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Effects of water-soluble vitamin supplementations on glycemic control and insulin resistance in adult type 2 diabetes: an umbrella review of meta-analyses of randomized controlled trials

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ABSTRACT

Background and Objectives: Growing evidence has explored the effects of water-soluble vitamins supplementation on glycemic control and insulin resistance in diabetic patients; however, the results of previous meta-analyses are inconsistent. In this regard, we performed an umbrella review to synthesize evidence on the effects of water-soluble vitamin supplementation on glycemic control and insulin resistance. Methods and Study Design: A systematic literature search in Web of science, PubMed, and Cochrane Database of Systematic Reviews was performed from 2012 to November 2022. Quality assessment of the meta-analyses was performed using AMSTAR-2 and GRADE. Results: Fourteen systematic reviews and meta-analyses were eligible, which studied the effects of five water-soluble vitamins (vitamin B-1, vitamin B-3, biotin, vitamin B-9, and vitamin C supplementation) supplements on glycemic control and insulin resistance. Results of the review suggest that vitamin C supplementations can improve glycemic control in type 2 diabetes indicated by reduced FBG and HbA1c, and having more significant effects with durations >30days on FBG. Conclusions: Insulin resistance is improved by folic acid supplementations. More welldesigned individual randomized controlled trials are needed in the future, as well as metaanalysis of higher quality.

Key Words: water-soluble vitamin, type 2 diabetes, glycemic control, insulin resistance, umbrella review, meta-analysis

INTRODUCTION

Diabetes mellitus, one of the leading causes of death and disability worldwide, is a significant public health issue. Until 2021, the number of adults with diabetes has reached 5.1 billion worldwide, with a prevalence rate of 10.5%; it is estimated that by 2045, the number will reach 6.4 billion worldwide, with a prevalence rate of 12.2%. In addition, diabetes is related to 6.7 million deaths and an expenditure of at least \$966 billion on healthcare in 2021.2 Type 2 diabetes mellitus (T2DM), the most common type of diabetes, accounted for more than 96% of diabetes cases globally in 2021. The mechanism of T2DM is mainly associated with impaired insulin sensitivity, namely insulin resistance, as well as pancreatic β-cell dysfunction. And inflammation generation may contribute to the progression of T2DM. For instance, MAPK signaling pathway, a key regulator for insulin signaling, has reported to be activated under oxidative stress, resulting in insulin resistance.

Mounting evidence suggests that glycemic control, the core target of treatment in diabetes, affects the development of its complications to a large extent.⁸⁻¹¹ It is widely accepted that glycosylated hemoglobin (HbA1c) is the most important indicator to reflect the long-term glycemic control status of diabetic patients, while fasting blood glucose (FBG) indicates a relatively short-term glycemic control status.¹² What's more, the variants of FBG and HbAc1 were strongly associated with the risk of developing retinopathy, nephropathy and all-cause mortality in diabetic patients.^{13, 14} Therefore, it is essential to maintain good glycemic management, especially for the purpose of decreasing the risk of various complications of diabetes mellitus.¹⁵ To find effective ways for glycemic control in diabetic patients has already become a central public health issue.

There are several recommended approaches in the existing diabetes guidelines to deal with the development of diabetes and its complications, 16, 17 such as exercise interventions, 18, 19 improvement of dietary pattern, 20, 21 as well as pharmacological control. In recent years, dietary supplements such as probiotics,²² soluble fiber,²³ resveratrol,²⁴ vitamins and minerals25 have aroused intensive interests in scientific field and been reported to exert good effects on diabetes control. Water-soluble vitamins, including B vitamins and vitamin C, mainly act as coenzymes or coenzymes component molecules involved in the body's metabolism, playing an important role in vital activities of the body including energy metabolism, antioxidation, etc.^{26, 27} We searched for all meta-analyses of water-soluble vitamin supplementation and assessed the quality of the meta-analyses and the randomized controlled trials (RCTs) they included, as well as counting the number of identical RCTs from different meta-analyses and calculating corrected coverage area (CCA) of RCTs vital activities of the body including energy metabolism, antioxidation, etc. It has been reported that water-soluble vitamins, such as vitamin C, folate, thiamine and biotin, had a significant impact on diabetes and its complications.²⁸ The possible underlying mechanisms were related to improve oxidative stress, inflammation, and insulin resistance. ²⁹ For instance, ascorbic acid (AA) has been reported to scavenge reactive oxygen and nitrogen species in vitro and in vivo, 30, 31 enhancing insulin sensitivity in skeletal muscle through ameliorating the oxidative stress;³² folic acid supplementation has been shown to reduce c-Jun N-terminal protein kinase (JNK) activation and TNF gene expression, thereby reducing glucose uptake and inhibiting inflammatory processes;^{33, 34} thiamine can activate glucose metabolism and insulin synthesis,³⁵ thus plays a role in blocking pathways that are responsible for hyperglycemia induced damage;³⁶ and biotin may compensate for low-concentration insulin exposure by inhibiting FOXO1 levels, increasing insulin expression and secretion. 37,38

There are several systematic reviews and meta-analyses (SRMAs) of RCTs summarizing the effects of water-soluble vitamin supplementations on insulin resistance and glycemic control; however, previous evidence of the pooled analysis shows inconsistent results. For example, two pooled studies showed that folic acid supplementation reduced FBG concentrations, ^{39, 40} but one study showed no such effect. ⁴¹ Three studies showed that vitamin C supplementation reduced HbAc1, ^{25, 42, 43} while two studies did not. ^{44, 45} As to the two niacin supplementation trials, 46, 47 no statistically significant effects on blood glucose were found neither. As for the effects of thiamine and biotin supplementations, 37, 48 there is only one SRMA for both, and no statistically significant effect was found on FBG. Umbrella review is primarily an analysis of the evidence given for different interventions for the same problem or disease condition, or evidence from multiple studies that synthesize studies that have investigated the same interventions and disease conditions but have addressed and reported different outcomes, providing a summary of the synthesis of existing studies related to a given topic or problem, rather than a re-synthesis.⁴⁹ There have been some umbrella reviews that describe the effects of probiotics, minerals, and individual vitamins such as vitamin C and vitamin D on glycemic control and insulin resistance.⁵⁰ However, umbrella review that specifically summarizes the effects of water-soluble vitamin supplementations on glycemic control and insulin resistance is still not available till now.

The purpose of this umbrella review is to re-evaluate SRMAs of the role of water-soluble vitamin supplementations in glycemic management in T2DM patients. The quality of the SRMAs was assessed by using the methodological quality assessment tool AMSTAR-2 and the quality of evidence evaluation tool GRADE to analyze the differences and associations of various water-soluble vitamins under different outcome indicators and to more comprehensively summarize the impact of water-soluble vitamin supplementation on glycemic control. Our study may provide important scientific evidence for proposing the nutritional recommendations targeting patients with type 2 diabetes.

MATERÍALS AND METHODS

Search strategy

We performed an extensive search of the SRMAs using three databases, Web of science, PubMed, and Cochrane Database of Systematic Reviews, including only English-language articles, with data search dates ending in November 2022. The search strategy is presented in Supplementary Table 1.

Study selection

Two researchers (Yin and Wang) independently completed the review of studies based on criteria for inclusion and exclusion. Firstly, relevant studies were selected based on the title and abstract of the studies. Secondly, selected studies were further screened by reading the full content of the included studies. Finally, disagreements were resolved by the judgment of the third author (Chen). We selected SRMAs by the appropriate inclusion criteria: (1) systematic reviews and meta-analysis of randomized controlled trials in adults aged 18 years or older; (2) reported supplementation with water-soluble vitamin as intervention, and compared with a control group; (3) reported weighted or standardized mean differences (MDs) and corresponding 95% confidence intervals (CIs) in glycemic control as the outcome of interest, the measured indices consisted of FBG, HbA1c, insulin, and HOMA-IR.

The criterion for exclusion includes: (1) the primary study was experimental in animals, *in vivo*, *in vitro* or *ex vivo*; (2) no summary effect size was reported in the systematic review and meta-analysis (e.g., systematic review without meta-analysis).

Quality assessment

We assessed the methodological quality of the SRMAs using AMSTAR 2,⁵¹ which is mainly used to assess systematic reviews that randomized or non-randomized studies of healthcare interventions, or both, and consists of 16 scored items, of which 7 are the critical items. AMSTAR 2 is concerned with the presence or absence of methodological flaws in critical items and rates the overall confidence in the results of the systematic reviews accordingly. Additionally, we used GRADE to assess the quality of evidence for the meta-analysis.^{52, 53} There are five main components that influence the downgrading of GRADE evaluations: (1) Risk of bias; (2) Imprecision; (3) Inconsistency; (4) Indirectness; (5) Publication bias. When a risk factor is present in the evidence, the certainty needs to be downgraded by one or two levels (e.g., from high to moderate).

Data extraction

Two investigators (Yin and Wang) independently extracted studies information for the metaanalysis that was eligible for inclusion. Information collected included the first author's name, years of publication, sample sizes (including the number of RCTs in the meta-analysis and the total number of participants in the intervention and control groups), type of study, vitamin species, doses and durations of interventions, study locations, and conflict of interest, etc. Besides, the pooled effect sizes and 95%CI for outcome indicators such as FBG, HbA1c, insulin, and HOMA-IR as well as the heterogeneity of the studies, p-values for heterogeneity and publication bias (p-values determined by Egger's test and Funnel plot) were extracted.

RESULTS

We searched a total of 2829 studies from three databases, and a total of 14 SRMAs of RCTs were included in our umbrella review (one of which was a network meta-analysis) after reading not only the titles and abstracts of the studies but also the full text according to the previously established exclusion criteria for inclusion in the studies (see Figure 1). The intervention trials in the SRMAs included the following 5 individual water-soluble vitamins: vitamin B-1 (N=1), vitamin B-3 (N=2), biotin (N=1), vitamin B-9 (N=4), and vitamin C (N=6).

Characteristics of the included systematic reviews and meta-analyses

The 14 included SRMAs were published between 2014 and 2022, the characteristics of which were summarized in Table 1. In this study, T2DM patients were the target population, also, persons with other metabolic disorders including obesity, polycystic ovary syndrome, metabolic syndrome, etc. were also included with the purpose to compare the effects. One systematic review reported thiamine intervention (dose: $100 \sim 900$ mg/day) for durations ranged from 1 to 3 months. Two systematic reviews reported niacin interventions for durations ranged from 8 to 64 weeks (dose: $150 \sim 4500$ mg/day). One systematic review reported biotin interventions with durations ranged from 4 weeks to 3 months (dose: 1.5-15 mg/day). Primary studies of the 4 systematic reviews that examined the effect of folic acid interventions for longer durations of 2weeks to 7.3years (dose: 0.5-15 mg/day). It worth noting that the duration of vitamin C interventions varied greatly between the primary studies, with durations ranged from 14 days to 9 years (dose: 72-6000 mg/day). All systematic reviews used random effects models for pooled estimation. Most of the primary RCTs used placebo controls, and a small proportion used blank controls.

There were 162 primary RCTs in the 14 included systematic reviews, and after excluding duplicate studies, there were totally 88 primary RCTs implemented in 89 regions, of which 4, 8, 5, 34, and 37 primary RCTs conducted vitamin B-1, vitamin B-3, biotin, vitamin B-9, and vitamin C supplementations, respectively (Supplementary Table 2). In addition, 17 RCTs were conducted in Iran, 11 of which had vitamin B-9 interventions, and 13 studies were conducted in the United States, with vitamin B-3 or vitamin C interventions in 5 studies each (Figure 2). We noticed that the quality of the primary RCTs was closely related to the

economic status of the places where the studies were conducted, which were significantly higher in countries with better economic status.

Estimating the degree of overlap or corrected coverage area (CCA) for the included SRMAs, high CCAs were found in the supplementation trials of vitamin B-3 (CCA=62.50%), vitamin B-9 (CCA=24.51%) and vitamin C (CCA=18.54%). If the meta-analysis were grouped according to the study outcomes, the degree of overlap or CCA) was calculated again, and the results showed that the CCAs remained high. (Table 2)

The corresponding authors of the systematic reviews were mainly from Iran (5/14), Australia (2/14), China (4/14), UK (1/14), Korea (1/14), and Thailand (1/14). The source of funding for the systematic reviews was mainly national foundation (3/14), and 64% of the systematic reviews did not report a source of funding. Most of the systematic reviews reported no conflict of interest.

Risk of bias and quality assessment of included meta-analyses

The assessment results of AMSTAR-2 for the studies are presented in Figure 3. One study was a network meta-analysis and AMSTAR-2 was not applicable. The remaining thirteen systematic reviews and meta-analyses were rated as high, moderate, and low at rates of 2 (3/13), 2 (2/13) and 8 (8/13), respectively. The most common critical flaw in the included studies was the failure to consider the risk of bias in the included studies when the investigator interpreted the results of each study (9/13). According to the assessment details of AMSTAR-2 and GRADE, most of the included SRMAs were low-quality articles with about 61.5% of the articles assessed as low by AMSTAR-2, mainly because the SRMAs did not consider quality assessment when interpreting the results; and about 31.6% and 26.3% of the articles assessed as low and very low by GRADE, mainly due to high heterogeneity among primary RCTs and publication bias also existed in meta-analysis studies.

The quality of evidence was assessed for 38 outcome indicators extracted from the included studies, resulting in three of high-quality evidence, thirteen of moderate quality evidence, twelve of low-quality evidence, and ten of very low-quality evidence. Inconsistency was the main factor affecting the downgrading, followed by risk of bias, indirectness, imprecision and publication bias (Figure 4, Supplementary Table 3). Also, Figure 4 shows the effects of water-soluble vitamin interventions on glycemic control and insulin resistance as reported in the included systematic reviews. In this review, we found that conclusions with significant differences were often derived from low-quality evidence. The inclusion of low and very low-quality evidence impacts the reliability and stability of the final results,

rendering the conclusions of the review potentially uncertain and insufficient to provide robust support for clinical practice. This underscores the need for further high-quality research to validate these findings.

We assessed the quality of the RCTs extracted from each meta-analysis with three quality assessment methods, namely JBI evidence-based center's quality assessment tool (N=1), Jadad scale (N=5), and Cochrane collaboration's tool for assessing risk of bias (N=8), and seven meta-analyses of vitamin B-3, folic acid and vitamin C having more than 50% of the primary RCTs of moderate and low quality (Figure 5).

The effect of water-soluble vitamin supplementation on FBG

Twelve systematic reviews explored the effects of the supplementation of five water-soluble vitamins including vitamin B-1, vitamin B-3, biotin, vitamin B-9, and vitamin C on FBG (Table 3, Figure 6).

There was only one meta-analysis targeting type 2 diabetic patients claiming that folic acid supplementation could reduce FBG,39 with pooled effect sizes -2.17 (95% CI: -3.69, -0.65). In agreement, another pooled analysis in metabolism-related diseases including T2DM, metabolic syndrome, overweight and obese, polycystic ovary syndrome, coronary artery disease also found folic acid supplementation could reduce FBG with pooled effect sizes ranging from -2.17 (95% CI: -3.69, -0.65) to -0.15 (95% CI: -0.29, -0.01).^{39,40} However, no statistically significant effects of folic acid on FBG were found by Maryam et al in the population with the same metabolism-related diseases aforementioned.⁴¹ There was consistent evidence that vitamin C supplementation could reduce FBG with pooled effect sizes ranging from -20.59 (95% CI: -40.77, -0.4) to -0.44 (95% CI: -0.81, -0.07);^{25, 42, 44, 45} and further subgroup analysis found that durations >30 days had a statistically more significant positive effect on FBG with pooled effect sizes ranging from -0.53 (95% CI: -0.97, -0.10).⁴⁴

There was consistent evidence that thiamine and biotin supplementation had no statistically significant effect on FBG.^{37, 48} As to the two niacin supplementation trials, no statistically significant effects on blood glucose were found neither; however, subgroup analysis found that high doses or >20 weeks' supplementation of niacin were significantly effective for FBG. ^{46, 47}

Totally, as to the influence of water-soluble vitamin on FBG, there were two SAMAs with high quality, three with intermediate quality, three with low quality, and four with very low quality (Figure 4).

The effect of water-soluble vitamin supplementation on HbA1c

Twelve meta-analyses explored the effect of the supplementation of five water-soluble vitamins including vitamin B-1, vitamin B-3, biotin, vitamin B-9, and vitamin C on HbA1c (Table 3, Figure 7). Two (50%) of the four meta-analyses found that vitamin C supplementation could reduce HbA1c with pooled effect sizes ranging from -0.54 (95% CI: -0.9, -0.17) to -0.37 (95% CI: -0.57, -0.17).^{25, 42} There was consistent evidence that thiamine, niacin and folic acid supplementation had no statistically significant effects on HbA1c;^{39, 54} however, subgroup analysis found that high-doses niacin intervention had a statistically significant positive effect on HbA1c with pooled effect sizes 0.90 (95% CI: 0.21, 2.41).⁴⁷ As to the one biotin supplementation trial, no statistically significant effect on HbA1c was found.³⁷ Overall, among the ten pooled studies, one SAMA provided evidence on HbA1c with high quality, four with moderate, two with low and three with very low quality. (See in Figure 4)

The effect of water-soluble vitamin supplementation on insulin resistance

Seven meta-analyses explored the effect of the supplementation of three water-soluble vitamins including biotin, folic acid, and vitamin C on fasting serum insulin (Table 3, Figure 8).

