Metabolomics – a novel window into inflammatory disease

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Summary

Inflammation is an important component of normal responses to infection and injury. However, chronic activation of the immune system, due to aberrant responses to normal stimuli, can lead to the establishment of a persistent inflammatory state. Such inflammatory conditions are often debilitating, and are associated with a number of important co-morbidities including cardiovascular disease. Resting non-proliferative tissues have distinctive metabolic activities and requirements, which differ considerably from those in infiltrating immune cells, which are undergoing proliferation and differentiation. Immune responses in tissues may therefore be modulated by the relative abundance of substrates in the inflamed site. In turn immune cell activity can feed back and affect metabolic behaviour of the tissues, as most clearly demonstrated in cachexia – the loss of cellular mass driven by tumour necrosis factor-alpha (TNF-α) a key mediator of the inflammatory response. Here we discuss the potential for metabolomic analysis to clarify the interactions between inflammation and metabolic changes underlying many diseases. We suggest that an increased understanding of the interaction between inflammation and cellular metabolism, energy substrate use, tissue breakdown markers, the microbiome and drug metabolites, may provide novel insight into the regulation of inflammatory diseases.

Key words: inflammation; metabolomics; rheumatoid arthritis; hypoxia; biomarkers

Metabolism and inflammation

Inflammation is a normal and important response to infection and injury. The cardinal features of inflammation – swelling, redness, stiffness and increased temperature – are outward indicators of significant local changes in metabolism. Increasing blood flow in turn affects local nutrient supply and in particular oxygenation of the tissues while infiltrating activated immune cells bring additional metabolic stressors that must be met. Furthermore, inflammation is associated with the rapid influx, differentiation and proliferation of leukocytes. Infiltrating cells have distinct metabolic requirements and, together with the increased cell number, result in a significant alteration in the metabolic profile of the inflamed tissues. In turn, the environment of the tissues may alter the activity, behaviour or differentiation of the invading cells. The activity of macrophages and neutrophils in both clearance of infection and tissue repair is particularly significant due to the production of cytokines and cytotoxic molecules, including reactive oxygen species and reactive nitrogen species, consuming considerable oxygen, adenosine 5’-triphosphate (ATP) and reduced nicotinamide adenine dinucleotide phosphate (NADPH) in the process. Reactive species, while essential for killing invading organisms, also put considerable stress on surrounding and distal tissues via loss of protective metabolites including reduced glutathione (GSH). In rheumatoid arthritis patients blood levels of GSH decline by 50% and this is associated with a 3-fold increase in lipid peroxides [1]. A similar picture is seen in healthy ageing and this is en-

Figure 1
Levels of plasma antioxidant glutathione decrease with age whereas markers of oxidative damage lipid hydroperoxides increase. This provides an example of the magnitude of changes in metabolite levels that can be observed in humans which, in this case, may be consequence of inflammaging. Adapted from Nuttall SL, Martin U, Sinclair AJ, Kendall MJ. Glutathione: in sickness and in health. Lancet. 1998;351(9103):645–6 [62].
hanced in individuals receiving medical attention (fig. 1) suggesting that the process of “inflamma ging” may inter-
act with pathological developments to promote changes in metabolism [2].

It is interesting to note that mechanisms of metabolic and immune control co-evolved, originating in single fat body
organ as still seen in Drosophila melanogaster [3]. This as-
association persists in higher organisms, where lymph nodes
are embedded in perinodal adipose tissue that may influence
immune responses [4]. In humans adipose tissue is
well infiltrated with macrophages, and the production of
inflammatory cytokines by both adipocytes and macro-
phages contributes to systemic inflammation [5]. This link
between inflammation and metabolism is well demon-
strated in cachexia, where the loss of cellular mass is driven
by tumour necrosis factor-alpha (TNF-α) a mediator of the
inflammatory response [5, 6]. Under normal circumstances
acute inflammatory triggers are cleared or repaired rapid-
ly, with subsequent homeostatic return. However, there are a number of chronic inflammatory diseases where aber-
rant immune activation results in a persistent inflammatory
state. The metabolic consequences of chronic inflamma-
tion extend beyond the local site of disease, driving im-
portant co-morbidities including accelerated atherosclero-
sis and cardiovascular disease. Understanding of the met-
bolic aspects may therefore be key to fully characterising
inflammatory disease. However, given the complexity of
the interlinking metabolic pathways in different organs, tis-
sues and cells a systems biology approach is needed to as-
sess and interpret these metabolic changes.

