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Evaluating Vitamin D levels in Rheumatic Heart Disease patients and matched controls: A case-control study from Nepal

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Abstract

Background

Diagnosis and treatment for Rheumatic Heart Disease (RHD) is inaccessible for many of the 33 million people in low and middle income countries living with this disease. More knowledge about risk factors and pathophysiologic mechanisms involved is needed in order to prevent disease and optimize treatment. This study investigated risk factors in a Nepalese population, with a special focus on Vitamin D deficiency because of its immunomodulatory effects.

Methods

Ninety-nine patients with confirmed RHD diagnosis and 97 matched, cardiac-healthy controls selected by echocardiography were recruited from hospitals in the Central and Western region of Nepal. Serum 25(OH)D concentrations were assessed using dried blood spots and anthropometric values measured to evaluate nutritional status. Conditional logistic regression analysis was used to define association between vitamin D deficiency and RHD.

Results

The mean age of RHD patients was 31 years (range 9–70) and for healthy controls 32 years (range 9–65), with a 4:1 female to male ratio. Vitamin D levels were lower than expected in both RDH and controls. RHD patients had lower vitamin D levels than controls with a mean s-25(OH)D concentration of 39 nmol/l (range 8.7–89.4) compared with controls 45 nmol/l (range 14.5–86.7) (p-value = 0.02). People with Vitamin D insufficiency had a higher risk (OR = 2.59; 95% CI: 1.04–6.50) of also having RHD compared to people with Vitamin D
concentrations >50 nmol/l. Body mass index was significantly lower in RHD patients (22.6; 95% CI, 21.5–23.2) compared to controls (24.2; 95% CI, 23.3–25.1).

**Conclusion**

RHD patients in Nepal have lower Vitamin D levels and overall poor nutritional status compared to the non-RHD controls. Longitudinal studies are needed to explore the causality between RHD and vitamin D level. Future research is also recommended among Nepali general population to confirm the low level of vitamin D as reported in our control group.

**Introduction**

Rheumatic heart disease (RHD) affects an estimated 33 million people globally [1, 2], making it the most commonly acquired heart disease in people <25 years and almost as prevalent as human immunodeficiency virus [3]. Although the incidence of RHD in high-income countries decreased markedly during the 20th century, it remains a major public health concern in many low-and middle-income countries and marginalized communities in high-income countries—such as such, three out of four children <15 years grow up in parts of the world where RHD is endemic [1].

RHD is characterized by chronic valvular lesions and is the result of Acute Rheumatic Fever (ARF), which develops as an autoimmune response to Group A Streptococcal (GAS) infection [4]. Currently, there is no curative treatment. The secondary prophylaxis consisting of benzathine penicillin G injections every 3–4 weeks is used to prevent recurrences, which can lead to new valvular lesions or to worsening of the previous one [5].

Despite knowing the etiologic agent, there are still many unanswered questions regarding pathophysiologic mechanisms involved in disease development and progression. The presence of residual autoreactive cells appears to play a role in the persistence and worsening of the valvular lesions in RHD patients [6, 7]. Not all GAS are rheumatogenic, and not all people are susceptible to developing ARF and RHD. While it is not known what makes a host susceptible [8], it is generally accepted that malnutrition has a great impact on the immune system affecting both innate and acquired immunity in children [9]. More specifically, hypovitaminosis D has been associated with an increased risk of infections such as GAS pharyngitis as well as the risk of developing autoimmune diseases [10, 11]. Links have also been demonstrated with ARF and rheumatic mitral stenosis [12, 13].

Finally, extravasation through the valvular endothelium seems to be an important step in the valvular lesions seen in RHD, and since s-25(OH)D regulates expression of vascular endothelial growth factor, this could explain the link between hypovitaminosis D and endothelial dysfunction, and subsequently ARF and RHD [14, 15].

Nepal is situated in a region with one of the highest prevalence rates of RHD in the world [2]. A school-based study estimated prevalence of subclinical RHD to be 10.2 per 1000 children [16]. Prevalence estimates for adults are not available from Nepal, despite peak prevalence usually occurring between 25–45 years of age. Nonetheless, RHD remains endemic in Nepal and the South-Asian region. Prevalence of insufficient Vitamin D levels in Nepal vary from 59.8% amongst new mothers to 17.2% in 6-8year olds [17, 18]. In healthy school-children in neighboring India, vitamin D deficiency prevalence is estimated to 35% [19]. To date, nutritional status in ARF/RHD patients in Nepal is unknown.
In this study we examined possible associations between s-25(OH)D levels and RHD status in Nepal. In addition, associations between socioeconomic factors and RHD is also investigated.

Methods and materials

Study design and area

This case-control study was carried out in the two largest cities in Nepal, Kathmandu and Pokhara, which are located in the Central and Western region of the country. Study participants were recruited from two hospitals; Western Regional Hospital (WRH) in Pokhara and Manmohan Cardiothoracic Vascular and Transplant Center (MCVTC) in Kathmandu, between March and July 2018. Both are governmental run health facilities, and have provided free prophylactic treatment to RHD patients through the national RHD control program funded by the Nepalese government from 2007 to 2018. MCVTC also provides free cardiac surgery for RHD patients, approximately 300 cases annually. The institutions are two of the largest governmental run health facilities in the country, and hence receive patients from all regions and districts.

