Cysteine and hydrogen sulfide in the regulation of metabolism

Citation for published version:

Digital Object Identifier (DOI):
10.1002/path.4659

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
The Journal of Pathology

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Cysteine and hydrogen sulphide in the regulation of metabolism: insights from genetics and pharmacology

Roderick N Carter and Nicholas M Morton*

Molecular Metabolism Group, University/BHF Centre for Cardiovascular Sciences, Queens Medical Research Institute, University of Edinburgh, UK

*Correspondence to: NM Morton, Molecular Metabolism Group, University/BHF Centre for Cardiovascular Sciences, Queens Medical Research Institute, University of Edinburgh, 47 Little France Crescent, Edinburgh EH16 4TJ, UK. E-mail: Nik.Morton@ed.ac.uk

Abstract

Obesity and diabetes represent a significant and escalating worldwide health burden. These conditions are characterized by abnormal nutrient homeostasis. One such perturbation is altered metabolism of the sulphur-containing amino acid cysteine. Obesity is associated with elevated plasma cysteine, whereas diabetes is associated with reduced cysteine levels. One mechanism by which cysteine may act is through its enzymatic breakdown to produce hydrogen sulphide (H₂S), a gasotransmitter that regulates glucose and lipid homeostasis. Here we review evidence from both pharmacological studies and transgenic models suggesting that cysteine and hydrogen sulphide play a role in the metabolic dysregulation underpinning obesity and diabetes. We then outline the growing evidence that regulation of hydrogen sulphide levels through its catabolism can impact metabolic health. By integrating hydrogen sulphide production and breakdown pathways, we re-assess current hypothetical models of cysteine and hydrogen sulphide metabolism, offering new insight into their roles in the pathogenesis of obesity and diabetes.

© 2015 The Authors. The Journal of Pathology published by John Wiley & Sons Ltd on behalf of Pathological Society of Great Britain and Ireland.

Keywords: cysteine; sulphide; obesity; diabetes; genetic models; metabolism; insulin resistance; adipose, liver

Obesity and diabetes

Obesity and type 2 diabetes are a major global health burdens that have dramatically increased in prevalence in the last few decades. According to the World Health Organization, worldwide obesity has more than doubled since 1980 [1]. As of 2014 it was estimated that 600 million worldwide could be classed obese, almost 10% of the global population [1]. Type 2 diabetes has become similarly common, with a global prevalence of 9% in those over 18 years of age [2]. The two conditions are strongly associated [3], largely due to obesity being a strong risk factor for the development of insulin resistance – a key underlying mechanism in type 2 diabetes [4]. This relationship is not absolute, and many individuals remain metabolically healthy in spite of having high fat mass, while others may suffer diabetes despite normal fat mass [5]. A cardinal feature of both obesity and type 2 diabetes is atypical carbohydrate and lipid metabolism. Nutritional excess eventually overloads the expanding adipose tissue, leading to ectopic accumulation of lipid in key metabolic tissues such as the liver, muscle and insulin-producing pancreatic islets [6,7]. This produces a lipotoxic stress [8] that leads to inflammation and insulin-resistance in the adipose tissue, liver and muscle, with a consequent dysregulation of glucose homeostasis that precipitates diabetes. Specifically, adipose tissue free fatty acid release is increased (a process normally suppressed by insulin after feeding) and glucose metabolism is skewed towards lipogenesis [9]. In the liver, insulin-resistance increases hepatic production of glucose by gluconeogenesis and glycogen breakdown (which is normally suppressed by post-prandial insulin action), contributing to hyperglycaemia. Of note, lipid synthesis and transport pathways remain sensitive to insulin in the liver, driving lipogenesis and altering lipoprotein profiles that ultimately contribute to atherosclerosis [10]. Insulin resistance in muscle leads to impaired glucose disposal and oxidation, thus exacerbating hyperglycaemia. Increased demand for insulin engendered by hyperglycaemia eventually leads to pancreatic β cell exhaustion, insulin deficiency and frank diabetes.

Links between cysteine and obesity and diabetes

An emerging factor associated with obesity is altered levels of the sulphur-containing amino acid cysteine. Plasma total cysteine correlates positively with obesity, as defined by body mass index [11,12], and in particular with fat mass [13]. This relationship appears to be specific to cysteine, rather than amino acids in general, as no...
other amino acid shares the same strength of association with obesity [14]. Plasma levels of other amino acids that associate with obesity, such as cystathionine and glutamate, normalize following gastric bypass surgery [15]; by contrast, plasma cysteine levels remain high, supporting the hypothesis that elevated cysteine is not merely the consequence of increased fat mass but may mechanistically underlie fat gain. This concept is supported by evidence that in some rodent models, increasing dietary cysteine levels can result in increased adiposity [16]. The mechanism by which cysteine could cause increased tissue fat accumulation remains unresolved; one possibility is that cysteine directly regulates energy expenditure and/or appetite. Conflicting with its proposed role in fat mass gain, there is evidence that cysteine reduces appetite in humans and in rodent models [17]. Alternatively, it has been reported that dietary cysteine reduces metabolic rate in mice, consistent with increased energy storage [18]. Understanding any mechanism of action of cysteine with respect to wider metabolic control must also take into account that, contrary to its relationship with obesity, type 2 diabetes has, in some studies, been associated with lower plasma cysteine levels [19].

