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30 Abstract

51	
32	Background: There is little information about serum phosphate levels among patients with pulmonary
33	tuberculosis (TB) and HIV infection.
34	Objective: We aimed to to assess the role of TB, HIV, inflammation and other correlates of on serum phosphate
35	levels.
36	Methods: A cross-sectional study was conducted among TB patients and age- and sex-matched non-TB controls.
37	Pulmonary TB patients were categorized as sputumnegative (TB-) and _positive (TB+)_ based on culture. Age-
38	and sex-matched non-TB controls were randomly selected among neighbors to TB+ sputum-positive TB patients.
39	Data on age, sex, alcohol and smoking habits were obtained. HIV status, serum phosphate, and the acute phase
40	reactants C-reactive protein (serum CRP) and α_1 -acid glycoprotein (serum AGP) were determined. Linear
41	regression analysis was used to identify correlates of serum phosphate.
42	Results: Of 1605 participants, 355 (22.1%) were controls and 1250 (77.9%) TB patients, of which 9.9% and
43	50.4% were HIV-infected. Serum phosphate was determined before start of TB treatment in 44%, and 1-14 days
44	after start of treatment in 56%. Serum phosphate was up to 0.10 mmol/L higher 1-3 days after start of TB
45	treatment, and lowest 4 days after treatment, after which it increased. In multivariable analysis, TB patients had
46	0.09 (95%CI: 0.05; 0.13) mmol/L higher serum phosphate than controls, and those with HIV had 0.05 (95%CI:
47	0.01; 0.08) mmol/L higher levels than those without. Smoking was also a positive correlate of serum phosphate,
48	whereas male sex and age were negative correlates.
49	Conclusion: While HIV and TB are associated with higher serum phosphate, our data suggest that TB treatment is
50	followed by transient reductions in serum phosphate, which may reflect hypophosphataemia in some patients.
51	
52	Key words: Serum phosphate, phosphorus, tuberculosis; HIV, acute phase response
53	
54	

5657 INTRODUCTION

58 Phosphorus is an essential mineral in the body, being, -a structural component of DNA and RNA and of 59 membranes, and important for metabolism and storage of energy (1). Phosphorus is likely to be a limiting nutrient 60 among individuals in low-income settings, since the typical diet is low in animal-source foods and high in cereals 61 (2). Although cereals, including maize, have a high content of phosphorus in the form of phytic acid (;ie-inositol 62 hexaphosphate), it is largely unabsorbable (2,3). 63 64 Data on phosphorus status are-is scarce, even among low-income individuals where phosphorus deficiency is 65 likely to be common. Although serum phosphate is known to be a poor marker of phosphorus status, it is the only 66 available (3), and low values are related to various clinical conditions. For example, serum phosphate is measured 67 to monitor patients at risk of refeeding hypophosphataemia, a potentially fatal condition that arises when high 68 energy feeding is initiated in as-phosphorus depleted individuals suddenly are fed high amounts of energy (4). As 69 refeeding restores metabolism, it also leads to further intracellular phosphorus depletion, due to utilization of 70 phosphorus for anabolic processes and storage of energy through phosphorylation of ADP to ATP-Such 71 refeeding hypophosphatemia potentially results in multi-organ failure and death (5). 72 73 Recently, there has been a renewed interest in serum phosphate in patients with HIV infection on antiretroviral 74 treatment (ART). Tenofovir has been shown to cause renal tubular function abnormalities (6), which is a known 75 cause of hypophosphataemia (7,8). Among HIV patients starting ART in Zambia, a case of acute 76 hypophosphataemia was described (9), and low serum phosphate prior to ART was found to be a predictor of 77 early mortality among those with low BMI (10,11). Fortification of a lipid-based nutritional supplement with 78 vitamins and minerals reduced renal wasting of phosphate among malnourished Zambian patients starting ART 79 (2).

80

As part of a larger nutrition study, we obtained cross-sectional data on serum phosphate among pulmonary TB
patients and age- and sex-matched neighbourhood controls, with an aim to assess the level of serum phosphate
and the role of pulmonary TB, HIV, the acute phase response and other potential correlates.

