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Citation: Vallianou, Natalia, Tsang, Catherine, Taghizadeh, Mohsen, Davoodvandi, Amirhossein and Jafarnejad, Sadegh (2019) Effect of cinnamon (Cinnamomum Zeylanicum) supplementation on serum C-reactive protein concentrations: A meta-analysis and systematic review. *Complementary Therapies in Medicine*, 42. pp. 271-278. ISSN 0965-2299

Published by: Elsevier

URL: <https://doi.org/10.1016/j.ctim.2018.12.005> <<https://doi.org/10.1016/j.ctim.2018.12.005>>

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1 **Effect of cinnamon (*Cinnamomum Zeylanicum*) supplementation on**
2 **serum C-reactive protein concentrations: A meta-analysis and**
3 **systematic review.**

4

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18 ***Running title: cinnamon on serum C-reactive protein concentrations***

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23 **Abstract**

24 **Objective:** The effect of cinnamon (*Cinnamomum Zeylanicum*) on serum
25 C-reactive protein (CRP), an acute phase protein commonly used as a
26 marker of inflammation, is uncertain. Therefore, the objective of the
27 present study was to conduct a systematic review and meta-analysis of
28 published randomised controlled trials (RCTs) of cinnamon to determine
29 the effect on levels of serum CRP, relative to controls.

30 **Design:** Studies were identified by a search of electronic databases
31 including PubMed, Cochrane Library, Google Scholar and Scopus before
32 August 2018. Combined and stratified analyses were used. Weighted
33 mean differences (WMD) and its 95% confidence interval were estimated
34 for net change in serum CRP by using random-effects model. The
35 heterogeneity of meta-analysis was assessed by χ^2 and I^2 test.

36 **Results:** Six studies were identified, and data from 285 participants were
37 included. Pooled analysis showed significant reductions in serum CRP
38 (WMD: -0.81 mg/L, 95% CI: -1.36 to -0.26 , $p=0.004$), with significant
39 heterogeneity between selected studies. Improvements in sub-group
40 analysis were observed when baseline CRP levels were greater than 3
41 mg/dL, and in trials of >12 weeks duration. Doses <1500 mg/day and
42 ≥ 1500 mg/day were effective in lowering serum CRP (WMD: -0.56
43 mg/dL, 95% CI: -1.01 to -0.10 , $p=0.02$ and WMD: -2.13 mg/dL, 95%

44 CI: -4.08 to -0.19, $p=0.03$), respectively, with significantly reduced
45 heterogeneity in trials with lower doses of cinnamon <1500 mg/day (test
46 for heterogeneity: $P=0.22$ and $I^2= 33\%$). No changes were found in
47 controls.

48 **Conclusion:** Cinnamon supplementation improves levels of serum CRP,
49 particularly in chronic conditions where basal CRP levels are raised.

50 **Key-words:** Anti-inflammatory; Cinnamon; CRP; meta-analysis; RCT.

51

52

53 **Introduction.**

54 Cinnamon (*Cinnamomum Zeylanicum*) belongs to the genus
55 *Cinnamomum* of the Lauraceae family, derived from the Hebraic and
56 Arabic term amomon, meaning fragrant spice plant. Comprising over 300
57 species, it is widely used for its culinary and medicinal properties with
58 Ceylon and Cassia cinnamon being the most abundant in the U.S and EU
59 markets¹⁻³. Cinnamon has attracted much attention due to their putative
60 health-related properties, which have been ascribed in part to their
61 polyphenolic content; a diverse group of secondary plant metabolites
62 classified as phenolic acids, flavonoids, stilbenes and lignans⁴. Evidence
63 from experimental studies have shown anti-inflammatory and
64 antioxidative properties, particularly in their ability to reduce reactive
65 oxygen species (ROS), and improve insulin sensitivity and carbohydrate
66 metabolism⁵⁻⁸. Clinical studies also indicate improvement in
67 anthropometric parameters, inflammatory mediators, glycemic indices
68 and lipid profiles in patients with type-2 diabetes mellitus (T2DM),
69 nonalcoholic fatty liver disease and rheumatoid arthritis, and those with a
70 BMI ≥ 27 kg/m², following cinnamon supplementation⁹.

