Environmental enteric dysfunction (EED) is a chronic subclinical condition, characterized by limited growth and absorption, and systemic inflammation [7]. EED pathology is concerned with small intestine structure and function, which affect the macro- and micronutrients absorption with consequent growth faltering.

AIM: This study aimed to evaluate some serum biomarkers involved in EED and determine their association with stunting and faltering growth in children; zonulin, endotoxin core antibody (EndoCAb), high-sensitive C-reactive protein (hsCRP), alpha-1-acid glycoprotein (AGP), and tumor necrosis factor (TNF), serum iron, and Vitamins A and D.

PATIENTS AND METHODS: This case–control study enrolled 105 children aged from 1 to 10 years old, having weight-for-age z-scores and height-for-age z-scores (WAZ or HAZ) ranging from −1.5 to −2. They were compared with control group consisted of 100 children having WAZ or HAZ > −1 of matched age and sex. Assessment of serum markers levels of enteric dysfunction (zonulin and EndoCAb), markers of systemic inflammation (hs CRP and AGP), along with serum micronutrients (vitamin A, vitamin D and iron) in children with malnutrition in comparison to controls.

RESULTS: There was a highly significant decrease as regarding the anthropometric measurements; weight, height, BMI, and arm circumference. Moreover, significant increase in serum zonulin, EndoCAb, HsCRP, and AGP and highly significant decrease of serum Vitamin D and iron in cases group as compared to control group. Height Z score showed negative correlation with zonulin, HsCRP, and AGP and positive correlation with Vitamin D. Weight Z score showed negative correlation with zonulin, HsCRP, and AGP and positive correlation with Vitamin D and Vitamin A. Regression analysis noted increase of zonulin and α1AGP as high associative markers with height Z score affection, however, increase of zonulin was high associative markers with weight Z score affection.

CONCLUSION: Faltering growth is associated with elevated serum systemic markers of intestinal inflammation (HsCRP and α1AGP), EED may be a cause of faltering growth.
with adulthood non-infectious diseases (hyperlipidemia, hypertension, insulin resistance, obesity, cardiovascular diseases, and diabetes mellitus [DM]) [10], [11].

There is evident link and relation between early stunting growth at early infant life and later consequence of impaired health, poor cognitive function, intellectual performance, and academic achievement [12], [13].

The criteria for diagnosis of EED have not been established yet. The golden diagnostic confirmatory method of EED is endoscopy and small intestinal biopsy which is considered as invasive, cost, and need technical experience, therefore, it is usually avoided measure [14]. Consequently, researches are challenging the usage of less invasive available biomarkers that emerged from the underlying mechanism of EED pathology to help in diagnosis.

Fecal and serum biomarkers of inflammation are available. However, no single biomarker or group of biomarkers has been thoroughly validated or documented to capture EED [7].

Highly sensitive C-reactive protein (hsCRP) and alpha-1-acid glycoprotein (AGP) and tumor necrosis factor (TNF) belong to acute-phase proteins family that produced from the liver in response of infectious inflammation and other non-infectious inflammatory states. These biomarkers are considered as systemic markers of intestinal inflammation [1], [15].

The antibody of endotoxin core is a marker of microbial translocation. It is produced by B cells in response to repeated bacterial enteric infections. It measures antibodies against endotoxin lipopolysaccharide (LPS) of Gram-negative bacteria that have translocated across a leaky gut [2], [5], [7].

Zonulin is a human analog of zonula occludens toxin which is known to affect tight junctions between enterocytes, it causes an intracellular signaling cascade that results in opening of tight junctions, and its elevation indicates increased intestinal permeability [2], [5], [16].

There is rising need to search for validated and approved biomarkers that can help in diagnosis and predict implications of EED as well as to evaluate management and enhance the prognosis.

The objectives of the study were to measure a group of potential available serum biomarkers involved in EED and determine their association with stunting and faltering growth in children.

