Effects of zinc, vitamin D, and their cosupplementation on mood, serum cortisol, and brain-derived neurotrophic factor in patients with obesity and mild to moderate depressive symptoms: A phase II, 12wk, 2 × 2 factorial design, double-blind, randomized, placebo-controlled trial Yosaee, S, soltani, S, Esteghamati, A, Motevalian, A, Tehrani-

Doost, M, Clark, C & Jazayeri, S Author post-print (accepted) deposited by Coventry University's Repository

Original citation & hyperlink:

Yosaee, S, soltani, S, Esteghamati, A, Motevalian, A, Tehrani-Doost, M, Clark, C & Jazayeri, S 2020, 'Effects of zinc, vitamin D, and their co-supplementation on mood, serum cortisol, and brain-derived neurotrophic factor in patients with obesity and mild to moderate depressive symptoms: A phase II, 12-wk, 2 × 2 factorial design, double-blind, randomized, placebo-controlled trial', Nutrition, vol. 71, 110601. https://dx.doi.org/10.1016/j.nut.2019.110601

DOI 10.1016/j.nut.2019.110601 ISSN 0899-9007

Publisher: Elsevier

NOTICE: this is the author's version of a work that was accepted for publication in Nutrition. Changes resulting from the publishing process, such as peer review, editing, corrections, structural formatting, and other quality control mechanisms may not be reflected in this document. Changes may have been made to this work since it was submitted for publication. A definitive version was subsequently published in Nutrition, 71, (2019) DOI: 10.1016/j.nut.2019.110601

© 2019, Elsevier. Licensed under the Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International <u>http://creativecommons.org/licenses/by-nc-nd/4.0/10.1016/j.nut.2019.110601</u>

Copyright © and Moral Rights are retained by the author(s) and/ or other copyright owners. A copy can be downloaded for personal non-commercial research or study, without prior permission or charge. This item cannot be reproduced or quoted extensively from without first obtaining permission in writing from the copyright holder(s). The content must not be changed in any way or sold commercially in any format or medium without the formal permission of the copyright holders.

This document is the author's post-print version, incorporating any revisions agreed during the peer-review process. Some differences between the published version and this version may remain and you are advised to consult the published version if you wish to cite from it.

Effects of zinc, vitamin D and their co-supplementation on mood, serum cortisol, and	1
brain-derived neurotrophic factor in patients with obesity and mild to moderate	2
depressive symptoms: A 12-week, 2×2 factorial design, double-blind, randomized,	3
placebo-controlled trial	4
Keywords: depression, obesity, BDNF, cortisol	5
Somaye Yosaee1,2,3, Sepideh Soltani4,5, Alireza Esteghamati6, Abbas Motevalian7, Mehdi	6
Tehrani-Doost8, Cain C. T. Clark9, Shima Jazayeri1,10*	7
	8
1 Department of Nutrition, School of Public Health, Iran University of Medical 7	9
Sciences, Tehran, Iran	10
2Department of Nutrition, School of Health, Larestan University of Medical Sciences,	11
Larestan, Iran	12
3 Department of Nutrition, Emam Reza Teaching Hospital, Larestan University of Medical	13
Sciences, Larestan, Iran	14
4Department of Nutrition, Faculty of Health, Shahid Sadoughi University of Medical	15
Sciences, Yazd, Iran	16
5Yazd Cardiovascular Research Center, Shahid Sadoughi University of Medical Sciences, 16	17
Yazd, Iran	18
6Endocrinology and Metabolism Research Center (EMRC), Vali-Asr Hospital, Tehran	19
University of Medical Sciences, Tehran, Iran.	20
7Department of Epidemiology, School of Public Health, Iran University of Medical Sciences, Tehran, Iran	21 22
8Department of Psychiatry School of Medicine, Roozbeh Psychiatry Hospital Tehran	23

University of Medical Sciences, Tehran, Iran.	24
9Centre for Sport, Exercise and Life Sciences, Coventry University, CV1 5FB, U.K.	25
10Research Center for Prevention of Cardiovascular Disease, Institute of Endocrinology &	26
Metabolism, Iran University of Medical Sciences, Tehran, Iran	27
	28

1. Introduction:

Mood disorders, particularly depression, are one of the most prevalent mental health symptoms 31 in patients with obesity (1-3). Several studies have reported that patients with obesity can suffer 32 with major depression (4-6). Although the direction of the association between depression and 33 obesity has been questioned (3, 7, 8), the co-occurrence of both may have a detrimental 34 synergistic effect on overall health and treatment response (9). A growing evidence base 35 indicates that there are abnormalities in the hypothalamic-pituitary-adrenal axis, particularly in 36 the regulation of cortisol, in clinically depressed patients, which is, in turn, related to obesity 37 (10-12). Moreover, empirical evidence has suggested that Brain-derived neurotrophic factor 38 (BDNF), a homo-dimer protein, has a critical role in the pathophysiology of depression (13). 39

Despite representing the most preponderant treatment for depression, antidepressant 40 medications have been reported to show resistance in depression associated with obesity (14, 41 15), whilst remission is only achieved in one-third of the patients after treatment with 42 antidepressant agents (16). Furthermore, pharmacotherapy is usually costly (16-18). The 43 expense and incongruent effectiveness in patients with obesity highlights the need to 44 investigate alternative preventive and treatment approaches to traditional antidepressant 45 medication. In recent years, it has been shown that nutritional intervention can be considered 46 as an effective alternative or adjunct, preventive or treatment strategy to pharmacotherapy in 47 depression (19). In particular, Zinc and vitamin D have been well linked with the treatment or 48 management of depression (20-23). 49

Zinc deficiency can induce depressive-like behavior, and in this instance, the symptom can be 50 effectively reversed by zinc supplementation (24, 25). Zinc may produce antidepressant-like 51 effects by modulating the functions of the hypothalamus-pituitary-adrenal (HPA) axis and 52 increasing serum BDNF (13, 26-28). Moreover, there is wealth of literature suggesting zinc 53 supplementation is an effective adjunct therapy for major depressive disorders (13, 29, 30). 54 Similar to zinc, inadequate vitamin D intake has been associated with depression (31, 32); 55 whilst it has also been demonstrated that multiple brain regions are associated with depressive 56 disorders, including the prefrontal cortex and hippocampus, and possess specific nuclear 57 receptors for 1,25(OH)D (32, 33). 58

Although there have been significant advances in understanding the potential role of zinc and
vitamin D in depression (13, 23), the literature, particularly in regard to vitamin D, is equivocal
(34-36). Whilst, to date, only one randomized control trial has examined the effects of zinc
61

monotherapy on BDI-II score in obese patients, which resulted in a decreased BDI-II score
(13). In addition, only one study has examined the effects of vitamin D monotherapy on BDIII in obese patients with pre-existing depressive symptoms, where the authors suggested that
supplementation of high doses of vitamin D can ameliorate depression symptoms (23).
However, the results other studies investigating the effect of vitamin D supplementation on
depression are divergent (37-39), and may be mediated by the inconsistent doses and duration
of vitamin D supplementation.