Subjects

Cases were selected from registries under the National ARF/RHD Prevention and Control Program when patients were seen for delivery of secondary prophylaxis, diagnostic and follow-up clinical evaluation or for surgical intervention. Patients were included based on a confirmed diagnosis of RHD by echocardiographic screening leading to registration in the national RHD Program Registry or new echocardiographic findings confirming RHD diagnosis, both in accordance with the World Heart Federation echocardiographic guidelines [5]. Exclusion criteria were issues which could interfere with Vitamin D metabolism or with the immunological conditions such as; patients below five years of age, recent hospitalization for more than one week, burn victims, chronic kidney disease and current tuberculosis or thyroid disease (self-reported).

Controls were selected by echocardiography and self-reported medical history among any people attending either institution as patients, relatives or acquaintants, matched on sex and age (maximum difference of 5 years). Exclusion criteria of controls were: diagnosis of ARF, RHD, congenital heart disease, any echocardiographic findings of valvular damage and otherwise the same exclusion criteria as for cases. Patients with congenital heart disease were excluded as they can display abnormal vitamin D concentrations [20, 21]. Cases were matched with controls from the same institution as themselves.

Collection of socio-demographic information

Face-to-face interviews were conducted with all participants using a structured questionnaire containing questions on household items and living circumstances (S1 File). Socioeconomic status was assessed by computing a wealth index using principal components analysis. This divided the participant into terciles; poorest, middle and richest. The components used in the wealth index were; ownership of a house, animals, vehicles and electronic equipment (electricity, radio, television, mobile phone, telephone, refrigerator), furniture (bed, sofa, cupboard, computer, table, chair, clock, fan, dhiki/janto), housing characteristics and fuel used for cooking. Thus, this study measures SES based on durable assets ownership and access to utilities, to accommodate the often very fluctuating income patterns seen in many low and middle income countries, including Nepal.
A food frequency questionnaire was also included (S1 File). Participants were asked to tick the number of times they had consumed the most commonly available fruit and meat, fish, soybean oil, and egg in the last week. The purpose of the fruit intake was to detect differences in variance of diet between cases and controls, and thus give an idea of the general nutritional state of the two groups. Questions regarding meat, fish, soybean oil, and egg were added due to their possible direct effect on vitamin D concentrations and RHD development, making an assessment of differences between the two groups of interest. Questions used in this study are adapted from the Nepal Demographic and Health Survey [22] and Piryani et al. [23] to ensure cultural appropriateness.

Anthropometric measurements
The following anthropometric measurements were determined: weight (kg), height (cm), age (years) and Mid-upper arm circumference (MUAC) (mm). The same weight scale and meter tape were used to measure cases and controls and measurements were performed by the same person. Shoes and heavy items of clothing were removed beforehand. Body mass index (BMI) was calculated as weight in kilogram divided by height squared in meter (kg/m$^2$). MUAC was measured on the non-dominant/left arm, except in patients with left-sided hemiplegia. Measurement was done at the mid-point between the olecranon and acromion on a relaxed arm. All participants were checked for pitting edema.

Biochemical measurements
Blood was collected as dried blood spots on Perkin Elmer 226 Five Spot RUO Card filter paper, and extracted samples analyzed at the Department of Clinical Biochemistry, Aarhus University Hospital. We measured serum 25-hydroxyvitamin D2 (25(OH)D2, ergocalciferol) and serum 25-hydroxyvitamin D3 (25(OH)D3, cholecalciferol) using liquid chromatography-tandem mass spectrometry on Sciex Triple Quad 5500 LCMSMS System calibrated using 25-OH-Vitamin D3/D2 Serum calibration standards (ChromSystems). The method is adapted from Kvaskoff et al. [24], and S-25(OH)D is expressed as the sum of 25(OH)D2 and 25(OH) D3. Vitamin D deficiency was divided into three categories following Danish national standards, defined as: Vitamin D insufficiency (VDI) as a serum concentration between 25–50 nmol/L, moderate vitamin D deficiency (VDD) as a serum concentration of 13–25 nmol/L and severe Vitamin D deficiency as concentrations <13 nmol/L. For some analysis the term hypovitaminosis was used, defined as all concentrations < 50 nmol/l.

Sample size
The prevalence of s-25(OH)D levels below 50 nmol/l was set to 77% in RHD patients based on previous studies [12]. Since data on vitamin D deficiency in Nepal is very inconsistent, we hypothesized hypovitaminosis D prevalence in non-RHD individuals to be 35%, based on results from a large study from Northern India [19]. We estimated sample size for two independent proportions with a 95% confidence level, 90% power and a margin of error (alpha) 5% to be 64; 32 cases and 32 controls.