Subjects and Methods

This is a case–control study conducted in 2020. It has randomly enrolled 105 children with moderate malnutrition, having weight-for-age z-scores and height-for-age z-scores (WAZ or HAZ) ranging from −1.5 to −2. They were of both sexes, aged from 1 to 10 years old, attending child health Clinic in Medical and Scientific Centre of Excellence, National Research Center (NRC), Cairo, Egypt. One hundred healthy children of matched age and sex, having WAZ and HAZ >-1 were enrolled in the study as controls.

Exclusion criteria

Children with chronic debilitating illness including (congenital heart diseases, chronic renal problems, and neurological or developmental disabilities), congenital abnormalities or genetic disorders, recent history or present case of diarrhea or hematochezia, and heavy parasitic infestation at time of inclusion (was excluded by stool analysis).

This study is a part of the in-house project funded by NRC and approved by the Medical Ethical Committee of the NRC (19/227). All participants were informed about the objectives of the study and ready to participate with obtaining a written informed consent from the parents.

All of them were subjected to; full history taking laying stress on personal data involving date of birth, sex, familial history of short stature or wasting, chronic illness such as DM, HTN, drug intake, and dietary history.

For all children participated in the study, complete clinical examination was done laying stress on signs of vitamins deficiency and anemia and local systemic examination of all body organs. All anthropometric measurements were taken according to techniques described in the Anthropometric Standardization Reference Manual [17]. Weight was measured using a calibrated Seca scale to the nearest 0.1 kg (Seca, Hamburg, Germany), height was measured using Seca 225 stadiometer to the nearest 0.1 cm [18]. Mid-arm circumference was measured using flexible graduated tape at the left upper arm in point midway between the tip of humorous and the tip of elbow. Height and weight, BMI were applied to AnthroPlus Pediatric calculator for personal computers to calculate Z-scores for many somatic parameters [19].

The children were defined as malnourished when they were underweight (WAZ) score < −2 or stunted growth (HAZ) score < −2 or wasted (WHZ) score < −2 and as nourished when the WAZ/HAZ/WHZ score was ≥ −2 based on the WHO standard [20].

Laboratory investigations

Three milliliters of fasting (8 h) venous blood samples will be withdrawn from each child. The venous sample was collected in a Vacutainer blood collection tube. The samples were divided into two parts: The first part was added to tube containing ethylenediaminetetraacetic acid for complete blood
count. The second part put in a serum separator tube and then centrifuged for 10 min at 3000 rpm to separate the serum from the remaining components. Serum was preserved in sterile Eppendorf tubes and stored in ~80°C.

Serum iron was quantified using the colorimetric CAB method. Kit was obtained from Egyptian Company for Biotechnology (S.A.E) Obour City industrial area, block 20008, Cairo, Egypt.

Serum zonulin, endotoxin core antibody (EndoCAb), hsCRP, AGP, and TNF were quantified in serum by an ELISA method using kits purchased from (SunLong Biotech Co., LTD). Zonulin kit detection range was 30–1500 pg/ml catalog number: SL2712Hu. The EndoCAb kit detection range was 3–120 pg/ml catalog number: SL3521Hu. The AGP kit detection range was 1–80 ng/ml catalog number: SL1845Hu. The TNF kit detection range was 20–400 ng/L catalog number: SL1761Hu.

Serum Vitamin A and Vitamin D were measured by ELISA technique using commercially available ELISA kits (SunLong Biotech Co., LTD). Vitamin A kit detection range was 50–1800 ng/ml catalog number: SL1826Hu. Vitamin D kit detection range was 0.8–50 ng/ml catalog number: SL1831Hu.