Several meta-analyses have evaluated effect of zinc/vitamin D on depression (40-43), 69 concluding that, although there is evidence supporting their use, available research needs to be 70 confirmed by larger RCT and prospective cohort studies (44-46). Furthermore, the effects of 71 vitamin D or zinc on depression have mostly been explored independently in previous studies 72 (13, 23), and to the author's knowledge, no study has examined the effects of zinc- vitamin D 73 co-supplementation on depression symptoms in obese subjects. Interestingly, there is evidence 74 to suggest that zinc homeostasis and function may be increased following vitamin D 75 supplementation, and that the control of zinc in systemic levels is regulated by vitamin D (47). 76 The current trial hypothesized that there may be additive benefits from combining zinc and 77 vitamin D. Thus, the present study sought to investigate the effects of zinc, vitamin D, and their 78 combination, on depression score, serum BDNF, and cortisol level in obese patients with mild 79 to moderate depression. 80

2. Material& method

2.1. participants

The present study was a 12-week 2×2 factorial design randomized double-blind placebo 83 controlled trial, and was conducted among 140 overweight/obese (BMI> 27 kg/m²) adult 84 subjects aged > 20 years with BDI-II≥10. Obese/overweight subjects were recruited from 85 patients who were referred to the Endocrinology and Metabolism Research Center (EMRC), 86 Vali-Asr, Emam khomeini Hospital in Tehran, Iran between July 2016 and February 2017. The 87 depression status was evaluated by Beck Depression Inventory-II (BDI-II) questionnaire, and 88 those who had a BDI-II score greater than 10 points were considered as eligible for current the 89 trial. We excluded patients who had a history of psychiatric and neurological disorders (such 90 as schizophrenia Parkinson's, Alzheimer's disease, anxiety, suicidal thoughts), coronary artery 91 disease, acute or chronic renal failure, acute or chronic hepatic failure, chronic inflammatory 92 and autoimmune disease, or any known malignancy, had been received antidepressant 93

82

medicines (in the preceding 3 months), or steroid or hormone-therapy. Other non-pathological
94
exclusion criteria included pregnancy, breastfeeding, post-menopause, smoking, professional
95
athlete, uncontrolled thyroid disorder, >3 kg weight change during the last 3 months, use of
96
medications for dyslipidemia or hypertension, hypnotics, sedatives and immunosuppressive,
97
any kind of supplements or following a special diet prescribed by the clinic dietitian. It was
98
explained to each participant that the dose and type of supplement(s) assigned must not be
99
changed during the intervention period, or they would be excluded from the study.

2.2. Ethics statements

This trial was performed according to the principles of the Declaration of Helsinki, and the102study protocol was approved by the institutional ethics committee of Iran University of Medical103Sciences (IR.IUMS.REC 1395.9221324205). The study protocol was carefully explained to all104subjects signing an informed consent form. This trial was registered in the Iranian Web site105(www.irct.ir) for registration of clinical trials (http://www.irct.ir: IRCT201601252394N31- 31-10610-2016). The project was financially supported by Iran University of Medical Sciences.107

2.3. Study design and intervention

Randomized assignment was performed using balanced the block randomization method. 109 Block randomization works by randomizing participants within blocks such that an equal 110 number are assigned to each treatment arm. Block randomization is a commonly used 111 technique in trials with small sample sizes (48). In the present trial, the block size was 8. 112 Participants were initially randomized to 'intervention vs Placebo,' and then assigned to one 113 of the four groups via the balanced block randomization method (in a 1:1:1:1 ratio), (1) subjects 114 received 2000 IU vitamin D3 daily plus a daily placebo for zinc; 2) subjects received 30 mg 115 zinc gluconate per day plus a daily placebo for vitamin D; 3) subjects received 2000 IU vitamin 116 D3 daily plus 30 mg zinc gluconate per day; or 4) subjects received identical matching placebos 117 for vitamin D and zinc for 12 weeks. At the beginning of the study, participants were requested 118 not to change their routine lifestyle throughout the study and not to consume any supplements 119 or medication other than that provided to them by the investigators. The doses of vitamin D3 120 and zinc were chosen after a comprehensive review of the available literature to achieve 121 optimal efficacy and safety (13, 49). The appearance of the placebo was indistinguishable in 122 color, shape, size, packaging, and taste from vitamin D and zinc tablets. The zinc supplements 123 were tablets manufactured by the Jalinus Pharmaceutical Company (Tabriz, Iran) and the 124 placebo (made from starch) was provided by the School of Pharmacy, Tehran University of 125 Medical Sciences. The vitamin D supplements and placebos (made from starch) were tablets 126

101

manufactured by Pars mino Company (Tehran, Iran). All research staff including investigators 127 and laboratory technicians, as well as participants, were blinded to the random allocation. 128

2.4. Treatment adherence

The tablets were supplied to the participants, fortnightly, by the trial staff. Each bottle contained 130 15 tablets, and a tablet count was performed fortnightly by the investigator to assess 131 compliance. To increase the adherence, all participants received short messages on their cell 132 phones to take the supplements every day. 133

2.5. Assessment of variables

All assessments and measurements were made at study baseline and after the intervention 135 period. Weight (in kilograms -kg) was measured using a calibrated Seca scale (Model 700, 136 USA) with participants in light clothing and unshod. Standing height was measured to nearest 137 1 cm using a Seca stadiometer (Model 700, USA) while subjects were unshod. Body mass 138 index (BMI) of each participant was calculated as body weight divided by height squared 139 (kg/m²). Waist circumference was measured using a flexible tape at the smallest circumference 140 around coastal margin. A professional nurse measured the systolic and diastolic blood pressure 141 on the non-dominant brachial artery, with the participants in a seated position, after having 142 rested at least for 10 minutes. Blood pressure was measured twice, and the average of 2 143 measurements was considered as the final systolic and diastolic blood pressure. A trained 144 researcher completed questionnaires on participants socio-demographic background and 145 physical activity, whilst a researcher provided a comprehensive explanation of how to complete 146 the self-rating BDI-II questionnaire. The trial used the validated Persian version of Beck 147 depression inventory-II (BDI-II) (50) 148

