Translating aetiological insight into sustainable management of type 2 diabetes

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Received: 3 April 2017 / Accepted: 11 August 2017 / Published online: 15 November 2017
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Abstract Using a low-energy diet as a tool, it has been possible to elucidate the sequence of pathophysiological changes that lead to the onset of type 2 diabetes. Negative energy balance in type 2 diabetes causes a profound fall in liver fat content resulting in normalisation of hepatic insulin sensitivity within 7 days. As the period of negative energy balance extends and liver fat levels fall to low normal, the rate of export of triacylglycerol from the liver falls. Consequent to this, the raised pancreas fat content falls and in early type 2 diabetes, normal first-phase insulin secretion becomes re-established with normal plasma glucose control. This research, driven by the predictions of the 2008 twin cycle hypothesis, has led to a paradigm shift in understanding. Studying the reversed sequence of pathophysiological changes, the linked abnormalities in liver and pancreas have been revealed. Early type 2 diabetes is a potentially reversible condition. Surprisingly, it was observed that the diet devised as an experimental tool was actually liked by research participants. It was associated neither with hunger nor tiredness in most people, but with rapidly increased wellbeing. A defined period of weight loss followed by carefully planned weight maintenance—the ‘One, Two’ approach—has since been applied in clinical practice.

Motivated individuals can reverse their type 2 diabetes and remain normoglycaemic over years. A large study is underway to evaluate the applicability of this general approach to routine primary care practice as a long-term management strategy.

Keywords Aetiology • Beta cell function • Liver fat • Low-energy diet • Management • Pancreas fat • Pathophysiology • Type 2 diabetes • Weight loss • Weight maintenance

Abbreviations

Counterpoint COUNTERacting Pancreatic inhibitiOn of INsulin secretion by Triglyceride
Counterbalance COUNTERacting BetA cell failure by Long term Action to Normalize Calorie intakE
DiRECT Diabetes Remission Clinical Trial

Introduction

A new concept of aetiology of type 2 diabetes resulted from several new research observations in 2006 [1]. Over the past decade the predictions of this twin cycle hypothesis have been tested using as a dynamic tool a low-energy diet [2, 3]. A new, simplified view of the physiological basis of type 2 diabetes has emerged, and this points to a novel approach to management. The nutritional intervention designed to evaluate the physiological basis of the twin cycle hypothesis was observed to be surprisingly well accepted by research participants, and therapeutic use has followed. Weight loss averaging 15% of body weight is reproducibly obtained with potential reversal of type 2 diabetes to a sustained non-diabetic state. Although the effect

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Electronic supplementary material The online version of this article (https://doi.org/10.1007/s00125-017-4504-z) contains a slideset of the figures for download, which is available to authorised users.
of weight loss in bringing about resolution of type 2 diabetes is not new, the easily explained pathophysiology enhances engagement. Additionally, the routinely reproducible method of bringing about and sustaining 15% weight loss permits clinical application. This review summarises the new information, bringing together metabolic knowledge of disease mechanisms with details of proven nutritional approaches.

The new understanding of type 2 diabetes

The new hypothesis predicted that there were metabolic vicious cycles operating in liver and pancreas, and that these should be able to be reversed. The primary driver of the metabolic problem was identified as positive energy balance, and reversing this was predicted to restore normal blood glucose control (Fig. 1) [1]. Each step shown in Fig. 1 has subsequently been tested and confirmed.

In individuals with a short duration of type 2 diabetes, an average of 15% weight loss allowed fasting plasma glucose to return to normal within 7 days [2]. The rapid return to normal was associated with a 30% decrease in liver fat content and normalisation of liver insulin sensitivity. The COUNTERacting Pancreatic inhibitOn of Insulin secretion by Triglyceride (Counterpoint) study also demonstrated that over the 8 week study period the level of fat in the pancreas gradually decreased and in step with this, first-phase insulin response returned to within the normal range [2]. In contrast, there was no change in muscle insulin resistance.

If the observed changes did represent normalisation of the underlying mechanisms of type 2 diabetes, rather than merely the effect of underfeeding, the beneficial changes should persist after return to normal eating. This was tested in the COUNTERacting BetA cell failure by Long term Action to Normalize Calorie intakE (Counterbalance) study. During a 6 month period of weight stability, levels of fat in the liver and pancreas remained constant, with normal function of these organs in those individuals returned to non-diabetic metabolic control (Fig. 2) [9].

Importantly, after the 15% weight loss in Counterbalance, half of the participants still had a BMI > 30 kg/m², but this had no impact upon their ability to avoid subsequent intra-organ fat re-accumulation. Each individual had dropped below a personal threshold permitting safe storage of fat in the subcutaneous adipose tissue.