### Statistical analysis

Data were collected, verified, coded, and analyzed using the Statistical Package for the Social Sciences (SPSS) version 23 (SSPS Inc., Pennsylvania, USA). Descriptive statistics were used to summarize baseline characteristics of the study population. Frequencies were reported for categorical variables while the mean ± standard deviation (SD) or standard error for continuous variables. Independent t-test was used to compare between two groups regarding quantitative data, respectively. Pearson correlation analysis was used to assess the relation between two quantitative parameters in the same group. Logistic regression was done to determine factors mostly affecting weight Z score and height Z score among cases using backward Wald. The p-values were two tailed, it was considered statistically significant at p ≤ 0.05 and highly significant at p ≤ 0.001.

### Results

This case–control study enrolled 105 children as cases compared with 100 healthy age- and sex-matched children as controls, as no significant difference between the two groups (p > 0.05) (Table 1).

There was a highly significant decrease in cases group (p < 0.001) as regarding the anthropometric measurements; weight, height, BMI, and arm circumference (Table 1).

Assessment of serum biomarkers of enteric dysfunction in EED showed significant increase in serum zonulin (biomarker of enteric permeability) and EndoCAb (biomarker of microbial translocation) in cases group as compared to control group (p ≤ 0.05). Meanwhile, there was significant increase in systemic inflammatory markers of hsCRP and AGP in cases group as compared to controls group (p ≤ 0.05). On the other hand, there was a highly significant decrease of serum Vitamin D and iron in cases group as compared to control group (p ≤ 0.001), while serum Vitamin A and Hb showed no significant difference between the two groups (p > 0.05) (Table 2).

### Coefficient correlation between laboratory markers and anthropometric parameter showed significant negative correlation between EndoCAb (biomarker of microbial translocation) and weight and height (p ≤ 0.05). Zonulin (biomarker of enteric permeability) has significant negative correlation with weight Z score and height Z score (p ≤ 0.05). hsCRP (systemic acute phase reactant) has significant negative correlation with weight, WAZ, and arm circumference (p ≤ 0.05), and highly significant negative correlation with height and HAZ (p ≤ 0.001). AGP (systemic acute-phase reactant) has significant negative correlation with weight Z score and height Z score (p ≤ 0.05). TNF-α has no significant correlation of any of the anthropometric parameter (p > 0.05). On the other hand, Vitamin A has significant positive correlation with weight Z score and BMI Z score (p ≤ 0.05). Vitamin D has significant positive correlation with weight Z score and height Z score (p ≤ 0.05), meanwhile, iron showed no significant correlation of any of the anthropometric parameter (p > 0.05) (Table 3).

### Table 1: Comparison between cases and controls groups regarding to age, sex, and anthropometric measurements

| Variable | Cases, n = 105 Mean ± SE | Control, n = 100 Mean ± SE | p-value |
|----------|--------------------------|----------------------------|---------|
| Age (months) | 79.71 ± 34.3 | 73.33 ± 39.5 | 0.85 |
| Sex (n %) | Male 47 (44.8%) | 46 (46.0%) | 0.859 |
| Weight (kg) | 17.56 ± 5.7 | 25.18 ± 11.0 | <0.001** |
| Weight Z score | -2.08 ± 0.8 | 0.4 ± 0.4 | <0.001** |
| Height (cm) | 107.94 ± 15.9 | 119.91 ± 21.3 | <0.001** |
| Height Z score | -2.18 ± 0.8 | 0.44 ± 0.4 | <0.001** |
| BMI | 15.14 ± 1.5 | 16.71 ± 1.5 | <0.001** |
| BMI Z score | -0.74 ± 1.2 | 0.34 ± 0.4 | <0.001** |
| Arm circumference (cm) | 16.56 ± 1.8 | 17.73 ± 2.0 | <0.001** |