71 C-reactive protein (CRP) is an acute phase protein commonly used as a
72 marker of inflammation, and is associated with early stages of several
73 chronic conditions including coronary artery disease (CAD), T2DM,

74 rheumatoid arthritis, pre-diabetes, obesity and nonalcoholic fatty liver
75 disease ¹⁰⁻¹². This increases greatly in inflammation processes and shows
76 specific responses in medical conditions such as polycythemia, anemia,
77 and congestive heart failure with no significant changes. However,
78 compared to conventional assessments of inflammation factors such as
79 erythrocyte sedimentation rate (ESR) test, CRP assessment is an ideal
80 indicator in inflammations ¹⁰⁻¹³. Effects of cinnamon supplementation on
81 serum CRP level have been investigated in clinical trial studies. However,
82 evidence from RCTs are limited and remain inconclusive. Therefore, the
83 aim of the present study was to conduct a systematic review and meta-
84 analysis to assess the efficacy of cinnamon supplementation on serum
85 CRP in several chronic inflammatory conditions.

86 **Methods and Materials.**

87 The present meta-analysis was conducted in accordance with PRISMA
88 (Preferred Reporting Items for Systematic reviews and Meta-Analysis)
89 requirements for interventional research ¹⁴.

90

91 *Search Strategy*

92 Four databases, including PubmedTM, Cochrane LibraryTM, Google
93 ScholarTM and ScopusTM were used to identify related publications.

94 Published RCTs were searched from inception to August 2018. Reference
95 lists from retrieved studies were also manually searched for additional
96 relevant publications. The following searches in titles, abstracts and
97 keywords: “CRP or C reactive protein” in combination with “cinnamon”
98 was performed. Studies were included if they followed a RCT study
99 design with cinnamon supplementation as the intervention. Those
100 published in English and/or Persian were included in the study.

101 *Inclusion and exclusion criteria*

102 The inclusion criteria for selected studies were based on the following;
103 RCTs of oral cinnamon supplementation, those with a duration of more
104 than one week and those reporting mean or median values of serum CRP
105 levels at baseline and by the end of supplementation in control and
106 intervention groups with SD, SEM or 95% CI. The exclusion criteria
107 included duplicated studies, those with no control or placebo group, those
108 with insufficient data at baseline and/or final levels of serum CRP in
109 control and treatment groups, studies with case-control, cohort or cross-
110 sectional design, in vitro and animal studies.

111 *Data extraction*

112 Data were extracted from published studies independently by three
113 reviewers, and any disagreements were resolved by consensus among the

114 researchers using the standardised extraction forms to guarantee accuracy
115 and consistency. The following key data were extracted: year of
116 publication, country where the intervention was conducted, sample size of
117 both intervention and control groups, clinical condition of subjects,
118 intervention/placebo details and composition including the dosage of
119 cinnamon supplementation (gram or mg per day), treatment duration and
120 significant outcomes. In addition, serum levels of CRP were reported as
121 mg/dL. For papers containing data in mmol/l, a numerical conversion to
122 mg/dL was carried out based on molecular weight. Corresponding authors
123 of trials with no reported mean and SD values for any outcomes of
124 interest were contacted to request their data. Only the studies providing
125 these data were included in the present meta-analysis¹⁵.

126 *Quality assessment*

127 We performed a systematic assessment of bias in the included study by
128 using the Cochrane criteria¹⁶. The items used for each included study
129 assessment were the following ones: adequacy of sequence generation,
130 the allocation concealment, blinding of participants, personnel and
131 outcome assessment, the addressing of drop-outs and incomplete
132 outcome data, selective outcome reporting and other potential sources of
133 bias. According to the recommendations of the Cochrane Handbook, the
134 included studies were rated on each of the items as 'L' indicating a

135 low risk of bias, 'h' indicating a high risk of bias or 'u' when the
136 risk of bias was unclear¹⁶.

