Mounting evidence indicates that inflammation may play a significant role in the development of depression. Patients with depression exhibit increased inflammatory markers, and administration of cytokines and other inflammatory stimuli can induce depressive symptoms. Mechanisms by which cytokines access the brain and influence neurotransmitter systems relevant to depression have also been described, as have preliminary findings indicating that antagonizing inflammatory pathways may improve depressive symptoms. One primary source of inflammation in depression appears to be adiposity. Adipose tissue is a rich source of inflammatory factors including adipokines, chemokines, and cytokines, and a bidirectional relationship between adiposity and depression has been revealed. Adiposity is associated with the development of depression, and depression is associated with adiposity, reflecting a potential vicious cycle between these two conditions which appears to center around inflammation. Treatments targeting this vicious cycle may be especially relevant for the treatment and prevention of depression as well as its multiple comorbid disorders such as cardiovascular disease, diabetes, and cancer, all of which have also been associated with both depression and inflammation.

An overwhelming amount of evidence indicates that depressed patients exhibit increased markers of innate immune system activation and inflammation. For example, in a meta-analysis of over 50 studies, Howren et al. found that the majority of studies show that depressed patients have elevations in the proinflammatory cytokines, interleukin (IL)-6, and IL-1β as well as the acute phase protein, C-reactive protein (CRP). A recent meta-analysis has revealed that the proinflammatory cytokine, tumor necrosis factor (TNF)-α, is also increased in patients with major depression. In addition to the simple association between depression and inflammatory markers, the administration of inflammatory cytokines such as the innate immune cytokine, interferon (IFN)-α, can induce depression in a high proportion of treated patients. In many ways this is parallel to what is referred to as sickness behavior in animals, which represents an adaptive response to acute infection and other sources of inflammation such as wounding. The sickness response can be induced in laboratory ani-
mals by the acute administration of proinflammatory cytokines such as IL-1β or TNF-α or indirectly via the induction of peripheral immune activation by stimuli such as bacterial endotoxin.12,13 Acute administration of endotoxin as well as other immune stimuli including typhoid vaccination causes a similar sickness syndrome in humans that includes depressed mood, decreased social interaction, sleep disturbance, and anhedonia.14,15 This constellation of symptoms, which parallels that found in major depression, has also been consistently observed during chronic administration of cytokines such as IFN-α and β for illnesses including hepatitis C, multiple sclerosis, and several types of cancers, including malignant melanoma.7 To explore the degree to which cytokine-induced depression parallels depression in ostensibly medically healthy individuals, Capuron et al8 compared 20 patients who were being treated with INFα for malignant melanoma with 28 medically healthy subjects with major depression using the Hamilton Rating Scale for Depression (HAM-D).8 Forty-five percent of the IFN-α-treated patients developed major depression during the 12-week follow-up period. There were minimal differences in the severity of individual depressive symptoms between patients who became depressed during IFN-α treatment versus medically healthy depressed individuals, although IFN-α-treated depressed patients did exhibit more psychomotor retardation and weight loss, and the medically healthy depressed group experienced greater feelings of guilt and thoughts of suicide.8 These results suggest that the depression induced by cytokines is remarkably similar to depression seen in medically healthy depressed patients. Of note, the link between inflammation and depression may explain the frequent association between medical illnesses and depression.17 As shown in Table I, while there are many medical conditions associated with increased rates of depression, the majority of these illnesses are also associated with increased inflammation, including not only infectious diseases and cancer but also cardiovascular disease and diabetes, both of which are now recognized to have an inflammatory component.18 Of note, when depression occurs in the context of medical illness, it has been associated with increased concentrations of inflammatory cytokines. For example, several studies have shown that depressed patients with cancer19-22 or cardiovascular disease23 have higher peripheral blood concentrations of IL-6 and CRP. Moreover, depression scores have been shown to be strongly correlated with blood cytokine concentrations in these patients.24

How do cytokines cause depression?

Access to the brain

Peripheral immune activation, such as that seen with local infection, wounding and/or psychological stress, induces release of IL-1α, IL-1β, IL-6, and TNF-α.5,25-27 However, these cytokines are too large to freely pass through the blood-brain barrier, which raises the question of how a centrally mediated behavioral effect is achieved. Several pathways by which cytokine signals can access the brain have been identified. Local release of cytokines can stimulate peripheral afferent nerve

| Noninflammatory diseases | Infectious diseases |
|-------------------------|--------------------|
| Hypothyroidism          | Neurological diseases |
| Cushing’s disease       | Cerebrovascular disease |
| Porphyria               | Multiple sclerosis |
| - Lewy body disease, etc |
| - Specific neoplasms    | - Lupus, psoriasis, etc |
| - Pancreas              | - Psoriasis |
| - Oropharynx*           | - Rheumatoid arthritis, etc |
| - Breast                | - Diabetes mellitus |
| - Melanoma              | - Inflammatory bowel diseases |
| - Lymphoma, etc         | - Crohn’s, ulcerative colitis |
| - Cardiovascular disease |
| - Connective tissue diseases |

Table I. Inflammatory and noninflammatory diseases associated with elevated rates of depression. *Particularly in the context of combined chemoradiation.
fibers such as the vagus that innervate peripheral tissues, ultimately leading to activation of microglia, which can produce cytokines in the brain. In addition, “leaky” regions in the blood brain barrier such as the circumventricular organs\(^{26-28}\) allow access of peripheral inflammatory mediators to the brain. Cytokines in the peripheral circulation can also cross the blood-brain barrier via saturable active transport molecules expressed on brain endothelial cells.\(^{29}\) Finally, in the context of chronic immune stimulation, microglia activated by peripheral TNF-\(\alpha\) can produce the chemokine, monocyte chemoattractant protein (MCP)-1, which in turn, can attract monocytes into the brain parenchyma.\(^{30}\)

Impact on neurotransmitter metabolism

Once cytokine signals reach the brain, there is a rich literature indicating that they can interact with virtually every pathophysiologic domain relevant to depression, including marked effects on brain monoamines, which are the target of conventional antidepressant medications. Indeed, cytokines have been shown to influence central monoamine synthesis, release, and synaptic reuptake.

Serotonin

Serotonin is synthesized from tryptophan by tryptophan hydroxylase (TH) and aromatic amino acid decarboxylase (AAAD), and the amount of serotonin in brain is highly dependent on tryptophan availability.\(^{31}\) Specifically, depletion of tryptophan rapidly leads to reduced brain serotonin levels, which in turn can precipitate depressive symptoms in vulnerable individuals.\(^{32}\) Activation of the enzyme idoleamine 2,3-dioxygenase—IDO (and the related liver enzyme tryptophan 2,3-dioxygenase) is an alternative pathway for tryptophan metabolism yielding kynurenine (KYN) and leading to tryptophan depletion and ultimately decreased serotonin in brain.\(^{32,33}\) Several cytokines and their signaling pathways have been shown to activate IDO\(^{34,35}\) (for a review see Shelton and Miller\(^{14}\)). Interestingly, peripheral administration of the cytokine-inducer, lipopolysaccharide (LPS) to mice activates IDO and is associated with depressive-like behavior.\(^{36}\) These LPS-induced behavioral changes can be reversed by IDO inhibition using the IDO antagonist 1-methyltryptophan.

IDO activation also has other effects that may be relevant to depression. For example, KYN is metabolized to kynurenic acid (KYNA), which antagonizes \(\alpha_7\) nicotinic acetylcholine receptors\(^{37}\) and can reduce striatal dopamine release (see below)\(^{38,39}\) KYN is also metabolized to quinolinic acid (QUIN); QUIN leads to the generation of toxic lipid peroxides and activates N-methyl-D-aspartic acid (NMDA) receptors and the release of glutamate, all of which can contribute to neurotoxicity.\(^{40}\) The impact of QUIN on neuronal integrity has been implicated in the pathophysiology of several degenerative neurological conditions including Alzheimer’s, Huntington’s, and Parkinson’s diseases, amyotrophic lateral sclerosis, and human immunodeficiency virus-related dementia.\(^{41-47}\) Of note, IFN-\(\alpha\) therapy has also been shown to increase KYN/tryptophan ratios in humans, and KYN has been found to access the brain in IFN-\(\alpha\)-treated patients where it is associated with increased cerebrospinal fluid (CSF) concentrations of both QUIN and KYNA.\(^{48,49}\) CSF KYN and QUIN were in turn correlated with depression in during IFN-\(\alpha\) treatment.

