

# Internal Clinical Guidelines Team

Draft for consultation

## Type 2 diabetes in adults

**Type 2 diabetes: management of type 2 diabetes in adults**

*Clinical Guideline Update (XXX)*

*Methods, evidence and recommendations*

*September 2015*

*Draft for Consultation*

*Commissioned by the National Institute for  
Health and Care Excellence*



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Healthcare professionals are expected to take NICE clinical guidelines fully into account when exercising their clinical judgement. However, the guidance does not override the responsibility of healthcare professionals to make decisions appropriate to the circumstances of each patient, in consultation with the patient and/or their guardian or carer.

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# 1 Summary Section

## 1.1.2 GDG membership and ICG technical team

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#### 1.25 Strength of recommendations

- 36 Some recommendations can be made with more certainty than others. The Guideline
- 37 Development Group makes a recommendation based on the trade-off between the benefits
- 38 and harms of an intervention, taking into account the quality of the underpinning evidence.
- 39 For some interventions, the Guideline Development Group is confident that, given the

1 information it has looked at, most patients would choose the intervention. The wording used  
2 in the recommendations in this guideline denotes the certainty with which the  
3 recommendation is made (the strength of the recommendation).

4 For all recommendations, NICE expects that there is discussion with the patient about the  
5 risks and benefits of the interventions, and their values and preferences. This discussion  
6 aims to help them to reach a fully informed decision (see also 'Patient-centred care').

#### 7 **Interventions that must (or must not) be used**

8 We usually use 'must' or 'must not' only if there is a legal duty to apply the recommendation.  
9 Occasionally we use 'must' (or 'must not') if the consequences of not following the  
10 recommendation could be extremely serious or potentially life threatening.

#### 11 **Interventions that should (or should not) be used – a 'strong' recommendation**

12 We use 'offer' (and similar words such as 'refer' or 'advise') when we are confident that, for  
13 the vast majority of patients, an intervention will do more good than harm, and be cost  
14 effective. We use similar forms of words (for example, 'Do not offer...') when we are  
15 confident that an intervention will not be of benefit for most patients.

#### 16 **Interventions that could be used**

17 We use 'consider' when we are confident that an intervention will do more good than harm  
18 for most patients, and be cost effective, but other options may be similarly cost effective. The  
19 choice of intervention, and whether or not to have the intervention at all, is more likely to  
20 depend on the patient's values and preferences than for a strong recommendation, and so  
21 the healthcare professional should spend more time considering and discussing the options  
22 with the patient.

#### 23 **Recommendation wording in guideline updates**

24 NICE began using this approach to denote the strength of recommendations in guidelines  
25 that started development after publication of the 2009 version of 'The guidelines manual'  
26 (January 2009). This does not apply to any recommendations shaded in grey and ending  
27 **[2009]** (see 'Update information' box below for details about how recommendations are  
28 labelled). In particular, for recommendations labelled **[2009]**, the word 'consider' may not  
29 necessarily be used to denote the strength of the recommendation.

## 1.3.1 Key Priorities for Implementation

### 1.3.1.2 Patient education

3 Offer structured education to adults with type 2 diabetes and/or their family members or  
4 carers (as appropriate) at and around the time of diagnosis, with annual reinforcement and  
5 review. Explain to people and their carers that structured education is an integral part of  
6 diabetes care. **[2009]**

7 Ensure that any structured education programme for adults with type 2 diabetes includes the  
8 following components<sup>a</sup>:

- 9 • It is evidence-based, and suits the needs of the person.
- 10 • It has specific aims and learning objectives, and supports the person and their family  
11 members and carers in developing attitudes, beliefs, knowledge and skills to self-manage  
12 diabetes.
- 13 • It has a structured curriculum that is theory-driven, evidence-based and resource-  
14 effective, has supporting materials, and is written down.
- 15 • It is delivered by trained educators who have an understanding of educational theory  
16 appropriate to the age and needs of the person, and who are trained and competent to  
17 deliver the principles and content of the programme.
- 18 • It is quality assured, and reviewed by trained, competent, independent assessors who  
19 measure it against criteria that ensure consistency.
- 20 • The outcomes are audited regularly. **[2015]**

Update 2015

### 1.3.2.1 Dietary advice

22 Integrate dietary advice with a personalised diabetes management plan, including other  
23 aspects of lifestyle modification, such as increasing physical activity and losing weight.  
24 **[2009]**

### 1.3.3.5 Blood pressure management

26 Add medications if lifestyle advice does not reduce blood pressure to below 140/80 mmHg  
27 (below 130/80 mmHg if there is kidney, eye or cerebrovascular damage). **[2009]**

28 Monitor blood pressure every 1–2 months, and intensify therapy if the person is already on  
29 antihypertensive drug treatment, until the blood pressure is consistently below 140/80 mmHg  
30 (below 130/80 mmHg if there is kidney, eye or cerebrovascular disease). **[2009]**

### 1.3.4.1 Blood glucose management

#### 1.3.4.1.2 Targets

33 Involve adults with type 2 diabetes in decisions about their individual HbA1c target.  
34 Encourage them to achieve the target and maintain it unless any resulting adverse effects  
35 (including hypoglycaemia), or their efforts to achieve their target, impair their quality of life.  
36 **[new 2015]**

Update 2015

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a [Structured patient education in diabetes: report from the patient education working group](#)

1 If HbA1c levels rise to 58 mmol/mol (7.5%) or higher, intensify drug treatment, set a target  
2 HbA1c level of 53 mmol/mol (7.0%), and reinforce advice about diet, lifestyle and adherence  
3 to drug treatment. See recommendations 8–16. For more information about supporting  
4 adherence, see the NICE guideline on medicines adherence. **[new 2015]**

#### 1.3.4.25 Self-monitoring of blood glucose

6 Do not routinely offer self-monitoring of blood glucose levels for adults with type 2 diabetes  
7 unless the person:

- 8 • is on insulin **or**
- 9 • experiences symptomatic hypoglycaemia **or**
- 10 • is on oral medication that may increase their risk of hypoglycaemia while driving or  
11 operating machinery **or**
- 12 • is pregnant, or is planning to become pregnant. For more information, see the NICE  
13 guideline on diabetes in pregnancy.

14 Consider short-term self-monitoring for adults with type 2 diabetes who start treatment with  
15 oral or intravenous corticosteroids. **[new 2015]**

#### 1.3.56 Drug treatment

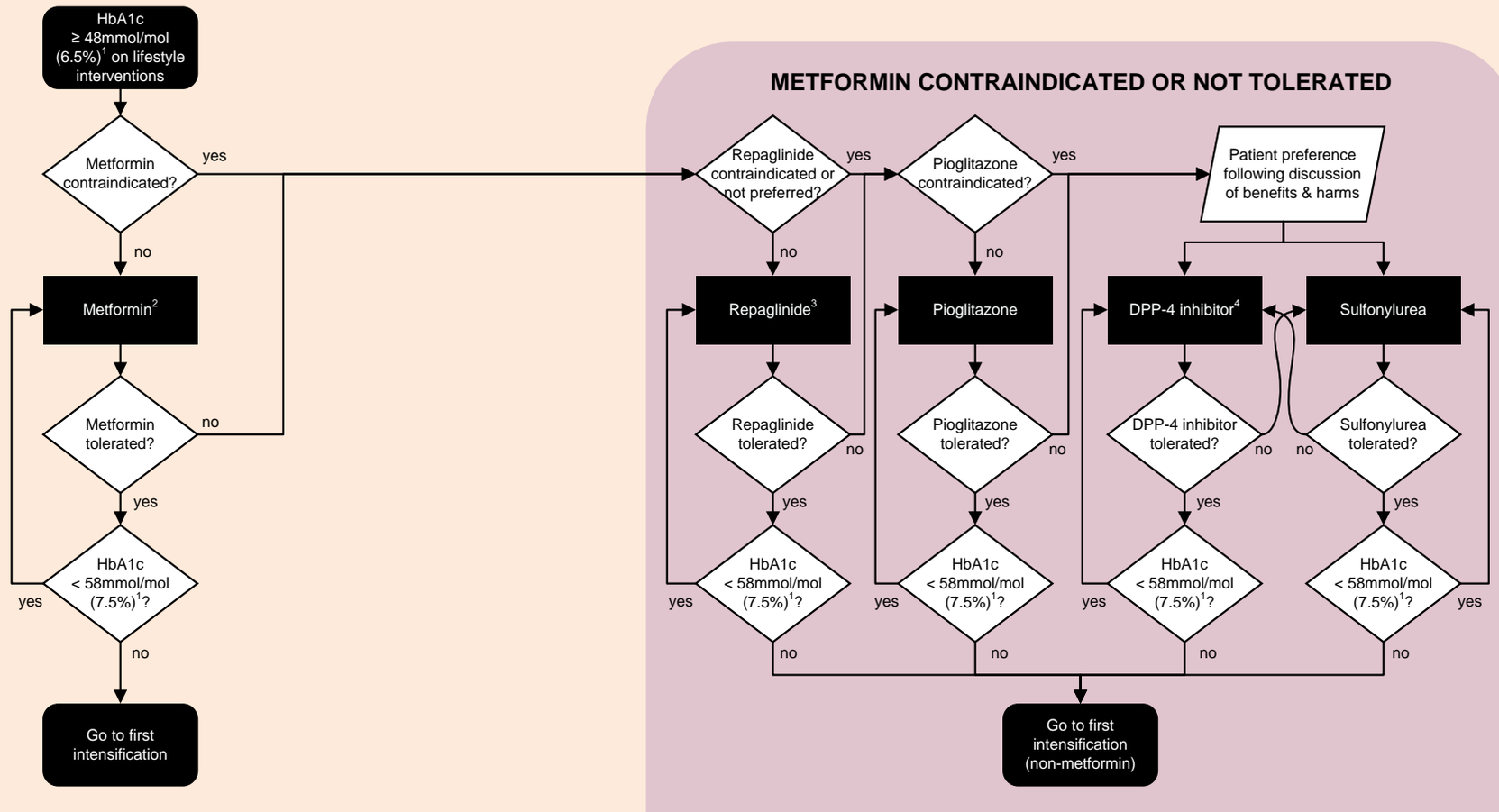
17 Offer standard-release metformin as the initial drug treatment for adults with type 2 diabetes.  
18 **[new 2015]**

19 If standard-release metformin is contraindicated or not tolerated, consider repaglinide as the  
20 initial drug treatment. Advise the person that if treatment with repaglinide does not control  
21 HbA1c, then the person would need to change to pioglitazone, a sulfonylurea or a dipeptidyl  
22 peptidase-4 (DPP-4) inhibitor before adding another treatment<sup>b</sup> (see First intensification of  
23 drug treatment). **[new 2015]**

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b Repaglinide has a marketing authorisation for use only as monotherapy or in combination with metformin. Therefore, for people who cannot take metformin, there is no licensed combination containing repaglinide that can be offered at first intensification. Patients should be made aware of this when initial therapy is being discussed.

### 1.4.1 Algorithm for blood glucose lowering therapy



**Notes**

<sup>1</sup> Or an alternative, individually agreed threshold for starting pharmacological treatment

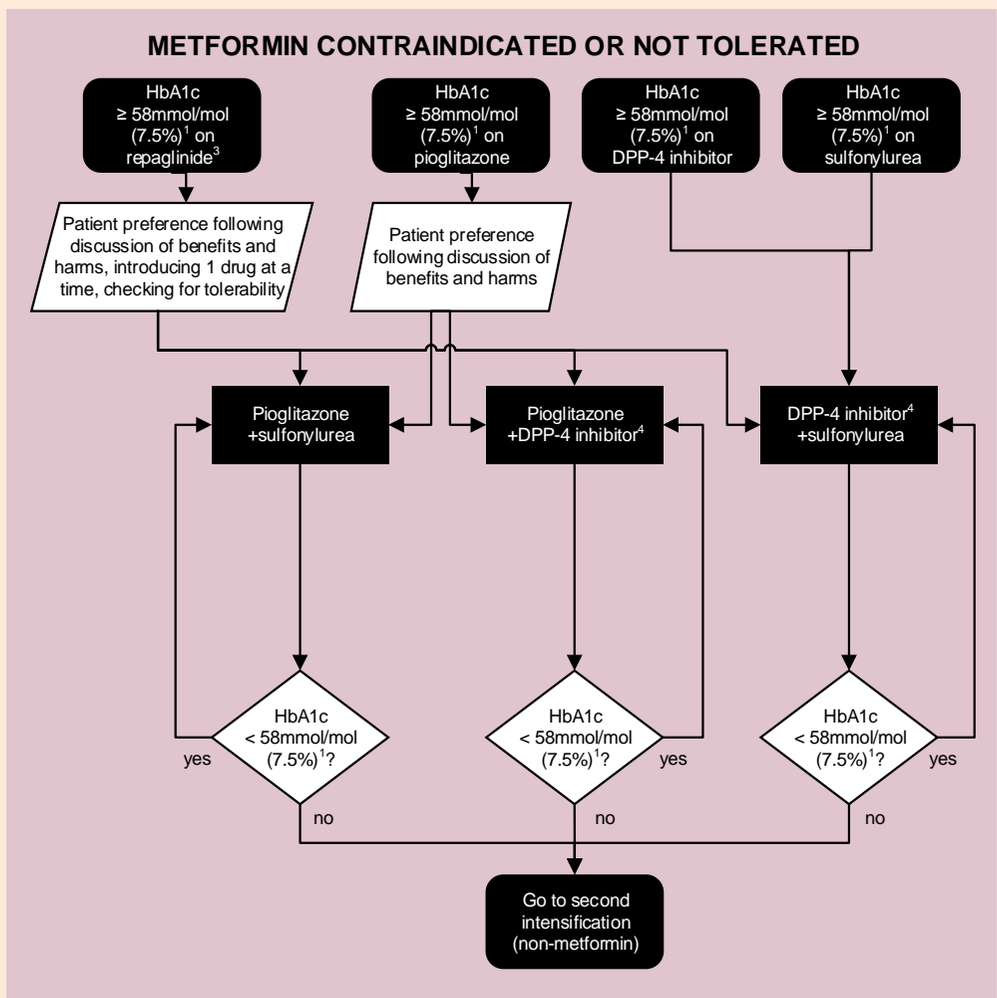
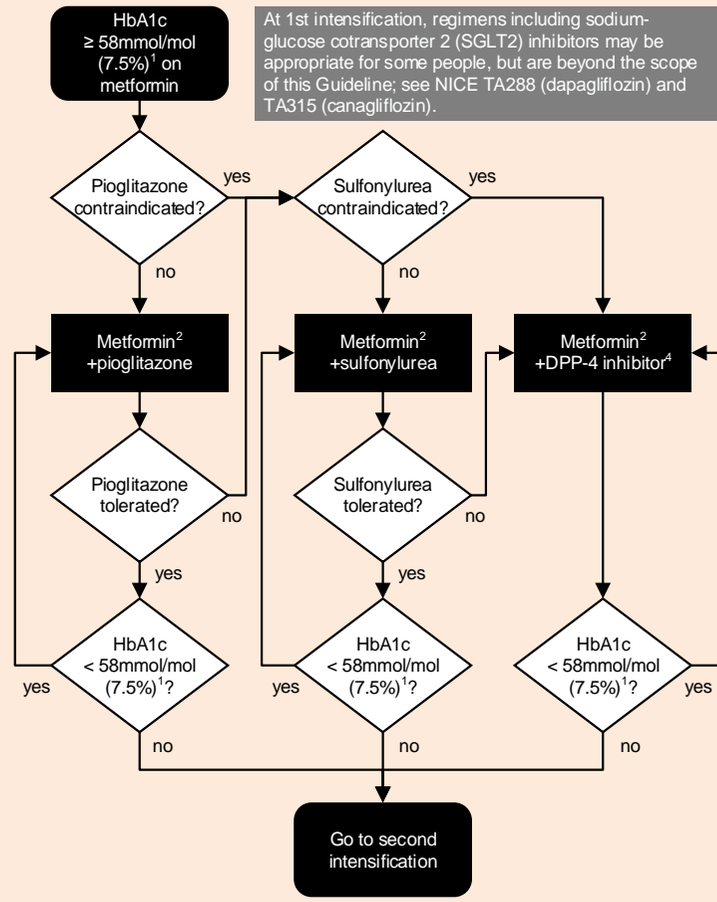
<sup>2</sup> Standard-release metformin

<sup>3</sup> For people who cannot take metformin, there is no licensed combination containing repaglinide that can be offered at first intensification. Patients should be made aware of this when initial therapy is being discussed

<sup>4</sup> Dipeptidyl peptidase-4 (DPP-4) inhibitor; choose the option with the lowest acquisition cost.

2

3 **Figure 1: Pharmacological treatment algorithm – initial therapy**



**Notes**

<sup>1</sup> Or an alternative, individually agreed threshold for intensification

<sup>2</sup> Standard-release metformin

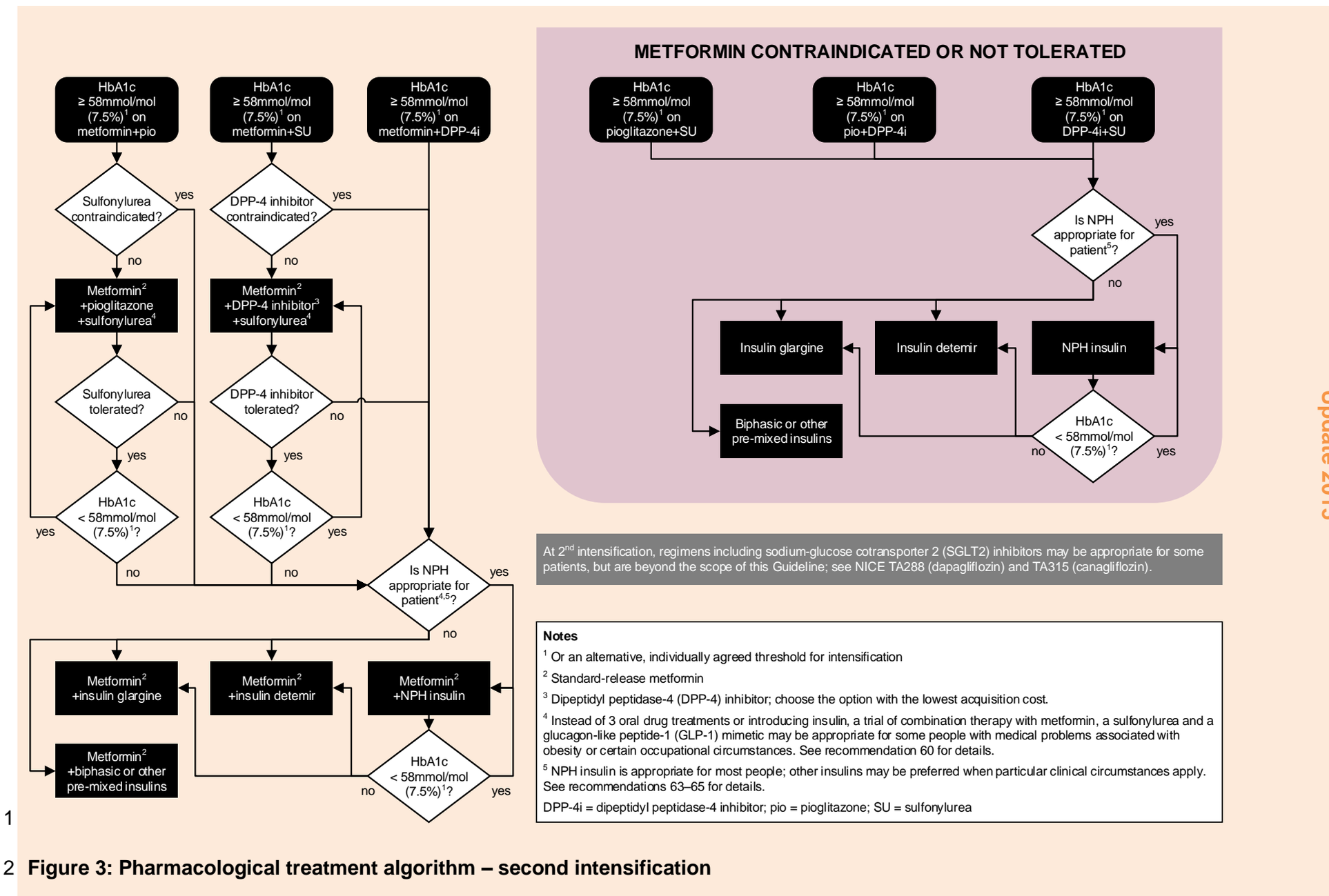
<sup>3</sup> For people who cannot take metformin, there is no licensed combination containing repaglinide that can be offered at first intensification.

<sup>4</sup> Dipeptidyl peptidase-4 (DPP-4) inhibitor; choose the option with the lowest acquisition cost.

Update 2015

1

2 **Figure 2: Pharmacological treatment algorithm – first intensification**



Update 2015

1

2 **Figure 3: Pharmacological treatment algorithm – second intensification**

## 1.5<sup>1</sup> Recommendations

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1. Adopt an individualised approach to diabetes care that is tailored to the person's needs and circumstances, taking into account their personal preferences, comorbidities, risks of polypharmacy, and their ability to benefit from long-term interventions due to reduced life expectancy. Such an approach is especially important in the context of multimorbidity. Reassess the person's needs and circumstances at each review and consider whether to stop any medicines that are not effective. [new 2015]

2. Offer structured education to adults with type 2 diabetes and/or their family members or carers (as appropriate) at and around the time of diagnosis, with annual reinforcement and review. Explain to people and their carers that structured education is an integral part of diabetes care. [2009]

3. Ensure that any structured education programme for adults with type 2 diabetes includes the following components:

- It is evidence-based, and suits the needs of the person.
- It has specific aims and learning objectives, and supports the person and their family members and carers in developing attitudes, beliefs, knowledge and skills to self-manage diabetes.
- It has a structured curriculum that is theory-driven, evidence-based and resource-effective, has supporting materials, and is written down.
- It is delivered by trained educators who have an understanding of educational theory appropriate to the age and needs of the person, and who are trained and competent to deliver the principles and content of the programme.
- It is quality assured, and reviewed by trained, competent, independent assessors who measure it against criteria that ensure consistency.
- The outcomes are audited regularly. [2015]

4. Ensure the patient-education programme provides the necessary resources to support the educators, and that educators are properly trained and given time to develop and maintain their skills. [2009]

5. Offer group education programmes as the preferred option. Provide an alternative of equal standard for a person unable or unwilling to participate in group education. [2009]

6. Ensure that the patient-education programmes available meet the cultural, linguistic, cognitive and literacy needs within the local area. [2009]

7. Ensure that all members of the diabetes healthcare team are familiar with the patient-education programmes available locally, that these programmes are integrated with the rest of the care pathway, and that adults with type 2 diabetes and their family members or carers (as appropriate) have the opportunity to contribute to the design and provision of local programmes. [2009]

8. Provide individualised and ongoing nutritional advice from a healthcare professional with specific expertise and competencies in nutrition. [2009]

- 1 9. Provide dietary advice in a form sensitive to the individual's needs, culture  
2 and beliefs, being sensitive to their willingness to change and the effects  
3 on their quality of life. [2009]
- 4 10. Emphasise advice on healthy balanced eating that is applicable to the  
5 general population when providing advice to adults with type 2 diabetes.  
6 Encourage high-fibre, low-glycaemic-index sources of carbohydrate in the  
7 diet, such as fruit, vegetables, wholegrains and pulses; include low-fat  
8 dairy products and oily fish; and control the intake of foods containing  
9 saturated and trans fatty acids. [2009]
- 10 11. Integrate dietary advice with a personalised diabetes management plan,  
11 including other aspects of lifestyle modification, such as increasing  
12 physical activity and losing weight. [2009]
- 13 12. For adults with type 2 diabetes who are overweight, set an initial body  
14 weight loss target of 5–10%. Remember that lesser degrees of weight  
15 loss may still be of benefit, and that larger degrees of weight loss in the  
16 longer term will have advantageous metabolic impact. [2009]
- 17 13. Individualise recommendations for carbohydrate and alcohol intake, and  
18 meal patterns. Reducing the risk of hypoglycaemia should be a particular  
19 aim for a person using insulin or an insulin secretagogue. [2009]
- 20 14. Advise adults with type 2 diabetes that limited substitution of sucrose-  
21 containing foods for other carbohydrate in the meal plan is allowable, but  
22 that they should take care to avoid excess energy intake. [2009]
- 23 15. Discourage the use of foods marketed specifically for people with  
24 diabetes. [2009]
- 25 16. When adults with type 2 diabetes are admitted to hospital as inpatients or  
26 to any other care setting, implement a meal planning system that provides  
27 consistency in the carbohydrate content of meals and snacks. [2009]
- 28 17. Measure blood pressure at least annually in an adult with type 2 diabetes  
29 without previously diagnosed hypertension or renal disease. Offer and  
30 reinforce preventive lifestyle advice. [2009]
- 31 18. For an adult with type 2 diabetes on antihypertensive drug treatment  
32 when diabetes is diagnosed, review blood pressure control and  
33 medications used. Make changes only if there is poor control or if current  
34 drug treatment is not appropriate because of microvascular complications  
35 or metabolic problems. [2009]
- 36 19. Repeat blood pressure measurements within:
  - 37 • 1 month if blood pressure is higher than 150/90 mmHg
  - 38 • 2 months if blood pressure is higher than 140/80 mmHg
  - 39 • 2 months if blood pressure is higher than 130/80 mmHg and  
40 there is kidney, eye or cerebrovascular damage.
- 41 Provide lifestyle advice (diet and exercise) at the same time. [2009]
- 42 20. Provide lifestyle advice (see section 5.1.6 and the lifestyle interventions  
43 section in 'Hypertension' [NICE clinical guideline 127]) if blood pressure is  
44 confirmed as being consistently above 140/80 mmHg (or above 130/80  
45 mmHg if there is kidney, eye or cerebrovascular damage). [2009]
- 46 21. Add medications if lifestyle advice does not reduce blood pressure to  
47 below 140/80 mmHg (below 130/80 mmHg if there is kidney, eye or  
48 cerebrovascular damage). [2009]

- 1 22. Monitor blood pressure every 1–2 months, and intensify therapy if the  
2 person is already on antihypertensive drug treatment, until the blood  
3 pressure is consistently below 140/80 mmHg (below 130/80 mmHg if  
4 there is kidney, eye or cerebrovascular disease). [2009]
- 5 23. First-line antihypertensive drug treatment should be a once-daily, generic  
6 angiotensin-converting enzyme (ACE) inhibitor. Exceptions to this are  
7 people of African or Caribbean family origin, or women for whom there is  
8 a possibility of becoming pregnant. [2009]
- 9 24. The first-line antihypertensive drug treatment for a person of African or  
10 Caribbean family origin should be an ACE inhibitor plus either a diuretic  
11 or a generic calcium-channel blocker. [2009]
- 12 25. A calcium-channel blocker should be the first-line antihypertensive drug  
13 treatment for a woman for whom, after an informed discussion, it is  
14 agreed there is a possibility of her becoming pregnant. [2009]
- 15 26. For a person with continuing intolerance to an ACE inhibitor (other than  
16 renal deterioration or hyperkalaemia), substitute an angiotensin II-  
17 receptor antagonist for the ACE inhibitor. [2009]
- 18 27. If the person's blood pressure is not reduced to the individually agreed  
19 target with first-line therapy, add a calcium-channel blocker or a diuretic  
20 (usually a thiazide or thiazide-related diuretic). Add the other drug (that is,  
21 the calcium-channel blocker or diuretic) if the target is not reached with  
22 dual therapy. [2009, amended 2015]
- 23 28. If the person's blood pressure is not reduced to the individually agreed  
24 target with triple therapy, add an alpha-blocker, a beta-blocker or a  
25 potassium-sparing diuretic (the last with caution if the individual is already  
26 taking an ACE inhibitor or an angiotensin II-receptor antagonist). [2009]
- 27 29. Monitor the blood pressure of a person who has attained and consistently  
28 remained at his or her blood pressure target every 4–6 months. Check for  
29 possible adverse effects of antihypertensive drug treatment – including  
30 the risks from unnecessarily low blood pressure. [2009]
- 31 30. Do not offer antiplatelet therapy (aspirin or clopidogrel) for adults with type  
32 2 diabetes without cardiovascular disease. [new 2015]
- 33 31. For guidance on the primary and secondary prevention of cardiovascular  
34 disease, see the NICE guidelines on lipid modification and myocardial  
35 infarction – secondary prevention. [new 2015]
- 36 32. Measure HbA1c levels at:
- 37
  - 3–6 monthly intervals (tailored to individual needs), until the  
38 HbA1c is stable on unchanging therapy
  - 6-monthly intervals once the HbA1c level and blood glucose  
40 lowering therapy are stable. [2015]
- 41 33. Calibrate HbA1c results according to International Federation of Clinical  
42 Chemistry (IFCC) standardisation. [new 2015]
- 43 34. If HbA1c monitoring is invalid (because of disturbed erythrocyte turnover  
44 or abnormal haemoglobin type), estimate trends in blood glucose control  
45 using one of the following:
- 46
  - fructosamine estimation
  - quality-controlled plasma glucose profiles
- 47

- 1                                   • total glycated haemoglobin estimation (if abnormal  
2                                   haemoglobins). **[2015]**
- 3           35. Investigate unexplained discrepancies between HbA1c and other glucose  
4           measurements. Seek advice from a team with specialist expertise in  
5           diabetes or clinical biochemistry. **[2015]**
- 6           36. Involve adults with type 2 diabetes in decisions about their individual  
7           HbA1c target. Encourage them to achieve the target and maintain it  
8           unless any resulting adverse effects (including hypoglycaemia), or their  
9           efforts to achieve their target, impair their quality of life. **[new 2015]**
- 10          37. Offer lifestyle advice and drug treatment to help adults with type 2  
11          diabetes achieve and maintain their HbA1c target. See recommendations  
12          8–16. For more information about supporting adherence, see the NICE  
13          guideline on medicines adherence. **[new 2015]**
- 14          38. Set a target HbA1c level of 48 mmol/mol (6.5%) for most adults with type  
15          2 diabetes that is managed either by lifestyle and diet, or by lifestyle and  
16          diet in combination with a single drug that is not associated with  
17          hypoglycaemia. **[new 2015]**
- 18          39. If HbA1c levels rise to 58 mmol/mol (7.5%) or higher, intensify drug  
19          treatment, set a target HbA1c level of 53 mmol/mol (7.0%), and reinforce  
20          advice about diet, lifestyle and adherence to drug treatment. See section  
21          5. For more information about supporting adherence, see the NICE  
22          guideline on medicines adherence. **[new 2015]**
- 23          40. Consider relaxing the target HbA1c level (see recommendations 38–39)  
24          on a case-by-case basis for adults with type 2 diabetes:
- 25                                   • who are unlikely to achieve longer-term risk-reduction benefits  
26                                   (for example, people with a reduced life expectancy)
- 27                                   • for whom tight glycaemic control poses risks
- 28                                   • with a high risk of the consequences of hypoglycaemia (for  
29                                   example, people who are at risk of falling, people who have  
30                                   impaired awareness of hypoglycaemia, and people who drive or  
31                                   operate machinery as part of their job)
- 32                                   • for whom intensive management would not be appropriate (for  
33                                   example, people taking multiple drugs and people with  
34                                   significant comorbidities).
- 35                                   These factors will need particular consideration for people who are  
36                                   older and frail. **[new 2015]**
- 37          41. If adults with type 2 diabetes achieve an HbA1c level that is lower than  
38          their target and they are not experiencing hypoglycaemia, encourage  
39          them to maintain it. **[new 2015]**
- 40          42. For guidance on HbA1c targets for women who are pregnant or planning  
41          to become pregnant, see the NICE guideline on diabetes in pregnancy.  
42          **[new 2015]**
- 43          43. Take the Driver and Vehicle Licensing Agency (DVLA) At a glance guide  
44          to the current medical standards of fitness to drive into account when  
45          offering self-monitoring of blood glucose levels. **[new 2015]**
- 46          44. Do not routinely offer self-monitoring of blood glucose levels for adults  
47          with type 2 diabetes unless the person:
- 48                                   • is on insulin **or**

- 1 • experiences symptomatic hypoglycaemia **or**
- 2 • is on oral medication that may increase their risk of
- 3 hypoglycaemia while driving or operating machinery **or**
- 4 • is pregnant, or is planning to become pregnant. For more
- 5 information, see the NICE guideline on diabetes in pregnancy.
- 6 Consider short-term self-monitoring for adults with type 2 diabetes who
- 7 start treatment with oral or intravenous corticosteroids. **[new**
- 8 **2015]**
- 9 45. If adults with type 2 diabetes are self-monitoring their blood glucose
- 10 levels, carry out a structured assessment at least annually. The
- 11 assessment should include:
  - 12 • the person's self-monitoring skills
  - 13 • the quality and frequency of testing
  - 14 • how the results are used
  - 15 • the impact on the person's quality of life
  - 16 • the continued benefit to the person
  - 17 • the equipment used. **[2015]**
- 18 46. If an adult with type 2 diabetes is symptomatically hyperglycaemic,
- 19 consider insulin (see recommendations 63–65) or a sulfonylurea, and
- 20 review treatment when blood glucose control has been achieved. **[new**
- 21 **2015]**
- 22 47. Offer standard-release metformin as the initial drug treatment for adults
- 23 with type 2 diabetes. **[new 2015]**
- 24 48. Gradually increase the dose of standard-release metformin over several
- 25 weeks to minimise the risk of gastrointestinal side effects. **[new 2015]**
- 26 49. Review the dose of standard-release metformin if the estimated
- 27 glomerular filtration rate (eGFR) is below 45 ml/minute/1.73m<sup>2</sup>:
  - 28 • Stop standard-release metformin if the eGFR is below
  - 29 30 ml/minute/1.73m<sup>2</sup>.
  - 30 • Prescribe standard-release metformin with caution for those at
  - 31 risk of a sudden deterioration in kidney function and those at risk
  - 32 of eGFR falling below 45 ml/minute/1.73m<sup>2</sup>. **[2015]**
- 33 50. If standard-release metformin is contraindicated or not tolerated, consider
- 34 repaglinide as the initial drug treatment. Advise the person that if
- 35 treatment with repaglinide does not control HbA1c, then the person would
- 36 need to change to pioglitazone, a sulfonylurea or a dipeptidyl peptidase-4
- 37 (DPP-4) inhibitor before adding another treatment (see First
- 38 intensification of drug treatment). **[new 2015]**
- 39 51. If both standard-release metformin and repaglinide are contraindicated or
- 40 not tolerated, or if repaglinide is not the preferred option, consider
- 41 pioglitazone as the initial drug treatment. **[new 2015]**
- 42 52. If both standard-release metformin and pioglitazone are contraindicated or
- 43 not tolerated, and repaglinide is contraindicated, not tolerated or not
- 44 preferred, consider initial drug treatment with either:
  - 45 • a DPP-4 inhibitor **or**
  - 46 • a sulfonylurea.

- 1 Base the choice on the person's preference after discussing the risks  
2 and benefits of each option. If a DPP-4 inhibitor is preferred,  
3 choose the option with the lowest acquisition cost. **[new 2015]**
- 4 53. If initial drug treatment with standard-release metformin has not controlled  
5 HbA1c to below the person's individually agreed threshold for  
6 intensification:
- 7 • Offer combination therapy with standard-release metformin and  
8 pioglitazone.
  - 9 • If pioglitazone is contraindicated or not tolerated, offer  
10 combination therapy with standard-release metformin and a  
11 sulfonylurea.
  - 12 • If both pioglitazone and sulfonylureas are contraindicated or not  
13 tolerated, offer combination therapy with standard-release  
14 metformin and a DPP-4 inhibitor (choose the option with the  
15 lowest acquisition cost). **[new 2015]**
- 16 54. If initial drug treatment with repaglinide has not controlled HbA1c to below  
17 the person's individually agreed threshold for intensification:
- 18 • Consider switching to combination therapy with either:
    - 19 ○ pioglitazone and a sulfonylurea **or**
    - 20 ○ pioglitazone and a DPP-4 inhibitor **or**
    - 21 ○ a sulfonylurea and a DPP-4 inhibitor.
- 22 Base the choice on the person's preference after discussing the risks  
23 and benefits of each combination. If a DPP-4 inhibitor is  
24 preferred, choose the option with the lowest acquisition cost.
- 25 • When switching from repaglinide to any of these combinations,  
26 introduce the 2 new medicines in a stepwise manner, checking  
27 for tolerability of each. **[new 2015]**
- 28 55. If initial drug treatment with pioglitazone has not controlled HbA1c to  
29 below the person's individually agreed threshold for intensification:
- 30 • Consider combination therapy with either:
    - 31 ○ pioglitazone and a sulfonylurea **or**
    - 32 ○ pioglitazone and a DPP-4 inhibitor.
- 33 Base the choice on the person's preference after discussing the risks  
34 and benefits of each option. If a DPP-4 inhibitor is preferred,  
35 choose the option with the lowest acquisition cost. **[new 2015]**
- 36 56. If initial drug treatment with a DPP-4 inhibitor has not controlled HbA1c to  
37 below the person's individually agreed threshold for intensification,  
38 consider combination therapy with the DPP-4 inhibitor and a sulfonylurea.  
39 **[new 2015]**
- 40 57. If initial drug treatment with a sulfonylurea has not controlled HbA1c to  
41 below the person's individually agreed threshold for intensification,  
42 consider combination therapy with the sulfonylurea and a DPP-4 inhibitor  
43 (choose the option with the lowest acquisition cost). **[new 2015]**
- 44 58. If combination therapy with either standard-release metformin and  
45 pioglitazone or standard-release metformin and a sulfonylurea has not

- 1 controlled HbA1c to below the person's individually agreed threshold for  
2 intensification:
- 3 • Consider combination therapy with standard-release metformin,  
4 pioglitazone and a sulfonylurea.
  - 5 • If pioglitazone is contraindicated or not tolerated, consider either:
    - 6 ○ combination therapy with standard-release metformin, a  
7 sulfonylurea and a DPP-4 inhibitor **or**
    - 8 ○ starting insulin-based treatment (see recommendations 63–65).
- 9 Base the choice on the person's preference after discussing the risks  
10 and benefits of each approach. If a DPP-4 inhibitor is preferred,  
11 choose the option with the lowest acquisition cost.
- 12 • If sulfonylureas are contraindicated or not tolerated, consider  
13 starting insulin-based treatment (see recommendations 63–65).  
14 **[new 2015]**
- 15 59. If standard-release metformin is contraindicated or not tolerated, and if  
16 combination therapy with 2 oral drug treatments has not controlled HbA1c  
17 to below the person's individually agreed threshold for intensification,  
18 consider starting insulin-based treatment (see recommendations 63–65).  
19 **[new 2015]**
- 20 60. If combination therapy with 2 oral drug treatments has not controlled  
21 HbA1c to below the person's individually agreed threshold for  
22 intensification, consider combination therapy with standard-release  
23 metformin, a sulfonylurea and a glucagon-like peptide-1 (GLP-1) mimetic  
24 (instead of 3 oral drug treatments or introducing insulin).
- 25 Consider this for adults with type 2 diabetes who:
- 26 • have a BMI of 35 kg/m<sup>2</sup> or higher **and** specific psychological or  
27 other medical problems associated with obesity, **or**
  - 28 • have a BMI lower than 35 kg/m<sup>2</sup> **and**
    - 29 ○ for whom insulin therapy would have significant occupational  
30 implications, **or**
    - 31 ○ weight loss would benefit other significant obesity-related  
32 comorbidities.
- 33 Base the choice of GLP-1 mimetic on the person's preference after  
34 discussing the risks and benefits of each licensed option. If more  
35 than 1 option is considered appropriate for the person, choose  
36 the GLP-1 mimetic with the lowest acquisition cost. **[new 2015]**
- 37 61. Only continue GLP-1 mimetic therapy if the person has had a beneficial  
38 metabolic response (a reduction of at least 11 mmol/mol [1%] in HbA1c  
39 and a weight loss of at least 3% of initial body weight in 6 months). **[2015]**
- 40 62. Only offer a GLP-1 mimetic in combination with insulin in a specialist care  
41 setting. **[new 2015]**
- 42 63. When starting insulin therapy, use a structured programme employing  
43 active insulin dose titration that encompasses:
- 44 • structured education
  - 45 • continuing telephone support
  - 46 • frequent self-monitoring

- 1 • dose titration to target
- 2 • dietary understanding
- 3 • management of hypoglycaemia
- 4 • management of acute changes in plasma glucose control
- 5 • support from an appropriately trained and experienced
- 6 healthcare professional. **[2015]**
- 7 64. When starting insulin therapy, continue to offer standard-release
- 8 metformin for people without contraindications or intolerance. [new 2015]
- 9 65. Initiate insulin therapy from a choice of a number of insulin types and
- 10 regimens:
- 11 • Offer human neutral protamine Hagedorn (NPH) insulin injected
- 12 at bed-time or twice daily according to need.
- 13 • Consider, as an alternative, using a long-acting insulin analogue
- 14 (insulin detemir, insulin glargine) if:
- 15 ○ the person needs assistance from a carer or healthcare
- 16 professional to inject insulin, and use of a long-acting insulin
- 17 analogue (insulin detemir, insulin glargine) would reduce the
- 18 frequency of injections from twice to once daily, **or**
- 19 ○ the person's lifestyle is restricted by recurrent symptomatic
- 20 hypoglycaemic episodes, **or**
- 21 ○ the person would otherwise need twice-daily NPH insulin
- 22 injections in combination with oral glucose-lowering drugs, **or**
- 23 ○ the person cannot use the device to inject NPH insulin.
- 24 • Consider twice-daily pre-mixed (biphasic) human insulin
- 25 (particularly if HbA1c is 75 mmol/mol [9.0%] or higher). A once-
- 26 daily regimen may be an option.
- 27 • Consider pre-mixed preparations that include short-acting insulin
- 28 analogues, rather than pre-mixed preparations that include
- 29 short-acting human insulin preparations, if:
- 30 ○ a person prefers injecting insulin immediately before a meal, **or**
- 31 ○ hypoglycaemia is a problem, **or**
- 32 ○ blood glucose levels rise markedly after meals. **[2015]**
- 33 66. Consider switching to a long-acting insulin analogue (insulin detemir,
- 34 insulin glargine) from NPH insulin in people:
- 35 • who do not reach their target HbA1c because of significant
- 36 hypoglycaemia, **or**
- 37 • who experience significant hypoglycaemia on NPH insulin
- 38 irrespective of the level of HbA1c reached, **or**
- 39 • who cannot use the device needed to inject NPH insulin but who
- 40 could administer their own insulin safely and accurately if a
- 41 switch to a long-acting insulin analogue were made, **or**
- 42 • who need help from a carer or healthcare professional to
- 43 administer insulin injections and for whom switching to a long
- 44 acting insulin analogue would reduce the number of daily
- 45 injections. **[2015]**

- 1 67. Monitor a person on a basal insulin regimen (NPH insulin or a long acting  
2 insulin analogue [insulin detemir, insulin glargine]) for the need for short-  
3 acting insulin before meals (or a pre-mixed insulin preparation). [2015]
- 4 68. Monitor a person on pre-mixed insulin for the need for a further injection  
5 of short-acting insulin before meals or for a change to a basal bolus  
6 regimen with NPH insulin or long-acting insulin analogues (insulin  
7 detemir, insulin glargine), if blood glucose control remains inadequate.  
8 [2015]
- 9 69. For guidance on insulin delivery, see the NICE guideline on type 1  
10 diabetes. [new 2015]
- 11 70. **Think about a** diagnosis of gastroparesis in adults with type 2 diabetes  
12 with erratic blood glucose control or unexplained gastric bloating or  
13 vomiting, taking into account possible alternative diagnoses. [2009,  
14 amended 2015]
- 15 71. Consider a trial of **metoclopramide, domperidone or erythromycin** for an  
16 adult with type 2 diabetes with **gastroparesis**. [2009, amended 2015]
- 17 72. If gastroparesis is suspected, consider referral to specialist services if:  
18
  - the differential diagnosis is in doubt **or**
  - persistent or severe vomiting occurs. **[2009]**
- 19
- 20 73. For guidance on painful neuropathy in adults with type 2 diabetes, see the  
21 NICE guideline on neuropathic pain – pharmacological management.  
22 [new 2015]
- 23 74. **Think about** the possibility of contributory sympathetic nervous system  
24 damage for adults with type 2 diabetes who lose the warning signs of  
25 hypoglycaemia. [2009, amended 2015]
- 26 75. **Think about** the possibility of autonomic neuropathy affecting the gut in  
27 adults with type 2 diabetes who have unexplained diarrhoea that happens  
28 particularly at night. [2009, amended 2015]
- 29 76. When using tricyclic drugs and antihypertensive drug treatments in adults  
30 with type 2 diabetes who have autonomic neuropathy, be aware of the  
31 increased likelihood of side effects such as orthostatic hypotension.  
32 [2009]
- 33 77. Investigate the possibility of autonomic neuropathy affecting the bladder in  
34 adults with type 2 diabetes who have unexplained bladder-emptying  
35 problems. [2009]
- 36 78. In managing autonomic neuropathy symptoms, include specific  
37 interventions indicated by the manifestations (for example, for abnormal  
38 sweating or nocturnal diarrhoea). [2009]
- 39 79. For guidance on managing foot problems in adults with type 2 diabetes,  
40 see the NICE guideline on diabetic foot problems. [new 2015]
- 41 80. For guidance on nephropathy in adults with type 2 diabetes, see the NICE  
42 guideline on chronic kidney disease. [new 2015]
- 43 81. Offer men with type 2 diabetes the opportunity to discuss erectile  
44 dysfunction as part of their annual review. [2015]
- 45 82. Carry out an assessment, and provide education and support for men with  
46 type 2 diabetes who have problematic erectile dysfunction, addressing  
47 contributory factors such as cardiovascular disease as well as possible  
48 treatment options. [2015]

- 1 83. Consider a phosphodiesterase-5 inhibitor to treat problematic erectile  
2 dysfunction, initially choosing the drug with the lowest acquisition cost  
3 and taking into account any contraindications. [new 2015]
- 4 84. Following discussion, refer men with type 2 diabetes to a service offering  
5 other medical, surgical, or psychological management of erectile  
6 dysfunction if treatment (including a phosphodiesterase-5 inhibitor, if  
7 appropriate) has been unsuccessful. [2015]
- 8 85. Arrange or perform eye screening at or around the time of diagnosis.  
9 Arrange repeat of structured eye screening annually. [2009]
- 10 86. Explain the reasons for, and success of, eye screening systems to adults  
11 with type 2 diabetes, so that attendance is not reduced by lack of  
12 knowledge or fear of outcome. [2009]
- 13 87. Use mydriasis with tropicamide when photographing the retina, after prior  
14 informed agreement following discussion of the advantages and  
15 disadvantages. Discussions should include precautions for driving. [2009]
- 16 88. Use a quality-assured digital retinal photography programme using  
17 appropriately trained staff. [2009]
- 18 89. Perform visual acuity testing as a routine part of eye screening  
19 programmes. [2009]
- 20 90. Depending on the findings, follow structured eye screening by:
- 21     • routine review in 1 year **or**
- 22     • earlier review **or**
- 23     • referral to an ophthalmologist. **[2009]**
- 24 91. Arrange emergency review by an ophthalmologist for:
- 25     • sudden loss of vision
- 26     • rubeosis iridis
- 27     • pre-retinal or vitreous haemorrhage
- 28     • retinal detachment. **[2009]**
- 29 92. Arrange rapid review by an ophthalmologist for new vessel formation.  
30 [2009]
- 31 93. Refer to an ophthalmologist in accordance with the National Screening  
32 Committee criteria and timelines if any of these features are present:
- 33     • referable maculopathy:
- 34         o exudate or retinal thickening within 1 disc diameter of the centre  
35             of the fovea
- 36         o circinate or group of exudates within the macula (the macula is  
37             defined here as a circle centred on the fovea, with a diameter  
38             the distance between the temporal border of the optic disc and  
39             the fovea)
- 40         o any microaneurysm or haemorrhage within 1 disc diameter of the  
41             centre of the fovea, only if associated with deterioration of best  
42             visual acuity to 6/12 or worse
- 43     • referable pre-proliferative retinopathy (if cotton wool spots are  
44             present, look carefully for the following features, but cotton wool  
45             spots themselves do not define pre-proliferative retinopathy):

- 1           o    any venous beading
- 2           o    any venous reduplication
- 3           o    any intraretinal microvascular abnormalities
- 4           o    multiple deep, round or blot haemorrhages
- 5           •    any large sudden unexplained drop in visual acuity. **[2009,**
- 6                    **amended 2015]**
- 7

## 1.6.1 Research recommendations

2 The Guideline Development Group has made the following recommendations for research,  
3 based on its review of evidence, to improve NICE guidance and patient care in the future.

4

5 1. What is the effectiveness of low carbohydrate diets in adults with type 2  
6 diabetes?

7

Why this is important

8

9 Type 2 diabetes is associated with obesity, and lifestyle interventions including  
10 diet and physical activity are thought to be useful in helping to control the  
11 condition and improve patient outcomes such as reducing the risk of long-  
12 term complications and increasing quality of life. Low carbohydrate diets  
13 have been a source of discussion over the past two decades and there is  
14 much debate regarding its effectiveness and safety in controlling blood  
15 glucose levels, particularly in the longer-term. Specifically, there is little  
16 consensus on the optimal intake of daily carbohydrates, where the risk of  
17 adverse effects such as hypoglycaemia is minimised. A double-blind  
18 randomised controlled trial addressing this clinical question would help to  
19 provide a better understanding of the effects of low carbohydrate diets on  
20 diabetes control and maintenance to inform appropriate management  
strategies.

21

22 2. What is the natural history of individuals who are diagnosed with type 2  
23 diabetes in childhood in terms of long-term complications/consequences  
in adulthood?

24

Why this is important

25

26 Type 2 diabetes has historically been associated with adults, with research  
27 largely focused on this population. However, there is growing concern of  
28 the increasing incidence of type 2 diabetes in younger people, thought to  
29 be linked to the rising levels of obesity. In order to improve clinical  
30 management of people diagnosed in childhood, a better understanding of  
31 the early progression of the condition is needed, particularly in terms of its  
32 effects on the long-term risks of developing microvascular and  
33 macrovascular complications. A prospective longitudinal 10 year cohort  
34 study of children diagnosed with type 2 diabetes would help improve  
35 understanding of whether diabetes spanning the growth spurt would  
36 result in long-term complications occurring at a different rate compared to  
individuals who are diagnosed during adulthood.

37

38 3. What is the optimal frequency for self-monitoring of blood glucose in  
adults with type 2 diabetes?

39

40 4. What are the optimal blood glucose targets for self-monitoring in adults  
with type 2 diabetes?

41

Why this is important

42

43 It is widely recognised that self-monitoring of blood glucose is a  
44 multicomponent intervention. As well as being educated about how to use  
45 a self-monitoring device to assess blood glucose levels, adults with type 2  
46 diabetes need to be able to understand their results and act on the  
47 observed readings. In adults for whom self-monitoring is appropriate,  
48 there is limited evidence to guide clinical practice in prescribing self-  
monitoring regimens, in terms of frequency of testing and optimal

glycaemic targets. Given the inconvenience and expense of self-monitoring, robust evidence from double-blind randomised controlled trials is needed to guide the optimal use of this intervention.

5. In adults with type 2 diabetes, what treatment combinations (for example, glucagon-like peptide-1 [GLP-1] mimetics and insulin) are most effective when initial drug treatment with non-metformin monotherapy fails to adequately control blood glucose levels?

Why this is important

Although it is recognised that metformin therapy is suitable for most adults with type 2 diabetes, its use is contraindicated or not tolerated in approximately 15% of individuals. To date, research evidence has largely focused on metformin-based treatment combinations. Given the progressive nature of the condition, in which intensification of blood glucose lowering drug therapies are indicated over time, there is little evidence, for some adults, to guide management strategies on treatment combinations that do not include metformin. Double-blind randomised controlled trials are therefore needed to better understand the treatment choices that are available which improve glycaemic control and long-term risks of complications associated with diabetes.

6. In adults with type 2 diabetes, what are the effects of early use of insulin and glucagon-like peptide-1 (GLP-1) mimetics?

Why this is important

Poor glycaemic control is associated with increased risk of vascular complications. Glucagon-like peptide-1 (GLP-1) mimetics are a new class of blood glucose lowering drugs that target the incretin system, regulating insulin and glucagon. It is associated with low rates of hypoglycaemia and some weight loss. Its effectiveness and safety in combination with insulin early on in the drug treatment pathway is unknown. Double-blind randomised controlled trials are needed to understand the short and long-term effects of early use of GLP-1 agonists with insulin in terms of glycaemic control, adverse effects, diabetes-related complications and mortality. Research on its use could have a significant impact on the management of adults with type 2 diabetes.

7. When third intensification of treatment is indicated, which blood glucose lowering therapies should be used to control blood glucose levels?

Why this is important

As the incidence of type 2 diabetes increases in the younger population and as glycaemic control declines naturally over time, it is likely that further intensification of therapies would be needed. Currently, there is evidence up to second intensification of drug therapies, that is, when 2 or more non-insulin based treatment combinations fail to adequately control blood glucose levels. Double-blind randomised controlled trials are needed to improve understanding of alternative treatment options for adults at second intensification who are inadequately controlled with insulin and/or triple non-insulin based drug therapies.

8. In adults with type 2 diabetes, what are the effects of stopping and/or switching drug treatments to control blood glucose levels, and what criteria should inform the decision?

Why this is important

1 There is a lack of evidence on the effects of stopping and/or switching drug  
2 treatments to control blood glucose levels. The current practice of  
3 'stopping rules' is typically motivated by either inadequate blood glucose  
4 control (rising HbA1c levels) or intolerable side effects. There is limited  
5 understanding of the short- and long-term effects of stopping a therapy  
6 and switching to another in terms of diabetes control (HbA1c levels),  
7 hypoglycaemic risk, weight gain, and cardiovascular morbidity and  
8 mortality. In addition, there is limited understanding of how quickly  
9 consideration should be given to stopping and switching to another drug  
10 treatment and, if stopping and switching may be needed, what the optimal  
11 sequencing is of drug treatments. Double-blind randomised controlled  
12 trials examining these different issues would help to improve diabetes  
13 care.

14 9. In adults with type 2 diabetes, what are the long-term effects of blood  
15 glucose lowering therapies such as dipeptidyl peptidase-4 (DPP-4)  
16 inhibitors, sodium–glucose co-transporter 2 (SGLT2) inhibitors and  
17 meglitinides?

18 Why this is important

19 There is limited evidence in relation to adverse events (for example,  
20 cardiovascular outcomes) on the long-term effects (at least 5 years) of  
21 blood glucose lowering therapies, particularly newer agents. Prospective  
22 longitudinal studies are needed to better understand the long-term safety  
23 issues surrounding these medicines.

24 10. In adults with type 2 diabetes, what patient characteristics predict  
25 response or non-response to pharmacological blood glucose lowering  
26 therapies?

27 Why this is important

28 There is little understanding of the prognostic characteristics that determine  
29 the likelihood that a person would benefit and respond or not respond to  
30 treatment. Increased understanding of important predictive criteria would  
31 better help clinicians target drug therapies and improve overall patient  
32 care. Prospective longitudinal cohort studies examining various types of  
33 prognostic factors such as demographic, disease-specific and comorbid  
34 are needed to identify characteristics that are likely to predict treatment  
35 response or non-response to blood glucose lowering therapies in adults  
36 with type 2 diabetes.

37 11. In adults with type 2 diabetes and multimorbidity, what are the optimal  
38 blood glucose lowering treatment strategies?

39 Why this is important

40 The evidence reviewed in this guideline commonly excluded participants with  
41 type 2 diabetes whose disease is complicated by significant coexisting  
42 conditions, although this is a common presentation in real-world practice.  
43 As a result, it is difficult to account for the impact of different comorbid  
44 conditions on the effectiveness of blood glucose lowering treatment  
45 strategies. A systematic review is needed to ascertain the optimal  
46 treatment strategies for glycaemic control in adults with type 2 diabetes  
47 and a range of comorbid conditions. Multimorbidity covers a wide range of  
48 conditions (for example, heart failure, chronic obstructive pulmonary  
49 disease and depression) and each would have different implications.  
50 Therefore, analyses should consider whether the optimal treatment  
51 strategies differ according to specific comorbid conditions.

- 1 12. What is the optimal dosing of different phosphodiesterase-5 (PDE-5)  
2 inhibitors for people with type 2 diabetes and erectile dysfunction?  
3 Why this is important  
4 Although phosphodiesterase-5 (PDE-5) inhibitors have been shown to be  
5 effective compared to placebo in improving erectile function in men with  
6 type 2 diabetes, there is little understanding of the optimal dosing  
7 strategies for the different drugs available in this class. Double-blind  
8 randomised controlled trials in this area could help inform clinical practice.
- 9 13. What is the effectiveness of pharmacological treatment strategies for  
10 people with type 2 diabetes and erectile dysfunction who do not respond  
11 to phosphodiesterase-5 (PDE-5) inhibitors, for example PDE-5 inhibitor  
12 plus prostaglandins?  
13 Why this is important  
14 There is limited understanding of alternative treatment strategies available to  
15 men who do not respond to phosphodiesterase-5 (PDE-5) inhibitors.  
16 Double-blind randomised controlled trials of combination therapies and  
17 other pharmacological treatments could help inform clinical practice.
- 18 14. What is the effectiveness of treatment strategies (pharmacological and  
19 non-pharmacological) for sexual dysfunction related to type 2 diabetes in  
20 women?  
21 Why this is important  
22 Sexual dysfunction affect women with type 2 diabetes and there is limited  
23 understanding of available effective treatment strategies. A systematic  
24 review is needed examining the clinical and cost-effectiveness of  
25 available treatment strategies for women with type 2 diabetes and sexual  
26 dysfunction.
- 27 15. What is the effectiveness of treatment strategies (pharmacological and  
28 non-pharmacological) for sexual dysfunction in adults with type 2 diabetes  
29 in same-sex relationships?  
30 Why this is important  
31 Sexual dysfunction in adults with type 2 diabetes in same-sex relationships is  
32 an important area, where there is a limited understanding about effective  
33 treatment strategies. A systematic review is needed examining the clinical  
34 and cost-effectiveness of available treatment strategies for adults with  
35 type 2 diabetes and sexual dysfunction in same-sex relationships.  
36

## 2<sub>1</sub> Overview

### 2.1.2 Introduction

3 Diabetes is a group of disorders with a number of common features, of which raised blood  
4 glucose (hyperglycaemia), by definition is the most evident. In England and Wales, the four  
5 commonest types of diabetes are:

6 • Type 1 diabetes

7 • Type 2 diabetes

8 • Secondary diabetes (from pancreatic damage, hepatic cirrhosis, endocrinological  
9 disease/therapy, or anti-viral/anti-psychotic therapy)

10 • Gestational diabetes (diabetes in pregnancy).

11 This guideline focuses on the management of type 2 diabetes in adults (18 years and over).

12 The World Health Organization's (WHO) definition of diabetes updated in 2011, was used in  
13 this guideline (International Diabetes Federation 2006). Although, no specific definition for  
14 type 2 diabetes is provided, the general definition refers to a state of high blood glucose  
15 levels that is sufficient to put the individual at risk of specific microvascular complications  
16 associated with the condition. In 2009, the WHO recommended that a glycated haemoglobin  
17 (HbA1c) threshold of 48 mmol/mol (6.5%) be used to diagnose diabetes. A person is  
18 normally thought to have type 2 diabetes if he or she does not have type 1 diabetes  
19 (characterised by a rapid onset, often in childhood, insulin-dependence, and ketoacidosis if  
20 neglected), monogenetic diabetes or other medical conditions or treatment suggestive of  
21 secondary diabetes. Diagnosis is not addressed in this guideline.

22 The underlying disorder of type 2 diabetes is usually that of a background of insulin  
23 insensitivity where the body is unable to respond to normal levels of insulin, and insulin  
24 deficiency where the pancreas is unable to secrete enough insulin to compensate for this  
25 resistance. Insulin insensitivity is usually evidenced by excess body weight or obesity, and is  
26 exacerbated by overeating and inactivity. It is commonly associated with raised blood  
27 pressure, a disturbance of blood lipid levels, and a tendency to develop thrombosis. This  
28 combination is often recognised as 'metabolic syndrome', and is associated with fatty liver  
29 and abdominal adiposity (increased waist circumference). Insulin deficiency is progressive  
30 over time, such that the high glucose levels usually worsen relentlessly over a period of  
31 years, requiring continued escalation of blood glucose lowering therapy.

32 Type 2 diabetes is associated with long-term complications, reduced quality of life and life  
33 expectancy. The UK Prospective Diabetes Study (UKPDS) found that approximately 50% of  
34 people newly diagnosed with type 2 diabetes already have complications. Type 2 diabetes is  
35 notable for the increased cardiovascular risk that it carries: coronary artery disease (leading  
36 to heart attacks, angina); peripheral artery disease (leg claudication, gangrene); and carotid  
37 artery disease (strokes, dementia). In addition, prolonged hyperglycaemia can lead to  
38 irreversible microvascular complications such as diabetic retinopathy, nephropathy and  
39 neuropathy (resulting in amputation, painful symptoms, erectile dysfunction and other  
40 problems).

41 Multiple vascular risk factors and wide-ranging complications make diabetes care complex  
42 and time-consuming, and many areas of healthcare services must be involved for optimal  
43 management. Necessary lifestyle changes, the complexities and possible side effects of  
44 therapy make patient education and self-management important aspects of diabetes care.

Update 2015

Update 2015

## 2.2.1 Prevalence

2 It is estimated that approximately 90% of adults currently diagnosed with diabetes have type  
3 2 diabetes. In 2013, almost 2.9 million people were diagnosed with diabetes, with prevalence  
4 rates of 6% and 6.7% in England and Wales respectively. In the UK, incidence rates are  
5 increasing, with more than 1 in 20 people estimated to have diagnosed or undiagnosed  
6 diabetes. While type 2 diabetes mainly develops in people aged over 40 years, it is usually  
7 diagnosed earlier in people of South Asian, Chinese, African or African Caribbean family  
8 origin. It can occur in all age groups and is increasingly being diagnosed in children. People  
9 who are overweight or obese, have inactive lifestyles or have a family history of diabetes are  
10 at risk. It is also more common in the less-affluent.

## 2.3.1 Health and resource burden

12 Type 2 diabetes can result in a wide range of complications with repercussions for both the  
13 individual and the NHS. The economic impact of this condition includes at least 3 factors:

- 14 • direct cost to the NHS and associated healthcare support services
- 15 • indirect cost to the economy, including the effects of early mortality and lost productivity
- 16 • personal impact of diabetes and subsequent complications on individuals and their  
17 families.

18 It is estimated that diabetes account for approximately 15 to 16% of deaths in England, with  
19 life expectancy for people with type 2 diabetes reduced by an average of up to 10 years.  
20 Diabetes care is estimated to account for at least 5% of UK healthcare expenditure, and up  
21 to 10% of NHS expenditure. The presence of diabetic complications can lead to a 5-fold  
22 increase in a patient's NHS costs and people with diabetes can experience prolonged stays  
23 in hospital.

24 This guideline contains recommendations for managing type 2 diabetes in adults and  
25 focuses on patient education, dietary advice, managing cardiovascular risk, managing blood  
26 glucose levels, and identifying and managing long-term complications. The guideline does  
27 not cover diagnosis, type 1 diabetes, diabetes in pregnancy and diabetes in children.

## 2.4.8 Reasons for the update

29 Since publication of the 2009 guideline, availability of new evidence and several key  
30 developments have prompted an update in the following areas: managing blood glucose  
31 levels, antiplatelet therapy and erectile dysfunction. In particular, reasons included safety  
32 concerns surrounding some blood glucose lowering medicines, new evidence on novel  
33 dipeptidyl peptidase-4 (DPP-4) inhibitors and glucagon-like peptide-1 (GLP-1) receptor  
34 agonists, new indications and combinations for licensed class members and the potential  
35 impact of drugs coming off patent on health-economic issues. In addition, new evidence and  
36 safety issues relating to the off-label use of antiplatelet therapy (aspirin and clopidogrel) in  
37 the primary prevention of cardiovascular disease motivated an update of this review.

## 2.5.8 Patient-centred care

39 This guideline offers best practice advice on the care of adults with type 2 diabetes.

40 When caring for older adults with type 2 diabetes, particular consideration should be given to  
41 their broader health and social care needs. Older people are more likely to have co-existing  
42 conditions and to be on a greater number of medicines. Their ability to benefit from risk-  
43 reduction interventions in the longer term may also be reduced.

1 Much of the evidence base used to inform this guideline has been generated from studies  
2 involving younger adults. While the Guideline Development Group (GDG) thought that the  
3 recommendations are applicable to a wider age group, they highlighted that there needs to  
4 be flexibility, to ensure that the care of older people with diabetes also addresses their  
5 broader health and social care needs.

6 Patients and healthcare professionals have rights and responsibilities as set out in the NHS  
7 Constitution for England – all NICE guidance is written to reflect these. Treatment and care  
8 should take into account individual needs and preferences. Patients should have the  
9 opportunity to make informed decisions about their care and treatment, in partnership with  
10 their healthcare professionals. If the patient is under 16, their family or carers should also be  
11 given information and support to help the child or young person to make decisions about  
12 their treatment. Healthcare professionals should follow the Department of Health's advice on  
13 consent. If someone does not have capacity to make decisions, healthcare professionals  
14 should follow the code of practice that accompanies the Mental Capacity Act and the  
15 supplementary code of practice on deprivation of liberty safeguards.

16 NICE has produced guidance on the components of good patient experience in adult NHS  
17 services. All healthcare professionals should follow the recommendations in Patient  
18 experience in adult NHS services.

## 3<sub>1</sub> Methods

2 This guideline update [2015] was developed in accordance with the process and methods  
3 outlined in 'The guidelines manual (2012)', which are different to those used to develop  
4 CG66 [2008] and CG87 [2009]. Chapters 7, 8, and 9.3 have been updated in 2015 and  
5 systematic reviews for each clinical question followed the review protocols (see Appendix C)  
6 agreed by the Guideline Development Group (GDG). GRADE (Grading of Recommendations  
7 Assessment, Development and Evaluation) methodology was used and/or adapted for  
8 appraising the quality of the evidence, and the Linking Evidence to Recommendations  
9 (LETR) framework was adopted to transparently document the GDG's decision making  
10 process. In instances where the guidelines manual does not provide advice, additional  
11 methods were used and are described in detail.

12 There is more information about how NICE clinical guidelines are developed on the NICE  
13 website. A booklet, 'How NICE clinical guidelines are developed: an overview for  
14 stakeholders, the public and the NHS' is available.

### 3.1<sub>5</sub> Population

16 The guideline focused on adults (aged 18 years and older) with type 2 diabetes. Studies with  
17 at least 85% of individuals with type 2 diabetes were included, unless otherwise stated.  
18 Evidence on specific patient subgroups for whom the management of type 2 diabetes may  
19 vary were considered where available. These included (but were not restricted to):

- 20 • adults aged 65 years and older
- 21 • people with renal impairment
- 22 • people in specific ethnic groups
- 23 • people in specific cardiovascular risk groups.

### 3.2<sub>4</sub> Outcomes

25 The outcomes prioritised in the review questions reflect the treatment objectives in the  
26 management of type 2 diabetes such as controlling blood glucose levels, reducing  
27 cardiovascular risk, minimising associated complications and improving life expectancy.  
28 Unless otherwise stated, the minimal important difference (MID) for dichotomous outcomes  
29 was defined as a relative risk reduction or an increase of 25% or more.

### 3.2.11 Change in blood glucose levels

2 Glycated haemoglobin (HbA1c) is commonly used in clinical practice to monitor glycaemic  
3 control as it provides a measure of average plasma glucose over the preceding 8 to 12  
4 weeks (Nathan et al. 2007), and therefore captures fluctuations including hypoglycaemic  
5 events. For this reason, the GDG agreed that change in HbA1c would be the main outcome  
6 measure used to reflect glycaemic control and a difference of 5 mmol/mol (0.5%) was  
7 considered to be clinically important. This blood test can be administered at any time and  
8 overcomes the issues of other tests (for example fasting and postprandial blood glucose)  
9 including day-to-day variability of glucose values and the inconvenience of special dietary  
10 preparation or fasting period. Since 1995, the International Federation of Clinical Chemistry  
11 (IFCC) has worked to standardise HbA1c analysis, establishing 2 reference methods – mass  
12 spectroscopy and capillary electrophoresis. Despite its advantages, HbA1c measurement  
13 may be affected by different factors such as haemoglobinopathies, illnesses like malaria that  
14 are associated with accelerated red blood cell turnover and certain anaemias.

15 The GDG agreed that fasting and postprandial blood glucose are rarely used in clinical  
16 practice and, therefore, were generally not prioritised as measures for change in blood  
17 glucose levels. However, they noted that a minimal important difference for both measures  
18 was 1 mmol/L (18 mg/dL).

### 3.2.29 Cardiovascular risk

20 Changes in blood pressure and lipid levels were included in the considered outcomes for the  
21 review question on drug treatments to control blood glucose (see review protocol in Appendix  
22 C). However, available data were too sparse or too different to allow for meaningful network  
23 meta-analyses to be undertaken and are therefore not reported.

### 3.2.34 Diabetes-related complications

25 Mortality, microvascular and macrovascular complications were prioritised by the GDG.  
26 These included cardiovascular disease, retinopathy, kidney damage, foot complications and  
27 erectile dysfunction specifically in men.

### 3.2.48 Adverse events

29 Across the included studies, adverse events were reported in many different ways. To allow  
30 for comparisons of studies and prevent double-counting of events, the following measures  
31 were prioritised for data extraction; total dropouts, dropouts due to adverse events and  
32 nausea. The GDG prioritised these measures because patients and clinicians are most  
33 interested in adverse events that affect treatment compliance and decisions.

### 3.2.4.34 Hypoglycaemia

35 Hypoglycaemia, although a common adverse event, was reported separately because of its  
36 significant negative impact on a person's wellbeing and quality of life and its influence on  
37 treatment decisions. Reporting of hypoglycaemia varied across the included trials in terms of  
38 definition of event and presentation of data. Both rate data (events per unit of person-time at  
39 risk) and dichotomous data (proportion of participants experiencing 1 or more event) were  
40 extracted. Where available, rate data were preferred to dichotomous data, because it is  
41 important to account for people who experience multiple events over time, and this  
42 information is lost when trial participants are split into those who have or have not  
43 experienced 1 or more event. Where rate data were not directly reported, they were  
44 sometimes estimable using the approach described in section 3.4.2.

1 The GDG also ranked the different types of hypoglycaemic data to reflect what they consider  
2 most clinically important. For the review question on drug treatments to control blood glucose  
3 (section 8.4), the highest ranking one reported in the trials was extracted. The hierarchy of  
4 hypoglycaemic data was:

- 5 • All hypoglycaemic events (number of events)
- 6 • All hypoglycaemic events (number of patients)
- 7 • Symptomatic hypoglycaemia
- 8 • Symptomatic (confirmed) hypoglycaemia
- 9 • Symptomatic (unconfirmed) hypoglycaemia
- 10 • Confirmed hypoglycaemia
- 11 • Minor hypoglycaemic events
- 12 • Minor (confirmed) hypoglycaemia
- 13 • Minor (unconfirmed) hypoglycaemia
- 14 • Moderate hypoglycaemia
- 15 • Moderate/severe hypoglycaemia
- 16 • Major/severe hypoglycaemic event
- 17 • Nocturnal hypoglycaemia
- 18 • Nocturnal (symptomatic) hypoglycaemia
- 19 • Nocturnal (confirmed) hypoglycaemia
- 20 • Nocturnal (mild) hypoglycaemia
- 21 • Nocturnal (moderate/severe) hypoglycaemia

#### 3.2.4.22 Change in body weight

23 Diabetes is related to obesity and some drug treatments are associated with weight gain.  
24 Change in body weight was considered separately from other adverse events and  
25 hypoglycaemia, because the GDG agreed that it is important to patients' quality of life and  
26 self-esteem, which may affect treatment compliance.

### 3.3 Data extraction

#### 3.3.18 Time-points

29 The included evidence reported a variety of follow-up periods. Given the number and  
30 heterogeneity of the time-points reported in the literature, it was important to prioritise which  
31 time-points were extracted. In order to enable the comparison of studies with different follow-  
32 up periods, the GDG considered it important to extract outcomes at common time-points.  
33 Based on clinical practice of 3-monthly medication review and the use of HbA1c as the main  
34 indicator of glycaemic control, the GDG agreed that the following time-points would provide  
35 clinically relevant evidence and enable comparisons across all studies for the review  
36 question focusing on drug treatments to lower blood glucose levels (section 8.4):

- 37 • 3 months (12 to 16 weeks)
- 38 • 6 months (22 to 30 weeks)
- 39 • 12 months (44 to 60 weeks)
- 40 • 24 months (96 to 112 weeks)

41 Data were extracted for each relevant timepoint that was reported in the included trials. If a  
42 study reported more than 1 data-point in the time ranges outlined above, the one closest to  
43 the central figure was extracted. For example, if data were reported at 25 and 28 weeks, the  
44 data-point closest to 6 months was extracted, that is 25 weeks. If data-points were

1 equidistant from the time-point, for example 24 and 28 weeks, the later time period, 28  
2 weeks was extracted. A minimum of 12 weeks' follow-up from start of treatment was agreed  
3 to be clinically relevant as it coincides with medicine reviews and HbA1c measurements.

4 For the supplementary review question on the long-term serious adverse effects of blood  
5 glucose lowering drug treatments (section 8.5), the GDG agreed that a minimum follow-up  
6 period of 2 years was sufficient to allow for adverse events and complications to occur.

7 For the review question on self-monitoring of blood glucose levels (section 8.3), the GDG  
8 agreed that a minimum follow-up period of 4 weeks would allow for important information on  
9 short-term outcomes such as hypoglycaemia to be captured.

10 No time restrictions were placed on the remaining review questions on optimal blood glucose  
11 targets (sections 8.1 and 8.2), use of antiplatelet therapy for primary prevention of  
12 cardiovascular disease (section 7) and management of erectile dysfunction (section 9.2).

13 For dichotomous outcomes such as adverse events, data were generally extracted at study  
14 end-point.

### 3.3.25 Conversion of continuous outcome data

16 Continuous outcomes which reported different units (for example, HbA1c in % or mmol/mol)  
17 were converted to a common unit prior to synthesis. Estimates of body weight in kilograms  
18 were calculated from studies which only reported body mass index (BMI). Where the mean  
19 height of the cohort was available, this was used to estimate weight; where no height data  
20 were available the mean height of people in the THIN dataset derived for the health  
21 economic model (168 cm; see 8.4.3.3) was used.

### 3.3.32 Process

23 Data were extracted by 1 reviewer and a second reviewer checked the studies included in  
24 the analyses. Where numerical data were not reported in tables or text, information was  
25 extracted from graphs by digitising the images and using a bespoke electronic ruler in  
26 Microsoft Excel. Data were typically extracted from graphs where relevant time-points were  
27 not reported (for example, the study reported outcomes at 1 year but provided a graph of  
28 changes over time with data-points at 3 and 6 months) and only if measures of dispersion  
29 were provided (for example, error bars from graphs were used to estimate standard  
30 deviations).

## 3.4 Data imputation

### 3.4.32 Estimating mean change from baseline

33 Where possible, mean difference from baseline to follow-up was the point of synthesis for  
34 continuous measures. If the study did not provide the mean difference, where possible, it  
35 was calculated from reported baseline and follow-up scores that is, follow-up score minus  
36 baseline value. However, the standard deviation (SD) of mean differences is also required for  
37 syntheses. To estimate this, it is necessary to specify the correlation between measurements  
38 at the 2 time-points. These were estimated from studies in the effectiveness evidence base.  
39 Where a study reports SD at baseline ( $\sigma_b$ ), SD at follow-up ( $\sigma_f$ ) and the SD of changes  
40 between baseline and follow-up ( $\sigma_c$ ), the correlation ( $C$ ) between baseline and follow-up for  
41 that study may be estimated by:

$$C = \frac{\sigma_b^2 + \sigma_f^2 - \sigma_c^2}{2 \times \sigma_b \times \sigma_f} \quad (1)$$

1 C was calculated for each arm (regardless of treatment assignment) in each study reporting  
2 the necessary information. These values were combined by a weighted average according to  
3 the number of people in the arm, and the resulting average C used to impute SDs of mean  
4 differences in studies that did not report them, using the formula:

$$\sigma_c = \sqrt{\sigma_b^2 + \sigma_f^2 - (2 \times C \times \sigma_b \times \sigma_f)} \quad (2)$$

5 In some instances, the correlation coefficient that was estimated from the evidence base was  
6 observed to be outside the acceptable values (that is, outside the range of -1 to 1) or were  
7 very close to perfect correlation. These were assumed to be a result of inaccuracies in the  
8 data, typos in the primary paper and unclear measures of reported variance (SD or standard  
9 error, SE), generally estimated from graphs. These estimated correlation coefficients were  
10 unlikely to represent true population values. In these cases, and also in syntheses where no  
11 studies provided sufficient evidence to estimate a correlation coefficient, a conservative value  
12 of 0.5 was used (Follmann et al. 1992).

### 3.4.23 Estimating person time at risk

14 When events are likely to occur to a person more than once (for example, hypoglycaemic  
15 events), it is preferable to use count or rate data. To calculate the rate of an event occurring,  
16 the total number of events and total person-time at risk are needed. However, papers did not  
17 commonly report person-time at risk.

18 Where papers reported the rate of events occurring and the total number of events, the  
19 corresponding person-time at risk was estimated. If studies provided data on specific timings  
20 of dropouts for individuals who withdrew from the trial, these durations were used to estimate  
21 the person-time at risk. Where these data were not reported, a crude estimate of person-time  
22 at risk for each arm in a trial was obtained from the number of participants ( $N$ ), the duration  
23 of the trial ( $D$ ) and the number of dropouts in the trial arm ( $y$ ) using the formula:

$$ND - 0.5Dy. \quad (3)$$

24 The accuracy of this crude estimation of person time at risk was tested by comparing values  
25 obtained using the equation above with values obtained using reported rates and total  
26 number of events. Although there were some differences in the values of person-time at risk,  
27 there was minimal impact on the overall rate of events.

### 3.4.38 Approach to missing data

29 Many of the included trials that used intention-to-treat (ITT) analyses used the last  
30 observation carried forward (LOCF) imputation, which is thought to overestimate treatment  
31 effects. Unfortunately, it is difficult to adequately deal with this data for continuous outcomes  
32 without individual patient data reported for each study.

## 3.5.3 Crossover trials

34 The incorporation of data from RCTs of parallel and crossover design in single quantitative  
35 syntheses is a subject of methodological debate (Elbourne et al. 2002). The following  
36 approaches were considered:

- 1 1. The optimal method is to include data from crossover studies in a way that exploits the  
2 increased precision the crossover design provides. This is straightforward where within-  
3 patient differences from a paired analysis are reported by authors; alternatively, methods  
4 are available that can impute these data if the correlation between treatment periods is  
5 known (or can be calculated) (Elbourne et al. 2002).
- 6 2. Another method sometimes used is to restrict attention to the first period of randomised  
7 treatment in each crossover trial only. In this way, a parallel trial of half the size is derived.  
8 This approach is suboptimal, as it discards data from the remainder of the trial, and relies  
9 on data being reported in a way that facilitates the extraction of data from the initial period  
10 only.
- 11 3. Another option is to exclude all crossover studies from consideration.
- 12 4. Finally, it is possible to ignore the crossover design of the trials, and analyse them as if  
13 they had a parallel design. This method is not generally recommended, as it ignores  
14 within-patient correlations and therefore discards the design advantages of crossover  
15 trials. However, this means that the approach is conservative, as it results in the trials  
16 having less weight in syntheses than they would have if paired data were used (or  
17 imputed).

18 The issue of washout period was discussed with the GDG and it was agreed that a minimum  
19 of 4 to 6 weeks would be adequate to minimise the influence of existing therapies. Therefore,  
20 the following decisions were taken relating to which data from crossover trials were  
21 extracted:

- 22 • If the trial reported analysis that is considered appropriate for crossover designs and a  
23 washout period of 4 to 6 weeks, then the end of treatment data were extracted.
- 24 • If the trial reported analysis that is considered appropriate for crossover designs but a  
25 washout period of less than 4 weeks, then data from the first treatment period only were  
26 extracted.
- 27 • If the trial did not report analysis that is considered appropriate for crossover designs, then  
28 data from the first treatment period only were extracted.

## 3.6.9 Evidence synthesis

### 3.6.10 Meta-analyses

31 Where possible, meta-analyses were conducted to combine the results of studies for each  
32 outcome. For continuous outcomes, where change from baseline data were reported in the  
33 trials and were accompanied by a measure of spread (for example standard deviation), these  
34 were extracted and used in the meta-analysis. Where measures of spread for change from  
35 baseline values were not reported, the corresponding values at study end were used and  
36 were combined with change from baseline values to produce summary estimates of effect.  
37 These studies were assessed to ensure that baseline values were balanced across the  
38 treatment groups; if there were differences at baseline these studies were not included in any  
39 meta-analysis and were reported separately.

### 3.6.20 Network meta-analyses

41 Network meta-analyses (NMAs) were conducted to simultaneously compare multiple  
42 treatments in a single meta-analysis, preserving the randomisation of the included trials in  
43 the reviews. This allows all evidence to be combined in a single internally consistent model.

44 An extensive series of NMAs was undertaken to synthesise evidence on pharmacological  
45 treatments to control blood glucose (see 8.4). The GDG's preferred approach to identifying  
46 and synthesising relevant evidence for these analyses relied on several critical assumptions  
47 that are discussed in 8.4.1.

1 Hierarchical Bayesian NMA was performed using the software WinBUGS version 1.4.3. The  
2 models were based on the approach and code provided in the NICE Decision Support Unit's  
3 Technical Support Documents on evidence synthesis, particularly Technical Support  
4 Document 2 ('A generalised linear modelling framework for pairwise and network meta-  
5 analysis of randomised controlled trials'; see <http://www.nicedsu.org.uk/>). Model code is  
6 provided in Appendix K.

### 3.6.2.17 Continuous outcomes

8 Identity-link models, which rely on a normal likelihood, were used for continuous outcomes. It  
9 should be emphasised that these models do not assume that the measures being  
10 synthesised are, themselves, normally distributed; rather, they assume that the sample  
11 means are normally distributed (given sufficiently large samples, this would be expected to  
12 be the case regardless of skewness in the underlying data, according to Central Limit  
13 Theorem; in the case in hand, many of the datasets are relatively small and convergence to  
14 a normal distribution of means may not have occurred; however, the same lack of data would  
15 make it difficult to select an alternative likelihood).

16 Mean difference from baseline to follow-up was the point of synthesis for continuous  
17 measures (see 3.4.1). We were unable to include the outcomes from studies where  
18 continuous data were reported in the form of median differences or as percentage change  
19 from baseline in syntheses as it is not possible to combine outcomes with these measures  
20 and mean differences (the point of synthesis chosen) without access to individual patient  
21 data.

22 The WinBUGS code used for this model is provided in Appendix K.

### 3.6.2.23 Dichotomous outcomes

24 As advised in NICE DSU TSD 2 (Dias et al. 2012a), dichotomous outcomes can be  
25 synthesised using 2 alternative models:

- 26 • The most straightforward model adopts a binomial likelihood with a logit link function, and  
27 generates output on a log-odds scale, with results transformed to odds ratios for  
28 presentation.
- 29 • An alternative model incorporates data on duration of follow-up in each underlying RCT,  
30 assuming a constant rate of events, to estimate the probability of events occurring over  
31 time. Again, a binomial likelihood is assumed, but a complementary log–log ('cloglog') link  
32 function is used, which results in outputs on a log-hazard scale (transformed into hazard  
33 ratios for presentation).

34 Where differences in follow-up in the underlying evidence were believed or shown to be  
35 minor and/or unimportant, the simpler logit-link model was preferred. Where duration of  
36 follow-up was believed to have a potential impact on outcomes, both models were explored,  
37 and the choice made on the basis of goodness of fit (see 3.6.2.7).

38 The WinBUGS code used for these models is provided in Appendix K.

### 39 Zero cells

40 In datasets containing studies with 'zero cells' (that is, trials in which no events occurred in 1  
41 or more arm), substantial instability was encountered when performing syntheses. To  
42 address this problem, a constant of 0.5 was added to all cell counts (effectively adding 0.5 to  
43 the numerator and 1 to the denominator of the proportion). The same approach was used to  
44 address instability for datasets containing studies with 100% events reported in all arms.

45 Studies reporting no events in any arms were excluded from NMAs, as they do not provide  
46 any information on the relative likelihood of events occurring.

### 3.6.2.31 Rate / count outcomes

2 For rate data (event per unit of person-time), a Poisson model with a log link function was  
3 used, to estimate the probability of events occurring over time. These models produce  
4 outputs on a log-hazard scale (transformed into hazard ratios for presentation).

### 3.6.2.45 Combining dichotomous and rate data

6 Because, as noted above, both rate data and dichotomous data (with an estimate of follow-  
7 up time) can be synthesised on a log-hazard scale, it is possible to combine both types of  
8 data in a hybrid model with appropriate likelihoods and link functions for each type of data.  
9 This assumes that, regardless of which way the data are reported, the incidence of events  
10 has the characteristics of a homogeneous Poisson process. Models of this type were run to  
11 combine heterogeneously reported data on incidence of hypoglycaemia (see 3.2.4.1).

12 The WinBUGS code used for the hybrid binomial–cloglog/Poisson–log model is provided in  
13 Appendix K.

### 3.6.2.54 Prior distributions

15 Non-informative prior distributions were used in all models. Trial baselines and treatment  
16 effects were assigned  $N(0, 100^2)$  priors. The between-trial standard deviations used in  
17 random-effects models were given  $U(0, 2)$  priors for dichotomous outcomes. It was felt that  
18 this standard deviation was appropriate as the upper limit of 2 represents a huge range of  
19 trial-specific treatment effects. This is recommended in NICE DSU Technical Support  
20 Document 2.  $U(0, 2)$  priors were also used for syntheses of continuous measures of HbA1c –  
21 given the relatively limited range in which HbA1c values fall, this was felt to be appropriately  
22 vague. Sensitivity analyses with broader priors demonstrated negligible impact.

### 3.6.2.63 Running the model

24 In the first instance, models were run with 50,000 burn-ins and 10,000 iterations. Three  
25 separate chains with different initial values were used. If models did not appear to converge  
26 well, they were re-run with more burn-ins and/or observations ‘thinned’ from a large number  
27 of posterior samples (for example, every 20<sup>th</sup> sample of 200,000 could be used to provide  
28 10,000 iterations with minimised autocorrelation).

29 Syntheses were assessed for any points that significantly deviated from the other data-points  
30 and the reasons for any deviate points were investigated.

### 3.6.2.71 Goodness of fit

32 Measures of model fit were scrutinised to assess appropriateness of each model. Particular  
33 attention was paid to:

- 34 • **Total residual deviance:** a calculation of the model’s ability to predict the individual data-  
35 points underlying it. In every iteration of the model sampling procedure, the amount each  
36 model-estimated data-point deviates from the observed evidence is calculated, summed  
37 and averaged over all iterations. Each data-point should contribute about 1 to the  
38 posterior mean deviance; therefore, the total residual deviance of a well-fitting model will  
39 be approximately the same as the number of independent data-points in the model
- 40 • **Deviance information criterion (DIC):** an estimate of deviance that is ‘penalised’  
41 according to the number of parameters in the model (adding parameters to a model  
42 should increase its ability to predict known data; however, this may come at the expense  
43 of reducing its ability to predict external datasets).
- 44 • **SD of random-effects term (tau):** where a random-effects model is fitted, the width of the  
45 inter-study heterogeneity distribution estimated by the model is a reflection of how well the

1 model accounts for heterogeneity in the underlying data. Therefore, while not a measure  
2 of goodness of fit *per se*, it is useful to consider as an indication of how broad a model is  
3 required to fit the data. Because inter-study heterogeneity is not modelled in fixed-effects  
4 models (that is, tau is assumed to be 0), there is no analogous quantity that can be used  
5 to compare different fixed-effects models.

### 3.6.2.86 Choice of model (random- versus fixed-effects)

7 For all syntheses, models were run as both random and fixed effects and model fit  
8 measurements were explored to select the most appropriate model for the specific outcome.  
9 If either model had clearly superior residual deviance and/or DIC, it was preferred; if there  
10 was little to choose between them, fixed-effects models were preferred for reasons of  
11 parsimony and interpretability. In practice, this led to a rule where fixed-effects models were  
12 preferred unless the corresponding random-effects model had a DIC that was 3 or more  
13 lower. Model fit statistics and selection decisions are shown in Appendix J.1.

14 An exception to this principle was in instances where there was only 1 study for each link in  
15 the network. In this case, no data are available to estimate the random-effects term;  
16 therefore, a fixed-effects model was used.

### 3.6.2.97 Meta-regression

18 For some larger datasets, the potential for heterogeneity of treatment effect to be explained  
19 by study-level covariates was explored in meta-regression (see NICE DSU TSD 3 [Dias et al.  
20 2012b]). In particular, for analyses of the relative effectiveness of pharmacological treatments  
21 (research question 1), it was considered important to account for baseline HbA1c level – it  
22 has been suggested that differences in baseline severity may account for some or all of  
23 observed differences in treatment effects (Chapell et al. 2009). However, none of these  
24 analyses produced models that provided a better fit to the data, as evident in the following  
25 characteristics:

- 26 • The regression coefficients were associated with broad credible intervals crossing 0
- 27 • In fixed-effects analyses, measures of goodness of fit were inferior for models including a  
28 covariate than for unadjusted models.
- 29 • In random-effects analyses, the heterogeneity term was not materially reduced.

30 For all these reasons, the approach was judged not to be informative, and results have not  
31 been reported here.

32 Although this was the case for the relative effect estimates presented here, it was not true of  
33 the absolute HbA1c effect estimates – to which relative effects are then applied – that are  
34 necessary for the health economic model (see Appendix F3.5.1 for a description of the  
35 adjustment of these analyses for baseline level).

### 3.6.2.106 Inconsistency between direct and indirect evidence

37 As suggested in NICE DSU TSD 4 [Dias et al. 2012c], an 'inconsistency' model was fitted to  
38 each dataset on which NMA was undertaken. The outputs of these models were compared  
39 with the relevant NMA ('consistency' model) to identify any discrepancies between direct and  
40 indirect evidence. In particular, the posterior mean of the residual deviance contribution of  
41 within-trial comparisons in each of the 2 models were plotted against each other and visually  
42 inspected to see if any inconsistency was suggested (any absolute discrepancy of greater  
43 than 0.5 was highlighted and investigated). In practice, few such inconsistencies were seen,  
44 and any that occurred were invariably easily explained (in particular, dichotomous syntheses  
45 in which zero events were observed in 1 or more trial-arm resulted in high and variable  
46 residual deviance estimates). For these reasons (and to avoid unnecessary multiplication of  
47 already-numerous results), outputs of the inconsistency models have not been reported. The

1 posterior estimates of effect have, however, been used to show direct evidence in the  
2 pairwise relative effect plots relating to dichotomous data (which relied on cloglog or hybrid  
3 models that do not lend themselves to simple pairwise frequentist meta-analysis).

### 3.6.2.114 Presentation of results for network meta-analyses

5 The results of the meta-analyses were presented in a number of ways.

- 6 • Network diagram, showing availability of evidence. These diagrams have the following  
7 features:
  - 8 ○ The size of each node is proportional to total number of participants randomised to  
9 receive the treatment in question across the evidence-base.
  - 10 ○ The width of connecting lines is proportional to number of trial-level comparisons  
11 available.
  - 12 ○ Where possible, arrowheads are added to the connecting lines to indicate direction of  
13 effect in pairwise data ( $a > b$  denotes  $a$  is more effective than  $b$ ) – filled arrowheads  
14 show comparisons where one option is significantly superior ( $p < 0.05$ ); outlined  
15 arrowheads show direction of trend where effect does not reach statistical significance.  
16 It has not been possible to add these for some analyses, as it is not straightforward to  
17 estimate direction of effect with more complex models.
- 18 • Plot of the relative effectiveness, including the results of the NMA of each regimen  
19 compared with the reference treatment (for example, see Figure 31) and any direct  
20 estimate available for the same comparison.
- 21 • Tabulated rank probabilities, giving the probability of each treatment being best (that is,  
22 ranked #1) and its median rank with 95% credible interval (CrI). In these outputs, higher  
23 ranking always reflect what is best for the patient (for example: higher rates of disease  
24 eradication, lower rates of adverse events, lower blood glucose levels, and so on).

25 More detailed model outputs and a summary of input data for each analysis are available in  
26 Appendix J.

## 3.7 Quality assessment

28 GRADE was used to assess the quality of evidence for the selected outcomes as specified in  
29 'The guidelines manual (2012)'.

### 3.7.10 GRADE for pairwise meta-analyses

31 The quality of the evidence base was downgraded for the reasons outlined in Table 1.

32 **Table 1: Rationale for downgrading quality of evidence in pairwise meta-analyses for**  
33 **intervention questions**

GRADE criteria	Example reasons for downgrading quality
Risk of bias	This includes limitations in the design or execution of the study, including concealment of allocation, blinding, loss to follow up (these can reduce the quality rating)
Inconsistency	Inconsistency of effects across studies: occurs when there is variability in the treatment effect demonstrated across studies (heterogeneity). This was assessed using the statistic, $I^2$ where ; $I^2 < 30$ was categorised as no inconsistency, $I^2$ between 30% and 60% was categorised as serious inconsistency and $I^2 > 60\%$ was categorised as very serious inconsistency (this can reduce the quality rating)
Indirectness	The extent to which the available evidence fails to address the specific review question (this can reduce the quality rating)
Imprecision	Present when there is uncertainty around the estimate of effect, for example

GRADE criteria	Example reasons for downgrading quality
	when the confidence intervals are wide and cross the 'imaginary' lines of clinically significant effect that is minimal important difference. This reflects the confidence in the estimate of effect. Minimal important differences are selected <i>a priori</i> by GDG consensus or from published estimates. For dichotomous outcomes, imprecision was assessed by use of minimal important difference of 0.25 (this can reduce the quality rating)
Other considerations	Large magnitude of effect, evidence of a dose-response relationship, or confounding variables likely to have reduced the magnitude of an effect; these can increase the quality ratings in observational studies, provided no downgrading for other features has occurred

### 3.7.21 Modified GRADE for network meta-analyses

2 The use of GRADE to assess the quality of studies addressing a particular review question  
3 for pairwise comparisons of interventions is relatively established. However, the use of  
4 GRADE to assess the quality of evidence across a NMA is still a developing methodology.  
5 While most criteria for pairwise meta-analyses still apply, it is important to adapt some of the  
6 criteria to take into consideration additional factors, such as how each 'link' or pairwise  
7 comparison within the network applies to the others. As a result, the following was used  
8 when applying modified GRADE to a NMA.

9 **Table 2: Rationale for downgrading quality of evidence in network meta-analyses**

GRADE criteria	Example reasons for downgrading quality
Risk of bias	Trials with large reductions in outcome measures were associated with high risk of bias for example if: <ul style="list-style-type: none"> <li>• There was a tendency for higher baseline HbA1c which may have had an undue effect (such as large trials with high baseline HbA1c levels of more than 69 mmol/mol (8.5%) for initial therapy may have had an impact on the overall rankings)</li> </ul>
Inconsistency	Evidence of any inconsistency between the direct and indirect estimates of effect was assessed using the residual deviance, deviance information criterion and the statistic tau. Downgrade if tau > 0.5
Indirectness	Trials were conducted in countries where dietary habits may differ and may not be representative of people with type 2 diabetes living in the UK (for example Japan and China). Evidence was only downgraded if this was likely to have a large impact on the overall rankings (that is, within smaller networks where there is a lack of evidence or within larger networks in large trials which show large reductions in outcomes)
Imprecision	This was assessed based on the overall distribution of the rankings, such that evidence was downgraded if no interventions had rank credible intervals ≤33% of total distribution of comparators

Update 2015

### 3.7.30 Modified GRADE for prognostic evidence

11 GRADE has not been developed for use with prognostic studies; therefore a modified  
12 approach was applied using the framework provided for GRADE in diagnostic studies. This  
13 assessment was used for evidence in the review question on optimal target values (see  
14 section 8.1).

15 Cohort studies within the non-modified GRADE approach start at the low quality level due to  
16 accepted inherent study design limitations. Within a modified approach it is acceptable to  
17 initially indicate a high quality level to this study type and to assess the quality of evidence  
18 from this point. The same criteria (risk of bias, inconsistency, imprecision and indirectness)  
19 were used to downgrade the quality of evidence. Quality ratings were downgraded further for  
20 risk of bias if there was evidence of selection bias. Indirectness was assessed by examining

1 any important differences in population, prognostic factor or outcome of the included  
2 evidence compared with those for whom the recommendation is intended. Imprecision was  
3 assessed by examining the sample size or the 95% confidence intervals around the estimate  
4 of effect. GRADE provides a guide when assessing imprecision in intervention questions  
5 (that is, where the total sample size is less than 400, the event rate is less than 300, or the  
6 95% confidence intervals cross the thresholds for appreciable benefit or harm or the minimal  
7 important difference). The evidence was downgraded for imprecision where the 95%  
8 confidence intervals were wide or the sample size was less than 400.

## 4<sub>1</sub> Education

### 4.1<sub>2</sub> Structured education

#### 4.1.1<sub>3</sub> Clinical introduction

4 Type 2 diabetes mellitus is a progressive long-term medical condition that is predominantly  
5 managed by the person with diabetes and/or their carer as part of their daily life. Accordingly,  
6 understanding of diabetes, informed choice of management opportunities, and the  
7 acquisition of relevant skills for successful self-management play an important role in  
8 achieving optimal outcomes. Delivery of these needs is not always assured by conventional  
9 clinical consultations. Structured programmes have been designed not only to improve  
10 people's knowledge and skills, but also to help motivate and sustain people with diabetes in  
11 taking control of their condition and in delivering effective self-management.

12 Information from the Health Commission survey in 2007 suggests that only 11% of people  
13 with type 2 diabetes report being offered structured education.<sup>8</sup> This suggests that the  
14 majority of healthcare providers have found it difficult to implement and resource quality  
15 education programmes that meet these standards. There appears to be an urgent need to  
16 ensure that all people with type 2 diabetes are offered high-quality structured education. The  
17 aims of structured education and self-management programmes are to improve outcomes  
18 through addressing the individual's health beliefs, optimising metabolic control, addressing  
19 cardiovascular risk factors (helping to reduce the risk of complications), facilitating behaviour  
20 change (such as increased physical activity), improving quality of life and reducing  
21 depression. An effective programme will also enhance the relationship between the person  
22 with diabetes and their healthcare professionals, thereby providing the basis of true  
23 partnership in diabetes management.

24 The clinical question that has been addressed is how to deliver such education, including  
25 what approaches deliver the intended benefits, and what components of the education  
26 process best deliver the surrogate, self-care, and quality of life outcomes.

#### 4.1.1<sub>27</sub> Methodological introduction and evidence statements

28 Please refer to the Technology Assessment Report 'The clinical effectiveness of diabetes  
29 education models for type 2 diabetes: a systematic review' commissioned by the NHS R&D  
30 Health Technology Assessment (HTA) programme on behalf of the NCC-CC. Available at  
31 [www.ncchta.org/project/1550.asp](http://www.ncchta.org/project/1550.asp)

#### 4.1.1<sub>32</sub> Health economic methodological introduction

33 Two papers were identified in the search for health economics. Neither study was conducted  
34 in the UK and the results were not generalisable to the UK setting so both were excluded.<sup>9,10</sup>

#### 4.1.1<sub>45</sub> Evidence to recommendations

36 The GDG noted that the last review of this area by a HTA on behalf of NICE in 2003 looked  
37 at the evidence for structured education. Little robust evidence of the effectiveness of any  
38 particular educational approach for people with type 2 diabetes was found. One conclusion  
39 was that further research was required, but meanwhile that educational programmes with a  
40 theoretical basis demonstrated improved outcomes, and that group education was a more  
41 effective use of resources and may have additional benefits.

42 Educational interventions are not only complex in themselves, but they also exist in a  
43 complex environment with other aspects of managing a chronic disease. Such interventions

1 will interact with, and support medical management directed at vascular risk factors and that  
2 of diabetes complications which have already developed. Their success is likely to depend  
3 on the individual's personal and cultural beliefs, the overall healthcare setting, their lifestyles,  
4 and perhaps their educational background.

5 It was noted that to address some of the difficulties in describing and implementing effective  
6 structured education and self-management programmes, a Patient Education Working Group  
7 (PEWG) had been convened by the Department of Health and Diabetes UK, and had laid out  
8 in detail the necessary requirements for developing high-quality patient education  
9 programmes. The key criteria had been endorsed by the recent HTA review. The 5 standards  
10 were as follows.