85 METHODOLOGY

86

87 Ethics Statement

- Ethical permission was obtained from the Medical Research Coordinating committee of the National Institute for
 Medical Research in Tanzania, and consultative approval was given by The Danish Central Medical Ethics
 Committee. Written and oral information was presented to all eligible participants by the health staff before
 written informed consent was obtained. Written consent was obtained from parents/legal guardians of any
 participant under 18 years of age.
- 93

94 Study setting and design

A cross-sectional study was conducted from April 2006 to March 2009 in Mwanza City, Tanzania, among TB patients recruited for a large nutrition intervention study and non-TB controls. Mwanza City is at the shores of Lake Victoria. The harvest is from May to July, and the staple foods are maize, cassava, sweet potato, rice, and millet. Fish is the most common animal-source food. and sSmall fish are often eaten whole, -but only 25% eat them >4 days per week (12).

100

101 Recruitment and management of TB patients

102 The TB patients were recruited at the four TB clinics under the TB treatment services, coordinated by the 103 National Tuberculosis and Leprosy Programme. If residents of Mwanza city, Both-both smearsputum-positive 104 (TB+) and smearsputum-negative (TB-) TB patients, based on culture, were enrolled in the study after giving 105 informed consent if they were residents of Mwanza city. Patients were excluded if pregnant, under the age of 15 106 years or were suffering from with extra-pulmonary TB .- pregnancy, age under 15 years, or a terminal illness-were 107 excluded. The diagnosis of TB followed the World Health Organization (WHO) guidelines (13) using the Ziehl-108 Neelsen staining technique (14). Briefly, all patients suspected of having TB were asked to bring three sputum 109 samples for microscopy, and chest X-rays were done as appropriate. Patients were considered to be smearsputum-110 positive, if two samples tested positive or one sample tested positive and a chest X-ray was suggestive of TB, and 111 to be smearsputum-negative TB patients if all the samples were negative, but chest X-ray and clinical suspicion 112 was suggestive of TB, and there was non-response to a course of broad-spectrum antibiotics. After diagnosis all 113 patients were started on a standardized TB treatment for 6-8 months based on existing national guidelines (15,16).

114 Those found to be HIV-infected were referred for management based on national guidelines at the time of the 115 study (17). At On the day TB treatment was started, the patients also started daily supplementation as part of two 116 nutrition intervention trials. In the energy-protein trial, those found TB+sputum-positive -and HIV co-infected 117 were randomized to receive one or six energy-protein biscuit bars daily (18). One of the biscuit bars given to the 118 experimental group and the one given to the control group contained additional micronutrients, so that the 119 micronutrient intake was similar in the two groups. All other TB patients were randomized to a daily biscuit bar 120 with or without additional micronutrients (19). Each biscuit bar weighed 30 g and contained 4.5 g protein, 615 kJ 121 energy, and 120 mg P.

122

123 Recruitment of non-TB controls

124 400 Four hundred consecutive smearsputum-positive participants were considered index cases for selection of 125 age- and sex-matched neighbourhood non-TB controls. Mwanza City is divided into wards, streets and communal 126 cells. Each cell has 10-20 households, and is headed by a ten-cell -leader. Each of the index patients was asked to 127 provide his/her residential address and the name of his/her ten-cell leader. Using this information, the study team 128 requested the ten-cell leader to provide the complete list of individuals in his/her jurisdiction meeting the age and 129 sex recruitment criteria. Of these, one was randomly selected using a lottery method and invited to participate in 130 the study as a non-TB control if meeting the following criteria: no history of previous TB exposure, active TB or 131 TB treatment, no evidence of current active TB (cough, intermittent fevers, and excessive night sweating in the 132 past two weeks and unexplained weight loss in the past month), same sex as index case, aged 15 years or above 133 and age difference from index case less than five years, had lived in the same street as index case for at least three 134 months, not pregnant, and consenting to participate in the study. Persons who were terminally ill were not invited. 135 The recruitment of non-TB controls was done in parallel with inclusion of cases from October 2006 to January 136 2009.

137

138 Data collection

For the purpose of the study, all TB patients provided an additional sputum sample for culture at the Zonal TB Reference Laboratory, and were subsequently categorized as <u>TBsputum+-positive</u> or <u>TBsputum-negative</u>, based on culture. For missing or contaminated culture samples, the initial evaluation from sputum smear microscopy was used. All TB patients and controls had data on demography, smoking, and alcohol intake collected using