There was only one meta-analysis targeting type 2 diabetic patients claiming that folic acid supplementation could reduce insulin, with pooled effect sizes ranging from -1.63 (95% CI: -2.53, -0.73).³⁹ In agreement, another pooled analysis in the previously mentioned metabolism-related diseases also found folic acid supplementation could reduce insulin, with pooled effect sizes ranging from -1.94 (95% CI: -3.28, -0.61) to -1.28 (95% CI: -1.99, -0.56).³⁹⁻⁴¹ As to the one biotin supplementation trials, no statistically significant effects on insulin were found.³⁷ For the two vitamin C supplementation trials, no statistically significant effects on insulin were found neither.^{42, 44} In conclusion, two SAMAs with moderate quality of evidence, three with low quality and one with very low quality (Figure 4).

We also analyzed the effects of these vitamins on HOMA-IR. Seven meta-analyses explored the effects of two water-soluble vitamins including folic acid and vitamin C on HOMA-IR (Table 3, Figure 9).

There was only one meta-analysis reporting that folic acid supplementation could reduce HOMA-IR, with pooled effect sizes -0.40 (95% CI: -0.70, -0.09).³⁹ In agreement, another pooled analysis in the metabolism-related diseases also found folic acid supplementation could reduce HOMA-IR, with pooled effect sizes ranging from -1.07 (95% CI: -1.80, -0.33)

to -0.40 (95% CI: -0.70, -0.09).³⁹⁻⁴¹ As to the three vitamin C supplementation trials, no statistically significant effects on insulin were found.^{25, 42, 43} In brief, as to insulin resistance, two SAMAs with moderate quality of evidence, four with low quality, and one with very low quality (Figure 4).

DISCUSSION

This umbrella review summarizes the effects of water-soluble vitamins on glycemic management in T2DM. We included a total of 14 manuscripts of systematic reviews and meta-analyses containing 92 primary RCTs of the effects of five water-soluble vitamin supplementations (vitamin B-1, vitamin B-3, biotin, folic acid, and vitamin C) on glycemic control and insulin resistance. We found that folic acid improved insulin concentrations and HOMA-IR and vitamin C supplementation improved FBG and HbA1c in T2DM.

Folic acid (vitamin B-9) significantly improved insulin resistance indicated by reduced serum/plasma insulin concentrations and HOMA-IR. Vitamin B-9 acts as a key one-carbon donor in the body that plays an essential role in cellular metabolism. Low concentrations of vitamin B-9 lead to hyperhomocysteinemia, which has been reported to be associated with the development of insulin resistance.⁵⁵⁻⁵⁷ The supplementation of folic acid could reduce serum homocysteine concentrations and improve glucose-induced oxidative stress and inflammation in T2DM.^{58, 59} This is consistent with our findings. As to FBG, there was one study implemented specifically in type 2 diabetes and found a statistically significant effect, while in the population of metabolism-related diseases including T2DM, metabolic syndrome, overweight and obese, polycystic ovary syndrome, coronary artery disease, there exists discrepancies in the pooled studies, two SAMAs showed that folic acid supplementation could reduce FBG, 39, 40 while one SAMA did not find the same effect; however, when sensitivity analysis was performed, the supplementation was found to decreased FBG again.⁴¹ Therefore, there may exist major confounding in the study. Besides, it did not show a significant effect of folic acid supplementation on HbA1c, probably because HbA1c tends to reflect an estimation of long-term glycemic control, which cannot be significantly modified in the case of a relatively short intervention period (duration <12 weeks) in the included studies. 60 Also, the number of RCTs investigating the possible role of folic acid on HbA1c in the SRMAs was relatively small. 40, 54

In the present umbrella review, vitamin C supplementation was discovered to have a significant effect on glycemic control indicated by FBG and HbA1c. Oxidative stress, predisposing to insulin resistance, beta-cell dysfunction, impaired glucose tolerance, as well

as mitochondrial dysfunction, is a major pathophysiological mechanism for diabetes and its complications.⁶¹ Ascorbic acid (AA), the most potent water-soluble antioxidants in the body, has been reported to scavenge reactive oxygen and nitrogen species *in vitro* and *in vivo*,^{30, 31} resulting in ameliorated oxidative stress.⁶² Therefore, the role of VC on glycemic control in our study mainly attributes to its potent antioxidant function in the body. For FBG, the results of the included meta-analysis were consistent. However, the discrepancy of the effects on HbA1c concentrations were found. The possible reason is that high concentrations of glucose in the blood lead to intracellular VC deficiency, in addition, VC bioavailability is affected by transport proteins, which is impaired in T2DM.⁴⁵ Besides, this may be also due to the small sample size and relatively early publication in some studies.⁴⁵

Ascorbic acid supplementation did not show significant effects on insulin resistance in the present study. The possible reason is the high risk of bias in some studies as reported by Kim et al.²⁵ In addition, the small number of included studies, high heterogeneity (I > 50%) among the studies and the high overlaps of the primary RCTs included in the three SRMAs may also contribute.

Mitochondria are the site of production of important metabolites that regulate insulin secretion, and ATP/ADP ratio is significantly associated with insulin secretion. 63, 64 Also, in subjects with T2DM, impaired secretory response to glucose in pancreatic beta cells was associated with significant alterations in mitochondrial function and morphology.65 As we all know, thiamine participates the process of energy production within mitochondria, affecting intracellular glucose metabolism. 66, 67 In addition, it was reported to regulate insulin secretion, when thiamine deficiency, insulin secretion is impaired by reduced glucose oxidation, leading to beta-cell dysfunction and impaired glucose tolerance. 68-70 Niacin, mainly present in the body as coenzyme 1 (NAD) and coenzyme 2 (NADP), also is an important substance involved in the process of mitochondrial ATP production. At present, although studies did not find that thiamine (vitamin B-1) and niacin (vitamin B-3) supplementations improve blood glucose control, in the context of hyperglycemia, thiamine and niacin supplementations were revealed to prevent diabetic complications.⁷¹⁻⁷³ The possible reason is the small number of included RCTs and populations and may be related to the early publication of the primary RCTs, the very low quality of the studies, and the very high degree of overlap between studies. Besides, one study even found that excess thiamine and niacin caused oxidative stress and insulin resistance in rats.⁷⁴ More rigorous studies are warranted in the future to investigate the effects of thiamine and niacin on glycemic control.

Also, we did not find a significant effect of biotin supplementation on glycemic management or insulin resistance. Unlikely, Zhang et al found that hyperglycemia and decreased insulin secretion and sensitivity was associated with biotin deficiency, and biotin supplementation was able to increase insulin secretion and increase the proportion of beta cells by expanding the size of the islets in rats. Considering the reason of the discrepancy, we found only one SRMA investigated the effects of biotin supplementation on glycemic control and insulin concentrations, and that study included only five RCTs and the pooled sample size of the RCTs was relatively small. In addition, by AMSTAR-2 and GRADE we found a low quality of the meta-analysis mainly due to not reporting publication bias. Therefore, more high-quality studies are needed in the future.

Strengths and limitations

Our study is the first umbrella review to systematically summarize the extensive evidence on the effects of water-soluble vitamin supplementation on glycemic control and insulin resistance. We searched for the effects of all water-soluble vitamin supplementation on glycemic control and insulin resistance and finally found 5 vitamins (vitamin B-1, vitamin B-3, biotin, vitamin B-9, and vitamin C supplementation). In our umbrella review, after categorizing the primary RCTs according to interventions and outcome indicators, we analyzed the quality and the overlap rate of included SRMAs, which is beneficial to the exploration of the reasons for inconsistencies among SRMAs. In addition, we mapped the locations where the primary RCTs were conducted, which may facilitate further studies to explore the potential impact of the region where the study was conducted on outcomes.

Nevertheless, there are still some shortcomings in our umbrella review. First, the degree of overlap or CCA in these included studies was very high and that the interventions in most of the primary RCTs were folic acid and vitamin C. Second, the quality assessment showed that the authors of these SRMAs did not consider the risk of bias in the included RCTs when interpreting the results; and the high heterogeneity of the SRMAs was one of the main factors influencing the downgrading of the quality of the GRADE evidence. Third, in our review, the interventions of RCTs included in the SRMAs were all supplementing single water-soluble vitamin, and thus future studies are needed to investigate the role and effects of multivitamin supplementation or vitamin supplementation in combination with other nutrients on glycemic control and insulin resistance. For instance, combined supplementation of vitamin C and vitamin E can improve glucose metabolism and oxidative stress in T2DM.77 Fourth, we only collected relevant information from the primary RCTs without subjecting them to a new

meta-analysis, and also only summarized the results of the included SRMAs and their quality assessment. Therefore, future studies should adopt a rigorous study design to improve the quality of the studies. Finally, we only visualized the study sites and did not consider or measure the regional differences when discussing and analyzing the results of each study. However, most of the primary RCTs were conducted in countries with unbalanced development, for which economic conditions and social factors had potential impacts on the studies.

Conclusion

Vitamin C supplementations can improve glycemic control in type 2 diabetes mellitus by reduced FBG and HbA1c, and folic acid supplementations improve insulin resistance. More well-designed individual RCTs were needed in the future. More well-designed individual randomized controlled trials are needed in the future, as well as meta-analysis of higher quality.

SUPPLEMENTARY MATERIALS

All supplementary tables and figures are available upon request.

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The all authors certify that they have no affiliations with or involvement in any organization or entity with any financial interest or non-financial interest in the subject matter or materials discussed in this manuscript.

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Table 1. Table 1 Characteristics of included systematic reviews and meta-analysis

SR Author and year	Primary studies,	Population				Age (years)	Intervention			
	n						Vitamin species	Dose (mg/day	y)	Duration
Arti Muley, 2022 ⁴⁸	6	T2DM				mean 52-65.3	B-1	100-900		1-3 months
Yi Ding, 2015 ⁴⁶	7	T2DM				59-67	B-3	150-4500		8-64 weeks
Maryam Akbari, 2018 ⁴¹	16	T2DM/metabolic	syndrome/0	Overweight a	nd obese	NR	B-9	1-10		2-12 weeks
		people/polycystic	ovary synd	rome						
Zhao JV, 2018 ⁴⁰	18	T2DM / Other me	etabolic dise	ases		24.6-67.3	B-9	0.15-10		2weeks-7.3years
Patcharaporn Sudchada, 2012 ⁵⁴	4	T2DM				mean 55-66	B-9	5		4 weeks-6 months
Omid Asbaghi, 2021 ³⁹	24	T2DM / metaboli	c syndrome/	Overweight	and obese	24-65	B-9	0.8-15		3-234 weeks
		people/polycystic	ovary synd	rome/ hypert	ension/		/			
		coronary artery di	isease							
Shaun A. Mason, 2021 ⁴²	28	T2DM				38-71	VC	500-1000		1-6 months
Yoonhye Kim, 2022 ²⁵	12	T2DM				NR	VC	200-1000		3-48weeks
AW Ashor, 2017 ⁴⁴	22	T2DM / healthy in	dividuals / T	1DM / coron	ary artery	22-60	VC	72-6000		14-120 days
AW Ashor, 2017	22	diseases patients					VC	72-0000		14-120 days
Asma Kazemi, 2022 ⁴³	19	T2DM / Diabetic I	Hyperlipidae:	mia		29.3-77 (median	VC	NR		2-52 weeks
						56.5)				
SR Author and year	Comparator	Outcome				Method of pooling	Funding	C	OI	Country of author
•	•	FBG	HbAc1	HOMA-	Insulin	estimates	C			•
				IR						
Arti Muley, 2022 ⁴⁸	placebo: 5, thiami	ine: 1 √	1		7	random effect	NO	N	R	Australia
Yi Ding, 2015 ⁴⁶	Placebo: 3	$\sqrt{}$				random effect	National Found	ation N	O	China
Maryam Akbari, 2018 ⁴¹	placebo	V	1		$\sqrt{}$	random effect	a grant from the	Vice- N	O	Iran
	_						chancellor for F	Research		
Zhao JV, 2018 ⁴⁰	placebo	$\sqrt{}$	V	V	$\sqrt{}$	random effect	NO	N	O	Hong Kong
Patcharaporn Sudchada, 2012 ⁵⁴	placebo		\checkmark			random effect	NO	N	O	Thailand
Omid Asbaghi, 2021 ³⁹	no intervention: 6	5, √	V	$\sqrt{}$	$\sqrt{}$	random effect	NO	N	O	Iran
-	Placebo: 18									
Shaun A. Mason, 2021 ⁴²	placebo	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	random effect	NR	N	O	Australia
Yoonhye Kim, 2022 ²⁵	placebo	V	$\sqrt{}$	\checkmark		random effect	National Founda	tion N	O	Korea
AW Ashor, 2017 ⁴⁴	placebo: 13	1	$\sqrt{}$		$\sqrt{}$	random effect	National Founda	tion N	O	UK
Asma Kazemi, 2022 ⁴³	no intervention: 1,	2	N	$\sqrt{}$	N	random effect	NR	N	\sim	Iran
	Placebo: 18	Y	٧	٧	٧	random enect	111/	11	J	11 411

FBG: fasting blood glucose, HbA1c: glycosylated hemoglobin, HOMA-IR: homeostatic model assessment for insulin resistance, COI: conflict of interest, NR: no report, SR: systematic review and meta-analysis.

Table 1. Table 1 Characteristics of included systematic reviews and meta-analysis (cont.)

	·	• • •				
SR Author and year	Primary studies,	Population	Age (years)	Intervention		
•	n			Vitamin species	Dose (mg/day)	Duration
Mehrnoosh Khodaeian, 2015 ⁷⁸	3	T2DM	20-75	VC	800-1000	4-16 weeks
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	12	T2DM	18-89	VC	120-2000	4weeks-9years
Yujia Zhang, 2022 ³⁷	5	T2DM	46-59	B-7	1.5-15	4 weeks-3 months
Dan Xiang, 2020 ⁴⁷	6	T2DM	mean 59-65	B-3	1500-4500	8 weeks-12 months

SR Author and year	Comparator	Outcome FBG	HbAc1	HOMA- IR	Insulin	Method of pooling estimates	Funding	COI	Country of author
Mehrnoosh Khodaeian, 2015 ⁷⁸	placebo			$\sqrt{}$		random effect	NO	NO	Iran
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	placebo	$\sqrt{}$	\checkmark			random effect	NO	NO	Iran
Yujia Zhang, 2022 ³⁷	placebo	$\sqrt{}$	\checkmark		$\sqrt{}$	random effect	Faculty Research Grants	NO	Macau
Dan Xiang, 2020 ⁴⁷	placebo: 3 statins:3	$\sqrt{}$	V			random effect	NR	NO	China

FBG: fasting blood glucose, HbA1c: glycosylated hemoglobin, HOMA-IR: homeostatic model assessment for insulin resistance, COI: conflict of interest, NR: no report, SR: systematic review and meta-analysis.

Table 2. The overlapping among included systematic reviews and meta-analyses

Vitamin species	Number of reviews	Number of included studies	CA statistic (%)	CCA statistic (%)	Degree of overlapping
Niacin	2	8	81.25%	62.50%	Very high
Folate	4	34	43.38%	24.51%	Very high
VC	6	41	32.11%	18.54%	Very high

CA: coverage area; CCA: corrected coverage area.