- 11 1. Any programme should have an underpinning philosophy, should be evidence-based, and  
12 suit the needs of the individual. The programme should have specific aims and learning  
13 objectives, and should support development of self-management attitudes, beliefs,  
14 knowledge and skills for the learner, their family and carers.
- 15 2. The programme should have a structured curriculum which is theory-driven, evidence-  
16 based, resource-effective, have supporting materials, and be written down.
- 17 3. It should be delivered by trained educators who should have an understanding of the  
18 educational theory appropriate to the age and needs of the programme learners, and be  
19 trained and competent in delivery of the principles and content of the specific programme  
20 they are offering.
- 21 4. The programme itself should be quality assured, be reviewed by trained, competent,  
22 independent assessors and be assessed against key criteria to ensure sustained  
23 consistency.
- 24 5. The outcomes from the programme should be regularly audited.

25 The GDG found no reason to diverge from these principles. The GDG noted and endorsed  
26 the importance of quality assurance and audit in this complex area.

27 As the intervention is complex, the measured outcomes of any particular programme are by  
28 nature multifaceted and will vary with such factors as the timing in relation to diagnosis,  
29 critical changes of therapy, or other critical clinical findings. Even then, appropriate study  
30 outcomes are for the most part interim surrogate measures; no studies included late  
31 complications. However, psychological outcomes as well as biomedical outcomes can be  
32 appropriately assessed, to include quality of life and change in healthcare behaviours, and  
33 aspects of depressed mood. More directly cognitive measures, knowledge, acquisition of  
34 skills, and changing health beliefs were found to be useful indicators of a programme's  
35 effectiveness.

36 The HTA commissioned for this review included 14 studies, of which 8 appeared to have  
37 been conducted since 2003, and most were for people with established (rather than newly  
38 diagnosed) type 2 diabetes. The GDG noted that, as expected, some studies showed effects  
39 on HbA1c, others improved body weight and other lifestyle changes, some improved quality  
40 of life or knowledge, and yet others changed health beliefs or reduced depression. This  
41 diversity was often a reflection of study aims and design. The HTA review acknowledged that  
42 health psychology approaches and some methods of health promotion have a good evidence  
43 base, but little is incorporated into studies of structured education, even though addressing  
44 health beliefs and motivating individuals to change behaviour is a cornerstone of any  
45 educational programme. Reported training for diabetes educators was poorly detailed in  
46 most studies.

47 The GDG was concerned that only 3 studies were UK-based. As cultural issues, patient  
48 health beliefs and attitudes are likely to differ from 1 country to another, applicability of the  
49 others may be limited. The GDG noted that the UK Diabetes Education and Self  
50 Management for Ongoing and Newly Diagnosed (DESMOND study) found changes in health  
51 beliefs, reduction in depression, and increases in self-reported physical activity, reduction in

- 1 weight and improvement in smoking status. In people with established diabetes there was  
2 useful evidence from the X-PERT programme with improvements in HbA1c, reduced  
3 diabetes medication, body weight, waist circumference, total serum cholesterol, diabetes  
4 knowledge and increase in self-reported physical activity and treatment satisfaction.
- 5 Overall the GDG then felt that well-designed and well-implemented programmes were likely  
6 to be effective and cost-effective interventions for people with type 2 diabetes, in line with the  
7 NICE TA. For those people in whom education delivered in a group setting is appropriate, it  
8 is evidently likely to be more cost effective.

#### 4.1.59 Recommendations and research recommendations

##### 4.1.5.10 Individualised care

- 11 **1. Adopt an individualised approach to diabetes care that is tailored to the person's**  
12 **needs and circumstances, taking into account their personal preferences,**  
13 **comorbidities, risks of polypharmacy, and their ability to benefit from long-term**  
14 **interventions due to reduced life expectancy. Such an approach is especially**  
15 **important in the context of multimorbidity. Reassess the person's needs and**  
16 **circumstances at each review and consider whether to stop any medicines that**  
17 **are not effective. [new 2015]**

##### 4.1.5.28 Patient education

- 19 **2. Offer structured education to adults with type 2 diabetes and/or their family**  
20 **members or carers (as appropriate) at and around the time of diagnosis, with**  
21 **annual reinforcement and review. Explain to people and their carers that**  
22 **structured education is an integral part of diabetes care. [2009]**
- 23 **3. Ensure that any structured education programme for adults with type 2 diabetes**  
24 **includes the following components<sup>c</sup>:**
- 25 • It is evidence-based, and suits the needs of the person.
  - 26 • It has specific aims and learning objectives, and supports the person  
27 and their family members and carers in developing attitudes, beliefs,  
28 knowledge and skills to self-manage diabetes.
  - 29 • It has a structured curriculum that is theory-driven, evidence-based and  
30 resource-effective, has supporting materials, and is written down.
  - 31 • It is delivered by trained educators who have an understanding of  
32 educational theory appropriate to the age and needs of the person, and  
33 who are trained and competent to deliver the principles and content of  
34 the programme.
  - 35 • It is quality assured, and reviewed by trained, competent, independent  
36 assessors who measure it against criteria that ensure consistency.
  - 37 • The outcomes are audited regularly. [2015]
- 38 **4. Ensure the patient-education programme provides the necessary resources to**  
39 **support the educators, and that educators are properly trained and given time to**  
40 **develop and maintain their skills. [2009]**

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<sup>c</sup> [Structured patient education in diabetes: report from the patient education working group](#)

- 1 **5. Offer group education programmes as the preferred option. Provide an alternative**  
2 **of equal standard for a person unable or unwilling to participate in group**  
3 **education. [2009]**
  
- 4 **6. Ensure that the patient-education programmes available meet the cultural,**  
5 **linguistic, cognitive and literacy needs within the local area. [2009]**
  
- 6 **7. Ensure that all members of the diabetes healthcare team are familiar with the**  
7 **patient-education programmes available locally, that these programmes are**  
8 **integrated with the rest of the care pathway, and that adults with type 2 diabetes**  
9 **and their family members or carers (as appropriate) have the opportunity to**  
10 **contribute to the design and provision of local programmes. [2009]**  
11

## 5<sub>1</sub> Lifestyle and non-pharmacological 2 management

### 5.1<sub>3</sub> Dietary advice

#### 5.1.1<sub>4</sub> Clinical introduction

5 All people with type 2 diabetes should be supported to:

- 6 • try to achieve and maintain blood glucose levels and blood pressure in the normal range
- 7 or as close to normal as is safely possible
- 8 • maintain a lipid and lipoprotein profile that reduces the risk of vascular disease.

9 Optimal dietary behaviours can contribute to all of these.

10 Dietary intervention should address the individual's nutritional needs, taking into account  
11 personal choices, cultural preferences and willingness to change, and to ensure that quality  
12 of life is optimised. It is usual that a registered dietician plays a key role in providing  
13 nutritional care advice within the multidisciplinary diabetes team. It is also recognised that all  
14 team members need to be knowledgeable about nutritional therapy, and give emphasis to  
15 consistent dietary and lifestyle advice.<sup>11</sup>

16 The management of obesity is not specifically addressed in the current guideline. Readers  
17 are referred to the NICE obesity guideline which addresses the area in some detail.<sup>12</sup>

18 Smoking cessation is not addressed in the current guideline. Readers are referred to the  
19 NICE public health programme guidance on smoking cessation services, including the use of  
20 pharmacotherapies, in primary care, pharmacies, local authorities and workplaces, with  
21 particular reference to manual working groups, pregnant smokers and hard to reach  
22 communities.

23 Clinical questions arise around the optimal strategies to reduce calorie intake (and thus  
24 improve sensitivity to endogenous insulin), to control exogenous delivery of free sugars into  
25 the circulation, to control blood pressure, and to optimise the blood lipid profile. Issues  
26 specifically related to people with kidney disease or of medical use of fish oils are not  
27 considered in this this guideline.

#### 5.1.2<sub>8</sub> Methodological introduction

29 The search attempted to identify RCTs and observational studies conducted in adults with  
30 type 2 diabetes which were assessing different forms of dietary advice targeting weight loss.  
31 A sample size threshold of N=50 and a follow-up of at least 3 months were established as  
32 cut-off points. Studies evaluating purely pharmacological interventions for weight reduction  
33 were excluded.

34 There were only 8 studies that addressed this question.<sup>13–20</sup> Two RCTs were excluded due to  
35 methodological limitations.<sup>d</sup> In all the studies, the intent was for participants to lose weight  
36 and thereby improve glycaemic, lipid and blood pressure control.<sup>e</sup> Among the remaining 6  
37 studies there were 4 RCTs and 2 observational studies. No major methodological limitations  
38 were identified across these studies.

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<sup>d</sup> One RCT comparing the effects of a high-protein with a low-protein diet<sup>15</sup> and another RCT comparing low-carbohydrate versus conventional weight loss diets in severely obese adults.<sup>18</sup>

<sup>e</sup> Four studies focused on the effects of diet in obese people with type 2 diabetes.

## 1 RCTs

- 2 One RCT<sup>17</sup> compared the effects of a combined intervention; low-calorie diet, sibutramine  
3 therapy and meal replacements with an individualised reduced calorie diet, and was the only  
4 study to include the use of weight-loss medication.
- 5 Two RCTs used the American Diabetes Association (ADA) guidelines as a comparison group  
6 to either a soy-based meal replacement intervention,<sup>13</sup> N=104 with a 1-year follow-up, or a  
7 low-fat vegan diet,<sup>14</sup> N=99 with a 22-week follow-up.
- 8 A further RCT compared a low-fat with a low-carbohydrate diet.<sup>16</sup>

## 9 Observational studies

- 10 A case series with a follow-up of 6.5 years investigated the onset of diabetic complications  
11 and adherence to ADA recommendations.<sup>19</sup> A prospective cohort study addressed the  
12 relationship between eating habits and long-term weight gain, following a group of patients  
13 being managed in primary care for a period of 4 years.<sup>20</sup>
- 14 It should be noted that the results of diet interventions aimed at patients with type 2 diabetes  
15 are difficult to interpret due to differences in the interventions, the populations, the study  
16 designs and the outcomes reported.
- 17 As is obvious, isolated diet interventions without adequate educational support and  
18 concomitant lifestyle changes are very unlikely to reduce risk factors and to improve clinical  
19 outcomes and quality of life for patients with type 2 diabetes.

### 5.1.30 Health economic methodological introduction

- 21 No health economic papers were identified.

### 5.1.42 Evidence statements

#### 5.1.4.23 Weight reduction and glycaemic control outcomes

##### 24 RCTs

##### 25 Studies that compared a meal replacement intervention with a reduced calorie diet

- 26 A RCT comparing a soy-based meal replacement with an individualised diet based on ADA  
27 recommendations in obese people with type 2 diabetes<sup>13</sup> found that average weight reduction  
28 in the meal replacement group was greater than that in the individualised diet group. At 6  
29 months, the meal replacement group had lost on average 5.24±0.60 kg, and the  
30 individualised diet group had lost an average of 2.85±0.67 kg (p=0.0031). At 1 year this  
31 difference was not significant with the meal replacement group losing on average 4.35±0.81  
32 kg and the individualised diet group losing an average of 2.36±0.76 kg (p=0.0670). **Level 1+**
- 33 The same RCT reported that similar changes were observed in the body mass index (BMI) at  
34 12 months with a reduction of 1.47±0.27 kg/m<sup>2</sup> in the meal replacement group and 0.77±0.25  
35 kg/m<sup>2</sup> in the individualised diet group. Although these values were significantly different from  
36 their baseline values, none were significantly different from each other (p=0.0687). **Level 1+**
- 37 With respect to glycaemic control, the RCT found that mean HbA1c levels were significantly  
38 lower in the meal replacement than in the individualised diet group, 0.49±0.22% (p=0.0291),  
39 for the entire study period. Plasma glucose concentrations were significantly lower in the  
40 meal replacement group than in the individualised diet group at 3 (p=0.04) and 6 (p=0.002)  
41 months, but not at 12 months (p=0.595). **Level 1+**

1 The study by Redmon<sup>17</sup> reported on a combination intervention including sibutramine, an  
2 intermittent low-calorie diet with the use of meal replacements for 1 week every 2 months,  
3 and the use of meal replacements between the low-calorie diet weeks. The comparison  
4 group received an individualised diet plan with a 500–1000 kcal energy deficit per day.

5 The study reported that at 1 year of follow-up, the combination therapy group had a  
6 significantly greater weight loss of  $7.3\pm 1.3$  kg than the standard therapy group  $0.8\pm 0.9$  kg  
7 ( $p<0.001$ ), with most weight loss occurring during the low-calorie weeks and some weight  
8 gain occurring in between the low-calorie weeks. **Level 1+**

9 In relation to glycaemic control, the study showed that at 1 year, HbA1c had declined from a  
10 baseline of  $8.1\pm 0.2\%$  to  $7.5\pm 0.3\%$  in the combination therapy group but had remained  
11 unchanged at  $8.2\pm 0.2\%$  in the standard therapy group, and this difference was significant  
12 ( $p=0.05$ ). After adjusting for medication changes, this difference remained significant. In an  
13 analysis of those participants whose medication had not changed, it was found that there  
14 was a significant positive linear association between change in weight at 1 year and change  
15 in HbA1c ( $r=0.53$ ;  $p=0.006$ ). A 5 kg decrease in weight at 1 year was associated with a 0.4%  
16 decrease in HbA1c. **Level 1+**

### 17 **Studies comparing a low carbohydrate with a low fat diet**

18 One RCT<sup>16</sup> examined the short-term effects, participants were followed up for 3 months, of a  
19 low-carbohydrate diet compared with a reduced portion low-fat diet in obese people with type  
20 2 diabetes. There was a significantly larger mean weight reduction in the low-carbohydrate  
21 arm ( $N=51$ ) of the RCT,  $3.55\pm 0.63$  kg, than in the low-fat arm ( $N=51$ ) which showed a mean  
22 reduction of  $0.92\pm 0.40$  kg ( $p=0.001$ ). **Level 1+**

23 The same RCT reported that glycaemic control improved in both arms of the trial.  
24 Improvements were greater in the low-carbohydrate arm, HbA1c decreased from a baseline  
25 of  $9.00\pm 0.20\%$ , by  $0.55\pm 0.17\%$ , but this did not reach statistical significance. In the low-fat  
26 arm HbA1c decreased from a baseline of  $9.11\pm 0.17\%$  by  $0.23\pm 0.13\%$  ( $p=0.132$ ). **Level 1+**

### 27 **Studies comparing low or modified fat diets with reduced calorie diets**

28 Barnard et al.<sup>14</sup> investigated the effects of a low-fat vegan diet compared with a diet based on  
29 ADA guidelines, on body weight and glycaemic control in a RCT with 99 people with type 2  
30 diabetes, followed up for 22 weeks. During the study period, 43% (21/49) of vegan  
31 participants and 26% (13/50) of ADA participants reduced their diabetic medications, mainly  
32 as a result of hypoglycaemia. Eight per cent in each group, 4/49 of the vegan group and 4/50  
33 of the ADA group, increased their medications.

34 The study concluded that for the whole sample, body weight was reduced in both groups by  
35 5.8 kg in the vegan group and 4.3 kg in the ADA group, but this difference was not  
36 statistically significant ( $p=0.082$ ). In those whose medication was stable this difference was  
37 significant with a 6.5 kg reduction in the vegan group, and 3.1 kg in the ADA group,  $p<0.001$ .  
38 BMI declined by  $2.1\pm 1.5$  kg/m<sup>2</sup> in the vegan group and by  $1.5\pm 1.5$  kg/m<sup>2</sup> in the ADA group  
39 ( $p=0.08$ ). The waist-to-hip ratio declined in the vegan group  $0.02\pm 0.01$  but not in the ADA  
40 group ( $p=0.003$ ). **Level 1+**

41 With respect to glycaemic control, the RCT stated that while the HbA1c decline in both  
42 groups was statistically significant from their baseline values with a decrease of 0.96%  
43 ( $p<0.0001$ ) in the vegan group and 0.56% ( $p=0.0009$ ) in the ADA group, there was no  
44 significant difference between the groups ( $p=0.089$ ). Again the results were different in those  
45 participants whose medication was unchanged. The HbA1c decline was greater in the vegan  
46 group,  $1.23\pm 1.38\%$ , than in the ADA group,  $0.38\pm 1.11\%$ , ( $p=0.01$ ). **Level 1+**

1 **Table 3: Summarised results for body weight reduction and glycaemic control across**  
2 **RCTs**

RCTs	Follow-up	Comparison	Comparison	Weight/BMI	Glycaemic control
Li (2005) <sup>13</sup>	1 year	Soy based meal replacement	Individualised diet	Weight and BMI=NS	HbA1c significantly lower in meal replacement arm
Redmon (2003) <sup>17</sup>	1 year	Sibutramine + low calorie diet + meal replacement	Individualised diet	Weight reduction significantly higher in combination arm	HbA1c significantly lower in combination arm*
Daly (2006) <sup>16</sup>	3 months	Low-carbohydrate diet	Reduced portion low-fat diet	Weight reduction significantly higher in combination arm	HbA1c=NS
Barnard (2006) <sup>14</sup>	22 weeks	Low-fat vegan diet	Diet based on ADA guidelines	Weight=NS	HbA1c=NS

\*A 5 kg decrease in weight at 1 year was associated with a 0.4% decrease in HbA1c  
NS not significant

### 3 **Observational studies**

4 In an observational study with 4 years of follow-up,<sup>20</sup> the authors investigated the association  
5 between eating behaviour and long-term weight gain. Ninety-seven people with type 2  
6 diabetes were recruited at diagnosis and after initial nutrition advice were followed up for a  
7 period of 4 years.

8 The study found that at the end of follow-up, mean body weight change in men was a gain of  
9 1.3±5.4 kg, whereas in women, there was a mean body weight reduction of -1.1±5.0 kg.  
10 These changes were not statistically significant (p values not given). Similarly, BMI increased  
11 in men by 0.42±1.76 kg/m<sup>2</sup> and decreased in women by 0.40±1.89 kg/m<sup>2</sup> (p values not  
12 given). Glycaemic outcomes were not reported. **Level 2+**

13 In the second observational study,<sup>19</sup> weight loss over the 6.5-year follow-up is not reported.  
14 However, metabolic control did improve in patients over the period, with the proportion of  
15 patients with HbA1c <7% increasing from 52.4% to 64.3% in men and from 43.9% to 50.9%  
16 in women. It was not reported whether or not this was significant. **Level 3**

#### 5.1.4.27 **Blood pressure and blood lipid control outcomes**

##### 18 **RCTs**

##### 19 **Studies that compared a meal replacement intervention with a reduced calorie diet**

20 The RCT by Li et al.<sup>13</sup> reporting on the comparison of a soy-based meal replacement plan  
21 with an individualised diet plan, did not report on changes in blood pressure during the study.

22 For the blood lipid control outcomes, while there were no significant differences between  
23 groups during the study for lipid parameters, there were differences within the groups when  
24 compared to baseline values. In the meal replacement group, there were decreases in total

1 cholesterol, triglycerol, low-density lipoprotein (LDL) and high-density lipoprotein (HDL) at the  
2 end of the study, however these changes were only significant in the triglycerol group with an  
3 overall decrease from baseline of 28.00 mg/dl ( $p=0.038$ ). Decreases in total cholesterol were  
4 significant at 3 ( $p<0.0001$ ) and 6 ( $p=0.0037$ ) months, but at 12 months with a reduction of  
5 10.76 mg/dl from baseline, this was not significant ( $p=0.084$ ). LDL decreased by 11.04 mg/dl  
6 at 3 months ( $p=0.024$ ), but at 12 months the change from baseline had reduced to 6.10 mg/dl  
7 ( $p=0.255$ ). HDL had decreased by 0.97 mg/dl at 12 months ( $p=0.345$ ). In the individualised  
8 diet plan group, after initial decreases at 3 or 6 months, at 12 months there were increases in  
9 total cholesterol by 5.26 mg/dl ( $p=0.396$ ), LDL by 8.76 mg/dl ( $p=0.129$ ) and HDL by 2.26  
10 mg/dl ( $p=0.012$ ). Only in triglycerol levels was there a sustained decrease at 12 months with  
11 a reduction from baseline of 28.89 mg/dl ( $p=0.119$ ). **Level 1+**

12 In the study by Redmon<sup>17</sup> which compared a combined intervention (described above) with  
13 an individualised diet plan, at 1 year there were reductions in systolic and diastolic blood  
14 pressure in both groups, although this did not differ between the groups. Systolic blood  
15 pressure reduced in the combination group by  $6\pm 3$  mmHg and by  $6\pm 2$  mmHg in the  
16 comparison group. Diastolic blood pressure reduced in the combination group by  $3\pm 1$  mmHg  
17 and by  $6\pm 2$  mmHg in the comparison group. **Level 1+**

18 At 1 year, changes in fasting cholesterol, HDL, LDL and fasting triglycerides did not differ  
19 between groups. There were reductions from baseline values in fasting cholesterol and LDL  
20 cholesterol in both groups, with a decrease in fasting cholesterol of  $6\pm 8$  mg/dl in the  
21 combination therapy group and  $17\pm 9$  mg/dl in the comparison group ( $p=0.90$ ). LDL  
22 decreased by  $12\pm 5$  mg/dl in the combination therapy group and  $13\pm 6$  mg/dl in the  
23 comparison group ( $p=0.89$ ). Fasting triglycerides decreased by  $46\pm 24$  mg/dl in the  
24 combination group compared to an increase of  $8\pm 18$  mg/dl in the comparison group, however  
25 this was not significant ( $p=0.07$ ). **Level 1+**

## 26 **Studies comparing a low-carbohydrate with low-fat diet**

27 At 12 weeks of follow-up, in the low-carbohydrate arm of this RCT<sup>16</sup> there was a reduction in  
28 systolic blood pressure of  $6.24\pm 2.96$  mmHg and a reduction of  $0.39\pm 2.64$  mmHg in the low-  
29 fat arm, with no significant difference between the arms ( $p=0.147$ ). **Level 1+**

30 With respect to lipid parameters, there was a greater reduction in the total cholesterol: HDL  
31 ratio in the low-carbohydrate arm, mean reduction of 0.48, than in the low-fat arm, mean  
32 reduction 0.10 ( $p=0.011$ ). There were also reductions in triglycerides in both arms, 0.67  
33 mmol/l in the low-carbohydrate arm and 0.25 in the low-fat arm, which did not approach  
34 statistical significance ( $p=0.223$ ). **Level 1+**

## 35 **Studies comparing low- or modified fat diets with reduced calorie diets**

36 In the RCT comparing the low-fat vegan diet with the ADA diet,<sup>14,20</sup> there were non-significant  
37 reductions in systolic and diastolic blood pressure in both groups. In the vegan group systolic  
38 blood pressure decreased by  $3.8\pm 12.6$  mmHg ( $p<0.05$ ) compared with baseline and in the  
39 ADA group by  $3.6\pm 13.7$  mmHg from baseline, with no significant difference between the  
40 groups ( $p=0.93$ ). Similarly the reduction in diastolic blood pressure was greater in the vegan  
41 group,  $5.1\pm 8.3$  mmHg ( $p<0.0001$ ) than in the ADA group  $3.3\pm 8.8$  mmHg ( $p<0.05$ ) although  
42 this was not different between groups ( $p=0.30$ ). **Level 1+**

43 For the entire sample, although lipid parameters decreased significantly from baseline  
44 values, there were no significant differences between groups. Among those whose lipid  
45 controlling medications remained constant (vegan  $N=39/49$ ; ADA  $N=41/50$ ), total cholesterol  
46 reduced in the vegan groups by  $33.5\pm 21.5$  mg/dl ( $p<0.0001$ ), in the ADA group by  $19.0\pm 28.5$   
47 mg/dl ( $p<0.0001$ ) and this was a significantly different between groups ( $p=0.01$ ). Reductions  
48 in HDL cholesterol were not significantly different between the groups.

- 1 Reductions in non-HDL cholesterol were significantly lower than baseline in the vegan  
2 groups  $27.6 \pm 21.1$  mg/dl ( $p < 0.0001$ ) and in the ADA group  $16.3 \pm 30.1$  mg/dl ( $p < 0.05$ ), but not  
3 significantly different between the groups ( $p = 0.05$ ).
- 4 LDL cholesterol reduced in the vegan group by  $22.6 \pm 22.0$  mg/dl ( $p < 0.0001$ ) and in the ADA  
5 group by  $10.7 \pm 23.3$  mg/dl ( $p < 0.05$ ), and was significantly different between the groups  
6 ( $p = 0.02$ ). The total-to-HDL cholesterol ratio and triglyceride concentrations fell for both  
7 groups, but there was no difference between the groups. **Level 1+**

8 **Table 4: Summarised results for blood pressure and lipid levels across RCTs**

RCTs	Follow-up	Comparison	Comparison	Blood pressure	Lipid levels
Li (2005) <sup>13</sup>	1 year	Soy-based meal replacement	Individualised diet	No changes	NS differences
Redmon (2003) <sup>17</sup>	1 year	Sibutramine + low calorie diet + meal replacement	Individualised diet	NS differences	NS differences
Daly (2006) <sup>16</sup>	3 months	Low carbohydrate diet	Reduced portion low-fat diet	NS differences	TC:HDL ratio significantly lower in carbohydrate arm
Barnard (2006) <sup>14</sup>	22 weeks	Low-fat vegan diet	Diet based on ADA guidelines	NS differences	NS differences

NS not significant

## 9 Observational studies

10 In the observational study investigating the effect of eating behaviours on weight,<sup>20</sup> changes  
11 in blood pressure or lipid profiles were not reported.

12 In the diabetes nutrition and complications trial<sup>19</sup> changes in blood pressure were reported as  
13 the proportion of patients who had a systolic blood pressure  $< 130$  mmHg, which decreased  
14 from 28.6% at baseline to 11.9% at the end of the study. Similarly in women there was a  
15 decrease from 15.8% at baseline to 8.8% after 6.5 years. The proportion of patients with a  
16 diastolic blood pressure of  $< 80$  mmHg decreased from 26.2% to 21.4% and from 31.6% to  
17 28.1% in men and women respectively.

18 In this study they reported the number of patients who were adherent to the ADA diet  
19 recommendations and were able to achieve the recommended intakes of various types of  
20 fats. They found that levels of adherence to the recommendations was low with only 26.6%  
21 of patients consuming the recommended amount of saturated fatty acids (SFAs), 13.0%  
22 consuming the recommended  $\geq 10\%$  of dietary energy from polyunsaturated fats, and 38.5%  
23 consuming the recommended  $\geq 60\%$  of dietary energy from carbohydrates and  
24 monounsaturated fats. They also estimated that 46.4% of patients consumed a ratio of  
25 polyunsaturated fatty acids (PUFAs)/SFAs  $> 0.4$  and 69% consumed a ratio of  
26 monounsaturated fats (MUFAs)/SFAs  $> 1.5$ . Patients who consumed MUFAs/SFAs  $< 1.5$  had  
27 a 3.6–4.7 times greater risk of developing diabetic complications (confidence intervals (CIs)  
28 not presented). Patients who consumed PUFAs/SFAs  $< 0.4$  were 3.4–8.2 times more at risk  
29 of developing diabetic complications. **Level 3**

### 5.1.51 Evidence to recommendations

- 2 The GDG noted that there was little new evidence to warrant any change to previous views  
3 in this field. The major consensus-based recommendations from the UK and USA emphasise  
4 sensible practical implementation of nutritional advice for people with type 2 diabetes..  
5 Management otherwise will concentrate on principles of healthy eating (essentially those for  
6 optimal cardiovascular risk protection), and reduction of high levels of free carbohydrate in  
7 foods that may cause hyperglycaemia in the presence of defective insulin secretory reserve.
- 8 If people are currently gaining weight, weight maintenance is advantageous.
- 9 The GDG noted that in some people with type 2 diabetes and weight problems it might be  
10 appropriate to consider pharmacotherapy, however this was not within the clinical questions  
11 addressed.
- 12 As with Patient Education delivery of dietary advice was noted to depend not only on specific  
13 skills, but also required all members of the diabetes care team to be familiar with local policy  
14 and thus delivering consistent advice.
- 15 Concerns continue to be noted over the promotion of 'diabetic foods' which may be low in  
16 classical sugars but high in calories and thus unsuitable as well as unnecessary for the  
17 overweight. While reduction in weight was clearly understood to be beneficial through  
18 improvements in insulin insensitivity (whether relying on endogenous or exogenous insulin),  
19 low-carbohydrate diets were noted to be of unproven safety in the long term and thus could  
20 not be endorsed. Similarly high-protein diets are acknowledged as promoting short-term  
21 weight loss, but cannot be recommended as safe in the long term.
- 22 A dietary plan for people with diabetes would follow the principles of healthy eating in the  
23 general population, and thus include carbohydrate from fruits, vegetables, wholegrains, and  
24 pulses (and thus high fibre and low glycaemic index), reduction in salt intake, the inclusion of  
25 low-fat milk and oily fish, and control of saturated and trans fatty acid intake.
- 26 The importance of advice on alcohol to the overweight and to those prone to hypoglycaemia  
27 through use of insulin secretagogues or insulin was judged important.

### 5.1.68 Recommendations and research recommendations

- 29 **8. Provide individualised and ongoing nutritional advice from a healthcare  
30 professional with specific expertise and competencies in nutrition. [2009]**
- 31 **9. Provide dietary advice in a form sensitive to the individual's needs, culture and  
32 beliefs, being sensitive to their willingness to change and the effects on their  
33 quality of life. [2009]**
- 34 **10. Emphasise advice on healthy balanced eating that is applicable to the general  
35 population when providing advice to adults with type 2 diabetes. Encourage high-  
36 fibre, low-glycaemic-index sources of carbohydrate in the diet, such as fruit,  
37 vegetables, wholegrains and pulses; include low-fat dairy products and oily fish;  
38 and control the intake of foods containing saturated and trans fatty acids. [2009]**
- 39 **11. Integrate dietary advice with a personalised diabetes management plan, including  
40 other aspects of lifestyle modification, such as increasing physical activity and  
41 losing weight. [2009]**
- 42 **12. For adults with type 2 diabetes who are overweight, set an initial body weight loss  
43 target of 5–10%. Remember that lesser degrees of weight loss may still be of**

- 1 **benefit, and that larger degrees of weight loss in the longer term will have**  
2 **advantageous metabolic impact. [2009]**
- 3 **13. Individualise recommendations for carbohydrate and alcohol intake, and meal**  
4 **patterns. Reducing the risk of hypoglycaemia should be a particular aim for a**  
5 **person using insulin or an insulin secretagogue. [2009]**
- 6 **14. Advise adults with type 2 diabetes that limited substitution of sucrose-containing**  
7 **foods for other carbohydrate in the meal plan is allowable, but that they should**  
8 **take care to avoid excess energy intake. [2009]**
- 9 **15. Discourage the use of foods marketed specifically for people with diabetes. [2009]**
- 10 **16. When adults with type 2 diabetes are admitted to hospital as inpatients or to any**  
11 **other care setting, implement a meal planning system that provides consistency**  
12 **in the carbohydrate content of meals and snacks. [2009]**

13

#### 14 **Research recommendations**

- 15 **1. What is the effectiveness of low carbohydrate diets in adults with type 2 diabetes?**

#### 16 **Why this is important**

17 Type 2 diabetes is associated with obesity, and lifestyle interventions including diet and  
18 physical activity are thought to be useful in helping to control the condition and improve  
19 patient outcomes such as reducing the risk of long-term complications and increasing  
20 quality of life. Low carbohydrate diets have been a source of discussion over the past  
21 two decades and there is much debate regarding its effectiveness and safety in  
22 controlling blood glucose levels, particularly in the longer-term. Specifically, there is little  
23 consensus on the optimal intake of daily carbohydrates, where the risk of adverse  
24 effects such as hypoglycaemia is minimised. A double-blind randomised controlled trial  
25 addressing this clinical question would help to provide a better understanding of the  
26 effects of low carbohydrate diets on diabetes control and maintenance to inform  
27 appropriate management strategies.  
28

Update 2015

## 6<sub>1</sub> Blood pressure therapy

### 6.1<sub>2</sub> Clinical introduction

3 People with type 2 diabetes are at high cardiovascular (CV) risk, high risk of diabetes eye  
4 damage, and high risk of renal disease. These adverse outcomes are known to be reduced  
5 by improved blood pressure (BP) control, which can be used to lower the risk of stroke, MI,  
6 blindness and renal failure.<sup>226</sup> Some other forms of diabetes associated microvascular  
7 damage, including peripheral nerve damage, are known to be associated with higher BP.<sup>227</sup>  
8 BP lowering is likely to be highly cost-effective in people with type 2 diabetes, more so than  
9 in the general population.

10 A number of clinical questions then face the person with diabetes and their advisors, these  
11 include:

- 12 • at what levels of BP to initiate therapy
- 13 • whether, and to what extent, those levels should be influenced by particular risk factors (in  
14 particular those involved in renal disease)
- 15 • what level of BP to aim for, and whether that should be modified by the presence of renal,  
16 eye, or macrovascular damage
- 17 • what lifestyle measures are effective and cost-effective in lowering BP
- 18 • what pharmacological interventions are effective and cost-effective in BP lowering
- 19 • how choice of agent might be modified by the presence of end organ damage.

20 Lifestyle measures (explored elsewhere) and monotherapy medication are known to have  
21 limited efficacy in lowering BP. Additional clinical questions arise over:

- 22 • the combinations of medications to be used after first-line therapy
- 23 • considerations including synergies of action, side effects of some combinations, and cost.

### 6.2<sub>4</sub> Blood pressure lowering – targets and intervention levels

#### 6.2.1<sub>5</sub> Methodological introduction

26 There were 8 papers identified as relevant to this question. These included 4 papers which  
27 further analysed data from large RCTs; 2 papers analysed data from the Irbesartan in  
28 Diabetic Nephropathy Trial (IDNT), N=1590, median follow-up 2.6 years,<sup>228</sup> and median  
29 follow-up 2.9 years.<sup>229</sup> One study analysed data from the UKPDS study,<sup>230</sup> N=1148, and a  
30 further study considered data from the Reduction of Endpoints in NIDDM with the  
31 Angiotensin II Antagonist Losartan (RENAAL) study, N=1513, median follow-up 3.4 years.<sup>231</sup>

32 Two RCTs considered the effects of intensive compared with moderate treatment, 1  
33 considered the effects of intensive treatment (valsartan) with moderate treatment (placebo)  
34 for BP control, mean follow-up <1–4 years (mean 1.9 years), N=129,<sup>232</sup> and the other, the  
35 Appropriate Blood Pressure Control in Diabetes (ABCD) trial, considered an intensive  
36 treatment with either enalapril or nisoldipine compared with moderate treatment (placebo),  
37 follow-up 5 years, N=480.<sup>233</sup>

38 A systematic review of several RCTs investigated the effects of different BP-lowering  
39 regimens on serious CV events in patients with and without diabetes.<sup>234</sup>

40 The final study was a 10 year observational study which considered a BP cut-off level for  
41 renal failure but not macrovascular complications, N=385.<sup>235</sup>

- 1 As with the papers considered for hypertension, studies which consider BP control have
- 2 flexibility in their design to allow for the introduction of further antihypertensive therapy during
- 3 the course of the study if required.

### 6.2.24 Health economic methodological introduction

- 5 No health economic papers were identified.

### 6.2.36 Evidence statements

7 Overall, an association could be established between low BP values and a lower incidence of  
8 CV events across 3 of the 4 studies looking at the relationship between BP levels and CV  
9 outcomes.<sup>229,232,233,235</sup> However, no clear BP threshold was identified as a potential  
10 therapeutic target.

11 An RCT<sup>233</sup> with a follow-up of 5 years concluded that intensive BP control (mean  
12 BP=28±0.8/75±0.3) in normotensive type 2 diabetes patients was associated with a  
13 significantly lower incidence of CV events compared with those in the moderate BP control  
14 group (mean BP=137±0.7/81±0.3). **Level 1**

15 Another RCT conducted in normotensive type 2 diabetes patients<sup>232</sup> showed non-significant  
16 differences in the incidence of CV events between the intensive blood control group (mean  
17 BP=118±10.9/75±5.7) and the moderate group (mean BP=124±10.9/80±6.5). **Level 1+**

18 The analysis completed on the IDNT data<sup>229</sup> identified a decreased risk in CV mortality and  
19 congestive heart failure (CHF) where the systolic blood pressure (SBP) decreased from >170  
20 to 120–130 mmHg, with a 20 mmHg lower SBP being associated with a 39% reduction in  
21 both. An achieved SBP ≤120 mmHg compared with >120 mmHg showed a greater risk of CV  
22 mortality and CHF (see Table 5). **Level 1+**

23 **Table 5: Post hoc analysis of the IDNT study – Berl<sup>229</sup>, N=1590**

CV Outcome	Size effect
CV mortality	A decrease in risk was observed where achieved SBP decreased from >170 to 120–130 mmHg. In this range a 20 mmHg lower SBP was associated with a 39% reduction in CV mortality, p<0.002
	An achieved SBP ≤120 showed a significantly greater risk of CV mortality compared to those with an achieved SBP >120 mmHg, RR 4.06 (2.11 to 7.80), p<0.0001
CHF	A decrease in risk was observed where achieved SBP decreased from >170 to 120–130 mmHg. In this range a 20 mmHg lower SBP was associated with a 39% reduction in CHF, p=0.001
	Those with an achieved SBP ≤120 had a significantly greater risk of CHF than those with an achieved SBP >120 mmHg, RR 1.80 (1.17 to 2.86), p=0.008
MI	A 10 mmHg lower mean achieved DBP was associated with a significantly higher risk of MI, RR 1.61 (1.28 to 2.02), p<0.0001
Stroke	A 10 mmHg lower mean achieved DBP was associated with a significantly lower risk of stroke, RR 0.65 (0.48 to 0.88), p=0.005
DBP diastolic blood pressure	

24 A systematic review<sup>234</sup> identified 27 trials which included 33,395 individuals with diabetes and  
25 125,314 without. Overall the analysis suggests that patients with diabetes achieved greater

- 1 reductions in the risk of total major CV events and CV death with regimens targeting lower  
2 BP goals<sup>f</sup> than those without diabetes (see Table 6). **Level 1+**

3 **Table 6: Systematic review – by the Blood Pressure Lowering Treatment Trialists’**  
4 **Collaboration (BPLTTC)<sup>234</sup>**

Stroke					
More vs less intensive	More intensive	Less intensive	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	63/1731	86/1868	-6.0/-4.6	0.64 (0.64 to 0.89)	NS differences
No diabetes	103/6303	204/12,080	-3.7/-3.3	0.89 (0.70 to 1.13)	NS differences
Coronary heart disease					
More vs less intensive	More intensive	Less intensive	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	63/1731	44/1868	-6.0/-4.6	0.69 (0.38 to 1.25)	NS differences
No diabetes	103/6303	31/12,080	-2.9/-3.0	1.10 (0.60 to 2.01)	NS differences
Heart failure					
More vs less intensive	More intensive	Less intensive	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	63/1731	44/1868	-6.0/-4.6	0.69 (0.38 to 1.25)	NS differences
No diabetes	103/6303	31/12,080	-3.7/-3.3	1.10 (0.60 to 2.01)	NS differences

- 5 The observational study<sup>235</sup> identified that baseline SBP was lower (141±19 mmHg) for those  
6 with no complications compared with those who had an MI (154±20 mmHg), p<0.01. SBP  
7 was also lower during the observation period for those with no complications (145±16  
8 mmHg) compared with those who had an MI (152±15 mmHg), p<0.05 and also those who  
9 had a stroke (153±15 mmHg), p<0.001. This study also noted that DBP was lower at  
10 baseline for those with no complications (84±9) compared with those who developed an MI  
11 (87±9 mmHg), p<0.05. **Level 2+**

### 6.2.3.12 Renal outcomes

- 13 Five studies<sup>228,231–233,235</sup> were identified looking at several renal outcomes and their relation  
14 with BP control. On the whole, it could be ascertained that high BP levels (SBP and/or DBP)  
15 in patients with type 2 diabetes were associated with a more rapid decline in renal function  
16 than in those with lower BP values.

### 6.2.3.27 RENAAL study

- 18 The RENAAL study<sup>231</sup> demonstrated that for SBP the baseline level of 160–179 mmHg or  
19 ≥180 mmHg compared with less than 130 mmHg had a significantly greater risk of reaching  
20 the primary end point (time to doubling of serum creatinine, end stage renal disease (ESRD)  
21 or death), risk of ESRD or death and risk of ESRD alone. Kaplan-Meier curve also showed  
22 that for those with a baseline SBP ≥140 compared with <140 mmHg there was a significantly  
23 higher risk of reaching the primary end point and risk of ESRD alone. For achieved SBP

f There were 5 studies comparing more intensive and less intensive regimes. The target BP levels (mmHg) for these studies were as follows: MAP £92 vs 102–107; DBP £75 vs £90; DBP 10 mmHg below baseline vs 80–89; DBP £80 vs £85 OR £90 and DBP <85 vs <105.

- 1 those who had a SBP of 140 to  $\geq 180$  mmHg compared with less than 130 mmHg had a  
 2 significantly greater risk of reaching the primary end point; for those with an achieved SBP of  
 3 140–159 mmHg compared with less than 130 mmHg there was a significantly greater risk of  
 4 ESRD or death and ESRD alone.
- 5 For achieved DBP those with a DBP from 90 to  $\geq 100$  mmHg compared with those with an  
 6 achieved DBP of  $< 70$  mmHg had a significantly greater risk of reaching the primary end point  
 7 (time to doubling of serum creatinine, ESRD or death), risk of ESRD or death and risk of  
 8 ESRD alone<sup>231</sup> (see Table 7 and Table 8). **Level 1+**

9 **Table 7: RENAAL study – systolic blood pressure in baseline**

SBP at baseline (mmHg)	Risk of doubling of SCr, ESRD or death (primary end point)	Risk of ESRD or death	Risk of ESRD alone
160–179 vs $< 130$	HR 1.28 (0.97 to 1.69) $p < 0.001$	HR 1.96 (1.40 to 2.74) $p < 0.001$	HR 2.13 (1.39 to 3.27) $p < 0.001$
$\geq 180$ vs $< 130$	HR 1.85 (1.33 TO 2.57) $p < 0.01^*$	HR 2.10 (1.44 to 3.06) $p < 0.01^{**}$	HR 2.02 (1.24 to 3.29) $p = 0.005^{***}$

\* Kaplan-Meier curve for baseline SBP  $< 140$  vs  $\geq 140$  mmHg, a significantly higher risk for those  $\geq 140$  mmHg (HR 1.66,  $p < 0.001$ )  
 \*\* Every 10 mmHg rise in baseline SBP increased the risk for ESRD or death by 6.7%,  $p = 0.007$  (multivariate model adjusted for urinary ACR (log scale), creatinine, albumin, haemoglobin)  
 \*\*\* Kaplan-Meier curve for baseline SBP  $< 140$  vs  $\geq 140$  mmHg, a significantly higher risk for those  $\geq 140$  mmHg (HR 1.72,  $p < 0.001$ )  
 SCr serum creatinine ratio

10 **Table 8: RENAAL study – systolic blood pressure achieved**

SBP achieved (mmHg)	Risk of doubling of SCr, ESRD or death (primary end point)	Risk of ESRD or death	Risk of ESRD alone
140–159 vs $< 130$	HR 1.49 (1.18 to 1.90) $p < 0.001$	HR 1.33 (1.02 to 1.72) $p = 0.03$	HR 1.52 (1.07 to 2.15) $p = 0.02$
90–99 vs $< 70$	HR 1.72 (1.32 to 2.23) $p < 0.001$	HR 1.55 (1.16 to 2.08) $p = 0.003$	HR 1.67 (1.15 to 2.44) $p = 0.008$
$\geq 100$ vs $< 70$	HR 2.54 (1.70 to 3.80) $p < 0.001$	HR 2.74 (1.78 to 4.24) $p < 0.001$	HR 3.26 (1.90 to 5.58) $p < 0.001$

\* Every 10 mmHg rise in baseline DBP decreased the risk for ESRD or death by 10.9% ( $p = 0.01$ ) (multivariate model adjusted for urinary ACR (log scale), creatinine, albumin, haemoglobin)

11 **Other studies reporting renal outcomes**

- 12 The 2 studies which used intensive and moderate control groups showed significant  
 13 differences between the groups only for adjusted log urinary albumin excretion rate (UAER)  
 14 findings.<sup>232,233</sup> **Level 1+**

- 15 The further analysis from the IDNT study identified that baseline BP correlated significantly  
 16 with doubling SCr or ESRD and that 36% of those with baseline SBP  $> 170$  mmHg compared  
 17 with 18% for those with baseline SBP  $< 145$  mmHg reached renal end point. Following  
 18 correction for estimated glomerular filtration rate (eGFR) and albumin:creatinine ratio (ACR)  
 19 each 20 mmHg decrease in SBP was associated with a 30% reduction in the risk of a renal  
 20 event. Though it should be noted that while there was an increasing risk for reaching a renal  
 21 end point with seated SBP, those with SBP  $< 120$  mmHg were not substantially better than  
 22 those between 120–130 mmHg.<sup>228</sup> **Level 1+**

- 1 The 10 year observational study identified that baseline SBP and DBP were significantly  
2 lower for those with no complications than those who developed renal failure, SBP was also  
3 lower for this during the observation period. A BP cut-off of >140 mmHg showed a NSx38.5  
4 increase in the risk of renal failure.<sup>235</sup> **Level 2+**

### 6.2.3.35 Retinopathy outcomes

- 6 The intensive (118±10.9/75±5.7) and moderate (124±10.9/80±6.5) groups found NS  
7 difference between the groups for progression or regression of retinopathy.<sup>232</sup> **Level 1+**
- 8 The other study which considered intensive (128±0.8/75±0.3) and moderate  
9 (137±0.7/81±0.3) groups identified less progression of retinopathy with the intensive group  
10 compared with the moderate group at both 2 years (13 vs 21%, p=0.046) and 5 years (34 vs  
11 46%, p=0.019).<sup>233</sup> **Level 1+**
- 12 The analysis completed on the data from the UKPDS study on retinopathy is detailed in the  
13 Table 9.<sup>230</sup> This considered the impact of tight blood pressure control (TBP) aiming for a BP  
14 less than 150/85 and less tight blood pressure control (LTBP) aiming for a BP of 180/105 or  
15 less. The TBP group had significantly lower microaneurysms, hard exudates and cotton wool  
16 spots than the LTBP group. This TBP group also had less retinopathy grading by the Early  
17 Treatment of Diabetic Retinopathy Study (ETDRS) grading and lower absolute risk events  
18 per 1000 patient years for photocoagulation and blindness in 1 eye. **Level 1+**
- 19 **Table 9: Retinopathy outcomes – Matthews study<sup>230</sup>**

Progression of retinopathy assessed by specific lesions	
MA % with ≥5 MA	<ul style="list-style-type: none"> <li>at 4.5 years; TBP vs LTBP (23.3% vs 33.5%) RR 0.7 (99% CI 0.51 to 0.95), p=0.003</li> <li>at 7.5 years; TBP vs LTBP (29.3% vs 44.8%) RR 0.66 (99% CI 0.48 to 0.90), p&lt;0.001</li> </ul>
Hard exudates	Overall increase 11.2% to 18.3% <ul style="list-style-type: none"> <li>at 4.5 years; TBP vs LTBP (12.5% vs 21.2%) RR 0.59 (99% CI 0.38 to 0.92), p=0.002</li> <li>at 7.5 years; TBP vs LTBP (14.1% vs 26.6%) RR 0.53 (99% CI 0.33 to 0.85), p&lt;0.001</li> </ul>
Cotton wool spots	Overall increase 14.0% to 22.4% <ul style="list-style-type: none"> <li>at 4.5 years; TBP vs LTBP (16.6% vs 17.4%) RR 0.69 (99% CI 0.47 to 1.02), p=0.02</li> <li>at 7.5 years; TBP vs LTBP (17.4% vs 32.5%) RR 0.53 (99% CI 0.35 to 0.81), p&lt;0.001</li> </ul>
Ocular end points	
Photocoagulation	<ul style="list-style-type: none"> <li>TBP vs LTBP had lower absolute risk events per 1000 patient years (11.0 vs 17.0) RR 0.63 (99% CI 0.39 to 1.07), p=0.03</li> <li>due to maculopathy, 7.6 vs 13.0 (TBP vs LTBP) RR 0.58 (99% CI 0.32 to 1.04), p=0.02</li> </ul>
Vision loss	
Blindness in 1 eye	<ul style="list-style-type: none"> <li>TBP group had lower absolute risk events per 1000 patient years than the LTBP group (3.1 vs 4.1) RR 0.76 (98% CI 0.29 to 1.99), p=0.046</li> </ul>
Retinopathy progression by ETDRS grading	<ul style="list-style-type: none"> <li>at 4.5 years 2-step or more deterioration; TBP vs LTBP (27.5% vs 36.7%) RR 0.75 (99% CI 0.50 to 0.89), p=0.02</li> <li>at 7.5 years 2-step or more deterioration; TBP vs LTBP (34.0% vs 51.3%) RR 0.66 (99% CI 0.50 to 0.89), p=0.001</li> <li>more than 1/3 (TBP) did not change compare with 1/5 (LTBP)</li> </ul>
MA microaneurysms	

#### 6.2.3.41 Nephropathy outcome

- 2 The intensive ( $118\pm 10.9/75\pm 5.7$ ) and moderate ( $124\pm 10.9/80\pm 6.5$ ) groups found NS  
3 difference between the groups for progression or regression of nephropathy.<sup>232</sup> **Level 1+**
- 4 The other study which considered intensive ( $128\pm 0.8/75\pm 0.3$ ) and moderate  
5 ( $137\pm 0.7/81\pm 0.3$ ) groups identified NS difference between the groups for progression of  
6 nephropathy.<sup>233</sup> **Level 1+**

#### 6.2.47 Evidence to recommendations

- 8 The GDG noted the problems in assigning BP lowering targets in this area, and in particular  
9 the:
- 10 • problem setting a cut-off where the evidence suggests the lower the blood pressure the  
11 better (without adverse effects)'
  - 12 • difficulties of achieving any reasonable target in some people
  - 13 • individual targets that should logically vary with individual risk
  - 14 • arbitrary dichotomy that arises immediately above and below any target level.
- 15 The results of some RCTs suggested that SBP well into the normal range (below usual target  
16 values) was both achievable and associated with benefit in people with type 2 diabetes,  
17 consistent with epidemiological evidence from other studies. In some other studies tight BP  
18 control seemed difficult to achieve, consistent with the group's clinical experience. This led  
19 the group to take a simple risk approach centered on a target level of  $<140/80$  mmHg for  
20 most people with type 2 diabetes, and  $<130/80$  mmHg for those at more particular risk. The  
21 latter group included people with raised albumin excretion rate (AER) (microalbuminuria or  
22 worse),  $eGFR < 60$  ml/min/1.73 m<sup>2</sup>, those with retinopathy, and those with prior stroke or  
23 transient ischaemic attack (TIA). The concern that more active prevention was being targeted  
24 at those who had already developed end-organ damage was recognised, but it was noted  
25 that for both microalbuminuria and early retinopathy the recommendations on annual  
26 surveillance meant that markers of damage would be detected many years before ill health  
27 ensued.

### 6.3 Blood pressure lowering medications

#### 6.3.19 Methodological introduction

- 30 The search identified a systematic review of several RCTs investigating the effects of  
31 different BP lowering therapies (that is, angiotensin-converting enzyme inhibitors (ACEI),  
32 angiotensin II receptor (A2RB) antagonists, calcium channel blockers (CCB), beta-blockers  
33 and diuretics) on serious CV events in patients with and without diabetes.<sup>234</sup>

#### 6.3.1.34 ACEI

- 35 There were 14 papers identified for this question, these included 2 Cochrane reviews,  
36 considering antihypertensive agents for preventing diabetic kidney disease<sup>236</sup> and ACEI and  
37 A2RB antagonists for preventing the progression of diabetic kidney disease.<sup>237</sup> There was  
38 also a meta-analysis which considered the effect of inhibitors of the renin-angiotensin system  
39 (RAS) and other antihypertensive drugs on renal outcomes.<sup>238</sup>

#### 40 ACEI vs placebo

- 41 Three studies compared ramipril with a placebo, they were sub-analysis of the 5-year Heart  
42 Outcomes and Prevention Evaluation (HOPE) study, considering the diabetic subgroup,

1 N=3577 (total study population, N=9297)<sup>239,240</sup> and an extension phase of 2.6 years,  
2 N=4528.<sup>241</sup>

### 3 ACEI vs A2RB

4 The DETAIL (Diabetics Exposed to Telmisartan and Enalapril) study considered telmisartan  
5 compared with enalapril over 5 years, N=250.<sup>242</sup> An open-label study considered lisinopril  
6 compared with telmisartan and compared with a combination of the 2 treatments over 52  
7 weeks, N=219.<sup>243</sup>

### 8 ACEI vs CCB

9 Three studies considered ACEI and CCB. One study considered lercanidipine compared with  
10 ramipril for 36–52 weeks, N=180.<sup>244</sup> An open-label study considered amlodipine compared  
11 with fosinopril and compared the combination of both drugs for 4 years, N=309.<sup>245</sup> A post hoc  
12 analysis of the Bergamo Nephrologic Diabetic Complications Trial (BENDICT) study was  
13 performed, this considered verapamil compared with trandopril compared with a combination  
14 of both drugs for 3.6 years, N=1204.<sup>246</sup>

### 15 ACEI vs CCB vs diuretic

16 One study considered lisinopril compared with amlodipine and chlorthalidone<sup>g</sup> with a type 2  
17 diabetes group analysis, mean follow-up 4.9 years, N=12,063 (total study population  
18 N=31,512); the Antihypertensive and Lipid-Lowering to Prevent Heart Attack Trial  
19 (ALLHAT).<sup>247</sup>

### 20 ACE1 + CCB vs ACEI + diuretic

21 One study considered verapamil + trandopril compared with enalapril + hydrochlorothiazide  
22 over 6 months, N=103.<sup>248</sup>

### 23 ACEI + CCB vs beta blocker + diuretic

24 Another study considered N=463 participants who were dosed with verapamil SR + ACE  
25 trandopril compared with atenolol + chlorthalidone for 20 weeks.<sup>249</sup>

26 All studies were either RCTs or subgroup analysis of RCTs, the majority of which were  
27 double- blinded (2 open-label studies).<sup>243,245</sup> All studies involved participants with type 2  
28 diabetes or considered a diabetic subgroup from a larger study. Many of the studies used BP  
29 target levels, if these were not achieved with the initial dose of the drug then either dose  
30 escalation or the introduction of other antihypertensive medication was allowed to ensure  
31 that target BP was maintained accordingly.

#### 6.3.1.22 A2RB

33 A total of 10 studies were found relevant to the question.<sup>237,250–258</sup>

34 The studies selected were RCTs with a follow-up of at least 6 months and with a sample size  
35 of more than 100. All studies involved participants with type 2 diabetes or considered a  
36 diabetic subgroup from a larger study. Many of the studies used BP target levels, if these  
37 were not achieved with the initial dose of the drug then either dose escalation or the

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g The ALLHAT study randomised patients to chlorthalidone 12.5–25.0 mg/day, amlodipine 2.5–10 mg/day or lisinopril 10–40 mg/day. The doses of these drugs were increased until a BP goal of <140/90 mmHg was achieved. In addition, other drugs could be added to the baseline treatments such as atenolol (25–100 mg/day), reserpine (0.1–0.2 mg/day) or clonidine (0.1–0.3 mg bid) at the discretion of the investigator. Also, hydralazine 25–100 mg bid could be added as a step three drug.

1 introduction of other antihypertensive medication was allowed to ensure that target BP was  
2 maintained according.

3 These 10 RCTs reviewed the evidence on the effectiveness and safety of A2RB blockers  
4 across several comparisons.

#### 5 **A2RB vs placebo**

6 One Cochrane review<sup>237</sup> was identified analysing data from 5 studies placebo-controlled trials  
7 that is Brenner et al. 2001 (RENAAL), Lewis et al. 2001 (Renal data – IDNT), Parving et al.  
8 2001 (IRMA), Tan et al. 2002 and, Berl et al. 2003 (CV data – IDNT).

9 Three post hoc analyses of large placebo-controlled trials were also identified: 2 post hoc  
10 studies of the RENAAL trial<sup>253,254</sup> and 1 post hoc study<sup>255</sup> of the IRMA study.

11 One post hoc analysis<sup>254</sup> analysed the impact of renal function at baseline on disease  
12 progression and response to treatment in 1513 patients who were enrolled in the RENAAL  
13 study.

14 Another post hoc analysis of the 1513 patients enrolled in the RENAAL study<sup>253</sup> analysed the  
15 effect of losartan versus placebo on long-term glycaemic control and serum potassium, uric  
16 acid, and lipid levels, as well as the relationship between these baseline metabolic factors  
17 and the composite end point (doubling of serum creatinine, ESRD, or death) or ESRD alone.

18 One post hoc analysis of the IRMA study<sup>255</sup> assessed the reversibility of kidney function  
19 changes after withdrawal of 2 years antihypertensive therapy with irbesartan on 133 type 2  
20 diabetes patients.

#### 21 **A2RB vs CCB**

22 Four studies looked at the comparison of an A2RB with a CCB. Irbesartan vs amlodipine,<sup>257</sup>  
23 valsartan vs amlodipine<sup>252,258</sup> and telmisartan vs nifedipine.<sup>251</sup> It should be noted that the  
24 study by Lewis<sup>257</sup> was included in the Cochrane review but no data on the head comparison  
25 between A2RB and CCB was reported.

#### 26 **A2RB vs sympatholytic agents**

27 One study<sup>256</sup> considered A2RB (losartan) compared with a beta-blocker agent (atenolol) and  
28 another study<sup>250</sup> compared A2RB (irbesartan) with an alpha-blocker drug (doxazosin).

29 Studies comparing ACEI with A2RB have been analysed under the ACEI section.

30 It should be noted that differing dosing and titration regimens and the differing populations  
31 included in the studies, may limit direct comparisons between studies.

### 6.3.1.32 **Beta blockers**

33 One paper was identified which considered carvedilol and metoprolol in N=1235 participants  
34 for 5 months.<sup>260</sup>

#### 35 **Beta-blockers vs CCB**

36 There were 3 papers identified for this. One paper was a sub-analysis of the Controlled  
37 Onset Verapamil Investigation of Cardiovascular End Points (CONVINCE) trial, which  
38 considered control-onset extended-release (COER) verapamil with atenolol or  
39 hydrochlorothiazide in N=16,476 (N=3239 type 2 diabetes) for 3 years.<sup>261</sup> A further paper  
40 considered a subgroup of the Anglo-Scandinavian Cardiac Outcomes Trial: Blood Pressure  
41 Lowering Arm (ASCOT: BPLA) trial, with N=19,257 (N=5145 with diabetes), which was  
42 stopped prematurely at 5.5 years.<sup>262</sup> The third paper reported on the International Verapamil-

- 1 Trandolapril Study (INVEST) trial which considered verapamil SR with atenolol for N=22,576
- 2 (N=6400 type 2 diabetes) participants over 24 months.<sup>259</sup>

### 6.3.23 Health economic methodological introduction

#### 6.3.2.14 ACEI

- 5 Three studies were identified, 2 based in the UK and 1 in Germany.
- 6 Beard et al. (2001)<sup>263</sup> and Schadlich et al. (2004)<sup>264</sup> used data from the HOPE and micro-
- 7 HOPE studies, which compared an ACEI, ramipril, to placebo. In both analyses the treatment
- 8 effects were not continued beyond the trial period of 5 years and the continued survival of
- 9 patients was considered.
- 10 Gray et al. (2001)<sup>265</sup> was based on UKPDS data, comparing an ACEI, captopril, to a beta-
- 11 blocker, atenolol. In this study a tight BP target of <150/<85 mmHg was set and other
- 12 antihypertensive treatments could be added on to achieve this target. After the trial period it
- 13 was assumed that beyond the trial period the 2 groups had identical hazard rates.
- 14 In all 3 studies the outcomes of interest were CV events.

#### 6.3.2.25 A2RB

- 16 The studies identified looked at the renal protection effect of angiotensin II receptor
- 17 antagonists (AR2B).
- 18 Three studies were based on the IDNT. Irbesartan 300 mg to amlodipine 10 mg and to a
- 19 control. All participants could take standard antihypertensive therapies which exclude ACEI,
- 20 AR2B, and CCBs. This study included type 2 diabetes patients with proteinuria. No
- 21 significant difference was found between irbesartan and amlodipine in reducing BP. The
- 22 control had an average of 3.3 mmHg increased BP.
- 23 The combined end point of the study was doubling of serum creatinine concentration, ESRD
- 24 or death from any cause. Irbesartan reduced this end point by 23% compared to amlodipine
- 25 and 20% compared to control.
- 26 Palmer et al. (2004)<sup>266</sup> was set in the UK, Rodby et al. (2003)<sup>267</sup> was set in the US, and Coyle
- 27 et al. (2004)<sup>268</sup> was set in Canada. In these studies various time horizons were used, where a
- 28 10-year time horizon was the base case, 25 years was tested in the sensitivity analysis.
- 29 Vora et al. (2005)<sup>269</sup> was based on the RENAAL study which compared losartan 50–100 mg
- 30 with a regimen of conventional antihypertensive treatment (CCBs, diuretics, alpha-blockers,
- 31 beta-blockers, and centrally acting agents). Patients had type 2 diabetes and nephropathy.
- 32 The same combined end point as the IDNT was used. Losartan was found to reduce this by
- 33 25% compared with control. This analysis was set in the UK and a lifetime time horizon was
- 34 used.
- 35 Smith et al. (2004)<sup>270</sup> was based on the Microalbuminuria Reduction with Valsartan
- 36 (MARVAL) study comparing the AR2B, to the CCB amlodipine. Patients with type 2 diabetes
- 37 and microalbuminuria were included. The study found that valsartan significantly reduced
- 38 urinary excretion rate compared to amlodipine. Similar reductions in BP were found. This
- 39 analysis was set in the US. An 8-year time horizon was used.

### 6.3.30 Evidence statements

- 41 A systematic review showed that for the outcome stroke, there was no evidence of
- 42 differences in the effects of the treatment regimens between patients with and without
- 43 diabetes except in the comparison that included A2RB-based regimens. In this comparison,

1 A2RB provided lesser protection to patients with diabetes compared with those without  
2 diabetes (see Table 10).<sup>234</sup>

3 For the outcomes coronary heart disease (CHD) and heart failure, the review did not show  
4 differences between patients with and without diabetes for any comparison, again except for  
5 the comparison that included A2RB. Diabetic patients treated with A2RB experienced a  
6 significantly greater protection compared to those without diabetes for the outcome heart  
7 failure.<sup>234</sup>

8 According to their review, there was also some evidence of a difference between the 2  
9 patient groups in protection against CV death and total mortality favouring patients with  
10 diabetes in the comparison of ACEI-based regimens vs placebo (see Table 13).<sup>234</sup>

11 **Table 10: Stroke – systematic review by the BPLTTC<sup>234</sup>**

ACEI	ACE	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	125/2378	174/2336	-3.6/-1.9	0.69 (0.55 to 0.86)	NS differences
No diabetes	347/6733	485/6782	-5.8/-2.7	0.73 (0.62 to 0.85)	
CCB	CCB	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	21/911	45/900	-6.3/-3.0	0.47 (0.28 to 0.78)	NS differences
No diabetes	52/2883	72/2788	-9.2/-3.7	0.70 (0.49 to 0.99)	
A2RB	ARB-based regimen	Control regimen	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	143/2226	173/2793	-2.1/-0.9	0.96 (0.77 to 1.19)	p=0.05 by X2 test of homogeneity
No diabetes	253/6186	342/6153	-1.4/-0.6	0.74 (0.63 to 0.86)	

12 **Table 11: Coronary heart disease – systematic review by the BPLTTC<sup>234</sup>**

ACEI	ACE	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	96/2378	105/2336	-3.6/-1.9	0.88 (0.67 to 1.16)	NS differences
No diabetes	123/6733	164/6782	-5.8/-2.7	0.78 (0.62 to 0.98)	
CCB	CCB	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	94/868	75/858	-6.3/-3.0	1.29 (0.97 to 1.72)	NS differences
No diabetes	10/2514	13/2416	-9.2/-3.7	1.07 (0.43 to 2.62)	
ARB	ARB-based regimen	Control regimen	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	150/2226	208/2793	-2.1/-0.9	0.92 (0.72 to 1.17)	NS differences

ACEI	ACE	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
No diabetes	285/6186	269/6153	-1.4/-0.6	1.05 (0.89 to 1.24)	

1 **Table 12: Heart failure – systematic review by the BPLTTC<sup>234</sup>**

ACEI	ACE	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	96/2378	105/2336	-3.6/-1.9	0.88 (0.67 to 1.16)	NS differences
No diabetes	123/6733	164/6782	-5.8/-2.7	0.78 (0.62 to 0.98)	
CCB	CCB	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	94/868	75/858	-6.3/-3.0	1.29 (0.97 to 1.72)	NS differences
No diabetes	10/2514	13/2416	-9.2/-3.7	1.07 (0.43 to 2.62)	
ARB	ARB-based regimen	Control regimen	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	150/2226	208/2793	-2.1/-0.9	0.92 (0.72 to 1.17)	NS differences
No diabetes	285/6186	269/6153	-1.4/-0.6	1.05 (0.89 to 1.24)	

2 **Table 13: CV Deaths – systematic review by the BPLTTC<sup>234</sup>**

ACEI	ACE	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	145/2378	211/2336	-3.6/-1.9	0.67 (0.55 to 0.82)	p=0.05 X2 test of homogeneity
No diabetes	330/6733	389/6782	-5.8/-2.7	0.86 (0.75 to 0.99)	
CCB	CCB	Placebo	⊗ BP mmHg	RR 95% CI	Diabetes vs no diabetes
Diabetes	42/868	62/858	-5.9/-3.1	0.54 (0.21 to 1.42)	NS differences
No diabetes	61/2514	73/2416	-9.3/-3.9	0.64 (0.24 to 1.68)	

3

4 Finally, the review did not report significant differences between different BP lowering  
5 regimens (that is, head-to-head comparisons) in terms of stroke, CHD, heart failure in  
6 patients with diabetes. The exception being CCBs, which were associated with a higher risk  
7 of heart failure when they were compared with diuretics or beta-blockers,<sup>234</sup> (see Table 14,  
8 Table 15 and Table 16). In the same way, no differences were seen in the head-to-head  
9 comparisons for total major CV events, CV deaths, and total mortality in patients with  
10 diabetes.

11 **Table 14: Head-to-head comparisons. Stroke – systematic review by the BPLTTC<sup>234</sup>**

ACE vs D/BB	ACE	D/BB	⊗ BP mmHg	RR 95% CI
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ACE vs D/BB	ACE	D/BB	⊗ BP mmHg	RR 95% CI
5 studies	282/4385	405/6614	2.2/0.3	1.02 (0.88 to 1.19)
CCB vs D/BB	CCB	D/BB	⊗ BP mmHg	RR 95% CI
8 studies	279/6276	427/8550	0.7/-0.8	0.94 (0.81 to 1.09)
ACE vs CCB	ACE	CCB	⊗ BP mmHg	RR 95% CI
5 studies	246/4101	227/4222	1.6/1.2	1.09 (0.88 to 1.36)

BB, beta-blocker, D, diuretics

1 **Table 15: Head-to-head comparisons. CHD – systematic review by the BPLTTC<sup>234</sup>**

ACE vs D/BB	ACE	D/BB	⊗ BP mmHg	RR 95% CI
5 studies	402/4385	623/6614	2.2/0	3 0.83 (0.62 to 1.12)
CCB vs D/BB	CCB	D/BB	⊗ BP mmHg	RR 95% CI
8 studies	431/6276	638/8550	0.7/-0.8	1.00 (0.89 to 1.13)
ACE vs CCB	ACE	CCB	⊗ BP mmHg	RR 95% CI
5 studies	358/4101	407/4222	1.6/1.2	0.76 (0.51 to 1.12)

2 **Table 16: Head-to-head comparisons. Heart failure – systematic review by the**  
3 **BPLTTC<sup>234</sup>**

ACE vs D/BB	ACE	D/BB	⊗ BP mmHg	RR 95% CI
4 studies	251/4076	384/6351	2.5/0.4	0.94 (0.55 to 1.59)
CCB vs D/BB	CCB	D/BB	⊗ BP mmHg	RR 95% CI
6 studies	337/5276	399/7521	0.5/-0.8	1.27 (1.01 to 1.61)
ACE vs CCB	ACE	CCB	⊗ BP mmHg	RR 95% CI
5 studies	263/4101	325/4222	1.6/1.2	0.92 (0.67 to 1.27)

### 6.3.3.14 ACEI

5 Overall, the evidence appraised showed no significant differences in terms of CV outcomes  
6 when treatment with ACEI was compared with other antihypertensive therapies or with  
7 placebo. ACEI also failed to demonstrate superiority over other agents on the basis of BP  
8 lowering power (unless combination therapy is compared with monotherapy). However, the  
9 evidence suggested that treatment with ACEI is related to greater benefits in terms of renal  
10 outcomes in patients with type 2 diabetes as compared with other BP lowering agents.

#### 6.3.3.1.11 Cardiovascular outcomes

##### 12 All-cause mortality

13 The Cochrane review on antihypertensives for preventing diabetic kidney disease found NS  
14 difference for ACEI vs placebo (3 trials, N=2683) and for ACEI vs CCBs (6 trials, N=1286).<sup>236</sup>  
15 These findings were supported by the Cochrane review on ACEI and A2RB for preventing

- 1 the progression of diabetic kidney disease for ACEI vs placebo (21 trials, N=7295)<sup>h</sup> and
- 2 ACEI vs A2RB (5 studies, N=3409).<sup>237</sup> **Level 1++**

### 3 **ACEI vs CCB vs diuretic**

- 4 The diabetes ALLHAT analysis showed NS difference between the treatments for the
- 5 incidence of total mortality.<sup>247</sup> **Level 1+**

#### 6.3.3.1.26 **Major cardiovascular events**

##### 7 **ACEI/placebo**

- 8 The extension phase of the HOPE study showed a NS trend towards reduction in major CV
- 9 events and risk of MI, with ramipril, stroke and CV death as NS. At follow-up of the study and
- 10 extension there was a significant risk reduction with ramipril for the outcomes of MI, stroke
- 11 and CV death.<sup>241</sup> **Level 1+**

##### 12 **ACEI vs CCB vs diuretic**

- 13 The diabetes analysis of ALLHAT identified NS difference in the incidence of fatal CHD and
- 14 non-fatal MI for lisinopril vs chlorthalidone in any of the 3 glycaemic strata that were analysed
- 15 diabetes mellitus, impaired fasting glucose and normoglycaemia. This was also evident for
- 16 diabetes mellitus and normoglycaemia for amlodipine vs chlorthalidone.<sup>247</sup> **Level 1+**

#### 6.3.3.1.37 **Blood pressure**

- 18 BP reduction with all hypertensive treatments was a consistent feature of the studies and
- 19 therefore only studies where there were significant differences between the treatments will
- 20 be highlighted.

##### 21 **ACEI/A2RB**

- 22 At the 52-week follow-up point, the combination of lisinopril and telmisartan showed
- 23 significantly greater reductions in both SBP and DBP than the individual monotherapies
- 24 (p=0.003 for both SBP and DBP).<sup>243</sup> **Level 1+**

##### 25 **ACEI/CCB + diuretic**

- 26 Similarly, the combination of amlodipine and fosinopril showed a reduction in sitting BP of
- 27 28.7/17.1 compared with 17.2/11.8 (fosinopril, p<0.01) and 19.9/12.8 (amlodipine, p<0.01).<sup>245</sup>
- 28 **Level 1+**

##### 29 **ACEI + CCB/beta-blocker + diuretic**

- 30 The study which compared verapamil + trandopril with atenolol + chlorthalidone identified
- 31 that while both treatments significantly reduced BP that comparison between the groups
- 32 showed a difference of 4.85 mmHg SBP (1.94 to 7.76, p=0.0011) and 1.79 mmHg DBP (0.26
- 33 to 3.32, p=0.0222) favouring atenolol + chlorthalidone.<sup>249</sup> **Level 1++**

##### 34 **ACEI/CCB**

- 35 A post hoc analysis of the BENEDICT<sup>246</sup> study considered the impact on BP control and
- 36 ACEI therapy on new-onset microalbuminuria. Baseline SBP, DBP, mean arterial pressure
- 37 (MAP) and pulse pressure did not predict the onset of microalbuminuria. Participants who

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h Though a subgroup analysis which used ACE at maximum tolerable dose did find a significant decrease vs placebo (5 trials; N=2034, RR 0.78, 0.61 to 0.98).

1 developed microalbuminuria had significantly lower reductions in SBP than those who did not  
2 develop microalbuminuria ( $7.9 \pm 11.5$  vs  $10.6 \pm 11.9$ ,  $p < 0.05$ ). This study also identified that  
3 those with follow-up BP below the medians or with BP reduction above the medians were  
4 more frequently on ACE therapy (particularly *trandopiril + verapamil*) and less frequently on  
5 concomitant treatment with diuretics, beta-blockers or CCBs.<sup>246</sup> **Level 1+**

### 6.3.3.1.46 Renal outcomes

7 The Cochrane review, ACEI and A2RB antagonists for preventing the progression of diabetic  
8 kidney disease, identified ACE compared with placebo reduced the progression from micro-  
9 to macroalbuminuria, increased the regression from micro- to normoalbuminuria, and  
10 reduced the risk of ESRD.<sup>237</sup>

11 The Cochrane review, antihypertensive agents for preventing diabetic kidney disease,  
12 identified that ACEI compared with placebo/no treatment reduced the development of  
13 microalbuminuria, and ACEI compared with CCB reduced the risk of developing kidney  
14 disease.<sup>236</sup>

15 The meta-analysis identified that an ACEI or A2RB compared with other treatments only  
16 showed significant reduction in UAER.<sup>238</sup>

17 The HOPE study identified that ramipril compared with placebo reduced the risk of new  
18 microalbuminuria and that both new microalbuminuria and progression of proteinuria was  
19 higher for the diabetic group than the non-diabetic group.<sup>240</sup>

### 20 Combination compared with monotherapy

21 The combination of lisinopril and telmisartan identified higher reduction with AER compared  
22 with the monotherapies.<sup>243</sup>

23 The combination of fosinopril + amlodipine reduced UAE compared with amlodipine  
24 monotherapy (all time points) and with fosinopril monotherapy (after 18 months).<sup>245</sup>

25 Renal outcomes are detailed in the Table 17, including study results which identified NS  
26 difference between treatments.

### 27 Table 17: ACEI – renal outcomes

Progression of proteinuria	
HOPE study <sup>240</sup> ACEI/placebo Level 1+	ACEI/placebo Progression higher with non-diabetic participants than diabetic (34% vs 17%, $p < 0.01$ ) Diabetes was the factor most strongly associated with the progression of proteinuria (OR 2.45, 2.148 to 2.75, $p < 0.05$ )* Ramipril vs placebo NS (adjustment for baseline reduced proteinuria by 22%, $p = 0.0495$ )
New microalbuminuria/risk of developing microalbuminuria	
Cochrane review <sup>236</sup> Level 1++	ACEI vs placebo/no treatment, reduced development of microalbuminuria (6 trials, N=3480, RR 0.58, 0.40 to 0.84) ACEI vs CCB reduced the risk of developing kidney disease (micro- or macroalbuminuria) (4 trials, N=1210, RR 0.58, 0.40 to 0.84) ACEI vs beta-blockers NS difference
Cochrane review <sup>237</sup> Level 1++	ACE vs placebo/no treatment significantly reduced the progression from micro- to macroalbuminuria (17 trials, N=2036, RR 0.49, 0.29 to 0.69) ACEI vs A2RB NS difference
HOPE study <sup>240</sup> Level 1+	ACEI/placebo New microalbuminuria was higher in diabetic than in non-diabetic participants (38.2% vs 18.1%)

<b>Progression of proteinuria</b>	
	Ramipril reduced the risk of new microalbuminuria by 10% p=0.046 vs placebo, in those with diabetes
<b>Regression from micro- to normoalbuminuria</b>	
Cochrane review <sup>237</sup> Level 1++	ACEI vs placebo/no treatment ACEI significantly increased regression (16 studies, N=1910, RR 3.06, 1.76 to 5.35) ACEI vs A2RB NS difference
Dalla (2004) <sup>244</sup> Level 1+	ACEI/CCB Ramipril vs lercanidipine NS for those who reverted to normoalbuminuria
Fogari (2002) <sup>245</sup> Level 1+	At 48 months 46% (fosinopril), 33% (amlodipine) and 67% (combination fosinopril + amlodipine) had moved to non-microalbuminuric status
<b>Doubling of creatinine</b>	
Cochrane review <sup>236</sup> Level 1++	ACEI vs placebo NS difference
Meta-analysis <sup>238</sup> Level 1+	ACEI or A2RB vs other active interventions NS, those with diabetes (6 trials, N=3044) and NS those without diabetes
<b>Serum creatinine</b>	
Meta-analysis <sup>238</sup> Level 1+	ACEI or A2RB vs other treatments NS, those with diabetes (18 trials, N=4615), those without diabetes, small reduction
HOPE study <sup>240</sup> Level 1+	ACEI/placebo No evidence of effect on ramipril on serum creatinine levels
Barnett (2004) <sup>242</sup> Level 1+	ACEI/A2RB Enalapril vs telmisartan NS difference
<b>GFR</b>	
Meta-analysis <sup>238</sup> Level 1+	ACEI or A2RB vs other treatments NS, those with diabetes (37 studies, N=15,742), NS those without diabetes
HOPE study <sup>240</sup> Level 1+	ACEI/placebo Ramipril vs placebo NS difference
Barnett (2004) <sup>242</sup> Level 1+	ACEI/A2RB Mean change in GFR: the lower treatment boundary in favour of enalapril was -7.6, greater than the pre-defined level of -10.0 indicating no difference between the treatments Enalapril vs telmisartan NS difference in annual decreases in GFR
<b>AER</b>	
Dalla (2004) <sup>244</sup> Level 1+	ACEI/CCB Ramipril vs lercanidipine NS difference Proportion of participants with reduction >50% was 22.2% with ramipril and 34.2% lercanidipine
Sengul (2006) <sup>243</sup> Level 1	ACEI/A2RB Lisinopril vs telmisartan NS difference Combination of lisinopril + telmisartan vs monotherapies AER reduction was significantly higher (p<0.001)
<b>ESRD</b>	
Cochrane review <sup>237</sup> Level 1++	ACEI vs placebo/no treatment reduction in the risk of ESRD (10 studies, N=6819, RR 0.68, 0.39 to 0.93)
Meta-analysis <sup>238</sup> Level 1+	ACEI or ARB vs other treatments, NS reduction in ESRD occurrence, those with diabetes (4 trials, N=14,437), those without diabetes there was a reduction with ACE or A2RB
Meta-analysis <sup>238</sup> Level 1+	ACEI or A2RB vs other treatments showed a reduction in UAER for those with diabetes, (34 trials, N=4772, RR -12.21, -21.68 to -2.74), for those without diabetes (44 trials, N=5266, RR -15.73, -24.75 to -6.74, p=0.001)

Progression of proteinuria	
Fogari (2002) <sup>245</sup> Level 1+	ACEI/CCB Combination of fosinopril + amlodipine showed significantly greater reduction vs amlodipine monotherapy at any time and vs fosinopril from 18 months onwards
Barnett (2004) <sup>242</sup> Level 1+	ACEI/A2RB Enalapril vs telmisartan, annual changes were small with large CI in both groups. % changes were NS difference
* The association with smoking, hypertension, male gender and peripheral vascular disease was less strong GFR, glomerular filtration rate	

### 6.3.3.1.51 Metabolic outcomes

#### 2 Risk of diabetes

3 The extended HOPE trial identified that at the end of the extension phase there was a  
4 significant further reduction in risk for diabetes for ramipril vs placebo (2.7% vs 4.0%, RR  
5 0.66, 0.46 to 0.95).<sup>241</sup> **Level 1+**

#### 6 HbA1c and glycaemic control

7 The study which considered fosinopril and amlodipine monotherapy, and in combination,  
8 found that HbA1c was NS changed by any treatments and body weight remained  
9 unchanged.<sup>245</sup> **Level 1+**

10 The study which compared verapamil SR + trandopril and atenolol + chlorthalidone found  
11 that HbA1c remained stable with verapamil SR + trandopril but increased with atenolol +  
12 chlorthalidone 7.8 (1.26) at baseline and 8.6 (1.77) at last visit, treatment difference,  
13  $p=0.0001$ ; fasting glucose and fructosamine treatment difference,  $p=0.0001$ .<sup>249</sup>

14 Similarly, fasting glucose and fructosamine remained stable with verapamil SR + trandopril  
15 but increased with atenolol + chlorthalidone, treatment difference  $p=0.0001$ .<sup>249</sup> **Level 1++**

16 The study which considered verapamil + trandopril vs enalapril + hydrochlorothiazide  
17 identified that HbA1c remained stable with verapamil + trandopril but increased with enalapril  
18 + hydrochlorothiazide (baseline  $5.96 \pm 1.25\%$  to final  $6.41 \pm 1.51\%$ ), difference between  
19 groups,  $p=0.040$ .<sup>248</sup> Crude blood glucose changes were  $23 \pm 69$  mg/dl for verapamil +  
20 trandopril (16.8% reduction) and  $1 \pm 32$  mg/dl (0.8% reduction) with enalapril +  
21 hydrochlorothiazide. The percentage of participants with glycaemic control ( $<126$  mg/dl)  
22 increased from 50% to 72% with verapamil + trandopril, but did not change with enalapril +  
23 hydrochlorothiazide.<sup>248</sup> **Level 1++**

### 6.3.3.1.54 Adverse events

25 Both Cochrane reviews identified an increased risk of cough with ACE vs placebo/no  
26 treatment (4 trials,  $N=3725$ , RR 1.79, 1.19 to 2.69),<sup>236</sup> (10 trials,  $N=7087$ , RR 3.17, 2.29 to  
27 4.38).<sup>237</sup> **Level 1++**

28 Throughout the other studies the incidence of discontinuation due to AEs was small and the  
29 AEs reported were mainly; progression of diabetes, unsatisfactory therapeutic response,  
30 hypotension, ankle oedema, tachycardia, headache, cough, nausea, stomach upset,  
31 respiratory infection, and dizziness. **Level 1+**

### 6.3.3.21 A2RB

2 In summary, A2RB therapy was associated with greater benefits for type 2 diabetes patients  
3 in terms of renal outcomes (e.g. progression to ESRD, doubling of serum creatinine,  
4 proteinuria) than treatment with placebo, CCB or sympatholytic agents. In addition, treatment  
5 with A2RB was also associated with a better metabolic and BP profile than sympatholytic  
6 therapy but non- significant differences were observed over those treated with CCB.

#### 7 A2RB vs placebo

### 6.3.3.2.18 Cardiovascular outcomes

#### 9 All-cause mortality

10 A Cochrane review<sup>237</sup> did not find a statistically significant reduction in the risk of all-cause  
11 mortality in the 5 studies (3409 patients) of A2RB vs placebo/no treatment. RR 0.99, 95% CI  
12 0.85 to 1.17. **Level 1++**

#### 13 Hospitalisations for heart failure

14 A post hoc analysis<sup>254</sup> compared the incidence of hospitalisation for heart failure within 3  
15 tertiles of baseline serum creatinine concentration (highest, 2.1 to 3.6 mg/dl; middle, 1.6 to  
16 2.0 mg/dl; lowest, 0.9 to 1.6 mg/dl). The study reported that the crude incidence of first  
17 hospitalisations for heart failure was higher in the highest (16.4%) and middle (15.0%) tertiles  
18 than in the lowest (11.1%) tertile (trend test across tertiles,  $p=0.02$ ).

19 The study concluded that losartan decreased the hospitalisations for heart failure by 50.2  
20 and 45.1, in the highest and middle tertile, respectively but was associated with a non-  
21 significant increased risk (42.5%) of hospitalisations in the lowest tertile. **Level 1+**

### 6.3.3.2.22 Renal outcomes

#### 23 Progression to ESRD

24 A Cochrane review<sup>237</sup> found a significant reduction in the risk of ESRD with A2RB compared  
25 to placebo/no treatment (3 studies,  $N=3251$ ): RR 0.78, 95% CI 0.67 to 0.91. **Level 1++**

26 A post hoc analysis<sup>254</sup> compared the incidence of ESRD within 3 tertiles of baseline serum  
27 creatinine concentration (highest, 2.1 to 3.6 mg/dl; middle, 1.6 to 2.0 mg/dl; lowest, 0.9 to 1.6  
28 mg/dl). The study reported that the observed crude incidence of ESRD was significantly  
29 higher in the highest (40.5%) and middle (19.3%) tertiles as compared with the lowest (7.3%)  
30 tertile (trend test across tertiles,  $p<0.0001$ ).

31 The study concluded that losartan decreased the risk of ESRD by 24.6, 26.3, and 35.3% in  
32 highest, middle, and lowest tertiles respectively. **Level 1+**

#### 33 Doubling of serum creatinine

34 A Cochrane review<sup>237</sup> found a significant reduction in the risk of doubling of serum creatinine  
35 concentration with A2RB compared to placebo/no treatment (3 studies, 3251 patients): RR  
36 0.79, 95% CI 0.67 to 0.93. **Level 1++**

#### 37 Progression from micro- to macroalbuminuria

38 A Cochrane review<sup>237</sup> showed that the use of A2RB versus placebo/no treatment was also  
39 associated with a significant reduction in the risk of progression from micro- to  
40 macroalbuminuria (3 studies, 761 patients); RR 0.45, 95% CI 0.32 to 0.75. **Level 1++**

## 1 Regression from micro- to normoalbuminuria

2 A Cochrane review<sup>237</sup> found a significant increase in regression from micro- to normo-  
3 albuminuria with A2RB versus placebo/no treatment (16 studies, 1910 patients) RR 1.42,  
4 95% CI 1.05 to 1.93. **Level 1++**

## 5 Proteinuria

6 A post hoc analysis<sup>254</sup> compared the median proteinuria reduction (%) within 3 tertiles of  
7 baseline serum creatinine concentration (highest, 2.1 to 3.6 mg/dl; middle, 1.6 to 2.0 mg/dl;  
8 lowest, 0.9 to 1.6 mg/dl). The study showed a significantly ( $p < 0.0001$ ) greater median  
9 percentage proteinuria reduction (versus baseline) on losartan than on placebo in the highest  
10 (24 vs -8%), middle (16 vs -8%), and lowest (15 vs -10%) tertiles respectively. **Level 1+**

11 A post hoc analysis of the IRMA study<sup>255</sup> reported that after 2 years of follow-up UAER  
12 decreased by 34% (95% CI 8 to 53), and 60% (95% CI 46 to 70) in the irbesartan 150 mg  
13 and irbesartan 300 mg groups respectively ( $p < 0.05$  vs baseline). No significant reductions in  
14 UAER were found in patients receiving placebo.

15 One month after withdrawal of irbesartan therapy, the same post hoc analysis<sup>255</sup> found no  
16 significant increases in UAER in patients receiving placebo or irbesartan 150 mg when  
17 compared with baseline values. However, the study reported that UAER remained  
18 persistently reduced by 47% (95% CI 24 to 63) in the irbesartan 300 mg group ( $p < 0.05$  vs  
19 baseline). This persistent reduction in the irbesartan 300 mg group, as compared with  
20 baseline, was highly significantly different from irbesartan 150 mg ( $p < 0.01$ ). This difference  
21 occurred although the regain in GFR between the 2 irbesartan groups were nearly identical.  
22 **Level 1+**

### 6.3.3.2.23 Blood pressure

24 A post hoc analysis of the IRMA study<sup>255</sup> found that after 2 years of treatment there were no  
25 significant differences in mean arterial blood pressure between patients treated with placebo  
26 or irbesartan (150 or 300 mg). However, 1 month after withdrawal of irbesartan therapy mean  
27 arterial blood pressure was unchanged in the placebo group, but increased significantly in  
28 the irbesartan groups to  $109 \pm 2$  and  $108 \pm 2$  in the 150 mg and 300 mg groups respectively  
29 ( $p < 0.01$ ). **Level 1+**

### 6.3.3.2.30 Metabolic outcomes

31 A post hoc analysis of the RENAAL study<sup>253</sup> found no significant differences between  
32 patients treated with losartan or placebo in terms of glycaemic levels, lipid profile or serum  
33 uric acid after 3.4 years of follow-up. **Level 1+**

### 6.3.3.2.54 Adverse events

35 A Cochrane review<sup>237</sup> found a significant increase in the risk of hyperkalaemia with A2RB  
36 compared to placebo/no treatment (2 studies, 194 patients); RR 4.93, 95% CI 1.87 to 15.65.  
37 A2RB were not found to be associated with an increased risk of cough compared to  
38 placebo/no treatment. **Level 1++**

### 6.3.3.39 A2RB vs CCB

### 6.3.3.3.40 Cardiovascular and renal outcomes

41 One RCT<sup>257</sup> with a follow-up of 2.6 years, found that treatment with irbesartan significantly  
42 reduced the risk of doubling serum creatinine concentration, development of ESRD, or death  
43 from any cause, by 23% compared to the amlodipine therapy ( $p = 0.006$ ). **Level 1++**

44 When individual end points were analysed the RCT<sup>257</sup> reported:

- 1 • A significantly lower risk of a doubling in the serum creatinine concentration in patients  
2 receiving irbesartan compared to amlodipine-treated patients (37% lower in the irbesartan  
3 group than in the amlodipine group,  $p < 0.001$ ).
- 4 • Non-significant differences in terms of progression to ESRD between irbesartan-treated  
5 patients and those receiving amlodipine (risk 23% lower in the irbesartan group  $p = 0.07$ ).
- 6 • Non-significant difference in the rates of death from any cause between patients treated  
7 with irbesartan and those treated with amlodipine. **Level 1++**
- 8 The same study<sup>257</sup> did not find a significant benefit associated with irbesartan as compared  
9 with amlodipine in reducing the secondary composite end point of death from CV causes,  
10 non-fatal MI, heart failure resulting in hospitalisation, a permanent neurologic deficit caused  
11 by a cerebrovascular event, or lower limb amputation above the ankle. **Level 1++**
- 12 An RCT<sup>258</sup> comparing therapy with valsartan and amlodipine reported results for a pre-  
13 specified subgroup of type 2 diabetes patients and found non-significant differences between  
14 the 2 treatment arms for the primary composite cardiac outcome which looked at cardiac  
15 mortality and morbidity.\*<sup>i</sup> **Level 1+**
- 16 Another RCT<sup>252</sup> which also compared treatment with valsartan and amlodipine, found that  
17 after 24 weeks there was a significant reduction in UAER in patients receiving valsartan as  
18 compared with those treated with amlodipine ( $p < 0.001$ ; 95% CI for ratio, 0.520 to 0.710). The  
19 UAER at 24 weeks with valsartan was 56% (95% CI, 49.6 to 63.0) of baseline, equivalent to  
20 a 44% reduction. The UAER for amlodipine at week 24 was 92% (95% CI, 81.7 to 103.7) of  
21 baseline, a reduction of only 8%. **Level 1++**
- 22 The same RCT<sup>252</sup> showed a significantly greater percentage of patients returning to normo-  
23 albuminuria status by week 24 with valsartan (29.9%) than with amlodipine (14.5%).  
24 Treatment difference 15.4%, 95% CI, 5.6 to 25.8,  $p < 0.001$ . **Level 1++**

#### 6.3.3.3.25 *Blood pressure*

- 26 One RCT<sup>257</sup> did not find significant differences in mean arterial pressure in patients treated  
27 with irbesartan and amlodipine after 2.6 years of follow-up. **Level 1++**

#### 6.3.3.3.38 *Metabolic outcomes*

- 29 One RCT<sup>251</sup> reported that at 12 months there were no significant changes from baseline in  
30 HbA1c, FPG, BMI, triglycerides and high-density lipoprotein cholesterol (HDL-C) in patients  
31 treated with telmisartan or nifedipine gastrointestinal therapeutic system (nifedipine GITS)  
32 and there were no significant differences in any of these parameters between treatments.  
33 **Level 1+**
- 34 The same RCT<sup>251</sup> showed that reduction in total cholesterol and low-density lipoprotein with  
35 telmisartan were significantly greater than those with nifedipine GITS ( $p < 0.05$ ). **Level 1+**

#### 6.3.3.3.36 *Adverse events*

- 37 One RCT<sup>257</sup> reported that the incidence of hyperkalaemia (necessitating discontinuation of  
38 the study medication) was significantly higher in patients receiving irbesartan as compared to  
39 those receiving amlodipine. **Level 1++**
- 40 One RCT<sup>252</sup> found that ankle oedema occurred significantly less frequently in valsartan-  
41 treated patients compared to those treated with amlodipine (1.2% vs 7.4%, difference -6.2%,  
42 95% CI -12.9% to -0.4%,  $p < 0.006$ ). **Level 1+**

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i The primary end point was time to first cardiac event (a composite of sudden cardiac death, fatal MI, death during or after percutaneous coronary intervention or coronary artery bypass graft, death as result of heart failure, and death associated with recent MI at autopsy, heart failure requiring hospital management, non-fatal MI, or emergency procedures to prevent MI).

#### 6.3.3.41 A2RB vs sympatholytic agents

##### 6.3.3.4.12 Cardiovascular outcomes

3 One RCT<sup>256</sup> with a follow-up of 4.7 years found that treatment with losartan significantly  
4 reduced the risk of CV death, stroke, or MI compared to atenolol therapy. RR 0.76 (95% CI  
5 0.58 to 0.98), p=0.031. **Level 1++**

6 When individual end points were analysed the RCT<sup>256</sup> reported:

- 7 • a statistically significant reduction in the risk of all-cause mortality in losartan-treated  
8 patients compared to those receiving atenolol. RR 0.61 (95% CI 0.45 to 0.84), p=0.002
- 9 • a statistically significant reduction in the risk of CV death favouring the losartan group. RR  
10 0.63 (95% CI 0.42 to 0.95), p=0.028
- 11 • a non-significant difference in the incidence of stroke or MI between patients treated with  
12 losartan and those treated with atenolol.

##### 6.3.3.4.23 Blood pressure<sup>j</sup>

14 One RCT<sup>250</sup> found that after 12 months, patients treated with irbesartan had significantly  
15 lower SBP and DBP levels as compared to those receiving doxazosin, (p<0.05). **Level 1+**

##### 6.3.3.4.36 Metabolic outcomes

17 One RCT<sup>250</sup> found significantly lower HbA1c levels in doxazosin-treated patients as  
18 compared to patients receiving irbesartan after 12 months of follow-up. **Level 1+**

19 The same RCT<sup>250</sup> found that patients treated with doxazosin had significantly higher levels of  
20 HDL-C as compared to those treated with irbesartan (p<0.05). **Level 1+**

##### 6.3.3.4.41 Adverse events

22 One RCT<sup>256</sup> showed that albuminuria was reported less frequently (p=0.002) as an AE in the  
23 losartan than in the atenolol group (losartan 7% vs atenolol 13%). **Level 1++**

24 The same RCT<sup>256</sup> found that chest pain was more frequently reported in the losartan arm  
25 (p=0.036) (losartan 2% vs atenolol 8%). **Level 1++**

##### 6.3.3.5.26 Beta-blockers

27 The evidence appraised suggested that treatment with beta-blockers in patients with type 2  
28 diabetes failed to demonstrate a better CV profile when compared with CCB therapy.  
29 Furthermore a landmark RCT showed a significant reduction in the incidence of CV  
30 outcomes in patients receiving CCB as compared with those treated with beta-blockers. In  
31 terms of BP control, the evidence did not demonstrate differences between beta-blocker  
32 therapy and other antihypertensives.

##### 6.3.3.5.33 Cardiovascular outcomes

34 All reported CV outcomes were for beta-blockers vs CCBs.

35 For the study considering COER verapamil and atenolol or hydrochlorothiazide there was NS  
36 difference between the groups for both the composite of acute MI, stroke or CV related death  
37 and also for the incidence of any component of the composite in the diabetes subgroup.<sup>261</sup>  
38 **Level 1+**

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j BP reduction with all hypertensive treatments was a consistent feature of the studies and therefore only studies where there were significant differences between the treatments will be highlighted.

1 The ASCOT-BPLA study found that for the diabetes subgroup for total CV events and  
2 procedures there was significantly lower occurrence with the amlodipine based group vs the  
3 atenolol based group (HR 0.87, 0.76 to 0.99,  $p=0.0283$ ), this was also found for the  
4 participants who did not have diabetes.<sup>262</sup> **Level 1++**

5 The INVEST study found NS difference in the treatments (verapamil SR and atenolol) for  
6 death or first occurrence of non-fatal MI or non-fatal stroke in both groups of patients with  
7 and without diabetes.<sup>259</sup> **Level 1+**

#### 6.3.3.5.28 *Blood pressure*

9 Within all the papers included that reported BP outcomes the treatments reduced BP and  
10 there was NS difference found between the treatment groups.<sup>260-262</sup>

#### 6.3.3.5.31 *Renal outcomes*

12 Only the study comparing 2 beta-blockers reported on renal outcomes.

13 The study considering carvedilol and metoprolol found that carvedilol reduced the  
14 albumin:creatinine ratio vs metoprolol (relative reduction 16%,  $p=0.003$ ).<sup>260</sup> This study also  
15 identified those with albuminuria of 30 mg or less at baseline, fewer in the carvedilol group vs  
16 the metoprolol group progressed to microalbuminuria (6.4%, 25/388 vs 10.3%, 56/542), or  
17 from carvedilol vs metoprolol, 0.60, 0.36 to 0.97,  $p=0.04$ ).<sup>260</sup> **Level 1++**

#### 6.3.3.5.48 *Metabolic outcomes*

19 Only the study comparing 2 beta-blockers reported on metabolic outcomes.

20 The study considering carvedilol and metoprolol found that carvedilol treatment had no  
21 HbA1c changes from baseline while metoprolol increased HbA1c. The mean difference was  
22 0.12%,  $p=0.006$ . More participants withdrew due to worsening glycaemic control with  
23 metoprolol (2.2%, 16/737) than with carvedilol (0.6%, 3/498),  $p=0.04$ .<sup>260</sup> **Level 1++**

#### 6.3.3.5.54 *Adverse events*

25 The study comparing COER verapamil with atenolol or hydrochlorothiazide<sup>261</sup> reported that  
26 participants assigned COER verapamil withdrew more often due to adverse signs or  
27 symptoms compared with those assigned atenolol or hydrochlorothiazide ( $p=0.02$ ); the most  
28 common reason was constipation (216 in the COER verapamil compared with 28 in the  
29 atenolol or hydrochlorothiazide group). However, fewer participants assigned COER  
30 verapamil (N=115) or atenolol or hydrochlorothiazide withdrew because of poor BP control  
31 compared with those assigned atenolol or hydrochlorothiazide (N=207) ( $p<0.001$  by log-  
32 rank). **Level 1+**

33 The INVEST study<sup>259</sup> showed that verapamil and atenolol were generally well tolerated in  
34 each treatment group. Patients in the verapamil group reported constipation and coughs  
35 more frequently than patients in the atenolol group, while atenolol-treated patients had more  
36 dyspnoea, light-headedness, symptomatic bradycardia, and wheezing. **Level 1+**

37 The RCT comparing carvedilol with metoprolol did not report significant differences between  
38 groups in overall safety profile. However, the study stated that no participant taking carvedilol  
39 had a respiratory event in contrast with 7 events in 6 participants taking metoprolol. **Level 1+**

40 The ASCOT-BPLA study concluded that the most frequent AEs found in the amlodipine  
41 based group were peripheral oedema 23%; cough 19%; joint swelling 14%; dizziness 12%;  
42 chest pain 8%; fatigue 8%. In the atenolol based group the most frequent AEs were  
43 dizziness 16%; fatigue 16%; dyspnoea 9%; cough 8%; erectile dysfunction 7%. **Level 1+**

### 6.3.4.1 Health economic evidence statements

#### 6.3.4.1.2 ACEI

3 Ramipril was found to be cost-effective compared to placebo, £2971/LYG<sup>263</sup> and  
4 €2486/LYG<sup>264</sup> (£1699/LYG, exchange rate 0.68, 13 March 2007).<sup>271</sup>

5 No statistically significant difference was found between captopril and atenolol. Atenolol had  
6 significantly lower mean costs.<sup>265</sup>

#### 6.3.4.2.7 A2RB

8 Irbesartan was found to be both more effective and cost saving than amlodipine and  
9 standard antihypertensive treatment.<sup>266-268</sup>

10 Losartan was found to be both more effective and cost saving than standard antihypertensive  
11 treatment.<sup>269</sup>

12 Valsartan was found to be both more effective and cost saving compared to amlodipine.<sup>270</sup>

### 6.3.5.3 Evidence to recommendations

14 The GDG used as its starting point the 2006 update of the NICE hypertension guidelines and  
15 the NICE type 2 diabetes hypertension guideline from 2002, available at [www.nice.org.uk](http://www.nice.org.uk).  
16 The group noted that the health economic model for the former did not include renal or  
17 retinopathy outcomes, both of particular importance when considering choice of therapies for  
18 use in people with type 2 diabetes. Thus 25% of people with type 2 diabetes develop diabetic  
19 nephropathy within 20 years of diagnosis, while the drugs studied in the UKPDS  
20 hypertension study had strong effects on retinopathy progression. Therefore, the GDG was  
21 particularly interested in reviewing the evidence as to whether there were any differential  
22 effects in terms of different classes of antihypertensive agent on microvascular as well as  
23 cardiovascular outcomes in people with type 2 diabetes.