Aside from its impact on tryptophan and serotonin synthesis, immune activation can also affect serotonin availability by acting on synaptic reuptake via the high-affinity serotonin transporter (5HTT).\(^{50}\) Activation of p38 mitogen activated protein kinase (MAPK) by both IL-1\(\beta\) and TNF-\(\alpha\) leads to phosphorylation of 5HTT and increased neuronal uptake of serotonin.\(^{51}\) Expression\(^{52}\) and trafficking of 5HTT to the cell surface\(^{53}\) is also increased by the activation of p38 MAPK. These effects of cytokines on 5HTT expression and function have been observed both in vitro and in vivo. Of note, polymorphisms in the 5HTT gene have also been associated with the development of depression during cytokine (IFN-\(\alpha\)) administration.\(^{54,55}\) The relevance of immune-serotonin interactions is further supported by the observation that serotonin reuptake inhibitors can block the development of depressive symptoms in the context of immune activation. For example, one study\(^{60}\) randomly assigned 40 patients undergoing IFN-\(\alpha\) therapy for malignant melanoma to treatment with the selective serotonin reuptake inhibitor (SSRI) paroxetine or placebo for 12 weeks. Eleven percent of the patients treated with paroxetine developed depression as compared to 45% of the placebo group. Almost all studies of SSRIs\(^{57,58}\) in the context of immune activation have demonstrated benefit in reversing or preventing immunotherapy-induced depressive symptoms.
Translational research

Dopamine

In addition to serotonin, cytokine effects on dopamine metabolism may also be important in the pathophysiology of inflammation-induced depression. Reduced prefrontal and striatal dopamine activity is thought to be associated with symptoms of depression such as decreased motivation, psychomotor slowing, fatigue, and lack of response to rewarding stimuli.68,69 Positron emission tomography imaging studies in humans undergoing IFN-α therapy show increased striatal resting state glucose metabolism,70,71 which is believed to represent increased oscillatory burst activity in neurons normally under tonic inhibition by dopamine. Increased striatal resting state glucose metabolism is also found in other dopamine depletion states including Parkinson’s disease.72,73 Animal studies show that immune stimulation by TNF-α and IFN-α reduce brain and CSF dopamine and its metabolites.74,75 In addition, prodopaminergic agents such as levodopa or psychostimulants improve fatigue and depression symptoms in patients undergoing IFN-α therapy as well as a variety of other conditions associated with inflammation including cancer and systemic HIV infection.76-79 There are several mechanisms by which dopamine may be depleted in the CNS during immune activation, aside from decreased dopamine release secondary to the α7 nicotinic acetylcholine receptor mechanism described above.8 For example, IFN-α9 administration to rodents has been associated with depletion of tetrahydrobiopterin (BH4), a cofactor for tyrosine hydroxylase, the rate-limiting enzyme in dopamine synthesis. Also, in a mechanism similar to the effects of immune activation on SHTT, phosphorylation of the dopamine transporter (DAT) by MAPK kinase (MEK) has been shown to increase cell surface expression of DAT and uptake of dopamine.90 Therefore, relative depletion of synaptic dopamine (via reduced synthesis and release and increased reuptake) may underlie some of the neurovegetative symptoms of sickness behavior and depression, such as low energy, reduced motivation, and reduced response to rewarding stimuli.89,91

The anti-inflammatory effects of antidepressant treatments and the antidepressant effects of anti-inflammatories

There have been a number of in vitro and in vivo studies of antidepressant medications82-85 and other antidepressant treatments such as electroconvulsive therapy86 indicating that antidepressant treatments can reduce proinflammatory factors including IL2, IL-6, TNF-α, and IFN-γ.1 In fact, the available evidence indicates that many antidepressant therapies induce a shift from a Th1 (proinflammatory) to a TH2/TH3 (anti-inflammatory) pattern.86-89,90-101 The IFN-γ to IL10 or IL4 ratio is a measure of relative TH1 to TH2-3 activity, and a number of studies indicate that antidepressants decrease this ratio.82-84,102 Because these effects have been observed both in vitro and in vivo, they do not appear to be dependent on the actions of these drugs on monoamines such as norepinephrine or serotonin, suggesting a direct impact of antidepressant medications on cytokines.93 Therefore, the mechanism of antidepressant action in the context of inflammation-induced depression may be a direct effect on inflammatory factors themselves. There is also a small but significant literature indicating that anti-inflammatory drugs may produce antidepressant effects. Cyclooxygenase 2 (COX-2) activity is increased by proinflammatory cytokines, particularly IL-6, and it, in turn, activates the release of IL-1β and TNF-α103 as well as prostaglandin E2 (PGE2), a central mediator of sickness behavior.104 COX-2 inhibitors have been shown to reverse depression-like behaviors in animal models.105 In addition, the COX-2 rofecoxib has been shown to reduce depressive symptoms in patients with osteoarthritis.106 Adjunctive treatment, the nonselective COX-1 and -2 antagonist acetylsalicylic acid (aspirin), increased remission rates in one open-label study of depressed patients previously nonresponsive to fluoxetine alone.107 A prospective, double-blind, placebo-controlled trial of the COX-2 antagonist celecoxib (400 mg. per day) added to the norepinephrine reuptake inhibitor antidepressant reboxetine (4-10 mg per day) for 6 weeks showed greater effects of the combination treatment than reboxetine alone.108 TNF receptor antagonists such as infliximab, adalimumab, golimumab, and certolizumab pegol, and the TNF receptor fusion protein etanercept have been developed in recent years to treat inflammatory and autoimmune diseases such as psoriasis, rheumatoid arthritis, and Crohn’s disease. Direct actions in depressed patients have not yet been reported. However, one study of etanercept treatment of psoriasis did examine antidepressant effects.109 Six hundred and eighteen patients with moderate to severe psoriasis received double-blind treatment with placebo or 50 mg twice weekly infusion treatment with etanercept for 12 weeks. Patients on
etanercept had greater improvements on measures of depression (as measured by Beck Depression Inventory) than those on placebo. Notably, these improvements were not associated with reduction in psoriatic plaques or joint pain, which indicates a primary effect of TNF antagonism on depression, not simply a cosmetic or analgesic effect. These effects were confirmed in subsequent longer term studies in psoriasis patients and in patients with rheumatoid arthritis. A similar effect has been shown with the TNF-κα monoclonal antibody infliximab.

**Adiposity as a possible causal pathway to depression**

In considering possible sources of inflammation leading to depression, there has been increasing interest in the role of obesity. Rates of overweight and obesity have increased tremendously in recent years in both adults and children. Along with this has been an epidemic of related metabolic conditions like type 2 diabetes, dyslipidemias, cardiovascular and fatty liver disease, and certain forms of cancer. The bulk of evidence links obesity and its attendant complications to inflammation. The possible relationship between depression and obesity appears to be bidirectional, as evidence indicates that being depressed also increases the risk for the subsequent development of obesity, probably mediated, in part, by inactivity.

**Obesity as an inflammatory state**

Adipose tissue is now understood as being a very complex organ system. White adipose tissue (WAT) is the main location for long-term fat storage in the body. WAT, particularly in the abdomen, is the main contributor to metabolic diseases. Adipocytes in WAT secrete a variety of hormones, inflammatory factors including cytokines (referred to as adipocytokines or adipokines). These factors include hormones traditionally associated with adipose tissue such as leptin, adiponectin, resistin, and visfatin; however, adipocytes can also secrete IL-6 and TNF-κα. Nevertheless, one of the primary mechanisms for the induction of inflammation in adipose tissue is the secretion of chemokines, particularly MCP-1, MCP-1 attracts leukocytes such as macrophages, T lymphocytes, and dendritic cells to adipose tissue, which in turn secrete cytokines including IL-1, IL-6, and TNF-κα. Thus, chemokines and cytokines produced by WAT may contribute to widespread immune activation, potentially causing or exacerbating diseases associated with inflammation such as type 2 diabetes, cardiovascular disease, cancer, and depression.

Leptin is another important peptide produced by adipocytes that regulates dietary intake. It regulates appetite by acting on leptin receptors in brain, particularly the hypothalamus. In the case of obesity, a state of leptin resistance develops in which circulating levels are actually increased but responsiveness is reduced. Excess calories in the diet lead to leptin resistance; however, high-fructose feeding is a major contributor. Leptin is a member of the type I cytokine superfamily; it is involved in the modulation of white blood cell response, including T-cell activation and a shift to Th1 cytokine production. Resistin is another proinflammatory adipocytokine produced by both WAT and monocytes. It sets up a positive inflammatory feedback system in which the secretion of resistin is increased by proinflammatory cytokines such as IL-1, IL-6, and TNF-κα, but it also increases the production of these same cytokines by macrophages. By contrast, adiponectin increases fatty acid oxidation and reduces the synthesis of glucose in the liver. Adiponectin, whose levels are reduced in obese persons, has a predominantly inhibitory role in Th1 immune responses, including the inhibition of IL-6 and TNF-κα production and an increase in the anti-inflammatory cytokine IL-10. Therefore, dietary excess, leading to expansion of WAT, produces a shift in the pro- and anti-inflammatory mediators such as leptin, resistin, adiponectin, and other adipocytokines, leading to a general proinflammatory state. This, then, contributes to metabolic derangements and disease such as dyslipidemias, cardiovascular disease, and type 2 diabetes.

The activation of inflammatory factors related to obesity also appears to induce the IDO–KYN pathway. Plasma tryptophan concentrations are reduced and the KYN/tryptophan ratio is increased in obese relative to lean individuals, indicating IDO activation. Weight reduction by diet or bariatric surgery restores a normal KYN/tryptophan balance. This is likely to be the result of a reduction in the proinflammatory state after weight loss. It, then, appears that, like other inflammatory diseases, the immune activation found in obesity may shift metabolism from tryptophan to KYN, which may contribute to depression.
Adiposity and depression

Both depression and obesity, then, are associated with Th1 activation. However, is there evidence of a causal link in either direction—ie, from depression to obesity of vice versa? Some larger-scale epidemiological studies have failed to find a strong association between obesity and depression. Nevertheless, while cross-sectional studies do not show strong correlations between depression and obesity, longitudinal studies tell a very different story. A recent meta-analysis of 15 longitudinal studies showed a bidirectional association between depression and obesity (especially abdominal adiposity) in which prior obesity increases the risk for depression and depression increases the likelihood of subsequent obesity.

