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ABSTRACT

Background: To accurately assess micronutrient status, it is necessary to characterize the effects of inflammation and the acute-phase response on nutrient biomarkers.

Objective: Within a norovirus human challenge study, we aimed to model the inflammatory response of C-reactive protein (CRP) and α-1-acid glycoprotein (AGP) by infection status, model kinetics of micronutrient biomarkers by inflammation status, and evaluate associations between inflammation and micronutrient biomarkers from 0 to 35 d post–norovirus exposure.

Methods: Fifty-two healthy adults were enrolled into challenge studies in a hospital setting and followed longitudinally; all were exposed to norovirus, half were infected. Post hoc analysis of inflammatory and nutritional biomarkers was performed. Subjects were stratified by inflammation resulting from norovirus exposure. Smoothed regression models analyzed the kinetics of CRP and AGP by infection status, and nutritional biomarkers by inflammation. Linear mixed-effects models were used to analyze the independent relations between CRP, AGP, and biomarkers for iron, vitamin A, vitamin D, vitamin B-12, and folate from 0 to 35 d post–norovirus exposure.

Results: Norovirus-infected subjects had median (IQR) peak concentrations for CRP [16.0 (7.9–29.5) mg/L] and AGP [0.9 (0.8–1.2) g/L] on day 3 and day 4 postexposure, respectively. Nutritional biomarkers that differed (P < 0.05) from baseline within the uninfamed group were ferritin (elevated day 3), hepcidin (elevated days 2, 3), serum iron (depressed days 2–4), transferrin saturation (depressed days 2–4), and retinol (depressed days 3, 4, and 7). Nutritional biomarker concentrations did not differ over time within the uninfamed group. In mixed models, CRP was associated with ferritin (positive) and serum iron and retinol (negative, P < 0.05).

Conclusion: Using an experimental infectious challenge model in healthy adults, norovirus infection elicited a time-limited inflammatory response associated with altered serum concentrations of certain iron and vitamin A biomarkers, confirming the need to consider adjustments of these biomarkers to account for inflammation when assessing nutritional status. These trials were registered at clinicaltrials.gov as NCT00313404 and NCT00674336.

Keywords: micronutrients, acute-phase response, inflammation, norovirus challenge, kinetics

Introduction

Describing the relation between acute-phase proteins (APPs) and nutritional biomarkers in settings with high infectious disease

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The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the CDC.

Supplemental Figure 1 and Supplemental Tables 1 and 2 are available from the “Supplementary data” link in the online posting of the article and from the same link in the online table of contents at https://academic.oup.com/ajcn/.

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Abreviations used: AGP, α-1-acid glycoprotein; APP, acute-phase protein; CRP, C-reactive protein; FUT2, fucosyltransferase 2; RBP, retinol-binding protein; RT-PCR, real-time PCR; sTfR, soluble transferrin receptor; 25(OH)D, 25-hydroxyvitamin D.

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burden is necessary for accurate assessment of micronutrient status (1). If underlying inflammation is ignored, iron and vitamin A status may be misclassified (2, 3). Within nationally representative surveys of children 6–59 mo old, misclassification ranged from underestimating iron deficiency by a median 25 percentage points (4) to overestimating vitamin A deficiency by a median 16 percentage points (5). Evaluation of nutrition programs or interventions may also be confounded by inflammation, specifically when they are combined with interventions targeted to reduce infection (6–8).

Currently, the only biomarker for which there is a consensus global guideline for adjustment in the context of inflammation is serum ferritin (9). The evidence base for adjusting nutritional biomarkers in the context of inflammation is limited to cross-sectional meta-analyses (10, 11), and a few longitudinal studies of adults undergoing surgery (12) or experiencing severe infections (13). Thus, an accurate assessment of micronutrient status depends on additional evidence quantifying time-varying interactions between nutritional biomarkers and infectious disease burden.

To assess patterns and interactions of inflammatory and nutritional biomarkers in response to an infectious challenge, we leveraged longitudinal data collected from 2 norovirus challenge trials (14, 15). Norovirus is the most common cause of diarrheal disease (16, 17), associated with 18% of diarrheal disease globally (18, 19). Although infections trigger an innate inflammatory response that temporarily alters circulating APPs (1, 11, 20), the kinetics of APPs triggered by exposure to norovirus has not been characterized. Iron and vitamin A biomarkers have been shown to be affected by subclinical levels of inflammation (5, 21–24), which may be better represented by the APP response to norovirus than a more severe infection, such as typhoid (13).