Principles of metabolomics

Metabolomics is a novel systems approach that can be used
to dissect the local and systemic metabolic consequences
of inflammation. Just as genomics studies the genetic basis
of phenotype, and transcriptomics and proteomics study
the products of these genes, metabolomics seeks to under-
stand the downstream effects caused by the action of these
proteins and enzymes in the context of energy and meta-
bolite consumption and regulation. A hypothesis-forming
approach, it is driven by the non-discriminant analysis of the
low molecular weight metabolite component of target
samples. The potential of metabolomics has shown promise
in the prediction and diagnosis of diseases including ulcer-
ative colitis, rheumatoid arthritis, multiple sclerosis (MS)
among others [7–9].

Metabolomic analysis starts with the acquisition of meta-
bolite data from samples. There are a number of variations
of techniques for this purpose, although the most com-
monly used are nuclear magnetic resonance (NMR) spec-
troscopy, or mass spectrometry (MS). Samples may be de-

erived from any sample suitably processed to be cell-free,

ingcluding urines, plasma, cerebrospinal fluid, faccial ex-
tracts and synovial fluid [7–9], and pH stable through addi-
tion of buffers. Alternatively cell extracts may be per-
formed into a suitable solvent, such as methanol with chlo-
roform extraction to remove protein components. Consistent
processing and buffering is key to the success of meta-
bolomic analysis due to the potential confounding factors
that may result from variations in pH during acquisition –

including shifted peaks and chemical reaction resulting in
loss or gain of metabolites. Acquisition of samples results
in spectra containing a number of peaks, representing pro-
ton resonance in $^1$H NMR or mass/charge (m/z) ratios in
mass spectrometry. In practise however the goal of analys-
is the same – using differences in metabolites to describe
the differences between sample groups. The large size of
the datasets and often numerous samples requires the use
of multivariate analysis to simplify and distil meaning from
the resulting data.

Statistical methods for the analysis of metabolomic data
may be broadly grouped into directed and undirected ap-
proaches [10]. In undirected analysis such as PCA, the
goal is to describe the maximum variation in the dataset it-
self without providing information to the model about the
class groupings. The output is a series of orthogonal prin-
cipal components, describing the maximum to minimum
variation in the data in turn through weighted peaks. If
the source of largest variation in the data is the difference
between control and test samples, principal components
analysis (PCA) alone is sufficient to differentiate class

![Figure 2](image)

**Figure 2** Metabolomic analysis of serum samples from old (○) and young ( □) healthy volunteers.

1D spectra were acquired in a 500MHz spectrometer and
processed with NMRlab/Metabolab and analysed with partial least
squares discriminant analysis (PLS-DA). The two groups (<25 and
>75 years) were well separated on PLSDA (A) with loadings (B)
showing amine derivatives and ketone bodies are raised in ageing,
which may be markers of underlying inflamming.
groupings. However, in more variable samples, as is often the case in biological sourced materials, other confounding or random data may mask the true differences in the samples. Here directed analysis techniques may be more useful. Partial least squares regression (PLS) may be used to describe variation in one dataset by reference to another – for example, using metabolite concentrations to predict disease activity, progression or severity. Similarly, the related technique PLS discriminant analysis (PLS-DA) uses multivariate metabolomic peak data to describe the assignment of samples to binary groups. By focusing on the variation that describes the differences between control and test samples, it is possible to tease out the clinically relevant variation in the face of significant unrelated noise. The output of these approaches is markedly similar: a statistical model from which it is possible to accurately predict disease state or patient group from similarly processed samples from new individuals. However, these models are complicated and include multiple correlated peaks. Forward selection regression analysis may therefore be employed to iteratively discard peaks resulting in a small number of metabolites while maintaining predictive capability – a requirement for the development of cheap specific bioassays. Selection of controls, accounting for confounding effects such as diet, smoking, and medication history, is key to the development of accurate models.

