The association of serum 25-hydroxyvitamin D\textsubscript{3} and D\textsubscript{2} with depressive symptoms in childhood – a prospective cohort study

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Background: Depression in adolescence is common and early onset predicts worse outcome in adulthood. Studies in adults have suggested a link between higher total 25-hydroxyvitamin D [25(OH)D] concentrations and lower risk of depression. \textbf{Objectives:} To investigate (a) the association between serum 25(OH)D\textsubscript{2} and 25(OH)D\textsubscript{3} concentrations and depressive symptoms in children, and (b) whether the associations of 25(OH)D\textsubscript{2} and 25(OH)D\textsubscript{3} are different from, and independent of, each other. \textbf{Methods:} Prospective cohort study with serum 25(OH)D\textsubscript{2} and 25(OH)D\textsubscript{3} concentrations measured at mean age of 9.8 years and depressive symptoms assessed with the Mood and Feelings Questionnaire by a trained interviewer at the mean ages of 10.6 years (n = 2,759) and 13.8 years (n = 2,752). \textbf{Results:} Higher concentrations of 25(OH)D\textsubscript{3} assessed at mean age 9.8 years were associated with lower levels of depressive symptoms at age 13.8 years [adjusted risk ratio (RR; 95% confidence interval (CI)): 0.90 (0.86–0.95)], but not at age 10.6 years [adjusted RR (95% CI): 0.98 (0.93–1.03)] and with increased odds of decreasing symptoms between age 10.6 and 13.8 years [adjusted RR (95% CI): 1.08 (1.01–1.16)]. Serum 25(OH)D\textsubscript{2} concentrations were not associated with depressive symptoms. \textbf{Conclusions:} This is the first study in children to suggest that the association between 25(OH)D\textsubscript{3} concentrations and depression emerges in childhood. The association is independent of a wide range of potential confounding factors, and appears to be stronger with greater time separation between assessment of 25(OH)D\textsubscript{3} and assessment of depressive symptoms. Confirmation of our findings in large prospective studies and trials would be valuable. \textbf{Keywords:} 25-Hydroxyvitamin D, calcium, parathyroid hormone, child, depression, ALSPAC.

Introduction
Depression affects 1–6% of adolescents worldwide and early onset often predicts more serious disease manifestation in later life (Thapar, Collishaw, Potter, & Thapar, 2010). Characterisation of modifiable risk factors that could be used to prevent or delay the early onset of depression is important. It has been suggested that higher concentrations of vitamin D may protect against depression in adults. Depression rates are higher in winter than in summer, which could support a role for vitamin D (Bertone-Johnson, 2009). Some (Armstrong et al., 2007; Berk et al., 2007; Eskandari et al., 2007; Ganji, Milone, Cody, McCarty, & Wang, 2010; Hoogetdijk et al., 2008; Jorde, Waterloo, Saleh, Haug, & Svartberg, 2006, Lee et al., 2010; Schneider, Weber, Fresch, Stein, & Fritz, 2000; Wilkins, Sheline, Roe, Birge, & Morris, 2006) but not all (Annweiler et al., 2010; Herran et al., 2000; Michelson et al., 1996; Nanni et al., 2009; Pan et al., 2009; Zhao, Ford, Li, & Balluz, 2010) cross-sectional studies in adults have found an association between higher serum concentrations of 25-hydroxyvitamin D (25(OH)D) and lower risk of depression in adults. Two prospective studies found that higher concentration of 25(OH)D were associated with lower risk of depression in older adults (May et al., 2010; Milaneschi et al., 2010) and three randomised controlled trials in selective populations had inconsistent findings, with a trial of vitamin D\textsubscript{3} supplements improving depression symptoms in overweight/obese adults (Jorde, Sneve, Figenschau, Svartberg, & Waterloo, 2008) but two further trials of D\textsubscript{3} showing no effect on symptoms in older women with seasonal affective disorder (Dumville et al., 2006) or on preventing depression in older women (Sanders et al., 2011).