Our goal was to characterize the inflammatory response after a norovirus challenge, and to characterize the time-varying nutritional biomarker fluctuations by inflammatory response status. Our specific objectives were to model the kinetics of C-reactive protein (CRP) and α-1-acid glycoprotein (AGP) stratified by infection status, and to model the kinetics of micronutrient biomarkers [ferritin, hepcidin, serum iron, soluble transferrin receptor (sTfR), transferrin concentration, transferrin saturation, retinol-binding protein (RBP), retinol, 25-hydroxyvitamin D [25(OH)D], vitamin B-12, and folate] stratified by inflammation status. We hypothesized that CRP and AGP concentrations would differ by infection status, and that some micronutrient biomarker concentrations would differ by inflammation status resultant from norovirus exposure [ferritin ↑, hepcidin ↑, serum iron ↓, RBP ↓, retinol ↓, 25(OH)D ↓], whereas others would remain static (sTfR, vitamin B-12, and folate). We further evaluated the independent associations between CRP, AGP, and micronutrient biomarkers from 0 to 35 d postexposure.

**Methods**

**Subjects and ethics**

The serum assayed for this study was collected from apparently healthy adults that participated in 1 of 2 independent longitudinal norovirus challenge trials (NCT00313404 and NCT00674336) conducted at a Clinical Research Center in the United States between 2006 and 2009, and outcomes from those trials have been published (14, 15). The trials shared a common design and had similar protocols for specimen collection and participant assessment; however, they had different inoculum doses of S/Flb Norwalk virus (HuNoV genogroup GL.1, ∼6.5 × 10⁷ genomic equivalent copies per milliliter from 10 mL spiked groundwater or 1 × 10⁴ genomic equivalent copies of virus from infected oysters) (14, 15). In brief, healthy volunteers were challenged with the virus, observed in controlled conditions in a hospital setting for 4 d, and then discharged with instructions to return on days 7, 14, 21, 28, and 35 postchallenge. Serum was collected every day during the hospital stay and at each follow-up visit. Serum was stored in −80°C freezers for 7–10 y until they were shipped on dry ice to laboratories for the biomarker assessments of this study. Emory University Institutional Review Board approved both trials.

Of 64 enrolled volunteers, 26 became infected [norovirus was detectable in stool by real-time PCR (RT-PCR)] and among the infected 19 were symptomatic. A subsequent case-control study was conducted to describe the cytokine response patterns from norovirus exposure in 52 of the volunteers (25). We analyzed biomarkers from the same 52 subjects (age-matched, balanced by infection status) that had been sampled for the cytokine study (Supplemental Figure 1). The repeated-visits study design enabled assessing kinetics of CRP, AGP, and selected nutritional biomarkers. The nutritional biomarkers are ordered throughout the article in ascending order of existing evidence pertaining to their relation between nutrient and inflammation (1): iron [ferritin, hepcidin, iron, transferrin (receptor, saturation, and concentration)], vitamin A (RBP and retinol), vitamin D [25(OH)D], folate, and vitamin B-12.

**Laboratory methods**

Sandwich ELISA was used to measure CRP, AGP, ferritin, sTfR, and RBP at the VitMin laboratory (26). Serum hepcidin was quantified using the Hecpandin-25 (Bioactive) HS ELISA kit (DRG International, Inc.) according to the manufacturer’s protocol in a laboratory at Oxford University. Serum iron and transferrin concentrations were quantified using automated assays on an Abbott Architect c16000 automated analyzer (Abbott Laboratories) in the same lab that assessed hepcidin. Serum retinol (200 μL) was measured using HPLC in isocratic mode with 95:5 methanol:water (10 mM ammonium acetate) at 1 mL/min in a laboratory at the University of Wisconsin-Madison. Serum 25(OH)D was measured using the Immunodiagnostic Systems iSYS chemiluminescent assay in a laboratory at Emory University participating in the vitamin D External Quality Assessment Scheme. Folate concentrations were measured in serum using the microbiological assay at the Instituto de Investigación Nutricional in Lima, Peru. Serum vitamin B-12 was analyzed using a Cobas e411 (Roche Diagnostics) with a competitive protein binding chemiluminescence immunoassay at the Western Human Nutrition Research Center in Davis, CA. Both the VitMin and Instituto de Investigación Nutricional laboratories participate in the US CDC’s external laboratory quality assurance program VITAL-EQA (27).
Variable descriptions

All biomarkers were used as continuous variables in the kinetic modeling. Micronutrient and inflammation cutoffs were applied to define deficiency and to categorize subjects with inflammation due to norovirus exposure. The following cutoffs applied to define deficiency and to categorize subjects with kinetic modeling. Micronutrient and inflammation cutoffs were natural log–transformed to model these associations. Linear mixed-effect models were constructed for each biomarker outcome separately. The temporal effects of inflammation resulting from norovirus exposure did not correlate perfectly with norovirus infection status. Therefore, lagged variables were created and fit up to \( t - 4 \). Each model included all time points, and was fit using backwards selection through assessing model fit statistics, i.e., Akaike Information Criterion and log likelihood, coefficients, and SEs.