To further investigate this bidirectional relationship especially as it pertains to inflammation, Miller et al conducted a mediational analysis evaluating the relationship between serum inflammatory markers (including IL-1β, IL-6, TNF-α, CRP, and MCP-1) in 50 physically healthy young adults with depression and 50 matched controls. IL-6, CRP, and BMI were elevated in the depressed sample compared with controls. When the relationship between depression and both IL-6 and CRP (but not IL-1β) were adjusted for BMI, the results became nonsignificant, indicating a mediational role for adiposity in the relationship between depression and IL-6 and CRP elevation. A separate analysis of the same dataset using structural equation modeling (SEM) estimated the relationship among depression, adiposity, leptin, and inflammation (IL-6 and CRP). The best fit model indicated that the primary causal pathway was from depression to adiposity to inflammation. This was interpreted as indicating that depression leads to increased adiposity (possibly through inactivity) which, in turn, leads to an increase in inflammatory markers.

Diet and depression

Diets in much of the world have shifted to high carbohydrates and a reduction in omega-3 (n-3) (unsaturated) compared with omega-6 (n-6) (saturated) fatty acids. The intake of fish and other sources of n-3 fatty acids appear to be somewhat protective from certain metabolic conditions and epidemiological studies have associated an increased relative intake of fish with a reduced risk for depression. However, it does not seem to be primarily intake of fish per se, but so-called fatty fish with high n-3 concentration (eg, anchovy, sea bass, carp, dogfish, eel, halibut, herring, mackerel, mullet, fish, roe, salmon, sardine, trout, and tuna) that lend protection against both metabolic diseases and depression.

The benefits of the Mediterranean diet pattern

Recent studies have found particular health benefits, including reduction in risk of depression, associated with the so-called Mediterranean Diet Pattern (MDP). As noted in the seminal work by Willett et al, this pattern of eating has been associated historically with good general health and longer life expectancy. This method “is based on food patterns typical of Crete, much of the rest of Greece, and southern Italy in the early 1960s” and “included regular physical activity... abundant plant foods (fruit, vegetables, breads, other forms of cereals, potatoes, beans, nuts, and seeds), fresh fruit as the typical daily dessert, olive oil as the principal source of fat, dairy products (principally cheese and yogurt), and fish and poultry consumed in low to moderate amounts, zero to four eggs consumed weekly, red meat consumed in low amounts, and wine consumed in low to moderate amounts, normally with meals.” This pattern of eating is characterized by lower saturated and total fat content. This manner of eating was shown recently to be associated with reduced risk for depression in a prospective study of the relationship between the MDP and health. A sample of 10,094 healthy persons in Spain were assessed using a validated 136-item item food frequency questionnaire to determine the relative adherence to the MDP, and followed for 4.4 years. Using the lowest adherence to the MDP as the reference condition, adjusted hazard ratios for depression for the higher categories of adherence ranged from 0.74 for modest adherence to 0.49. These results indicate a strong prospective protective effect for the MDP. Of relevance, earlier research found a strong inverse relationship between adherence to the MDP and serum IL-6 with a trend for CRP.
refined sugars such as cane sugar has declined over the last 40 years, the total caloric load from sweeteners has increased; this has primarily been in the form of fructose, particularly in the form of high-fructose corn syrup (also known as “corn sugar”). A high level of fructose intake is associated with obesity and metabolic diseases. Although the specific role of fructose intake, as opposed to increased total calories, has been questioned, it is increasingly clear that high intake of fructose contributes uniquely to problems of obesity and metabolic diseases such as cardiovascular disease, dyslipidemia, and type 2 diabetes. Fructose has a very high extraction ratio by the liver, and does not contribute significantly to problems of obesity and metabolic diseases such as cardiovascular disease, dyslipidemia, and type 2 diabetes. Fructose has a very high extraction ratio by the liver, and does not contribute significantly to increases in insulin or satiety signaling. High levels of fructose loading in the liver leads to the synthesis of triglycerides, which contribute to liver and abdominal fat. The shift in intake from proteins and “healthy” fats to saturated fats and carbohydrates, particularly fructose, has contributed to the worldwide epidemic of obesity.

**Does n-3 fatty acid supplementation reduce depression?**

A recent study indicates that not all n-3 fatty acids reduce inflammation; this study actually showed that docosahexaenoic acid, one constituent of fish oil, may actually increased the ratio of interferon gamma to IL-10, indicating a proinflammatory effect. However, eicosapentaenoic acid (EPA) did not show this effect; EPA has shown to reduce depressive symptoms in a few, small-scale studies. One study randomized 70 persons with major depression not responsive to antidepressants to ethyl-eicosapentaenoic acid (e-EPA) (a specific n-3 fatty acid) 1, 2, or 4 g per day or placebo as add-on therapy. Curiously, the 1 mg per day, but not 2 or 4 mg./day doses was significantly better than placebo. Subsequent studies have supported these results. Of note, a polymorphism in the gene for phospholipase A2, a key enzyme in the metabolism of polyunsaturated fatty acids, was associated with a 3-fold increase in the likelihood of developing major depression during IFN-α treatment as well as lower blood concentrations of EPA.

**Diet, adiposity, and risk for depression in children**

The increase in obesity in adults has been paralleled in children and adolescents, along with an increase in inflammatory diseases previously thought to occur mostly in adults: type 2 diabetes, fatty liver disease, cardiovascular disease, and dyslipidemia. As described earlier for adults, the current evidence suggests a bidirectional relationship between obesity and depression in children. Prior depression in childhood is a relatively strong predictor of the subsequent development of obesity, metabolic syndrome, and related diseases in adult life. Depression may increase risk by changes in diet, eating behavior, and inactivity. Alternatively, baseline obesity may increase risk for depression via increases in inflammation as well as cultural aspects of beauty. Obesity negatively impacts self-esteem based on cultural aspects of beauty and desirability. Obesity also may contribute to risk for depression via effects on physical activity, sleep, and eating behavior.

**Summary and conclusions**

It seems clear at this point that inflammatory mediators, whether they are generated by specific diseases or administered exogenously (as with IFN therapy) can lead to depression. It also appears that a significant subset of depressed patients without known inflammatory disease have inherent upregulation of inflammatory factors, particularly IL-6, TNF-α, and CRP, without other known inflammatory disease. As posited in this paper, one causal pathway for this increased inflammation may be overweight and obesity. Therefore, depression (and the inactivity and diet changes associated with it), obesity, and inflammation may represent a “vicious cycle” (Figure 1). A person may enter this cycle at any point—obesity may lead to inflammation which leads to depression; depression may lead to inactivity and dietary changes, which lead to obesity leading to inflammation; inflammatory diseases may lead to both depression and inactivity, resulting in obesity. Western high-fat, high-carbohydrate diets, as lower blood concentrations of EPA.
bohydrate diets and inactivity may lead to obesity, inflammation, and depression. This cycle may also explain the common association between inflammatory diseases such as lupus or fibromyalgia and both depression and obesity. Therefore, multiple, interacting factors may lead to a general decline in mental and physical health. However, this cycle also provides multiple nodal points for both treatment and prevention. For example, children and adolescents at risk for depression (ie, with positive family history or those who have been traumatized) may represent a group for whom targeted diet and exercise programs would be beneficial to help to prevent or reduce risk for depression. In addition, recent data indicate that overweight and obese patients have reduced response to antidepressant treatments. For example, a recent combined analysis of outcomes in three clinical trials of marketed antidepressants divided participants into normal weight (BMI <25), overweight (BMI 25-<30), and obese (BMI >30). The results indicated progressive resistance to antidepressant therapies from normal weight to obesity. Future interventions could target overweight and obesity as a possible remediable cause of treatment resistance. Depression is a complex condition with many potential causal pathways; two, possibly interrelated mechanisms, diet-associated overweight and obesity and inflammation have been reviewed. Although these mechanisms represent only two among many causal paths, they potentially explain many features, such as the common association between inflammatory diseases and depression. Nevertheless, there is cause for optimism for possible intervention strategies given the evidence for success of lifestyle modifications such as exercise, diet, and other weight loss approaches to inflammatory diseases and obesity.