Statistical analysis
Data were collected in hard copies and entered using REDCap electronic data capture tools. Statistical analysis was performed using Stata Statistical Software IC 15.1 (StataCorp LP, TX). Two-tailed p-values $p \leq 0.05$ were considered statistically significant.
Normality in distribution was tested using q-q plots. Normally distributed continuous variables were compared between groups using Student’s t-test and reported as means with standard deviation (SD). Differences between groups of categorical variables were compared using Chi-square test. SES was calculated by creating wealth index scores for each participant using principal components analysis (PCA), following the Measuring Equity with Nationally Representative Wealth Quintiles guide [25]. Due to the smaller sample size in this study compared to large epidemiological surveys, quintiles were converted into terciles. Multivariate conditional logistic regression analysis between sufficient Vitamin D concentrations and hypovitaminosis D were performed, adjusting for potential confounders; age, sex, BMI, education, and SES [26–32]. Potential confounders were identified a priori through a literature review. Paired 1:1 matching between cases and controls was performed after data collection, except 5 cases and 1 control who were 2:1 matched, because of the post hoc exclusion of 12 subjects (Fig 1). A sensitivity analysis of available variables before and after excluding the 12 subjects revealed no difference in major (or other) variables.

Ethical considerations
This study received ethical approval from the ethical review board of the Nepal Health Research Council (ref. no. 2398). Written information was handed out to participants and relatives in Nepal’s official language Nepali. Prior to data collection, an informed consent form had to be signed. If illiterate, oral information was given and fingerprint was used as signature. Informed assent form was obtained from participants under 18 years of age. After biochemical analysis, all results were reported back to the participants along with advice of treatment if needed. This was done by local health personnel in Nepali language.

Results
A flowchart illustrating exclusions for the different steps in data analysis is illustrated in Fig 1. The study excluded 6 cases because they were misclassified as having RHD when in fact their echocardiography report revealed no significant disease. Later, an additional 6 people were excluded before any analysis regarding s-25(OH)D concentrations were performed because these 6 had comorbidities assessed to potentially affect their s-25(OH)D concentrations. They could still be part of SES calculation, dietary intake, and other demographic measures since their comorbidities would not influence these parameters.

In total, 99 patients with confirmed RHD diagnosis and 97 matched controls were included from two different sites in Nepal. A comparison of unadjusted sociodemographic, anthropometric and socioeconomic differences is presented in Table 1.

The mean age of all participants was 31.71 years ± 11.5 and 75.5% were female. There was no significant difference between cases and control on gender and age.

BMI was significantly lower in cases than in controls, 22.4 ± 4.5 kg/m² and 24.2 ± 4.6 kg/m², respectively. Mean MUAC for cases were 253 ±34 mm and 266 ±43 mm for controls.

Almost half (47%) of cases belonged to the lowest possible SES class. For controls, this number was 21%. In the highest SES class, 25% of cases were represented compared to 40% of controls. The cases did not differ significantly on any other sociodemographic variables including overcrowding, compared to their controls.

We observed VDI and VDD in 78% of cases and mean concentration was significantly lower compared to controls. The prevalence of VDI and VDD in controls were 70%. Fig 2 displays the distribution of s-25(OH)D concentrations in all groups.

Univariate conditional logistic regression analysis revealed only association between Vitamin D deficiency (<25 nmol/l) and RHD (OR = 3.14; 95% CI: 1.02–9.64) but not for Vitamin
D insufficiency. However, after adjusting for potential confounders, multivariate analysis found a significant association (OR = 2.59; 95% CI: 1.04–6.50) for Vitamin D insufficiency but a non-significant association for Vitamin D deficiency (Table 2).

Detailed echocardiography report was available from 41 patients. Upon Chi-squared test of the report from the 41 patients available, there was no correlation between the disease stage and the presence of Vitamin D insufficiency.
Table 1. Sociodemographic characteristics of cases and controls.