A simple example of the output of a metabolomic study is shown in figure 2, where we assessed NMR-derived metabolic fingerprints of serum from a small cohort of healthy elderly individuals (age >75 years) in comparison to young donors (<25 years). The groups are clearly separated in a PLS-DA model and analysis of the loadings plot (which indicates which regions of the NMR spectra contribute to the separation) indicates several distinguishing metabolites including methionine and ketone bodies which have been associated with ageing in rats probably as a result in changes in pyruvate and energy metabolism [11]. Ketone bodies are also generated by inflammatory leukocytes and have been observed in the synovium in RA [2]. Therefore, here they may be present as a result of low grade inflammation which is characteristic of inflamming and so may provide novel markers of this process [12]. By applying these metabolomics techniques to inflammatory disease the goal is to determine the role of metabolic processes in the development of normal and aberrant inflammatory states. Metabolite markers, assayed from the various tissues and fluids, are accessible and simple to obtain from patients while having the potential to provide deeper indications of underlying disease processes.

**Metabolic markers**

In the following sections we take a systematic approach to outline key markers identified by metabolomic studies to date which are summarised in table 1.

In spite of the early state of the field, there already exist a number of key areas of investigation where metabolic patterns are emerging, both in general inflammatory disease and in specific pathologies. Here we focus on the most important of these which include energy sources, hypoxia, regulation of inflammation, tissue degradation, waste and drug metabolites.

Table 1: Key metabolites and their association with inflammatory diseases in current literature arranged according to the major themes discussed.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Associations</th>
<th>Fluids</th>
<th>Inflammatory diseases</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactate</td>
<td>Warburg and the hypoxic environment</td>
<td></td>
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<tr>
<td>Lacatate</td>
<td></td>
<td>SF, Urine, CSF</td>
<td>▲RA, ▲OA†, ▲IIH, ▲MS</td>
<td>[8, 30, 32]</td>
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<tr>
<td>Citric acid cycle intermediates</td>
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<tr>
<td>Oxaloacetate</td>
<td></td>
<td>CSF, VF</td>
<td>▲IIH, ▲LIU</td>
<td>[9]</td>
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<tr>
<td>Citrate</td>
<td></td>
<td>CSF</td>
<td>▲IIH</td>
<td>[9]</td>
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<tr>
<td>Malate</td>
<td></td>
<td>Urine</td>
<td>▲OA†</td>
<td>[32]</td>
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<tr>
<td>Alternative energy sources</td>
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<td>Amino acid metabolites</td>
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<td>Xanthurenic acid</td>
<td></td>
<td></td>
<td>▲CD</td>
<td>[48]</td>
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<tr>
<td>Uracil, xanthine, glycine</td>
<td></td>
<td></td>
<td>▲RA†</td>
<td>[50]</td>
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<tr>
<td>Ketone bodies</td>
<td></td>
<td>SF, CSF</td>
<td>▲RA, ▲MS</td>
<td>[12, 42, 43]</td>
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<tr>
<td>Lipoprotein-associated fatty acid breakdown</td>
<td></td>
<td>SF, Blood</td>
<td>▲IA, ▲OA</td>
<td>[41]</td>
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<td>Tissue degradation and waste</td>
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<tr>
<td>Essential amino acids</td>
<td></td>
<td>Protein breakdown</td>
<td>▲OA</td>
<td>[46]</td>
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<td>Hyaluronic acid</td>
<td></td>
<td>Serum</td>
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<tr>
<td>Trimethylamine</td>
<td></td>
<td>Urine</td>
<td>▲IBD, ▲IBD†, ▼CD</td>
<td>[47, 53]</td>
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<tr>
<td>Urea</td>
<td></td>
<td>VF</td>
<td>▲LIU</td>
<td>[52]</td>
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<td>Xenobiotics and the Microbiome</td>
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<tr>
<td>Oxylipids</td>
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<td>NSAIDs</td>
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<td>[54, 61]</td>
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<td>Simvastatin</td>
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<td>Faecal</td>
<td>▼CD</td>
<td>[47]</td>
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<td>Faecal</td>
<td>▼CD, ▼T1D</td>
<td>[47, 58]</td>
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<tr>
<td>Formate</td>
<td></td>
<td>Urine</td>
<td>▲CD</td>
<td>[47]</td>
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<tr>
<td>Others</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Creatinine</td>
<td></td>
<td>CSF</td>
<td>▲MS</td>
<td>[30]</td>
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RA = rheumatoid arthritis; IA = inflammatory arthritis; OA = osteoarthritis; IIH = idiopathic intracranial hypertension; MS = multiple sclerosis; LIU = lens induced uveitis; CD = Crohn’s disease. CSF = cerebrospinal fluid;VF = vitreous fluid, SF = synovial fluid; † animal model.
Lactate: the Warburg effect and hypoxia