Fig. 1 The 2008 twin cycle hypothesis. During chronic intake of more energy than is expended each day, any carbohydrate in excess of requirement must undergo conversion to fat in the liver to permit storage of the metabolic energy as fat [4]. As this process is sensitive to endogenous insulin, individuals with a degree of insulin resistance (and hence higher plasma insulin levels) will tend to accumulate liver fat more readily than others. If the subcutaneous adipose tissue stores have reached capacity, the newly synthesised fat will accumulate in the liver together with excess dietary fat. There it inhibits insulin suppression of glucose production by the liver, and a vicious cycle of hyperinsulinaemia and increased glucose production becomes established. Too much fat in the liver leads to increased export of fat in the form of VLDL-triacylglycerol [5] (shown in lighter red to indicate this is secondary to loss of insulin suppression of liver glucose production). When subcutaneous fat storage capacity is exhausted (i.e. exceeds the personal fat threshold), this will increase fat delivery to all tissues, and the pancreatic islets take up fat avidly [6]. Postprandial hyperglycaemia causes increased and prolonged insulin secretion with further stimulation of de novo lipogenesis. This second vicious cycle thus further increases de novo lipogenesis, and fat delivery to the pancreas. Over many years, the excess pancreas fat brings about loss of specialised function and de-differentiation of the beta cell [7]. Eventually, the inhibitory effects of fatty acids and glucose on the islets reach a trigger level leading to a relatively sudden onset of clinical diabetes. The twin cycle hypothesis predicted that both vicious cycles could be reversed by inducing negative energy balance. Figure adapted from [1] and [8].
compartment, and this was equally so whether their BMI dropped from 40 to 36 kg/m² or from 28 to 24 kg/m². This illustrates the important concept of the personal fat threshold [10]. The ability to store fat safely in adipose tissue varies considerably between individuals with otherwise normal metabolism [11, 12], and once that capacity has been exceeded the body has to use other storage sites, including liver and pancreas. Individuals can exceed their personal fat threshold at BMIs well within the normal range, as illustrated by the finding that the type 2 diabetes prevalence for women with a BMI of < 22 kg/m² is 4.3-fold that for women with a BMI of 23–24 kg/m² [13]. That is a remarkable increase in relative risk for a state of overweight or obesity. In the responders, pancreas fat decreased to low levels, but in the non-responders only fell to levels equivalent to baseline responder levels. At baseline, the first-phase insulin response was higher in responders and increased to normal levels, whereas the grossly deficient baseline level in non-responders hardly changed. Although non-responders normalise both liver fat content and liver insulin sensitivity, plasma glucose remains elevated in the absence of normalised beta cell function and low fasting plasma insulin levels [9]. *p < 0.05; **p < 0.01 vs baseline. To convert HbA1c values from % to mmol/mol, subtract 2.15 and multiply by 10.929. Triglyceride is elsewhere referred to as triacylglycerol. Copyright: ADA [9]. Copyright and all rights reserved.

Fig. 2 The Counterbalance study. Thirty people with type 2 diabetes of up to 23 years duration lost approximately 15 kg in weight then maintained steady weight for 6 months. Those achieving non-diabetic fasting plasma glucose levels were classified as responders and those whose levels remained in the diabetic range, as non-responders. In (b-d) the three columns for each group represent (from left to right) baseline on usual drug therapy; after weight loss, on an isoenergetic diet; and after a further 6 months of weight stability. (a) The change in HbA1c in responders (circles) and non-responders (triangles). (b) There was a similar and sustained decrease in liver fat content in both groups despite ongoing overweight or obesity. (c) In the responders, pancreas fat decreased to low levels, but in the non-responders only fell to levels equivalent to baseline responder levels. (d) At baseline, the first-phase insulin response was higher in responders and increased to normal levels, whereas the grossly deficient baseline level in non-responders hardly changed. Although non-responders normalise both liver fat content and liver insulin sensitivity, plasma glucose remains elevated in the absence of normalised beta cell function and low fasting plasma insulin levels [9]. *p < 0.05; **p < 0.01 vs baseline. To convert HbA1c values from % to mmol/mol, subtract 2.15 and multiply by 10.929. Triglyceride is elsewhere referred to as triacylglycerol. Copyright: ADA [9]. Copyright and all rights reserved. Material from this publication has been used with the permission of ADA.

The observation that type 2 diabetes appeared to be a state of excess fat in the pancreas causing the pathognomonic defect in beta cell function, raises the question of how the fat could be interfering with beta cell function. A series of studies has demonstrated that excess saturated fat produces endoplasmic reticulum stress, and that this is associated with de-differentiation of the beta cell [16–19]. It appears to enter a survival mode in which its specialised functions are shut down. Removal of the fat allows re-expression of the insulin gene and resumption of acute glucose-mediated insulin secretion [7].

Understanding what is happening in the pancreas is the key to type 2 diabetes, as the condition never occurs without a major decrease in acute insulin secretion [20, 21]. The volume of the pancreas in type 2 diabetes is only a half to two-thirds that of people with normal glucose tolerance [22, 23]. Also, the pancreas in type 2 diabetes has a markedly irregular border. As a consequence, when selecting a volume of tissue by imaging to be within the pancreas it is more difficult to include only pancreas parenchyma and not the visceral fat that...
Type 2 diabetes is commonly regarded as being heterogeneous, with possible differing aetiologies in different groups requiring specific approaches to therapy [28]. Individuals vary markedly with respect to any physiological variable, and both genetic and environmental heterogeneity is clear [29]. However, it must be considered that the observed heterogeneity resides in the individuals, not in the basic mechanisms of disease. A parallel could be drawn with observations on the effect of changing an indisputably single mechanism. For example, lisinopril exerts very different effects on blood pressure in different individuals [30]. Observation of the heterogeneity of response to antihypertensive therapy (due to ethnic, genetic and environmental factors affecting each individual) may be interpreted by an observer blinded to the nature of the therapy to suggest that several different drugs are being used. In reality, a single drug is being applied to individuals who respond in a heterogeneous manner. In the same way, the single cause of ectopic fat affecting beta cell function, revealed by the studies discussed above, could be misinterpreted as different specific causes. These considerations apply only to type 2 diabetes itself—a very common condition. There are some clear-cut rare conditions such as monogenic, slow-onset type 1 and pancreatic diabetes that must be considered when making a diagnosis, and there may be other rare conditions that are as yet unrecognised. However, type 2 diabetes itself is a disease so simple that prevalence increases dramatically when a population has ready access to cheap food and decreases when food is scarce [31, 32]. When the Pima Indians were living as subsistence farmers, a comprehensive health survey reported only one case of diabetes, and Joslin famously observed no diabetes at that time [33, 34]. Yet subsequent overnutrition revealed susceptibility to type 2 diabetes in around 40% of the population [35]. Added to this is the steadily increasing beta cell defect as disease duration increases in any one individual, indicating that duration must be taken into account in comparing groups. The lack of precise knowledge of duration of disease in most people adds to apparent heterogeneity. Hence, it is postulated that anyone who develops true type 2 diabetes has accumulated more fat in the pancreas than they individually can tolerate. The cause of this common disease is not heterogeneous, unlike the individuals.