|                              | RHD Patients [n = 99] | Controls [n = 97] | P for difference |
|------------------------------|-----------------------|-------------------|------------------|
| Female %                     | 77%                   | 71%               | 0.500            |
| Age (y), n (%)               |                       |                   |                  |
| 5–19                         | 13 (13)               | 12 (12)           |                  |
| 20–29                        | 28 (28)               | 28 (29)           |                  |
| 30–39                        | 40 (41)               | 36 (37)           |                  |
| 40–49                        | 10 (10)               | 12 (12)           |                  |
| >50                          | 8 (8)                 | 9 (10)            |                  |
| Mean age (y) ±SD             | 31 ± 11.4             | 32 ± 11.6         | 0.700            |
| Mean BMI (kg/m²) ±SD         | 22.2 ± 4.4            | 24.2 ± 4.6        | 0.002            |
| Mean MUAC mm ± SD            | 252 ± 34              | 266 ± 43          | 0.013            |
| Socioeconomic status, n (%)  |                       |                   | 0.001            |
| Poorest                      | 46 (47)               | 20 (21)           |                  |
| Middle                       | 27 (28)               | 38 (39)           |                  |
| Richest                      | 25 (25)               | 39 (40)           |                  |
| Number of siblings mean ± SD | 4.2 ± 2.4             | 3.8 ± 2.1         | 0.300            |
| Sisters                      | 2.3 ± 2.0             | 1.9 ± 1.5         | 0.120            |
| Brothers                     | 1.8 ± 1.4             | 1.9 ± 1.4         | 0.770            |
| Sleeping arrangement         |                       |                   | 0.959            |
| Bed                          | 83                    | 82                |                  |
| Blanket/mattress on floor    | 14                    | 15                |                  |
| People sleeping in same room (overcrowding) |                 |                   | 0.867            |
| 1–2                          | 14                    | 15                |                  |
| 3–6                          | 82                    | 79                |                  |
| 7–10                         | 2                     | 3                 |                  |
| Heart disease in family      | 11                    | 11                | 0.301            |
| RHD                          | 5                     | 5                 |                  |
| CHD                          | 2                     | 0                 |                  |
| Other                        | 4                     | 6                 |                  |
| ARF in family                | 10                    | 15                | 0.590            |
| Family type                  |                       |                   |                  |
| Nuclear                      | 29                    | 34                |                  |
| Joint                        | 61                    | 60                |                  |
| Mean s-25(OH)D (range) [n]   | 38.7 (8.7–89.4) [97]  | 44.7 (14.5–86.7) [93] | 0.014 |

Characteristics of 99 rheumatic heart disease patients and 97 controls. SD: Standard Deviation; BMI: body-mass-index, kg/m²; MUAC: middle-upper-arm circumference, mm; RHD: rheumatic heart disease; CHD: congenital heart disease; RF: rheumatic fever.

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and lower s-25(OH)D concentrations. The same was applicable when the patients without detailed echocardiographic reports were distributed into the s-25(OH)D categories considered sufficient, VDI and VDD and showed the same pattern of distribution.

Analysis of the food-frequency questionnaire revealed no statistically significant differences in intake between cases and controls (Table 3).

Discussion

Both RHD patients and controls in Nepal have a high prevalence of vitamin D deficiency and insufficiency. RHD patients on average have a significantly lower s-25(OH)D concentration compared to their controls. Especially, the prevalence of VDD was higher in RHD patients. This is to date the largest study on Vitamin D and RHD. Three other studies have examined the relationship between vitamin D concentrations and RF or RHD [12, 13, 33]. The previous studies showed a 27–59% lower concentration in patients whereas we only demonstrated a

| s-25(OH)D | OR 95% CI | P value | OR 95% CI | P value |
|-----------|-----------|---------|-----------|---------|
| >50 nmol/l | Reference |         | Reference |         |
| 50–25 nmol/l | 1.61 0.78–3.33 | 0.20 | 2.59 1.04–6.50 | 0.04* |
| <25 nmol/l | 3.14 1.02–9.64 | 0.05* | 3.62 0.93–14.09 | 0.06 |

* Statistically significant p-values.
* Adjusted for: age, BMI, sex, education, and SES.

Unadjusted (left) and adjusted (right) Odds Ratio of RHD diagnosis in relation to s-25(OH)D status. Sufficient Vitamin D status used as a reference value.
12.9% difference. However, Yusuf et al. only included RHD patients, no healthy controls, and only looked at calcification (Wilkins calcium score) of exclusively stenotic mitral valves [33]. Onan et al. only included patients with rheumatic mitral stenosis [13] and the third study only included ARF patients and none with RHD [12]. Since inclusion criteria vary between studies, it is not surprising that results also vary. Worth noting is the fact that they all have the same conclusion: s-25(OH)D is significantly lower in patients with RHD. However, it is not possible to report on the causality of this relationship. Vitamin D status can change throughout time, making it difficult to say if the lower concentration was present before the patient developed RHD. For instance, the chronically ill RHD patient may not be active and going outside as much as before falling ill, thus being less exposed to sunlight which might explain a lower s-25 (OH)D concentration. To address these questions a longitudinal study is required. Unfortunately, such data is not available from the study area, and current status is here accepted as a compromise.

Table 3. Dietary intake in the last week stratified by cases and controls.

