

1 **The role of sleep duration in diabetes and glucose control**

2 **Abstract**

3 Sleep curtailment is common in the westernised world and coincides with an increase in the prevalence
4 of Type 2 diabetes (T2DM). This review considers the recently published evidence for whether sleep
5 duration is involved in the development of T2DM in humans and whether sleep has a role to play in
6 glucose control in people who have diabetes. Data from large, prospective studies indicates a U-shaped
7 relationship between sleep duration and the development of T2DM. Smaller, cross-sectional studies
8 also support a relationship between short sleep duration and the development of both insulin resistance
9 and T2DM. Intervention studies show that sleep restriction leads to insulin resistance, with recent sleep
10 extension studies offering tantalising data showing a potential benefit of sleep extension on glucose
11 control and insulin sensitivity. In people with established diabetes the published literature shows an
12 association between poor glucose control and both short and long sleep duration. However, there are
13 currently no studies that determine the causal direction of this relationship, nor whether sleep
14 interventions are likely to offer benefit for people with diabetes to help them achieve tighter glucose
15 control.

16 **Introduction**

17 Life on earth is governed by the 24-hour cycle of light and darkness associated with the rotation of the
18 earth. Normally metabolic and physiological pathways are coordinated to this 24-hour cycle by an
19 endogenous clock enabling our bodies to coordinate appropriate physiological processes to the time of
20 day. Research in animals and humans suggests that the advent of a 24/7 lifestyle that disrupts our
21 natural sleep cycles and their alignment to the external light/dark cycle is important in the regulation of
22 energy balance and glucose metabolism^(1; 2; 3). Sleep curtailment has become a prevalent behaviour in
23 the Western and developing world where it is estimated that average sleep duration has declined by
24 almost 2 hours in the past 50 years⁽⁴⁾. In the USA and UK, a third of the population report getting less
25 than 7 hours sleep per night^(5; 6). Coinciding with this there has been an explosion in the prevalence of
26 type 2 diabetes (T2DM), raising the important question of whether there is a causal link between the
27 two. This review will specifically consider the recently published evidence for whether sleep duration
28 is involved in the development of T2DM in humans. It will not appraise the considerable number of
29 studies looking predominantly at the role of sleep in the development of obesity. Nor will it address the
30 separate, but related, literature linking sleep related breathing disorders to obesity and T2DM. Having

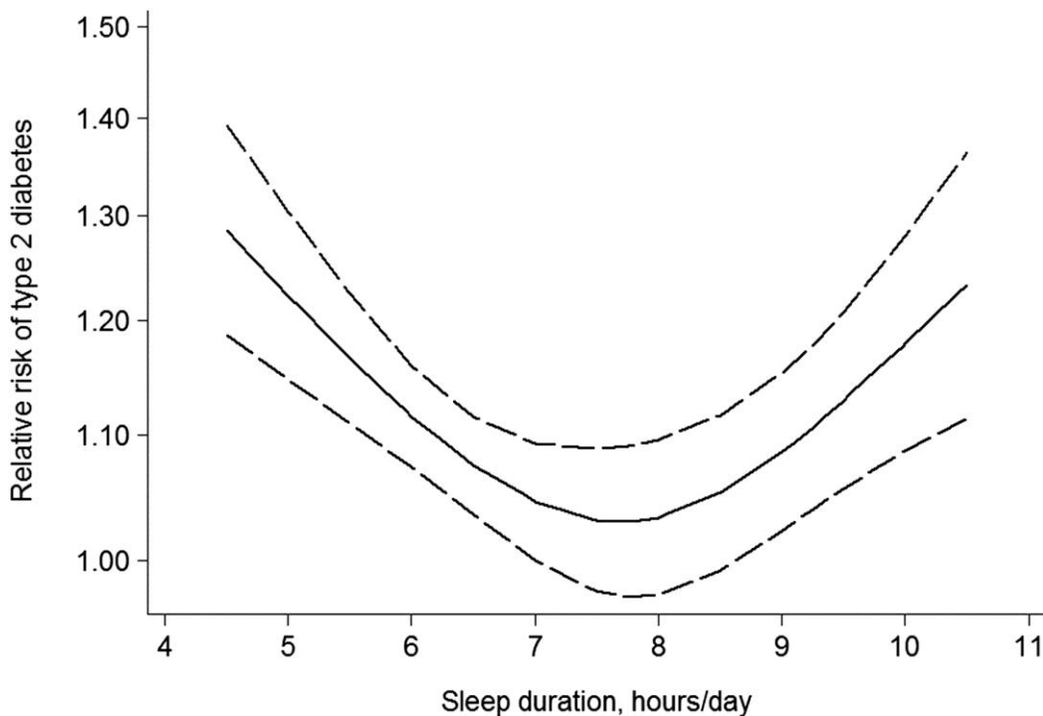
31 considered the role of sleep duration per se in the development of diabetes, the evidence for whether
32 sleep duration has a role to play in glucose control in people who have diabetes is reviewed.