The hypothesis-driven sequence of studies revealing the pathogenesis of type 2 diabetes by observing changes during the return to normal glucose metabolism is illustrated in Fig. 3.

Testing the twin cycle hypothesis by inducing 15% weight loss

Conventional dietetic approaches to achieving weight loss are not notably successful. The prolonged nature of most interventions is a drawback, and the two major adverse factors associated with this approach are hunger and daily decisions about what and how much to eat. These potentially modifiable factors were accounted for in the design of the Counterpoint and Counterbalance studies with a short, time-limited weight loss phase and a subsequent weight maintenance phase—a ‘One, Two’ approach.

To induce weight loss, a liquid formula diet provided in individual meal sachets was used (2.51 MJ [600 kcal]/day; three sachets). To minimise constipation, up to 240 g/day of non-starchy vegetables were allowed. The total energy intake was thus around 2.93 MJ (700 kcal)/day. A relatively high sugar content was necessary for palatability, but this did not prevent normalisation of fasting plasma glucose within 7 days despite withdrawal of oral hypoglycaemic agents [2]. Unexpectedly, the dietary approach adopted to allow the hypothesis to be tested was actually liked by the participants.

Counterbalance demonstrated that the individuals who did not return to non-diabetic blood glucose control typically had a longer disease duration, but notably already had severely impaired beta cell function at baseline [9]. Weight loss produced normalisation of liver fat content with normalisation of hepatic insulin sensitivity in all, but this alone was insufficient to normalise glucose control. The apparent heterogeneity of response lay in the duration-dependent progression to complete beta cell de-differentiation, whereby those individuals with beta cells still at the reversible stages could achieve complete re-differentiation and resumption of beta cell specialist function [7].
All participants reported almost complete lack of hunger within 1–2 days of commencing the diet. This is striking during achievement of ~15 kg weight loss. Notable wellbeing was reported [37]. Several participants wished to continue the low-energy liquid diet after the 8 week period of the study to meet their own weight target. A recent systematic review exposes the misconception that rapid weight gain inevitably follows rapid weight loss [38]. If weight loss by any regimen is not followed up by a supportive programme it is unlikely to be sustained. In Counterbalance, participants were seen at monthly intervals for the 6 month follow-up period and weight remained stable. Importantly, they were advised in advance to expect to be eating only two-thirds of their usual intake of food. The concept that weight loss is more difficult to achieve in people with type 2 diabetes is incorrect [38].

Following media interest in the Counterpoint results, a large number of emails and letters were received from people with diabetes, and full how-to-do-it information was placed on the Magnetic Resonance Centre website [39]. Analysis of the initial email responses to the information showed that half had used a liquid diet replacement, and half had used small portions of ordinary foods [40]. Weight loss was approximately 15 kg in both groups, illustrating the practical possibility of achieving this degree of weight loss by any energy-restricted approach that is sustainable. There was clearly an important matter of individual preference.

**Nature of the ongoing support required after reversal of diabetes**

The liquid low-energy diet is prescriptive, rendering it reassuringly straightforward to follow. In contrast, the prospect of returning to normal eating after the liquid diet, with inherent decisions about what and how much to eat, is often a time of
great anxiety. RCT evidence confirms that a gradual transition to the weight maintenance diet following weight loss with a low- or very-low-energy diet over a period of weeks is associated with improved weight maintenance at 12 months [41, 42]. A gradual, stepwise change from the low-energy liquid diet to normal foodstuffs was employed in the subsequent studies. During this phase of Counterbalance, weekly face-to-face review was undertaken, with directive advice on what and how much to eat, consideration being given to individual dietary preferences and calculated energy requirements. The energy prescription for weight stabilisation was estimated from achieved body weight using predictive equations [43]. Additional telephone support was also available. Monthly review and weekly self-weighing was then undertaken [44], resulting in weight stability over 6 months. One critical ingredient for success was observed to be family support. If the spouse or partner was not supportive, then the chance of long-term success was low. Notably, in both Counterpoint and Counterbalance spouses/partners reported losing weight as well. Changing the obesogenic microenvironment of the home is critical.

While some individuals can successfully maintain a new lower body weight under their own direction [40], most are likely to require ongoing support to limit weight regain over time [45]. The support that may be required during weight loss maintenance is summarised in Fig. 4.

**Nature of ongoing diet**

An individual who has achieved remission of type 2 diabetes through weight loss remains inherently predisposed to recurrence if there is a return to chronic energy excess. It is therefore of utmost importance to maintain the body weight at a level below the personal fat threshold. However, long-term maintenance of lost weight is recognised as the most challenging aspect of obesity treatment [46]. With appropriate support, significant weight loss can be maintained over the longer term [47].