137 *Statistical analysis*

138 The statistical analyses were performed using Review Manager Software
139 (RevMan 5.3; Cochrane Collaboration, Oxford, England) and
140 Comprehensive Meta-Analysis (version 3.2; Biostat). The pooled
141 weighted mean difference (WMD) and its 95% confidence interval (CI)
142 were estimated to assess the effects of cinnamon on levels of serum CRP.
143 The mean and standard deviation (SD) of levels of serum CRP at baseline
144 and after supplementation in both intervention and control groups were
145 used. Based on the method of Hozo *et al.* all reported median values with
146 their confidence intervals (CI) or their ranges were converted to mean and
147 SD¹⁷. Existence of heterogeneity and the percentage of total variation
148 between studies was assessed by the Cochran's Q-test at $P < 0.05$ level of
149 significance and I² test (I² < 50%). Based on the results (present
150 significant heterogeneity with $p < 0.05$ from χ^2 test), a random effects
151 model was used if I² > 50% and $P < 0.1$. A fixed effects model was used if
152 I² < 50% and $P > 0.1$. To identify the influence of modulators, pre-defined
153 subgroup analyses were conducted according to the Cochrane guidelines
154 including treatment duration, dose of intervention, measuring serum
155 CRP/hs-CRP and baseline CRP level. Sensitivity analysis was performed

156 to estimate the effects of each trial on the pooled effect size, in which a
157 single trial was omitted each time and the effect size was re-calculated to
158 assess the influence on the overall effect size. In order to examine
159 potential publication bias, the funnel plot test was performed. If
160 publication bias exists, the funnel plot shows an asymmetric shape.
161 Additionally, Begg's rank correlation test and Egger's weighted
162 regression test were used to elucidate possible bias. A P-value <0.05 was
163 considered statistically significant.

164

165 **Results.**

166 *Search results and study selection*

167 A flow chart depicting the process of selection and literature search is
168 presented in Figure 1. The literature search of electronic databases
169 identified 205 potential relevant articles. After removing duplicates
170 (n=112), titles and abstracts were screened and sixty-four studies were
171 excluded, as they were not relevant to our analysis or were not in English
172 language. A further 23 studies were excluded after further evaluation due
173 to molecular or animal experiments (n=11), observational studies (n=2),
174 reviews or editorial papers (n=5), not enough data for characterisation of
175 subjects or insufficient reporting of baseline and/or follow-up serum CRP

176 levels in the cinnamon and/or control group (n=2), and studies with no
177 control group (n=3). Finally, a total of 6 RCTs were included in this
178 meta-analysis.

179 *Description of the studies*

180 All trials were published between 2014 to 2018 and were conducted in
181 France, India, Iran and the USA ¹⁸⁻²³. A total of 285 adult participants
182 were re-analysed in the study, of which 144 were allocated to receive
183 cinnamon supplementation and 141 to a control group. Cinnamon dosage
184 ranged from 1200 mg/day to 3000 mg/day, with a median dose of 1850
185 mg/day ¹⁸⁻²³. Cinnamon capsules, stick and extracts were the formulations
186 used in these trials. Duration of supplementation ranged from 8 weeks to
187 24 weeks with a median duration of 14 weeks ¹⁸⁻²³. Selected studies
188 enrolled patients with non-alcoholic fatty liver disease, T2DM, metabolic
189 syndrome, obesity, pre-diabetes and rheumatoid arthritis ¹⁸⁻²³.

190 Baseline level of serum CRP ranged from 1.69 mg/dL to 5.74 mg/dL with
191 a median level of 3.76 mg/dL in the intervention and 3.75 mg/dL in the
192 control groups, respectively. Five of the 6 studies were conducted in both
193 males and females, with one study conducted only in female participants
194 ¹⁸⁻²³. All included trials followed a parallel study design. Three trials
195 evaluated cinnamon in combination with black tea, L-carnosine plus
196 chromium guanylate and a multiple dietary supplement containing

197 cinnamon powder ^{19, 22, 23} (Table 1). Cinnamon supplementation was
198 apparently safe and well tolerated by participants in all of the included
199 studies, and no adverse effects were reported.

