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Iodine supplementation of lactating women and assessment of infant visual information processing and maternal and infant thyroid function: A randomized trial

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Abstract

Iodine deficiency is one of the major causes of brain damage in childhood. However, iodine supplementation during early pregnancy and lactation can prevent the ill effects of iodine deficiency. This study evaluated maternal and infant thyroid function and infant visual information processing (VIP) in the context of maternal iodine supplementation. A community-based, randomized, supplementation trial was conducted. Mother infant dyads (n = 106) were enrolled within the first 10 days after delivery to participate in this study. Mothers were randomly assigned either to receive a potassium iodide capsule (225 μg iodine) daily for 26 weeks or iodized salt weekly for 26 weeks. Maternal thyroxine (T4), triiodothyronine (T3), thyroid stimulating hormone (TSH), thyroglobulin (Tg), urinary iodine concentration (UIC), breast milk iodine concentration (BMIC) and infant T4, TSH, UIC and VIP were measured as outcome variables. At baseline, neither mothers nor infants in the two groups were significantly different in any of the biomarkers or anthropometric measurements. Maternal TSH and goiter prevalence significantly decreased following iodine supplementation. The percentage of infants who preferentially remembered the familiar face was 26% in the capsule and 51% in the I-salt groups. Infant sex, length for age Z score, BMIC, maternal education and household food security were strong predictors of novelty quotient. In conclusion supplementation daily for six months with an iodine capsule or the use of appropriately iodized salt for an equivalent time was sufficient to reduce goiter and TSH in lactating women. Higher BMIC and LAZ as well as better household food security, maternal education, and male sex predicted higher novelty quotient scores in the VIP paradigm.
Introduction

Iodine is essential for the synthesis of thyroid hormones that regulate the metabolic processes of most cells and play important roles in human growth and development [1]. Iodine deficiency disorders (IDDs) are a major global public health problem and one of the most severe causes of preventable brain damage in childhood [2].

IDDs can be prevented and controlled by providing iodine and iodine can be provided in different ways. However, the United Nations Children's Fund and the World Health Organization jointly recommended salt iodization where iodized salt is accessible. Iodized salt is a safe, cost effective and sustainable strategy to ensure sufficient intake of iodine by all individuals and to improve maternal and infant health [3–6]. Globally, over 70% of households use iodized salt [7]. In places where iodized salt is not accessible, the recommendation from WHO/UNICEF/ICCIDD has been a daily dose of 250 μg iodine supplement for pregnant and lactating women or a single annual dose of 400 mg iodine as an iodized oil supplement [4]. However, one recent study has reported that 150 μg of iodine daily was enough to reduce the number of women with insufficiency [8]. For children below 2 years of age, the recommendation is a daily dose of 90 μg iodine or a single annual dose of 200 mg iodine as an iodized oil supplement [4]. For infants 0 to 6 months of age, because a significant amount of iodine is secreted into breast milk, iodine should be available through breast milk, provided that the lactating mother is receiving sufficient iodine and her child is exclusively breastfed [9].

Although there are some studies that confirmed the effect of iodine supplement on thyroid status, thyroid gland and brain function [6, 10], little has been done to assess the efficacy of iodized salt in relation to these outcomes. In a prospective study in Denmark four years after a mandatory salt iodization program (13 mg/kg iodine), a lower median thyroid volume was observed in women aged 18 to 65 years. A larger relative decline of thyroid volume was observed in the younger compared to older females from a site where iodine deficiency was moderate [11]. In our study, women supplemented with adequately iodized salt had breast milk iodine concentrations that were not different from women who received a daily capsule containing 225 ug iodine; infant urinary iodine concentrations also were not significantly different based on source of maternal iodine supplement [12].

Ethiopia launched a national salt iodization program in 2012. However salt has been iodized manually using knapsack sprayers which makes it difficult to get a homogenous product. According to the Ethiopian Public Health Institute, of the 5605 salt samples tested nationally in 2014 only 42.7% had iodine content above 15 mg/kg, the rest varied between 0 and < 15 mg/kg [13].