So which dietary strategy is most appropriate? A low-fat diet (< 30% total energy from fat) has long been the mainstay of dietary advice. This is traceable to the association of high fat intake with cardiovascular death reported in the seven countries study [48]. Epidemiological associations from cross-sectional studies have been shown repeatedly not to represent cause and effect [49], but the belief is ingrained in practice and reflected in current guidelines for type 2 diabetes. However, it is now well established that it is adherence to any strategy rather than a particular macronutrient composition which is likely to result in long-term weight maintenance. Over recent years nutrition guidelines have moved away from prescriptive macronutrient content towards a more person-centred philosophy, acknowledging that there is no ‘one best diet’ for diabetes [50]. Three evidenced approaches (low carbohydrate, Mediterranean and intermittent energy restriction) are discussed in the following section as potential options for a maintenance diet following the initial rapid weight loss period.

For many Europeans, decreasing or omitting the carbohydrate portion of the main meal, with or without change in other meals, is a simple change, easily achievable in the context of family eating. Low-carbohydrate diets (less than 120 g per day of total carbohydrate) and very-low-carbohydrate diets (20–50 g total carbohydrate per day) for diabetes management continue to excite vehement debate [49, 51]. A carbohydrate-restricted diet implies an increase in the ratio of fat to carbohydrate, conflicting strategies where such support is absent. All of these points require exploration at ongoing follow-up visits. Not illustrated in the diagram is input of outside agencies as required and the crucial potential for the whole obesogenic environment to be influenced by policy makers
with long-held beliefs about the risks of higher-fat diets. There is insufficient evidence to recommend an ideal proportion of total energy from carbohydrate in diabetes management [52]. For equivalent weight loss, different compositions of diet do not affect liver fat content nor any other aspect of fat distribution [53]. Despite incorporation into evidence-based nutrition guidelines in 2011 [50], UK clinical practice relating to carbohydrate restriction has not kept pace with the evidence base [54]. Low-carbohydrate diets have been reported to be superior to low-fat diets for short to medium term use in type 2 diabetes, with comparable longer-term (> 12 months) outcomes [55].

The Mediterranean diet is based on a combination of food-stuffs high in monounsaturated fats (primarily olive oil), legumes and vegetables, and with restricted red meat, processed foods and refined carbohydrates. Such diets consistently give benefit in terms of weight control and cardiovascular health [56–58]. A decrease in diabetes incidence independent of weight has also been observed [59]. Given the current evidence, the Mediterranean diet could be recommended as an option for a weight maintenance diet regardless of diabetes remission status. A combined Mediterranean and carbohydrate-restricted diet may be particularly beneficial for those for whom weight loss has not resulted in diabetes remission, almost halving the need for diabetes drugs over 4 years following diagnosis of type 2 diabetes [60].

Novel time-limited approaches to eating (alternate day fasting, time-restricted eating and meal timing during the day) provide an alternative to daily energy restriction. Intermittent energy restriction achieved by a variety of methods is as effective as daily energy restriction in achieving weight loss and maintenance for up to 12 months [61]. Weight loss outcomes were better in studies that applied an energy prescription on the non-fasting (‘feed’) days in order to ensure an overall weekly deficit. In some studies mean weight losses were similar between groups but the proportion of participants losing a clinically significant amount of weight (≥ 65%) was higher with intermittent energy reduction (60–65% at 12 weeks) compared with the daily energy restriction group (37%) [62]. One day of energy restriction per week maintained the improvements in weight and insulin sensitivity (achieved using a 5:2 approach) over 6 months of follow-up in overweight or obese women. The first published results using the 5:2 approach in a population with type 2 diabetes demonstrate comparable reductions in weight and HbA1c between daily and intermittent energy-restricted groups with no adverse effects on exercise levels or appetite [63, 64]. Evidence relating to the use of this approach for longer-term weight maintenance is lacking and warrants further study.

Studies reporting an association of omission of breakfast with increased weight usually have a clear commercial bias [65]. Prospective study reveals no disbenefit of omitting breakfast in terms of eating more later in the day and demonstrates the potential major energy advantage [66]. For some individuals omission of breakfast may usefully contribute to long-term weight control if they do not like breakfast and can readily extend the overnight fast to lunchtime [67].

As different dietary approaches are likely to suit the needs and preferences of different individuals (especially those who have had little success with conventional low-fat diets), a decision aid such as that developed to facilitate informed choices about diabetes medications may be a useful tool. Long-term weight control will be most effectively achieved by finding an approach to eating that an individual, and their family, can sustain. The composition (restricted carbohydrate/restricted fat) is far less important than the overall quantity of food, and there are varying, equally valid ways of achieving this goal.

**Physical activity**

There is clear-cut evidence for encouraging sustainable, daily physical activity during the long-term weight maintenance phase, alongside energy control [68]. This is of great importance. However, little attention has been paid to the compensatory eating phenomenon, which is a major hazard of the weight loss phase itself [69–71]. If overweight or obese individuals embark upon an exercise regimen, there tends to be an increase in weight due to an increase in energy intake, partly conscious and partly subconscious. This varies between individuals [70], but extra exercise can be avoided with no disbenefit to substantial weight loss. The energy expenditure during exercise that is tolerated by most overweight, older people is modest and easily cancelled out by a snack. During the weight loss phase of both Counterpoint and Counterbalance, participants were asked to continue usual activities but to avoid any increase in physical activity. This appeared to contribute to the achievement of the target ~ 15% weight loss in 8 weeks. In contrast, the more widely used approach of advising an increase in exercise from the beginning of a longer-term period of a diet with modest energy restriction may contribute to falling short of the weight loss target [72]. A randomised study testing the effect of avoiding additional prescribed exercise during the weight loss phase of any energy-restricted diet may be helpful in clarifying the effectiveness of this approach.