33 **Association between sleep duration and the risk of T2DM**

34 **Evidence from prospective studies**

35 Several large cohort studies have investigated the association between sleep duration and the risk of
36 subsequently developing T2DM, studied over varying lengths of follow-up^(7; 8; 9; 10; 11; 12; 13; 14; 15; 16).
37 Details of their study design and outcomes are shown in Table 1a. Several big studies in the USA and
38 Germany have shown a U-shaped association between sleep duration and increased risk of T2DM^{(7; 8;}
39 ⁹⁾. The studies relied on subjective measures, with sleep duration self-reported at baseline, and T2DM
40 incidence was mainly by self-report of physician's diagnosis. Using 7 hours of sleep duration per night
41 as a reference category, those with shorter and longer sleep duration were significantly more likely to
42 develop T2DM over the follow-up period of 5-15 years (risk estimates ranging between 1.47-1.95 for
43 short sleep duration, and between 1.40-3.12 for long sleep duration) . Regression models employed in
44 the analysis were adjusted for many potential confounders, mainly; age, physical activity, BMI, alcohol
45 consumption, ethnicity, education, marital status, depression and history of hypertension. Not all
46 studies have shown this U-shaped relationship however. A large Australian study of >192,000 adults,
47 used information recorded in medical insurance records⁽¹⁰⁾ and reported a positive association between
48 short (but not long) sleep duration and subsequent incidence of T2DM. However, T2DM incidence was
49 determined from hospital admission records and so those who were not admitted to hospital during the
50 follow-up period could not be identified as T2DM which might have led to underestimation of the
51 actual diabetes incidence. In addition, the follow up period was relatively short (mean duration 2.3
52 years). Another prospective study examined the association of sleep duration with development of
53 impaired fasting glucose (IFG) over six years of follow-up⁽¹¹⁾, with 6-8 hour sleep duration as a
54 reference category, short (but not long) sleepers had higher odds of developing IFG (OR 3.0, 95% CI
55 1.05-8.59; OR 1.6, 95% CI 0.45-5.42: for short and long sleep duration respectively). Whereas a
56 Finnish study in overweight individuals with impaired glucose tolerance found an increased risk of
57 T2DM only in participants with long sleep duration ≥ 9 hours(HR 2.29, 95% CI 1.38–3.80)⁽¹⁴⁾. Finally
58 two recent meta-analyses of nine⁽¹⁷⁾ and fourteen⁽¹⁸⁾ prospective cohort studies have also confirmed the
59 U-shaped relationship (Figure 1).A couple of other studies have investigated the association between
60 short compared to normal sleep duration and the risk of T2DM without examining for a U-shaped
61 relationship. The first showed that sleeping ≤ 7 hours per night was associated with higher odds of

62 developing T2DM after 6 years follow- (OR 1.96, 95% CI 1.10-3.50)⁽¹²⁾. Models were adjusted for age,
63 sex, physical activity, smoking habit, weight gain, and abnormal glucose regulation at baseline. The
64 study also found that the odds of becoming obese were significantly higher in subjects who slept ≤ 7
65 hours per night (OR 1.99, 95% CI 1.12-3.55). There was a lack of association between sleep duration
66 and TD2M at 11 years follow-up that could be related to the attrition in study population over time and
67 to the mediation effect exhibited by adjusting for weight gain in the model. The second study found a
68 higher odds of T2DM after 2 years follow-up, sleeping ≤ 5 hours compared to >7 hours sleep duration
69 (OR 5.37, 95% CI 1.38-20.91)⁽¹³⁾. But the logistic regression models were adjusted for fasting plasma
70 glucose, an integral feature of the outcome measure - T2DM. This leads to a statistical phenomenon
71 known as mathematical coupling thus rendering the result spurious ^(19; 20).