|                        | RHD patients | Controls | P value |
|------------------------|--------------|----------|---------|
| Fruit intake, n (%)    |              |          |         |
| Not consumed           | 19 (19)      | 11 (11)  |         |
| 1 time/week            | 7 (7)        | 14 (15)  |         |
| 2–6 times/week         | 47 (48)      | 46 (47)  |         |
| > 7 times/week         | 26 (26)      | 26 (27)  | 0.216   |
| Meat, n (%)            |              |          |         |
| Not consumed           | 10 (10)      | 7 (7)    |         |
| 1 time/week            | 37 (38)      | 35 (36)  |         |
| 2–6 times/week         | 44 (45)      | 46 (47)  |         |
| > 7 times/week         | 7 (7)        | 9 (10)   | 0.832   |
| Egg, n (%)             |              |          |         |
| Not consumed           | 30 (31)      | 27 (28)  |         |
| 1 time/week            | 26 (26)      | 19 (20)  |         |
| 2–6 times/week         | 32 (33)      | 41 (42)  |         |
| > 7 times/week         | 10 (10)      | 10 (10)  | 0.503   |
| Soybean Oil, n (%)     |              |          |         |
| Not consumed           | 44 (45)      | 40 (41)  |         |
| 1 time/week            | 8 (8)        | 3 (3)    |         |
| 2–6 times/week         | 19 (19)      | 18 (19)  |         |
| > 7 times/week         | 27 (28)      | 36 (37)  | 0.287   |
| Fish, n (%)            |              |          |         |
| Not consumed           | 59 (60)      | 62 (64)  |         |
| 1 time/week            | 30 (31)      | 25 (26)  |         |
| 2–6 times/week         | 9 (9)        | 10 (10)  |         |
| > 7 times/week         | 0 (0)        | 0 (0)    | 0.750   |

Dietary intake in last one week, in rheumatic heart disease patients and controls.

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To make sense of these findings, an understanding of the pleiotropic nature of Vitamin D is important. Vitamin D receptors (VDR) and the activating enzyme 1α-hydroxylase are identified in over 30 target organs including many cells of the immune system [34]. In the innate immune system, vitamin D enhances macrophages’ phagocytic abilities and monocytes increase expression of 1α-hydroxylase and VDR through toll-like receptor signaling when encountering pathogens. The VDR complexes, in turn, activate transcription of antimicrobial cytokines [10]. Vitamin D also affects the adaptive immune system by modulation of antigen presenting cells to a more immature state, expressing fewer major histocompatibility complex class II molecules and thereby presenting fewer antigens and producing less interleukin-12, resulting in reduced activation of B and T cells. Furthermore, the modulated antigen presenting cells produce more interleukin-10—a tolerogenic cytokine [35]. Altogether, Vitamin D enhances the body’s resistance against infections while decreasing the risk of an inappropriate autoimmune response. In general, RHD patients to a greater extent suffer from malnutrition with both lower BMI and MUAC. Whether they were malnourished before becoming ill or malnourished because they were ill, is not possible to determine from this study. It could be part of a two-way causal relationship as undernutrition lower immune responses, thereby increasing the risk of infection, and infection consequently aggravating undernutrition by an increase in demands of nutrients while simultaneously decreasing appetite in the affected individual [9].

Overcrowding and social disadvantage are two of the most commonly reported risk factors for GAS infection leading to the first episode of ARF in susceptible individuals as well as resulting in recurrences in RHD patients [36]. This study also demonstrated a clear association between low SES and RHD but did not find an association with overcrowding. This could be because the mean age of patients was 31 years. Since the first acute episode of ARF occurs more frequently in the age group of 5–14 years [37], this means their housing situation could have changed since their childhood and adolescent years when overcrowding contributed for a higher risk of exposure to GAS and triggered ARF in those susceptible individuals. On the other hand, in Nepal, it is not unusual to live as a joint family throughout life, as demonstrated in Table 1.

We found slightly higher s-25(OH)D concentrations in samples collected from MCVTC compared to samples from WRH. However, geographical plottings showed that participants from both institutions represent both central and peripheral districts, and the difference between patients and controls was still present when comparing the groups within the two institutions.

No association was found between dietary intake and RHD (Table 3). This is in contrast to the general acceptance of fish and meat products as an important source of Vitamin D [38], as well as previous studies where consumption of soybean oil and egg, has been shown to suppress the rheumatic process because of their high concentrations of phospholipids and palmtamid [39].

Finally, we would like to highlight that the majority of RHD patients were young women. Though not surprising, it is still an important aspect that raises concern especially since the age of the women coincides with childbearing age. Having a heart disease, including RHD, whilst becoming pregnant can lead to serious complications [40]—particularly when access to health care is limited. In fact, RHD has been suggested a leading cause of indirect obstetric death in some sub-Saharan countries [30]. Yet no one has been able to explain why women are more often affected by RHD than men [27, 28, 30]. It is worth noting that this difference is not present when comparing ARF prevalence. Explanations to this sex-based difference could include extrinsic factors such as reduced access to primary and secondary ARF prophylaxis for girls and culturally rooted disadvantages affecting female health in general. Additionally,
intrinsic factors increasing host susceptibility such as immunogenic, genetic and in particular hormonal differences, should also be considered and investigated further.

This study has some limitations. The major limitation being the usage of current vitamin D status as a proxy for previous concentrations. Furthermore, the higher than expected prevalence of hypovitaminosis D in the background population reduces the power of the study. However, the impact was limited by including 3 times as many participants than required in the sample size calculations. Nonetheless, the sample size is relatively small. This should also be considered when understanding the measurement of associations. While some are statistically significant, they are largely marginal. This reduces the immediate clinical applicability of the results, but should be seen in light of the aforementioned smaller sample size. Increasing the sample size for future studies could help make the associations clearer.