### Table 2: Comparison between cases and controls regarding to laboratory results

| Variable | Cases, n = 105 Mean ± SE | Control, n = 100 Mean ± SE | p-value |
|----------|--------------------------|----------------------------|---------|
| Hb | 10.9 ± 0.6 | 11.2 ± 0.9 | 0.4 |
| Iron | 78.24 ± 3.1 | 104.78 ± 5.4 | <0.001** |
| Vitamin A (Pg/ml) | 63.26 ± 2.2 | 68.02 ± 1.3 | 0.884 |
| Vitamin D (Pg/ml) | 10.19 ± 0.9 | 13.5 ± 0.6 | <0.001** |
| hsCRP (mg/L) | 4.73 ± 0.4 | 1.61 ± 0.2 | 0.031* |
| AGP (mg/dl) | 17.11 ± 1.4 | 12.48 ± 0.9 | 0.924* |
| TNF (ng/L) | 44.45 ± 2.6 | 39.2 ± 2.2 | 0.567 |
| Zonulin (pg/ml) | 619.3 ± 15.4 | 463.2 ± 12.2 | 0.02* |
| EndoCAb (Pg/ml) | 15.19 ± 1.3 | 10.48 ± 0.5 | 0.05* |

Independent t-test and Chi-square test. **p ≤ 0.001 (highly significant), *p ≤ 0.05 (significant). hsCRP: High-sensitive C-reactive protein, AGP: Alpha-1-acid glycoprotein, TNF: Tumor necrosis factor.
Table 3: Correlation between anthropometric parameter and laboratory markers

| Variable | hsCRP (mg/l) | Zonulin (ng/ml) | AGP (ng/ml) | TNF (ng/mL) | Vitamin A (Pg/ml) | EndoCAb (Pg/ml) | Vitamin D (Pg/ml) | Iron (µg/dl) |
|----------|--------------|-----------------|-------------|-------------|-------------------|----------------|------------------|-------------|
| Weight (kg) |              |                 |             |             |                   |                |                  |             |
| R        | -0.473*      | -0.135          | -0.138      | -0.137      | -0.150            | -0.276**       | 0.117            | 0.023       |
| P        | 0.002        | 0.141           | 0.160       | 0.162       | 0.126             | 0.004          | 0.236            | 0.817       |
| Weight z score |            |                 |             |             |                   |                |                  |             |
| R        | -0.375*      | -0.412*         | -0.279*     | 0.017       | 0.273*            | -0.143         | 0.300**          | -0.032      |
| P        | 0.008        | 0.002           | 0.004       | 0.863       | 0.005             | 0.144          | 0.002            | 0.749       |
| Height (cm) |             |                 |             |             |                   |                |                  |             |
| R        | -0.447**     | -0.127          | -0.109      | -0.089      | -0.181            | -0.275*        | 0.099            | -0.009      |
| P        | 0.000        | 0.273           | 0.269       | 0.368       | 0.064             | 0.004          | 0.313            | 0.929       |
| Height z score |          |                 |             |             |                   |                |                  |             |
| R        | -0.385**     | -0.358*         | -0.289*     | 0.098       | 0.065             | -0.177         | 0.237*           | -0.133      |
| P        | 0.001        | 0.041           | 0.003       | 0.321       | 0.508             | 0.071          | 0.015            | 0.176       |
| BMI      |              |                 |             |             |                   |                |                  |             |
| R        | -0.172       | -0.182          | -0.144      | -0.159      | 0.134             | -0.118         | 0.105            | 0.042       |
| P        | 0.056        | 0.054           | 0.143       | 0.105       | 0.172             | 0.232          | 0.287            | 0.669       |
| BMI z score |             |                 |             |             |                   |                |                  |             |
| R        | -0.037       | -0.162          | -0.076      | -0.124      | 0.201*            | -0.084         | 0.021            | 0.020       |
| P        | 0.463        | 0.115           | 0.441       | 0.206       | 0.040             | 0.396          | 0.834            | 0.837       |
| Arm circumference (cm) |            |                 |             |             |                   |                |                  |             |
| R        | -0.395*      | -0.062          | -0.086      | -0.130      | -0.150            | -0.095         | 0.121            | -0.108      |
| P        | 0.050        | 0.713           | 0.384       | 0.187       | 0.126             | 0.334          | 0.218            | 0.272       |

**p ≤ 0.001 (highly significant), *p ≤ 0.05 (significant). AGP: Alpha-1-acid glycoprotein.**

Logistic regression analysis for predicting factors for weight Z score (at level <=-2) revealed association with zonulin, α1AGP, and Vitamin D. Its high lightened zonulin as the most significant predictor marker for weight Z score affection (adjusted odds ratio [AOR] 1.065; 95% C.I.: 1.025, 1.060) (Table 4).