72
73 **Figure 1. U-shaped relationship between sleep duration and risk of T2DM (adapted from Shan, Ma et al.**
74 **2015⁽¹⁷⁾)**

75
76 Extending the understanding of the relationship between sleep duration and risk of T2DM the impact of
77 a change in sleep duration over time has also been investigated. In the Whitehall II study in the UK,
78 change in sleep duration was calculated for participants without diabetes at the beginning and end of
79 four 5-year cycles and T2DM incidence was observed at the end of the subsequent cycle⁽¹⁵⁾. Another,
80 rather convoluted, prospective study (the Nurses' Health Study) examined whether historic changes in

81 women's sleep duration over the preceding 14 years was associated with developing T2DM over the
82 subsequent 12 year follow up ⁽¹⁶⁾. In both studies logistic regression models showed higher risk of
83 developing T2DM in participants with chronic short sleep duration (≤ 5.5 -6 hours) (Whitehall II study:
84 OR 1.35, 95% CI 1.04- 1.76; Nurses' Health Study: HR 1.10, 95%CI 1.001, 1.21) and in those with an
85 increase of ≥ 2 hours sleep duration (Whitehall II study: OR 1.65 , 95% CI 1.15- 2.37; Nurses' Health
86 Study: HR 1.15, 95% CI 1.01- 1.30) compared to those who maintained a 7-8 hour sleep duration.
87 After adjusting for body mass index (BMI) both associations were attenuated, suggesting that BMI is a
88 mediator in the association. These studies suggest that the adverse metabolic influence of short sleep
89 duration may not be ameliorated by increasing sleep duration later in life.

90 Taken together, these large, prospective studies which include both men and women, and a wide range
91 of ages, show that a U-shaped relationship exists between self-reported sleep duration and the
92 development of T2DM. Given the increasing societal pressures to sleep less this data is very persuasive
93 of short sleep duration being implicated in the co-existent T2DM epidemic.

94 **Evidence from cross- sectional studies**

95 Whilst they do not carry the same weight as prospective studies, there have been several recent cross-
96 sectional studies that suggest there may be some key sociodemographic factors that influence the
97 relationship between sleep duration and T2DM, and these are summarised in Table 1-b. Again a U-
98 shaped association between sleep duration (over a 24 hour period) and T2DM was observed in 130,943
99 adults, aged 18-85 years, using data from the National Health Interview Survey (NHIS) from 2004 to
100 2011 ⁽²¹⁾. However, short and long sleep durations were more strongly associated with T2DM in white
101 participants than in black. Adjustment for socioeconomic status and other health behavioural factors
102 attenuated the associations in both groups and remained significant only in white participants. An
103 additional cross-sectional study using the NHIS data, from years 2004-2005, also reported a U-shaped
104 association between sleep duration and T2DM ⁽²²⁾. A Chinese study showed that longer sleep duration
105 over a 24 hour period was positively associated with having the metabolic syndrome and T2DM, but
106 only in women. ⁽²³⁾ Objectively measured sleep duration (using wrist actigraphy) showed a significant
107 association between shorter sleep duration and having IFG and diabetes, but did not find any ethnic
108 differences in a multi ethnic study ⁽²⁴⁾.

109 **Association between sleep duration and the risk of insulin resistance**

110 One of the key features of T2DM is impaired insulin mediated glucose uptake, otherwise known as
111 insulin resistance. This precedes the glucose abnormalities and clinical manifestation of T2DM, often
112 by many years. A selection of both observational and intervention studies have recently explored the
113 relationship between sleep duration and measures of insulin resistance and these are summarised in
114 Table 2.

115 **Evidence from cross sectional studies**

116 A small study compared self-reported sleep duration in insulin-resistant individuals (n= 35) with that
117 seen in insulin-sensitive individuals (n=21). Those with insulin resistance slept 43 minutes less/night
118 (p-value = 0.018). The study also found that 60% of insulin-resistant participants slept less than 7 hours
119 in comparison to only 24% only of insulin-sensitive participants (p-value 0.013), though no adjustment
120 for potential confounders was performed ⁽²⁵⁾.

121 Whilst observational studies are of interest, it is well designed interventional studies that really help us
122 to understand the role of sleep duration in the development of insulin resistance and glucose tolerance.
123 There have been a cluster of these published recently looking at the metabolic effects of both sleep
124 restriction and sleep extension.

