Don’t neglect nutrition in rheumatoid arthritis!

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STATE OF KNOWLEDGE
Nutrients and immune-inflammatory response in RA

Rheumatoid arthritis (RA) is characterised by a systemic immune-inflammatory response, in genetically susceptible individuals exposed to environmental and endogenous triggers, including specific nutrients.1 The major pathways in RA are characterised by an intense inflammatory response, involving impaired immunoregulatory processes and the production of different proinflammatory mediators.2 Research on possible risk factors has traditionally focused on triggers setting off disease, such as microbial/viral agents, cigarette smoking and environmental pollution, hormonal imbalance and chronic stress, but with less focus in the past few decades on nutritional factors that can influence disease onset, progression and outcomes, possibly through epigenetic mechanisms.2

More recently, simple and daily dietary factors have been implicated in the development of RA, even directly through triggering inflammatory pathways: for example, the recent evidence that increased sodium chloride salt (figure 1), activates proinflammatory macrophages (M1), Th17 cells and decrease T-regulator cells, all crucial players in RA pathogenesis.3 In addition, sodium excretion was recently found higher in patients with early RA than in matched controls.4

Other interferences exerted by nutrients like cocoa, ginseng or capsaicin (pepper) on the RA pathways and mediators are reported in figure 1 and discussed in greater detail below.

Despite accumulating evidence over time, the important role that the diet plays on human health in general and more specifically in chronic conditions such as RA, has been subject to much controversy. This has had direct impact on the most important stakeholders, the patients, who are the most frequent active seekers and ‘consumers’ of this crucial information.

Evidence for the possible role of different ‘fatty diets’ in models and patients with RA

An important reason for rheumatologists needing to pay attention to nutrition is in order to help control inflammation by encouraging the use of anti-inflammatory diets and decreasing the use of proinflammatory ones. The notion of ‘inflammatory foods’ is becoming increasingly recognised across chronic diseases such as RA. The ‘modern diet’5 especially practised in Western cultures could be viewed as the greatest enemy of chronic inflammatory conditions like RA, whereby the increased consumption of refined carbohydrates, vegetable oils rich in omega-6 fatty acids and decreased consumption of long-chain omega-3 fatty acids represent ‘the perfect nutritional storm’.6 Supporting these observations, animal data have confirmed that a low ratio of n-6/n-3 polyunsaturated fatty acids (PUFA) reduces adjuvant-induced arthritis in rats.7

Since the 1980s and early 1990s where health education initiatives advocated the consumption of antifat diets, in more recent times dietary fats have been increasingly recognised to have a positive impact on health and arthritis.8 Animal studies support that by inducing a collagen-induced arthritis (CIA), in a model of RA in mice consuming high-fat diet (HFD), a risk factor for RA, is related to inflammation but responds minimally to medication. HFD-CIA mice had a high level of α2-glycoprotein 1 (Azgp1), a soluble protein that stimulates lipolysis and fat loss that causes increased IL-17; therefore, those mice showed more severe CIA. The same findings are observed in patients with RA.9 Of particular interest, a recent study showed that obese mice fed with HFD had an earlier onset of CIA compared with mice on regular diet, with even a more sustained joint inflammation in obese mice.10 As a matter of fact, despite established RA disease being unaffected by obesity, the early and the resolution

To cite: Cutolo M, Nikiphorou E. Don’t neglect nutrition in rheumatoid arthritis. RMD Open 2018;4:e000591. doi:10.1136/rmdopen-2017-000591

Prepublication history for this paper is available online. To view these files, please visit the journal online (http://dx.doi.org/10.1136/bmjopen-2017-000591).