Table 4: Logistic regression analysis for factors affecting weight Z score at level of <=-2

| Model | OR    | S.E. | Wald | p   | AOR 95% C.I. for AOR* | Lower | Upper |
|-------|-------|------|------|-----|-----------------------|-------|-------|
| Zonulin | -0.031 | 0.009 | 12.947 | 0.000** | 1.065 | 1.025 | 1.060 |
| AGP   | -0.049 | 0.020 | 5.937  | 0.05**  | 0.952 | 0.915 | 0.990 |
| Vitamin D | 0.118 | 0.035 | 11.425 | 0.001** | 1.125 | 1.051 | 1.204 |
| Iron  | 0.025  | 0.003 | 2.934  | 0.143 | 0.737 | 0.711 | 0.756 |
| EndoCAb | -0.035 | 0.006 | 2.473  | 0.156 | 0.852 | 0.833 | 0.871 |
| Constant | -0.841 | 0.714 | 1.388  | 0.239 | 0.431 |       |       |

**p ≤ 0.001 (highly significant), *p ≤ 0.05 (significant). AGP: Alpha-1-acid glycoprotein.**

Meanwhile, linear stepwise regression analysis showed zonulin and α1AGP as the most significant predictor marker for height Z score affection (at level <=-2) (AOR 1.042; 95% C.I.: 1.020, 1.065) (AOR 0.935; 95% C.I.: 0.900, 0.970), respectively (Table 5).

Table 5: Logistic regression analysis for factors affecting height Z score at level of <=-2

| Model | OR    | S.E. | Wald | p   | AOR 95% C.I. for AOR* | Lower | Upper |
|-------|-------|------|------|-----|-----------------------|-------|-------|
| Zonulin | -0.037 | 0.005 | 16.184 | 0.000** | 1.042 | 1.020 | 1.055 |
| AGP   | -0.066 | 0.019 | 12.613 | 0.000** | 0.935 | 0.900 | 0.970 |
| Vitamin D | 0.053 | 0.028 | 3.768  | 0.052 | 1.055 | 0.999 | 1.113 |
| Iron  | 0.0362 | 0.009 | 2.152  | 0.110 | 0.755 | 0.730 | 0.780 |
| EndoCAb | -0.043 | 0.017 | 2.376  | 0.098 | 0.843 | 0.821 | 0.860 |
| Constant | -1.674 | 0.859 | 6.447  | 0.019 | 0.970 |       |       |

**p ≤ 0.001 (highly significant), *p ≤ 0.05 (significant). AGP: Alpha-1-acid glycoprotein.**

Discussion

EED is subclinical chronic condition, often without gastrointestinal symptoms. It created a state of inefficient nutrients absorption and chronic longstanding inflammation which negatively worse EED and aggravates the condition. Causes and consequences are acting bidirectional in vicious circle pattern initiating and maintaining EED.

Malnutrition in young children has long term comorbidities in old children and later adulthood. Stunting is impaired linear growth, it is defined as a height-for-age z-score ≤ -2 SD of the median height of the WHO reference population "wasting, is defined as reduction of the tissue mass ≤ -2 SD of the mean in weight-for-height z scores [21]." Stunting usually presents in the early infancy period that may passed clinically neglected until late childhood. Untreated or improper nutrition may lead to permanent irreversible condition.