The full model took the form:

\[
\ln Y_t = \alpha + \theta + \text{CRP}_1 + \text{CRP}_{t-1} + \text{CRP}_{t-2} + \text{CRP}_{t-3} + \text{CRP}_{t-4} + \text{AGP}_1 + \text{AGP}_{t-1} + \text{AGP}_{t-2} + \text{AGP}_{t-3} + \text{AGP}_{t-4} + \ln Y_{t-1} + \varepsilon \quad (1)
\]

where \( Y_t \) is the nutritional biomarker concentration at measurement time \( t \), \( \alpha \) is the fixed intercept, \( \theta \) is the subject random-effect intercept as \( N(r^2 = 0) \), and \( \varepsilon \) is the residual model error as \( N(\sigma^2 = 0) \). The acute-phase inflammation biomarkers CRP and AGP are included at time \( t \) as CRP, and AGP. Lags are denoted as subtractions from \( t \), e.g., CRP

Statistical analysis

All data processing and analyses were conducted in R version 3.4.3 (R Core Team) using the “dplyr,” “nlme,” “dunn.test,” and “ggplot2” packages. Baseline characteristics were compared across infection and inflammation status, separately, using \( \chi^2 \) and Kruskal–Wallis tests for categorical and continuous measures, respectively. To characterize the inflammatory response of norovirus infection, the Kruskal–Wallis test was used to compare differences in median APP concentrations at days 1, 2, 3, 4, 7, 14, 21, and 35 by infection status. If any of the median tests indicated significant differences, then pairwise comparisons between each time point and baseline concentration were compared using Dunn’s test with correction for multiple comparisons. To characterize the nutritional biomarker fluctuations by inflammatory response status, the Kruskal–Wallis test was used to compare differences in median concentration of nutritional biomarkers at each time point postinfection, stratified by inflammation status. Similarly, Dunn’s test was used to compare differences between baseline and time points that differed between inflamed and uninflamed groups. Concentrations of biomarkers over time were modeled using local regressions (locally estimated scatterplot smoothing) to provide a smooth visual of temporal trends.

To evaluate the independent associations between APPs and each nutritional biomarker, we constructed time-lagged models of inflammation effects and autocorrelation of the prior measurement of nutritional biomarkers. All nutritional biomarkers were natural log–transformed to model these associations. Linear mixed-effect models were constructed for each nutritional biomarker outcome separately. The temporal effects and autocorrelation were explicitly modeled using time-lagged concentrations, i.e., \( \text{CRP}_{t-4} \), where \( t \) is the current measurement of a biomarker concentration and \( i \) represents a previous measure. Initially, autocorrelation was assessed within subjects using the function “acf” for \( \leq 10 \) previous measurements. A majority of biomarkers had the strongest correlations within days 1–4; therefore, lagged variables were created and fit up to \( t - 4 \). Each model included all time points, and was fit using backwards selection through assessing model fit statistics, i.e., Akaike Information Criterion and log likelihood, coefficients, and SEs.

Results

Participant characteristics

One infected subject was dropped owing to incomplete data. The median (IQR) age of study subjects was 25 (21–28) y old, and baseline temperature, pulse, respiration, and blood pressure indicated no evidence of acute illness. Baseline micronutrient deficiencies were absent for retinol (<0.7 \( \mu \)mol/L) and vitamin B-12 (<150 pmol/L). Six percent of subjects had folate deficiency (<10 nmol/L), 14% were at risk of vitamin D deficiency [25(OH)D <12 ng/mL], and 16% had low iron stores (serum ferritin <15 \( \mu \)g/L). Median vitamin B-12 concentration at baseline was significantly higher among subjects that became infected with norovirus than among noninfected subjects (Supplemental Table 1). Baseline nutritional profiles and APP concentrations did not differ between subjects stratified by clinical inflammation resulting from norovirus exposure (Table 1).