149

To assess serum 25(OH) D, zinc, BDNF and cortisol levels, blood samples (10 cc) were 150 obtained after a 12-hour overnight fast at study baseline (week 0) and after intervention (week 151 12). All blood samples were centrifuged at 3000g for 10 minutes, the serum was separated into 152 clean tube aliquots, and were stored at -80°C until analysis at the Iran University of Medical 153 Sciences Reference Laboratory. Serum 25(OH)D concentrations were quantified using the 154 enzyme-linked immunosorbent assay (ELISA) method (Euroimmun,). The inter- and intra-155 assay coefficients of variation of this method were 8.6% and 3.2%. respectively. A serum 156 25(OH) D level <75 nmol/l (<30 ng/ml) was considered insufficient. ELISA methods were 157 used to measure serum cortisol (diametra, Italy). The inter- and intra-assay coefficients of 158

6

variation of this method were 11% and 5.1%. respectively. Serum BDNF was quantified using
the ELISA method (crystalday, china). The inter- and intra-assay coefficients of variation of
this method were 10% and 8% respectively.

2.6. Sample size

To calculate sample size, we used the standard formula suggested for factorial design, clinical 164 trials by considering type I error (a) of 0.05 and type II error (b) of 0.20. Based on a previous 165 study (13), we used 5.698 as the standard deviation and 2.92 as the mean difference in BDI-II 166 score as a variable primary outcome. Based on this, we required 33 persons in each group. 167 Accounting for 2 dropouts (effect size: 0.52) in each group, the final sample size was 168 determined to be 35 persons per group. 169

2.7. Statistical methods

171

187

170

162

163

We evaluated changes in depression symptoms as a primary outcome, during an average of 12 172 weeks of follow-up, in those randomized to zinc-vitamin D and their combined 173 supplementation compared to placebo. The Kolmogorov-Smirnov test was used to examine 174 and confirm the normal distribution of variables. The analyses were performed based on a per-175 protocol approach. One-way analysis of variance (ANOVA) was used to detect differences in 176 general characteristics, blood pressure and anthropometric measurement, at the study baseline 177 and post-intervention between the groups. To estimate the effect of zinc, vitamin D and their 178 combination on outcomes, we first computed the changes from baseline by subtracting the 179 baseline value from the end-of-trial value, and then applied analysis of co-variance (ANCOVA; 180 adjusted by baseline serum zinc, vitamin D, cortisol, BDNF and beck depression score). To 181 ensure that baseline characteristics balance was achieved by the randomization, analyses was 182 conducted for baseline characteristics (zinc, vitamin D, age, sex, BMI, weight, height). In cases 183 of minor imbalances, adjustment for these variables will be made our analyses. For all tests, 184 statistical significance was accepted at p≤0.05. All analyses were performed using the 185 Statistical Package for Social Sciences version 21 (SPSS, Chicago, IL, USA). 186

3.Results:

The present study was conducted using 140 obese patients with mild to moderate depression. 188 In total, 15 patients declined to complete the intervention, and were therefore excluded from 189

subsequent analysis (diagram 1). Dropout rate was not significant between the treatment arms 190 (P=0.785). Serum levels and tablet counts of zinc and vitamin D at the end of the intervention 191 suggested that the compliance was excellent. No significant differences in demographic 192 characteristics were observed among the 4 intervention arms at baseline. Moreover, there was 193 no significant difference in anthropometric measurements, such as weight, BMI, waist 194 circumference (WC) and blood pressure, in zinc, vitamin D and co-supplementation groups 195 compared to placebo (p>0.05). We found a significant decrease in blood pressure among those 196 who received vitamin D (systolic blood pressure: 121.61±13.81 vs 118.00±15.76, P=0.0001. 197 diastolic blood pressure: 81.96±10.00 vs 84.58±9.79, P=0.014) or joint zinc-vitamin D 198 (systolic blood pressure: 119.26±14.07 vs 114.73±14.66, P=0.0001. diastolic blood pressure: 199 80.30±10.80 vs 78.73±11.94, P=0.0001) supplements. A significant decrease in WC was 200 shown in vitamin D (106.91±11.39 vs 105.33±11.69, P=0.0001), zinc (103.63±5.84 vs 201 103.14±6.38, P=0.0001) and combined zinc-vitamin D (105.25±8.95 vs 103.85±9.20, 202 P=0.0001) groups. The baseline and post-intervention characteristics of the study population 203 are presented in Table 1. There was no evidence that supplementation elicited any injurious or 204 negative side effect(s). 205

The effects of vitamin D, zinc and combined zinc-vitamin D supplementation on BDI-II, serum 206 cortisol and BDNF are detailed in Table 2. No significant differences in BDI-II score were 207 observed among the 4 intervention arms at baseline. The baseline mean of the BDI-II score 208 was 19.21±7.34, for the whole population. All subjects had BDI-II≥10 at baseline, and after 12 209 weeks intervention, 45.2%, 66.7%, 59.4%, and 86.2% of participants in zinc, vitamin D, zinc-210 vitamin D and placebo groups, respectively. had BDI-II≥10. Zinc, vitamin D, and their co-211 supplementation yielded a significant reduction in depression score (p<0.0001). However, a 212 greater reduction in depression score was observed in the zinc group compared with vitamin D 213 group (p<0.001). No significant changes in BDI-II score were observed in placebo group 214 (p=0.396). 215

We found significant differences in serum zinc (p<0.001), vitamin D (table 3) (p<0.001), 216 cortisol (p=0.049) and BDNF (p=0.004) (table 2) between the study groups at baseline. The 217 baseline mean of BDNF, cortisol, zinc, and 25-(OH) D levels were 2.54 ± 1.61 ng/ml, 218 16.52 ± 5.87 ng/ml, 74.85 ± 36.36 mg/dl, and 18.99 ± 12.02 ng/ml respectively, for the whole 219 sample population. We found a significant increase in serum zinc, and 25-(OH) D levels among 220 those who received zinc, vitamin D or joint zinc–vitamin D supplements. However serum 25-(OH) D levels was also significantly reduced in the placebo group (p< 0.001). 222 Zinc, vitamin D or zinc+vitamin D had no significant effects on serum cortisol level. There 223 was a significant decrease in serum BDNF levels in the zinc (p=0.035) and placebo (p=0.016) 224 groups, respectively; whilst there was no significant change in the vitamin D group. 225

