Capsaicin may have important potential for promoting vascular and metabolic health

Mark F McCarty,1 James J DiNicolantonio,2 James H O’Keefe2

ABSTRACT
Capsaicin, the phytochemical responsible for the spiciness of peppers, has the potential to modulate metabolism via activation of transient receptor potential vanilloid 1 (TRPV1) receptors, which are found not only on nociceptive sensory neurons, but also in a range of other tissues. TRPV1 activation induces calcium influx, and in certain tissues this is associated with increased activation or expression of key proteins such as endothelial nitric oxide synthase (eNOS), uncoupling protein 2 (UCP2), KLF2, PPARdelta, PPARgamma, and LXRα. The calcium influx triggered by TRPV1 activation in endothelial cells mimics the impact of shear stress in this regard, activating and increasing the expression of eNOS—but also increasing expression of cox-2, thrombomodulin, and nrf2-responsive antioxidant enzymes, while decreasing expression of proinflammatory proteins. Hence, dietary capsaicin has favourably impacted endothelium-dependent vasodilation in rodents. TRPV1-mediated induction of LXRα in foam cells promotes cholesterol export, antagonising plaque formation. Capsaicin-mediated activation of TRPV1-expressing neurons in the gastrointestinal tract promotes sympathetically mediated stimulation of brown fat, raising metabolic rate. The increased expression of UCP2 induced by TRPV1 activation exerts a protective antioxidant effect on the liver in non-alcoholic fatty liver disease, and on vascular endothelium in the context of hyperglycaemia. In rodent studies, capsaicin-rich diets have shown favourable effects on atherosclerosis, metabolic syndrome, diabetes, obesity, non-alcoholic fatty liver, cardiac hypertrophy, hypertension and stroke risk. Clinically, ingestion of capsaicin—or its less stable non-pungent analogue capsiate—has been shown to boost metabolic rate modestly. Topical application of capsaicin via patch was found to increase exercise time to ischaemic threshold in patients with angina. Further clinical studies with capsaicin administered in food, capsules, or via patch, are needed to establish protocols that are tolerable for most patients, and to evaluate the potential of capsaicin for promoting vascular and metabolic health.

CAPSAICIN STIMULATES THE TRPV1 RECEPTOR
Transient receptor potential vanilloid 1 (TRPV1) is a membrane receptor that, when activated, acts as a non-specific cation channel, allowing influx of calcium. Endogenous activators of TRPV1 include heat, low pH, and certain lipid metabolites; the best known exogenous activator is the phytochemical capsaicin, responsible for the spiciness in peppers.1–3 Inasmuch as nanomolar concentrations of capsaicin can activate this receptor (EC50=99 nM4), and capsaicin is efficiently absorbed,5 a sufficiently high oral intake of capsaicin can induce systemic activation of TRPV1.

Few studies have evaluated the clinical pharmacokinetics of orally administered capsaicin.6 After acute ingestion of 5 g of a capsaicin-rich hot pepper extract, a peak serum capsaicin level of 8.2 nM was observed after 45 min; an hour later, capsaicin was no longer detectible, presumably owing to rapid hepatic metabolism.7 In mice given a bolus dose of 10 mg/kg capsaicin—far higher than humans could be expected to use—the peak serum concentration was about 3 µM; after 8 h, capsaicin was undetectable in serum. It is therefore reasonable to expect that clinically tolerable intakes of capsaicin will achieve serum concentrations in the nanomolar range. Although capsaicin can inhibit certain voltage-sensitive calcium channels with an EC50 of 5 µM or higher,8 9 it does not appear likely that this effect would be germane with feasible oral intakes of capsaicin in humans.

TRPV1 is expressed by many nociceptive sensory neurons, and its activation triggers pain sensations. However, the massive neuronal calcium influx triggered by topical exposure to sufficient concentrations of capsaicin is potentially cytotoxic, and triggers a reflex down-regulation of TRPV1 activity.10 Hence, these neurons become less responsive to endogenous agonists for TRPV1, resulting in analgesia.11 12 Capsaicin skin patches are currently employed clinically for local pain control.13

TRPV1 is also expressed by vascular endothelial cells, hepatocytes, adipocytes, smooth
muscle cells, fibroblasts, various epithelia, T cells, mast cells, and by neurons and astrocytes in the brain and spinal column. Hence, TRPV1 has the potential to modulate the function of these cells by boosting their intracellular-free calcium levels (Ca$_i$). At present, there does not appear to be any evidence that the desensitisation phenomenon evoked by capsaicin in sensory neurons is pertinent to these other tissues; no down-regulation of TRPV1 expression or function was noted in the vasculature of newborn rats that had been injected with potent doses of capsaicin for 5 days.