The general objective of this study was to assess the efficacy of adequately iodized salt (30–40 mg/kg) for lactating women and for their breast-fed infants to the age of six months compared to a daily 225 μg iodine capsule. The specific objective of this study was to evaluate maternal and infant thyroid function and infant visual information processing (VIP) in the context of maternal iodine supplementation.

Materials and methods

Subjects and study design

The study was a community-based, randomized, supplementation trial conducted in 2013 in Sidama zone, southern Ethiopia. The study population depended on subsistence farming for their livelihood. Their major staple food was enset (Enset ventricosum) followed by unrefined maize [14]. The area was known for severe iodine deficiency prior to national implementation of salt iodization in 2012 [15–17]. Breast feeding behavior, household salt use, and compliance
of subjects was reported in our previously published manuscript [12]. This study is registered with ClinicalTrials.gov, number NCT03889431.

**Subject selection methodology**

This study enrolled all lactating women in the participating communities who delivered between January and February 2013. A total of 106 healthy mother-infant dyads were included in the study and data collection was completed by August 30, 2013. Eligibility criteria were: the women must be lactating and must have delivered a full-term single infant within the prior 10 days. Mothers and infants must have no history of illness, thyroid dysfunction or iodine supplementation. All mother-infant dyads who met the criteria were invited to volunteer for the study, and enrollment was continued until the required sample size was attained.

**Ethical clearance.** Ethical approval was obtained from Oklahoma State University (OSU), Hawassa University, the Ethiopian Ministry of Science and Technology, and the Food, Medicine, and Health Care Administration and Control Authority of Ethiopia. Following receipt of ethical clearance, study participants were given a detailed explanation of the objectives of the research and consent was obtained from the women for themselves and their infants before data collection was started. Women were enrolled with the help of health extension workers working in the study area.

**Randomization.** The 106 mothers with their infants were randomly assigned either to the iodine capsule group or to the iodized salt (I-salt) group using random numbers (Fig 1).

**Procedures.** The capsule group (n = 53) received 225 μg of iodine daily as a capsule of potassium iodide (Pure Encapsulations, Inc., Boston, MA) for 26 weeks. Iodine capsules were ingested daily by mothers under supervision of the health extension worker. The I-salt group was provided weekly with a 450 g packet of iodized salt (30 to 40 mg I as KIO$_3$ /kg) for their household for 26 weeks (Guts Agro Industry PLC, Addis Ababa, Ethiopia) by the health extension workers. The 450 g of salt provided weekly for each family was planned to be enough that they would not purchase any additional (perhaps inappropriately iodized) salt from the market. There was no requirement to use all of the provided salt.

A national survey conducted in 2005 with a multistage cluster sampling design estimated salt consumption per person per day using 24-hour dietary recalls. The national mean (±SD) for salt consumption was 8.4 (5.9) g/day. However, based on the 1,677 households sampled in the SNNPR region (our study region), salt consumption was 17.2 (13.8) g per person per day, or more than twice the national average. One reason for high salt consumption in the region is the custom for use of salt rather than sugar in coffee [18].

Prior to beginning supplementation, baseline data were collected from both mothers and infants. Collection and analysis of breast milk and urine has been described in our previous paper [12]. Maternal blood samples were collected using a disposable 10 cc syringe coated with lithium heparin with a 21 gauge needle (Sarstedt, Inc., Newton, N.C.). The blood was centrifuged and plasma was separated immediately. Plasma was frozen at – 20°C and analyzed for concentrations of triiodothyronine (T$_3$), thyroxine (T$_4$), thyroid stimulating hormone (TSH) and thyroglobulin (Tg).

Each woman was examined for goiter at baseline and end point by a single health officer using palpation based on the following grades: grade 0, no palpable or visible goiter; grade 1, palpable goiter but not visible when neck is in the normal position; grade 2, visible goiter when neck is in the normal position [4]. For mothers weight and height to calculate body mass index (BMI = Wt$_{kg}$/Ht$_{m2}$), and mid upper arm circumference (MUAC) were measured. Infant anthropometry including head circumference, weight and length were measured. Infants’ weight-for-age (WAZ), length-for-age (LAZ), weight-for-length (WLZ), and head circumference-for-age...
(HCAZ) were calculated using WHO Anthro software (Version 2.0.4). Anthropometric measurements were done in duplicate.