Table 3. Efficacy of water-soluble vitamin supplementation on glycemic control and insulin resistance

SR author and year (number of studies)	I/C	Outcomes	Relative effect (95% CI)	$I^{2}(\%)$	Publicatior bias
Vitamin B1					
Arti Muley, 2022 ⁴⁸					
2	24/24	FBG	MD=-0.20 (-0.69, 0.29)	0	YES
		(<3 Mon)	(, ,		
1	40/40	FBG	MD=1.30 (-0.12,2.72)	NR	YES
		(>3 Mon)			
2	51/55	HbA1c	MD=-0.02% (-0.35, 0.31)	0	YES
		(<3 Mon)			
2	79/83	HbA1c	MD=0.19% (-0.17,0.55)	0	YES
2	17/05	(>3 Mon)	1112 = 0.1570 (0.17,0.55)		123
Vitamin B3		(> 5 1/1011)			
Yi Ding,2015 ⁴⁶					
7	452/386	FBG	WMD=-0.07 (-0.44, 0.29)	68.50	NO
Dan Xiang, 2020 ⁴⁷	432/300	100	WWD= 0.07 (0.44, 0.25)	00.50	110
6	658/615	FBG	WMD=0.18 (-0.14, 0.50)	5.20	NO
5	646/603	HbAc1	WMD=0.39 (-0.15, 0.94)	57.60	NO
Vitamin B7	040/003	HUACI	WWD=0.37 (-0.13, 0.74)	37.00	110
Yujia Zhang, 2022 ³⁷					
5	284/161	FBG	MD=-1.21(-2.73, 0.31)	0.00	NR
1	226	HbAc1	MD=-0.18 (-0.39, 0.03)	NR	NR NR
4	266/151	insulin	MD=1.88(-13.44, 17.21)	58.00	NR NR
Vitamin B9	200/131	IIISUIIII	NID=1.88(-13.44, 17.21)	38.00	NK
Omid Asbaghi, 2021 ³⁹					
O ,	17270/17225	EDC	WMD- 217 (260, 065)	91.50	VEC
27	17379/17235	FBG	WMD=-2.17 (-3.69, -0.65)	81.50	YES
4	85/85	HbAc1	WMD=-0.27 (-0.73, 0.18)	74.90	NO
12	322/295	HOMA-IR	WMD=-0.40 (-0.70, -0.09)	80.90	NO
12	315/291	insulin	WMD=-1.63 (-2.53, -0.73)	65.80	NO
Maryam Akbari, 2018 ⁴¹					
10	254/257	FBG	SMD=-0.30 (-0.63, 0.02)	69.10	NO
6	144/134	HbAc1	SMD=-0.29 (-0.61, 0.03)	40.60	NO
8	226/227	insulin	SMD= -1.28 (-1.99, -0.56)	91.50	NO
9	240/244	HOMA-IR	SMD = -1.07 (-1.80, -0.33)	92.50	NO
Zhao JV, 2018 ⁴⁰					
15	8369/8399	FBG	MD=-0.15 (-0.29, -0.01)	53.30	NO
4	157/156	HbAc1	MD=-0.17 (-0.49, 0.16)	77.80	NO
8	190/190	insulin	MD=-1.94 (-3.28, -0.61)	66.10	NO
9	221/214	HOMA-IR	MD=-0.83 (-1.31, -0.34)	80.90	NO
Patcharaporn Sudchada,					
2012 ⁵⁴					
3	71/71	HbAc1	WMD=-0.37 (-1.10, 0.35)	83.80	NO
Vitamin C					
AW Ashor, 2017 ⁴⁴					
13	NR	FBG	WMD=-0.44 (-0.81, -0.07)	NR	NR
10	NR	HbAc1	WMD=-0.02 (-0.19, 0.15)	0.00%	NR
6	NR	insulin	WMD=-13.63 (-22.73, -4.54)	NR	NR
Shaun A. Mason, 2021 ⁴²					
20	670/635	FBG	MD=-0.74 (-1.17, -0.31)	74.95%	NO
16	570/563	HbAc1	MD=-0.54% (-0.9, -0.17)	88.70%	NO
5	222/214	HOMA-IR	MD=-1.43 (-2.88, 0.01)	60.98%	NO
9	133/130	insulin	MD=-0.74 (-2.09, 0.61)	85.44%	NO
Ozra Tabatabaei-Malazy,		-	(,,		-
2014 ⁴⁵					
5	184/181	FBG	MD=-20.59 (-40.77, -0.4)	NR	NO
5	184/181	HbAc1	MD=-0.46 (-1.75, 0.84)	NR	YES
Asma Kazemi, 2022 ⁴³	10 1/101	1107101	2.22 - 0.10 (1.73, 0.0 1)	1121	120
19 (18) [†]	676/610	FBG	MD=-12.03 (-19.43, -4.63)	93.30%	YES
15 (16)	543/538	HbAc1	MD=-0.48 (-0.75, -0.21)	83%	YES
5 (4) [†]	131/126	HOMA-IR	MD=-0.48 (-0.75, -0.21) MD=-0.06 (-1.15, 1.02)	75.30%	NO
8 (7) [†]	215/207	insulin	MD=-1.164 (-3.21,0.86)	71.20%	YES

CA: coverage area; CCA: corrected coverage area.

Table 3. Efficacy of water-soluble vitamin supplementation on glycemic control and insulin resistance (cont.)

SR author and year (number of studies)	I/C	Outcomes	Relative effect (95% CI)	I ² (%)	Publication bias
Vitamin C					_
Mehrnoosh Khodaeian,					
2015^{78}					
3	92	HOMA-IR	SMD=- 0.15 (- 0.49, 0.19)	35.40%	NO
Yoonhye Kim, 2022 ²⁵					
12	318/318	FBG	MD=-11.96 (-19.94, -3.97)	60%	NO
8	225/224	HbAc1	MD=-0.37 (-0.57, -0.17)	0%	NO
3	75/77	HOMA-IR	MD=-1.86 (-4.10, 0.39)	61%	NO

SR: systematic reviews and meta-analyses; FBG: fasting blood glucose, HbA1c: glycosylated hemoglobin, HOMA-IR: homeostatic model assessment for insulin resistance; I/C: intervention/comparison; NR: no report; MD: mean difference; SMD: standard mean difference; WMD: weighted mean difference.

[†]The number of RCTs actually found in the meta-analysis.

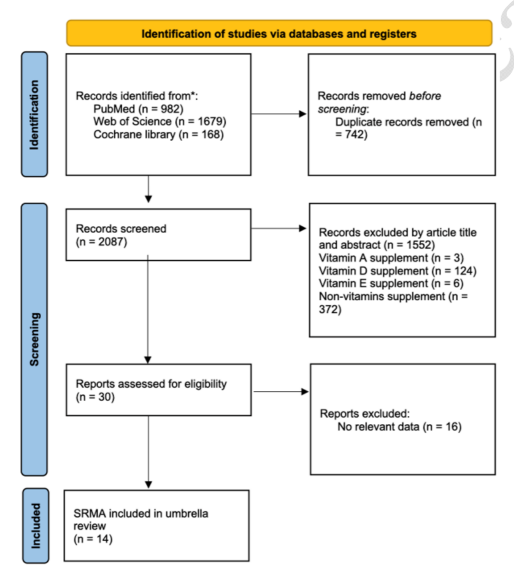


Figure 1. PRISMA Flow chart for search strategy exploring the effects of water-soluble on glycemic control and insulin resistance

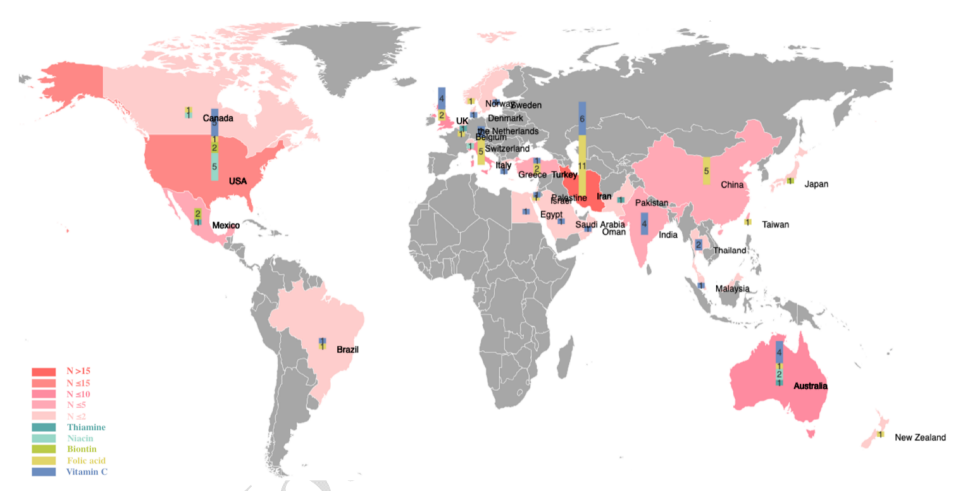


Figure 2. The locations where randomized controlled trials of water-soluble vitamin interventions were conducted

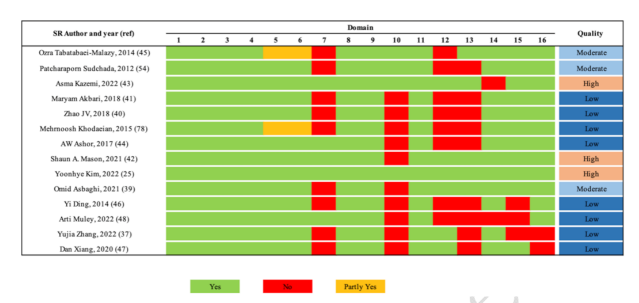


Figure 3. Results of assess the methodological quality of meta-analysis

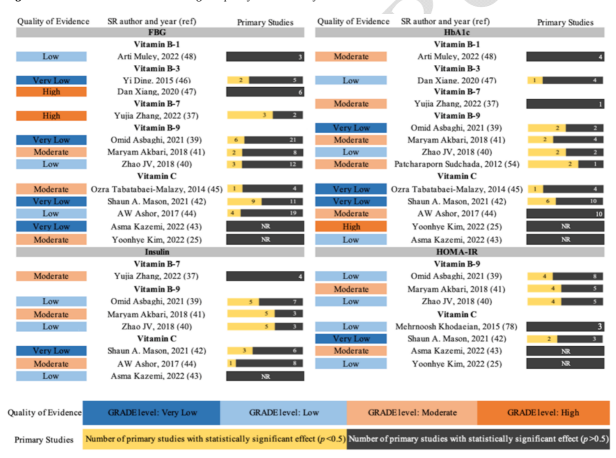


Figure 4. Summary of the strength of evidence for the effects of water-soluble vitamin supplementations. The left column indicates the meta-analyses with GRADE ratings that were very low, low, moderate, or high. Numbers in the right column indicate the modified consistency rating (number of primary randomized controlled trials with a statistically significantly positive effect or no statistically significant effect for each outcome).

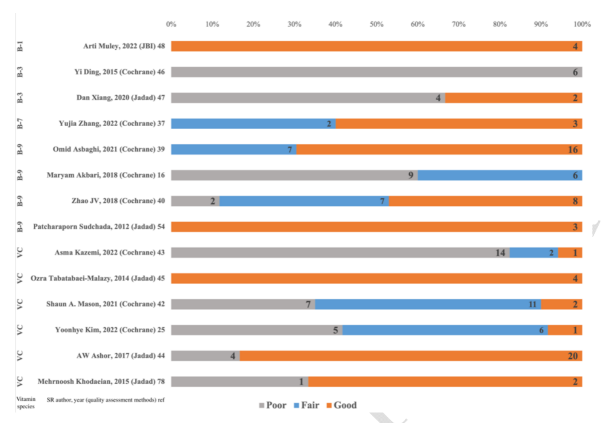


Figure 5. The quality of primary randomized controlled trials in meta-analysis

study	N	IC		ES (95% CI)
B-3				
Yi Ding, 2015 (46)	7	452/386	HH	-0.07 (-0.44 to 0.29)
Dan Xiang, 2020 (47)	6	658/615	He-I	0.18 (-0.14 to 0.50)
B-7				
Yujia Zhan, 2022 (37)	5	284/161		-1.21 (-2.73 to 0.31)
B-9				
Omid Asbaghi, 2021 (39)	27	17379/17235	──	-2.17 (-3.69 to -0.65)
VC				
AW Ashor, 2017 (44)	13	NR	++√	-0.44 (-0.81 to -0.07)
Shaun A. Mason, 2021 (42)	20	670/635	HH-1	-0.74 (-1.17 to -0.31)
Ozra Tabatabaei-Malazy, 2014 (45)	5	184/181		-20.59 (-40.77 to -0.40)
Yoonhye Kim, 2022 (25) N: number of primary studies, IC: intervention/Comparison (): reference	12	636	-5 -1 0 1 ES (95% CI)	-11.96 (-19.94 to -3.97)

 $\textbf{Figure 6.} \ \textbf{The effects of water-soluble vitamin supplementation on FBG}$

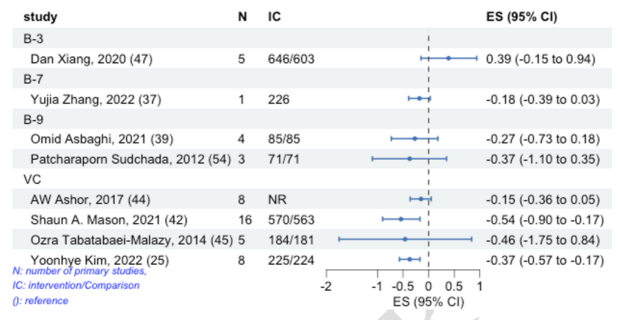


Figure 7. The effects of water-soluble vitamin supplementation on HbA1c

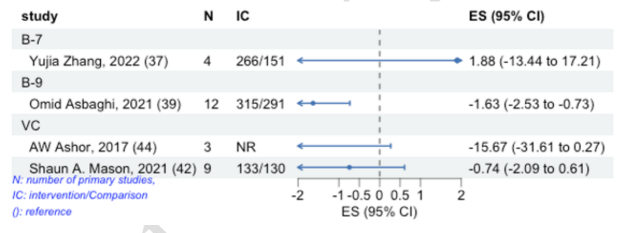


Figure 8. The effects of water-soluble vitamin supplementation on insulin

study	N	IC	ES (95% CI)
B-9			
Omid Asbaghi, 2021 (39)	12	322/295	→ -0.40 (-0.70 to -0.09)
VC			
Shaun A. Mason, 2021 (42)	5	436 (222/214)	-1.43 (-2.88 to 0.01)
Mehrnoosh Khodaeian, 2015 (78)	3	92	-0.15 (-0.49 to 0.19)
Yoonhye Kim, 2022 (25) N: number of primary studies, IC: intervention/Comparison (): reference	3	152 (75/77)	-1.86 (-4.10 to 0.39) -4 -1-0.5 0 0.5 ES (95% CI)

Figure 9. The effects of water-soluble vitamin supplementation on HOMA-IR

Supplementary Tables

Supplementary Table 1. Search strategy

Database	
PubMed	
#1	(((("Vitamins"[Mesh] OR "Vitamins" [Pharmacological Action]) OR "Vitamin B Complex"[Mesh]) OR "Dietary Supplements"[Mesh]) OR "Provitamins"[Mesh])
#2	((((((((((((((((((((((((((((((((((((((
	Acid"[Mesh]) OR "Vitamin B 6"[Mesh]) OR "Biotin"[Mesh]) OR "Folic Acid"[Mesh]) OR
	"Hydroxocobalamin"[Mesh]) OR "Inositol"[Mesh]) OR "Levoleucovorin"[Mesh]) OR "Pyridoxal
	Phosphate"[Mesh]) OR "Pyridoxamine"[Mesh]) OR "Pyridoxine"[Mesh]) OR "Tetrahydrofolates"[Mesh])
	OR "Thioctic Acid" [Mesh]) OR "Vitamin B 12" [Mesh]) OR "Cobamides" [Mesh]) OR "Ascorbic
	Acid"[Mesh]
"0	W 07 19
#3	#1 OR #2
#4	((((((((((((((((((((((((((((((((((((((
	supplements[Title/Abstract])) OR (vitamin* supplement*[Title/Abstract])) OR (vitamin B[Title/Abstract])) OR (B Vitamins[Title/Abstract])) OR (B Vitamins[Title/Abstract])) OR (Vitamin, B[Title/Abstract])) OR
	(Neurobion[Title/Abstract])) OR (Thiamin[Title/Abstract])) OR (Vitamin B1[Title/Abstract])) OR
	(Aneurin[Title/Abstract])) OR (Vitamin B 1[Title/Abstract])) OR (Thiamine Mononitrate[Title/Abstract]))
	OR (Mononitrate, Thiamine[Title/Abstract])) OR (Vitamin G[Title/Abstract])) OR (Vitamin
	B2[Title/Abstract])) OR (Vitamin B 2[Title/Abstract])) OR (Riboflavin[Title/Abstract])) OR (Vitamin
	B3[Title/Abstract])) OR (B3, Vitamin[Title/Abstract])) OR (Vitamin PP[Title/Abstract])) OR
	(Nicotinamide[Title/Abstract])) OR (3-Pyridinecarboxamide[Title/Abstract])) OR (3
	Pyridinecarboxamide[Title/Abstract])) OR (Vitamin B 3[Title/Abstract])) OR (B 3,
	Vitamin[Title/Abstract])) OR (Nicotinsäureamid Jenapharm[Title/Abstract])) OR (Jenapharm,
	Nicotinsäureamid[Title/Abstract])) OR (Nicobion[Title/Abstract])) OR (Vitamin B 5[Title/Abstract])) OR
	(B 5, Vitamin[Title/Abstract])) OR (Vitamin B5[Title/Abstract])) OR (B5, Vitamin[Title/Abstract])) OR
	(Pantothenate[Title/Abstract])) OR (Pantothenic Acid[Title/Abstract])) OR (Vitamin B6[Title/Abstract])) OR (Vitamin H[Title/Abstract])) OR (Medebiotin[Title/Abstract])) OR (Biodermatin[Title/Abstract])) OR
	(Biokur[Title/Abstract])) OR (Biotin Gelfert[Title/Abstract])) OR (Rombellin[Title/Abstract])) OR
	(Vitamin M[Title/Abstract])) OR (Vitamin B9[Title/Abstract])) OR (B9, Vitamin[Title/Abstract])) OR
	(Pteroylglutamic Acid[Title/Abstract])) OR (Folic Acid, Monopotassium Salt[Title/Abstract])) OR (Folic
	Acid, Monosodium Salt[Title/Abstract])) OR (Folic Acid, Potassium Salt[Title/Abstract])) OR (Folic Acid,
	(DL)-Isomer[Title/Abstract])) OR (Folvite[Title/Abstract])) OR (Folacin[Title/Abstract])) OR
	(Folate[Title/Abstract])) OR (B 12, Vitamin[Title/Abstract])) OR (Vitamin B12[Title/Abstract])) OR (B12,
	Vitamin[Title/Abstract])) OR (Cyanocobalamin[Title/Abstract])) OR (Cobalamins[Title/Abstract])) OR
	(Cobalamin[Title/Abstract])) OR (Acid, Ascorbic[Title/Abstract])) OR (L-Ascorbic Acid[Title/Abstract]))
	OR (Acid, L-Ascorbic[Title/Abstract])) OR (L Ascorbic Acid[Title/Abstract])) OR (Vitamin
μ г	C[Title/Abstract])
#5 #6	#3 OR #4 (((((("Diabetes Mellitus"[Mesh] OR "Diabetes Mellitus, Type 2"[Mesh] OR "Diabetes Mellitus, Type
πΟ	1"[Mesh] OR "Diabetes Mellitus, Lipoatrophic"[Mesh] OR "Diabetes, Gestational"[Mesh]) OR
	"Hyperglycemia" [Mesh]) OR ("Blood Glucose" [Mesh] OR "Glycemic Control" [Mesh])) OR "Glucose
	Tolerance Test" [Mesh]) OR "Glycated Hemoglobin A" [Mesh]) OR "Insulin" [Mesh]) OR "Insulin"
	Resistance"[Mesh]
#7	((((((((((((((((((((((((((((((((((((((
	(T2DM[Title/Abstract])) OR (hyperglycemia[Title/Abstract])) OR (hyperglycaemia[Title/Abstract])) OR
	(glucose[Title/Abstract])) OR (HbA1c[Title/Abstract])) OR (hemoglobin A1c[Title/Abstract])) OR
	(glycated hemoglobin[Title/Abstract])) OR (insulin resistance[Title/Abstract])) OR (insulin
	sensitivity[Title/Abstract])) OR (HOMA[Title/Abstract])) OR (HOMA-IR[Title/Abstract])) OR (glucose
	homeostasis[Title/Abstract])) OR (insulin secretion[Title/Abstract])) OR (insulin[Title/Abstract])) OR
	(beta-cell function[Title/Abstract])) OR (glycemic control[Title/Abstract])) OR (glucose
	tolerance[Title/Abstract])) OR (glucose metabolism[Title/Abstract])) OR (homeostatic model assessment[Title/Abstract])) OR (fasting blood sugar[Title/Abstract])) OR (FBS[Title/Abstract])) OR
	(OGTT[Title/Abstract])
	(0011[1.110/1.1004.001])
#8	#6 OR #7
#9	#5 AND #8
#10	Filters: Meta-Analysis; Systematic Reviews; published in the last 10 years

