Obesity and chronic disease: always offender or often just accomplice?

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Over a decade ago, the finding of a form of low-grade systemic inflammation (‘metaflammation’) associated with obesity, insulin resistance and chronic disease proffered a causal explanation for the latter. However, recent work has shown that metaflammation is also associated with several modern lifestyle-related and environmental inducers, with or without obesity. Here, we present accumulating data to show a link between metaflammation and a number of non-microbial environmental and lifestyle stimulants, both with and without obesity. This implies that obesity may often be an accomplice to, as much as an offender in, major metabolic disease. The real (albeit distal) cause of such a disease appears to lie in aspects of the modern techno-industrial environment driving unhealthy lifestyle behaviours. If true, this suggests that while individual weight loss may be a component of chronic disease management, it may be neither ‘necessary’ nor ‘sufficient’ to reduce the problem at a population level. Greater multidisciplinary and policy input is needed to modify the economic and political drivers of the modern, obesogenic environment.

Obesity: Inflammation: Environment: Lifestyle

Obesity is known to be a cause of chronic disease, although the exact mechanisms for this are unclear. Over a decade ago(1), the discovery of a form of subclinical, low-grade systemic inflammation, later called ‘metaflammation’(2) (referring to metabolically triggered inflammation) associated with both obesity and chronic disease, raised hopes of an improved understanding of the causal links between the two, as this form of inflammation is linked with insulin resistance and chronic disease, and obesity seemed to be the driving cause(2). However, increasing knowledge about metaflammation, and its connection with a range of lifestyle-related and environmental factors (diet, inactivity, smoking, sleep, stress, pollution, etc), in the absence, as well as in the presence of obesity, suggests an alternative view; that is, obesity may often be just an accomplice to, as much as a perpetrator of, many metabolic diseases. The real (albeit distal) determinants of such diseases appear to lie in aspects of the modern techno-industrial environment enabling and encouraging such lifestyle-related immune stimuli, with an outcome potential in chronic diseases ranging from metabolic disorders(3) to certain forms of cancer (4). If this is the case, weight loss, while being a useful component of chronic disease management at the clinical level, may not always be either ‘necessary’ or ‘sufficient’, thus changing the nature of any proposed intervention to manage, at the population level, what should be seen as a natural physiological response to an unnatural, obesogenic(5) environment.

Inflammation and ‘metaflammation’

Classical inflammation represents an acute immune reaction to infection or injury. Metaflammation(2) differs from this in that:

(a) it does not involve the classical symptoms of inflammation (tumour, rubor, dolour and calor); (b) it causes only a small rise in immune system markers (i.e. 4–6-fold v. several 100-fold); (c) it results in chronic, rather than acute, allostasis; (d) it has its effects systemically; (e) its antigens are less apparent as foreign agents or microbial organisms (and hence may be better referred to as ‘inducers’(6)); (f) it appears to perpetuate, rather than resolve, a disease(1–4,6). Fig. 1 is a proposed graphical representation of the differences between the two forms of an inflammation.

There may be a number of agents associated with inducers of the metaflammatory response (shown on the right-hand side of Fig. 1), oxidised LDL being a known example. As well as aiding the atherogenic process, the leakage of foam cells from the phagocytosis of oxidised LDL formed in the arterial intima, through unstable plaques into the bloodstream, can have a thrombotic impact(3). What is unclear to date, however, is the type and range of initial inducers of this and other processes stimulating metaflammation.

Metaflammatory inducers

Obesity is a cause of oxidative stress and insulin resistance(3) and hence has been proposed as the primary causative factor in chronic and metabolic diseases such as type 2 diabetes(7). However, depending largely on the site of fat storage, obesity can be relatively benign, with little negative impact on health(8), or metabolic, with apparent links to a range of metabolic and other disorders(9). Also, a significant proportion of obese individuals suffer none of the dysmetabolism expected from obesity, and a significant proportion of lean

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fats(19), and excessive fructose- (20) and glucose-rich responses to a range of other stimuli(14), has enabled researchers to identify a number of inducers of inflammation (both pro- and anti-) in the body. A list of those with supporting evidence is shown in Table 1.

There is a known link between obesity and metaflammation(13), but the causal relationship is not clear. Metaflammation is also associated with lifestyle and environmental factors that in some instances, but not always, cause obesity(12). Awareness of the relationship between postprandial glucose and fat excursions and inflammatory markers(13), and inflammatory responses to a range of other stimuli(14), has enabled researchers to identify a number of inducers of inflammation (both pro- and anti-) in the body. A list of those with supporting evidence is shown in Table 1.