Accepted 8 February 2018

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phases of RA are impacted by obesity through different mechanisms. For example, conditioned media from RA adipose tissue can transform RA and wild-type naïve myeloid cells into M1 proinflammatory macrophages.\(^\text{10}\)

On the other hand, the benefits of omega-3 fatty acids (Figure 1) and of monounsaturated fatty acids (MUFA) (key components of the Mediterranean diet) in controlling disease activity in RA have been published\(^\text{11}\) and also shown in human clinical trials.\(^\text{12,13}\) In addition, obesity, a global health problem, represents an important and rising comorbidity even on first presentation of RA\(^\text{14}\) and appears to be a key determinant of insulin resistance, even more so than circulating proinflammatory cytokines.\(^\text{15}\)

What has been specifically shown for patients with RA and what has not

Further important nutritional aspects have been specifically shown for patients with RA. Taking a focus on the nutritional effects of RA pharmacotherapy, one cannot ignore the established side effects related to the disease-modifying treatments (DMARDs) including the anchor drug in RA, that is methotrexate,\(^\text{16}\) causing some kind of 'iatrogenic malnutrition', whether this is due to nausea, stomatitis, upset stomach, diarrhoea and other.

For some of the DMARDs, the gastrointestinal side effects can be particularly prominent with consequences on nutritional status and thus, indirectly, RA disease outcomes. Furthermore, the widespread and non-optimised use of glucocorticoids (GCs) (high dosages, at the wrong time and of prolonged duration) can be 'blamed' at least for the well assessed increased weight gain/body mass index and diabetes. The further interaction between some of these conditions adds to the disease burden in RA and requires specific dietary/pharmacological management.\(^\text{17}\)

Therefore, as rheumatologists we should at the very least encourage reduction or elimination of carbohydrates and high sugar content foods and beverages in our patients with concomitant GC use.

Another important aspect in RA relates to the effects of pregnancy, where there exists strong evidence on the epigenetic/‘therapeutic’ effects of pregnancy states, due to the intense steroid hormonal changes; adherence to the Mediterranean diet during fetal development are key factors in the protection from metabolic syndrome (MS9).\(^\text{18}\) On the other hand, it is supportable, but not specifically shown, that omega-3 PUFA (n-3 PUFA) supplementation of the maternal diet in pregnancy may provide a non-invasive intervention with significant potential to prevent the development of allergic and possibly other immune-mediated diseases, including RA.\(^\text{19}\)

Furthermore, revolutionary treatments for RA, such as the TNF inhibitors, are becoming standard practice...
for most of the 21st century and despite their impressive potential to reduce or even halt overexpression of proinflammatory cytokines, they are not effective by themselves, for example, in increasing muscle mass. In fact, they increase fat mass and related metabolic/nutritional consequences.  

Microbiome, diet and RA

The emerging role of the gut microbiome in RA must also be considered as evidence supports its impact on nutrition and disease progression. The human body contains millions of commensal bacteria (the microbiome), with the bowel being the most prevalent site of colonisation. The process of colonisation begins at birth, and despite interfering factors such as diet and drug use affecting the microbiome composition, by adulthood the gut bacteria are relatively consistent across local populations. Manipulation of the microbiome in inflammatory arthritis, both in animal and human models, offers a potential therapeutic target.

For example, probiotics have been shown to lower the proinflammatory cytokine IL-6 in RA, although how this translates to clinically apparent effects remains unclear, emphasising the need for high-quality trials to investigate these links and prove or disprove the effects on clinically apparent disease. Probiotics contain living healthy bacteria such as Bifidobacteria, Bacteroides-Porphyromonas-Prevotella, Bacteroides fragilis and the Eubacterium rectale-Clostridium cocoides species, that are significantly reduced in the gut microbiome of patients with RA. In contrast, bacteria such as Prevotella copri are found in 75% of people with new, untreated RA and are considered a possible risk factor for triggering disease.

A more healthy diet (fibres) may lead to a more healthy gut microbiota, less active immune system and inflammatory reactions in the gut and finally leading to less inflammation systemically.

As matter of fact, a healthy gut microbiota also releases food metabolites that are anti-inflammatory for the gut immune system and epithelium such as short-chain fatty acids (SCFA). The SCFA are regarded as one of the major microbial metabolites formed by microbial fermentation of dietary fibres, which can improve intestinal mucosal immunity.