200 *Risk of bias assessment*

201 An unclear risk of bias was observed in some of the items including
202 allocation concealment and other potential sources of bias. However,
203 most of the included studies were characterized by adequate information
204 regarding sequence generation, allocation concealment and blinding of
205 participants and personnel. The incomplete outcome data and selective
206 outcome reporting showed a low risk of bias. Details of the quality of bias
207 assessment are presented in Table 2.

208 *Pooled estimate of the effect of cinnamon supplementation on serum CRP*

209 Significant reductions in the levels of serum CRP were observed
210 following cinnamon supplementation in 3 studies ^{20, 22, 23}. Weighted mean
211 difference (WMD) of studies with random effects model analysis showed
212 a significant improvement in serum CRP (WMD: -0.81 mg/L, 95% CI:
213 -1.36 to -0.26, p=0.004) with a significant heterogeneity between the
214 included trials (test for heterogeneity: $P < 0.0002$ and $I^2 = 79\%$)(Figure
215 2).

216 *Subgroup analyses*

217 Subgroup analysis was performed to determine the potential source of
218 heterogeneity, based on study duration, cinnamon dose, serum CRP
219 and/or high sensitivity CRP (hs-CRP) and baseline CRP following
220 supplementation (Table 3). Results showed that cinnamon
221 supplementation significantly reduced serum CRP levels in participants
222 when the duration of the study was >12 weeks (WMD: -0.42 mg/L, 95%
223 CI: -0.65 to -0.20, p=0.0002). The heterogeneity significantly decreased
224 after subgroup analysis by duration of study (test for heterogeneity: P =
225 0.96 and I²= 0%). Subgroup analysis on studies with cinnamon doses of
226 <1500 mg/day and ≥1500 mg/day also significantly influenced levels of
227 serum CRP (WMD: -0.56 mg/dL, 95% CI: -1.01 to -0.10, p=0.02 and
228 WMD: -2.13 mg/dL, 95% CI: -4.08 to -0.19, p=0.03), respectively.
229 There was significantly reduced heterogeneity in studies with lower doses
230 of cinnamon supplementation (test for heterogeneity: P=0.22 and I²=
231 33%). Results of subgroup analysis based on baseline serum CRP also
232 showed that cinnamon supplementation decreased levels of CRP in those
233 with baseline CRP levels of more than 3 mg/dL (WMD: -0.42 mg/L,
234 95% CI: -0.65 to -0.20, p=0.0002). Moreover, the heterogeneity
235 decreased significantly after subgroup analysis by trials with baseline
236 CRP levels of more than 3 mg/dL.

237 *Sensitivity analysis*

238 Sensitivity analysis was performed to determine the effect of each study
239 on the estimated pooled effect size. Results of omitting each study on the
240 effect size ranged from -0.55 mg/L (95% CI=-0.98, -0.11) to -1.07 mg/L
241 (95% CI=-1.80,-0.35)(Figure 3).

242 *Publication bias*

243 The publication bias of this meta-analysis was assessed by examination of
244 funnel plot. The symmetrical funnel plots suggested that the selection of
245 publication was not a possible source of bias (Figure 4). The absence of
246 publication bias was confirmed by Egger's linear regression (intercept: -
247 3.9; standard error: 3.82; 95% CI: -5.91, 1.94; t= 1.4, df=4; two-tailed
248 p=0.23). Moreover, Begg's rank correlation did not highlight any
249 publication bias (Kendall's Tau with continuity correction:-0.4; z=1.12;
250 two-tailed p=0.25).

251

252 **Discussion**

253 The present meta-analysis included a total of 285 adults presenting with
254 non-alcoholic fatty liver disease, T2DM, metabolic syndrome, obesity,
255 pre-diabetes and rheumatoid arthritis from 6 RCTs. Despite considerable
256 heterogeneity among the studies, our findings indicate improvement in
257 the levels of serum CRP following cinnamon supplementation. To our

258 knowledge, this is the first systematic review that has assessed the effects
259 of cinnamon supplementation on serum CRP.