The left-hand side of Table 1 lists inducers with evidence of a pro-inflammatory (metaflammatory) response. As well as obesity and weight gain, this includes excessive alcohol(15), acute excess energy intake(16), a Western-style diet(17) and a range of nutritive factors including saturated(18) and trans-fats(19) and excessive fructose(20) and glucose-rich foods(21). Non-nutritive factors include inadequate sleep(22), smoking(23), stress and depression(24). While some of these (e.g. inactivity, excess energy intake) can cause weight gain, this is not a prerequisite for metaflammation to occur. Nutrient overload from acute excess energy intake, for example, even in the absence of weight gain(25), can abnormally tax the intracellular metabolism, cause acute oxidative stress, possibly disrupt normal protein folding in the endoplasmic reticulum(26) and lead to the accumulation of intracellular metabolites, activating inflammatory pathways and inducing insulin resistance(26). Similarly, a high glycaemic index load, or even an excess of otherwise benign low glycaemic index foods, can have an inflammatory effect in the absence of obesity(27). At the other extreme, a similar response results from extended fasting(28) (probably because of the protective effects of insulin resistance in reducing energy losses). Paradoxically, there are also similar pro-inflammatory effects of both inactivity(29) and excessive exercise(30), suggesting a healthy range of certain lifestyle actions, above or below which there is a negative metabolic outcome.

Fig. 2, which has been expanded from a previous article discussing lifestyle-related determinants of inflammation in adolescence(31), distinguishes between those factors with an aetiological effect on inflammation through obesity, and those with an effect ‘independent’ of it.

Obesity, while often present, may thus not be a necessary condition for metaflammation, or the chronic diseases associated with this. Many of the causes of obesity, on the other hand, as well as other non-obesogenic lifestyle-related inducers, seem to be more directly related aetio-logically to chronic disease outcomes through the development of metaflammation. We have discussed this in more detail elsewhere(12), pointing out that while the metaflammatory process appears to be a persistent causal factor, the association of inflammation with oxidative stress and insulin resistance in metabolic breakdown (described in Fig. 1 as the metabolic ‘milieu’) is not always clear.

### Anti-inflammatory inducers

The pro-inflammatory inducers shown in Table 1 provide some suggestion of a distal causality, as all of these are relatively new to the human environment. This is supported by the identification of the inducers with evidence of an anti-inflammatory effect listed on the right-hand side of Table 1. Anti-inflammatory responses (or neutral responses to corrections of those conditions causing a pro-inflammatory reaction on the left-hand side of Table 1 such as obesity and smoking)

<table>
<thead>
<tr>
<th>Met探flammatory inducers</th>
<th>Anti-inflammatory/neutral</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exercise (excessive)</td>
<td>Exercise/physical activity/CV fitness</td>
</tr>
<tr>
<td>Inactivity/low CV fitness</td>
<td>Walking/yoga/resistance, etc</td>
</tr>
<tr>
<td>Acute nutrient overload</td>
<td>Alcohol (moderate)</td>
</tr>
<tr>
<td>Alcohol (excessive)</td>
<td>Cocoa/dark chocolate</td>
</tr>
<tr>
<td>Energy intake (excessive)</td>
<td>Energy intake (restricted)</td>
</tr>
<tr>
<td>‘Fast’ food/Western-style diet</td>
<td>Mediterranean/varied diet</td>
</tr>
<tr>
<td>Fasting</td>
<td>Fibre (high intake)</td>
</tr>
<tr>
<td>Fat</td>
<td>Fish</td>
</tr>
<tr>
<td>Saturated fat</td>
<td>Fruits/vegetables</td>
</tr>
<tr>
<td>Trans fat</td>
<td>Garlic</td>
</tr>
<tr>
<td>High-fat diet</td>
<td>Herbs and spices</td>
</tr>
<tr>
<td>Low n-3/n-6 fatty acids</td>
<td>Lean game meats</td>
</tr>
<tr>
<td>Fructose</td>
<td>Low-GI foods</td>
</tr>
<tr>
<td>Glucose</td>
<td>Nuts</td>
</tr>
<tr>
<td>High glycaemic/GL foods</td>
<td>Raisins</td>
</tr>
<tr>
<td>Glycaemic load</td>
<td>Tea/green tea</td>
</tr>
<tr>
<td>Refined carbohydrate</td>
<td>Vinegar</td>
</tr>
<tr>
<td>Salt (excessive)</td>
<td>Smoking cessation</td>
</tr>
<tr>
<td>Obesity/weight gain</td>
<td>Weight loss</td>
</tr>
<tr>
<td>Sleep deprivation</td>
<td>Smoking</td>
</tr>
<tr>
<td>Stress</td>
<td>‘burnout’/anxiety/depression</td>
</tr>
</tbody>
</table>

CV, cardiovascular; GI, glycaemic index.