Supplementary Table 1. Search strategy (cont.)

```
Database
Web of Science
   #1
             (((((TS=(Vitamins)) OR TS=(Vitamin B Complex)) OR TS=(Antioxidants)) OR TS=(Multivitamins)) OR
             TS=(Multivitamins)) OR TS=(vitamin* supplement*)
   #2
             TS=(Thiamine Mononitrate)) OR TS=(Vitamin G)) OR TS=(Vitamin B2)) OR TS=(Riboflavin)) OR
             TS=(Vitamin B3)) OR TS=(Vitamin PP)) OR TS=(Nicotinamide)) OR TS=(3-Pyridinecarboxamide)) OR
             TS=(Papulex)) OR TS=(Papulex)) OR TS=(Nicotinsäureamid Jenapharm)) OR TS=(Enduramide)) OR
             TS=(Nicobion)) OR TS=(Vitamin B 5)) OR TS=(Zinc Pantothenate)) OR TS=(Calcium Pantothenate)) OR
             TS=(Pantothenic Acid)) OR TS=(Vitamin B6)) OR TS=(Vitamin H)) OR TS=(Deacura)) OR
             TS=(Gabunat)) OR TS=(Medebiotin)) OR TS=(Biodermatin)) OR TS=(Biotin Gelfert)) OR TS=(Biotin
             Hermes)) OR TS=(Rombellin)) OR TS=(Vitamin M)) OR TS=(Vitamin B9)) OR TS=(Pteroylglutamic
             Acid)) OR TS=(Folic Acid)) OR TS=(Folvite)) OR TS=(Folacin)) OR TS=(Folacin)) OR TS=(Vitamin B12))
             OR TS=(Cyanocobalamin)) OR TS=(Cobalamin)) OR TS=(Eritron)) OR TS=(Ascorbic Acid)) OR
             TS=(Vitamin C)) OR TS=(Hybrin)) OR TS=(Magnorbin)) OR TS=(Sodium Ascorbate)) OR TS=(Ferrous
             Ascorbate)) OR TS=(Magnesium Ascorbate)) OR TS=(Magnesium di-L-Ascorbate)
   #3
             #1 OR #2
   #4
             OR TS=(hyperglycaemia)) OR TS=(glucose)) OR TS=(HbA1c)) OR TS=(hemoglobin A1c)) OR
             TS=(glycated hemoglobin)) OR TS=(insulin resistance)) OR TS=(insulin sensitivity)) OR TS=(HOMA))
             OR TS=(HOMA-IR)) OR TS=(glucose homeostasis)) OR TS=(insulin secretion)) OR TS=( insulin)) OR
             TS=(beta-cell function)) OR TS=(glycemic control)) OR TS=(glucose tolerance)) OR TS=(glucose
             metabolism)) OR TS=(homeostatic model assessment)) OR TS=(fasting blood sugar)) OR TS=(FBS)) OR
             TS=(OGTT)
   #5
             #3 AND #4
   #6
             (TS=(meta analyses*)) OR TS=(systematic review*)
   #7
             #5 AND #6
Cochrane Library
   #1
             MeSH descriptor: [Vitamins] explode all trees
   #2
             MeSH descriptor: [Vitamin B Complex] explode all trees
   #3
             MeSH descriptor: [Antioxidants] explode all trees
   #4
             MeSH descriptor: [Biotin] explode all trees
   #5
             MeSH descriptor: [Folic Acid] explode all trees
   #6
             MeSH descriptor: [Formyltetrahydrofolates] explode all trees
   #7
             MeSH descriptor: [Inositol] explode all trees
   #8
             MeSH descriptor: [Leucovorin] explode all trees
   #9
             MeSH descriptor: [Niacin] explode all trees
   #10
             MeSH descriptor: [Niacinamide] explode all trees
   #11
             MeSH descriptor: [Nicorandil] explode all trees
   #12
             MeSH descriptor: [Nicotinic Acids] explode all trees
   #13
             MeSH descriptor: [Pyridoxal] explode all trees
             MeSH descriptor: [Pyridoxal Phosphate] explode all trees
   #14
             MeSH descriptor: [Pyridoxamine] explode all trees
   #15
   #16
             MeSH descriptor: [Pyridoxine] explode all trees
   #17
             MeSH descriptor: [Riboflavin] explode all trees
   #18
             MeSH descriptor: [Tetrahydrofolates] explode all trees
   #19
             MeSH descriptor: [Thiamine] explode all trees
   #20
             MeSH descriptor: [Thioctic Acid] explode all trees
   #21
             MeSH descriptor: [Vitamin B 12] explode all trees
   #22
             MeSH descriptor: [Vitamin B 6] explode all trees
   #23
             MeSH descriptor: [Ascorbic Acid] explode all trees
   #24
             #1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16 or #17
             or #18 or #19 or #20 or #21 or #22 or #23
   #25
             ((thiamin*) or (niacin*) or (riboflavin*) or (folic acid) or (folate*) or (cobalamin*) or (biotin*) or
             (neurobion*) or (pantothenic acid*) or (pyridox*) or (vitamin b*)):ti,ab,kw AND ("ascorbic acid" or
             "vitamin C" or "L-Ascorbic Acid" or "Acid, L-Ascorbic" or "L Ascorbic Acid" or "Hybrin" or
             "Magnorbin" or "Sodium Ascorbate" or "Ascorbate, Sodium" or "Ascorbic Acid, Monosodium Salt' or
             "Ferrous Ascorbate' or " Ascorbate, Ferrous" or "Magnesium Ascorbate" or "Ascorbate, Magnesium" or
             "Magnesium di-L-Ascorbate" or "Magnesium di L Ascorbate" or "di-L-Ascorbate, Magnesium" or
             "Magnesium Ascorbicum"):ti,ab,kw (Word variations have been searched)
   #26
             #24 or #25
   #27
             MeSH descriptor: [Diabetes Mellitus] explode all trees
   #28
             MeSH descriptor: [Hyperglycemia] explode all trees
```

Supplementary Table 1. Search strategy (cont.)

Database	
Cochrane I	<i>i</i> brary
#29	MeSH descriptor: [Glycemic Control] explode all trees
#30	MeSH descriptor: [Blood Glucose] explode all trees
#31	MeSH descriptor: [Glucose Tolerance Test] explode all trees
#32	MeSH descriptor: [Glycated Hemoglobin A] explode all trees
#33	#26 or #27 or #28 or #29 or #30 or #31 or #32
#34	("diabetes" OR "diabetes mellitus" OR "T2DM" OR "hyperglycemia" OR "hyperglycaemia glucose" OR "HbA1c" OR "hemoglobin A1c" OR "glycated hemoglobin" OR "insulin resistance" OR "insulin sensitivity" OR "HOMA" OR "HOMA-IR" OR "glucose homeostasis" OR "insulin secretion" OR "insulin" OR "beta-cell function" OR "glycemic control" OR "glucose tolerance" OR "glucose metabolism" OR
#35	"homeostatic model assessment" OR "fasting blood sugar" OR "FBS" OR "OGTT"):ti,ab,kw #33 or #34
#36	#34 and #35
#37	Filters: Reviews; published in the last 10 years

TWI: total water intake; TDF: total drinking fluids; WFF: water from food; EFI: exercise-related fluid intake; NEFI: non-exercise-related fluid intake.

Values were shown as medians (QR).

*p<0.05 there were statistically significant differences between different PAEE or MET groups; **p<0.05 there was statistically significant trend with the PAEE or MET level increase.

 $^{\dagger}p$ <0.05 compared with Gp1; $^{\ddagger}p$ <0.05 compared with Gp2; $^{\$}p$ <0.05 compared with Gm1; $^{\dagger}p$ <0.05 compared with Gm3.





Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	n			Total, n (interventio	n/comparison)
FBG						_/\`	
Vitamin B-1 Arti Muley, 2022 ⁴⁸	Rabbani N, 2009 González-Ortiz M, 201 Alaei Shahmiri F, 2013		overweight or ob	esity		40 24 (12/12) 17	
Vitamin B-3	ruaci Shailinii 1, 2013	nypergryc	cime subjects			17	
Yi Ding, 2015 ⁴⁶	Pang, 2014 MacLean, 2011 Hamilton, 2010	T2DM T2DM T2DM		0		(12/12) (298/277) (7/8)	
	Sorrentino, 2010	T2DM				(15/15)	
	Fazio, 2010	MetS				(58/31)	
	Elam, 2000	DM				(49/55)	
	Garg, 1990	T2DM				(13/13)	
SR Author and year	Male / female	Intervention Dose	Duration	Comparator	Study design	Setting	Quality assessment
FBG		Dosc	Duration				assessment
Vitamin B-1							
Arti Muley, 2022 ⁴⁸	Unequal distribution Unequal distribution	300 mg/day 150 mg/day	3months 1months	placebo placebo	parallel parallel	Pakistan Mexico	JBI, 24/26 JBI, 23/26
Vitamin B-3	Unequal distribution	100 mg/day	3weeks	placebo	crossover	Australia	JBI, 25/26
Yi Ding, 2015 46	NR	1-2 g/day	12weeks	Rosuvastatin	Crossover	Australia	Cochrane, poor
11 Ding, 2013	NR NR	1-3 g/day	36weeks	Placebo with lipid- modifying regimen	Parallel; DB	USA	Cochrane, poor
	NR	1500 mg/day	20weeks	Statin	Parallel; SB	Australia	Cochrane, poor
	NR	500-1500 mg/day	3months	placebo	Parallel	Switzerland	Cochrane, poor
	NR	500-2000 mg/day	64weeks	E/S (10/20 mg)	Parallel; DB	USA	Cochrane, poor
	NR	1500-3000 mg/day	18weeks	placebo	Parallel; DB	USA	NR
	NR	150-4500 mg/day	8weeks	placebo	Crossover	USA	Cochrane, poor

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's author	and year Population	1			Total, n	, , ,
(SR Author and year (ref))						(interventio	n/comparison)
FBG						-/ *	
Vitamin B-3							
Dan Xiang, 2020 47	Garg, 1990	T2DM				13/13	
	Elam 2000	T2DM				64/61	
	Hamilton 2010	T2DM				7/8	
	Sorrentino 2010	T2DM				15/15	
	Pang 2014	T2DM				12/12	
	Goldberg 2016	T2DM				547/506	
Vitamin B-7							
Yujia Zhang, 2022 ³⁷	Cristina. 2006	T2MD				18 (10/8)	
	Cesar . 2007	T2MD				348 (226/12	22)
	Armida,2004	T2MD				15 (10/5)	
	Gregory,2006	T2MD				36 (20/16)	
	Masaru,1993	T2MD				28 (18/10)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
·		Dose	Duration	_		· ·	assessment
FBG							
Vitamin B-3							
Dan Xiang, 2020 47	26M/F	4.5 g/d	8.0wk	Placebo	Crossover	US	Jadad, poor
	109M/16F	3000 mg/d	18.0weeks	Placebo	Parallel; DB	US	Jadad, good
	NR	1500 mg/d	20.0weeks	Statin	Parallel; DB	Australia	Jadad, poor
	0M/30F	1500 mg/d	3.0months	Placebo	Parallel; DB	Switzerland	Jadad, poor
	58.8%M	NR	12.0week	Rosuvastatin	Crossover	Australia	Jadad, poor
	1053M	NR	12.0months	Simvastatin/ezetimibe	Parallel; DB	USA and Canada	Jadad, good
Vitamin B-7					,		, &
Yujia Zhang, 2022 ³⁷	11/7	15mg/day	28days	PC	parallel	Mexico	Cochrane, good
<i>y 5</i> ,	140/208	2mg/day	90days	PC	parallel	United States	Cochrane, good
	NR	6.14µmol/d	28days	PC	parallel	Mexico	Cochrane, good
	NR	2mg/day	4weeks	PC	parallel	USA	Cochrane, fair
	NR	9mg/day	NR	PC	parallel	Japan	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's author and year		Population			Total, n		
(SR Author and year (ref))						(intervention	on/comparison)	
FBG								
Vitamin B-9								
Omid Asbaghi, 2021 39	Gargari, 2011		Overweight and obese men with type 2 diabetes			48 (24/24)		
	Cagnacci, 2009		Postmenopausal		30 (15/15)			
	Mangoni, 2005		T2DM			26 (13/13)		
	Moens, A.L, 2007		Acute myocardial infarction			40 (20/20)		
	Aarsand, 1998		T2DM			28 (14/14)		
	Doshi, 2001		Coronary artery disease	50 (50/50)				
	Doshi, 2002		Coronary artery disease		33			
	Sheu, 2005		Obese women			74 (36/38)		
	Villa, 2005	Postmenopausal				20 (10/10)		
	Moat, 2006 (A)		Coronary artery disease			59 (30/15)		
	Moat, 2006 (B)		Coronary artery disease			54 (25/14)		
	Solini, 2006		Overweight subjects			60 (30/30)		
	Title, 2006		T2DM		19 (19/19)			
_	Mao, 2008 (A) Mild to moderate primary hypertension						5)	
SR Author and year	Male / female	Interventio		Comparator	Study design	Setting	Quality	
		Dose	Duration				assessment	
FBG								
Vitamin B-9	103.5	- /.		20		_		
Omid Asbaghi, 2021 ³⁹	48M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good	
	30F	15mg/d	3weeks	PC	parallel; DB	Italy	Cochrane, good	
	14/12	5mg/d	4weeks	PC	parallel; DB	Australia	Cochrane, fair	
	35/5	10mg/d	6weeks	PC	crossover; DB	Belgium	Cochrane, good	
	21/7	0.25mg/d	12weeks	PC	parallel; DB	Norway	Cochrane, fair	
	44/6	5mg/d	// 6weeks	PC	parallel	United Kingdom	Cochrane, fair	
	30/3	5mg/d	6weeks	PC	crossover	United Kingdom	Cochrane, fair	
	74F	5mg/d	12weeks	PC	parallel; DB	Taiwan	Cochrane, good	
	20F	7.5mg/d	8weeks	PC	parallel	Italy	Cochrane, fair	
	52/7	0.4mg/d	6weeks	PC	parallel; DB	USA	Cochrane, good	
	46/8	5mg/d	6weeks	PC	parallel; DB	USA	Cochrane, good	
	19/41	2.5mg/d	12weeks	PC	parallel	Italy	Cochrane, fair	
	9/10	10mg/d	2weeks	PC	crossover; DB	Canada	Cochrane, good	
	120/175	0.4mg/d	8weeks	No intervention	parallel; DB	China	Cochrane, good	