Six subjects with inflammatory biomarkers above thresholds commonly used to define inflammation at baseline (4 with elevated CRP, 2 with elevated AGP) were excluded from all analyses beyond the descriptive statistics. Clinical inflammation resulting from norovirus exposure did not correlate perfectly with norovirus infection, defined by norovirus detectable in stool by RT-PCR. The majority of subjects that demonstrated clinical inflammation resulting from norovirus exposure were infected with norovirus (\( n = 20 \) of 21, or 95%; Table 2). However, of the 24 noninfected subjects, 3 people were norovirus infected (detectable in stool by RT-PCR), although they were asymptomatic.

APP response to norovirus infection

The inflammatory response of CRP and AGP serum concentrations was modeled by norovirus infection status. There were similarities between CRP and AGP peak concentrations as well as normalization time among norovirus-infected individuals (Figure 1). CRP peaked with a median (IQR) concentration of 16.0 (7.9–29.5) mg/L at day 3, and AGP peaked at 0.9
Norovirus inflammation and micronutrient kinetics

TABLE 1 Baseline characteristics of norovirus challenge subjects, stratified by inflammation resulting from norovirus exposure

<table>
<thead>
<tr>
<th>Demographics</th>
<th>Uninflamed (n = 24)</th>
<th>Inflamed(^1) (n = 21)</th>
<th>Inflamed at baseline (n = 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>24.0 (20.8–27.0)</td>
<td>26.0 (22.0–28.0)</td>
<td>23.5 (23.0–36.0)</td>
</tr>
<tr>
<td>Female</td>
<td>64</td>
<td>57</td>
<td>33</td>
</tr>
<tr>
<td>Inflammation</td>
<td>CRP, mg/L</td>
<td>1.0 (0.5–1.5)</td>
<td>0.6 (0.3–1.8)</td>
</tr>
<tr>
<td></td>
<td>AGP, g/L</td>
<td>0.6 (0.5–0.8)</td>
<td>0.6 (0.4–0.7)</td>
</tr>
<tr>
<td>Nutrition</td>
<td>Ferritin, µg/L</td>
<td>50.8 (15.4–96.9)</td>
<td>65.0 (34.1–107.0)</td>
</tr>
<tr>
<td></td>
<td>Haptoglobin, ng/mL</td>
<td>5.9 (2.0–15.0)</td>
<td>10.0 (7.6–17.4)</td>
</tr>
<tr>
<td></td>
<td>Serum iron, µmol/L</td>
<td>12.4 (10.6–17.8)</td>
<td>15.2 (12.2–19.1)</td>
</tr>
<tr>
<td></td>
<td>sTfR, mg/L</td>
<td>5.1 (4.1–6.7)</td>
<td>5.0 (4.2–5.6)</td>
</tr>
<tr>
<td></td>
<td>Transferrin, g/L</td>
<td>2.7 (2.5–3.3)</td>
<td>2.7 (2.5–2.9)</td>
</tr>
<tr>
<td></td>
<td>TSAT, %</td>
<td>20.0 (16.1–30.7)</td>
<td>27.6 (21.3–31.7)</td>
</tr>
<tr>
<td>RBP, µmol/L</td>
<td>1.7 (1.6–2.0)</td>
<td>1.9 (1.8–2.2)</td>
<td>2.2 (1.9–2.4)</td>
</tr>
<tr>
<td>Retinol, µmol/L</td>
<td>1.3 (1.0–1.4)</td>
<td>1.3 (1.2–1.6)</td>
<td>1.6 (1.1–1.7)</td>
</tr>
<tr>
<td>25(OH)D, ng/mL</td>
<td>20.2 (14.9–27.4)</td>
<td>20.0 (17.4–23.0)</td>
<td>22.4 (16.0–24.8)</td>
</tr>
<tr>
<td>Vitamin B-12, pmol/L</td>
<td>315.3 (269.9–388.5)</td>
<td>467.8 (307.9–609.1)</td>
<td>315.2 (277.6–329.6)</td>
</tr>
<tr>
<td>Folate, ng/mL</td>
<td>14.3 (8.6–18.8)</td>
<td>12.9 (10.2–21.3)</td>
<td>6.1 (4.4–16.9)</td>
</tr>
<tr>
<td>Vital signs</td>
<td>Temp, °C</td>
<td>36.6 (36.1–36.7)</td>
<td>36.0 (36.0–36.8)</td>
</tr>
<tr>
<td></td>
<td>Pulse, bpm</td>
<td>72 (66–78)</td>
<td>71 (63–83)</td>
</tr>
<tr>
<td></td>
<td>Diastolic blood pressure, mm Hg</td>
<td>73 (67–80)</td>
<td>69 (59–76)</td>
</tr>
<tr>
<td></td>
<td>Systolic blood pressure, mm Hg</td>
<td>115 (110–124)</td>
<td>116 (111–123)</td>
</tr>
<tr>
<td></td>
<td>Respiration, brpm</td>
<td>18 (18–18)</td>
<td>16 (16–18)</td>
</tr>
</tbody>
</table>