Therefore, recent randomised controlled clinical trials (RCTs) seem to provide evidence that specific probiotic supplementation exhibits anti-inflammatory effects, helps to increase daily activities and alleviates symptoms in patients with RA.

WHY RHEUMATOLOGISTS SHOULD CARE ABOUT DIET IN RA

Direct and indirect therapeutic effects of specific nutrients in RA

Referring to a recent example of potential therapeutic effects of nutrients in RA, red hot chili peppers (capsaicin) (figure 1) have been suggested to play anti-inflammatory roles by increasing anti-inflammatory macrophages (M2), modulating the neuroimmune response and decreasing neurogenic pain (topic effect). Recent research has focused on the evaluation of the efficacy of dietary antioxidants such as the phytomolecules and Coenzyme Q10 (CoQ10), an endogenous antioxidant with positive effects. Cocoa (figure 1) represents another nutrient of increasing interest because of its antioxidant properties, which are mainly attributed to the content of flavonoids such as (-)-epicatechin, catechin and procyanidins. In addition, regulatory activity on the secretion of inflammatory mediators from macrophages and other leucocytes in vitro has been proven to be exerted by cocoa.

Interestingly, nanopowdered red ginseng (NRG) (figure 1) used together with methotrexate in arthritic mice significantly reduced cytokines including TNF-a, IL-6 and IL-1b and IgM and IgG1 and suggested the effectiveness of NRG at least in preventing type II collagen-induced RA in mice. Such observations about specific nutrients also trigger questions around the value of supplementation in chronic inflammatory diseases such as RA.

Keeping a focus on evidence-based medicine, one important nutritional aspect in chronic immune/inflammatory diseases such as RA is related to the role and level of vitamin D, or better-said, the D hormone, since it is a true steroid hormone synthesised in the skin from 7-dh-cholesterol under the action of the ultraviolet (UV) sun radiations. Since only 20% of the daily need of vitamin D can be obtained by the diet (80% from UV), vitamin D supplementation is a more accepted practice with important control (as steroid hormone) of both innate and adaptive immune response, especially with increasing recognition that vitamin D insufficiency/deficiency is a frequent observation in RA (figure 1).

Patient-reported outcomes in RA appear to be of value in detecting/predicting by clinical symptoms, the effects of the epidemic deficiency of vitamin D/D hormone in Europe and especially during the winter. Further adding to the evidence base, data from a recent large study suggest that higher intake of dietary vitamin D as well as omega-3 fatty acids, during the year preceding DMARD initiation may be associated with better treatment results in patients with early RA.

Rheumatoid cachexia

Another important reason nutrition should not be neglected in RA is in order to prevent and/or treat ‘rheumatoid cachexia’. Evidence suggests that any ongoing, uncontrolled and chronic inflammatory process in RA, involves adverse effects on body composition, including in particular reduced muscle and increased fat mass. Rheumatoid cachexia refers to these effects, which although rarely apparent, due to the loss of lean body mass being counter-balanced by the maintenance or gain in fat mass, is associated with poor prognosis.

Evidence also suggests that nutrition should be part of routine care in patients with RA with muscle
wasting disorders. More specifically, up to 75% of patients with RA believe that food and nutrition play an important role in their symptom severity, with 50% of patients with RA reportedly trying some form of dietary manipulation in an attempt to attenuate symptomology. A previous study investigated the effects of a daily mixture of β-hydroxy-β-methylbutyrate, glucose, and arginine (HMB/GLN/ARG) protein 12-week supplementation in 40 patients with RA with rheumatoid cachexia. The results showed that both HMB/GLN/ARG and a control mixture of other non-essential amino acids (alanine, glutamic acid, glycine and serine) were both equally effective in increasing lean mass (~0.4 kg) and improving some measures of physical function and strength.