**Hydrogen sulphide as a metabolite of cysteine; clinical correlates with obesity/diabetes**

In addition to the potential effects of cysteine *per se*, it is possible that metabolites of cysteine could be crucial. In this regard, cysteine is a critical substrate for the intracellular generation of hydrogen sulphide (H$_2$S), an enzymatically produced physiologically active gasotransmitter. Hydrogen sulphide has emerged as an important factor in the modulation of insulin action in tissues such as liver, adipose tissue and islets of Langerhans. In contrast to plasma total cysteine, which has a positive correlation to obesity and adiposity [14], plasma hydrogen sulphide has been found to be negatively correlated with measures of adiposity, in particular waist circumference and waist:hip ratio [20]. In this study, hydrogen sulphide was lowest in subjects with obesity and type 2 diabetes, although regression analysis suggested that the adiposity was the main driver for predicting low plasma hydrogen sulphide. Lower plasma hydrogen sulphide levels were confirmed independently in another cohort of type 2 diabetes subjects [21]. Thus, cysteine and hydrogen sulphide have opposing relationships in obesity but share the same negative correlation with type 2 diabetes. It should be noted that, as accurate hydrogen sulphide measurements from plasma are technically challenging [22], these findings remain preliminary.

**Hydrogen sulphide: production, breakdown and mechanisms of action**

Hydrogen sulphide was first discovered to be an endogenously produced, physiologically active compound in rodent brain [23,24], and previously was regarded as a lethal respiratory toxin. A significant route of hydrogen sulphide production is the reverse transulphuration system, whereby homocysteine is converted to cysteine via cystathionine. The two steps of this metabolic conversion are carried out by the predominantly cytosolic enzymes cystathionine β-synthase (CBS) and cystathionine γ-lyase (CSE; also referred to as CTH) [25,26]. Both CBS and CSE are capable of a number of related but distinct enzymatic conversions, with hydrogen sulphide formed as a product of some of these reactions [27,28] (Figure 1). The relative contribution of these two enzymes to hydrogen sulphide production are dependent on tissue-specific enzyme levels, substrate and co-factor availability, and activators such as S-adenosyl methionine [27,28]. Mercaptopyruvate sulphur transferase (MPST; also referred to as MST) may also generate hydrogen sulphide through the cysteine transamination product 3-mercaptoppyruvate [29]. MPST acquires the sulphur atom from mercaptopyruvate to form a persulphide intermediate, which is released as hydrogen sulphide in the presence of thioredoxin or other reducing agents [30].

Significantly, hydrogen sulphide breakdown is often overlooked, but has the potential to regulate physiological levels and may be important in preventing toxic accumulation in tissues. Clearance of hydrogen sulphide can occur by at least two mechanisms; the most characterized is oxidation by mitochondria through a respiratory route (Figure 2). This pathway has established hydrogen sulphide as the first non-carbon-based respiratory fuel in mammalian systems [31]. The proposed canonical breakdown pathway involves sulphide quinone oxidoreductase (SQR; also referred to as SQRDL), persulphide dioxygenase (also referred to as Ethel1, or sulphite dioxygenase), and thiosulphate sulphur transferase (TST; also referred to as rhodanese), [32,33]. The main products of this oxidation appear to be sulphite and thiosulphate. Sulphite itself can be further oxidized to sulphate by mitochondria by the action of sulphite oxidase (SOX; also referred to as SOXO), in a mechanism that can contribute electrons to the electron transport chain via cytochrome c [34,35]. A less established route for hydrogen sulphide catabolism may involve reaction with iron-heme species present, for example, in red blood cells and the mitochondrial cytochrome c oxidase [36,37]. In *vitro*, the products of heme-mediated oxidation are polysulphides and thiosulphate, rather than sulphite [38]. It is noteworthy that sulphate, sulphite, thiosulphate and polysulphides all have significant bioactive properties, extending the potential physiological impact of cysteine metabolism and hydrogen sulphide production to the activities of these metabolites [39–42].