24 The GDG noted a wealth of new evidence in this area since the hypertension guideline 2002  
25 was published, and were cognisant of the early revision of the NICE hypertension guidelines  
26 2006, albeit these applying to people without diabetes. Much of the new evidence seemed to  
27 be driven by studies in people with diabetes with increased AER (microalbuminuria or  
28 worse). The high known prevalence of renal damage in people with type 2 diabetes and the  
29 need to prevent this and its progression were noted to emphasise the importance of BP  
30 control. Little evidence on retinopathy prevention was available to the GDG, but it was aware  
31 of the positive data previously assessed for ACEI and a beta-adrenergic blocker. Published  
32 CV outcome data was noted to be of limited quality in some studies due to under powering in  
33 studies with other primary end points, even when combined for meta-analysis.

34 The GDG noted that the evidence did not distinguish between medications on the basis of  
35 degree of BP lowering. The issues of importance revolved around differences of evidence of  
36 effectiveness in renal related outcomes and metabolic worsening. Some classes of  
37 medications, notably A2RB and alpha-adrenergic blockers, were only available in more  
38 expensive proprietary form, and thus without added evidence of efficacy would not be cost-  
39 effective compared to older drugs.

40 Overall it was felt that the best evidence for prevention of renal disease and limitation of  
41 metabolic worsening related to the renin angiotensin system-blockers (RAS-blockers) (ACEI  
42 and A2RB) as a class.

43 With regard to non-renal outcomes, no evidence was identified that caused the GDG to  
44 reach any different conclusions from the review of the evidence carried out for the NICE

1 hypertension guideline 2006. The GDG recognised there was good evidence of efficacy for  
2 thiazide diuretics and CCBs, including when used in combination with RAS-blockers.

3 Given the benefits in terms of reno-protection and retinopathy of RAS blockade, it was felt  
4 appropriate to recommend RAS-blockers as first-line medication in the treatment of  
5 hypertension in type 2 diabetes. This was the only change in sequencing that the GDG felt  
6 was appropriate to make to the NICE hypertension guidelines 2006. On the grounds of cost a  
7 generic 24-hour ACEI should be used first line. A2RB (also selected on grounds of cost)  
8 should only be substituted in the event of significant ACEI intolerance, usually troublesome  
9 chronic cough (and not if hyperkalaemia or decreased renal function is the problem). An  
10 exception was highlighted in the NICE hypertension guideline 2006, where people of African-  
11 Caribbean descent are noted to respond less well to RAS- blockers, and for someone in this  
12 group either combination ACEI + diuretic therapy or CCB was thought appropriate first line  
13 therapy. Little specific information was available for other ethnic groups.

14 Thiazide diuretics and CCBs are recommended as second-line medications, though it was  
15 noted that it would be usual to need at least 2 drugs or more, so these would be added to a  
16 RAS-blocker and each other for the most part. There was some concern about the adverse  
17 metabolic effects of thiazides (in contrast to the positive effects of RAS-blockers and neutral  
18 effects of CCB), though the standard dose of bendroflumethiazide was thought not to be a  
19 problem in this regard.

20 Many people with diabetes do require 4 or even 5 antihypertensive agents to approach target  
21 levels. After 3 classes of medication had been used the GDG felt that reasons for  
22 distinguishing between other drug classes were poor. It was felt that any alpha-blocker, beta-  
23 blocker, or potassium-sparing diuretic could be added at this stage. If an RAS-blocker is  
24 used with a potassium-sparing diuretic, the potassium levels should be carefully monitored,  
25 the clinician being alert to the possibility of hyperkalaemia.

26 While in general this was felt to be the appropriate positioning of the beta-blockers,  
27 particularly because of their metabolic effects when used in combination with thiazides, it  
28 was recognised that some people would have a clearer indication for these drugs through  
29 having angina, heart failure, or previous heart attack. In these circumstances the drugs would  
30 already be being prescribed. One study suggested that carvedilol was superior to metoprolol  
31 both in metabolic terms and for renal protection. The GDG found the evidence interesting but  
32 incomplete in regard of target groups and active comparisons with the RAS-blockers;  
33 accordingly no out-of-class recommendations are made.

34 There is a need to emphasise caution over the use of some drug classes in the increasing  
35 numbers of women with type 2 diabetes who might become pregnant. The GDG felt  
36 comfortable that the decision to use, or not use such drugs should be one of informed  
37 agreement between each woman and their professional advisor.

38 Issues of adherence and the use of fixed-dose combination therapy were considered. The  
39 evidence was not formally available to the GDG, but clinical experience over the combined  
40 burden of medications faced by many people with type 2 diabetes led to an overall view that  
41 combination tablets could be appropriate in reducing that burden, and possibly improving  
42 outcomes through better adherence. No formal recommendations could be made.

43 The GDG were aware of the issues that arose from the burden of use of multiple therapies.  
44 In this area in particular it was therefore felt appropriate to further emphasise communication,  
45 discussion and agreement about medication use.

46 An issue considered of importance, but not covered in the evidence review was that of BP  
47 monitoring, including the role of self-monitoring and of ambulatory BP monitoring. The GDG  
48 was happy to defer to the NICE hypertension guideline 2006 (now update by the NICE  
49 hypertension guideline 2011) on these issues.

### 6.3.61 Recommendations

- 2 **17. Measure blood pressure at least annually in an adult with type 2 diabetes without**  
3 **previously diagnosed hypertension or renal disease. Offer and reinforce**  
4 **preventive lifestyle advice. [2009]**
- 5 **18. For an adult with type 2 diabetes on antihypertensive drug treatment when**  
6 **diabetes is diagnosed, review blood pressure control and medications used. Make**  
7 **changes only if there is poor control or if current drug treatment is not**  
8 **appropriate because of microvascular complications or metabolic problems.**  
9 **[2009]**
- 10 **19. Repeat blood pressure measurements within:**
- 11                   • 1 month if blood pressure is higher than 150/90 mmHg  
12                   • 2 months if blood pressure is higher than 140/80 mmHg  
13                   • 2 months if blood pressure is higher than 130/80 mmHg and there is  
14                   kidney, eye or cerebrovascular damage.
- 15 **Provide lifestyle advice (diet and exercise) at the same time. [2009]**
- 16 **20. Provide lifestyle advice (see section 5.1.6 and the [lifestyle interventions](#) section**  
17 **in ‘Hypertension’ [NICE clinical guideline 127]) if blood pressure is confirmed as**  
18 **being consistently above 140/80 mmHg (or above 130/80 mmHg if there is kidney,**  
19 **eye or cerebrovascular damage). [2009]**
- 20 **21. Add medications if lifestyle advice does not reduce blood pressure to below**  
21 **140/80 mmHg (below 130/80 mmHg if there is kidney, eye or cerebrovascular**  
22 **damage). [2009]**
- 23 **22. Monitor blood pressure every 1–2 months, and intensify therapy if the person is**  
24 **already on antihypertensive drug treatment, until the blood pressure is**  
25 **consistently below 140/80 mmHg (below 130/80 mmHg if there is kidney, eye or**  
26 **cerebrovascular disease).[2009]**
- 27 **23. First-line antihypertensive drug treatment should be a once-daily, generic**  
28 **angiotensin-converting enzyme (ACE) inhibitor. Exceptions to this are people of**  
29 **African or Caribbean family origin, or women for whom there is a possibility of**  
30 **becoming pregnant. [2009]**
- 31 **24. The first-line antihypertensive drug treatment for a person of African or Caribbean**  
32 **family origin should be an ACE inhibitor plus either a diuretic or a generic**  
33 **calcium-channel blocker. [2009]**
- 34 **25. A calcium-channel blocker should be the first-line antihypertensive drug**  
35 **treatment for a woman for whom, after an informed discussion, it is agreed there**  
36 **is a possibility of her becoming pregnant. [2009]**
- 37 **26. For a person with continuing intolerance to an ACE inhibitor (other than renal**  
38 **deterioration or hyperkalaemia), substitute an angiotensin II-receptor antagonist**  
39 **for the ACE inhibitor. [2009]**
- 40 **27. If the person’s blood pressure is not reduced to the individually agreed target with**  
41 **first-line therapy, add a calcium-channel blocker or a diuretic (usually a thiazide or**

- 1 **thiazide-related diuretic). Add the other drug (that is, the calcium-channel blocker**  
 2 **or diuretic) if the target is not reached with dual therapy. [2009, amended 2015]**
- 3 **28. If the person's blood pressure is not reduced to the individually agreed target with**  
 4 **triple therapy, add an alpha-blocker, a beta-blocker or a potassium-sparing**  
 5 **diuretic (the last with caution if the individual is already taking an ACE inhibitor or**  
 6 **an angiotensin II-receptor antagonist). [2009]**
- 7 **29. Monitor the blood pressure of a person who has attained and consistently**  
 8 **remained at his or her blood pressure target every 4–6 months. Check for**  
 9 **possible adverse effects of antihypertensive drug treatment – including the risks**  
 10 **from unnecessarily low blood pressure. [2009]**

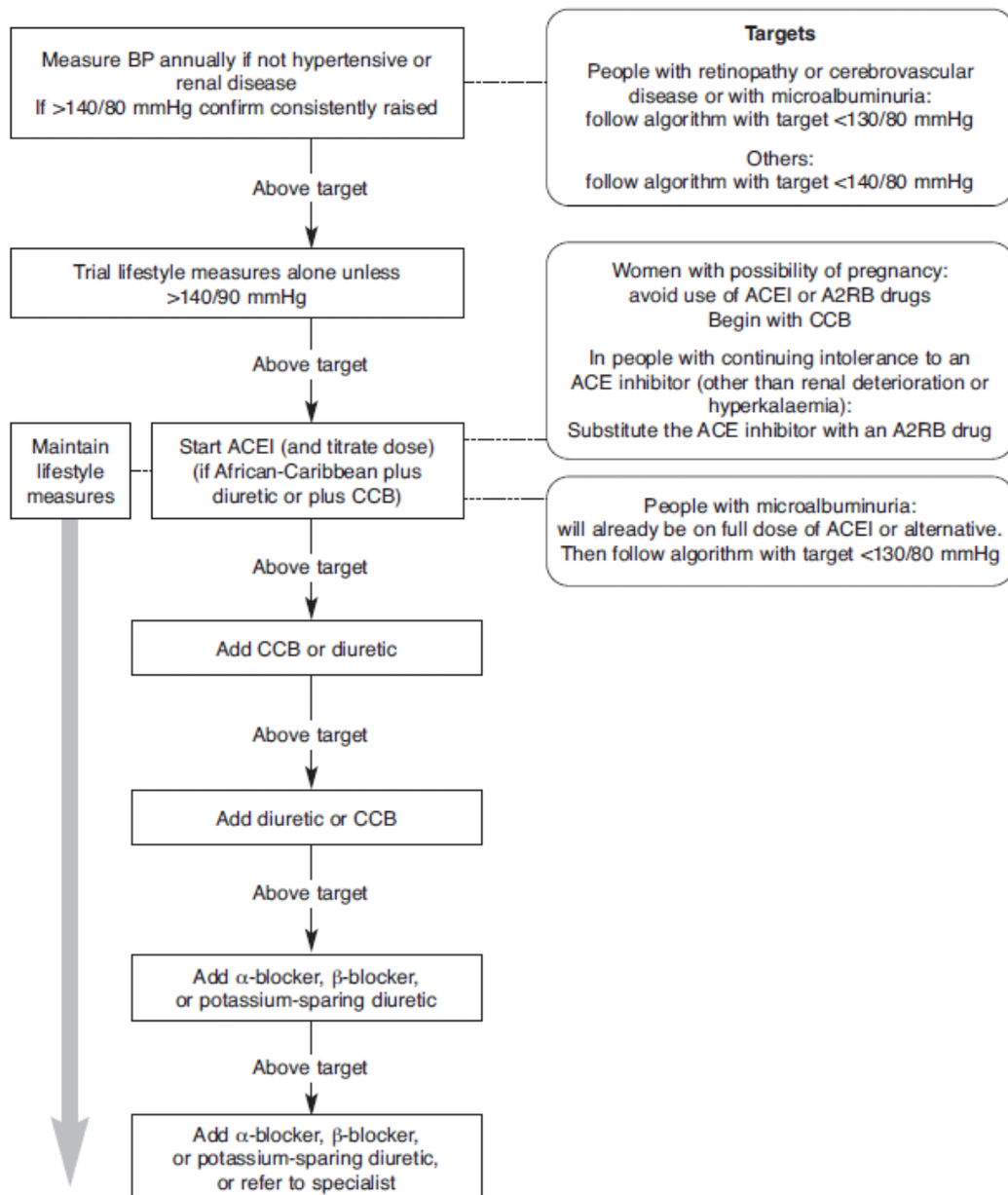


Figure 12.1 Scheme for the management of blood pressure (BP) for people with Type 2 diabetes  
 ACEI, angiotensin-converting enzyme inhibitor; A2RB, angiotensin 2 receptor blocker (sartan); CCB, calcium channel blocker

# 7.1 Antiplatelet therapy for primary prevention of cardiovascular disease

## 7.1.3 Clinical introduction

4 Antiplatelet therapy has an established role in the management of people with cardiovascular  
5 disease. However, its role in primary prevention for people without existing cardiovascular  
6 disease is less clear. This review question addressed whether aspirin or clopidogrel (either  
7 alone or in combination) should be used for the prevention of cardiovascular events in people  
8 with type 2 diabetes who do not have existing cardiovascular disease, that is for primary  
9 prevention. This question also covered whether their use should be restricted to specific  
10 subgroups of the population, when these treatments should be used and what adverse  
11 events are associated with their use.

### 7.1.12 Antiplatelet therapy in Clinical Guideline 66

13 Antiplatelet therapy was originally covered as part of CG66. The original searches were  
14 conducted from 2001 to 2007 (see Appendix G for search strategies from CG66). Update  
15 searches have been carried out for this topic with a date restriction of 2007 to June 2014  
16 (see Appendix C for updates search strategies). Although the focus in CG66 was primary  
17 prevention of cardiovascular disease, the evidence also included studies on secondary  
18 prevention. In total, 8 RCTs were originally included for this review question.

### 7.1.29 Antiplatelet therapy in the update (2015)

20 Although aspirin and clopidogrel are not licensed for primary prevention of cardiovascular  
21 disease, the GDG considered that an updated evidence review was important as such off-  
22 label use of these particular drugs is common in current clinical practice. The group agreed  
23 that only studies on adults with type 2 diabetes who did not have established cardiovascular  
24 disease should be included, to ensure that the findings of the review are specific to primary  
25 prevention. The GDG considered that people with type 2 diabetes and established  
26 cardiovascular disease are inherently different in terms of risk factors, and therefore findings  
27 from secondary prevention studies could not credibly be extrapolated to those without  
28 cardiovascular disease. In addition, the GDG recognised that the evidence supporting the  
29 role of antiplatelet therapy in secondary prevention is established, whereas there is debate  
30 surrounding its use in primary prevention, and changes in the evidence base would likely  
31 impact on clinical practice.

### 7.1.32 Evidence review

#### 7.1.3.33 Review question

34 Should aspirin and/or clopidogrel be used for primary prevention of cardiovascular disease in  
35 people with type 2 diabetes?

#### 36 Table 18: PICO table

Population	Adults (18 years and over) with type 2 diabetes without established cardiovascular disease
Interventions	Aspirin, clopidogrel, aspirin plus clopidogrel
Comparators	Placebo, listed interventions
Outcomes	Development of cardiovascular disease (myocardial infarction, heart failure, ischaemic stroke, acute coronary syndrome, transient ischaemic attack,

revascularisation and stenting)  
Adverse events such as any bleeding including gastrointestinal bleeding,  
haemorrhagic stroke  
Mortality  
Health-related quality of life

- 1
- 2 RCTs examining the use of aspirin or clopidogrel in people with type 2 diabetes were  
3 included. Papers were excluded if they:
- 4 • were non-randomised studies (such as observational studies, narrative reviews and  
5 conference abstracts)
- 6 • included a mixed population of people with type 1 and 2 diabetes and either did not report  
7 subgroup analyses, or less than 85% of the study population had type 2 diabetes
- 8 • focused on the use of aspirin or clopidogrel after acute cardiological events, cardiac  
9 interventions or cerebrovascular events (that is, secondary prevention)
- 10 • focused on the use of antiplatelet drugs other than aspirin or clopidogrel (such as  
11 dipyridamole, prasugrel, ticagrelor as these are generally used for secondary prevention).  
12 For the full excluded list, see Appendix L.
- 13 The main outcomes for this review question were the development of cardiovascular disease  
14 and adverse events specifically any bleeding including gastrointestinal bleeding. The detailed  
15 protocol is available in Appendix C.

#### 7.1.3.26 Clinical evidence

17 The evidence that was originally included as part of CG66 was re-reviewed as part of the  
18 update. Six trials examining clopidogrel (either alone or in combination with aspirin) were  
19 excluded as they were conducted in people with established cardiovascular disease (Bhatt et  
20 al. 2002; Diener et al. 2004; Mehta et al. 2001; Steinhubl et al. 2002; Yusuf et al. 2001) or did  
21 not provide separate results for primary prevention (Bhatt et al. 2006). Another trial from  
22 CG66 (Khajehdehi et al. 2002) was also excluded as it reported data on kidney damage but  
23 no cardiovascular outcomes, and was limited to only 2 months of treatment. The final study,  
24 the Primary Prevention Project (PPP) trial (Sacco et al 2003) met the revised inclusion  
25 criteria for this update.

26 In total, 1204 references were found in the update searches and 1 RCT was included  
27 (Ogawa et al. 2008). The GDG was also aware of a post hoc analysis of cardiovascular  
28 outcomes that was being undertaken on the Early Treatment Diabetic Retinopathy Study  
29 (ETDRS Investigators 1992), and requested unpublished data from the authors on adults  
30 with type 2 diabetes without a history of cardiovascular disease.

31 Data from all 3 trials focused on the use of aspirin therapy compared with no aspirin. No trials  
32 were identified that examined the use of clopidogrel (alone or in combination with aspirin) in  
33 people with type 2 diabetes without existing cardiovascular disease.

34 Pooling of studies using meta-analysis was not possible because the definitions of  
35 cardiovascular outcomes varied across the studies and different estimates of effect were  
36 used that is, hazard ratios and risk ratios.

#### 7.1.3.37 Description of included studies

38 The 3 RCTs including a total of 7281 participants were carried out in the USA (ETDRS:  
39 unpublished data 2013), Italy (Sacco et al. 2003) and Japan (Ogawa et al. 2008). All trials  
40 randomised participants to aspirin or no aspirin (placebo or vitamin E), with doses of aspirin  
41 ranging from 81 (Ogawa et al. 2008) to 650 mg (ETDRS: unpublished data 2013). The mean  
42 age of participants in 2 trials ranged from 64 to 65 years, while the last study did not provide

- 1 this information (ETDRS: unpublished data 2013). Mean HbA1c at baseline ranged from 53
- 2 to 54 mmol/mol (7.0% to 7.1%) in 2 trials, with 1 study reporting that about 33% of the
- 3 participants had baseline HbA1c greater than 86 mmol/mol (10%) (ETDRS: unpublished data
- 4 2013). The median follow-up ranged from 3.7 to 5 years. Details of the included studies are
- 5 found in the evidence tables (see Appendix E).
  
- 6 A summary GRADE table is presented for this review question (see Appendix D for full
- 7 GRADE tables).

1 Table 19: Summary GRADE profile for aspirin therapy for primary prevention of cardiovascular disease

Number of RCTs	Number of people		Relative effect (95% CI)	Quality
	Aspirin	Control		
<b>All-cause mortality; follow-up for up to 5 years</b>				
1 (ETDRS)†	587	565	HR 0.99 (0.83 to 1.17)	Moderate
1 (Sacco 2003)-PPP	25/519	20/512	RR 1.23 (0.69 to 2.19)	Very low
<b>Cardiovascular mortality; follow-up for up to 5 years</b>				
1 (ETDRS)†	587	565	CV death: HR 0.97 (0.79 to 1.19)	Moderate
1 (Sacco 2003)-PPP	10/519	8/512	CV mortality: RR 1.23 (0.49 to 3.10)	Very low
1 (Ogawa 2008)-JPAD	0/1262	5/1277	Fatal MI: HR not estimable due to no events in aspirin group	Low
<b>Cerebrovascular mortality; follow-up for median 4.4 years</b>				
1 (Ogawa 2008)-JPAD	1/1262	5/1277	Fatal stroke: HR 0.20 (0.024 to 1.74)	Low
<b>Coronary and cerebrovascular mortality; follow-up for median 4.4 years</b>				
1 (Ogawa 2008)-JPAD	1/1262	10/1277	HR 0.10 (0.01 to 0.79)	Low
<b>Non-cardiovascular mortality; follow-up to median 3.7 years</b>				
1 (Sacco 2003)-PPP	15/519	12/512	RR 1.23 (0.58 to 2.61)	Very low
<b>Any atherosclerotic event<sup>a</sup>; follow-up from median 3.7 to 4.4 years</b>				
1 (Sacco 2003)-PPP	20/519	22/512	RR 0.90 (0.50 to 1.62)	Very low

Number of RCTs	Number of people		Relative effect (95% CI)	Quality
	Aspirin	Control		
1 (Ogawa 2008)-JPAD	68/1262	86/1277	<p>HR 0.80 (0.58 to 1.10)</p> <p><u>Subgroup: age</u>                      ≥ 65 years: HR 0.68 (0.46 to 0.99)                      &lt; 65 years: HR 1.00 (0.57 to 1.70)</p> <p><u>Subgroup: sex</u>                      Male: HR 0.74 (0.49 to 1.12)                      Female: HR 0.88 (0.53 to 1.44)</p> <p><u>Subgroup: cardiovascular risk factors</u>                      Hypertensive: HR 0.88 (0.60 to 1.30)                      Normotensive: HR 0.64 (0.36 to 1.13)                      Dyslipidaemia: HR 0.88 (0.57 to 1.37)                      Normolipidaemia: HR 0.71 (0.45 to 1.14)                      Current/past smoking: HR 0.73 (0.47 to 1.14)                      Non-smoker: HR 0.83 (0.53 to 1.31)</p> <p><u>Subgroup: renal function</u>                      eGFR ≥ 90: HR 0.87 (0.36 to 2.12)<sup>d</sup>                      eGFR 60-89: HR 0.53 (0.34 to 0.83)<sup>d</sup>                      eGFR &lt; 60: HR 1.24 (0.69 to 2.23)<sup>d</sup></p> <p><u>Subgroup: existing therapies</u>                      Insulin: HR 1.00 (0.50 to 2.00)<sup>d</sup>                      OHA: HR 0.77 (0.52 to 1.14)<sup>d</sup>                      Diet alone: HR 0.20 (0.06 to 0.68)<sup>d</sup></p>	Low
<b>Coronary heart disease events; follow-up from median 3.7 to 5 years</b>				
1 (ETDRS)†	587	565	<p>MI: HR 0.85 (0.70 to 1.05)</p> <p>CV event<sup>b</sup>: HR 0.97 (0.82 to 1.15)</p>	Moderate
1 (Sacco 2003)-PPP	53/519	59/512	Total CV events: RR 0.89 (0.62 to 1.26)	Very low

Number of RCTs	Number of people		Relative effect (95% CI)	Quality
	Aspirin	Control		
1 (Ogawa 2008)-JPAD	5/519	10/512	All MI: RR 0.49 (0.17 to 1.40)	Low
	13/519	16/512	Angina: RR 0.80 (0.39 to 1.64)	
	28/1262	35/1277	Any fatal or nonfatal event: HR 0.81 (0.49 to 1.33)	
	12/1262	9/1277	Nonfatal MI: HR 1.34 (0.57 to 3.19)	
	12/1262	11/1277	Stable angina: HR 1.10 (0.49 to 2.50)	
	4/1262	10/1277	Unstable angina: HR 0.40 (0.13 to 1.29)	
			<u>Cardiovascular events subgrouped by cardiovascular risk:</u> In low risk group: HR 0.53 (0.23 to 1.21) In high risk group: HR 0.78 (0.55 to 1.11)	
<b>Cerebrovascular events; follow-up from median 3.7 to 5 years</b>				
1 (ETDRS)†	587	565	Stroke: HR 1.09 (0.78 to 1.53)	Low
1 (Sacco 2003)-PPP	9/519	10/512	All stroke: RR 0.89 (0.36 to 2.17)	Very low
	7/519	10/512	Transient ischaemic attack: RR 0.69 (0.27 to 1.79)	
1 (Ogawa 2008)-JPAD	28/1262	32/1277	Any fatal or nonfatal event: HR 0.84 (0.53 to 1.32)	Low
	22/1262	24/1277	Nonfatal ischaemic stroke: HR 0.93 (0.52 to 1.66)	
	5/1262	3/1277	Nonfatal haemorrhagic stroke: HR 1.68 (0.40 to 7.04)	
	5/1262	8/1277	Transient ischaemic attack: HR 0.63 (0.21 to 1.93)	
			<u>Cerebrovascular events subgrouped by blood pressure control<sup>c</sup>:</u> In non-aspirin group: HR 2.84 (1.52 to 5.52) indicating higher incidence in unattained group In aspirin group: HR 1.64 (0.83 to 3.29) indicating no difference in incidence in unattained vs. attained No HR reported for aspirin vs. non-aspirin but reported as not significant	
<b>Peripheral artery disease; follow-up from median 3.7 to 4.4 years</b>				
1 (Sacco 2003)-PPP	11/519	13/512	RR 0.83 (0.38 to 1.84)	Very low
1 (Ogawa 2008)-JPAD	7/1262	11/1277	HR 0.64 (0.25 to 1.65)	Low
<b>Revascularisation; follow-up to median 3.7 years</b>				
1 (Sacco 2003)-PPP	8/519	10/512	RR 0.79 (0.31 to 1.97)	Very low
			Creatinine clearance: MD -2.30 (-5.42 to 0.82)	

Number of RCTs	Number of people		Relative effect (95% CI)	Quality
	Aspirin	Control		
			Urine protein:creatinine ratio: MD -0.30 (-0.53 to -0.07)	
			% proteinuria change: MD -17.80 (-22.95 to -12.65)	
<b>Adverse events: Any bleeding; follow-up for median 4.4 years</b>				
1 (ETDRS 1992)	587	565	Only a few patients (2%) in both groups had some indication of bleeding <sup>‡</sup>	Low
1 (Ogawa 2008)-JPAD	1251	1272	Haemorrhagic events subgrouped by renal function: eGFR ≥ 90: HR not estimable eGFR 60-89: HR 1.03 (0.24 to 4.35) eGFR < 60: HR: 0.87 (0.10 to 7.27)	Low
	21/1262	6/1277	Other bleeding: RR 3.54 (1.43 to 8.75)	
	12/1262	4/1277	Gastrointestinal bleeding: RR 3.04 (0.98 to 9.39)	
<b>Non-bleeding gastrointestinal event; follow-up for median 4.4 years</b>				
1 (Ogawa 2008)-JPAD	47/1262	4/1277	RR 11.89 (4.30 to 32.90)	Moderate
<b>Other adverse event<sup>e</sup>; follow-up for median 4.4 years</b>				
1 (Ogawa 2008)-JPAD	5/1262	0/1277	RR 11.13 (0.62 to 201.08)	Low

Abbreviations: BP blood pressure; CV cardiovascular; eGFR estimated glomerular filtration rate; HR hazard ratio; MD mean difference; MI myocardial infarction; OHA Oral hypoglycaemic agents; RCT randomised controlled trial; RR relative risk

NB: data from ETDRS (unpublished 2013) are from multivariate analysis; data from the JPAD trial (Ogawa et al. 2008) are from Cox proportional hazards model (not specified as multivariate) in multiple publications; data from the PPP trial (Sacco et al. 2003) are relative risks as multivariate analyses using Cox regression are not reported for people with diabetes

<sup>a</sup> any atherosclerotic event was defined as a composite of sudden death, death from coronary, cerebrovascular and aortic causes, nonfatal acute MI, unstable angina, newly developed exertional angina, nonfatal ischaemic and haemorrhagic stroke, transient ischaemic attack or nonfatal aortic and peripheral vascular disease

<sup>b</sup> CV event was defined as CV death, myocardial infarction or stroke

<sup>c</sup> unattained group had systolic BP ≥ 140 mmHg and/or diastolic BP ≥ 90 mmHg and the attained group had systolic BP < 140mmHg and/or diastolic BP < 90mmHg

<sup>d</sup> adjusted for age, hypertension, dyslipidaemia and history of smoking

<sup>e</sup> Anaemia and asthma

<sup>†</sup> Unpublished subgroup analysis for people with type 2 diabetes without a history of cardiovascular disease from the ETDRS trial was provided by the authors

<sup>‡</sup> haemoglobin < 100 g/L or haematocrit < 0.30, haematuria, or blood in the stool

#### 7.1.3.41 Health economic evidence

- 2 Literature searches were undertaken to find any existing cost–utility analyses (CUAs) of  
3 using clopidogrel or aspirin for the primary prevention of cardiovascular disease in people  
4 with type 2 diabetes (see appendix C for details of the search strategies). In total, 537  
5 articles were found and 2 CUAs were returned that met the NICE reference case (National  
6 Institute for Health and Social Care, 2012).
- 7 One CUA (Li et al. 2010) used an existing diabetes health economic model (CDC-RTI model]  
8 [CDC Diabetes Cost-Effectiveness Group 2002) to compare daily aspirin use with no aspirin  
9 in a population of people with newly diagnosed type 2 diabetes. The treatment effect was  
10 taken from a non-diabetes-specific meta-analysis but other parameters (including costs and  
11 utilities) were specific to people with type 2 diabetes.
- 12 Another CUA (Lamotte et al. 2006) created a Markov model to assess the impact of daily  
13 aspirin use to no aspirin over 10 years in 4 countries (including UK). This model was not  
14 diabetes specific but used varying prespecified annual risks of CVD events. Costs were  
15 taken from UK reference costs and the UKPDS trial; utility sources were unclear.
- 16 One CUA (Lamotte et al. 2006) found that for the UK, daily aspirin use dominated no aspirin  
17 at baseline risks of CVD greater than 0.24% per year, whilst the other CUA (Li et al. 2010)  
18 found that, for America, daily aspirin use was cost effective compared to no aspirin (ICER  
19 \$8800 per QALY). Both results were unchanged under both deterministic and probabilistic  
20 sensitivity analyses.
- 21 No CUAs were found that assessed the use of clopidogrel for primary prevention of  
22 cardiovascular disease in people with type 2 diabetes.
- 23 This question was not prioritised by the GDG for de novo economic modelling.

1 Table 20: Economic evidence for aspirin use to prevent cardiovascular events

Study, Population, Comparators and Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Cost	Effect	ICER		
<b>Lamotte et al. (2006)</b> People without CHD history, annual baseline risk 1.5% per annum 4 countries (UK, Italy, Germany, Spain) Aspirin 75mg daily <b>Partly applicable</b> <sup>a,b</sup> <b>Potentially serious limitations</b> <sup>c,h</sup>	<u>Effects:</u> 2 meta-analyses (same trials), reasons for selection not given. Not UK specific <u>Costs:</u> Country specific. UK reference costs and UKPDS for complications (€, 2003, country specific discounting) <u>Utilities:</u> from literature. Not UK, limited detail	Markov model with 10 year time horizon Baseline annual CHD risk 1.5% 5 states – no CVD, MI, stroke, CVD, death. TIA, PAD and stable angina not considered. Only 2/5 trials contain women Funded by industry	UK -€201	UK 0.04 QALYs	UK: dominant	In the UK, aspirin is likely to be cost saving given baseline annual CHD risk > 0.24% Results over 10 years at baseline risk of 1.5% per annum ICER sensitive to risk GI bleeding and stroke risk in Italy In PSA, aspirin is dominant in 97% of replications Country cost comparisons differ due to ratio between aspirin and complication costs Country utility comparisons differ due to country specific discount rates	
			Germany -€281	Germany 0.02 QALYs	Germany: dominant		
			Spain -€797	Spain 0.03 QALYs	Spain: dominant		
<b>Li et al. (2010)</b> US residents aged 40-94 with newly diagnosed type 2 diabetes Aspirin 80mg daily <b>Partly applicable</b> <sup>d,e</sup> <b>Potentially serious limitations</b> <sup>c,f,g</sup>	<u>Effects:</u> US age-gender specific, non-diabetes specific meta-analysis <u>Costs:</u> other US studies, health system perspective (\$, 2006, discount rate 3% for cost and utilities) <u>Utilities:</u> from literature. QWB scale from US type 2 diabetes attending hospital clinic	Existing Markov model with lifetime horizon Only RRs for major events and strokes statistically significant Diabetes specific meta-analysis found effect of aspirin in primary prevention unproven Base case models ischaemic and haemorrhagic strokes together People with newly diagnosed diabetes only but utilities from longstanding diabetes	\$1700	0.19 QALYs	\$8801/ QALY	Daily aspirin appears very cost effective for newly diagnosed people with type 2 diabetes aged 40+ years at \$50,000/QALY threshold ICER sensitive to gender, primary and secondary effectiveness, but ICERs remain < \$23,000/QALY In PSA, all iterations gave ICERs < \$27,000/QALY (not all parameters varied) Cardiac events avoided offset the cost and risk of bleeding Probably not cost-saving due to aspirin extending life (and potential for complications)	

Study, Population, Comparators and Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Cost	Effect	ICER		
a Not specific to type 2 diabetes b Source of utility value unclear c No evidence of systematic review for selection of clinical effects d Not 3.5% discount rate e Not UK based f May not include all relevant complications g Not all parameters varied in PSA h Potential conflict of interest CDC-RTI: Centre for Disease Control and Prevention Research Triangle Institute CHD: coronary heart disease CVD: cardiovascular disease ICER: incremental cost effectiveness ratio MI: myocardial infarction PAD: peripheral arterial disease PSA: probabilistic sensitivity analysis QALY: quality adjusted life years QoL: quality of life RR: relative risks TIA: transient ischaemic attack UK: United Kingdom UKPDS: United Kingdom Prospective Diabetes Study US: United States							
1							

#### 7.1.41 Evidence statements

##### 7.1.4.12 Clinical evidence

3 No trials were identified examining the use of clopidogrel alone or in combination with aspirin  
4 for primary prevention of cardiovascular disease in people with type 2 diabetes.

##### 7.1.4.1.15 Mortality

6 One trial found a clinically important reduction in the rate of combined coronary and  
7 cerebrovascular mortality in people with type 2 diabetes who were assigned aspirin therapy  
8 compared with people who were assigned to no aspirin therapy. The quality of the evidence  
9 is low.

10 There was no clinically important difference in the risk of mortality by any other definition in  
11 the 3 trials. The evidence ranged from moderate to very low.

##### 7.1.4.1.22 Atherosclerotic events with or without ischaemic events

13 Pre-specified subgroup analyses of 1 trial found a clinically important reduction in the  
14 incidence of atherosclerotic events in people aged 65 years and older who were assigned to  
15 aspirin therapy compared with those who were not.

16 Post hoc subgroup analyses from 1 trial found clinically important reductions in the rate of  
17 any atherosclerotic event in those who were assigned to aspirin therapy compared with those  
18 who did not receive aspirin in the following subgroups:

- 19 • people with mild renal dysfunction (eGFR between 60 and 89)
- 20 • people managed by diet alone

21 The quality of the evidence was low.

22 There was no clinically important difference in the risk of overall atherosclerotic events in 2  
23 trials. The evidence ranged from moderate to very low.

##### 7.1.4.1.34 Coronary heart disease

25 Evidence from 2 trials found no clinically important differences between those who received  
26 aspirin and those who did not for the following conditions: myocardial infarction, angina  
27 (stable or unstable), transient ischaemic attack, peripheral artery disease, revascularisation  
28 and any cardiovascular event. The quality of the evidence ranged from moderate to very low.

##### 7.1.4.1.29 Adverse events

30 Evidence from 1 trial found no clinically important differences in the rates of any bleeding  
31 between those who received aspirin therapy compared to people who did not. The quality of  
32 the evidence was low.

33 Evidence from 1 trial found no clinically important differences in the rates of gastrointestinal  
34 bleeding between people who received aspirin therapy and those who did not. The quality of  
35 the evidence was low. Evidence from the same trial found a clinically important difference in  
36 those receiving aspirin who were at greater risk of 'other' bleeding (non-gastrointestinal)  
37 compared with those who did not receive aspirin. The quality of the evidence was moderate.

38 Evidence from 1 trial found a clinically important difference in the rates of non-bleeding  
39 gastrointestinal events in those receiving aspirin who were at greater risk compared with  
40 those who did not receive aspirin. The quality of the evidence was moderate.

- 1 Evidence from 1 trial found no clinically important differences between the groups receiving
- 2 aspirin or no aspirin, in terms of 'other adverse events' and haemorrhagic events. The quality
- 3 of the evidence was low.

#### 7.1.4.24 Health economic evidence

- 5 Two cost–utility analyses assessed the impact of taking aspirin compared with not taking
- 6 aspirin for the primary prevention of cardiovascular events. While they were based on
- 7 different assumptions and data, they found that aspirin was likely to be cost effective
- 8 compared with not taking aspirin in both deterministic and probabilistic analyses.

## 7.2<sup>9</sup> Evidence to recommendations

### 10 Table 21: Linking evidence to recommendations

Relative value of different outcomes	<p>The GDG noted that, although reducing the risk of mortality or developing cardiovascular disease was important in improving life expectancy and quality of life, the relative impact of adverse events associated with the off-label use of aspirin and clopidogrel (such as bleeding) for primary prevention was also important in determining the safety and acceptability of treatment to the patient. Thus all outcomes were considered equally critical in decision making.</p>
Trade-off between benefits and harms	<p>In clinical practice, the decision to offer aspirin or clopidogrel depends on the net benefit to the patient. Potential benefits must be balanced against the possible harms from adverse effects, such as bleeding and gastrointestinal symptoms. It is particularly important to know the risk of these adverse effects when aspirin is used as primary prevention in people as yet free of, but at risk of developing, cardiovascular disease.</p> <p>The GDG considered that, overall, there was limited evidence to indicate that aspirin was beneficial in reducing the incidence of mortality, with only 1 study showing a reduction in a specific combined outcome of coronary and cerebrovascular mortality. Overall, there was no benefit in reducing the risk of developing a cardiovascular event, except in certain subgroups, such as those aged 65 years or older, people with mild renal dysfunction and those managing their diabetes using dietary changes alone.</p> <p>The GDG agreed that there was evidence to support an increased risk of harm associated with the use of aspirin, in terms of an increased risk of non-gastrointestinal bleeding and non-bleeding related gastrointestinal events. The Group discussed that any bleeding events would have a large negative impact on a patient's quality of life and anxiety levels. The GDG also agreed that the treatment of this adverse event may be costly. Non-bleeding gastrointestinal events can also have a negative impact on an individual's quality of life.</p> <p>The Group considered all the evidence and agreed that the increased risk of bleeding outweighed the potential benefits of taking aspirin.</p>
Consideration of health benefits and resource use	<p>The GDG thought that neither of the 2 CUAs reviewed accurately reflected the decision problem, and that both had serious limitations, but agreed that they both lent some value to the question.</p>

	<p>The GDG acknowledged that neither CUAs used diabetes-specific treatment effects, but the UK-based study used a range of baseline cardiovascular event risks, the higher values of which could be seen as approximating the baseline cardiovascular event risks of people with type 2 diabetes.</p> <p>The GDG agreed that, if the clinical review had found aspirin use to be effective, then it was likely to have been cost effective. However, the GDG noted that both the CUAs could be underestimating rates of adverse events compared with the clinical review. Underestimating adverse event rates would make the intervention appear more cost effective than it is.</p>
<p>Quality of evidence</p>	<p>The Group noted that there was uncertainty surrounding the majority of the outcome data as indicated by confidence intervals that generally crossed the line of no effect.</p> <p>The GDG noted that all of the included studies examined aspirin and agreed that the overall quality of evidence was low to very low. The GDG expressed concern about some methodological and clinical issues with the evidence base. The baseline HbA1c levels (approximately 53 mmol/mol [7%]) of people in the included trials were relatively low compared to the UK, which encouraged the GDG to question the generalisability of these findings to clinical practice in the UK. None of the studies had a follow-up period longer than 10 years, which is the typical timeframe by which the risk of developing cardiovascular disease is defined.</p> <p>The Group noted that the aspirin doses (81–650 mg) used in the studies were above the recommended UK maintenance dose of 75 mg. This cast further doubts on the generalisability of the findings to the UK clinical population. The GDG thought that higher doses could explain the increased risk of adverse effects found in the studies, but not the relative lack of benefit that one might expect to see.</p> <p>The GDG also noted that all of the studies included ‘any type of stroke’ in their composite outcomes of cerebrovascular and atherosclerotic events. However, the Group thought that different types of stroke should be considered separately, with prevention of ischaemic stroke being classified as a beneficial effect, but haemorrhagic stroke classified as an adverse event. The GDG recognised that the findings of trials which report both outcomes, that is, development of cardiovascular disease and adverse events, would not be affected by the combined reporting of all types of stroke events.</p> <p>The GDG recognised that the majority of data were derived from the JPAD trial, which was conducted in Japan, and questioned the generalisability of the findings to western countries. The GDG noted that the overall incidence of cardiovascular events was generally lower – possibly explained by different dietary habits, particularly fish consumption – in Japan, compared with western countries. The Group noted that the significant findings for the 3 different subgroups in the post hoc analyses were from this trial and also commented on the overall significant difference observed in favour of aspirin for the composite outcome of coronary (fatal myocardial infarction) and</p>

	<p>cerebrovascular (fatal stroke) mortality. The GDG thought that because of the low event rate, this single significant finding was likely to be fragile and very small changes in the event numbers would have a large impact on the estimate of effect.</p>
<p>Other considerations</p>	<p>The GDG noted that there were 2 ongoing trials that should provide more direct and applicable evidence to answer this review question in the future.</p> <p>The GDG was aware of an ongoing trial (ASCEND), which is fully recruited, randomised and includes 15,480 people with either type 1 or type 2 diabetes without occlusive arterial disease. The trial is being conducted in the UK and is scheduled to continue until 2017. The purpose of this 2x2 factorial, double-dummy study is to determine whether 100 mg of aspirin daily, with or without supplementation of 1 g of omega-3 fatty acid daily, prevents serious vascular events compared with placebo or supplementation of 1 g of omega-3 fatty acid daily only. The primary outcome measure is the combination of non-fatal myocardial infarction, non-fatal stroke or vascular death, excluding confirmed cerebral haemorrhage. The study also aims to assess serious bleeding and other adverse events.</p> <p>Another ongoing trial (ACCEPT-D) aims to assess the effects of low-dose aspirin on the incidence of major vascular events in people with type 1 or type 2 diabetes with no clinical evidence of vascular disease. The trial is being conducted in Italy and is scheduled to end in 2015.</p> <p>When making recommendations for the use of antiplatelet therapy (aspirin and clopidogrel), the GDG considered the following points:</p> <ul style="list-style-type: none"> <li>• Although the evidence base is small, the included evidence supported an increased risk of harm (including bleeding events), which was associated with the use of aspirin.</li> <li>• There was uncertainty around whether aspirin reduced the incidence of cardiovascular events.</li> </ul> <p>A strong ‘do not do’ recommendation was made for this review question because, despite the small amount of evidence, the GDG was confident that aspirin would not be of sufficient benefit for the majority of patients with type 2 diabetes who had not previously experienced a cardiovascular event. A strong recommendation was felt to be justified because the potential harm associated with the off-label use of aspirin (such as bleeding) outweighed the benefits (such as reduction in cardiovascular events). Although it was acknowledged that the review only identified studies on aspirin, the Group thought that the recommendation should be extended to include all off-label use of antiplatelet therapy, as it had seen no evidence of the effectiveness and safety of other drugs. The Group agreed that the most appropriate thing to do, in this circumstance, was to assume that all options have similar benefits and harms. The Group discussed the possibility of making no recommendation on the use of clopidogrel; however, the concern was expressed that, when set against the ‘do not do’ recommendation for aspirin, this might be read as tacit approval of clopidogrel, which the GDG was keen to avoid.</p>

The GDG noted that a cross-reference to other NICE guidance addressing the use of antiplatelet medicines for secondary prevention of cardiovascular disease was important to ensure healthcare professionals used these drugs as appropriate when caring for patients who have experienced a cardiovascular event.

## 7.3.1 Recommendations and research recommendations

- 2 **30. Do not offer antiplatelet therapy (aspirin or clopidogrel) for adults with type 2**  
3 **diabetes without cardiovascular disease. [new 2015]**
- 4 **31. For guidance on the primary and secondary prevention of cardiovascular disease,**  
5 **see the NICE guidelines on lipid modification and myocardial infarction –**  
6 **secondary prevention. [new 2015]**
- 7 **Research recommendations**
- 8 No research recommendations were made in relation to this review question.

Update 2015

## 8<sub>1</sub> Blood glucose management

2 The risks of arterial disease and microvascular complications in people with diabetes are  
3 thought to be related to the extent of hyperglycaemia over time. A comprehensive approach  
4 to blood glucose management incorporating education, assessment, self-monitoring and  
5 pharmacological strategies is required to facilitate optimal care. The chapter focuses on  
6 these key areas to provide guidance on optimal target values for blood glucose measures  
7 (sections 8.1 and 8.2), the use of self-monitoring to improve glycaemic control (section 8.3)  
8 and the effectiveness of different pharmacological interventions (section 8.4 and 8.5).

### 8.1<sub>9</sub> Optimal target values for blood glucose measures

#### 8.1.1<sub>0</sub> Clinical introduction

11 This section addresses the clinical question of what blood glucose values should be targeted  
12 to reduce the risk of future vascular damage from diabetes. It also aims to explore the impact  
13 of different blood glucose lowering drug treatments on optimal target values and the nature of  
14 the relationship between target values and specific subgroups of the population.

#### 8.1.1.1<sub>5</sub> Target values in Clinical Guideline 66

16 Target values for HbA1c were covered as part of CG66. However, fasting blood glucose and  
17 postprandial blood glucose target values were not included in CG66. The original searches  
18 were conducted from 2001 to 2007 to include systematic reviews, RCTs with sample sizes of  
19 at least 2000 and observational studies. CG66 included 1 meta-analysis (Selvin et al. 2004),  
20 1 RCT (UK Prospective Diabetes Study, UKPDS; Adler et al. 1999) and 2 observational  
21 studies (Gerstein et al. 2005; Iribarren et al. 2001).

#### 8.1.1.2<sub>2</sub> Target values in the update (2015)

23 For this update, several amendments were made to the review strategy. The sample size  
24 threshold applied in CG66 was removed as it was considered arbitrary and possibly  
25 inappropriate for specific population subgroups where participant numbers may be lower  
26 such as older people and different ethnic groups. As the question focused on elucidating the  
27 optimal blood glucose targets to reduce long-term macrovascular and microvascular  
28 complications in people with type 2 diabetes, studies which included rosiglitazone were  
29 excluded, as its association with cardiovascular mortality has the potential to confound the  
30 review findings. Similarly, studies with mixed populations of type 1 and type 2 diabetes  
31 patients were excluded as small numbers of people with type 1 diabetes may bias  
32 findings: the interventions used to manage diabetes are different and the long-term risk of  
33 cardiovascular disease may be different between type 1 and type 2 diabetes. Only  
34 prospective cohort studies that examined the development of long-term complications and its  
35 association with blood glucose measures were included.

36 The update review searches were completed in June 2014 with no date restriction for the  
37 following glycaemic measures: HbA1c, fasting blood glucose and postprandial blood glucose.

#### 8.1.2<sub>8</sub> Evidence review

#### 8.1.2.1<sub>9</sub> Review question

40 What are the optimal target values for HbA1c, fasting blood glucose and postprandial blood  
41 glucose in people with type 2 diabetes?

1 **Table 22: PICO table**

Population	Adults (18 years and over) with type 2 diabetes
Predictors	HbA1c, fasting blood glucose, postprandial blood glucose
Outcomes	Development of microvascular and macrovascular complications: <ul style="list-style-type: none"> <li>• retinopathy (specific lesions or macular changes, referable retinopathy, blindness/loss of vision, visual acuity)</li> <li>• kidney damage (eGFR, serum creatinine, proteinuria, microalbuminuria, dialysis)</li> <li>• cardiovascular disease (myocardial infarction, heart failure, stroke, ACS, TIA, revascularisation and stenting)</li> <li>• foot complications (amputations, diabetic foot ulcers, Charcot osteoarthropathy, diabetic foot infection)</li> </ul> Mortality

- 2 Prospective, longitudinal, cohort studies focusing on the development of microvascular or  
3 macrovascular complications and its association with blood glucose measures were  
4 included. Papers were excluded if they:
- 5 • were cross-sectional, case series and retrospective observational studies or conference  
6 abstracts, letters and editorials
  - 7 • exploratory prognostic studies that examined blood glucose measures as one of many risk  
8 factors for diabetes-related complications
  - 9 • focused on an association between blood glucose measures and microvascular or  
10 macrovascular complications without giving further information about the association
  - 11 • focused only on an association between the variability of blood glucose measures (for  
12 example HbA1c-coefficient of variation, HbA1c-standard deviation) and long-term  
13 complications
  - 14 • included a mixed population of people with type 1 and 2 diabetes, unless relevant  
15 subgroup analyses were reported
  - 16 • included rosiglitazone as part of the drug treatment strategy.
- 17 For the full excluded list, see Appendix L. The detailed protocol is also available in Appendix  
18 C.

### 8.1.2.29 Clinical evidence

- 20 The evidence that was originally included in CG66 was re-reviewed as part of the update. All  
21 included studies in the meta-analysis (Selvin et al. 2004) were checked against the update  
22 protocol resulting in only 1 relevant study (Adler et al. 1999) which was also identified in the  
23 update search. Consequently, original publications of the UKPDS study were used and the  
24 meta-analysis was excluded. The 2 observational studies in CG66 were excluded as they  
25 included people with both type 1 and type 2 diabetes (Iribarren et al. 2001) or did not specify  
26 the type of diabetes (Gerstein et al. 2005).
- 27 In total, 14,660 references were found in the update searches and 14 studies were included  
28 (Adler et al. 1999; Drechsler et al. 2009; Eeg-Olofsson et al. 2010; Hsu et al. 2012; Hunt et  
29 al. 2013; Landman et al. 2010; Molyneaux et al. 1998; Morisaki et al. 1994; Nakagami et al.  
30 1997; Salinero-Fort et al. 2013; Schulze et al. 2004; Torffvit and Agardh 2001; Zhao et al.  
31 2013; Zoungas et al. 2012).
- 32 The evidence included studies that reported outcomes in specific subgroup populations:
- 33 • 4 studies provided data for older people [60 years and over (Morisaki et al. 1994; Zhao et  
34 al. 2013), 65 years and over (Zoungas et al. 2012) and over 75 years (Landman et al.  
35 2010)]
  - 36 • 2 studies reported data based on sex (Zhao et al. 2013; Zoungas et al. 2012)

- 1 • 1 study reported data based on ethnicity (Hunt et al. 2013)
  - 2 • 2 studies reported data based on duration of diabetes [7 years and over (Eeg-Olofsson et  
3 al. 2010; Zoungas et al. 2012)]
  - 4 • 2 studies reported data based on microvascular or macrovascular disease status  
5 [cardiovascular disease (Eeg-Olofsson et al. 2010); microvascular and macrovascular  
6 disease (Zoungas et al. 2012)]
  - 7 • 1 study included people who were receiving dialysis (Drechsler et al. 2009)
- 8 The included studies all reported HbA1c as the main blood glucose measure or indicator.  
9 Outcomes were reported in relation to varying aspects of HbA1c including HbA1c at baseline  
10 and updated mean baseline HbA1c across the entire follow-up period. Where available,  
11 HbA1c at baseline was preferred. The associated risks of outcomes occurring depending on  
12 HbA1c were explored as a categorical variable, that is, at different threshold values of  
13 HbA1c. For example, Dreschler et al. 2009 and Zhao et al. 2013 reported results using  
14 reference HbA1c of 42 mmol/mol (6.0%) or less, Adler et al. 1999 used 45 mmol/mol (6.3%)  
15 or less, Eeg-Olofsson et al. 2010 used 42 to 52 mmol/mol (6.0 to 6.9%), Landman et al. 2010  
16 used 48 to 53 mmol/mol (6.5 to 7.0%), Salinero-Fort et al. 2013 used 53 mmol/mol (7.0%) or  
17 less and Hunt et al. 2013 used 53 to 64 mmol/mol (7.0 to 8.0%). Other studies explored the  
18 association of risks of outcomes with a continuous variable (for example 11 mmol/mol (1%)  
19 increase or decrease in HbA1c). Owing to the different reference HbA1c values and analyses  
20 used to address confounding variables in the included studies, pooling of data was not  
21 possible and individual studies were assessed using the modified GRADE approach (see  
22 section 3.7.3).
- 23 Two studies also explored the identification of specific threshold values for HbA1c. Zoungas  
24 et al. 2012 examined the non-linear relationship between HbA1c and risk of the outcomes of  
25 all-cause mortality, microvascular and macrovascular events and identified HbA1c thresholds  
26 above which risk increased; this was considered to be 48 to 53 mmol/mol (6.5 to 7.0%) for  
27 macrovascular disease and for mortality, and 42 to 48 mmol/mol (6.0 to 6.5%) for  
28 microvascular disease. Analysis of the UKPDS trial (Adler et al. 1999) found no indication of  
29 a threshold for mortality or any complication below which risk no longer decreased or a level  
30 above which risk no longer increased.
- 31 One study (Adler et al. 1999) reported on fasting blood glucose but no studies reported on  
32 postprandial blood glucose.

### 8.1.2.2.33 **Description of included studies**

34 A total of 968,656 people (study size ranged from 114 to 892,223) were included from 14  
35 prospective cohort studies, carried out in the UK (Adler et al. 1999; Zoungas et al. 2012), the  
36 Netherlands (Landman et al. 2010), Spain (Salinero-Fort et al. 2013), Germany (Drechsler et  
37 al. 2009), Sweden (Eeg-Olofsson et al. 2010; Torffvit and Agardh 2001), USA (Hunt et al.  
38 2013; Schulze et al. 2004; Zhao et al. 2013), Australia (Molyneaux et al. 1998), Japan  
39 (Morisaki et al. 1994; Nakagami et al. 1997) and Taiwan (Hsu et al. 2012). The mean age in  
40 13 studies ranged from 49.9 to 68.7 years, with 1 study not reporting this information (Adler  
41 et al. 1999). Mean HbA1c at baseline in 13 studies ranged from 50 to 81 mmol/mol (6.7% to  
42 9.6%), with 1 study not reporting this information (Adler et al. 1999). The median follow-up in  
43 the studies ranged from 28 months to 10.4 years. Details of the included studies are found in  
44 the evidence tables (see Appendix E).

45 Summary GRADE tables for this review question are presented below (see Appendix D for  
46 full GRADE tables).

47

1 Table 23: Summary GRADE profile for optimal target values for HbA1c in relation to mortality

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
<b>All-cause mortality</b>			
1 (Landman 2010) – ZODIAC 5 to 10 year follow-up	1145	<p>Categorical with 6.5-7.0% as a reference:            &lt; 6.5% HR 1.11 (0.71, 1.74)            7 to 8% HR 1.40 (0.99, 1.97)            8 to 9% HR 1.43 (0.97, 2.10)            ≥ 9% HR 2.26 (1.39, 3.67)</p> <p>Per 1% HbA1c decrease:            updated mean baseline HbA1c: HR 1.21 (1.07, 1.36)</p> <p><u>Subgroup</u>: age &gt;75 years (n=374)            Per 1% HbA1c increase:            &lt; 5 years diabetes duration: HR 1.51 (1.17, 1.95)            5 to 11 years diabetes duration: HR 1.04 (0.84, 1.28)            ≥ 11years diabetes duration: HR 1.05 (0.85, 1.30)</p>	High
1 (Adler 1999) – UKPDS Median 10.4 year follow-up	3642	<p>Per 1% HbA1c decrease:            Risk reduction baseline HbA1c: 6% (2, 10)</p>	High
1 (Zoungas 2012) – ADVANCE Mean 4.5 year follow-up	11,086 (event rate not reported)	<p>&lt; 7%: HR 1.01 (0.85, 1.21)            &gt; 7%: HR 1.38 (1.29, 1.48)</p> <p>Per 1% HbA1c increase:            6.0%: HR 1.35 (1.27, 1.43)            6.5%: HR 1.38 (1.29, 1.46)            7.0%: HR 1.38 (1.29, 1.48)            7.5%: HR 1.38 (1.27, 1.49)</p> <p>Per 1% HbA1c decrease:            6.0%: HR 0.36 (.21, 0.62)</p>	Moderate

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
		<p>6.5%: HR 0.73 (0.55, 0.96)            7.0%: HR 1.01 (0.85, 1.21)            7.5%: HR 1.16 (1.02, 1.32)</p> <p><u>Subgroup</u>: age &lt;65 years (<i>n</i> not reported)            Per 1% HbA1c increase:            &gt; 7%: HR 1.33 (1.16, 1.53)</p> <p><u>Subgroup</u>: age ≥65 years (<i>n</i> not reported)            Per 1% HbA1c increase:            &gt; 7%: HR 1.40 (1.30, 1.52)</p> <p><u>Subgroup</u>: male (<i>n</i>=6383)            Per 1% HbA1c increase:            &gt; 7%: HR 1.32 (1.20, 1.44)</p> <p><u>Subgroup</u>: female (<i>n</i>=4703)            Per 1% HbA1c increase:            &gt; 7%: HR 1.45 (1.31, 1.61)</p> <p><u>Subgroup</u>: duration of diabetes &lt;7 years (<i>n</i> not reported)            Per 1% HbA1c increase:            &gt; 7%: HR 1.51 (1.33, 1.71)</p> <p><u>Subgroup</u>: duration of diabetes ≥7 years (<i>n</i> not reported)            Per 1% HbA1c increase:            &gt; 7%: HR 1.33 (1.22, 1.45)</p> <p><u>Subgroup</u>: no macrovascular disease (<i>n</i>~7514)            Per 1% HbA1c increase:            &gt; 7%: HR 1.35 (1.24, 1.47)</p>	

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
		<p><u>Subgroup:</u> macrovascular disease (n=3572) Per 1% HbA1c increase: &gt; 7%: HR 1.42 (1.27, 1.59)</p> <p><u>Subgroup:</u> no microvascular disease (n=9933) Per 1% HbA1c increase: &gt; 7%: HR 1.37 (1.26, 1.49)</p> <p><u>Subgroup:</u> microvascular disease (n=1153) Per 1% HbA1c increase: &gt; 7%: HR 1.42 (1.25, 1.62)</p>	
<p>1 (Eeg-Olofsson 2010) 5 to 6 year follow-up</p>	<p>18,334</p>	<p>Categorical with 6.0-6.9% as a reference: 7.0 to 7.9% HR 1.08 (0.95 to 1.23) 8.0 to 8.9% HR 1.19 (1.03 to 1.38), p=0.02</p> <p>Per 1% HbA1c increase: Baseline HbA1c: HR 1.09 (1.05, 1.14), p&lt;0.001</p> <p><u>Subgroup:</u> duration of diabetes ≤7 years (n=10,016) Per 1% HbA1c increase: Baseline HbA1c: HR 1.13 (1.05, 1.21)</p> <p><u>Subgroup:</u> duration of diabetes &gt;7 years (n=8318) Per 1% HbA1c increase: Baseline HbA1c: HR 1.07 (1.01, 1.13)</p> <p><u>Subgroup:</u> previous cardiovascular disease (n=3276) Per 1% HbA1c increase: Baseline HbA1c: HR 1.08 (1.01, 1.15)</p>	<p>Moderate</p>

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
		<p><u>Subgroup</u>: no previous cardiovascular disease (n=15,058) Per 1% HbA1c increase: Baseline HbA1c: HR 1.10 (1.04, 1.16)</p>	
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	<p>Categorical with ≤6% as a reference: &gt; 6 to ≤8% HR 1.34 (1.10, 1.63) &gt; 8% HR 1.34 (1.02, 1.76)</p> <p>Per unit increase in HbA1c: HR 1.09 (1.02 to 1.17)</p>	Moderate
1 (Hunt 2013) Mean 4.4 year follow-up	892,223	<p><b>Non-Hispanic White (n=548,808)</b> Categorical with 7.0 to 8.0% as a reference: &lt; 7.0% HR 0.99 (0.97, 1.00) 8.0 to 9.0% HR 1.10 (1.08, 1.13) ≥ 9.0% HR 1.17 (1.14, 1.20)</p> <p><b>Non-Hispanic Black (n=108,356)</b> Categorical with 7.0 to 8.0% as a reference: &lt; 7.0% HR 1.07 (1.02, 1.12) 8.0-9.0% HR 1.00 (0.94, 1.06) ≥ 9.0% HR 1.09 (1.03, 1.15)</p> <p><b>Hispanic (n=123,670)</b> Categorical with 7.0 to 8.0% as a reference: &lt; 7.0% HR 1.02 (0.95, 1.09) 8.0-9.0% HR 1.09 (1.00, 1.19) ≥ 9.0% HR 1.15 (1.06, 1.25)</p> <p><b>Other (n=111,389)</b> Categorical with 7.0 to 8.0% as a reference: &lt; 7.0% HR 0.92 (0.87, 0.97) 8.0-9.0% HR 1.25 (1.16, 1.35)</p>	Moderate

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
		≥ 9.0% HR 1.30 (1.20, 1.40)	
<b>Mortality related to diabetes</b>			
1 (Adler 1999) – UKPDS Median 10.4 year follow-up	3642	Per 1% HbA1c decrease: Risk reduction baseline HbA1c: 9% (3, 14)	High
<b>Sudden death</b>			
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	Categorical with ≤6% as a reference: > 6 to ≤8% HR 1.85 (1.22, 2.81) > 8% HR 2.26 (1.33, 3.85)  Per unit increase in HbA1c: HR 1.21 (1.06 to 1.38)	Moderate
<b>Mortality except for sudden death</b>			
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	Categorical with ≤6% as a reference: > 6 to ≤ 8% HR 1.19 (0.96, 1.50) > 8% HR 1.10 (0.80, 1.52)  Per unit increase in HbA1c: HR 1.04 (0.96 to 1.13)	Moderate
<b>Cardiovascular mortality</b>			
1 (Landman 2010) – ZODIAC 5 to 10 year follow-up	1145	Categorical with 6.5 to 7.0% as a reference: < 6.5% HR 0.94 (0.47, 1.91) 7 to 8% HR 1.40 (0.84, 2.31) 8 to 9% HR 1.71 (0.99, 2.96) ≥ 9% HR 3.13 (1.62, 6.05)  <u>Subgroup:</u> age >75 years (n=374) Per 1% HbA1c increase: < 5 years diabetes duration: HR 1.72 (1.19, 2.48) 5 to 11 years diabetes duration: HR 1.18 (0.87, 1.60) ≥ 11 years diabetes duration: HR 1.16 (0.86, 1.58)	Moderate

Number of cohort studies	Number of people	Relative effect (95% CI)	Quality
1 (Eeg-Olofsson 2010) 5 to 6 year follow-up	18,334	<p>Categorical with 6.0 to 6.9% as a reference: 7.0 to 7.9% HR 1.11 (0.96 to 1.29) 8.0 to 8.9% HR 1.27 (1.07 to 1.50)</p> <p>Per 1% HbA1c increase: HR baseline HbA1c: 1.10 (1.05, 1.16)</p> <p><u>Subgroup</u>: duration of diabetes ≤7 years (n=10,016) Per 1% HbA1c increase: Baseline HbA1c: HR 1.14 (1.05, 1.24)</p> <p><u>Subgroup</u>: duration of diabetes &gt;7 years (n=8318) Per 1% HbA1c increase: Baseline HbA1c: HR 1.07 (1.01, 1.14)</p> <p><u>Subgroup</u>: previous cardiovascular disease (n=3276) Per 1% HbA1c increase: Baseline HbA1c: HR 1.09 (1.01, 1.17)</p> <p><u>Subgroup</u>: no previous cardiovascular disease (n=15,058) Per 1% HbA1c increase: Baseline HbA1c: HR 1.11 (1.04, 1.19)</p>	Moderate
1 (Drechsler 2009) - 4D study Heart failure death Median 4 year follow-up	1255	<p>Categorical with ≤6% as a reference: &gt; 6 to ≤ 8% HR 1.53 (0.70, 3.33) &gt; 8% HR 2.12 (0.75, 5.98)</p> <p>Per unit increase in HbA1c: HR 1.30 (1.00 to 1.68)</p>	Low

Abbreviations: HR hazard ratio; n number of people

1 **Table 24: Summary GRADE profile for optimal target values for HbA1c in relation to macrovascular complications**

Number of cohort studies	Number of people	Effect (95% CI)	Quality
<b>Composite of combined cardiovascular events</b>			
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	Categorical with ≤6% as a reference: > 6 to ≤ 8% HR 1.31 (1.05, 1.65) > 8% HR 1.37 (1.00, 1.87)  Per unit increase in HbA1c: HR 1.09 (1.01 to 1.18)	Moderate
<b>Macrovascular events</b>			
1 (Zoungas 2012) – ADVANCE Mean 4.5 year follow-up	11,086 (event rate not reported)	< 7%: HR 1.02 (0.86, 1.21) > 7%: HR 1.38 (1.30, 1.47)  Per 1% HbA1c increase: 6.0%: HR 1.35 (1.27, 1.42) 6.5%: HR 1.37 (1.29, 1.45) 7.0%: HR 1.38 (1.30, 1.47) 7.5%: HR 1.39 (1.29, 1.50)  Per 1% HbA1c decrease: 6.0%: HR 0.41 (0.25, 0.68) 6.5%: HR 0.77 (0.59, 1.00) 7.0%: HR 1.02 (0.86, 1.21) 7.5%: HR 1.13 (1.00, 1.28)  <u>Subgroup</u> : age <65 years ( <i>n</i> not reported) Per 1% HbA1c increase: > 7%: HR 1.34 (1.19, 1.50)  <u>Subgroup</u> : age ≥65 years ( <i>n</i> not reported) Per 1% HbA1c increase: > 7%: HR 1.40 (1.30, 1.51)	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p><u>Subgroup: male (n=6383)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.38 (1.27, 1.50)</p> <p><u>Subgroup: female (n=4703)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.35 (1.23, 1.48)</p> <p><u>Subgroup: duration of diabetes &lt;7 years (n not reported)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.54 (1.38, 1.72)</p> <p><u>Subgroup: duration of diabetes ≥7 years (n not reported)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.30 (1.21, 1.41)</p> <p><u>Subgroup: no macrovascular disease (n~7514)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.37 (1.26, 1.49)</p> <p><u>Subgroup: macrovascular disease (n=3572)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.38 (1.25, 1.52)</p> <p><u>Subgroup: no microvascular disease (n~9933)</u> Per 1% HbA1c increase: &gt; 7%: HR 1.37 (1.27, 1.48)</p> <p><u>Subgroup: microvascular disease (n=1153)</u> Per 1% HbA1c increase:</p>	

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		> 7%: HR 1.44 (1.27, 1.62)	
<b>Cardiovascular disease (fatal/non-fatal)</b>			
1 (Eeg-Olofsson 2010) 5 to 6 year follow-up	18,334	<p>Categorical with 6.0 to 6.9% as a reference: 7.0 to 7.9% HR 1.18 (1.08 to 1.29) 8.0 to 8.9% HR 1.31 (1.18 to 1.45)</p> <p>Per 1% HbA1c increase: Baseline HbA1c: HR 1.10 (1.07, 1.13)</p> <p><u>Subgroup</u>: duration of diabetes ≤7 years (n=10,016) Per 1% HbA1c increase: Baseline HbA1c: HR 1.08 (1.03, 1.13)</p> <p><u>Subgroup</u>: duration of diabetes &gt;7 years (n=8318) Per 1% HbA1c increase: Baseline HbA1c: HR 1.10 (1.06, 1.14)</p> <p><u>Subgroup</u>: previous cardiovascular disease (n=3276) Per 1% HbA1c increase: Baseline HbA1c: HR 1.10 (1.05, 1.16)</p> <p><u>Subgroup</u>: no previous cardiovascular disease (n=15,058) Per 1% HbA1c increase: Baseline HbA1c: HR 1.09 (1.06, 1.13)</p>	Moderate
<b>Myocardial infarction (fatal and non-fatal)</b>			
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	<p>Categorical with ≤6% as a reference: &gt; 6 to ≤ 8% HR 0.94 (0.68, 1.30) &gt; 8% HR 0.77 (0.47, 1.26)</p> <p>Per unit increase in HbA1c:</p>	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		HR 0.94 (0.83 to 1.07)	
1 (Adler 1999) – UKPDS Median 10 to 10.4 year follow-up	3845	Categorical with $\leq 6.3\%$ as a reference: > 6.3 to $\leq 7.6$ HR 1.2 (0.9, 1.5) > 7.6 HR 1.5 (1.2, 1.8)  Per 1% HbA1c decrease (n=3642): Risk reduction baseline HbA1c: 5% (0, 9)	High
<b>Coronary heart disease (fatal/non-fatal)</b>			
1 (Eeg-Olofsson 2010) 5 to 6 year follow-up	18,334	Categorical with 6.0 to 6.9% as a reference: 7.0 to 7.9% HR 1.25 (1.11 to 1.39) 8.0 to 8.9% HR 1.36 (1.20 to 1.55)  Per 1% HbA1c increase: HR baseline HbA1c: 1.11 (1.07, 1.15)  <u>Subgroup</u> : duration of diabetes $\leq 7$ years (n=10,016) Per 1% HbA1c increase: Baseline HbA1c: HR 1.09 (1.03, 1.15)  <u>Subgroup</u> : duration of diabetes >7 years (n=8318) Per 1% HbA1c increase: Baseline HbA1c: HR 1.11 (1.06, 1.16)  <u>Subgroup</u> : previous cardiovascular disease (n=3276) Per 1% HbA1c increase: Baseline HbA1c: HR 1.08 (1.02, 1.15)  <u>Subgroup</u> : no previous cardiovascular disease (n=15,058) Per 1% HbA1c increase: Baseline HbA1c: HR 1.12 (1.07, 1.16)	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
1 (Schulze 2004) Mean 7.4 year follow-up	921	Categorical into quartiles of median HbA1c with 5.21% as a reference: 5.80% RR 2.49 (1.19, 5.23) 6.90% RR 3.19 (1.56, 6.53) 8.97% RR 4.92 (2.46, 9.85)	Very low
<b>Heart failure</b>			
1 (Adler 1999) – UKPDS Median 10.4 years	3642	Per 1% HbA1c decrease: Risk reduction baseline HbA1c: 0% (-12, 11)	High
<b>Newly diagnosed angina</b>			
1 (Adler 1999) – UKPDS Median 10 to 10.3 years	3836	Categorical with ≤6.3% as a reference: > 6.3 to ≤ 7.6 HR 1.5 (1.1, 2.0) > 7.6 HR 1.6 (1.1, 2.1)	High
<b>Stroke (fatal and non-fatal)</b>			
1 (Drechsler 2009) - 4D study Median 4 year follow-up	1255	Categorical with ≤6% as a reference: > 6 to ≤ 8% HR 1.56 (0.93, 2.62) > 8% HR 1.67 (0.84, 3.30)  Per unit increase in HbA1c: HR 1.11 (0.93 to 1.32)	Low
1 (Eeg-Olofsson 2010) 5 to 6 year follow-up	18,334	Per 1% HbA1c increase: HR baseline HbA1c: 1.08 (1.03, 1.13)  <u>Subgroup</u> : duration of diabetes ≤7 years (n=10,016) Per 1% HbA1c increase: Baseline HbA1c: HR 1.06 (0.98, 1.14)  <u>Subgroup</u> : duration of diabetes >7 years (n=8318) Per 1% HbA1c increase: Baseline HbA1c: HR 1.07 (1.01, 1.14)  <u>Subgroup</u> : previous cardiovascular disease (n=3276)	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p>Per 1% HbA1c increase: Baseline HbA1c: HR 1.11 (1.03, 1.20)</p> <p><u>Subgroup</u>: no previous cardiovascular disease (n=15,058) Per 1% HbA1c increase: Baseline HbA1c: HR 1.06 (1.00, 1.12)</p>	
1 (Adler 1999) – UKPDS Median 10 to 10.3 years	3670	<p>Categorical with ≤6.3% as a reference: &gt; 6.3 to ≤ 7.6 HR 1.2 (0.8, 1.7) &gt; 7.6 HR 1.1 (0.7, 1.6)</p> <p>Per 1% HbA1c decrease (n=3642): Risk reduction baseline HbA1c: -4% (-14, 6)</p>	High
<b>Peripheral vascular disease</b>			
1 (Adler 1999) – UKPDS Median 10.4 years	2398	<p>Per 1% HbA1c increase: OR 1.28 (1.12, 1.46)</p> <p><u>Amputation or peripheral vascular disease death (n=3642) :</u> Per 1% HbA1c decrease: Risk reduction baseline HbA1c: 28% (18, 37)</p>	High
1 (Zhao 2013) – LSUHLS study Lower limb amputation Mean 6.83 year follow-up	35,368	<p><u>African Americans (n=19,808)</u> Categorical with &lt;6% as a reference and baseline HbA1c: 6.0 to 6.9% HR 1.73 (1.07, 2.80) 7.0 to 7.9% HR 1.65 (0.99, 2.77) 8.0 to 8.9% HR 1.96 (1.14, 3.36) 9.0 to 9.9% HR 3.02 (1.81, 5.04) ≥ 10% HR 3.30 (2.10, 5.20)</p> <p>Per 1% HbA1c increase: Baseline HbA1c: HR 1.12 (1.08, 1.17)</p>	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p><u>Caucasians (n=15,560)</u> Categorical with &lt;6% as a reference and baseline HbA1c: 6.0 to 6.9% HR 1.16 (0.66, 2.02) 7.0 to 7.9% HR 2.28 (1.35, 3.85) 8.0 to 8.9% HR 2.38 (1.36, 4.18) 9.0 to 9.9% HR 2.99 (1.71, 5.22) ≥10% HR 3.25 (1.98, 5.33)</p> <p>Per 1% HbA1c increase: Baseline HbA1c: HR 1.15 (1.09, 1.21)</p> <p><u>Subgroup: male (n=13,363 at baseline)</u> Categorical with &lt;6% as a reference and baseline HbA1c: 6.0 to 6.9% HR 1.48 (0.95, 2.26) 7.0 to 7.9% HR 1.85 (1.20, 2.85) 8.0 to 8.9% HR 2.19 (1.40, 3.42) 9.0 to 9.9% HR 3.15 (2.04, 4.85) ≥ 10% HR 2.84 (1.93, 4.17)</p> <p><u>Subgroup: female (n=22,005 at baseline)</u> Categorical with &lt;6% as a reference and baseline HbA1c: 6.0 to 6.9% HR 1.63 (0.80, 3.32) 7.0 to 7.9% HR 2.37 (1.17, 4.80) 8.0 to 8.9% HR 2.26 (1.04, 4.91) 9.0 to 9.9% HR 3.43 (1.63, 7.24) ≥ 10% HR 4.96 (2.50, 9.71)</p> <p><u>Subgroup: age 60-94 years (n not reported)</u> Categorical with &lt;6% as a reference and baseline HbA1c: 6.0 to 6.9% HR 2.02 (0.94, 4.35) 7.0 to 7.9% HR 3.19 (1.42, 7.18)</p>	

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p>8.0 to 8.9% HR 3.06 (1.18, 7.95)            9.0 to 9.9% HR 2.37 (0.80, 7.01)            ≥ 10% HR 3.19 (1.27, 8.00)</p> <p><u>Subgroup:</u> age 50-59 years (<i>n</i> not reported)            Categorical with &lt;6% as a reference and baseline HbA1c:            6.0 to 6.9% HR 1.13 (0.66, 1.94)            7.0 to 7.9% HR 1.50 (0.86, 2.63)            8.0 to 8.9% HR 2.26 (1.22, 4.18)            9.0 to 9.9% HR 3.69 (2.10, 6.47)            ≥ 10% HR 2.89 (1.73, 4.82)</p> <p><u>Subgroup:</u> age &lt;50 years (<i>n</i> not reported)            Categorical with &lt;6% as a reference and baseline HbA1c:            6.0 to 6.9% HR 1.80 (0.95, 3.43)            7.0 to 7.9% HR 2.41 (1.27, 4.57)            8.0 to 8.9% HR 2.34 (1.25, 4.38)            9.0 to 9.9% HR 3.01 (1.63, 5.57)            ≥ 10% HR 3.93 (2.26, 6.84)</p> <p><u>Subgroup:</u> previous use of blood glucose lowering medication (<i>n</i>=12,788)            Categorical with &lt;6% as a reference and baseline HbA1c:            6.0 to 6.9% HR 1.30 (0.72, 2.33)            7.0 to 7.9% HR 2.24 (1.26, 3.98)            8.0 to 8.9% HR 1.94 (0.97, 3.88)            9.0 to 9.9% HR 2.81 (1.43, 5.51)            ≥ 10% HR 2.73 (1.55, 4.82)</p> <p><u>Subgroup:</u> no previous use of blood glucose lowering medication (<i>n</i>=22,580)            Categorical with &lt;6% as a reference and baseline HbA1c:            6.0 to 6.9% HR 1.62 (1.02, 2.59)</p>	

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		7.0 to 7.9% HR 1.93 (1.20, 3.12) 8.0 to 8.9% HR 2.20 (1.36, 3.58) 9.0 to 9.9% HR 3.41 (2.14, 5.45) ≥ 10% HR 3.50 (2.28, 5.36)	

Abbreviations: HR hazard ratio; n number of people; OR odds ratio; RR relative risk

1 **Table 25: Summary GRADE profile for optimal target values for HbA1c in relation to microvascular complications**

Number of cohort studies	Number of people	Effect (95% CI)	Quality
<b>Microvascular end points</b>			
1 (Adler 1999) – UKPDS Median 10.4 years	3642	Per 1% HbA1c decrease: Risk reduction baseline HbA1c: 23% (20, 27)	High
1 (Zoungas 2012) – ADVANCE Mean 4.5 year follow-up	11,086 (event rate not reported)	HR < 6.5%: 1.02 (0.76, 1.39) HR > 6.5%: 1.40 (1.33, 1.47)  Per 1% HbA1c increase: 6.0%: HR 1.39 (1.32, 1.46) 6.5%: HR 1.40 (1.33, 1.47) 7.0%: HR 1.38 (1.30, 1.46) 7.5%: HR 1.33 (1.24, 1.42)  Per 1% HbA1c decrease: 6.0%: HR 0.67 (0.36, 1.23) 6.5%: HR 1.02 (0.76, 1.02) 7.0%: HR 1.33 (1.10, 1.60) 7.5%: HR 1.51 (1.32, 1.72)  <u>Subgroup</u> : age <65 years (n not reported) Per 1% HbA1c increase:	Moderate

Update 2015

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p>&gt; 6.5%: HR 1.40 (1.30, 1.50)</p> <p><u>Subgroup</u>: age ≥65 years (<i>n</i> not reported) Per 1% HbA1c increase: &gt; 6.5%: HR 1.39 (1.29, 1.50)</p> <p><u>Subgroup</u>: male (<i>n</i>=6383) Per 1% HbA1c increase: &gt; 6.5%: HR 1.42 (1.33, 1.52)</p> <p><u>Subgroup</u>: female (<i>n</i>=4703) Per 1% HbA1c increase: &gt; 6.5%: HR 1.39 (1.29, 1.50)</p> <p><u>Subgroup</u>: duration of diabetes &lt;7 years (<i>n</i> not reported) Per 1% HbA1c increase: &gt; 6.5%: HR 1.27 (1.14, 1.40)</p> <p><u>Subgroup</u>: duration of diabetes ≥7 years (<i>n</i> not reported) Per 1% HbA1c increase: &gt; 6.5%: HR 1.45 (1.36, 1.54)</p> <p><u>Subgroup</u>: no macrovascular disease (<i>n</i>~7514) Per 1% HbA1c increase: &gt; 6.5%: HR 1.44 (1.35, 1.53)</p> <p><u>Subgroup</u>: macrovascular disease (<i>n</i>=3572) Per 1% HbA1c increase: &gt; 6.5%: HR 1.30 (1.17, 1.43)</p> <p><u>Subgroup</u>: no microvascular disease (<i>n</i>~9933)</p>	

Number of cohort studies	Number of people	Effect (95% CI)	Quality
		<p>Per 1% HbA1c increase: &gt; 6.5%: HR 1.40 (1.32, 1.49)</p> <p><u>Subgroup</u>: microvascular disease (n=1153) Per 1% HbA1c increase: &gt; 6.5%: HR 1.36 (1.23, 1.50)</p>	
<b>Retinopathy</b>			
1 (Molyneaux 1998) Median 28 month follow-up	963	<p>Per 10% HbA1c decrease: Relative risk reduction: 24% (16, 32)</p>	Moderate
1 (Morisaki 1994) 5 year follow-up	114	<p>Multivariate logistic regression analysis showed that HbA1c was the only significant predictor of retinopathy</p> <p>Retinopathy prevalence at HbA1c: &lt; 7%: 2% ≥ 7 to &lt; 8%: 20% ≥ 8 to &lt; 9%: 40% ≥ 9%: 61%</p> <p>With retinopathy HbA1c 8.8±1.1 Without retinopathy HbA1c 7.1±1.2</p>	Very low
1 (Nakagami 1997) 10 year follow-up	137	<p>Retinopathy prevalence at HbA1c: &lt; 6%: 0% 6 to 6.9%: 17.2% 7 to 7.9%: 14.3% 8 to 8.9%: 41.9% ≥ 9%: 54.8%</p> <p>Multivariate logistic regression analysis showed that mean HbA1c over 10 year follow-up period was the only significant predictor of retinopathy</p>	Very low
1 (Salinero-Fort 2013) – MADIABETES	2405	<p>Categorical with &lt;7% as a reference: 7 to 8% HR 1.39 (1.01, 1.92)</p>	Moderate

Number of cohort studies	Number of people	Effect (95% CI)	Quality
4 year follow-up		> 8% HR 1.90 (1.30, 2.77)	
<b>Cataract extraction</b>			
1 (Adler 1999) – UKPDS Median 10.4 years	3642	Per 1% HbA1c decrease: Risk reduction baseline HbA1c: 9% (2, 16)	High
<b>Nephropathy</b>			
1 (Molyneaux 1998) Microalbuminuria Median 28 month follow-up	399	Per 10% HbA1c decrease: Relative risk reduction: 9% (-2, 19)	Very low
1 (Torffvit and Agardh 2001) Albuminuria Median 9 year follow-up	385	Cox regression analysis showed that HbA1c significantly predicted greater fractional albumin clearance ( $p < 0.01$ ) and development of renal failure ( $p < 0.05$ )  Normoalbuminuria mean HbA1c $7.8 \pm 1.5$ Micro/macro-albuminuria HbA1c $8.5 \pm 1.6$	Very low
1 (Hsu 2012) Microalbuminuria 5 to 7 year follow-up	821	Per 1% HbA1c decrease: Baseline HbA1c $\leq 8\%$ : HR 1.13 (0.91, 1.39) Baseline HbA1c $> 8\%$ : HR 1.18 (1.04, 1.34)	Moderate
<i>Abbreviations: HR hazard ratio; n number of people</i>			

1 **Table 26: Summary GRADE profile for optimal target values for fasting blood glucose in relation to macrovascular complications**

Number of cohort studies	Number of people	Effect (95% CI)	Quality
<b>Myocardial infarction (fatal and non-fatal)</b>			
1 (Adler 1999, UKPDS)† Median 10 to 10.3 year follow-up up	5045	Categorical with $\leq 9.7$ mmol/L as a reference: > 9.7 to $\leq 13.4$ HR 1.1 (0.9, 1.4) > 13.4 HR 1.3 (1.1, 1.6)	High
<b>Newly diagnosed angina</b>			
1 (Adler 1999, UKPDS)† Median 10 to 10.3 year follow-up	5036	Categorical with $\leq 9.7$ mmol/L as a reference: > 9.7 to $\leq 13.4$ HR 1.3 (1.0, 1.7) > 13.4 HR 1.2 (0.9, 1.5)	High

Number of cohort studies	Number of people	Effect (95% CI)	Quality
<b>Stroke (fatal and non-fatal)</b>			
1 (Adler 1999, UKPDS)‡ Median 10 to 10.3 year follow-up	5040	Categorical with $\leq 9.7$ mmol/L as a reference: > 9.7 to $\leq 13.4$ HR 1.3 (0.9, 1.7) > 13.4 HR 1.3 (1.0, 1.8)	High
<p><i>Abbreviations: HR hazard ratio</i>  <i>‡ Baseline data extracted at diagnosis only, not after dietary run-in. Model controlled for age at diabetes diagnosis, sex and ethnicity</i></p>			

Update 2015

### 8.1.2.31 Health economic evidence

2 Literature searches were undertaken to find any existing cost-utility analyses (CUAs) of  
3 intensive versus conventional blood glucose targets. Due to the similarity of the literature  
4 base, health economic evidence for review question 3 (target values) and review question 4  
5 (intensive versus conventional regimens) are presented together. The GDG discussed and  
6 noted that, given the nature of the CUAs presented, it was difficult to categorise them to  
7 either review question. See section 8.2.2.3 for the health economic evidence for this review  
8 question and section 8.2.2.4 for the health economic evidence statement.

### 8.1.2.49 Evidence statements

#### 8.1.2.4.10 Clinical evidence

##### 11 Optimal target values

12 One study found that risk significantly increased above HbA1c levels of 48 mmol/mol (6.5%)  
13 for microvascular complications and 53 mmol/mol (7%) for mortality and macrovascular  
14 complications. The quality of the evidence was moderate. The second study did not find a  
15 specific threshold for which risk increased or decreased for mortality or any diabetes-related  
16 complication. The evidence was of high quality.

##### 17 Mortality

18 Evidence from 6 studies found that all-cause mortality risk rose with increasing baseline  
19 levels of HbA1c. The quality of the evidence was moderate to high. Evidence from 3 studies  
20 found that a 11 mmol/mol (1%) decrease in HbA1c led to a lower risk of all-cause mortality,  
21 while a 11 mmol/mol (1%) increase was associated with an increased risk of all-cause  
22 mortality.

##### 23 Macrovascular complications

24 Evidence from 6 studies found that the risk of macrovascular complications (defined as a  
25 composite of combined cardiovascular end points, macrovascular events, cardiovascular  
26 disease, myocardial infarction, coronary heart disease, heart failure, newly diagnosed  
27 angina, stroke and peripheral vascular disease) rose with increasing levels of baseline  
28 HbA1c. The quality of the evidence ranged from high to very low.

29 Evidence from 1 study found that in general, people aged 60 to 94 years were at greater risk  
30 of lower limb amputations at the same baseline HbA1c compared to people aged less than  
31 59 years. The quality of the evidence was moderate.

32 Evidence from 1 study found that risk of myocardial infarction rose with increasing FBG  
33 levels, but there was no difference in the risk of stroke and angina with increasing FBG  
34 levels. The quality of the evidence was high.

##### 35 Microvascular complications

36 Evidence from 8 studies found that the risk of microvascular complications (defined as a  
37 composite of microvascular end points, retinopathy, cataract extraction and renal outcomes)  
38 rose with increasing levels of baseline HbA1c, or that study participants who developed the  
39 specified end point had higher levels of HbA1c than those who did not. The quality of the  
40 evidence ranged from high to very low.

#### 8.1.2.4.21 **Health economic evidence**

2 See section 8.2.2.4 for the health economic evidence statement.

### 8.1.33 **Evidence to recommendations**

#### 4 **Table 27: Linking evidence to recommendations**

Relative value of different outcomes	<p>The GDG agreed that the critical outcomes in determining the optimal target values for blood glucose measures are the risk of developing long-term diabetic complications (macrovascular and microvascular) and all-cause mortality.</p> <p>The GDG agreed that all outcomes should be weighted equally when deciding the optimal target values.</p>
Trade-off between benefits and harms	<p>The GDG recognised the trade-off between the increased benefits of setting target values for blood glucose to protect against long-term complications, and the possible associated harms (for example hypoglycaemia). The GDG agreed that overall, the evidence showed that rising levels of HbA1c increase the risk of mortality and developing macrovascular and microvascular complications, with critical thresholds ranging from 42 to 53 mmol/mol (6 to 7%). The GDG agreed that it was not possible to provide guidance on HbA1c levels less than 42 mmol/mol (6%), as only 1 very-low-quality study reported data for values ranging from 33 to 38 mmol/mol (5.21 to 5.8%). The Group discussed optimal target values for HbA1c, and agreed that a mid-range value of 48 mmol/mol (6.5%) would be achievable for most adults with type 2 diabetes that was managed by lifestyle and diet and/or 1 oral anti-diabetic drug. However, the GDG agreed that people who achieve this target HbA1c level using diet and exercise alone with no hypoglycaemic risk should be encouraged to safely attain lower levels if possible. The GDG discussed the progressive nature of the condition, and agreed that drug treatment should be intensified if HbA1c levels rose to 58 mmol/mol (7.5%) and considering the risk of hypoglycaemia, a realistic target of 53 mmol/mol (7%) should be set to achieve glycaemic control.</p> <p>The GDG also considered that, while guidance on target values was important, the complexities of individual patient needs should predominate. In particular, the Group thought that special consideration of appropriate target values should be given to people at risk of hypoglycaemia, to achieve an acceptable balance between good glycaemic control and the likely negative impact on quality of life of this adverse event. The GDG also discussed groups for whom the target levels may not be appropriate, such as people with renal failure, people for whom the target level may necessitate increased medication that may cause adverse events or decreased medication compliance, or people who would probably not benefit from the long-term impact on macrovascular and/or microvascular complications.</p>
Consideration of health benefits and resource use	<p>The GDG found the health economic evidence on optimal target values and intensive versus conventional control hard to distinguish. No CUAs gave direct evidence on whether a particular HbA1c target was more cost effective than another HbA1c target, but all the CUAs</p>

	<p>found intensive control at lower HbA1c targets to be more cost effective than less intensive control at higher HbA1c targets.</p>
<p>Quality of evidence</p>	<p>The GDG agreed that the evidence ranged from high to very low quality. The Group discussed that in the majority of studies, HbA1c categorical levels started from 42 to 48 mmol/mol (6 to 6.5%), but that in routine clinical practice, target levels less than 42 mmol/mol (6%) would not be set. The GDG noted that one of the studies on HbA1c included patients who were on dialysis, which was a specified subgroup of interest. However, the GDG agreed that patients with advanced complications were not a true representation of the average type 2 diabetes population, and that dialysis may affect the accuracy of HbA1c measurements.</p> <p>The GDG discussed the findings of the clinical review and noted that there was little or no evidence on fasting and postprandial blood glucose measures, and therefore agreed that it was not possible to set target values for these tests. However, the Group recognised the importance of these measures as they directly influence HbA1c levels.</p>
<p>Other considerations</p>	<p>The GDG noted that the mean age in the included studies ranged from 50 to 69 years and agreed that there was no evidence for younger adults with type 2 diabetes and limited evidence for those over the age of 70. The Group discussed whether there were different considerations in reviewing target values for these groups. The GDG thought that, when agreeing target values with adults with type 2 diabetes, it is more important to consider the nature of the individual's current medical condition – that is, diabetes, its complications and any other comorbidities – rather than age alone.</p> <p>The GDG agreed the importance of ensuring that all adults with type 2 diabetes are aware of the benefits associated with lowering HbA1c levels and achieving appropriate blood glucose targets with minimal fluctuation to maintain good glycaemic control. The Group agreed that target values, appropriate to the individual's situation, should be discussed and agreed with the patient to optimise care.</p> <p>The Group also noted that the Quality and Outcomes Framework (QOF) refer to 3 levels of glucose control in people with type 2 diabetes: HbA1c of 75 mmol/mol (9%) or less, 64 mmol/mol (8%) or less and 59 mmol/mol (7.5%) or less.</p>

#### 8.1.4.1 Recommendations and research recommendations

##### 8.1.4.1.2 HbA1c measurement and targets

##### 8.1.4.1.13 Measurement

#### 4 32. Measure HbA1c levels at:

- 5 • 3–6 monthly intervals (tailored to individual needs), until the HbA1c is
- 6 stable on unchanging therapy
- 7 • 6-monthly intervals once the HbA1c level and blood glucose lowering
- 8 therapy are stable. **[2015]**

- 1 **33. Calibrate HbA1c results according to International Federation of Clinical**  
2 **Chemistry (IFCC) standardisation. [new 2015]**
- 3 **34. If HbA1c monitoring is invalid (because of disturbed erythrocyte turnover or**  
4 **abnormal haemoglobin type), estimate trends in blood glucose control using one**  
5 **of the following:**
- 6 • fructosamine estimation
  - 7 • quality-controlled plasma glucose profiles
  - 8 • total glycated haemoglobin estimation (if abnormal haemoglobins).  
9 **[2015]**
- 10 **35. Investigate unexplained discrepancies between HbA1c and other glucose**  
11 **measurements. Seek advice from a team with specialist expertise in diabetes or**  
12 **clinical biochemistry. [2015]**

#### **8.1.4.1.23 Targets**

- 14 **36. Involve adults with type 2 diabetes in decisions about their individual HbA1c**  
15 **target. Encourage them to achieve the target and maintain it unless any resulting**  
16 **adverse effects (including hypoglycaemia), or their efforts to achieve their target,**  
17 **impair their quality of life. [new 2015]**
- 18 **37. Offer lifestyle advice and drug treatment to help adults with type 2 diabetes**  
19 **achieve and maintain their HbA1c target. See recommendations 8–16. For more**  
20 **information about supporting adherence, see the NICE guideline on medicines**  
21 **adherence. [new 2015]**
- 22 **38. Set a target HbA1c level of 48 mmol/mol (6.5%) for most adults with type 2**  
23 **diabetes that is managed either by lifestyle and diet, or by lifestyle and diet in**  
24 **combination with a single drug that is not associated with hypoglycaemia. [new**  
25 **2015]**
- 26 **39. If HbA1c levels rise to 58 mmol/mol (7.5%) or higher, intensify drug treatment, set**  
27 **a target HbA1c level of 53 mmol/mol (7.0%), and reinforce advice about diet,**  
28 **lifestyle and adherence to drug treatment. See section 5. For more information**  
29 **about supporting adherence, see the NICE guideline on medicines adherence.**  
30 **[new 2015]**
- 31 **40. Consider relaxing the target HbA1c level (see recommendations 38–39) on a case-**  
32 **by-case basis for adults with type 2 diabetes:**
- 33 • who are unlikely to achieve longer-term risk-reduction benefits (for  
34 example, people with a reduced life expectancy)
  - 35 • for whom tight glycaemic control poses risks
  - 36 • with a high risk of the consequences of hypoglycaemia (for example,  
37 people who are at risk of falling, people who have impaired awareness  
38 of hypoglycaemia, and people who drive or operate machinery as part of  
39 their job)
  - 40 • for whom intensive management would not be appropriate (for example,  
41 people taking multiple drugs and people with significant comorbidities).  
42 These factors will need particular consideration for people who are older  
43 and frail. **[new 2015]**

1 **41. If adults with type 2 diabetes achieve an HbA1c level that is lower than their target**  
2 **and they are not experiencing hypoglycaemia, encourage them to maintain it.**  
3 **[new 2015]**

4 **42. For guidance on HbA1c targets for women who are pregnant or planning to**  
5 **become pregnant, see the NICE guideline on diabetes in pregnancy. [new 2015]**

## 6 **Research recommendations**

7 **2. What is the natural history of individuals who are diagnosed with type 2 diabetes**  
8 **in childhood in terms of long-term complications/consequences in adulthood?**

### 9 **Why this is important**

10 Type 2 diabetes has historically been associated with adults, with research largely focused  
11 on this population. However, there is growing concern of the increasing incidence of type  
12 2 diabetes in younger people, thought to be linked to the rising levels of obesity. In order  
13 to improve clinical management of people diagnosed in childhood, a better  
14 understanding of the early progression of the condition is needed, particularly in terms of  
15 its effects on the long-term risks of developing microvascular and macrovascular  
16 complications. A prospective longitudinal 10 year cohort study of children diagnosed with  
17 type 2 diabetes would help improve understanding of whether diabetes spanning the  
18 growth spurt would result in long-term complications occurring at a different rate  
19 compared to individuals who are diagnosed during adulthood.

Update 2015

## 8.2.1 Intensive and conventional blood glucose targets

### 8.2.1.2 Clinical introduction

3 There has been a general acceptance that tight glycaemic control is beneficial in reducing  
4 the risk of cardiovascular disease. Evidence reported in the previous section (see 8.1)  
5 identified an increased risk of long-term complications associated with higher baseline  
6 HbA1c levels. The risk increased with each 11 mmol/mol (1%) rise in HbA1c levels and  
7 correspondingly decreased with each 11 mmol/mol (1%) fall in HbA1c levels. However, the  
8 impact of intensive control at lower target values on other outcomes such as hypoglycaemia  
9 compared to conventional control at higher targets is unclear.

10 This section addresses the clinical question of whether intensive strategies to lower target  
11 values are more effective than conventional strategies to higher targets in reducing long-term  
12 complications. It also aims to explore situations in which intensive strategies should be used  
13 and whether the effect of intensive strategies differs in specific subgroups of the population.

### 8.2.1.14 Intensive and conventional blood glucose targets in Clinical Guideline 66

15 CG66 did not report on the effectiveness of intensive glycaemic control compared to  
16 conventional glycaemic control.

### 8.2.1.27 Intensive and conventional blood glucose targets in the update (2015)

18 This is a new question in this update and therefore searches have been carried out for this  
19 topic without any date restrictions.

20 This review compared the use of intensive glycaemic control against conventional glycaemic  
21 control. The strategies used to achieve intensive and conventional glycaemic control could  
22 include oral antidiabetic agents and/or insulin. Outcomes of interest to the GDG included  
23 hypoglycaemic episodes, development of macrovascular and microvascular complications  
24 (retinopathy, kidney damage, cardiovascular disease, foot complications), mortality and  
25 changes in body weight.

### 8.2.26 Evidence review

#### 8.2.2.27 Review question

28 Should intensive or conventional target values be used to control blood glucose levels in  
29 people with type 2 diabetes?

#### 30 Table 28: PICO table

Population	Adults (18 years and over) with type 2 diabetes
Intervention	Intensive blood glucose control (using pharmacological blood glucose lowering therapies) with target blood glucose levels lower than conventional values
Comparator	Conventional target values (targets that would be considered to be in the normal range for adults with type 2 diabetes)
Outcomes	Hypoglycaemic events Development of microvascular and macrovascular complications: <ul style="list-style-type: none"> <li>• retinopathy (specific lesions or macular changes, referable retinopathy, blindness/loss of vision, visual acuity)</li> <li>• kidney damage (eGFR, serum creatinine, proteinuria, microalbuminuria, dialysis)</li> <li>• cardiovascular disease (myocardial infarction, heart failure, stroke, ACS, TIA,</li> </ul>

revascularisation and stenting)  
• foot complications (amputations, diabetic foot ulcers, Charcot osteoarthropathy, diabetic foot infection)  
Mortality  
Changes in body weight

1 <Insert Note here>

2 Systematic reviews and RCTs focusing on the use of intensive blood glucose control  
3 compared to conventional strategies were included. Papers were excluded if they:

- 4 • were non-randomised studies (including cohort, case–control and case series) or narrative  
5 reviews, conference abstracts, letters and editorials  
6 • included a mixed population of people with type 1 and 2 diabetes, unless relevant  
7 subgroup analyses were reported  
8 • included rosiglitazone as part of the drug treatment strategy. For the full excluded list, see  
9 Appendix L.

10 The main outcomes for this review question were the development of microvascular and  
11 macrovascular complications and adverse events. The detailed protocol is available in  
12 Appendix C.

### 8.2.2.23 Clinical evidence

14 This topic was not covered in CG66 so no date restrictions were placed on the search  
15 strategy (see Appendix C for update search strategies). A total of 1782 references were  
16 identified for this question, including a number of systematic reviews and meta-analyses. A  
17 recent Cochrane systematic review (Hemmingsen et al. 2013) included all relevant RCTs  
18 and therefore was the primary source of evidence for this question.

19 For the purposes of this question, the studies in the Cochrane review were assessed for  
20 relevance. RCTs where intensive and conventional glycaemic control groups had significant  
21 baseline differences in adjunctive treatment for cardiovascular risk factors were excluded.  
22 This led to the exclusion of 8 RCTs included in the Cochrane systematic review: ACCORD  
23 2008, ADDITION 2011, ADVANCE 2008, Araki 2012, Guo 2008, Steno-2 2008, VADT 2009  
24 and Yang 2007.

### 8.2.2.2.15 Description of included studies

26 Data from a Cochrane review was used to answer this question. This review included studies  
27 of adults (aged 18 years and older) with type 2 diabetes. The intensive control groups  
28 targeted HbA1c values ranging from 42 mmol/mol (6%) or less and up to 58 mmol/mol  
29 (7.5%) while the conventional control groups either had no target values or targeted HbA1c  
30 values above 42 mmol/mol (6%). The mean duration of the intervention period varied from 3  
31 days to 12.5 years. Details of the included review are found in the evidence tables (see  
32 Appendix E).