**Previous studies of weight loss as management for type 2 diabetes**

Observation of the beneficial effect of weight loss on blood glucose control is not new. Ayurvedic scriptures from
2000 years ago recognised fat and thin forms of diabetes and recommended fasting and exercise [73]. Bouchardat described the resolution of glycosuria in diabetes during the famine caused by the 1870 siege of Paris [74]. Allen’s low-energy diet also permitted prolonged survival of individuals with what is now recognised as type 1 diabetes [75]. Energy restriction brought about by bariatric surgery has long been known to return blood glucose control to normal in some people [76]. A 6 day very-low-energy diet supervised in hospital was shown to improve fasting plasma glucose from 18.6 to 11.2 mmol/l, but in common with all such studies did not consider the effect of long duration of diabetes [77]. Two studies are notable in using a low-energy diet with follow-up over 1 year [78] and 5 years [79] to demonstrate the beneficial effect on glucose control. In the latter study, one out of 15 remained normoglycaemic off all medications, giving a very favourable ‘number needed to treat’ for the complete reversal of diabetes.

The only older study to have used a very-low-energy diet as a tool to evaluate mechanisms of disease was published in 1986 [80]. In-patient supervision of a 1.38–2.51 MJ/day (330–600 kcal/day) diet for up to 40 days was followed by 60–380 days of outpatient follow-up. This study specifically avoided additional exercise in the weight loss phase, and is one of the few in which participants achieved weight loss of around 15%. This decrease caused a fall in fasting plasma glucose from 15.4 to 6.8 mmol/l, which correlated with a fall in hepatic glucose output. However, the study group of eight included people with a long duration of diabetes, and no change was observed in average insulin response to oral glucose.

### Lifelong reversal of type 2 diabetes?

Information from observational studies on large groups of people with type 2 diabetes suggest an inexorable worsening of blood glucose control with a need for increasing numbers of tablets and, eventually, insulin [81, 82]. The associated demonstration of a steady decline in beta cell function [83] has given rise to the widespread belief that this is inevitable. However, these observational studies were carried out only in the situation of maintained or increasing body weight [84].

The demonstration that the non-diabetic control of glucose metabolism is fully maintained over 6 months is useful in confirming the mechanistic basis of long-term remission of type 2 diabetes [9]. In those who had achieved a post-weight-loss fasting plasma glucose of < 7 mmol/l, liver and pancreas fat content fell to normal levels. First-phase insulin response became and remained normal. There was no accumulation of fat in either pancreas or liver, even though mean BMI was 30 kg/m² [9]. These data are important in clarifying the physiological basis of continuing reversal of type 2 diabetes, separate from the well-recognised human problem of avoiding weight regain in the face of unchanged environmental pressures. Longer-term data are required from formal studies, but non-diabetic metabolic control can be maintained over several years, providing that weight regain is avoided [72, 79, 85].

The Diabetes Remission Clinical Trial (DiRECT) is a large randomised trial that will determine the effectiveness of rapid weight loss followed by supportive follow-up in primary care, head-to-head with conventional management [36]. It will report 12 month outcomes at the end of 2017. The intervention consists of 8 h of training for the practice nurse (or practice dietitian if available), with continuing specialist dietitian support. All oral glucose-lowering agents are withdrawn. The dietary and behavioural approach known as CounterWeight has been trialled for obesity management in primary care with 1 year follow-up demonstrating maintenance of ≥ 15 kg weight loss in 30% [47]. DiRECT will address the question of selection bias and degree of self-motivation that follows from the smaller, pathophysiological studies. It will also provide further evidence of the basic mechanisms of disease. Rapid recruitment to DiRECT reflects the extreme dislike of type 2 diabetes by many people. This is generally underestimated by professionals and, given the previous lack of means to restore normal metabolism, has led to a situation of learned helplessness. The outcomes of DiRECT may indicate an important way forward.

### Summary

1. A single cause of common type 2 diabetes can be postulated
2. Type 2 diabetes appears to be caused by reversible de-differentiation of beta cells following excess fat exposure in those susceptible
3. Heterogeneity lies in the individuals, not the basic mechanism of disease
4. Evidence-based means of achieving 15% weight loss in clinical practice are described
5. Low-carbohydrate, Mediterranean and intermittent fasting diets have an evidence base to justify use to minimise weight regain
6. A supportive approach to these dietary strategies in routine care is required to maximise favourable long-term outcomes
Acknowledgements The research work of E. L. Lim, S. Steven, C. Peters, A. Al-Mrabeh, K. Hollingsworth and S. Zhyzhneuskaya (all of the Magnetic Resonance Centre, University of Newcastle) is gratefully acknowledged.

Duality of interest The authors declare that there is no duality of interest associated with this manuscript.

Contribution statement Both authors were responsible for drafting the article and revising it critically for important intellectual content. Both authors approved the version to be published.