125 **Evidence from sleep restriction clinical studies**

126 Multiple small lab-based crossover studies on healthy young participants have looked at the effect of
127 sleep restriction on insulin sensitivity and glucose tolerance ^(26; 27; 28; 29; 30). Sleep restriction was
128 associated with reduced insulin sensitivity in all the studies (except one⁽³⁰⁾), with reduced glucose
129 tolerance in one of them ⁽²⁷⁾. The study that found no effect on glucose or insulin aimed to determine
130 the hormonal effects of restricting sleep duration under controlled feeding conditions ⁽³⁰⁾. A controlled
131 diet was provided and the participants lost weight in both the habitual and short sleep phases. It is
132 possible that in the context of negative energy balance, acute short sleep duration does not lead to a
133 state of increased insulin resistance. These studies were all performed in controlled laboratory based
134 environments and explored acute and often severe sleep restriction. One recent study has examined
135 participants in their home environment to determine if milder and more chronic sleep restriction, akin
136 to daily life, has a role to play ⁽³¹⁾. Nineteen healthy, young, normal-weight men with habitual sleep
137 durations of 7.0–7.5 hours and no sleep disturbances were randomised to either study arm (1.5 hours
138 reduction in habitual bedtime) or control arm (habitual bedtime) for three weeks. Sleep restriction led
139 to a decrease in insulin sensitivity at the end of first week but then recovered to baseline levels.

140 In summary these intervention studies are showing that sleep restriction is associated with the
141 development of insulin resistance and impairment of glucose tolerance. Whether these effects are short
142 lived adaptive responses to an acute stress, or whether they persist longer term and contribute to the
143 risk of T2DM remains unclear.

144 **Evidence from sleep extension clinical studies**

145 Given that short sleep duration and sleep restriction are linked to the development of insulin resistance
146 it is timely that a couple of studies are starting to address whether sleep extension has beneficial effects
147 on insulin and glucose metabolism.

148 The first, a crossover study ⁽³²⁾ showed that whilst insulin sensitivity deteriorates after acute sleep
149 restriction it recovers after two days of catch-up sleep. Under lab-controlled conditions participants had
150 up to 8.5 hours of sleep per night for 4 consecutive nights and up to 4.5 hours of sleep for another 4
151 consecutive nights in a randomised order. After the nights of restricted sleep participants had 2 ‘catch-
152 up’ nights of 10-12 hours of sleep. Participants had a 23% decrease in insulin sensitivity after 4 days of
153 sleep curtailment compared to normal sleep. However, insulin sensitivity was restored after 2 days of
154 catch-up sleep. Although the study showed that catch-up sleep may reverse the negative impact of
155 short-term sleep deprivation, the long-term impact of a repeated sleep deprivation and catch-up sleep
156 cycles on diabetes risk is not known.

157 The second study investigated whether sleep extension in the home environment has a positive impact
158 on glucose metabolism in healthy adults with chronic sleep curtailment ⁽³³⁾. Sixteen young healthy non-
159 obese adults, mostly females, had two weeks of habitual time in bed followed by 6 weeks of one hour
160 per day extension time in bed. Glucose and insulin were assayed at the end of the two periods. During
161 the intervention phase; participants mostly went to bed an hour earlier and had higher sleep duration
162 during weekdays but maintained the same sleep duration during weekends. The study indicated no
163 significant difference between pre- and post-intervention fasting glucose and insulin levels, though no
164 statistics were shown ⁽³⁴⁾. A moderate linear relationship was reported between the relative change in
165 sleep duration and the relative change in fasting glucose ($r = +0.65$, $P = 0.017$) and insulin levels ($r =$
166 -0.57 , $P = 0.053$), however we cannot quantify these relationships nor state if they were statistically
167 significant, without a reported estimate measure of associations.

