidoreductase, a three redox centred flavoprotein, generally believed to play a part in purine metabolism, is transcriptionally induced by hypoxia.9 The functional significance of this enzyme, specifically in synovitis, is still far from clear. However, several theories based on hypoxia/reperfusion injury, have attributed a pivotal role for this enzyme in pathology because of its capacity to generate reactive oxygen species. Recent observations relating to the signalling role of such ROS have indicated a “signal transducer” role for the enzyme, particularly in hypoxia/reperfusion related pathology.

The relevance of this enzyme in hypoxia driven pathology has been the subject of much debate since xanthine was considered to be the essential substrate for the enzyme mediated generation of ROS via reduction of molecular oxygen. It is now clear that the enzyme is also capable of using NADH as a substrate that facilitates the reduction of molecular oxygen to ROS.28,29 We discussed the evidence for the accumulation of NADH in hypoxic tissues. Under physiological conditions, the enzyme is believed to metabolise xanthine/hypoxanthine to urate with the aid of NAD+ as the electron acceptor. It follows therefore, that this redox active enzyme (having three intact redox centres) is capable of maintaining the redox balance via alternately using NAD+ as an electron acceptor and generating NADH as well as using NADH as a substrate during hypoxia, leading to ROS generation. The former mechanism feeds into the “salvage pathway” of purine metabolism, maintaining the nucleotide pool whereas the latter mechanism may act as a “signal transducer” for redox changes resulting from hypoxia. Such a versatile mechanism has previously been described for a bacterial Put A protein with a dual role in enzymatic metabolism as well as transcriptional control.30 Furthermore, we have recently reported that under conditions of hypoxia, this enzyme is also capable of generating nitric oxide from nitrite/nitrite,31 which presents the enzyme as a suitable candidate for being a “hypoxia sensor” with redox mediated signalling capacity.

How can xanthine oxidoreductase act as a “redox sensor” and modulate transcriptional events downstream? Firstly, the predominant endothelial localisation of the enzyme indicates a strategic locality for it to act as a sensor of any PO2 changes as well as related metabolic alterations, such as NADH/NAD+ redox balance. Secondly, the enzyme has specific high affinity for sulphated glycosaminoglycans (GAGs), integral structural components of extracellular matrix. The sulphation pattern of GAGs is particularly influenced by hypoxia.32 In this context, xanthine oxidoreductase is a putative sensor for environmental changes transduced via the extracellular matrix GAGs. The presence of high circulating levels of this enzyme in various conditions, including rheumatoid arthritis,34 may also indicate the GAG mediated “trapping” or “recruitment” of this putative sensor where it is required to transduce the signal from an extracellular to intracellular environment via ROS production.

How can hypoxia/redox activated XO and XO derived ROS modulate subsequent intracellular events? The activation of a number of specifically hypoxia controlled transcription factors, such as Hif-1,9 AP-1, and NF-kB, are all modulated by ROS driven redox events.35,36 The localisation and functional capacity of xanthine oxidase to generate signalling molecules such as hydrogen peroxide and nitric oxide that can modulate hypoxia related transcription factors, has provided the basis for our current investigations on xanthine oxidase as an “oxygen sensor”.

The net change in PO2 is signalled and transduced via the pathway described above, which leads to the production of associated proteins required for the achievement of homeostasis. At this level, the cell types that are most sensitive to PO2 changes go through necrosis or apoptosis. Altered cellular proteins, which are potentially toxic or antigenic, can be cleared in two ways. One pathway is via the induction of “heat shock” or “stress proteins”, which either correct the protein alterations or act as chaperones to clear/recycle these potential toxins. Many of these stress proteins have hypoxia response elements (HRE) in their promoters, facilitating Hif-1 mediated activation.38