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References

1. Taylor R (2008) Pathogenesis of type 2 diabetes: tracing the reverse route from cure to cause. Diabetologia 51:1781–1789
2. Lim EL, Hollingsworth KG, Arbisala BS, Chen MJ, Mathers JC, Taylor R (2011) Reversal of type 2 diabetes: normalisation of beta cell function in association with decreased pancreas and liver triacylglycerol. Diabetologia 54:2506–2514
3. Steven S, Hollingsworth KG, Small P et al (2016) Weight loss decreases excess pancreatic triacylglycerol specifically in type 2 diabetes. Diabetes Care 39:158–165
4. Schwarz JM, Linfoot P, Dare D, Aghajanian K (2003) Hepatic de novo lipogenesis in normoinsulinemic and hyperinsulinemic subjects consuming high-fat, low-carbohydrate and low-fat, high-carbohydrate isoenergetic diets. Am J Clin Nutr 77:43–50
5. Adiels M, Taskinen MR, Packard C et al (2006) Overproduction of large VLDL particles is driven by increased liver fat content in man. Diabetologia 49:755–765
6. Lalloyer F, Vandewalle B, Percevault F et al (2006) Peroxisome proliferator-activated receptor alpha improves pancreatic adaptation to insulin resistance in obese mice and reduces lipotoxicity in human islets. Diabetes 55:1605–1613
7. White MG, Shaw JAM, Taylor R (2016) Type 2 diabetes: the pathologic basis of reversible beta-cell dysfunction. Diabetes Care 39:2080–2088
8. Taylor R (2013) Type 2 diabetes: etiology and reversibility. Diabetes Care 36:1047–1055
9. Steven S, Hollingsworth KG, Al-Mrabeh A et al (2016) Very low calorie diet and 6 months of weight stability in type 2 diabetes: pathophysiological changes in responders and nonresponders. Diabetes Care 39:808–815
10. Taylor R, Holman R (2015) Normal weight individuals who develop type 2 diabetes: the personal fat threshold. Clin Sci 128:405–410
11. Robbins AL, Savage DB (2015) The genetics of lipid storage and human lipodystrophies. Trends Mol Med 21:433–438
12. Virtue S, Vidal-Puig A (2010) Adipose tissue expandability, lipotoxicity and the metabolic syndrome—an allostatic perspective. Biochim Biophys Acta 1801:338–349
13. Colditz GA, Willett WC, Rotnitzky A, Manson JE (1995) Weight gain as a risk factor for clinical diabetes mellitus in women. Ann Intern Med 122:481–486
14. UKPDS (1991) UK Prospective Diabetes Study (UKPDS). VIII. Study design, progress and performance, Diabetologia 34:877–890
15. Rosenbaum S, Skinner RK, Knight IB, Garrow JS (1985) A survey of heights and weights of adults in Great Britain, 1980. Ann Hum Biol 12:115–127
16. Pinnick K, Neville M, Clark A, Fielding B (2010) Reversibility of metabolic and morphological changes associated with chronic exposure of pancreatic islet beta-cells to fatty acids. J Cell Biochem 109:683–692
17. Pinnick KE, Collins SC, Londos C, Gaugudier D, Clark A, Fielding BA (2008) Pancreatic ectopic fat is characterized by adipocyte infiltration and altered lipid composition. Obesity 16:522–530
18. Talchai C, Yuan S, Lin HV, Sussel L, Accili D (2012) Pancreatic beta cell dedifferentiation as a mechanism of diabetic beta cell failure. Cell 150:1223–1234
19. White MG, Marshall HL, Rigby R et al (2013) Expression of mesenchymal and alpha-cell phenotypic markers in islet beta-cells in recently diagnosed diabetes. Diabetes Care 36:3818–3820
20. Cali AM, Man CD, Cobelli C et al (2009) Primary defects in beta-cell function further exacerbated by worsening of insulin resistance mark the development of impaired glucose tolerance in obese adolescents. Diabetes Care 32:456–461
21. Ferrannini E, Nannipieri M, Williams K, Gonzales C, Haffner SM, Stern MP (2004) Mode of onset of type 2 diabetes from normal or impaired glucose tolerance. Diabetes 53:160–165
22. Macauley M, Pereival K, Thelwall PE, Hollingsworth KG, Taylor R (2015) Altered volume, morphology and composition of the pancreas in type 2 diabetes. PLoS One 10:e0126825
23. Al-Mrabeh A, Hollingsworth KG, Steven S, Taylor R (2016) Morphology of the pancreas in type 2 diabetes: effect of weight loss with or without normalisation of insulin secretory capacity. Diabetologia 59:1753–1759
24. Yamazaki H, Tsuboya T, Kat anuma A et al (2016) Lack of independent association between fatty pancreas and incidence of type 2 diabetes: 5-year Japanese cohort study. Diabetes Care 39:1677–1683
25. van der Zijl NJ, Goossens GH, Moors CC et al (2011) Ectopic fat storage in the pancreas, liver, and abdominal fat depots: impact on beta-cell function in individuals with impaired glucose metabolism. J Clin Endocrinol Metab 96:459–467
26. Kuhn JP, Berthold F, Mayerle J et al (2015) Pancreatic steatosis demonstrated at MR imaging in the general population: clinical relevance. Radiology 276:129–136
27. Al-Mrabeh A, Hollingsworth KG, Steven S, Tiniakos D, Taylor R (2017) Quantification of intrapancreatic fat in type 2 diabetes. PLoS One 12:e0174660
28. Skyler JS, Bakris GL, Bonifacio E et al (2017) Differentiation of diabetes by pathophysiology, natural history, and prognosis. diabetes 66:241–255