A number of distinct mechanisms of hydrogen sulphide action have been proposed for regulating cellular function. One mechanism of action is regulation of the anti-oxidant status of cells. The low micromolar concentrations of endogenous hydrogen sulphide, compared to the millimolar concentrations of more established thiol
Cysteine-sulfide in metabolism: genes and pharmacology

Transulfuration Enzymes: Production of Cysteine and Sulphide

Cysteine from Homocysteine

Homocysteine + Serine

H₂O

CBS

Cystathionine

H₂O

NH₄⁺

CSE

α-Ketoglutarate + Cysteine

+ Serine

CBS

Cysteine

2 x Cysteine

CSE

Homocysteine + Cysteine

CBS

Cysteine

(SH₂/SH⁻)

CSE

Sulphide

Lanthionine

Figure 1. Transulphuration – a source of cysteine and sulphide. CBS and CSE link cysteine generation from homocysteine, an intermediate of the methionine cycle, with sulphide generation. CBS and CSE can be regulated at substrate, protein and RNA levels, and thus dynamic changes in the flux of both cysteine and sulphide production is predicted to be possible in tissues.

Antioxidants such as glutathione and cysteine (free and protein-bound), suggest that hydrogen sulphide likely plays an indirect role in the antioxidant capacity of cells through its signalling functions [43]. In an early study of this property, primary cortical neurons were protected from glutamate-induced cell death by administration of 100 μM sodium hydrogen sulphide (NaHS), an inorganic source of hydrogen sulphide [44]. In that study hydrogen sulphide increased cysteine import and glutathione production. There is also emerging evidence that hydrogen sulphide interacts with nitric oxide signalling [45,46]. Hydrogen sulphide reacts with nitric oxide to form nitrosothiol, and in some circumstances may stimulate cGMP signalling as may nitric oxide. Endogenous production of nitrosothiol was also measured in that study, suggesting a physiological role for direct sulphide–nitric oxide reactions. Modification of proteins by sulphydration (or persulphidation) is a further mechanism by which hydrogen sulphide may alter protein function. This modification refers to the presence of an additional sulphur atom attached to a thiol, such as cysteine, essentially a persulphide group. Formation of persulphides through hydrogen sulphide requires either oxidation of the hydrogen sulphide prior to reaction with a thiol, or reaction of hydrogen sulphide with an already oxidized species, such as a disulphide group [47–49]. Sulphydration modification regulates the activity of ATP sensitive potassium channels, other membrane ion channels, kinases and transcription factors [48,49] to elicit diverse biological effects. A persulphide cysteine is a known intermediate of the mercaptopyruvate sulphurtransferase enzyme catalytic cycle [50], suggesting that persulphide formation may increase reactivity of other active site cysteines.

Hydrogen sulphide in the liver: enhancing glucose production and lipid oxidation?

Investigation into whether, and how, sulphide influences hepatic function is at an early stage. Existing data support the view that sulphide promotes glucose production, enhances lipid oxidation and inhibits insulin action [51,52].

In the study by Zhang et al [51], treatment of clonal HepG2 hepatocyte cells with 10–100 μM NaHS reduced glucose consumption and reduced glycogen levels in association with reduced gluconeogenesis activity. In contrast, phosphoenol pyruvate carboxykinase (PEPCK) activity and gluconeogenesis rates were increased by similar concentrations of NaHS in these cells. Notably, NaHS blocked stimulation of glucose consumption following insulin treatment, and this was associated with a reduction in phosphorylation (activation) of the insulin-signalling intermediate AKT. A feedback mechanism between hydrogen sulphide and insulin signalling was uncovered, with insulin exposure reducing CSE levels and hydrogen sulphide production. Reversal of this regulatory loop was observed when the HepG2 cells were rendered insulin-resistant through chronic culture in high glucose. In these cells, CSE protein was increased, as was hydrogen sulphide production.

In vivo, Wu et al [52] reported that daily injections of 50 μmoles/kg NaHS administration during a 16 week high-fat diet regime prevented diet-induced liver damage and increased markers of lipid oxidation. Specifically, hepatic expression of fatty acid synthase was reduced and expression of carnitine palmitoyltransferase-1 was increased. High-fat diet-induced oxidative damage, as assessed by measuring...
hepatic malondialdehyde, was reduced in the NaHS-injected mice, consistent with the findings of increased superoxide dismutase and glutathione peroxidase activity.

Taken together, hydrogen sulphide appears to oppose insulin action in the liver, and may mechanistically contribute to hepatic insulin resistance. Intriguingly, by enhancing glucose production and lipid oxidation, hydrogen sulphide mimics the actions of glucagon and/or adrenaline. This property fits with the increased production of hydrogen sulphide found in mice under dietary restriction [53] and clinical findings that hydrogen sulphide levels are low in obesity and type 2 diabetes [20,21]. This is consistent with the loss of potentially beneficial lipid oxidation effects and dysregulated hepatic lipid metabolism in these states. However, from the mechanistic studies above, low hydrogen sulphide levels would predict improved insulin sensitivity with respect to hepatic glucose homeostasis, which is clearly not the case in human type 2 diabetes. A full understanding of the potentially distinct regulation of glucose and lipid homeostasis by hydrogen sulphide signalling will be important to elucidate. A clearer understanding of the relationship between enzyme levels for hydrogen sulphide production, breakdown and the consequent steady-state levels in liver and other key metabolic tissues in diabetes and obesity is also of critical importance.