Mothers were interviewed at baseline using an individual questionnaire adapted from the Ethiopian Demographic and Health Survey. The questionnaire included socioeconomic status and demographic characteristics of the women [19]. Additionally, data were collected using the Household Food Insecurity Access Scale (HFIAS) developed by the Food and Nutrition Technical Assistance (FANTA) project of the United States Agency for International Development (USAID) [20].

After 26 weeks, collection of biological samples and anthropometric measurements were repeated. Additionally, at 26 weeks infant blood samples were collected on specimen paper by finger prick for TSH and T4 analysis, and the Visual Information Processing (VIP) test was
administered to infants. All the participating mothers lactated throughout the study; thus no mother was excluded for having stopped lactation after enrollment.

**Administration of visual information processing (VIP) test**

**Materials.** Testing of infant VIP was modified from an earlier report with Ethiopian infants [21] and from the work of Rose and colleagues [22]. For the VIP, eight pictures of young adult Ethiopian faces without emotional expression were used. One laptop computer controlled the presentation of stimuli. A second computer was used to follow and record the infant’s looking behavior. The infant’s looking behavior was coded live, but a camera in the second computer recorded the VIP test for later reliability testing. VIP measurements were always conducted by the same person.

**VIP test procedures.** For the VIP test, the infant sat on the mother’s lap half a meter from a 15” laptop screen in a darkened room at the health center. The tester (who was blind to the study and out of sight of the infant) focused the camera in the second laptop on the infant’s eyes. The habituation phase began with the presentation of one face randomly selected from the eight faces. When the infant looked at a picture, its reflection was visible in the infant’s pupil and the experimenter pressed a key on his computer. Response latency (time between stimulus onset and the look) and the duration of the look were recorded by the computer software. When the infant looked away, the experimenter released the key; if the look away lasted 1s or longer, the face disappeared from the screen for 2s and then reappeared. This procedure continued until the criterion for habituation was achieved, namely two consecutive looks that were less than one half the mean of the two longest looks.

After habituation, the comparison phase began without interruption. In the comparison phase, the face that appeared during habituation (the “familiar” face) appeared again on one side of the screen and a novel face of the opposite sex appeared on the other side of the screen. The faces remained on the screen regardless of whether or not the infant was looking at one of them. Once the infant accrued a total looking time at the faces of 5s, the faces disappeared for 1s, and then reappeared but on opposite sides of the screen. They remained until an additional 5s of looking had accrued. The infant’s look duration, shifts between faces, and looks away from the pictures were compiled simultaneously by the software.

Visual recognition memory is composed of various constructs [22]. During habituation, look duration measures processing speed and attention [23] and average look duration and number of looks reflect the speed at which recognition was achieved [24]. For the comparison phase, the novelty quotient (NQ; preference for the new face) measures recognition memory (21) and total shifts identify how active the infant was in comparing the two faces.

A university student, blind to the study, coded recorded videos for 30 randomly selected infants to test coding reliability. For longest look, the correlation was 0.78; correlation for novelty preference was low probably because distinguishing between infant’s looks at the right and left side of the screen was more difficult in the taped version than in person.

**Measurement of biomarkers**

The $T_3$, $T_4$, TSH and Tg were quantitatively determined by ELISA assay (ALPCO Diagnostics, Salem, NH). Quality control for $T_3$ and $T_4$ were obtained from Bio-Rad Laboratories (Irvine, Ca). Infant’s $T_4$ and TSH also were determined by ELISA assay (Diagnostic Automation, Calabasas, CA). Reference ranges for thyroid hormone biomarkers of mothers and infants are included as supplied by the manufacturers. Hence, for mothers TSH: 0.4–4.2 μIU/mL, $T_3$: 0.75–2.2 ng/mL, $T_4$: 5–13 μg/dL and Tg: 2–50 ng/mL. Reference ranges for infants were: TSH: 0.7–34 μIU/mL and $T_4$: 8–23 μg/dL.
**Statistical analysis**

Percentages, frequency distributions, means (SD) and medians (IQR) were used to describe the socio-economic status, demographic characteristics, and iodine status of lactating mothers and their infants. All data were checked for normal distribution and skewed data were log-transformed before analysis. To compare the means between the two treatment groups independent-sample t-tests were used for normally distributed data and the Mann-Whitney U test for non-normally distributed data. For comparison of samples at baseline and 26 weeks, paired-sample t-tests were used for normally distributed data and Wilcoxon signed-Rank tests were used for non-normally distributed data.