4.Discussion:

226

In the present study, we found that 3 months of supplementation with 30 mg/day of zinc 227 gluconate, and 2000 IU vitamin D, either individually or in combination, significantly 228 improved depression status compared to placebo in obese subjects who had BDI-II≥10. 229 However, among these participants we found that, monotherapy with 30 mg zinc gluconate 230 outperformed 2000 IU vitamin D in improving the depression-related symptoms over 12 weeks 231 of follow-up. There was no significant differences in BDI-II score among the 4 intervention 232 arms at baseline. Our results suggest that depressed patients could take zinc, vitamin D, either 233 individually or concurrently, to improve mood and depression severity. Previous studies have 234 shown that vitamin D or zinc supplementation, individually, may improve depression severity 235 (13, 23), although this has been disputed in some empirical work (34-36). 236

Although favorable effects from zinc supplementation on depression-related symptoms have 237 been demonstrated in previous clinical trials (13, 51, 52), some work has reported that there is 238 no evidence for the significant association between zinc and depression (34). Importantly, it 239 should be noted that the aforementioned clinical trials were generally conducted in people who 240 were not affected by obesity. Thus, the role of supplemental zinc in obese patients with 241 depression remains controversial. One meta-analysis showed that zinc concentrations were 242 approximately -1.85mmol/L lower in depressed subjects vs. control subjects (46). Similarly, 243 Zongyao Li noted a significant inverse association between dietary zinc intake and risk of 244 depression (44). The current trial suggests a causal association between zinc status and 245 depression in obese/overweight subjects, and that greater depression improvement is manifest 246 in zinc, compared to vitamin D, supplementation. 247

In the present study, we found that vitamin D elicited a favorable effect on depression scores 248 in overweight/obese subjects. Findings in some earlier clinical trials regarding the effect of 249 vitamin D supplementation on depression have been inconsistent (23, 35, 53). A large 250 randomized trial in older women assessed the effect of a single annual dose of 500 kIU vitamin 251 D for 3–5 y, and did not find any effect of vitamin D supplementation on depression symptom 252 (35); whilst further work reported that weekly supplementation with 40000 IU vitamin D for 6 253 months had no significant effect on depressive symptom scores, when compared with placebo 254 (54). A recent meta-analysis, which included nine trials with a total of 4923 participants, 255 concluded that no significant reduction in depression was seen after vitamin D supplementation 256 (45). In a double-blinded, placebo controlled prospective trial, involving 489 postmenopausal 257 elderly women, there was no effect of hormone therapy and calcitriol, either individually or in 258 combination, on depression (55). Supplementation with 400 IU of vitamin D combined with 259 1,000 mg of elemental calcium on measures of depression in a randomized, double-blinded US 260 trial comprising 36,282 postmenopausal women did not affect depression symptoms (56). 261 However, contrastingly, some studies have reported a beneficial effect of vitamin D 262 supplementation on depression (23). For instance, a trial conducted in elderly patients with 263 MDD showed that a single 300-kIU dose of vitamin D results in a decreased score of depression 264 (57); whilst in a study of 441 obese subjects, it was indicated that supplementation with 20.000 265 or 40.000 IU vitamin D per week for 1 year ameliorates depression symptoms (23). Congruent 266 with previous meta-analyses (58, 59), and a RCT study (23), our study found a significant 267 reduction in depressive symptoms after 3 months vitamin D supplementation. It is possible that 268 vitamin D is clinically beneficial for individuals who are depressed, but not in healthy 269 participants; thus, variability in depression stage could be considered as an explanatory factor 270 for this finding, but clearly necessitates further investigation. In the current trial, zinc 271 supplementation outperformed vitamin D for improving depression score. 272

Several biological mechanisms for the beneficial effects of zinc and vitamin D on depression 273 have been proposed (44, 60). 'Neurotrophin hypothesis of depression', mainly based on the 274 inverse association between stress and brain-derived neurotrophic factor (BDNF) levels, is one 275 of the most popular hypotheses in the pathophysiology of depression (61). Expression of BDNF 276 has been reported to be down-regulated in depressed patients, compared to non-depressed 277 matched controls (62), however, it is shown to increase after chronic antidepressant 278 administration (62). In Solati, et al., zinc supplementation resulted in increased serum BDNF 279 levels (13). However, in the present study, after 12 weeks of treatment with zinc, serum levels 280 of BDNF dropped significantly compared to baseline, but were not statistically different as 281 compared to the placebo group. This may conceivably be related to a placebo effect, because 282 the placebo group had a significant reduction in levels of BDNF, while no concurrent reduction 283 in depression was noted. In addition, the regression to mean phenomenon, where if a variable 284 is extreme on its first measurement, it will tend to be closer to the average on its second 285 measurement—and if it is extreme on its second measurement, it will tend to have been closer 286 to the average on its first, may be considered as an alternative explanation for the changes in 287

BDNF in the zinc and placebo groups. At baseline, the zinc group had a significantly higher288level of serum BDNF compare to all other groups. In Ranjbar et al, it was reported that serum289levels of BDNF in depressed patients receiving zinc supplement did not increase (51). In the290present study, to the best of our knowledge, we are the first to report vitamin D and joint zinc-291vitamin D supplementation has no effect on serum BDNF.292

Other mechanisms have been proposed to mediate the effects of zinc and vitamin D on 293 depression, including modulation of the hypothalamus-pituitary-adrenal (HPA) axis (63). Zinc 294 and vitamin D, an anti-inflammatory element, help to maintain endocrine homeostasis and 295 regulation of the hippocampal and cortical glutamatergic circuits that subserve affective 296 regulation and cognitive function (64, 65). However, in the current trial, serum levels of cortisol 297 in obese patients with depression receiving zinc, vitamin D or their co-supplementation were 298 not decreased. Several hypotheses exist to explain why no beneficial effects of zinc-vitamin D 299 joint supplementation, compared to zinc or vitamin D individually, reduced depression related 300 symptom. The dose of vitamin D we tested (2000 IU/day) may not have been sufficient to 301 affect the depression score in obese patients. Participants in our study were obese individuals 302 with MDD, who might have an increased need to vitamin D. In addition, normal status in zinc 303 at baseline and continuity of vitamin D deficiency at study cessation in the zinc-vitamin D 304 group could explain this finding. 305