260 Significant reductions in serum CRP levels by -0.81 mg/dL were
261 observed following cinnamon supplementation with no detectable
262 changes in the control group. These findings were consistent across four
263 of the individual six RCTs assessed in this study^{18, 20-22}. Reductions in the
264 levels of serum CRP, as observed in the present study, are clinically
265 important because levels <1 mg/dL are associated with a lower risk of
266 cardiovascular events, with concentrations > 3 mg/dL exacerbating the
267 risk of coronary heart disease up to 58%^{24, 25}.

268 There was significant heterogeneity between studies in this meta-analysis,
269 and subgroup analysis indicated that cinnamon supplementation could
270 lower the levels of serum CRP when the trial duration was >12 weeks.
271 Evidence from other meta-analyses assessing the anti-inflammatory
272 properties of complex medicinal herbs (cinnamon, ginger and other
273 traditional herbs) have also demonstrated significant improvements in
274 serum CRP levels with study durations exceeding 6 and 10 weeks^{26, 27}.

275 Subgroup analysis on studies with cinnamon doses of <1500 mg/day and
276 ≥ 1500 mg/day found significant reductions in the levels of serum CRP. It
277 therefore seems likely that lower doses are effective and may be better
278 than using larger doses of cinnamon, which have been associated with

279 certain adverse effects including diarrhea and headache ²⁸. However,
280 there were no reported adverse effects observed in the included studies in
281 the present meta-analysis. Similar studies have also failed to report any
282 adverse effect or reaction following cinnamon supplementation. Talaei *et*
283 *al.* reported beneficial effects of 1000 mg/day cinnamon (*Cinnamomum*
284 *zeylanicum*) without side effects ²⁹, and Tjandrawinata *et al.* reported a
285 lower risk of hypoglycemic episodes with no effect on gastrointestinal
286 symptoms [27]

287 Moreover, subgroup analysis based on baseline levels showed that
288 cinnamon improved serum CRP levels in those with a higher baseline
289 value (i.e. > 3 mg/dL), with heterogeneity decreasing significantly after
290 subgroup analysis. This finding is in agreement with another studies in
291 which vitamin E supplementation significantly reduced circulating levels
292 of serum CRP only in those with a baseline value of > 3 mg/dL³⁰.
293 Therefore, the duration of the study and baseline serum CRP levels were
294 considered to be important and potential sources of observed
295 heterogeneity.

296 CRP, and indeed hs-CRP, is one of the most common and frequently used
297 biomarkers for inflammation status with predictive values for several
298 chronic diseases including CVD ³¹⁻³⁹. The anti-inflammatory properties of
299 cinnamon has been reviewed extensively, and several mechanisms of

300 action have been described⁴⁰⁻⁴⁴. In vitro and in vivo studies have reported
301 inhibition of nuclear factor kappa B (NF- κ B) by 2'-
302 hydroxycinnamaldehyde isolated from *C. cassia* bark⁴¹, and tumor
303 necrosis factor- α (TNF- α) with extracts of cinnamon in a
304 lipopolysaccharide (LPS) model⁴². Inhibition of TNF- α genes by
305 cinnamon water extract via modulation of JNK, p38, ERK1/2 activation
306 and I κ -B α degradation have also been demonstrated⁴³. Hong *et al.*
307 reported the inhibition of the expression of TNF- α by polyphenol-rich
308 cinnamon water extract (CWE) fraction containing procyanidins,
309 catechin, epicatechin and ellagic acid⁴⁴. Cinnamon may also
310 downregulate the expression of various NF- κ B-regulated pro-
311 inflammatory adipo-cytokines, (i.e. MCP-1, MCP-4, and eotaxin and
312 interleukins)^{45, 46}, in addition to plasminogen activator inhibitor type-1
313 (PAI-1), through the inhibition of the transcription factor early growth
314 response (Egr)-1 gene product, which has been closely linked with insulin
315 resistance and obesity^{47, 48}.