CAPSAICIN CAN INCREASE EXPRESSION AND ACTIVATION OF ENOS

The impact of TRPV1 activation on vascular endothelium is of particular interest, since an increase in Ca$_i$ is a key mediator of the protective impact of pulsatile shear stress—and of aerobic exercise—on endothelial function. This increase in Ca$_i$ acts rapidly to stimulate endothelial nitric oxide synthase (eNOS) activity via binding of the Ca$^{2+}$/calmodulin complex; in addition, Ca$_i$-mediated activation of AMPK and Sirt1 stimulates eNOS activity by modifying its phosphorylation and acetylation status. In the longer term, expression of eNOS increases as well. Increased Ca$_i$ acts to boost the expression and activity of the endothelium-specific transcription factor KLF2 via a complex chain of events involving activation of Ca$_{2+}$/calmodulin-dependent kinase kinase-$\beta$ and downstream phosphorylations of AMPK, ERK2, HDAC5, and the transcription factor MEF2. KLF2, in turn, promotes transcription of the eNOS, thrombomodulin, and Nrf2-responsive antioxidant enzymes, and works indirectly to suppress transcription of various proinflammatory proteins.

As might be expected, treatment of endothelial cells with capsaicin leads to increased expression and activation of eNOS. Consistent with this, in wild-type, but not TRPV1-knockout mice, dietary capsaicin enhances endothelium-dependent vasodilation. In spontaneously hypertensive stroke-prone rats, dietary capsaicin increases activation and expression of eNOS in the cerebrovasculature, an effect associated with a reduction of arteriolar hypertrophy, a delay in stroke occurrence, and an increase in mean survival time. In atherosclerosis-prone ApoE knockout mice, dietary capsaicin slows atherogenesis, an effect which may reflect improved endothelial function, but also a favourable impact of TRPV1 activation on foam cells, increasing the expression of membrane transporters that mediate cholesterol efflux; this latter effect is contingent on increased expression of the transcription factor LXR$\alpha$. The potential clinical relevance of these findings is demonstrated by a controlled crossover study in which patients with stable coronary disease and angina were treated with capsaicin skin patches (typically employed for control of lower back pain) or placebo patches. During exercise testing, average time until ischaemic threshold (1 mm ST segment depression) was significantly higher during capsaicin administration (424 s vs 372 s, p=0.027). Notably, serum NO levels (assessed by measuring its stable metabolites nitrate and nitrite) were found to be significantly higher when the patients were using the capsaicin patches, suggesting that increased NO production within the coronary tree may have been responsible for the improved exercise tolerance associated with capsaicin.

Capsaicin feeding has shown an antihypertensive effect in rats genetically prone to this disorder, and this compound also blunts the nocturnal rise in blood pressure or development of hypertension in mice fed a high salt diet. Conceivably, improved NO function may underlie these effects. Capsaicin dilates the coronary arteries of pigs ex vivo, an effect that is half-maximal at 116 nM; this effect is blocked by endothelial denudation and inhibitors of eNOS, and is less notable with coronary arteries from pigs experiencing metabolic syndrome, which disrupts eNOS function via oxidative stress. Release of CGRP from perivascular sensory neurons may also contribute to the vasodilatory impact of capsaicin. Paradoxically, the direct impact of capsaicin on vascular smooth muscle is to provoke constriction, owing to increased calcium influx. Hence, the net impact of capsaicin on vascular tone and blood pressure may reflect complex interactions and countervailing effects. Several case histories of acute hypertensive crisis provoked by very high intakes of chilli peppers have appeared; down-regulated function of CGRP-producing neurons owing to acute high capsaicin exposure has been suggested as an explanation for this effect. On the other hand, a more moderate capsaicin exposure associated with the use of capsaicin patches—sufficient to alleviate angina pain—did not alter plasma levels of CGRP, but plasma levels of NO metabolites increased. Whether and how moderate, clinically tolerable dosing with capsaicin would influence human hypertension has not yet been assessed.

The antihypertensive effect of dietary capsaicin in salt-fed rats may reflect, in part, an inhibitory effect on renal sodium retention. In the kidney, cortical collecting duct epithelium expresses TRPV1, and its activation decreases the function and expression of epithelial sodium channels in these cells, resulting in increased urinary sodium loss.