Hydrogen sulphide in adipose tissue: enhancing glucose disposal and increasing conditions for lipid storage?

Early studies with primary rat adipocytes suggested that hydrogen sulphide prepared as a saturated gas solution, and used effectively at doses in the range 10–1000 μM, could inhibit glucose uptake, thereby opposing insulin action [55]. This finding has not been consistently replicated [56,57], albeit some of the work in these later studies used NaHS, or sodium sulphide (Na2S) as a means of introducing hydrogen sulphide to their systems.

Studies of differentiated 3T3-L1 adipocytes have generally supported the pro-insulin and pro-fat storage effects of hydrogen sulphide. High-glucose culture...
conditions of differentiating 3T3-L1s, associated with insulin resistance, was found to reduce protein levels of molecular markers of the insulin-signalling pathway, including PI3KInase and phospho-AKT as well as PIP3 levels [56]. These effects were reversed by 10–100 μM Na₂S, 100–1000 μM cysteine or the addition of insulin to the medium. The same authors also showed that reducing the insulin receptor level using siRNA blocked these effects of Na₂S on signal transduction. In another study, high-glucose medium conditions were found to reduce CSE mRNA, coincident with reduced production of the insulin sensitizer adiponectin and increased production of the pro-inflammatory factor MCP-1 [57]. The addition of 10–50 μM NaHS or forced expression of CSE reversed these effects. Finally, treatment with the compound GYY4137 (50 μM) or NaHS (50 μM) throughout the differentiation of 3T3-L1 cells led to the increased formation of lipid droplets and inhibition of adrenergic receptor-stimulated lipolysis [58]. Consistent with these findings, administration of GYY4137 at 200 μmole/kg/day to mice reduced lipolysis, although no change in adiposity was observed [59]. GYY4137, formally (p-methoxyphenyl) morpholino-phosphinodithioic acid, is claimed to release hydrogen sulphide slowly, and is used in an effort to mimic longer-term, and potentially more physiological, levels of the gas. However, the backbone structure of a donor such as GYY4137 could contribute to cellular effects, and how this is controlled for is not clear in this and similar studies. There are concerns over how to manipulate hydrogen sulphide levels in a physiologically appropriate manner, and the emergence of new donors will help future studies in this regard [60].

Overall, a model whereby hydrogen sulphide promotes insulin effects in the adipocyte could be proposed through its anti-lipolytic action, although with respect to glucose uptake and lipogenesis its role is less clear. Hydrogen sulphide may also contribute to fat storage by indirectly maintaining or supporting insulin sensitivity in this tissue. Cysteine, when it has been studied in the context of adipose function, largely mimics the effects of hydrogen sulphide [54,55,58] and supports the hypothesis that it is a pro-obesogenic factor. The notion that hydrogen sulphide may mediate some of these effects is perhaps not itself surprising; however, this does not easily fit with the initial findings that obesity is correlated with low plasma levels of hydrogen sulphide.

Cysteine, hydrogen sulphide and insulin-secreting pancreatic β cells

Over the last decade, hydrogen sulphide and/or cysteine have been shown to regulate insulin secretion from pancreatic islets, or clonal insulin-secreting cells such as MIN6, INS-1E and HIT-T15. A consistent finding is that either cysteine or hydrogen sulphide can inhibit insulin release from β cells, particularly following stimulation with glucose [61–66]. It has been proposed that cysteine may mediate its effects, in part, via production of hydrogen sulphide, as CSE and CBS are expressed in the whole pancreas [67]. Indeed CSE is specifically induced in β cells by high glucose [65]. Hydrogen sulphide, added as gas-saturated solution to 100 μM, enhances K-ATP channel opening, whereas NaHS at 100 μM inhibits L-type VDCC calcium channels, both mechanistically consistent with reduction of insulin release [61,68]. Sulphhydration of K-ATP channels has been demonstrated [69] and may contribute to the effect of hydrogen sulphide on insulin secretion. Regulation of mitochondrial respiratory function per se by hydrogen sulphide may be of further importance, given the key role of the ADP:ATP ratio on the secretory function of islet cells. Apart from a role in regulating insulin secretion per se, hydrogen sulphide may regulate other aspects of β cell physiology, including viability. In the streptozotocin-induced diabetes model, elevated hydrogen sulphide was implicated in contributing to β cell failure [70] and hydrogen sulphide prepared as saturated gas solution, administered at doses of 50–200 μM, were found to decrease the viability of INS-1E cells in culture [71]. In contrast, NaHS used at 100 μM was found to have a protective role in maintaining islet function and viability following high-fat feeding in rats [72]. Differences in the dose and duration of exposure may be a factor determining the overall effect of hydrogen sulphide on islet function, as in other tissues. Regulation of endogenous hydrogen sulphide production is also a complex process. Exposure to 20 mM glucose decreased hydrogen sulphide production in INS-1E insulinoma cells [61], whereas similar glucose exposure stimulated expression of CSE in primary mouse islets [65]. Plasma levels of hydrogen sulphide in a variety of diabetic models also reveals a complex picture. Lowered [73–75] and elevated [76,77] levels have been reported in blood or plasma of type 1 and type 2 diabetes models. Even considering type 2 diabetes models exclusively, the aetiology of the diabetic state is different across the studies and the methodologies used to derive blood or plasma measurements of hydrogen sulphide are discordant, and thus interpretation remains somewhat controversial. If the current consensus, that hydrogen sulphide suppresses insulin secretion, holds, then the low hydrogen sulphide levels found in the plasma of type 2 diabetic patients [21,22] may be a compensatory response. This may be permissive for the development of hyperinsulinaemia, which is required to maintain normal glucose levels in an insulin-resistant state. It appears a critical issue to clarify the effects of hydrogen sulphide on islet function, development and dysfunction in diabetes.