29. Tuomi T, Santoro N, Caprio S, Cai M, Weng J, Groop L (2014) The many faces of diabetes: a disease with increasing heterogeneity. Lancet 383:1084–1094
30. van Rijn-Bikker PC, Mairuhu G, van Montfrans GA et al (2009) Self-reported and measured height and weight of heights and weights of adults in Great Britain, 1980. Ann Hum Biol 12:115–127
31. Franco M, Bilal U, Ordonez P et al (2013) Population-wide weight loss and regain in relation to diabetes burden and cardiovascular mortality in Cuba 1980-2010: repeated cross sectional surveys and ecological comparison of secular trends. BMJ 1515:346
32. Himsworth HP (1949) Diet in the aetiology of human diabetes. Proc R Soc Med 42:323–326
33. Hrdlička A (1908) Physiological and medical observations amongst the Indians of southwestern United States and north Mexico. Washington Government Printing Office, Washington DC
34. Joslin EP (1940) The universality of diabetes: a survey of diabetes mortality in Arizona. JAMA 115:203–2038
35. Knowler WC, Bennett PH, Hamman RF, Miller M (1978) Diabetes incidence and prevalence in Pima Indians: a 19 fold greater incidence than in Rochester Minnesota. Am J Epidemiol 108:497–505
36. Leslie WS, Ford I, Sattar N et al (2016) The Diabetes Remission Clinical Trial (DiRECT): protocol for a cluster randomised trial. BMC Fam Pract 17:20
37. Rehakova L, Armijo-Soares V, Adamson AJ, Stevens S, Sniehotta FF (2017) Acceptability of a Very Low Energy Diet in Type 2 Diabetes: Patient Experiences and Behaviour Regulation. Diabet Med. https://doi.org/10.1111/dme.13426
38. Leslie WS, Taylor R, Harris L, Lean ME (2017) Weight losses with low-energy formula diets in obese patients with and without type 2 diabetes: systematic review and meta-analysis. Int J Obes 41:96–101
39. Taylor R (2011) Reversing type 2 diabetes. Available from http://www.ncl.ac.uk/magres/research/diabetes/reversal/#publicinformation. Accessed 19 May 2017
40. Steven S, Lim E, Taylor R (2013) Population response to information on reversibility of type 2 diabetes. Diabet Med 30:e135–e138
41. Johansson K, Neovius M, Herrmannsson E (2014) Effects of anti-obesity drugs, diet, and exercise on weight-loss maintenance after a very-low-calorie diet or low-calorie diet: a systematic review and meta-analysis of randomized controlled trials. Am J Clin Nutr 99:14–23
42. Gripepetz L, Torgerson J, Karlsson J, Lindroos AK (2010) Prolonged refeeding improves weight maintenance after weight loss with very-low-energy diets. Br J Nutr 103:141–148
43. Mifflin MD, St Jeor ST, Hill LA, Scott BJ, Daugherty SA, Koh YO (1990) A new predictive equation for resting energy expenditure in healthy individuals. Am J Clin Nutr 51:241–247
44. Shieh C, Knisely MR, Clark D, Carpenter JS (2016) Self-weighing in weight management interventions: A systematic review of literature. Obes Res Clin Pract 10:493–519
45. Middleton KMR, Patidar SM, Perri MG (2012) The impact of extended care on the long-term maintenance of weight loss: a systematic review and meta-analysis. Obes Rev 13:509–517
46. MacLean PS, Wing RR, Davidson T et al (2015) NIH Working Group Report: innovative research to improve maintenance of weight loss. Obesity 23:7–15
47. Lean M, Borsnahan N, McLoone P et al (2013) Feasibility and indicative results from a 12-month low-energy liquid diet treatment and maintenance programme for severe obesity. Br J Gen Pract 63:e115–e124
48. Keys A (1953) Atherosclerosis: a problem in newer public health. J Mt Sinai Hosp N Y 20:118–139
49. Feinman RD, Pogozelski WK, Astrup A et al (2015) Dietary carbohydrate restriction as the first approach in diabetes management: critical review and evidence base. Nutrition 31:1–13
50. Dyson PA, Kelly T, Deakin T et al (2011) Diabetes UK evidence-based nutrition guidelines for the prevention and management of diabetes. Diabet Med 28:1282–1288
51. Spiro A, Stanner S (2016) The National Obesity Forum report is an opinion piece not a scientific review. Nutr Bull 41:257–269
52. van Wyk HJ, Davis RE, Davies JS (2016) A critical review of low-carbohydrate diets in people with type 2 diabetes. Diabet Med 33:148–157
53. de Souza RJ, Bray GA, Carey VJ et al (2012) Effects of 4 weight-loss diets differing in fat, protein, and carbohydrate on fat mass, lean mass, visceral adipose tissue, and hepatic fat: results from the POUNDS LOST trial. Am J Clin Nutr 95:614–625
54. Mc Ardle PD, Greenfield SM, Avery A, Adams GG, Gill PS (2017) Dietitians’ practice in giving carbohydrate advice in the management of type 2 diabetes: a mixed methods study. J Hum Nutr Diet 30:385–393
55. Dyson P (2015) Low carbohydrate diets and type 2 diabetes: what is the latest Evidence? Diabetes Ther 6:411–424
56. Estruch R, Martinez-Gonzalez MA, Corella D et al (2016) Effect of a high-fat Mediterranean diet on bodyweight and waist circumference: a prespecified secondary outcomes analysis of the PREDIMED randomised controlled trial. Lancet Diabetes Endocrinol 4:666–676