In the presence of adequate oxygen, differentiated cells metabolise glucose through mitochondrial oxidation of glycolytic pyruvate in the tricarboxylic acid (TCA) cycle, with resulting NADH fuelling oxidative phosphorylation to produce 36 ATP per glucose molecule. Under hypoxic conditions the oxidative pathways are inhibited and glycolytic pyruvate is instead excreted as lactate, with a decreased 2 ATP output per glucose molecules in the cell. Physiological levels of oxygen range between 5%–12% depending on tissue location. Structures in the body held naturally at the lower end of this range, by virtue of low perfusion or vascularisation, include the lymph nodes, eye and synovium. In normal inflammatory conditions hypoxia results as a side effect of blood vessel occlusion due to inflammation or as a result of increased cell density following infiltration. Hypoxia is a key driver of cell migration into the tissues and perfusion may act as a signalling mechanism to invading cells [13]. Reductions in oxygen lead to stabilisation of transcription factor hypoxia-inducible factor (HIF) providing an elegant cellular oxygen detection system. Stabilisation of HIF drives major changes in cellular gene expression including the promotion of glycolytic enzymes. Peripheral blood CD4+ T cells exposed to hypoxia show dramatic induction of genes involved in metabolism and homeostasis [14]. Innate immune cells, including neutrophils and macrophages are thought better adapted to function at low oxygen tensions with preferential use of glycolysis to provide ATP even under normal physiological oxygen [15]. Macrophages tend to accumulate in the hypoxic sites of chronic inflammation and hypoxia is associated with the activation of infiltrating and tissue-resident macrophages resulting in up-regulation of pro-inflammatory IL-1, IL-6, IFN-γ and TNF-α [16–20]. Most infections or injuries are cleared or repaired quickly with perfusion and metabolic homeostasis soon restored. However, in poorly perfused tissues such as the rheumatoid synovium and sclerotic tissues oxygen perfusion can fall as low as 0.5%–2.5% and this regulation may be perturbed leading to continued recruitment [21, 22]. Lactate is a waste product of normal metabolism reversibly converted from glycolytic pyruvate via the enzyme lactate dehydrogenase. Excess cellular lactate is excreted and re-circulated via conversion in the live to glucose (the Cori cycle). In sufficiently oxygenated cells it may also be converted back to pyruvate via reaction with intracellular NAD+ to fuel the citric acid cycle. However, in poorly perfused tissues the production of lactate outstrips this homeostatic control and lactate accumulates. In hypoxic conditions the HIF-mediated up-regulation of lactate dehydrogenase A further promotes this accumulation [23].