\(1^*\) = 51. Values are median (IQR) or %. *Significant at \( P < 0.05 \) using either \(\chi^2\) or Kruskal–Wallis test to compare inflamed and uninflamed subjects for categorical and continuous data, respectively. AGP, α-1-acid glycoprotein; bpm, beats per minute; brpm, breaths per minute; CRP, C-reactive protein; RBP, retinol-binding protein; sTfR, soluble transferrin receptor; TSAT, transferrin saturation; 25(OH)D, 25-hydroxyvitamin D.

\(2^\)Clinical inflammation resulting from norovirus exposure was defined as CRP > 5 mg/L or AGP > 1 g/L at any time 1–3 d postexposure, given CRP ≤ 5 mg/L and AGP ≤ 1 g/L on day 0.

(0.8–1.2) g/L at day 4 for the infected group. At day 3, both APPs were significantly different from the uninflamed group (CRP: \(\chi^2_{\text{K-W}} = 21.5, P < 0.01\); AGP: \(\chi^2_{\text{K-W}} = 4.6, P < 0.01\)). CRP and AGP medians of the infected group were indistinguishable from the uninflamed group by day 7 (\(\chi^2_{\text{K-W}} = 1.5, P = 0.21\); \(\chi^2_{\text{K-W}} = 1.0, P = 0.30\)). Within the uninflamed group, neither CRP nor AGP concentrations deviated from baseline (\(\chi^2_{\text{K-W}} = 6.9, P = 0.64\); \(\chi^2_{\text{K-W}} = 11.3, P = 0.24\)) over the study course (Figure 1).

**TABLE 2** Comparison of subject classifications for inflamed, norovirus infected, and symptomatic within a norovirus human challenge study

<table>
<thead>
<tr>
<th>Classification</th>
<th>Inflamed</th>
<th>Uninfected</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Norovirus infected and symptomatic</td>
<td>16</td>
<td>0</td>
<td>16</td>
</tr>
<tr>
<td>Norovirus infected and asymptomatic</td>
<td>4</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Norovirus uninfected and symptomatic</td>
<td>0</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Norovirus uninfected and asymptomatic</td>
<td>1</td>
<td>19</td>
<td>20</td>
</tr>
<tr>
<td>Total</td>
<td>21</td>
<td>24</td>
<td>45</td>
</tr>
</tbody>
</table>

\(1^*\)Inflammation resulting from norovirus exposure was defined as CRP > 5 mg/L or AGP > 1 g/L at any time 1–3 d postexposure, given CRP ≤ 5 mg/L and AGP ≤ 1 g/L on day 0. Six subjects were excluded based on CRP > 5 mg/L or AGP > 1 g/L on day 0. Norovirus infection was defined as norovirus detectable in stool by real-time PCR. Symptomatic was defined as diarrhea, vomiting, or ≥ 1 of the following self-reported characteristics: nausea, abdominal cramps, headache, chills, myalgia, or fatigue. AGP, α-1-acid glycoprotein; CRP, C-reactive protein.

**Nutritional biomarker responses to inflammation caused by norovirus exposure**

The kinetics of micronutrient biomarkers [ferritin, haptoglobin, serum iron, sTfR, transferrin concentration, transferrin saturation, RBP, retinol, 25(OH)D, folate, and vitamin B-12] were stratified by clinical inflammation resulting from norovirus exposure. Concentrations of the following nutrient biomarkers differed significantly compared with baseline concentrations among subjects that experienced inflammation from norovirus exposure: ferritin (elevated day 3, \(P < 0.05\)), haptoglobin (elevated days 2–3, \(P < 0.01\)), serum iron (depressed days 2–4, \(P < 0.01\)), tranferrin saturation (depressed days 2–4, day 2 \(P < 0.05\), day 3–4 \(P < 0.01\)), and retinol (depressed days 3, 4, and 7, \(P < 0.05\); Figure 2). Although fluctuations in concentrations of the remaining nutritional biomarkers can be observed visually, there were no time points that differed in statistical significance from baseline concentration for sTfR, transferrin concentration, RBP, 25(OH)D, folate, and vitamin B-12 among the inflamed group (Figure 2). Among the uninflamed group, regardless of infection status, no nutritional biomarkers had concentrations that were statistically significantly different from baseline.