**REFERENCES**

1. Miller AH, Maletic V, Raison CL. Inflammation and its discontents: the role of cytokines in the pathophysiology of major depression. Biol Psychiatry. 2009;65:732-741.
2. Howren MB, Lamkin DM, Suls J. Associations of depression with C-reactive protein, IL-1, and IL-6: a meta-analysis. Psychosom Med. 2009;71:171-186.
3. Raison CL, Capuron L, Miller AH. Cytokines sing the blues: inflammation and the pathogenesis of depression. Trends Immunol. 2006;27:24-31.
4. Hart BL. Biological basis of the behavior of sick animals. Neurousci. Biobehav Rev. 1988;12:123-137.
5. Dantzer R, O’Connor JC, Freund GG, Johnson RW, Kelley KW. From sickness behaviour to sickness behavior: a study with interleukin-1 type I receptor-deficient mice. Eur J Neurosci. 2000;12:4447-4456.
6. Dantzer R, Cytokine-induced sickness behavior: mechanisms and implications. Annu Rev Med. 2003;54:67-92.
7. Dantzer R, Kelley KW. Twenty years of research on cytokine-induced sickness behavior. Brain Behav Immun. 2007;21:153-160.
8. Hamilton M. A rating scale for depression. J Clin Psychiatry. 1985;9:350-357.
9. Bluthe RM, Laye S, Michaud B, Combe C, Dantzer R, Pernet R. Role of interleukin-1beta and tumour necrosis factor-alpha in lipopolysaccharide-induced sickness behaviour: a study with interleukin-1 type I receptor-deficient mice. Eur J Neurosci. 2000;12:4447-4456.
10. Dantzer R. Cytokine-induced sickness behavior: mechanisms and implications. Annu Rev Med. 2003;54:67-92.
11. Miller AH, Maletic V, Raison CL. Inflammation and its discontents: the role of cytokines in the pathophysiology of major depression. Biol Psychiatry. 2009;65:732-741.
12. Howren MB, Lamkin DM, Suls J. Associations of depression with C-reactive protein, IL-1, and IL-6: a meta-analysis. Psychosom Med. 2009;71:171-186.
13. Raison CL, Capuron L, Miller AH. Cytokines sing the blues: inflammation and the pathogenesis of depression. Trends Immunol. 2006;27:24-31.
14. Hart BL. Biological basis of the behavior of sick animals. Neurousci. Biobehav Rev. 1988;12:123-137.
15. Dantzer R, O’Connor JC, Freund GG, Johnson RW, Kelley KW. From sickness behaviour to sickness behavior: a study with interleukin-1 type I receptor-deficient mice. Eur J Neurosci. 2000;12:4447-4456.
16. Dantzer R, Kelley KW. Twenty years of research on cytokine-induced sickness behavior. Brain Behav Immun. 2007;21:153-160.
17. Hamilton M. A rating scale for depression. J Clin Psychiatry. 1985;9:350-357.
18. Hevener AL, Febbraio MA. The 2009 Stock Conference Report: Inflammation, Obesity and Metabolic Disease. Obes Rev. 2010;11:635-644.
19. Musselelman DL, Miller AH, Porter MR, et al. Higher than normal plasma interleukin-6 concentrations in cancer patients with depression: preliminary findings. Am J Psychiatry. 2001;158:1252-1257.
20. Soygar H, Palaoglu O, Akarsu ES, et al. Interleukin-6 levels and HPA axis activity in breast cancer patients with major depressive disorder. Prog Neuropsychopharmacol Biol Psychiatry. 2007;31:1242-1247.
21. Lutgendorf SK, Weinrib AZ, Penedo F, et al. Interleukin-6, cortisol, and depressive symptoms in ovarian cancer patients. J Clin Oncol. 2008;26:4820-4827.
22. Juhn CF, Kuehnhardt D, Bartholomae A, et al. Biomarkers of depression in cancer patients. Cancer. 2006;107:2723-2729.
23. Miller GE, Freedland KE, Duntley S, Carney RM. Relation of depressive symptoms to C-reactive protein and pathogen burden (cytomegalovirus, herpes simplex virus, Epstein-Barr virus) in patients with earlier acute coronary syndromes. Am J Cardiol. 2005;95:317-321.
24. Jacobson CM, Rosenfeld B, Pessin H, Breitbart W. Depression and IL-6 blood plasma concentrations in advanced cancer patients. Psychosomatics. 2008;49:64-66.
25. Dantzer R, Kelley KW. Twenty years of research on cytokine-induced sickness behavior. Brain Behav Immun. 2007;21:153-160.
26. Bierhaus A, Wolf J, Andrassy M, et al. A mechanism converting psycosocial stress into mononuclear cell activation. Proc Natl Acad Sci U S A. 2003;100:1920-1925.
27. Pace TW, Mietzko TC, Alaghe O, et al. Increased stress-induced inflammatory responses in male patients with major depression and increased early life stress. Am J Psychiatry. 2006;163:1630-1633.
28. Ermisch A, Ruhle HJ, Landgraf R, Hess J. Blood-brain barrier and peptides. J Cereb Blood Flow Metab. 1985;5:350-357.
La inflamación en la depresión: ¿es la adiposidad una causa?

Existe una evidencia creciente que señala que la inflamación puede jugar un papel significativo en el desarrollo de la depresión. Los pacientes con depresión muestran aumentados marcadores inflamatorios, y la administración de citocinas y otros estimuladores inflamatorios pueden inducir síntomas depresivos. También se han descrito mecanismos a través de los cuales las citocinas tienen acceso al cerebro y afectan los sistemas de neurotransmisión importantes en la depresión, y se cuenta con hallazgos preliminares que indican que el antagonizar las vías inflamatorias puede mejorar los síntomas depresivos. Una fuente primaria de inflamación en la depresión parece ser la adiposidad. El tejido adiposo es una fuente fuente de factores inflamatorios que incluyen las adipokinas, las quemoquinas y las citocinas, y también se ha revelado una relación bidireccional entre adiposidad y depresión. La adiposidad está asociada con el desarrollo de la depresión y la depresión está asociada con la adiposidad, lo que refleja un potencial círculo vicioso entre estas dos condiciones que parece estar centrado en la inflamación. Los tratamientos que se enfocan en este círculo vicioso pueden ser especialmente relevantes para el tratamiento y prevención de la depresión como de sus múltiples trastornos comórbidos como la enfermedad cardiovascular, la diabetes y el cáncer, todos los cuales también se han asociado con la depresión y la inflamación.

L’inflammation dans la dépression : l’adiposité en est-elle une cause ?

Un faisceau d’arguments sont en faveur d’un rôle significatif de l’inflammation dans le développement de la dépression. En effet, les patients déprimés présentent une augmentation des marqueurs inflammatoires, et l’administration de cytokines et d’autres stimuli inflammatoires peut induire des symptômes dépressifs. Des mécanismes par lesquels les cytokines ont accès au cerveau et influent sur les systèmes neurotransmetteurs liés à la dépression ont aussi été décrits, des résultats préliminaires ayant indiqué que le fait d’antagoniser des voies inflammatoires pouvait améliorer les symptômes dépressifs. L’adiposité semble une des premières sources d’inflammation dans la dépression. Le tissu adipeux est une source importante de facteurs inflammatoires comme les adipokines, les chémo- kines et les cytokines et il existe une relation bidirectionnelle entre adiposité et dépression. En effet, l’adiposité est associée au développement de la dépression et la dépression est associée à l’adiposité, traduisant un cercle vicieux potentiel entre ces pathologies centrées autour de l’inflammation. Des traitements visant ce cercle vicieux peuvent être particulièrement pertinents dans le traitement et la prévention de la dépression et de ses multiples comorbidités comme la maladie cardiovasculaire, le diabète et le cancer, qui sont aussi associés à la dépression et à l’inflammation.

29. Quan N, Banks WA. Brain-immune communication pathways. Brain Behav Immun. 2007;21:727-735.
30. D’Mello C, Le T, Swain MG. Cerebral microglia recruit monocytes into the brain in response to tumor necrosis factoralpha signaling during peripheral organ inflammation. J Neurosci. 2009;29:2089-2102.
31. Delgado PL, Chamey DS, Price LH, Aghajanian GK, Landis H, Heninger GR. Serotonin function and the mechanism of antidepressant action. Reversal of antidepressant-induced remission by rapid depletion of plasma tryptophan. Arch Gen Psychiatry. 1990;47:411-418.
32. Schwarz R, Pellicciani R. Manipulation of brain kynurenines: gial targets, neuronal effects, and clinical opportunities. J Pharmacol Exp Ther. 2002;303:1-10.
33. Schrooksadel K, Wirleitner B, Winkler C, Fuchs D. Monitoring tryptophan metabolism in chronic immune activation. Curr Chin Acta. 2006;364:82-90.
34. Popov A, Abraham Z, Wickenhauser C, et al. Indoleamine 2,3-dioxygenase-expressing dendritic cells form suppurative granulomas following Listeria monocytogenes infection. J Clin Invest. 2006;116:3160-3170.
35. Takikawa O, Tagawa Y, Iwakura Y, Yoshida R, Truscott RJ. Interferon-gamma-dependent/independent expression of indoleamine 2,3-dioxygenase. Studies with interferon-gamma-knockout mice. Adv Exp Med Biol. 1999;467:553-557.
36. O’Connor JC, Lawson MA, Andre C, et al. Lipopolysaccharide-induced depressive-like behavior is mediated by indoleamine 2,3-dioxygenase activation in mice. Mol Psychiatry. 2009;14:511-522.
37. Rassoulpour A, Wu HQ, Ferre S, Schwarz R. Nanomolar concentrations of kynurenine acid reduce extracellular dopamine levels in the striatum. J Neurochem. 2005;93:762-765.
38. Amori L, Wu HQ, Marinozzi M, Pellicciari R, Guidetti P, Schwarz R. Specific inhibition of kynurenate synthesis enhances extracellular dopamine levels in the rodent striatum. Neurosci. 2009;159:196-203.
39. Jang S, Jeong HS, Park JS, et al. Neuroprotective effects of (-)-epigallocatechin-3-gallate against quinolinic acid-induced excitotoxicity via the pi3k pathway and NO inhibition. Brain Res. 2010;1313:25-33.
40. Oswe-Young R, Webster NL, Mukhtar M, et al. Kynurenine pathway metabolism in human blood-brain-barrier cells: implications for immune tolerance and neurotoxicity. J Neurochem. 2008;105:1346-1357.
41. Kwidzinski E, Bechmann I. IDO expression in the brain: a double-edged sword. J Mol Med. 2007;85:1351-1359.
42. Brew BJ, Halman M, Catalan J, et al. Factors in AIDS dementia complex trial design: results and lessons from the abacavir trial. PLoS Clin Trials. 2007;2:e13.
43. Guillemín GJ, Kerr SJ, Brew BJ. Involvement of quinolinic acid in AIDS dementia complex. Neurotox Res. 2005;7:103-123.
44. Zadori D, Klivenyi P, Vamos E, Fulep F, Toldi J, Vecsei L. Kynurenines in chronic neurodegenerative disorders: future therapeutic strategies. J Neural Transm. 2009;116:1403-1409.