143	questionnaires, while data on ART were-was retrieved from ART-use databases in ART clinics. Morning venous
144	blood was collected in a 10 ml plain vacutainer tube for HIV testing and a 5 ml EDTA vacutainer tube for CD4
145	count. This was preferably done immediately prior to start of TB treatment, but could for logistical reasons be
146	delayed; median delay was - As previously reported, the median (range) delay of blood sampling was 1 (range 0-
147	14) days after initiation of TB treatment (20). All tubes were cooled on dry ice before transported to the
148	laboratory, where they were centrifuged and serum samples were stored at -80°C. HIV status was determined
149	using Capillus HIV-1/HIV-2 (Trinity Biotech Plc., Wicklow, Ireland) and Determine HIV-1/HIV-2 (Inverness
150	Medical Innovations, Inc., Delaware, U.S.A.) tests in parallel. HIV infection was diagnosed if both tests gave a
151	positive result and HIV negative diagnosis was made if both tests produced a negative result. Indeterminate
152	results were resolved using ELISA- Organon Uniform II (Organon Teknia Ltd, Boxtel, Netherlands). CD4 count
153	was determined as cells/µl using a Partec Cyflow Counter (Partec GmbH, Münster, Germany). The biochemical
154	analyses, ie acute phase reactants and phosphate, were conducted at Aalborg University Hospital, FBE Clinical
155	Biochemistry South. Serum phosphate was determined using Phosphate (Inorganic) ver.2 (PHOS2) on a Cobas
156	6000 instrument from Roche (Roche Diagnostics GmbH, Mannheim, Germany) according to the manufacturer's
157	instructions. The cut-offs used to define low and high serum phosphate were 0.80 and 1.60 mmol/L (3). Serum α_{1} -
158	acid glycoprotein (AGP) was determined with a standard Alpha1-Acid Glycoprotein Kit using Beckman Coulter
159	ImageH Immunochemistry Systems (Beckman Coulter, Galway, Ireland) and C-reactive protein (CRP) was
160	determined with Tina-quant C-Reactive Protein Gen.3 (CRPL3) on a Roche COBAS 6000 instrument (Roche
161	Diagnostics GmbH, Mannheim, Germany).
162	
163	Statistical analysis
164	Normal probability plots were used to assess the distribution of continuous variables. Chi-square test was used to
165	test for differences in proportions. To assess and adjust for the effect of TB treatment and nutritional interventions
166	on serum phosphate in case blood sampling was delayed, a variable was created to express the number of days

167 delay after start of TB treatment. For controls and those treated the same day or the day after blood sampling this

- 168 variable was given the value 0. The two-sample t test or oneway ANOVA were used to test for differences in
- 169 means between two or more groups, respectively, and Scheffe post hoc tests were used to adjust for multiple
- 170 comparisons. Linear regression analysis was used to identify correlates of serum phosphate. The variables
- 171 assessed were age, sex, smoking, consumption of alcohol, TB and HIV status, and serum CRP or AGP. Age and

- 172 sex, and variables found significant in the univariate analyses were assessed in a final multivariable analysis, with
- 173 all variables included, with and without adjustment for elevated levels of either serum CRP or AGP. Year and
- 174 month of recruitment and delay in blood sampling since initiation of TB treatment were adjusted for. We
- 175 examined normal and residual-vs.-fitted plots to assess normality and homoscedasticity of residuals. Stata version
- 176 12.1 (StataCorp, Texas, USA) was used for all analyses.
- 177