**Independent effects of inflammation on nutritional biomarkers**

Based on the most conservative models, adjusting for lagged inflammation effects and autocorrelation of the prior measurement of nutritional biomarkers for all subjects across
FIGURE 1 Baseline (day 0) to day 35 postexposure CRP (A) and AGP (B) measurements plotted using box and whiskers, displaying median and IQR (box range) with whiskers representing 1.5 × the 25% and 75% quartile and outliers depicted by dots (n = 45 subjects with repeated measures). Pairwise differences between baseline and other days within infection grouping tested using Dunn’s test. Box plot locally estimated scatterplot smoothing fit with 95% prediction interval in black and gray banding. AGP, α-1-acid glycoprotein; CRP, C-reactive protein.

Discussion

Using an experimental infectious challenge model, we have demonstrated a clear causative, time-limited relation between acute inflammation and perturbations in micronutrient biomarkers of iron and vitamin A status, confirming the need to consider adjustments of these biomarkers to account for inflammation when assessing nutritional status. Norovirus induced an inflammatory response (demonstrated by elevations in CRP and AGP) 3–4 d after exposure, which caused elevations in serum ferritin and hepcidin, and reductions in serum iron and retinol concentrations. These perturbations were supported by the independent positive association between ferritin and CRP, and the independent negative associations between serum iron, retinol, and CRP in linear mixed models. No nutritional biomarker perturbations were observed over the 35-d study duration among uninflamed subjects, further supporting that the concentration changes seen among the inflamed group were likely artificial and not reflecting true nutrition status alterations.

Inflammatory protein response to norovirus infection

CRP has been described as the “first line of innate host defense” for decades (33–36). Although animal models have

time (Y_{t-1}), ferritin_t, serum iron_t, sTfR_t, retinol_t, RBP_t, and vitamin B-12, were independently associated with CRP_t or AGP_t (Table 3). The iron biomarkers ferritin_t (β: 0.11; 95% CI: 0.06, 0.15) and serum iron_t (β: −0.10; 95% CI: −0.19, −0.01) were significantly associated with CRP_t, sTfR_t was significantly positively associated with AGP_t (0.10; 95% CI: 0.08, 0.13) and negatively associated with CRP_t (β: −0.06; 95% CI: −0.08, −0.03). The vitamin A biomarkers retinol_t (β: −0.05; 95% CI: −0.07, −0.04) and RBP_t (β: −0.07; 95% CI: −0.09, −0.05) were significantly negatively associated with CRP_t, RBP_t was also significantly positively associated with AGP_t (β: 0.04; 95% CI: 0.02, 0.06). Vitamin B-12 was significantly positively and negatively associated, respectively, with AGP_t (β: 0.03; 95% CI: 0.01, 0.05) and CRP_t (β: −0.02; 95% CI: −0.04, −0.003) (Table 3). The coefficients for the lagged inflammation effects are included in Supplemental Table 2.
characterized CRP kinetics (37), there are limited studies in humans. Furthermore, most investigations have modeled CRP kinetics under extreme conditions, such as surgeries (12, 38). As evident from recent meta-analyses, nutritional biomarkers are affected by even low levels of inflammation in both high and low infection burden settings (5, 21–24), raising the question of APP patterns during common, self-limiting infections. In our trial, norovirus infection triggered an acute-phase response as indicated by median peak concentrations of 16.0 mg CRP/L at day 3 postexposure, >3 times the usual CRP cutoff (5 mg/L). However, the median peak concentration of AGP (0.9 g/L) at day 4 was lower than the traditionally used cutoff of AGP of 1 g/L. In a controlled typhoid experiment among apparently healthy adults, CRP peaked 2 d after diagnosis, with a median concentration >100 mg/L (13). Two to 3 d after elective orthopedic surgery, CRP reached a maximum concentration, ranging from 48 to 140 mg/L (38). CRP has demonstrated increases >1000-fold post–inflammatory stimuli (39). Evidence of AGP kinetics during infection or in response to tissue injury is limited, compared with CRP. However, there has been a general recognition that AGP rises more slowly than CRP and stays elevated for a longer period of time (40), although this pattern was not observed in this study.