Lessons from genetic models with altered cysteine and sulphide metabolism

Complementing pharmacological and association studies, genetic models relevant to the production and
metabolism of cysteine have been informative. So far, there has been a tendency to find that in those models resulting in lowered plasma total cysteine levels, lower adiposity is also observed whereas lean mass is largely unchanged [78]. Notably, few of these models have been studied with respect to their hydrogen sulphide levels. The models summarized below provide insight into the regulation of, and phenotypic consequences of, cysteine and hydrogen sulphide perturbation.

The trans sulphur pathway: CBS and CSE

Insights into the metabolism and physiology of cysteine have been obtained from the study of transgenic manipulation of the genes Cbs and Cse. Mice homozygous for a knockout of the Cbs gene (Cbs−/−) rarely survive after age 5 weeks [79]. For this reason, most studies report on Cbs deficiency, rather than full knockout of the gene. By studying heterozygotes [80] or Cbs−/− that transgenically express a hypermorphic human CBS enzyme [81], Cbs−/− have been studied but, by necessity, experiments are restricted to very young mice [82]. Cse gene knockout (Cse−/−) is not associated with gross abnormality or lethality unless cysteine is limited in the diet [82,83]. Genetic backgrounds can significantly influence the phenotypic observations made with these genetic models. Lethality before adulthood of the Cbs−/− is observed on a C57BL/6 J, DBA/2 J or BALB/cA background, whereas on a C3H/HeJ background significant numbers of pups survive [84]. Indeed, Cbs−/− mice that survive on the C3H/HeJ genetic background lose many of the phenotypic aspects considered typical for this model, suggesting induction of a compensatory system.

In spite of these caveats, some consistent findings relating to cysteine metabolism are summarized here. Knockout or deficiency of either Cbs or Cse in mice alters cysteine metabolism, consistent with their role in cysteine production from methionine. In some studies, this is reflected by lowered plasma or tissue levels of cysteine in Cbs [85a] and Cse-knockout or -deficient mice [83,86]. Hyperhomocysteinaemia is another consistent finding in these genetic models [80,82,87]. Notably, homocysteine is a long-established risk factor for vascular disease when elevated [88]. Another prediction of lowered cysteine production in tissue is reduced tau- rine. Cysteine dioxygenase (CDO) activity in liver, a first step in taurine synthesis, is increased when cysteine is elevated [89]. Consistent with this, Cse−/− and Cbs−/− mice have low plasma taurine compared to control mice [82]. Also consistent with a deficiency in cysteine, glutathione levels are lower in the liver of Cbs−/− [85] and Cse−/− [83] mice. Moreover, in the Cse−/− model, glutathione levels were normalized when cysteine was supplemented in the diet [83].

An expectation from Cbs−/− and Cse−/− deficient models is reduced hydrogen sulphide synthesis. In some studies this is the case. Lower plasma hydrogen sulphide levels have been reported in Cse−/− [83,86,87] and Cbs heterozygote mouse models [80]. The expected reduction in hydrogen sulphide levels is not always observed in these models, for example liver hydrogen sulphide levels were no different from controls in Cse−/− [87]. Furthermore, in that study protein levels of MPST, capable of hydrogen sulphide production, were increased. Moreover, sulphide quinone oxidoreductase (SQR) levels, important for oxidation of hydrogen sulphide, were elevated in this Cse−/− model, not expected in a model of hydrogen sulphide insufficiency. These findings highlight that a complex interplay and feedback mechanism(s) exists to regulate hydrogen sulphide signalling and the resultant phenotypic consequences.