In addition, common mental health comorbidities in RA such as depression and anxiety might influence the nutritional status by inducing anorexia-cachexia. Mental health comorbidities can affect lifestyle and nutritional status with detrimental outcomes. In this respect, supplements containing amino acids are believed to be beneficial, since they are converted to neurotransmitters which in turn alleviate depression and other mental health problems. These observations further highlight the value of personalising dietary regimes for patients with RA and their respective comorbidities.

**Diet and cardiovascular risk in RA**

Several clinical, epidemiological and experimental evidence suggest that consumption of the Mediterranean Diet reduces the incidence of pathologies related to the immune system, oxidative stress and chronic inflammation including atherosclerosis and cardiovascular disease. These reductions can be partially attributed to extra virgin olive oil (EVOO) consumption which has been described as a key bioactive food because of its high nutritional quality and its particular composition of fatty acids, vitamins and polyphenols. Indeed, the beneficial effects of EVOO have been linked to its fatty acid composition, which is very rich in MUFA, and has moderate saturated and PUFA.

Several RCTs to assess potential changes in RA inflammation and related cardiovascular (CV) risk after oral intake of ω-3 PUFA have come to light. A meta-analysis evaluating 20 RCTs, involving 717 patients with RA in the intervention group and 535 patients with RA in the control group was recently published. Despite the evidence of overall low quality trials, consumption of ω-3 fatty acids was found to significantly improve eight disease-activity-related markers. Regarding inflammation, only leukotriene B4 was reduced (five trials, P<0.001), whereas a significant amelioration was found for blood triacylglycerol levels (three trials, P=0.012). The beneficial properties of ω-3 PUFA on RA disease activity confirm the results of previous meta-analyses. On the other hand, a large recent study demonstrated that statin therapy is associated with a lower event rate of new-onset acute coronary syndrome in patients with RA, with the beneficial effect being dose-responsive.

**Prevention of metabolic effects of glucocorticoids in RA**

Patients who are taking exogenous GCs might also be more susceptible to poor food choices; however, the effect of increased fat consumption in combination with elevated exogenous GCs has only recently been investigated. These studies have shown that the metabolic effects initiated through exogenous GC treatment are significantly amplified when combined with a HFD. Animal data confirm that rodents on a HFD and elevated GCs demonstrate more glucose intolerance, hyperinsulinaemia, visceral adiposity and skeletal muscle lipolysis deposition when compared with rodents subjected to either treatment on its own. Exercise has recently been shown to be a viable therapeutic option for GC-treated, high-fat fed rodents. Clinically, these mechanistic studies again underscore the importance of a low-fat diet and increased physical activity levels when patients, like in the case of patients with RA, are given a course of GC treatment. In fact, even at low doses, prednisolone exerts adverse effects on fat metabolism, which could exacerbate insulin resistance and increase CV risk.

**CONCLUSION**

All these observations lead us to further stress and conclude that nutrition matters and importantly in RA, it plays a role in disease progression and outcomes. Although origins of our understanding of diet and disease stem back to palaeontological times, the subject seems to be receiving some ‘revival’ in modern times. Despite this, nutrition and the impact on chronic musculoskeletal disease including RA remain a poorly taught subject, both in medical schools and in postgraduate rheumatology training. One would argue that addressing nutrition in our patients is not the ‘job’ of a rheumatologist, but instead of a dietician. Whereas this may be partly true, rheumatologists are the ones who come face to face with patients and their families and access to a dietician may not always be easily and readily available or at all possible. Similarly, working closely in a multidisciplinary team setting with dieticians is certainly optimal, but not always possible. We therefore advocate that at least some basic knowledge on the subject is warranted by rheumatologists in order to appropriately guide on the best ‘recipe’ for their patients with RA.

**Acknowledgements** This editorial introduces the recent interest from EULAR to educate rheumatologists and postgraduate to the problem of nutrition in RMD. Supported by the EULAR study group on neuroendocrine immunology of rheumatic diseases (NEIRD).

**Provenance and peer review** Commissioned; externally peer reviewed.

**Data sharing statement** Data are elaborated by the authors.

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References