25(OH)D\textsubscript{3} is a robust and reliable indicator of vitamin D status, reflecting both dietary intake and synthesis in skin, which normally accounts for most of the vitamin D in humans (Seamans & Cashman, 2009). Circulatory total 25(OH)D consists of 25(OH)D\textsubscript{3} [metabolite of vitamin D\textsubscript{3} synthesised in skin after ultraviolet B (UVB) exposure and obtained from animal food sources] and 25(OH)D\textsubscript{2} (synthesised from vitamin D\textsubscript{2} obtained from plant sources). 25(OH)D\textsubscript{3} and 25(OH)D\textsubscript{2} are converted to 1,25-dihydroxyvitamin D\textsubscript{3} and D\textsubscript{2}, the steroid hormones...
that mediate the biological actions of vitamin D. The former is known to have higher affinity to vitamin D binding protein and receptor (Glendenning et al., 2009; Houghton & Vieth, 2006) and with respect to bone health, vitamin D₃ has been suggested to be more potent than D₂ (Finch, Brown, & Slatopolsky, 1999). However, to date no studies have examined whether associations between these two differ with respect to depression. Vitamin D, together with parathyroid hormone (PTH), regulates calcium and phosphate homeostasis (Brown, Dusso, & Slatopolsky, 1999; Mundy & Guise, 1999) and some (Eskandari et al., 2007; Herran et al., 2000, Hoo- gendijk et al., 2008; Jorde et al., 2006; May et al., 2010) but not all (Michelson et al., 1996; Schneider et al., 2000) studies have reported higher serum PTH among adults with depression. As the previous studies have included only adults, it is unknown if serum 25(OH)D concentrations are associated with mood in childhood or early adolescence. Examining this association in childhood/adolescence is important because confounding by alcohol, smoking and mood-altering drugs is somewhat less likely than in adult studies and because it is increasingly recognised that depression can emerge in childhood/adolescence (Thapar et al., 2010) and its prevention may be best started at this age.

The aims of this study were (a) to investigate the prospective association between serum 25(OH)D₂ and 25(OH)D₃ concentrations and depressive symptoms in children, (b) to investigate if the associations of 25(OH)D₂ and 25(OH)D₃ are different, and (c) to examine whether any associations were independent of serum PTH, phosphate and calcium concentrations.

Participants and methods

Population

The Avon Longitudinal Study of Parents and Children (ALSPAC) is a population-based birth cohort from South West England. The cohort consisted of 14,062 live births from 14,541 enrolled pregnant women who were expected to give birth between 1 April 1991 and 31 December 1992 (Golding, Pembrey, Jones and ALSPAC Study Team, 2001). From age 7, all children were invited for an annual assessment of physical and psychological development. Parents gave informed consent at enrolment, and ethical approval was obtained from the ALSPAC Law and Ethics Research Committee and the National Health Service (NHS) local research ethics committee.

Single and twin births were included in this study; the very small number of triplets and quadruplets were not included for reasons of confidentiality. Figure 1 shows how the numbers included in the analyses presented here was derived. Complete data on outcomes, exposures and confounders were available from 2,759 and 2,752 children, respectively, for assessment with outcomes at 10.6 years and with outcomes at 13.8 years.

Outcome

The prevalence of depressive symptoms was evaluated with the Mood and Feelings Questionnaire (MFQ) by a trained interviewer at the mean ages of 10.6 and 13.8 years. The MFQ is a 13-item depression inventory validated for use in 6- to 18-year-olds. Each item is scored between 0 and 2, resulting to a maximum score of 26 with higher scores indicating presence of symptoms of depression. MFQ correlates highly with more extensive evaluations like the Children’s Depression Inventory and the Diagnostic Interview Schedule for Children (Costello & Angold, 1988).

The MFQ scores were positively skewed even after various transformations so we generated three categories of the score (0–2, 3–5, 6), representing approximate thirds of the distribution. An ordinal categorical variable was derived to indicate change in MFQ score category (increase by two categories/increase by one category/no change/decrease by one category/decrease by two categories) between ages 10.6 and 13.8 years.