The second wave of reparative responses to hypoxia mediated localised cell death and related release of altered proteins, is the inflammatory process. The initial step in the recruitment of immunomodulatory cells is the induction of a whole host of adhesion molecules (integrins), which mediate the capture, rolling, adhesion, and transmigration of these cells from the circulation. The transcriptional induction of the genes for these integrin molecules, for the most part, involve hypoxia or redox controlled transcription factors.39

The recruited immunomodulatory cells, as well as dealing with the clearance of the antigenic proteins, release a range of growth factors and cytokines that lead to proliferative responses in fibroblasts. Macrophages also contribute to this process by releasing potent cytokines such as TNFa and TGFβ.40,41 Both hypoxia and TGFβ are known to induce ET-1 production from endothelial cells, although differential expression is observed in different vascular beds.42 Fibroblastic cells are then primed by ET-1 to produce factors that initiate extracellular matrix production and angiogenic factors, collagen, FGF, PDGF, and VEGF, thus aiding in the reorganisation and repair of the damaged tissue.43,44 The genes for these factors are controlled by hypoxia/redox sensitive transcription factors such as Hif-1, AP-1, and NFκB.45

At the whole body level, hypoxia influences the regulation of ventilation and red cell mass via transcriptional modulation of erythropoietin and tyrosine hydroxylase genes, which are also controlled by the transcription factor Hif-1.46 This represents a hypoxia driven global response for re-establishing normal perfusion.
Tissue Responses to Anoxia

Anoxia results in the activation of the “anoxic response system” that plays an important part in clearing debris from a wound, before release of angiogenic factors and restoration of blood supply. As the oxygen tension rises, proteolytic enzyme release ceases and anoxic response gives way to matrix resynthesis leading to restitution of tissue lost during the wounding or trauma. In parallel with hypoxia mediated induction of Hif-1, anoxia induces a separate transcription regulatory element anoxia inducible factor (Aif), with a distinct set of target genes separate from those regulated by Hif-1. In rodents, the anoxic response system is characterised by the upregulation of retroviral inserts, VL30 retrotransposon elements, with unknown functions. These are only seen during wound repair and invasion of certain tumours in rodents, leading Anderson and Stoler to suggest that some tumours have “borrowed” the wound anoxic response system to invade normal tissue. It is conceivable that erosion of cartilage and bone in RA may involve this same anoxic response system. Interestingly, the MRL/lpr strain of mice that develop spontaneous erosive arthritis have a retrotransposon inserted in the fas gene, thereby interfering with apoptosis.

We suggest that the slow, intermittent progression of cartilage and bone erosion is consistent with intermittent on/off cycles of the anoxic response system in a wound environment, but more studies are required to prove this.

In contrast, hypoxia induced events occurring on the other side of the synovial pannus, we believe, are responsible for the synovitis driven by induced inflammatory cytokines. See Table 1 for a summary of hypoxia and anoxia induced proteins.

Transcriptional and Phenotypic Changes in Rheumatoid Synovium Reflect Genetic Responses to a Hypoxic/Anoxic Metabolism

We have reviewed above the collective evidence supporting the chronically hypoxic nature of the inflamed rheumatoid synovium. Upregulation of the glycolytic enzymes glyceraldehyde-3-phosphate dehydrogenase and lactate dehydrogenase in synovial tissues clearly reflect a hypoxia mediated metabolic change because the transcriptional control of such enzymes rely predominantly on the activation of Hif-1. In fact, Hif-1 is probably the most directly involved transcription factor in the activation of hypoxia responsive genes that are believed to play significant parts in the pathology of synovitis. The prime example of genes controlled by hypoxia through Hif-1 activation is the gene for VEGF. VEGF is a cytokine with strong influence on the regulation of wound healing, response to hypoxic injury, and tumour pathogenesis (the common element being the induction of angiogenesis). Recent evidence suggests a role for this cytokine in bone remodelling. Significant transcriptional upregulation of VEGF expression in rheumatoid synovial fibroblastic lining cells has also been reported. These observations, we suggest, again reflect a hypoxia driven attempt to establish the much required vascularisation to adequately perfuse the rheumatoid synovium.