Pearson’s correlation coefficient (r) or Spearman’s rho were used to examine relations between variables. Variables predicting VIP were analyzed by multiple regression. All variables were checked for collinearity using variance inflation factors (VIF). All of the biomarkers analyzed and infants’ anthropometric indices with VIF < 4 were entered into the models.

Multiple classification analysis (MCA) was used for predicting novelty quotient (NQ). MCA examines the contribution of each category of the predetermined predictor variables before and after adjustment for the control variables. MCA is effective to control multicollinearity [25]. Post hoc power calculation was made using STATA software for four outcomes (Tg, T₃, T₄ and TSH) and the power was adequate (>80%) for the Tg and TSH outcomes. However, it was lower than 80% for T₃ and T₄.

**Results**

A total of 106 mother-infant pairs were recruited between Jan and Feb 10, 2013 and data collection was completed by Aug 30, 2013. Three mother-infant pairs were lost from the capsule group because of moving to other places and two mother-infant pairs were lost from the I-salt group because of problems during testing. The baseline characteristics of treatment groups are presented in Table 1. There were no significant differences between groups in any measured variables for mothers or infants at baseline. The median maternal TSH, T₃, T₄ and Tg were all within the reference ranges. Visible goiter was seen in 52% of women at baseline and an additional 23% had palpable goiter.

The median mothers’ age was 22 (20, 25) years, mean body mass index (BMI) was 21.5 (2.3) and mid-upper-arm-circumference (MUAC) was 23.3 (1.7) cm. The median infant age was 5 (3, 7) days. Excluding the five subjects who dropped out the study, there were 50 (49.9%) male and 51 (50.5%) female infants participating in the study. Infants did not differ in any of the anthropometric indices.

As shown in Table 1, there were no significant differences between mothers in the capsule group and I-salt group for TSH, T₃ and Tg at 26 weeks, but T₄ was significantly lower in the I-salt group. Only T₄ (p = 0.014) showed a treatment by time interaction. Combined prevalence of visible and palpable goiter at 26 weeks tended to be higher in the I-salt group although not significant (p = 0.14).

Infants’ TSH and T₄ at 26 weeks were within reference ranges in both groups (Table 2). However unlike mothers, infants’ T₄ was higher in the I-salt group. Similar to baseline, infants were not significantly different in any of the anthropometry indices at 26 weeks (Table 1).

In the VIP tests infants were different (p < 0.02) by treatment for novelty quotient but not for the other VIP measures. A novelty quotient > 0.55 indicates that the infant remembers the familiar face [22, 23]. Based on this classification, 26% of the infants in the capsule and 51% in the I-salt group remembered the familiar face (Table 2).

As shown in Fig 2, maternal goiter decreased from 76% at baseline to 30% at 26 weeks (X² = 7.9, p = 0.02) in the capsule group and from 74% at baseline to 43% at 26 weeks (X² = 21.1,
Maternal TSH was associated with goiter ($X^2 = 5.8$, $p = 0.016$) at baseline but not at 26 weeks when all TSH values were much lower. Because there were few significant differences between groups at 26 weeks based on iodine source, data from both supplementation groups were combined to examine effects of either source of iodine in lactating women and their infants (Table 3). For the group as a whole, the

Table 1. Socio-demographic, thyroid hormones and goiter of mothers and anthropometry of infants at baseline and at 26 weeks.