Exposures and phlebotomy-based covariables

Serum 25(OH)D₃, 25(OH)D₂, PTH, phosphate and calcium were assayed on nonfasting blood samples.
collected at mean age 9.9 years for the majority of participants \((n = 2,130)\) for MFQ assessed at age 13.8 years and \(n = 2,493\) for MFQ assessed at age 10.6 years). If no samples were available from the 9.9-year assessment, samples from mean age 11.8 years (\(n = 416\)) or, second, the 7.6-year assessments \((n = 206)\) for MFQ assessed at age 13.8 years and \(n = 266\) for MFQ assessed at age 10.6 years) were used. The mean age at sample collection in the whole study sample was 9.8 years (standard deviation: \(SD = 0.74\)). To keep the analyses prospective, we excluded exposure measurements that were taken at 11.8-year clinic when the outcome was measured at age 10.6 years.

Following collection, samples were immediately spun, frozen and stored at −80 °C. Assays were performed after a maximum of 12 years in storage with no previous freeze-thaw cycles. 25(OH)D3, 25(OH)D2 and deuterated internal standard were extracted from serum samples, following protein precipitation, using Isolute C18 solid phase extraction cartridges. Potential interfering compounds were removed by initial elution with 50% methanol followed by elution of the vitamins using 10% tetra-hydrofuran in acetonitrile. Dried extracts were reconstituted prior to injection into a high-pressure liquid chromatography tandem mass spectrometer (Waters Acuity, Manchester, UK). The following transitions (mass to charge ratio) in multiple reaction mode were used: 413.2 > 395.3, 401.1 > 383.3 and 407.5 > 107.2 for 25(OH)D2, 25(OH)D3, and hexa-deuterated 25(OH)D3 respectively. Interassay coefficients of variation for the assay were < 10% across a working range of 1–250 ng/ml for both 25(OH)D2 and 25(OH)D3. Data on head of household occupational social class, ethnicity, parents’ education and family history of depression and schizophrenia were obtained from parent-completed questionnaires. Time spent outdoors during summer months on school days, school weekends and holidays was reported as None, 1 hr/day, 1–2 hr/day and 3 or more hr/day in parent-completed questionnaires at mean age of 8.5 years. Responses were coded as follows: None = 0, 1 = 1, 1–2 = 1.5 and 3 = 5. Average hours spent outdoors per summer day (1 June–31 August) were calculated using term dates from Bristol City Council’s Education Committee term dates for 2001–2002 (summer term 1 June–24 July, holidays 24 July–31 August). Information on protection from UVB exposure (use of sunblock, covering clothing or hat and avoidance of midday sun) were obtained from the same questionnaires. A summary variable for UVB protection score was derived by scoring the responses to questions on use of sunblock, covering clothing or hat and avoidance of midday sun as Always = 3, Usually = 2, Sometimes = 1, Never = 0 and summing these scores. This gives a single variable that ranges from 0 to 12, with 0 indicating the least meticulous protection from UVB.

Height and weight were measured at the same time as blood samples for obtaining 25(OH)D3 and other assays and were used to calculate BMI. Total IQ score in Wechsler Intelligence Scale for Children (WISC–III UK version) was assessed at mean age 8.5. Puberty stage was assessed by parental report using Tanner staging (Tanner, 1962) of pubic hair, breast and genitalia development on repeat occasions. In our analyses, we used data from the questionnaire closest to the time of phlebotomy for the exposures for each child.