33 A summary GRADE table is presented for this review question (see Appendix D for full  
34 GRADE tables).

1 **Table 29: Summary GRADE profile for intensive versus conventional target values**

Number of studies	Number of people		Measure of effect	Quality
	Intensive	Conventional		
<b>All-cause mortality</b>				
1 systematic review (Hemmingsen 2013) including 16 RCTs (Bagg 2001, Cao 2011, DIGAMI 2 2005, Fantin 2011, IDA 2009, Jaber 1996, Kumamoto 2000, Melidonis 2000, Natarajan 2012, REMBO 2008, Service 1983, Stefanidis 2003, UGDP 1975, UKPDS 1998, VA CSDM 1995, Zhang 2011)	762/4296	381/2208	RR 0.98 (0.88 to 1.09)	High
<b>Cardiovascular mortality</b>				
1 systematic review (Hemmingsen 2013) including 14 RCTs (Bagg 2001, Cao 2011, DIGAMI 2 2005, IDA 2009, Jaber 1996, Kumamoto 2000, Melidonis 2000, REMBO 2008, Service 1983, Stefanidis 2003, UGDP 1975, UKPDS 1998, VA CSDM 1995, Zhang 2011)	445/4225	195/2131	RR 1.15 (0.98 to 1.35)	Moderate
<b>Macrovascular complications</b>				
1 systematic review (Hemmingsen 2013) including 8 RCTs (Bagg 2001, Becker 2003, DIGAMI 2 2005, Fantin 2011, Kumamoto 2000, UKPDS 1998, VA CSDM 1995, Zhang 2011)	394/3543	235/1791	RR 0.98 (0.74 to 1.30)	Low
<b>Non-fatal myocardial infarction</b>				
1 systematic review (Hemmingsen 2013) including 9 RCTs (Bagg 2001, DIGAMI 2 2005, Fantin 2011, Kumamoto 2000, Melidonis 2000, Stefanidis 2003, UGDP 1975, UKPDS 1998, VA CSDM 1995)	342/3995	187/1907	RR 0.92 (0.78 to 1.09)	High
<b>Congestive heart failure</b>				

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Number of studies	Number of people		Measure of effect	Quality
	Intensive	Conventional		
1 systematic review (Hemmingsen 2013) including 8 RCTs (Bagg 2001, DIGAMI 2 2005, Fantin 2011, Melidonis 2000, REMBO 2008, Stefanidis 2003, UKPDS 1998, VA CSDM 1995)	120/3777	75/1683	RR 0.82 (0.62 to 1.08)	Moderate
<b>Non-fatal stroke</b>				
1 systematic review (Hemmingsen 2013) including 8 RCTs (Bagg 2001, DIGAMI 2 2005, Fantin 2011, Kumamoto 2000, Melidonis 2000, Stefanidis 2003, UKPDS 1998, VA CSDM 1995)	156/3791	65/1697	RR 1.06 (0.80 to 1.41)	Moderate
<b>Amputation of lower extremity</b>				
1 systematic review (Hemmingsen 2013) including 7 RCTs (Fantin 2011, Kumamoto 2000, Melidonis 2000, Stefanidis 2003, UGDP 1975, UKPDS 1998, VA CSDM 1995)	36/3500	20/1579	RR 0.73 (0.42 to 1.25)	Moderate
<b>Microvascular complications</b>				
1 systematic review (Hemmingsen 2013) including 3 RCTs (Fantin 2011, UKPDS 1998, Zhang 2011)	253/3154	130/1222	RR 0.75 (0.61 to 0.92)	Moderate
<b>Nephropathy</b>				
1 systematic review (Hemmingsen 2013) including 7 RCTs (Bagg 2001, Fantin 2011, Kumamoto 2000, UGDP 1975, UKPDS 1998, VA CSDM 1995, Zhang 2011)	45/3167	66/1587	RR 0.64 (0.32 to 1.29)	Low
<b>End-stage renal disease</b>				
1 systematic review (Hemmingsen 2013) including 4 RCTs (Fantin 2011, Kumamoto 2000, UGDP 1975, UKPDS 1998)	28/3365	11/1438	RR 0.94 (0.47 to 1.89)	Low
<b>Retinopathy</b>				

Number of studies	Number of people		Measure of effect	Quality
	Intensive	Conventional		
1 systematic review (Hemmingsen 2013) including 5 RCTs (Fantin 2011, Kumamoto 2000, UGDP 1975, UKPDS 1998, VA CSDM 1995)	441/3098	273/1516	RR 0.79 (0.56 to 1.11)	Low
<b>Severe hypoglycaemia</b>				
1 systematic review (Hemmingsen 2013) including 13 RCTs (Bagg 2001, Blonde 2009, Cao 2011, Fantin 2011, IDA 2009, Jaber 1996, Kumamoto 2000, Melidonis 2000, Natarajan 2012, Stefanidis 2003, UKPDS 1998, VA CSDM 1995, Zhang 2011)	53/3688	11/1764	RR 2.23 (1.22 to 4.08)	Moderate
<b>Mild hypoglycaemia</b>				
1 systematic review (Hemmingsen 2013) including 12 RCTs (Bagg 2001, Blonde 2009, DIGAMI 2 2005, Fantin 2011, Kumamoto 2000, Melidonis 2000, Natarajan 2012, Stefanidis 2003, UGDP 1975, UKPDS 1998, VA CSDM 1995, Zhang 2011)	791/4200	263/2120	RR 1.85 (1.53 to 2.25)	Moderate
<b>Changes in body weight</b>				
No studies identified for this outcome				
<i>Abbreviations: RR relative risk</i>				

Update 2015

1  
2

### 8.2.2.31 Health economic evidence

2 Literature searches were undertaken to find any existing cost–utility analyses (CUAs) of  
3 optimal target values for blood glucose measures. Due to the similarity of the literature base,  
4 health economic evidence for review question 3 (target values) and review question 4  
5 (intensive versus conventional regimens) are presented together. The GDG discussed and  
6 noted that, given the nature of the CUAs presented, it was difficult to categorise them to  
7 either review question.

8 In total, 1680 references were returned for review question 3 and 421 references were  
9 returned for review question 4. Five CUAs were returned that met the NICE reference case  
10 (CDC Diabetes Cost-effectiveness Group 2002; Clarke et al. 2005; Eastman et al. 1997;  
11 Palmer et al. 2004; Valentine et al. 2006). Details of the 5 included studies are given in Table  
12 30.

13 One CUA (Clarke et al. 2005) was a lifetime modelled analysis based on a UK RCT for newly  
14 diagnosed people with type 2 diabetes. The intensive arm received insulin and sulfonylureas;  
15 the conventional arm received dietary advice and pharmacological treatments if necessary. A  
16 further CUA (CDC Diabetes Cost-effectiveness Group 2002) used the same RCT but a  
17 different model to produce a lifetime analysis based on American costs and utilities.

18 One CUA (Eastman et al. 1997) gave a lifetime modelled analysis for newly diagnosed  
19 people with type 2 diabetes, using clinical data that was not specific to people with type 2  
20 diabetes and that pre-dated the UKPDS RCT. The intensive arm received maximum doses of  
21 pharmacological agents (including insulin if necessary); the conventional arm received  
22 average doses of pharmacological agents (including insulin if necessary). This CUA only  
23 modelled microvascular and not macrovascular complications.

24 Two CUAs (Palmer et al. 2004; Valentine et al. 2006) used the CORE diabetes model to give  
25 lifetime modelled analyses for people with existing type 2 diabetes of unspecified  
26 interventions to achieve pre-specified reductions in HbA1c. These 2 CUAs did not include  
27 intervention costs or the costs of day-to-day diabetes management. Palmer et al. (2004) did  
28 not apply a discount rate to QALYs gained and Valentine et al. (2006) did not model adverse  
29 events.

30 Three of the CUAs (Eastman et al. 1997; Palmer et al. 2004; Valentine et al. 2006) assumed  
31 that treatment effects could be maintained for life. The GDG thought this assumption was  
32 unrealistic.

33 The CUAs used a range of baseline HbA1c values (between 48 and 86 mmol/mol (6.5% and  
34 10%)) and a range of HbA1c improvements (between 10 and 31 mmol/mol (0.9% and  
35 2.8%)). The highest quality and most applicable evidence was for people with newly  
36 diagnosed rather than with existing type 2 diabetes.

37 The CUA based on the UKPDS RCT was the most applicable and had the fewest limitations  
38 (Clarke et al. 2005). Other CUAs were limited by their use of non-UK (CDC Diabetes Cost-  
39 effectiveness Group 2002) and non-type-2-specific data (Eastman et al. 1997). CUAs that  
40 assumed a lifetime treatment effect and/or did not include the costs of the intervention were  
41 viewed to have very serious limitations (Eastman et al. 1997; Palmer et al. 2004; Valentine et  
42 al. 2006).

43 Most CUAs (CDC et al. 2002 (ICER \$41,400 per QALY); Clarke et al. 2005 (ICER £6000 per  
44 QALY); Eastman et al. 1997 (ICER \$16,000 per QALY)) found interventions that intensively  
45 reduce HbA1c to a given target to provide good value for money (at cost-effectiveness  
46 thresholds common in the relevant jurisdiction). The CUAs that did not include the cost of  
47 such treatment (Palmer et al. 2004; Valentine et al. 2006) found the intervention to be

1 dominant. These 2 CUAs were also, at least in part, industry funded. Notably, no CUAs  
2 modelled the impact of differential rates of hypoglycaemia between treatment arms.

3 No health economic evidence was found to comment on the cost-effectiveness of different  
4 treatment regimens or target values. The most applicable and least limited CUAs (CDC et al.  
5 2002; Clarke et al. 2005) were both based on the intervention used in the UKPDS RCT.

6 In sensitivity analysis, all CUAs noted the need for a long period of treatment (or young  
7 enough age at diagnosis) to enable costs of treatment to be recouped via complications  
8 avoided and utility to be accumulated.

9 It was noted that, whilst the clinical evidence was equivocal, the health economic evidence  
10 suggested intensive regimens to achieve lower HbA1c targets were cost effective. The GDG  
11 were presented with evidence on the uncertainty of the CUA results, as represented by  
12 probabilistic sensitivity analysis. Clarke et al. (2005) provided PSA details which showed the  
13 intervention to have a 74% probability of cost effectiveness, assuming QALYs are valued at  
14 £20,000 each.

1 Table 30: Economic evidence table for intensive control of blood glucose in people with type 2 diabetes

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p><b>CDC Diabetes Cost Effectiveness Group (2002)</b></p> <p>People with newly diagnosed type 2 diabetes aged 25+ Intensive HbA1c, hypertension and cholesterol control USA</p> <p><b>Partially applicable</b><sup>a,f,i</sup></p> <p><b>Potentially serious limitations</b><sup>j,m,n</sup></p>	<p><u>Effects:</u> UKPDS and other literature.</p> <p><u>Costs:</u> various literature sources (\$USA, 1997)</p> <p><u>Utilities:</u> various USA sources (including type 1 RCTs)</p>	<p>CDC-RTI Markov model with lifetime horizon</p> <p>Intensive HbA1c UKPDS (insulin and/or SU)</p> <p>Baseline HbA1c: 6.8%</p> <p>Conventional HbA1c based on UKPDS at 7.9%</p> <p>Hypertension and cholesterol baselines taken from NHANES</p> <p>Discounted at 3%</p> <p>Also models intensive control of hypertension and cholesterol</p>	\$7927	0.1915 QALYs	\$41,384/QALY	<p>Intensive blood glucose control seems cost effective compared with other interventions funded in the health care system.</p> <p>ICERs increase with age at diagnosis</p>	<p>ICERs for blood glucose and cholesterol increased with age at diagnosis whereas hypertension ICERs did not</p> <p>Blood glucose ICERs only less than \$50,000/QALY for those aged under 55 at diagnosis and between 45-84 for cholesterol</p> <p>Reducing incremental HbA1c costs reduce ICERs</p> <p>No PSA reported, limited OSA</p>
<p><b>Clarke et al. (2005)</b></p> <p>People with newly diagnosed type 2 diabetes aged 25-65</p> <p>Intensive control of blood glucose UK</p> <p><b>Directly applicable</b></p> <p><b>Minor limitations</b><sup>p</sup></p>	<p><u>Effects:</u> UKPDS RCT based</p> <p><u>Costs:</u> UKPDS RCT based (£UK, 2004)</p> <p><u>Utilities:</u> UKPDS RCT based</p>	<p>UKPDS model with lifetime horizon</p> <p>Intensive aimed for FPG &lt; 6mmol/l (with insulin and/or SU), conventional FPG &lt; 15mmol/l</p> <p>Intervention effect only lasts for RCT duration (11 years) – then all patients set to mean HbA1c</p> <p>UKPDS found no utility difference by regime</p> <p>Discounted at 3.5%</p>	£884	0.15 QALYs	£6028/QALY	<p>Although point estimates of cost effectiveness fall within the acceptable range, cannot be confident that the interventions are cost effective</p>	<p>ICER sensitive to primary care costs and benefit duration, but remained cost-effective under wide range of assumptions</p> <p>In PSA, 10% chance of being cost-saving 74% change of being cost-effective at £20,000/QALY</p> <p>Changes to standard care may mean benefits reported may no longer be achievable</p>

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p><b>Eastman et al. (1997)</b> People with newly diagnosed non-insulin type 2 diabetes Intensive versus conventional blood glucose control. USA</p> <p><b>Partially applicable</b><sup>a,c,i</sup></p> <p><b>Potentially serious limitations</b><sup>d,j,m,n</sup></p>	<p><u>Effects:</u> USA WESDR study Some extrapolation from type 1 data <u>Costs:</u> Medicare rates (\$USA, 1994) <u>Utilities:</u> literature (some type 1)</p>	<p>Eastman model with lifetime horizon (unspecified) Intensive HbA1c: 7.2% Conventional HbA1c aim: 10.0% Intervention HbA1c assumed to last for patient lifetime Only models impact on microvascular complications Discounted at 3%</p>	\$13,922	0.87 QALYs	\$16,002/QALY	Intensive HbA1c control appears cost effective compared to conventional control	ICERs sensitive to age at diagnosis and only remain cost effective up to around age 60 at diagnosis ICER sensitive to baseline HbA1c – cost effective to around 9% No PSA reported
<p><b>Palmer et al. (2004)</b> People with existing type 2 diabetes, treatment unspecified, aged 52 Compares 10 % improvement in HbA1c to no improvement USA</p> <p><b>Partially applicable</b><sup>a,b,i</sup></p> <p><b>Very serious limitations</b><sup>g,h,k,l,m,n,o</sup></p>	<p><u>Effects:</u> 10% HbA1c decrease HbA1c assumed, no details of how achieved <u>Costs:</u> only complication costs included, daily management and intervention costs excluded (\$USA, 2003) <u>Utilities:</u> no details given, assumed CDM standard</p>	<p>CDM with lifetime horizon (unspecified) Baseline HbA1c: 9.1% Assumes 10% improvement lasts for patient lifetime. Intensive HbA1c control not specified or costed Costs discounted at 3%; QALYs not discounted Also models intensive control of hypertension and cholesterol (10% improvement) individually and all 4 combined Funded by industry</p>	HbA1c: -\$10,800	HbA1c 0.81 QALYs	Improved HbA1c dominates no change	Improved blood glucose increase QALYs and reduce costs, meaning improved blood glucose dominates no change Cost savings driven by decreased end stage renal disease	Due to lack of intervention and day to day management costs, results may underestimate lifetime treatment costs Very limited OSA, no PSA Results insensitive to deterministic changes in discount rates or costs.

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<b>Valentine et al. (2006)</b> People with poorly controlled type 2 diabetes USA <b>Partially applicable</b> <sup>a,i</sup> <b>Very serious limitations</b> <sup>h,k,m,n,o</sup>	<u>Effects:</u> assumed, to represent various targets, no details given of how achieved	CDM with lifetime horizon 3 stepwise HbA1c reductions (all versus no reduction from that base) 9.5% to 8.0%, 8.0% to 7.0%, 7.0% to 6.5% Baseline characteristics from NHANES and RCT No adverse events modelled (hypos, weight gains) Discounted at 3% Funded by industry	9.5%-8.0%	9.5%-8.0%	9.5%- 8.0%	Improving HbA1c dominates no change in HbA1c in all cases	ICERs sensitive to time horizon – effect benefits only apparent after 2 years and cost savings after 10 years  No other details given
	<u>Costs:</u> various sources for complications.		8.0%-7.0%	8.0%-7.0%	8.0%-7.0%		
	<u>Utilities:</u> no details given, assumed CORE model standard		7.0%-6.5%	7.0%-6.5%	7.0%-6.5%		

- a Not UK based
- b Very limited details given of baseline population
- c Clinical data predates UKPDS study
- d Effectiveness data not type 2 diabetes specific
- e Assumed effectiveness remains for patient lifetime
- f Multifactorial intervention, impact not limited to impact on HbA1c
- g Day to day diabetes care costs not included
- h Costs of intervention not included
- i Costs and outcomes not discounted at 3.5%
- j Utilities not type 2 diabetes specific
- k Utility values not detailed
- l QALYs not discounted
- m Limited or no deterministic sensitivity analyses
- n Limited or no probabilistic sensitivity analyses
- o Potential conflict of interest
- p Changes to current day standard care may mean benefits reported may no longer be achievable

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p>CDC: Centre for Disease Control            CDC-RTI: Centre for Disease Control Research Triangle Institute            CDM: Centre for Outcomes Research Diabetes Model            FPG: fasting plasma glucose            HbA1c: glycosylated haemoglobin            ICER: incremental cost effectiveness ratio            mmol/l: millimoles/litre            NHANES: National Health and Nutrition Examination Survey            OSA: one-way sensitivity analysis            PSA: probabilistic sensitivity analysis            QALY: quality adjusted life year            RCT: randomised controlled trial            SU: sulfonylureas            UK: United Kingdom            UKPDS: United Kingdom Prospective Diabetes Study            USA: United States of America            WESDR: Wisconsin Epidemiologic Study of Diabetic Retinopathy</p>							

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#### 8.2.2.42 Evidence statements

##### 8.2.2.4.13 Clinical evidence

#### 4 Mortality

5 Evidence from 16 RCTs showed a trend of decreased risk of all-cause mortality with  
6 intensive target levels compared with conventional target levels. The quality of the evidence  
7 was high. However, 14 RCTs of moderate quality showed a trend of increased risk of  
8 cardiovascular mortality with intensive compared to conventional target levels. There was  
9 uncertainty surrounding these findings as all of the associated 95% confidence intervals  
10 crossed the line of minimal important difference and/or no effect.

#### 11 Macrovascular complications

12 Evidence from 9 RCTs showed trends of decreased risk of macrovascular complications  
13 (composite macrovascular end point, myocardial infarction, congestive heart failure, lower  
14 extremity amputation) with intensive compared to conventional target levels. However, 8  
15 RCTs of moderate quality showed trends of increased risk of stroke with intensive compared  
16 to conventional target levels. There was uncertainty surrounding these findings as all of the  
17 associated 95% confidence intervals crossed the line of minimal important difference and/or  
18 no effect. The quality of the evidence ranged from high to low.

#### 19 Microvascular complications

20 Evidence from 7 RCTs showed trends of decreased risk of microvascular complications  
21 (nephropathy, end-stage renal disease, retinopathy) with intensive compared to conventional  
22 target levels. The quality of the evidence was low. For the composite outcome of  
23 microvascular complications, 3 RCTs of moderate quality found a significant decrease in risk  
24 with intensive compared to conventional target levels. Generally, there was uncertainty  
25 surrounding these findings as all of the associated 95% confidence intervals crossed the line  
26 of minimal important difference and/or no effect.

#### 27 Hypoglycaemia

28 Evidence from 15 RCTs showed significant increased risk of hypoglycaemic events (mild and  
29 severe) with intensive target levels compared to conventional target levels. The quality of the  
30 evidence was moderate.

#### 31 Changes in body weight

32 No studies were identified for this outcome.

##### 8.2.2.4.23 Health economic evidence

34 One directly applicable CUA with minor limitations found that, for people newly diagnosed  
35 with type 2 diabetes, intensive control at lower HbA1c targets was cost effective compared to  
36 conventional control at higher HbA1c targets. Four partially applicable CUAs with potentially  
37 or very serious limitations found intensive control to be cost effective or dominant compared  
38 to conventional control.

### 8.2.31 Evidence to recommendations

2 **Table 31: Linking evidence to recommendations**

Relative value of different outcomes	<p>The development of macrovascular and microvascular complications, mortality and hypoglycaemic events were considered critical in decision making.</p> <p>The GDG noted that while reducing the risk of developing diabetes-related complications to improve life expectancy and quality of life, the relatively high impact of hypoglycaemic events that are associated with tight glycaemic control was also important in determining the safety and acceptability of treatment to the patient.</p> <p>The GDG agreed that all outcomes were weighted equally in deciding the optimal target values.</p>
Trade-off between benefits and harms	<p>The GDG discussed the presented evidence relating to the intensive and conventional target values. The Group noted that there was a lack of consistency in the definition of intensive and conventional targets, as this differed considerably between the included studies, which may have changed over time.</p> <p>However, the GDG agreed that there is tentative evidence to suggest that intensive target levels may be beneficial in improving risks associated with mortality, macrovascular and microvascular complications compared with conventional target levels. The GDG also recognised that intensive target levels are associated with increased risk of hypoglycaemia compared with conventional target levels. In addition, the Group acknowledged that there was a statistically non-significant trend for increased risk of cardiovascular mortality and non-fatal stroke for people receiving intensive treatment compared to conventional strategies, but agreed that the findings were uncertain.</p> <p>The GDG agreed overall that there was evidence to support the setting of target values, but considered it important to ensure that a person's risk of hypoglycaemia is evaluated when setting appropriate target levels.</p>
Consideration of health benefits and resource use	<p>The GDG found the health economic evidence on optimal target values and intensive versus conventional control hard to distinguish. No CUAs gave direct evidence on whether a particular HbA1c target was more cost effective than another HbA1c target, but all the CUAs found intensive control at lower HbA1c targets to be more cost effective than less intensive control at higher HbA1c targets.</p>
Quality of evidence	<p>The GDG agreed that overall the quality of the evidence ranged from high to low. The GDG noted that there was considerable heterogeneity in the target HbA1c levels used in the intensive control arms, as these ranged between 42 mmol/mol (6%) or lower to less than 58 mmol/mol (7.5%). There was also no restriction placed on which interventions could be used to achieve these targets. Both of these issues served to raise some doubt over the findings.</p>
Other considerations	<p>The GDG also discussed the differences in the strategies and target values used, and the potential for confusion for patients by the</p>

indeterminate nature of the intensive and conventional terminology. The GDG agreed that it would be better to provide recommendations on appropriate target values without classifying whether they are considered to be intensive or conventional.

#### **8.2.41 Recommendations and research recommendations**

- 2 See section 8.1.4 for recommendations.

## 8.3.1 Self-monitoring of blood glucose

### 8.3.1.2 Clinical introduction

3 Self-monitoring is a direct method by which a person with diabetes can be made aware of  
4 their level of blood glucose control. It is useful in people on drug treatments that require dose  
5 adjustments (such as insulin), have erratic effects or increase the risk of hypoglycaemia.  
6 There is debate surrounding the routine use of self-monitoring in people with type 2 diabetes  
7 as part of an overall educational package designed to enhance self-care and provide  
8 feedback on the impact of lifestyle measures on blood glucose control. Indirect monitoring  
9 using urine glucose tests is cheaper, but is less informative than blood glucose monitoring.

10 This section addresses the use of self-monitoring of blood glucose (SMBG) to manage  
11 glycaemic control in people with type 2 diabetes treated with diet alone or in combination with  
12 any blood glucose lowering therapies including insulin. In addition, the review looked at  
13 whether the use of self-monitoring should be restricted to specific subgroups of the  
14 population, how often and when people should self-monitor, and where on the body tests  
15 should be carried out. The review also looked at the comparative effects of different types of  
16 SMBG.

### 8.3.1.17 Self-monitoring in Clinical Guideline 66

18 Self-monitoring was originally covered as part of CG66. The original searches were  
19 conducted from 2001 to 2007 (see Appendix G for search strategies from CG66). Update  
20 searches have been carried out for this topic with a date restriction of 2007 to June 2014  
21 (see Appendix C for update search strategies). The evidence considered in this review  
22 question in CG66 included 4 systematic reviews, 2 RCTs, 4 cohort studies, 1 cross-sectional  
23 study, 1 case-series and 2 qualitative studies.

### 8.3.1.24 Self-monitoring in the update (2015)

25 For this review question, the GDG agreed that at this update, there was sufficient evidence  
26 from systematic reviews and RCTs to warrant excluding other study designs. In addition, the  
27 group expanded the scope of the review to include comparisons of different types of SMBG.

### 8.3.2.8 Evidence review

### 8.3.2.19 Review question

30 Should self-monitoring be used to manage blood glucose levels in people with type 2  
31 diabetes?

#### 32 Table 32: PICO table

Population	Adults (aged 18 years and over) with type 2 diabetes
Intervention	Self-monitoring of blood glucose using lancets
Comparators	No self-monitoring of blood glucose, standard or usual care, self-monitoring of urine glucose, other types of self-monitoring of blood glucose (such as augmentation via education, telecare, continuous glucose monitoring; or different aspects of treatment for example frequency and location of testing)
Outcomes	Changes in blood glucose levels (HbA1c, fasting and postprandial blood glucose) Hypoglycaemic events Development of microvascular and macrovascular complications: <ul style="list-style-type: none"> <li>• retinopathy (specific lesions or macular changes, referable retinopathy, blindness/loss of vision, visual acuity)</li> </ul>

- kidney damage (eGFR, serum creatinine, proteinuria, microalbuminuria, dialysis)
  - cardiovascular disease (myocardial infarction, heart failure, stroke, ACS, TIA, revascularisation and stenting)
  - foot complications (amputations, diabetic foot ulcers, Charcot osteoarthropathy, diabetic foot infection)
- Health-related quality of life

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2 RCTs that focused on the use of SMBG in people with type 2 diabetes with a minimum  
3 follow-up of 4 weeks were included. Papers were excluded if they:  
4 • were non-randomised (such as observational studies, narrative reviews and conference  
5 abstracts)  
6 • included a mixed population of people with type 1 and 2 diabetes and either did not report  
7 subgroup analyses, or less than 85% of the study population had type 2 diabetes  
8 • focused on testing of fructosamine  
9 • did not provide adequate details of standard/usual care or sufficient data for extraction on  
10 relevant outcomes. For the full excluded list, see Appendix L.

11 The main outcomes for this review question were changes in blood glucose levels,  
12 hypoglycaemia, diabetes-related complications and adverse events. The detailed protocol is  
13 available in Appendix C.

14 In the comparison on the effectiveness of SMBG versus no SMBG (including usual care and  
15 self-monitoring of urine glucose, SMUG), any type of SMBG reported in the studies was  
16 included in the meta-analysis. However, studies were excluded if they did not clearly specify  
17 that usual or standard care in the control group did not involve SMBG. Subgroup analyses on  
18 3 characteristics were undertaken:

- 19 • current diabetes treatment that is diet, oral antidiabetic and/or insulin medicines
- 20 • type of SMBG that is, standard or enhanced
- 21 • frequency of SMBG that is, less than once a day, 1 to 2 times a day or more than twice a  
22 day. Frequency was taken as the average number of tests per day and calculated based  
23 on the trial prescription described in the study methods, or if reported, the actual  
24 frequency of SMBG that was applied by the study participants.

25 For the comparison of different types of SMBG, SMBG was categorised according to the  
26 main defining feature such as enhanced education, use of phone or web-based applications  
27 to transmit blood glucose readings with or without automated and/or personalised feedback  
28 (telecare) and use of continuous glucose monitoring.

### 8.3.2.29 Clinical evidence

30 The evidence that was originally included in CG66 was re-reviewed as part of the update.  
31 Eight studies that did not meet the updated study design inclusion criteria were excluded. In  
32 addition, 1 RCT in CG66 (Moreland et al. 2006) did not meet the revised population inclusion  
33 criteria for this update as only 65% of the study participants had type 2 diabetes. One of the  
34 systematic reviews (Welschen et al. 2005) in CG66 has since been updated (Malanda et al.  
35 2012), so the more recent version was used in this question. This Cochrane review was  
36 restricted to comparisons involving SMBG and usual care (Malanda et al. 2012). However, all  
37 of the RCTs included in the Cochrane review were also identified in the update searches and  
38 where possible, the original papers were preferentially used.

39 In total, 1808 references were found for this review question and 29 unique trials were  
40 included ((Allen et al. 1990; Barnett et al. 2008; Bonomo et al. 2010; Bosi et al. 2013; Cho et  
41 al. 2009; Davidson et al. 2005; Del Prato et al. 2012; Farmer et al. 2007; Fontbonne et al.

1 1989; Franciosi et al. 2011; Guerci et al. 2003; Ismail et al. 2013; Kleefstra et al. 2010;  
2 Knapp et al. 2009; Kwon et al. 2004; Lim et al. 2011; Lu et al. 2011; Muchmore et al. 1994;  
3 Nauck et al. 2014; O’Kane et al. 2008; Pimazoni-Netto et al. 2011; Polonsky et al. 2011;  
4 Quinn et al. 2011; Scherbaum et al. 2008; Schwedes et al. 2002; Tildesley et al. 2010;  
5 Vigersky et al. 2012; Wing et al. 1986; Yoo et al. 2008). There were 2 cluster RCTs  
6 (Polonsky et al. 2011; Quinn et al. 2011), 1 of which was a 4-armed study with 2 groups  
7 relevant to this review (Quinn et al. 2011); 1 RCT applied a 2x2 factorial design (Nauck et al.  
8 2014); while 4 RCTs involved 3 treatment arms (Farmer et al. 2007; Fontbonne et al. 1989;  
9 Lim et al. 2011; Lu et al. 2011). The following comparisons were included as part of this  
10 review question:

- 11 • SMBG versus no SMBG (including standard/usual care and SMUG) – 17 trials (Allen et al.  
12 1990; Barnett et al. 2008; Bosi et al. 2013; Davidson et al. 2005; Farmer et al. 2007;  
13 Fontbonne et al. 1989; Franciosi et al. 2011; Guerci et al. 2003; Ismail et al. 2013;  
14 Kleefstra et al. 2010; Lim et al. 2011; Lu et al. 2011; Muchmore et al. 1994; Nauck et al.  
15 2014; O’Kane et al. 2008; Schwedes et al. 2002; Wing et al. 1986)
- 16 • SMBG plus education versus conventional SMBG – 3 trials (Farmer et al. 2007; Pimazoni-  
17 Netto et al. 2011; Polonsky et al. 2011)
- 18 • SMBG plus telecare via phone or internet with tailored or automated feedback versus  
19 conventional SMBG – 5 trials (Del Prato et al. 2012; Kwon et al. 2004; Lim et al. 2011;  
20 Quinn et al. 2011; Tildesley et al. 2010)
- 21 • Different mechanisms of exporting glucose readings that is using an automated mobile  
22 phone connected glucometer versus standard glucometer requiring web log in to enter  
23 data – 1 trial (Cho et al. 2009)
- 24 • SMBG plus continuous glucose monitoring (CGM) versus conventional SMBG – 2 trials  
25 (Vigersky et al. 2012; Yoo et al. 2008)
- 26 • Frequency of SMBG testing – 2 trials (Bonomo et al. 2010, Scherbaum et al. 2008)
- 27 • Location of SMBG testing – 1 trial (Knapp et al. 2009)

### 8.3.2.2.18 *Description of included studies*

29 Details of the included studies are found in the evidence tables (see Appendix E).

#### 30 **SMBG versus no SMBG**

31 A total of 4710 people (study size ranged from 23 to 1024) were included from 17 RCTs,  
32 carried out in the UK (Farmer et al. 2007; O’Kane et al. 2008), the Netherlands (Kleefstra et  
33 al. 2010), France (Fontbonne et al. 1989; Guerci et al. 2003), Germany (Nauck et al. 2014),  
34 Italy (Bosi et al. 2013; Franciosi et al. 2011), USA (Allen et al. 1990; Davidson et al. 2005;  
35 Muchmore et al. 1994; Wing et al. 1986), Korea (Lim et al. 2011; Lu et al. 2011) and  
36 Malaysia (Ismail et al. 2013); the remaining 2 were multinational studies (Barnett et al. 2008;  
37 Schwedes et al. 2002). The mean age ranged from 48.9 to 67.5 years. The mean duration of  
38 diabetes in 15 studies ranged from 2.7 to 15.4 years; 2 trials did not report this information  
39 (O’Kane et al. 2008; Wing et al. 1986). Mean HbA1c at baseline ranged from 56 to 108  
40 mmol/mol (7.3% to 12%). Mean BMI ranged from 25 to 34.2 kg/m<sup>2</sup>, with 7 studies not  
41 reporting this information (Allen et al. 1990; Barnett et al. 2008; Davidson et al. 2005;  
42 Franciosi et al. 2011; Guerci et al. 2003; Schwedes et al. 2002; Wing et al. 1986). People  
43 taking insulin were included in 5 studies (Barnett et al. 2008; Ismail et al. 2013; Lim et al.  
44 2011; Nauck et al. 2014; Wing et al. 1986), 1 study included people managed on diet alone  
45 (O’Kane et al. 2008), while the participants in the remaining trials were managed on diet  
46 and/or oral antidiabetic medicines. Follow-up periods ranged from 24 to 208 weeks.

#### 47 **SMBG plus education versus conventional SMBG**

48 A total of 1015 people (study size ranged from 63 to 499) were included from 3 RCTs,  
49 carried out in the UK (Farmer et al. 2007), USA (Polonsky et al. 2011) and Brazil (Pimazoni-

1 Netto et al. 2011). The mean age ranged from 56 to 65.6 years. The mean duration of  
2 diabetes ranged from 3 to 12 years. Mean HbA1c at baseline ranged from 58 to 86 mmol/mol  
3 (7.5% to 10%). Mean BMI ranged from 31.3 to 35.1 kg/m<sup>2</sup>, with 1 study not reporting this  
4 information (Pimazoni-Netto et al. 2011). One study included people taking insulin (Pimazoni-  
5 Netto et al. 2011), while the other 2 studies included participants managed on non-insulin  
6 based therapies. Follow-up periods ranged from 12 to 208 weeks.

#### 7 **SMBG plus telecare versus conventional SMBG**

8 A total of 768 people (study size ranged from 50 to 291) were included from 5 RCTs, carried  
9 out in the USA (Quinn et al. 2011), Canada (Tildesley et al. 2010), Italy (Del Prato et al.  
10 2012) and Korea (Kwon et al. 2004; Lim et al. 2011). The mean age ranged from 53 to 67.5  
11 years. The mean duration of diabetes ranged from 6.8 to 18.8 years. Mean HbA1c at  
12 baseline ranged from 57 to 79 mmol/mol (7.4% to 9.4%). Mean BMI in 4 studies ranged from  
13 24 to 35.6 kg/m<sup>2</sup>, with 1 study not reporting this information (Del Prato et al. 2012). Three  
14 studies included people taking insulin (Del Prato et al. 2012; Lim et al. 2011; Tildesley et al.  
15 2010), while the other 2 studies did not specify whether participants were on existing  
16 therapies (Kwon et al. 2004; Quinn et al. 2011). Follow-up periods ranged from 12 to 52  
17 weeks.

#### 18 **Automated mobile phone glucometer versus standard glucometer**

19 One 12 week trial conducted in Korea including 75 people (mean age 48 years; mean  
20 duration of diabetes 6.8 years; mean HbA1c at baseline 64 mmol/mol (8%); mean BMI 24.5  
21 kg/m<sup>2</sup>) with unspecified existing therapies was analysed in this comparison (Cho et al. 2009).

#### 22 **SMBG plus continuous glucose monitoring (CGM) versus conventional SMBG**

23 A total of 165 people (study sizes 65 and 100) were included from 2 RCTs, carried out in the  
24 USA (Vigersky et al. 2012) and Korea (Yoo et al. 2008). The mean ages were 56 and 58  
25 years. The mean duration of diabetes was reported in 1 study as 13 years (Yoo et al. 2008).  
26 Mean HbA1c levels at baseline were 67 mmol/mol (8.3%) and 74 mmol/mol (8.9%). Mean  
27 BMI was not reported in either studies, while both studies included people taking insulin.  
28 Follow-up periods were 12 and 52 weeks.

#### 29 **Frequency of SMBG testing**

30 A total of 475 people (study sizes 202 and 273) were included from 2 RCTs, carried out in  
31 Italy (Bonomo et al. 2010) and Germany (Scherbaum et al. 2008). The mean ages were 61  
32 and 64 years. The mean duration of diabetes was 8 and 10.6 years. Mean HbA1c levels at  
33 baseline were 55 mmol/mol (7.2%) and 64 mmol/mol (8%). Mean BMI was reported in 1  
34 study as 29 kg/m<sup>2</sup> (Bonomo et al. 2010). Both studies included people managed on diet  
35 and/or oral antidiabetic medicines. Follow-up periods were 26 and 52 weeks.

#### 36 **Location of SMBG testing**

37 One 30 week trial conducted in the USA including 174 people (mean age 53 years; mean  
38 duration of diabetes 12 years; mean HbA1c at baseline 73 mmol/mol (8.8%); mean BMI 36  
39 kg/m<sup>2</sup>), some of whom were managed on insulin was analysed in the comparison of SMBG  
40 administered on the fingertip or on the forearm (Knapp et al. 2009).

41 The summary GRADE tables are presented for this review question (see Appendix D for full  
42 GRADE tables).

1 Table 33: Summary GRADE profile for SMBG versus no SMBG

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG	no SMBG		
<b>HbA1c (%) at 24 to 52 week follow-up</b>				
17 (Allen 1990; Barnett 2008; Bosi 2013; Davidson 2005; Farmer 2007; Fontbonne 1989; Franciosi 2011; Guerci 2003; Ismail 2013; Kleefstra 2010; Lim 2011; Lu 2011; Muchmore 1994; Nauck 2014; O’Kane 2008; Schwedes 2002; Wing 1986)	2217	2084	MD -0.22 (-0.31 to -0.13)  <u>Subgroup analysis based on current medication:</u> Diet alone: MD -0.2 (-0.8 to 0.4) Diet ± OADs: MD -0.21 (-0.29 to -0.13) Diet, OADs ± insulin: MD -0.38 (-0.86 to 0.10), I <sup>2</sup> =84%  <u>Subgroup analysis based on type of SMBG:</u> Standard SMBG: MD -0.21 (-0.31 to -0.11) Enhanced SMBG: MD -0.29 (-0.49 to -0.09)  <u>Subgroup analysis based on frequency of SMBG:</u> <1 per day: MD -0.31 (-0.55 to -0.07), I <sup>2</sup> =68% 1-2 times per day: MD -0.19 (-0.29 to -0.10) >2 per day: MD -0.20 (-0.73 to 0.32)	Low
<b>Change in HbA1c (%) by prespecified subgroups at 1 year follow-up</b>				
1 (Farmer 2007) – DiGEM	151†	152	Diet alone: MD -0.12 (-0.29 to 0.05) Oral antidiabetic therapy: MD -0.19 (-0.40 to 0.02) Diabetes duration <36 months: MD -0.17 (-0.37 to 0.03) >36 months: MD -0.17 (-0.37 to 0.03) No diabetic complications: MD -0.23 (-0.43 to -0.03) With complications: MD -0.36 (-0.55 to -0.17)	Moderate
<b>Fasting blood glucose (mmol/L) at 24 to 52 week follow-up</b>				

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG	no SMBG		
6 (Allen 1990; Barnett 2008; Guerci 2003; Lim 2011; Lu 2011; Wing 1986)	835	810	MD -0.38 (-0.68 to -0.07)  <u>Subgroup analysis based on current medication:</u> Diet ± OADs: MD -0.26 (-0.59 to 0.07) Diet, OADs ± insulin: MD -1.33 (-2.27 to -0.38)  <u>Subgroup analysis based on type of SMBG:</u> Standard SMBG: MD -0.31 (-0.63 to 0.00) Enhanced SMBG: MD -1.57 (-2.94 to -0.20)  <u>Subgroup analysis based on frequency of SMBG:</u> <1 per day: MD -0.20 (-0.86 to 0.47) 1-2 times per day: MD -0.55 (-1.30 to 0.20), I <sup>2</sup> =54% >2 per day: MD -0.51 (-2.01 to 0.99)	Low
<b>Postprandial blood glucose (mg/dL) at 6 months in older adults with type 2 diabetes treated with diet, oral antidiabetic and/or insulin medicines</b>				
1 (Lim 2011)	96	48	MD -71.78 (-96.62 to -46.94)  <u>Subgroup analysis based on type of SMBG:</u> Standard SMBG: MD -61.30 (-97.61 to -24.99) Enhanced SMBG: MD -81.00 (-111.05 to -46.95)	Low
<b>Any hypoglycaemia* at 6 to 12 month follow-up (measured as the number of patients experiencing 1 or more events)</b>				

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG	no SMBG		
6 (Barnett 2008; Farmer 2007; Guerci 2003; Lim 2011; Lu 2011; O'Kane 2008;)	203/1354	88/1138	RR 1.62 (1.19 to 2.22), I <sup>2</sup> =34%  <u>Subgroup analysis based on current medication:</u> Diet alone: RR 1.27 (0.66 to 2.44) Diet ± OADs: RR 1.80 (1.16 to 2.79), I <sup>2</sup> =47% Diet, OADs ± insulin: RR 1.30 (0.70 to 2.39)  <u>Subgroup analysis based on frequency of SMBG:</u> <1 per day: RR 2.28 (1.61 to 3.23) 1-2 times per day: RR 1.26 (0.89 to 1.79) >2 per day: RR 0.51 (0.06 to 4.37)	Low
<b>Severe hypoglycaemia at 6 to 12 month follow-up (measured as the number of patients experiencing 1 or more events)</b>				
3 (Bosi 2013; Farmer 2007; Lim 2011)	1/853	4/727	RR 0.35 (0.07 to 1.77)  <u>Subgroup analysis based on current medication:</u> Diet ± OADs: RR 0.17 (0.01 to 4.12) Diet, OADs ± insulin: RR 0.45 (0.07 to 2.99)  <u>Subgroup analysis based on frequency of SMBG:</u> <1 per day: RR 0.17 (0.01 to 4.12) 1-2 times per day: RR 0.45 (0.07 to 2.99)	Low
<b>Any adverse events<sup>a</sup> at 6 month follow-up in people treated with diet and/or oral antidiabetic medicines (majority of events were of mild or moderate severity)</b>				
1 (Barnett 2008)	41/311	45/299	RR 0.88 (0.59 to 1.30)	Moderate
<p>Abbreviations: CI confidence interval; MD mean difference; OADs oral antidiabetic drugs; RR relative risk</p> <p><sup>a</sup>Definitions of hypoglycaemia differed across the included studies. Overall Barnett (2008) and Lu (2011) used grades of hypoglycaemia with grades 1 and 2 relating to mild or moderate episodes and grades 3 and 4 referring to more severe episodes that require medical assistance. Barnett (2008) did not refer to specific blood glucose values but reported that 11/51 events in 27 patients in the SMBG group were SMBG confirmed hypoglycaemia. In both studies no patients experienced more than grade 3. Guerci (2003) referred to symptomatic or asymptomatic hypoglycaemia with no further definition but it was noted that no serious episode of hypoglycaemia was reported. Lim (2011) defined minor symptomatic hypoglycaemia as symptoms with blood glucose levels &lt;3.5 mmol/L, major symptomatic hypoglycaemia as blood glucose levels &lt;2.8 mmol/L and an episode requiring medical intervention or markedly depressed levels of consciousness or seizure and nocturnal hypoglycaemia as events occurring while asleep</p> <p><sup>†</sup> intervention group relates to more intensive SMBG (this has not been combined with less intensive monitoring)</p>				

1 **Table 34: Summary GRADE profile for SMBG plus education versus conventional SMBG**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG plus education	Conventional SMBG		
<b>HbA1c (%) at 3 to 12 month follow-up</b>				
3 (Farmer 2007; Pimazoni-Netto 2011; Polonsky 2011)	439	408	MD -0.31 (-0.67 to 0.05), I <sup>2</sup> =79%  Subgroup analysis based on current medication: Diet ± OADs: MD -0.15 (-0.42 to 0.11), I <sup>2</sup> =69% Diet, OADs ± insulin: RR -0.97 (-1.62 to -0.32)	Low
<b>Any hypoglycaemia at 12 month follow-up in people treated with diet and/or oral antidiabetic medicines</b>				
2 (Farmer 2007; Polonsky 2011)	48/407	37/377	RR 1.28 (0.88 to 1.86)	Low
1 (Pimazoni-Netto 2011)	32	31	The frequency of events was not significantly higher in intervention (4.11± 0.96%) vs. control (2.24 ± 0.64%, p>0.05)	Low
<i>Abbreviations: CI confidence interval; MD mean difference; OADs oral antidiabetic drugs; RR relative risk</i>				

2 **Table 35: Summary GRADE profile for SMBG plus telecare (phone or internet with feedback) versus conventional SMBG**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG plus telecare	Conventional SMBG		
<b>HbA1c (%) at 12 to 52 week follow-up</b>				
5 (Del Prato 2012; Kwon 2004; Lim 2011; Quinn 2011; Tildesley 2010)	260	295	MD -0.57 (-1.06 to -0.08), I <sup>2</sup> =85%  Subgroup analysis based on current medication: Insulin: MD -0.27 (-0.68 to 0.13), I <sup>2</sup> =71% Not specified: MD -1.04 (-1.42 to -0.65)	Low
<b>Fasting blood glucose (mmol/L) at 26 and 44 week follow-up in people treated with oral hypoglycaemic drugs and/or insulin</b>				
2 (Del Prato 2012; Lim 2011)	164	171	MD -0.19 (-0.61 to 0.24), I <sup>2</sup> =40%	Low
<b>Postprandial blood glucose (mg/dL) at 26 week follow-up in older adults treated with oral hypoglycaemic drugs and/or insulin</b>				
1 (Lim 2011)	49	47	MD -19.07 (-42.84 to 3.44)	Low
<b>Any hypoglycaemia at 26 week follow-up in people treated with oral hypoglycaemic drugs and/or insulin</b>				

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG plus telecare	Conventional SMBG		
1 (Lim 2011)	16/51	12/51	RR 1.33 (0.70 to 2.53)	Low
<b>Total symptomatic hypoglycaemia at 44 week follow-up in people treated with insulin therapy</b>				
1 (Del Prato 2012) – ELEONER	1.89 events per patient year	1.76 events per patient year	Rate ratio <sup>‡</sup> 1.07 (0.89 to 1.29)	Very low
<b>Severe nocturnal hypoglycaemia at 44 week follow-up in people treated with insulin therapy</b>				
1 (Del Prato 2012) – ELEONER	0.04 events per patient year	0.02 events per patient year	Rate ratio 2.00 (0.44 to 9.06)	Very low

Abbreviations: CI confidence interval; MD mean difference; RR relative risk  
<sup>‡</sup> Estimated using likely patient years to calculate number of events as only rates reported in full paper

1 **Table 36: Summary GRADE profile for Automated mobile phone glucometer versus standard glucometer**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	Mobile phone glucometer	Standard glucometer		
<b>HbA1c (%) at 3 month follow-up in people with unspecified current therapy</b>				
1 (Cho 2011)	35	34	MD 0.29 (-0.25 to 0.83)	Low
<b>Fasting blood glucose (mmol/L) at 3 month follow-up in people with unspecified current therapy</b>				
1 (Cho 2011)	35	34	MD -0.33 (-1.64 to 0.99)	Low
<b>Postprandial blood glucose (mg/dL) at 3 month follow-up in people with unspecified current therapy</b>				
1 (Cho 2011)	35	34	MD -11.57 (-46.55 to 23.41)	Very low

Abbreviations: CI confidence interval; MD mean difference

2 **Table 37: Summary GRADE profile for SMBG plus continuous glucose monitoring (CGM) versus conventional SMBG**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG plus CGM	SMBG		
<b>HbA1c (%) up to 52 week follow-up in people treated with oral antidiabetic and/or insulin medicines</b>				

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG plus CGM	SMBG		
2 (Vigersky 2012; Yoo 2008)	79	78	MD -0.46 (-0.87 to -0.06)	Very low
<b>Fasting blood glucose (mmol/L) at 12 week follow-up in people treated with oral antidiabetic and/or insulin medicines</b>				
1 (Yoo 2008)	29	28	MD -0.70 (-1.62 to 0.22)	Low
<b>Postprandial blood glucose (mmol/L) at 12 week follow-up in people treated with oral antidiabetic and/or insulin medicines</b>				
1 (Yoo 2008)	29	28	MD -0.90 (-2.67 to 0.87)	Low

Abbreviations: CI confidence interval; MD mean difference

1 Table 38: Summary GRADE profile for frequency of SMBG testing (monthly versus fortnightly and 4 times weekly versus once weekly)

Number of RCTs	Number of people		Effect (85% CI)	Quality
	SMBG more frequently	SMBG less frequently		
	SMBG fortnightly	SMBG monthly		
<b>HbA1c (%) at 6 month follow-up in people not on insulin</b>				
1 (Bonomo 2010)	177	96	MD 0.04 (-0.20 to 0.28)  Subgroup: people compliant with SMBG MD -0.31 (-0.59 to -0.03)	Moderate
<b>Hypoglycaemia at 6 month follow-up in people not on insulin (defined as blood glucose &lt;3.3 mmol/L)</b>				
1 (Bonomo 2010)	177	96	RR 0.30 (0.03 to 2.86)	Low
	SMBG 4 times weekly	SMBG once weekly		
<b>HbA1c (%) at study end in people not treated with insulin</b>				
1 (Scherbaum 2008)	95	93	3 months: MD 0.00 (-0.28 to 0.28) 6 months: MD 0.10 (-0.20 to 0.40) 12 months: MD 0.20 (-0.10 to 0.50)	Moderate
<b>Hypoglycaemia at 12 month follow-up in people not treated with insulin (1 event of SMBG &lt;3.2mmol/L or several events)</b>				

1 (Scherbaum 2008)	18/102	5/100	RR 3.53 (1.36 to 9.14)	Moderate
<b>Adverse events at 12 month follow-up in people not treated with insulin (hyperglycaemia, deteriorating neuropathy, retinopathy or nephropathy, multiple events or other events)</b>				
1 (Scherbaum 2008)	8/102	14/100	RR 0.56 (0.25 to 1.28)	Low
<b>Serious adverse events at 12 month follow-up in people not treated with insulin (hypoglycaemic shock, hyperosmolar coma, inpatient stay or death)</b>				
1 (Scherbaum 2008)	15/102	20/100	RR 0.74 (0.40 to 1.35)	Low

Abbreviations: CI confidence interval; MD mean difference; RR relative risk

**Table 39: Summary GRADE profile for location of SMBG testing (forearm versus fingertip)**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	SMBG at forearm	SMBG at fingertip		
<b>Change in HbA1c (%) at 6 month follow-up in people treated with insulin</b>				
1 (Knapp 2009)	89	85	MD 0.10 (-0.29 to 0.49)  Subgroup analysis based on baseline HbA1c levels: ≤7%: MD 0.00 (-0.41 to 0.41) 7.0-8.5%: MD 0.00 (-0.52 to 0.52) >8.5%: MD 0.20 (-0.45 to 0.85)	High
<b>Hypoglycaemia at 6 month follow-up in people treated with insulin (more than 1 episode per month)</b>				
1 (Knapp 2009)	3/89	3/85	RR 0.96 (0.20 to 4.60)	Moderate
<b>Severe hypoglycaemia at 6 month follow-up in people treated with insulin (requiring urgent medical attention)</b>				
1 (Knapp 2009)	3/89	1/85	RR 2.87 (0.30 to 27.01)	Moderate

Abbreviations: CI confidence interval; MD mean difference; RR relative risk

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### 8.3.2.31 Health economic evidence

2 Literature searches were undertaken to find any existing cost–utility analyses (CUAs) of self-  
3 monitoring of blood glucose in people with type 2 diabetes (see appendix C for detail of the  
4 search strategies). In total, 838 articles were found and 8 CUAs were returned (Cameron et  
5 al. 2010; Farmer et al. 2009; Palmer et al. 2006; Pollock et al. 2010; Simon et al. 2008; Tunis  
6 et al. 2010; Tunis and Minshall 2008; Tunis and Minshall 2010) that met the NICE reference  
7 case (National Institute for Health and Care Excellence 2012).

8 In addition, 1 recent UK health technology assessment report (HTA) was found (Clar et al.  
9 2010). The HTA report did not undertake de novo modelling but reviewed the existing health  
10 economic evidence. Four of the CUAs included for this guideline (Farmer et al. 2009; Palmer  
11 et al. 2006; Simon et al. 2008; Tunis and Minshall 2008b) were included in the HTA report,  
12 with 4 CUAs (Cameron et al. 2010; Pollock et al. 2010; Tunis et al. 2010; Tunis and Minshall  
13 2010) published after the HTA report.

14 With the exception of 1 CUA (Simon et al. 2008), all the CUAs were lifetime-modelled  
15 analyses using either the CDM (Centre for Outcomes Research Diabetes Model) (Palmer et  
16 al. 2004) or the UKPDS Outcomes Model (Clarke et al. 2004). One CUA (Simon et al. 2008)  
17 was an RCT-based economic evaluation that was later extended to a lifetime-modelled  
18 analysis (Farmer et al. 2009).

19 Three CUAs (Farmer et al. 2009; Palmer et al. 2006; Simon et al. 2008) were based on  
20 mainly UK data, 3 CUAs (Cameron et al. 2010; Tunis and Minshall 2008; Tunis and Minshall  
21 2010) were based on mainly American or Canadian data and 2 CUAs (Pollock et al. 2010;  
22 Tunis et al. 2010) were based on data mainly from European countries.

23 The CUAs all compared self-monitoring of blood glucose to no monitoring, but contained  
24 different self-monitoring comparisons. Two CUAs (Farmer et al. 2009; Simon et al. 2008)  
25 incrementally compared more and less intensive testing regimes to no self-monitoring. One  
26 CUA (Cameron et al. 2010) compared self-monitoring at 9 tests per week to no self-  
27 monitoring. Five CUAs (Palmer et al. 2006; Pollock et al. 2010; Tunis et al. 2010; Tunis and  
28 Minshall 2008; Tunis and Minshall 2010) compared self-monitoring 1, 2 or 3 times per day to  
29 no self-monitoring but not as incremental comparisons against the marginal value of each  
30 extra test.

31 All the CUAs included or modelled people with existing type 2 diabetes. Most of the CUAs  
32 considered patients new to self-monitoring; only 1 CUA (Tunis and Minshall 2010)  
33 considered patients who had previously used self-monitoring.

34 Cohorts of people with type 2 diabetes covered a variety of generic diabetes treatment  
35 regimens. Three CUAs (Cameron et al. 2010; Farmer et al. 2009; Simon et al. 2008)  
36 modelled patients treated with diet and OADs together, 4 CUAs (Pollock et al. 2010; Tunis et  
37 al. 2010; Tunis and Minshall 2008; Tunis and Minshall 2010) modelled only patients treated  
38 with OADs and 1 CUA (Palmer et al. 2006) presented analyses separately for patients  
39 treated with diet, OADs and insulin. In their sensitivity analyses, 1 CUA (Cameron et al.  
40 2010) also presented results separately for patients treated with diet, OADs and insulin.

41 The level of HbA1c change used within the health economic modelling ranged from a 1.5  
42 mmol/mol (0.14%) reduction (Farmer et al. 2009; Simon et al. 2008) to a 11 mmol/mol  
43 (1.02%) reduction (Pollock et al. 2010; Tunis et al. 2010). Only 3 CUAs (Cameron et al.  
44 2010; Farmer et al. 2009; Simon et al. 2008) sourced their HbA1c change from RCT or  
45 systematic reviews; 4 CUAs (Pollock et al. 2010; Tunis et al. 2010; Tunis and Minshall 2008;  
46 Tunis and Minshall 2010) sourced their HbA1c change from an American observational study  
47 (Karter et al. 2006) and 1 CUA (Palmer et al. 2006) assumed their HbA1c change. One CUA  
48 (Cameron et al. 2010) employed a level of change (3 mmol/mol (0.25%) reduction) that was  
49 closest to that found in the clinical evidence review. Five CUAs (Palmer et al. 2006; Pollock

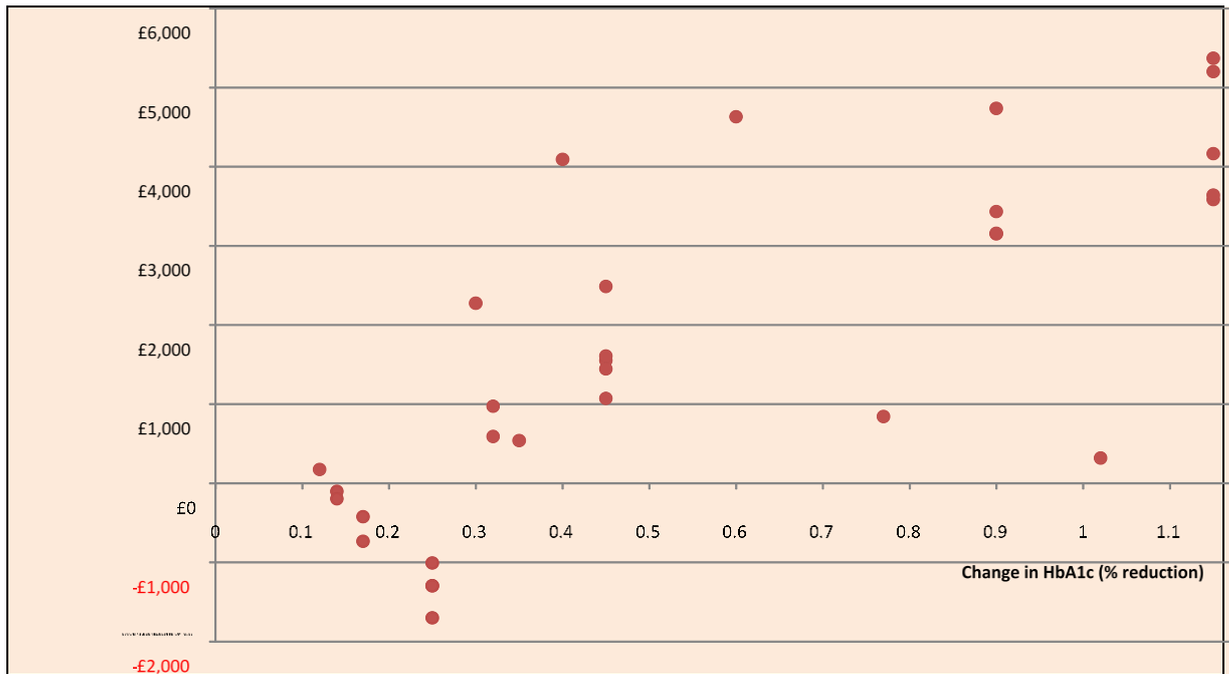
1 et al. 2010; Tunis et al. 2010; Tunis and Minshall 2008; Tunis and Minshall 2010) modelled  
2 an increase in testing frequency linked with assumed (but not evidence based) greater  
3 HbA1c reductions.

4 The included CUAs differed in their assumptions of how long their change in HbA1c levels  
5 due to self-monitoring would be maintained. The 2 CUAs based on the DiGEM RCT (Farmer  
6 et al. 2009; Simon et al. 2008), the Canadian HTA (Cameron et al. 2010) and the Swiss-  
7 based CUA (Pollock et al. 2010) assumed the change in HbA1c resulting from self-  
8 monitoring would last for 1 year. The other 4 CUAs (Palmer et al. 2006; Tunis et al. 2010;  
9 Tunis and Minshall 2008; Tunis and Minshall 2010) assumed the change in HbA1c resulting  
10 from self-monitoring would last for a patient lifetime.

11 The 2 CUAs based on the DiGEM RCT (Farmer et al. 2009; Simon et al. 2008) were most  
12 applicable and had the fewest limitations. The Canadian HTA (Cameron et al. 2010) had few  
13 limitations but was not directly applicable as it used non-UK costs; 1 CUA (Palmer et al.  
14 2006) was directly applicable but was limited by assuming levels of HbA1c change and  
15 lacking adequate details of costs and utilities used (as noted in previous NICE guidance  
16 CG66). Four CUAs (Pollock et al. 2010; Tunis et al. 2010; Tunis and Minshall 2008b; Tunis  
17 and Minshall 2010) that were not based in the UK and used an HbA1c change from  
18 American observational data were only partially applicable and had serious limitations.

19 The included CUAs gave heterogeneous cost-effectiveness results. The most applicable  
20 CUAs with fewest limitations (Farmer et al. 2009; Simon et al. 2008) found self-monitoring to  
21 be not cost effective (dominated) compared with no self-monitoring. The Canadian HTA  
22 (Cameron et al. 2010) found self-monitoring not to be cost effective compared with no self-  
23 monitoring using Canadian costs and thresholds (ICER \$114,000 per QALY); 1 UK-based  
24 CUA (Palmer et al. 2006) found self-monitoring to be cost effective compared with no self-  
25 monitoring (ICERs ranged from £4600 per QALY for people on OADs or insulin to £15,per  
26 QALY 600 for people on diet alone). Using a variety of thresholds, 4 partially applicable  
27 CUAs with serious limitations (Pollock et al. 2010 (ICERs CHF9200 per QALY for 1 test per  
28 day, CHF13,000 per QALY for 2 tests per day and CHF17,300 per QALY or for 3 tests per  
29 day); Tunis et al. 2010 (ICERs between €1600 per QALY and €15,400 per QALY depending  
30 on country and testing frequency); Tunis and Minshall 2008 (ICERs \$7900 per QALY for 1  
31 test per day and \$6600 per QALY for 3 tests per day); Tunis and Minshall 2010 (ICERs  
32 \$26,200 per QALY for 1 test per day, \$18,600 per QALY for 2 tests per day and £25,400 per  
33 QALY for 3 tests per day) found self-monitoring to be cost effective compared with no self-  
34 monitoring . No CUAs found self-monitoring to be cost saving compared with no self-  
35 monitoring, meaning that the trade-offs between cost and benefits and the opportunity costs  
36 of self-monitoring to the rest of the NHS had to be considered.

37 An association between modelled HbA1c change and incremental net monetary benefit (at  
38 £20,000/QALY threshold) seemed apparent in the included CUAs (see figure 4). Greater  
39 HbA1c reductions assumed to be achievable by self-monitoring of blood glucose led to  
40 higher HbA1c gains.



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**1 Figure 4: Self-monitoring of blood glucose - HbA1c reduction used in included CUAs**

2 The CUAs displayed high levels of uncertainty in their cost effectiveness results. The 2 CUAs  
 3 that found self-monitoring to be not cost effective compared with no self-monitoring found the  
 4 reverse would be true in fewer than 40% of PSA iterations (Cameron et al. 2010; Farmer et  
 5 al. 2009). Three of the 5 CUAs (Palmer et al. 2006; Tunis et al. 2010; Tunis and Minshall  
 6 2008) that found self-monitoring to be cost effective compared with no self-monitoring  
 7 showed this to be the case in fewer than 60% of replications; 1 of the 5 CUAs (Tunis and  
 8 Minshall 2010) did not report uncertainty and the least applicable study (Pollock et al. 2010)  
 9 reported self-monitoring to be cost effective compared to no self-monitoring in fewer than  
 10 72% of replications.

11 The 2 CUAs based on the DiGEM RCT (Farmer et al. 2009; Simon et al. 2008) included a  
 12 disutility associated with performing self-monitoring. Two other CUAs (Cameron et al. 2010;  
 13 Palmer et al. 2006) included such a disutility in their sensitivity analyses and found doing so  
 14 decreased the cost effectiveness of self-monitoring compared with no self-monitoring  
 15 (increased the incremental cost effectiveness ratios).

1 **Table 40: Economic evidence table for self monitoring of blood glucose**

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p><b>Cameron et al. (2010)</b> People with existing type 2 diabetes treated with diet and exercise or OADs SMBG v none Canada</p>	<p><u>Effects:</u> systematic review <u>Costs:</u> Canadian standard sources (\$Can, 2008) <u>Utilities:</u> US confounder-controlled EQ-5D catalogue. No disutility from SMBG</p>	<p>UKPDS model with lifetime horizon (40 years) 1.3 strips/day (9/week) HbA1c baseline: 8.4% HbA1c effect: -0.25%, lasts for 1 year Discounted at 5%  Assumed no pre-existing complications</p>	\$2711	0.024 QALYs	\$114,000/QALY	<p>SMBG not cost effective compared to no SMBG at \$100,000/QALY threshold  Clinical benefits of SMBG and associated savings do not offset the cost of the strips  Monitoring once or twice per week could be cost effective</p>	<p>ICERs sensitive to treatment: diet \$292,000/QALY; OADs: \$92,000/QALY; insulin: \$92,000/QALY ICERs sensitive to strip cost and testing frequency. 50% reduction in either makes SMBG cost effective ICERs sensitive to HbA1c change. If HbA1c reduction is doubled, ICER is around \$50,000/QALY ICERs sensitive to disutility from SMBG – if applied, ICER is around \$180,000/QALY In PSA, SMBG is cost effective in &lt;10% of replications at \$CAN50,000/QALY and 40% of replications at \$CAN100,000/QALY</p>
<p><b>Partly applicable</b><sup>a,b,c,e</sup></p>							
<p><b>Minor limitations</b><sup>a</sup></p>							
<p><b>Farmer et al. (2009)</b> People with existing type 2 diabetes on diet and exercise or OADs More or less intense SMBG v usual care England</p>	<p><u>Effects:</u> DiGEM RCT <u>Costs:</u> UKPDS model (£UK, 2005) <u>Utilities:</u> EQ-5D from UKPDS. Includes disutility from SMBG</p>	<p>UKPDS model with lifetime horizon. Lifetime model of Simon et al. (2008) 0.9 strips/day (6/week) HbA1c baseline: 7.5% HbA1c effect: no impact post RCT period Control: usual care “Less” intensive: SMBG with no intervention “More” intensive: SMBG with intervention and education</p>	<p>Less £59</p> <p>More £56</p>	<p>Less -0.004 QALYs</p> <p>More -0.020 QALYs</p>	<p>Less Dominated</p> <p>More Dominated</p>	<p>SMBG dominated by no SMBG. Lifetime QALY gains are outweighed by initial negative impacts of SMBG; lifetime savings did not offset SMBG costs</p>	<p>Lifetime cost effectiveness results provide no convincing evidence for routine SMBG in people with type 2 diabetes not treated with insulin In PSA SMBG is cost effective at £20,000/QALY threshold in less than 40% of replication for the less intense arm and less than 15% for the more intense arm</p>
<p><b>Directly applicable</b><sup>c</sup></p>							
<p><b>Minor limitations</b><sup>f,n</sup></p>							

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p><b>Palmer et al. (2006)</b> People with existing type 2 diabetes, by treatment type SMBG v no SMBG England</p> <p><b>Directly applicable</b></p> <p><b>Some limitations<sup>g,i,j</sup></b></p>	<p><u>Effects:</u> HbA1c assumed</p> <p><u>Costs:</u> SMBG published acquisition costs, diabetic complication costs (£UK, 2004)</p> <p><u>Utilities:</u> sources not given. No disutility from SMBG</p>	<p>CDM model with lifetime horizon (length not stated)</p> <p>Test frequency: Diet and exercise 1/day, OADs 2/day Insulin 3/day</p> <p>HbA1c baseline: Diet and exercise 7.9% OADs 8.6% Insulin 8.5%</p> <p>HbA1c change: 1 test/day -0.3% 2 tests/day -0.4% 3 tests/day -0.6% Effect lasts for lifetime</p>	<p>Diet £2564</p> <p>OADs £1013</p> <p>Insulin £1171</p>	<p>Diet 0.165 QALYs</p> <p>OADs 0.225 QALYs</p> <p>Insulin 0.255 QALYs</p>	<p>Diet £15,515/ QALY</p> <p>OADs £4508/ QALY</p> <p>Insulin £4593/ QALY</p>	<p>ICERs for SMBG v no SMBG all fell well below the accepted threshold. However, uncertainty is large</p>	<p>ICERs sensitive to time horizon. (only insulin cost effective at 10 years), length of effect (diet and exercise not cost effective if only 5 years) and applying SMBG disutility (diet and exercise not CE).</p> <p>In PSA, SMBG cost effective at £30,000 in the following %s: Diet and exercise 51% OADs 51% Insulin 55%</p>
<p><b>Pollock et al. (2010)</b> People with existing type 2 diabetes treated with OADs SMBG 1, 2 or 3 times/day v none Switzerland</p> <p><b>Partly applicable<sup>a,b,d,e</sup></b></p> <p><b>Some limitations<sup>h,k</sup></b></p>	<p><u>Effects:</u> HbA1c observational study</p> <p><u>Costs:</u> Swiss unit costs used where available. No other details given (CHF, 2006)</p> <p><u>Utilities:</u> UKPDS and other literature. No disutility from SMBG</p>	<p>CDM with lifetime horizon (30 years) HbA1c baseline: 8.6%</p> <p>HbA1c change: 1 test/day -0.32% 2 tests/day -0.77% 3 tests/day -1.02%</p> <p>Lasts for 1 year Discounted at 3%</p> <p>Analysis not incremental, all compared to 0/day</p>	<p>1/day 528</p> <p>2/day 1650</p> <p>3/day 2899</p> <p>(all CHF)</p>	<p>1/day 0.058 .QALYs</p> <p>2/day 0.128 QALYs</p> <p>3/day 0.167 QALYs</p>	<p>1/day CHF9177/ QALY</p> <p>2/day CHF12,928 / QALY</p> <p>3/day CHF17,342 / QALY</p>	<p>SMBG is cost effective compared to no SMBG, as ICERs are below accepted Swiss thresholds</p>	<p>ICERs sensitive to HbA1c effect (cost effective as long as hbA1c effect of 0.08% or more), time horizon (not cost effective at 5 years) and using Swiss cohort baseline data</p> <p>In PSA, SMBG cost effective at CHF thresholds in following %s: CHF30,000 CHF80,000 1/day 60% 67% 2/day 68% 81% 3/day 72% 84%</p>

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty																											
			Costs	Effects	ICER																													
<p><b>Simon et al. (2008)</b> People with existing type 2 diabetes treated with diet exercise or OADs SMBG v none England</p> <p><b>Directly applicable<sup>c</sup></b></p> <p><b>Minor limitations<sup>n,o</sup></b></p>	<p><u>Effects:</u> DiGEM RCT</p> <p><u>Costs:</u> person level data from DiGEM RCT and standard UK unit costs (£UK, 2005)</p> <p><u>Utilities:</u> EQ-5D from DiGEM RCT; UK tariff. No disutility from SMBG</p>	<p>RCT – 1 year time horizon 0.9 strips/day (6/week) HbA1c baseline: 7.5% HbA1c effect for 1 year: Less -0.14% More -0.17% Control: standardised usual care “Less” intensive: SMBG with no intervention “More” intensive: SMBG with intervention and education</p>	<p>Less £92</p> <p>More £84</p>	<p>Less -0.008 QALYs</p> <p>More -0.036 QALYs</p>	<p>Less Dominated</p> <p>More Dominated</p>	<p>SMBG is dominated by SMBG.</p> <p>SMBG significantly more expensive than no SMBG and negative impact on QoL</p>	<p>Missing data techniques analysed, results do not change</p>																											
<p><b>Tunis and Minshall (2008)</b> People with existing type 2 diabetes treated with OADs SMBG v none USA</p> <p><b>Partly applicable<sup>a,b,d,e</sup></b></p> <p><b>Some limitations<sup>h,k,l</sup></b></p>	<p><u>Effects:</u> HbA1c observational study</p> <p><u>Costs:</u> relevant literature (\$US, 2006)</p> <p><u>Utilities:</u> UKPDS and other literature. No disutility from SMBG</p>	<p>CDM with lifetime horizon HbA1c baseline: 8.6% HbA1c change 0 tests/day 0.13% 1 test/day -0.32% 3 tests/day -1.02% Lasts for lifetime Discounted at 3% New SMBG users only Analysis not incremental Funded by industry</p>	<p>1/day \$808</p> <p>3/day \$2161</p>	<p>1/day 0.103 QALYs</p> <p>3/day 0.327 QALYs</p>	<p>1/day \$7856/ QALY</p> <p>3/day \$6601/ QALY</p>	<p>SMBG appears cost effective compared to no SMBG at accepted US thresholds but uncertainty is large.</p> <p>Some costs are offset by reduced complications and small QALY increases</p>	<p>ICERs sensitive to time horizon</p> <table border="1"> <tr> <td></td> <td>1/day</td> <td>3/day</td> </tr> <tr> <td>5 years</td> <td>\$23,380</td> <td>\$29,137</td> </tr> <tr> <td>10 years</td> <td>\$9346</td> <td>\$518</td> </tr> <tr> <td>Lifetime</td> <td>\$7856</td> <td>\$6601</td> </tr> </table> <p>ICERs sensitive to compliance (assessed via strip cost):</p> <table border="1"> <tr> <td>100%</td> <td>\$6601</td> </tr> <tr> <td>66%</td> <td>\$10362</td> </tr> <tr> <td>33%</td> <td>\$28676</td> </tr> </table> <p>In PSA, SMBG cost effective at US thresholds in the following %</p> <table border="1"> <tr> <td></td> <td>\$20k</td> <td>\$50k</td> </tr> <tr> <td>1/day</td> <td>51.6%</td> <td>52.6%</td> </tr> <tr> <td>3/day</td> <td>56.7%</td> <td>60.7%</td> </tr> </table>		1/day	3/day	5 years	\$23,380	\$29,137	10 years	\$9346	\$518	Lifetime	\$7856	\$6601	100%	\$6601	66%	\$10362	33%	\$28676		\$20k	\$50k	1/day	51.6%	52.6%	3/day	56.7%	60.7%
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Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty																				
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<p><b>Tunis et al. (2010)</b> People with existing type 2 diabetes treated with OADs SMBG 1, 2 or 3 times/day v none France, Germany, Italy, Spain</p> <p><b>Partly applicable</b><sup>a,b,d,e</sup></p> <p><b>Some limitations</b><sup>h,i,m</sup></p>	<p><u>Effects:</u> HbA1c observational study. Country specific for other therapies, treatment programmes, ESRD and mortality</p> <p><u>Costs:</u> Country specific SMBG costs and complication costs (€2007, country specific discounting)</p> <p><u>Utilities:</u> UKPDS and other literature. No disutility from SMBG</p>	<p>CDM with lifetime horizon (40 years) HbA1c baseline: 8.6% HbA1c change: 0 tests /day +0.13% 1 test/day -0.32% 2 tests /day -0.77% 3 tests /day -1.02%</p> <p>Lasts for lifetime Discounted at country specific rates</p> <p>Analysis not incremental, all compared to 0/day Funded by industry</p>	France	France	France	<p>SMBG at any frequency up to 3/day appears cost effective in all countries, compared to no SMBG (at accepted thresholds). However uncertainty is large</p> <p>Cost differences driven by different SMBG acquisition costs</p>	<p>ICERs sensitive to time horizon – not cost effective at 5 years for any countries or tests/day</p> <p>ICERs not sensitive to SMBG disutility (-0.036 in year 1 only). ICERs modestly increased, but remained within thresholds.</p> <p>In PSA, SMBG cost effective at €30,000/QALY threshold (€10,000 and €50,000 also given, similar %s)</p> <table border="1"> <thead> <tr> <th>%</th> <th>Fr</th> <th>Ger</th> <th>It</th> <th>Sp</th> </tr> </thead> <tbody> <tr> <td>1/day</td> <td>53</td> <td>55</td> <td>53</td> <td>54</td> </tr> <tr> <td>2/day</td> <td>56</td> <td>58</td> <td>54</td> <td>58</td> </tr> <tr> <td>3/day</td> <td>58</td> <td>59</td> <td>55</td> <td>59</td> </tr> </tbody> </table> <p>Different ICERs by country highlight the need for country specific analyses</p>	%	Fr	Ger	It	Sp	1/day	53	55	53	54	2/day	56	58	54	58	3/day	58	59	55	59
			%	Fr	Ger			It	Sp																		
			1/day	53	55			53	54																		
			2/day	56	58			54	58																		
			3/day	58	59			55	59																		
			1/day	1/day	1/day			€959	0.079	€12,114																	
			2/day	2/day	2/day			€1296	0.206	€6282																	
			3/day	3/day	3/day			€2101	0.264	€7958																	
			Ger	Ger	Germany			1/day	1/day	1/day	€213	0.130	€1633														
			2/day	2/day	2/day			€493	0.250	€1974																	
			3/day	3/day	3/day			€1561	0.309	€5045																	
			Italy	Italy	Italy			1/day	1/day	1/day	€1386	0.109	€12,694														
2/day	2/day	2/day	€2766	0.232	€11,934																						
3/day	3/day	3/day	€4660	0.303	€15,368																						
Spain	Spain	Spain	1/day	1/day	1/day	€325	0.089	€3661																			
2/day	2/day	2/day	€532	0.172	€3101																						
3/day	3/day	3/day	€1237	0.215	€5751																						

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<b>Tunis and Minshall (2010)</b> People with existing type 2 diabetes treated with OADs SMBG 1, 2 or 3 times/day v none USA <b>Partly applicable</b> <sup>a,b,d,e</sup> <b>Some limitations</b> <sup>h,k,l</sup>	<u>Effects:</u> HbA1c observational study <u>Costs:</u> relevant literature (\$US, 2006) <u>Utilities:</u> UKPDS and other literature. No disutility from SMBG	CORE model with lifetime horizon (40 years) HbA1c baseline: 7.6% HbA1c change: 0 tests/day -0.02% 1 test/day -0.14% 2 tests/day -0.34% 3 tests/day -0.37% Lasts for lifetime (assumed) Previous SMBG users Analysis not incremental, Funded by industry	1/day \$1225	1/day 0.047 QALYs	1/day \$26,208/ QALY	SMBG appears cost effective compared to no SMBG at US thresholds. However, uncertainty is large. Some costs are offset by reduced complications and small QALY increases	ICERs sensitive to time horizons. No option cost effective at 5 years, only 2/day cost effective at 10 years.  ICERs are worse than for new users (Tunis and Minshall, 2008) due to smaller treatment gain and lower baseline HbA1c  In PSA, percentages not presented as assume would not be favourable for SMBG
			2/day \$2147	2/day 0.116 QALYs	2/day \$18,572/ QALY		
			3/day \$3349	3/day 0.132 QALYs	3/day \$25,436/ QALY		

- a Not UK based analysis
- b Not UK baseline characteristics or treatment values
- c Only includes patients on diet and exercise and OADs
- d Only includes patients on OADs
- e Not UK discount rates
- f Limited details reported on modelling of future HbA1c trajectories
- g Baseline characteristics from observational study, rather than meta-analysis or RCT
- h Treatment effect from observational study, rather than meta-analysis or RCT
- i Treatment effect not source sourced systematically
- j Utility sources not specified
- k Limited cost details reported
- l Potential conflict of interest
- m PSA not well reported
- n Standard care arm may be better quality than real life
- o Limited time horizon

Study, Population, Comparators, Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Costs	Effects	ICER		
<p>CHF: Swiss francs            CDM: Centre for Outcomes Research Diabetes Model            DiGEM: diabetes glycaemic education and monitoring trial            EQ-5D: EuroQoL five dimension health-related quality of life questionnaire            Fr: France            Ger: Germany            HbA1c: glycosylated haemoglobin            ICER: incremental cost effectiveness ratio            It: Italy            OADs: oral anti-diabetic drugs            PSA: probabilistic sensitivity analysis            QALY: quality adjusted life years            QoL: quality of life            RCT: randomised controlled trial            SMBG: self-monitoring of blood glucose            Sp: Spain            UKPDS: United Kingdom Prospective Diabetes Study            UK: United Kingdom            US: United States of America</p>							

1

### 8.3.2.41 Evidence statements

#### 8.3.2.4.12 Clinical evidence

3 None of the studies reported evidence on diabetes-related complications.

#### 4 SMBG versus no SMBG

5 Evidence from a meta-analysis of 17 trials showed a small, clinically unimportant reduction in  
6 HbA1c levels with SMBG compared to no SMBG at up to 1 year. None of the subgroup  
7 analyses based on existing treatment (that is diet alone or combined with oral antidiabetic  
8 and/or insulin medicines), type of SMBG (standard or enhanced) or overall prescribed  
9 frequency of SMBG testing (that is less than once a day, 1 to 2 times a day or more than  
10 twice a day) showed a clinically important reduction in HbA1c levels. The quality of the  
11 evidence was low.

12 Evidence from a meta-analysis of the 6 trials reporting data on fasting blood glucose showed  
13 no significant changes in the 5 trials that included people who were treated with diet and/or  
14 oral antidiabetic medicines up to 1 year, but a significant reduction at 6 months in a trial of  
15 older adults who were on insulin therapy and undertaking SMBG (standard or enhanced)  
16 compared to no SMBG. Subgroup analyses based on overall prescribed frequency of SMBG  
17 testing showed no significant differences in fasting blood glucose in people undertaking  
18 SMBG compared to no SMBG. The quality of the evidence was low.

19 The low quality trial including older adults on insulin therapy also reported data on  
20 postprandial blood glucose levels and found a significant reduction in those undertaking  
21 SMBG (standard or enhanced) compared to no SMBG at 6 months.

22 A meta-analysis of 6 trials that reported any hypoglycaemic event showed a significantly  
23 increased risk in those undertaking SMBG compared to no SMBG for people on diet and/or  
24 oral antidiabetic medicines (4 studies), but no difference in risk for people on diet alone (1  
25 low quality study) or on diet, oral antidiabetic and/or insulin medicines (1 low quality study) up  
26 to 1 year. Subgroup analyses based on overall prescribed frequency of SMBG testing only  
27 showed a significantly increased risk in those undertaking SMBG less than once a day  
28 compared to no SMBG (2 studies). Overall, the quality of the evidence was low. A meta-  
29 analysis of the 3 trials that reported severe hypoglycaemic events showed low event rates  
30 and no significant difference in risk in those undertaking SMBG compared to no SMBG. One  
31 moderate quality trial showed no significant difference in risk in adverse events in people  
32 undertaking SMBG compared to no SMBG. The quality of the evidence was low.

#### 33 Different forms of SMBG

##### 34 *SMBG plus education versus conventional SMBG*

35 Overall, 2 meta-analyses were conducted on HbA1c levels and any hypoglycaemic events  
36 for 3 studies that examined SMBG plus education compared to standard SMBG on people  
37 treated with diet and/or oral antidiabetic and/or insulin medicines up to 1 year. Overall, no  
38 significant differences in HbA1c levels and hypoglycaemic events were observed in people  
39 undertaking SMBG plus education compared to SMBG alone. However, 1 very low quality  
40 trial showed a significant clinically relevant reduction in HbA1c levels at 3 months in people  
41 on oral antidiabetic and/or insulin medicines who were undertaking SMBG plus education  
42 compared to SMBG alone. Overall, the quality of the evidence was low.

1 *SMBG plus telecare versus conventional SMBG*

2 A meta-analysis of 5 trials showed a non-significant reduction in HbA1c levels up to 44  
3 weeks in people on diet, oral antidiabetic and/or insulin undertaking SMBG plus telecare  
4 compared to SMBG only (3 studies), but a significant and clinically important reduction in  
5 HbA1c levels was observed in favour of SMBG plus telecare compared to SMBG alone in 2  
6 trials that did not specify the diabetes treatment that people were receiving. Overall, the  
7 quality of the evidence was low. Two low quality trials also reported data on fasting blood  
8 glucose up to 44 weeks which showed no significant differences in people on diet, oral  
9 antidiabetic medicines and/or insulin undertaking SMBG plus telecare compared to SMBG.  
10 One low quality trial additionally reported data on postprandial blood glucose levels and any  
11 hypoglycaemic events, and showed no significant differences at 26 weeks between people  
12 on diet, oral antidiabetic and/or insulin undertaking SMBG plus telecare compared to SMBG  
13 alone in either of these outcomes.

14 *Automated mobile phone glucometer versus standard glucometer*

15 One small, low quality trial showed no significant differences in blood glucose measures  
16 (HbA1c, fasting and postprandial blood glucose) at 3 months in SMBG using an automated  
17 glucometer compared to a standard glucometer in people with unspecified current diabetes  
18 treatments.

19 *SMBG plus continuous glucose monitoring (CGM) versus conventional SMBG*

20 Overall, a meta-analysis of 2 trials showed a significant and clinically important reduction in  
21 HbA1c levels in people on insulin undertaking SMBG plus CGM compared to those on  
22 SMBG alone up to 12 months. The quality of the evidence was very low. One low quality trial  
23 reported no significant differences in fasting and postprandial blood glucose at 3 months in  
24 people on insulin undertaking SMBG plus CGM compared to those on SMBG alone.

25 *Frequency and location of SMBG testing*

26 Two moderate to low quality trials showed no clinically important differences in HbA1c levels  
27 in people treated with oral antidiabetic medicines undertaking monthly versus fortnightly self-  
28 monitoring or 4 times weekly versus once weekly monitoring. There was an increased risk of  
29 any hypoglycaemic event with increased monitoring.

30 High to moderate quality evidence from 1 trial in people with type 2 diabetes treated with  
31 insulin showed that there were no clinically important differences in HbA1c levels or  
32 hypoglycaemia associated with forearm versus fingertip testing.

**8.3.2.4.23 Health economic evidence**

34 Two directly applicable CUAs with minor limitations found that, for people with type 2  
35 diabetes treated with diet or OADs, SMBG was more costly and produced less QALYs than  
36 no SMBG.

37 Four partly applicable CUAs with potentially serious limitations that based their treatment  
38 effect on the same US observational study found SMBG to be cost effective, though there  
39 was substantial uncertainty in their results.

**8.3.30 Evidence to recommendations**

41 **Table 41: Linking evidence to recommendations**

Relative value of  
different outcomes

The GDG agreed that impact on blood glucose levels,  
hypoglycaemia and diabetes-related complications were critical

	<p>to decision making.</p> <p>The GDG noted that while SMBG provides the potential for tight glycaemic control and therefore reduced risk of diabetes-related complications, the possible impact of such control on hypoglycaemic events is important in determining the safety and acceptability to patients.</p> <p>The GDG agreed that all outcomes were weighted equally, and noted their importance in decision making with respect to treatment choices and associated patient compliance, safety and costs. However, specific to blood glucose measures, the Group agreed that HbA1c was more important than fasting and postprandial blood glucose.</p>
Trade-off between benefits and harms	<p>The GDG discussed the evidence presented for SMBG compared to no SMBG and agreed that overall, while a difference was observed in HbA1c levels in favour of SMBG, the small reduction at less than 5 mmol/mol (0.5%; the threshold for minimal important difference) was not clinically meaningful. In addition, the GDG noted that no specific subgroup in terms of current diabetes treatment, type or frequency of SMBG, was shown to have a clinically meaningful reduction in HbA1c levels.</p> <p>The GDG discussed the higher incidence of any hypoglycaemia observed in the SMBG group compared to no SMBG, and agreed that most of the reported events in the studies were minor or asymptomatic. The GDG thought it likely that the greater occurrence of hypoglycaemic events in the SMBG group was related to increased detection, rather than an increased risk of events associated with self-monitoring. The Group noted that asymptomatic hypoglycaemia also occurs in people who do not have diabetes, and discussed the relative importance of these events compared to symptomatic hypoglycaemia. The GDG noted the low numbers of severe hypoglycaemic events that were reported in the studies in both SMBG and no SMBG groups. The GDG discussed the role of baseline HbA1c levels and its possible association with hypoglycaemic events, and noted that hypoglycaemia can occur for various reasons at different baseline HbA1c levels.</p> <p>The GDG discussed the evidence presented for the different forms of SMBG, and noted that generally there was no difference in HbA1c levels and hypoglycaemic events between enhanced SMBG (education, telecare, automated glucometer) and conventional SMBG. The GDG noted that there was little evidence on frequency and location of SMBG testing, but findings from the 3 included studies also showed no difference in HbA1c levels and hypoglycaemic events between the groups comparing more frequent (every 2 weeks or 4 times a week) and less frequent (monthly or once a week) SMBG and different sites of testing (forearm or fingertip). The GDG discussed the conflicting evidence presented for continuous glucose monitoring compared to standard SMBG from 2 small, low quality trials in people on insulin, where 1 trial showed no difference in HbA1c levels at 3 months while the second trial showed a clinically important reduction in HbA1c levels at 12 months. The Group</p>

	<p>agreed that there was still uncertainty regarding the effectiveness of continuous glucose monitoring.</p> <p>The GDG noted the overall lack of evidence on diabetes-related complications.</p>
<p>Consideration of health benefits and resource use</p>	<p>All of the modelled CUAs were based on existing economic models, meaning that many of the underlying assumptions, probabilities and utilities in the CUAs were the same. Given this, the GDG agreed that key factors in assessing the quality of the evidence were the country costs used and the source of the HbA1c change estimates.</p> <p>The GDG thought the CUAs based on UK evidence that took their HbA1c change level from RCTs were the most applicable evidence with fewest limitations. These studies found self-monitoring to be not cost effective compared with no self-monitoring.</p> <p>CUAs based on observational American evidence used a higher level of HbA1c change, but the GDG considered that the potential role of confounders in the observational evidence rendered that evidence too unreliable to be used in CUAs.</p> <p>Evidence was presented that showed a roughly linear increase in net monetary benefit with increasing HbA1c change modelled. The GDG did not think that the larger HbA1c changes modelled in some CUAs were likely to be achievable. They noted the clinical evidence review found a 2 mmol/mol (0.22%) decrease in HbA1c associated with self-monitoring and this was unlikely to be cost effective compared with modelled changes in HbA1c. Also, the GDG felt the CUAs that assumed the HbA1c impact of self-monitoring would last for a patient lifetime were unrealistic.</p> <p>The GDG considered that the high degree of uncertainty displayed by CUAs that found self-monitoring to be cost effective compared with no self-monitoring meant that they could not conclude such studies gave convincing evidence of cost effectiveness of the intervention. Also, the GDG noted the correlation between industry funding and positive cost-effectiveness conclusions compared to the negative cost-effectiveness conclusions of non-industry funded studies.</p> <p>The GDG highlighted a number of gaps in the economic evidence. Few CUAs reported results for people with type 2 diabetes using insulin and no CUAs reported results for newly diagnosed patients. No health economic evidence of a quality level high enough to be included was found to assess the marginal benefits of increasing the frequency of self-monitoring.</p> <p>Overall, the GDG thought the economic evidence did not make it possible to state conclusively that self-monitoring is or is not likely to be cost effective compared to no self-monitoring, but the most applicable evidence with least limitations suggested that self-monitoring is not likely to be cost effective compared to no self-monitoring.</p>
<p>Quality of evidence</p>	<p>The GDG noted that the quality of the evidence varied from high</p>

to very low, but agreed overall that the quality was low.

Specific to the comparison of SMBG and no SMBG, the GDG noted that although most of the trials were based in western countries, only 1 study was conducted in the UK and that most participants were on diet and/or oral antidiabetic medicines, rather than diet alone or insulin. The GDG noted that while people on insulin therapy are able to titrate their dose based on what they eat, this is not normally the case for people who are controlled by oral blood glucose lowering therapies, and this may have an impact on their compliance with the use of self-monitoring.

For the comparisons on different forms of SMBG, 4 of the 14 trials were conducted in Korea, where people with type 2 diabetes are generally slimmer and may have different diet and lifestyles compared with people living in the UK. In addition, some trials reported mean HbA1c levels at baseline close to 53 to 58 mmol/mol (7 to 7.5%) showing good blood glucose control, which the GDG agreed may not be representative of people with type 2 diabetes in the UK. The GDG also noted that 1 of these trials restricted inclusion to people aged 60 years and over, and agreed that older adults tend to have more comorbidities and therefore drug therapy selection may vary.

The GDG noted that people who are recruited into trials may be more likely to be motivated to carry out self-monitoring. In addition, the group agreed that the differing quality of information across the trials may have influenced the results. The GDG agreed that it would be difficult to draw conclusions based on current treatments and intensity of treatment regimens, as the evidence base largely covered mixed populations of people on several different blood glucose lowering therapies without any subgroup analyses.

#### Other considerations

The GDG discussed the recommendations from the DVLA, and noted that accidents involving driving were not an outcome for this review question. The GDG noted that the DVLA recommends that people driving cars and motorcycles need to inform the DVLA if they start insulin therapy. More recent guidance is available for people driving buses and lorries, which states that drivers on insulin need to show adequate control by regular self-monitoring. Currently there are no specific details on self-monitoring for people who are taking oral blood glucose lowering therapies. The GDG discussed people treated with sulfonylureas in particular, and noted the increased risk of hypoglycaemia, similar to insulin therapy. Because of the different pharmacodynamics of these drug classes, it was suggested that hypoglycaemic events experienced with sulfonylureas may be slower to occur, and come on so gradually that people are less aware of them, and in addition they may last for longer compared with insulin-induced events. The GDG discussed the negative impact this would have on people, for example, falls, and also other implications, in terms of medication selection and target setting for HbA1c levels.

The GDG discussed the effects of corticosteroids, including

increased insulin resistance, which may lead to higher plasma glucose levels. The Group discussed the greater risks of hyperglycaemia for people with type 2 diabetes who start on corticosteroid therapy and agreed that these individuals would benefit from short-term self-monitoring.

The GDG noted the limited evidence on people on insulin, and agreed that it may not be appropriate to extrapolate the evidence base from people with type 1 diabetes due to differences in the characteristics of this group (for example people with type 1 diabetes will have been testing blood glucose levels for several years, as the age of onset is much younger, and they may also be more familiar with the effect of dietary intake on blood glucose levels, that is, glycaemic index and carbohydrate counting).

The GDG noted that SMUG is not used in clinical practice. In particular, it was noted that for people treated with newer SGLT-2 inhibitors such as dapagliflozin, sugars are excreted through the urine and so testing urine glucose levels would not be appropriate. In addition, it was noted that continuous glucose monitoring is not routinely used for people with type 2 diabetes.

The GDG discussed the importance of individual preferences for SMBG because, while some people may find it useful, others may find it has a negative impact on quality of life.

The GDG also noted the lack of evidence concerning the frequency of SMBG and specific target values when SMBG is used. The GDG felt unable to make any recommendations on these issues and chose instead to draft 2 research recommendations.

When making recommendations for the use of self-monitoring, the GDG considered the following points:

- Overall, the evidence showed a small reduction in HbA1c levels that was not clinically important.
- There was uncertainty around whether self-monitoring was cost-effective, but the GDG felt it was unlikely to be at the magnitude of HbA1c changes reported.
- Some medications have been shown to increase the risk of hypoglycaemia.

Overall, a strong 'do not do' recommendation was made for the majority of people with type 2 diabetes, because the GDG agreed that self-monitoring would not be of sufficient benefit for most people. However, exception groups were added to this recommendation, because the GDG agreed it was important to offer targeted self-monitoring to people at higher risk of experiencing hypoglycaemic events. This included people who are taking insulin therapy, other oral medication or those who experienced symptomatic hypoglycaemia. The GDG also added a further recommendation for healthcare professionals to refer to the DVLA to ensure that targeted self-monitoring was carried out in accordance with legislative guidance.

### 8.3.4.1 Recommendations and research recommendations

- 2 **43. Take the Driver and Vehicle Licensing Agency (DVLA) [At a glance guide to the](#)**  
3 **[current medical standards of fitness to drive](#) into account when offering self-**  
4 **monitoring of blood glucose levels. [new 2015]**
- 5 **44. Do not routinely offer self-monitoring of blood glucose levels for adults with type**  
6 **2 diabetes unless the person:**
- 7 • is on insulin **or**
  - 8 • experiences symptomatic hypoglycaemia **or**
  - 9 • is on oral medication that may increase their risk of hypoglycaemia while  
10 driving or operating machinery **or**
  - 11 • is pregnant, or is planning to become pregnant. For more information,  
12 see the NICE guideline on diabetes in pregnancy.
- 13 Consider short-term self-monitoring for adults with type 2 diabetes who start treatment  
14 with oral or intravenous corticosteroids. **[new 2015]**
- 15 **45. If adults with type 2 diabetes are self-monitoring their blood glucose levels, carry**  
16 **out a structured assessment at least annually. The assessment should include:**
- 17 • the person's self-monitoring skills
  - 18 • the quality and frequency of testing
  - 19 • how the results are used
  - 20 • the impact on the person's quality of life
  - 21 • the continued benefit to the person
  - 22 • the equipment used. **[2015]**

### 23 Research recommendations

- 24 **3. What is the optimal frequency for self-monitoring of blood glucose in adults with**  
25 **type 2 diabetes?**
- 26 **4. What are the optimal blood glucose targets for self-monitoring in adults with type**  
27 **2 diabetes?**

### 28 Why this is important

29 It is widely recognised that self-monitoring of blood glucose is a multicomponent  
30 intervention. As well as being educated about how to use a self-monitoring device to  
31 assess blood glucose levels, adults with type 2 diabetes need to be able to understand  
32 their results and act on the observed readings. In adults for whom self-monitoring is  
33 appropriate, there is limited evidence to guide clinical practice in prescribing self-  
34 monitoring regimens, in terms of frequency of testing and optimal glycaemic targets.  
35 Given the inconvenience and expense of self-monitoring, robust evidence from double-  
36 blind randomised controlled trials is needed to guide the optimal use of this intervention.

## 8.4.1 Drug treatment to control blood glucose

2 Lifestyle interventions such as diet and physical activity are commonly used to initially  
3 manage type 2 diabetes. However, it is uncommon for individuals to maintain glycaemic  
4 control to target levels for extended periods of time using only these interventions. Because  
5 type 2 diabetes is a progressive condition, with secretion of insulin decreasing over time,  
6 blood glucose lowering medicines are often indicated. The choice, order and combination in  
7 which these treatments are used will reflect consideration of the following:

- 8 • prevention of microvascular and arterial damage
- 9 • glycaemic control
- 10 • assessment of the inconvenience
- 11 • risks of side effects.

12 The benefits, side effects and relative cost-effectiveness differ among pharmacological  
13 classes, and to a lesser extent between individual drugs within the same class. The clinical  
14 questions covered in this section are concerned with the selection of optimal drug treatment  
15 strategies for people with type 2 diabetes, taking into consideration individual characteristics  
16 such as occupation and body mass index.

### 8.4.1.7 Clinical introduction

#### 8.4.1.18 Approach to drug treatment

19 The approach to drug treatment to control blood glucose levels in people with type 2 diabetes  
20 was discussed in detail with the GDG. Due to the progressive nature of the condition, the  
21 main assumption underpinning the analysis for this review question, is that augmenting  
22 existing drug treatments with additional medicines will provide better glycaemic control (see  
23 Figure 5: Overview of intensification of drug treatment as blood glucose control  
24 declines). The rationale for this is that the added medicines will have a different mode of  
25 action that is complementary to the existing drug treatment.

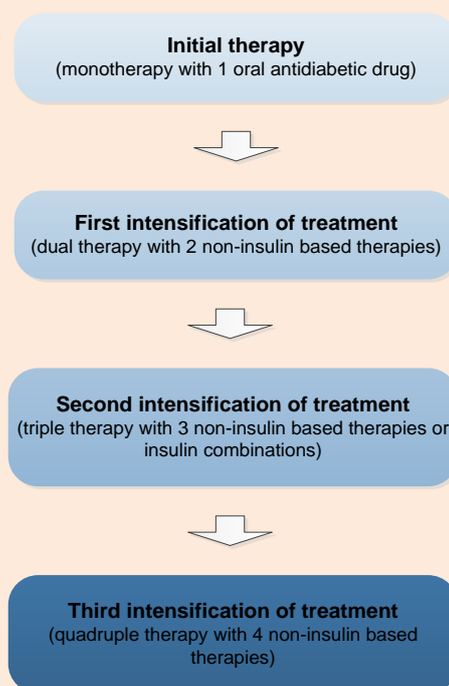


Figure 5: Overview of intensification of drug treatment as blood glucose control

**declines**

**8.4.1.21 Drug treatment to control blood glucose in Clinical Guidelines 66 and 87**

2 Pharmacological management of blood glucose levels was originally covered as part of  
3 CG66 and CG87. The searches in CG66 were conducted from 2001 to 2007 (see Appendix  
4 G for search strategies from CG66) and searches in CG87 were conducted from 1990 to  
5 2008 (see Appendix I for search strategies from CG87).

6 In previous versions of this guideline, the GDG also prioritised a series of pairwise  
7 comparisons that were of particular clinical interest and where possible, meta-analyses were  
8 conducted to combine the results of studies for different outcomes.

**8.4.1.39 Drug treatment to control blood glucose in the update (2015)**

10 For the current update, further searches have been carried out for drug treatments previously  
11 reviewed in CG66 (metformin, sulfonylureas, acarbose and insulin) with a date restriction of  
12 2007 to June 2014; for drugs covered in CG87 (DPP-4 inhibitors that is saxagliptin, sitagliptin  
13 and vildagliptin; GLP-1 mimetics that is exenatide and liraglutide; insulin that is glargine and  
14 detemir; thiazolidinediones that is pioglitazone) with a date restriction of 2008 to June 2014;  
15 and for interventions not previously covered (DPP-4 inhibitors that is linagliptin; GLP-1  
16 mimetics that is lixisenatide) with no date restrictions (see Appendix C for update search  
17 strategies).