178 **Results**

179	Of the 1605 study participants, 355 (22.1%) were controls and 1250 (77.9%) TB patients. Culture data were
180	available on 1142 (91.4%) of the 1250 TB patients. In the remaining 108 (8.6%) cultures were
181	contaminated or missing, and the categorization of TB patients as sputum-negative and sputum-positive
182	therefore based on microscopy. Thus, of the 1250 TB patients, of which 427 (34.2%) were TBsputum-
183	negative and 823 (65.8%) TB+- sputum-positive (Table 1). As previously reported (19), the HIV prevalence was
184	higher among TB patients compared to controls (50.4 vs 9.9%, p<0.001), and higher among sputum-negative TB-
185	compared to TB+sputum-positive patients (64.4 vs 43.1%, p<0.001). The mean BMI was 18.8 among TB patients
186	and 22.6 among controls (p<0.001). Data on serum phosphate were available for 1522 (94.8%) of the 1605
187	participants. Among 349 controls, mean (±SD) serum phosphate was 1.14 (±0.28) mmol/L with 4.3% (n=15)
188	having values below 0.80 mmol/L, and 4.9% (n=17) above 1.6 mmol/L, respectively. Among 1173 TB patients
189	the mean (\pm SD) serum phosphate was 1.27 (\pm 0.29) mmol/L, and 2.2% (n=26) had values below 0.80 mmol/L, and
190	9.2% (n=108) above 1.6 mmol/L. Of these, 518 (44%) had blood samples taken before start of TB treatment,
191	while 218 (18.6%) were bled with 1 day delay, and the remaining with 2-14 days delay. Those bled with delay
192	were 2.3 (95% CI: 0.8; 3.7) years older, had a higher prevalence of HIV (54.7 vs 45.5%, p=0.01), whereas there
193	was no difference in sex distribution (p>0.30). As seen in the Figure 1, unadjusted mean serum phosphate was
194	up to 0.10 mmol/L higher in those bled 1-3 days after start of TB treatment, and lowest in those bled with 4 days
195	delay, after which it seemed to increase with number of days delay. Numbers were too small to allow
196	stratification by nutritional intervention.
197	
198	Mean serum phosphate by category of sex, age, smoking, alcohol consumption, pulmonary TB and HIV is shown
199	in Table 12, with TB patients and controls combined. There were no differences by sex and age in univariate
200	analyses, but serum phosphate were higher in those smoking or taking alcohol. There was no difference in serum
201	phosphate between TB-sputum-negative and TB+sputum-positive TB patients (1.26 vs 1.27 mmol/L, p=0.41).
202	However, TB patients together had higher serum phosphate than controls (1.27 vs 1.14 mmol/L, p<0.0001). The
203	difference was similar if tested only among the index cases and controls (1.26 vs 1.14 mmol/L, p<0.0001; not
204	shown in table). HIV+ patients had higher serum phosphate than HIV- (1.29 vs 1.21 mmol/L, p<0.0001),
205	irrespective of ART status. The association between HIV status and serum phosphate was not different between

206 TB patients and controls (interaction, p=0.56, data not shown). While serum phosphate was higher among those

with HIV, it was lower in those with CD4 counts below 250 compared to above 500 cells/µL, although the
difference was only marginally significant (p=0.08, Scheffe post-hoc). Elevated serum CRP or AGP were both
associated with higher serum phosphate.

210

211 The results of a multivariable analysis, with adjustment for year and months of recruitment and delay in blood 212 sampling, are shown in Table 23. The relationship between age, sex, HIV, smoking and serum phosphate were 213 not different between TB patients and controls (interaction, p>0.10). Without adjustment for elevated serum AGP 214 (model 1), serum phosphate was lower in males compared to females, and lower in those above 25 years of age. 215 Alcohol intake was not associated with serum phosphate, but current smoking was associated with higher levels. 216 TB patients had 0.09 (95% CI: 0.05; 0.13) mmol/L higher serum phosphate compared to the non-TB controls, 217 whereas there was no difference between sputum-positive TB+ and TB-sputum-negative patients. Finally, those 218 with HIV infection had 0.05 (95% CI: 0.01; 0.08) mmol/L higher levels than those without. The associations with 219 delayed bleeding, assessed in this multivariable model, was similar to what was shown in the Figure 1. As such, 220 delays for 1 to 3 days were associated with 0.10 (95% CI: 0.06; 0.15), 0.01 (95% CI: -0.06; 0.09) and 0.08 (95% 221 CI: 0.02; 014) higher serum phosphate, while delay to day 4 was associated with 0.06 (95% CI: -0.01; 0.13) lower 222 serum phosphate. If days since TB treatment were not adjusted for, then the regression coefficient for TB was 223 0.12 (95% CI: 0.08; 0.15).

224

225 Elevated serum AGP was a strong positive correlate of serum phosphate, while elevated serum CRP was not, in 226 multivariable analysis. Adjustment for elevated serum AGP (Table 23, model 2) considerably reduced the 227 regression coefficient of TB+sputum-positive TB (from 0.09 to 0.03 mmol/L), whereas that of HIV and other 228 correlates did not change considerably. Compared to the overall mean serum phosphate of 1.24 (95% CI: 1.23; 229 1.25), the intercept was 1.20 (95% CI: 1.13; 1.26), and reflects the mean among individuals in all reference 230 categories, ie young, non-smoking females without TB, HIV and elevated serum AGP. While no interaction 231 between age and sex was found (p=0.40), there was an interaction between age and sex among controls (p=0.01). 232 The interaction reflected a decline in serum phosphate per 10 year increase in age among males (-0.04, 95% CI: -233 0.08; -0.010, p=0.01), but not among females (0.01, 95% CI: -0.03; .05, p=0.56).