25(OH)D3 concentrations had displayed a sinusoidal seasonal variation (Figure 2). In order to derive a value of 25(OH)D3 for each participant that was accurately adjusted for the seasonal effects of when the sample was taken, we used linear regression with date of blood sampling as the independent variable and loge 25(OH)D3 as the outcome (dependent) variable and with trigonometric sine and cosine functions. 25(OH)D3 was loge transformed in this regression model to ensure that the residuals in the sine–cosine regression were approximately normally distributed. The residuals from this regression are the participants logged 25(OH)D3 having adjusted for seasonal differences in when the sample was taken. This seasonal adjusted 25(OH)D3 variable represents a participants average 25(OH)D3 across seasons and is the main 25(OH)D3 exposure measurement used in our analyses [we also present associations without this adjustment]. There was no mass index (BMI) and cognitive function to be important confounders because of their known associations with 25(OH)D3 concentrations and depressive symptoms. We also adjusted for pubertal stage as this might affect depressive symptoms and 25(OH)D3. Data on head of household occupational social class, ethnicity, parents’ education and family history of depression and schizophrenia were obtained from parent-completed questionnaires.

Confounding factors

We considered gender, age, ethnicity (white, non-white), head of household occupational social class, maternal and paternal education, family history of depression or schizophrenia, UVB exposure, body

The association of potential confounders with serum 25(OH)D$_3$, 25(OH)D$_2$, calcium, phosphate and PTH concentrations was assessed with linear regression and associations of confounders with PTH concentrations was assessed with linear regression. Serum 25(OH)D$_3$, 25(OH)D$_2$, calcium, phosphate and PTH were age- and gender-standardised using the internal cohort data with age in 1-month categories.

The association of potential confounders with serum 25(OH)D$_3$, 25(OH)D$_2$, calcium, phosphate and PTH concentrations was assessed with linear regression and associations of confounders with depressive symptoms with ordered logistic regression. We investigated the linearity of associations between 25(OH)D$_3$ and 25(OH)D$_2$ and depressive symptoms by splitting the 25(OH)D variables into fifths of their distribution and graphically examining the odds of depressive symptoms across these fifths. We statistically tested for linearity by obtaining a quintile effect on the log odds of depressive symptoms at age 13 years and higher BMI with higher depressive symptoms at ages 10.6 and 13.8 years. Higher socioeconomic position and IQ founders and depressive symptoms at ages 10.6 and 13.8 years. Higher socioeconomic position and IQ.

The main analyses assessing associations of seasonally adjusted 25(OH)D$_3$ and 25(OH)D$_2$ with outcomes were done with a nonparametric bootstrap procedure (10,000 replications) in conjunction with ordered logistic regression using `bsample` and `ologit` commands in Stata. The bootstrapping procedure (Efron & Tibshirani, 1986) enabled us to statistically compare associations of 25(OH)D$_3$ to those of 25(OH)D$_2$. The difference between the effect of 25(OH)D$_3$ and 25(OH)D$_2$ was calculated from the bootstrap replicate distribution. To numerically compare the associations of two forms of 25(OH)D, we scaled them the same by multiplying the beta coefficients from the regression models by log$_2$(2). The results are interpreted as the difference in odds of MFQ between the lowest and middle or middle and highest MFQ score per doubling of exposure.

In addition to examining associations with 25(OH)D$_2$ and 25(OH)D$_3$ on a continuous scale, we also examined the association between total 25(OH)D deficiency and insufficiency and depressive symptoms. Total 25(OH)D was calculated by summing 25(OH)D$_2$ and 25(OH)D$_3$ and insufficiency was defined as a value below 30 ng/ml and deficiency below 20 ng/ml (Holick, 2007). Finally, in a sensitivity analysis we examined the association of 25(OH)D$_3$ that was not adjusted for seasonal variation with outcomes. All association analyses were performed for both genders combined as there was no strong statistical evidence for Gender × Exposure interaction (p ≥ .20).

**Results**

The mean (interquartile range) serum concentrations of season-adjusted 25(OH)D$_3$ and 25(OH)D$_2$ were 24.9 (24.7–25.1) and 1.3 (0.5–2.7) ng/ml, respectively. The median (interquartile range) of serum phosphate, calcium and PTH were 1.54 (1.43–1.64) mmol/L, 2.38 (2.31–2.44) mmol/L and 4.5 (3.4–5.8) pmol/L respectively. Other characteristics of the participants are shown in Table S1.