|                     | Capsule group (n = 50) | I-salt group (n = 51) | p—value | Capsule group (n = 50) | I-salt group (n = 51) | p—value |
|---------------------|------------------------|-----------------------|---------|------------------------|-----------------------|---------|
| **Mothers**         |                        |                       |         |                        |                       |         |
| - Age (years)       | 23 (20, 27)            | 21 (20, 25)           | 0.14    |                       |                       |         |
| - MUAC (cm)         | 23.2 (1.8)             | 23.3 (1.7)            | 0.62    |                       |                       |         |
| - BMI (kg/m$^2$)    | 21.2 (2.5)             | 21.8 (2.0)            | 0.15    |                       |                       |         |
| - Gravidity         | 2 (1.7, 4)             | 2 (1, 5)              | 0.41    |                       |                       |         |
| - Parity            | 2 (1, 4)               | 2 (1, 4)              | 0.38    |                       |                       |         |
| - School years      | 2 (2, 3)               | 2 (1, 3)              | 0.42    |                       |                       |         |
| - $T_3$ (ng/mL)     | 1.3 (0.3)              | 1.4 (0.3)             | 0.06    | 1.14 (0.4)            | 1.25 (0.4)            | 0.17    |
| - $T_4$ (μg/dL)     | 6.7 (5.9, 7.2)         | 6.8 (5.4, 7.2)        | 0.96    | 6.7 (5.5, 8)          | 5.4 (4.8, 6.9)        | 0.02    |
| - $Tg$ (ng/mL)      | 4.2 (1.6, 9.3)         | 4.2 (2.3, 9.5)        | 0.73    | 4.2 (1.8, 9)          | 4.2 (2, 8)            | 0.61    |
| - TSH (μIU/mL)      | 2.6 (1.1, 4.4)         | 1.9 (0.8, 3.2)        | 0.29    | 1.2 (0.9, 2.4)        | 1.1 (0.7, 1.8)        | 0.16    |
| - Goiter            |                        |                       |         | 0.96                  |                       | 0.14    |
| - Grade 0           | 12 (24%)               | 13 (26%)              |         | 35 (70%)              | 29 (57%)              |         |
| - Grade 1           | 11 (22%)               | 12 (23%)              |         | 15 (30%)              | 21 (41%)              |         |
| - Grade 2           | 27 (54%)               | 26 (51%)              |         | 0                     | 1 (2%)                |         |
| **Infants**         |                        |                       |         |                       |                       |         |
| - Age (days)        | 5 (3, 7)               | 6 (5, 8)              | 0.06    |                       |                       |         |
| - WAZ               | -0.29 (1.23)           | -0.43 (1.37)          | 0.59    | -0.35 (1.3)           | -0.4 (1.1)            | 0.82    |
| - WLZ               | -0.38 (1.9)            | -0.44 (1.14)          | 0.80    | 0.38 (1.1)            | 0.46 (1.2)            | 0.72    |
| - LAZ               | -0.28 (1.44)           | -0.41 (1.47)          | 0.66    | -0.8 (1.3)            | -1.1 (1.1)            | 0.35    |
| - HCAZ              | 1.45 (0.4, 1.9)        | 1.09 (0.1, 1.7)       | 0.16    | 0.73 (0.24, 1.62)     | 0.78 (0.24, 1.60)     | 0.34    |

Data are median, mean, or percentage. Continuous data were analyzed using T-Test and Mann-Whitney U test, categorical variable was analyzed using Chi-square test and normality was checked using Kolmogorov-Smirnov. $T_3 =$ triiodothyronine, $T_4 =$ Thyroxin, $Tg =$ thyroglobulin, TSH = thyroid stimulating hormone, MUAC = mid- upper-circumference, BMI = body mass index

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p < 0.001) in the I-salt group. Maternal TSH was associated with goiter ($X^2 = 5.8$, $p = 0.016$) at baseline but not at 26 weeks when all TSH values were much lower.

Because there were few significant differences between groups at 26 weeks based on iodine source, data from both supplementation groups were combined to examine effects of either source of iodine in lactating women and their infants (Table 3). For the group as a whole, the

Table 2. Thyroid hormones and visual information processing variables of infants at 26 weeks.

|                     | Capsule group (50) | I-salt group (51) | p—value |
|---------------------|--------------------|-------------------|---------|
| TSH (μIU/mL)        | 0.56 (0.1, 1.6)    | 0.51 (0.1, 1.3)   | 0.92    |
| $T_4$ (μg/dL)       | 10.8 (8.7, 13.6)   | 13.9 (10.6, 17.6) | 0.003   |
| VIP variables of infants |                 |                   |         |
| Total shift         | 4 (2, 6)           | 4 (2, 6)          | 0.92    |
| No. of looks        | 6 (5, 7)           | 6 (5, 8)         | 0.09    |
| Average look        | 9.6 (5.8)          | 8.0 (5.2)        | 0.18    |
| Longest look        | 22.5 (17.8)        | 19.5 (19.1)      | 0.21    |
| Total novelty quotient (> 0.55) |        |                   | 0.02    |
| - Yes               | 13 (26%)           | 26 (51%)         |         |
| - No                | 37 (74%)           | 25 (49%)         |         |