The current trial has several strengths. Mood disorders, particularly depression, are among the 306 leading causes of morbidity in obese subjects (4, 5). Current modalities for treatment of 307 depression are insufficient and expensive (15, 16), and the prevalence of zinc and vitamin D 308 insufficiency is high (66, 67). Furthermore, our study was conducted on patients with 309 depressive symptoms with limited physical activity levels and sun exposure, which, in turn, 310 would contribute to vitamin D insufficiency (53). This study highlights the need for adequately 311 powered randomized clinical trials to establish whether there is a causal relation between zinc, 312 vitamin D status and depression severity in overweight/obese subjects. We believe that this 313 study is the first clinical trial to quantitatively evaluate the joint effects of zinc and vitamin D 314 monotherapy on serum BDNF, cortisol levels and depression severity in overweight/obese 315 subjects with depressive symptoms. In light of the current evidence represents a valuable 316 addition to the current findings on the efficacy of zinc and vitamin D monotherapy on 317 depression symptoms. 318

However, there are some limitations to the present RCT which must be considered. Firstly, the 319 study population was comprised of obese/overweight subjects with BDI-II≥10 who received 320 zinc, vitamin D supplementation in the absence of anti-depressant medications, thus, these 321 findings may not be generalizable to severely depressed subjects already prescribed anti-322 depressant medications. Second, our sample size calculation was based on the variability in 323 depression score, and may not have been adequate for the analysis of serum cortisol as a 324 secondary outcome. Third, serum cortisol levels were reported in this trial, however, it is 325 arguable that urine measurement is the preferred method for cortisol assessment. Furthermore, 326 daily physiological and behavioral rhythms including sleep, and body temperature can 327 significantly influence cortisol concentration, which we could not control in the present study. 328 Finally, in the current trial, despite the randomization, serum zinc and vitamin D level at 329 baseline were unequal, however, these variables were adjusted accordingly in the applied 330 analysis. 331

Supplementation with zinc, vitamin D or joint zinc-vitamin D can improve BDI-II score in 332 obese patients with depressive symptoms. However, the zinc, vitamin D and depression 333 improvements appear to be independent from serum cortisol and BDNF. To confirm the 334 veracity of the findings of present trial, further trials with longer durations and larger sample 335 sizes are needed. 336

Acknowledge:

Our research group would like to thank all subjects who took part in current study.	338
Author statement contributor:	339
The authors' responsibilities were as follows: SY, AE and SJ: designed the project, SS and SY	340
wrote the first draft of the manuscript; AM and SJ data analysis and interpreted the data, SJ,	341
AE, CC and MT: revised the subsequent drafts for important intellectual content, and approved	342
the final version of the manuscript to be published.	343
Conflict of interest:	344
SY, SS, AE, AM, MT, CC, and SJ declared that there were no conflicts of interest. This	345
research did not receive any specific grant from funding agencies in the public, commercial, or	346
not-for-profit sectors.	347
	348
	349
Role of funding source:	350

The project was financially supported by Iran University of Medical Sciences.	351
Submission declaration:	352
All authors have seen and approved the final manuscript. Neither the article nor any part it has	353
been published and is not under consideration elsewhere before appearing in the journal.	354
	355
	356

References:	357
1. Simon GE, Von Korff M, Saunders K, Miglioretti DL, Crane PK, Van Belle G, et al. Association	358
between obesity and psychiatric disorders in the US adult population. Archives of general psychiatry.	359
2006;63(7):824-30.	360
2. Marazziti D, Rutigliano G, Baroni S, Landi P, Dell'Osso L. Metabolic syndrome and major	361
depression. CNS spectrums. 2014;19(4):293-304.	362
3. Green AJ, Bazata DD, Fox KM, Grandy S. Quality of life, depression, and healthcare resource	363
utilization among adults with type 2 diabetes mellitus and concomitant hypertension and obesity: a	364
prospective survey. Cardiology research and practice. 2012;2012.	365
4. Faith M, Butryn M, Wadden T, Fabricatore A, Nguyen A, Heymsfield S. Evidence for	366
prospective associations among depression and obesity in population-based studies. Obesity	367
Reviews. 2011;12(5):e438-e53.	368
5. Zhao G, Ford ES, Li C, Tsai J, Dhingra S, Balluz LS. Waist circumference, abdominal obesity,	369
and depression among overweight and obese US adults: National Health and Nutrition Examination	370
Survey 2005-2006. BMC psychiatry. 2011;11(1):130.	371
6. Chen Y, Jiang Y, Mao Y. Association between obesity and depression in Canadians. Journal of	372
Women's Health. 2009;18(10):1687-92.	373
7. Harrison DL, Miller MJ, Schmitt MR, Touchet BK. Variations in the probability of depression	374
screening at community-based physician practice visits. Primary care companion to the Journal of	375
clinical psychiatry. 2010;12(5).	376
8. Pan A, Sun Q, Czernichow S, Kivimaki M, Okereke OI, Lucas M, et al. Bidirectional association	377
between depression and obesity in middle-aged and older women. International journal of obesity. 2012:595.	378 379
 Ma J, Xiao L. Obesity and depression in US women: results from the 2005–2006 National 	380
Health and Nutritional Examination Survey. Obesity. 2010;18(2):347-53.	381
10. Rosmond R, Björntorp P. The hypothalamic–pituitary–adrenal axis activity as a predictor of	382
cardiovascular disease, type 2 diabetes and stroke. Journal of internal medicine. 2000;247(2):188-97.	383
11. Joseph JJ, Wang X, Spanakis E, Seeman T, Wand G, Needham B, et al. Diurnal salivary	384
cortisol, glycemia and insulin resistance: the multi-ethnic study of atherosclerosis.	385
Psychoneuroendocrinology. 2015;62:327-35.	386
12. Stetler C, Miller GE. Depression and hypothalamic-pituitary-adrenal activation: a quantitative	387
summary of four decades of research. Psychosomatic medicine. 2011;73(2):114-26.	388
13. Solati Z, Jazayeri S, Tehrani-Doost M, Mahmoodianfard S, Gohari MR. Zinc monotherapy	389
increases serum brain-derived neurotrophic factor (BDNF) levels and decreases depressive	390
symptoms in overweight or obese subjects: a double-blind, randomized, placebo-controlled trial.	391
Nutritional neuroscience. 2015;18(4):162-8.	392
14. Isingrini E, Camus V, Le Guisquet A-M, Pingaud M, Devers S, Belzung C. Association between	393
repeated unpredictable chronic mild stress (UCMS) procedures with a high fat diet: a model of	394
fluoxetine resistance in mice. PLoS One. 2010;5(4):e10404.	395
15. Gartlehner G, Hansen RA, Morgan LC, Thaler K, Lux L, Van Noord M, et al. Second-generation	396
antidepressants in the pharmacologic treatment of adult depression: an update of the 2007	397
comparative effectiveness review [Internet]2011.	398
16. Rush AJ, Trivedi MH, Wisniewski SR, Nierenberg AA, Stewart JW, Warden D, et al. Acute and	399
longer-term outcomes in depressed outpatients requiring one or several treatment steps: a STAR* D	400
report. American Journal of Psychiatry. 2006;163(11):1905-17.	401
17. Trivedi MH, Rush AJ, Wisniewski SR, Nierenberg AA, Warden D, Ritz L, et al. Evaluation of	402
outcomes with citalopram for depression using measurement-based care in STAR* D: implications	403
for clinical practice. American journal of Psychiatry. 2006;163(1):28-40.	404
18. Andrews G, Sanderson K, Corry J, Lapsley HM. Using epidemiological data to model	405
efficiency in reducing the burden of depression. The journal of mental health policy and economics.	406
2000;3(4):175-86.	407