Table S2 shows the associations of 25(OH)D$_3$, 25(OH)D$_2$, phosphate, calcium and PTH concentrations with potential confounders. Those of nonwhite ethnicity had higher PTH and lower 25(OH)D$_3$ concentrations. BMI was negatively associated with serum 25(OH)D$_3$ and 25(OH)D$_2$ concentrations and positively with PTH concentrations. Higher socioeconomic position of parents was associated with lower concentrations of calcium and 25(OH)D$_2$ and higher concentrations of 25(OH)D$_3$. Less meticulous protection from UVB was associated with lower PTH concentrations and higher 25(OH)D$_3$ concentrations. Children who spent more time outdoors during summer had higher 25(OH)D$_3$ and 25(OH)D$_2$ concentrations. Children with family history of psychiatric problems had lower 25(OH)D$_3$ and calcium concentrations. Those with more advanced puberty stage had lower 25(OH)D$_3$ concentrations and higher calcium concentrations.

Table S3 summarises the associations of confounders and depressive symptoms at ages 10.6 and 13.8 years. Higher socioeconomic position and IQ were associated with lower risk of depressive symptoms at age 10.6 years and higher BMI with higher risk of depressive symptoms at age 13 years. Children who spent more time outdoors during summer...
had lower risk of depressive symptoms at age 13.8 years. Children with family history of psychiatric problems had higher risk of depressive symptoms at both ages.

The associations of 25(OH)D3 and 25(OH)D2 with depressive symptoms were either linear or null, with no strong evidence of deviation from linearity or suggestion of a threshold association (Figure 3).

Table 1 shows the prospective associations between serum 25(OH)D3, 25(OH)D2, phosphate, calcium and PTH concentrations and depressive symptoms at ages 10.6 and 13.8 years. Higher concentrations of 25(OH)D3 were associated with lower risk of depressive symptoms at age 13.8 years but not at age 10.6 years. 25(OH)D2 was not associated with depressive symptoms at either age. There was statistical evidence that the association of 25(OH)D3 with depressive symptoms at age 13.8 years differed from the association of 25(OH)D2 with the same outcome ($p \leq .001$), but there was no evidence of difference in associations of 25(OH)D3 and 25(OH)D2 with symptoms at age 10.6 years ($p = .81$). There was some suggestion that higher concentrations of PTH were associated with lower risk of depressive symptoms at age 13.8 years after adjusting for all confounders and other measured analytes, but

Figure 3 Odds of depressive symptoms at age 10.6 years (A and B) and age 13.8 years (C and D) by fifths of the distribution of 25(OH)D3 (A and C) and 25(OH)D2 (B and D)