18 The evidence that was originally included in CG66 and CG87 was re-reviewed as part of the  
19 update. For this update, a series of 4 network meta-analyses (NMAs) were proposed rather  
20 than a series of pairwise comparisons. Details of the definitions and included drug  
21 comparisons for each phase of treatment are set out in Table 43. Overall, the following  
22 phases of clinical treatment were agreed and formed the area of 4 sub-review questions:

- 23 • initial therapy (monotherapy)
- 24 • first intensification (dual therapy)
- 25 • second intensification (triple therapy and treatment combinations containing insulin)
- 26 • third intensification (quadruple therapy compared with triple therapy)

27 The aim of these sub-review questions was to identify which medicines were most effective  
28 in each phase, once treatment initiation or intensification was considered to be clinically  
29 indicated. Importantly, this meant that drug comparisons across the phases of treatment (for  
30 example initial therapy with metformin compared to first intensification with metformin plus  
31 sulfonylurea) were not included in this review question. Table 42 provides information on the  
32 different drug treatments that were considered for this review question. For each treatment  
33 phase, the review also focused on the specific drug comparisons listed in Table 43 that the  
34 GDG prioritised as clinically important.

35 The evidence for each treatment phase (that is initial therapy, followed by first, second and  
36 third intensification) is reviewed and analysed separately, although the results from each  
37 sub-review question will be used to inform a single treatment algorithm for people with type 2  
38 diabetes (see 1.4).

**8.4.1.49 Assumptions underpinning analytical approach**

40 With regard to both the decision problems adopted and the evidence considered relevant to  
41 those problems, the approach adopted by the GDG had important assumptions and  
42 implications that should be made explicit.

43 The GDG advised that differences in previous treatment history in cohorts recruited to RCTs  
44 are likely to reflect prescriber preferences, rather than fundamental clinical differences.  
45 Therefore, it was assumed that the treatment effects observed in trials solely reflect the

1 regimens to which people had been randomised in each RCT, and not the treatments they  
2 had previously received. This assumption was especially relevant for first intensification and  
3 second intensification. The assumption implies that, as long as trials met the eligibility criteria  
4 for the relevant decision problem, the identity of the treatments that had failed to control  
5 participants' HbA1c prior to recruitment should not be considered a material determinant of  
6 treatment effect. By this logic, the GDG was content to assume that, for example, it was valid  
7 to pool trials in which metformin+sulfonylurea were given to people for whom metformin  
8 monotherapy had proved inadequate in controlling HbA1c with trials in which  
9 metformin+sulfonylurea were given to people for whom sulfonylurea monotherapy had  
10 proved inadequate.

11 The GDG recognised that the prevailing approach to pharmacotherapy in type 2 diabetes  
12 contrasts with that adopted in many other clinical areas, where people for whom 1 treatment  
13 proves inadequate would commonly discontinue that treatment and switch to another, with  
14 the result that people may be on different monotherapies or combinations at different phases  
15 of the treatment pathway. In those contexts, treatment history may become a critical  
16 component of potentially important clinical heterogeneity. In contrast, the common clinical  
17 pathways in type 2 diabetes are well established, and this is reflected in many of the RCTs  
18 providing evidence for this review. In particular:

- 19 • Most people meeting the GDG's definition of first intensification of pharmacotherapy have  
20 poorly controlled HbA1c despite prior treatment with an appropriate dose of metformin  
21 monotherapy;
- 22 • Many participants in second intensification trials have, prior to recruitment, experienced  
23 suboptimal blood glucose control on combination treatment with metformin-sulfonylurea.

24 The fair degree of homogeneity apparent in this evidence has advantages and  
25 disadvantages, from the perspective of evidence synthesis. It is positive because it reinforces  
26 the appropriateness of pooling the data (the so-called 'transitivity' of treatment effects). On  
27 the other hand, it is unhelpful that any inferences drawn beyond the common pathway  
28 invariably rely on a degree of extrapolation. For example, the GDG considered it would be  
29 helpful to make recommendations for people for whom metformin is contraindicated, but no  
30 RCTs were identified of treatments in this population. Therefore, the GDG had little option  
31 but to assume that the best options for those who cannot take metformin are the non-  
32 metformin options that provide greatest effects in trials in the broader population (even  
33 though that population predominantly comprises metformin-tolerant people).

34 One important implication of this approach was that the GDG believed it was appropriate to  
35 exclude RCTs from 2 categories:

- 36 •  $a$  versus  $a+b$  (commonly  $a+placebo$  versus  $a+b$ ). These trials were not considered  
37 relevant because, from the perspective of the specified decision problems, they conflate  
38 different phases of treatment (that is, people who require 1 treatment and those who  
39 need 2). The GDG believed it was reasonable to take it as given that intensification of  
40 therapy has effects, and the question of the appropriate point in the treatment pathway at  
41 which to intensify treatment should be examined separately (see 8.1 and 8.2).
- 42 •  $a+(c, d \text{ or } e)$  versus  $b+(c, d \text{ or } e)$ . Because the GDG's interest was in the particular  
43 combination of medicines that may be given, experimental designs in which a single agent  
44 was added to a heterogeneous collection of 'background' therapies were not considered  
45 informative, unless they contained enough detail to isolate the effect of particular  
46 combinations (in this example, a trial would only be considered to provide relevant  
47 evidence if it reported subgroup results for  $a+c$  versus  $b+c$ ,  $a+d$  versus  $b+d$  and  $a+e$   
48 versus  $b+e$ ; such comparisons would be entered into synthesis as independent  
49 observations. In practice, no such trials were identified).

### 8.4.21 Review question

2 The overarching review question for this section is “Which pharmacological blood glucose  
3 lowering therapies should be used to control blood glucose levels in people with type 2  
4 diabetes?”

5 The overall review question was broken down into 4 further sub-questions:

- 6 • Which pharmacological blood glucose lowering therapies should be used initially to control  
7 blood glucose levels in people with type 2 diabetes?
- 8 • When first intensification of treatment is indicated, which blood glucose lowering therapies  
9 should be used to control blood glucose levels?
- 10 • When second intensification of treatment is indicated, which blood glucose lowering  
11 therapies should be used to control blood glucose levels?
- 12 • When third intensification of treatment is indicated, which blood glucose lowering  
13 therapies should be used to control blood glucose levels?

14 **Table 42: Blood glucose lowering drug treatments included in the review**

Drug class	Drug	Route of administration	Recommended daily doses
Alpha-glucosidase inhibitors	Acarbose	Oral	50 to 600 mg
Biguanides	Metformin	Oral	500 to 3000 mg
	Metformin modified release		500 to 2000 mg
Dipeptidyl-peptidase 4 inhibitors (DPP-4 inhibitors)	Linagliptin	Oral	5 mg
	Saxagliptin		5 mg
	Sitagliptin		100 mg
	Vildagliptin		100 mg
Meglitinides	Nateglinide*	Oral	180 to 540 mg
	Repaglinide		0.5 to 16 mg
Sulfonylureas	Glibenclamide/ Glyburide	Oral	2.5 to 15 mg
	Gliclazide		40 to 320 mg
	Gliclazide modified release		30 to 120 mg
	Glimepiride		1 to 6 mg
	Glipizide		2.5 to 20 mg
	Tolbutamide		500 to 2000 mg
Thiazolidinediones	Pioglitazone	Oral	15 to 45 mg
Glucagon-like peptide-1 agonists (GLP-1 mimetics)	Exenatide*	Subcutaneous	10 to 20 mcg
	Exenatide modified release*		
	Liraglutide* Lixisenatide*		0.6 to 1.8 mg 10 to 20 mcg
Insulin	Biphasic insulin aspart	Subcutaneous	variable
	Insulin aspart		
	Insulin degludec		
	Insulin detemir		
	Insulin glargine		
	Insulin lispro		
	Neutral protamine		
	Hagedorn insulin (NPH)		

Drug class	Drug	Route of administration	Recommended daily doses
	insulin)		
<i>Information taken from the British National Formulary and summary of product characteristics; * not licensed for monotherapy</i>			

1 **Table 43: Definitions of treatment phases and included drug comparisons to control blood glucose**

Phase of clinical treatment	Definition	Included drug comparisons
Initial therapy (monotherapy)	This phase refers to treatment with a single non-insulin based blood glucose lowering therapy. This is generally appropriate for people who are newly diagnosed with type 2 diabetes or who are at an early stage and have previously been treated on diet and exercise alone	The following drug comparisons were included for initial therapy: <ul style="list-style-type: none"> <li>• 1 oral antidiabetic versus 1 oral antidiabetic</li> <li>• 1 oral antidiabetic versus placebo</li> </ul>
First intensification (dual therapy)	This phase refers to treatment with 2 non-insulin based blood glucose lowering therapies in combination. This is generally appropriate for people with type 2 diabetes who are not achieving adequate glycaemic control with a single non-insulin based therapy	The following drug comparisons were included for first intensification: <ul style="list-style-type: none"> <li>• 2 non-insulin therapies versus 2 non-insulin therapies</li> </ul>
Second intensification (triple therapy)	This phase refers to treatment with either 3 non-insulin based blood glucose lowering therapies (triple therapy) or any treatment combination containing insulin. This is generally appropriate for people with type 2 diabetes who are not achieving adequate glycaemic control with 2 non-insulin based therapies	The following drug comparisons were included for second intensification: <ul style="list-style-type: none"> <li>• 3 non-insulin therapies versus 3 non-insulin therapies</li> <li>• Insulin versus 3 non-insulin therapies</li> <li>• Insulin + 1 non-insulin therapy versus 3 non-insulin therapies</li> <li>• Insulin + 2 non-insulin therapies versus 3 non-insulin therapies</li> <li>• Insulin versus insulin + 1 non-insulin therapy</li> <li>• Insulin versus insulin + 2 non-insulin therapies</li> <li>• Insulin + 1 non-insulin therapy versus insulin + 1 non-insulin therapy</li> <li>• Insulin + 2 non-insulin therapies versus insulin + 2 non-insulin therapies</li> <li>• Insulin + 1 non-insulin therapy versus insulin + 2 non-insulin therapies</li> </ul>
Third intensification (quadruple therapy)	This phase refers to possible treatment with 4 non-insulin based blood glucose lowering therapies in combination. This is generally appropriate for people with type 2 diabetes who are not achieving adequate glycaemic control with therapies considered at second intensification	The following drug comparisons were included for third intensification: <ul style="list-style-type: none"> <li>• 4 non-insulin therapies versus 3 non-insulin therapies</li> </ul>
<i>Non-insulin therapy includes both oral and injectable non-insulin agents; all included drug comparisons followed current summary of product characteristics (SPC) and licensed indications</i>		

1 RCTs with a minimum of 12 weeks of treatment and follow-up in people with type 2 diabetes  
2 were included for this review question. Several main exclusion criteria were used across all  
3 sub-review questions and these are outlined below:

- 4 • Non-randomised evidence (including observational, cohort, case-control and case series  
5 studies, uncontrolled or single arm trials), narrative reviews, conference abstracts, letters,  
6 editorials and trial protocols.
- 7 • Studies including a mixed population of people with type 1 and 2 diabetes, unless  
8 subgroup analyses were reported or 85% or more of the study population have type 2  
9 diabetes.
- 10 • Comparisons with unlicensed indications (for example, GLP-1 mimetics for use in initial  
11 therapy), unlicensed modes of delivery (for example, inhaled insulin), drugs not included  
12 in the scope and drug comparisons not of interest (for example, comparisons across  
13 treatment phases).
- 14 • Studies focusing on markers of cardiovascular disease or other diabetic complications  
15 without any blood glucose measures (HbA1c).
- 16 • Unclear washout of existing drug treatments, where a proportion or all participants  
17 continued previous medicines that will likely confound study results (papers were  
18 excluded unless this represented a small proportion of patients that is less than 5%).
- 19 • Unclear if analyses were adjusted in trials where rescue medication was given.

20 Further specific criteria are reported in the evidence review for each sub-question. For the full  
21 excluded list, see Appendix L.

22 The outcomes that were selected by the GDG as critical and important to decision making for  
23 the clinical evidence review are listed below.

24 **Table 44: Critical and important outcomes**

Critical outcomes	Important outcomes
<ul style="list-style-type: none"> <li>• Change in blood glucose levels (HbA1c)*</li> <li>• Hypoglycaemia*</li> <li>• Adverse events (total dropouts, dropouts due to adverse events*, nausea)</li> </ul>	<ul style="list-style-type: none"> <li>• Change in body weight*</li> </ul>
*Studies reporting all of these 4 outcomes were included in health economic model	

25 The detailed protocol is available in Appendix C.

26 Sensitivity analyses to determine whether participants' previous exposure to blood glucose  
27 lowering therapies affected the network meta-analyses results, for change in HbA1c at 12  
28 months and hypoglycaemia at study end point were undertaken for each treatment phase.  
29 These critical outcomes were selected as they represented the more important outcomes  
30 and provided evidence for benefits and harms. One-year follow-up was prioritised for HbA1c  
31 as this was used in the health economic model. Table 43 describes the typical population  
32 characteristics for each treatment phase, which were used to inform the sensitivity analyses.

- 33 • For initial therapy, people are usually drug naïve and are managed using dietary changes  
34 only, with no previous experience of taking blood glucose lowering pharmacological  
35 treatments. Some of the included studies for initial therapy had participants who were  
36 previously on drug treatments. Therefore, sensitivity analyses on individuals who were  
37 completely drug naïve were undertaken. The sensitivity analyses showed that, overall,  
38 there was little difference in the direction of effect for changes in HbA1c and  
39 hypoglycaemia, between drug naïve individuals and the full population which included  
40 individuals who were previously exposed and "washed-off" of prior anti-hyperglycaemic  
41 medications (see Appendix J). Therefore, the full analyses were used and reported in  
42 section 8.4.4.2.

- 1 • For first intensification of treatment, sensitivity analyses were undertaken on the typical  
2 population for this phase of treatment, that is, people who were previously on 1 oral  
3 antidiabetic medicine, including those whose medication had failed to adequately control  
4 blood glucose levels. No major differences were observed in the direction of effect for  
5 changes in HbA1c and hypoglycaemia, between people on 1 oral antidiabetic medicine  
6 and the full population which included studies of mixed populations of people who were  
7 drug naïve, or on 1 or more oral antidiabetic medicines at screening (see Appendix J).  
8 Therefore, the full analyses were used and reported in section 8.4.8.2.
- 9 • For second intensification of treatment, sensitivity analyses were undertaken on the  
10 typical population for this phase of treatment, that is, people who were previously on 2  
11 non-insulin based therapies, including those whose medication had failed to adequately  
12 control blood glucose levels. No major differences in the direction of effect for changes in  
13 HbA1c and hypoglycaemia, between people on 2 antidiabetic medicines and the full  
14 population which included studies of mixed populations of people who did not necessarily  
15 fail on/or were previously exposed to 2 drugs, or studies of people who failed on 1 oral  
16 antidiabetic drug were observed (see Appendix J). Therefore, the full analyses were used  
17 and reported in section 8.4.12.2.

### 8.4.38 Health economic methods

#### 8.4.3.19 Health economic evidence – search methodology

20 Previous guidelines (CG66 and CG87) have conducted health economic literature searches  
21 focused on specific drug comparisons that did not include initial therapy comparisons. For  
22 the current guideline, 1 systematic literature review with no date restrictions was undertaken  
23 to identify all existing cost–utility analyses (CUAs) addressing all 3 review sub–questions and  
24 yielded 3963 citations (see Appendix C for the search strategy).

25 In total 81 CUAs of pharmacological management of type 2 diabetes were found. Of these 81  
26 CUAs, 79 were funded by a pharmaceutical manufacturer and found the sponsor’s drug to be  
27 cost effective (see appendix F for a full list of the 81 CUAs). Two HTA-type studies found that  
28 the older, less expensive drugs provided better value for money than newer drugs.

29 For this guideline, in addition to meeting the NICE reference case (National Institute for  
30 Health and Care Excellence 2012) and covering included drug comparisons, 2 additional  
31 exclusion criteria were agreed by the GDG:

- 32 • Trial-based evaluations (that is, not extrapolated to lifetime outcomes) were excluded
- 33 • Non UK based CUAs were excluded.

34 As no directly applicable studies with only minor limitations were found that covered all the  
35 comparators under consideration for each sub-question for this guideline, an original  
36 economic analysis was undertaken.

#### 8.4.3.27 Original health economic modelling – methods

38 A full description of the health economic model can be found in Appendix F; a summary is  
39 presented here. The model was developed in line with the NICE reference case (National  
40 Institute for Health and Care Excellence 2012). A single health economic model structure  
41 was developed to address all 3 sub-questions for review question 1.

42 Along with the option of building a completely new model, a number of health economic  
43 diabetes models already exist (Mount Hood 4 Modeling Group 2007, Yi et al, 2010). The  
44 GDG selected the UKPDS Outcomes Model version 1 (UKPDS OM1, Clarke et al. 2004) as it  
45 matched the NICE reference case (National Institute for Health and Care Excellence 2012),  
46 was internally and externally validated and allowed greatest flexibility for modelling of  
47 additional short term outcomes.

1 The UKPDS OM1 does not directly allow the modelling of outcomes that the GDG  
 2 considered important (weight changes, hypoglycaemia and treatment dropouts due to  
 3 intolerance). Therefore, original functionality was added to the UKPDS OM1 HbA1c profiling  
 4 (see figure 6). The UKPDS OM1 has annual model cycles; therefore, for HbA1c and weight,  
 5 only treatment effect data at 12 months were used. Only treatments for which data on all 4  
 6 outcomes at the given time-points were available could be included in the health economic  
 7 model. The model was built in Microsoft Excel 2010 (32 bit). Base-case models were run  
 8 separately for each review sub-question and used 50,000 generated people run through  
 9 1000 loops of the UKPDS OM1.

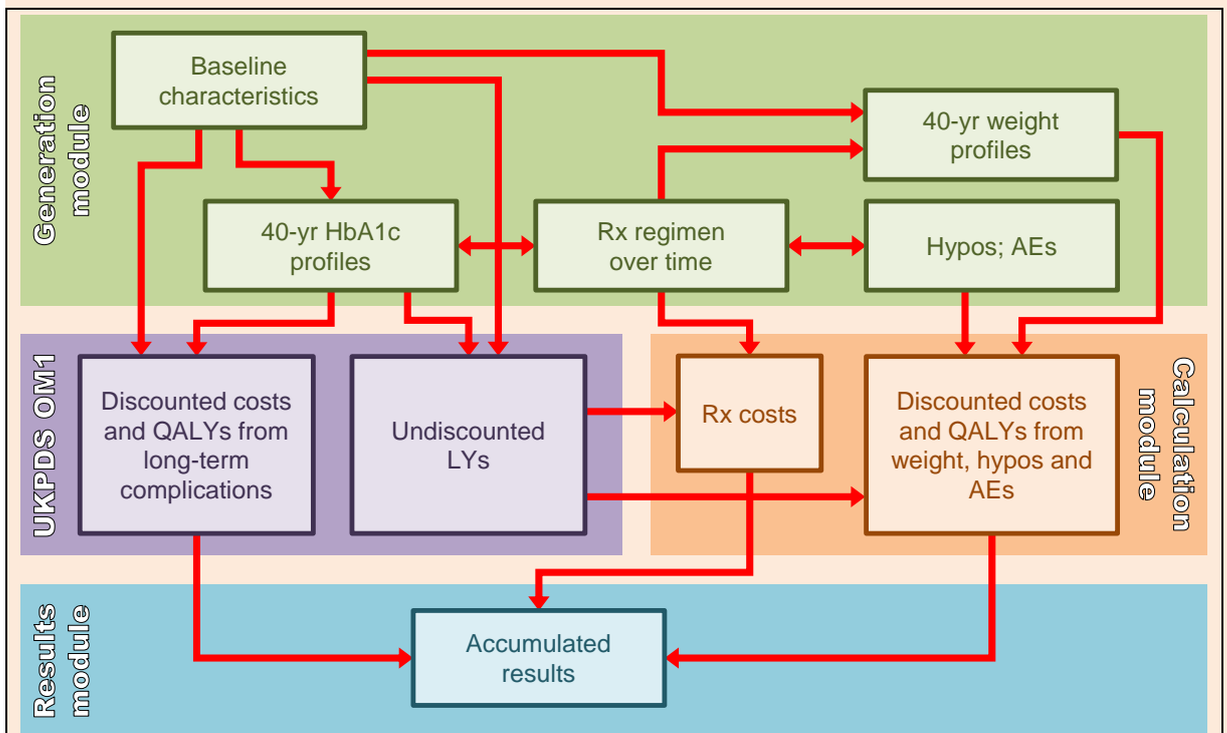
10 Following the initial, 1-year treatment effect, HbA1c was modelled to follow the UKPDS risk  
 11 equations (Clarke et al. 2004). For initial therapy and first intensification, people intensified  
 12 treatment to pre-specified higher therapy levels when their HbA1c rose above 58 mmol/mol  
 13 (7.5%).

14 Treatment dropouts due to intolerance led to pre-specified treatment switches that were  
 15 limited to 2 further treatments within the same level of therapy. The model retained no  
 16 memory of a person's intolerances between therapy levels. When a person was modelled to  
 17 switch treatments, the HbA1c treatment effect for the new treatment was not applied, but the  
 18 treatment effects for weight, hypoglycaemia and dropouts due to intolerance were modelled.

19 Body weight was assumed to increase at a rate of 0.1kg per year for all people. In line with  
 20 the available clinical evidence, treatment-related weight losses were modelled to only last 1  
 21 year, after which the weight loss was regained. However, the GDG advised that treatment-  
 22 related weight gains would remain indefinitely.

23 Rates of all hypoglycaemic episodes were modelled, of which the same proportion (2%,  
 24 Donnelly et al. 2005) were assumed to be severe events. For each therapy level, relative  
 25 treatment effects were taken from the clinical NMA and applied to baseline rates for given  
 26 treatments from epidemiological studies.

27



28 **Figure 6: Schematic representation of original health economic model**

### 8.4.3.31 Baseline data

2 Baseline population characteristics were different for each therapy level (see table 45) and  
3 were largely taken from a large UK database of people registered with GPs (The Health  
4 Improvement Network (THIN) 2014). For the initial therapy analysis, characteristics were  
5 taken from when people were prescribed their first non-insulin anti-diabetes agent (British  
6 National Formulary section 6.1.2, Joint Formulary Committee 2014). Based on an analysis of  
7 diabetes duration in included RCTs, first intensification characteristics were taken when  
8 people had a diabetes duration of 4.5 years and second intensification characteristics when  
9 duration was 8.5 years.

10 To allow baseline population heterogeneity to be accurately modelled through the UKPDS  
11 OM1, individual person data were randomly sampled from a multivariate distribution taking  
12 account of the correlations between variables.

13 Ethnicity is not well recorded at an individual level in the THIN dataset, so this characteristic  
14 was taken from type 2 diabetes respondents in the Health Survey for England (Health and  
15 Social Care Information Centre (HSCIC) 2012). Ethnicity correlation data were taken from  
16 the same data set for the variables that were available. It was not possible to source ethnicity  
17 data specific to each therapy level.

18 In addition to baseline population characteristics, HbA1c profiles were generated for each  
19 person. A floor of 6% was applied to sampling distribution for the generation of baseline  
20 HbA1c.

21 **Table 45: Baseline characteristics used to populate UKPDS OM1**

Category	Characteristic	Initial Therapy	First Intensification	Second Intensification
Demographics	Number of people	90,219	74,144	43,075
	Ethnicity – white	94.6%	94.6%	94.6%
	Ethnicity – African-Caribbean	2.7%	2.7%	2.7%
	Ethnicity – Asian Indian	2.7%	2.7%	2.7%
	Gender (% male)	57.1%	55.9%	55.8%
	Age (years)	59.8	62.7	65.4
	Duration of diabetes (years)	2.0	4.5	8.5
	Weight (kg)	89.9	87.7	86.7
	Height (cm)	168	168	168
Risk Factors at Diagnosis	Atrial fibrillation	0.81%	0.78%	0.63%
	Peripheral vascular disease	0.51%	0.53%	0.47%
	Smoking – current smoker	19.1%	18.0%	19.0%
	Smoking – past smoker	33.2%	33.6%	30.7%
	Total cholesterol (mmol/l)	5.3	5.3	5.5
	HDL (mmol/l)	1.17	1.21	1.21
	Systolic blood pressure (mmHg)	139.6	141.3	143.2
	HbA1c	8.2%	7.8%	7.9%
Current Risk Factors	Smoking – current smoker	18.1%	15.1%	13.4%
	Smoking – past smoker	34.0%	35.8%	36.4%
	Total cholesterol (mmol/l)	5.0	4.5	4.4
	HDL (mmol/l)	1.18	1.23	1.23
	Systolic blood pressure (mmHg)	137.5	136.3	136.2

Category	Characteristic	Initial Therapy	First Intensification	Second Intensification
	HbA1c	8.4%	7.3%	7.6%
Years since pre-existing complications (% of people)	IHD	3.2 (2.7%)	2.8 (5.2%)	5.3 (9.7%)
	CHF	2.5 (0.5%)	2.4 (1.2%)	3.9 (2.3%)
	Amputation	2.0 (0.1%)	2.4 (0.2%)	3.8 (0.4%)
	Blindness	2.3 (0.4%)	2.5 (1.4%)	4.8 (2.2%)
	Renal	3 (0.2%)	2.3 (0.5%)	3.8 (1.0%)
	Stroke	2.7 (0.5%)	2.5 (0.9%)	4.2 (1.8%)
	MI	2.9 (0.8%)	2.6 (1.4%)	4.6 (2.5%)

1 (a) Not all variables are recorded for all people. Therefore, whilst the total number of people in the dataset is  
2 shown, each variable may have a different denominator

3 (b) Ethnicity data source: Health Survey for England 2009-2011

4 (c) THIN data as at 31 August 2013

5 (d) For definitions of variables, see appendix F

#### 8.4.3.46 Resource use and costs

7 NHS inpatient and primary care consultation resource use associated with long-term  
8 complications were modelled by the UKPDS OM1 and costed using the UKPDS costs  
9 (Clarke et al. 2003). Treatment switches due to intolerance were assumed to incur the cost of  
10 1 GP appointment. Severe hypoglycaemic episodes were costed at £380 per episode (ref  
11 Hammer et al. 2009). Weight change incurred no cost.

12 Weighted average doses of the drugs used in the included RCTs were used to calculate the  
13 drug resource use for each arm. Drug unit costs were based on published prices (NHS Drug  
14 Tariff 2014). Consumables and staff time resource used were agreed by the GDG; unit costs  
15 were based on current average usage (Health and Social Care Information Centre 2014) and  
16 published prices (Curtis 2013).

17 All resource use and costs were measured from an NHS and PSS perspective (National  
18 Institute for Health and Care Excellence 2014) and inflated to 2012–13 prices (Curtis 2013).

#### 8.4.3.59 Utilities

20 Baseline utility (0.785) and utility decrements associated with modelled long-term  
21 complications were taken from the UKPDS RCT (Clarke et al. 2002). Treatment switches due  
22 to intolerance assumed an annual utility decrement equivalent to 6 weeks of nausea (-0.005,  
23 Matza et al. 2007). Symptomatic hypoglycaemic episode utility decrements (-0.014) were  
24 modelled on a natural logarithmic scale; severe hypoglycaemic episodes utility decrements (-  
25 0.047) were modelled on a binomial scale. Both were taken from Currie et al. (2006). Utility  
26 decrements associated with weight change (-0.0061 per kg) are applied for BMIs above 27.7  
27 kg/m<sup>2</sup> (Bagust and Beale 2005).

#### 8.4.3.68 Results and sensitivity analyses

29 Results reported were the means of the probabilistic sensitivity analyses. Probabilistic  
30 sensitivity analyses were run for each sub review question, using 1000 iterations of 50,000  
31 people run through 100 UKPDS OM1 loops. One-way sensitivity analyses were run for key  
32 variables and results were based on 50,000 people run through 1000 UKPDS OM1 loops.

#### 8.4.3.73 Model limitations

34 The health economic modelling has addressed a number of limitations of previous analyses,  
35 including the use of detailed, appropriate baseline population data, the use of 12-month  
36 treatment-effect data, a fully incremental analyses of relevant options and the presentation of

1 a thoroughgoing, valid PSA. However, a number of limitations remain. All type 2 diabetes  
2 health economic models rely on extrapolating short-term biological markers to predict long-  
3 term outcomes. Treatment-related weight change and hypoglycaemia effects were key  
4 model drivers that are based on extrapolations of short-term trial-based data; moreover,  
5 these effects are assumed to have quality-of-life impacts that are informed by a small,  
6 methodologically limited evidence-base. In these respects, the analysis presented here is no  
7 more susceptible to bias than any other health economic analysis of its type; however, it is  
8 acknowledged that, if these shortcomings were addressed, this and other analyses might  
9 reach different conclusions.

#### 8.4.40 Clinical evidence review for initial therapy

11 In total 17,037 references were found for the main review question and 122 papers were  
12 included for initial therapy which relate to 114 trials.

13 This sub-review question addressed which initial non-insulin based treatment option is most  
14 effective when people with type 2 diabetes have inadequate blood glucose control. Most  
15 people are at an early stage in diabetes and are generally drug naïve, having been treated  
16 with dietary changes alone.

17 RCTs of at least 12 week treatment duration examining either any oral antidiabetic drug  
18 (OAD) compared to each other or any OAD compared to placebo were included (see section  
19 8.4.2 for main exclusion criteria). As people are more likely to be drug naïve when they start  
20 initial therapy, it was important to ensure included trials used current licensed doses.

21 Therefore, the following additional exclusion criteria were applied:

- 22 • Trials of monotherapy using only doses of blood glucose lowering therapies above the  
23 recommended daily dose.
- 24 • Trials reporting no information relating to doses.
- 25 • Trials termed monotherapy with individuals who were not drug naïve or had washout  
26 periods of less than 4 weeks.

#### 8.4.4.27 Description of included studies for initial therapy

28 A total of 36,938 participants from 114 RCTs were included. The majority of studies were  
29 carried out in multiple centres across different countries. The mean age ranged from 45.6 to  
30 74.4 years, with 6 studies not reporting this information. Mean HbA1c levels at baseline  
31 ranged from 42 to 107 mmol/mol (6% to 11.9%), with 5 studies not reporting this information.  
32 The mean BMI ranged from 23.2 to 39.8 kg/m<sup>2</sup>, with 8 studies not reporting this information.  
33 Mean duration of diabetes ranged from 10.4 weeks to 17.3 years, with 51 studies not  
34 reporting this information. Follow-up periods ranged from 12 to 260 weeks. For full details of  
35 the included studies, see Appendix E.

#### 8.4.4.26 Network meta-analyses for initial therapy

37 To facilitate comparison across all available treatment options, 10 network meta-analyses  
38 were performed for all 3 critical and 1 important outcomes – change in HbA1c at 3, 6, 12 and  
39 24 months, hypoglycaemia at study end point, adverse events (that is, dropouts due to  
40 adverse events, total dropouts and nausea) at study end point and change in body weight at  
41 12 and 24 months. Placebo was selected as the reference treatment as it was the most  
42 common comparator. Full details of methods and additional NMA outputs are provided in  
43 Appendix J.

44 Generally, well-connected networks were produced for shorter follow-up times although  
45 these tended to be sparser and contained fewer treatment options at 12 and 24 months.  
46 Pairwise comparisons that did not form part of the main network were not presented as they  
47 would not add to the GDG decision making.

1 On the whole, the quality of the evidence was moderate to low as networks were generally  
 2 well connected. However, some included trials were not double-blind and did not report  
 3 adequate details of randomisation and allocation concealment methods. It was noted that  
 4 random-effects models tended to estimate a fairly large inter-study heterogeneity term, which  
 5 will reduce the precision of effect estimates.

6 **Table 46: GRADE profile for network meta-analyses for initial therapy**

Assessment time points/ Measure	Number of RCTs	Risk of bias	Inconsistency	Indirectness	Imprecision	Quality
<b>Change in blood glucose (HbA1c)</b>						
3 months	68	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	not serious	Moderate
6 months	62	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	not serious	Moderate
12 months	21	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
24 months	6	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	not serious	Moderate
<b>Hypoglycaemia at study end point</b>						
Study end point	44	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<b>Adverse events at study end point</b>						
Dropouts due to adverse events	73	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
Total dropouts	73	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
Nausea	29	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<b>Change in body weight</b>						
12 months	12	serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low <sup>6</sup>
24 months	6	serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low <sup>6</sup>
<sup>1</sup> Downgrade 1 level: baseline HbA1c ranged from 5.3 to 12.7%						
<sup>2</sup> Assessed based on residual deviance, deviance information criterion and $\tau^2$ ( $\tau^2 < 0.5$ )						
<sup>3</sup> Considered not serious as population, interventions, comparator and outcomes are as defined in protocol						
<sup>4</sup> Downgrade 1 level: no interventions had probability of being best and worse $\geq 0.5$						
<sup>5</sup> Downgrade 1 level: $\tau^2 \geq 0.5$						
<sup>6</sup> Maximum downgrade by 2 levels						

Update 2015

#### 8.4.4.37 Change in blood glucose (HbA1c) at 3, 6, 12 and 24 months

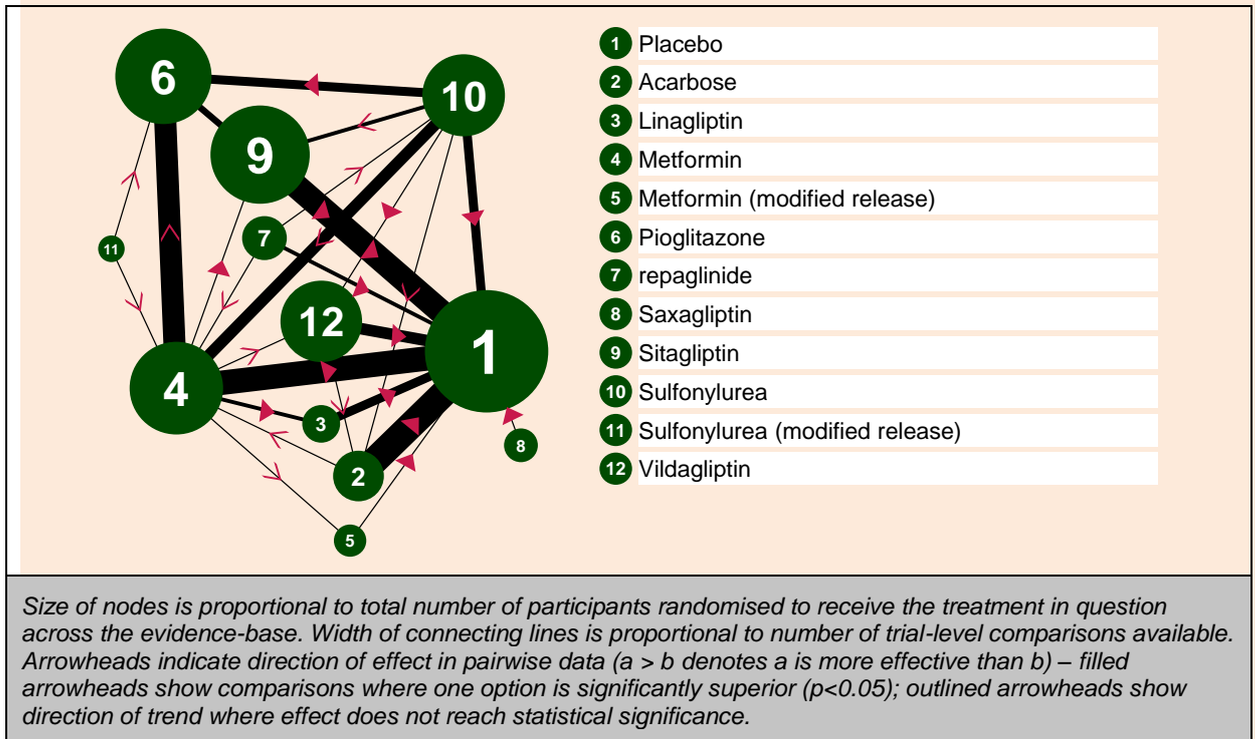
8 Results of the NMAs are summarised below for the 11 treatment options that were compared  
 9 with placebo at 3 and 6 months and the 8 and 6 treatment options assessed at 12 and 24  
 10 months respectively.

11 At the 4 follow-up time points, all treatments are shown to be consistently more effective than  
 12 placebo, though with varying levels of precision. At longer follow-up periods, the 95%  
 13 credible intervals are generally wider, and at 12 months in particular, they tend to overlap  
 14 making it difficult to distinguish between the various treatments. Where available, there is  
 15 reasonable agreement between the NMA evidence and direct pairwise treatment effect  
 16 estimates which compared different options with placebo in the underlying evidence. The  
 17 inclusion of indirect evidence alongside direct evidence slightly reduces uncertainty, and also  
 18 results in some small changes in effect estimates. However, there is substantial overlap  
 19 between the 95% credible/confidence intervals, suggesting reasonable consistency between  
 20 direct and indirect evidence.

21 The rankings of each treatment option, summarised in the tables similarly support the  
 22 conclusion that the option that is least likely to be effective is placebo. At 3 and 6 months,  
 23 repaglinide and sulfonylurea demonstrated consistently high rankings with narrow credible

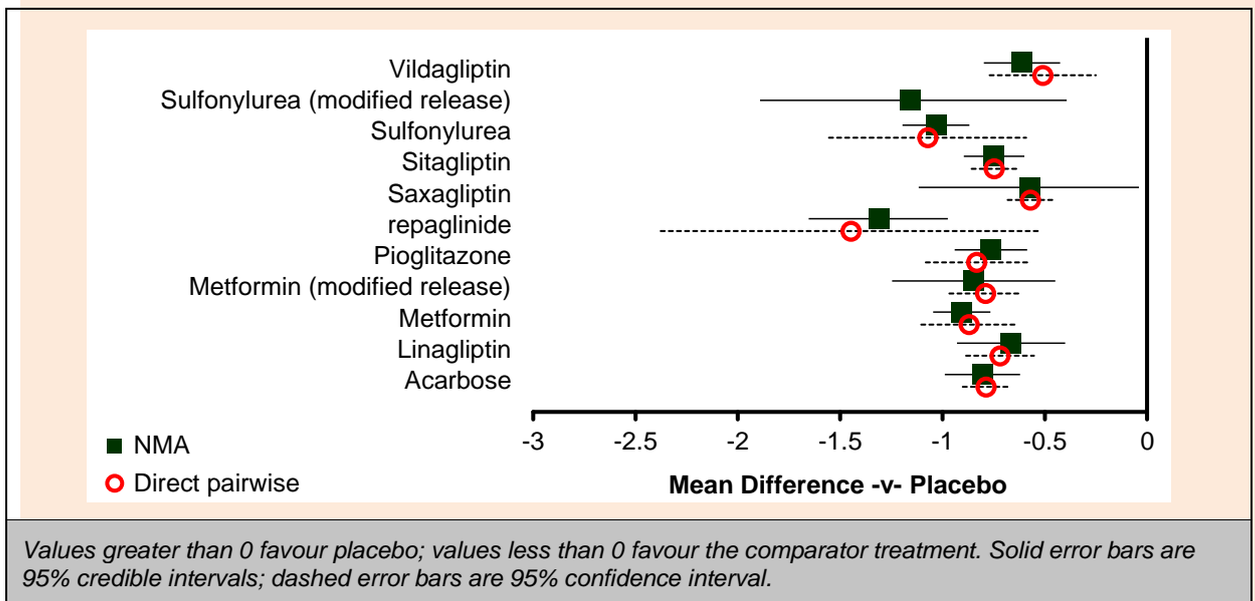
1 intervals. Repaglinide also had the highest ranking at 12 months, though with a lower  
2 probability and wider credible intervals. The option with the highest individual probability of  
3 maximum effectiveness is pioglitazone at 24 months.

4



5 **Figure 7: Network meta-analysis of change in HbA1c (3 months) – evidence network**

6



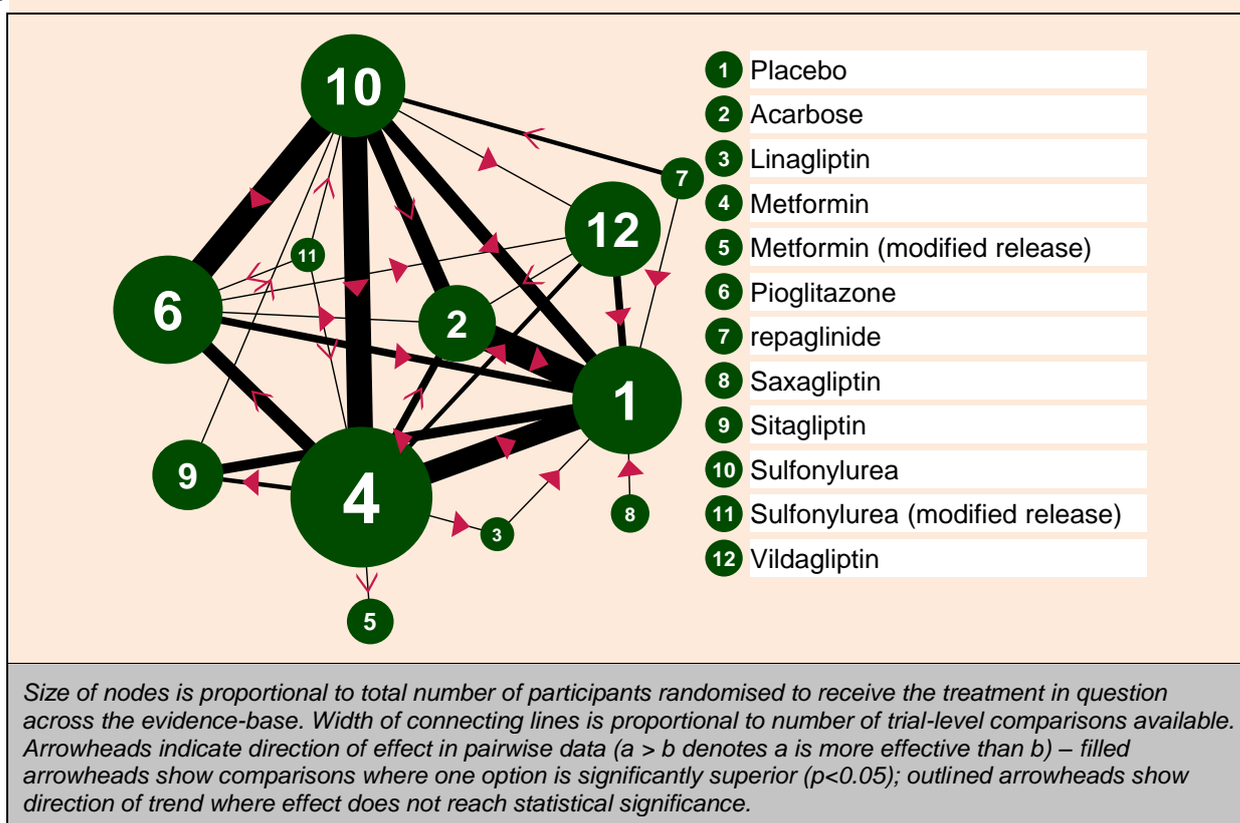
7 **Figure 8: Network meta-analysis of change in HbA1c (3 months) – relative effect of all**  
8 **options compared with common comparator (placebo)**

1  
2  
3

**Table 47: Network meta-analysis of change in HbA1c (3 months) – rankings for each comparator**

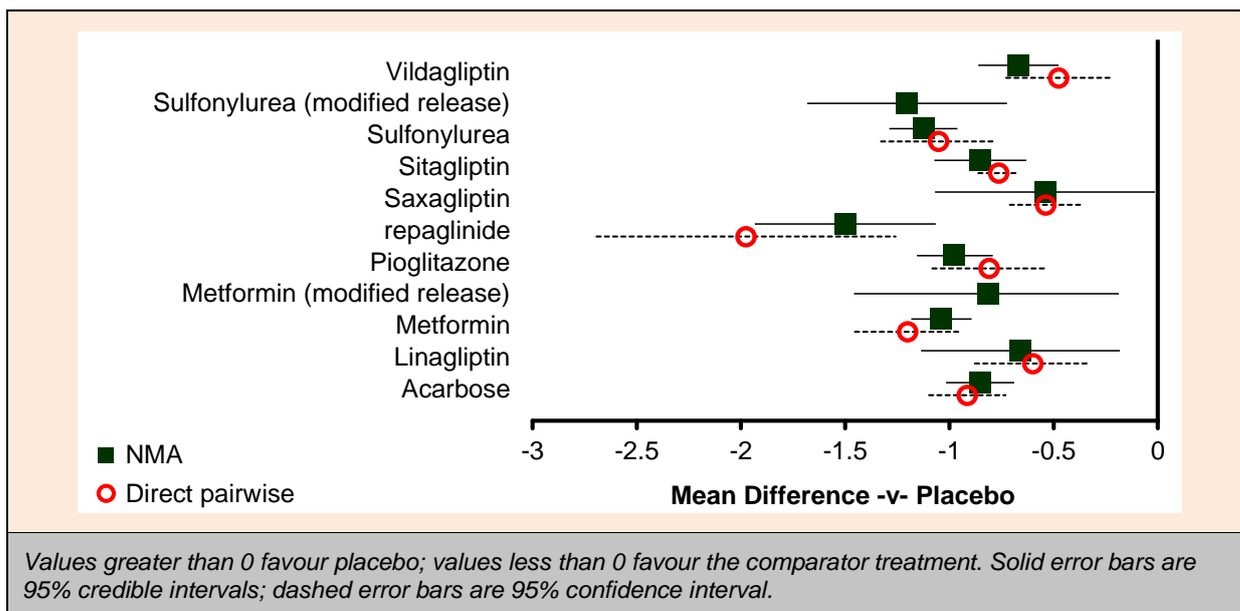
	Probability best	Median rank (95%CrI)
Placebo	0.000	12 (12, 12)
Acarbose	0.000	6 (3, 10)
Linagliptin	0.000	9 (4, 11)
Metformin	0.000	4 (3, 7)
Metformin (modified release)	0.020	5 (2, 11)
Pioglitazone	0.000	7 (4, 10)
repaglinide	0.611	1 (1, 3)
Saxagliptin	0.005	10 (2, 11)
Sitagliptin	0.000	7 (4, 10)
Sulfonylurea	0.020	3 (2, 5)
Sulfonylurea (modified release)	0.344	2 (1, 11)
Vildagliptin	0.000	10 (6, 11)

4



**Figure 9: Network meta-analysis of change in HbA1c (6 months) – evidence network**

5  
6

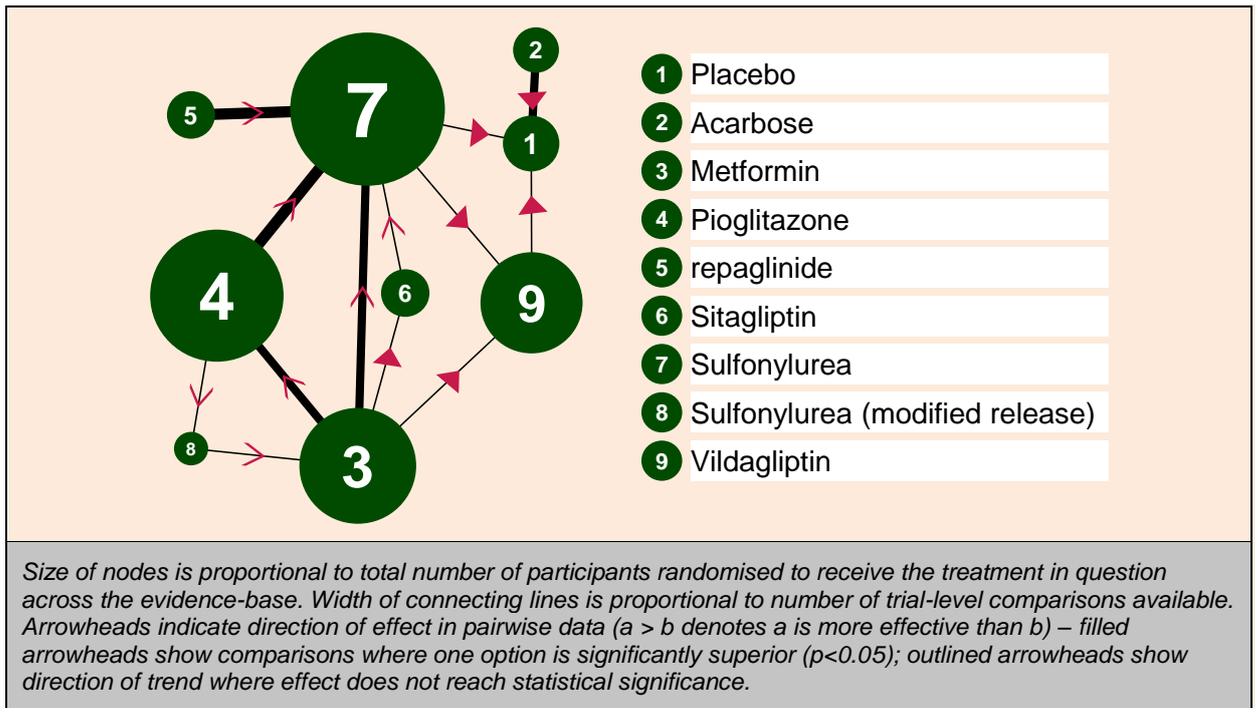


1 **Figure 10: Network meta-analysis of change in HbA1c (6 months) – relative effect of**  
 2 **all options compared with common comparator (placebo)**

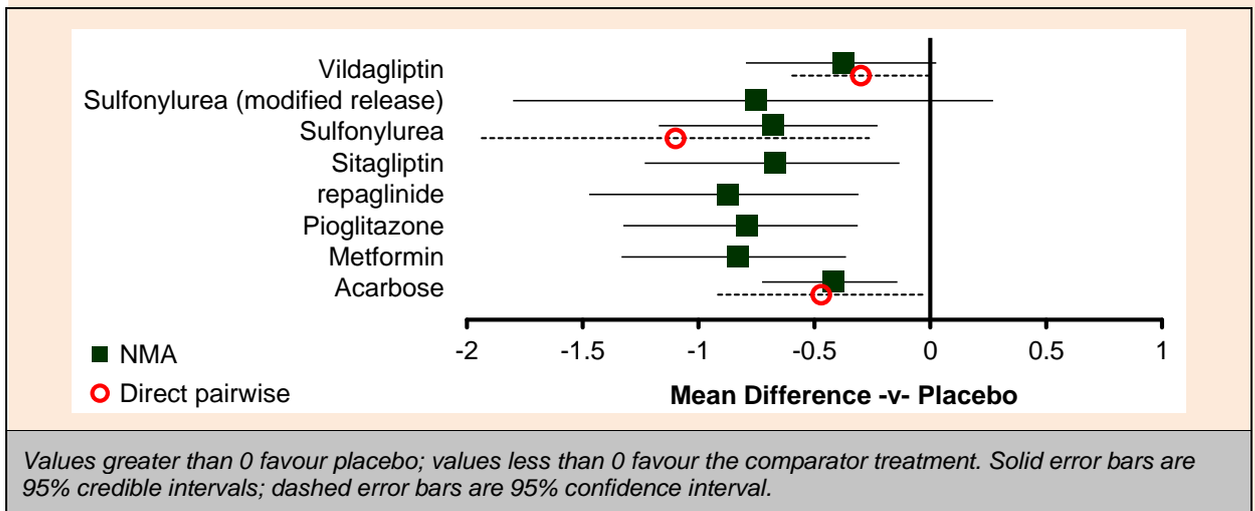
4 **Table 48: Network meta-analysis of change in HbA1c (6 months) – rankings for each**  
 5 **comparator**

	Probability best	Median rank (95%CrI)
Placebo	0.000	12 (11, 12)
Acarbose	0.000	7 (5, 9)
Linagliptin	0.002	9 (3, 11)
Metformin	0.002	4 (2, 6)
Metformin (modified release)	0.027	8 (1, 11)
Pioglitazone	0.000	5 (3, 8)
repaglinide	0.797	1 (1, 3)
Saxagliptin	0.002	10 (4, 11)
Sitagliptin	0.000	7 (4, 10)
Sulfonylurea	0.010	3 (2, 5)
Sulfonylurea (modified release)	0.160	2 (1, 8)
Vildagliptin	0.000	9 (7, 11)

6



1 **Figure 11: Network meta-analysis of change in HbA1c (12 months) – evidence**  
 2 **network**



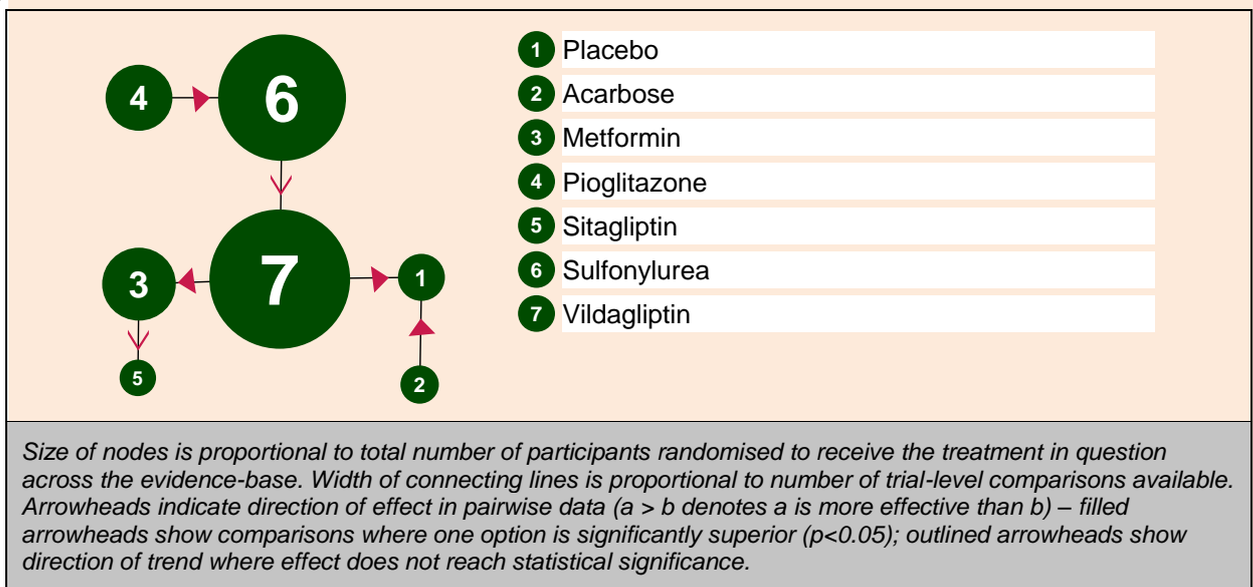
4 **Figure 12: Network meta-analysis of change in HbA1c (12 months) – relative effect**  
 5 **of all options compared with common comparator (placebo)**

Update 2015

1 **Table 49: Network meta-analysis of change in HbA1c (12 months) – rankings for each**  
2 **comparator**

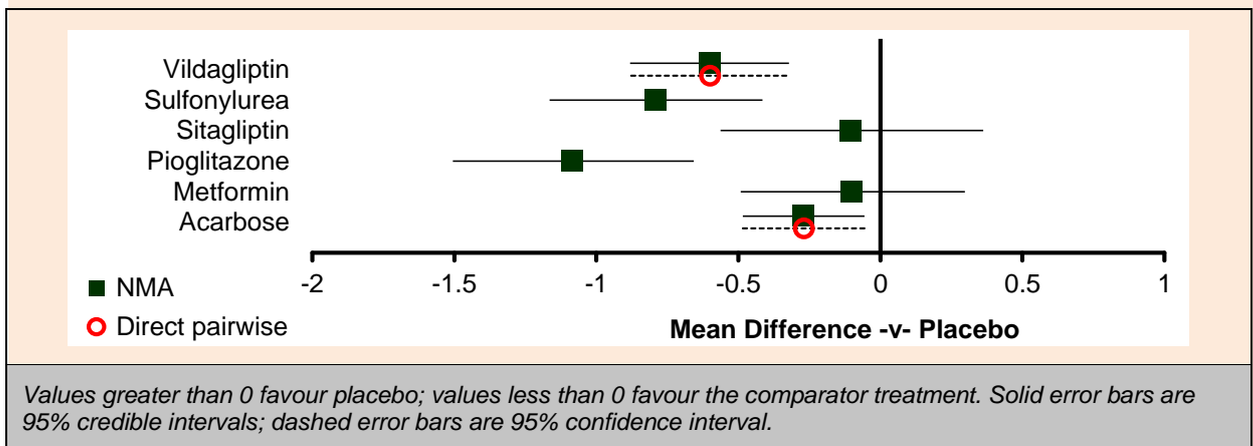
	Probability best	Median rank (95%CrI)
Placebo	0.000	9 (8, 9)
Acarbose	0.023	7 (2, 8)
Metformin	0.159	3 (1, 5)
Pioglitazone	0.086	3 (1, 6)
repaglinide	0.364	2 (1, 6)
Sitagliptin	0.030	5 (1, 8)
Sulfonylurea	0.002	5 (3, 7)
Sulfonylurea (modified release)	0.335	4 (1, 9)
Vildagliptin	0.001	7 (5, 8)

3



4 **Figure 13: Network meta-analysis of change in HbA1c (24 months) – evidence**  
5 **network**

6



7 **Figure 14: Network meta-analysis of change in HbA1c (24 months) – relative effect**  
8 **of all options compared with common comparator (placebo)**

9

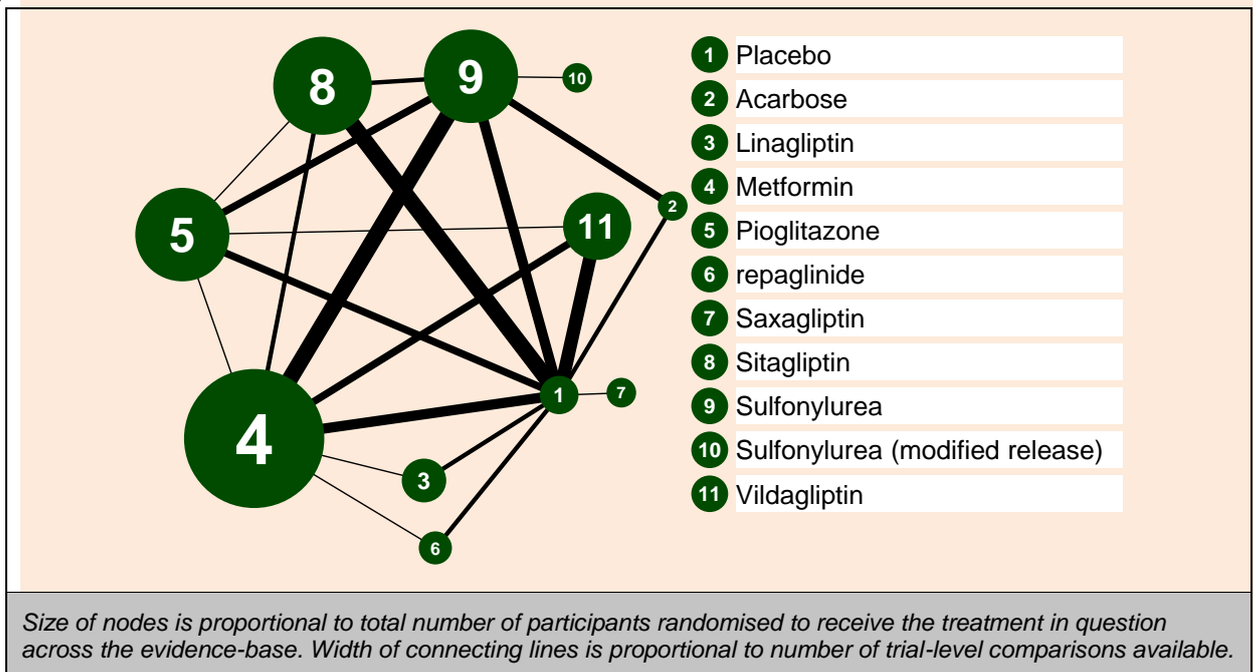
1 **Table 50: Network meta-analysis of change in HbA1c (24 months) – rankings for each**  
2 **comparator**

	Probability best	Median rank (95%CrI)
Placebo	0.000	7 (5, 7)
Acarbose	0.000	4 (3, 6)
Metformin	0.000	6 (4, 7)
Pioglitazone	0.996	1 (1, 1)
Sitagliptin	0.000	6 (4, 7)
Sulfonylurea	0.002	2 (2, 3)
Vildagliptin	0.001	3 (2, 4)

#### 8.4.4.43 Hypoglycaemia at study end point

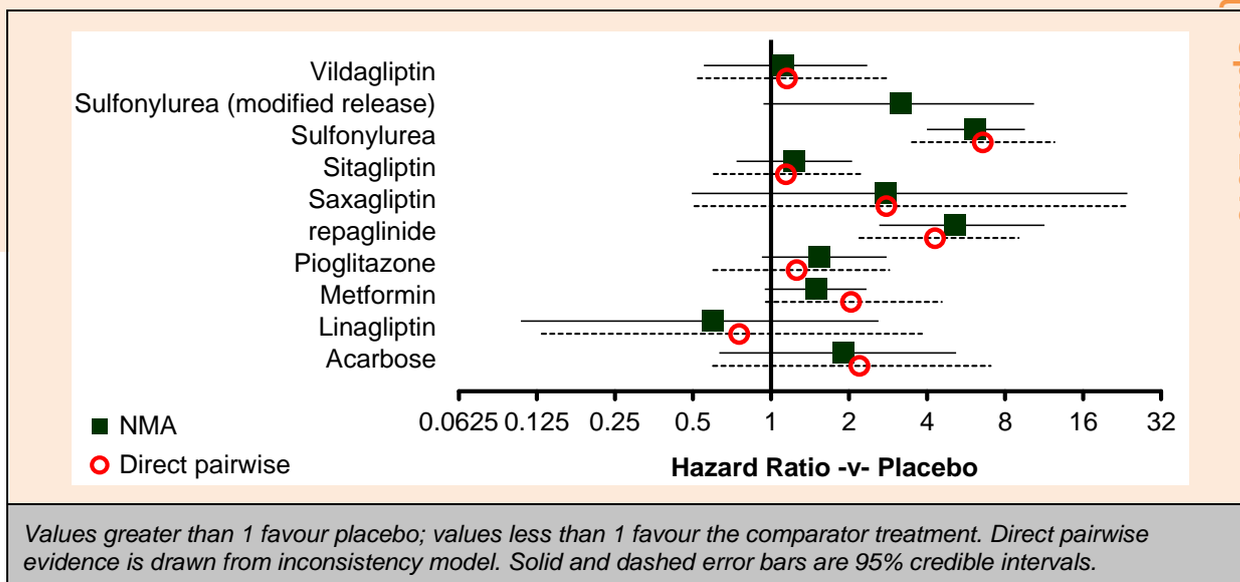
4 Results of the NMA are summarised below for the 10 treatment options that were compared  
5 with placebo. There is reasonable agreement between the NMA evidence and direct pairwise  
6 treatment effect estimates as demonstrated by the substantial overlap between the  
7 credible/confidence intervals. In general, there was lower incidence of hypoglycaemic events  
8 in the placebo group compared to the active interventions. While linagliptin had the highest  
9 individual probability of maximum effectiveness, it was associated with wide credible intervals  
10 (ranging from 1 to 8), indicating that this treatment option could credibly be ranked as low as  
11 8<sup>th</sup>.

12



13 **Figure 15: Network meta-analysis of hypoglycaemic events (study end point) –**  
14 **evidence network**

15



1 **Figure 16: Network meta-analysis of hypoglycaemic events (study end point) –**  
 2 **relative effect of all options compared with common comparator (placebo)**

4 **Table 51: Network meta-analysis of hypoglycaemic events (study end point) – rankings**  
 5 **for each comparator**

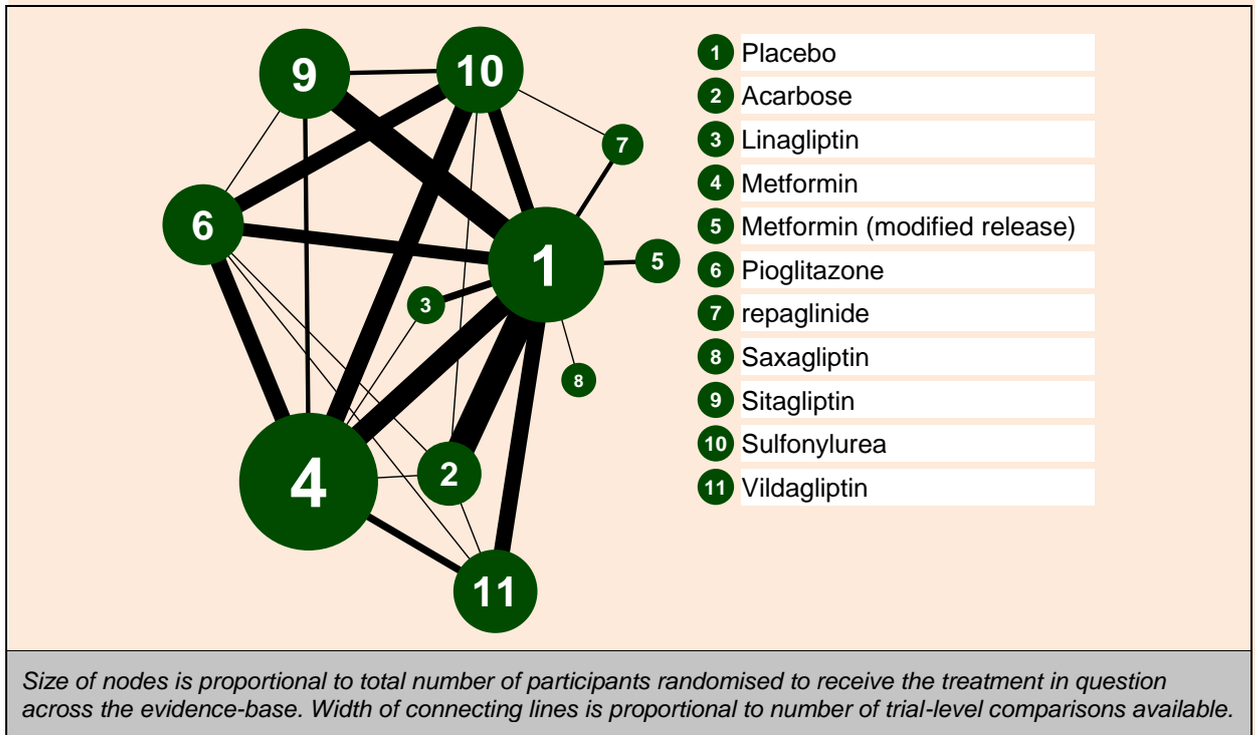
	Probability best	Median rank (95%CrI)
Placebo	0.100	3 (1, 5)
Acarbose	0.032	7 (1, 10)
Linagliptin	0.663	1 (1, 8)
Metformin	0.002	6 (3, 8)
Pioglitazone	0.005	6 (2, 8)
repaglinide	0.000	10 (8, 11)
Saxagliptin	0.052	8 (1, 11)
Sitagliptin	0.033	4 (1, 7)
Sulfonylurea	0.000	10 (9, 11)
Sulfonylurea (modified release)	0.008	8 (3, 11)
Vildagliptin	0.106	3 (1, 8)

#### 8.4.4.56 Adverse events at study end point

- 7 Results of the 3 NMAs are summarised below. For dropouts due to adverse events and total  
 8 dropouts, 10 treatment options were compared with placebo, while 8 treatment options were  
 9 compared with placebo for nausea.
- 10 There is moderate agreement between the NMA evidence and direct pairwise treatment  
 11 effect estimates. Whilst there is substantial overlap between the credible/confidence intervals  
 12 suggesting reasonable consistency between the direct and indirect evidence, for nausea in  
 13 particular, the point estimates for the direct and indirect evidence were in opposite directions  
 14 for pioglitazone.
- 15 In general, active treatment options were effective at preventing total dropouts. However,  
 16 active treatment options in the main were associated with higher dropouts due to adverse  
 17 events and nausea when compared to placebo. Repaglinide and sulfonylurea (modified  
 18 release) were associated with the highest probability of maximum effectiveness and highest

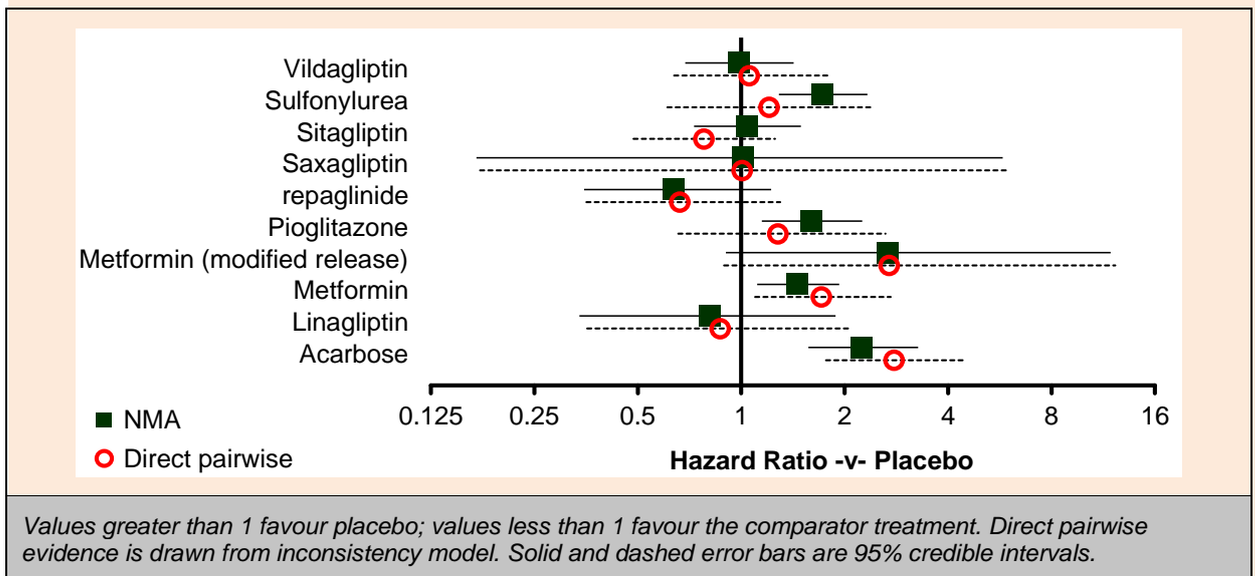
1 median ranks for dropouts due to adverse events and total dropouts respectively, but these  
2 rankings were associated with wide credible intervals (1 to 6 and 1 to 11 respectively).  
3 Similarly, placebo was associated with lower incidence of nausea when compared to active  
4 treatment options.

5



6 **Figure 17: Network meta-analysis of dropouts due to adverse events (study end**  
7 **point) – evidence network**

8



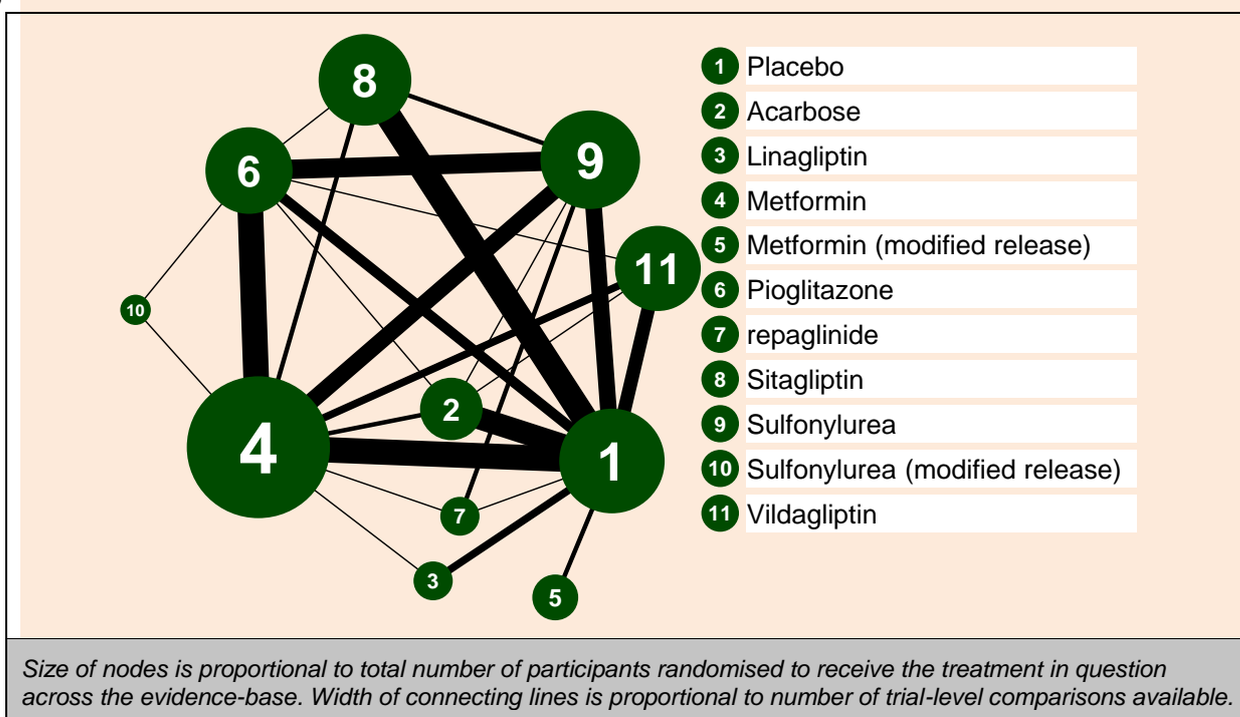
9 **Figure 18: Network meta-analysis of dropouts due to adverse events (study end**  
10 **point) – relative effect of all options compared with common comparator**  
11 **(placebo)**

12

1 **Table 52: Network meta-analysis of dropouts due to adverse events (study end point) –**  
2 **rankings for each comparator**

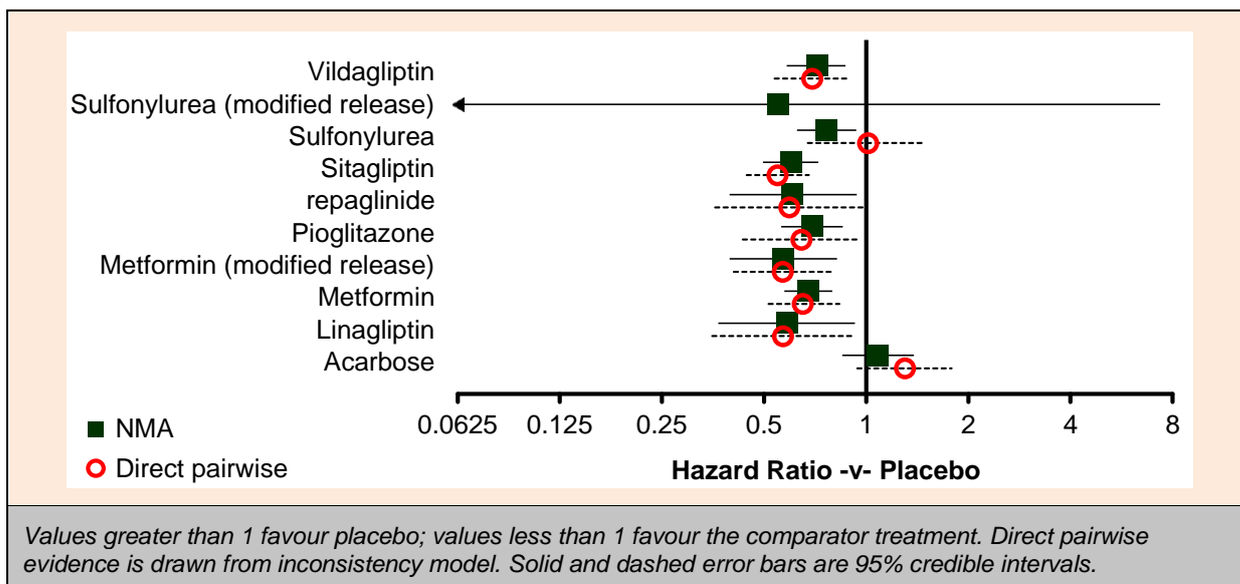
	Probability best	Median rank (95%CrI)
Placebo	0.004	4 (2, 6)
Acarbose	0.000	10 (8, 11)
Linagliptin	0.231	3 (1, 9)
Metformin	0.000	7 (5, 9)
Metformin (modified release)	0.003	11 (3, 11)
Pioglitazone	0.000	8 (6, 10)
repaglinide	0.462	2 (1, 6)
Saxagliptin	0.261	4 (1, 11)
Sitagliptin	0.013	5 (2, 7)
Sulfonylurea	0.000	9 (7, 10)
Vildagliptin	0.026	4 (1, 7)

3



4 **Figure 19: Network meta-analysis of total dropouts (study end point) – evidence**  
5 **network**

6

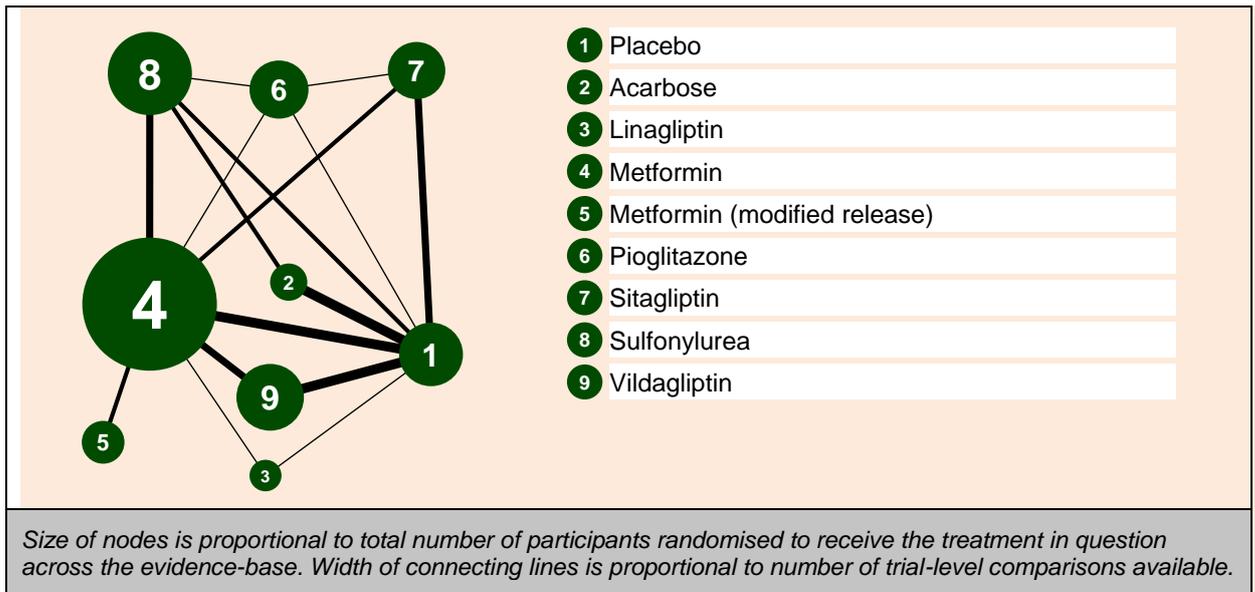


1 **Figure 20: Network meta-analysis of total dropouts (study end point) – relative**  
 2 **effect of all options compared with common comparator (placebo)**

4 **Table 53: Network meta-analysis of total dropouts (study end point) – rankings for**  
 5 **each comparator**

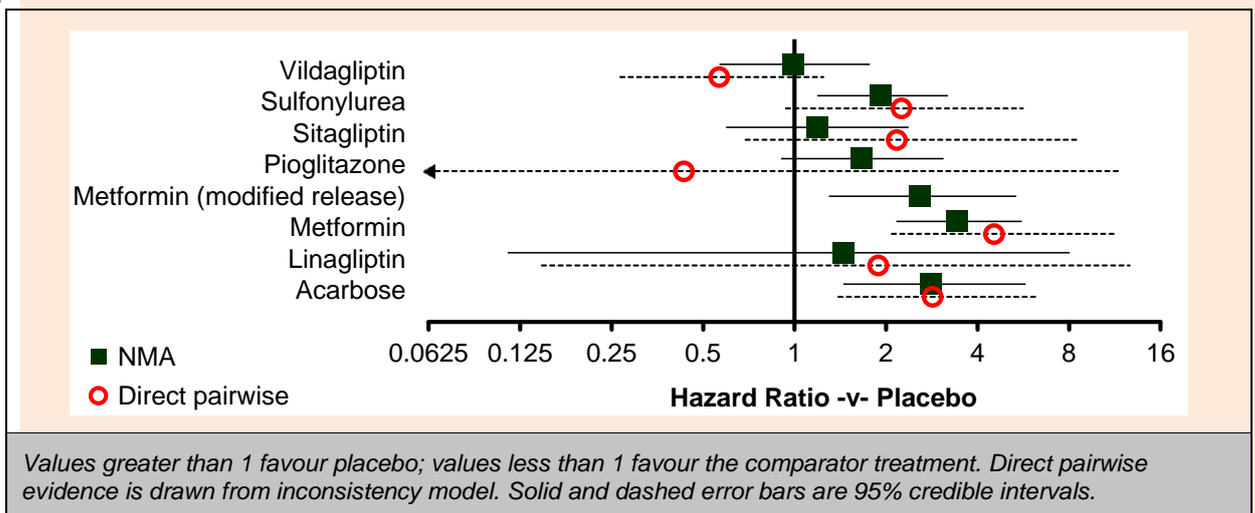
	Probability best	Median rank (95%CrI)
Placebo	0.000	10 (9, 11)
Acarbose	0.000	11 (9, 11)
Linagliptin	0.167	3 (1, 9)
Metformin	0.003	5 (3, 8)
Metformin (modified release)	0.173	3 (1, 9)
Pioglitazone	0.004	6 (2, 9)
repaglinide	0.125	4 (1, 9)
Sitagliptin	0.061	3 (1, 7)
Sulfonylurea	0.000	8 (5, 9)
Sulfonylurea (modified release)	0.465	2 (1, 11)
Vildagliptin	0.003	7 (3, 9)

6



1 **Figure 21: Network meta-analysis of nausea (study end point) – evidence network**

2



3 **Figure 22: Network meta-analysis of nausea (study end point) – relative effect of all options compared with common comparator (placebo)**

4

5

1 **Table 54: Network meta-analysis of nausea (study end point) – rankings for each**  
2 **comparator**

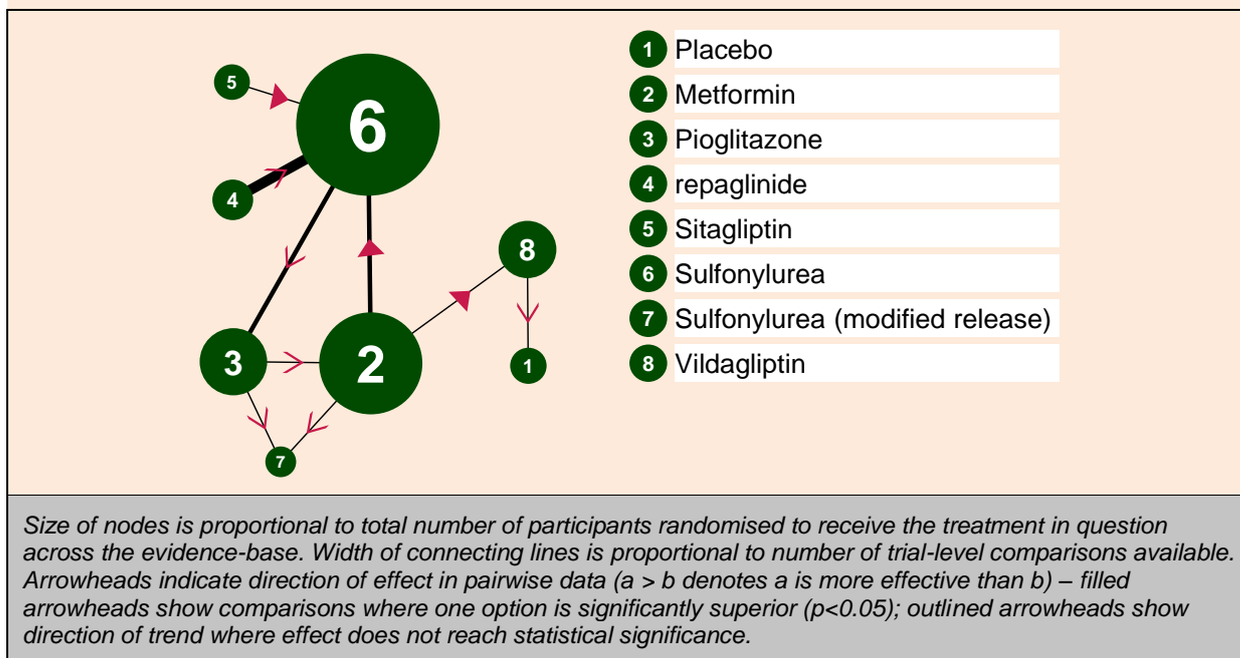
	Probability best	Median rank (95%CrI)
Placebo	0.256	2 (1, 4)
Acarbose	0.000	8 (4, 9)
Linagliptin	0.306	4 (1, 9)
Metformin	0.000	8 (7, 9)
Metformin (modified release)	0.000	7 (4, 9)
Pioglitazone	0.006	5 (2, 7)
Sitagliptin	0.139	3 (1, 6)
Sulfonylurea	0.000	6 (4, 7)
Vildagliptin	0.292	2 (1, 4)

3

#### 8.4.4.64 Change in body weight at 12 and 24 months

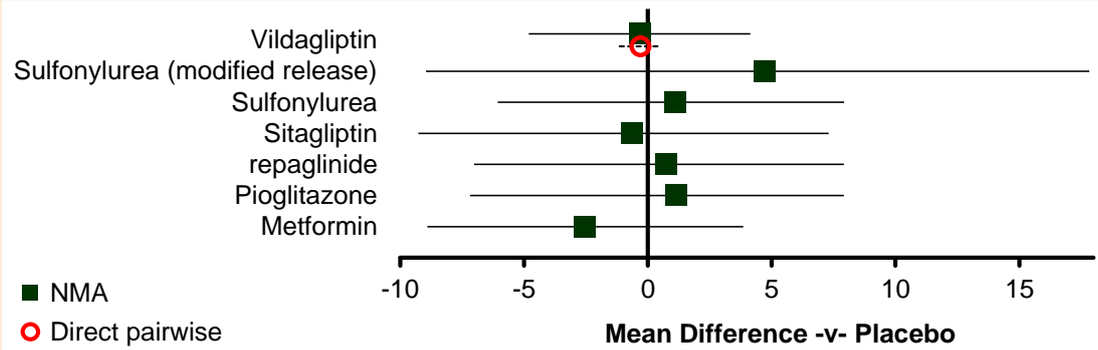
5 Results of the 2 NMAs are summarised below for the 7 and 5 treatment options that were  
6 compared with placebo at 12 and 24 months respectively. Where available, there was  
7 reasonable agreement in the indirect evidence and direct pairwise treatment effect estimates,  
8 with substantial overlap between the credible/confidence intervals. In general, metformin and  
9 sitagliptin (at 24 months only) were shown to be most effective at weight loss compared to  
10 placebo. However, the credible intervals associated with these relative effects were  
11 considerably wide. Metformin had the highest individual probability of maximum effectiveness  
12 and highest ranking at 12 and 24 months, with consistently narrow credible intervals  
13 surrounding the rankings (1 to 3 and 1 to 2 respectively).

14



15 **Figure 23: Network meta-analysis of change in body weight (12 months) – evidence**  
16 **network**

17

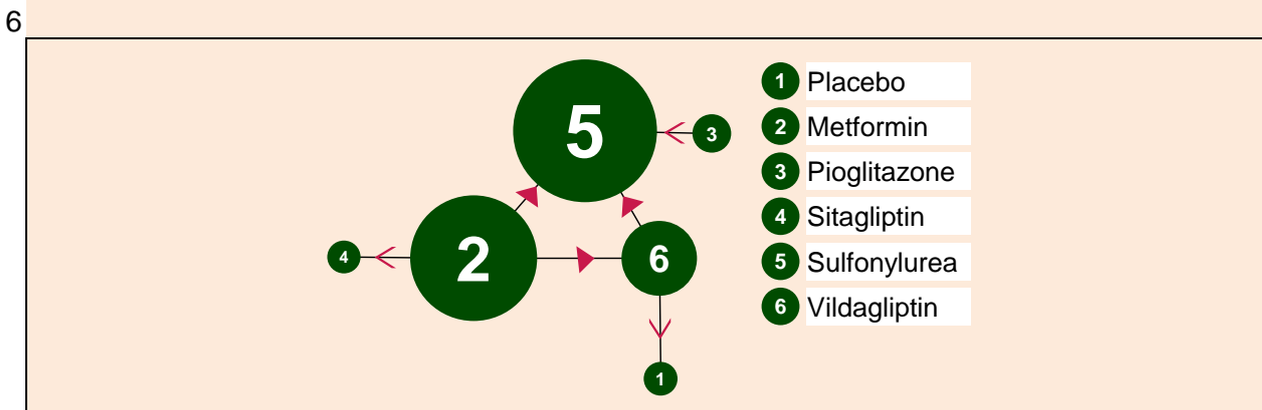


Values greater than 0 favour placebo; values less than 0 favour the comparator treatment. Solid error bars are 95% credible intervals; dashed error bars are 95% confidence interval.

1 **Figure 24: Network meta-analysis of change in body weight (12 months) – relative**  
 2 **effect of all options compared with common comparator (placebo)**

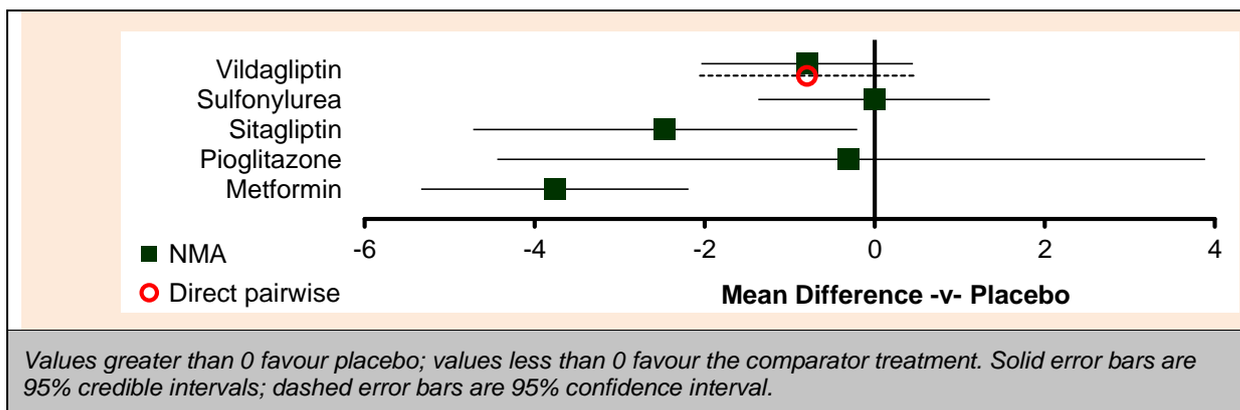
4 **Table 55: Network meta-analysis of change in body weight (12 months) – rankings for**  
 5 **each comparator**

	Probability best	Median rank (95%CrI)
Placebo	0.103	4 (1, 8)
Metformin	0.576	1 (1, 4)
Pioglitazone	0.018	6 (2, 8)
repaglinide	0.018	5 (2, 8)
Sitagliptin	0.141	3 (1, 8)
Sulfonylurea	0.001	6 (3, 8)
Sulfonylurea (modified release)	0.101	8 (1, 8)
Vildagliptin	0.043	4 (1, 8)



Size of nodes is proportional to total number of participants randomised to receive the treatment in question across the evidence-base. Width of connecting lines is proportional to number of trial-level comparisons available. Arrowheads indicate direction of effect in pairwise data ( $a > b$  denotes  $a$  is more effective than  $b$ ) – filled arrowheads show comparisons where one option is significantly superior ( $p < 0.05$ ); outlined arrowheads show direction of trend where effect does not reach statistical significance.

7 **Figure 25: Network meta-analysis of change in body weight (24 months) – evidence**  
 8 **network**



1 **Figure 26: Network meta-analysis of change in body weight (24 months) – relative**  
 2 **effect of all options compared with common comparator (placebo)**

4 **Table 56: Network meta-analysis of change in body weight (24 months) – rankings for**  
 5 **each comparator**

	Probability best	Median rank (95%CrI)
Placebo	0.000	5 (3, 6)
Metformin	0.893	1 (1, 2)
Pioglitazone	0.047	4 (1, 6)
Sitagliptin	0.060	2 (1, 4)
Sulfonylurea	0.000	5 (4, 6)
Vildagliptin	0.000	3 (3, 5)

#### 8.4.56 Health economic evidence for initial therapy

7 For initial therapy, no CUAs met the UK inclusion criteria and only 2 studies were found  
 8 worldwide. Therefore, an original economic analysis was undertaken.

9 For initial therapy, 7 treatments could be modelled. People accrued an average of 18.3  
 10 undiscounted life years, of which 3.4 years were spent on initial therapy and 3.1 were spent  
 11 on first intensification therapy. There was little difference in lifetime complication rates, due to  
 12 small differences in HbA1c treatment effects and the normalising effects of treatment  
 13 intensification.

14 People accumulated an average of 9.0 lifetime discounted QALYs, with most loss coming  
 15 from weight profiles and differences driven by weight treatment effects. Treatment-related  
 16 costs accounted for most variation in lifetime discounted costs.

17 Initial therapy with metformin incurred the lowest lifetime discounted costs and gained most  
 18 lifetime discounted QALYs and therefore metformin dominated all other treatment options  
 19 (see table 57).

1 **Table 57: Mean lifetime incremental cost–utility results for initial therapy**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Metformin -> Met-SU -> Met-I(NPH)	£19,250	9.033			
Repaglinide -> Met-SU -> Met-I(NPH)	£19,298	8.974	£48	-0.059	Dominated
Pioglitazone -> Met-SU -> Met-I(NPH)	£19,412	8.973	£163	-0.060	Dominated
Sulfonylurea -> Met-SU -> Met-I(NPH)	£19,580	8.950	£330	-0.082	Dominated
Placebo -> Met-SU -> Met-I(NPH)	£20,043	8.912	£794	-0.121	Dominated
Sitagliptin -> Met-SU -> Met-I(NPH)	£20,457	8.990	£1207	-0.043	Dominated
Vildagliptin -> Met-SU -> Met-I(NPH)	£20,627	8.954	£1377	-0.078	Dominated

2 (a) Met-SU = Metformin-Sulfonylurea

3 (b) Met-I(NPH) = Metformin-NPH insulin

4 For people who could not tolerate metformin, repaglinide was the most cost-effective  
5 treatment option (see table 58). If people were unwilling to take repaglinide at initial therapy  
6 (as it would require switching to 2 different drugs at first intensification), pioglitazone was the  
7 cheapest treatment option; sitagliptin had an ICER of £62,500 per QALY compared with  
8 pioglitazone (see table 59).

9 If people could not tolerate metformin, could not tolerate or did not wish to take repaglinide  
10 and were contraindicated for pioglitazone, sulfonylurea was the cheapest treatment option  
11 and sitagliptin had an ICER of £22,300 per QALY (see Table 60).

12 **Table 58: Mean lifetime incremental cost–utility results for initial therapy when**  
13 **metformin is not a treatment option**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Repaglinide -> Met-SU -> Met-I(NPH)	£19,298	8.974			
Pioglitazone -> Met-SU -> Met-I(NPH)	£19,412	8.973	£115	-0.001	Dominated
Sulfonylurea -> Met-SU -> Met-I(NPH)	£19,580	8.950	£282	-0.024	Dominated
Placebo -> Met-SU -> Met-I(NPH)	£20,043	8.912	£746	-0.062	Dominated
Sitagliptin -> Met-SU -> Met-I(NPH)	£20,457	8.990	£1159	0.016	£73,287
Vildagliptin -> Met-SU -> Met-I(NPH)	£20,627	8.954	£170	-0.035	Dominated

14 (a) Met-SU = Metformin-Sulfonylurea

15 (b) Met-I(NPH) = Metformin-NPH insulin

16 **Table 59: Mean lifetime incremental cost–utility results for initial therapy when**  
17 **metformin and repaglinide are not treatment options**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Pioglitazone -> Met-SU -> Met-I(NPH)	£19,412	8.973			
Sulfonylurea -> Met-SU -> Met-I(NPH)	£19,580	8.950	£167	-0.023	Dominated
Placebo -> Met-SU -> Met-I(NPH)	£20,043	8.912	£631	-0.061	Dominated
Sitagliptin -> Met-SU -> Met-I(NPH)	£20,457	8.990	£1044	0.017	£62,473
Vildagliptin -> Met-SU -> Met-I(NPH)	£20,627	8.954	£170	-0.035	Dominated

18 (a) Met-SU = Metformin-Sulfonylurea

19 (b) Met-I(NPH) = Metformin-NPH insulin

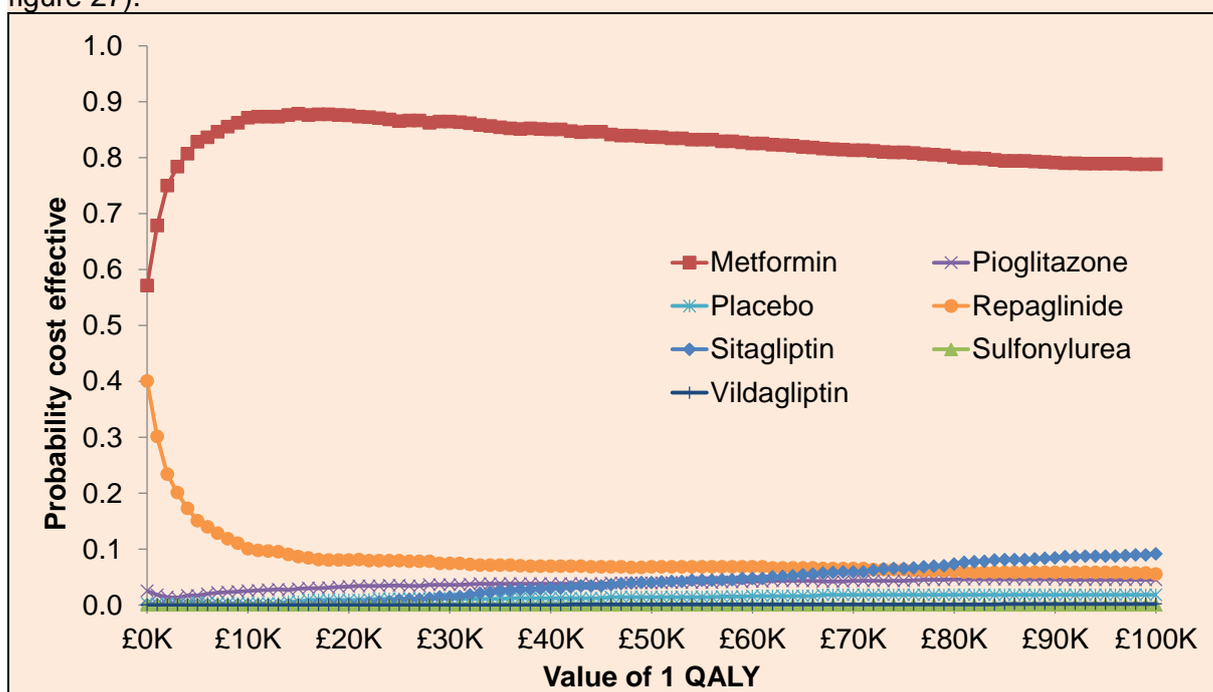
1 **Table 60: Mean lifetime incremental cost–utility results for initial therapy when neither**  
2 **metformin, repaglinide nor pioglitazone are treatment options**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Sulfonylurea -> Met-SU -> Met-I(NPH)	£19,580	8.950			
Placebo -> Met-SU -> Met-I(NPH)	£20,043	8.912	£464	-0.039	Dominated
Sitagliptin -> Met-SU -> Met-I(NPH)	£20,457	8.990	£877	0.039	£22,256
Vildagliptin -> Met-SU -> Met-I(NPH)	£20,627	8.954	£170	-0.035	Dominated

3 (a) *Met-SU = Metformin-Sulfonylurea*

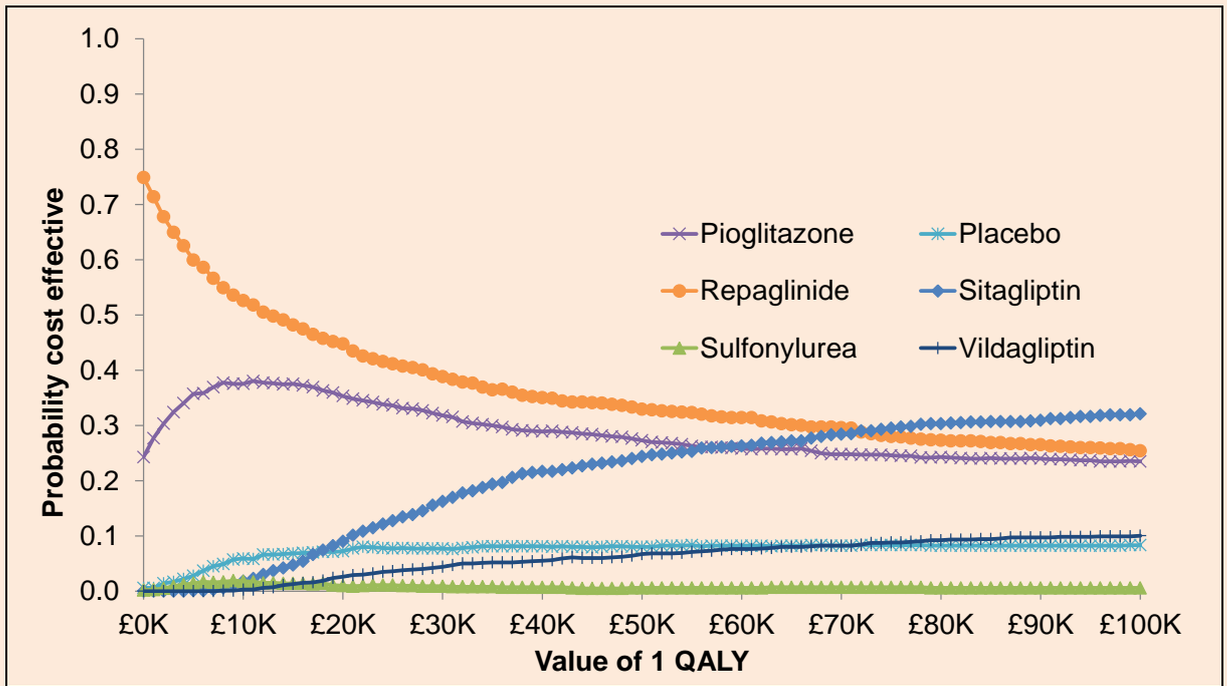
4 (b) *Met-I(NPH) = Metformin-NPH insulin*

5 Over 1000 PSA iterations, metformin was the most cost effective of the initial therapy  
6 treatments in 88% of iterations at a maximum acceptable ICER of £20,000 per QALY (see  
7 figure 27).