Of relevance to adiposity, in one study of Cbs−/− hypomorphic mice, weight gain and in particular fat mass were lower than controls [90]. When maintained on a high-cysteine diet, Cse−/− mice are typically of normal body weight. On a cysteine-restricted diet, however, weight loss is apparent [82,83], including reduced fat mass [83]. Supplementing drinking water with cysteine (1 mg/ml) partly reversed this weight loss, whereas daily intraperitoneal injection of NaHS (39 μg/kg), did not [83]. A more recent study of Cse−/− mice on an atherogenic diet also found reduced weight gain compared to controls [91].

Little is known about glycaemic control in the Cbs and Cse knockout models. Very young (2 week-old) Cbs−/− mice have lower plasma glucose levels, whereas comparably young Cse−/− mice do not [92], a finding that may simply reflect the severity of the Cbs−/− phenotype and makes assessment of the direct effects of hydrogen sulphide on glucose homeostasis difficult. Hepatic glycogen was higher in fed or mildly fasted Cse−/− mice, implying enhanced glucose storage in this tissue [51]. Indeed, primary hepatocytes from these Cse−/− mice exhibited increased glucose uptake and reduced gluconeogenesis when compared to control hepatocytes. Moreover, islets from Cse−/− mice secreted more insulin in response to glucose [68]. In contrast to these metabolically favourable phenotypes, another group reported that Cse−/− mice fed a high-fat diet developed more severe glucose intolerance after 6 months [72]. Given the complexity of changes attributable to lack of CSE enzyme activity, it remains unclear whether a change in hydrogen sulphide production is a primary mediator of altered glucose homeostasis.

More established, perhaps, are the profound effects on lipid metabolism and inflammatory pathology found in the liver of Cbs and Cse genetic models. Livers of Cbs−/− hypomorphic mice develop hepatic fibrosis and steatosis [79,93]. In another study, Cbs−/− hypomorphic mice showed increased hepatic mRNA levels for genes encoding lipid synthesis enzymes, and genes involved in the endoplasmic stress response [94]. Cbs−/− hypomorphic and Cbs−/− mice are reported to have lower plasma levels of HDL and higher LDL, non-esterified fatty acids and triglycerides [82,95]. A reduction in β-oxidation of fatty acids is part of the mechanism for these phenotypes [95]. By contrast, Cse−/− mice are protected from oxidative damage in the liver following lipopolysaccharide and galactosamine challenge [87].
Cse\textsuperscript{−/−} mice in some models appear normal with respect to lipid phenotypes [82]. However, Cse\textsuperscript{−/−} mice exposed to an atherogenic diet had higher total cholesterol levels, lower plasma triglycerides [91] and developed more extensive atherosclerotic lesions in their vessels. These phenotypes were reversed, in part, by intraperitoneal injection of NaHS (39 μg/kg), supporting that attenuated production of hydrogen sulphide in Cse\textsuperscript{−/−} mice is an important factor in disease development. Of note, the correction of lipid abnormalities in this study did not prevent development of lesions in Cse\textsuperscript{−/−} mice, further implicating vessel-specific hydrogen sulphide signalling in atherosclerosis.

It is clear that CBS and CSE have roles relevant to glucose and lipid metabolism in the liver, and perhaps throughout the body, and have the potential to influence adiposity and diabetes. The relationship of reduced adiposity found in the gene deficiency models fits the clinical relationship with plasma cysteine, but less so with that reported for plasma hydrogen sulphide. Further work is required to clarify these important relationships.

Reductive sulphide production: MPST

Another established source of endogenous sulphide is from the reductive removal of persulphide at the active site of 3-mercaptoppyruvate sulphur transferase (MPST) [96]. 3-Mercaptoppyruvate (3-MP) is a product of the transamination of cysteine. MPST then acquires the sulphur from 3-MP as a persulphide into its active site, which, in combination with reduced thioredoxin, can be released as hydrogen sulphide. A Mpst\textsuperscript{−/−} mouse has been generated, but no reports exist from this model describing cysteine, sulphide or phenotypes relating to adiposity, carbohydrate or lipid metabolism [97].

Methionine cycle: PEMT and BHMT

The methionine cycle, upstream of the transphosphatidylethanolamine N-methyl transferase (PEMT) and BHMT. Knockout mice for both Pemt and Bhmt have been generated. Pemt\textsuperscript{−/−} mice show a protected metabolic phenotype, resisting the weight gain and deterioration of glucose metabolism caused by a high-fat diet [98,99]. Apart from having lower homocysteine levels in plasma [100], no data exist regarding tissue or plasma cysteine in Pemt\textsuperscript{−/−} mice. The Bhmt\textsuperscript{−/−} mouse exhibits lower plasma cysteine, reduced fat mass, smaller adipocytes and improved glucose disposal following insulin injection [101]. The involvement of hydrogen sulphide in these phenotypes remains to be investigated. However, methionine cycle intermediates are known to activate CBS activity [102], consistent with a potential impact of the methionine cycle on sulphide generation and its effects.