19. Lai JS, Hiles S, Bisquera A, Hure AJ, McEvoy M, Attia J. A systematic review and meta-analysis	408
of dietary patterns and depression in community-dwelling adults–. The American journal of clinical	409
nutrition. 2013;99(1):181-97.	410
20. Jacka FN, Maes M, Pasco JA, Williams LJ, Berk M. Nutrient intakes and the common mental	411
disorders in women. Journal of affective disorders. 2012;141(1):79-85.	412
21. Jacka FN, Pasco JA, Williams LJ, Mann N, Hodge A, Brazionis L, et al. Red meat consumption	413
and mood and anxiety disorders. Psychotherapy and psychosomatics. 2012;81(3):196-8.	414
22. Szewczyk B, Poleszak E, Sowa-Kucma M, Siwek M, Dudek D, Ryszewska-Pokrasniewicz B, et	415
alAntidepressant activity of zinc and magnesium in view of the current hypotheses of	416
antidepressant action. Pharmacological Reports. 2008;60(5):588.	417
	417
23. Jorde R, Sneve M, Figenschau Y, Svartberg J, Waterloo K. Effects of vitamin D	
supplementation on symptoms of depression in overweight and obese subjects: randomized double	419
blind trial. Journal of internal medicine. 2008;264(6):599-609.	420
24. Młyniec K, Nowak G. Zinc deficiency induces behavioral alterations in the tail suspension test	421
in mice. Effect of antidepressants. Pharmacological Reports. 2012;64(2):249-55.	422
25. Młyniec K, Davies CL, Budziszewska B, Opoka W, Reczyński W, Sowa-Kućma M, et al. Time	423
course of zinc deprivation-induced alterations of mice behavior in the forced swim test.	424
Pharmacological Reports. 2012;64(3):567-75.	425
26. Huang EP. Metal ions and synaptic transmission: think zinc. Proceedings of the National	426
Academy of Sciences. 1997;94(25):13386-7.	427
27. Szewczyk B, Kubera M, Nowak G. The role of zinc in neurodegenerative inflammatory	428
pathways in depression. Progress in Neuro-Psychopharmacology and Biological Psychiatry.	429
2011;35(3):693-701.	430
28. Szewczyk B, Poleszak E, Wlaź P, Wróbel A, Blicharska E, Cichy A, et al. The involvement of	431
serotonergic system in the antidepressant effect of zinc in the forced swim test. Progress in Neuro-	432
Psychopharmacology and Biological Psychiatry. 2009;33(2):323-9.	433
29. Nowak G, Siwek M, Dudek D, Ziêba A, Pilc A. Effect of zinc supplementation on	434
antidepressant therapy in unipolar depression: a preliminary placebo-controlled study. Polish journal	435
of pharmacology. 2003;55(6):1143-8.	436
30. Siwek M, Dudek D, Paul IA, Sowa-Kućma M, Zięba A, Popik P, et al. Zinc supplementation	437
augments efficacy of imipramine in treatment resistant patients: a double blind, placebo-controlled	438
study. Journal of affective disorders. 2009;118(1-3):187-95.	439
31. Garcion E, Wion-Barbot N, Montero-Menei CN, Berger F, Wion D. New clues about vitamin D	440
functions in the nervous system. Trends in Endocrinology & Metabolism. 2002;13(3):100-5.	441
32. Eyles DW, Smith S, Kinobe R, Hewison M, McGrath JJ. Distribution of the vitamin D receptor	442
and 1α -hydroxylase in human brain. Journal of chemical neuroanatomy. 2005;29(1):21-30.	443
33. Stumpf WE, Sar M, Clark SA, DeLuca HF. Brain target sites for 1, 25-dihydroxyvitamin D3.	444
Science. 1982;215(4538):1403-5.	445
34. Lehto SM, Ruusunen A, Tolmunen T, Voutilainen S, Tuomainen T-P, Kauhanen J. Dietary zinc	446
intake and the risk of depression in middle-aged men: a 20-year prospective follow-up study. Journal	447
of affective disorders. 2013;150(2):682-5.	448
35. Sanders KM, Stuart AL, Williamson EJ, Jacka FN, Dodd S, Nicholson G, et al. Annual high-dose	449
vitamin D 3 and mental well-being: randomised controlled trial. The British Journal of Psychiatry.	450
2011;198(5):357-64.	451
36. Zhao G, Ford ES, Li C, Balluz LS. No associations between serum concentrations of 25-	452
hydroxyvitamin D and parathyroid hormone and depression among US adults. British Journal of	453
Nutrition. 2010;104(11):1696-702.	454
37. Gloth FM, 3rd, Alam W, Hollis B. Vitamin D vs broad spectrum phototherapy in the	454
	455
treatment of seasonal affective disorder. The journal of nutrition, health & aging. 1999;3(1):5-7.	450
PubMed PMID: 10888476. Epub 2000/07/11. eng.	437