235 DISCUSSION

236

237 Hypophosphataemia

238 We found that serum phosphate, compared to those examined before TB treatment start, was higher in those 239 examined 1-3 days after start of TB treatment, but lower in those examined 4 days after. While selection bias 240 cannot be excluded, this pattern more likely reflects changes in phosphate metabolism due to the commencement 241 of TB treatment with regain in appetite, and increased food intake, from the diet as well as from the supplements 242 provided as part of the trials. The nadir at day 4 may reflect that some TB patients could have refeeding 243 hypophosphataemia. In the classical description of refeeding syndrome, starved individuals refed with high 244 amounts of glucose and amino acids developed hypophosphataemia accompanied by cardio-pulmonary failure. 245 The existence of a similar syndrome among HIV patients starting ART has been suggested (9-11), whereas it 246 does not seem to have been studied among TB patients. Nevertheless, the risk and magnitude of refeeding 247 hypophosphataemia after initiation of TB-treatment may depend on the initial phosphorus status, as well as the 248 intake of energy and bioavailable phosphorus and other bulk minerals and probably vitamins. There is currently 249 increasing awareness that patients with TB need nutritional care and support (21), and it is important to ensure 250 adequate intake of phosphorus to not only to prevent refeeding syndrome, but also to support regain in lean mass 251 and body functions. 252

253 After adjustment for the effect of delayed blood sampling, TB was associated with 0.09 mmol/L higher serum 254 phosphate, compared to no TB, much of which was explained by elevated serum acute phase reactants. Yet, there 255 are several reasons to believe that phosphorus status was low. First, the staple food is maize and the intake of 256 animal source foods is limited. This is supported by a relatively low mean serum phosphate among controls (1.14 257 mmol/L), although within the reference interval was 0.80 and 1.60 mmol/L, with 4.3% having low values. 258 Second, the TB patients in the study had typically been ill for some time, and we have previously shown that they 259 have an average weight deficit of 9 kg (22). The weight deficit is due to lower habitual weight as well as reduced 260 food intake and increased utilization of energy as a result of the TB disease itself. Since inflammation-induced 261 wasting to a large extent is due to catabolism of lean mass, this may result in increased serum phosphate. It has 262 been shown in children with inflammatory bowel disease, another condition involving systemic inflammation, 263 that flare-up leads to a sustained upregulation (reduced degradation) of the phosphatonin Fibroblast Growth

Factor 23 (FBG23), which increases renal phosphorus excretion (23). Hence, <u>although speculative</u>, it is likely that patients with TB, despite the elevated serum phosphate, are phosphorus depleted, and may continue to have high urinary phosphorus excretion for some time. This will contribute to increase the phosphorus requirements during the critical period of convalescence when there is a need to rebuild lean mass, ie organ and muscle.

268

269 HIV infection

270 We found 0.05 mmol/L higher serum phosphate in those with compared to without HIV infection, both among 271 TB patients and controls. Among HIV patients, serum phosphate was not different in the 76 ART-treated 272 compared to the 558 ART-naïve patients. The main ART regimen used was stavudine/lamivudine/nevirapine, 273 whereas tenofovir, known to cause hypophosphataemia as part of Fanconi's syndrome (6,7), was not used. In 274 contrast to TB, only a minor part of the association between HIV and serum phosphate was explained by the acute 275 phase response. In a trial among malnourished Zambian and Tanzanian HIV patients starting ART, there were 276 complex interrelationships between serum phosphate and early mortality which were accentuated by vitamin and 277 mineral fortification of a lipid-based supplement (24,25)(Woodd et al., submitted for publication; Rehman et al., 278 submitted for publication) although the supplement improved renal phosphate retention (2). The results suggested 279 it was variability of serum phosphate, possibly due to poor metabolic control among malnourished, seriously ill 280 patients, which was associated with mortality risk (25) (Rehman et al., submitted for publication). 281 282 Serum phosphate was lower in males compared to females and in the higher compared to lower age groups in the 283 multivariable models. However, among controls only, we found an interaction between age and sex, due to a 284 decline in serum phosphate with increasing age in males, but not in females. The overall decline of serum 285 phosphate with age has been reported from several population based studies (26,27). A large US-study from the 286 USA found a decline with age in men. However, in women, there was that the lack of a consistent a decline in 287 serum phosphate with age among females concealed a decline up to 44 years as for males, and then a transient 288 increase with the onset of menopause, in parallel with changes in tubular phosphate reabsorption (28). While this 289 age-sex pattern was also seen among healthy adults in Tanzanians, it disappeared among TB patients, even after

adjustment for other factors. The higher serum phosphate in smokers is also in accordance with previous studies

291 (29), and may be explained by greater bone loss in smokers.