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Exposure</th>
<th>Model 1a</th>
<th>Model 2b</th>
<th>Model 3c</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depressive symptoms at age 10.6 years</td>
<td>25(OH)D3</td>
<td>0.98 (0.93, 1.03)</td>
<td>0.98 (0.93, 1.03)</td>
<td>0.98 (0.93, 1.03)</td>
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<tr>
<td></td>
<td>25(OH)D2</td>
<td>1.02 (0.94, 1.10)</td>
<td>1.01 (0.93, 1.10)</td>
<td>0.99 (0.94, 1.04)</td>
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<tr>
<td></td>
<td>Albumin-adjusted calcium</td>
<td>1.02 (0.97, 1.07)</td>
<td>1.01 (0.97, 1.06)</td>
<td>1.00 (0.95, 1.05)</td>
</tr>
<tr>
<td></td>
<td>Phosphate</td>
<td>1.03 (0.98, 1.08)</td>
<td>1.04 (0.99, 1.09)</td>
<td>1.04 (0.98, 1.09)</td>
</tr>
<tr>
<td></td>
<td>Parathyroid hormone</td>
<td>0.98 (0.93, 1.03)</td>
<td>0.98 (0.94, 1.03)</td>
<td>0.97 (0.93, 1.03)</td>
</tr>
<tr>
<td>Depressive symptoms at age 13.8 years</td>
<td>25(OH)D3</td>
<td>0.90 (0.86, 0.94)</td>
<td>0.91 (0.86, 0.95)</td>
<td>0.90 (0.86, 0.95)</td>
</tr>
<tr>
<td></td>
<td>25(OH)D2</td>
<td>1.02 (0.95, 1.11)</td>
<td>1.03 (0.95, 1.12)</td>
<td>1.02 (0.97, 1.08)</td>
</tr>
<tr>
<td></td>
<td>Albumin-adjusted calcium</td>
<td>0.99 (0.94, 1.04)</td>
<td>1.00 (0.95, 1.04)</td>
<td>0.99 (0.94, 1.04)</td>
</tr>
<tr>
<td></td>
<td>Phosphate</td>
<td>1.01 (0.97, 1.06)</td>
<td>1.02 (0.97, 1.06)</td>
<td>1.02 (0.98, 1.07)</td>
</tr>
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<td></td>
<td>Parathyroid hormone</td>
<td>0.99 (0.94, 1.03)</td>
<td>0.98 (0.93, 1.02)</td>
<td>0.96 (0.91, 1.01)</td>
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</tbody>
</table>

OR, odds ratio.

aModel 1 is unadjusted (the exposures are standardised for age and gender and 25OHD3 is adjusted for season and ethnicity).
bModel 2 is adjusted for ethnicity, head of household social class, mothers and partners education, time spent outdoors during summer (age 8.5 years), UVB protection score, WISC IQ score at 8.5 years, BMI, family history of psychiatric problems and puberty stage.
cModel 3 is adjusted for Model 2 plus serum concentrations of other hormones/metabolites which are related vitamin D homeostasis [e.g. association of 25(OH)D3 is adjusted for 25(OH)D2, phosphate, albumin-adjusted calcium and parathyroid hormone].
Table 2  Prospective association of total 25(OH)D deficiency and insufficiency with depressive symptoms assessed by short Mood and Feelings Questionnaire at age 10.6 years (n = 2,759; vitamin D status assessed at mean age 9.2 years) and age 13.8 years (n = 2,752; vitamin D status assessed at mean age 9.8 years).

<table>
<thead>
<tr>
<th>Outcome \ Exposure</th>
<th>OR for category decrease per doubling of exposure (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Model 1(^a)</td>
</tr>
<tr>
<td>Depressive symptoms at age 10.6 years</td>
<td>0.97 (0.87, 1.08)</td>
</tr>
<tr>
<td>Depressive symptoms at age 13.8 years</td>
<td>1.00 (0.90, 1.11)</td>
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</table>

OR, odds ratio.
\(^a\)Model 1 is unadjusted. \(^b\)Model 2 is adjusted for ethnicity, age, gender, head of household social class, mothers and partners education, time spent outdoors during summer (age 8.5 years), UVB protection score, WISC IQ score at 8.5 years, BMI, maternal history of psychiatric problems and puberty stage. \(^c\)Model 3 is adjusted for Model 2 plus serum concentrations of phosphate, albumin-adjusted calcium and parathyroid hormone.

Discussion

In our prospective study of children, we have found higher concentrations of season-adjusted 25(OH)D\(_3\) assessed at mean age 9.8 years to be associated with lower levels of depressive symptoms at age 13.8 years and with increased odds of decreasing symptoms between age 10.6 and 13.8 years. These associations were independent of a wide range of potential confounding factors, as well as of 25(OH)D\(_2\), calcium, phosphate and PTH concentrations, which were not strongly associated with depressive symptoms at either age. We also found statistical evidence that the association of 25(OH)D\(_3\) with depressive symptoms was stronger than that of 25(OH)D\(_2\). The association of 25(OH)D\(_3\) with depressive symptoms was linear across the distribution of 25(OH)D\(_2\) concentrations, suggesting that even amongst those with what would be considered normal concentrations, an increase might result in somewhat lower depressive symptoms (if our findings are in future studies shown to represent causal effects). Consistent with these findings, and reflecting the fact that 25(OH)D\(_3\) is the biggest contributor to total 25(OH)D, we found that risk of depressive symptoms was greater at 13.8 years in those with total 25(OH)D deficiency or total 25(OH)D insufficiency.