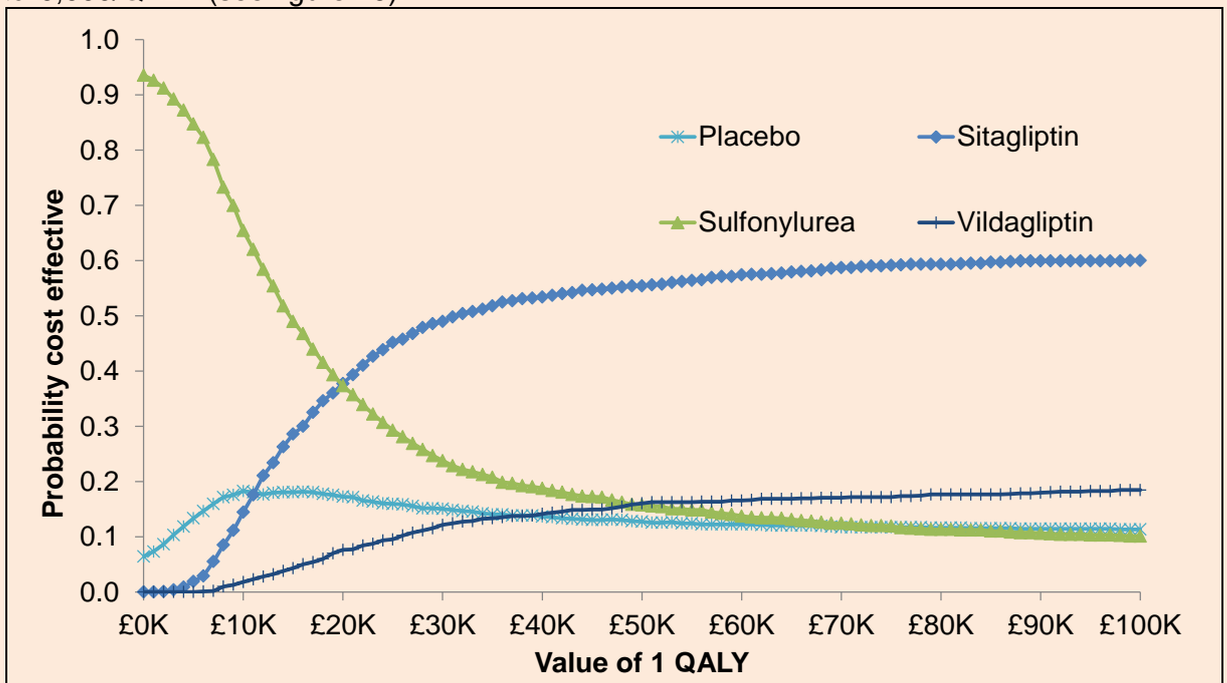
8 **Figure 27: Cost-effectiveness acceptability curve for initial therapy**

9 For people who could not tolerate metformin, repaglinide was the most cost-effective initial  
10 therapy at a maximum acceptable ICER of £20,000 per QALY in 45% of iterations, with  
11 pioglitazone the most cost-effective initial therapy in 35% of iterations (see figure 28).



1 **Figure 28: Cost-effectiveness acceptability curve for initial therapy when metformin is not a**  
2 **treatment option**

3 For people who could not tolerate metformin, could not tolerate or choose not to initiate  
4 therapy with repaglinide and were contraindicated for pioglitazone, sitagliptin (most cost-  
5 effective in 38% of iterations) and sulfonyleurea (most cost-effective in 37% of iterations) were  
6 the most cost-effective initial therapy treatment options at a maximum acceptable ICER of  
7 £20,000/QALY (see figure 29).



8 **Figure 29: Cost-effectiveness acceptability curve for initial therapy when neither metformin,**  
9 **repaglinide nor pioglitazone are treatment options**

## 8.4.6.1 Evidence statements for initial therapy

### 8.4.6.1.2 Clinical evidence

#### 8.4.6.1.1.3 Change in blood glucose

4 Evidence from 4 network meta-analyses including data from 68, 62, 21 and 6 RCTs at 3, 6,  
5 12 and 24 months respectively for HbA1c levels showed that repaglinide was consistently  
6 associated with higher rankings at 3 (median rank 1 [95% credible interval 1 to 3]), 6 (median  
7 rank 1 [1 to 3]) and 12 months of follow-up (median rank 2 [1 to 6]). Sulfonylurea  
8 demonstrated high rankings at 3 (median rank 3 [2 to 5]), 6 (median rank 3 [2 to 5]) and 24  
9 months (median rank 2 [2 to 3]). Pioglitazone had the highest ranking at 24 months (median  
10 rank 1 [1, 1]). The quality of the evidence was moderate to low.

#### 8.4.6.1.2.1 Hypoglycaemia at study end point

12 Evidence from a single network meta-analysis of 44 RCTs showed that repaglinide (median  
13 rank 10 [8 to 11]) and sulfonylurea (median rank 10 [9 to 11]) were associated with low  
14 rankings which may suggest higher rates of hypoglycaemia. In contrast, linagliptin (median  
15 rank 1 [1 to 8]) had the highest ranking but was associated with wide credible intervals. The  
16 quality of the evidence was low.

#### 8.4.6.1.3.7 Adverse events at study end point

18 Evidence from 3 network meta-analyses incorporating data from 73, 73 and 29 RCTs for  
19 adverse events, total dropouts and nausea respectively showed that repaglinide,  
20 sulfonylurea (modified release) and vildagliptin were associated with the highest rankings for  
21 dropouts due to adverse events, total dropouts and nausea respectively. However, these  
22 rankings were associated with wide credible intervals. The quality of the evidence was low.

#### 8.4.6.1.4.3 Change in body weight

24 Evidence from 2 network meta-analyses incorporating 12 and 6 RCTs at 12 and 24 months  
25 respectively showed that metformin was associated with the highest ranking at 12 (median  
26 rank 1 [1 to 3]) and 24 (median rank 1 [1 to 2]) months, suggesting that it is effective in  
27 weight loss. Sulfonylurea (standard and modified release) and pioglitazone were associated  
28 with lower rankings which may suggest worse weight related outcomes. The quality of the  
29 evidence was low.

### 8.4.6.2.0 Health economic evidence

31 A directly applicable health economic model with potentially serious limitations found  
32 metformin was less costly and more effective than all other modelled options for initial  
33 therapy. For people who could not take metformin, repaglinide was the most cost-effective  
34 option. If people were unwilling to take repaglinide, pioglitazone was the most cost-effective  
35 option; for people who could not take metformin, repaglinide or pioglitazone, sulfonylurea or  
36 sitagliptin were cost-effective modelled options.

## 8.4.7.7 Evidence to recommendations for initial therapy

### 38 Table 61: Linking evidence to recommendations

Relative value of different outcomes

The GDG agreed that the critical outcomes to consider were glycaemic control (change in HbA1c), hypoglycaemia and adverse events. The GDG agreed that an important outcome to consider was change in body weight.

Oral anti-diabetic medicines are only used when diet and lifestyle

	<p>have not demonstrated an optimal benefit, in terms of lowering or maintaining HbA1c levels. These interventions are important as type 2 diabetes is a progressive condition and review in previous chapters has shown the increased risk of complications and mortality associated with higher levels of HbA1c.</p> <p>While the GDG noted that glycaemic control was important in mitigating the increased risk of microvascular and macrovascular complications associated with hyperglycaemia, they also acknowledged that tight glycaemic control may be associated with increased risk of hypoglycaemia. Increased rates of hypoglycaemia can lead to non-compliance with therapy and increased stress and anxiety levels of the individual, associated with a detrimental effect on quality of life. It may also increase the risk of hypoglycaemia unawareness leading to more severe hypoglycaemia.</p> <p>Drug intolerability (due to adverse effects) and change in body weight have a negative impact on overall diabetes management and on the individual's quality of life. Type 2 diabetes is associated with clinical obesity and medication that results in weight gain will likely further impact on the individual's self-esteem and negatively affect quality of life.</p> <p>The relative importance of each outcome is further dependent on several factors:</p> <ul style="list-style-type: none"> <li>• Short-term (3 and 6 months) versus long-term (12 and 24 months) evaluation. For example, glucose levels are important at 3 and 6 months, but at 12 and 24 months both glucose levels and adverse events are important. Adverse events and change in body weight are also likely to be reflected at longer time points.</li> <li>• Severity of hyperglycaemia.</li> <li>• Individual circumstances, such as comorbidities.</li> </ul> <p>As medicine reviews for new treatments are usually at 3 and 6 months and maintenance reviews are held annually, these are important time points to note initially the presence of adverse effects and the effect of the medicine on glycaemic control.</p>	
<p>Trade-off between benefits and harms</p>	<p>The GDG discussed the results of the NMAs and noted that there were more data available at 3, 6 and 12 months, whereas at 24 months there was less evidence resulting in sparser networks and a limited number of interventions.</p> <p>Overall, the networks included 12 comparators including placebo. Of these 12 comparators, 7 included studies which reported data for all required outcomes in the health economic model. The 5 interventions that were not included in the health economic model were acarbose, metformin (modified release), sulfonylurea (modified release), linagliptin and saxagliptin. Of these, the GDG referred to their experience that linagliptin and saxagliptin would be expected to perform well if data were available for inclusion in these analyses.</p> <p>The GDG agreed that, while standard-release metformin was not associated with the greatest reduction in HbA1c in the reviewed evidence, the additional cardiovascular benefits associated with metformin use are very important in the overall long-term management of the individual with type 2 diabetes. Moreover, metformin was associated with fewer hypoglycaemic events, and</p>	<p>Update 2015</p>

weight loss at 12 and 24 months which are considered important for individual's quality of life. The Group also discussed the use of gradual dosing and titration of metformin which may help to reduce gastrointestinal adverse events. The GDG noted that there was limited evidence on alternative forms of metformin for individuals who cannot tolerate standard-release metformin.

The GDG discussed the evidence surrounding the remaining drug interventions available to people for whom metformin is contraindicated or not tolerated.

The GDG noted that, although sulfonylureas were associated with clinically important reductions in HbA1c in the short-term at 3 and 6 months, they were consistently associated with greater hypoglycaemic events and weight gain at 12 and 24 months. The Group noted that the occurrence of hypoglycaemic events was consistent with their experiences in clinical practice. The GDG discussed the value of using sulfonylureas to achieve rapid blood glucose control (rescue therapy) in clinical practice, but thought that the use of sulfonylureas as an immediate second option if metformin is contraindicated or not tolerated was not supported by the evidence base, due to the short-term efficacy in change in HbA1c and associated increased risks of adverse events including hypoglycaemia. The Group agreed that use of sulfonylurea as rescue therapy should consider the balance of good glycaemic control and the risk of poor weight outcomes and hypoglycaemia in discussion with patients and therefore, treatment should be reviewed once agreed targets have been met.

The GDG then considered repaglinide which was shown to be consistently associated with the largest reduction in HbA1c at 3, 6 and 12 months, but also with a greater number of hypoglycaemic events. The Group also noted that the occurrence of hypoglycaemic events was consistent with their experience in clinical practice. The GDG considered the change in body weight associated with repaglinide, and thought that while it was associated with weight gain it fared better than sulfonylureas for this outcome. The GDG recognised that repaglinide is a secretagogue not widely used in current UK clinical practice and that a recommendation to offer repaglinide as an alternative initial therapy when metformin is contraindicated or not tolerated would lead to a large change in practice but considered that the consistent findings of significantly large clinically important reductions in HbA1c up to 1 year shown in the evidence justified the recommendation.

Moreover, the high likelihood that treatment intensification would become necessary meant that the potential role of repaglinide as an initial therapy option is constrained by the fact that it is presently only licensed in combination with metformin. This means that if repaglinide does not lead to optimal results as initial therapy, then there are no licensed options to intensify with another antihyperglycaemic medicine. The GDG thought that people should be made aware of this constraint prior to starting drug treatment. The Group discussed the impact that these constraints may have on implementation and clinical practice and drafted a recommendation that when repaglinide is offered as initial therapy, the patient should be advised that if it does not have the desired results, then it will have to be stopped and another oral anti-diabetic medicine offered, prior to intensifying treatment.

	<p>The GDG discussed the evidence on the use of pioglitazone and sitagliptin, which showed similar profiles in terms of change in HbA1c and adverse events. While pioglitazone was associated with the greater reduction in HbA1c at 24 months, sitagliptin was associated with fewer hypoglycaemia and weight loss at 12 and 24 months. The Group discussed the long-term safety concerns associated with the use of pioglitazone and DPP-4 inhibitors, and agreed that MHRA guidance and patient suitability should be considered, for example, pioglitazone is not recommended for people with active bladder cancer, a history of bladder cancer, or uninvestigated haematuria, or for people with heart failure or a risk of osteoporosis. The GDG noted that there was limited information on the long-term safety of DPP-4 inhibitors but considered the evidence was strong enough to recommend these as treatment options if both metformin and repaglinide were contraindicated, not tolerated or not preferred. The GDG suggested that a cross reference to appropriate MHRA publications would also be appropriate.</p> <p>While vildagliptin generally showed less reduction for change in HbA1c at 3, 6 and 12 months, a relatively greater reduction was observed at 24 months. High to middle rankings were observed for hypoglycaemia, dropouts due to adverse events, nausea and weight at 12 and 24 months. However, overall, many point estimates were associated with large credible intervals indicating uncertainty around the data.</p>
<p>Consideration of health benefits and resource use</p>	<p>The GDG were happy to recommend metformin as initial therapy for people with type 2 diabetes, as it clearly dominated the other treatments that could be modelled. The GDG noted that, if metformin is associated with longer-term cardiovascular benefits over and above those associated with reduction of HbA1c, these would not be reflected in the economic model. While such future outcomes would be discounted, the GDG noted their inclusion may further improve the cost effectiveness of metformin. Equally, the model did not reflect potential long-term safety concerns of pioglitazone and DPP-4 inhibitors that could decrease their cost effectiveness.</p> <p>For people not able to take metformin, either because it is contraindicated or not tolerated, repaglinide was the most cost-effective option. However, the GDG noted that repaglinide is not licensed for combination use with any drug other than metformin. This means that future intensifications of treatment – that is, when HbA1c is no longer controlled by monotherapy alone – would not be straightforward for people taking repaglinide. For this reason, the Group queried what the most cost-effective initial therapy was for people who could not take metformin and did not wish to take repaglinide. Of the remaining drugs modelled, pioglitazone was the cheapest. The Group discussed the known contraindications for pioglitazone and that the vast majority of people would be taking metformin or repaglinide; the GDG were happy to consider either sulfonylurea or sitagliptin as a fourth choice for initial therapy.</p> <p>Differences between drugs at initial therapy were small due to the normalising effect of future intensifications in the economic model – simulated people were only on their initial therapies for an average of 3.4 years. QALY differences were driven by differences in weight gained, both from initial therapy itself and differences in time until intensification. Cost differences were largely due to the costs of the drugs themselves.</p>

	<p>The economic model used a 1-year cycle and the GDG noted that this may not fully reflect the clinical utility of treatments such as sulfonylurea and repaglinide that may achieve shorter-term HbA1c benefits that may not be sustained at 1 year. In contrast, the economic model did reflect the low rankings at 1 year for hypoglycaemia and body weight for these treatments. The GDG appreciated the ability of the model to combine all modelled outcomes.</p> <p>The GDG queried whether dosing differences may have driven different uptake patterns, as sulfonylureas are generally taken once daily whereas repaglinide is taken multiple times daily. However, it was noted that metformin is also taken multiple times daily so the GDG felt it was unlikely that any disutility or increased dropout rate would be associated with repaglinide due to multiple daily tablets.</p>
Quality of the evidence	<p>The GDG agreed that the overall quality of the evidence for initial therapy was generally moderate.</p>

#### 8.4.8.1 Clinical evidence review for first intensification

- 2 In total 17,037 references were found for the main review question and 47 papers were  
3 included for first intensification which relate to 34 trials.
- 4 This sub-review question addressed which treatment combination of 2 non-insulin based  
5 options is most effective when people with type 2 diabetes have inadequate blood glucose  
6 control, typically following management with diet and a single oral antidiabetic medicine.
- 7 RCTs of at least 12 week treatment duration comparing dual therapies of 2 non-insulin based  
8 treatments were included. In contrast to initial therapy, it was assumed that most patients  
9 would be titrated to the maximal tolerated doses of previous oral therapy before starting a  
10 trial. Therefore, trials that did not report specific doses of continued previous therapy were  
11 still included (see section 8.4.2 for the main exclusion criteria).

#### 8.4.8.12 Description of included studies for first intensification

13 A total of 17,835 participants in 34 RCTs were included. The majority of studies were carried  
14 out in multiple centres across different countries. The mean age ranged from 50.8 to 63.2  
15 years, with 3 studies not reporting this information. Mean HbA1c levels at baseline ranged  
16 from 54 to 77 mmol/mol (7.1% to 9.2 %), with 1 study not reporting this information. The  
17 mean BMI ranged from 22.9 to 51.5 kg/m<sup>2</sup>, with 1 study not reporting this information. Mean  
18 duration of diabetes ranged from 1.9 to 8.6 years, with 8 studies not reporting this  
19 information. Follow-up periods ranged from 12 to 156 weeks. For full details of the included  
20 studies see Appendix E.

#### 8.4.8.21 Network meta-analyses for first intensification

- 22 To facilitate comparison across all available treatment options, 10 network meta-analyses  
23 were performed for all 3 critical and 1 important outcomes – change in HbA1c at 3, 6, 12 and  
24 24 months, hypoglycaemia at study end point, adverse events (that is, dropouts due to  
25 adverse events, total dropouts and nausea) at study end point and change in body weight at  
26 12 and 24 months. Metformin-sulfonylurea was selected as the reference treatment option as  
27 this combination was considered to reflect current standard clinical practice. Full details of  
28 methods and additional NMA outputs are provided in Appendix J.
- 29 Generally, well-connected networks were produced for shorter follow-up times although  
30 these tended to be sparser and contained fewer treatment options at 12 and 24 months.

1 Pairwise comparisons that did not form part of the main network were not presented as they  
2 would not add to the GDG decision making.

3 On the whole, the quality of the evidence was moderate to low as networks were relatively  
4 well-connected by a star shaped network with metformin-sulfonylurea treatment in the  
5 middle. However some included trials were not double-blind and did not report adequate  
6 details of randomisation and allocation concealment methods. It was noted that random-  
7 effects models tended to estimate a fairly large inter-study heterogeneity term, which will  
8 reduce the precision of effect estimates.

9 **Table 62: GRADE profile for network meta-analyses for first intensification**

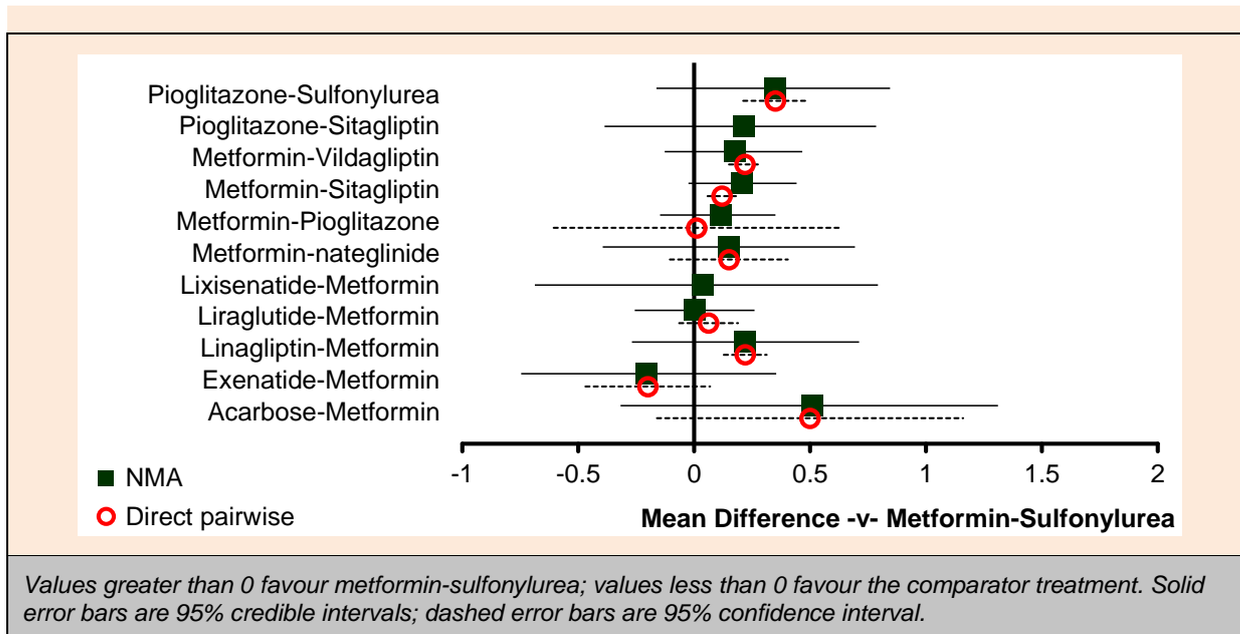
Assessment time points/ Measure	Number of RCTs	Risk of bias	Inconsistency	Indirectness	Imprecision	Quality
<b>Change in blood glucose (HbA1c)</b>						
3 months	20	not serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Moderate
6 months	22	not serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Moderate
12 months	16	not serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Moderate
24 months	6	not serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Moderate
<b>Hypoglycaemia at study end point</b>						
Study end point	21	not serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<b>Adverse events at study end point</b>						
Dropouts due to adverse events	27	not serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
Total dropouts	29	not serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Moderate
Nausea	11	not serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<b>Change in body weight</b>						
12 months	8	not serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
24 months	8	not serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<sup>1</sup> Baseline HbA1c ranged from 7.1 to 9.9%						
<sup>2</sup> Assessed based on residual deviance, deviance information criterion and $\tau^2$ ( $\tau^2 < 0.5$ )						
<sup>3</sup> Considered not serious as population, interventions, comparator and outcomes are as defined in protocol						
<sup>4</sup> Downgrade 1 level: no interventions had probability of being best and worse $\geq 0.5$						
<sup>5</sup> Downgrade 1 level: $\tau^2 \geq 0.5$						

#### 8.4.8.30 Change in blood glucose (HbA1c) at 3, 6, 12 and 24 months

11 Results of the NMAs are summarised below for the 11 treatment options that were compared  
12 with metformin-sulfonylurea at 3 and 6 months, and the 10 and 6 treatment options assessed  
13 at 12 and 24 months respectively.

14 Across all 4 follow-up time points, metformin-based combinations were shown to be the most  
15 effective in reducing HbA1c levels when compared to metformin-sulfonylurea. However,  
16 these relative effects were generally associated with wide credible intervals, which except for  
17 2 treatment combinations at 6 months, crossed the line of no effect. At 3 months, metformin-



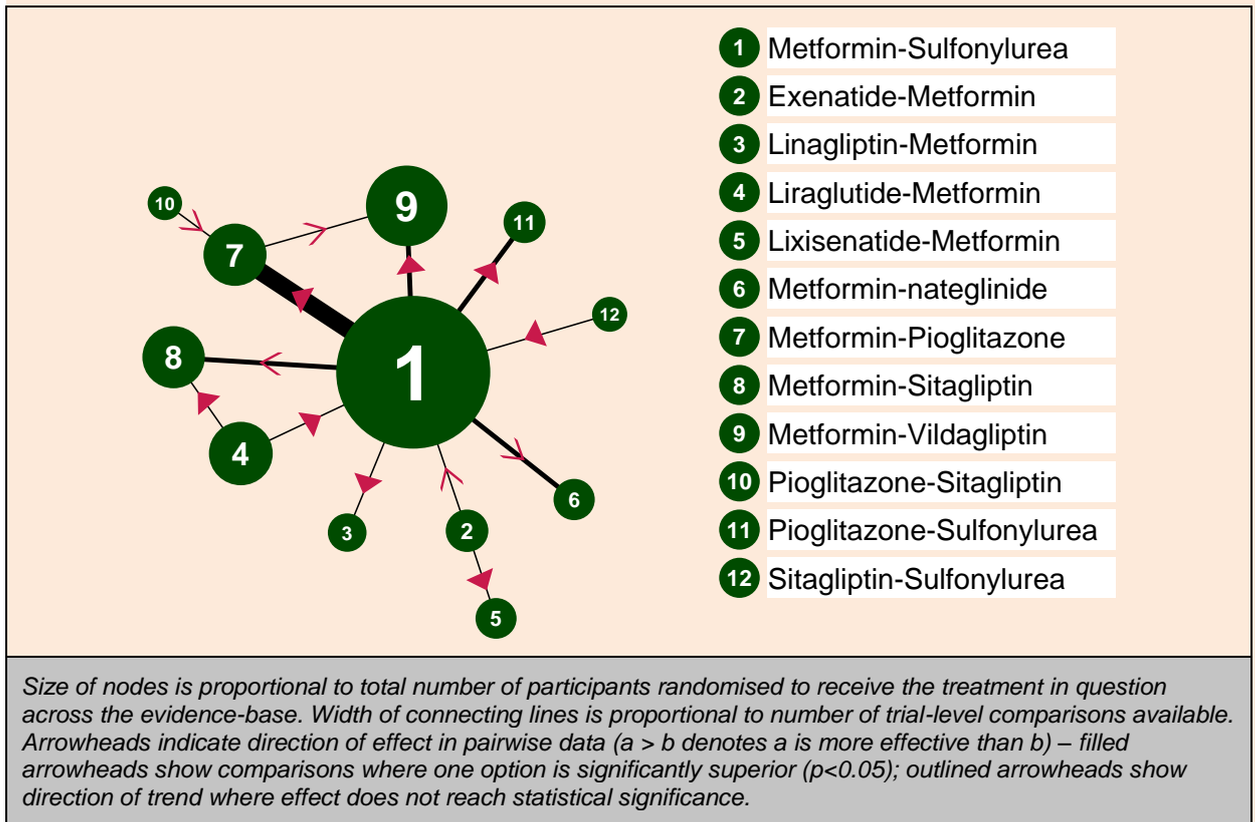


1 **Figure 31: Network meta-analysis of change in HbA1c (3 months) – relative effect of**  
 2 **all options compared with common comparator (metformin-sulfonylurea)**

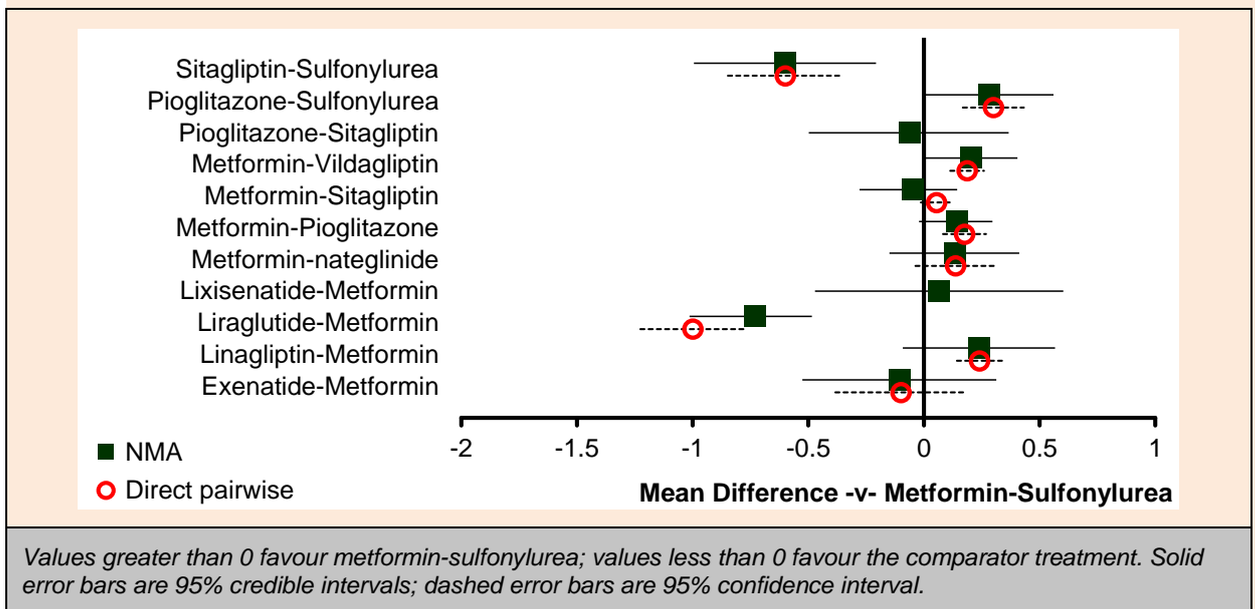
3  
 4 **Table 63: Network meta-analysis of change in HbA1c (3 months) – rankings for each**  
 5 **comparator**

	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.030	4 (1, 7)
Acarbose-Metformin	0.036	11 (1, 12)
Exenatide-Metformin	0.512	1 (1, 9)
Linagliptin-Metformin	0.038	8 (1, 12)
Liraglutide-Metformin	0.094	4 (1, 9)
Lixisenatide-Metformin	0.103	5 (1, 12)
Metformin-nateglinide	0.082	7 (1, 12)
Metformin-Pioglitazone	0.014	6 (2, 10)
Metformin-Sitagliptin	0.001	8 (4, 11)
Metformin-Vildagliptin	0.012	7 (2, 11)
Pioglitazone-Sitagliptin	0.064	8 (1, 12)
Pioglitazone-Sulfonylurea	0.014	10 (2, 12)

6  
7



1 **Figure 32: Network meta-analysis of change in HbA1c (6 months) – evidence**  
2 **network**

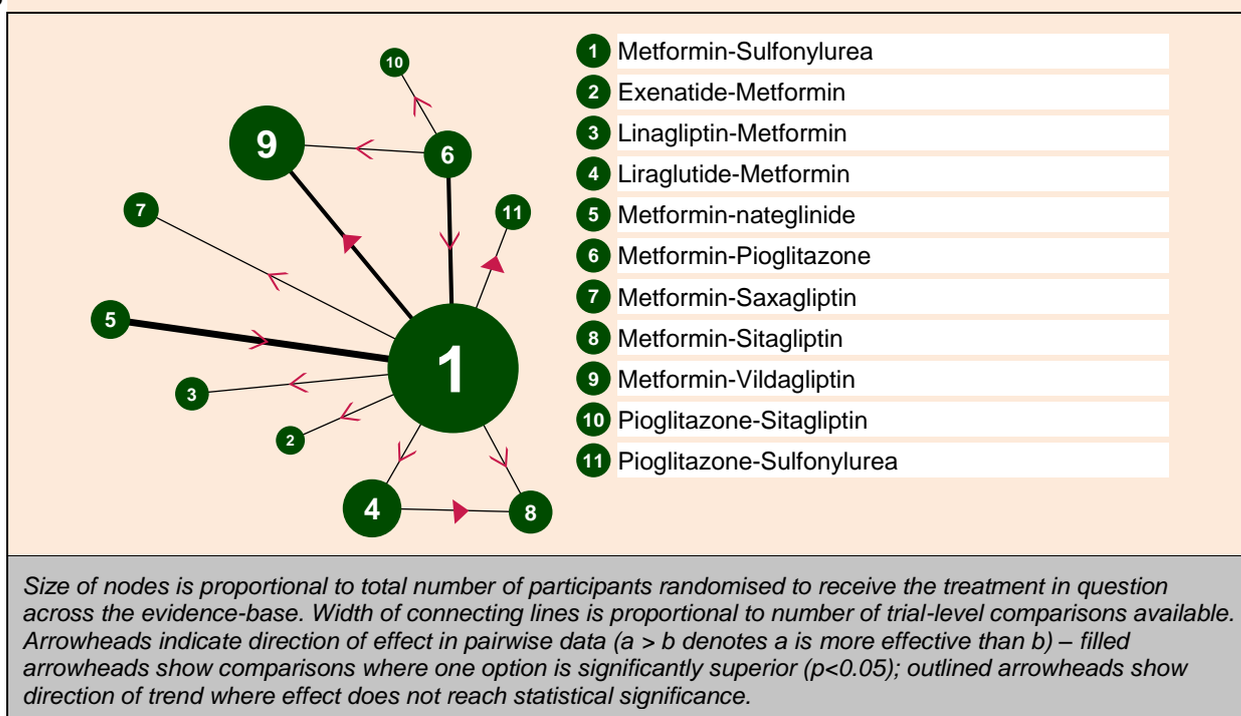


4 **Figure 33: Network meta-analysis of change in HbA1c (6 months) – relative effect of**  
5 **all options compared with common comparator (metformin-sulfonylurea)**

1 **Table 64: Network meta-analysis of change in HbA1c (6 months) – rankings for each**  
2 **comparator**

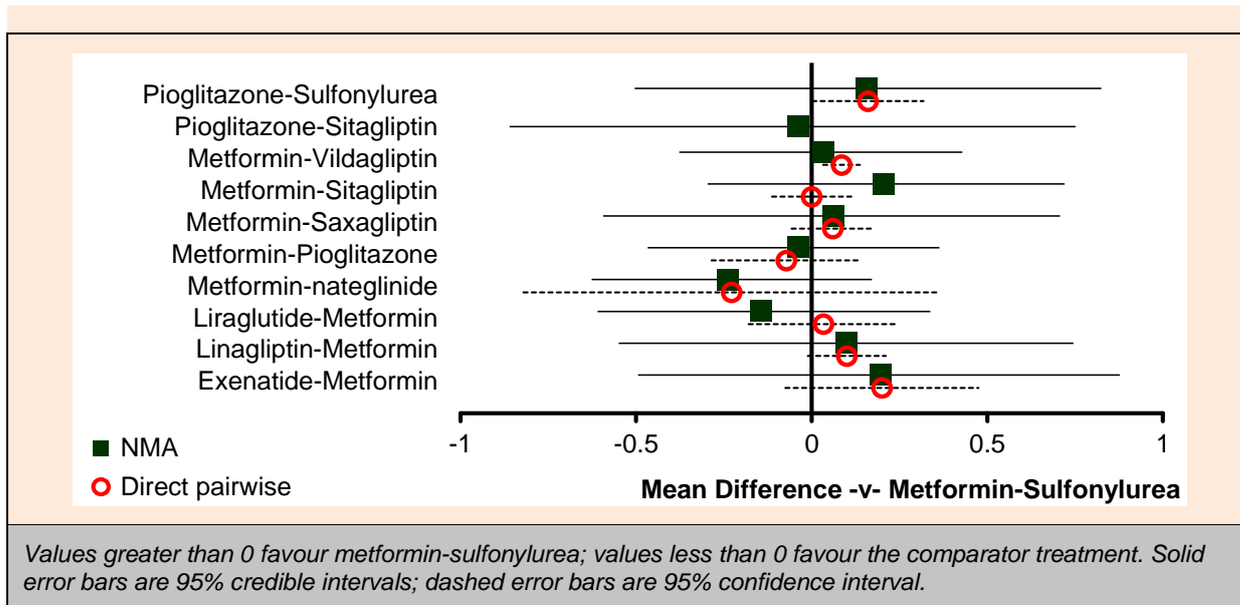
	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	6 (4, 8)
Exenatide-Metformin	0.003	4 (2, 10)
Linagliptin-Metformin	0.000	10 (4, 12)
Liraglutide-Metformin	0.712	1 (1, 2)
Lixisenatide-Metformin	0.003	7 (3, 12)
Metformin-nateglinide	0.000	8 (3, 12)
Metformin-Pioglitazone	0.000	9 (5, 11)
Metformin-Sitagliptin	0.000	5 (3, 9)
Metformin-Vildagliptin	0.000	10 (6, 12)
Pioglitazone-Sitagliptin	0.003	5 (2, 12)
Pioglitazone-Sulfonylurea	0.000	11 (6, 12)
Sitagliptin-Sulfonylurea	0.278	2 (1, 4)

3



4 **Figure 34: Network meta-analysis of change in HbA1c (12 months) – evidence**  
5 **network**

6

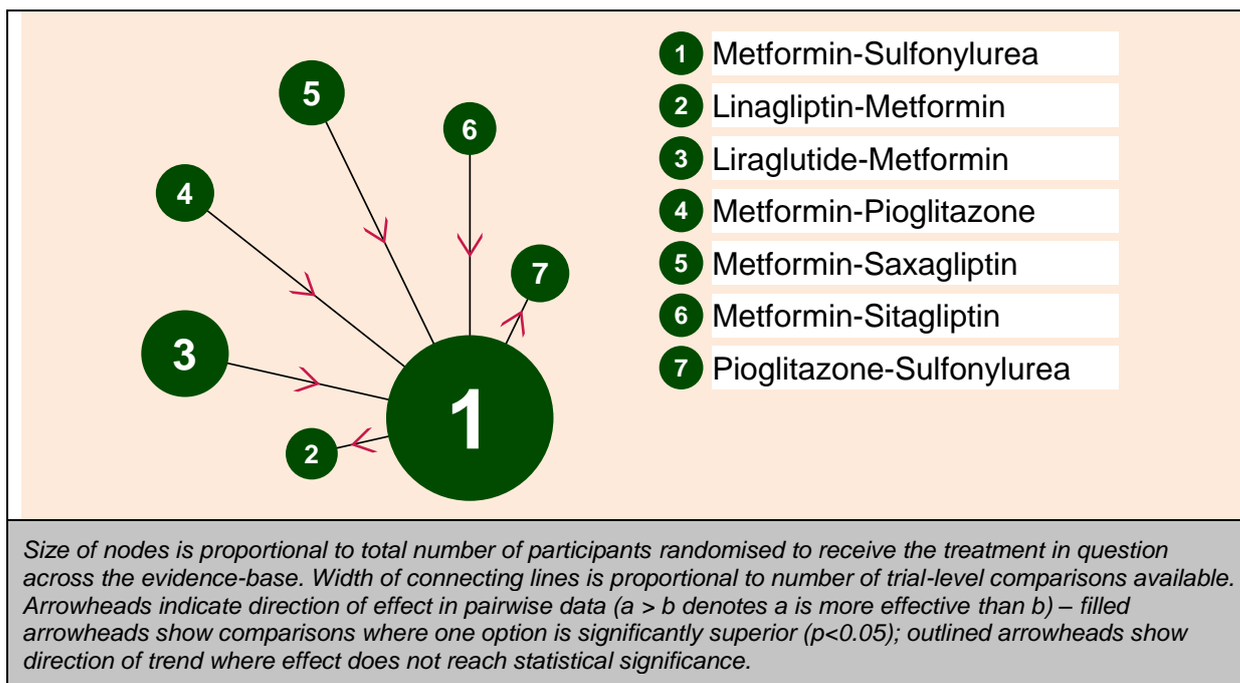


1 **Figure 35: Network meta-analysis of change in HbA1c (12 months) – relative effect**  
 2 **of all options compared with common comparator (metformin-sulfonylurea)**

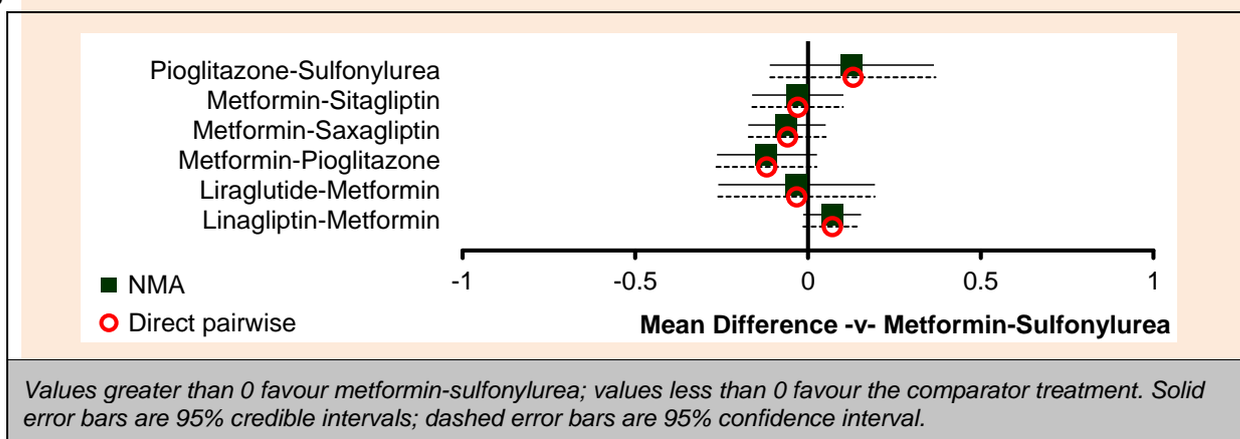
3  
 4 **Table 65: Network meta-analysis of change in HbA1c (12 months) – rankings for each**  
 5 **comparator**

	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.001	6 (3, 9)
Exenatide-Metformin	0.044	9 (1, 11)
Linagliptin-Metformin	0.061	7 (1, 11)
Liraglutide-Metformin	0.179	3 (1, 9)
Metformin-nateglinide	0.327	2 (1, 8)
Metformin-Pioglitazone	0.039	5 (1, 10)
Metformin-Saxagliptin	0.081	7 (1, 11)
Metformin-Sitagliptin	0.007	9 (2, 11)
Metformin-Vildagliptin	0.025	6 (2, 11)
Pioglitazone-Sitagliptin	0.191	5 (1, 11)
Pioglitazone-Sulfonylurea	0.047	8 (1, 11)

6



1 **Figure 36: Network meta-analysis of change in HbA1c (24 months) – evidence**  
2 **network**



4 **Figure 37: Network meta-analysis of change in HbA1c (24 months) – relative effect**  
5 **of all options compared with common comparator (metformin-sulfonylurea)**

7 **Table 66: Network meta-analysis of change in HbA1c (24 months) – rankings for each**  
8 **comparator**

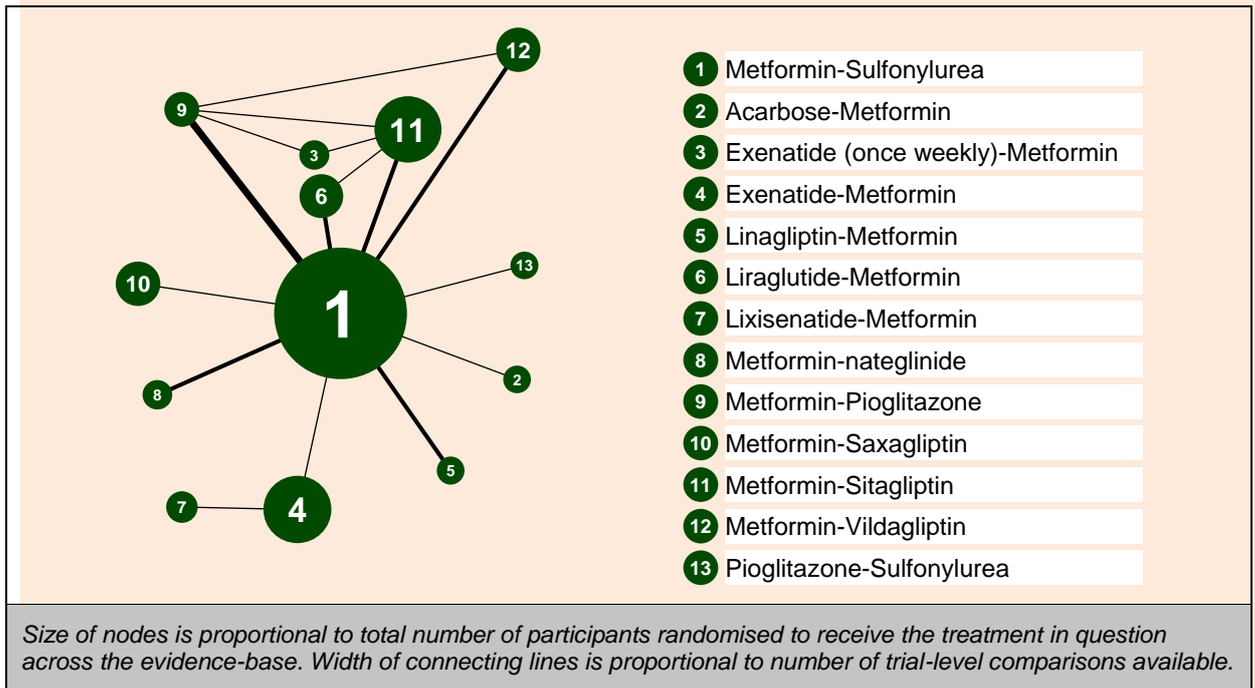
	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.001	4 (3, 6)
Linagliptin-Metformin	0.000	6 (4, 7)
Liraglutide-Metformin	0.201	3 (1, 7)
Metformin-Pioglitazone	0.538	1 (1, 5)
Metformin-Saxagliptin	0.155	3 (1, 6)
Metformin-Sitagliptin	0.088	3 (1, 7)
Pioglitazone-Sulfonylurea	0.018	7 (2, 7)

1

#### 8.4.8.42 Hypoglycaemia at study end point

3 Results of the NMA are summarised below for the 11 treatment combinations that were  
 4 compared with metformin-sulfonylurea. In general, all treatment combinations were more  
 5 effective at preventing hypoglycaemic events than metformin-sulfonylurea which had the  
 6 lowest ranking (median rank 12 [10 to 12]), followed by pioglitazone-sulfonylurea (median  
 7 rank 11 [6 to 12]). Metformin-acarbose (median rank 2 [1 to 10]), metformin-lixisenatide  
 8 (median rank 2 [1 to 7]) and metformin-saxagliptin (median rank 2 [1 to 6]) shared the highest  
 9 ranking position, though metformin-saxagliptin had the narrowest credible intervals.

10

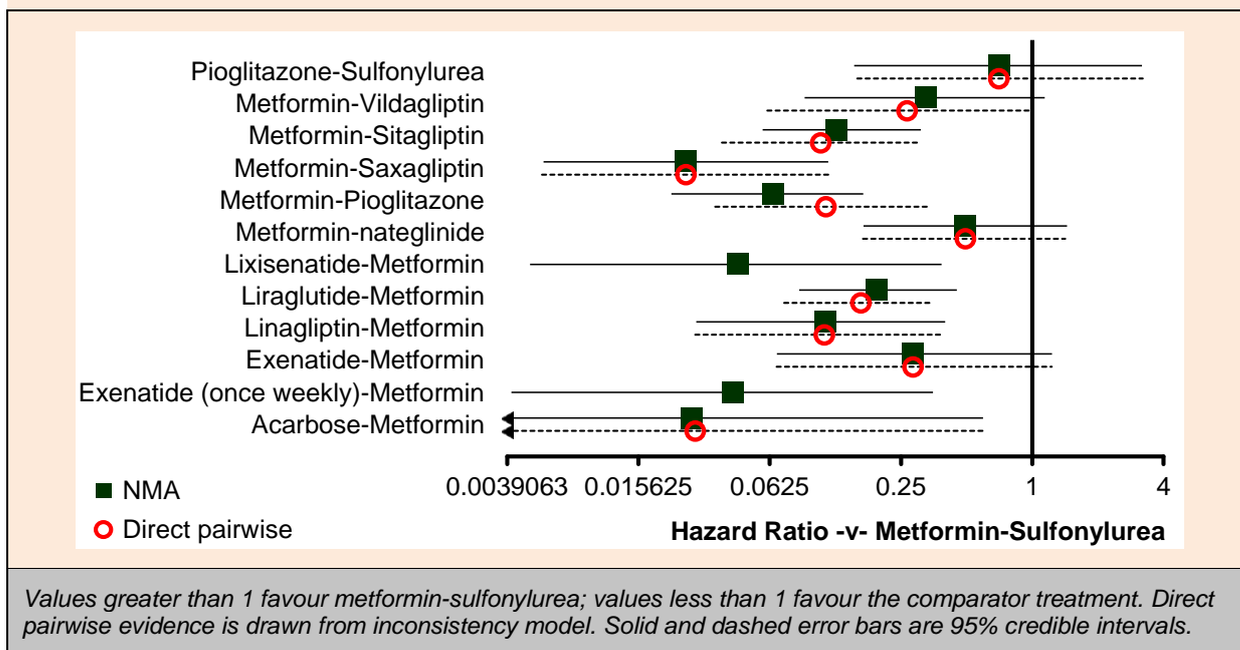


Update 2015

11 **Figure 38: Network meta-analysis of hypoglycaemic events (study end point) –**  
 12 **evidence network**

13

1



2 **Figure 39: Network meta-analysis of hypoglycaemic events (study end point) –**  
 3 **relative effect of all options compared with common comparator (metformin-**  
 4 **sulfonylurea)**

5 **Table 67: Network meta-analysis of hypoglycaemic events (study end point) – rankings**  
 6 **for each comparator**

	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	13 (11, 13)
Acarbose-Metformin	0.401	2 (1, 11)
Exenatide (once weekly)-Metformin	0.167	3 (1, 10)
Exenatide-Metformin	0.000	9 (5, 13)
Linagliptin-Metformin	0.009	6 (2, 10)
Liraglutide-Metformin	0.000	8 (5, 11)
Lixisenatide-Metformin	0.140	3 (1, 9)
Metformin-nateglinide	0.000	11 (7, 13)
Metformin-Pioglitazone	0.016	4 (2, 7)
Metformin-Saxagliptin	0.267	2 (1, 6)
Metformin-Sitagliptin	0.000	6 (4, 9)
Metformin-Vildagliptin	0.000	10 (5, 13)
Pioglitazone-Sulfonylurea	0.000	12 (7, 13)

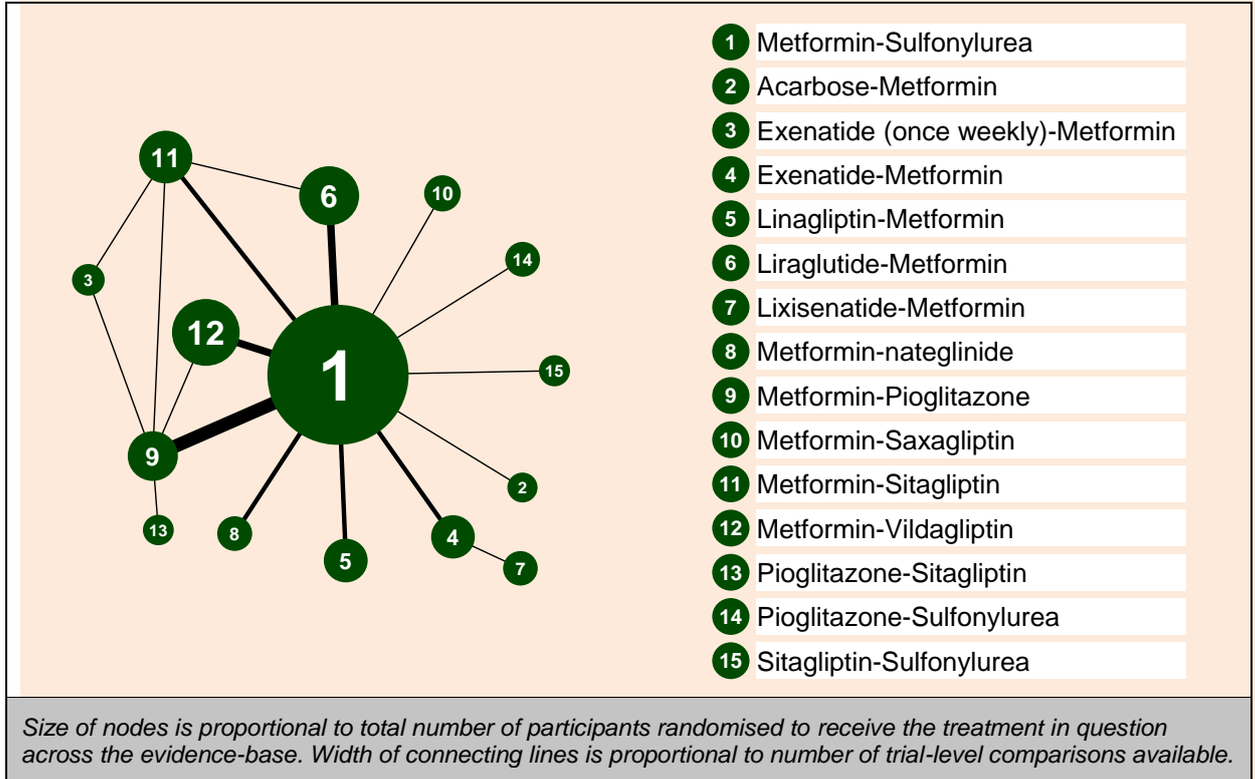
#### 8.4.8.57 Adverse events at study end point

8 Results of the 3 NMAs are summarised below. For dropouts due to adverse events and total  
 9 dropouts, 12 treatment combinations were compared with metformin-sulfonylurea, while 7  
 10 treatment combinations were compared with metformin-sulfonylurea for nausea.

11 There is reasonable agreement between the NMA evidence and direct pairwise treatment  
 12 effect estimates, as demonstrated by the substantial overlap between the  
 13 credible/confidence intervals. In general, across all 3 measures, there were wide credible  
 14 intervals which crossed the line of no effect. However, for all 3 measures, there was a trend

1 for metformin-GLP1 mimetics (exenatide, liraglutide and lixisenatide) to be less effective at  
 2 preventing attrition and nausea than metformin-sulfonylurea.  
 3 Pioglitazone-sitagliptin (median rank 3 [1 to 12]), metformin-nateglinide (median rank 2 [1 to  
 4 10]) and metformin-pioglitazone or sulfonylurea (median rank 2 [1 to 5] or median rank 2 [1 to  
 5 4] respectively) were associated with the highest rankings for dropouts due to adverse  
 6 events, total dropouts and nausea respectively but the associated credible intervals were  
 7 generally appreciably wide.

8

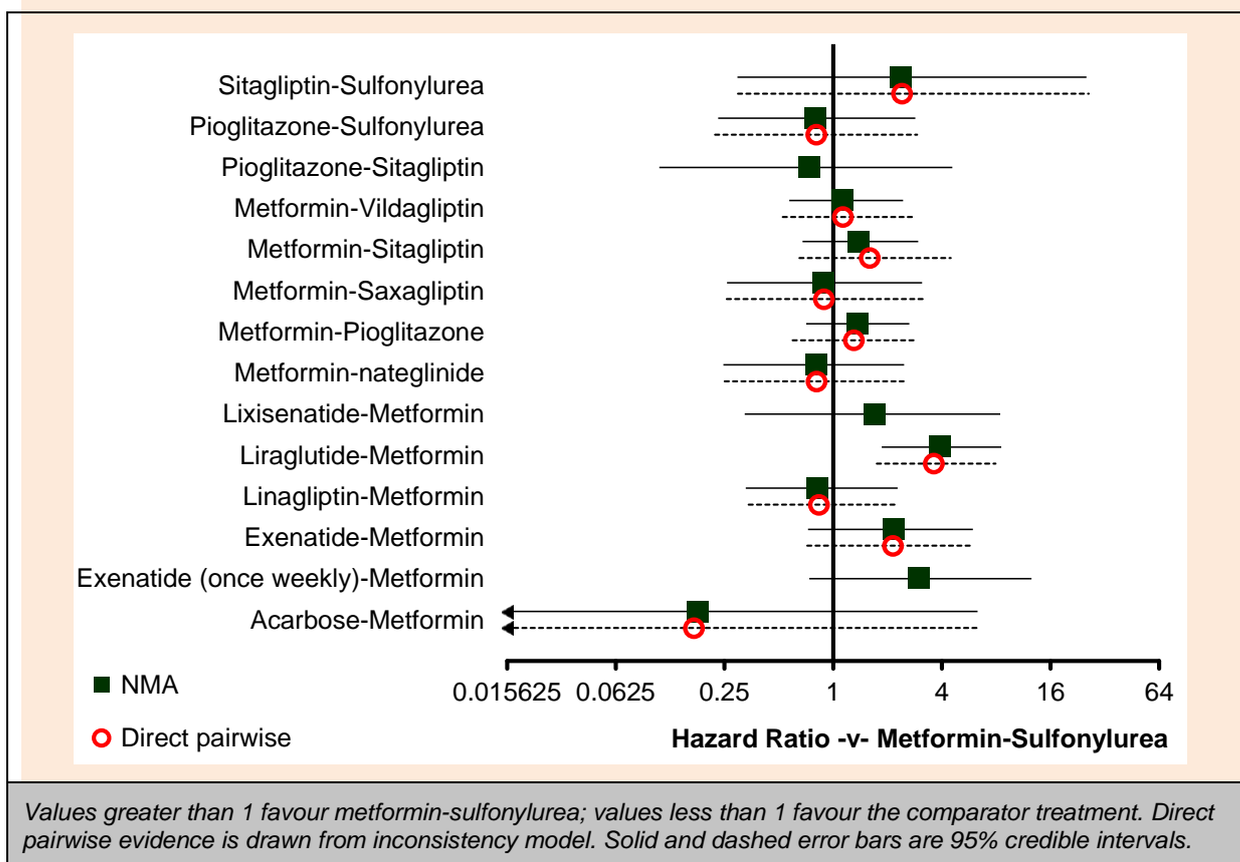


Update 2015

9 **Figure 40: Network meta-analysis of dropouts due to adverse events (study end**  
 10 **point) – evidence network**

11

1



2 **Figure 41: Network meta-analysis of dropouts due to adverse events (study end**  
 3 **point) – relative effect of all options compared with common comparator**  
 4 **(metformin-sulfonylurea)**

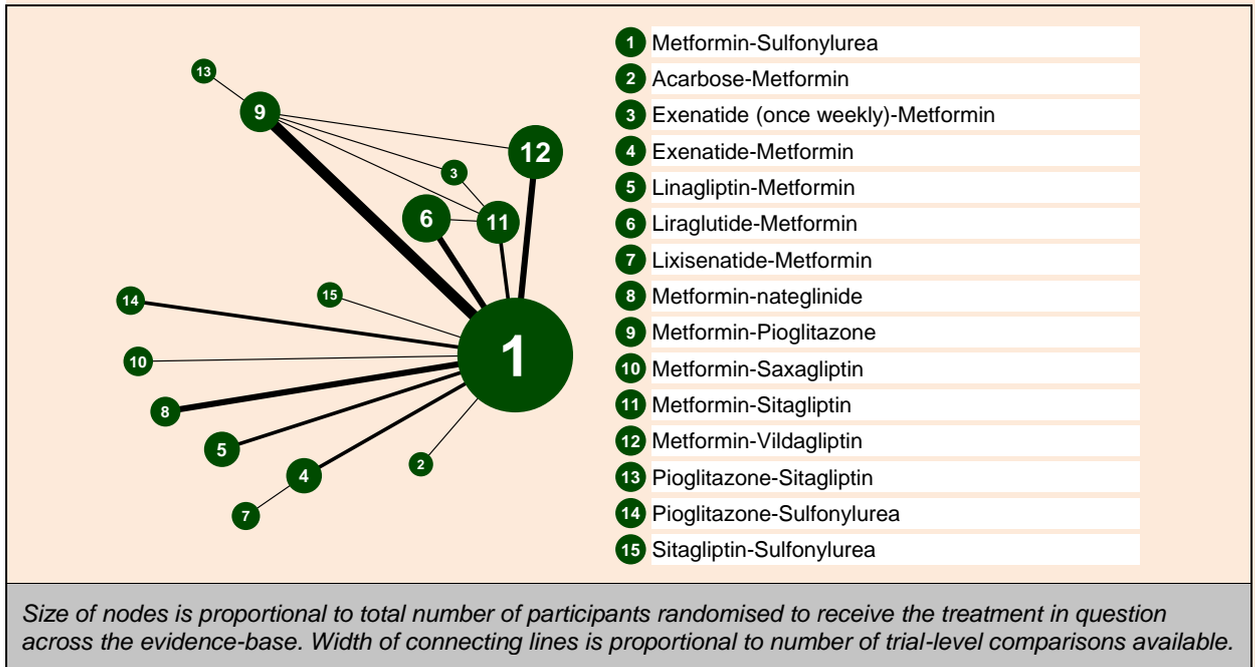
5

6 **Table 68: Network meta-analysis of dropouts due to adverse events (study end point) –**  
 7 **rankings for each comparator**

	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	6 (3, 9)
Acarbose-Metformin	0.638	1 (1, 15)
Exenatide (once weekly)-Metformin	0.001	13 (4, 15)
Exenatide-Metformin	0.001	12 (4, 15)
Linagliptin-Metformin	0.036	5 (1, 12)
Liraglutide-Metformin	0.000	14 (11, 15)
Lixisenatide-Metformin	0.017	11 (2, 15)
Metformin-nateglinide	0.054	5 (1, 13)
Metformin-Pioglitazone	0.000	9 (4, 13)
Metformin-Saxagliptin	0.043	5 (1, 13)
Metformin-Sitagliptin	0.001	9 (3, 13)
Metformin-Vildagliptin	0.003	7 (3, 13)
Pioglitazone-Sitagliptin	0.124	4 (1, 14)
Pioglitazone-Sulfonylurea	0.060	5 (1, 13)
Sitagliptin-Sulfonylurea	0.020	12 (2, 15)

Update 2015

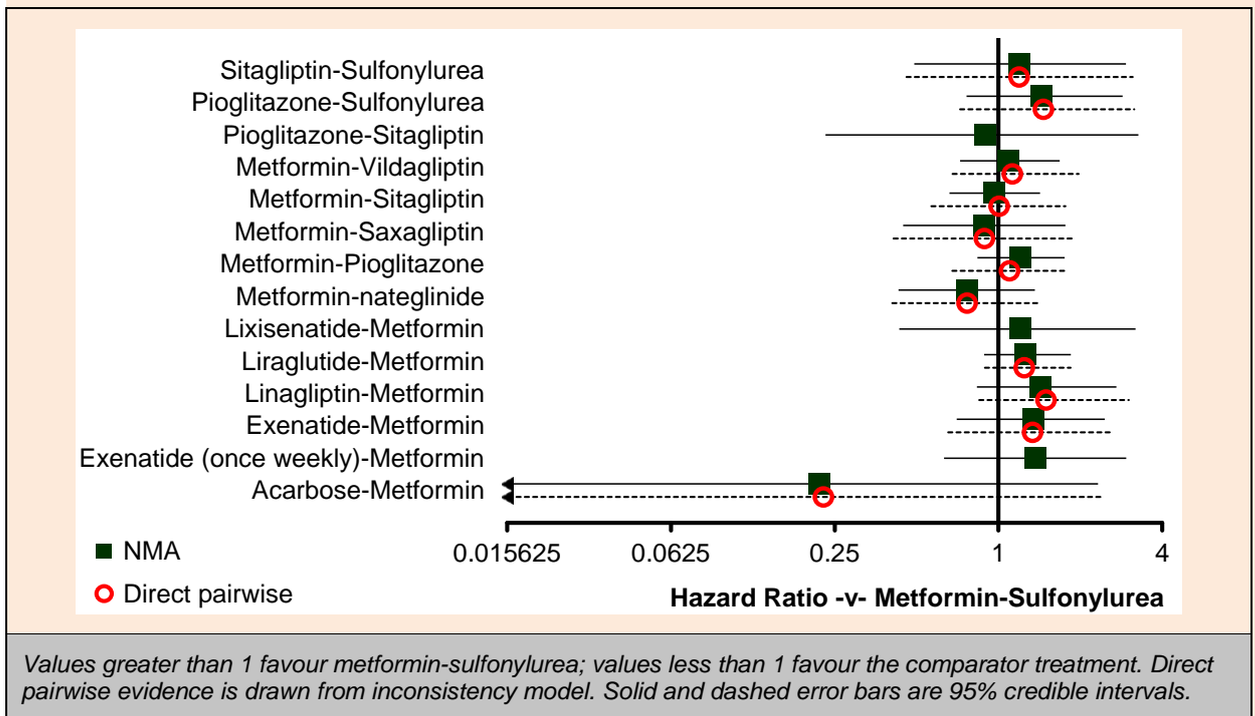
1



2 **Figure 42: Network meta-analysis of total dropouts (study end point) – evidence**  
3 **network**

4

5

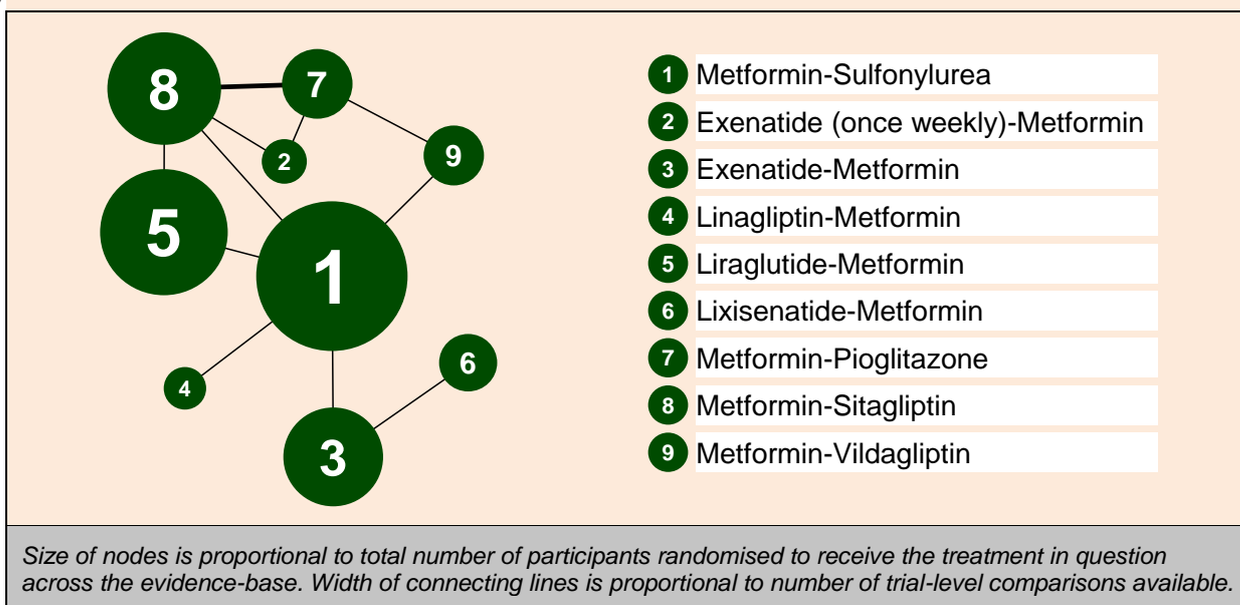


6 **Figure 43: Network meta-analysis of total dropouts (study end point) – relative**  
7 **effect of all options compared with common comparator (metformin-**  
8 **sulfonylurea)**

1 **Table 69: Network meta-analysis of total dropouts (study end point) – rankings for**  
2 **each comparator**

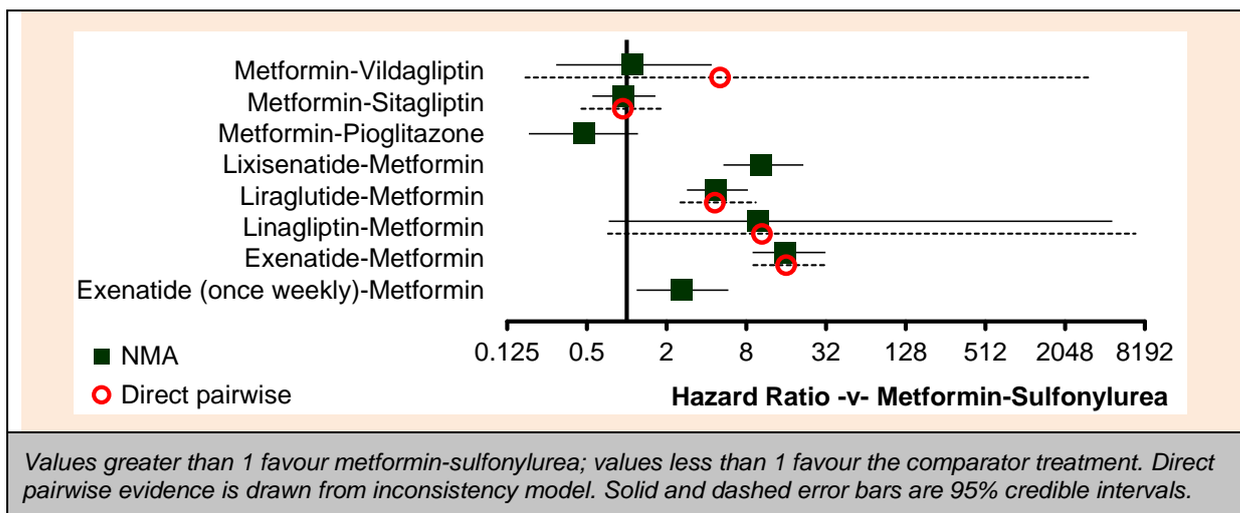
	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	6 (3, 10)
Acarbose-Metformin	0.754	1 (1, 15)
Exenatide (once weekly)-Metformin	0.006	11 (2, 15)
Exenatide-Metformin	0.001	11 (3, 15)
Linagliptin-Metformin	0.001	12 (4, 15)
Liraglutide-Metformin	0.000	10 (5, 14)
Lixisenatide-Metformin	0.022	10 (2, 15)
Metformin-nateglinide	0.059	3 (1, 12)
Metformin-Pioglitazone	0.000	10 (4, 14)
Metformin-Saxagliptin	0.034	5 (1, 14)
Metformin-Sitagliptin	0.006	6 (2, 12)
Metformin-Vildagliptin	0.002	8 (3, 14)
Pioglitazone-Sitagliptin	0.096	5 (1, 15)
Pioglitazone-Sulfonylurea	0.001	12 (3, 15)
Sitagliptin-Sulfonylurea	0.018	9 (2, 15)

3



4 **Figure 44: Network meta-analysis of nausea (study end point) – evidence network**

5



1 **Figure 45: Network meta-analysis of nausea (study end point) – relative effect of all**  
 2 **options compared with common comparator (metformin-sulfonylurea)**

3 **Table 70: Network meta-analysis of nausea (study end point) – rankings for each**  
 4 **comparator**

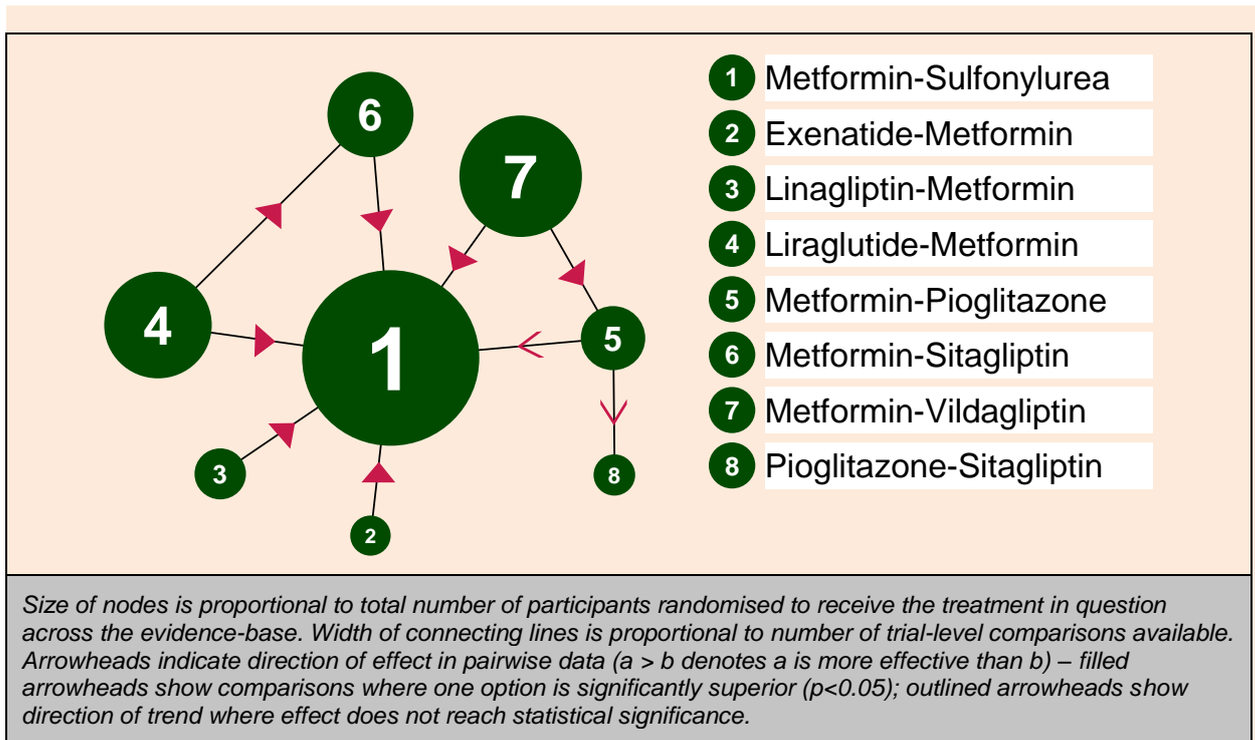
	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.048	3 (1, 4)
Exenatide (once weekly)-Metformin	0.000	5 (4, 6)
Exenatide-Metformin	0.000	9 (8, 9)
Linagliptin-Metformin	0.014	7 (2, 9)
Liraglutide-Metformin	0.000	6 (5, 7)
Lixisenatide-Metformin	0.000	7 (6, 8)
Metformin-Pioglitazone	0.863	1 (1, 3)
Metformin-Sitagliptin	0.023	3 (2, 4)
Metformin-Vildagliptin	0.053	4 (1, 6)

#### 8.4.8.65 Change in body weight at 12 and 24 months

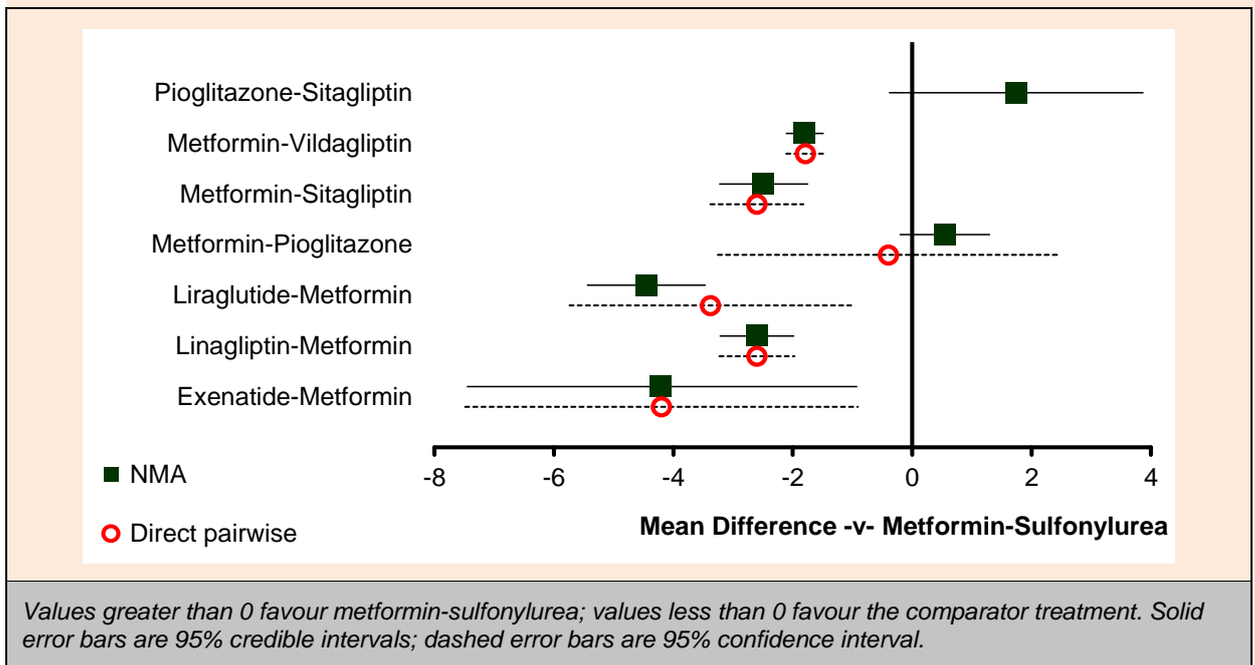
6 Results of the 2 NMAs are summarised below for the 7 and 8 treatment combinations that  
 7 were compared with metformin-sulfonylurea at 12 and 24 months respectively. There was  
 8 reasonable agreement in the NMA evidence and direct pairwise treatment effect estimates,  
 9 with substantial overlap between the credible/confidence intervals.

10 In general, metformin combined with a DPP-4 inhibitor (linagliptin, sitagliptin and vildagliptin)  
 11 or a GLP-1 mimetic (exenatide and liraglutide) were effective at weight loss when compared  
 12 to metformin-sulfonylurea at 12 and 24 months. Metformin-exenatide and metformin-  
 13 liraglutide had the highest ranking position at 12 months (median rank 2 [1 to 6] and median  
 14 rank 2 [1 to 4] respectively) while metformin-liraglutide and metformin-linagliptin had the  
 15 highest ranking position at 24 months (median rank 2 [1 to 5] and median rank 2 [1 to 6]  
 16 respectively). Pioglitazone combined with sitagliptin or metformin at 12 months (median rank  
 17 8 [5 to 8] or median rank 7[5 to 8] respectively) and pioglitazone combined with sulfonylurea  
 18 at 24 months (median rank 9 [9 to 9]) had the lowest ranking, suggesting weight gain.

19



1 **Figure 46: Network meta-analysis of change in body weight (12 months) – evidence**  
2 **network**  
3

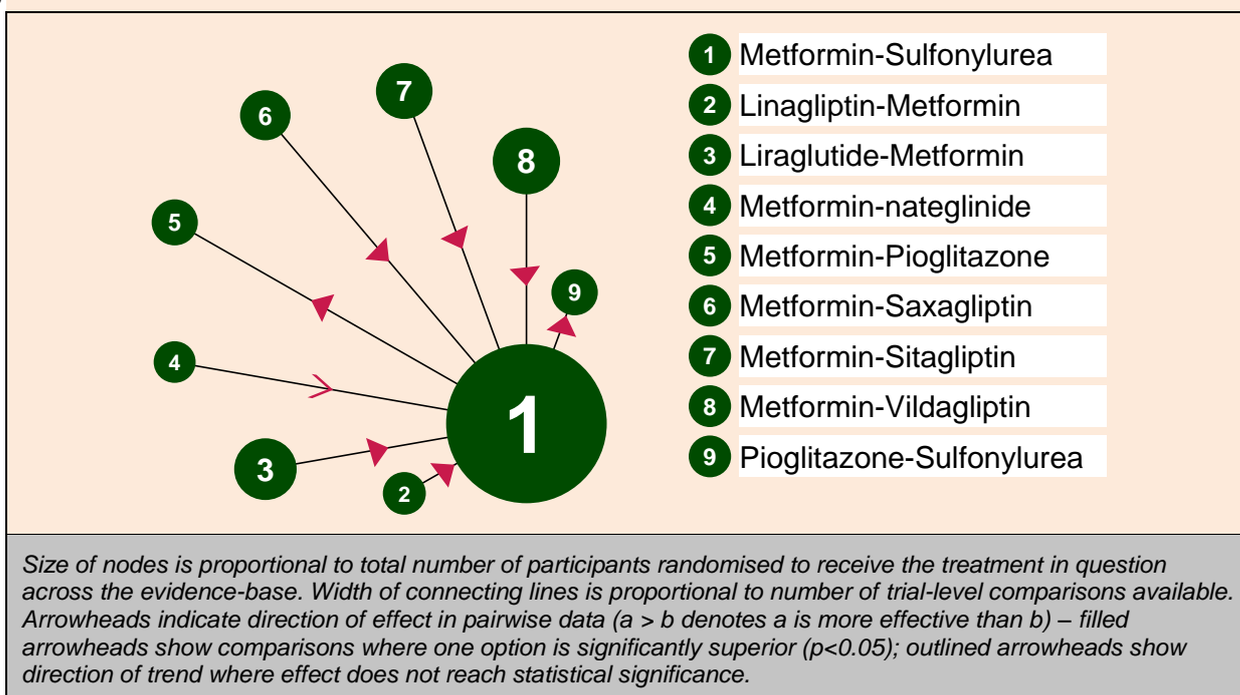


4 **Figure 47: Network meta-analysis of change in body weight (12 months) – relative**  
5 **effect of all options compared with common comparator (metformin-**  
6 **sulfonylurea)**  
7

1 **Table 71: Network meta-analysis of change in body weight (12 months) – rankings for**  
2 **each comparator**

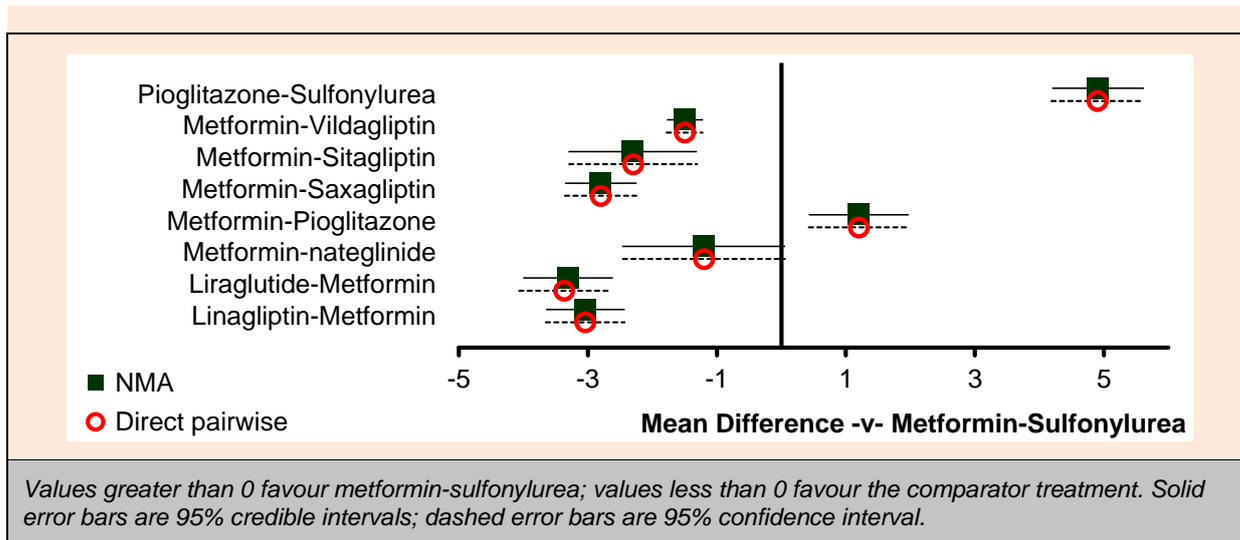
	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	6 (6, 7)
Exenatide-Metformin	0.445	2 (1, 5)
Linagliptin-Metformin	0.000	3 (2, 4)
Liraglutide-Metformin	0.555	1 (1, 2)
Metformin-Pioglitazone	0.000	7 (6, 8)
Metformin-Sitagliptin	0.000	4 (2, 5)
Metformin-Vildagliptin	0.000	5 (4, 5)
Pioglitazone-Sitagliptin	0.000	8 (6, 8)

3



4 **Figure 48: Network meta-analysis of change in body weight (24 months) – evidence**  
5 **network**

6



1 **Figure 49: Network meta-analysis of change in body weight (24 months) – relative**  
 2 **effect of all options compared with common comparator (metformin-**  
 3 **sulfonylurea)**

5 **Table 72: Network meta-analysis of change in body weight (24 months) – rankings for**  
 6 **each comparator**

	Probability best	Median rank (95%CrI)
Metformin-Sulfonylurea	0.000	7 (6, 7)
Linagliptin-Metformin	0.249	2 (1, 4)
Liraglutide-Metformin	0.652	1 (1, 3)
Metformin-nateglinide	0.001	6 (4, 7)
Metformin-Pioglitazone	0.000	8 (8, 8)
Metformin-Saxagliptin	0.068	3 (1, 4)
Metformin-Sitagliptin	0.030	4 (1, 5)
Metformin-Vildagliptin	0.000	5 (4, 6)
Pioglitazone-Sulfonylurea	0.000	9 (9, 9)

Update 2015

## 8.4.97 Health economic evidence for first intensification

### 8.4.9.18 Systematic review of published cost–utility analyses

9 For first intensification, 2 UK studies were included covering 3 comparisons (Davies et al.  
 10 2012; Schwartz et al. 2008). Davies et al. (2012) found liraglutide-metformin to be cost  
 11 effective compared with both metformin-sulfonylurea (liraglutide 1.2mg ICER £9400 per  
 12 QALY, liraglutide 1.8mg ICER £16,500 per QALY) and metformin-sitagliptin (liraglutide  
 13 1.2mg ICER £9900 per QALY, liraglutide 1.8mg ICER £10,500 per QALY). Schwartz et al.  
 14 (2008) found metformin-sitagliptin to be cost effective compared with metformin-sulfonylurea  
 15 in Scotland (ICER €11,600 per QALY). Both papers included treatment effects for systolic  
 16 blood pressure and cholesterol (as well as HbA1c, weight and hypoglycaemia). Some  
 17 assumptions and data sources used in these CUAs were unclear. Both used relatively large  
 18 utility decrements for weight gain and hypoglycaemia.

19 As no directly applicable studies with only minor limitations were found that covered all the  
 20 comparators under consideration for each sub-question for this guideline, an original  
 21 economic analysis was undertaken.

### 8.4.9.21 Original health economic analysis

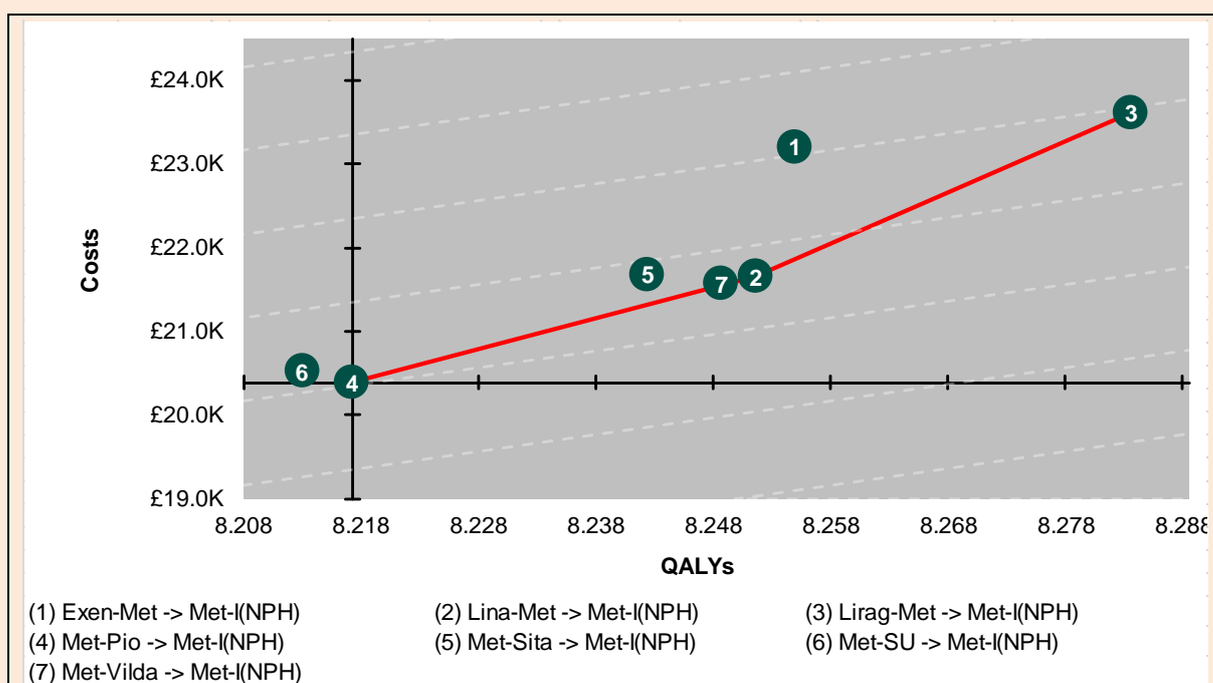
- 2 For first intensification, 7 treatments could be modelled – all the modelled combinations  
 3 contained metformin and none contained a meglitinide. People accrued an average of 16.3  
 4 undiscounted life years, of which 3.7 years were spent on first intensification therapy. As for  
 5 initial therapy, there was little difference in lifetime complication rates as the differences in  
 6 HbA1c treatment effects were even smaller.
- 7 People accumulated an average of 8.2 lifetime discounted QALYs, with most losses and  
 8 differences coming from weight profiles and some from hypoglycaemic episodes. Treatment-  
 9 related costs accounted for most variation in lifetime discounted costs.
- 10 First intensification therapy with metformin-pioglitazone was the cheapest and most cost-  
 11 effective treatment option (see table 73). All DPP4 inhibitor-metformin combinations  
 12 produced very similar lifetime discounted QALYs and costs and the GDG were happy to  
 13 consider the 3 combinations to be equivalent, particularly if people could not take metformin-  
 14 pioglitazone and metformin-sulfonylurea (see figure 50).

15 **Table 73: Mean lifetime incremental cost–utility results for first intensification therapy**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Metformin-pioglitazone -> Met-I(NPH)	£20,390	8.217			
Metformin-sulfonylurea -> Met-I(NPH)	£20,522	8.213	£132	-0.004	Dominated
Metformin-vildagliptin -> Met-I(NPH)	£21,569	8.249	£1179	0.031	Ext. dom.
Linagliptin-metformin -> Met-I(NPH)	£21,654	8.252	£1264	0.034	£36,788
Metformin-sitagliptin -> Met-I(NPH)	£21,685	8.243	£31	-0.009	Dominated
Exenatide-metformin -> Met-I(NPH)	£23,213	8.255	£1560	0.003	Ext. dom.
Liraglutide-metformin -> Met-I(NPH)	£23,614	8.284	£1960	0.032	£61,381

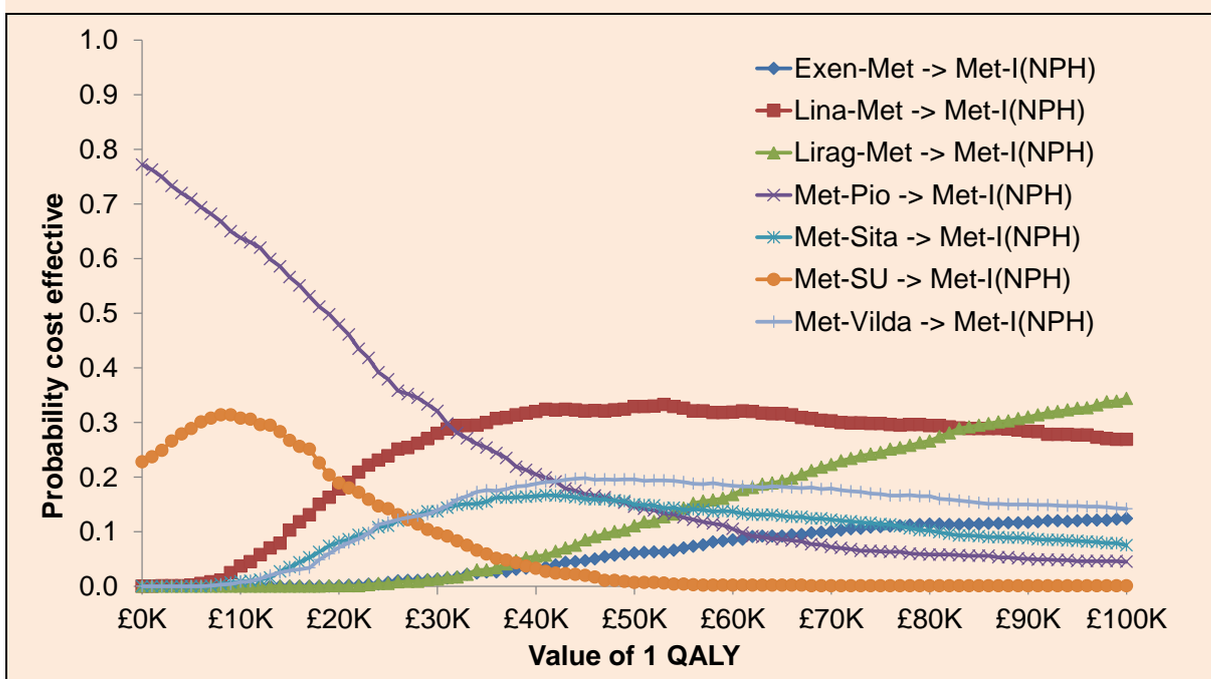
- 16 (a) *Met-I(NPH)* = Metformin-NPH insulin  
 17 (b) *Ext. dom.* = extendedly dominated

18



1 **Figure 50: Cost–utility plane for first intensification**

2 At first intensification, metformin-pioglitazone was the most cost-effective treatment  
3 combination at a maximum acceptable ICER of £20,000 per QALY in 48% of iterations (see  
4 figure 51). While metformin-pioglitazone and metformin-sulfonylurea showed only small  
5 incremental differences in the base case (see table 73), the superiority of metformin-  
6 pioglitazone was maintained in most probabilistic iterations.



7 **Figure 51: Cost-effectiveness acceptability curve for first intensification (class level)**

8

8.4.10.9 **Evidence statements for first intensification**

8.4.10.10 **Clinical evidence**

8.4.10.1.11 **Change in blood glucose**

12 Evidence from 4 network meta-analyses including data from 20, 22, 16 and 6 RCTs at 3, 6,  
13 12 and 24 months respectively for HbA1c levels showed that metformin-based combinations  
14 were generally associated with higher rankings at all 4 follow-up timepoints. At 3 and 6  
15 months, metformin combined with a GLP-1 mimetic (exenatide, liraglutide) was most  
16 effective in reducing HbA1c levels, while metformin combined with nateglinide or pioglitazone  
17 were shown to be most effective at 12 and 24 months respectively. In general, the credible  
18 intervals surrounding these ranking were considerably wide, except at 6 months where there  
19 was greater certainty in the data. The quality of the evidence was moderate.

#### 8.4.10.1.21 **Hypoglycaemia at study end point**

2 Evidence from a single network meta-analysis including data from 21 RCTs showed that  
3 sulfonylurea combined with metformin (median rank 12 [10 to 12]) or pioglitazone (median  
4 rank 11 [6 to 12]) were least effective in preventing hypoglycaemic events. Metformin-  
5 saxagliptin, metformin-lixisenatide and metformin-acarbose had the highest ranking  
6 suggesting these treatment combinations are effective in preventing hypoglycaemic events.  
7 The quality of the evidence was low.

#### 8.4.10.1.38 **Adverse events at study end point**

9 Evidence from 3 network meta-analyses including data from 27, 29 and 11 RCTs for  
10 dropouts due to adverse events, total dropouts and nausea respectively showed that in  
11 general, metformin combined with a GLP-1 mimetic (exenatide, liraglutide and lixisenatide) is  
12 less effective in preventing dropouts and nausea compared to metformin-sulfonylurea.  
13 Metformin combined with pioglitazone or sulfonylurea were shown to most effective at  
14 preventing nausea. There was generally some uncertainty around the results demonstrated  
15 by wide credible intervals which in the main crossed the line of no effect. The quality of the  
16 evidence was moderate to low.

#### 8.4.10.1.47 **Change in body weight**

18 Evidence from 2 network meta-analyses including data from 8 RCTs at 12 and 24 months  
19 showed that metformin combined with a DPP-4 inhibitor (linagliptin) or GLP-1 mimetic  
20 (exenatide and liraglutide) were most effective at promoting weight loss at 12 and 24 months.  
21 Whereas, pioglitazone combined with sitagliptin and metformin or sulfonylurea were  
22 associated with weight gain at 12 and 24 months respectively. The quality of the evidence  
23 was low.

#### 8.4.10.24 **Health economic evidence**

25 A directly applicable health economic model with potentially serious limitations found that a  
26 combination of metformin–pioglitazone was the most cost-effective modelled option for first  
27 intensification therapy.