Erythropoietin (EPO) production is another significant hypoxia regulated event that is controlled by the activation of Hif-1. Studies analysing EPO production in RA have shown increased concentrations, again implying a possible hypoxia mediated impairment of the normal relation between EPO concentrations and the degree of anaemia.

Several other genes that are transcriptionally modulated by hypoxia have significant contribution to the pathology of synovitis. These include the genes for cytokines such as TGFβ, which mediate both inflammatory and bone resorptive events, and the vasoactive, fibroelastic mitogen ET-1. Hypoxia induces TGFβ through pathways common to hypoxia sensing mechanism(s) described for EPO. TGFβ, in turn, is known to regulate ET-1 expression via Fos and Jun oncoproteins (the transcription factor AP-1). TGFβ also mediates expression of collagen genes, which contain the AP-1 binding element. AP-1 is regulated directly by hypoxia/redox-balance. NFκB, also a redox controlled transcription factor, regulates the transcription of the integrins. Inflamed synovial endothelial cells show increased surface expression of a series of integrin molecules (ELAM-1/ICAM-1). Lymphocytes and neutrophils within inflamed synovial tissues also show increased expression of the associated genes.

Table 1 Proteins and implicated transcription factors induced by hypoxia and anoxia

<table>
<thead>
<tr>
<th>Protein</th>
<th>Transcription factor</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Xanthine oxidoreductase (XOR)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haem oxygenase-1 (HO-1)</td>
<td>Hif-1, AP-1</td>
<td>27</td>
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<tr>
<td>Glyceroldehyde-3-phosphate dehydrogenase</td>
<td>Hif-1, AP-1</td>
<td>25</td>
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<tr>
<td>Lactate dehydrogenase</td>
<td>Hif-1, AP-1</td>
<td>28</td>
</tr>
<tr>
<td>Erythropoietin (EPO)</td>
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<td></td>
</tr>
<tr>
<td>Vascular endothelial growth factor (VEGF)</td>
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<td></td>
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<tr>
<td>Transforming growth factor β (TGFβ)</td>
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<tr>
<td>Endothelin-1 (ET-1)</td>
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<td></td>
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<tr>
<td>Human α(1) collagen</td>
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<td></td>
</tr>
<tr>
<td>TNFα, IL1, IL8, IL6</td>
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<td></td>
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<tr>
<td>Anoxia induced</td>
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<td></td>
</tr>
<tr>
<td>Lactate dehydrogenase, Cathepsin D, Cathepsin L, Endonuclease NX</td>
<td>? AIF</td>
<td>48</td>
</tr>
</tbody>
</table>

Table 1 for a summary of hypoxia and anoxia induced proteins.
influence of hypoxia in inflammatory synovitis

sections has also been reported.59 These proto-oncogenes in inflamed synovial c-jun factor AP-1, is the protein product of c-jun proto-oncogene. Increased expression of c-jun has been reported in RA synoviocytes.77 Within the chronically hypoxic innermost layer (pannus tissue) and erosive sites, fibroblastic lining cells have a transformed morphology, show anchorage independent growth characteristics and increased proliferation.58 This, we correlate with the increased secretion of PDGF from the fibroblastic cells, as an autocrine factor within the inflamed synovium. Similarly, the transcription factor AP-1, is the protein product of c-fos and c-jun proto-oncogenes. Increased expression of these proto-oncogenes in inflamed synovial sections has also been reported.78

Other proto-oncogene related transcription factors, such as NFkB, are modulated by hypoxia mediated downstream events associated with redox balance and oxidant stress. A functionally significant component of NFkB, is the proto-oncogene c-rel. Increased activation of this hypoxia/redox controlled transcription factor and its predominant vascular localisation within the inflamed synovium further emphasises the pivotal influence of hypoxia in the disease.69 We suggest therefore, that hypoxia/redox controlled oncogenic activity and the control of associated transcription factors and their target genes, largely explain the structural and phenotypic changes within the inflamed synovium.