CAPSAICIN BOOSTS UCP2 EXPRESSION IN CERTAIN TISSUES

TRPV1 activation has also been shown to increase expression of uncoupling protein 2 (UCP2) in endothelial cells, hepatocytes and cardiac tissue. In the heart, this effect may be downstream from increased expression of PPARDelta, a factor which opposes cardiac hypertrophy and fibrosis. Hence, dietary capsaicin was found to oppose the cardiac hypertrophy induced by a high salt diet in mice—an effect not seen in TRPV1
knockout mice. With respect to UCP2, this function as a mitochondrial uncoupling protein when mitochondrial substrate oxidation is high and superoxide generation is elevated; by diminishing the proton gradient across the mitochondrial inner membrane, UCP2 relieves the resistance to electron flow down the respiratory chain and hence decreases the rate at which electrons are shunted to superoxide generation at complexes I and III. 

UCP2 can be of particular value when cells that are constitutively permeable to glucose—such as vascular endothelium—are subjected to hyperglycaemia. Under these circumstances, elevated glucose oxidation in the Krebs cycle tends to boost mitochondrial superoxide generation, an effect opposed by UCP2. In diabetic mice, capsaicin administration was shown to alleviate vascular oxidative stress and improve endothelium—mice, capsaicin administration was shown to alleviate vas-

In men with diabetes, a polymorphism in the UCP2 promoter (-866G>A), linked to increased expression of UCP2 in some studies, was found to be associated with significantly lower risk for developing coronary disease. 

The beneficial effects of capsaicin on metabolic syndrome in mice may be mediated in part by increased secretion of glucagon-like peptide-1 (GLP-1). Indeed, gastric administration of capsaicin has been shown to evoke increased secretion of GLP-1 by the gastrointestinal (GI) tract, and to raise plasma levels of this factor. This effect is absent in TRPV1 knockout mice. Increased calcium influx into intestinal L cells may mediate this impact on GLP-1 secretion.

How activation of the TRPV1 receptor manages to increase the expression of various regulatory factors—UCP2, PPARalpha and PPARdelta, LXRe—remains obscure; calcium influx per se seems unlikely to mediate all these effects. Perhaps TRPV1 has a distinctive micro-environment reflecting binding affinities to other proteins, such that influxing calcium tends to preferentially activate certain calcium-binding proteins in this micro-environment. It is also conceivable that some of TRPV1’s signalling effects are independent of calcium influx.

**THERMOGENIC AND APPETITE CONTROL EFFECTS OF CAPSAICIN AND CAPSIATE**

Another intriguing TRPV1-dependent effect of capsaicin ingestion is activation of brown adipose tissue. Activation of TRPV1-expressing neurons in the digestive tract sends a signal to the brain via the vagal nerve; this in turn evokes an activation of sympathetic neurons that is selective for brown fat—that is, the heart rate is not impacted. Many clinical trials have evaluated the impact of capsin ingestion on metabolic rate, respiratory quotient and appetite; these conclude that capsaicin can modestly enhance energy expenditure, while boosting fat oxidation (lower RQ) and diminishing appetite—effects conducive to weight control.

Similar effects are seen with a non-spicy analogue of capsaicin, capsiate, which owing to lower stability does not induce pain in the oral cavity and appears to have limited systemic availability. Capsiate is found in certain sweet peppers; it is very similar in structure to capsaicin, and can activate TRPV1, with an affinity about one-third that of capsaicin (EC50=290 nM). Whereas capsaicin contains an amide linkage that is relatively weak and subject to rapid catabolism, capsiate has an ester linkage that is more stable. Capsiate may have additional advantages over capsaicin, in particular the ability to activate TRPV1 in the oral mucosa, in the gut, and in adipose tissue.
stable, capsiate contains an ester that is readily cleaved; when administered orally, intact capsiate fails to reach oral TPRV1-expressing neurons, but does manage to stimulate such neurons lower in the GI tract. No intact capsiate appears in the portal blood after oral administration, but its hydrolysis products are detectible, implying that capsiate is hydrolysed during the process of absorption.65 Hence, the effects of capsiate attributable to TPRV1 agonism appear to be mediated by stimulation of GI sensory neurons.

Both capsaicin and capsiate may have modest utility as adjuvants to weight control programmes. Supplementation with capsiate (9 mg daily) for 12 weeks in a double-blind study was shown to decrease abdominal fat mass relative to placebo, albeit to a modest extent.66 (Over the 12 weeks, the capsiate group, on average, lost 0.4 kg of weight and 1 cm of waist girth beyond that achieved with placebo—not an effect of much practical importance unless it persists and increases over time.) Not surprisingly, the effects of capsaicin or capsiate on thermogenesis are most notable in humans bearing detectible amounts of brown fat.6566 However, there is some evidence that prolonged ingestion of these agents may lead to recruitment of brown fat in humans.67 These effects on thermogenesis are modest in magnitude; there do not appear to be any reports of clinically significant hyperthermia with ingestion of capsaicin or capsiate.