45. Vamos E, Pardutz A, Klivenyi P, Toldi J, Vecsei L. The role of kynurenines in disorders of the central nervous system: possibilities for neuroprotection. J Neural Sci. 2009;283:21-27.

46. Mosley RL, Benner EJ, Kadiu I, et al. Neuroinflammation, oxidative stress and the pathogenesis of parkinson’s disease. Clin Neurosci Res. 2006;6:261-281.

47. Sas K, Robotka H, Toldi J, Vecsei L. Mitochondria, metabolic disturbances, oxidative stress and the kynurenine system, with focus on neurodegenerative disorders. J Neuro Sci. 2007;257:221-239.

48. Wickers MC, Koek GH, Robaey G, Verkerk R, Scharpe S, Maes M. IDO and interferon-alpha-induced depressive symptoms: a shift in hypothesis from tryptophan depletion to neurotoxicity. Mol Psychiatry. 2005;10:538-544.

49. Raison CL, Dantzer R, Kelley KW, et al. CSF concentrations of brain tryptophan and kynurenines during immune stimulation with IFN-alpha: relationship to CNS immune responses and depression. Mol Psychiatry. In press.

50. Blakely RD, Berson HE. Molecular biology of serotonin receptors and transporters. Clin Neuropharmacol. 1992;15(suppl 1 Pt A):35A-352A.

51. Zadori D, Klivenyi P, Dostmann WR, Hewlett WA, Blakely RD. p38 MAPK activation elevates serotonin transport activity via a trafficking-independent, protein phosphatase 2A-dependent process. J Biol Chem. 2005;280:15649-15658.

52. Tsao CW, Lin YS, Cheng JT et al. Interferon-alpha-induced serotonin uptake in Jurkat T cells via mitogen-activated protein kinase and transcriptional regulation of the serotonin transporter. J Pharmacol Exp Ther. 2008;22:753-768.

53. Samuel DJ, Jayanthi LD, Bhat NR, Ramamoorthy S. A role for p38 mitogen-activated protein kinase in the regulation of the serotonin transporter: evidence for distinct cellular mechanisms involved in transporter surface expression. J Neurosci. 2005;25:29-41.

54. Lotrich FE, Ferrell RE, Rabinvitz M, Pollock BG. Risk for depression during interferon-alpha treatment is affected by the serotonin transporter polymorphism. Biol Psychiatry. 2009;65:344-348.

55. Muller DJ, Hueso-Diaz P, Binder EB, et al. Functional polymorphisms in the interleukin-6 and serotonin transporter genes, and depression and fatigue induced by interferon-alpha and ribavirin treatment. Mol Psychiatry. 2009;14:1095-1104.

56. Musumeci DL, Lawson DH, Gumnick JF, et al. Paroxetine for the prevention of depression induced by high-dose interferon alfa. N Engl J Med. 2001;344:961-966.

57. Raison CL, Woolwine BJ, Demetrashvili MF, et al. Paroxetine for prevention of major depressive symptoms induced by interferon-alpha and ribavirin for hepatitis C. Aliment Pharmacol Ther. 2007;25:1163-1174.

58. Morasco BJ, Rifai MA, Loftis JM, Indest DW, Hu F, et al. Interferon and interleukin-10 secretion. Neuropharmacology. 2005;45:108-114.

59. Shuto H, Kataoka Y, Horikawa T, Fujihara N, Oishi R. Repeated interferon-alpha treatment of interferon-alpha-induced anhedonia. Psychiatry Clin Neurosci. 2008;62:1108-1114.

60. Schwartz AL, Thompson JA, Masood N. Interferon-induced fatigue in patients with hepatitis C. Mol Psychiatry. 2002;7:942-947.

61. Salamone JD, Correa M. Dopamine/adenosine interactions involved in effort-related aspects of food motivation. Appetite. 2009;53:422-425.

62. Dunlop BW, Nemiroff CB. The role of dopamine in the pathophysiology of depression. Arch Gen Psychiatry. 2007;64:327-337.

63. Capuron L, Pagnoni G, Demetrashvili MF, et al. Prefrontal cortical hypometabolism during low-dose interferon alpha treatment. Psychopharmacology. 2000;152:383-389.

64. Spetsiers PG, Moeller JR, Dhawan V, Ishikawa T, Eidelberg D. Visualizing interleukin-6 and serotonin transporter genes, and depression and fatigue induced by interferon-alpha and ribavirin treatment. J Neurosci. 2009;344:961-966.

65. Bassi A, Bassi F, Onori M, et al. Antidepressant reversal of interferon-alpha associated depression in psychiatric risk patients with chronic hepatitis C. Neuropsychopharmacology. 2007;32:2384-2392.

66. Loftis JM, Socherman RE, Howell CD, et al. Association of interferon-alpha induced depression and improved response treatment in patients with hepatitis C. Neurosci Lett. 2004;365:87-91.

67. Hauser P, Khosla J, Aurora H, et al. A prospective study of the incidence and open-label treatment of interferon-induced major depressive disorder in patients with hepatitis C. Mol Psychiatry. 2002;7:942-947.

68. Capuron L, Pagnoni G, Demetrashvili MF, et al. Basal ganglia hypermetabolism and symptoms of fatigue during interferon-alpha therapy. Neuropsychopharmacology. 2007;32:2384-2392.

69. Juengling FD, Ebert D, Gut O, et al. Preferential cortical hypometabolism during low-dose interferon alpha treatment. Psychopharmacology. 2000;152:383-389.

70. Spetsiers PG, Moeller JR, Dhawan V, Ishikawa T, Eidelberg D. Visualizing the evolution of abnormal metabolic networks in the brain using PET. Computerized Medical Imaging and Graphics. 2005;19:295-306.

71. Felgin A, Fukuda M, Dhawan V, et al. Metabolic correlates of levodopa response in Parkinson’s disease. Neurology. 2001;57:2083-2088.

72. Felger JC, Alagbe O, Hu F, et al. Effects of interferon-alpha on rhesus monkeys: a nonhuman primate model of cytokine-induced depression. Biol Psychiatry. 2007;62:1324-1333.

73. Shuto H, Kataoka Y, Horikawa T, Fujihara N, Oishi R. Repeated interferon-alpha administration inhibits dopaminergic neural activity in the mouse brain. Brain Research. 1997;747:348-351.

74. Lou JS, Kearsn G, Benice T, Oken B, Sexton G, Nutt J. Levodopa improves physical fatigue in Parkinson’s disease: a double-blind, placebo-controlled, crossover study. Mov Disord. 2003;18:1108-1114.

75. Schwartz AL, Thompson JA, Masood N. Interferon-induced fatigue in patients with melanoma: a pilot study of exercise and methylphenidate. Oncol Nurs Forum. 2002;29:885-900.

76. Breitbart W, Rosenfeld B, Kaim M, Funesti-Esch J. A randomized, double-blind, placebo-controlled trial of psychostimulants for the treatment of fatigue in ambulatory patients with human immunodeficiency virus disease. Arch Intern Med. 2001;161:411-420.

77. Kitagami T, Yamada K, Miura H, Hashimoto R, Nabeshima T, Ohta T. Mechanism of systemically injected interferon-alpha impeding monoamine biosynthesis in rats: role of nitric oxide as a signal crossing the blood-brain barrier. Brain Research. 2003;978:104-114.

78. Moron JA, Zakharaov I, Ferrer JV, et al. Mitogen-activated protein kinase regulates dopamine transporter surface expression and dopamine transport capacity. J Neurosci. 2003;23:8480-8488.

79. Brustolim D, Ribeiro-dos-Santos R, Kast RE, Alschuler EL, Soares MB. A new chapter opens in anti-inflammatory treatments: the antidepressant bupropion lowers production of tumor necrosis factor-alpha and interferon-gamma in mice. Int Immunopharmacol. 2006;6:903-907.

80. Szuster-Ciesielka A, Tustanowska-Stachura A, Slotwinska M, Marmurwos-Michalowska H, Kandefer-Szerszen M. In vitro immunoregulatory effects of antidepressants in healthy volunteers. Pol J Pharmacol. 2003;55:353-362.

81. Bengtson BO, Zhu J, Thorell LH, Olsson T, Link H, Waldner J. Effects of zimeldine and its metabolites, clomipramine, imipramine and maprotiline in experimental allergic neuritis in Lewis rats. J Neuroimmunol. 1992;39:109-122.

82. Song C, Dinan T, Leonard BE. Changes in immunoglobulin, complement and acute phase protein levels in the depressed patients and normal controls. J Affect Disord. 1994;30:283-288.