Nutritional biomarker responses to inflammation caused by norovirus exposure

Although the kinetics we observed for CRP and AGP in response to norovirus infection were distinct from typical thresholds used to describe the acute-phase response (10, 22, 41, 42), the iron and vitamin A responses to inflammation were as hypothesized. Marked reductions in circulating iron (serum iron and transferrin saturation) (43–45) were likely caused by the observed induction of the hormonal iron regulator, hepcidin, and the iron storage protein ferritin also increased. The hepcidin concentration peaked (day 2 postexposure) before the ferritin peak (day 3 postexposure). Similar to other APPs, hepcidin is upregulated during the innate response to infection via the IL6/signal transducer and activator of transcription 3 pathway (46, 47). An increase in hepcidin concentrations after pathogen exposure has been likewise demonstrated in other human infection challenge studies, including typhoid (13) and malaria (48, 49). The vitamin A biomarkers, RBP and retinol, demonstrated similar responses to inflammation: retinol concentrations were statistically lower than baseline on days 3, 4, and 7 post–exposure to norovirus, and RBP followed a similar pattern that was observed visually, but was not statistically significant,
possibly owing to small sample size or a modest inflammatory response. This depression of vitamin A biomarkers coinciding with elevated inflammatory proteins has been demonstrated in numerous studies (50–52). Concentrations of sTfR, transferrin, 25(OH)D, folate, and vitamin B-12 did not differ over time in the inflamed group. Among the uninflamed group, none of the nutritional biomarker concentrations differed over time, indicating that the nutritional biomarker perturbations seen among the inflamed group were likely artificial and not reflecting true nutrient status alterations.

The lower vitamin B-12 concentrations at baseline that appeared protective of norovirus infection may have a genetic explanation, such as a single nucleotide polymorphism. Although secretor-positive adults were enrolled into these challenge studies (32), genetic variants of fucosyltransferase 2 (FUT2) have been associated with lower vitamin B-12 status (53, 54). Homozygous FUT2 nonsecretor genotype is characterized as resistant to norovirus infection (55, 56).

**Independent effects of inflammation on nutritional biomarkers**

The independent associations of ferritin, serum iron, and retinol with CRP at time \( t \) support the causal framework that inflammation affects iron and vitamin A metabolism. However, the combination of assessing CRP and AGP simultaneously adds complexity to this analysis and interpretation. For example, sTfR, RBP, and vitamin B-12 all had significant direct associations with CRP and significant inverse associations with AGP at time \( t \), possibly due to distinct roles of the APPs that result in nutrient sequestration that varies across time. As demonstrated by Wessells et al. (57), accounting for the temporal effects of inflammation resulted in more subtle corrections for inflammation than cross-sectional-based techniques. Having access to repeated measures on individuals within surveys to estimate the effects of inflammation on micronutrient biomarkers may be rare, but additional analysis of existing cohort data sets that contain

**TABLE 3** Time \( t \) \( \beta \) (95% CI) estimates of associations between inflammatory markers and nutritional biomarkers, accounting for measurements at all times and the lagged effects of inflammation

<table>
<thead>
<tr>
<th>( Y_t )</th>
<th>CRP ( \beta )</th>
<th>AGP ( \beta )</th>
<th>( Y_{t-1} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ferritin</td>
<td>0.11 (0.06, 0.15)*</td>
<td>0.02 (−0.02, 0.06)</td>
<td>0.91 (0.865, 0.95)*</td>
</tr>
<tr>
<td>Hepcidin</td>
<td>0.09 (−0.13, 0.30)</td>
<td>0.06 (−0.09, 0.21)</td>
<td>0.54 (0.43, 0.64)*</td>
</tr>
<tr>
<td>Serum iron</td>
<td>−0.10 (−0.19, −0.01)*</td>
<td>—</td>
<td>0.24 (0.12, 0.35)*</td>
</tr>
<tr>
<td>sTfR</td>
<td>−0.06 (−0.08, −0.03)*</td>
<td>0.10 (0.08, 0.13)*</td>
<td>0.15 (0.05, 0.25)*</td>
</tr>
<tr>
<td>Transferrin</td>
<td>−0.001 (−0.02, 0.02)</td>
<td>—</td>
<td>0.89 (0.83, 0.95)*</td>
</tr>
<tr>
<td>TSAT</td>
<td>−0.08 (−0.17, 0.02)</td>
<td>—</td>
<td>0.225 (0.105, 0.34)*</td>
</tr>
<tr>
<td>RBP</td>
<td>−0.07 (−0.09, −0.05)*</td>
<td>0.04 (0.02, 0.06)*</td>
<td>0.29 (0.19, 0.385)*</td>
</tr>
<tr>
<td>Retinol</td>
<td>−0.05 (−0.07, −0.04)*</td>
<td>—</td>
<td>0.26 (0.16, 0.36)*</td>
</tr>
<tr>
<td>25(OH)D</td>
<td>—</td>
<td>0.03 (0.00, 0.05)</td>
<td>0.18 (0.07, 0.30)*</td>
</tr>
<tr>
<td>Vitamin B-12</td>
<td>−0.02 (−0.04, −0.003)*</td>
<td>0.03 (0.01, 0.05)*</td>
<td>0.84 (0.79, 0.89)*</td>
</tr>
<tr>
<td>Folate</td>
<td>0.00 (−0.04, 0.04)</td>
<td>−0.02 (−0.07, 0.03)</td>
<td>0.485 (0.40, 0.57)*</td>
</tr>
</tbody>
</table>