57. Garcia-Fernandez E, Rico-Cabanas L, Rosgaard N, Estruch R, Bach-Faig A (2014) Mediterranean diet and cardiobiadeity: a review. Nutrients 6:3474–3500
58. Martinez-Gonzalez MA, Martin-Calvo N (2016) Mediterranean diet and life expectancy; beyond olive oil, fruits, and vegetables. Curr Opin Clin Nutr Metab Care 19:401–407
59. Salas-Salvado J, Bullo M, Babio N et al (2011) Reduction in the incidence of type 2 diabetes with the Mediterranean diet: results of the PREDIMED- Reus nutrition intervention randomized trial. Diabetes Care 34:14–19
60. Esposito K, Maiorino MI, Petrizzi M, Bellastella G, Giugliano D (2014) The effects of a Mediterranean diet on the need for diabetes drugs and remission of newly diagnosed type 2 diabetes: follow-up of a randomized trial. Diabetes Care 37:1824–1830
61. Davis CS, Clarke RE, Coulter SN et al (2016) Intermittent energy restriction and weight loss: a systematic review. Eur J Clin Nutr 70:292–299
62. Harvie M, Wright C, Peganin M et al (2013) The effect of intermittent energy and carbohydrate restriction v. daily energy restriction on weight loss and metabolic disease risk markers in overweight women. Br J Nutr 110:1534–1547
63. Carter S, Clifton PM, Keogh JB (2016) The effects of intermittent compared to continuous energy restriction on glycemic control in type 2 diabetes; a pragmatic pilot trial. Diabetes Res Clin Pract 122:106–112
64. Harvie MN, Howell T (2016) Could intermittent energy restriction and intermittent fasting reduce rates of cancer in obese, overweight, and normal-weight subjects? A summary of evidence. Advances in Nutrition 7:690–705
65. Brown AW, Bohan Brown MM, Allison DB (2013) Belief beyond the evidence: using the proposed effect of breakfast on obesity to show 2 practices that distort scientific evidence. Am J Clin Nutr 98:1298–1308
66. Clayton DJ, Stensel DJ, James LJ (2016) Effect of breakfast omission on subjective appetite, metabolism, acylated ghrelin and GLP-17-36 during rest and exercise. Nutrition 32:179–185
67. Kealey T (2016) Breakfast is a dangerous meal. 4th Estate, London
68. Wing RR, Phelan S (2005) Long-term weight loss maintenance. Am J Clin Nutr 82:2225S–2255S
69. Finlayson G, Bryant E, Blundell JE, King NA (2009) Acute compensatory eating following exercise is associated with implicit hedonic wanting for food. Physiol Behav 97:62–67
70. Hopkins M, Blundell JE, King NA (2014) Individual variability in compensatory eating following acute exercise in overweight and obese women. Br J Sports Med 48:1472–1476
71. King NA, Homer K, Hills AP et al (2012) Exercise, appetite and weight management: understanding the compensatory responses in eating behaviour and how they contribute to variability in exercise-induced weight loss. Br J Sports Med 46:315–322
72. LookAhead (2014) Eight-year weight losses with an intensive lifestyle intervention and maintenance of type 2 diabetes: a mixed methods study. J Hum Nutr Diet 30:385–393
73. Sheth H, Sheth J, Sheth F, Burn J (2016) The poor patient with diabetes ‘should live like a saint’. Diabet Med 33:134–135
74. Bouchardat A (1875) De la glycosurie ou diabète sucré. Librarie G. Bailliére, Paris [document in French]
75. Allen FM (1914) Studies concerning diabetes. J Am Med Assoc 63: 939–943
76. Friedman MN, Sancetta AJ, Magovern GJ (1955) The amelioration of diabetes mellitus following subtotal gastrectomy. Surg Gynecol Obstet 100:201–204
77. Lara-Castro C, Newcomer BR, Rowell J et al (2008) Effects of short-term very low-calorie diet on intramyocellular lipid and insulin sensitivity in nondiabetic and type 2 diabetic subjects. Metab Clin Exp 57:1–8
78. Wing RR, Marcus MD, Salata R, Epstein LH, Miaskiewicz S, Blair EH (1991) Effects of a very-low-calorie diet on long-term glycemic control in obese type 2 diabetic subjects. Arch Intern Med 151:1334–1340
79. Paisey RB, Frost J, Harvey P et al (2002) Five year results of a prospective very low calorie diet or conventional weight loss programme in type 2 diabetes. J Hum Nutr Diet 15:121–127
80. Henry RR, Wallace P, Olefsky JM (1986) Effects of weight loss on mechanisms of hyperglycaemia in obese non-insulin dependent diabetes mellitus. Diabetes 35:990–998
81. UKPDS (1999) Intensive blood glucose control with sulphonylureas or insulin compared with conventional treatment and risk of complications in patients with type 2 diabetes (UKPDS 33). Lancet 352: 837–853
82. Holman RR, Paul SK, Bethel MA, Matthews DR, Neil HA (2008) 10-year follow-up of intensive glucose control in type 2 diabetes. N Engl J Med 359:1577–1589
83. UKPDS (1995) UK prospective diabetes study 16. Overview of 6 years’ therapy of type II diabetes: a progressive disease. Diabetes 44:1249–1258
84. UKPDS (1998) Effect of intensive blood-glucose control with metformin on complications in overweight patients with type 2 diabetes (UKPDS 34). UK Prospective Diabetes Study (UKPDS) Group. Lancet 352:854–865
85. Taylor R (2016) Calorie restriction and reversal of type 2 diabetes. Expert Rev Endocrinol Metab 11:521–528