The dried blood spot sampling procedure for vitamin D concentration was validated twice—once before and once after data collection—at Aarhus University Hospital, using the exact same method as in this study, hence the sampling procedure should not be a considerable source of error. Furthermore, all anthropometric data were measured upon inclusion and not self reported or taken from previous examinations, adding value to the results.

However, the lack of systematic record keeping in the Nepalese health system made identification of comorbidities a challenge. All participants were questioned about their medical history and whenever possible, their record books were examined thoroughly, but on many occasions, the patients did not bring their record books, which could result in underreporting of comorbidities that might affect s-25(OH)D concentrations. Additionally, there is a risk of recall bias when conducting a survey, especially the food frequency intake should be interpreted with caution.

In light of the abovementioned limitations and the very nature of the study design, we suggest caution when generalizing to other populations, but instead advise more research in the field of vitamin D deficiencies especially in low income countries, and how it affects immune responses.

Conclusion

Both Vitamin D Insufficiency (VDI) and Vitamin D Deficiency (VDD) is highly prevalent among Rheumatic Heart Disease (RHD) patients in Nepal. RHD patients presented with significantly lower mean s-25(OH)D concentrations and overall poor nutritional status compared to the non-RHD controls. People with Vitamin D insufficiency had a higher risk (OR = 2.59) of also having RHD, underlining the potential of hypovitaminosis D being either a risk factor or feature of RHD, but longitudinal studies are needed to explore the causality of this relationship further. Larger studies among the Nepali population is also recommended to confirm the high prevalence of hypovitaminosis D found in our control group.

Other potential risk factors found in this study include low BMI, low mid upper arm circumference, low socioeconomic status, and female sex.

Supporting information

S1 Checklist. STROBE checklist. (DOCX)

S1 File. Questionnaire used to collect information on demographic variables and for wealth index score analysis. (DOCX)
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References

1. Watkins DA, Johnson CO, Colquhoun SM, Karthikeyan G, Beaton A, Bukhman G, et al. Global, Regional, and National Burden of Rheumatic Heart Disease, 1990–2015. N Engl J Med. 2017; 377 (8):713–22. Epub 2017/08/24. https://doi.org/10.1056/NEJMoa1603693 PMID: 28834488
2. World Health Organization. Rheumatic fever and rheumatic heart disease—resolution A71/25 2018. https://apps.who.intiris/bitstream/handle/10665/276479/A71_25-en.pdf?sequence=1&isAllowed=y.
3. UNAIDS. Prevention Gap Report 2016. http://www.unaids.org/sites/default/files/media_asset/2016-prevention-gap-report_en.pdf.
4. Sika-Paotonu D, Beaton A, Raghu A, Steer A, Carapetis J. Acute Rheumatic Fever and Rheumatic Heart Disease. In: Ferretti JJ, Stevens DL, Fischetti VA, editors. Streptococcus pyogenes: Basic Biology to Clinical Manifestations. Oklahoma City (OK)2016.
5. Remenyi B, Wilson N, Steer A, Ferreira B, Kado J, Kumar K, et al. World Heart Federation criteria for echocardiographic diagnosis of rheumatic heart disease—an evidence-based guideline. Nat Rev Cardiol. 2012; 9(5):297–309. Epub 2012/03/01. https://doi.org/10.1038/nrcardio.2012.7 PMID: 22371105
6. Guilherme L, Kaill J. Rheumatic fever and rheumatic heart disease: cellular mechanisms leading auto-immune reactivity and disease. J Clin Immunol. 2010; 30(1):17–23. Epub 2009/10/06. https://doi.org/10.1007/s10875-009-9332-9 PMID: 19802690.

7. Fae KC, da Silva DD, Oshiro SE, Tanaka AC, Pomerantzeff PM, Douay C, et al. Mimicry in recognition of cardiac myosin peptides by heart-intralesional T cell clones from rheumatic heart disease. J Immunol. 2006; 176(9):5662–70. Epub 2006/04/20. https://doi.org/10.4049/jimmunol.176.9.5662 PMID: 16622036.

8. Kerdemelidis M, Lennon DR, Arroll B, Peat B, Jarman J. The primary prevention of rheumatic fever. J Paediatr Child Health. 2010; 46(9):534–48. Epub 2010/09/22. https://doi.org/10.1111/j.1440-1754.2010.01854.x PMID: 20854326.

9. Ryter MJ, Kolte L, Briand A, Fries H, Christensen VB. The immune system in children with malnutrition—a systematic review. PLoS One. 2014; 9(8):e105017. Epub 2014/08/26. https://doi.org/10.1371/journal.pone.0105017 PMID: 25153531.

10. Prietl B, Treiber G, Pieber TR, Amrein K. Vitamin D and immune function. Nutrients. 2013; 5(7):2502–10. https://doi.org/10.3390/nu5072502 PMID: 23857223.