#### 8.4.1128 **Evidence to recommendations for first intensification**

##### 29 **Table 74: Linking evidence to recommendations**

<p>Relative value of different outcomes</p>	<p>The following outcomes were considered critical to decision making; glycaemic control (change in HbA1c), hypoglycaemic events and adverse events. Change in body weight was considered important to decision making.</p> <p>The GDG noted that glycaemic control was important in mitigating the increased risk of microvascular and macrovascular complications associated with increased levels of hyperglycaemia, necessitating intensification of drug therapy. However, the GDG acknowledged that tight glycaemic control may be associated with an increased risk of hypoglycaemia, which may negatively impact on quality of life. Drug tolerability and change in body weight were considered important in determining the acceptability of treatment to the patient.</p> <p>The relative importance of each outcome is variable depending on several factors:</p> <ul style="list-style-type: none"> <li>• Short-term (3 and 6 months) versus long-term (12 and 24 months) evaluation. For example, adverse events and change in body</li> </ul>
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	<p>weight are reflected at longer time points (12 and 24 months).</p> <ul style="list-style-type: none"> <li>• Severity of hyperglycaemia.</li> <li>• Individual circumstances, such as comorbidities.</li> </ul>
<p>Trade-off between benefits and harms</p>	<p>The GDG acknowledged that there was generally less evidence at this treatment level, resulting in sparser networks. The GDG noted that there was greater uncertainty in the evidence at this intensification level as demonstrated by the wide credible intervals that surrounded many of the point estimates across all outcomes. Moreover, the GDG recognised that the current evidence base was biased towards metformin-based combinations such that of the 14 available treatment options, 3 did not include metformin (pioglitazone plus sitagliptin, pioglitazone plus sulfonylurea and sitagliptin plus sulfonylurea).</p> <p>Of the 14 treatment combinations, 7 included studies that reported data for all required outcomes in the health economic model. The 7 interventions that were not included in the main health economic model were 4 metformin-based combinations (metformin-acarbose, metformin-lixisenatide, metformin-nateglinide and metformin-saxagliptin) and the 3 aforementioned combinations that did not include metformin.</p> <p>The GDG considered the evidence surrounding the intensification options for individuals inadequately controlled by metformin alone.</p> <p>The GDG agreed that, while metformin combined with a DPP-4 inhibitor (linagliptin, saxagliptin sitagliptin or vildagliptin) was moderately effective in controlling blood glucose levels, this treatment combination was associated with less hypoglycaemic events and weight loss.</p> <p>The GDG discussed the evidence surrounding the use of metformin in combination with pioglitazone and noted that it was most effective at reducing HbA1c levels at 24 months and preventing nausea, but was associated with weight gain.</p> <p>The GDG discussed the long-term safety concerns associated with the use of pioglitazone and DPP-4 inhibitors, and agreed that MHRA guidance and patient suitability should be considered. For example, pioglitazone is not recommended for people with active bladder cancer, a history of bladder cancer, or uninvestigated haematuria, or people with heart failure or who are at risk of osteoporosis. The GDG noted that there was limited information on long-term safety of DPP-4 inhibitors.</p> <p>The GDG agreed that there was tentative evidence that metformin-sulfonylurea was moderately effective in reducing HbA1c levels, but this treatment combination was strongly associated with greater hypoglycaemic events. However, the GDG noted that the point estimates were associated with large credible intervals, indicating some measure of uncertainty around the data.</p> <p>The GDG recognised that there was evidence to indicate that metformin combined with a GLP-1 mimetic (exenatide or liraglutide) may be effective in reducing HbA1c levels in the short term (up to 6 months), preventing hypoglycaemic events and promoting weight loss. The GDG discussed the long-term safety risks associated with the use of GLP-1 mimetics and the evidence from the health</p>

	<p>economic model, which they thought were important in the decision making. The GDG considered that there was strong evidence from the health economic model that showed that this treatment combination was not cost effective and agreed not to recommend this option routinely.</p> <p>The GDG recognised that there was limited evidence for treatment intensification options for individuals where metformin is contraindicated or not tolerated. They noted that sitagliptin-sulfonylurea was associated with high rankings in change in HbA1c at 6 months, whereas pioglitazone-sulfonylurea was associated with weight gain at 24 months.</p>
<p>Consideration of health benefits and resource use</p>	<p>Economic model results showed a clustering of treatments, with all 3 modelled DPP-4 inhibitor-metformin combinations showing similar lifetime discounted costs and QALYs. In the base case, linagliptin-metformin produced an ICER of £36,800 per QALY compared with metformin-pioglitazone. In the PSA, metformin-pioglitazone had a 48% chance of being the most cost-effective treatment option.</p> <p>Metformin-pioglitazone and metformin-sulfonylurea showed similar lifetime discounted costs and QALYs. However, metformin-pioglitazone provided better value for money than metformin-sulfonylurea in most iterations of the PSA, meaning that, while the differences may be small, the superiority of metformin-pioglitazone appears to be a relatively robust finding.</p> <p>Given the differences in lifetime discounted costs were mainly in treatment costs, the GDG chose to recommend the DPP-4 inhibitor-metformin treatment options with the lowest DPP-4 inhibitor acquisition cost.</p> <p>Differences between drugs at first intensification were small, partly due to the normalising effect of future intensification in the economic model – patients were only on their first intensification therapies for an average of 3.7 years. QALY differences were driven by differences in weight gained; cost differences were predominantly due to the costs of the drugs themselves.</p> <p>The GDG noted that the economic model was not able to take account of the stopping rules from CG87 for GLP-1 combinations. The treatment effects for HbA1c and weight from the current guideline analysis were substantially less than those required by the CG87 stopping rules.</p> <p>The economic model did not provide any evidence for combinations that did not contain metformin. As the economic results were driven primarily by body weight and hypoglycaemia, the GDG felt it was highly unlikely combinations including pioglitazone and/or sulfonylurea for patients not taking metformin would appear cost effective compared with metformin-based combinations. However, it was unclear which combinations would be cost effective in a decision space that only contained non-metformin combinations. The GDG noted this would be a small subgroup of patients.</p>
<p>Quality of the evidence</p>	<p>The GDG agreed that the overall quality of the evidence for first intensification was moderate to low.</p>
<p>Other considerations</p>	<p>When defining the decision problem for this question, the GDG preferred not to make an <i>a priori</i> assumption of class effect across DPP-4 inhibitors. Therefore, each individual option for which</p>

evidence was available was analysed separately. Having reviewed the assembled evidence for each phase of treatment, the GDG noted that it was difficult to judge whether the different DPP-4 inhibitors could, in fact, be considered interchangeable:

- In a few areas, a case could be made for the superiority of one option over another (for example, as initial therapy, sitagliptin seemed to have somewhat superior benefits to vildagliptin at similar net costs).
- In other areas, all the DPP-4 inhibitors for which evidence was available appeared to have very similar benefits, harms and costs (for example, in combination with metformin at first intensification).
- Elsewhere in the treatment pathway, evidence was extremely limited (for example, sitagliptin-metformin-sulfonylurea was the only treatment combination for which evidence was available at second intensification) or absent (for example, at first intensification, there was no evidence that could be used to assess the relative clinical effectiveness and cost effectiveness of DPP-4 inhibitors in combination with pioglitazone or sulfonylureas).

Having considered these different situations, the GDG concluded that the most helpful recommendations would be ones that treated DPP-4 inhibitors as a class. Had it been presented with evidence that suggested that 1 or more of the options was superior to others across all phases of treatment, the GDG would clearly have been inclined to favour such option(s) in its recommendations. However, the picture that had emerged was much more sporadic, and the Group was not confident that any apparent dissimilarities between options represented real differences that would be expected in clinical practice. Moreover, the GDG was mindful that a series of recommendations that alternated between treating DPP-4 inhibitors as a class, in some parts of the treatment pathway, and focusing on individual options in others would be confusing to readers of the guideline, even if those recommendations could be directly allied with the available evidence. For all of these reasons, the GDG took the view that recommendations should consistently refer to DPP-4 inhibitors as a class. It was a natural extension of this principle that prescribers should be encouraged to select the individual DPP-4 inhibitor with the lowest acquisition cost available to them.

The GDG noted that the mean age in the included studies was about 57 years and agreed that these trials are biased towards younger and fitter participants, who are less likely to experience significant comorbidities than the majority of people with type 2 diabetes seen in clinical practice. The GDG thought that the treatment effects observed in trials are likely to generalise to a population facing more comorbidities and other challenges to effective management of their disease. However, the Group agreed that the balance of benefits and harms may be different in such cases, and there are specific issues based on clinical experience which may require particular attention that should be highlighted in the recommendations.

It was noted that reporting of hypoglycaemia differed across the included studies. All categories of hypoglycaemia (for example, confirmed hypoglycaemia) were generally a subset of 'any hypoglycaemia', which was the most commonly reported category of hypoglycaemia across the included studies. The GDG discussed the risk of bias associated with reported hypoglycaemia and noted

that self-reported hypoglycaemia may not be a reliable measure as an individual's perception of hypoglycaemia is variable at different glucose levels.

The GDG noted that the results from the sensitivity analyses of individuals who had previous experience of using 1 oral antidiabetic medicine were similar to the full dataset which included studies of mixed populations of people who were drug naïve, or on 1 or more oral antidiabetic medicines at screening.

#### 8.4.12.1 Clinical evidence review for second intensification

2 In total 17,037 references were found for the main review question and 45 papers were  
3 included for second intensification which relate to 42 trials.

4 This review question addressed which treatment combination is most effective when people  
5 with type 2 diabetes who are treated with diet and a combination of 2 non-insulin based  
6 therapies have inadequate blood glucose control. The GDG agreed that both triple non-  
7 insulin based therapies and insulin based medicines are potential treatment options at  
8 second intensification. Due to the large volume of evidence relating to insulin therapy, the  
9 GDG prioritised the drug comparisons listed in Table 43 for second intensification, which  
10 were of particular clinical interest:

- 11 • 3 non-insulin based therapies versus 3 non-insulin based therapies
- 12 • Insulin versus 3 non-insulin based therapies
- 13 • Insulin + 1 non-insulin based therapy versus 3 non-insulin based therapies
- 14 • Insulin + 2 non-insulin based therapies versus 3 non-insulin based therapies
- 15 • Insulin versus insulin + 1 non-insulin based therapy
- 16 • Insulin versus insulin + 2 non-insulin based therapies
- 17 • Insulin + 1 non-insulin based therapy versus insulin + 1 non-insulin based therapy
- 18 • Insulin + 2 non-insulin based therapies versus insulin + 2 non-insulin based therapies
- 19 • Insulin + 1 non-insulin based therapy versus insulin + 2 non-insulin based therapies

20 RCTs of at least 12 week treatment duration examining the drug comparisons above were  
21 included. In contrast to initial therapy, it was assumed that most patients would be titrated to  
22 the maximal tolerated doses of previous oral therapy before starting a trial. Therefore, trials  
23 that did not report specific doses of continued previous therapy were still included (see  
24 section 8.4.2 for the main exclusion criteria).

#### 8.4.12.25 Description of included studies for second intensification

26 A total of 10,170 participants from 39 RCTs were included. The majority of studies were  
27 carried out in multiple centres across different countries. The mean age ranged from 52.6 to  
28 64.8 years. Mean HbA1c levels at baseline ranged from 62 to 97 mmol/mol (7.8% to 11%).  
29 The mean BMI ranged from 24.7 to 36.08 kg/m<sup>2</sup>. Mean duration of diabetes ranged from 3.5  
30 to 13.7 years, with 2 studies not reporting this information. Follow-up periods ranged from 12  
31 weeks to 104 weeks. For full details of the included studies, see Appendix E.

#### 8.4.12.22 Network meta-analyses for second intensification

33 To facilitate comparison across all available treatment options, 6 network meta-analyses  
34 were performed for all 3 critical and 1 important outcomes – change in HbA1c up to 12  
35 months, hypoglycaemia at study end point, adverse events (that is, dropouts due to adverse  
36 events, total dropouts and nausea) at study end point and change in body weight up to 12  
37 months. Where available, metformin-neutral protamine Hagedorn (NPH) insulin was selected

1 as the reference treatment option as this combination was considered to reflect current  
 2 standard clinical practice. For nausea only, metformin-biphasic insulin aspart was used as  
 3 the reference treatment as no studies included metformin-NPH insulin. Full details of  
 4 methods and additional NMA outputs are provided in Appendix J.

5 For continuous outcomes, measurements up to 1 year follow-up were pooled together. This  
 6 is related to the way in which HbA1c levels varies as type 2 diabetes progresses.  
 7 Specifically, although initial reductions in HbA1c levels are observed following treatment,  
 8 these levels will eventually drift back up over time. Further exploration of the included HbA1c  
 9 data showed that there was little difference between measurements at 6 months and 12  
 10 months. Furthermore, as 3-month measurements were likely to be more conservative (that  
 11 is, not bias in favour of the intervention) due to the J-shaped curve, pooling of these  
 12 timepoints was considered appropriate. Where included trials reported more than one  
 13 timepoint between 12 weeks and 1 year, only the latest timepoint was included in synthesis.  
 14 A sparse but relatively connected network was produced for change in HbA1c levels. Only 1  
 15 trial (Gram et al. 2011, Holman et al. 1999) reported outcomes at 2 years follow-up and over.  
 16 These results have not been presented because this did not form a network.

17 On the whole, the quality of the evidence was low as many of the connections were limited to  
 18 single trials, the majority of studies were open label and some included RCTs may not have  
 19 been representative of UK clinical population with type 2 diabetes who require second  
 20 intensification of drug therapy. It was noted that random-effects models tended to estimate a  
 21 fairly large inter-study heterogeneity term, which will reduce the precision of effect estimates.

22 **Table 75: GRADE profile for network meta-analyses for second intensification**

Assessment time points/ Measure	Number of RCTs	Risk of bias	Inconsistency	Indirectness	Imprecision	Quality
<b>Change in blood glucose (HbA1c)</b>						
Up to 12 months	37	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	not serious	Moderate
<b>Hypoglycaemia at study end point</b>						
Study end point	34	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<b>Adverse events at study end point</b>						
Dropouts due to adverse events	25	serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low <sup>6</sup>
Total dropouts	25	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
Nausea	4	serious <sup>1</sup>	serious <sup>5</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low <sup>6</sup>
<b>Change in body weight</b>						
Up to 12 months	27	serious <sup>1</sup>	not serious <sup>2</sup>	not serious <sup>3</sup>	serious <sup>4</sup>	Low
<sup>1</sup> Downgrade 1 level: baseline HbA1c ranged from 7.8 to 11%						
<sup>2</sup> Assessed based on residual deviance, deviance information criterion and $\tau^2$ ( $\tau^2 < 0.5$ )						
<sup>3</sup> Considered not serious as population, interventions, comparator and outcomes are as defined in protocol						
<sup>4</sup> Downgrade 1 level: no interventions had probability of being best and worse $\geq 0.5$						
<sup>5</sup> Downgrade 1 level: $\tau^2 \geq 0.5$						
<sup>6</sup> Maximum downgrade by 2 levels						

Update 2015

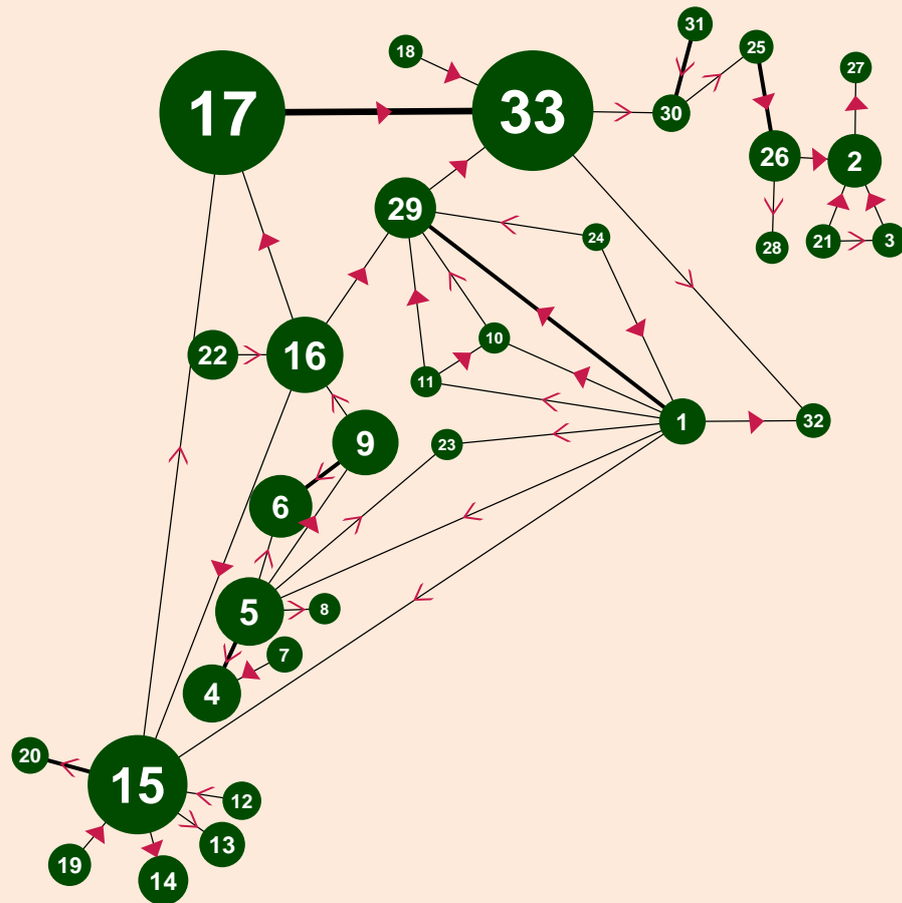
#### 8.4.12.23 Change in blood glucose (HbA1c) up to 12 months

24 Results of the NMA are summarised below for the 32 treatment combinations that were  
 25 compared with metformin-NPH insulin up to 12 months. Of the 32 treatment combinations, 6

1 were 3 non-insulin based drug combinations, 6 were insulin only, 16 were insulin + 1 non-  
2 insulin based drug combinations and 4 were insulin + 2 non-insulin based drug combinations.

3 Where available, there is reasonable agreement between the NMA evidence and direct  
4 pairwise treatment effect estimates as demonstrated by the substantial overlap between the  
5 credible/confidence intervals. Overall, credible intervals crossed the line of no effect.

6 However, in general, compared to metformin-NPH insulin, 3 non-insulin based drug  
7 combinations, insulin only and insulin + 1 non-insulin based drug were shown to be less  
8 effective in reducing HbA1c levels. Of the 4 insulin + 2 non-insulin based drug combinations,  
9 only NPH insulin-metformin-repaglinide were shown to be more effective in reducing HbA1c  
10 levels than metformin-NPH insulin. This treatment combination had the highest ranking  
11 (median rank 1 [95% credible interval 1 to 7]), whereas metformin-repaglinide-sulfonylurea  
12 had the lowest ranking (median rank 33 [26 to 33]). The combination with the second highest  
13 ranking was biphasic insulin aspart-pioglitazone (median rank 2 [1 to 15]).

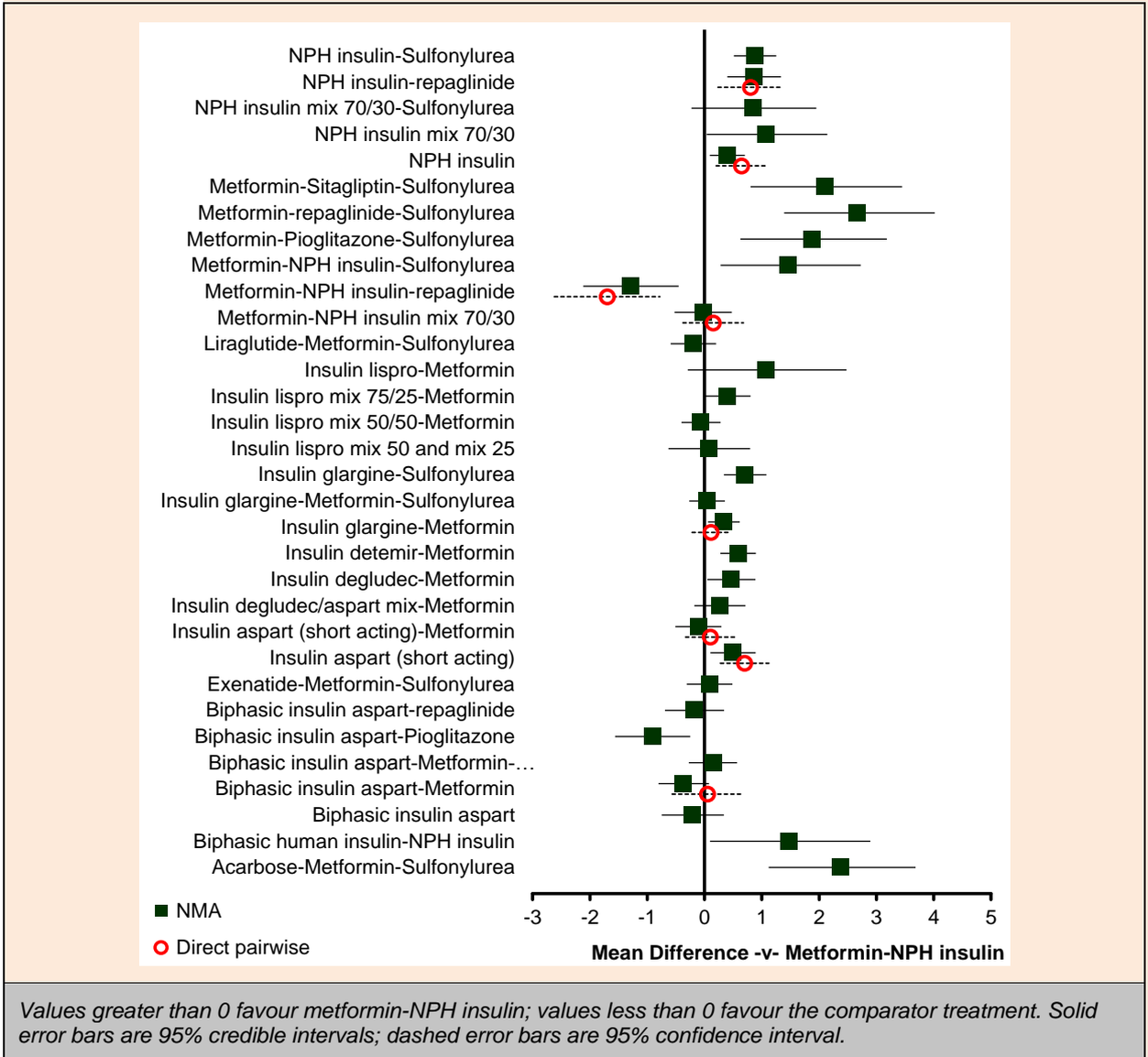


1 Metformin-NPH insulin	17 Insulin glargine-Sulfonylurea
2 Acarbose-Metformin-Sulfonylurea	18 Insulin lispro mix 50 and mix 25
3 Biphasic human insulin-NPH insulin	19 Insulin lispro mix 50/50-Metformin
4 Biphasic insulin aspart	20 Insulin lispro mix 75/25-Metformin
5 Biphasic insulin aspart-Metformin	21 Insulin lispro-Metformin
6 Biphasic insulin aspart-Metformin-Sulfonylurea	22 Liraglutide-Metformin-Sulfonylurea
7 Biphasic insulin aspart-Pioglitazone	23 Metformin-NPH insulin mix 70/30
8 Biphasic insulin aspart-repaglinide	24 Metformin-NPH insulin-repaglinide
9 Exenatide-Metformin-Sulfonylurea	25 Metformin-NPH insulin-Sulfonylurea
10 Insulin aspart (short acting)	26 Metformin-Pioglitazone-Sulfonylurea
11 Insulin aspart (short acting)-Metformin	27 Metformin-repaglinide-Sulfonylurea
12 Insulin degludec/aspart mix-Metformin	28 Metformin-Sitagliptin-Sulfonylurea
13 Insulin degludec-Metformin	29 NPH insulin
14 Insulin detemir-Metformin	30 NPH insulin mix 70/30
15 Insulin glargine-Metformin	31 NPH insulin mix 70/30-Sulfonylurea
16 Insulin glargine-Metformin-Sulfonylurea	32 NPH insulin-repaglinide
	33 NPH insulin-Sulfonylurea

Size of nodes is proportional to total number of participants randomised to receive the treatment in question across the evidence-base. Width of connecting lines is proportional to number of trial-level comparisons available. Arrowheads indicate direction of effect in pairwise data ( $a > b$  denotes  $a$  is more effective than  $b$ ) – filled arrowheads show comparisons where one option is significantly superior ( $p < 0.05$ ); outlined arrowheads show direction of trend where effect does not reach statistical significance.

1 **Figure 52: Network meta-analysis of change in HbA1c (up to 12 months) – evidence**  
2 **network**

1



Update 2015

2 **Figure 53: Network meta-analysis of change in HbA1c (up to 12 months) – relative**  
 3 **effect of all options compared with common comparator (metformin-NPH**  
 4 **insulin)**

1 **Table 76: Network meta-analysis of change in HbA1c (up to 12 months) – rankings for**  
2 **each comparator**

	Probability best	Median rank (95%CrI)
Metformin-NPH insulin	0.000	10 (5, 15)
Acarbose-Metformin-Sulfonylurea	0.000	32 (31, 32)
Biphasic human insulin-NPH insulin	0.000	28 (14, 30)
Biphasic insulin aspart	0.000	6 (3, 16)
Biphasic insulin aspart-Metformin	0.000	4 (3, 9)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.000	13 (7, 21)
Biphasic insulin aspart-Pioglitazone	0.236	2 (1, 3)
Biphasic insulin aspart-repaglinide	0.000	6 (3, 16)
Exenatide-Metformin-Sulfonylurea	0.000	12 (6, 18)
Insulin aspart (short acting)	0.000	20 (12, 27)
Insulin aspart (short acting)-Metformin	0.000	8 (3, 16)
Insulin degludec/aspart mix-Metformin	0.000	15 (7, 23)
Insulin degludec-Metformin	0.000	19 (12, 26)
Insulin detemir-Metformin	0.000	21 (17, 27)
Insulin glargine-Metformin	0.000	17 (12, 21)
Insulin glargine-Metformin-Sulfonylurea	0.000	11 (6, 16)
Insulin glargine-Sulfonylurea	0.000	22 (18, 28)
Insulin lispro mix 50 and mix 25	0.001	12 (3, 23)
Insulin lispro mix 50/50-Metformin	0.000	8 (3, 15)
Insulin lispro mix 75/25-Metformin	0.000	18 (10, 25)
Insulin lispro-Metformin	0.001	26 (5, 28)
Liraglutide-Metformin-Sulfonylurea	0.000	6 (3, 12)
Metformin-NPH insulin mix 70/30	0.000	9 (3, 20)
Metformin-NPH insulin-repaglinide	0.761	1 (1, 3)
Metformin-NPH insulin-Sulfonylurea	0.000	28 (19, 29)
Metformin-Pioglitazone-Sulfonylurea	0.000	30 (26, 31)
Metformin-repaglinide-Sulfonylurea	0.000	33 (32, 33)
Metformin-Sitagliptin-Sulfonylurea	0.000	31 (29, 32)
NPH insulin	0.000	18 (13, 23)
NPH insulin mix 70/30	0.000	26 (13, 29)
NPH insulin mix 70/30-Sulfonylurea	0.000	24 (6, 27)
NPH insulin-repaglinide	0.000	24 (18, 31)
NPH insulin-Sulfonylurea	0.000	24 (22, 30)

Update 2015

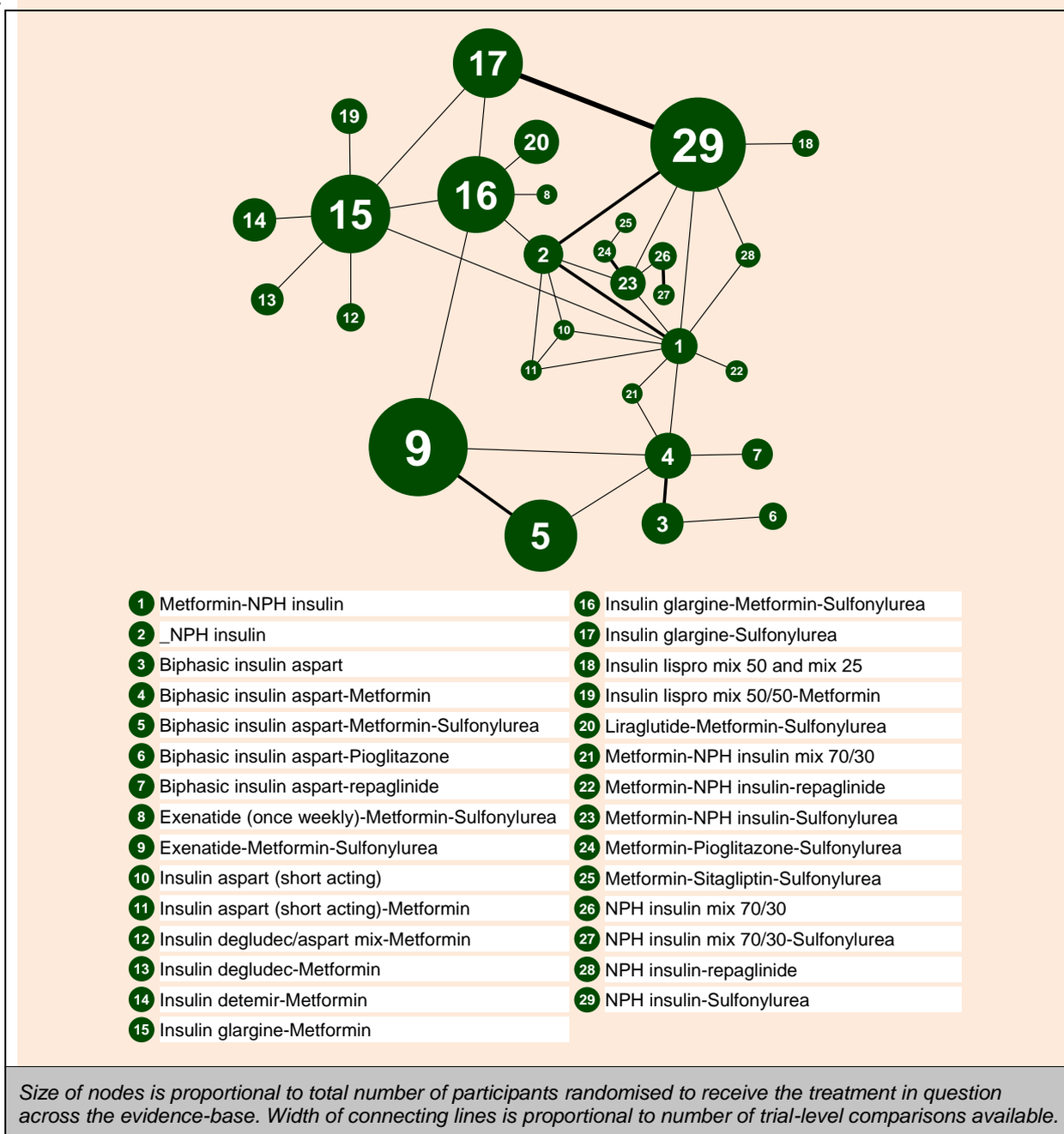
#### 8.4.12.43 Hypoglycaemia at study end point

- 4 Results of the NMA are summarised below for the 28 treatment combinations that were  
5 compared with metformin-NPH insulin. Of the 28 treatment combinations, 5 were 3 non-  
6 insulin based drug combinations, 5 were insulin only, 14 were insulin + 1 non-insulin based  
7 drug combinations and 4 were insulin + 2 non-insulin based drug combinations.
- 8 There is reasonable agreement between the NMA evidence and direct pairwise treatment  
9 effect estimates as demonstrated by the substantial overlap between the credible/confidence  
10 intervals. In the main, credible intervals were wide and crossed the line of no effect.

1 However, in general, compared to metformin-NPH insulin, insulin only and insulin + 2 non-  
 2 insulin based drug combinations were shown to be associated with greater hypoglycaemic  
 3 events, whereas, 3 non-insulin based drug combinations were generally associated with less  
 4 hypoglycaemic events. Insulin + 1 non-insulin based drug combination were generally  
 5 associated with greater hypoglycaemic events when compared to metformin-NPH insulin  
 6 except for metformin combined with insulin glargine, detemir or degludec and NPH-insulin  
 7 combined with repaglinide.

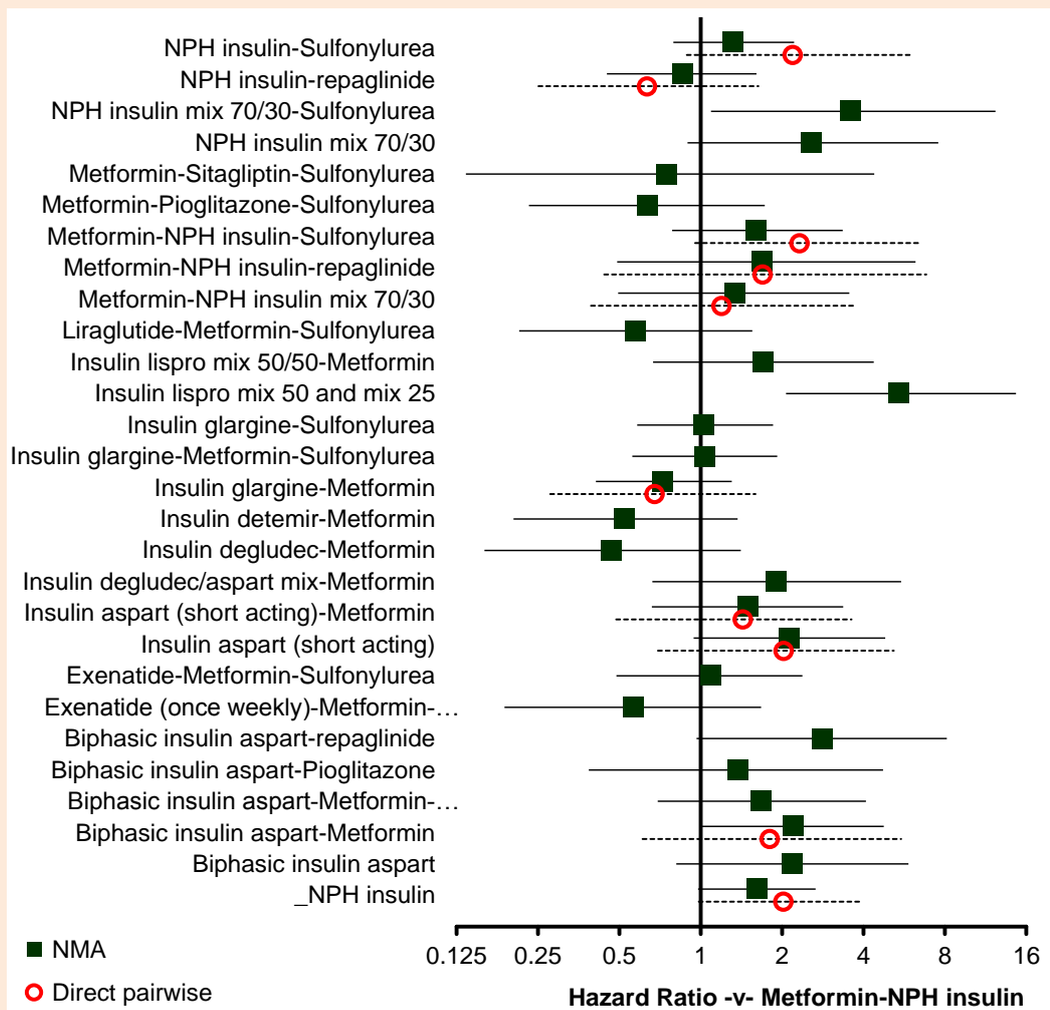
8 The treatment combinations with the highest ranking were metformin-insulin degludec  
 9 (median rank 3 [95% credible interval 1 to 16]) and metformin-insulin detemir (median rank 3  
 10 [1 to 15]) though the associated credible intervals were wide. Insulin lispro mix 50 and mix 25  
 11 was associated with the lowest ranking (median rank 29 [22 to 29]).

12



13 **Figure 54: Network meta-analysis of hypoglycaemic events (study end point) –**  
 14 **evidence network**

1



Values greater than 1 favour metformin-NPH insulin; values less than 1 favour the comparator treatment. Direct pairwise evidence is drawn from inconsistency model. Solid and dashed error bars are 95% credible intervals.

2 **Figure 55: Network meta-analysis of hypoglycaemic events (study end point) –**  
 3 **relative effect of all options compared with common comparator (metformin-**  
 4 **NPH insulin)**

5

Update 2015

1 **Table 77: Network meta-analysis of hypoglycaemic events (study end point) – rankings**  
2 **for each comparator**

	Probability best	Median rank (95%CrI)
Metformin-NPH insulin	0.000	10 (5, 17)
Biphasic insulin aspart	0.000	23 (9, 28)
Biphasic insulin aspart-Metformin	0.000	23 (13, 27)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.000	19 (8, 27)
Biphasic insulin aspart-Pioglitazone	0.015	16 (2, 27)
Biphasic insulin aspart-repaglinide	0.000	26 (12, 29)
Exenatide (once weekly)-Metformin-Sulfonylurea	0.156	4 (1, 17)
Exenatide-Metformin-Sulfonylurea	0.001	12 (4, 21)
Insulin aspart (short acting)	0.000	22 (10, 28)
Insulin aspart (short acting)-Metformin	0.001	17 (5, 27)
Insulin degludec/aspart mix-Metformin	0.000	21 (7, 28)
Insulin degludec-Metformin	0.284	3 (1, 16)
Insulin detemir-Metformin	0.157	3 (1, 15)
Insulin glargine-Metformin	0.001	6 (3, 14)
Insulin glargine-Metformin-Sulfonylurea	0.000	11 (5, 19)
Insulin glargine-Sulfonylurea	0.000	11 (5, 19)
Insulin lispro mix 50 and mix 25	0.000	29 (22, 29)
Insulin lispro mix 50/50-Metformin	0.000	19 (7, 28)
Liraglutide-Metformin-Sulfonylurea	0.120	4 (1, 16)
Metformin-NPH insulin mix 70/30	0.005	15 (3, 26)
Metformin-NPH insulin-repaglinide	0.009	19 (3, 29)
Metformin-NPH insulin-Sulfonylurea	0.000	18 (8, 26)
Metformin-Pioglitazone-Sulfonylurea	0.075	5 (1, 18)
Metformin-Sitagliptin-Sulfonylurea	0.164	7 (1, 27)
NPH insulin	0.000	19 (12, 25)
NPH insulin mix 70/30	0.000	24 (10, 28)
NPH insulin mix 70/30-Sulfonylurea	0.000	27 (14, 29)
NPH insulin-repaglinide	0.009	8 (2, 19)
NPH insulin-Sulfonylurea	0.000	15 (9, 22)

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#### 8.4.12.53 Adverse events at study end point

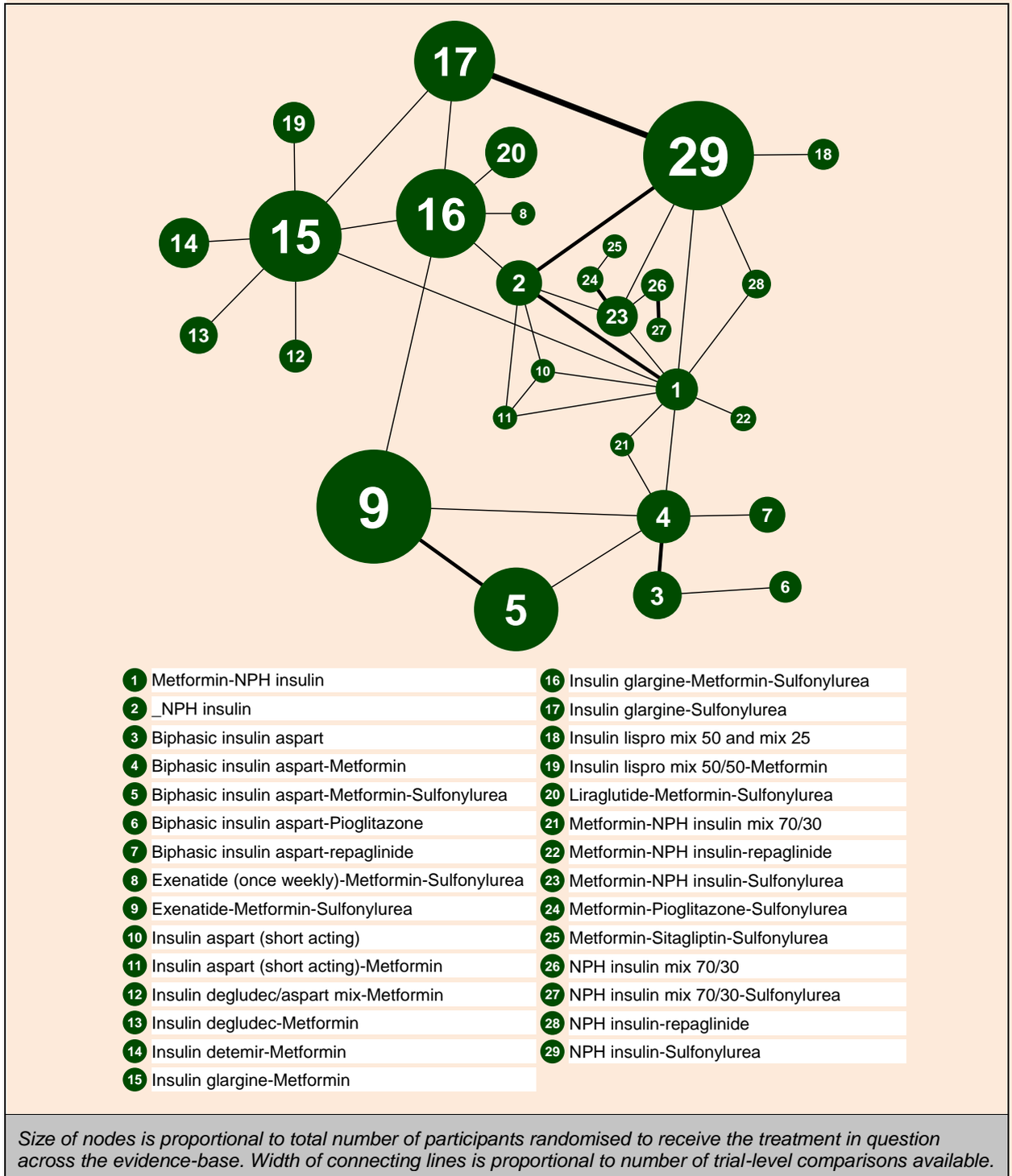
4 Results of the 3 NMAs are summarised below. For dropouts due to adverse events and total  
5 dropouts, 26 and 25 treatment combinations were compared with metformin-NPH insulin  
6 respectively, while 4 treatment combinations were compared with metformin-biphasic insulin  
7 aspart for nausea.

8 In general, there is reasonable agreement between the NMA evidence and direct pairwise  
9 treatment effect estimates, with substantial overlap between the credible/confidence  
10 intervals. However, there is substantial uncertainty in the data as the relative estimates are  
11 associated with considerably wide credible intervals with all crossing the line of no effect.

12 Insulin lispro mix 50 and mix 25 had the highest ranking (median rank 4 [1 to 19]) for  
13 dropouts due to adverse events, whereas a 3 non-insulin based drug combination  
14 (metformin-repaglinide-sulfonylurea) had the highest ranking for total dropouts (median rank  
15 1 [1 to 23]) and 3 of the insulin combinations shared the highest ranking for nausea;

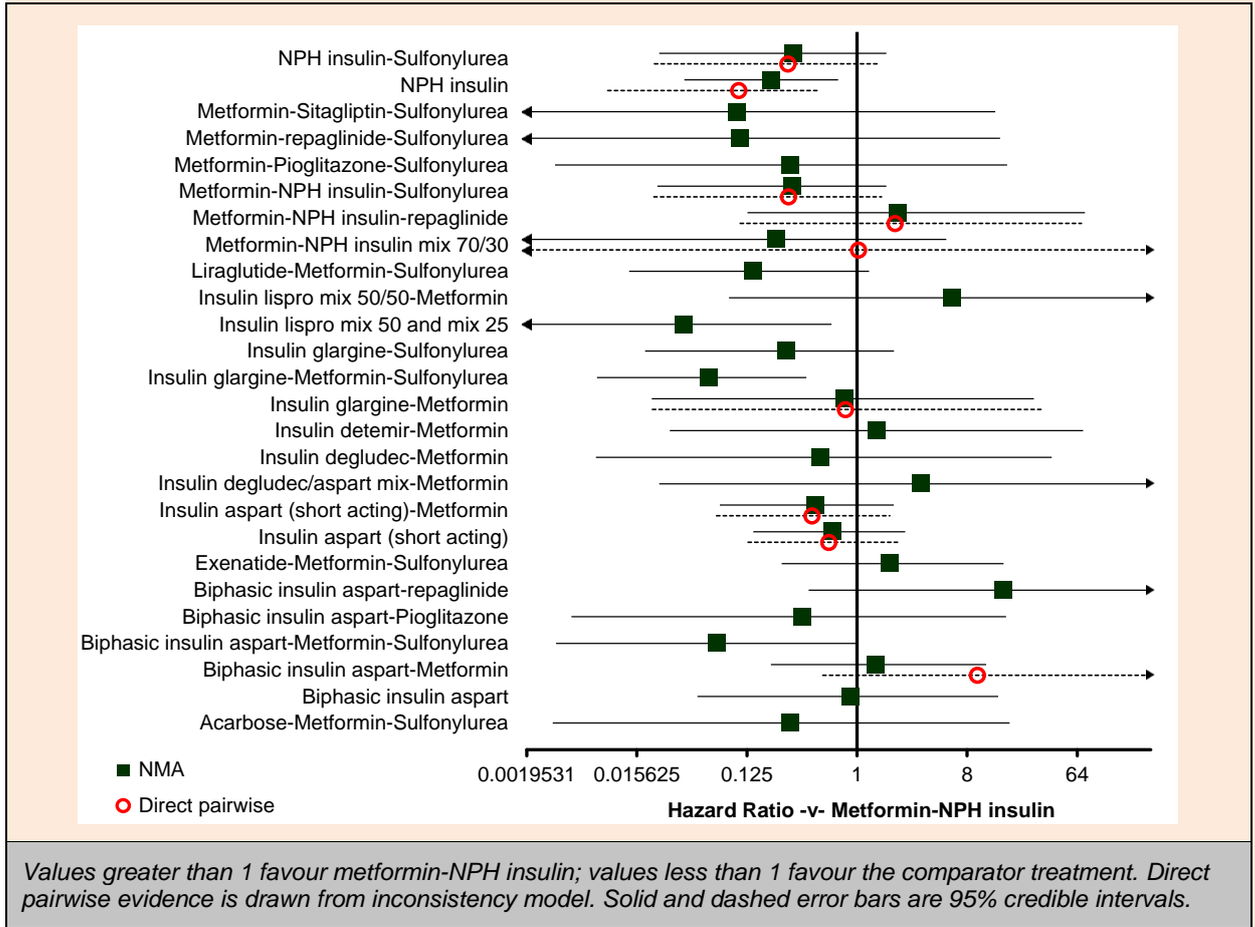
- 1 metformin-biphasic insulin aspart (median rank 2 [1 to 5]), metformin-sulfonylurea-biphasic  
2 insulin aspart (median rank 2 [1 to 5]) and metformin-sulfonylurea-insulin glargine (median  
3 rank 2 [1 to 5]).
- 4 Biphasic insulin aspart-repaglinide (median rank 26 [12 to 27]) and insulin aspart (short  
5 acting) (median rank 25 [13 to 26]) were ranked lowest for dropouts due to adverse events  
6 and total dropouts respectively.

7



8 **Figure 56: Network meta-analysis of dropouts due to adverse events (study end**  
9 **point) – evidence network**

1



2

**Figure 57: Network meta-analysis of dropouts due to adverse events (study end point) – relative effect of all options compared with common comparator (metformin-NPH insulin)**

3

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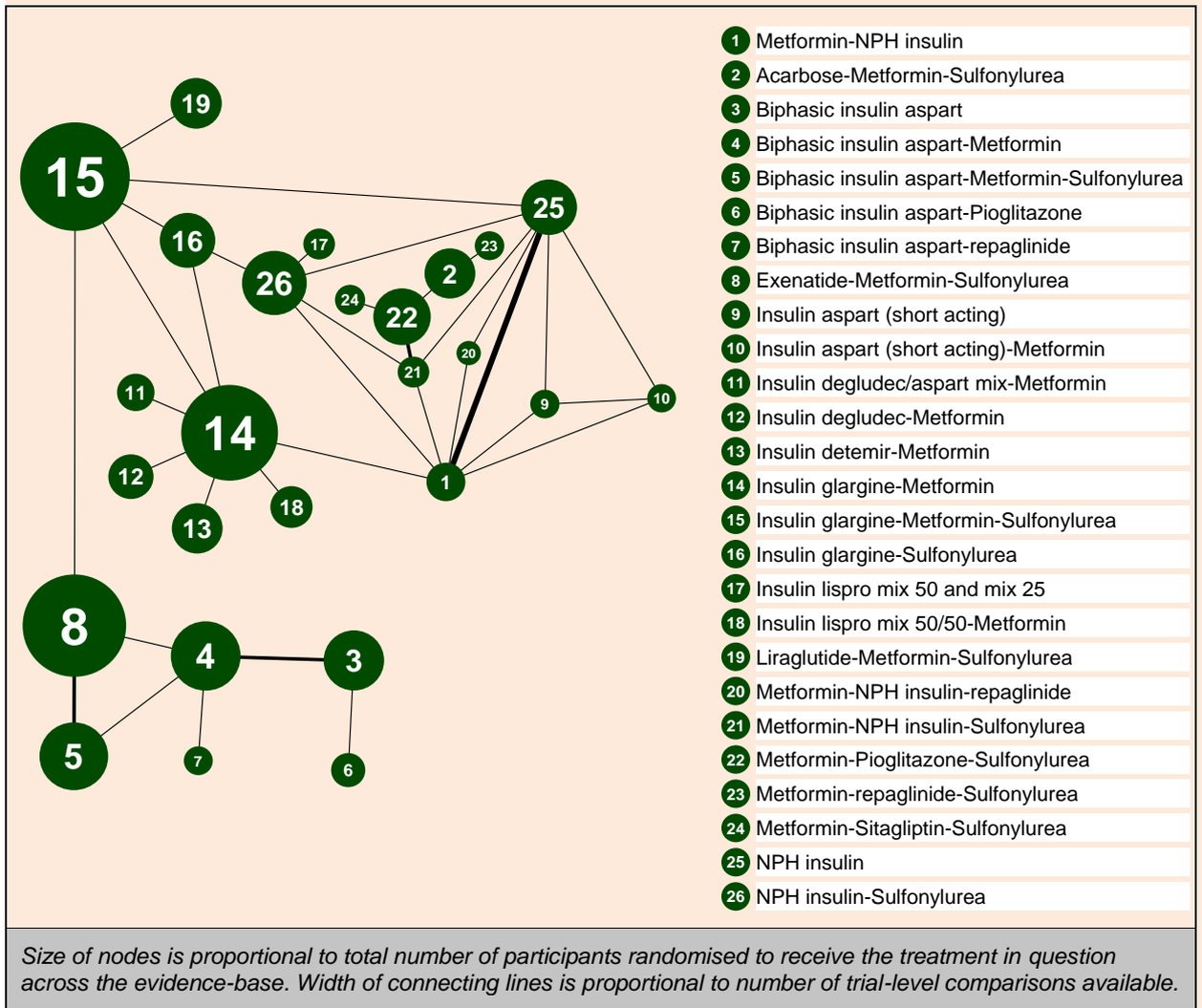
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Update 2015

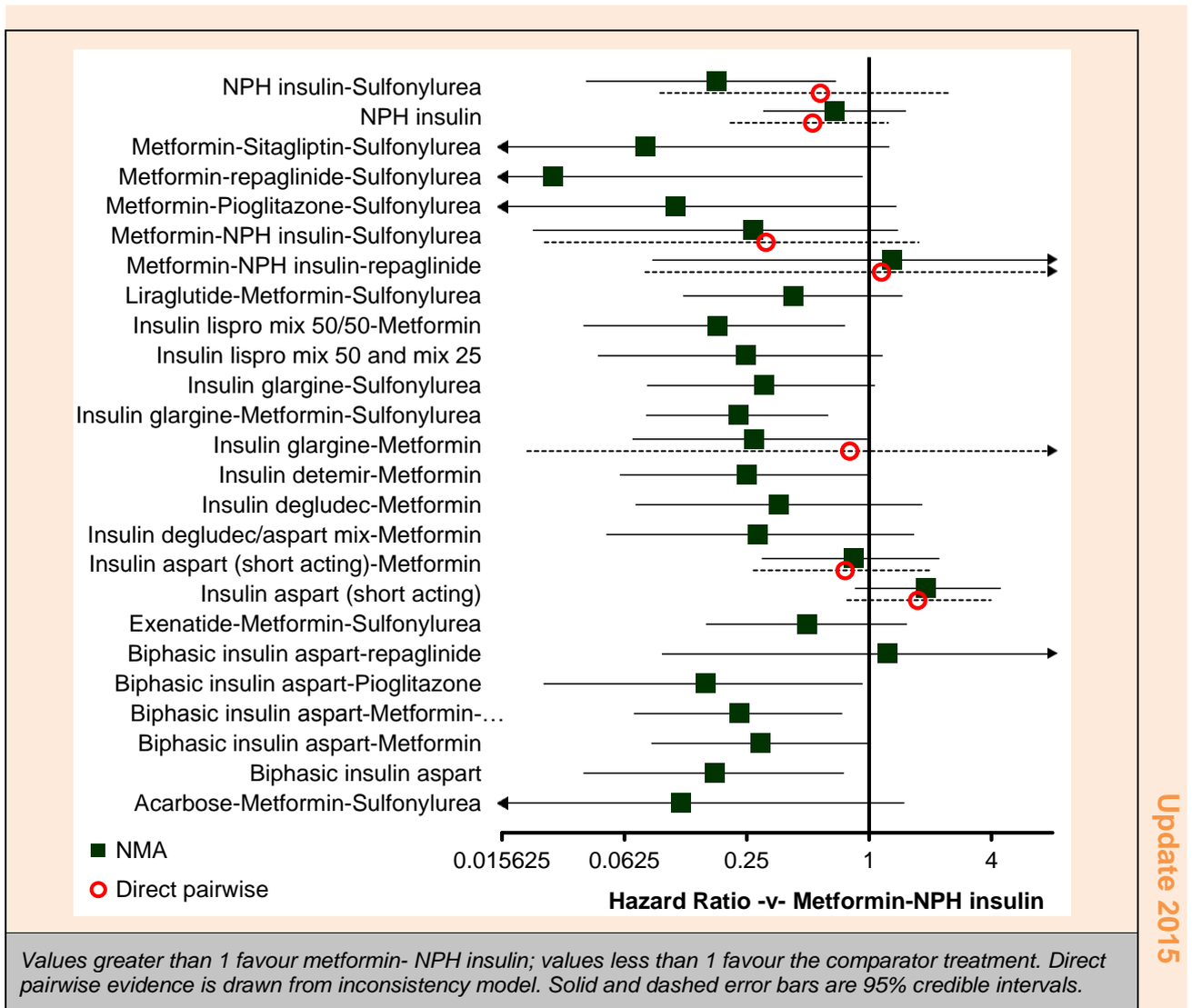
1 **Table 78: Network meta-analysis of dropouts due to adverse events (study end point) –**  
2 **rankings for each comparator**

	Probability best	Median rank (95%CrI)
Metformin-NPH insulin	0.000	19 (12, 24)
Acarbose-Metformin-Sulfonylurea	0.004	12 (2, 26)
Biphasic insulin aspart	0.000	18 (5, 26)
Biphasic insulin aspart-Metformin	0.000	20 (10, 25)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.114	5 (1, 17)
Biphasic insulin aspart-Pioglitazone	0.050	13 (1, 26)
Biphasic insulin aspart-repaglinide	0.000	26 (14, 27)
Exenatide-Metformin-Sulfonylurea	0.000	22 (12, 26)
Insulin aspart (short acting)	0.000	16 (6, 24)
Insulin aspart (short acting)-Metformin	0.003	14 (4, 23)
Insulin degludec/aspart mix-Metformin	0.008	23 (3, 27)
Insulin degludec-Metformin	0.036	14 (1, 26)
Insulin detemir-Metformin	0.003	20 (4, 26)
Insulin glargine-Metformin	0.004	17 (3, 24)
Insulin glargine-Metformin-Sulfonylurea	0.082	4 (1, 12)
Insulin glargine-Sulfonylurea	0.003	11 (2, 22)
Insulin lispro mix 50 and mix 25	0.268	3 (1, 15)
Insulin lispro mix 50/50-Metformin	0.000	25 (8, 27)
Liraglutide-Metformin-Sulfonylurea	0.006	8 (2, 19)
Metformin-NPH insulin mix 70/30	0.119	10 (1, 25)
Metformin-NPH insulin-repaglinide	0.002	22 (7, 27)
Metformin-NPH insulin-Sulfonylurea	0.005	12 (3, 21)
Metformin-Pioglitazone-Sulfonylurea	0.003	12 (3, 26)
Metformin-repaglinide-Sulfonylurea	0.146	7 (1, 26)
Metformin-Sitagliptin-Sulfonylurea	0.143	6 (1, 25)
NPH insulin	0.001	9 (3, 17)
NPH insulin-Sulfonylurea	0.000	12 (3, 22)

3



1 **Figure 58: Network meta-analysis of total dropouts (study end point) – evidence**  
 2 **network**  
 3



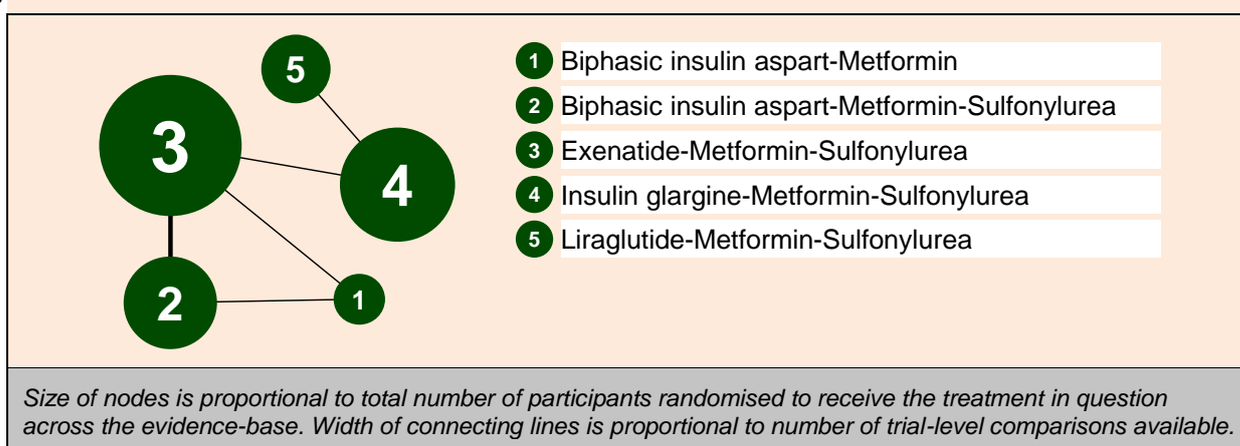
Update 2015

1 **Figure 59: Network meta-analysis of total dropouts (study end point) – relative**  
 2 **effect of all options compared with common comparator (metformin-NPH**  
 3 **insulin)**

1 **Table 79: Network meta-analysis of total dropouts (study end point) – rankings for**  
2 **each comparator**

	Probability best	Median rank (95%CrI)
Metformin-NPH insulin	0.000	23 (17, 25)
Acarbose-Metformin-Sulfonylurea	0.009	5 (2, 24)
Biphasic insulin aspart	0.025	8 (2, 18)
Biphasic insulin aspart-Metformin	0.000	14 (5, 20)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.003	10 (3, 18)
Biphasic insulin aspart-Pioglitazone	0.081	7 (1, 21)
Biphasic insulin aspart-repaglinide	0.005	24 (5, 26)
Exenatide-Metformin-Sulfonylurea	0.000	19 (11, 24)
Insulin aspart (short acting)	0.000	25 (22, 26)
Insulin aspart (short acting)-Metformin	0.000	22 (11, 25)
Insulin degludec/aspart mix-Metformin	0.014	13 (2, 24)
Insulin degludec-Metformin	0.003	16 (4, 25)
Insulin detemir-Metformin	0.003	11 (3, 21)
Insulin glargine-Metformin	0.000	13 (4, 20)
Insulin glargine-Metformin-Sulfonylurea	0.002	10 (4, 17)
Insulin glargine-Sulfonylurea	0.001	14 (5, 22)
Insulin lispro mix 50 and mix 25	0.015	11 (2, 23)
Insulin lispro mix 50/50-Metformin	0.038	8 (1, 18)
Liraglutide-Metformin-Sulfonylurea	0.000	17 (7, 24)
Metformin-NPH insulin-repaglinide	0.010	24 (3, 26)
Metformin-NPH insulin-Sulfonylurea	0.005	12 (3, 24)
Metformin-Pioglitazone-Sulfonylurea	0.013	4 (2, 23)
Metformin-repaglinide-Sulfonylurea	0.610	1 (1, 21)
Metformin-Sitagliptin-Sulfonylurea	0.130	3 (1, 22)
NPH insulin	0.000	21 (14, 24)
NPH insulin-Sulfonylurea	0.033	8 (1, 18)

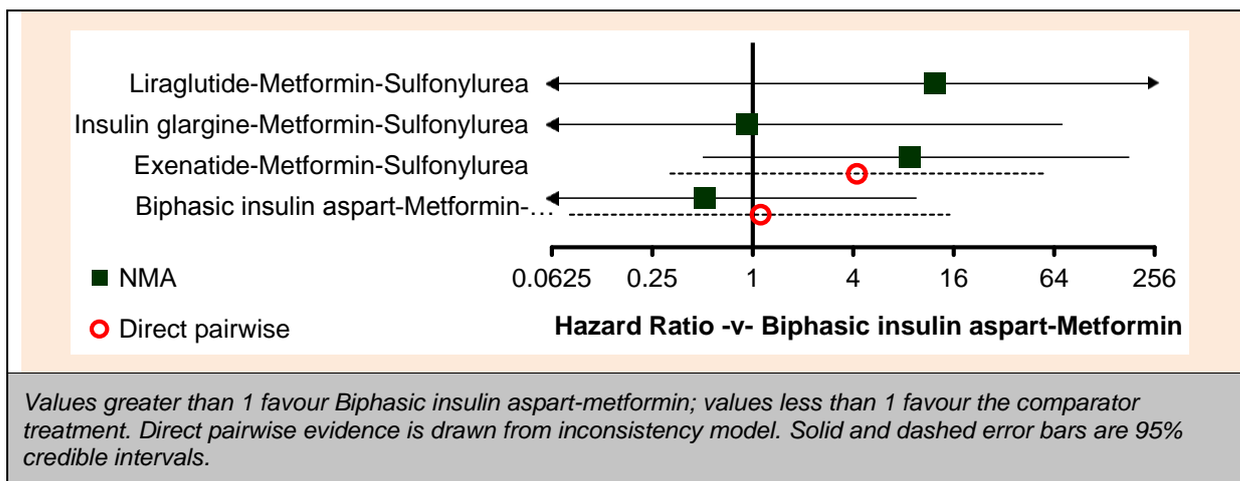
3



4 **Figure 60: Network meta-analysis of nausea (study end point) – evidence network**

5

6



1 **Figure 61: Network meta-analysis of nausea (study end point) – relative effect of all**  
 2 **options compared with common comparator (metformin-biphasic insulin**  
 3 **aspart)**

5 **Table 80: Network meta-analysis of Nausea (study end point) – rankings for each**  
 6 **comparator**

	Probability best	Median rank (95%CrI)
Biphasic insulin aspart-Metformin	0.222	2 (1, 5)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.460	2 (1, 4)
Exenatide-Metformin-Sulfonylurea	0.000	4 (3, 5)
Insulin glargine-Metformin-Sulfonylurea	0.292	2 (1, 4)
Liraglutide-Metformin-Sulfonylurea	0.025	5 (1, 5)

Update 2015

#### 8.4.12.67 Change in body weight up to 12 months

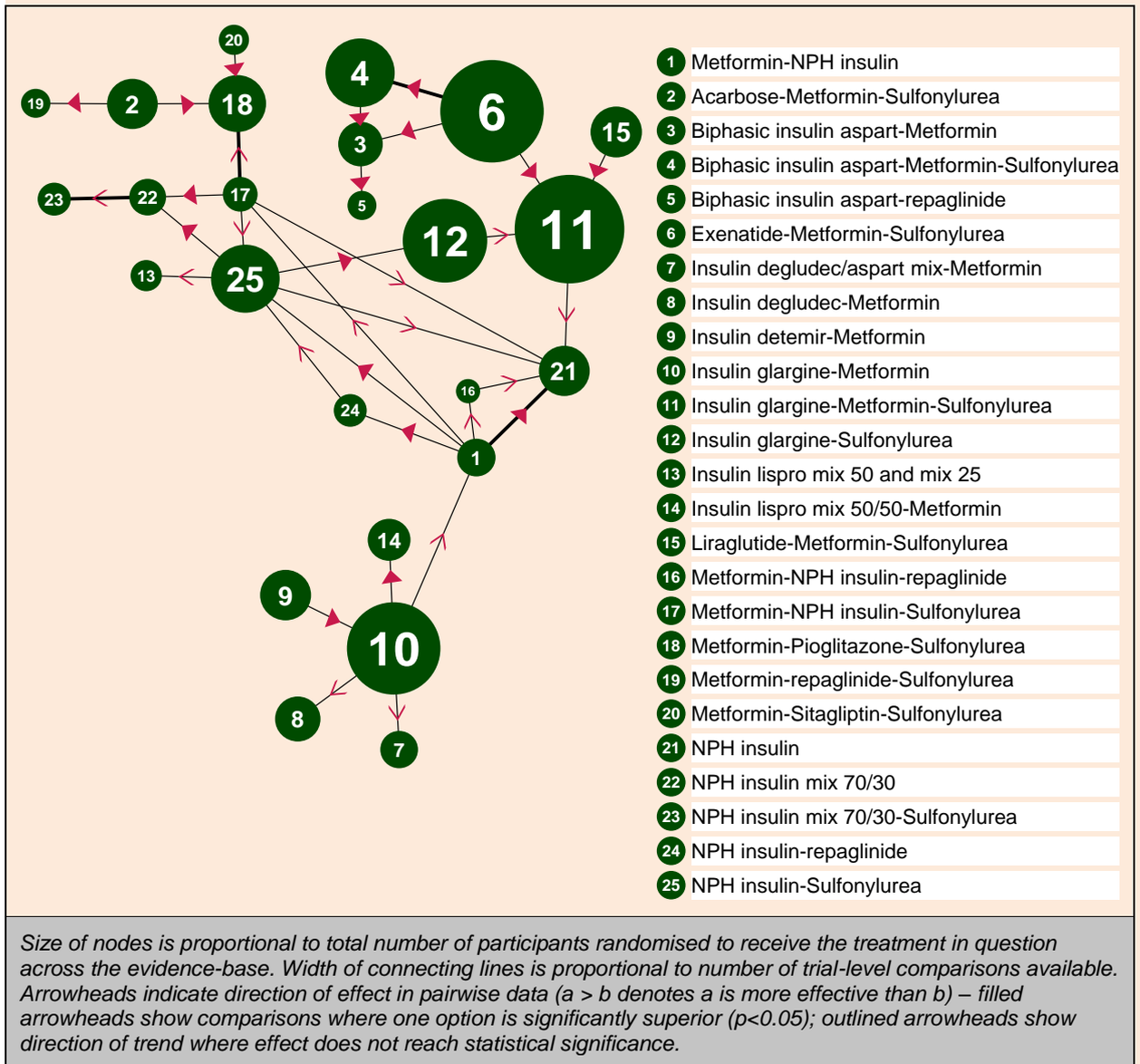
8 Results of the NMA are summarised below for the 24 treatment combinations that were  
 9 compared with metformin-NPH insulin. Of the 24 treatment combinations, 6 were 3 non-  
 10 insulin based drug combinations, 3 were insulin only, 11 were insulin + 1 non-insulin based  
 11 drug combinations and 4 were insulin + 2 non-insulin based drug combinations.

12 There is reasonable agreement between the NMA evidence and direct pairwise treatment  
 13 effect estimates as demonstrated by the substantial overlap between the credible/confidence  
 14 intervals.

15 In general, compared to metformin-NPH insulin, insulin only and insulin + 2 non-insulin based  
 16 drug combination were shown to be associated with weight gain. Combinations of 3 non-  
 17 insulin based drug combinations were generally associated with weight gain except for  
 18 combinations with GLP-1 mimetics (exenatide, liraglutide) with metformin and sulfonylurea  
 19 which showed a trend for weight loss compared to metformin-NPH insulin, though credible  
 20 intervals crossed the line of no effect. Insulin + 1 non-insulin based drug combinations were  
 21 generally associated with weight gain, except for metformin-insulin detemir which was  
 22 associated with weight loss.

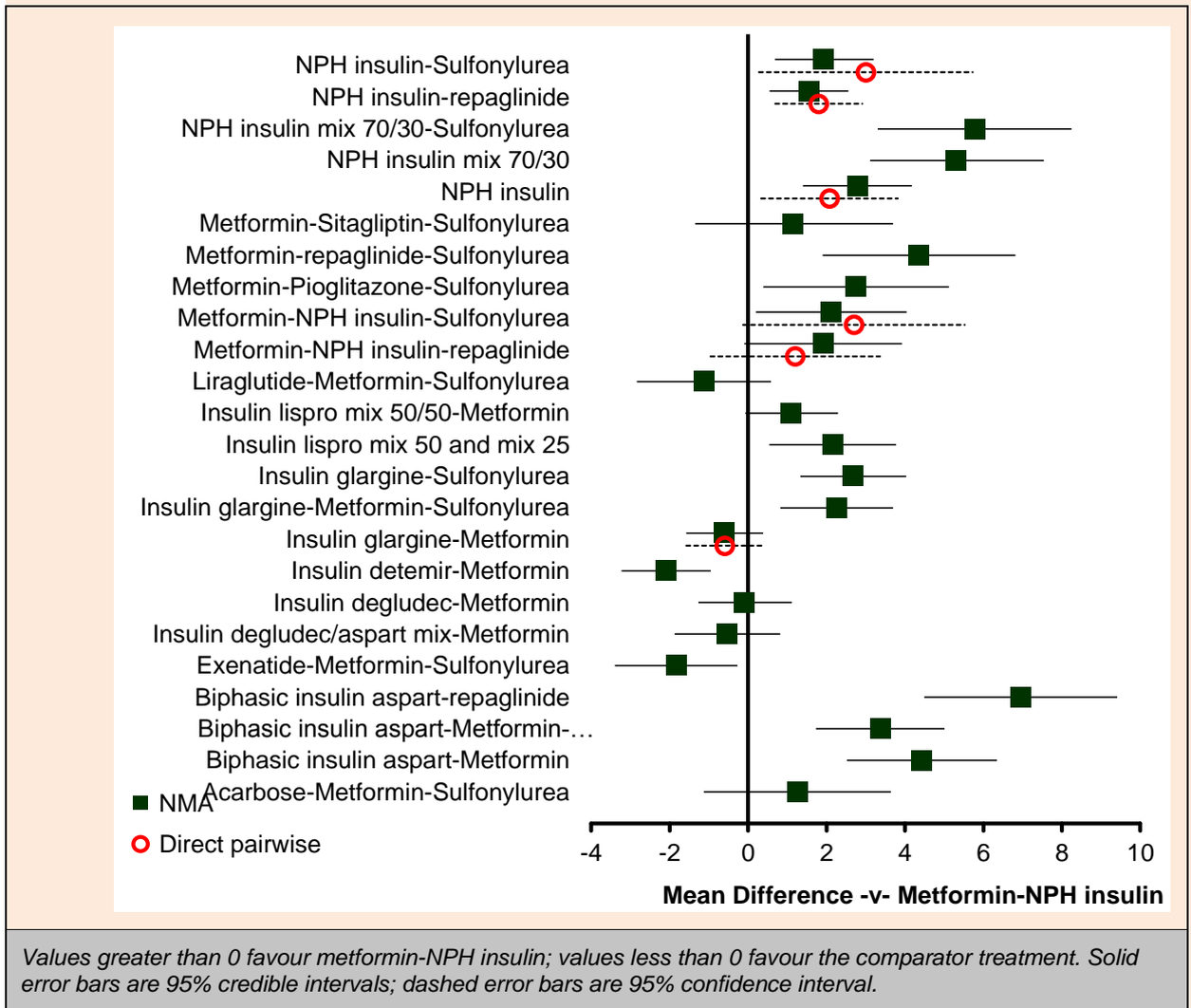
23 The treatment combinations with the highest ranking were exenatide-metformin-sulfonylurea  
 24 (median rank 2 [1 to 7]) and metformin-insulin detemir (median rank 2 [1 to 4]). Biphasic  
 25 insulin aspart-repaglinide was associated with the lowest ranking (median rank 25 [22 to 25]).

26



1 **Figure 62: Network meta-analysis of change in body weight (up to 12 months) –**  
 2 **evidence network**

3



1 **Figure 63: Network meta-analysis of change in body weight (up to 12 months) –**  
 2 **relative effect of all options compared with common comparator (metformin-**  
 3 **NPH insulin)**  
 4

1 **Table 81: Network meta-analysis of change in body weight (12 months) – rankings for**  
2 **each comparator**

	Probability best	Median rank (95%CrI)
Metformin-NPH insulin	0.000	7 (4, 9)
Acarbose-Metformin-Sulfonylurea	0.001	10 (4, 18)
Biphasic insulin aspart-Metformin	0.000	22 (19, 24)
Biphasic insulin aspart-Metformin-Sulfonylurea	0.000	20 (14, 22)
Biphasic insulin aspart-repaglinide	0.000	25 (22, 25)
Exenatide-Metformin-Sulfonylurea	0.366	2 (1, 5)
Insulin degludec/aspart mix-Metformin	0.001	5 (2, 9)
Insulin degludec-Metformin	0.000	6 (3, 10)
Insulin detemir-Metformin	0.589	1 (1, 3)
Insulin glargine-Metformin	0.000	4 (2, 7)
Insulin glargine-Metformin-Sulfonylurea	0.000	15 (9, 19)
Insulin glargine-Sulfonylurea	0.000	17 (12, 20)
Insulin lispro mix 50 and mix 25	0.000	14 (8, 20)
Insulin lispro mix 50/50-Metformin	0.000	10 (7, 17)
Liraglutide-Metformin-Sulfonylurea	0.041	3 (1, 8)
Metformin-NPH insulin-repaglinide	0.000	13 (7, 21)
Metformin-NPH insulin-Sulfonylurea	0.000	14 (9, 20)
Metformin-Pioglitazone-Sulfonylurea	0.000	18 (10, 21)
Metformin-repaglinide-Sulfonylurea	0.000	22 (16, 24)
Metformin-Sitagliptin-Sulfonylurea	0.003	10 (3, 18)
NPH insulin	0.000	18 (12, 20)
NPH insulin mix 70/30	0.000	23 (20, 25)
NPH insulin mix 70/30-Sulfonylurea	0.000	24 (21, 25)
NPH insulin-repaglinide	0.000	11 (8, 17)
NPH insulin-Sulfonylurea	0.000	13 (9, 17)

Update 2015

### 8.4.133 Health economic evidence for second intensification

#### 8.4.13.14 Systematic review of published cost–utility analyses

5 For second intensification, 7 UK studies were included covering 4 broad comparisons  
6 (Beaudet et al. 2011; McEwan et al. 2007; Pollock et al. 2012; Ray et al. 2007; Valentine et  
7 al. 2005; Waugh et al. 2010; Woehl et al. 2008), none of which covered all the comparators  
8 included in this guideline. Ray et al. (2007), Waugh et al. (2010) and Woehl et al. (2008) all  
9 compared exenatide with insulin glargine. All were based on the same RCT evidence (Heine  
10 et al. 2005) but found different results, due to differing treatment effect assumptions, drug  
11 price assumptions and weight loss utilities/profiles. Ray et al. (2007) thought exenatide-  
12 metformin-sulfonylurea was cost-effective compared to insulin glargine-metformin-  
13 sulfonylurea (ICER £22,400 per QALY); Waugh et al. (2010) found similar ICERs (ICERs  
14 19,900 per QALY for males and £18,400 for females). Woehl et al. (2008) found insulin  
15 glargine-metformin-sulfonylurea dominated exenatide-metformin-sulfonylurea. Beaudet et al.  
16 (2011) compared exenatide once weekly with insulin glargine twice daily and found  
17 exenatide to be cost effective (ICER £10,600 per QALY), but did not model treatment  
18 withdrawals.

19 Two studies compared different biphasic insulins with insulin glargine. Pollock et al. (2012)  
20 found insulin lispro 50/50 to be dominant compared with insulin glargine, but assumed

1 people were not taking concomitant oral medications and did not list their cost and utility  
2 sources. Valentine et al. (2005) found insulin aspart 70/30 to be cost effective compared with  
3 insulin glargine (ICER £7000 per QALY), but used a non-UK population and did not appear  
4 to model hypoglycaemia.

5 Two studies compared NPH insulin with insulin glargine, but came to opposing conclusions.  
6 McEwan et al. (2007) modelled either HbA1c or hypoglycaemia using unpublished treatment  
7 effect data and found insulin glargine to be cost effective compared with NPH insulin (ICER  
8 £13,900 per QALY for HbA1c reduction only, £10,000 per QALY for hypoglycaemia reduction  
9 only). Waugh et al. (2010) found insulin glargine-metformin-sulfonylurea was not cost  
10 effective compared with metformin-NPH insulin-sulfonylurea (ICERs £281,300 per QALY for  
11 males and £178,000 per QALY for females). Waugh et al. (2010) also found insulin detemir  
12 was not cost-effective compared with NPH insulin (ICER £187,700 per QALY for males and  
13 £102,000 per QALY for females), but their analysis did not cover all the comparators  
14 included in this guideline.

15 As no directly applicable studies with only minor limitations were found that covered all the  
16 comparators under consideration for each sub-question for this guideline, an original  
17 economic analysis was undertaken.

#### 8.4.13.28 Original health economic analysis

19 For second intensification, 20 treatments could be modelled. People accrued an average of  
20 13.9 undiscounted life years. Due to slightly greater differences in HbA1c treatment effects  
21 and a lack of further intensification, second intensification showed larger differences in  
22 lifetime complication rates than initial therapy and first intensification.

23 People accumulated between 6.8 and 7.4 lifetime discounted QALYs, with losses due to  
24 weight changes of between 0.3 and 0.5 QALYs and losses due to hypoglycaemic episodes  
25 of between 0.2 and 0.6 QALYs.

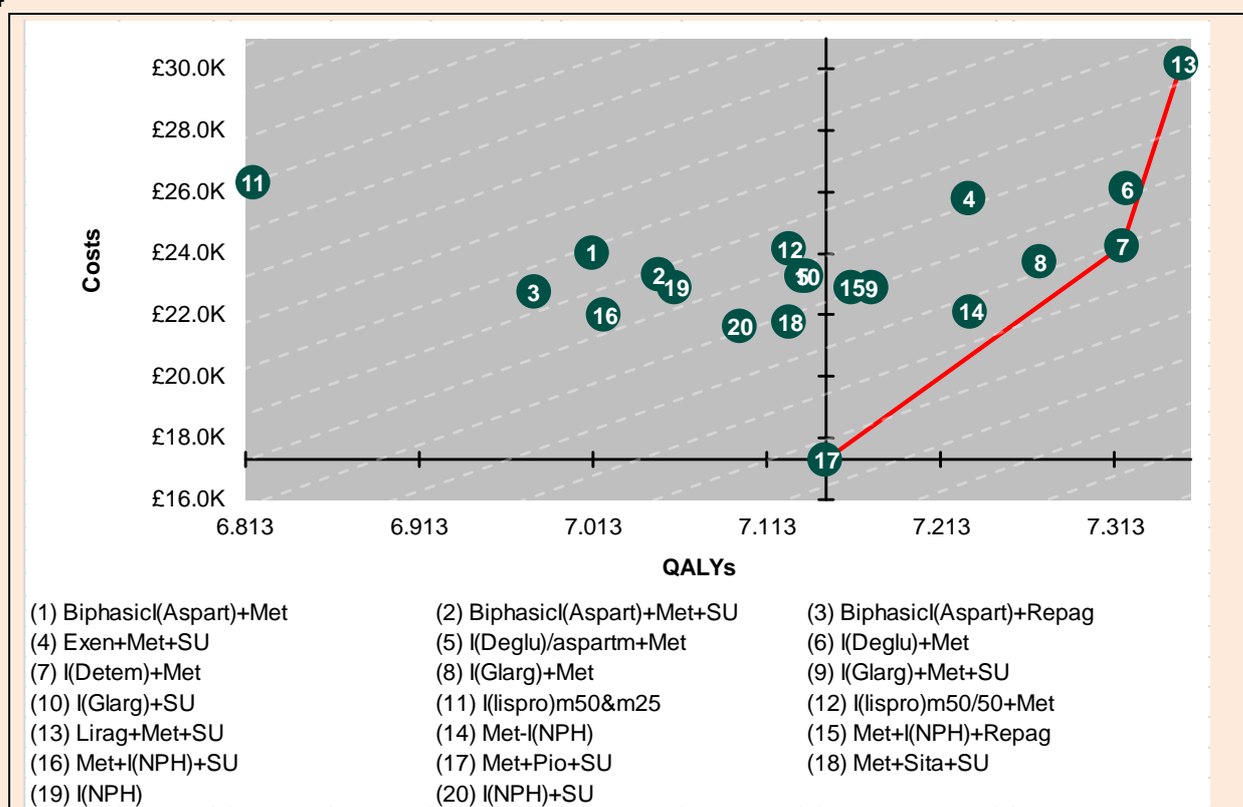
26 Second intensification therapy with metformin-pioglitazone-sulfonylurea was the cheapest  
27 and most cost-effective treatment option (see table 82). Compared with this option, all other  
28 treatment options were subject to dominance or extended dominance, with the exceptions of  
29 insulin detemir-metformin (ICER £40,800 per QALY) and liraglutide-metformin-sulfonylurea  
30 (ICER £172,900 per QALY compared with insulin detemir-metformin).

1 **Table 82: Mean lifetime incremental cost–utility results for second intensification**  
2 **therapy**

Therapy	Lifetime discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Metformin-pioglitazone-sulfonylurea	£17,279	7.147			
NPH insulin-sulfonylurea	£21,636	7.097	£4358	-0.050	Dominated
Metformin-sitagliptin-sulfonylurea	£21,763	7.126	£4484	-0.021	Dominated
Metformin-NPH insulin-sulfonylurea	£22,000	7.020	£4721	-0.127	Dominated
Metformin-NPH insulin	£22,108	7.230	£4829	0.083	Ext. dom.
Biphasic insulin aspart-repaglinide	£22,738	6.979	£5460	-0.168	Dominated
Insulin glargine-metformin-sulfonylurea	£22,870	7.173	£5591	0.026	Dominated
NPH insulin	£22,896	7.060	£5617	-0.086	Dominated
Metformin-NPH insulin-repaglinide	£22,899	7.161	£5620	0.015	Dominated
Insulin glargine-sulfonylurea	£23,260	7.135	£5982	-0.011	Dominated
Insulin degludec/aspart mix-metformin	£23,263	7.134	£5984	-0.013	Dominated
Biphasic insulin aspart-metformin-sulfonylurea	£23,303	7.051	£6025	-0.096	Dominated
Insulin glargine-metformin	£23,716	7.270	£6437	0.123	Ext. dom.
Biphasic insulin aspart-metformin	£24,028	7.013	£6750	-0.134	Dominated
Insulin lispro mix 50/50-metformin	£24,136	7.126	£6858	-0.021	Dominated
Insulin detemir-metformin	£24,228	7.317	£6950	0.170	£40,778
Exenatide-metformin-sulfonylurea	£25,795	7.229	£1567	-0.088	Dominated
Insulin degludec-metformin	£26,097	7.320	£1869	0.003	Ext. dom.
Insulin lispro mix 50 and mix 25	£26,307	6.818	£2078	-0.499	Dominated
Liraglutide-metformin-sulfonylurea	£30,166	7.352	£5937	0.034	£172,890

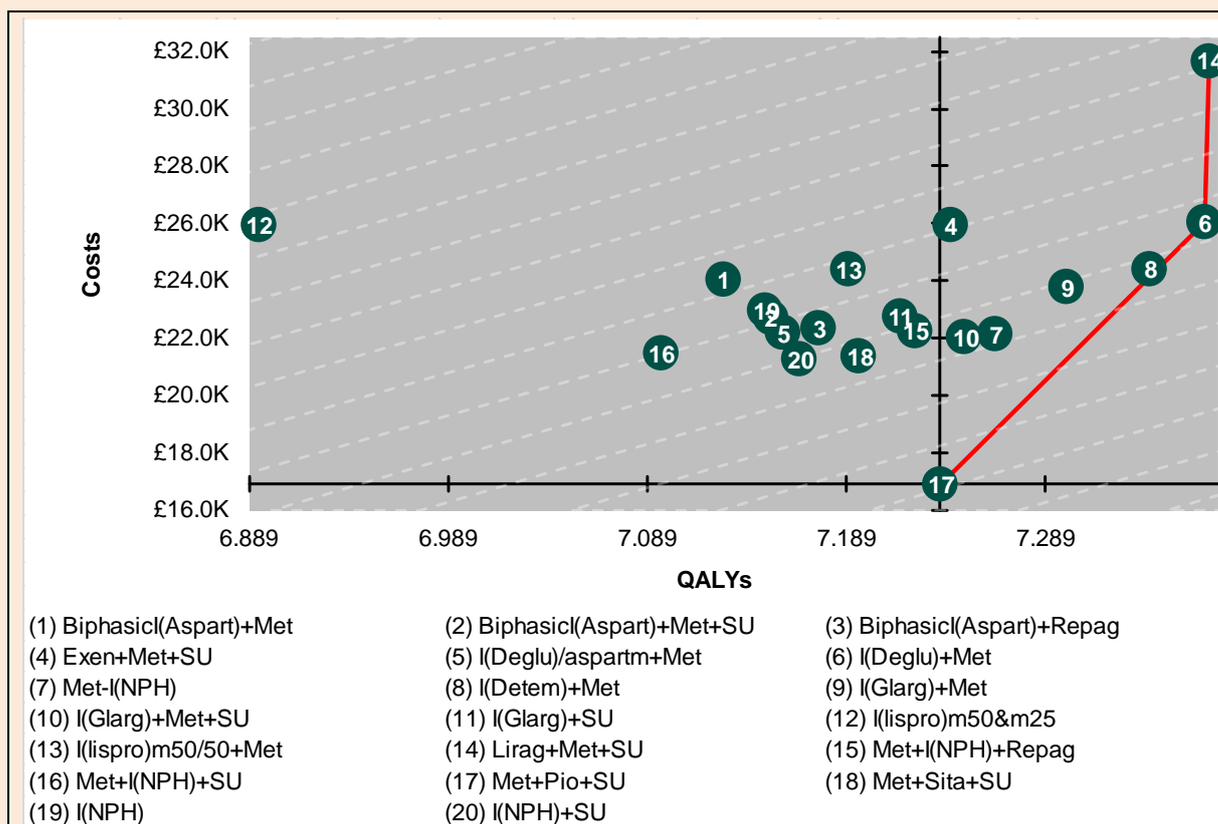
3 (a) *Ext. dom.* = extendedly dominated

4



5 **Figure 64: Cost–utility plane for 2nd intensification of therapy**

1 There were a number of treatments with similar QALY gains and costs to insulin detemir-  
2 metformin (see figure 64). Of the group, insulin detemir-metformin gained the most QALYs  
3 due to its superior weight treatment effect, despite having the worst HbA1c/UKPDS QALYs  
4 of the group of treatment options. However, the GDG expressed concern as to whether such  
5 a weight change was achievable and sustainable in practice. A sensitivity analysis was  
6 undertaken where the weight change assumptions were changed to both weight loss and  
7 weight gain only lasting for 1 year (as per the clinical evidence) – in the base case, weight  
8 loss only lasted 1 year but weight gained remained forever. In the sensitivity analysis, insulin  
9 detemir-metformin was dominated by metformin-pioglitazone-sulfonylurea, indicating insulin  
10 detemir-metformin was highly sensitive to the weight profile assumptions applied (see figure  
11 65).



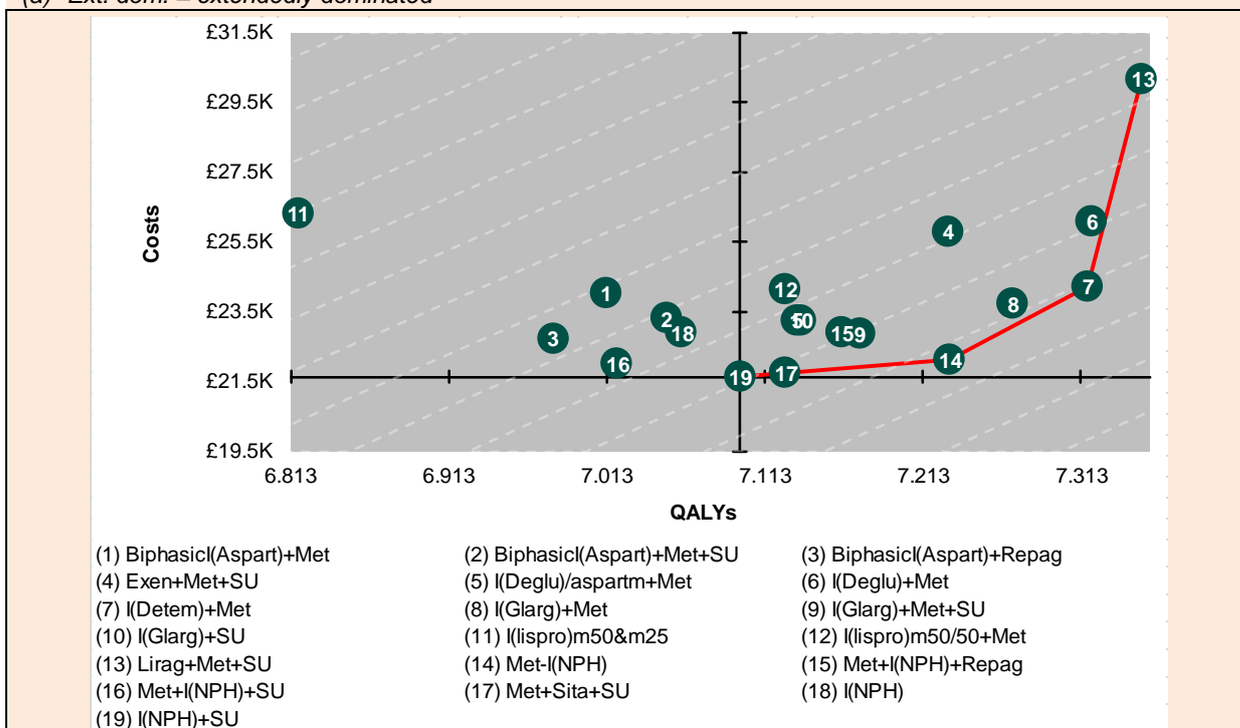
12 **Figure 65: Cost-utility plane for 2nd intensification of therapy alternative weight**  
13 **profile sensitivity analysis**

14 Given the many contraindications for prescribing pioglitazone, a further analysis was  
15 undertaken for a decision space without metformin-pioglitazone-sulfonylurea. In this analysis,  
16 Metformin-NPH insulin was a cost-effective option, resulting in QALY gains of 0.133  
17 compared with the least expensive option, NPH insulin-sulfonylurea, at an ICER of around  
18 £3600 per QALY gained. Insulin detemir-metformin was associated with an ICER of £24,300  
19 per QALY when compared with metformin-NPH insulin (see table 85). Metformin-sitagliptin-  
20 sulfonylurea was extendedly dominated by NPH insulin-sulfonylurea and metformin-NPH  
21 insulin (see figure 66), but represented the only remaining non-injectable based treatment  
22 option.

1 **Table 83: Mean lifetime incremental cost–utility results for second intensification**  
 2 **therapy – when metformin-pioglitazone-sulfonylurea is not within the**  
 3 **decision space**

Therapy	Lifetime discounted		Incremental		ICER
	Costs	QALYs	Costs	QALYs	
NPH insulin-sulfonylurea	£21,636	7.097			
Metformin-sitagliptin-sulfonylurea	£21,763	7.126	£127	0.029	Ext. dom.
Metformin-NPH insulin-sulfonylurea	£22,000	7.020	£364	-0.077	Dominated
Metformin-NPH insulin	£22,108	7.230	£472	0.133	£3552
Biphasic insulin aspart-repaglinide	£22,738	6.979	£631	-0.251	Dominated
Insulin glargine-metformin-sulfonylurea	£22,870	7.173	£762	-0.057	Dominated
NPH insulin	£22,896	7.060	£788	-0.169	Dominated
Metformin-NPH insulin-repaglinide	£22,899	7.161	£791	-0.068	Dominated
Insulin glargine-sulfonylurea	£23,260	7.135	£1153	-0.094	Dominated
Insulin degludec/aspart mix-metformin	£23,263	7.134	£1155	-0.096	Dominated
Biphasic insulin aspart-metformin-sulfonylurea	£23,303	7.051	£1196	-0.179	Dominated
Insulin glargine-metformin	£23,716	7.270	£1608	0.040	Ext. dom.
Biphasic insulin aspart-metformin	£24,028	7.013	£1921	-0.217	Dominated
Insulin lispro mix 50/50-metformin	£24,136	7.126	£2028	-0.104	Dominated
Insulin detemir-metformin	£24,228	7.317	£2121	0.087	£24,260
Exenatide-metformin-sulfonylurea	£25,795	7.229	£1567	-0.088	Dominated
Insulin degludec-metformin	£26,097	7.320	£1869	0.003	Ext. dom.
Insulin lispro mix 50 and mix 25	£26,307	6.818	£2078	-0.499	Dominated

4 (a) *Ext. dom. = extendedly dominated*



5 **Figure 66: Cost–utility plane for 2nd intensification of therapy where metformin-**  
 6 **pioglitazone-sulfonylurea is not a treatment option**

7 If, following treatment with 3 oral anti-diabetic agents, NPH insulin-based treatment options  
 8 fail to control a person's HbA1c, insulin glargine-metformin had an ICER of £8700 compared

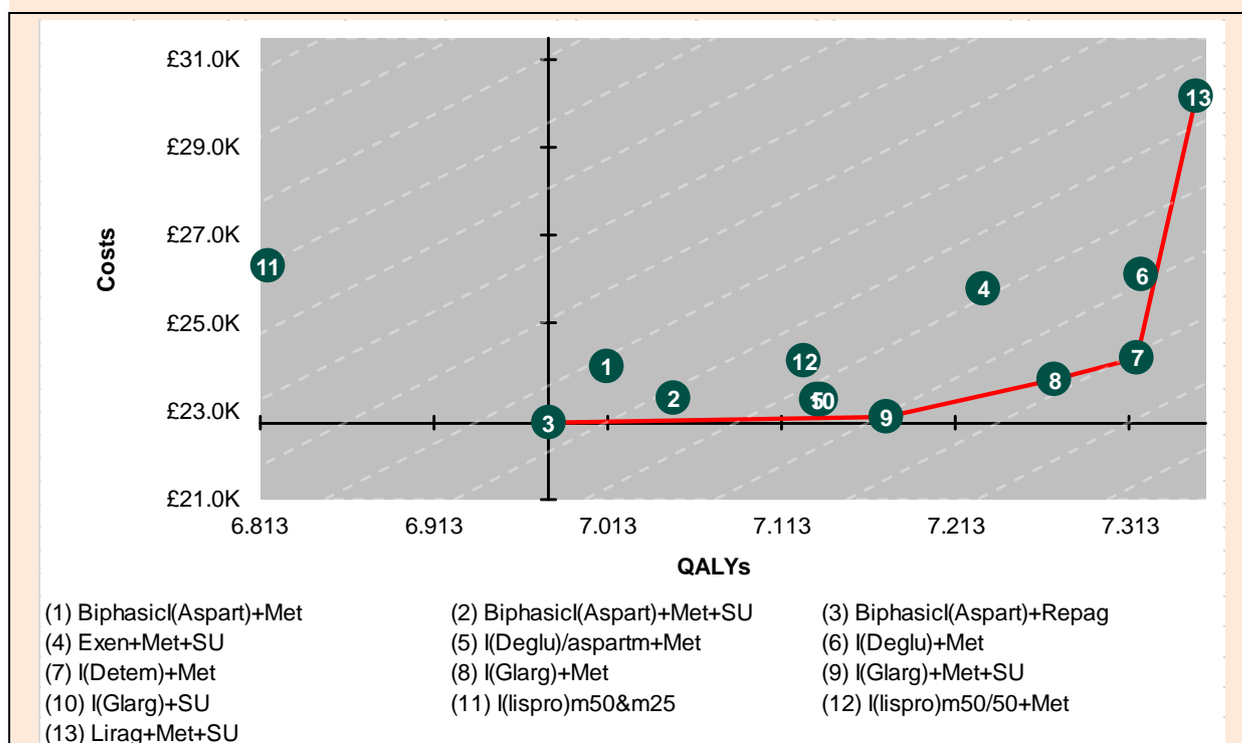
1 with insulin glargine-metformin-sulfonylurea and insulin detemir-metformin had an ICER of  
2 £10,800/QALY compared with insulin glargine-metformin (see table 84 and figure 67).

3 **Table 84: Mean lifetime incremental cost–utility results for second intensification of**  
4 **therapy when metformin-pioglitazone-sulfonylurea and NPH insulin is not a**  
5 **treatment option**

Therapy	Lifetime Discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
Biphasic insulin aspart-repaglinide	£22,738	6.979			
Insulin glargine-metformin-sulfonylurea	£22,870	7.173	£132	0.194	£678
Insulin glargine-sulfonylurea	£23,260	7.135	£391	-0.038	Dominated
Insulin degludec/aspart mix-metformin	£23,263	7.134	£393	-0.039	Dominated
Biphasic insulin aspart-metformin-sulfonylurea	£23,303	7.051	£434	-0.122	Dominated
Insulin glargine-metformin	£23,716	7.270	£846	0.097	£8,740
Biphasic insulin aspart-metformin	£24,028	7.013	£313	-0.257	Dominated
Insulin lispro mix 50/50-metformin	£24,136	7.126	£420	-0.144	Dominated
Insulin detemir-metformin	£24,228	7.317	£513	0.047	£10,795
Exenatide-metformin-sulfonylurea	£25,795	7.229	£1567	-0.088	Dominated
Insulin degludec-metformin	£26,097	7.320	£1869	0.003	Ext. dom.
Insulin lispro mix 50 and mix 25	£26,307	6.818	£2078	-0.499	Dominated
Liraglutide-metformin-sulfonylurea	£30,166	7.352	£5937	0.034	£180.982

6 (a) *Ext. dom.* = extendedly dominated

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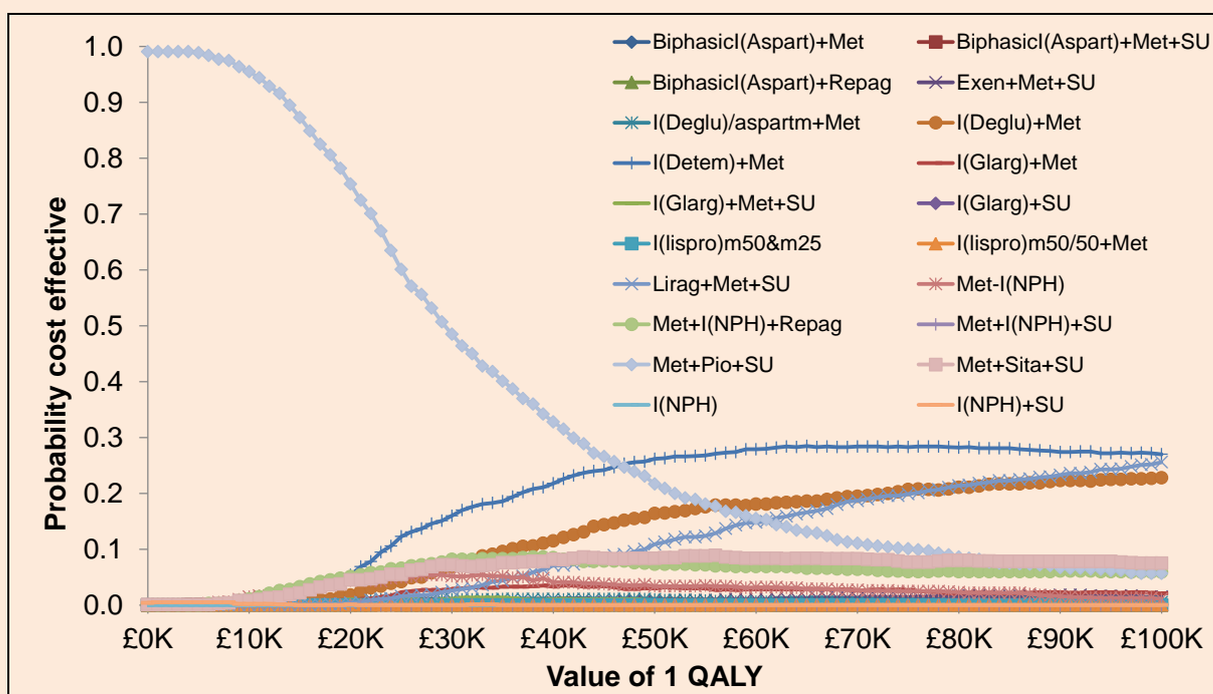
8 **Figure 67: Cost–utility plane for 2nd intensification of therapy where 3 oral anti-**  
9 **diabetic agent treatment options followed by metformin-NPH insulin have**  
10 **failed to control HbA1c**

1 For people who could not tolerate metformin, NPH insulin-sulfonylurea was the cheapest and  
2 most cost effective of the 5 treatment options (see table 85).

3 **Table 85: Mean lifetime incremental cost-utility results for second intensification of**  
4 **therapy when metformin cannot be tolerated**

Therapy	Lifetime Discounted		Incremental		
	Costs	QALYs	Costs	QALYs	ICER
NPH insulin-sulfonylurea	£21,636	7.097			
Biphasic insulin aspart-repaglinide	£22,738	6.979	£1102	-0.118	Dominated
NPH insulin	£22,896	7.060	£1260	-0.037	Dominated
Insulin glargine-sulfonylurea	£23,260	7.135	£1624	0.038	£42,369
Insulin lispro mix 50 and mix 25	£26,307	6.818	£3046	-0.317	Dominated

5 At second intensification, metformin-pioglitazone-sulfonylurea was the most cost-effective  
6 treatment option at a maximum acceptable ICER of £20,000 per QALY in 75% of iterations  
7 (see figure 68).



8 **Figure 68: Cost-effectiveness acceptability curve for second intensification**

9

### 8.4.140 Evidence statements for second intensification

#### 8.4.14.11 Clinical evidence

##### 8.4.14.1.12 Change in blood glucose up to 12 months

13 Evidence from a single network meta-analysis including data from 37 RCTs for HbA1c levels  
14 showed that NPH-insulin combined with metformin and repaglinide were most effective in  
15 blood glucose control, followed by biphasic insulin aspart-pioglitazone. Metformin-  
16 sulfonylurea-repaglinide was ranked lowest suggesting that this combination was least  
17 effective in decreasing HbA1c levels. The quality of the evidence was moderate.