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- 292 Despite the limitations of serum phosphate as a marker of phosphorus status, and of our cross-sectional
- 293 design to draw conclusions about cause-effect relationship, the study suggests that some patients may experience
- 294 refeeding hypophosphataemia a few days after start of treatment.

Table 1. Diagnosis of 1250 tuberculosis	natients as s	putum-negative or s	nutum-positive
Tuble It Blaghosis of 1250 tubereulosis	patiento ao o	putum negutive of 5	putum positive

	Sputu	Sputum status ¹			
	Positive ²	Negative ³	Total (%)		
Culture	754 ⁴	388	1142 (91,4)		
Microscopy	69	39	108 (8,6)		
Total (%)	823 (65.8)	427 (34,2)	1250 (100%)		

 10tal (%)
 025 (05.0)
 427 (34,2)
 1250 (105.0)

 1
 Sputum status was based on culture, if available, otherwise microscopy.

 2
 Patients were considered to have sputum-positive tuberculosis, if two samples tested positive or one sample tested positive and a chest X-ray was suggestive of tuberculosis.

 3
 Patients were considered to have sputum-negative tuberculosis if all the samples were negative, but chest X-ray and clinical suspicion was suggestive of tuberculosis, and there was non-response to a course of brack-spectrum antibiotics.

 of broad-spectrum antibiotics.

⁴ Based on 400 consecutive sputum-positive index cases 400 age- and sex-matched neighbourhood non-TB controls were selected

L	% (n)	Mean (SD)	95% CI	Р
Sex				
Females	42.0 (639)	1.25 (0.28)	1.23; 1.28	0.11
Males	58.0 (883)	1.23 (0.31)	1.21; 1.25	
Age (y)				
<25	21.9 (333)	1.27 (0.28)	1.23; 1.30	0.16
25-45	58.2 (886)	1.24 (0.30)	1.22; 1.25	
45+	19.9 (303)	1.23 (0.31)	1.19; 1.26	
Smoking				
Never	71.3 (1073)	1.23 (0.30)	1.21; 1.25	0.04
Previously	8.9 (134)	1.27 (0.27)	1.22; 1.31	
Currently	19.8 (297)	1.27 (0.28)	1.24; 1.30	
Alcohol intake				
No	58.0 (883)	1.23 (0.29)	1.21; 1.24	0.03
Yes	42.0 (638)	1.26 (0.31)		
Pulmonary TB status ¹				
Non-TB control	22.9 (349)	1.14 (0.28)	1.11; 1.17	< 0.0001
TB-Sputum-negative TB	26.7 (406)	1.26 (0.27)	1.23; 1.29	
TB+Sputum-positive TB	50.4 (767)	1.27 (0.31)	1.25; 1.30	
HIV and ART status				
HIV-	58.3 (888)	1.21 (0.26)	1.19; 1.22	< 0.0001
HIV+ not on ART	36.7 (558)	1.29 (0.34)	1.26; 1.31	
HIV+ on ART	5.0 (76)	1.28 (0.29)	1.22; 1.35	
CD4 count (cells/µL)				
HIV-	58.4 (888)	1.21 (0.26)		< 0.0001
500+	6.4 (97)	1.35 (0.32)	1.28; 1.41	
250-500	12.6 (191)	1.30 (0.31)	1.26; 1.34	
<250	22.7 (345)	1.26 (0.35)	1.22; 1.30	
Serum C-Reactive Protein (mg/L)				
≤ 2	20.1 (305)	1.16 (0.28)	1.13; 1.20	< 0.0001
2-10	13.3 (201)	1.19 (0.24)	1.15; 1.22	
10-50	18.2 (275)	1.24 (0.26)	1.21; 1.27	
50-100	25.7 (390)	1.26 (0.27)	1.23; 1.28	
100+	22.7 (344)	1.32 (0.36)	1.28; 1.35	
Serum a1-Acid Glycoprotein (mg/L)				
≤ 1	25.6 (389)	1.15 (0.27)	1.13; 1.18	< 0.0001
1-2	20.1 (305)	1.22 (0.27)	1.19; 1.25	
2-3	37.4 (567)	1.26 (0.26)	1.24; 1.28	
3+	16.9 (257)	1.35 (0.39)	1.30; 1.40	