11. Nseir W, Mograb J, Abu-Rahmeh Z, Mahamid M, Abu-Elheja O, Shalata A. The association between vitamin D levels and recurrent group A streptococcal tonsillitis in adults. Int J Infect Dis. 2012; 16(10):e735–8. Epub 2012/07/31. https://doi.org/10.1016/j.ijid.2012.05.1036 PMID: 22841558.

12. Onan SH, Demirbilek H, Aldudak B, Bilici M, Demir F, Yilmazer MM. Evaluation of vitamin D levels in patients with acute rheumatic fever. Anatol J Cardiol. 2017; 18(1):75–6. Epub 2017/07/07. https://doi.org/10.14744/anatoljcardiol.2017.7720 PMID: 28680013.

13. Yavuz B, Sen O, Deveci OS, Akin KO, Dal K, Ata N, et al. Serum 25-hydroxy vitamin D levels are correlated with mitral valve calcification score in patients with rheumatic mitral stenosis. J Heart Valve Dis. 2012; 21(5):570–5. Epub 2012/11/22. PMID: 23167220.

14. Roberts S, Kosanke S, Terrence Dunn S, Jankelow D, Duran CM, Cunningham MW. Pathogenic mechanisms in rheumatic carditis: focus on valvular endothelium. J Infect Dis. 2001; 183(3):507–11. Epub 2001/01/03. https://doi.org/10.1086/318076 PMID: 11133385.

15. Sarkar S, Chopra S, Rohit MK, Banerjee D, Chakraborti A. Vitamin D regulates the production of vascular endothelial growth factor: A triggering cause in the pathogenesis of rheumatic heart disease? Med Hypotheses. 2016; 95:62–6. Epub 2016/10/04. https://doi.org/10.1016/j.mehy.2016.09.001 PMID: 27692170.

16. Shrestha NR, Karki P, Mahro R, Gurung K, Pandey N, Agrawal K, et al. Prevalence of Subclinical Rheumatic Heart Disease in Eastern Nepal: A School-Based Cross-sectional Study. JAMA Cardiol. 2016; 1 (1):89–96. Epub 2016/07/22. https://doi.org/10.1001/jamacardio.2015.0292 PMID: 27437661.

17. Haugen J, Ulak M, Chandyi D, Henjum S, Thorne-Lyman AL, Ueland PM, et al. Low Prevalence of Vitamin D Insufficiency among Nepalese Infants Despite High Prevalence of Vitamin D Insufficiency among Their Mothers. Nutrients. 2016; 8(12). Epub 2016/12/24. https://doi.org/10.3390/nu8120825 PMID: 28008810.

18. Schulze KJ, Christian P, Wu LS, Arguello M, Cui H, Nanayakkara-Bind A, et al. Micronutrient deficiencies are common in 6- to 8-year-old children of rural Nepal, with prevalence estimates modestly affected by inflammation. J Nutr. 2014; 144(6):979–87. Epub 2014/04/20. https://doi.org/10.3945/jn.114.192336 PMID: 24744314.

19. Marwaha RK, Tandon N, Reddy DR, Aggarwal R, Singh R, Sawhney RC, et al. Vitamin D and bone mineral density status of healthy schoolchildren in northern India. Am J Clin Nutr. 2005; 82(2):477–82. Epub 2005/08/10. https://doi.org/10.1093/ajcn.82.2.477 PMID: 16087996.

20. Izumi G, Inai K, Shimada E, Nakanishi T. Vitamin D Kinetics and Parathyroid Gland Function in Patients with Congenital Heart Disease. Congenit Heart Dis. 2016; 11(6):700–6. Epub 2016/07/05. https://doi.org/10.1111/chd.12389 PMID: 27375053.

21. Passeri E, Rigolini R, Costa E, Verdelli C, Arcidiacono C, Carminati M, et al. Serum NT-proBNP Levels Are Not Related to Vitamin D Status in Young Patients with Congenital Heart Defects. Dis Markers. 2016; 2016:3970284. Epub 2016/03/10. https://doi.org/10.1155/2016/3970284 PMID: 26955207.

22. Ministry of Health Nepal. Nepal Demographic and Health Survey 2016 2017. https://www.dhsprogram.com/pubs/pdf/fr336/fr336.pdf.

23. Piryani S, Baral KP, Pradhan B, Poudyal AK, Piryani RM. Overweight and its associated risk factors among urban school adolescents in Nepal: a cross-sectional study. BMJ Open. 2016; 6(5):e010335. Epub 2016/05/22. https://doi.org/10.1136/bmjopen-2015-010335 PMID: 27207624.