#### 8.4.14.1.21 **Hypoglycaemia at study end point**

2 Evidence from a single network meta-analysis including data from 34 RCTs showed that  
3 metformin combined with insulin degludec or insulin detemir were associated with high  
4 rankings (median rank 3 [1 to 16] and median rank 3 [1 to 15] respectively) indicating lower  
5 hypoglycaemic events. Insulin lispro mix 50 and mix 25 was associated with the lowest  
6 ranking suggesting higher hypoglycaemic events. However, there was greater uncertainty  
7 surrounding the evidence as the credible intervals were generally wide and crossed the line  
8 of no effect. The quality of the evidence was low.

#### 8.4.14.1.39 **Adverse events at study end point**

10 Evidence from 3 network meta-analyses including data from 25, 25 and 4 RCTs for dropouts  
11 due to adverse events, total dropouts and nausea respectively, showed that insulin lispro 50  
12 and 25 mix had highest ranking for dropouts due to adverse events, whereas a triple oral  
13 combination (metformin-sulfonylurea-repaglinide) had the highest ranking for total dropouts.  
14 Insulin combinations rather than triple non-insulin based drug combinations demonstrated  
15 comparatively higher rankings indicating lower nausea events (biphasic aspart-metformin,  
16 biphasic aspart-metformin-sulfonylurea and glargine-metformin-sulfonylurea). However, there  
17 was considerable uncertainty around the network meta-analyses demonstrated by wide  
18 credible intervals which in the main crossed the line of no effect. The quality of the evidence  
19 was low.

#### 8.4.14.1.40 **Change in body weight up to 12 months**

21 Evidence from a single network meta-analysis including data from 27 RCTs showed that  
22 metformin-insulin detemir and a triple non-insulin based drug combination of metformin-  
23 sulfonylurea and a GLP-1 mimetic (exenatide, liraglutide) were associated with weight loss.  
24 Biphasic insulin aspart-repaglinide was associated with lowest ranking. The quality of the  
25 evidence was low.

#### 8.4.14.26 **Health economic evidence**

27 A directly applicable health economic model with potentially serious limitations found  
28 metformin-pioglitazone-sulfonylurea was the most cost-effective modelled option for second  
29 intensification therapy. A further analysis found metformin-NPH insulin to be the most cost-  
30 effective treatment option when pioglitazone is not a treatment option. NPH insulin-  
31 sulfonylurea was the most cost-effective combination that did not contain metformin.

### 8.4.152 **Evidence to recommendations for second intensification**

#### 33 **Table 86: Linking evidence to recommendations**

Relative value of different outcomes

The following outcomes were considered critical to decision making; glycaemic control (HbA1c), hypoglycaemic events and adverse events. Change in body weight was considered important to decision making.

The GDG noted that glycaemic control was important in mitigating the much increased risk of microvascular and macrovascular complications associated with high levels of hyperglycaemia at this intensification level. However, the GDG acknowledged that tight glycaemic control may be associated with increased risk of hypoglycaemia, which may negatively impact on quality of life. Drug tolerability and change in body weight were considered important in determining the acceptability of treatment to the patient.

The relative importance of each outcome is variable depending on

	<p>several factors:</p> <ul style="list-style-type: none"> <li>• Severity of hyperglycaemia.</li> <li>• Individual circumstances such as comorbidities and body mass index.</li> </ul>
<p>Trade-off between benefits and harms</p>	<p>The GDG acknowledged that there was generally less evidence at this treatment level, resulting in sparser networks. The GDG noted that there was some uncertainty in the evidence at this intensification level as demonstrated by the wide credible intervals that surrounded many of the point estimates particularly related to adverse events. The GDG noted all 6 triple non-insulin based drug combinations included metformin.</p> <p>Of the 32 treatment combinations, 20 included studies which reported data for all required outcomes in the health economic model. The 12 treatment combinations that were not included in the health economic model were 2 triple oral therapies (metformin-sulfonylurea-acarbose and metformin-sulfonylurea-repaglinide), 4 insulin-only combinations (biphasic insulin aspart, insulin aspart (short acting), NPH insulin 70/30 and biphasic insulin-NPH insulin) and 6 combinations of insulin + 1 oral antidiabetic drug (biphasic aspart-pioglitazone, insulin aspart-metformin, lispro 75/25-metformin, NPH 70/30-metformin, NPH 70/30-sulfonylurea and NPH insulin-repaglinide). The GDG noted that many of the triple oral antidiabetic drug combinations are not commonly used in clinical practice, such as acarbose and sulfonylurea-repaglinide which are both secretagogues that act by stimulating the pancreas.</p> <p>The GDG discussed that many patients are generally unwilling to start insulin therapy due to a fear of injections, hypoglycaemia and its potential impact on quality of life. The GDG discussed the evidence surrounding 3 non-insulin based drug combinations and noted that while they were not the most effective in decreasing HbA1c levels, they were associated with less hypoglycaemic events and, for some combinations, weight loss.</p> <p>The GDG discussed the evidence of combinations including GLP-1 mimetics and noted that while triple non-insulin based drug combinations including GLP-1 mimetics had better weight profiles, there was uncertainty in the data due to relatively wide credible intervals that crossed the line of no effect. Hence, they thought that this combination should be available to people for whom obesity is a concern. The GDG noted that there was a lack of evidence for combinations of GLP-1 mimetics and insulin, and therefore agreed that this option should only be offered in a specialist care setting.</p> <p>The GDG discussed alternative options available if triple non-insulin based drug combinations failed to adequately control blood glucose levels. They noted that metformin-NPH insulin was ranked in at least the top third for reducing HbA1c levels, hypoglycaemic events and change in body weight. The GDG recognised that there were other insulin-metformin combinations which had variable degrees of effectiveness across the 3 outcomes such as metformin-detemir ranked in bottom third for change in HbA1c levels but highest third for hypoglycaemic events and change in body weight.</p> <p>The GDG discussed the evidence surrounding the relative benefit of weight loss compared to other treatments in the metformin-detemir combination, and noted that this was predominantly due to the comparative weight gain observed by all other insulin-based</p>

	<p>treatment combinations, rather than the marginal weight decrease in individuals receiving metformin-detemir observed in 1 trial (weight reduction of 0.5 kg).</p> <p>The GDG discussed intensification options for individuals where metformin is contraindicated or not tolerated, and noted that there was some evidence for sulfonylurea-insulin combinations and insulin only combinations. They noted that the evidence profile for NPH insulin-sulfonylurea, NPH insulin 70/30-sulfonylurea and insulin glargine-sulfonylurea were similar.</p> <p>The GDG discussed the value of using insulin to achieve rapid blood glucose control (rescue therapy) in clinical practice in patients who are symptomatically hyperglycaemic but agreed that treatment should be reviewed once blood glucose targets have been achieved.</p>
<p>Consideration of health benefits and resource use</p>	<p>Compared with earlier therapy levels, treatments at second intensification showed slightly greater differences in lifetime complication rates, as there were slightly greater HbA1c differences and no further intensifications of treatment.</p> <p>The economic model made explicit the trade-offs between the higher costs, benefits (HbA1c) and harms (hypoglycaemia and body weight) of insulin-based therapies against triple non-insulin based drug combinations, which are cheaper and associated with less harm but relatively ineffective at controlling HbA1c. Metformin-pioglitazone-sulfonylurea dominated all other treatment combinations, except for insulin detemir-metformin (ICER £40,800 per QALY), and liraglutide-metformin-sulfonylurea (ICER £172,900 per QALY compared with insulin degludec-metformin).</p> <p>Insulin detemir-metformin showed a smaller QALY loss due to lower weight gain and lower hypoglycaemia rates than other treatments. The GDG expressed strong reservations as to whether these lower weight gains were seen in clinical practice and noted the very low quality of the clinical network supporting this evidence. It was also mindful that, in the base case, the model sustained the weight gains for other treatments for life and the group were unsure that a sustained weight difference between treatments would occur.</p> <p>The GDG felt that many people would be contraindicated for prescribing metformin-pioglitazone-sulfonylurea. The GDG considered a decision space without metformin-pioglitazone-sulfonylurea, which showed a cluster of longer-acting insulins combined with metformin to have similar lifetime discounted costs and QALYs. Compared with NPH insulin-sulfonylurea, metformin-NPH insulin produced an ICER of £3600 per QALY. Compared with metformin-NPH insulin, insulin detemir-metformin produced an ICER of £24,300 per QALY; the GDG were not convinced the lower weight gain associated with detemir-metformin was clinically realistic.</p> <p>Whilst metformin-sitagliptin-sulfonylurea was extendedly dominated by NPH insulin-sulfonylurea and metformin-NPH insulin, the GDG felt the dominance was marginal (the option was extremely close to the cost-effectiveness frontier) and there was value in recommending metformin-sitagliptin-sulfonylurea as a non-injectable alternative to metformin-NPH insulin. The GDG noted there was no evidence for other DPP4-inhibitor-metformin-</p>

	<p>sulfonylurea combinations.</p> <p>The GDG felt GLP-1 mimetic combinations may be a cost effective option for patients with high BMIs who would require high doses (and therefore costs) of insulin. They also felt that, in patients for whom using insulin would have significant occupational implications, this could have a catastrophic impact on the person's quality of life. As a result, the health economic model might critically undervalue the benefits that would be associated with a treatment that forestalled the need for insulin. For these reasons, the GDG chose to retain the GLP-1 mimetic combination options with their eligibility criteria from CG87.</p> <p>The GDG considered the small subset of modelled treatments that did not consider metformin. Whilst NPH insulin and sulfonylurea was the most cost-effective option, the GDG felt they had little clinical experience of using this combination and felt patients might prefer to use NPH insulin alone.</p>
<p>Quality of the evidence</p>	<p>The GDG agreed that the overall quality of the evidence for second intensification was low. This was generally because the network was sparse with many connections limited to a single trial which led to some uncertainty around the results. In addition, for some outcomes, such as weight loss in 1 study, the results were not consistent with clinical experience.</p> <p>The GDG commented that the Derosa et al. (2013) trial was conducted in patients who were drug naïve at study baseline and may not be representative of the clinical population who require second intensification for glycaemic control. Therefore, this trial was excluded from the evidence base for second intensification.</p> <p>The GDG highlighted that outcomes, in particular hypoglycaemia would be affected by the patients' stage of the condition. Specifically, it was suggested that patients with early type 2 diabetes will have relatively tight glycaemic control, are more likely to be using long acting insulin and may be less likely to experience hypoglycaemia. In contrast, patients who are at a later stage of the condition may have higher glycaemic targets, are more likely to require biphasic insulin and are therefore more likely to experience hypoglycaemia.</p>
<p>Other considerations</p>	<p>When defining the decision problem for this question, the GDG preferred not to make an <i>a priori</i> assumption of class effect across DPP-4 inhibitors. Therefore, each individual option for which evidence was available was analysed separately. Having reviewed the assembled evidence for each phase of treatment, the GDG noted that it was difficult to judge whether the different DPP-4 inhibitors could, in fact, be considered interchangeable:</p> <ul style="list-style-type: none"> <li>• In a few areas, a case could be made for the superiority of one option over another (for example, as initial therapy, sitagliptin seemed to have somewhat superior benefits to vildagliptin at similar net costs).</li> <li>• In other areas, all the DPP-4 inhibitors for which evidence was available appeared to have very similar benefits, harms and costs (for example, in combination with metformin at first intensification).</li> <li>• Elsewhere in the treatment pathway, evidence was extremely limited (for example, sitagliptin-metformin-sulfonylurea was the only treatment combination for which evidence was available at second intensification) or absent (for example, at first</li> </ul>

intensification, there was no evidence that could be used to assess the relative clinical effectiveness and cost effectiveness of DPP-4 inhibitors in combination with pioglitazone or sulfonylureas).

Having considered these different situations, the GDG concluded that the most helpful recommendations would be ones that treated DPP-4 inhibitors as a class. Had it been presented with evidence that suggested that 1 or more of the options was superior to others across all phases of treatment, the GDG would clearly have been inclined to favour such option(s) in its recommendations. However, the picture that had emerged was much more sporadic, and the Group was not confident that any apparent dissimilarities between options represented real differences that would be expected in clinical practice. Moreover, the GDG was mindful that a series of recommendations that alternated between treating DPP-4 inhibitors as a class, in some parts of the treatment pathway, and focusing on individual options in others would be confusing to readers of the guideline, even if those recommendations could be directly allied with the available evidence. For all of these reasons, the GDG took the view that recommendations should consistently refer to DPP-4 inhibitors as a class. It was a natural extension of this principle that prescribers should be encouraged to select the individual DPP-4 inhibitor with the lowest acquisition cost available to them.

The GDG noted that the mean age in the included studies was about 57 years and agreed that these trials are biased towards younger and fitter participants, who are less likely to experience significant comorbidities than the majority of people with type 2 diabetes seen in clinical practice. The GDG thought that the treatment effects observed in trials are likely to generalise to a population facing more comorbidities and other challenges to effective management of their disease. However, the Group agreed that the balance of benefits and harms may be different in such cases, and there are specific issues based on clinical experience which may require particular attention that should be highlighted in the recommendations.

The GDG discussed that in clinical practice the use of triple non-insulin based drug combinations is preferred as patients are unwilling to start insulin therapy. The Group noted that insulin therapy may not be appropriate for some patients. The GDG discussed that progress and individual care plans should be reassessed in people for whom insulin therapy may not be appropriate. Based on their clinical experience and expertise, the GDG agreed that this should be carried out after 6 months. This duration was agreed to maximise the accuracy of HbA1c measurements. Specifically, it was discussed that the accuracy of HbA1c measurements taken before 6 months may vary with some treatments taking longer to have an effect and missed doses having a larger impact.

It was noted that reporting of hypoglycaemia differed across the included studies. All categories of hypoglycaemia (for example, confirmed hypoglycaemia) were generally a subset of 'any hypoglycaemia', which was the most commonly reported category of hypoglycaemia across the included studies. The GDG discussed the risk of bias associated with reported hypoglycaemia and noted that self-reported hypoglycaemia may not be a reliable measure as an individual's perception of hypoglycaemia is variable at different

glucose levels.

The GDG noted that the results from the sensitivity analyses of individuals who had previously failed to adequately control blood glucose levels on 2 or more non-insulin based drug combinations were similar to the full dataset which included studies of mixed populations of people who did not necessarily fail on/or were previously exposed to 2 drugs, or studies of people who failed on 1 oral antidiabetic drug.

Based on the health economic evidence of the associated cost-effectiveness of triple non-insulin based drug combinations, a strong 'offer' recommendation was made to intensify treatment by adding a sulfonylurea for individuals who had failed to achieve blood glucose targets on 2 non-insulin based drug combinations. This was assumed (as part of the overall structure of the pharmacological therapy review question) to provide better glycaemic control than dual therapy. However, it was agreed that this trial should be stopped if target HbA1c levels are not achieved. Where treatment has not been effective, a patient's individual risks and benefits should be reassessed after 6 months and appropriate changes to their treatment plan should be made. This may involve discussing the risks and benefits associated with insulin based therapy, ensuring any issues such as changes in employment are taken into account.

Based on the lack of research evidence on combinations of insulin-GLP1 mimetics, a strong 'only offer' recommendation was made to provide this treatment combination in a specialist care setting.

#### 8.4.161 Evidence review for third intensification

2 In total 17,037 references were found for the main review question, but no trials were  
3 identified for inclusion for third intensification.

#### 8.4.174 Recommendations

##### 8.4.17.15 Rescue therapy at any phase of treatment

6 **46. If an adult with type 2 diabetes is symptomatically hyperglycaemic, consider**  
7 **insulin (see recommendations 63–65) or a sulfonylurea, and review treatment**  
8 **when blood glucose control has been achieved. [new 2015]**

##### 8.4.17.29 Initial drug treatment

10 **47. Offer standard-release metformin as the initial drug treatment for adults with type**  
11 **2 diabetes. [new 2015]**

12 **48. Gradually increase the dose of standard-release metformin over several weeks to**  
13 **minimise the risk of gastrointestinal side effects. [new 2015]**

14 **49. Review the dose of standard-release metformin if the estimated glomerular**  
15 **filtration rate (eGFR) is below 45 ml/minute/1.73m<sup>2</sup>:**

- 16       • Stop standard-release metformin if the eGFR is below  
17       30 ml/minute/1.73m<sup>2</sup>.





1 **60. If combination therapy with 2 oral drug treatments has not controlled HbA1c to**  
2 **below the person's individually agreed threshold for intensification, consider**  
3 **combination therapy with standard-release metformin, a sulfonylurea and a**  
4 **glucagon-like peptide-1 (GLP-1) mimetic (instead of 3 oral drug treatments or**  
5 **introducing insulin).**

6 **Consider this for adults with type 2 diabetes who:**

- 7 • have a BMI of 35 kg/m<sup>2</sup> or higher **and** specific psychological or other
- 8 medical problems associated with obesity, **or**
- 9 • have a BMI lower than 35 kg/m<sup>2</sup> **and**
- 10 ○ for whom insulin therapy would have significant occupational
- 11 implications, **or**
- 12 ○ weight loss would benefit other significant obesity-related
- 13 comorbidities.
- 14 Base the choice of GLP-1 mimetic on the person's preference after
- 15 discussing the risks and benefits of each licensed option. If more than
- 16 1 option is considered appropriate for the person, choose the GLP-1
- 17 mimetic with the lowest acquisition cost. **[new 2015]**

18 **61. Only continue GLP-1 mimetic therapy if the person has had a beneficial metabolic**  
19 **response (a reduction of at least 11 mmol/mol [1%] in HbA1c and a weight loss of**  
20 **at least 3% of initial body weight in 6 months). [2015]**

21 **62. Only offer a GLP-1 mimetic in combination with insulin in a specialist care setting.**  
22 **[new 2015]**

23 **Treatment with combinations of medicines including SGLT2 inhibitors may be**  
24 **appropriate for some people; see the NICE guidance on dapagliflozin in combination**  
25 **therapy for treating type 2 diabetes and canagliflozin in combination therapy for**  
26 **treating type 2 diabetes<sup>m</sup>.**

#### 8.4.17.27 Insulin-based treatments

28 **63. When starting insulin therapy, use a structured programme employing active**  
29 **insulin dose titration that encompasses:**

- 30 • structured education
- 31 • continuing telephone support
- 32 • frequent self-monitoring
- 33 • dose titration to target
- 34 • dietary understanding
- 35 • management of hypoglycaemia
- 36 • management of acute changes in plasma glucose control
- 37 • support from an appropriately trained and experienced healthcare
- 38 professional. **[2015]**

39 **64. When starting insulin therapy, continue to offer standard-release metformin for**  
40 **people without contraindications or intolerance. [new 2015]**

m NICE guidance on empagliflozin for treating type 2 diabetes is in development and is due to be published in March 2015.

- 1 **65. Initiate insulin therapy from a choice of a number of insulin types and regimens:**  
2  
3
  - Offer human neutral protamine Hagedorn (NPH) insulin injected at bed-  
4 time or twice daily according to need.
  - Consider, as an alternative, using a long-acting insulin analogue (insulin  
5 detemir, insulin glargine) if:
    - 6 o the person needs assistance from a carer or healthcare professional  
7 to inject insulin, and use of a long-acting insulin analogue (insulin  
8 detemir, insulin glargine) would reduce the frequency of injections  
9 from twice to once daily, **or**
    - 10 o the person's lifestyle is restricted by recurrent symptomatic  
11 hypoglycaemic episodes, **or**
    - 12 o the person would otherwise need twice-daily NPH insulin injections in  
13 combination with oral glucose-lowering drugs, **or**
    - 14 o the person cannot use the device to inject NPH insulin.
  - 15 • Consider twice-daily pre-mixed (biphasic) human insulin (particularly if  
16 HbA1c is 75 mmol/mol [9.0%] or higher). A once-daily regimen may be  
17 an option.
  - 18 • Consider pre-mixed preparations that include short-acting insulin  
19 analogues, rather than pre-mixed preparations that include short-acting  
20 human insulin preparations, if:
    - 21 o a person prefers injecting insulin immediately before a meal, **or**
    - 22 o hypoglycaemia is a problem, **or**
    - 23 o blood glucose levels rise markedly after meals. **[2015]**

24 **66. Consider switching to a long-acting insulin analogue (insulin detemir, insulin**  
25 **glargine) from NPH insulin in people:**
  - 26 • who do not reach their target HbA1c because of significant  
27 hypoglycaemia, **or**
  - 28 • who experience significant hypoglycaemia on NPH insulin irrespective of  
29 the level of HbA1c reached, **or**
  - 30 • who cannot use the device needed to inject NPH insulin but who could  
31 administer their own insulin safely and accurately if a switch to a long-  
32 acting insulin analogue were made, **or**
  - 33 • who need help from a carer or healthcare professional to administer  
34 insulin injections and for whom switching to a long acting insulin  
35 analogue would reduce the number of daily injections. **[2015]**

36 **67. Monitor a person on a basal insulin regimen (NPH insulin or a long acting insulin**  
37 **analogue [insulin detemir, insulin glargine]) for the need for short-acting insulin**  
38 **before meals (or a pre-mixed insulin preparation). [2015]**  
39 **68. Monitor a person on pre-mixed insulin for the need for a further injection of short-**  
40 **acting insulin before meals or for a change to a basal bolus regimen with NPH**  
41 **insulin or long-acting insulin analogues (insulin detemir, insulin glargine), if blood**  
42 **glucose control remains inadequate. [2015]**

#### 8.4.17.83 Insulin delivery

- 44 **69. For guidance on insulin delivery, see the NICE guideline on type 1 diabetes. [new**  
45 **2015]**

#### 8.4.181 Research recommendations

- 2 **5. In adults with type 2 diabetes, what treatment combinations (for example,**  
3 **glucagon-like peptide-1 [GLP-1] mimetics and insulin) are most effective when**  
4 **initial drug treatment with non-metformin monotherapy fails to adequately control**  
5 **blood glucose levels?**

##### 6 **Why this is important**

7 Although it is recognised that metformin therapy is suitable for most adults with type 2  
8 diabetes, its use is contraindicated or not tolerated in approximately 15% of individuals.  
9 To date, research evidence has largely focused on metformin-based treatment  
10 combinations. Given the progressive nature of the condition, in which intensification of  
11 blood glucose lowering drug therapies are indicated over time, there is little evidence, for  
12 some adults, to guide management strategies on treatment combinations that do not  
13 include metformin. Double-blind randomised controlled trials are therefore needed to  
14 better understand the treatment choices that are available which improve glycaemic  
15 control and long-term risks of complications associated with diabetes.

- 16 **6. In adults with type 2 diabetes, what are the effects of early use of insulin and**  
17 **glucagon-like peptide-1 (GLP-1) mimetics?**

##### 18 **Why this is important**

19 Poor glycaemic control is associated with increased risk of vascular complications.  
20 Glucagon-like peptide-1 (GLP-1) mimetics are a new class of blood glucose lowering  
21 drugs that target the incretin system, regulating insulin and glucagon. It is associated  
22 with low rates of hypoglycaemia and some weight loss. Its effectiveness and safety in  
23 combination with insulin early on in the drug treatment pathway is unknown. Double-  
24 blind randomised controlled trials are needed to understand the short and long-term  
25 effects of early use of GLP-1 agonists with insulin in terms of glycaemic control, adverse  
26 effects, diabetes-related complications and mortality. Research on its use could have a  
27 significant impact on the management of adults with type 2 diabetes.

- 28 **7. When third intensification of treatment is indicated, which blood glucose lowering**  
29 **therapies should be used to control blood glucose levels?**

##### 30 **Why this is important**

31 As the incidence of type 2 diabetes increases in the younger population and as  
32 glycaemic control declines naturally over time, it is likely that further intensification of  
33 therapies would be needed. Currently, there is evidence up to second intensification of  
34 drug therapies, that is, when 2 or more non-insulin based treatment combinations fail to  
35 adequately control blood glucose levels. Double-blind randomised controlled trials are  
36 needed to improve understanding of alternative treatment options for adults at second  
37 intensification who are inadequately controlled with insulin and/or triple non-insulin  
38 based drug therapies.

- 39 **8. In adults with type 2 diabetes, what are the effects of stopping and/or switching**  
40 **drug treatments to control blood glucose levels, and what criteria should inform**  
41 **the decision?**

##### 42 **Why this is important**

1 There is a lack of evidence on the effects of stopping and/or switching drug treatments to  
2 control blood glucose levels. The current practice of 'stopping rules' is typically motivated  
3 by either inadequate blood glucose control (rising HbA1c levels) or intolerable side  
4 effects. There is limited understanding of the short- and long-term effects of stopping a  
5 therapy and switching to another in terms of diabetes control (HbA1c levels),  
6 hypoglycaemic risk, weight gain, and cardiovascular morbidity and mortality. In addition,  
7 there is limited understanding of how quickly consideration should be given to stopping  
8 and switching to another drug treatment and, if stopping and switching may be needed,  
9 what the optimal sequencing is of drug treatments. Double-blind randomised controlled  
10 trials examining these different issues would help to improve diabetes care.

11 **9. In adults with type 2 diabetes, what are the long-term effects of blood glucose**  
12 **lowering therapies such as dipeptidyl peptidase-4 (DPP-4) inhibitors, sodium-**  
13 **glucose co-transporter 2 (SGLT2) inhibitors and meglitinides?**

14 **Why this is important**

15 There is limited evidence in relation to adverse events (for example, cardiovascular  
16 outcomes) on the long-term effects (at least 5 years) of blood glucose lowering  
17 therapies, particularly newer agents. Prospective longitudinal studies are needed to  
18 better understand the long-term safety issues surrounding these medicines.

19 **10. In adults with type 2 diabetes, what patient characteristics predict response or**  
20 **non-response to pharmacological blood glucose lowering therapies?**

21 **Why this is important**

22 There is little understanding of the prognostic characteristics that determine the  
23 likelihood that a person would benefit and respond or not respond to treatment.  
24 Increased understanding of important predictive criteria would better help clinicians  
25 target drug therapies and improve overall patient care. Prospective longitudinal cohort  
26 studies examining various types of prognostic factors such as demographic, disease-  
27 specific and comorbid are needed to identify characteristics that are likely to predict  
28 treatment response or non-response to blood glucose lowering therapies in adults with  
29 type 2 diabetes.

30 **11. In adults with type 2 diabetes and multimorbidity, what are the optimal blood**  
31 **glucose lowering treatment strategies?**

32 **Why this is important**

33 The evidence reviewed in this guideline commonly excluded participants with type 2  
34 diabetes whose disease is complicated by significant coexisting conditions, although this  
35 is a common presentation in real-world practice. As a result, it is difficult to account for  
36 the impact of different comorbid conditions on the effectiveness of blood glucose  
37 lowering treatment strategies. A systematic review is needed to ascertain the optimal  
38 treatment strategies for glycaemic control in adults with type 2 diabetes and a range of  
39 comorbid conditions. Multimorbidity covers a wide range of conditions (for example,  
40 heart failure, chronic obstructive pulmonary disease and depression) and each would  
41 have different implications. Therefore, analyses should consider whether the optimal  
42 treatment strategies differ according to specific comorbid conditions.

43

## 8.5.1 Long-term serious adverse effects of blood glucose lowering drug treatments

### 8.5.1.3 Clinical introduction

4 The aim of this review is to provide supplementary information on the long-term serious  
5 adverse effects of the blood glucose lowering drug treatments that were assessed in section  
6 8.4. For cohesiveness, included RCTs in section 8.4 that had relevant data at 2 or more  
7 years are reported in this review. In addition, this review links to the work undertaken by the  
8 Medicines and Healthcare products Regulatory Authority (MHRA) which has a role in  
9 ensuring that medicines such as those for controlling blood glucose are safe for use.

#### 8.5.1.10 Long-term serious adverse effects of drug treatments in Clinical Guideline 66

11 CG66 did not cover the long-term serious adverse effects associated with blood glucose  
12 lowering drug treatments.

#### 8.5.1.23 Long-term serious adverse effects of drug treatments in the update (2015)

14 This is a new question in this update and therefore searches have been carried out for this  
15 topic without any date restrictions (see Appendix C for update search strategies).

### 8.5.26 Evidence review

#### 8.5.2.17 Review question

18 What are the serious adverse effects of long-term use of pharmacological interventions to  
19 control blood glucose in people with type 2 diabetes?

#### 20 Table 87: PICO table

Population	Adults (18 years and over) with type 2 diabetes
Interventions	Acarbose Dipeptidyl peptidase-4 inhibitors (linagliptin, saxagliptin, sitagliptin and vildagliptin) Glucagon-like peptide-1 receptor agonists (conventional and prolonged release exenatide, liraglutide and lixisenatide) Insulin Meglitinides Metformin Sulfonylureas Thiazolidinediones (pioglitazone)
Comparators	Placebo/no treatment or other treatment (including combinations)
Outcomes	Cancer Cardiovascular disease (myocardial infarction, heart failure, stroke, ACS, TIA, revascularisation and stenting) Cognitive impairment Fracture Pancreatic disease Morbidity Mortality

21 Prospective, longitudinal, cohort studies focusing on the development of long-term safety  
22 issues such as renal failure, severe pancreatitis, cancer (for example bladder, thyroid),  
23 cardiac failure and other microvascular or macrovascular complications were considered.

- 1 Studies were included if they had at least 200 participants and a minimum follow-up period of  
2 2 years. Papers were excluded if they:
- 3 • were conference abstracts, letters, editorials and other non-prospective observational  
4 studies (evidence from registries and healthcare databases were considered to be  
5 retrospective)
  - 6 • included a mixed population of people with type 1 and 2 diabetes and either did not report  
7 subgroup analyses, or less than 85% of the study population had type 2 diabetes
  - 8 • included treatment groups that had mixed pharmacological interventions, for example  
9 intensive strategies
  - 10 • included rosiglitazone as part of the drug treatment strategy
  - 11 • did not include comparative data on the exposure to drug treatments
  - 12 • did not report on the incidence of the safety outcomes.
- 13 For the full excluded list, see Appendix L. The detailed protocol is also available in Appendix  
14 C.

### 8.5.2.25 Clinical evidence

16 From the evidence review in section 8.4, 2 included RCTs (Gallwitz et al. 2012; Holman et al.  
17 1999) provided long-term safety data and are reported here.

18 In total, 4669 references were found in the update searches and 5 prospective cohort studies  
19 were included (Aas et al. 2009; Bruno et al. 1999; Fisman et al 2001; Henricsson et al. 1997;  
20 Landman et al. 2010).

21 Studies focused on comparing glucose lowering therapies to each other (and/or dietary  
22 management), either in isolation or in combination with other pharmacological interventions,  
23 or to placebo. Evidence was available on acarbose (Holman et al. 1999), linagliptin (a  
24 dipeptidyl peptidase-4 (DPP-4) inhibitor; Gallwitz et al. 2012), insulin (Aas et al. 2009; Bruno  
25 et al. 1999; Henricsson et al. 1997), metformin (Fisman et al. 2001, Landman et al. 2010)  
26 and sulfonylurea (Bruno et al. 1999; Fisman et al. 2001). No relevant studies were identified  
27 for glucagon-like peptide-1 (GLP-1) receptor agonists, meglitinides and pioglitazone.

28 Pooling of data using meta-analysis was not possible because of differences in the reported  
29 outcomes and/or study designs. Cohort data were also adjusted for confounding factors,  
30 which were not consistent across the included studies. Therefore, results were presented in  
31 modified GRADE profiles, where individual studies rather than outcomes were assessed.

### 8.5.2.2.32 Description of included studies

33 Details of the included studies are found in the evidence tables (see Appendix E).

#### 34 Acarbose

35 One 3-year RCT conducted in the UK including 1946 people (mean age 60 years; mean  
36 duration of diabetes 8 years; mean HbA1c at baseline 72 mmol/mol (8.7%); mean BMI not  
37 reported) provided data for acarbose compared to placebo (UKPDS; Holman et al. 1999).

#### 38 Linagliptin (DPP-4 inhibitor)

39 One 2-year RCT conducted in multiple countries including 1552 people (mean age 59.8  
40 years; mean HbA1c at baseline 61 mmol/mol (7.7%); mean BMI 30.3 kg/m<sup>2</sup>), more than half  
41 of whom had diabetes for at least 5 years provided data for metformin compared to  
42 sulfonylurea (Gallwitz et al. 2012).

## 1 **Insulin**

2 A total of 4208 people (study size ranged from 865 to 1965) were included from 3  
3 prospective cohort studies, carried out in Sweden (Henricsson et al. 1997) and in multiple  
4 countries (Aas et al. 2009; Bruno et al. 1999). The mean age ranged from 54 to 66. Mean  
5 duration of diabetes was reported in 2 studies as 8.5 years; the other study did not report this  
6 information (Aas et al. 2009). Mean HbA1c at baseline was reported in 1 study ranging from  
7 53 to 57 mmol/mol (ranged from 7% to 7.4%) in the different groups (Aas et al. 2009). No  
8 studies reported BMI. Follow-up periods ranged from 3 to 7 years.

## 9 **Metformin**

10 A total of 3628 people (study sizes 1353 and 2275) were included from 2 cohort studies,  
11 carried out in the Netherlands (ZODIAC; Landman et al. 2010) and Israel (Fisman et al.  
12 2001). The mean ages were 67.8 and 60 years. The mean duration of diabetes was reported  
13 in 1 study as 6 years (Landman et al. 2010); the other study did not report this information.  
14 Mean HbA1c levels at baseline was 58 mmol/mol (7.5%) in 1 study (Landman et al. 2010);  
15 the other study did not report this information. Mean BMI was 28.9 and 27.5 kg/m<sup>2</sup>. Follow-up  
16 periods were 7.7 and 10 years.

## 17 **Sulfonylureas**

18 A total of 4240 people (study sizes 1965 and 2275) were included from 2 cohort studies,  
19 carried out in Italy (Bruno et al. 1999) and Israel (Fisman et al. 2001). The mean ages were  
20 66 and 60 years. The mean duration of diabetes was reported in 1 study as 8.5 years (Bruno  
21 et al. 1999); the other study did not report this information. Mean HbA1c levels at baseline  
22 were not reported in either study. Mean BMI was 27.5 kg/m<sup>2</sup> in 1 study (Fisman et al. 2001);  
23 the other study did not report this information. Follow-up periods were 7 and 7.7 years.

24 The summary GRADE tables are presented for this review question (see Appendix D for full  
25 GRADE tables).

26

1 **Table 88: Summary GRADE profile for acarbose**

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
<b>Acarbose plus existing therapy (n=973) compared to placebo plus existing therapy (n=973); mean 3 years follow-up; subgroup of the UKPDS study</b>				
1 (Holman 1999) –UKPDS	RCT	Any diabetes related end point Microvascular disease	RR 1.00 (0.81 to 1.23) RR 0.91 (0.61 to 1.35)	Moderate
<i>Abbreviations: CI confidence intervals; RR relative risk</i>				

2 **Table 89: Summary GRADE profile for linagliptin (dipeptidyl-peptidase 4 inhibitor)**

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
<b>DPP-4 inhibitor (linagliptin) plus metformin (n=776) compared to sulfonylurea (glimepiride) plus metformin (n=775); mean 2 year follow-up; people with type 2 diabetes on a stable dose of metformin</b>				
1 (Gallwitz 2012)	RCT	All-cause mortality Any cardiovascular event <sup>†</sup> Cardiovascular death Myocardial infarction Stroke Admission due to unstable angina	RR not significant RR 0.46 (0.23 to 0.91) RR 1.00 (0.14 to 7.07) RR 0.60 (0.22 to 1.64) RR 0.27 (0.08 to 0.97) RR 1.00 (0.20 to 4.93)	Moderate
<i>Abbreviations: CI confidence intervals; RR relative risk; <sup>†</sup> Any cardiovascular event defined as cardiovascular death, myocardial infarction, stroke and admission due to unstable angina</i>				

3 **Table 90: Summary GRADE profile for insulin**

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
<b>Insulin compared to diet alone (overall n=1941); mean 7 year follow-up; people with type 2 diabetes</b>				

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
1 (Bruno 1999)	cohort	All-cause mortality Cardiovascular mortality Ischaemic heart mortality Cerebrovascular mortality Chronic renal failure	Adj RR 1.71 (1.18 to 2.48) Adj RR 1.35 (0.79 to 2.32) Adj RR 2.95 (1.07 to 8.10) Adj RR 1.00 (0.41 to 2.45) Adj RR 2.26 (0.82 to 6.19)	Very low
<b>Insulin (n=333) compared to oral antidiabetic medication (n=unclear, up to 1045); median 3.1 year follow-up; people with type 2 diabetes attending retinopathy screening</b>				
1 (Henricsson 1997)	cohort	<u>People who changed from oral medication to insulin compared to those remaining on oral medication</u> - Blindness/visual impairment - Progression of retinopathy 3 or more levels	Adj RR 2.7 (1.8 to 4.0) Adj RR 1.6 (1.3 to 1.9)	Very low
<b>Diet alone (n=99) compared to oral antidiabetic drugs (n=250) compared to new insulin users (n=245) compared to existing insulin users (n=271); mean 3 year follow-up; people with type 2 diabetes and suspected myocardial infarction who took part in the DIGAMI RCT (24 hour insulin infusion compared to conventional management)</b>				
1 (Aas 2009) – DIGAMI	cohort	<u>Existing insulin users compared to other groups</u> - cardiovascular death <u>New insulin users compared to other groups</u> - Reinfarction	HR 2.38 (1.34 to 4.22) HR 2.49 (1.23 to 5.03)	Very low
<i>Abbreviations: CI confidence intervals; HR hazard ratio; RR relative risk ; Adj RR adjusted relative risk – see evidence tables for details of individual adjustments that were applied</i>				

**1 Table 91: Summary GRADE profile for metformin**

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
<b>Metformin (n=79) compared to diet alone (n=990); mean 7.7 year follow-up; people with type 2 diabetes and coronary artery disease</b>				
1 (Fisman 2001)	cohort	All-cause mortality	Adj HR 1.19 (0.76 to 1.84)	Very low
<b>Metformin plus existing diabetes therapy (n=289) compared to existing diabetes therapy alone (n=1064); mean 10 year follow-up; unclear population, part of ZODIAC study</b>				

1 (Landman 2010) – ZODIAC	cohort	All-cause mortality Cancer mortality Cardiovascular mortality	Adj HR 0.94 (0.73 to 1.22) Adj HR 0.43 (0.23 to 0.80) Adj HR 2.27 (1.36 to 3.78)	Very low
<b>Metformin plus sulfonylurea (glyburide) (n=253) compared to diet alone (n=990); mean 7.7 year follow-up mean; people with type 2 diabetes and coronary artery disease</b>				
1 (Fisman 2001)	cohort	All-cause mortality	Adj HR 1.53 (1.20 to 1.96)	Very low
<i>Abbreviations: CI confidence intervals; HR hazard ratio; RR relative risk; Adj RR adjusted relative risk – see evidence tables for details of individual adjustments that were applied</i>				

1 **Table 92: Summary GRADE profile for sulfonylurea**

Number of studies	Design	Effect (95% CI)		Quality
		Outcome	Estimate	
<b>Sulfonylurea compared to diet alone (overall n=1941); mean 7 year follow-up; people with type 2 diabetes</b>				
1 (Bruno 1999)	cohort	All-cause mortality Cardiovascular mortality Ischaemic heart mortality Cerebrovascular mortality	Adj RR 1.14 (0.82 to 1.58) Adj RR 1.02 (0.64 to 1.63) Adj RR 1.63 (0.64 to 1.14) Adj RR 1.09 (0.52 to 2.32)	Very low
<b>Glyburide (n=953) compared to diet alone (n=990); mean 7.7 year follow up; people with type 2 diabetes and coronary artery disease</b>				
1 (Fisman 2001)	cohort	All-cause mortality	Adj HR 1.21 (1.02 to 1.44)	Very low
<b>Sulfonylurea plus biguanides compared to diet alone (overall n=1941); mean 7 year follow-up; people with type 2 diabetes</b>				
1 (Bruno 1999)	cohort	All-cause mortality Cardiovascular mortality Ischaemic heart mortality Cerebrovascular mortality	Adj RR 1.13 (0.79 to 1.62) Adj RR 1.04 (0.62 to 1.75) Adj RR 2.49 (0.96 to 6.50) Adj RR 0.91 (0.39 to 2.12)	Very low
<i>Abbreviations: CI confidence intervals; HR hazard ratio; RR relative risk</i>				

### 8.5.2.31 Health economic evidence

- 2 No health economic evidence was found for this question. It was noted that most type 2  
3 diabetes health economic analyses are based on projections of long-term outcomes from  
4 short-term clinical biomarkers, but these do not take account of long-term safety concerns.

### 8.5.35 Evidence statements

#### 8.5.3.16 Clinical evidence

- 7 No relevant studies on glucagon-like peptide-1 receptor agonists, meglitinides and  
8 pioglitazone were identified.
- 9 Evidence on the long-term serious adverse effects associated with other blood glucose  
10 lowering medicines (acarbose, linagliptin, insulin, metformin and sulfonylurea) was provided  
11 by 7 studies (2 randomised controlled trials and 5 prospective cohort studies). The quality of  
12 the evidence ranged from high to very low.
- 13 The overall effects of the reviewed drug treatments on long-term safety outcomes were  
14 unclear. This is because studies were often underpowered to detect differences between the  
15 intervention and comparator groups; it is likely that confounding factors were present in the  
16 data and outcomes were not reported consistently across the included studies. Therefore,  
17 there is uncertainty in the results of the individual studies, and no conclusions can be drawn  
18 about the long-term serious adverse effects of the pharmacological interventions that were  
19 reviewed.

### 8.5.40 Evidence to recommendations

#### 21 Table 93: Linking evidence to recommendations

Relative value of different outcomes	All long-term safety outcomes were considered critical to decision making.  Equal value was placed on all outcomes, since the risk of any serious adverse events was considered to be clinically important.
Trade-off between benefits and harms	People with type 2 diabetes are at risk of long-term microvascular and macrovascular complications. Blood glucose lowering drug treatments that aim to reduce the likelihood of these complications by improving glycaemic control, are also associated with potential harms.  Consideration of the balance between pharmacological benefits and harms against the long-term complications of diabetes is required.  The review question only focused on the serious adverse effects caused by the long-term use of drug treatments. The relative benefits of the pharmacological interventions were reviewed in section 8.4 of this guideline, where the trade-off between benefits and harms was considered in more detail, along with the evidence from this question.
Consideration of health benefits and resource use	No health economic evidence was discussed for this question. The GDG noted that most type 2 diabetes health economic analyses are based on projections of long-term outcomes from short-term clinical biomarkers, but these do not take account of long-term safety concerns.
Quality of evidence	The GDG noted the limited amount of evidence that was identified in the review and discussed the possibility of expanding the study design criteria to include data from registries and databases that were set up to prospectively collect data. The Group agreed that these designs do not address pre-specified hypotheses, have significant methodological limitations such as enrolment biases and are inherently retrospective as

	<p>the study is developed once observations of interest have been made. Therefore, the GDG agreed that such evidence would not add to the review and should not be included. In addition, the Group noted that the MHRA whose specific remit is to examine the benefits and harms of pharmacological interventions and issue regulatory action when necessary, considers all available evidence such as those from databases and registries and therefore, the inclusion of such evidence would also duplicate work already carried out.</p> <p>The GDG discussed the relative quality of RCTs and prospective cohort studies. The Group agreed that the lack of randomisation in cohort studies means that results are likely to be confounded and need to be appropriately adjusted. Although all of the studies did adjust data to take identified confounding factors into consideration, the GDG thought that unidentified confounding factors were likely to be present in the evidence, which cannot be adjusted for by the studies. The Group noted that confounding factors were adjusted inconsistently across the studies such that some adjusted for all known factors, while others to varying degrees. Therefore, the GDG agreed that overall, they could not be confident of the findings of the studies derived from the cohort studies.</p> <p>The Group noted that trials with appropriate randomisation methods are less likely to be affected by biases from confounding factors, since adequate randomisation should lead to an equal balance of known and unknown factors in the randomised arms. However, the GDG expressed concern regarding the reporting of outcomes from the included RCTs such that multiple variations of outcome combinations were used as composite outcomes, which undermines the credibility of the findings.</p> <p>The GDG noted that in most of the studies, the natural progression of diabetes to worsen over time was not addressed, such that it was not clear whether changes or augmentation of drug treatments were considered, which are likely to confound the results.</p> <p>The GDG agreed that overall, the quality of the evidence was low and noted the lack of studies on some pharmacological interventions. In particular, the Group noted that the PROActive trial on pioglitazone was excluded but agreed that long-term serious adverse effects are identified in the MHRA safety alerts. The Group noted the lack of evidence in some serious outcomes such as bone fracture and renal cancer which are new concerns for people using these drug treatments to control blood glucose levels.</p> <p>The GDG agreed that there was insufficient evidence to inform making recommendations regarding the long-term safety of the pharmacological interventions included in the review. The Group thought that the MHRA, with its remit to look at the ongoing safety of pharmacological interventions, is able to provide the most up to date information in this area.</p>
Other considerations	<p>The GDG agreed that the evidence in this review would be considered with the findings on the benefits and shorter term adverse effects in the pharmacological review question in section 8.4, to develop the overall recommendations for these interventions.</p>

### 8.5.51 Recommendations and research recommendations

- 2 See sections 8.4.17 and 8.4.18 for recommendations.

## 9<sub>1</sub> Managing complications

### 9.1.2 Autonomic neuropathy

#### 9.1.1.3 Clinical introduction

4 There are many manifestations of autonomic neuropathy as a complication of long-term  
5 hyperglycaemia. These include gastroparesis, diarrhoea, faecal incontinence, erectile  
6 dysfunction, bladder disturbance, orthostatic hypotension, gustatory and other sweating  
7 disorders, dry feet, and unexplained ankle oedema.

8 Gastroparesis can be one of the more devastating complications of autonomic neuropathy.  
9 While it can present as bloating, nausea and fullness on eating, severe intermittent  
10 hypoglycaemia can be a major problem for people on glucose lowering therapy, while  
11 vomiting may be intermittent and sudden or occasionally severe and protracted.

12 The clinical questions addressed include in whom to suspect gastroparesis might be present,  
13 what medications might help, and what other measures might be taken.

#### 9.1.2.4 Methodological introduction

15 Eight studies were identified in this area all of which involved domperidone, metoclopramide  
16 or erythromycin. Two studies were excluded for methodological reasons.<sup>381,382</sup>

17 The remaining 6 studies comprised 4 RCTs of the drug against placebo; erythromycin vs  
18 placebo,<sup>383</sup> metoclopramide vs placebo,<sup>384,385</sup> domperidone vs placebo,<sup>386</sup> and 2 direct drug  
19 RCT comparisons; metoclopramide vs erythromycin,<sup>387</sup> and domperidone vs  
20 metoclopramide.<sup>388</sup>

21 There were methodological quality issues with these studies, which often involved small  
22 numbers of participants with a range of demographic and clinical details. Furthermore,  
23 although symptom scores were used as measures in 3 studies,<sup>384,385,388</sup> these were not based  
24 on a recognised or validated scale and were not consistent in the measures they recorded or  
25 in the scoring system allotted to the measures. The remaining 3 studies used the SF-36  
26 health-related quality of life tool,<sup>386</sup> gastric emptying using a  $\gamma$ -camera<sup>387</sup> and scintigraphic  
27 studies.<sup>383</sup>

#### 9.1.2.28 Health economic methodological introduction

29 No health economic papers were identified.

#### 9.1.2.20 Evidence statements

#### 9.1.2.31 Drug vs placebo

#### 9.1.2.3.32 *Erythromycin*

33 One crossover study with 10 participants with diabetes and known prolonged gastric  
34 emptying were given 200 mg of IV erythromycin or IV placebo.<sup>383</sup> Ten age and sex matched  
35 health participants were also used as a comparator group. This study used scintigraphic  
36 studies and found that for 60 and 120 minutes IV erythromycin significantly increased gastric  
37 emptying, (measured as the mean percentage simultaneously ingested food retained in the  
38 stomach, for solids), compared with placebo (21±5 vs 85±7, p<0.0005 and 4±1 vs 63±9,  
39 p<0.0005 respectively).

- 1 For liquids the mean percentage retained was significantly lower for the IV erythromycin
- 2 compared with placebo again at both 60 and 120 minutes (22±5 vs 54±5,  $p<0.0005$  and 9±3
- 3 vs 32±4,  $p<0.005$  respectively).
- 4 IV erythromycin was also found to have increased gastric emptying for solids at 60 minutes
- 5 when compared with healthy subjects in the comparator group ( $p<0.05$ ).
- 6 There were no AEs found with this study, this study had a further open-label phase with oral
- 7 erythromycin, not reported here. **Level 1+**

#### 9.1.2.3.28 *Metoclopramide*

9 Two studies,<sup>384,385</sup> one of which was a crossover study,<sup>384</sup> were identified comparing oral  
10 metoclopramide 10 mg QID and placebo, both studies used the diary recording of symptoms  
11 and though the scales used were broadly similar they were not identical, there were no major  
12 AEs identified in either study.

13 One study identified that the mean symptom scores for the 3-week treatment phase was  
14 significantly less for metoclopramide than for placebo; 26.5±3.7 vs 45.3±7.8,  $p<0.01$ . This  
15 study also found that the mean individual scores for 4/5 symptoms (fullness, pressure and  
16 bloating, nausea, vomiting, anorexia) showed that metoclopramide significantly reduced the  
17 symptoms compared with placebo ( $p<0.05$ ).<sup>385</sup>

18 The crossover study found that symptom improvement was significantly greater for  
19 metoclopramide than placebo for nausea at weeks 1 and 3 ( $p<0.05$ ). This was also found for  
20 fullness at weeks 2 and 3 ( $p<0.05$ ). Changes found for other symptoms were not significantly  
21 improved for metoclopramide compared with placebo.<sup>384</sup> **Level 1+**

#### 9.1.2.3.32 *Domperidone*

23 One study<sup>386</sup> considered domperidone vs placebo, this study combined a 4-week period  
24 where participants took 20 mg domperidone QID (single-blind phase) orally, followed by a 4-  
25 week period of 20 mg domperidone QID or placebo (double-blind phase). Entry into the  
26 second phase was dependent on a decrease on the baseline symptom score, those classed  
27 as responders, following completion of the single-blind phase.

28 Single-blind phase: significant symptomatic improvement was found at the end of the single-  
29 blind phase ( $p<0.0001$ ). Improvements were also noted in the health-related quality of life  
30 measured on the SF-36 scale (all domains  $p<0.001$ , except physical functioning,  $p<0.01$ ).

31 Double-blind phase: symptom severity increased with both domperidone and placebo,  
32 though they did not return to baseline levels, this increase in severity was greater for placebo  
33 compared with domperidone ( $p<0.05$ ). AEs were not reported. **Level 1+**

#### 9.1.2.4 **Head-to-head comparisons**

##### 9.1.2.4.35 *Metoclopramide vs erythromycin*

36 One crossover study with 13 participants considered erythromycin 250 mg TID with  
37 metoclopramide 10 mg TID.

38 Gastric emptying was considered at 60 and 90 minutes and while significant improvements  
39 were found for both drugs there was no significant difference found between the effects  
40 between erythromycin and metoclopramide.

41 The symptom score was significantly less for erythromycin; 2(0–5), than for metoclopramide;  
42 3(0–11),  $p<0.05$ .

43 No serious AEs were noted, though N=2 of the patients did have weakness, sedation and leg  
44 cramps with metoclopramide. **Level 1+**

#### 9.1.2.4.21 *Domperidone vs metoclopramide*

2 One study with 95 participants considered domperidone 20 mg QID with metoclopramide 10  
3 mg QID. Gastroparetic symptoms and tolerability were assessed, it should be noted for  
4 tolerability assessment participants were specifically asked about central nervous system  
5 (CNS) associated side effects; these have previously been identified in association with  
6 metoclopramide.

7 Although significant reductions in symptoms were found with both domperidone and  
8 metoclopramide, there was no significant difference found between the 2 treatments.

9 For tolerability, at week 2 the severity of somnolence ( $p < 0.001$ ), akathisia ( $p = 0.03$ ), anxiety  
10 ( $p = 0.02$ ) and depression ( $p = 0.05$ ) were significantly greater for metoclopramide than for  
11 domperidone ( $p < 0.001-0.05$ ). While at week 4 this was found for severity of somnolence  
12 ( $p = 0.03$ ) and reduced mental acuity ( $p = 0.04$ ). **Level 1+**

#### 9.1.33 Evidence to recommendations

14 The evidence reported had methodological limitations, notably studies of small sample sizes.  
15 The GDG agreed that there is a poor evidence base for the treatment of gastroparesis.  
16 Nevertheless they noted that the evidence reported suggested that the prokinetic drugs,  
17 metoclopramide, domperidone, along with erythromycin, were all effective in at least some  
18 people with gastroparesis resulting from autonomic neuropathy. On consideration of the  
19 evidence it was not possible to distinguish usefully between the prokinetic drugs. The group  
20 agreed that choice of initial therapy should be based on tolerability issues, including drug  
21 interactions. It was noted that differential diagnosis can be difficult, and the diagnostic tests  
22 not secure, while serious prolonged vomiting could become a medical emergency.  
23 Accordingly referral beyond diabetes services is sometimes indicated.

24 While the group gave priority to medication for the management of this condition, clinical  
25 experience suggested that non-pharmacological approaches including postural advice and  
26 timing of ingestion of fluids and solids could prove useful to some people.

#### 9.1.47 Recommendations

28 **70. Think about a diagnosis of gastroparesis in adults with type 2 diabetes with**  
29 **erratic blood glucose control or unexplained gastric bloating or vomiting, taking**  
30 **into account possible alternative diagnoses. [2009, amended 2015]**

31 **71. Consider a trial of metoclopramide, domperidone or erythromycin<sup>n</sup> for an adult**  
32 **with type 2 diabetes with gastroparesis<sup>o</sup>. [2009, amended 2015]**

33 **72. If gastroparesis is suspected, consider referral to specialist services if:**

- 34                   • the differential diagnosis is in doubt **or**  
35                   • persistent or severe vomiting occurs. **[2009]**

36

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<sup>n</sup> At the time of consultation (January 2015), metoclopramide, domperidone and erythromycin did not have a UK marketing authorisation for this indication. The prescriber should follow relevant professional guidance, taking full responsibility for the decision. Informed consent should be obtained and documented. See the General Medical Council's Good practice in prescribing and managing medicines and devices for further information.  
<sup>o</sup> Diagnosis of gastroparesis needing specific therapy can only be made in the absence of hyperglycaemia at the time of testing, because hyperglycaemia induces a physiological delay in gastric emptying.

## 9.2<sub>1</sub> Nerve damage

### 9.2.1<sub>2</sub> Other aspects of autonomic neuropathy

#### 9.2.1.1<sub>3</sub> Clinical introduction

4 Other aspects of autonomic neuropathy, including diarrhoea, faecal incontinence, bladder  
5 disturbance, orthostatic hypotension, gustatory and other sweating disorders, dry feet, and  
6 unexplained ankle oedema, can offer diagnostic and management problems, and on  
7 occasion be very disabling.

8 Alternatively symptoms may be vague and may present insidiously without realisation that  
9 they are diabetes-related, while nerve damage can be also be found in asymptomatic people.  
10 A mixed presentation is common, may be exacerbated by other drug therapy (e.g. tricyclic  
11 drugs), and may give troublesome hypoglycaemia. People with advanced autonomic  
12 neuropathy may also have advanced retinopathy, nephropathy, and somatic neuropathy.

#### 9.2.1.2<sub>3</sub> Evidence to recommendations

14 The GDG reviewed the opinion-based recommendations made in the NICE type 1 diabetes  
15 guideline 2004.<sup>26</sup> They were found for the most part appropriate, and are reproduced with  
16 some editorial change only. It was recognised that these recommendations are for the most  
17 part identification and diagnostic issues, and that specialist management where required  
18 would often lie outside diabetes services.

#### 9.2.1.3<sub>9</sub> Recommendations

##### 9.2.1.3.2<sub>0</sub> Painful neuropathy

21 **73. For guidance on painful neuropathy in adults with type 2 diabetes, see the NICE**  
22 **guideline on neuropathic pain – pharmacological management. [new 2015]**

##### 9.2.1.3.2<sub>3</sub> Autonomic neuropathy

24 **74. Think about the possibility of contributory sympathetic nervous system damage**  
25 **for adults with type 2 diabetes who lose the warning signs of hypoglycaemia.**  
26 **[2009, amended 2015]**

27 **75. Think about the possibility of autonomic neuropathy affecting the gut in adults**  
28 **with type 2 diabetes who have unexplained diarrhoea that happens particularly at**  
29 **night. [2009, amended 2015]**

30 **76. When using tricyclic drugs and antihypertensive drug treatments in adults with**  
31 **type 2 diabetes who have autonomic neuropathy, be aware of the increased**  
32 **likelihood of side effects such as orthostatic hypotension. [2009]**

33 **77. Investigate the possibility of autonomic neuropathy affecting the bladder in adults**  
34 **with type 2 diabetes who have unexplained bladder-emptying problems. [2009]**

35 **78. In managing autonomic neuropathy symptoms, include specific interventions**  
36 **indicated by the manifestations (for example, for abnormal sweating or nocturnal**  
37 **diarrhoea). [2009]**

**9.2.1.3.31 *Diabetic foot problems***

- 2 **79. For guidance on managing foot problems in adults with type 2 diabetes, see the**  
3 **NICE guideline on diabetic foot problems. [new 2015]**

**9.2.1.3.44 *Diabetic kidney disease***

- 5 **80. For guidance on nephropathy in adults with type 2 diabetes, see the NICE**  
6 **guideline on chronic kidney disease. [new 2015]**

7

## 9.3.1 Erectile dysfunction

### 9.3.1.2 Clinical introduction

3 People with type 2 diabetes have an increased risk of microvascular complications, and  
4 damage to small blood vessels and autonomic nerves may affect sexual stimulation and  
5 response, leading to erectile dysfunction in men.

6 This section addressed whether pharmacological treatment (either alone or in combination)  
7 should be used to manage erectile dysfunction. This review also looked at whether the use of  
8 pharmacological treatments should be restricted to specific subgroups of the population and  
9 what adverse events are associated with their use.

#### 9.3.1.10 Erectile dysfunction in Clinical Guideline 66

11 The pharmacological management of erectile dysfunction was originally covered as part of  
12 CG66 and included men with both type 1 and type 2 diabetes. The original searches were  
13 conducted from 2001 to 2007 (see Appendix G for search strategies from CG66). Update  
14 searches have been carried out for this topic with a date restriction of 2007 to June 2014 for  
15 phosphodiesterase 5 (PDE-5) inhibitors, and no date restrictions for alprostadil and  
16 testosterone therapy (see Appendix C for update search strategies) as these terms had not  
17 previously been searched. The evidence considered in this review question in CG66 included  
18 1 systematic review and 9 RCTs.

#### 9.3.1.29 Erectile dysfunction in the update (2015)

20 CG66 focused on the use of PDE-5 inhibitors for the management of erectile dysfunction. For  
21 this update, the review question has been expanded to cover the use of alprostadil and  
22 testosterone therapy (see Appendix C for full review protocols).

### 9.3.2.3 Evidence review

#### 9.3.2.14 Review question

25 What pharmacological treatment should be used to manage erectile dysfunction in men with  
26 type 2 diabetes?

#### 27 Table 94: PICO table

Population	Men (18 years and over) with diabetes (including type 1 and type 2)
Interventions	Testosterone therapy, phosphodiesterase 5 (PDE-5) inhibitors and alprostadil (alone or in combination)
Comparators	Placebo, standard care (or other treatment)
Outcomes	Erectile function (assessed using validated scale/measure such as International Index of Erectile Function; IIEF) Adverse events Health-related quality of life

28 RCTs examining the use of alprostadil, PDE-5 inhibitors and testosterone therapy (alone or  
29 in combination) for the management of erectile dysfunction in men with diabetes were  
30 included. Papers were excluded if they:

- 31 • were non-randomised (including cohort, case-control and case series studies), narrative
- 32 reviews, conference abstracts, letters and editorials
- 33 • focused on the diagnosis of erectile dysfunction

- 1 • assessed the use of testosterone therapy in men who did not have erectile dysfunction.
- 2 For the full excluded list, see Appendix L. The detailed protocol is also available in Appendix
- 3 C.
- 4 The main outcomes for this review question were erectile function and adverse events.
- 5 Erectile function was assessed using 4 main measures:
- 6 • Erectile function (EF) domain of the international index of erectile function (IIEF)
  - 7 questionnaire
  - 8 • Question 2 from the sexual encounter profile (SEP-2) relating to success in penetration
  - 9 • Question 3 from the sexual encounter profile (SEP-3) relating to success in intercourse
  - 10 • Global efficacy question (GEQ) relating to whether treatment improved erections.
- 11 Where possible, studies were pooled using meta-analysis techniques (pairwise
- 12 comparisons). The GDG agreed that it was not clinically appropriate to undertake a network
- 13 meta-analysis for the available evidence on PDE-5 inhibitors because of the heterogeneity of
- 14 the studies in terms of population, interventions, outcomes and quality.

### 9.3.2.25 Clinical evidence

16 The evidence that was originally included in CG66 was re-reviewed as part of the update,

17 and all were found to be relevant. The Cochrane systematic review included in CG66 on

18 PDE-5 inhibitors had not been updated (Vardi and Nini 2007). Full text papers of the relevant

19 RCTs included in the Cochrane review were obtained and these were preferentially used.

20 Data for Escobar-Jimenez (2002) was taken from the Cochrane systematic review.

21 In total, 349 references were found for this review question and 15 RCTs were included

22 (Boulton et al. 2001; Buvat et al. 2006; Deyoung et al. 2012; Escobar-Jimenez 2002;

23 Goldstein et al. 2003, 2012; Hackett et al. 2013; Hatzichristou et al. 2008; Ishii et al. 2006;

24 Kamenov 2011; Rendell et al. 1999; Saenz de Tejada et al. 2002; Safarinejad 2004; Stuckey

25 et al. 2003; Ziegler et al. 2006). One trial used a crossover design (Buvat et al. 2006). Four

26 studies included people with type 2 diabetes only (Boulton et al. 2001; Deyoung et al. 2012;

27 Escobar-Jimenez 2002; Hackett et al. 2013), 2 studies included people with type 1 diabetes

28 only (Stuckey et al. 2003; Ziegler et al. 2006), 1 study did not report the proportion of people

29 with type 1 and 2 diabetes (Ishii et al. 2006) and the remaining 8 studies included

30 populations with type 2 diabetes ranging from 80% to 90.7%. All but 3 studies (Escobar-

31 Jimenez 2002; Hackett et al. 2013; Kamenov 2004) specified in the inclusion criteria that

32 participants should be heterosexual males or with a female partner. No relevant studies on

33 alprostadil were identified.

34 The following comparisons were included as part of this review question:

- 35 • PDE-5 inhibitors versus placebo – 12 trials; 1 on avanafil (Goldstein et al. 2012), 6 on
- 36 sildenafil (Boulton et al. 2001; Deyoung et al. 2012; Escobar-Jimenez 2002; Rendell et al.
- 37 1999; Safarinejad 2004; Stuckey et al. 2003), 2 on tadalafil (Hatzichristou et al. 2008;
- 38 Saenz de Tejada et al. 2002) and 3 on vardenafil (Goldstein et al. 2003; Ishii et al. 2006;
- 39 Ziegler et al. 2006)
- 40 • PDE-5 inhibitors versus PDE-5 inhibitors – 2 trials (Buvat et al. 2006; Kamenov 2011)
- 41 • Testosterone replacement therapy versus placebo – 1 trial (Hackett et al. 2013)

### 9.3.2.2.42 Description of included studies

43 Details of the included studies are found in the evidence tables (see Appendix G).

### 1 **PDE-5 inhibitors versus placebo**

2 A total of 3513 people (study size ranged from 24 to 778) were included from 12 RCTs,  
3 carried out in the USA (Goldstein et al. 2003, 2012; Rendell et al. 1999), Canada (Deyoung  
4 et al. 2012), Germany (Ziegler et al. 2006), Spain (Escobar-Jimenez 2002; Saenz de Tejada  
5 et al. 2002), Iran (Safarinejad 2004), Japan (Ishii et al. 2006) and multiple countries (Boulton  
6 et al. 2001; Hatzichristou et al. 2008); 1 study did not report this information (Stuckey et al.  
7 2003). The mean age ranged from 46 to 59 years, with 1 study not reporting this information  
8 (Escobar-Jimenez 2002). The mean duration of diabetes in 4 studies ranged from 11 to 12  
9 years, with the remaining 8 studies not reporting this information. Mean HbA1c levels at  
10 baseline were not reported by any studies. Mean BMI in 4 studies ranged from 27.1 to 30.7  
11 kg/m<sup>2</sup>, with 8 studies not reporting this information. Follow-up periods ranged from 10 to 16  
12 weeks.

### 13 **PDE-5 inhibitors versus PDE-5 inhibitors**

14 A total of 811 people (study sizes 49 and 762) were included from 2 RCTs, carried out in the  
15 Bulgaria (Kamenov 2004) and in different countries (Buvat et al. 2006). The mean ages were  
16 50 and 57 years. The mean duration of diabetes was 9.5 and 10.8 13 years. Mean HbA1c  
17 levels at baseline were not reported by either study. Mean BMI was 29.2 and 28.7 kg/m<sup>2</sup>. A  
18 follow-up period of 12 weeks was reported by 1 study (Buvat et al. 2006), but this information  
19 was unclear in the other study. One trial compared different treatment regimens of the same  
20 drug (Buvat et al. 2006), while the other compared two different drugs (Kamenov 2004).

### 21 **Testosterone replacement therapy versus placebo**

22 One 30 week trial conducted in the UK including 199 people diagnosed with type 2 diabetes  
23 and hypogonadism (mean age 61.6 years; mean duration of diabetes and HbA1c levels not  
24 reported; mean BMI 32.7 kg/m<sup>2</sup>), 84.% of whom were diagnosed with erectile dysfunction,  
25 compared intramuscular testosterone undecanoate with placebo (Hackett et al. 2013).

26 The summary GRADE tables are presented for this review question, the full versions can be  
27 found in Appendix D.

28

1 **Table 95: Summary GRADE profile for PDE-5 inhibitors versus placebo**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	PDE-5 inhibitor	Placebo		
<b>Erectile Function using the International Index of Erectile Function [IIEF] mean score on EF domain; 12 to 16 week follow-up</b>				
11 (Boulton 2001; Escobar-Jimenez 2002; Goldstein 2003, 2012; Hatzichristou 2008; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003; Ziegler 2006)	2142	1174	MD 5.58 (4.48 to 6.68)	Low
<b>Erectile function using the Sexual Encounter Profile mean scores of SEP Q2 (successful insertion); 12 week follow-up</b>				
5 (Goldstein 2003, 2012; Hatzichristou 2008; Ishii 2006; Ziegler 2006)	1059/1559	274/616	RR 1.47 (1.33 to 1.61)	Low
<b>Erectile function using the Sexual Encounter Profile mean scores of SEP Q3 (successful intercourse); 12 week follow-up</b>				
5 (Goldstein 2003, 2012; Hatzichristou 2008; Ishii 2006; Ziegler 2006)	800/1551	160/618	RR 1.87 (1.61 to 2.16)	Low
<b>Erectile function-using the Global Efficacy Question mean scores of GEQ (global improvement); 12 to 16 week follow-up</b>				
8 (Boulton 2001; Escobar-Jimenez 2002; Goldstein 2003; Hatzichristou 2008; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003)	623/1064	116/743	RR 3.62 (2.57 to 5.09)	Moderate
<b>Any adverse events; 12 to 16 week follow-up</b>				
11 (Boulton 2001; Escobar-Jimenez 2002; Goldstein 2003, 2012; Hatzichristou 2008; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003; Ziegler 2006)	610/9064	115/5249	RR 2.69 (1.87 to 3.86)	Low
<b>Headache; 12 to 16 week follow-up</b>				
10 (Boulton 2001; Escobar-Jimenez 2002; Goldstein 2003, 2012; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003; Ziegler 2006)	185/2065	43/1126	RR 3.08 (1.46 to 6.48)	Low
<b>Flushing; 12 to 16 week follow-up</b>				

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Number of RCTs	Number of people		Effect (95% CI)	Quality
	PDE-5 inhibitor	Placebo		
10 (Boulton 2001; Escobar-Jimenez 2002; Goldstein 2003, 2012; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003; Ziegler 2006)	191/2065	6/1126	RR 8.65 (4.5 to 16.66)	Low
<b>Bronchitis; 12 to 16 week follow-up</b>				
1 (Ziegler 2006)	3/163	4/155	RR 0.71 (0.16 to 3.14)	Moderate
<b>Upper respiratory tract infections; 12 to 16 week follow-up</b>				
7 (Goldstein 2003, 2012; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Ziegler 2006)	147/1814	43/875	RR 1.12 (0.57 to 2.2)	Low
<b>Discontinuation due to adverse events; 12 to 16 week follow-up</b>				
9 (Goldstein 2003, 2012; Hatzichristou 2008; Ishii 2006; Rendell 1999; Saenz de Tejada 2002; Safarinejad 2004; Stuckey 2003; Ziegler 2006)	46/2013	14/1167	RR 1.67 (0.89 to 3.13)	Low
<b>Dyspepsia; 12 to 16 week follow-up</b>				
4 (Boulton 2001; Goldstein 2012; Rendell 1999; Stuckey 2003)	26/601	2/465	RR 6.09 (1.77 to 20.94)	Moderate
<b>Abnormal vision; 12 to 16 week follow-up</b>				
3 (Boulton 2001; Rendell 1999; Stuckey 2003)	12/343	3/335	RR 2.92 (0.71 to 11.99)	Moderate

Abbreviations: 95%CI, 95% confidence interval; IIEF International Index of Erectile Function questionnaire; EF Erectile function domain of IIEF; MD mean difference; SEP Sexual Encounter Profile (diary questions regarding sexual encounter); GEQ Global Efficacy Question; RR risk ratio

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1 Table 96: Summary GRADE profile of subgroup analyses by baseline HbA1c level for PDE-5 inhibitors versus placebo

Number of RCTs	Number of people		Measure of effect	Quality
	Intervention	Placebo		
<b>Erectile Function (measured with International Index of Erectile Function [IIEF] mean score on EF domain</b>				

Number of RCTs	Number of people		Measure of effect	Quality
	Intervention	Placebo		
<b>Sildenafil versus placebo</b>				
1 (Boulton 2001)	47	47	Mean change from baseline stratified by baseline HbA1c level: <ul style="list-style-type: none"> <li>• &lt;8.3%: 8.9* with sildenafil versus 0.6 with placebo</li> <li>• ≥8.3%: 8.2* with sildenafil versus -0.5 with placebo</li> </ul>	Moderate
<b>Vardenafil versus placebo</b>				
1 (Ziegler 2006)	154	149	Mean end point stratified by baseline HbA1c level: <ul style="list-style-type: none"> <li>• Good (&lt;7%): 21* with vardenafil versus 15 with placebo</li> <li>• Moderate (7-8%): 21* with vardenafil versus 14 with placebo</li> <li>• Poor (&gt;8%): 18* with vardenafil versus 16 with placebo</li> </ul> Interaction term between treatment and level of glycaemic control was not statistically significant	Moderate
<b>Tadalafil versus placebo</b>				
2 (Hatzichristou 2008; Saenz de Tejada 2002)	339	169	Mean change from baseline stratified by baseline HbA1c level <ul style="list-style-type: none"> <li>• Good (&lt;7%): 3.8 (2.5 mg), 6.6 (5 mg) 9.7 (10 mg), 8.3 (20 mg) with tadalafil versus -1.0, 1.4 with placebo</li> <li>• Fair (7-9.5%): 7.3 (2.5 mg), 3.2 (5 mg), 6.0 (10 mg), 6.7 (20 mg) with tadalafil versus -0.9, 1.4 with placebo</li> <li>• Poor (&gt;9.5%): 1.4 (2.5 mg), 4.7 (5 mg), 3.8 (10 mg), 8.3 (20 mg) with tadalafil versus 3.9, 0.5 with placebo</li> </ul>	Very low
* <i>p</i> <0.0001 versus placebo				

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1 Table 97: Summary GRADE profile for PDE-5 inhibitor versus PDE-5 inhibitor

Number of RCTs	Number of people		Measure of effect	Quality
	Intervention	Comparator		
	Tadalafil on demand	Tadalafil 3 times per week		
<b>Erectile Function-using the International Index of Erectile Function [IIEF] mean score on EF domain; 12 week follow-up</b>				
1 (Buvat 2006)	762	762	Mean change from baseline 8.9 (SE 0.3) on demand versus 9.1 (SE 0.3) for 3 times per week	Low
<b>Erectile function using the Sexual Encounter Profile mean scores of SEP Q2 (successful insertion); 12 week follow-up</b>				

Number of RCTs	Number of people		Measure of effect	Quality
	Intervention	Comparator		
1 (Buvat 2006)	762	762	Percentage of people answering 'yes' at end point was 73.0% on demand versus 74.9% for 3 times per week (p<0.05)	Low
<b>Erectile function using the Sexual Encounter Profile mean scores of SEP Q3 (successful intercourse); 12 week follow-up</b>				
1 (Buvat 2006)	762	762	Percentage of people answering 'yes' at end point was 58.0% on demand and 60.5% for 3 times per week (p<0.05).	Low
<b>Treatment emergent adverse events</b>				
1 (Buvat 2006)	762	762	<ul style="list-style-type: none"> <li>• Back pain: 2.5% on demand versus 2.1% 3 times per week</li> <li>• Dyspepsia: 5.9% on demand versus 5.8% 3 times per week</li> <li>• Flushing: 1.6% on demand versus 2.1% 3 times per week</li> <li>• Headache: 4.7% on demand versus 5.6% 3 times per week</li> <li>• Myalgia: 1.4% on demand versus 2% 3 times per week</li> </ul>	Low
	<b>Tadalafil</b>	<b>Vardenafil</b>		
<b>Any adverse events</b>				
1 (Kamenov 2004)	7/24	6/25	<ul style="list-style-type: none"> <li>• Dyspepsia: 8.4% with tadalafil versus 4% with vardenafil</li> <li>• Flushing: 4.2% with tadalafil versus 8% with vardenafil</li> <li>• Headache: 8.3% with tadalafil versus 8% with vardenafil</li> <li>• Myalgia: 8.4% with tadalafil versus 0% with vardenafil</li> <li>• Nasal congestion: 0% with tadalafil versus 8% with vardenafil</li> </ul>	Low

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1 **Table 98: Summary GRADE profile for testosterone therapy versus placebo**

Number of RCTs	Number of people		Effect (95% CI)	Quality
	Testosterone	Placebo		
<b>Erectile Function domain of IIEF questionnaire</b>				
1 (Hackett 2013)	91	95	Mean difference 3.47 (0.40 to 6.54)	Low
<b>Adverse event (total dropouts)</b>				
1 (Hackett 2013)	4/97	5/102	Relative risk 0.84 (0.23 to 3.04)	Low

2

### 9.3.2.31 Health economic evidence

2 Literature searches were carried out to find any existing cost utility analyses (CUAs) of the  
3 pharmacological management of erectile dysfunction in people with type 2 diabetes (see  
4 appendix C for search strategies). In total 88 articles were returned, and 2 CUAs were  
5 retained (Smith and Roberts 2000; Stolk et al. 2000). However neither of these studies was  
6 specific to a diabetic population. The GDG felt it might be possible to extrapolate from the  
7 general erectile dysfunction population to the type 2 diabetes erectile dysfunction population,  
8 so the searches were re-run without the type 2 diabetes search terms. This produced a  
9 further 1 CUA which again was not specific to the type 2 diabetes population (Aspinall et al.  
10 2011) but it did specify that no difference in clinical effectiveness by risk factor (including type  
11 2 diabetes) had been found.

12 None of the 3 studies compared different pharmacological treatments for erectile dysfunction  
13 – 1 study compared sildenafil to no treatment (Smith and Roberts 2000), 1 study compared  
14 sildenafil to injection therapy (Stolk et al. 2000) and 1 study compared different doses of  
15 vardenafil (Aspinall et al. 2011). No studies included avanafil or tadalafil.

16 None of the 3 studies were specific to the UK setting; 2 studies were model based analyses  
17 (Smith and Roberts 2000), (Stolk et al. 2000) whilst the third study (Aspinall et al. 2011) was  
18 a decision tree based on an RCT.

19 Two of the studies (Smith and Roberts 2000), (Aspinall et al. 2011) used a 3% discount rate  
20 (instead of a 3.5% discount rate) and 1 study (Stolk et al. 2000) did not specify whether a  
21 discount rate was used.

22 All studies used similar utility decrements for the erectile dysfunction state. 1 study (Aspinall  
23 et al. 2011) used a utility decrement of 0.13 that was taken from another included study  
24 (Smith and Roberts 2000) which in turn was taken from an American time trade off study in  
25 the context of prostate cancer. 1 study (Stolk et al. 2000) undertook their own population  
26 based time trade off study which produced a utility decrement of 0.13. It is not known  
27 whether the utility decrement for erectile dysfunction in people with type 2 diabetes is likely to  
28 differ from that in the general population.

29 All 3 studies found that the new treatment (sildenafil (Smith and Roberts 2000), (Stolk et al.  
30 2000) or vardenafil (Aspinall et al. 2011)) was cost effective compared to the alternative  
31 chosen, at the usually accepted thresholds. For sildenafil, Smith and Roberts (2000) reported  
32 incremental cost effectiveness ratios (ICERs) less than \$12,000/QALY; Stolk et al. (2000)  
33 reported ICERs less than £4000/QALY. For vardenafil, Aspinall et al. (2011) reported ICERs  
34 below \$6000/QALY. All 3 studies assessed the uncertainty in the ICERs and found that,  
35 whilst the results were sensitive to some inputs, the ICERs were likely to remain below  
36 conventional thresholds in the majority of cases.

37 This question was not prioritised by the GDG for de novo economic modelling.

38

1 Table 99: Economic evidence table for erectile dysfunction in the general population

Study, Population, Comparators and Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Cost	Effect	ICER		
<p><b>Aspinall et al, 2011</b> Hypothetical cohort of USA male veterans aged 60 with ED Vardenafil compared 0 doses to 4, 6 or 8 doses per month</p> <p><b>Partly applicable</b><sup>a,d,e</sup></p> <p><b>Potentially serious limitations</b><sup>a,b,c</sup></p>	<p><u>Effects:</u> Systematic review (PDE5s) <u>Costs:</u> VA pharmacy data (\$, 2009) <u>Utilities:</u> Baseline and ED as Smith (2000); increased gain by dose assumed</p>	<p>Markov model with lifetime time horizon Systematic review found no difference in efficacy for diabetes Estimates of extra doses utilities were conservative No mortality effect, no loss of treatment effect AEs not modelled, states same rates as placebo</p>	<p>0/month \$0 4/month \$707.70 6/month \$353.90 8/month \$353.90</p>	<p>0/month 0 QALYs 4/month 1.23 6/month 0.14 8/month 0.07</p>	<p>0/month Not applicable 4/month \$576 /QALY 6/month \$2585 /QALY 8/month \$5169 /QALY</p>	<p>Providing extra monthly doses of Vardenafil is cost effective compared with less monthly doses at \$50,000/QALY threshold</p>	<p>ICER sensitive to utility for 6/8 month and drug cost. ICER remains &lt; \$50,000 if 6/8 month QALY gain &gt; 0.001 (base case 0.01), drug cost &lt; \$15 (base case \$2, UK equivalent £3.50). In PSA, 6 doses was favoured 84% and 8 doses 61% at \$50,000/ QALY</p>
<p><b>Smith, 2000</b> Hypothetical cohort of USA males aged 60 with ED Sildenafil compared to no treatment</p> <p><b>Partly applicable</b><sup>a,d,e,g</sup></p> <p><b>Potentially serious limitations</b><sup>a,b,f</sup></p>	<p><u>Effects:</u> RCTs <u>Costs:</u> US\$ 1998; drugs wholesale price; AEs estimated <u>Utilities:</u> Baseline US TTO; ED from prostate cancer screening study; AEs estimated</p>	<p>Markov model with lifetime time horizon No treatment assumed to incur no costs AEs: assumed No external funding listed Supported by VA Centre for Medication Safety</p>	<p>Treated v not \$3970<sup>h</sup>  \$3950<sup>i</sup></p>	<p>Treated v not 0.3519<sup>h</sup>  0.351<sup>i</sup></p>	<p>Treated v not  \$11,290/ QALY<sup>h</sup>  \$11,230/ QALY<sup>i</sup></p>	<p>Sildenafil treatment is cost effective compared with no treatment at \$50,000/ QALY threshold</p>	<p>Remains cost effective when assumptions biased against treatment<sup>g</sup>. If utility gain &gt; 0.05 (base case 0.13), ICER remained &lt; \$50,000/ QALY<sup>g</sup> In PSA, sildenafil was favoured 98% of times at \$50k/QALY<sup>g</sup></p>
<p><b>Stolk et al. (2000)</b> RCT (n=532) Sildenafil compared to usual treatment (injection therapy)</p> <p><b>Partly applicable</b><sup>a,d,e</sup></p> <p><b>Potentially serious limitations</b><sup>a,f,j</sup></p>	<p><u>Effects:</u> RCT, expert opinion. Uptake assumed <u>Costs:</u> 1999, drugs data, resource use estimated <u>Utilities:</u> Dutch population TTO, assumed same both treatments</p>	<p>RCT based decision tree with 5 year time horizon Likely to under estimate utility gain due to RCT ITT and QoL assumptions Funded by industry</p>	<p>Year 1 £28,368  Year 5 £89,226</p>	<p>Year 1 7.79 QALYs  Year 5 33.92</p>	<p>Year 1 £3639/ QALY  Year 5 £2630/ QALY</p>	<p>Sildenafil is cost effective compared to usual care at £20,000/QALY threshold</p>	<p>ICER sensitive to dosing frequency (base case 1/week), utility gain and effectiveness. But in worst case scenario, ICER remains &lt;£10k/QALY</p>

Study, Population, Comparators and Quality	Data Sources	Other Comments	Incremental			Conclusions	Uncertainty
			Cost	Effect	ICER		
<p>a Not UK based                      b Not 3.5% discount rate                      c No non-drug costs included                      d Does not compare relevant treatments                      e Not diabetes specific                      f Drug costs appear high                      g Analysis mainly from societal not NHS perspective                      h Societal perspective                      i Third party payer perspective                      j Potential conflict of interest</p> <p>AEs: adverse events                      ED: erectile dysfunction                      ICER: incremental cost effectiveness ratio                      PDE5s:                      QALY: quality adjusted life year                      QoL: quality of life                      USA: United States of America                      VA: Veterans Association</p>							
1							
2							

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#### 9.3.2.41 Evidence statements

#### 9.3.2.52 Clinical evidence

3 No relevant studies on alprostredil were identified.

#### 9.3.2.5.14 PDE-5 inhibitors versus placebo

5 Overall, evidence from 4 meta-analyses including data from up to 11 trials showed on 4  
6 different assessment scales, a significant improvement in erectile function with PDE-5  
7 inhibitors compared to placebo up to 16 weeks. The quality of the evidence ranged from  
8 moderate to low. Four RCTs showed no difference in erectile function outcomes based on  
9 baseline HbA1c levels. The quality of the evidence ranged from moderate to very low.

10 Evidence from 4 meta-analyses including data from up to 11 trials showed a significant  
11 increase in risk of any adverse events, dyspepsia, flushing and headache with PDE-5  
12 inhibitors compared to placebo up to 16 weeks. The quality of the evidence ranged from  
13 moderate to low.

#### 9.3.2.5.24 PDE-5 inhibitors versus PDE-5 inhibitors

15 Two small trials provided no conclusive findings regarding different regimens of tadalafil that  
16 is, on demand versus 3 times per week or different drugs that is, tadalafil versus vardenafil.  
17 The quality of the evidence was low.

#### 9.3.2.5.38 Testosterone replacement therapy versus placebo

19 Evidence from a single RCT showed that the use of long-acting intramuscular testosterone  
20 therapy was associated with improvement in erectile function in people with type 2 diabetes  
21 diagnosed with hypogonadism. There is limited data on the associated adverse effects of  
22 testosterone therapy. The quality of the evidence was low.

#### 9.3.2.63 Health economic evidence

24 No CUAs were found that directly compare the 4 PDE-inhibitor treatments under  
25 consideration and no CUAs were found that were specific to people with diabetes. Three  
26 CUAs found that 2 of the treatments (sildenafil and vardenafil) for erectile dysfunction are  
27 likely to be cost effective at the appropriate thresholds, but used different comparators (no  
28 treatment or injection therapy). No CUAs were found for tadalafil or avanafil. While none  
29 were undertaken in diabetic specific or UK populations, all the CUAs used similar utility gains  
30 for successful erectile dysfunction treatment and produced base case ICERs that are likely to  
31 be below the £20,000 per QALY threshold. All 3 CUAs contained assumptions that are  
32 conservative or biased towards the alternative treatment but under sensitivity analysis the  
33 treatment option remained likely to be cost effective.

#### 9.3.34 Evidence to recommendations

#### 35 Table 100: Linking evidence to recommendations

Relative value of  
different outcomes

Topic experts were invited to the GDG meeting to inform the clinical discussions prior to making recommendations. The Group agreed that the critical outcomes for decision making were change in erectile function and adverse events, and that both outcomes were weighted equally.

The GDG acknowledged that for PDE-5 inhibitors, adverse effect profiles may differ according to the specific drug, but agreed that it

	<p>was not possible to weight the severity of the events as most side effects are mild and may be individualised.</p>
<p>Trade-off between benefits and harms</p>	<p>The GDG discussed the benefits associated with PDE-5 inhibitors in improving erectile function, self-esteem and quality of life for patients and their partners.</p> <p>The Group noted that the use of PDE-5 inhibitors was associated with relatively mild side effects including headaches and flushing, which may reduce over time. The GDG agreed that it is unlikely that these reductions in adverse events would have been observed in the presented evidence because of the trials' short follow-up periods which ranged from 10 to 16 weeks. The Group noted the different side effects that are associated with individual drugs, for example, tadalafil with backaches and sildenafil with blue-green vision, and agreed that it was not possible to differentiate between the severity of these generally mild adverse events and associated impact on individuals which may be variable.</p>
<p>Consideration of health benefits and resource use</p>	<p>The 3 CUAs found did not meet the NICE reference case, but the GDG felt they showed that effective treatments were likely to increase utility by an extent that would offset reasonable costs. Although no economic evidence was found for using PDE-5 inhibitors to treat erectile dysfunction in people with type 2 diabetes, the GDG felt it was possible to extrapolate from evidence in the general population.</p> <p>The GDG noted that men with type 2 diabetes and erectile dysfunction are likely to be on the higher doses of PDE-5 inhibitor drugs but, even with this in mind, felt that the CUAs presented indicated that effective treatments were likely to increase utility by an extent that would offset reasonable costs.</p>
<p>Quality of evidence</p>	<p>The GDG discussed the overall quality of the evidence for the PDE-5 inhibitors and agreed that it was low to very low.</p> <p>The Group discussed the characteristics of people who were included in the trials and noted that some studies excluded individuals who had cardiovascular disease, hypertension and vascular impairment. Therefore, the GDG agreed that the studies may not be representative of the clinical type 2 diabetes population. The Group also noted that patients taking nitrates (for example, for ischaemic heart disease) would not be able to participate because the use of PDE-5 inhibitors is specifically contraindicated in these individuals. The GDG discussed the inclusion of the 2 studies where all participants were men with type 1 diabetes and noted that these studies may also have underestimated treatment effects, as these patients were younger and may have had different baseline characteristics compared with people with type 2 diabetes.</p> <p>The GDG discussed the 2 trials examining testosterone therapy. One trial was considered to be very low quality evidence because oral testosterone is not used in clinical practice, the small sample of men included in the study had symptoms of andropause or erectile dysfunction, and the trial was open label, with no treatment used as a comparison group rather than placebo. The GDG agreed that this trial should be excluded (Boyanov et al. 2003).</p>

	<p>The second placebo-controlled trial on intramuscular testosterone therapy was not considered to be generalisable to men with type 2 diabetes because the study included a specific subgroup of men who were purposely screened for hypogonadism. The GDG noted that there was little evidence on the safety issues associated with testosterone therapy.</p> <p>The GDG noted the lack of evidence on alprostadil.</p>
Other considerations	<p>The GDG also discussed contributory risk factors and generally agreed that this would include cardiovascular risk so this was added to the existing recommendation about assessment and education. The GDG discussed the other recommendations that were included in CG66, and agreed that these were still relevant.</p> <p>The GDG noted that the majority of studies were conducted in heterosexual couples. The Group thought that a research recommendation would be useful given that it is not clear from the limited evidence base whether the effectiveness of therapies would be similar for men with type 2 diabetes who are in same-sex relationships.</p> <p>When making recommendations for the use of testosterone therapy, the GDG considered the following points:</p> <ul style="list-style-type: none"><li>• There were 2 low quality trials that were not relevant to clinical practice and were associated with several methodological limitations.</li><li>• Therefore, the GDG did not think that there was sufficient evidence to make any recommendations for the use of testosterone therapy.</li></ul> <p>When making recommendations for the use of PDE-5 inhibitors, the GDG considered the following points:</p> <ul style="list-style-type: none"><li>• Overall, it was agreed that the included evidence was of low quality and involved a heterogeneous population, which may not be representative of patients with type 2 diabetes.</li><li>• Alternative treatment options were not considered as part of the evidence review.</li><li>• Treatment of erectile dysfunction that patients consider to be problematic should be discussed with patients and be treated on an individual basis.</li></ul> <p>Therefore, the GDG changed the wording of the recommendation from 'offer' to 'consider'. Although they were confident that PDE-5 inhibitors will do more good than harm for most men with type 2 diabetes and are likely to be cost effective, it was also agreed that alternative options (which were not reviewed as part of this question) may be similarly cost effective. The GDG also added the word 'initially' to reflect that in clinical practice, drugs and doses are chosen but may be altered depending on the progress of the individual patient.</p> <p>The GDG also agreed that there was a lack of evidence for the use of PDE-5 inhibitors in specific subgroups of the population and as a result, no specific recommendations were made.</p>

1

### 9.3.4.1 Recommendations and research recommendations

2 **81. Offer men with type 2 diabetes the opportunity to discuss erectile dysfunction as**  
3 **part of their annual review. [2015]**

4 **82. Carry out an assessment, and provide education and support for men with type 2**  
5 **diabetes who have problematic erectile dysfunction, addressing contributory**  
6 **factors such as cardiovascular disease as well as possible treatment options.**  
7 **[2015]**

8 **83. Consider a phosphodiesterase-5 inhibitor to treat problematic erectile**  
9 **dysfunction, initially choosing the drug with the lowest acquisition cost and**  
10 **taking into account any contraindications. [new 2015]**

11 **84. Following discussion, refer men with type 2 diabetes to a service offering other**  
12 **medical, surgical, or psychological management of erectile dysfunction if**  
13 **treatment (including a phosphodiesterase-5 inhibitor, if appropriate) has been**  
14 **unsuccessful. [2015]**

#### 15 **Research recommendations**

16 **12. What is the optimal dosing of different phosphodiesterase-5 (PDE-5) inhibitors for**  
17 **people with type 2 diabetes and erectile dysfunction?**

#### 18 **Why this is important**

19 Although phosphodiesterase-5 (PDE-5) inhibitors have been shown to be effective  
20 compared to placebo in improving erectile function in men with type 2 diabetes, there is  
21 little understanding of the optimal dosing strategies for the different drugs available in  
22 this class. Double-blind randomised controlled trials in this area could help inform clinical  
23 practice.

24 **13. What is the effectiveness of pharmacological treatment strategies for people with**  
25 **type 2 diabetes and erectile dysfunction who do not respond to**  
26 **phosphodiesterase-5 (PDE-5) inhibitors, for example PDE-5 inhibitor plus**  
27 **prostaglandins?**

#### 28 **Why this is important**

29 There is limited understanding of alternative treatment strategies available to men who  
30 do not respond to phosphodiesterase-5 (PDE-5) inhibitors. Double-blind randomised  
31 controlled trials of combination therapies and other pharmacological treatments could  
32 help inform clinical practice.

33 **14. What is the effectiveness of treatment strategies (pharmacological and non-**  
34 **pharmacological) for sexual dysfunction related to type 2 diabetes in women?**

#### 35 **Why this is important**

36 Sexual dysfunction affect women with type 2 diabetes and there is limited understanding  
37 of available effective treatment strategies. A systematic review is needed examining the  
38 clinical and cost-effectiveness of available treatment strategies for women with type 2  
39 diabetes and sexual dysfunction.

1 **15. What is the effectiveness of treatment strategies (pharmacological and non-**  
2 **pharmacological) for sexual dysfunction in adults with type 2 diabetes in same-**  
3 **sex relationships?**

4 **Why this is important**

5 Sexual dysfunction in adults with type 2 diabetes in same-sex relationships is an  
6 important area, where there is a limited understanding about effective treatment  
7 strategies. A systematic review is needed examining the clinical and cost-effectiveness  
8 of available treatment strategies for adults with type 2 diabetes and sexual dysfunction in  
9 same-sex relationships.  
10

## 9.4.1 Eye damage

2 Diabetes eye damage is the single largest cause of blindness before old age with a  
3 progressive incidence in people with type 2 diabetes.<sup>346</sup> The success of laser therapy in the  
4 treatment of sight-threatening retinopathy is an accepted part of ophthalmological care and  
5 has not been assessed for this guideline.

6 Appropriate clinical questions to be addressed are, however, how people with developing  
7 retinopathy can be selected for ophthalmological referral in time for optimal treatment, and  
8 whether preventative therapy other than good blood glucose, good blood pressure, and good  
9 blood lipid control can be useful in people with type 2 diabetes.

### 9.4.10 Methodological introduction

11 It was noted that management in this area was largely determined by practice for all people  
12 with diabetes and not just those with type 2 diabetes. Indeed retinopathy screening  
13 programmes to be provided on a local community basis were a key early target of the  
14 National Service Framework (NSF) for diabetes, and since that time the UK National  
15 Screening Programme has published and updated a workbook on 'Essential elements in  
16 developing a diabetic retinopathy screening programme' for the guidance of health  
17 authorities and primary care trusts in England.<sup>347</sup>

18 These observations, and a lack of awareness amongst experts of new publications that might  
19 affect recommendations on retinopathy screening, led to the conclusion that  
20 recommendations for people with type 2 diabetes should closely follow those for type 1  
21 diabetes (NICE guideline 2004),<sup>26</sup> which themselves were largely based on generic evidence  
22 independent of type of diabetes.

23 Accordingly the recommendations of the type 1 diabetes guidelines, and the evidence  
24 statements underlying them were reviewed, together with the national screening document.  
25 There are no significant changes from the type 1 diabetes recommendations.

### 9.4.26 Recommendations

27 **85. Arrange or perform eye screening at or around the time of diagnosis. Arrange**  
28 **repeat of structured eye screening annually. [2009]**

29 **86. Explain the reasons for, and success of, eye screening systems to adults with**  
30 **type 2 diabetes, so that attendance is not reduced by lack of knowledge or fear of**  
31 **outcome. [2009]**

32 **87. Use mydriasis with tropicamide when photographing the retina, after prior**  
33 **informed agreement following discussion of the advantages and disadvantages.**  
34 **Discussions should include precautions for driving. [2009]**

35 **88. Use a quality-assured digital retinal photography programme using appropriately**  
36 **trained staff. [2009]**

37 **89. Perform visual acuity testing as a routine part of eye screening programmes.**  
38 **[2009]**

39 **90. Depending on the findings, follow structured eye screening by:**

- 40                   • routine review in 1 year **or**  
41                   • earlier review **or**

- 1                                   • referral to an ophthalmologist. **[2009]**
- 2 **91. Arrange emergency review by an ophthalmologist for:**
- 3                                   • sudden loss of vision
- 4                                   • rubeosis iridis
- 5                                   • pre-retinal or vitreous haemorrhage
- 6                                   • retinal detachment. **[2009]**
- 7 **92. Arrange rapid review by an ophthalmologist for new vessel formation. [2009]**
- 8 **93. Refer to an ophthalmologist in accordance with the National Screening Committee**
- 9 **criteria and timelines if any of these features are present:**
- 10                                  • referable maculopathy:
- 11                                   o exudate or retinal thickening within 1 disc diameter of the centre of
- 12                                   the fovea
- 13                                   o circinate or group of exudates within the macula (the macula is
- 14                                   defined here as a circle centred on the fovea, with a diameter the
- 15                                   distance between the temporal border of the optic disc and the fovea)
- 16                                   o any microaneurysm or haemorrhage within 1 disc diameter of the
- 17                                   centre of the fovea, only if associated with deterioration of best visual
- 18                                   acuity to 6/12 or worse
- 19                                  • referable pre-proliferative retinopathy (if cotton wool spots are present,
- 20                                   look carefully for the following features, but cotton wool spots
- 21                                   themselves do not define pre-proliferative retinopathy):
- 22                                   o any venous beading
- 23                                   o any venous reduplication
- 24                                   o any intraretinal microvascular abnormalities
- 25                                   o multiple deep, round or blot haemorrhages
- 26                                  • any large sudden unexplained drop in visual acuity. **[2009, amended**
- 27                                   **2015]**
- 28

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# 11.1 Glossary and Abbreviations

## 11.1.2 Glossary

### 3 Cohort study

4 (also known as follow-up, incidence, longitudinal, or prospective study): an observational  
5 study in which a defined group of people (the cohort) is followed over time. Outcomes are  
6 compared in subsets of the cohort who were exposed or not exposed (or exposed at different  
7 levels) to an intervention or other factor of interest.

### 8 Comorbidity

9 Two or more diseases or conditions occurring at the same time, such as depression and  
10 anxiety.

### 11 Confidence interval (CI)

12 The range within which the 'true' values (for example, size of effect of an intervention) are  
13 expected to lie with a given degree of certainty (for example, 95% or 99%). (Note: confidence  
14 intervals represent the probability of random errors, but not systematic errors or bias.)

### 15 Cost-effectiveness analysis (CEA)

16 An economic evaluation that compares alternative options for a specific patient group looking  
17 at a single effectiveness dimension measured in a non-monetary (natural) unit. It expresses  
18 the result in the form of an incremental (or average or marginal) cost-effectiveness ratio  
19 (ICER).

### 20 Economic evaluation

21 Technique developed to assess both costs and consequences of alternative health strategies  
22 and to provide a decision-making framework.

### 23 Guideline Development Group (GDG)

24 A group of healthcare professionals, patients, carers and members of the Short Clinical  
25 Guidelines Technical Team who develop the recommendations for a clinical guideline. The  
26 group writes draft guidance, and then revises it after a consultation with organisations  
27 registered as stakeholders.

### 28 Generalisability

29 The degree to which the results of a study or systematic review can be extrapolated to other  
30 circumstances, particularly routine healthcare situations in the NHS in England and Wales.

### 31 Heterogeneity

32 A term used to illustrate the variability or differences between studies in the estimates of  
33 effects.

1 **Odds ratio (OR)**

2 A measure of treatment effectiveness. The odds of an event happening in the intervention  
3 group, divided by the odds of it happening in the control group. The 'odds' is the ratio of non-  
4 events to events.

5 **Quality-adjusted life year (QALY)**

6 A statistical measure, representing 1 year of life with full quality of life.

7 **Randomised controlled trial**

8 A form of clinical trial to assess the effectiveness of medicines or procedures. Considered  
9 reliable because it tends not to be biased.

10 **Relative risk (RR)**

11 Also known as risk ratio; the ratio of risk in the intervention group to the risk in the control  
12 group. The risk (proportion, probability or rate) is the ratio of people with an event in a group  
13 to the total in the group. An RR of 1 indicates no difference between comparison groups. For  
14 undesirable outcomes, an RR that is less than 1 indicates that the intervention was effective  
15 in reducing the risk of that outcome.

16 **Systematic review**

17 Research that summarises the evidence on a clearly formulated question according to a pre-  
18 defined protocol using systematic and explicit methods to identify, select and appraise  
19 relevant studies, and to extract, collate and report their findings. It may or may not use  
20 statistical meta-analysis.

21

## 11.21 Abbreviations

2 **Table 101: Abbreviations**

Abbreviation	Term
BMI	body mass index
CI	confidence interval
CrI	credible intervals
CUA	cost-utility analysis
DIC	deviance information criterion
DPP-4	dipeptidyl peptidase-4
GDG	guideline development group
GEQ	global efficacy question
GLP-1	glucagon-like peptide-1
GRADE	Grading of Recommendations, Assessment, Development and Evaluation
HbA1c	glycated haemoglobin
HDL	high-density lipoprotein
ICER	incremental cost-effectiveness ratio
IFCC	International Federation of Clinical Chemistry
IIEF	International index of erectile dysfunction
IIEF-EF	erectile function domain of the IIEF
ITT	intention-to-treat
LOCF	last observation carried forward
MHRA	Medicines and Healthcare products Regulatory Authority
MID	minimal important difference
NICE DSU TSD	NICE Decision Support Unit's Technical Support Documents
NIT	non-insulin based therapy
NMA	network meta-analysis
OAD	oral antidiabetic drug
OR	odds ratio
QALY	quality-adjusted life year
RR	relative risk
SD	standard deviation
SE	standard error
SEP	Sexual encounter profile
SPC	summary of product characteristics
UKPDS	UK Prospective Diabetes Study

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## 12.1 Appendices A–K are in a separate file