Glutathione turnover: GGT and GCLM

Glutathione offers a major source of cysteine in the cell, with steady-state concentrations generally far greater than that of cysteine. The production of glutathione from cysteine is contributed to by glutamate–cysteine ligase (GCL). A subunit of this complex (GCLM) has been knocked out in mice and results, as expected, in lower plasma and tissue glutathione but also in greatly reduced plasma cysteine [103]. Strikingly, these Gclm\textsuperscript{−/−} mice appear significantly affected in their adipose and liver metabolism; displaying a higher metabolic rate despite impaired whole-body lipid oxidation, reduced fat storage in adipose, reduced fat accumulation in liver, greater oxidative stress, yet maintained insulin sensitivity on a high fat diet. Another study of this model found lower mRNA levels for genes involved in fatty acid oxidation and synthesis, as well as a profound protection from the development of hepatic steatosis [104].

Another enzyme relevant to glutathione turnover is γ-glutamyl transferase (GGT), which is involved in recycling glutathione to cysteine. The enzyme participates in a number of other reactions and cannot be considered uniquely related to cysteine metabolism. Ggr\textsuperscript{−/−} mice display alterations in glutathione and lower cysteine in plasma, suggesting that recycling of cysteine from glutathione represents an important source of the amino acid in tissues. Fat mass changes were not reported, but the mice were significantly reduced in body weight [105].

Taurine biosynthesis: CDO and CSAD

Taurine is an essential and abundant metabolite of cysteine. The first step in its synthesis is by oxidation of a cysteine with molecular oxygen by the enzyme cysteine dioxygenase. The phenotype of mice with knock-out of the cysteine dioxygenase gene (CDO) does not follow the predictions relating to cysteine levels and adiposity. Thus, while Cdo\textsuperscript{−/−} mice have elevated cysteine levels, as expected, the mice exhibit lowered fat mass [106]. However, hydrogen sulphide levels are elevated in Cdo\textsuperscript{−/−} mice, which does fit the reported clinical relationship between hydrogen sulphide and obesity. The phenotype of Cdo\textsuperscript{−/−} mice may be due to the toxicity of hydrogen sulphide and other altered metabolites, rather than a metabolic influence on fat mass per se [107]. Cysteine sulphinic acid decarboxylase (CSAD) follows on from CDO in the taurine biosynthesis pathway, converting cysteine sulphinate to hypotaurine. CSAD may prevent accumulation of sulphite – a spontaneous breakdown product of cysteine sulphinate. A knockout mouse has been generated [108] but the result is lethal unless neonates are supplemented with taurine, and few data exist regarding any effects on metabolism in general.
Genetic modification of the sulphide oxidation unit (SOU) represents a target for manipulating sulphide levels

While the genetic models discussed above largely concern cysteine metabolism or cytoplasmic sulphide generation, mitochondria are known to oxidize hydrogen sulphide, in part contributing to respiration, but also to prevent toxic build-up. The known components of this system have been referred to as the sulphide-oxidizing unit (SOU). The SOU consists of the sulphide quinone reductase (SQRDL), sulphur dioxygenase (SDO/ETHE1) and thiosulphate sulphur transferase (TST/rhodanese) [32,33]. In addition, sulphite is oxidized further to sulphate through sulphite oxidase (SUOX) [34]. Together, this system is capable of oxidizing sulphide into sulphate and thiosulphate [109]. As the system is predicted to lower hydrogen sulphide levels by oxidation via respiration, it is not surprising that the Ether+/− mouse, a model of ethylmalonic encephalopathy, is characterized by hydrogen sulphide levels in liver and muscle >10-fold higher than in control mice, and by a short life span [110]. As yet, no reports exist for genetic manipulation of either the SQRDL or SUOX genes, although they have been studied in other ways and are suggested to be essential for the detoxification of sulphide and sulphite, respectively [111,112]. New studies on mice with gene deficiency of Tsr are emerging, with phenotypes in the context of ischaemia–reperfusion injury in the heart (outlined in Abstract form [113]). A full analysis of phenotypes related to metabolic disease are currently ongoing in our laboratory and will add to the increasingly recognized role for enzymes in the hydrogen sulphide breakdown pathway in many physiological processes.

Summary of findings in genetic models of altered sulphur metabolism with respect to obesity

Genetic and biochemical investigations have delineated many of the important enzymes involved in the metabolism of cysteine and hydrogen sulphide (Figure 2). In the context of obesity, a number of the gene knockout models presented above mirror the clinical findings that obesity is associated with high plasma total cysteine. Gene knockout of enzymes of the methionine cycle (in particular Bhmt), the transulfuration pathway (Cbs and Cse) and glutathione metabolism (Gclm and Ggt) all reduce body weight, usually along with reduced adiposity, and present with lowered plasma cysteine. As yet, no genetic model has been described linking elevated cysteine to higher adiposity. Indeed, the Cdo knockout mouse has been shown to have significantly higher plasma cysteine and yet reduced body weight.