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author and year		Population				Total, n (intervention/comparison)	
FBG Vitamin B-9							(mer vention comparison)	
Omid Asbaghi, 2021 ³⁹	Mao, 2008 (B)		Mild to moderate primary hypertension				297 (148/74)	
	Palomba, 2010 Aghamohammad, 2011 Grigoletti, 2013 Asemi, 2014 (A) Asemi, 2014 (B) Asemi, 2016		Polycystic ovary syndrome T2DM HIV-infected individuals Overweight women with polycystic ovary syndrome Overweight women with polycystic ovary syndrome Cervical intraepithelial neoplasia grade 1				47 (23/24) 68 (34/34) 30 (15/15) 81 (27/14) 81 (27/13) 58 (29/29)	
	Hashemi, 2016		Pre-eclamptic patients				85 (43/42)	
	Qin, 2016		Hypertension				20030 (10014/10016)	
	Talari, 2016		Metabolic syndrome				60 (30/30)	
	Li Y, 2017 (A)		Diabetics				1636 (800/836)	
	Li Y, 2017 (B)		Nondiabetics				11435 (5711/5724)	
	Bahmani, 2018		Endometrial hyperplasia				60 (30/30)	
SR Author and year	Male / female	Intervent	ion	Comparator	Study design	Setting	Quality	
		Dose	Duration				assessment	
FBG								
Vitamin B-9								
Omid Asbaghi, 2021 ³⁹	126/171	0.8mg/d	8weeks	No intervention	parallel; DB	China	Cochrane, good	
	47F	0.4mg/d	25weeks	PC	parallel; DB; non-random	Italy	Cochrane, good	
	68M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good	
	14/16	5mg/d	4weeks	PC	parallel; DB	Brazil	Cochrane, good	
	81F	1mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good	
	81F	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good	
	58F	5mg/d	25weeks	PC	parallel; DB	Iran	Cochrane, good	
	85F	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good	
	8295/11735	0.8mg/d	234 days	No intervention	parallel; DB	China	Cochrane, good	
	26/34	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, good	
	585/1051	0.8mg/d	229days	No intervention	parallel; DB	China	NR	
	4444/6991	0.8mg/d	229days	No intervention	parallel; DB	China	NR	
	60F	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, good	

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	and year Population	Population				Total, n		
(SR Author and year (ref))						(intervention	(intervention/comparison)	
FBG						/ ·		
Vitamin B-9						7		
Maryam Akbari, 2018 ¹⁶	Gargari BP, 2011		Overweight and obese men with type 2 diabetes				48 (24/24) 26 (13/13)	
	Mangoni AA, 2005			T2DM				
	Asemi Z, 2014		Women with polycystic ovary syndrome				54 (27/27)	
	Talari HR, 2016		Patients with metabolic syndrome				60 (30/30)	
	Khiavi A, 2011		T2DM				64 (34/34) 50 (25/25)	
	Setola E, 2004		Patients with metabolic syndrome					
	Solini A, 2006		Overweight subjects				60 (30/30)	
	Title LM, 2006	T2DM	T2DM				38 (19/19)	
	Doshi SN, 2002 Patients with coronary artery disease				33 (16/17)	33 (16/17)		
	Sheu WH-H, 2005	Obese women				74 (36/38)	74 (36/38)	
Zhao JV, 2018 40	Talari, 2016	With type	2 diabetes at base	eline; Overweight and	stable CHD	60 (30/30)		
	Qin, 2016	Hypertens	Hypertension				15951 (7960/7991)	
	Asemi, 2016		Cervical intraepithelial neoplasia grade 1				58 (29/29)	
	Asemi, 2014		Overweight or obesity, and PCOS				54 (27/27)	
						. .		
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality	
		Dose	Duration				assessment	
FBG			00					
Vitamin B-9								
Maryam Akbari, 2018 ¹⁶	NR	5mg/d	8weeks	PC	r ,	Iran	Cochrane, fair	
	NR	5mg/d	4weeks	PC	parallel; DB	Australia	Cochrane, poor	
	NR	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, poor	
	NR	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, poor	
	NR	5mg/d	8weeks	PC	NR	Iran	Cochrane, fair	
	NR	Folate plus vitamins	8weeks	PC	parallel; DB	Italy	Cochrane, fair	
		B6 or B12, 5mg/d						
	NR	2.5mg/d	12weeks	PC	NR	Italy	Cochrane, poor	
	NR	10mg/d	2weeks	PC	crossover	Canada	Cochrane, poor	
	NR	5mg/d	6weeks	PC	NR	UK	Cochrane, fair	
	NR	5mg/d	12weeks	PC	parallel; DB	Taiwan	Cochrane, fair	
Zhao JV, 2018 ⁴⁰	both	5mg/d	12weeks	placebo	parallel	Iran	Cochrane, good	
	both	0.8mg/d	4.5years	placebo	parallel	China	Cochrane, fair	
	58F	5mg/d	6months	placebo	parallel	Iran	Cochrane, fair	
	54F	1mg/d	8weeks	placebo	parallel	Iran	Cochrane, fair	

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's autho	or and year	Population		Total, n				
(SR Author and year (ref))							(intervention	comparison)	
FBG						_/			
Vitamin B-9									
Zhao JV, 2018 40	Gargari, 2011		With type 2 diabetes at base	eline Overweight			48 (24/24)		
	Liu, 2011		With type 2 diabetes at base	eline、BMI≥22 kg/m2			182 (92/90)		
	Kurt, 2010		Vitamin B12 deficiency				44 (24/20)		
	Mashavi, 2008		T2DM			57 (28/29)			
	Mao, 2008		Baseline fasting glucose≥6.			60 (28/32)			
	Gu, 2008		T2DM				60 (30/30)		
	Solini, 2006		NO				60 (30/30)		
	Title, 2006		T2DM		38 (19/19)				
	Mangoni, 2005		type 2 diabetes, microalbum	ninuria			26 (13/13)		
	Villa, 2005		NO				20 (10/10)		
	Setola, 2004		metabolic syndrome, hyperi	nsulinemia			50 (25/25)		
	Masaru,1993		T2MD				28 (18/10)		
SR Author and year	Male / female	Intervent	ion	Comparator	Study design	Setting	5	Quality	
		Dose	Duration					assessment	
FBG									
Vitamin B-9									
Zhao JV, 2018 40	48M	5mg/d	8weeks	placebo	parallel	Iran		Cochrane, fair	
	both	0.15mg/d	l 6months	placebo	parallel	China		Cochrane, good	
	both	5mg/d	8weeks	placebo	parallel	Turkey	7	Cochrane, fair	
	both	1mg/d	4months	placebo	parallel	Israel		Cochrane, good	
	both	0.8 mg/d	8weeks	placebo	parallel	China		Cochrane, good	
	both	5mg/d	2weeks	placebo	parallel	China		Cochrane, fair	
	both	2.5mg/d	12weeks	placebo	parallel	Italy		Cochrane, poor	
	both	10mg/d	2 weeks	placebo	Crossover	Canada	a	Cochrane, good	
	both	5mg/d	4weeks	placebo	parallel	UK		Cochrane, good	
	20F	7.5mg/d	8weeks	placebo	parallel	Italy		Cochrane, poor	
	both	5mg/day	2months	placebo	parallel	Italy		Cochrane, fair	

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	or and year	Population			Total, n (interventi	on/comparison)
FBG							-
Vitamin C							
Asma Kazemi, 2022 43	Tousoulis, 2007	,	T2DM			(13/13)	
	Nayaka, 2013	ŗ	T2DM			30	
	Ghaffari, 2015	ŗ	T2DM			(17/14)	
	Bishop, 1984]	Diabetic Hyperlipidemia	,25/25		25/25	
	Dakhale, 2011	,	T2DM			33	
	Siavash, 2014	,	T2DM			15/15	
	Lu, 2005	,	T2DM	· · · · · · · · · · · · · · · · · · ·		17	
	Gillani, 2017	ŗ	T2DM			139/142	
	Bhatt, 2012		T2DM			30/29	
	Devanandan, 2020	ŗ	T2DM			68/67	
	Kunsongkeit, 2019	ŗ	T2DM			15/16	
	Mason, 2018		T2DM			27/27/27	
	El-Aal, 2018	ŗ	T2DM			10/10	
	Ramzy Ragheb, 2020	,	T2DM			20/13	_
SR Author and year	Male / female	Intervention Dose	n Duration	Comparator	Study design	Setting	Quality assessment
FBG		Dosc	Duration				assessment
Vitamin C		- /-				_	
Asma Kazemi, 2022 43	NR	2g/day	4weeks	No intervention	Parallel	Greece	Cochrane, poor
	NR	1g/d	8weeks	Placebo	Parallel	India	Cochrane, poor
	NR	800 mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	NR	500mg/d	52weeks	Placebo	Cross-over	UK	Cochrane, poor
	NR	1000mg/d	12weeks	Placebo	Parallel	India	Cochrane, fair
	NR	1000mg	// 6weeks	No intervention	Parallel	Iran	Cochrane, poor
	NR	3g/d	2weeks	Placebo	Cross-over	Sweden	Cochrane, poor
	NR	500mg/d	52weeks	Placebo	Parallel	Saudi Arabia	Cochrane, poor
	NR	500mg/d	12weeks	Placebo	Parallel	Oman	Cochrane, poor
	NR	500mg/d	36weeks	Placebo	Parallel	India	Cochrane, poor
	NR	500mg/d	8weeks	Placebo	Parallel	Thailand	Cochrane, poor
	NR	1000mg/d	17weeks	Placebo	Parallel	Australia	Cochrane, good
	20M	800mg/d	12weeks	Placebo	Parallel	Palestine	Cochrane, poor
	NR	500mg/d	8weeks	placebo	Parallel	Egypt	Cochrane

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	n			Total, n	n/comparison)
FBG							, , , , , , , , , , , , , , , , , , ,
Vitamin C							
Asma Kazemi, 2022 43	Sanguanwong, 2016	T2DM				50	
	Froghi, 2018	T2DM				21/21	
	Chen, 2006	T2DM				15/17	
	Paolisso, 1995	T2DM			V. A	40	
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	Bhatt J, 2012	T2DM				30/29	
	Shakouri, 2011	T2DM				32/33	
	Delvarianzadeh M, 200	8 T2DM				68/68	
	Farvid M, 2000 (A)	diabetics				28/28	
	Farvid M, 2000 (B)	diabetics				26/23	
Shaun A. Mason, 2021 42	Bhatt, 2012	T2DM				59 (30/29)	
	Hui Chen, 2006	T2DM				32(15/17)	
	Dakhale, 2011	T2DM	/ ·			70(35/35)	
	Devanandan, 2020	T2DM				135(68/67)	
SR Author and year	Male / female	Intervention	D. Air	Comparator	Study design	Setting	Quality
FBG		Dose	Duration				assessment
Vitamin C							
Asma Kazemi, 2022 43	NR	1000mg/d	8weeks	Placebo	Parallel	Thailand	Cochrane, fair
	NR	500mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	NR	800mg/d	4weeks	Placebo	Parallel	USA	Cochrane, poor
	both	1000 mg/d	12 weeks	Placebo	Cross-over	Italy	Cochrane, poor
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	NR	500mg/d, AA	3month	placebo	open label; cross over	NR	Jadad, good
	65M	200mg/d, AA	8weeks	500mg/d; EPA	DB; cross over	Iran	Jadad, good
	NR	1250mg/d, AA	3month	placebo	DB; cross over	NR	Jadad, good
	NR	500mg/d, AA	4weeks	placebo, VE	crossover	NR	Jadad, good
	NR	500mg/d, AA	9weeks	placebo, VE	crossover	NR	Jadad, good
Shaun A. Mason, 2021 42	42/17	500mg/day	90days	active control	parallel	India	Cochrane, poor
	13/19	800mg/day	28days	placebo	parallel; DB	US	Cochrane, fair
	28/38	1000mg/day	84days	placebo	parallel; DB	India	Cochrane, good
	84/51	1000mg/day	270days	placebo	parallel	India	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	l			Total,	n ention/comparison)
FBG						(IIICI)	entron comparison)
Vitamin C							
Shaun A. Mason, 2021 42	El-Aal, 2018	T2DM				40(10/	10/10/10)
	Foroghi, 2018	T2DM				78(38/	
	Ghaffari, 2015	T2DM				31(17/	
	Gillani, 2017	T2DM				,	52/152)
	Kunsongkeit, 2019	T2DM				31(15/	16)
	Lu, 2005	T2DM			,	(17/17	
	Mahmoudabadi, 2011	T2DM				34(17/	17)
	Mason, 2016	T2DM				(7/7)	,
	Mason, 2019	T2DM				(27/27))
	Paolisso, 1995	T2DM				(40/4	(0)
	Rafighi, 2013	T2DM				84(44/	40)
	Dakhale, 2011	T2DM				70(35/	
	Devanandan, 2020	T2DM				135(68	3/67)
	Ragheb, 2020	T2DM				33(20/	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
Sit ridiior and year	Wate / Temale	Dose	Duration	Comparator	Study design	Setting	assessment
FBG							
Vitamin C							
Shaun A. Mason, 2021 42	40M	1000mg/day	90days	placebo	parallel	Palestine	Cochrane, poor
	41/37	500mg/day	60days	placebo	parallel; DB	Iran	Cochrane, fair
	13/18	800mg/day	60days	placebo	parallel	Iran	Cochrane, fair
	183/121	500mg/day	365days	placebo	parallel	Malaysia	Cochrane, poor
	9/22	500mg/day	60days	placebo	crossover; DB	Thailand	Cochrane, poor
	12/5	3000mg/day	14days	placebo	crossover; DB	Sweden	Cochrane, fair
	34M	200mg/day	56days	placebo	parallel; DB	Iran	Cochrane, fair
	12/1	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, fair
	26/5	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, good
	19/21	1000mg/day	120days	placebo	crossover; DB	Italy	Cochrane, fair
	44/40	800mg/day	90 days	placebo	parallel	Iran	Cochrane, fair
	28/38	1000mg/day	84days	placebo	parallel; DB	India	Cochrane, good
	84/51	1000mg/day	270days	placebo	parallel	India	Cochrane, fair
	10/23	500mg/day	56days	only received anti-	parallel	Egypt	Cochrane, poor
				diabetes treatment			

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's author	and year Population	1			Tota	
(SR Author and year (ref))						(inte	ervention/comparison)
FBG							
Vitamin C	D 11 0010	map					• 0
Shaun A. Mason, 2021 42	Rekha, 2013	T2DM				(55//	
	Sanguanwong, 2016	T2DM				(50/:	,
	Siavash, 2014	T2DM				`	5/15)
77 1 77 000025	Tousoulis, 2007	T2DM				`	3/13)
Yoonhye Kim, 2022 ²⁵	Hui Chen, 2006	T2DM				(15/	
	Ali Abd El-Aal, 2018	T2DM				(10/	The state of the s
	Ganesh, 2011	T2DM				(35/:	,
	M Evans, 2003	T2DM					0/10)
	Ghaffari, 2015	T2DM					7/14)
	Mahmoudabadi, 2014	T2DM					20/20)
	Mason, 2019	T2DM				(27/	
	Paolisso, 1995	T2DM				(40/4	
	Rekha, 2013	T2DM				(30/	
	Sanguanwong, 2016	T2DM				(50/:	50)
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
FBG			0151				
Vitamin C							
Shaun A. Mason, 2021 42	NS	1000 or	56days	active cotrol	parallel	India	Cochrane, poor
		2000mg/day	-0.1				
	NS	1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
	12/18	1000mg/day	42days	active cotrol	parallel	Iran	Cochrane, fair
M. 1 M. 2022 ²⁵	14/12	200mg/day	28days	active cotrol	parallel	Greece	Cochrane, poor
Yoonhye Kim, 2022 ²⁵	NR	800mg/day	4weeks	PC	parallel; DB	USA	Cochrane, poor
	NR NB	1000mg/day	12weeks	PC	parallel	USA	Cochrane, fair
	NR	1000mg/day	12weeks	PC	parallel; DB	India	Cochrane, good
	17/3	1000mg/day	6weeks	PC	parallel	UK	Cochrane, fair
	NR 40M	800mg/day	8weeks	placebo	parallel	NR	Cochrane, poor
	40M	200mg/day	8weeks	placebo	parallel; DB	Iran	Cochrane, fair
	NR	1000mg/day	16weeks	placebo	crossover; DB	Australia	Cochrane, fair
	NR ND	1000mg/day	16weeks	placebo	crossover; DB	Italy	Cochrane, fair
	NR	1000mg/day	8weeks	placebo	parallel	NR NB	Cochrane, poor
	NR	1000mg/day	8weeks	placebo	parallel; DB	NR	Cochrane, poor