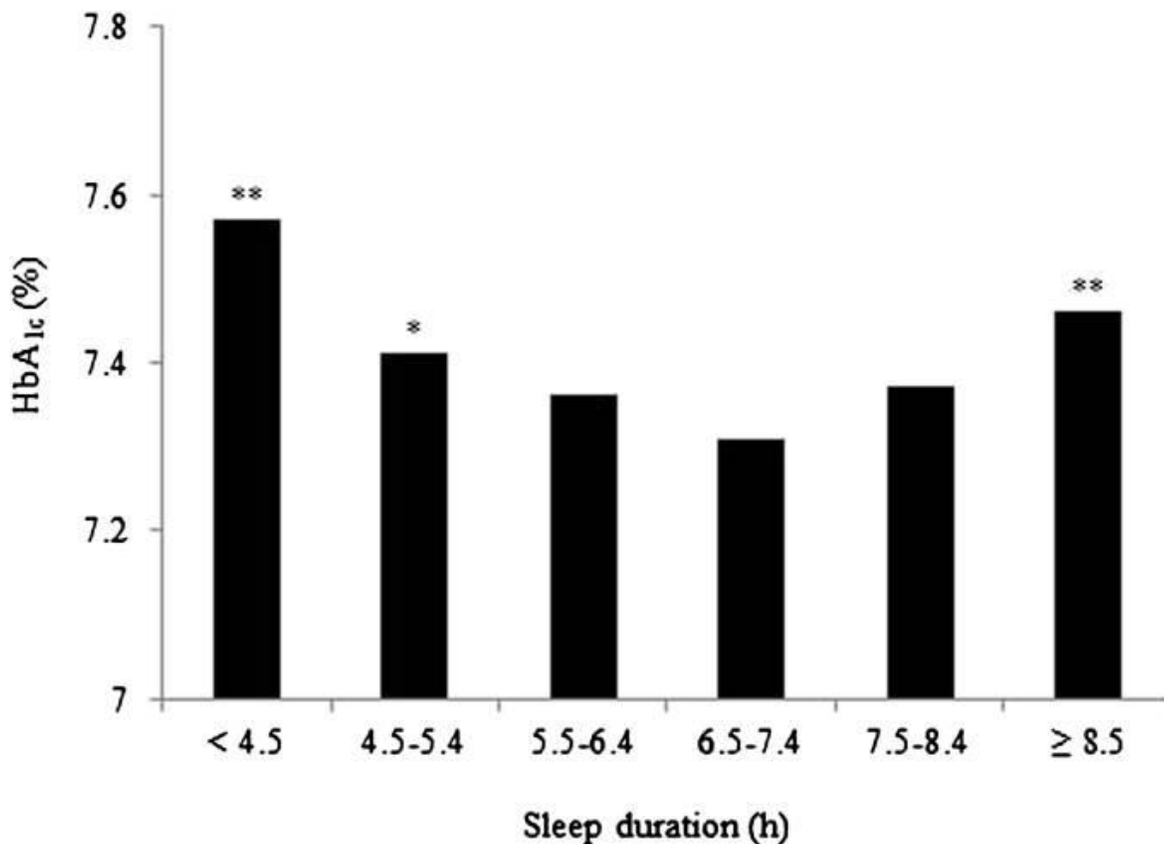
168 In conclusion, these sleep extension studies, offer tantalising data supporting a potential benefit of sleep
169 extension on glucose control and insulin sensitivity. Much more work is clearly needed in this area.

170 **Association between sleep duration and glycaemic control in patients with diabetes**

171 Given the mounting evidence supporting a relationship between sleep duration and the development of
172 insulin resistance and T2DM, it is relevant to consider whether sleep duration has an impact on
173 glycaemic control in people with established diabetes. Most studies to date are cross sectional with
174 sample size ranging from as low as 18 participants to as high as 8543 participants, and these are
175 summarised in Table 3. Most of these studies assessed glycaemic control using HbA1c except one that
176 used capillary glucose levels ⁽³⁵⁾. Sleep parameters were mainly self-reported except for three studies
177 that measured sleep objectively using wrist actigraphy ^(35; 36; 37).

178 **Evidence from cross-sectional studies using subjectively reported sleep duration**

179 Okhuma et al showed that shorter and longer sleep durations were associated with a higher HbA1c
180 level compared with a sleep duration of 6.5–7.4 hours in T2DM patients (Figure 2)⁽³⁸⁾. Sleep duration
181 including naps was self-reported. This U-shaped association was not attenuated after adjusting for
182 BMI, total energy intake and depressive symptoms. A similar U-shaped relation was reported in a large
183 Korean study which included participants with both T1DM and T2DM ⁽³⁹⁾. The association between
184 short sleep duration and poor glycaemic control was strongest for females and participants below the
185 age of 65 years. However, these associations was attenuated after adjusting for BMI and waist
186 circumference and no association was observed after further adjustment for treatment status, duration
187 of diabetes, and daily caloric intake. On the other hand, only longer sleep duration was associated with
188 poor glycaemic control in T2DM patients in a large Chinese ⁽⁴⁰⁾ and a smaller Taiwanese study ⁽⁴¹⁾.
189 Perceived sleep debt but not sleep duration was positively associated with HbA1c in African
190 Americans with T2DM without diabetes complications and not using insulin⁽⁴²⁾.