\[ Y_t = \alpha + \beta \cdot \text{CRP}_t + \beta \cdot \text{CRP}_{t-1} + \beta \cdot \text{CRP}_{t-2} + \beta \cdot \text{CRP}_{t-3} + \beta \cdot \text{CRP}_{t-4} + \beta \cdot \text{AGP}_t + \beta \cdot \text{AGP}_{t-1} + \beta \cdot \text{AGP}_{t-2} + \beta \cdot \text{AGP}_{t-3} + \beta \cdot \text{AGP}_{t-4} + \ln(Y_{t-1}) + \varepsilon, \]

where \( Y_t \) is the nutritional biomarker concentration at measurement time \( t \), \( \alpha \) is the fixed intercept, \( \beta \) is the subject random-effect intercept as \( \alpha \sim N(0^2, 0) \), and \( \varepsilon \) is the residual model error as \( \varepsilon \sim N(0, 0) \). The inflammation biomarkers CRP and AGP are included at time \( t \) as CRP, and AGP, as well as lagged terms up to \( t - 4 \). Values are standardized \( t \) (95% CI) for CRP, and AGP. Models adjusted for subject and temporal effects using mixed effects. CRP, AGP, 1, CRP, and AGP, were not included in any final models. Significant at \( \alpha < 0.05 \); \( n = 45 \) subjects at 10 time points. Supplemental Table 2 presents \( t \) (95% CI) estimates for the lagged inflammation variables. AGP, \( \alpha \)-1-acylglycoprotein; CRP, C-reactive protein; RBP, retinol-binding protein; sTfR, soluble transferrin receptor; TSAT, transferrin saturation; 25(OH)D, 25-hydroxyvitamin D; —, term was not included in the final model.

**Strengths and limitations**

Our study is one of the first to assess dynamic changes in inflammation and micronutrient biomarkers after an experimental enteric viral challenge. We quantified the kinetics of APPs given norovirus infection, and the subsequent effects of inflammation on nutrient biomarkers, providing clear indication that nutrient biomarkers fluctuate during relatively mild infections among adults. The strengths of this analysis were the repeated-measures design and the controlled norovirus challenge with serum samples utilized at 10 time points. A limitation of this analysis was that results from adults cannot be directly transferred to children. The population group of highest concern for nutritional status misclassification and norovirus infections are children <5 y of age; however, our challenge studies were performed on apparently healthy adults. The immune system maturation likely influences the inflammation and nutritional biomarker kinetic response to norovirus exposure. Therefore, patterns might be more pronounced in children. Another limitation was the small sample size. In some cases such as RBP, changes in nutritional biomarker concentrations seemed apparent graphically, yet the studies were likely underpowered and did not reach statistical significance. The inflammation lag terms in the linear mixed model may have been easier to interpret within a larger sample as well.

**Conclusions**

The confounding effects of inflammation on micronutrient biomarkers have been identified as a critical research gap (1), and efforts to adjust nutrition biomarkers for inflammation have been limited to cross-sectional data (11, 22, 42, 58). This study demonstrates that some people exposed to norovirus do not experience virus in their stools or elevated APPs. Inflammation
from norovirus affects the measurement concentrations of some iron and vitamin A biomarkers, and therefore should be interpreted accordingly.

We recognize input from Dr Christine Clewes, PhD, independent consultant on the interpretation of results.

The authors’ responsibilities were as follows—AMW, JSL, BAL, PSS, and RCF-A: conceptualized this research; AMW and CKNL: had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis; VT, RDW, SAT, AEA, AM, S-RP, SS-F, KW, and LA: all assisted in laboratory analysis and interpretation of the results; DT: provided insights on the interpretation of the results; and all authors: read the manuscript and provided intellectual content. None of the authors reported a conflict of interest related to the study.

References