24. Kvaskoff D, Ko P, Simila HA, Eyles DW. Distribution of 25-hydroxyvitamin D3 in dried blood spots and implications for its quantitation by tandem mass spectrometry. J Chromatogr B Analyst Technol Biomed Life Sci. 2012; 901:47–52. Epub 2012/06/26. https://doi.org/10.1016/j.chrombi.2012.05.040 PMID: 22727750.
25. Fry K FR, Chakraborty N.M. Measuring Equity with Nationally Representative Wealth Quintiles Washington DC: PSI2014. https://www.psi.org/wp-content/uploads/2014/10/Wealth-Quintile-Guide.pdf.
26. Parva NR, Tadepalli S, Singh P, Qian A, Joshi R, Kandala H, et al. Prevalence of Vitamin D Deficiency and Associated Risk Factors in the US Population (2011–2012). Cureus. 2018; 10(6):e2741. Epub 2018/08/09. https://doi.org/10.7759/cureus.2741 PMID: 30087817
27. Zuhlke L, Engel ME, Karthikeyan G, Rangarajan S, Mackie P, Cupido B, et al. Characteristics, complications, and gaps in evidence-based interventions in rheumatic heart disease: the Global Rheumatic Heart Disease Registry (the REMEDY study). Eur Heart J. 2015; 36(18):1115–22a. Epub 2014/11/27. https://doi.org/10.1093/eurheartj/ehu449 PMID: 25425448
28. Marjon E, Mirabel M, Celermajer DS, Jouven X. Rheumatic heart disease. Lancet. 2012; 379(9819):953–64. Epub 2012/03/13. https://doi.org/10.1016/S0140-6736(11)61171-9 PMID: 22405798.
29. Gurney JK, Stanley J, Baker MG, Wilson NJ, Sarfati D. Estimating the risk of acute rheumatic fever in New Zealand by age, ethnicity and deprivation. Epidemiol Infect. 2016; 144(14):3058–67. Epub 2016/06/06. https://doi.org/10.1017/S0950268816001291 PMID: 27311633.
30. Carapetis JR, Beaton A, Cunningham MW, Guilherme L, Karthikeyan G, Mayosi BM, et al. Acute rheumatic fever and rheumatic heart disease. Nat Rev Dis Primers. 2016; 2:15084. Epub 2016/05/18. https://doi.org/10.1038/nrdp.2015.84 PMID: 27188830
31. Hansel L, Tjonneland A, Koster B, Brot C, Andersen R, Cohen AS, et al. Vitamin D Status and Seasonal Variation among Danish Children and Adults: A Descriptive Study. Nutrients. 2018; 10(11). Epub 2018/11/23. https://doi.org/10.3390/nu10111801 PMID: 30463277
32. Pereira-Santos M, Costa PRF, Assis AMO, Santos CAST, Santos DB. Obesity and vitamin D deficiency: a systematic review and meta-analysis. Obes Rev. 2015; 16(4):341–9. https://doi.org/10.1111/obr.12239 PMID: 25688659
33. Yusuf J, P J, Mukhopadhayya S, Vignesh V, Tyagi S. Evaluation of serum 25-hydroxyvitamin D levels in calcific rheumatic mitral stenosis- A cross sectional study. Indian Heart J. 2018; 70(2):206–13. Epub 2018/05/03. https://doi.org/10.1016/j.ihj.2017.06.010 PMID: 29716693.
34. Reichel H, Koefler HP, Norman AW. The role of the vitamin D endocrine system in health and disease. N Engl J Med. 1989; 320(15):980–91. Epub 1989/04/13. https://doi.org/10.1056/NEJM198904133201506 PMID: 2648151.
35. Baeke F, Takiishi T, Korf H, Gysemans C, Mathieu C. Vitamin D: modulator of the immune system. Curr Opin Pharmacol. 2010; 10(4):482–96. Epub 2010/04/30. https://doi.org/10.1016/j.coph.2010.04.001 PMID: 20427238.
36. Coffey PM, Ralph AP, Krause VL. The role of social determinants of health in the risk and prevention of group A streptococcal infection, acute rheumatic fever and rheumatic heart disease: A systematic review. PLoS Negl Trop Dis. 2018; 12(6):e0006577. Epub 2018/06/14. https://doi.org/10.1371/journal.pntd.0006577 PMID: 29897915
37. Lawrence JG, Carapetis JR, Griffiths K, Edwards K, Condon JR. Acute rheumatic fever and rheumatic heart disease: incidence and progression in the Northern Territory of Australia, 1997 to 2010. Circulation. 2013; 128(5):492–501. Epub 2013/06/25. https://doi.org/10.1161/CIRCULATIONAHA.113.001477 PMID: 23794730.
38. Schmid A, Walther B. Natural vitamin D content in animal products. Adv Nutr. 2013; 4(4):453–62. Epub 2013/07/17. https://doi.org/10.3945/an.113.003780 PMID: 23858093
39. Zaman MM, Yoshiike N, Chowdhury AH, Nakayama T, Yokoyama T, Faruque GM, et al. Nutritional factors associated with rheumatic fever. J Trop Pediatr. 1998; 44(3):142–7. Epub 1998/07/29. https://doi.org/10.1093/tropej/44.3.142 PMID: 9680778.
40. French KA, Poppas A. Rheumatic Heart Disease in Pregnancy: Global Challenges and Clear Opportunities. Circulation. 2018; 137(8):817–9. Epub 2018/02/21. https://doi.org/10.1161/CIRCULATIONAHA.118.033465 PMID: 29459467.