Lactate therefore is a key marker of inflammatory sites under hypoxia. However, this relationship is not restricted to inflammation. In 1924 Otto Warburg first observed that cancer cells metabolise glucose by glycolysis even under aerobic conditions [24]. It has been suggested that all differentiating and replicating cells must use this same mechanism in order to produce the requisite biomass for cellular replication [25]. By this mechanism, glucose and glutamine can adequately supply the majority of carbon, nitrogen and energy for cell growth and division. Indeed, under growth conditions, metabolising glucose to CO2 via oxidative phosphorylation runs counter to the needs of the cell. Therefore, non-proliferative, highly ATP-dependent organs such as the brain favour free-energy production via oxidative phosphorylation of glucose [26], while in contrast, under resting conditions, skeletal muscle metabolism is predominantly fatty acid dependent, switching to glycolysis and oxidative phosphorylation under activity [27]. In proliferative tissues such as the lymph nodes, there is instead a tendency towards metabolism of glucose to lactate, pyruvate and CO2, and glutamine to ammonia, glutamate and aspartate. Importantly this may also apply to cells, such as polymorphonuclear leukocytes, that invade inflammatory sites [28, 29]. The switch to aerobic glycolysis therefore compounds the production of lactate, locally and distally – including where hypoxia is absent.

The role for lactate as a universal marker of inflammation is borne out clinically. Increases in the ratio of lactate: glucose in blood plasma and synovial fluid in patients with rheumatoid arthritis has been shown to correlate with active inflammation and oxidative damage to the joint, for example [8]. Meanwhile, in a wide-ranging study of cerebrospinal fluid (CSF) from patients with a number of neurological conditions, high lactate was a distinguishing feature of idiopathic intracranial hypertension (IH) hinting to a potential inflammatory basis for the disease with pressure occlusion of the intracranial vasculature. Increased CSF lactate and creatinine are also seen in multiple sclerosis patients versus controls, potentially reflecting leukocyte infiltration into the lesions [30]. Such alterations in pyruvate and lactate can in turn drive downstream effects on the citric acid cycle. Pyruvate may be converted to intermediates of the citric acid cycle, malate and oxaloacetate [31] by pyruvate carboxylase and malic enzyme respectively. Raised urine malate is a strong predictor of disease activity in the Hartley guinea pig model of OA [32]. Raised oxaloacetate and reduced citrate are also associated with disease in idiopathic intracranial hypertension together with increases in ketone bodies 3-hydroxybutyrate, suggestive of an anaerobic carbohydrate-deficient environment [9].

Alternative energy sources

The base relationship between resting energy expenditure and body mass is described by the Kleiber formula (REE = 293M0.75). Organ-specific metabolic rates and preferred substrates are also well established (fig. 3) with the heart and kidney consuming the major proportion of energy in the resting adult, followed by the brain and liver [33, 34]. However, these basal rates are significantly altered by malnutrition, illness, smoking, and inflammation and changes in substrate availability can have disproportionate effects [34]. Recent work on a cohort of RA patients found that their global basal metabolic rate (BMR) was 6,276 kJoules/day, which was 8% higher that the value from a control group. Importantly a group of RA patients who smoked had a 20% higher BMR of 7,188 kJoules/day (fig. 4) [35]. This indicates how large the effects of inflammation and factors promoting it can be and explains why changes in blood and...
metabolites are easily observed. Inflammation is a highly energy-dependent process and acutely reliant on adequate energy availability to function correctly [36]. Fever is associated with an 11% increase in energy consumption per 1°C, while phagocyte production during an infectious episode consumes approximately 790 kJoules [37, 38]. In sepsis the metabolic rate can reach up to 30%–60% above baseline levels [39]. The converse is also true, with the energy surplus associated with obesity and other metabolic syndromes driving altered immune responses and chronic inflammation [40].