83. Zhou J, Bengtsson BO, Mix E, et al. Clomipramine and imipramine suppress clinical signs and T and B cell response to myelin proteins in experimental autoimmune neuritis in Lewis rats. J Autoimmun. 1998;11:319-327.

84. Zhu J, Bengtsson BO, Mix E, Thorell LH, Olsson T, Link H. Effect of monoamine reuptake inhibiting antidepressants on major histocompatibility complex expression on macrophages in normal rats and rats with experimental allergic neuritis (EAN). Immunopharmacology. 1994;27:225-244.

85. Maes M, Song C, Lin AH, et al. Negative immunoregulatory effects of antidepressants: inhibition of interferon-gamma and stimulation of interleukin-10 secretion. Neuropsychopharmacology. 1999;20:370-379.
88. Kubera M, Lin AH, Kenis G, Bosmans E, van BD, Maes M. Anti-inflammatory effects of antidepressants through suppression of the interferon-gamma/ interleukin-10 production ratio. J Clin Psychopharmacol. 2001;21:199-206.

89. Kubera M, Kenis G, Bosmans E, et al. Suppressive effect of TRH and imipramine on human interferon-gamma and interleukin-10 production in vitro. Pol J Pharmacol. 2000;52:481-486.

90. Kubera M, Maes M, Holan V, Basta-Kaim A, Roman A, Shani J. Prolonged desipramine treatment increases the production of interleukin-10, an anti-inflammatory cytokine, in C57BL/6 mice subjected to the chronic mild stress model of depression. J Affect Disord. 2001;63:171-178.

91. Kubera M, Simbirtsev A, Mathison R, Maes M. Effects of repeated fluoxetine and citalopram administration on cytokine release in C57BL/6 mice. Psychiatry Res. 2000;20:96:255-266.

92. Kubera M, Kenis G, Bosmans E, et al. Plasma levels of interleukin-6, interleukin-10, and interleukin-1 receptor antagonist in depression: comparison between the acute state and after remission. Pol J Pharmacol. 2000;52:237-241.

93. Mohr DC, Goodkin DE, Isler J, Hauser SL, Genain CP. Treatment of depression is associated with suppression of nonspecific and antigen-specific TH1 responses in multiple sclerosis. Arch Neurol. 2001;58:1081-1086.

94. Seidel T, Arolt V, Hunstiger M, Rink L, Behnisch A, Kirchner H. Cytokine production and serum proteins in depression. Scand J Immunol. 1995;41:534-538.

95. Obuchowicz E, Kowalski J, Labuzek K, Krysik S, Rendicz J, Herman ZS. Amitriptyline and nortriptyline inhibit interleukin-1 release by rat mixed glial and microglial cell cultures. Int J Psychopharmacol. 2006;9:27-35.

96. Lanquillon S, Krieger JC, Bening-Abu-Shach U, Vedder H. Cytokine production and treatment result in major depressive disorder. Neuropsychopharmacology. 2002;27:370-379.

97. Słuzewska A, Rybakowski JK, Lacik M, Mackiewicz A, Sobieska M, Wiktoryczk K. Interleukin-6 serum levels in depressed patients before and after treatment with fluoxetine. Ann N Y Acad Sci. 1995;762:474-476.

98. Basterzi AD, Aydemir C, Kisa C, et al. IL-6 levels decrease with SRI treatment in patients with major depression. Hum Psychopharmacol. 2000;15:473-476.

99. Hestad KA, Tonseth S, Stoen CD, Ueland T, Aukrust P. Raised plasma levels of tumor necrosis factor alpha in patients with depression: normalization during electroconvulsive therapy. J ECT. 2003;19:183-188.

100. Muller N, Schwarz MJ. The immune-mediated alteration of serotonin and glutamate: towards an integrated view of depression. Mol Psychiatry. 2007;12:984-991.

101. Myint AM, Leonard BE, Steinbusch HW, Kim YK. Th1, Th2, and Th3 cytokine alterations in major depression. J Affect Disord. 2005;88:167-173.

102. Myint AM, Steinbusch HW, Goehgen H, Luchtman D, Kim YK. Leukocyte cytokine profiles in major depression: a pilot study. J Neuropsychopharmacol. 2006;29:969-707.

103. Collantes-Estevez E, Fernandez-Perez C. Improved control of osteoarthritis pain and self-reported health status in non-responders to celecoxib switched to rofecoxib: results of PAVIA, an open-label post-market-ing survey in Spain. Curr Med Res Opin. 2003;19:402-410.

104. Kumbhakar SS, Garber AJ, Regier DA, Bucholz K, H ser, et al. The association of substance use and HIV infection with trajectories of cigarette smoking in young adult HIV seronegative men and women. J Clin Psychiatry. 2006;67:680-684.

105. Tybring S, Gottlieb A, Papp K, et al. Etanercept and clinical outcomes, fatigue, and depression in psoriasis: double-blind placebo-controlled randomised phase III trial. Lancet. 2006;367:29-35.

106. Dauden E, Griffiths C, Ortonne JP, et al. Improvements in patient-reported outcomes in moderate-to-severe psoriasis patients receiving continuous or paused etanercept treatment over 54 weeks: the CRYSTEL study. J Eur Acad Dermatol Venereol. 2009;23:1374-1382.

107. Krishnan R, Cella D, Leonardi C, et al. Effects of etanercept therapy on fatigue and symptoms of depression in subjects treated for moderate to severe plaque psoriasis for up to 96 weeks. Br J Dermatol. 2007;157:1275-1277.

108. Kekow J, Moots RJ, Emery P, et al. Patient-reported outcomes improve with etanercept plus methotrexate in early active rheumatoid arthritis and the improvement is strongly associated with remission: the COMET trial. Ann Rheum Dis. 2010;69:222-225.

109. Feldman SR, Gottlieb AB, Bala M, et al. Infliximab improves health-related quality of life in the presence of comorbidities among patients with moderate-to-severe psoriasis. Br J Dermatol. 2008;159:704-710.

110. Lichtenstein EHR, Bala M, Han C, DeWoody K, Schaible T. Infliximab improves quality of life in patients with Crohn’s disease. Inflamm Bowel Dis. 2002;8:237-243.

111. Bornstein SR, Ehgart-Bornstein M, Wong ML, Licinio J. Is the worldwide epidemic of obesity a communicable feature of globalization? Exp Clin Endocrinol Diabetes. 2008;116(suppl 1):S30-S32.

112. Cumming BE, O’Ziurt E, Ethan I, Demir S, Karllidag R. Total antioxidant capacity and total oxidant status in patients with major depression: impact of antidepressant treatment. Psychiatr Clin Neurosci. 2009;63:639-645.

113. Wilborn C, Beckham J, Campbell B, et al. Obesity: prevalence, theories, medical consequences, management, and research directions. Int J Soc Sports Nutr. 2005;2:4:314-31.

114. Wang Y, Lobstein T. Worldwide trends in childhood overweight and obesity. Int J Pediatr Obes. 2006;1:11-25.

115. MacPhee M. Global childhood obesity: how to curb an epidemic. J Pediatr Nurs. 2008;23:1-4.

116. Ben-Sefer E, Ben-Natan M, Ehrenfeld M. Childhood obesity: current literature, policy and implications for practice. Int Nurs Rev. 2009;56:166-173.

117. Bellantoni S, Marino M. Epidemiology and natural history of non-alcoholic fatty liver disease (NAFLD). Ann Hepatol. 2009;8(suppl 1):58-54-S58.

118. Golden SH, Robinson KA, Saldanha I, Anton B, Ladenson PW. Clinical review: prevalence and incidence of endocrine and metabolic disorders in the United States: a comprehensive review. J Clin Endocrinol Metab. 2009;94:1853-1878.

119. Hevener AL, Febbraio MA. The 2009 Stock Conference Report: inflammation, Obesity and Metabolic Disease. Obes Rev. 2010;11:635-644.

120. Shoelson SE, Herrero L, Naaz A. Obesity, inflammation, and insulin resistance. Gastroenterology. 2007;132:2169-2180.

121. Sutherland JP, McKinley B, Erkel RH. The metabolic syndrome and inflammation. Metab Syndr Relat Disord. 2004;2:82-104.

122. Dandoma P, Alajda A, Chaudhuri A, Mohanty P, Garg R. Metabolic syndrome: a comprehensive perspective based on interactions between obesity, diabetes, and inflammation. Circulation. 2005;111:1488-1454.

123. Miller GE, Freedland KE, Carney RM, Stetler CA, Banks WA. Pathways linking depression, adiposity, and inflammatory markers in healthy young adults. Brain Behav Immun. 2003;17:276-285.

124. Mathieu P, Poirier P, Pibarot P, Lemieux I, Despres JP. Visceral obesity: the link among inflammation, hypertension, and cardiovascular disease. Hypertension. 2009;53:577-584.

125. Calabro P, Yeh ET. Intra-abdominal adiposity, inflammation, and cardiovascular risk: new insight into global cardiometabolic risk. Curr Hypertens Rep. 2008;10:32-38.

126. Despres JP, Arsenault BJ, Cote M, Cartier A, Lemieux I. Abdominal obesity: the cholesterol of the 21st century? Can J Cardiol. 2008;24(suppl D):70-D12.