To our knowledge, this is the first prospective study to examine this association in children. Our findings in this cohort of children are consistent with findings from the two prospective studies in adults (May et al., 2010; Milaneschi et al., 2010) and from...
The association of 25(OH)D3 with depressive symptoms in children only emerged with symptoms measured 3 years after exposure assessment, and was not present when symptoms were assessed just 1 year after exposure assessment. One might expect a stronger association with the earlier age, possibly explained by differences in outcome measurement and these previous studies in adults could be explained by differences in outcome measurement and study sample or could reflect real differences in these associations by age.

Study strengths and limitations

To our knowledge, this is the first study to examine these associations in children and is one of the few studies to examine them prospectively. We had a large sample size and were able to examine potential confounding by a wide range of characteristics and study the different effects of 25(OH)D3 and 25(OH)D2. We also used self-reported, rather than parent-reported depressive symptoms. This is important because parent-reported scores do not reveal depressive symptoms as early as self-reported measures (Cole et al., 2002). Consistent with other prospective cohort studies there has been substantial attrition over time with those who continued to attend the follow-up clinics being more likely to be from higher socioeconomic backgrounds (Golding, Pembrey, Jones and ALSPAC Study Team, 2001). Twenty-seven per cent of our study population had one randomised controlled trial that examined the effect of vitamin D3 supplementation on depressive symptoms in adults (Jorde et al., 2008).

The association of 25(OH)D3 with depressive symptoms in children only emerged with symptoms measured 3 years after exposure assessment, and was not present when symptoms were assessed just 1 year after exposure assessment. One might expect a stronger association with the earlier age, possibly in part because of reverse causality (i.e. depressive symptoms resulting in less outdoor activity and hence reduced vitamin 25(OH)D3 concentrations). It is possible that children with higher concentrations of 25(OH)D3 at age 9.8 years have on average higher concentrations over the subsequent 3 years and that the inverse association with depressive symptoms requires an accumulation of consistent concentrations. Alternatively, factors other than 25(OH)D3 but that are associated with it and accumulate over time (e.g. outdoor physical activity) might explain the association. The risk factors for childhood–onset depression may also be different from those of adolescence–onset depression (Jaffee et al., 2002). Lastly, it is possible that the biological pathways linking 25(OH)D3 to depression involve a chain of effects that take some time to emerge.

The stronger association of 25(OH)D3 compared to 25(OH)D2 could be a chance finding. The difference could also reflect possible greater residual confounding by, for example, different dietary patterns associated with 25(OH)D2 and 25(OH)D3 or outdoor physical activity, which will affect D3 more than D2; whilst we have attempted to adjust for a wide range of potential confounding factors residual confounding is possible. Lastly, the different associations could be explained by D3 being truly more potent at preventing depressive symptoms than D2. This finding requires further replication in other studies before we can conclude that D3 is more strongly associated with depressive symptoms than is D2.

Vitamin D receptors are expressed throughout the brain and both 25(OH)D3 and 25(OH)D2 cross the blood–brain barrier (Eyles, Smith, Kinobe, Hewison, & McGrath, 2005). Animal studies have shown that vitamin D is essential for normal neurogenesis (Cui, McGrath, Burne, Mackay-Sim, & Eyles, 2007), learning ability and behaviour in rodents (Becker, Eyles, McGrath, & Grecksch, 2005; Eyles et al., 2006; Harms, Eyles, McGrath, Mackay-Sim, & Burne, 2008), but currently it is unknown if the neural actions of vitamin D metabolites affect monoamine concentrations, hypothalamic–pituitary–adrenal axis responsiveness to stress or other mechanisms involved in depression (Belmaker & Agam, 2008).