38. Harris S, Dawson-Hughes B. Seasonal mood changes in 250 normal women. Psychiatry Res.	458
1993 Oct;49(1):77-87. PubMed PMID: 8140183. Epub 1993/10/01. eng.	459
39. Vieth R, Kimball S, Hu A, Walfish PG. Randomized comparison of the effects of the vitamin	460
D3 adequate intake versus 100 mcg (4000 IU) per day on biochemical responses and the wellbeing of	461
patients. Nutrition journal. 2004 Jul 19;3:8. PubMed PMID: 15260882. Pubmed Central PMCID:	462
PMC506781. Epub 2004/07/21. eng.	463
40. Li G, Mbuagbaw L, Samaan Z, Falavigna M, Zhang S, Adachi JD, et al. Efficacy of vitamin D	464
supplementation in depression in adults: a systematic review. The Journal of clinical endocrinology	465
and metabolism. 2014 Mar;99(3):757-67. PubMed PMID: 24423304. Pubmed Central PMCID:	466
PMC5112012. Epub 2014/01/16. eng.	467
41. Anglin RE, Samaan Z, Walter SD, McDonald SD. Vitamin D deficiency and depression in	468
adults: systematic review and meta-analysis. The British journal of psychiatry : the journal of mental	469
science. 2013 Feb;202:100-7. PubMed PMID: 23377209. Epub 2013/02/05. eng.	470
42. Li Z, Li B, Song X, Zhang D. Dietary zinc and iron intake and risk of depression: A meta-	471
analysis. Psychiatry Res. 2017 May;251:41-7. PubMed PMID: 28189077. Epub 2017/02/12. eng.	472
43. Swardfager W, Herrmann N, Mazereeuw G, Goldberger K, Harimoto T, Lanctot KL. Zinc in	473
depression: a meta-analysis. Biol Psychiatry. 2013 Dec 15;74(12):872-8. PubMed PMID: 23806573.	474
Epub 2013/06/29. eng.	475
44. Li Z, Li B, Song X, Zhang D. Dietary zinc and iron intake and risk of depression: a meta-	476
analysis. Psychiatry research. 2017;251:41-7.	477
45. Gowda U, Mutowo MP, Smith BJ, Wluka AE, Renzaho AM. Vitamin D supplementation to	478
reduce depression in adults: meta-analysis of randomized controlled trials. Nutrition.	479
2015;31(3):421-9.	480
46. Swardfager W, Herrmann N, Mazereeuw G, Goldberger K, Harimoto T, Lanctôt KL. Zinc in	481
depression: a meta-analysis. Biological psychiatry. 2013;74(12):872-8.	482
47. Claro da Silva T, Hiller C, Gai Z, Kullak-Ublick GA. Vitamin D3 transactivates the zinc and	483
manganese transporter SLC30A10 via the Vitamin D receptor. The Journal of steroid biochemistry	484
and molecular biology. 2016 Oct;163:77-87. PubMed PMID: 27107558. Epub 2016/04/25. eng.	485
48. Efird J. Blocked randomization with randomly selected block sizes. International journal of	486
environmental research and public health. 2011 Jan;8(1):15-20. PubMed PMID: 21318011. Pubmed	487
Central PMCID: PMC3037057. Epub 2011/02/15. eng.	488
49. Pirotta S, Kidgell D, Daly R. Effects of vitamin D supplementation on neuroplasticity in older	489
adults: a double-blinded, placebo-controlled randomised trial. Osteoporosis international.	490
2015;26(1):131-40.	491
50. Ghassemzadeh H, Mojtabai R, Karamghadiri N, Ebrahimkhani N. Psychometric properties of	492
a Persian-language version of the Beck Depression Inventory-Second edition: BDI-II-PERSIAN.	493
Depression and anxiety. 2005;21(4):185-92.	494
51. Ranjbar E, Shams J, Sabetkasaei M, M-Shirazi M, Rashidkhani B, Mostafavi A, et al. Effects of	495
zinc supplementation on efficacy of antidepressant therapy, inflammatory cytokines, and brain-	496
derived neurotrophic factor in patients with major depression. Nutritional neuroscience.	497
2014;17(2):65-71.	498
52. Salari S, Khomand P, Arasteh M, Yousefzamani B, Hassanzadeh K. Zinc sulphate: a reasonable	499
choice for depression management in patients with multiple sclerosis: a randomized, double-blind,	500
placebo-controlled clinical trial. Pharmacological Reports. 2015;67(3):606-9.	501
53. Sepehrmanesh Z, Kolahdooz F, Abedi F, Mazroii N, Assarian A, Asemi Z, et al. Vitamin D	502
Supplementation Affects the Beck Depression Inventory, Insulin Resistance, and Biomarkers of	503
Oxidative Stress in Patients with Major Depressive Disorder: A Randomized, Controlled Clinical Trial,	504
2. The Journal of nutrition. 2015;146(2):243-8.	505
54. Kjærgaard M, Waterloo K, Wang CE, Almås B, Figenschau Y, Hutchinson MS, et al. Effect of	506
vitamin D supplement on depression scores in people with low levels of serum 25-hydroxyvitamin D:	507

nested case—control study and randomised clinical trial. The British Journal of Psychiatry.	508
2012;201(5):360-8.	509
55. Yalamanchili V, Gallagher JC. Treatment with hormone therapy and calcitriol did not affect	510
depression in elderly postmenopausal women: no interaction with estrogen and vitamin D receptor	511
genotype polymorphisms. Menopause (New York, NY). 2012;19(6):697.	512
56. Bertone-Johnson ER, Powers SI, Spangler L, Larson J, Michael YL, Millen AE, et al. Vitamin D	513
supplementation and depression in the women's health initiative calcium and vitamin D trial.	514
American journal of epidemiology. 2012;176(1):1-13.	515
57. Zanetidou S, Murri MB, Buffa A, Malavolta N, Anzivino F, Bertakis K. Vitamin D supplements in geriatric major depression. International journal of geriatric psychiatry. 2011;26(11):1209-10.	516 517
58. Shaffer JA, Edmondson D, Wasson LT, Falzon L, Homma K, Ezeokoli N, et al. Vitamin D	517
supplementation for depressive symptoms: a systematic review and meta-analysis of randomized	518
controlled trials. Psychosomatic medicine. 2014;76(3):190.	520
59. Li G, Mbuagbaw L, Samaan Z, Falavigna M, Zhang S, Adachi JD, et al. Efficacy of vitamin D	520
supplementation in depression in adults: a systematic review. The Journal of Clinical Endocrinology	522
& Metabolism. 2014;99(3):757-67.	523
60. Bertone-Johnson ER. Vitamin D and the occurrence of depression: causal association or	524
circumstantial evidence? Nutrition reviews. 2009;67(8):481-92.	525
61. Martinowich K, Manji H, Lu B. New insights into BDNF function in depression and anxiety.	526
Nature neuroscience. 2007;10(9):1089.	527
62. Duman R. Pathophysiology of depression: the concept of synaptic plasticity. European	528
psychiatry. 2002;17:306-10.	529
63. Dallman MF, Pecoraro NC, La Fleur SE, Warne JP, Ginsberg AB, Akana SF, et al.	530
Glucocorticoids, chronic stress, and obesity. Progress in brain research. 2006;153:75-105.	531
64. Swardfager W, Herrmann N, McIntyre RS, Mazereeuw G, Goldberger K, Cha DS, et al.	532
Potential roles of zinc in the pathophysiology and treatment of major depressive disorder. Neuroscience & biobehavioral reviews. 2013;37(5):911-29.	533 534
65. Al-Dujaili EA, Munir N, Iniesta RR. Effect of vitamin D supplementation on cardiovascular	535 535
disease risk factors and exercise performance in healthy participants: a randomized placebo-	536
controlled preliminary study. Therapeutic advances in endocrinology and metabolism.	537
2016;7(4):153-65.	538
66. Marreiro DDN, Fisberg M, Cozzolino SMF. Zinc nutritional status in obese children and	539
adolescents. Biological trace element research. 2002;86(2):107-22.	540
67. Mithal A, Wahl DA, Bonjour J-P, Burckhardt P, Dawson-Hughes B, Eisman JA, et al. Global	541
vitamin D status and determinants of hypovitaminosis D. Osteoporosis international.	542
2009;20(11):1807-20.	543
	544
	544
	545
	546
	F 47
	547
	548
	549
	550
	551