127. Tils H, Moschen AR. Adipocytokines: mediators linking adipose tissue, inflammation and immunity. Nat Rev Immunol. 2006;6:772-783.

128. Tils H, Moschen AR. Inflammatory mechanisms in the regulation of insulin resistance. Mol Med. 2008;14:222-231.

129. Carr MW, Roth SJ, Luther E, Rose SS, Springer TA. Monocyte chemoattractant protein 1 acts as a T-lymphocyte chemoattractant. Proc Natl Acad Sci U S A. 1994;91:3652-3656.
133. Xu LL, Warren MK, Rose WL, Gong W, Wang JM. Human recombinant monocyte chemotactic protein and other C-C-chemokines bind and induce directional migration of dendritic cells in vitro. J Leukoc Biol. 1996;60:365-371.

134. Brennan AM, Mantzoros CS. Drug insight: the role of leptin in human physiology and pathophysiology—emerging clinical applications. Nat Clin Pract Endocrinol Metab. 2006;2:318-327.

135. Shapiro A, Mu W, Roncal C, Cheng KY, Johnson RJ, Scarpie PJ. Fructose-induced leptin resistance exacerbates weight gain in response to subsequent high-fat feeding. Am J Physiol Regul Integr Comp Physiol. 2008;295:R1370-R1375.

136. Vasselli JR. Fructose-induced leptin resistance: discovery of an unsuspected form of the phenomenon and its significance. Focus on "fructose-induced leptin resistance exacerbates weight gain in response to subsequent high-fat feeding," by Shapiro, et al. Am J Physiol Regul Integr Comp Physiol. 2008;295:R1365-R1369.

137. Lago F, Dieguez C, Gomez-Reino J, Gualillo O. Adipokines as emerging mediators of immune response and inflammation. Nat Clin Pract Rheumatol. 2007;3:716-724.

138. Lago F, Gomez R, Gomez-Reino JJ, Dieguez C, Gualillo O. Adipokines as novel modulators of lipid metabolism. Trends Biochem Sci. 2009;34:500-511.

139. Silwao N, Singh AK, Aruna B, Mukhopadhyay S, Ghosh S, Ehtesham NZ. Human resistin stimulates the pro-inflammatory cytokines TNF-[alpha] and IL-12 in macrophages by NF-[kappa]B-dependent pathway. Biochem Biophys Res Commun. 2005;334:1092-1101.

140. Bays HE, Gonzalez-Campoy JM, Bray GA, et al. Pathogenic potential of adipose tissue and metabolic consequences of adipocyte hypertrophy and increased visceral adiposity. Exp Rev Cardiaco Ther. 2008;6:343-368.

141. Gustafson B. Adipose tissue, inflammation and atherosclerosis. J Atheroscler Thromb. 2010;17:332-342.

142. Bremme L, Rasmussen MH, Hilsted J, Fernstrom JD. Twenty-four-hour plasma tryptophan concentrations and ratios are below normal in obese subjects and are not normalized by substantial weight reduction. Am J Clin Nutr. 2003;77:1112-1118.

143. Brandacher G, Winkler C, Aigner F, et al. Bariatric surgery cannot prevent tryptophan depletion due to chronic immune activation in morbidly obese patients. Obes Surg. 2006;16:541-548.

144. Murphy JM, Horton NJ, Burke JD, Jr., et al. Obesity and weight gain in relation to depression: findings from the Stirling County Study. Int J Obes (Lond). 2009;33:335-341.

145. Atlants E, Baker M. Obesity effects on depression: systematic review of epidemiological studies. Int J Obes (Lond). 2008;32:881-891.

146. Vogelzangs N, Kritchevsky SB, Beekman AT, et al. Obesity and onset of depressive episodes in middle-aged French men and women. Prostaglandins, Leukotrienes Essential Fatty Acids. 2008;78:171-182.

147. Malhotra AK, Goldman D, Mazzanti C, Clifton A, Breier A, Pickar D. A functional serotonin transporter (5-HTT) polymorphism is associated with psychosis in neuroleptic-free schizophrenics. J Psychosom Res. 1998;33:328-332.

148. Willett WC, Sacks F, Trichopoulou A, et al. Mediterranean diet pyramid: a cultural model for healthy eating. Am J Clin Nutr. 1995;61:1402S-1406S.

149. Tortsa A, Bes-Rastrollo M, Sanchez-Villegas A, et al. Mediterranean diet inversely associated with the incidence of metabolic syndrome. Diabetes Care. 2007;30:2957-2959.

150. Vogelzangs N, Delgado-Rodriguez M, Alonso A, et al. Association of the Mediterranean dietary pattern with the incidence of depression: the Seguimiento Universidad de Navarra (SUN) Cohort. Arch Gen Psychiatry. 2009;66:1090-1098.

151. Dai J, Miller AH, Bremer JD, et al. Adherence to the Mediterranean diet is inversely associated with circulating interleukin-6 among middle-aged men: a twin study. Circulation. 2008;117:169-175.

152. Marriott BP, Cole N, Lee E. National estimates of dietary fructose intake increased from 1977 to 2004 in the United States. J Nutr. 2009;139:12385-12395.

153. Johnson RJ, Segal MS, Sautin Y, et al. Potential role of sugar (fructose) in the epidemic of hypertension, obesity and the metabolic syndrome, diabetes, kidney disease, and cardiovascular disease. Am J Clin Nutr. 2007;86:899-906.

154. Zimmermann MB, Aeberli I. Dietary determinants of subclinical inflammation, dyslipidaemia and components of the metabolic syndrome in overweight children: a review. Int J Obes (Lond). 2008;32(suppl 6):S11-518.

155. Rodriguez-Leyva D, Dupasquier C, McCullough R, Pierce GN. The cardiovascular effects of fructose and its omega-3 fatty acid, alpha-linolenic acid. Can J Cardiol. 2010;26:489-496.

156. Buchhun-Bedient S, Carpenter DO. Benefits versus risks associated with consumption of fish and other seafood. Rev Environ Health. 2010;25:161-191.

157. Chang CL, Seo T, Du CB, Accili D, Deckelbaum RJ. n-3 Fatty acids decrease arterial low-density lipoprotein cholesterol delivery and lipoprotein lipase levels in insulin-resistant mice. Arterioscler Thromb Vasc Biol. 2010;30:2510-2519.

158. Hassan KS, Hassan SK, Hijazi EG, Khazim KO. Effects of omega-3 on lipid profile and inflammation markers in peritoneal dialysis patients. Ren Fail. 2010;32:1031-1035.

159. Defilippis AP, Blaha MJ, Jacobson TA. Omega-3 fatty acids for cardiovascular disease prevention. Curr Treat Options Cardiovasc Med. 2010;12:365-380.

160. Harris WS. The omega-3 index: clinical utility for therapeutic intervention. Curr Cardiol Rep. 2010;12:503-508.

161. Mazza M, Pomponi M, Janiri L, Bria P, Mazza S. Omega-3 fatty acids and antioxidants in neurological and psychiatric diseases: an overview. Prog Neuropharmacol Biol Psychiatry. 2010;31:26-27.

162. Juturu V. Omega-3 fatty acids and the cardiometabolic syndrome. J Cardiometab Syndr. 2008;3:244-253.

163. Ruxton CH, Reed SC, Simpson MJ, Millington JK. The health benefits of omega-3 polyunsaturated fatty acids: a review of the evidence. J Hum Nutr Diet. 2004;17:449-459.

164. Kraguljac NV, Montori VM, Pavuluri M, Chai HS, Wilson BS, Uenal SS. Efficacy of omega-3 fatty acids in mood disorders - a systematic review and metaanalysis. Psychopharmac Bull. 2009;42:39-54.

165. Astorg P, Couthouis A, Bertrais S, et al. Association of fish and long-chain n-3 polyunsaturated fatty acid intake with the occurrence of depressive episodes in middle-aged French men and women. Prostaglandins, Leukotrienes Essential Fatty Acids. 2008;78:171-182.

166. Malhotra AK, Goldman D, Mazzanti C, Clifton A, Breier A, Pickar D. A functional serotonin transporter (5-HTT) polymorphism is associated with psychosis in neuroleptic-free schizophrenics. J Psychosom Res. 1998;33:328-332.

167. Willett WC, Sacks F, Trichopoulou A, et al. Mediterranean diet pyramid: a cultural model for healthy eating. Am J Clin Nutr. 1995;61:1402S-1406S.

168. Tortsa A, Bes-Rastrollo M, Sanchez-Villegas A, et al. Mediterranean diet inversely associated with the incidence of metabolic syndrome. Diabetes Care. 2007;30:2957-2959.

169. Vogelzangs N, Delgado-Rodriguez M, Alonso A, et al. Association of the Mediterranean dietary pattern with the incidence of depression: the Seguimiento Universidad de Navarra (SUN) Cohort. Arch Gen Psychiatry. 2009;66:1090-1098.

170. Dai J, Miller AH, Bremer JD, et al. Adherence to the Mediterranean diet is inversely associated with circulating interleukin-6 among middle-aged men: a twin study. Circulation. 2008;117:169-175.

171. Marriott BP, Cole N, Lee E. National estimates of dietary fructose intake increased from 1977 to 2004 in the United States. J Nutr. 2009;139:12385-12395.