*Stimuli shown on the left-hand side can cause rises in pro-inflammatory markers, often without obesity, whereas those on the right-hand side can cause an opposite reaction, or a reduction of a pro-inflammatory to neutral response (e.g. in the case of smoking cessation and weight loss) often without weight loss. See text and Egger & Dixon(12) for detailed references (also at www.lifestylemedicine.net.au/staging/health-information/lifestyle-medicine-evidence-base/inflammation-database/index.html).
Implications for health policy

From the research evidence accumulated over the last decade, cited here and elsewhere(14,6,12,14), it appears that metaflammation could be the link between a range of lifestyle and environmental factors (including, but not limited to, obesity), and many, if not most, modern chronic diseases. Metaflammation thus becomes to chronic disease what classical inflammation is to injury or microbial invasion, albeit with a different outcome, the latter leading usually to a resolution to homeostasis, but the former leading to dysmetabolism and chronic allostasis (see Fig. 1). Seen in this light, attempts to clinically manage obesity through diets, exercise programmes or medication are unlikely to change chronic disease rates in the community. This by no means negates the importance of weight loss in disease management, as strategies for doing this include changes to the lifestyle causes discussed here. Indeed, a range of simple lifestyle interventions can have a profound effect to just obesity, as many of these, and other metaflammatory inducers: (a) can have acute effects; (b) can lead to a metaflammatory reaction in the absence of obesity (e.g. smoking, sleep loss, depression); or (c) are not, in themselves, causes of obesity (e.g. smoking, environmental pollution).

Factoring pro- and anti-inflammatory causes

A significant factor distinguishing both sides of Table 1 could be labelled ‘modernity’, associated with the modern techno-industrial environment, arising in the main since around the time of the industrial revolution of the late nineteenth century, with inducers on the left-hand side being largely post-industrial revolution (modern), and thus relatively new to the human repertoire. Those on the right-hand side (anti-inflammatory) with obesity, in the dysmetabolism often ascribed to just obesity, as many of these, and other metaflammatory inducers: (a) can have acute effects; (b) can lead to a metaflammatory reaction in the absence of obesity (e.g. smoking, sleep loss, depression); or (c) are not, in themselves, causes of obesity (e.g. smoking, environmental pollution).
the single biggest driving force behind improved health and well-being since the start of the industrial revolution\(^{50}\). The world economic crisis beginning in 2007–8 has signalled to many, however, not just a need for a reordering of the current system, but for a paradigm shift in the growth system of economics\(^{51,52}\), thus echoing the views of its early architects such as Mill\(^{53}\) and Keynes\(^{54}\) that growth would need to be pursued for a time, but eventually would need to be replaced by an alternative system that focused more on human improvement\(^{55}\) (presumably including health). This is because, as recognised then, but often overlooked in the current growth-dominated political environment, nothing can grow forever. As stated by one observer: ‘...after maturity, continued growth is either obesity or cancer’\(^{55}\). This also signals the need for health scientists to become involved in the economic debate.

Second, what is the connection between obesity and chronic disease in the absence of damaging lifestyle cofactors? Does the fit-but-fat phenomenon reduce the need for a population emphasis on weight control in favour of an increased emphasis on other aspects of lifestyle (such as activity levels, sleep, stress management, etc) and the inflammatory environment, with possible beneficial impact on both the biological and ecological environments\(^{14,56}\)? Health and environmental issues, such as pollution and climate change, have recently been shown to be intimately associated through the use of non-renewable fossil fuels in transport, also leading to reduced personal energy expenditure and increased obesity\(^{14,56}\).

Finally, what can be done to counteract this environment-initiated epidemic, given that a return to a pre-industrial society is unlikely? It is not the intention here to detail what will obviously be a complex and multidisciplinary response, which is often considered outside the realm of health scientists, but which we propose is integral to the management of chronic disease. There is a need for an array of workable solutions at many levels of modern society, and away from an isolated view of health and obesity as simply a pharmaceutical problem. While a major economic paradigm shift such as that suggested previously is unlikely to happen in the immediate future, interim initiatives such as corporate and personal carbon trading have begun to be put in place to moderate the effects of unlimited growth, albeit serendipitously through rising interest in climate change\(^{14,56}\), but also with a potential benefit in managing the chronic disease epidemic\(^{12,14,57}\). There are still many answers to be provided, however, and undoubtedly many more questions arising from these suggestions, for which health scientists will need to expand their horizons.

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References