This case–control study enrolled 100 cases aged 1–10 years, suffering from malnutrition, and compared with 105 healthy children as controls. The mean age of our study group ranged between 6 and 7 years old, anthropometric measurements were highly significantly affected in cases group compared to controls group.

Several biomarkers have been carried out in EED diagnosis and prognosis. This study concerned with serum zonulin as indicator for intestinal permeability and absorption, serum EndoCAb as indicator for microbial translocation. Also, some serum markers for systemic inflammation as hsCRP, AGP, and TNF have been investigated.

Zonulin affects tight junctions between enterocytes producing increased intestinal permeability [5]. EndoCAb is antibody released against the LPS of Gram-negative bacteria [7].

This study revealed significant increase in serum zonulin and EndoCAb biomarkers in cases group more than the controls group.

Correlating the zonulin and EndoCAb in cases group with anthropometric parameters showed that both of them had significant negative correlation with weight z score and height z score, which is similar to the findings of Guerrant et al. [22] who found negative correlation between serum zonulin and stunting in a study done in Northeast Brazil on 375 child aged 6–26 months suffering from wasting or stunting. Meanwhile, logistic regression analysis in our study noted increase of zonulin as high associative markers with weight Z score and height Z score affection.
Furthermore, in a study done by Uddin et al., 2021, the basic level of EndoCAb was assessed in children at 3 months of age with follow-up at 18 months, they found that the level of EndoCAb is elevated at the age of 3 months in the group of children who were subsequently underweight or stunted more than those who were well nourished. The same was similar to the finding of Campbell et al., 2003, who found EndoCAb correlated with growth retardation in the follow-up of the infants from 2 to 15 months on rural Gambian infants [23], [24].

In another study, EndoCAb has been assessed in multiplicity of diarrheal attacks and enteropathogenic infection as there was an association between serum EndoCAb antibodies with stunting at 1 year of age [25].

However, other results were not in line with ours, the increased EndoCAb was directly related with increased baseline HAZ [26]; no difference in EndoCAb levels was found in cases and controls groups that followed up in infancy period. Others [27] reported that EndoCAb titers were not correlated with measurements of growth HAZ or $\Delta$HAZ (change of initial HAZ from the time of assessment and times of follow-up) or intestinal permeability. Lin et al., 2013, notified that, however, EndoCAb titers in children from clean are lower than children from contaminated households, no strong association has found between EndoCAb and anthropometric measurements.

Overt diarrhea is not pathognomonic feature of EED. It has been associated with high variety and frequencies of infections even if not associated with clinical symptom of diarrhea [24], [28], [29]. EED may associated with subclinical infection which may be present apart from manifested diarrhea, however, other histopathological abnormalities and associated environmental and nutritional risk factors are present [30].

Regarding the systemic inflammatory biomarkers in this study, there was significant increase of serum hsCRP and AGP in cases group compared to controls group, however, TNF increased in cases group but with insignificant value. Correlating the systemic inflammatory biomarkers in cases group with anthropometric parameters revealed significant negative correlation between weight Z score and both hsCRP and AGP. Furthermore, we detected significant negative correlation between height Z score and hsCRP and AGP. Moreover, hsCRP showed highly significant negative correlation with height and significant negative correlation with weight and arm circumference. However, no significant correlations were found between TNF and anthropometric measurements. Our results were supported by the findings of Syed et al. [31], Iqbal et al. [32] where systemic inflammatory biomarkers, CRP and AGP, were positively correlated with stunting in Young Tanzanian Children that assessed at 6 weeks and 6 months.

In a study done on 202 Zimbabwean infants by Prendergast et al., 2014, the levels of CRP and AGP were consistently higher from 6 weeks to 12 months of age, which indicate that early inflammation at 6 weeks as measured by both AGP and CRP was associated with an increased risk of stunting. Furthermore, children who were stunted at 18 months of age had significantly high CRP and reduced insulin-like growth factor-1 which is a marker for linear growth [27].