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author and	year Population	1			Total, n	n/comparison)
FBG						(intervention	i/companison)
Vitamin C							
Yoonhye Kim, 2022 ²⁵	Bhatt JK, 2012	T2DM				(33/32)	
	Ellulu MS, 2015	T2DM				(36/36)	
AW Ashor, 2017 44	Ganesh, 2011	T2DM				(33/33)	
	Ellulu, 2015	T2DM				(31/33)	
	Tousoulis, 2007	T2DM				26 (13/13)	
	Hui Chen, 2006	T2DM			4	32 (17/15)	
	Mahmoudabadi, 2011	T2DM				34 (17/17)	
	Zahra Rafighi, 2013	T2DM				170	
	Mansour Siavash, 2014	T2DM				35 (20/15)	
	Shaun A Mason, 2016	T2DM				14 (7/7)	
	Davison, 2008 (B)	T1DM				26	
	F Klein, 1995	T1DM				24 (12/12)	
SR Author and year	Male / female In	ntervention		Comparator	Study design	Setting	Quality
•	D	ose	Duration			, and the second	assessment
FBG							
Vitamin C							
Yoonhye Kim, 2022 ²⁵	NR 50	00mg/day	12weeks	PC	parallel	NR	Cochrane, poor
	NR 10	000mg/day	8weeks	PC	parallel	Malaysia	Cochrane, fair
AW Ashor, 2017 44		000mg/day	84days	placebo	parallel; DB	India	Jadad, 3
	22/50	000mg/day	56days	No intervention	parallel	Malaysia	Jadad, 4
		000mg/day	30days	No intervention	parallel	Athens, Greece	Jadad, 3
		00mg/day	28days	placebo	parallel; DB	USA	Jadad, 5
		00mg/day	56days	placebo	parallel; DB	Iran	Jadad, 3
		C: 800mg/day;	90 days	placebo	parallel	Iran	Jadad, 4
	Vi	itamin C (266.7					
		ng), vitamin E (300					
		J), vitamin C+E					
		300IU+266.7mg)					
		000mg/day	42days	600 mg gemfibrozil	parallel	Iran	Jadad, 2
		000mg/day	120days	placebo	crossover; DB	Australia	Jadad, 5
		000mg/day	1days	placebo	parallel; DB	UK	Jadad, 3
	24M 60	000mg/day	28days	placebo	parallel; DB	Denmark	Jadad, 3

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author a	nd year Population	l			Total, n (intervention	n/comparison)
FBG							•
Vitamin C							
AW Ashor, 2017 44	Bhatt JK, 2012	T2DM				59	
	Gutierrez AD, 2013	Healthy				28	
	Ghaffari , 2015	T2DM		()		31	
	N Bishop, 1985 (B)	T2DM				25	
	N Bishop, 1985 (A)	T2DM				50	
	C S Johnston, 1994	Healthy			4	9	
	L Pirbudak, 2004	Healthy				22 (11/11)	
	G W Davison, 2008 (A)	Healthy				26	
	Johannes Pleiner, 2002	Healthy				10	
	Simona Bo, 2007	Healthy				78 (40/38)	
	N Gokce, 1999	CAD				46 (21/25)	
	Brian A Mullan, 2005	Healthy				9 ` ′	
	David C Nieman, 1985	Healthy				28 (15/13)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
FBG							
Vitamin C							
AW Ashor, 2017 44	17/42	500mg/day	90 days	placebo	parallel	NR	Jadad, 2
	5/9	1000 mg/day	120days	placebo	Parallel	USA	Jadad, 3
	13/17	800mg/day	56days	placebo	Parallel	Iran	Jadad, 2
	11/14	500mg/day	60days	placebo	crossover; DB	UK	Jadad, 3
	13/12	500mg/day	60days	placebo	crossover; DB	UK	Jadad, 3
	2/7	1000mg/day	14days	placebo	crossover; DB	USA	Jadad, 5
	22F	AA 500 mg,	1days	fentanyl 1–2 mg/kg and	parallel	Turkey	Jadad, 2
		fentanyl 1-2 mg/kg		etomidate 0.3–0.4			
		and etomidate 0.3-		mg/kg			
		0.4 mg/kg					
	12M	1000mg/day	1days	placebo	parallel; DB	UK	Jadad, 3
	10M	72mg/day	1days	placebo	crossover; DB	Australia	Jadad, 3
	24/54	2000mg/day	14days	No intervention	parallel	Italy	Jadad, 3
	42/4	500mg/day	30days	placebo	DB	USA	Jadad, 3
	9M	2000mg/day	1days	placebo	crossover; DB	UK	Jadad, 3
	NR	1500mg/day	1days	placebo	parallel; DB	USA	Jadad, 4

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author a	and year Population	l			Total, n (intervention	n/comparison)
FBG							·
Vitamin C							
AW Ashor, 2017 44	Bhatt JK, 2012	T2DM			6/33	59	
	Gutierrez AD, 2013	Healthy				28	
	Ghaffari, 2015	T2DM				31	
	N Bishop, 1985 (B)	T2DM				25	
	N Bishop, 1985 (A)	T2DM				50	
	C S Johnston, 1994	Healthy			1	9	
	L Pirbudak, 2004	Healthy				22 (11/11)	
	G W Davison, 2008 (A)	Healthy				26	
	Johannes Pleiner, 2002	Healthy				10	
	Simona Bo, 2007	Healthy				78 (40/38)	
	N Gokce, 1999	CAD				46 (21/25)	
	Brian A Mullan, 2005	Healthy				9	
	David C Nieman, 1985	Healthy				28 (15/13)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
FBG							
Vitamin C							
AW Ashor, 2017 44	17/42	500mg/day	90 days	placebo	parallel	NR	Jadad, 2
	5/9	1000 mg/day	120days	placebo	Parallel	USA	Jadad, 3
	13/17	800mg/day	56days	placebo	Parallel	Iran	Jadad, 2
	11/14	500mg/day	60days	placebo	crossover; DB	UK	Jadad, 3
	13/12	500mg/day	60days	placebo	crossover; DB	UK	Jadad, 3
	2/7	1000mg/day	14days	placebo	crossover; DB	USA	Jadad, 5
	22F	AA 500 mg,	1days	fentanyl 1–2 mg/kg and	parallel	Turkey	Jadad, 2
		fentanyl 1-2 mg/kg		etomidate 0.3–0.4			
		and etomidate 0.3-		mg/kg			
	, ,	0.4 mg/kg					
	12M	1000mg/day	1days	placebo	parallel; DB	UK	Jadad, 3
	10M	72mg/day	1days	placebo	crossover; DB	Australia	Jadad, 3
	24/54	2000mg/day	14days	No intervention	parallel	Italy	Jadad, 3
	42/4	500mg/day	30days	placebo	DB	USA	Jadad, 3
	9M	2000mg/day	1days	placebo	crossover; DB	UK	Jadad, 3
	NR	1500mg/day	1days	placebo	parallel; DB	USA	Jadad, 4

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Popul	ation			Total, n	on/comparison)
HbA1C						(intervention	ni/companson)
Vitamin B-1							
Arti Muley, 2022 ⁴⁸	Alkhalaf, 2010	T2MI)			82	
,,	González-Ortiz, 2011		I or overweight or ob	esity		24 (12/12)	
	Rabbani, 2009	T2MI				40	
	Alkhalaf, 2010	T2MI				82	
Vitamin B-3	,						
Dan Xiang, 2020 ⁴⁷	Garg, 1990	T2DN	1		4	13/13	
8, 1	Elam, 2000	T2DN	1			64/61	
	Hamilton, 2010	T2DN				7, 8	
	Sorrentino, 2010	T2DN	1			15/15	
	Pang, 2014	T2DN				547/506	
Vitamin B-7							
Yujia Zhang, 2022 ³⁷	Cesar, 2007	T2MI)	* * *		348 (226/12	22)
Vitamin B-9	2007	12				2.0 (220/1	/
Omid Asbaghi, 2021 39	Gargari, 2011	Overv	weight and obese men	with type 2 diabetes		48 (24/24)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Ouality
Sit Hathor and your	Triale / Telliale	Dose	Duration	Comparator	Study design	Setting	assessment
HbA1C							
Vitamin B-1							
Arti Muley, 2022 ⁴⁸	77%/33%	900mg/day	12weeks	placebo	parallel	the Netherlands	JBI, 23/26
•	NR	150mg/day	1 months	placebo	parallel	Mexico	JBI, 23/26
	NR	300mg/day	3months	placebo	parallel	Pakistan	JBI, 24/26
	77%/33%	900mg/day	12weeks	placebo	parallel	the Netherlands	JBI, 23/26
Vitamin B-3		<i>U</i> ,		1	1		,
Dan Xiang, 2020 ⁴⁷	26M	4.5 g/d	8weeks	placebo	Crossover	US	Jadad, poor
٥,	109M/16F	3000mg/d	18weeks	placebo	Parallel; DB	US	Jadad, good
	15F	1500mg/d	20weeks	Statin	Parallel: DB	Australia	Jadad, poor
	25M/5F	1501mg/d	3month	Placebo	Parallel; DB	Switzerland	Jadad, poor
	874M/179F	NR	12month	Simvastatin/ezetimibe	Parallel; DB	USA and Canada	Jadad, poor
Vitamin B-3					,		, I
Yujia Zhang, 2022 37	140/208	2mg/day	90 days	PC	parallel	United States	Cochrane, good
Vitamin B-9		<i>U</i> ,	•				, 8
Omid Asbaghi, 2021 39	48M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's author and	d year Populati	on			Total, n	. ,
(SR Author and year (ref))						(intervent	ion/comparison)
HbA1C Vitamin B-9							
	Managari 2005	TODM				26 (12/12)	\
Omid Asbaghi, 2021 39	Mangoni, 2005	T2DM T2DM				26 (13/13)	
	Aarsand, 1998					28 (14/14)	
M	Aghamohammadi Khiavi,		. 1 . 1 . 1	2.1 . 2.11 .		68 (34/34)	
Maryam Akbari, 2018 ¹⁶	Gargari BP, 2011		ight and obese me	n with type 2 diabetes		48 (24/24)	
	Mangoni AA, 2005	T2DM				26 (13/13)	
	Khiavi A, 2011	T2DM				64 (34/34)	
	Alian Z, 2012	T1DM				55 (34/21)	
	Mosavi Z, 2015	T2DM			/	45 (24/21)	
TI 111 0010 10	Peña AS, 2004	T1DM				36 (15/21)	
Zhao JV, 2018 40	Gargari, 2011	• •		seline, Overweight		48 (24/24)	
	Liu, 2011	• •	be 2 diabetes at ba	seline、BMI≥22 kg/m2		182 (92/9	•
	Mashavi, 2008	T2DM				57 (28/29)	
	Mangoni, 2005	With typ	be 2 diabetes at ba	seline; Hypertension in 1	6 patients; microalbumin	uria in 8 26 (13/13))
		patients					
SR Author and year	Male / female I	ntervention		Comparator	Study design	Setting	Quality
	Ī	Oose	Duration				assessment
HbA1C							
Vitamin B-9							
Omid Asbaghi, 2021 39	14/12 5	img/d	4weeks	PC	parallel; DB	Australia	Cochrane, fair
	21/7	0.25mg/d	12weeks	PC	parallel; DB	Norway	Cochrane, fair
	68M 5	img/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
Maryam Akbari, 2018 ¹⁶	NR 5	img/d	8weeks	PC	parallel; DB	Iran	Cochrane, fair
	NR 5	img/d	4weeks	PC	parallel; DB	Australia	Cochrane, poor
	NR 5	img/d	8weeks	PC	NR	Iran	Cochrane, fair
	NR 5	img/d	8weeks	PC	crossover; DB	Iran	Cochrane, poor
		mg/d	12weeks	PC	NR	Iran	Cochrane, poor
		omg/d	8weeks	PC	crossover; DB	New Zealand	Cochrane, poor
Zhao JV, 2018 40		img/d	8weeks	placebo	parallel	Iran	Cochrane, fair
).15mg/d	6months	placebo	parallel	China	Cochrane, good
		mg/d	4months	placebo	parallel	Israel	Cochrane, good
		5mg/d	4weeks	placebo	parallel	UK	Cochrane, good

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's auth	or and year P	Population			Total, n (intervent	tion/comparison)
HbA1C							
Vitamin B-9							
Patcharaporn Sudchada, 2012 ⁵⁴	Bahram Pourghasser 2011	n Gargari, T	T2DM			24/24	
	Vahide Aghamoham	madi, 2011 T	T2DM			34/34	
	Mangoni AA, 2005		T2DM			13/13	
Vitamin C	,						
Asma Kazemi, 2022 43	Bishop, 1984	Γ	Diabetic Hyperlipidemia	a	4	25/25	
,	Dakhale, 2011		T2DM			33	
	Siavash, 2014	T	T2DM			15/ 15	
	Lu, 2005		T2DM			17	
	Gillani, 2017	Т	T2DM			139/ 142	
	Bhatt, 2012		T2DM			30/29	
	Devanandan et al, 20		T2DM			68/67	
	Kunsongkeit, 2019		T2DM			15/16	
	Mason, 2018		T2DM			27/ 27/ 27	7
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration			8	assessment
HbA1C				7			
Vitamin B-9							
Patcharaporn Sudchada,	48M	5mg/day	8weeks	Placebo	parallel	Iran	Jadad, good
2012 ⁵⁴		2 2 3			r		, 8
2012	68M	5mg/day	8weeks	Placebo	parallel; DB	Iran	Jadad, good
	14M/12F	5mg/day	4weeks	Placebo	parallel; DB	Australia	Jadad, good
Vitamin C	1 11/2 121	emg any	· · · · · · · · ·	1140000	paraner, 22	1100114114	vadad, good
Asma Kazemi, 2022 ⁴³	NR	VC, 500 mg	/d 52weeks	Placebo	Cross-over	UK	Cochrane, poor
1 1311111 111111111, 2322	NR	VC, 1000 m		Placebo	Parallel	India	Cochrane, fair
	NR	VC, 1000 m		No intervention	Parallel	Iran	Cochrane, poor
	NR /	VC, 1000 III VC, 3 g/d	2weeks	Placebo	Cross-over	Sweden	Cochrane, poor
	NR NR	VC, 500 mg		Placebo	Parallel	Saudi Arabia	Cochrane, poor
	NR	VC, 500 mg		Placebo	Parallel	Oman	Cochrane, poor
	NR	VC, 500 mg		Placebo	Parallel	India	Cochrane, poor
	NR	VC, 500 mg		Placebo	Parallel	Thailand	Cochrane, poor
	NR	VC, 1000 m		Placebo	Parallel	Australia	Cochrane, good

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	n			Total, n (intervention	/comparison)
HbA1C						(inter , entire)	tompungon)
Vitamin C							
Asma Kazemi, 2022 43	El-Aal, 2018	T2DM				10,10	
,	Sanguanwong, 2016	T2DM				50	
	Froghi, 2018	T2DM				21/21	
	Chen, 2006	T2DM			V A	15/17	
	Paolisso, 1995	T2DM				40	
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	Bhatt J, 2012	T2DM				30/29	
	Shakouri Mahmoudaba	adi, 2011 T2DM				32/33	
	Delvarianzadeh M, 200	08 T2DM				68/68	
	Farvid M, 2000 (A)	diabetics				28/28	
	Farvid M, 2000 (B)	diabetics				26/23	
Shaun A. Mason, 2022 42	Bhatt, 2012	T2DM				59 (30/29)	
•	Dakhale, 2011	T2DM				70 (35/35)	
	Devanandan, 2020	T2DM				135 (68/67)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
HbA1C							
Vitamin C							
Asma Kazemi, 2022 43	20 Male	VC, 800 mg/d	12weeks	Placebo	Parallel	Palestine	Cochrane, poor
	NR	VC, 1000 mg/d	8weeks	Placebo	Parallel	Thailand	Cochrane, fair
	NR	VC, 500 mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	NR	VC, 800 mg/d	4weeks	Placebo	Parallel	USA	Cochrane, poor
	both	1000 mg/d	12 weeks	Placebo	Cross-over	Italy	Cochrane, poor
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	NR	500mg/d; AA	3month	placebo	open label; cross over	NR	Jadad, good
	65M	200mg/d; AA	8weeks	500mg/d; EPA	DB; cross over	Iran	Jadad, good
	NR	1250mg/d; AA	3month	placebo	DB; cross over	NR	Jadad, good
	NR	500mg/d; AA	4weeks	placebo, VE	cross over	NR	Jadad, good
	NR	500mg/d; AA	9weeks	placebo, VE	cross over	NR	Jadad, good
Shaun A. Mason, 2022 42	42/17	500mg/day	90days	active cotrol	parallel	India	Cochrane, poor
	28/38	1000mg/day	84days	placebo	parallel; DB	India	Cochrane, good
	84/51	1000mg/day	270days	placebo	parallel	India	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	l			Total, (interv	n ention/comparison)
HbA1C							
Vitamin C							
Shaun A. Mason, 2022 42	El-Aal, 2018	T2DM				,	/10/10/10)
	Foroghi, 2018	T2DM				78 (38)	
	Gillani, 2017	T2DM				,	52/152)
	Kunsongkeit, 2019	T2DM				31 (15)	,
	Lu, 2005	T2DM				(17/17)	,
	Mahmoudabadi, 2011	T2DM				34(17/	17)
	Mason, 2016	T2DM				(7/7)	
	Mason, 2019	T2DM			,	(27/27)	
	Paolisso, 1995	T2DM				(40/40)	
	Rafighi, 2013	T2DM				84 (44,	,
	Ragheb, 2020	T2DM				33 (20)	
	Sanguanwong, 2016	T2DM				(50/50)	
	Siavash, 2014	T2DM				30 (15)	/15)
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
HbA1C							
Vitamin C				<i>y</i>			
Shaun A. Mason, 2022 42	40M	1000mg/day	90days	placebo	parallel	Palestine	Cochrane, poor
	41/37	500mg/day	60days	placebo	parallel; DB	Iran	Cochrane, fair
	9/22	500mg/day	365days	placebo	parallel	Malaysia	Cochrane, poor
	45191	500mg/day	60days	placebo	crossover; DB	Thailand	Cochrane, poor
	12/5	3000mg/day	14days	placebo	crossover; DB	Sweden	Cochrane, fair
	34M	200mg/day	56days	placebo	parallel; DB	Iran	Cochrane, fair
	12/1	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, fair
	26/5	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, good
	19/21	1000mg/day	120days	placebo	crossover; DB	Italy	Cochrane, fair
	44/40	800mg/day	90days	placebo	parallel	Iran	Cochrane, fair
	10/23	500mg/day	56days	No	parallel	Egypt	Cochrane, poor
	NS	1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
	12/18	1000mg/day	42days	active control	parallel	Iran	Cochrane, poor