Energy availability is therefore a key regulator of immune activation and resolution. Resting energy requirements for non-brain and non-proliferative tissues, particularly the skeletal muscle and heart, are predominantly met by the oxidation of free fatty acids [27]. Due to the oxidation requirement their use is typically limited to well-perfused tissues, however damage to the joint in inflammatory and osteoarthritis appears to be associated with an increased perfusion of macromolecular fatty acids into the synovium. In osteoarthritis increased lipoprotein associated fatty acids, together with metabolism products glycerol, ketones and pyruvate, are all indicative of lipolysis occurring as an energy source. Similarly, a study of RA synovial fluid samples has shown reduced chylomicron and very-low-density-lipoprotein associated triglycerides compared to controls, suggesting increased utilisation of fats as an energy source in the joint in spite of the hypoxia environment. Blood plasma levels of cholesterol, acetylated glycoprotein and lipids were also raised in RA [41]. Fatty acids are additionally broken down in the liver and kidney, producing ketone bodies for recirculation to the tissues. Ketone bodies acetoacetate and 3-hydroxybutyrate provide a key energy source for the heart and brain, through reconversion to acetyl-CoA to feed the citric acid cycle in the tissues, while acetone is readily excreted as waste. In normal fasting conditions plasma levels of ketone bodies are low, as tissues with lower metabolic needs sustain themselves on local reserves of fatty acids, but these sources are not sufficient for active, highly energy-dependent tissues and ketone body production rises significantly. After 3.5 days of glucose starvation the brain can obtain as much as 25% of total energy from ketone bodies [26]. However, alterations in ketone body production and metabolism have also been observed in inflammatory diseases in the absence of glucose restriction. Elevated 3-hydroxybutyrate has also been seen in MS with samples containing as much as 32 μM concentration of this metabolite [42, 43]. As previously described, chronic inflammation may drive metabolic changes, deriving in particular from cachexia-associated catabolism and ketogenesis. The presence of the ketone body 3-hydroxybutyrate in urine has been linked to respiratory chain deficiency leading to impaired NADH oxidation [44]. Local changes may also be informative: raised synovial fluid concentrations of ketone bodies, particularly 3-hydroxybutyrate, in patients with rheumatoid arthritis are suggested to reflect the degradation of local acetyl-CoA under hypoxic conditions [12]. This was also suggested by the low glucose NMR signal in synovial fluid which was 20% lower than paired sera, which is a direct measure of the decrease in this metabolite.

Tissue degradation and waste

Normal tissue homeostasis requires the turnover of both cellular and acellular biomass. In the inflammatory state however, these processes are often altered offering the possibility of characteristic excess or paucity of specific biomarkers. Specific tissue-derived metabolites may provide the clearest markers of gross tissue destruction. Hyaluronic acid for example, is a major component of articular cartilage proteoglycan aggregate and essential for the functional integrity of extracellular matrix. Hyaluronidase activity is absent in both normal and inflamed synovial fluid, and therefore breakdown does not occur. However, in rheumatoid arthritis the action of reactive oxygen species results
Xenobiotics are chemicals found in an organism that are not derived from normal biological processes or dietary intake – a classification that includes most drug treatments. Metabolism of xenobiotics is a complex process, predominantly occurring in the liver and typically involving multiple steps of oxidation, reduction, hydrolysis, cyclisation, and decyclisation with the end goal of detoxification and excretion. Metabolism is associated with characteristic profile of intermediates and excretion products that are unique to the drug or class. In a clinical context these metabolites simply restate what is already known – the treatment regimen of the patient. More interesting are the effects of drug metabolism on disease progression and resolution. For example, in a study of diclofenac significantly altered levels of oxylipids were seen in those patients that responded to the drug, with arachadonic acid metabolite 5,6-DHET identified as a novel marker of inflammation [54]. Similarly, analysis of patient responses to simvastatin successfully discriminated responders and non-responders by metabolic profiles of cholesterol esters and phospholipid metabolites [55]. Interestingly, this study was also able to discriminate between metabolites associated with drug effects on LDL-C or C-reactive protein independently – characterising mechanism of action within an individual. Potentially, the baseline metabolic characteristics of an individual may have important implications for drug lifetime, and effectiveness, and provide a route to tailoring dosing or drug choice to an individual in accordance to not just genetic variability but also lifestyle and diet.