The clinical finding that plasma hydrogen sulphide correlates negatively with obesity is less well supported by the evidence from transgenic models. Of the models discussed, the Cbs, Cse, Cdo and Sdo genetic models have, in some studies at least, reported changes in plasma or tissue hydrogen sulphide, in the direction that would be predicted by their biochemical functions. However, in models where lower plasma hydrogen sulphide are found, such as the Cse- and Cbs-deficient or -knockout mice, lower rather than higher fat mass has been observed. The Cdo- and Sdo-knockout mice, on the face of it, support the clinical relationship, as they have elevations in hydrogen sulphide and lower body weight. These elevations in hydrogen sulphide are, however, reported in tissue, not plasma, and little has been reported about fat mass per se. Regardless of the similarity or otherwise to the clinical findings, caution must be exercised in translating adipose and whole body weight phenotypes of genetic models. Phenotypes that result from toxicity, as has been suggested for both the Sdo and Cdo knockout, and some Cbs and Cse models, greatly hamper interpretation.

The majority of models discussed above have a major impact on sulphur amino acid metabolism, including of cysteine and homocysteine. Genetic manipulation of components of the SOU may allow assessment of the contribution of hydrogen sulphide, and potentially sulphite, thiosulphate and sulphate, to lipid and carbohydrate metabolism, without necessarily interfering with cysteine metabolism directly.

Summary of findings in genetic models of altered sulphur metabolism with respect to diabetes

Some of the above findings, in agreement with the existing clinical data, suggest that lowered cysteine is associated with impaired glucose homeostasis or insulin resistance. Few of the studies of the gene knockout models discussed above test this concept directly. Perhaps significantly, Bhmt−/− mice show lower cysteine levels and reduced adiposity, but improved glucose tolerance and insulin sensitivity. This association between lower plasma cysteine does not support the clinical data. Knockout of the transulfuration enzymes CSE and CBS present with pathology in the liver that might be considered diabetogenic; however, strict assessment of glucose homeostasis and insulin sensitivity is still lacking. Similarly, GCLM knockout mice, while presenting with lower cysteine, glutathione and increased oxidative stress, are reported to be comparable to controls regarding insulin sensitivity. With respect to hydrogen sulphide level, as data are lacking from the some of the above models, little can be said regarding its role in the observed phenotypes relating to diabetes or insulin function. In those knockout models where hydrogen sulphide levels are reported (Cbs, Cse, Cdo and Ethe1), a full investigation of potential diabetic phenotypes has not been reported. There therefore remains considerable uncertainty regarding the genetics of cysteine/sulphide-metabolizing enzymes with respect to insulin resistance or diabetes.
Conclusions and future directions

The association between obesity and cysteine in humans appears robust. However, it remains possible that the true mechanism underlying the association is a factor(s) other than cysteine, which concurrently drives elevations in plasma cysteine. Data on the dietary effects of cysteine and fat gain across a wide range of models is lacking. Differences between rodent strains, dietary sources and manipulation, the impact on microbiota and interactions with the non-sulphur components of the diet may all influence how dietary cysteine effects adiposity. Similar issues apply to hydrogen sulphide’s effects, which are obscured by a lack of standardization in measurement and administration methods. The field would benefit from a systematic analysis of tissue and blood/plasma sulphur metabolites across the key transgenic models. Robust methods for the measurement of hydrogen sulphide will be required in order to reflect true endogenous levels. The existence in blood or tissue samples of polysulphides [83], acid-labile sulphur pools [114] and possible chemical interference by other factors can confound measurement techniques and result in wide variation in estimates of hydrogen sulphide [115]. Tissue-specific or inducible manipulation may be particularly informative to minimize toxicity and lethality that might result from whole-body gene knockouts.

A full understanding of the impact of variation within the homologous human genes of the cysteine–sulphur metabolism pathway on metabolic parameters would also be informative. Nevertheless, exogenous administration of cysteine or hydrogen sulphide inhibits adrenergic-stimulated lipolysis from adipocytes [14,59,60] and glucose-stimulated insulin release from β cells [61–66]. Cysteine has further important effects on glutathione and taurine production in the tissue and lethality that might result from whole-body gene knockouts.

Acknowledgements

We would like to thank members of the Molecular Metabolism Group for insights, namely Martin Barrios-Llerena, Clare McFadden and Matthew Gibbins. This study was supported by a Wellcome Trust Investigator Award (Grant No. WT100981MA) to NMM.

Author contributions

RC wrote the manuscript and NMM edited it.

References