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author an	d year Population	1			Total, n	n/comparison)
HbA1C						(Intervention	i/comparison)
Vitamin C							
Yoonhye Kim, 2022 ²⁵	Ali Abd El-Aal, 2018	T2DM				(10/10)	
	Ganesh N Dakhale, 2011	T2DM				(35/35)	
	Mahmoudabadi, 2014	T2DM				40 (20/20)	
	Mason, 2019	T2DM				(27/27)	
	Paolisso, 1995	T2DM				(40/40)	
	Bhatt JK, 2012	T2DM			,	(33/32)	
	M Evans, 2003	T2DM				20 (10/10)	
	Sanguanwong, 2016	T2DM				(50/50)	
AW Ashor, 2017 ⁴⁴	Ganesh N Dakhale, 2011	T2DM				(33/33)	
	Mahmoudabadi, 2011	T2DM				34 (17/17)	
	Zahra Rafighi, 2011	T2DM				170	
	Mansour Siavash, 2014	T2DM				35 (20/15)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
•		Dose	Duration			_	assessment
HbA1C							
Vitamin C							
Yoonhye Kim, 2022 ²⁵		1000mg/day	12weeks	PC	parallel	NR	Cochrane, fair
		1000mg/day	12weeks	PC	parallel; DB	India	Cochrane, good
		200mg/day	8weeks	placebo	parallel; DB	Iran	Cochrane, fair
		1000mg/day	16weeks	placebo	crossover; DB	Australia	Cochrane, fair
		1000mg/day	16weeks	placebo	crossover; DB	Italy	Cochrane, fair
		500mg/day	12weeks	PC	parallel	NR	Cochrane, poor
		1000mg/day	6weeks	PC	parallel	UK	Cochrane, fair
		1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
AW Ashor, 2017 ⁴⁴		1000mg/day	84days	placebo	parallel; DB	India	Jadad, 3
		200mg/day	56days	placebo	parallel; DB	Iran	Jadad, 3
		VC: 800mg/day;	90days	placebo	parallel	Iran	Jadad, 4
		vitamin C was					
		(266.7 mg), vitamin					
		C+E (300 IU+266.7					
		mg)	40.1				
	12/23	1000mg/day	42days	600 mg gemfibrozil	parallel	Iran	Jadad, 2

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year 1	Population			Total, n	ention/comparison)
HbA1C						(Interve	ention/comparison)
Vitamin C							
AW Ashor, 2017 ⁴⁴	Shaun A Mason, 2016	-	Т2DM			14 (7/7))
71 W 715HO1, 2017	Bhatt JK, 2012		T2DM			59	,
	N Bishop, 1985 (A)		T2DM			25	
	N Bishop, 1985 (B)		T2DM			25	
	F Klein. 1995		T1DM			24 (12/	12)
	Joíza L Camargo, 2006		Healthy		4	14 (7/7)	
HOMA-IR	Joiza L Camargo, 2000		rically			14 (1/1)	,
Vitamin B-9							
Omid Asbaghi, 2021 ³⁹	Kilicdag, 2005	j	Polycystic ovarian sy	ndrome patients		31(17/1	4)
0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Sheu, 2005		Obese women	F		74(36/3	
	Solini, 2006	(Overweight subjects			60(30/3	
	Cagnacci, 2009		Postmenopausal			30(15/1	
	Palomba, 2010		Polycystic ovary sync	rome		47(23/2	(4)
SR Author and year	Male / female	Intervention		Componenton	Study design	Catting	Quality
SK Author and year	Maie / Temaie	Dose	Duration	Comparator	Study design	Setting	assessment
HbA1C							
Vitamin C				<i></i>			
AW Ashor, 2017 ⁴⁴	12/2	1000mg/day	y 120days	placebo	crossover; DB	Australia	Jadad, 5
	17/42	500mg/day	90days	NR	parallel	NR	Jadad, 2
	11/14	500mg/day		placebo	crossover; DB	UK	Jadad, 3
	13/12	500mg/day		placebo	crossover; DB	UK	Jadad, 3
	24M	6000mg/day		placebo	parallel; DB	Denmark	Jadad, 3
	5/9	1000mg/day	y 120days	No intervention	parallel	Brazil	Jadad, 5
HOMA-IR			/				
Vitamin B-9							
Omid Asbaghi, 2021 39	31F	0.348mg/d	12weeks	No intervention	parallel	Turkey	Jadad, 3
	74F	5mg/d	12weeks	PC	parallel; DB	Taiwan	Jadad, 3
	19/41	2.5mg/d	12weeks	PC	parallel	Italy	Jadad, 4
	30F	15mg/d	3weeks	PC	parallel; DB	Italy	Jadad, 2
	47F	0.4mg/d	25weeks	PC	parallel; DB;	Italy	
					non-random		

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's auth	or and year Population	on			Total, n	, , ,
(SR Author and year (ref))						(intervention	/comparison)
HOMA-IR							
Vitamin B-9						10(01/01)	
Omid Asbaghi, 2021 ³⁹	Gargari, 2011		ght and obese me	en with type 2 diabetes	48(24/24)		
	Aghamohammadi Kl					68(34/34)	
	Asemi, 2014 (A)			polycystic ovary syndro		81(27/14)	
	Asemi, 2014 (B)			polycystic ovary syndro	ome	81(27/13)	
	Asemi, 2016		intraepithelial ne	eoplasia grade 1		58(29/29)	
	Talari, 2016		c syndrome			60(30/30)	
	Bahmani, 2018		rial hyperplasia			60(30/30)	
Maryam Akbari, 2018 ¹⁶	Gargari BP, 2011	Overweig	ght and obese me	en with type 2 diabetes		48(24/24)	
	Asemi Z, 2014	Women v	with polycystic o	ovary syndrome	Y	54(27/27)	
	Talari HR, 2016	Patients v	with metabolic sy	yndrome //		60(30/30)	
	Khiavi A, 2011	T2DM				64(34/34)	
	Setola E, 2004	Patients v	with metabolic sy	yndrome		50(25/25)	
	Solini A, 2006	Overweig	ght subjects			60(30/30)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
HOMA-IR							
Vitamin B-9							
Omid Asbaghi, 2021 39	48M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	68M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	81F	1mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	81F	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	58F	5mg/d	25weeks	PC	parallel; DB	Iran	Cochrane, good
	26/34	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, good
	60F	5mg/d	12weeks	PC		Iran	Cochrane, good
Maryam Akbari, 2018 16	NR	5mg/d	8weeks	PC		Iran	Cochrane, poor
•	NR	5mg/d	8weeks	PC		Iran	Cochrane, poor
	NR	5mg/d	12weeks	PC		Iran	Cochrane, poor
	NR	5mg/d	8weeks	PC		Iran	Cochrane, fair
	NR	Folate plus vitamins		PC		Italy	Cochrane, fair
		B6 or B12, 5mg/d	32.2-2-2		r,	3	,
	NR	2.5mg/d	12weeks	PC	NR	Italy	Cochrane, poor

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species	Primary study's author	r and year	Population	1				Total, n	,
(SR Author and year (ref))								(intervention	/comparison)
HOMA-IR Vitamin B-9									
Maryam Akbari, 2018 16	Sheu WH-H, 2005		Obese wor	men				74(36/38)	
•	Dehkordi EH, 2016		Overweigh	nt and obese chi	ldren and adolescents			39(20/19)	
	Kilicdag EB, 2005			ith polycystic o				31(14/17)	
Zhao JV, 2018 ⁴⁰	Talari, 2016				rweight and stable CHD			60(30/30)	
	Asemi, 2016			ntraepithelial ne				58(29/29)	
	Asemi, 2014		Overweigh	nt or obesity, an	d PCOS			54(27/27)	
	Gargari, 2011		With type	2 diabetes at ba	seline; Overweight			48(24/24)	
	Kurt, 2010		Vitamin B	12 deficiency	_			44(24/20)	
	Solini, 2006		NO					60(30/30)	
	Setola, 2004		With meta	bolic syndrome	and hyperinsulinemia			50(25/25)	
	Cagnacci, 2015		NO					30(15/15)	
	Kilicdag, 2005		PCOS					40(20/20)	
SR Author and year	Male / female	Interventi	ion		Comparator	Study design	Setting		Quality
		Dose		Duration					assessment
HOMA-IR									
Vitamin B-9									
Maryam Akbari, 2018 ¹⁶	NR	5mg/d		12weeks	PC	parallel; DB	Taiwan		Cochrane, fair
	NR	5mg/d	A	8weeks	PC	parallel; DB	Iran		Cochrane, fair
	NR	2.5mg/d		12weeks	PC	NR	Turkey		Cochrane, poor
Zhao JV, 2018 ⁴⁰	both	5mg/d		12weeks	placebo	parallel	Iran		Cochrane, good
	58F	5mg/d		6months	placebo	parallel	Iran		Cochrane, fair
	54F	1mg/d		8weeks	placebo	parallel	Iran		Cochrane, fair
	48M	5mg/d		8weeks	placebo	parallel	Iran		Cochrane, fair
	both	5mg/d		8weeks	placebo	parallel	Turkey		Cochrane, fair
	both	2.5mg/d		12weeks	placebo	parallel	Italy		Cochrane, poor
	both	5mg/day		2months	placebo	parallel	Italy		Cochrane, fair
	30F	15mg/d		3weeks	placebo	parallel	Italy		Cochrane, good
	40F	0.35mg/d	ay	3months	placebo	parallel	Turkey		Cochrane, good

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's a	author and year Pop	ılation			Total, n	on/comparison)
HOMA-IR						(Interventi	on/companson/
Vitamin C							
Mehrnoosh Khodaeian, 2015 ⁷⁸	Chen, 2006	T2D	M			NR	
	Evans, 2003	T2D	M			NR	
	Paolisso, 1995	T2D				NR	
Asma Kazemi, 2022 43	Ramzy Ragheb, 2					20/13	
	Sanguanwong, 20				4	50	
	Froghi, 2018	T2D				21/21	
	Chen, 2006	T2D				15/17	
Shaun A. Mason, 2022 42	Ragheb, 2020	T2D				33(20/13)	
5114411 11 11145011, 2022	El-Aal, 2018	T2D				40(10/10/	
	Foroghi, 2018	T2D				78(38/40)	/
	Sanguanwong, 20			· · · · · · · · · · · · · · · · · · ·		(50/50)	
	Hui Chen, 2006	T2D				32(15/17)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
HOMA-IR							
Vitamin C							
Mehrnoosh Khodaeian, 2015 ⁷⁸	both	800 mg/d	4weeks	Placebo	DB	USA	Jadad, 4 good
	both	1000 mg/d VC + 0.2	6weeks	Placebo + 0.2 IU/kg	DB	UK	Jadad, 1 poor
		IU/kg insulin Lispro		insulin Lispro			, I
	both	1000 mg /d	16w/4 w wash out	Placebo	DB	Italy	Jadad, 3 good
Asma Kazemi, 2022 43	NR	VC, 500 mg/d	8weeks	placebo	Parallel	Egypt	Cochrane
,	NR	VC, 1000 mg/d	8weeks	Placebo	Parallel	Thailand	Cochrane, fair
	NR	VC, 500 mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	NR	VC, 800 mg/d	4weeks	Placebo	Parallel	USA	Cochrane, poor
Shaun A. Mason, 2022 42	10/23	500mg/day	56days	only received anti-	parallel	Egypt	Cochrane, poor
			•	diabetes treatment	•		
	40M	1000mg/day	90days	placebo	parallel	Palestine	Cochrane, poor
	41/37	500mg/day	60days	placebo	parallel; DB	Iran	Cochrane, fair
	NS	1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
	13/19	800mg/day	28days	placebo:500 mg citric	parallel; DB	US	Cochrane, fair
			•	acid/25 ml			

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Pop	ulation			Total, n (interven	tion/comparison)
HOMA-IR							· .
Vitamin C							
Yoonhye Kim, 2022 ²⁵	Ali Abd El-Aal, 2018	T2E				(10/10)	
	Hui Chen, 2006	T2D				(15/17)	
	Sanguanwong, 2016	T2D)M			(50/50)	
Fasting insulin							
Vitamin B-7							
Yujia Zhang, 2022 ³⁷	Cristina, 2006	T2N	MD .			18 (10/8)	ı
	Cesar, 2007	T2N	/ID			348 (226)	/122)
	Armida, 2004	T2N	/ID			15 (10/5)	
	Gregory, 2006	T2N	1D		1	36 (20/16	5)
Vitamin B-9	2 2,					`	
Omid Asbaghi, 2021 39	Gargari, 2011	Ove	rweight and obese me	n with type 2 diabetes		48 (24/24	1)
8 /	Cagnacci, 2015		tmenopausal	5.		30 (15/15	
	Sheu, 2005		se women			74 (36/38	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
·		Dose	Duration		, ,	· ·	assessment
HOMA-IR							
Vitamin C							
Yoonhye Kim, 2022 25	NR	1000mg/day	12weeks	PC	parallel	NR	Cochrane, fair
	170	000 (1		D.C.	11 1 55	TTG 4	(unclear)
	NR	800mg/day	4weeks	PC	parallel; DB	USA	Cochrane, poor
	NR	1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
Fasting insulin							
Vitamin B-7			•••	D.C.			
Yujia Zhang, 2022 ³⁷	11/7	15mg/day	28days	PC	parallel	Mexico	Cochrane, good
	140/208	2mg/day	90days	PC	parallel	United States	Cochrane, good
	NR	6.14µmol/d	28days	PC	parallel	Mexico	Cochrane, good
	NR	2mg/day	4weeks	PC	parallel	USA	Cochrane, fair
Vitamin B-9							
Omid Asbaghi, 2021 39	48M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	30F	15mg/d	3weeks	PC	parallel; DB	Italy	Cochrane, good
	74F	5mg/d	12weeks	PC	parallel; DB	Taiwan	Cochrane, good