The microbiome is a collective term for the totality of commensal bacteria within our bodies. Bacteria outnumber our own cells by a ratio of 10 to 1 [56], are responsible for many essential metabolic processes for our health, and respond to both inflammatory and immune processes. In an analysis of faecal extracts from patients with Crohn’s disease (CD) and ulcerative colitis (UC) reduced levels of butyrate, acetate, methylamine and TMA were detected compared to controls [47]. Urinary metabolites derived from alterations in gut metabolism also allow distinction to be drawn between the two diseases with hippurate and 4-cresol low in CD versus UC and controls, and formate higher in CD [57]. While it is perhaps unsurprising that inflammatory gut conditions would drive alterations in gut bacterial metabolism, changes have also been seen in other conditions with no known gut involvement. In a case-control study of type-1 diabetes a switch in gut bacteria from mucin-degrading, butyrate producing bacteria, to non-butyrate and lactate-utilising bacteria has been associated with development of the disease [58]. The role of the microbiome in cardiovascular conditions was likewise demonstrated by the finding that metabolism of dietary lecithin by gut flora leads to absorption and accumulation of derivatives of choline which in turn promotes disease [59]. This close relationship between the metabolism of gut bacteria and elsewhere in the body provides a key challenge in our understanding of disease processes through analysis of body fluids.
Conclusion

Metabolomics is, a novel systems approach offering insight into the underlying mechanisms of inflammatory disease. As outlined here, seemingly disparate metabolic markers can be broadly arranged into functional groupings that together describe local and global disease processes. The key difficulty is in separating these local and global effects when assaying through a single bio-fluid. However, in the context of diagnostic or prognostic markers, this is not necessarily required. Metabolic processes do not occur in isolation, and through the noise the interaction of the multiple systems involved in an inflammatory response emerges as a characteristic fingerprint of disease. In developing diagnostic assays we are aided by the fact that these fingerprints frequently consist of the same relatively small subset of metabolites, yet maintain their discriminative capacity, sensitivity and specificity. Metabolomics offers a novel non-invasive approach to rapid diagnosis, prognosis and prediction of inflammatory disease outcomes.

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References


Figure 1
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Figure 2
Metabolomic analysis of serum samples from old (●) and young (○) healthy volunteers. 1D spectra were acquired in a 500MHz spectrometer and processed with NMRLab/Metabolab and analysed with partial least squares discriminant analysis (PLS-DA). The two groups (<25 and >75 years) were well separated on PLSDA (A) with loadings (B) showing amine derivatives and ketone bodies are raised in ageing, which may be markers of underlying inflamming.
Figure 3
Metabolic requirements of the major tissues.
Figure 4
Smoking has a significant effect on the basal metabolic rate of rheumatoid arthritis patients (RA). Metabolic rate is higher in RA compared to healthy controls and this is enhanced even further by smoking, indicating the complexity of interactions between metabolism and disease. Adapted from Metsios GS, Stavropoulos-Kalinoglou A, Nevill M, Douglas KMJ, Koutedakis Y, Kitas GD. Cigarette smoking significantly increases basal metabolic rate in patients with rheumatoid arthritis. Ann Rheum Dis. 2008;67(1):70–3 [35].