191

192 **Figure 2. Higher HbA1c observed in shorter and longer sleep duration in Japanese T2DM compared to**
 193 **6.5–7.4 hour sleep duration(*P < 0.05; **P < 0.01), adapted from Okhuma, T et al. 2013 ⁽³⁸⁾**
 194

194

195 **Evidence from cross-sectional studies using objectively measured sleep duration**

196 Using wrist actigraphy to objectively measure sleep parameters for three consecutive days in the home
 197 environment Trento et al. found no difference in sleep duration between T2DM and control subjects
 198 ⁽³⁶⁾. T2DM sleep quality was found to correlate slightly with glycaemic control, while no correlation
 199 with sleep duration was reported. On the other hand, Borel et al. shown that T1DM patients with
 200 shorter sleep duration (< 6.5 hours) had a higher HbA1c than those with longer sleep duration (> 6.5
 201 hours), (mean 8.5% vs. mean 7.7%; p-value = 0.001) ⁽³⁷⁾. In an adjusted regression model shorter sleep
 202 duration was associated with 0.64% increase in mean HbA1c level compared to longer sleep duration
 203 but no 95% CI or p-value were reported. Participants with short sleep duration were also more likely to
 204 have obstructive sleep apnoea and as this was not considered in the analysis, part of the difference
 205 reported could be attributed to it.

206 Lastly Barone et al. assessed the association between objectively measured sleep parameters using
 207 wrist actigraphy and glycemic control using capillary glucose levels from glucometer in a group of 18

208 young adults with T1DM ⁽³⁵⁾. They showed a positive correlation between the average amount of rest at
209 night and both average glucose levels ($r = 0.5404$; $p\text{-value} = 0.0697$) and glycemic variability ($r =$
210 0.5706 ; $p\text{-value} = 0.0527$). No estimate measure of the association nor adjustment for any potential
211 confounders were made. Moreover, the average night rest duration was calculated from 10 days
212 actigraphy which may have included two weekends for some participants and only one weekend for
213 others, and thus it will be inevitably incomparable between participants.

214 In summary, in people with diabetes there also appears to be evidence of an association between both
215 short and long sleep duration and worse glucose control. However, these studies are cross-sectional and
216 so evidence of causality cannot be inferred. A complicating factor when interpreting the relationship
217 between glucose and sleep in people with diabetes is that extremely poor glucose control is well
218 recognised to cause polyuria, polydypsia and nocturia meaning they are awake during the night.
219 Reverse causality cannot be excluded in these studies as they do not distinguish the severity of glucose
220 control and are likely to include participants experiencing some of these osmotic symptoms.
221 Randomised clinical trials with exposure to sleep duration modification (restriction or extending) and
222 robust methods of assessing temporal glucose across the 24 hour day and night ⁽⁴³⁾ are needed to yield
223 more definitive answers.

224 **Conclusions**

225 The recently published literature of both observational and interventional studies strongly supports a
226 role for both short and long sleep duration in the development of insulin resistance and T2DM. There
227 are insufficient sleep intervention studies to determine whether this risk is modifiable long term. In
228 people with established diabetes the published literature supports an association between poor glucose
229 control and both short and long sleep duration. However, there are no studies that determine the causal
230 direction of this relationship in people with diabetes nor whether sleep interventions may offer benefit
231 in achieving glucose control. This is a fertile field for future research.

Table 1-a: Prospective studies on sleep duration and the risk of developing T2DM.