172. Johnson RJ, Segal MS, Sautin Y, et al. Potential role of sugar (fructose) in the epidemic of hypertension, obesity and the metabolic syndrome, diabetes, kidney disease, and cardiovascular disease. Am J Clin Nutr. 2007;86:899-906.

173. Stanhope KL, Havel PJ. Endocrine and metabolic effects of consuming beverages sweetened with fructose, glucose, sucrose, or high-fructose corn syrup. Am J Clin Nutr. 2008;88:1733S-1737S.

174. Schaefer EJ, Gleason JA, Dansinger ML. Dietary fructose and glucose differentially affect lipid and glucose homeostasis. J Nutr. 2009;139:1257S-1262S.

175. Stanhope KL, Havel PJ. Fructose consumption: considerations for future research on its effects on adipose distribution, lipid metabolism, and insulin sensitivity in humans. J Nutr. 2009;139:12385-12415.

176. Dolan LC, Potter SM, Burdock GA. Evidence-based review on the effect of normal dietary consumption of fructose on development of hyperlipidemia and obesity in healthy, normal weight individuals. Crit Rev Food Sci Nutr. 2010;50:53-84.
Adiposity and depression - Shelton and Miller

Discourses in Clinical Neuroscience - Vol 13 - No. 1 - 2011

177. Tappy L, Le KA. Metabolic effects of fructose and the worldwide increase in obesity. Physiol Rev. 2010;90:23-46.

178. White JS. Misconceptions about high-fructose corn syrup: is it uniquely responsible for obesity, reactive dicarbonyl compounds, and advanced glycation endproducts? J Nutr. 2009;139:1219S-1225S.

179. Bocarsly ME, Powell ES, Avena NM, Hoebel BG. High-fructose corn syrup causes characteristics of obesity in rats: Increased body weight, body fat and triglyceride levels. Pharmacol Biochem Behav. 2010;97:101-106.

180. Tappy L, Le KA. Metabolic effects of fructose and the worldwide increase in obesity. Physiol Rev. 2010;90:23-46.

181. Figlewicz DP, Ioannou G, Bennett JJ, Kittleson S, Savard C, Roth CL. Effect of moderate intake of sweeteners on metabolic health in the rat. Physiol Behav. 2009;8:613-618.

182. Angelloupolou TJ, Lowndes J, Zukley L, et al. The effect of high-fructose corn syrup consumption on triglycerides and uric acid. J Nutr. 2009;139:1242S-1245S.

183. Schaerfe EJ, Gleson JA, Dansinger ML. Dietary fructose and glucose differentially affect lipid and glucose homeostasis. J Nutr. 2009;139:1257S-1262S.

184. Basciano H, Federico L, Adelli K. Fructose, insulin resistance, and metabolic dyslipidia. Nutr Metab Cardiovasc Dis. 2005;2:5-24.

185. Moran TH. Fructose and satiety. J Nutr. 2009;139:1253S-1256S.

186. Stanhope KL, Havel PJ. Fructose consumption: potential mechanisms for its effects to increase visceral adiposity and induce dyslipidemia and insulin resistance. Curr Opin Lipidol. 2008;19:16-24.

187. Peet M. Eicosapentaenoic acid in the treatment of schizophrenia and depression: rationale and preliminary double-blind clinical trial results. Prostaglandins Leukot Essent Fatty Acids. 2003;69:477-485.

188. Frangou S, Lewis M, McPone P. Efficacy of ethyl-eicosapentaenoic acid in bipolar depression: randomised double-blind placebo-controlled study. Br J Psychiatry. 2006;188:46-50.

189. Mischoulion D, Papakostas GI, Dording CM, et al. A double-blind, randomized controlled trial of ethyl-eicosapentaenate for major depressive disorder. J Clin Psychiatry. 2010;71:1636-1644.

190. Nembt B, Stahl Z, Belmaker RH. Addition of omega-3 fatty acid to maintenance medication treatment for recurrent unipolar depressive disorder. Am J Psychiatry. 2002;159:477-479.

191. Su KP. Mind-body interface: the role of n-3 fatty acids in psychiatry and depression - Shelton and Miller Dialogues in Clinical Neuroscience - Vol 13 - No. 1 - 2011. 2009;39:301-311.

192. Hasler G, Pine DS, Gamma A, et al. The associations between psychopharmacology and being overweight: a 20-year prospective study. Psychol Med. 2004;34:1047-1057.

193. Pulkkki-Raback L, Elvaniovi M, Kivimaki M, et al. Depressive symptoms and the metabolic syndrome in childhood and adulthood: a prospective cohort study. Health Psychol. 2009;28:108-115.

194. Reeves GM, Postolache TT, Snitker S. Childhood obesity and depression: connection between these growing problems in growing children. Int J Child Health Dev. 2008;1:103-114.

195. Okifuji A, Donaldson GW, Barck L, Fine PG. Relationship between fibromyalgia and obesity in pain, function, mood, and sleep. J Pain. 2010;11:1329-1337.

196. Arranz LI, Canela MA, Rafecas M. Fibromyalgia and nutrition, what do we know? Rheumatol Int. 2010;30:1417-1427.

197. Wright LJ, Schur E, Nooan C, Ahumada S, Buchwald D, Afari N. Chronic pain, overweight, and obesity: findings from a community-based twin registry. J Pain. 2010;11:628-635.

198. Mark PJ, Vasseljen O, Nilsen T. Association between physical exercise, body mass index, and risk of fibromyalgia: longitudinal data from the Norwegian Nord-Trondelag Health Study. Arthritis Care Res (Hoboken). 2010;62:611-617.

199. Oliver JE, Silman AJ. What epidemiology has told us about risk factors and aetiopathogenesis in rheumatic diseases. Arthritis Rheum. 2005;11:223-233.

200. Okifuji A, Bradshaw DH, Olson C. Evaluating obesity in fibromyalgia: neuroendocrine biomarkers, symptoms, and functions. Clin Rheumatol. 2009;28:475-478.

201. Trock D. Tired, achy, and overweight, the inflammatory nature of obesity. J Clin Rheumatol. 2009;15:50-54.

202. Neumann L, Lerner E, Glazer Y, Bolotin A, Shefer A, Bukula D. A cross-sectional study of the relationship between body mass index and clinical characteristics, tenderness measures, quality of life, and physical functioning in fibromyalgia patients. Clin Rheumatol. 2008;27:1543-1547.

203. Zhang Z, Cherryholmes G, Mao A, et al. High plasma levels of MCP-1 and eotaxin provide evidence for an immunological basis of fibromyalgia. Exp Biol Med (Maywood). 2008;233:1171-1180.

204. Bennett RM, Jones J, Turk DC, Russell JJ, Mataallana L. An internet survey of 2,596 people with fibromyalgia. BMC Musculoskelet Disord. 2007;8:2-7.

205. Jones KD, Adams D, Winters-Stone K, Burckhardt CS. A comprehensive review of 46 exercise treatment studies in fibromyalgia (1988-2005). Health Qual Life Outcomes. 2006;4:67.

206. Katz P, Gregorich S, Yazdany J, et al. Obesity and its measurement in a community-based sample of women with systemic lupus erythematosus. Arthritis Care Res (Hoboken). 2010.

207. Hahn BH, Lourencco EV, McMahon M, et al. Pro-inflammatory high-density lipoproteins and atherosclerosis are induced in lupus-prone mice by a high-fat diet and leptin. Lupus. 2010;19:913-917.

208. Paras ML, Murad MH, Chen LP, et al. Sexual abuse and lifetime diagnosis of somatic disorders: a systematic review and meta-analysis. JAMA. 2009;302:550-561.

209. Papakostas GI, Petersen T, Lisofsky DV, et al. Obesity among outpatients with major depressive disorder. Int J Neuropsychopharmacol. 2005;8:59-63.

210. Osikoilait C, Wilcox CS, Tung ML, Grosz DE. Body mass index and response to antidepressants in depressed research subjects. J Clin Psychiatry. 2009;70:1609-1610.

211. Markowitz S, Friedman FA, Arent SM. Understanding the relation between obesity and depression: causal mechanisms and implications for treatment. Clin Psychol Sci Pract. 2008;15:1-20.

212. Cicero AF, Deroga S, Bove M, Di G, V, Gaddi AV, Borghi C. Effect of a sequential training programme on inflammatory, prothrombotic and vascular remodelling biomarkers in hypertensive overweight patients with or without metabolic syndrome. Eur J Cardiovasc Prev Rehabil. 2009;16:698-704.

213. Forstye LK, Wallace JM, Livingstone MB. Obesity and inflammation: the effects of weight loss. 2008;21:117-133.

214. Kelishadi R, Hashemi M, Mohammadiard N, Asgary S, Khavarian N. Association of changes in oxidative and proinflammatory states with major depressive disorder. Arthritis Care Res (Hoboken). 2010.

215. Paras ML, Murad MH, Chen LP, et al. Sexual abuse and lifetime diagnosis of somatic disorders: a systematic review and meta-analysis. JAMA. 2009;302:550-561.

216. Papakostas GI, Petersen T, Lisofsky DV, et al. Obesity among outpatients with major depressive disorder. Int J Neuropsychopharmacol. 2005;8:59-63.

217. Osikoilait C, Wilcox CS, Tung ML, Grosz DE. Body mass index and response to antidepressants in depressed research subjects. J Clin Psychiatry. 2009;70:1609-1610.