In parallel to hypoxia controlled genes within the inflamed synovium, anoxia plays a significant part in transcriptional events at bone and cartilage erosive sites—that is, the pannus tissue. Descriptions by Gay and his colleagues of a so called “transformed fibroblast phenotype” within pannus tissue with the property of eroding cartilage and bone in RA,50 are very similar to those of the activation state of the wound fibroblast of the “anoxic response system”. At the eroding interface of the pannus, transformed fibroblasts have been shown to release cathepsins B, L and, importantly, K, which is the isotype released by osteoclasts during bone resorption.62 These cells express the adhesion molecule VCAM-1, which is thought to play a part in adhesion of the transformed fibroblast to the tissue being resorbed. Similar studies to compare wound fibroblasts remain to be undertaken because the process of erosion has overtones of the process of wound repair by the fibroblast in anoxia and may be a quite separate process to the hypoxia driven cytokine and immune events on the cavity side of the synovial membrane. Telling evidence for this comment derives from observation on patients with AIDS who also have the misfortune to suffer from RA. At the time CD4+ lymphocytes fall precipitously, symptoms of the arthritis disappear, however, at necropsy evidence was found that erosion was still continuing in the absence of cytokine stimulation of the eroding fibroblasts.

Ingenious transfection studies of RA transformed fibroblasts with the IL1 inhibitor IRAP (IL1 receptor antagonist protein), and coimplantation of these with human cartilage in a SCID mouse system, demonstrated that the erosive process through cartilage is not driven by the cytokine IL1. The transfected cells eroded cartilage at the same rate as those with an irrelevant construct, however, the accompanying chondrocytic chondrolysis was inhibited.63 These cells are able to produce low levels of cytokines, but do not seem to respond to such cytokines. Finally, the first cells within the joint to proliferate proliferate in the MRL/lpr mouse model of erosive arthritis are the synovial fibroblasts, and the erosive process is underway before any inflammatory and immune cell involvement.64

The “physiological” cell and tissue responses to hypoxia/anoxia are adaptive and organised, and lead to the resolution of injury and homeostasis. Although the pathological characteristics of synovitis parallel these physiological responses to hypoxia/anoxia, lack of resolution of injury implies a specific disruption of such organised and tightly controlled responses in synovitis. We argue that movement of inflamed joints introduce mechanical factors as well as pressure changes that contribute to chronic hypoxia and localised anoxia (particularly at erosive sites), preventing the completion of much required angiogenesis. Thus, the net effect of hypoxia and/or anoxia induced changes within the rheumatoid joints is persistent, unresolved inflammation and cartilage/bone destruction.

therapeutic approaches

An extensive array of therapeutic strategies have been suggested and tested for the treatment of RA, albeit with no definitive outcome. Some approaches have been based on the antioxidative imbalance in inflamed synovial tissues. Supplementation with folic acid and the antioxidant vitamin E, have been one such approach. Previous studies have not conclusively demonstrated that the rheumatoid joint is locally folate deficient, or that, if such is the case, folate gets into the joint when given as an oral supplement. Folate given intrarticularly however, was found to suppress some of the hyaluronan induced flares. Further pharmacokinetic studies are required for the elucidation of the role of folate in rheumatoid synovitis.

In contrast, it is known that patients with RA are locally deficient of vitamin E and that the latter gets into the joint when given orally.70 However, a significant anti-inflammatory effect has not yet been reported. Antioxidants and vitamins act in concert with each other and supplementing one without the other may actually deplete tissues of certain vitamins. It is
also possible that other antioxidants are depleted in patients with RA. Indeed, it is known that patients with RA are vitamin C deficient. Clearly, if vitamin C regenerates vitamin E in vivo as it does in vitro, patients who are deficient in both vitamins have little chance of benefiting if one is given without the other. Vitamin E does however have an analgesic effect that seems to be independent of its antioxidant properties. The interactions between the various antioxidant enzymes in vivo have yet to be fully elucidated, and there remains much scope for clinical trials using combinations of these compounds.