316 Some limitations of this meta-analysis include not controlling for
317 confounding factors (i.e. dietary intake, physical activity and medications
318 for several chronic conditions), which may have influenced our results.
319 Most of the RCTs included were of a relatively small sample size, and the
320 characteristics of the study population was heterogeneous (i.e. patients

321 with non-alcoholic fatty liver disease, T2DM, metabolic syndrome,
322 obesity, pre-diabetes and rheumatoid arthritis). Despite these limitations,
323 there were several strengths to this study. Firstly, it is to our knowledge,
324 the first time a systematic review and meta-analysis has been performed
325 in the evaluation of cinnamon supplementation on serum CRP levels. A
326 random effects model was used for assessing heterogeneity between
327 studies, and RCTs were assessed using subgroup analysis with performed
328 sensitivity and meta-regression analyses.

329 **Conclusion**

330 In conclusion, the findings from this meta-analysis suggest some
331 improvement in serum CRP levels following cinnamon supplementation,
332 especially in patients with higher baseline CRP levels. However, due to
333 the limited availability of RCTs further investigation is warranted.

334 *Acknowledgements*

335 We are grateful to our colleagues for their patience and their advice on
336 searching the papers.

337 *Funding*

338 There is no funding sources in the writing of the manuscript, data
339 collection, analysis, or interpretation; or the decision to submit it for
340 publication. We have not been paid to write this article by a
341 pharmaceutical company or other agency

342

343 *Authors' contributions*

344 N.V. and S.J. conceived and planned the experiments. S.J. and A.D.
345 carried out the literature search in databases. S.J. and M.T. and A.D.
346 contributed to quality assessment and statistical analysis. C.T. and N.V.
347 contributed to the interpretation of the results. C.T. took the lead in
348 writing the manuscript. All authors provided critical feedback and helped
349 shape the research, analysis and manuscript.

350 *Conflict of interest*

351 There is no conflict of interest regarding this manuscript.

352

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501 **Figure 1:** *Meta-analysis Flow Diagram*

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503 **Figure 2:** *Forest plots showing the pooled effect size of cinnamon supplementation on serum*
504 *C-reactive protein (mg/L). Random effects model was used to pool the mean change of*
505 *indicators. CI, confidence interval; I-squared inconsistency.*

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507 **Figure 3:** *Sensitivity analysis for the effect of cinnamon supplementation on serum CRP*
508 *levels.*

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510 **Figure 4:** *Funnel plot of included studies measured serum CRP level. MD = Mean*
511 *Difference, SE = standard error.*

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520 **Table 1:** General characteristics of the included studies in the meta-analysis