Some studies have also evaluated the impact of oral capsaicin or capsiate on appetite and subsequent food consumption in various contexts. The findings of these studies have been inconsistent, though an overview of these studies by Ludy et al63 concludes that, on balance, consumption of these agents tends to decrease orexigenic sensations. In positive studies, capsaicin-treated subjects reported less desire to consume fatty foods, sweet foods, salty foods and food overall, and achieved significant weight loss with less food intake than in the placebo group.40 In a 12-week study, capsiate-treated volunteers took 45 mg capsaicin three times daily with meals, when Lejeune et al77 had study volunteers take 45 mg capsaicin three times daily with meals, 24% of them experienced significant stomach discomfort and were allowed to cut this dose in half; however, this dose regimen seems likely to be a higher dose than would be required for metabolic benefits.

IMPACT ON GASTRIC PATHOLOGY

Ironically, many laypeople are under the impression that spicy foods can cause ulcers; to the contrary, there is evidence that capsaicin tends to prevent and accelerate healing of gastric ulcers.68–70 This phenomenon reflects capsaicin’s ability to inhibit gastric acid secretion, boost secretion of alkali and mucous, and stimulate gastric blood flow. A clinical study found that the gastric tissue damage and microbleeding induced acutely by indomethacin or ethanol ingestion was blunted if capsaicin was administered concurrently.69 These findings have prompted the suggestion that capsaicin could be used as a protective adjuvant to non-steroidal anti-inflammatory drug therapy.6970 Limited epidemiology suggests that gastric ulcers may be less common in ethnic groups that prefer spicy foods.68

With respect to risk of gastric cancer, the epidemiology on spicy foods is rather perplexing. A recent meta-analysis of pertinent studies in Korea and Mexico, where heavy consumption of spicy foods is common, concludes that moderate daily intakes of capsaicin (less than 30 mg daily) are associated with a significant decrease in gastric cancer risk (OR=0.55, p=0.003) relative to non-consumption—perhaps reflecting the gastroprotective effects of capsaicin—whereas, heavy daily consumption is associated with a notable increase in risk (OR=1.94, p=0.0004).71 Bley et al72 suggest that the increase in risk associated with heavy consumption of spicy traditional foods might reflect mutagens present in these foods, rather than an effect of capsaicin per se. Aflatoxins, pesticides and nitrosamines or their precursors have been detected in chillies sold for human consumption.72 The traditional Korean dish kimchi, a salty pickled cabbage usually fermented with red pepper and linked to increased risk of gastric cancer, is typically high in nitrate and contains N-nitroso compounds with mutagenic potential; the high salt content of this food may act as gastric co-carcinogen.73–76 Studies with high-purity capsaicin indicate that it is not genotoxic; in animal studies, capsaicin lacks carcinogenicity, and opposes the carcinogenicity of certain mutagens.72 Further clarification of this situation will be desirable if in the future people are encouraged to consume more capsaicin for potential health benefits.

DOSAGE CONSIDERATIONS

In rodents, large metabolic effects have been reported with dietary capsaicin intakes in the range of 0.01–0.02% of diet. If a human were to eat (say) 400 g dry weight of food daily, 0.01% of diet would correspond to 40 mg capsaicin. Oral administration of capsaicin represents a clinical challenge—many people, especially those not acclimated to a spicy diet, do not enjoy the oral pain associated with capsaicin-laced foods, and capsaicin capsules may cause GI distress in some persons; this latter effect is mitigated somewhat by ingesting capsaicin capsules with meals. When Lejeune et al77 had study volunteers take 45 mg capsaicin three times daily with meals, 24% of them experienced significant stomach discomfort and were allowed to cut this dose in half; however, this dose regimen seems likely to be a higher dose than would be required for metabolic benefits.