552
553
554
555
556
557
558
559
560
561

		Zinc (n=24)	Vitamin D (n=27)	Zinc-vitaminD (n=25)	Placebo (n=22)	P-value
Age (years))	38.71±7.16	38.28±7.28	38.93±5.39	37.31±7.12	0.823
Blood	Baseline	121.20±12.72	121.61±13.81	119.26±14.07	116.79±11.75	0.625
pressure	After 12	117.40±11.28	118.00±15.76	114.73±14.66	118.17±12.58	0.790
systolic	week					
	\mathbf{P} -value ^{\mathbf{I}}	0.117	< 0.0001	< 0.0001	0.613	
Blood	Baseline	80.68±7.91	84.58±9.79	80.30±10.80	80.21±6.07	0.882
pressure	After 12					
diastolic	week	81.55±7.48	81.96±10.00	78.73±11.94	79.08±10.92	0.127
	P-value [†]	0.055	0.014	<0.0001	0.212	
Weight	Baseline	86.40±10.13	90.42±16.51	86.31±14.36	87.20±12.86	0.625
(Kg)	After 12	86.62±9.89	90.61±16.01	85.42±14.88	87.67±12.62	0.493
	week					
	P-value [†]	0.884	0.636	0.284	0.770	
BMI(kg/	Baseline	29.08±2.96	30.59±4.08	29.59±3.64	30.11±3.68	0.428
m ²)	After 12	29.14±2.99	30.63±3.93	29.38±3.48	30.13±3.73	0.373
	week					
	P-value [†]	0.866	0.717	0.154	0.850	
Gender(ma	le/female)	26/8	27/7	24/9	26/6	0.760
Waist	Baseline	103.63±5.84	106.91±11.39	105.25±8.95	103.84±8.58	0.513
circumfer	After 12	103.14±6.38	105.33±11.69	103.85±9.20	104.43±9.53	0.846
ence	week	<0.0001	< 0.0001	< 0.0001	< 0.0001	
	P-value ^H					

Table 1: characterization of participants at baseline and after 12 week

Note: BMI=Body Mass Index ‡: Values are analyzed by one-way analysis of variance; I: Values are analyzed by paired-samples T test values are mean ± SD

Table 2: change in BDI-II, serum BDNF and cortisol level from baseline to 12 weeks

	Zinc (mean) (95% CI)	Vitamin D (mean)	Zinc-vitaminD(mean)	Placebo (mean) (95%	P-value [‡]	ß
		(95% CI)	(95% CI)	CI)		
BDI-II score	-7.02 (-9.13, -4.74)	-3.87 (-6.15, -1.59)	-7.62 (-10.63, -4.61)	-0.76 (-3.05, 1.52)	0.0001*	0.969
BDNF	-0.19 (-0.36, -0.02)	-0.11 (-0.28, 0.05)	0.19 (-0.02, 0.41)	-0.17 (-0.34, -0.003)	0.08	0.589
Cortisol	-1.37 (-3.43, 0.67)	-1.66 (-3.71, 0.38)	-0.82 (-3.52. 1.88)	-1.19 (-3.29, 0.90)	0.974	0.056

BDI-II: Beck Depression Score-II

 [‡]: Values are analyzed by ANCOVA (adjusted by baseline serum zinc, vitamin D, cortisol and BDNF and beck depression score);
 *Significant differences between zinc and placebo groups, vitamin D and placebo groups, zinc-vitamin D and placebo groups, zinc and vitamin D groups values are mean \pm SD

571

		Zinc	Vitamin D	Zinc-vitamin D	Placebo	P-value [‡]	ß
Zinc µg/dl	Baseline	57.13±21.91	60.25±17.74	115.37±22.45	62.48±19.01	>0.001*	
	After 12 week	77.03±47.26	58.60±13.30	127.90±38.65	56.80±14.36	>0.001**	0.969
	P-value ¹	0.008	0.588	0.068	0.213		
Vitamin D	Baseline	15.86±9.03	26.07±13.27	10.44±5.23	20.51±10.43	>0.001***	
ng/ml	After 12 week	14.00±8.06	36.29±11.28	17.93±7.28	16.85±10.07	>0.001****	1.000
	P-value ¹	0.141	>0.001	>0.001	>0.001		

 *: Values are analyzed by ANCOVA (adjusted by baseline serum zinc, vitamin D); I: Values are analyzed by paired-samples T test *Significan differences between zinc-vitamin D and placebo groups, zinc-vitamin D and zinc groups, zinc-vitamin D and vitamin D groups ** Significan differences between zinc-vitamin D and placebo groups, zinc-vitamin D and vitamin D groups, zinc and vitamin D groups. 	577 578 579 580
*** significant differences between vitamin D and zinc groups, zinc- vitamin D and placebo groups, zinc- vitamin D and vitamin D groups. **** significant differences between zinc-vitamin D and vitamin D groups, zinc-vitamin D and placebo groups, zinc-vitamin D and zinc groups, vitamin D and zinc groups, zinc-vitamin D and placebo groups, zinc-vitamin D and zinc groups, vitamin D	580 581 582

Flow diagram1: depicting progress through different phases of the clinical trial