Author, year	country	participant	Study design	exposure	outcome	result	comment
Gangwisch et al., 2007 ⁽⁸⁾	USA	8992 adult aged 32-68 years from the NHANES I	Prospective cohort	Subjective nighttime sleep duration	T2DM incidence over 8-10 year follow-up period	U-shaped associations	
Yaggi et al., 2006 ⁽⁷⁾	USA	1564 men aged 40-70 years from the Massachusetts Male Aging Study	Prospective cohort study	Subjective nighttime sleep duration	T2DM incidence over 15 year follow-up period	U-shaped associations	
Kowall et al., 2016 ⁽⁹⁾	Germany	4814 adults aged 45-75 years from the Heinz Nixdorf Recall study	Prospective cohort study	Subjective nighttime sleep duration	T2DM incidence over 5 year follow-up period	U-shaped associations	
Holliday et al., 2013 ⁽¹⁰⁾	Australia	192728 adults aged \geq 45 selected from medical insurance database	Prospective cohort study	Subjective sleep duration	T2DM incidence over a mean follow up period of 2.3 years	Positive association, only short sleep duration	Diabetes incidents extracted from hospital admission or mortality electronic records, short follow up period
Rafalson et al., 2010 ⁽¹¹⁾	USA	363 participants; 91 cases, 272 controls, aged 35-79 years	nested case-control	Subjective sleep duration (weekdays only)	Impaired fasting glucose	Positive association, only short sleep duration	
Tuomilehto et al., 2009 ⁽¹⁴⁾	Finland	522 participants aged 40-64 years without diabetes randomly allocated either to a study arm or to a control arm.	Two Prospective cohorts based on arms of a randomised controlled trial	Subjective sleep duration	T2DM incidence over 7 year follow up period	Positive association, only long sleep duration	Only in the control arm cohort
Gutiérrez-Repiso et al., 2014 ⁽¹²⁾	Spain	1145 randomly selected participants aged 16-65 years from the Pizzara study	Prospective cohort	Subjective nighttime sleep duration	T2DM incidence at 6 and 11 years follow up	Positive association only at 6 year follow up	Change in sleep duration over the 11 year follow up
Kita et al., 2012 ⁽¹³⁾	Japan	3570 adults aged 35-55 years	Prospective cohort	Subjective sleep duration and sleep quality	T2DM incidence after 2 year follow up)	Positive spurious association	Statistical issues

Ferrie et al., 2015 ⁽¹⁵⁾	UK	5613 adults aged 35-55 years from the Whitehall II study	Prospective cohort, four 5-year cycles	Change in nighttime sleep duration in the following cycle	T2DM incidence at the end of subsequent cycle	Positive association, increase \geq 2 hours	Association could be mediated by weight gain
Cespedes et al., 2016 ⁽¹⁶⁾	USA	59031 middle aged to old women without diabetes	Prospective cohort study	Change in sleep duration over 14 years	T2DM incidence over 12 year follow up period	Positive association, increase \geq 2 hours	Association only with increase in sleep duration \geq 2 h/day, Change in sleep duration from a historic baseline to time of enrolment

Table 1-b: Cross-sectional studies on sleep duration and the risk of developing T2DM.

Author, year	country	participant	Study design	exposure	outcome	result	comment
Jackson et al., 2013 ⁽²¹⁾	USA	130,943 adults aged 18-85 years from the NHIS (years 2004 to 2011)	Cross sectional	Subjective sleep duration in a 24 hours period	self-reported T2DM status	U-shaped associations	Stronger association in white population
Buxton and Marcelli, 2010 ⁽²²⁾	USA	56507 adults from the NHIS (years 2004 to 2005)	Cross sectional	Subjective sleep duration in a 24 hours period	self-reported chronic diseases including T2DM	U-shaped associations	multilevel logistic regression
Wu et al., 2015 ⁽²³⁾	China	25184 adults mean age 63 years from the Dongfeng-Tongji Cohort study	Cross sectional	Subjective sleep duration	Risk of metabolic syndrome including T2DM	No association with nighttime sleep duration	Positive association with daytime napping duration
Bakker et al., 2015 ⁽²⁴⁾	USA	2151 participant aged 45-84 years from the Multi-Ethnic Study of Atherosclerosis	Cross sectional	Objective sleep duration	Diabetes	No association	Model adjusted for OSA