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's auth	nor and year	Population			Total, n	n ention/comparison)
Fasting insulin						(Interve	ention/companson)
Vitamin B-9							
Omid Asbaghi, 2021 39	Villa, 2005		Postmenopausal			20 (10/	10)
	Solini, 2006		Overweight subjects			60 (30/	*
	Palomba, 2010		Polycystic ovary syndrome	e		47 (23/	,
	Aghamohammadi K		T2DM			68 (34/	,
	Asemi, 2014 (A)		Overweight women with p	olycystic ovary syndro	me	81 (27/	,
	Asemi, 2014 (B)		Overweight women with p			81 (27/	,
	Asemi, 2016		Cervical intraepithelial neo	, , , , , , , , , , , , , , , , , , ,		58 (29//	,
	Talari, 2016		Metabolic syndrome	. L		60 (30/	,
	Bahmani, 2018		Endometrial hyperplasia		\mathbf{M}	60 (30/	,
Zhao JV, 2018 ⁴⁰	Talari, 2016		With type 2 diabetes at bas	seline. Overweight and	d stable CHD	60 (30/	,
	Asemi, 2016		Cervical intraepithelial neo		a stable CIID	58 (29/2	
	Asemi, 2014		Overweight or obesity, and			54 (27/2	
	Gargari, 2011		With type 2 diabetes at bas			48 (24/2	•
	Gurguri, 2011		With type 2 diabetes at bas	Scrincs Over weight		40 (247)	2-1)
SR Author and year	Male / female	Intervention	1	Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
Fasting insulin							
Vitamin B-9							
Omid Asbaghi, 2021 39	20F	7.5mg/d	8weeks	PC	parallel	Italy	Cochrane, Fair
	19/41	2.5mg/d	12weeks	PC	parallel	Italy	Cochrane, Fair
	47F	0.4mg/d	25weeks	PC	parallel; DB;	Italy	Cochrane, good
					non-random		
	68M	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	81F	1mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	81F	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, good
	58F	5mg/d	25weeks	PC	parallel; DB	Iran	Cochrane, good
	60F	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, good
	40M	1000mg/da	y 90days	placebo	parallel	Palestine	Cochrane, poor
Zhao JV, 2018 ⁴⁰	both	5mg/d	12weeks	placebo	parallel	Iran	Cochrane, good
	58F	5mg/d	6months	placebo	parallel	Iran	Cochrane, fair
	54F	1mg/d	8weeks	placebo	parallel	Iran	Cochrane, fair
	48M	5mg/d	8weeks	placebo	parallel	Iran	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's autho	r and year Populat	and year Population				n vention/comparison)
Fasting insulin							
Vitamin B-9	a 11 1 2006	210					V(2.0)
Zhao JV, 2018 ⁴⁰	Solini, 2006	NO				60 (30	
	Villa, 2005	NO				20 (10	
	Setola, 2004		etabolic syndrome	and hyperinsulinemia		50 (25	
		Cagnacci, 2015 NO				30 (15	· ·
Maryam Akbari, 2018 ¹⁶	Gargari BP, 2011			n with type 2 diabetes		48 (24	
	Asemi Z, 2014		with polycystic o			54 (27	
	Talari HR, 2016		with metabolic sy	ndrome		60 (30	
	Khiavi A, 2011	T2DM				64 (34	· ·
	Setola E, 2004		with metabolic sy	ndrome		50 (25	
	Solini A, 2006		ight subjects			60 (30	· ·
	Sheu WH-H, 2005	Obese v				74 (36	,
	Dehkordi EH, 2016	Overwe	ight and obesity			39 (20	0/19)
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
Fasting insulin							
Vitamin B-9							
Zhao JV, 2018 ⁴⁰	both	2.5mg/d	12weeks	placebo	parallel	Italy	Cochrane, poor
	20F	7.5mg/d	8weeks	placebo	parallel	Italy	Cochrane, poor
	both	5mg/day	2months	placebo	parallel	Italy	Cochrane, fair
	30F	15mg/d	3weeks	placebo	parallel	Italy	Cochrane, good
Maryam Akbari, 2018 ¹⁶	NR	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, fair
	NR	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, poor
	NR	5mg/d	12weeks	PC	parallel; DB	Iran	Cochrane, poor
	NR	5mg/d	8weeks	PC	NR	Iran	Cochrane, fair
	NR	Folate + vitamins	8weeks	PC	parallel; DB	Italy	Cochrane, fair
		B6 or B12, 5mg/d			-		
	NR	2.5mg/d	12weeks	PC	NR	Italy	Cochrane, poor
	NR	5mg/d	12weeks	PC	parallel; DB	Taiwan	Cochrane, fair
	NR	5mg/d	8weeks	PC	parallel; DB	Iran	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author	and year Population	1			Tota (inte	ıl, n ervention/comparison)
Fasting insulin						_/_	
Vitamin C							
Asma Kazemi, 2022 43	Mason, 2018	T2DM					27/ 27
	El-Aal, 2018	T2DM				10,	
	Ramzy Ragheb, 2020	T2DM				20/1	3
	Sanguanwong, 2016	T2DM				50	
	Froghi, 2018	T2DM				21/2	
	Chen, 2006	T2DM				15/1	
	Ghaffari, 2015	T2DM				(17/	14)
	Paolisso, 1995	T2DM				40	
Shaun A. Mason, 2022 42	Paolisso, 1995	T2DM				(40/	40)
	Mason, 2016	T2DM				(7/7)
	Mason, 2019	T2DM				(27/	27)
	Ragheb, 2020	T2DM					20/13)
	El-Aal, 2018	T2DM				40 (10/10/10/10)
	Foroghi, 2018	T2DM	()			78 (38/40)
SR Author and year	Male / female	Intervention	D.	Comparator	Study design	Setting	Quality
		Dose	Duration				assessment
Fasting insulin Vitamin C							
Asma Kazemi, 2022 43	NR	VC, 1000mg/d	17weeks	Placebo	Parallel	Australia	Cochrane, good
,	20 Male	VC, 800mg/d	12weeks	Placebo	Parallel	Palestine	Cochrane, poor
	NR	VC, 500mg/d	8weeks	placebo	Parallel	Egypt	Cochrane
	NR	VC, 1000mg/d	8weeks	Placebo	Parallel	Thailand	Cochrane, fair
	NR	VC, 500mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	NR	VC, 800mg/d	4weeks	Placebo	Parallel	USA	Cochrane, poor
	NR	VC, 800mg/d	8weeks	Placebo	Parallel	Iran	Cochrane, poor
	both	1000 mg/d	12 weeks	Placebo	Cross-over	Italy	Cochrane, poor
Shaun A. Mason, 2022 42	19/21	1000mg/day	120days	placebo	crossover; DB	Italy	Cochrane, fair
, -	12/1	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, fair
	26/5	1000mg/day	120days	placebo	crossover; DB	Australia	Cochrane, good
	10/23	500mg/day	56days	PC	parallel	Egypt	Cochrane, poor
	40M	1000mg/day	90days	placebo	parallel	Palestine	Cochrane, poor
	41/37	500mg/day	60days	placebo	parallel; DB	Iran	Cochrane, fair

Supplementary Table 2. Characteristics of included randomized controlled trials of meta-analysis exploring the effects of water-soluble vitamin on glycemic control and insulin resistance (cont.)

Variables and vitamin species (SR Author and year (ref))	Primary study's author a	and year Population	ı			Total, n (intervention	n/comparison)
Fasting insulin Vitamin C							
Shaun A. Mason, 2022 ⁴²	Sanguanwong, 2016 Hui Chen, 2006	T2DM T2DM				(50/50) 32 (15/17)	
	Ghaffari, 2015	T2DM				31 (17/14)	
AW Ashor, 2017 44	Hui Chen, 2006	T2DM				32 (17/15)	
,	L Pirbudak, 2004	Healthy				22 (11/11)	
	Johannes Pleiner, 2002	Healthy			4	10	
	Simona Bo, 2007	Healthy				78 (40/38)	
	Shaun A Mason, 2016	T2DM				14 (7/7)	
	Gaffari, 2015	T2DM				31	
	C S Johnston, 1994	Healthy				9	
	Brian A Mullan, 2005	Healthy				9	
	David C Nieman, 2002	Healthy				(15/13)	
SR Author and year	Male / female	Intervention		Comparator	Study design	Setting	Quality
		Dose	Duration	<u>"</u>			assessment
Fasting insulin							
Vitamin C							
Shaun A. Mason, 2022 42	NS	1000mg/day	60days	placebo	parallel; DB	Thailand	Cochrane, fair
	13/19	800mg/day	28days	placebo:	parallel; DB	US	Cochrane, fair
	13/18	800mg/day	60days	placebo	parallel	Iran	Cochrane, fair
AW Ashor, 2017 44	13/19	800mg/day	28days	placebo	parallel; DB	USA	Jadad, 5
	22F	AA 500 mg,	1days	fentanyl 1–2 mg/kg and	parallel	Turkey	Jadad, 2
		fentanyl 1–2 mg/kg		etomidate 0.3–0.4			
		and etomidate 0.3		mg/kg			
	102.5	0.4 mg/kg					
	10M	72mg/day	1days	placebo	crossover; DB	Australia	Jadad, 3
	24/54	2000mg/day	14days	no intervention	parallel	Italy	Jadad, 3
	12/2	1000mg/day	120days	placebo	crossover; DB	Australia	Jadad, 5
	13/17	800mg/ day	56days	placebo	parallel	Iran	Jadad, 2
	2/7	1000mg/day	14days	placebo	crossover; DB	USA	Jadad, 5
	9M	2000mg/day	1days	placebo	crossover; DB	UK	Jadad, 3
	NR	1500mg/day	1days	placebo	parallel; DB	USA	Jadad, 4

Supplementary Table 3. Results of assess quality of evidence in meta-analysis

SR author and year (ref)	Vitamin species	Outcomes	Risk of bias	Inconsistency
Arti Muley, 2022 48	Thiamine	FBG	No serious limitations	Serious limitations b
		HbAc1	No serious limitations	No serious limitations
Yi Ding, 2014 46	Niacin	FBG	Serious limitations a1	Serious limitations b
Dan Xiang, 2020 47	Niacin	BG	No serious limitations	No serious limitations
		HbAc1	No serious limitations	Serious limitations b
Yujia Zhang, 2022 37	Biotin	FBG	No serious limitations	No serious limitations
-		HbAc1	No serious limitations	No serious limitations
		insulin	No serious limitations	No serious limitations
Omid Asbaghi, 2021 ³⁹	Folic acid	FBG	No serious limitations	Very serious limitations b
		HbAc1	No serious limitations	Serious limitations b
		HOMA-IR	No serious limitations	Very serious limitations b
		insulin	No serious limitations	Serious limitations b
Maryam Akbari, 2018 ¹⁶	Folic acid	FBG	No serious limitations	Serious limitations b
		HbAc1	No serious limitations	No serious limitations
		HOMA-IR	No serious limitations	Serious limitations b
		insulin	No serious limitations	Serious limitations b
SR author and year (ref)	Indirectness	Imprecision	Publication bias	Quality
Arti Muley, 2022 48	No serious limitations	No serious limitations	Serious limitations e1	Low
	No serious limitations	No serious limitations	Serious limitations e1	Moderate
Yi Ding, 2014 46	No serious limitations	No serious limitations	Serious limitations e1	Very low
Dan Xiang, 2020 47	No serious limitations	No serious limitations	No serious limitations	High
	No serious limitations	serious limitations ^{d3}	No serious limitations	Low
Yujia Zhang, 2022 ³⁷	No serious limitations	No serious limitations	No serious limitations	High
	No serious limitations	Serious limitations d2	No serious limitations	Moderate
	No serious limitations	Serious limitations d3	No serious limitations	Moderate
Omid Asbaghi, 2021 ³⁹	Serious limitations c1	No serious limitations	Serious limitations e2	Very low
	Serious limitations c1	Serious limitations d1	No serious limitations	Very low
	Serious limitations c1	No serious limitations	No serious limitations	Low
	Serious limitations c1	No serious limitations	No serious limitations	Low
Maryam Akbari, 2018 ¹⁶	No serious limitations	No serious limitations	No serious limitations	Moderate
	No serious limitations	Serious limitations d1	No serious limitations	Moderate
	No serious limitations	No serious limitations	No serious limitations	Moderate
	No serious limitations	No serious limitations	No serious limitations	Moderate

a1: high risk of bias regarding allocation concealment. a2: Bias risk was low for 17 studies, whereas a high risk of bias was found in five studies. a3: Of 12 trials, only 4 trials had score equal to 4 (high-quality studies) and the others were categorized as low-quality studies. a4: 93.75% of studies were at high risk. a5: 10 studies (77%) were at high risk. a6: 6 studies were at high risk. b: The test for heterogeneity is significant, and the I is moderate, >50%. b2: The Cochrane Q test for heterogeneity indicated that the studies are heterogeneous (p < 0.0001). c1: Studies conducted subject to various conditions. c2: Surrogate outcome measure, not a patient-important end point. d1: Values are distributed within opposite direction across studies. d2: The sample size is small. d3: Upper bound 95% CI of estimate outside of clinical meaningfulness. e1: The risk of publication bias is high. e2: The Egger's test for publication bias. is significant(p=0.039). e3: The Egger's test for publication bias, is significant(p=0.01).

Supplementary Table 3. Results of assess quality of evidence in meta-analysis (cont.)

SR author and year (ref)	Vitamin species	Outcomes	Risk of bias	Inconsistency
Zhao JV, 2018 ⁴⁰	Folic acid	FBG	No serious limitations	Serious limitations b
		HbAc1	No serious limitations	Serious limitations b
		HOMA-IR	No serious limitations	Very serious limitations b
		insulin	No serious limitations	Serious limitations b
Patcharaporn Sudchada, 2012 54	Folic acid	HbAc1	No serious limitations	Serious limitations b
AW Ashor, 2017 44	Vitamin C	FBG	Serious limitations a2	Serious limitations b
		HbAc1	Serious limitations a2	No serious limitations
		insulin	Serious limitations a2	No serious limitations
Shaun A. Mason, 2021 42	Vitamin C	FBG	Serious limitations	Serious limitations b
		HbAc1	Serious limitations	Serious limitations b
		PPG	Serious limitations	Serious limitations b
		HOMA-IR	Serious limitations	Serious limitations b
		insulin	Serious limitations	Serious limitations b
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	Vitamin C	FBG	Serious limitations a3	No serious limitations
		HbAc1	Serious limitations a3	Serious limitations b2
SR author and year (ref)	Indirectness	Imprecision	Publication bias	Quality
Zhao JV, 2018 40	No serious limitations	No serious limitations	Serious limitations e1	Low
	No serious limitations	No serious limitations	Serious limitations e1	Low
	No serious limitations	No serious limitations	No serious limitations	Low
	No serious limitations	No serious limitations	Serious limitations e1	Low
Patcharaporn Sudchada, 2012 54	No serious limitations	No serious limitations	No serious limitations	Moderate
AW Ashor, 2017 44	No serious limitations	No serious limitations	No serious limitations	Low
	No serious limitations	No serious limitations	No serious limitations	Moderate
	No serious limitations	No serious limitations	No serious limitations	Moderate
naun A. Mason, 2021 42	Serious limitations c2	Serious limitations d3	No serious limitations	Very low
	Serious limitations c2	Serious limitations d3	No serious limitations	Very low
	No serious limitations	Serious limitations d3	No serious limitations	Very low
	No serious limitations	Serious limitations d3	No serious limitations	Very low
	No serious limitations	Serious limitations d3	No serious limitations	Very low
Ozra Tabatabaei-Malazy, 2014 ⁴⁵	No serious limitations	No serious limitations	No serious limitations	Moderate
	No serious limitations	No serious limitations	Serious limitations e3	Very low

a1: high risk of bias regarding allocation concealment. a2: Bias risk was low for 17 studies, whereas a high risk of bias was found in five studies. a3: Of 12 trials, only 4 trials had score equal to 4 (high-quality studies) and the others were categorized as low-quality studies. a4: 93.75% of studies were at high risk. a5: 10 studies (77%) were at high risk. a6: 6 studies were at high risk. b: The test for heterogeneity is significant, and the I is moderate, >50%. b2: The Cochrane Q test for heterogeneity indicated that the studies are heterogeneous (p < 0.0001). c1: Studies conducted subject to various conditions. c2: Surrogate outcome measure, not a patient-important end point. d1: Values are distributed within opposite direction across studies. d2: The sample size is small. d3: Upper bound 95% CI of estimate outside of clinical meaningfulness. e1: The risk of publication bias is high. e2: The Egger's test for publication bias. is significant(p=0.039). e3: The Egger's test for publication bias, is significant(p=0.01).

Supplementary Table 3. Results of assess quality of evidence in meta-analysis (cont.)

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SR author and year (ref)	Vitamin species	Outcomes	Risk of bias	Inconsistency
Asma Kazemi, 2022 ⁴³	Vitamin C	FBG	Very serious a4	Very serious
		HbAc1	Serious limitations a5	Serious limitations b
		insulin	Serious limitations a6	Serious limitations b
		HOMA-IR	No serious limitations	Serious limitations b
Mehrnoosh Khodaeian, 2015 ⁷⁸	Vitamin C	HOMA-IR	Serious limitations a1	No serious limitations
Yoonhye Kim, 202 ²⁵	Vitamin C	FBG	No serious limitations	Serious limitations b
		HbAc1	No serious limitations	No serious limitations
		HOMA-IR	No serious limitations	Serious limitations b
SR author and year (ref)	Indirectness	Imprecision	Publication bias	Quality
Asma Kazemi, 2022 ⁴³	No serious limitations	No serious limitations	No serious limitations	Very low
	No serious limitations	No serious limitations	No serious limitations	Low
	No serious limitations	No serious limitations	No serious limitations	Low
	No serious limitations	No serious limitations	No serious limitations	Moderate
Mehrnoosh Khodaeian, 2015 ⁷⁸	No serious limitations	Serious limitations d2	No serious limitations	Low
Yoonhye Kim, 202 ²⁵	No serious limitations	No serious limitations	No serious limitations	Moderate
	No serious limitations	No serious limitations	No serious limitations	High
	No serious limitations	Serious limitations d2	No serious limitations	Low

a1: high risk of bias regarding allocation concealment. a2: Bias risk was low for 17 studies, whereas a high risk of bias was found in five studies. a3: Of 12 trials, only 4 trials had score equal to 4 (high-quality studies) and the others were categorized as low-quality studies. a4: 93.75% of studies were at high risk. a5: 10 studies (77%) were at high risk. a6: 6 studies were at high risk. b: The test for heterogeneity is significant, and the I is moderate, >50%. b2: The Cochrane Q test for heterogeneity indicated that the studies are heterogeneous (p < 0.0001). c1: Studies conducted subject to various conditions. c2: Surrogate outcome measure, not a patient-important end point. d1: Values are distributed within opposite direction across studies. d2: The sample size is small. d3: Upper bound 95% CI of estimate outside of clinical meaningfulness. e1: The risk of publication bias is high. e2: The Egger's test for publication bias. is significant(p=0.039). e3: The Egger's test for publication bias, is significant(p=0.01).