Surveillance of CRP and AGP was clearly associated with decreasing LAZ score at 9 months of age. Meanwhile, CRP and AGP were both negatively correlated with IGF-1 at 6 and 9 months of age, which support the connection pathway between enteric biomarkers and systemic inflammatory markers with growth retardation [32].

A study ran on rural Ugandan infants based on established cutoffs values, about 46% of enrolled children had elevated AGP and about 30% had elevated CRP. Higher AGP and CRP concentrations were significantly associated with lower LAZ at 6 months of age but not WAZ or WLZ [33]. Similarly, study survey according to the cutoff point in 18 months aged infants revealed 56% having elevated AGP and 20% having elevated CRP [26], while others found that no significant correlations were noted with hsCRP levels and HAZ or WAZ at study start or with delta HAZ of impairing linear growth [22].

Tumor necrotic factor is pro-inflammatory cytokine that produced from T lymphocytes, in EED, the ratio of TNF is increased in relation to other protective cytokines [3], [24]. It activates and drives differentiation of immune cells on infection [7]. It is one of the systemic inflammatory cytokines that detected with small intestine bacterial overgrowth in EED [34], TNF has role and affects on the tight junction barrier between enterocytes in the gut, TNF pathological effect in the gut is known to be mediated by disruption of myosin light chain kinase which leading to affection of tight junction and gut permeability [35].

Limited data present about TNF assessment in EED or associative relations with the anthropometric parameters of growth were available. Meanwhile, this study showed no significant difference between cases and controls groups, or significant correlating relations with anthropometric measurements.

There is a logical comprehensive link between the implicated pathology in EED and intestinal absorption and bioavailability of essential micronutrients.

Vitamins A and D are belonging to fat-soluble group of vitamins. Vitamin A enhances gut mucosal lining and control permeability leading to defendant barrier against microbial infestation [36], [37]. In addition, it has a modulating role in T lymphocytes production, which plays a role controlling the chronic condition of EED. Moreover, Vitamin A deficiency may alter the intestinal microbiota which may compromise enteric pathogen
clearance [38]. Vitamin D is involved in the processes of bone mineralization and growth through the regulation of calcium and phosphorus homeostasis [39]. The main site of Vitamins A and D absorption is small intestine which is the place of EED pathology [40].

Moreover, assessment of some micronutrients states revealed highly significant decrease of Vitamin D in cases group (p ≤ 0.001), in contrast to Vitamin A where no significant difference between the two groups was present (p > 0.05). Vitamin A showed significant positive correlation with weight Z score and BMI Z score (p ≤ 0.05). Likewise, Vitamin D exhibited significant positive correlation with weight Z score and height Z score (p ≤ 0.05).

No significant statistical difference was found in this study throughout comparing the hemoglobin level between cases and controls groups. However, there was highly significant decrease of iron level in cases group (p ≤ 0.001).

Applied interventional studies have shown that micronutrients, vitamins, and mineral supplementation improve intestinal morphology, reduce intestinal permeability, ameliorate gut functionality, and enhance linear growth [37].

Bacterial over growth and imbalance of the gut microbiota can both explain the state of subclinical infection and inflammatory markers imbalance that found in our studied cases group. It resulted in intestinal impermeability and elevated zonulin level and elevated antibodies against core bacteria (EndoCAb) creating a state of systemic inflammation denoted by elevated hsCRP and AGP, all that affect the absorptive function of the intestine affect the nutritional state that cleared in decreased Vitamin D micronutrient and manifested by affection of anthropometric parameters of weight and height.

Conclusion

Faltering growth is associated with elevated serum markers of Environmental enteric dysfunction such as biomarkers of enteric permeability (zonulin) and biomarkers of microbial translocation (EndoCAb), as well as biomarkers of systemic inflammation (HsCRP and AGP). EED may be a potential cause of faltering growth.

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