Hot peppers typically contain capsaicin in conjunction with lesser amounts of its analogues, dihydrocapsaicin and nordihydrocapsaicin; the latter is a very minor component, but dihydrocapsaicin may constitute as much as 40% of total capsaicinoids. The relative proportion of...
capsaicin and dihydrocapsaicin in a food is of little practical import, as the abilities of these compounds to activate TRPV1 are roughly equivalent. Commercial capsules of cayenne pepper are available that provide 40 000–100 000 Scoville heat units per capsule. The Scoville scale quantifies the spicy heat (or pungency) of foods which contain capsaicinoids; a gram of capsaicin corresponds to 16 million Scoville heat units; a gram of dihydrocapsaicin to 15 million units; and a gram of nor-dihydrocapsaicin to 9.1 million units. Therefore, a capsule containing 100 000 Scoville heat units can be expected to contain about 6.6 mg of capsaicinoids. Consuming three of these daily with meals will provide about 20 mg, and those who enjoy spicy foods could supplement this with peppers, pepper sauces, or cayenne powder added to foods. Perhaps this would be an appropriate ‘baseline’ regimen to study clinically. Topical administration of capsaicin in patches may represent a reasonable alternative in people unable to tolerate it orally—albeit this will be a more expensive option, and local pain is commonly experienced for an hour or more after patch application.78

EXPLORING THE HEALTH POTENTIAL OF CAPSAICIN

This brief overview should make it clear that dietary capsaicin—and, likely to a more limited degree, non-pungent capsiate—has intriguing potential for health promotion. Rodent studies suggest that capsaicin may merit clinical evaluation with respect to endothelial function, progression of atherosclerosis (most notably in diabetics), angina, non-alcoholic fatty liver disease, cardiac hypertrophy, metabolic syndrome, hypertension, obesity and gastric ulceration. (See table 1 for a summary of these potential benefits and the mechanisms that may underlie them.) In addition to the many studies assessing capsaicin’s impact on metabolic rate and adiposity, the trial of topical capsaicin in patients with angina, and the studies documenting capsaicin’s gastroprotective effects, represent initial efforts in this regard. A study examining endothelium-dependent vasodilation in diabetics might be particularly useful, as a systematically adequate dose of capsaicin could be expected to have a notably favourable impact on this parameter. Assessment of the dose-dependency of this effect could provide useful insight into capsaicin clinical dosage schedules which could provide systemic metabolic benefits. Both oral and topical application of capsaicin could be tested in this regard. The rodent literature is sufficiently intriguing that serious efforts to evaluate the feasibility of capsaicin administration as a clinical or lifestyle strategy appear to be warranted. However, owing to the fact that TRPV1 receptors are expressed on a wide range of tissues, the possibility that high-dose capsaicin might exert unanticipated or unwanted physiological effects should be borne in mind.

Contributors MFM conceived of this essay, and wrote the initial draft. JJD and JHO’K suggested revisions and wrote portions of the revised draft.

Competing interests JJD works for a company that sells capsaicin products, but he has no direct role in marketing or selling them. JHO’K and MFM have ownership interests in companies that make nutritional supplements, but these companies do not sell capsaicin products.

Provenance and peer review Not commissioned; externally peer reviewed.

Open Access This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

REFERENCES

Table 1: Health benefits of capsaicin administration suggested by preclinical and clinical research

<table>
<thead>
<tr>
<th>Condition benefited</th>
<th>Likely mechanisms of action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atherosclerosis26 29 30</td>
<td>Improved endothelial function, including eNOS activation/induction; induction of LXRxAlpha in foam cells, promoting cholesterol export</td>
</tr>
<tr>
<td>Diabetic vasculopathy39</td>
<td>Induction of UCP2 and eNOS in endothelium</td>
</tr>
<tr>
<td>Stroke58</td>
<td>Improved endothelial function, including eNOS activation/induction</td>
</tr>
<tr>
<td>Angina31</td>
<td>Improved endothelium-dependent vasodilation of coronary arteries</td>
</tr>
<tr>
<td>Hypertension27 32 33</td>
<td>Activation/induction of eNOS; decreased renal sodium retention</td>
</tr>
<tr>
<td>Metabolic syndrome54–58</td>
<td>Decreased adipose inflammation—reflecting PPARgamma induction</td>
</tr>
<tr>
<td>Cardiac hypertrophy41</td>
<td>Induction of PPARdelta</td>
</tr>
<tr>
<td>Fatty liver59 52</td>
<td>Induction of UCP2 in hepatocytes; decreased adipose inflammation</td>
</tr>
<tr>
<td>Obesity56 57 61 62 66</td>
<td>Increased GLP-1 secretion</td>
</tr>
<tr>
<td>Gastric ulceration68–70</td>
<td>Decreased acid secretion; increased alkalii; increased gastric blood flow</td>
</tr>
</tbody>
</table>

eNOS, endothelial nitric oxide synthase; GLP-1, glucagon-like peptide-1; UCP2, uncoupling protein 2