To our knowledge, this is the first study to examine these associations in children and is one of the few studies to examine them prospectively. We had a large sample size and were able to examine potential confounding by a wide range of characteristics and study the different effects of 25(OH)D3 and 25(OH)D2. We also used self-reported, rather than parent-reported depressive symptoms. This is important because parent-reported scores do not reveal depressive symptoms as early as self-reported measures (Cole et al., 2002). Consistent with other prospective cohort studies there has been substantial attrition over time with those who continued to attend the follow-up clinics being more likely to be from higher socioeconomic backgrounds (Golding, Pembrey, Jones and ALSPAC Study Team, 2001). Twenty-seven per cent of our study population had
total 25(OH)D concentration below 20 ng/ml so the results are likely to apply to other populations with high prevalence of low 25(OH)D concentrations (Lips, 2010; Mansbach, Ginde, & Camargo, 2009).

Depressive symptoms were analysed as a categorical variable instead of a continuous score. This might have lost some refinement, but this was necessary due to highly skewed distribution. Serum 25(OH)D$_3$, D$_2$, phosphate, calcium and PTH were measured on a single occasion and this may not accurately reflect usual status (Schram et al., 2007). However, previous epidemiological studies in adults (including with bone phenotypes for which these exposures have established biological relationships) also use single measurements and for season-specific vitamin D status over a longer time, a single measure is likely to be adequate (Hofmann, Yu, Horst, Hayes, & Purdue, 2010) and serum calcium concentrations are normally maintained within relatively narrow limits in humans (Parfitt, 1987). As noted above, our results do not imply causality and the association of 25(OH)D$_3$ with depressive symptoms 3 years later might be explained by residual confounding.

Conclusions

Our results suggest that the association between 25(OH)D$_3$ concentrations and depression emerges in childhood. The association is independent of a wide range of potential confounding factors, and appears to be stronger with greater time separation between assessment of 25(OH)D$_3$ and that of depressive symptoms. Given the importance of depression in childhood and adolescence and the relative ease with which 25(OH)D$_3$ could be increased through supplementation, randomised controlled trials to assess the effectiveness of this for prevention of depressive symptoms in this age group would be appropriate.

Supporting information

Additional Supporting Information may be found in the online version of this article:

Table S1 Sample characteristics (only complete cases for depressive symptoms at age 13.8 years are included)

Table S2 Univariable associations between potential confounders and age- and gender-standardised serum 25-hydroxyvitamin D$_3$, D$_2$, phosphate, calcium and PTH concentrations

Table S3 Univariable associations between potential confounders, exposures and depressive symptoms

Table S4 Prospective association of unadjusted 25(OH)D$_3$ with depressive symptoms assessed by short Mood and Feelings Questionnaire at age 10.6 years ($n = 2,759$; exposures assessed at mean age 9.2 years) and age 13.8 years ($n = 2,752$; exposures assessed at mean age 9.8 years)

Table S5 Prospective association of total 25(OH)D [25(OH)D$_3$ + 25(OH)D$_2$] with depressive symptoms assessed by short Mood and Feelings Questionnaire at age 10.6 years ($n = 2,759$; exposures assessed at mean age 9.2 years) and age 13.8 years ($n = 2,752$; exposures assessed at mean age 9.8 years)

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Key points

- Depression in adolescence is common and early onset predicts worse outcome in adulthood.
- Previous, mainly cross-sectional studies in adults have suggested a link between higher total 25-hydroxyvitamin D [25(OH)D] concentrations and lower risk of depression.
- Findings from this first prospective study in children suggest that the linear association between 25(OH)D concentrations and depression emerges in childhood/early adolescence and is driven by the 25(OH)D$_3$ form.
- 25(OH)D$_2$ concentrations were not associated with depressive symptoms.
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