Table 2: Studies on sleep duration and development of insulin resistance

Author, year	country	participant	Study design	exposure	outcome	result	comment
Liu et al., 2013 ⁽²⁵⁾	USA	56 non-diabetic overweight-obese participants	Cross sectional	Subjective sleep duration	Insulin sensitivity	Positive association	Only P-values reported.
Broussard et al., 2012 ⁽²⁶⁾	USA	7 young healthy participants	Crossover clinical study	Sleep restriction	Insulin sensitivity	Positive association	
Nedeltcheva et al., 2009 ⁽²⁷⁾	USA	11 young-middle aged healthy participants	Crossover clinical study	Sleep restriction	Insulin sensitivity and glucose tolerance	positive association	
Wang et al., online 2016 ⁽²⁸⁾	USA	15 young healthy non-obese participants	Crossover clinical study	time-in-bed restriction by 1 to 3 hours for 3 nights	Insulin sensitivity and glucose tolerance	Positive association with insulin sensitivity	No association with glucose tolerance
Donga et al., 2010(1) ⁽²⁹⁾	The Netherlands	9 healthy participants, mean age 44.6 years	Crossover clinical study	Sleep restriction	Insulin sensitivity	Positive association	
St-Onge et al., 2012 ⁽³⁰⁾	USA	27 healthy young non-obese adults	Crossover clinical study	Time in bed restricted to 4 hours	insulin sensitivity	No association	Participants had controlled diet and lost weight during the study
Robertson et al., 2013 ⁽³¹⁾	UK	19 healthy young lean men	Randomised controlled trial	Around 1.5 hours sleep restriction per night for 3 weeks	insulin sensitivity	Positive association only at the end of first week	absence of an overall effect of sleep restriction on insulin sensitivity
Broussard et al., 2016 ⁽³²⁾	USA	19 healthy young lean men	Crossover clinical study	two days of catch-up sleep	Recovery of insulin sensitivity	Positive association	
Leproult et al., 2015 ⁽³³⁾	Belgium	16 healthy young non-obese adults with chronic sleep restriction	Crossover clinical study	Around one hour sleep extension per night for 6 weeks	Fasting glucose and insulin levels	No difference in pre- and post-intervention levels	Moderate correlation between relative change in sleep duration and relative change in fasting glucose and insulin levels

Table 3: Studies on sleep and glycemic control in patients with diabetes

Author, year	country	participant	Study design	exposure	outcome	result	comment
Ohkuma et al., 2013 ⁽³⁸⁾	Japan	4870 adults, aged ≥ 20 years with T2DM	Cross-sectional	Subjective sleep duration including naps	Glycemic control (HbA1c)	U-shaped associations	
Kim et al., 2013 ⁽³⁹⁾	Korea	2134 adults, aged > 20 years with T1DM or T2DM	Cross-sectional	Subjective daily sleep duration	Glycemic control (HbA1c)	positive associations	J-shaped trend with HbA1c; stronger in females and in the younger age group (< 65 years). Association disappear after adjusting for more covariate in the logistic regression model.
Zheng et al., 2015 ⁽⁴⁰⁾	China	8543 adults, aged ≥ 40 years with T2DM or impaired glucose tolerance	Cross-sectional	Subjective nighttime sleep duration	Glycemic control (HbA1c, FPG, PPG)	Positive association with long sleep duration	Only adjusted means and p-values reported but no estimate of association
Tsai et al., 2012 ⁽⁴¹⁾	Taiwan	46 adults, aged 43-83 years with T2DM	Cross-sectional	Subjective sleep duration and quality (PSQI)	Glycemic control, HbA1c	Positive association	Participants with diabetic complication or major co-morbidities were excluded. association only with sleep efficiency and PSQI score of 8 or more but not sleep duration
Knutson et al., 2006 ⁽⁴²⁾	USA	161 African Americans, mean age 57 years with T2DM	Cross sectional	Subjective sleep duration and sleep quality, modified PSQI, and perceived sleep debt	Glycemic control (HbA1c)	Positive association	Sleep debt association only in participants without diabetic complication or not using insulin. sleep quality only in participants with diabetic complication or using insulin association
Trento et al., 2008 ⁽³⁶⁾	Italy	47 middle aged adults with T2DM and 23 healthy controls	Cross sectional study	Objective sleep parameters; duration and quality using wrist actigraphy	glycemic control (HbA1c) in T2DM group	Positive association	Weak negative correlation with sleep efficiency and mild positive correlation with moving time while asleep. No estimate measures of association reported
Borel et al., 2013 ⁽³⁷⁾	France	79 adults, median age 40 years with T1DM	Cross sectional study	Objective sleep parameters using wrist actigraphy	glycemic control (HbA1c)	Positive association	
Barone et	Brazil	18 young adult,	Cross	Objective sleep	Glycemic control	No	Methodological issues

al., 2015 ⁽³⁵⁾		aged 20-38 years with T1DM	sectional	measures using wrist actigraphy	(average glucose from glucometer reading)	association	
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