Table 2. Serum phosphate (mmol/L) among <u>1173 of</u> 1250 pulmonary TB patients and <u>349 of</u> 355 non-TB neighbourhood controls by categories of sex, age, smoking, pulmonary TB, HIV and serum acute phase reactants 1

¹ Pulmonary TB status was based on culture, except where culture data were not available. For 355 consecutively recruited sputum_positive TB patients a control was randomly selected among individuals

14

from the neighbourhood with same sex and age. Serum phosphate data were available on 1522, but n may sum up to less, due to missing data on smoking (n=18), alcohol intake (n=1), CD4 count (n=1), serum C-Reactive Protein (n=7), serum α 1-Acid Glycoprotein (n=4). P-values were based on t-test and oneway. TB is tuberculosis, HIV is human immunodeficiency syndrome, ART is antiretroviral treatment

297 298

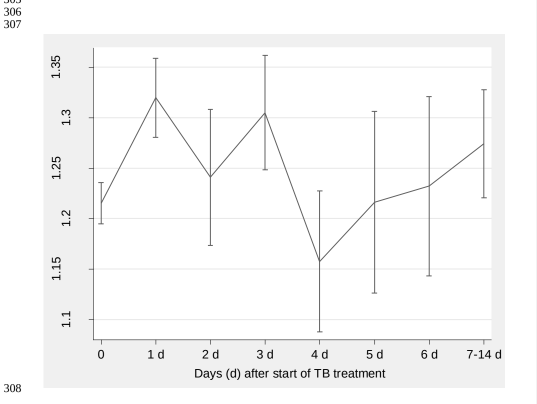
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Table 3. Multivariable models with correlates of serum phosphate in among 1173 of 1250 pulmonary TB
patients and <u>349 of</u> 355 non-TB neighbourhood controls with regression coefficient B, 95% confidence
interval (CI) and P-values ¹

		Model 1 ²		Model 2 ³		
	В	95% CI	Р	В	95% CI	Р
Sex						
Female	-					
Male	-0.04	-0.07; -0.005	0.02	-0.04	-0.07; -0.01	0.02
Age (years)						
<25	-					
25-45	-0.06	-0.10; -0.03	0.001	-0.06	-0.10; -0.02	0.001
45+	-0.07	-0.12; -0.02	0.003	-0.06	-0.10; -0.01	0.02
Smoking						
Never	-					
Previously	0.04	-0.01; 0.10	0.12	0.04	-0.01; 0.10	0.13
Currently	0.06	0.02; 0.10	0.004	0.06	0.02; 0.10	0.004
TB status ¹						
Non-TB control	-			-		
ТВ	0.09	0.05; 0.13	<0.000 1	0.03	-0.03; 0.09	0.34
HIV status						
HIV-	-					
HIV+	0.05	0.01; 0.08	0.004	0.04	0.01; 0.07	0.02
Serum α_1 -acid glycoprotein (mg/L)						
<1				-		
1-2				0.02	-0.04;0.08	0.56
2-3				0.05	-0.01; 11	0.10
3+				0.14	0.07; 0.20	< 0.001

¹ Pulmonary TB status was based on culture, and microscopy only if culture data were not available. For 355 consecutively recruited sputum positive TB patients a control was randomly selected among individuals with same sex and age from the neighbourhood. ² Model 1: N=1491, adjusted R²=0.07 and intercept=1.20 (95% CI: 1.14; 1.27). ³ Model 2: N=1487, adjusted R²=0.09 and intercept=1.20 (95% CI: 1.13; 1.26). Both models contained all the variables and were adjusted for year and months of recruitment and days since start of TB treatment. TB is tuberculosis, HIV is human immunodeficiency syndrome

\$01 302 303 304 305 Figure 1. Serum phosphate by day after start of of TB treatment. Based on linear regression, with adjustment for TB, and non-TB controls and TB patients commencing TB treatment before or at the day of blood sampling coded as 0. Number of participants: day 0 (n=867), 1 (n=218), 2 (n=72), 3 (n=102), 4 (n=67), 5 (n=40), 6 (n=41) and 7-14 (n=115).



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