Alternative recent strategies involve anti-cytokine treatment. Clinical trials using chimeric anti-TNFα monoclonal antibodies, in open label and randomised placebo controlled studies, have demonstrated a dose dependent efficacy with significant improvement in disease activity. However, the possible long term effects of such treatment are largely unknown.

The alternative to counteracting oxidative stress or inhibiting specific cytokines, we believe, is to use the established and substantiated observations that the rheumatoid synovium is hypoxic. On this basis, we suggest development of novel therapeutic strategies that target hypoxia directly.

The direct killing of hypoxic cells has long been the concern of radiotherapists and oncologists, as hypoxic cells are radiation resistant compared with oxic cells, and this resistance is an important factor influencing local tumour control by radiation. The hypoxic intracellular environment is a reducing one, as we eluded to earlier. This reductive metabolism can be used by certain drugs that are reduced to more toxic metabolites than the parent drug. Such “bioreductive” drugs have considerably more toxic effects on hypoxic than oxic mammalian cells, and their cytotoxicity correlates with their electron affinity. The basis for their “selective” toxicity towards hypoxic tissue is ascribed to the activation of the drug to a free radical intermediate. In fully oxygenated tissues, the free electron from the reduced drug is rapidly received by cellular oxygen via the process of “futile cycling”. The leakage of superoxide anion (O₂⁻) is of potential concern but is suppressible by the antioxidant mechanisms of normal cells.

The critical step in hypoxic cytotoxicity is enzyme mediated reductive activation. A decrease in cellular PO₂ is known to induce the synthesis of reductase enzymes such as xanthine oxidoreductase, NADPH:cytochrome P₄₅₀ reductase, and i-NOS. Indeed, hypoxia has been identified to be a significant factor in the transcriptional regulation of these particular enzymes. In addition, other factors prevalent within the hypoxic environment of the inflamed rheumatoid joint, such as calcium mobilisation and cytokine release, are also known to activate these enzymes. We have previously reviewed the significance of these enzymes in RA synovium.

One particular bioreductive drug, metronidazole, has previously been used in RA with reports on positive therapeutic effects. This was based on the postulate that RA may be caused by anaerobic bacteria. However, these studies were not fully controlled. Furthermore, a double blind study in a mixed rheumatoid population did not confirm such a therapeutic action. Metronidazole is, however, the least active of the nitro-heterocyclic compounds, and it is reasonable to speculate that the rheumatoid synovium is a potential target for other bioreductives with a greater electron affinity. A range of bioreductive drugs are yet to be tested in rheumatoid patients. Our current in vivo studies have positive primary indications. The possibility of targeted, intra-articular administration of such cytotoxic bioreductive drugs would result in “auto-synovectomy” and avoid the need for painful and lengthy surgery. An additional advantage of such a treatment regimen would be to minimise the use of potentially more toxic drugs that are currently the mainstay of RA treatment.

Our recent strategy is based on the potential use of bioreductives as “carrier” molecules to specifically target other drugs to the synovium. This would not only have the effect of reducing the risk of systemic side effects, but also increase the therapeutic effect of the “targeted drug” linked to the “carrier”. For example, the weak acidic based NSAIDs, which undergo ion trapping in acidic tissue could be linked to a bioreductive agent and targeted to hypoxic or acidic cells, achieving improved efficacy and minimal toxicity. Corticosteroids may also be covalently linked to bioreductive compounds, again improving the therapeutic index and minimising systemic corticosteroid effects. We are currently designing the synthesis of such bioreductively linked novel therapeutic agents to be tested in our model systems.

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References:
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