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Author	Year	Country	Design	No. of Subjects in case group	No. of controls	Gender	Age(mean)-case group	Age(mean)-control	Inclusion criteria	Clinical Condition of Subjects	Follow-up Duration (weeks)	Dosage (mg/d)	Co-Supplements or other drugs	Significant Outcome	Baseline CRP level (mg/l)
Askari	2014	Iran	R, DB, PC	23	22	F/M	44.8 ± 8.5	45.4 ± 8.2	Age between 20 and 65 ; ALT not greater than 60 U/L; no alcohol or drug abuse;no chemotherapy	Nonalcoholic fatty liver disease (NAFLD)	12	1500	No co-supplement	FBS, HOMA index, QUICKI index, total cholesterol, triglycerides, ALT, AST, GGT and hs-CRP changed significantly in the treatment group.	5mg/l
Azimi	2015	Iran	R, SB, PC	40	39	F/M	54.15± 1.0	53.64 ± 1.3	Subjects with T2D, aged ≥30 years, overweight, not on insulin therapy, not taking medications except for oral hypoglycemic agents.	Type 2 diabetes	8	3000	Three glasses of black tea	Significant reduction in total cholesterol, LDL, and elevation in HDL levels.	5.74mg/l
Jain	2017	India	R, DB, PC	58	58	F/M	44.3 ± 7.2	45.1 ± 8.4	Subjects suffering from other chronic diseases (except for metabolic syndrome) or those on medication of lipid lowering drugs were excluded.	Metabolic syndrome	16	3000	No co-supplement	Significantly decrease in weight, WC, WHR, percentage body fat, total cholesterol, serum triglycerides, LDL-C, SBP, DBP and significant increase in HDL-C.	2.8mg/l
Liu	2015	France	R, DB, PC	26	26	F/M	Not Mentioned	Not Mentioned	Subjects aged between 25 and 65 years, overweight and unwilling to change their usual dietary and activity were included for randomization.	Overweight or Obese Pre-Diabetic	16	456	200 mg/day L-carnosine 2.5 mg/day Chromium guanylate	Insulin secretion, evaluated by HOMA-B%, increased significantly in supplement group.	4mg/l
Shishehbor	2018	Iran	R, DB, PC	18	18	F	44.66 ± 11.22	49.11 ± 7.45	Having active Rheumatoid Arthritis, being under treatment with DMARDs, not receiving NSAIDs or cytokine inhibitors.	Rheumatoid Arthritis	8	2000	No co-supplement	There was a significant decrease of serum levels of CRP and TNF-a in the cinnamon group. Diastolic blood pressure was also significantly lower in the intervention group.	3.53mg/l
Soare	2014	USA	R, SB, PC	28	26	F/M	47±5	44±6	Participants were free of chronic disease. Exclusion criteria included chronic use of medications or dietary supplements, tobacco use, alcohol abuse, and habitual vigorous exercise.	Healthy adults	24	1700	100 mg of resveratrol, 800 mg of green, black, and white tea, 250 mg of pomegranate, 650 mg of quercetin, 500 mg of l carnitine, 600 mg of lipoic acid, 900 mg of curcumin, 1 g of sesamin and fish oil.	No significant outcomes.	1.69mg/l

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Table 2. *Quality of bias assessment of the included trials according to the Cochrane guidelines.*

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Studies ,Year	Sequence Generation	Allocation Concealment	Blinding of participants and personnel	Blinding of outcome assessment	Incomplete outcome data	selective outcome reporting	other potential sources of bias
Askari;2014	L	L	L	L	L	L	U
Azimi;2015	L	L	L	H	L	L	L
Jain;2017	L	U	L	H	L	L	H
Liu;2015	L	U	L	H	L	L	L
Shishehbor;2018	L	L	L	L	L	H	U
Soare;2014	L	U	H	L	L	L	L

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L, low risk of bias; H, high risk of bias; U, unclear risk of bias.

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535 **Table 3: Subgroup analysis of cinnamon supplementation on serum CRP level**

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Subgroup	No	WMD (95% CI)	Test for overall effect	Test for heterogeneity	I²(%)
Duration of study, weeks					
≤12 weeks	3	-1.37 [-2.86, 0.12]	p=0.07	p<0.0001	91
>12 weeks	3	-0.42 [-0.65, -0.20]	P = 0.0002	P = 0.96	0
Cinnamon dose, mg/day					
≥1500 mg/d	4	-0.56 [-1.01, -0.10]	P = 0.02	P = 0.007	76
<1500 mg/d	2	-2.13 [-4.08, -0.19]	P = 0.03	P = 0.22	33
CRP/hsCRP					
CRP	2	-0.96 [-2.14, 0.22]	P = 0.11	P = 0.003	89
hsCRP	4	-0.83 [-1.78, 0.11]	P = 0.08	P = 0.002	79
Baseline CRP, mg/L					
<3	2	-1.26 [-2.62, 0.10]	P = 0.07	P < 0.0001	87
≥3	4	-0.42 [-0.65, -0.20]	P = 0.0002	P = 0.77	0

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538 *: Abbreviations: CRP, C-reactive protein; hsCRP, high-sensitivity C-reactive protein; WMD, weighted mean difference; CI, confidence interval.