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median maternal TSH and prevalence of maternal goiter decreased significantly after 26 weeks of access to iodine. Total goiter declined from 76% at baseline to 37% at 26 weeks (p < 0.001).

Infants in the combined groups (Table 3) did not differ in WAZ and HCAZ but there were significant differences in WLZ and LAZ at 26 weeks compared to baseline. The means of these anthropometric Z-scores were above the cutoff level set for nutritional insult (≤ -2 Z score); however, 14% were stunted, 13% were underweight and 8% were wasted at baseline and at 26 weeks, 18% were stunted, 11% were underweight and 3% were wasted. Males had significantly better scores than females for total novelty quotient ($X^2 = 4.54$, p = 0.033) and number of looks ($X^2 = 6.6$, p = 0.011).

Table 4 shows significant predictors of the VIP variables using stepwise multiple linear regression analysis. One unit increase in infant urinary iodine adjusted for creatinine at 26 weeks predicted 0.24 more total shifts by the infant. Maternal T3 at 26 weeks predicted increased total shifts of 0.21 units and maternal T4 at baseline predicted increased total shifts of 0.23 units. About 12% of the variation in total shift was explained by the model.

A one unit increase in maternal T3 at baseline predicted a 0.26 unit decrease in average look and one unit increase in Tg at baseline was associated with a 0.24 unit decrease in average look. The model explained 10.4% of the variation in average look.

For longest look, one unit increase in maternal Tg at baseline was associated with a 0.25 unit decrease in longest look and one unit higher maternal T3 at baseline was associated with a 0.24 unit decrease in longest look; 10% of the variation in longest look was explained by the model.

Multiple classification analysis results (Table 5) showed that several variables significantly predicted infants NQ including sex (p = .027), LAZ (p = .015), BMIC (p = .042), maternal education (p = .004) and HFIAS score (p = .042). Taking the beta ($\beta$) coefficients as indicators, household food security, women education and LAZ were strongest predictors compared with the other four predictors included in the model. Infants from food secure households had

![Fig 2. Prevalence of goiter in women at baseline and 26 weeks in the capsule vs salt groups (n = 101).](https://doi.org/10.1371/journal.pone.0223348.g002)
significantly higher NQ (0.638) compared to infants from food insecure households and infants from literate mothers had significantly higher NQ (0.55) compared to infants from illiterate mothers. Likewise, infants whose LAZ score was above -1 SD had significantly higher NQ (0.55) compared to infants with lower LAZ scores. Moreover, infants whose mothers had BMI \(< 120 \mu g/L and infants who were male had significantly higher NQ than their counterparts.

### Discussion

We examined the effects of 26 weeks of iodine supplementation on thyroid hormones, thyroglobulin and goiter of lactating women and on T4, TSH and VIP in infants. At baseline mothers and infants in the two supplemented groups (capsule and I-salt) were not significantly different in any of the biomarkers or other characteristics.

The study population had been known to have severe iodine deficiency for decades [15–18, 26]. The current study was conducted in 2013, following the rapid implementation of a salt

| Table 3. Thyroid hormone biomarkers of mothers and anthropometry of infants at baseline and at 26 weeks in participants who received either iodine capsule or iodized salt (n = 101). |
|----------------------------------|-----------------|-----------------|-----------------|
|                                   | Baseline        | 26 weeks        | p-value         |
| **Mothers**                      |                 |                 |                 |
| TSH (μIU/mL)                     | 1.9 (0.8, 3.7)  | 1.2 (0.7, 2.1)  | 0.001           |
| T4 (μg/dL)                       | 6.8 (5.7, 7)    | 5.8 (5.1, 7.5)  | 0.14            |
| T3 (ng/mL)                       | 1.33 (0.3)      | 1.27 (0.3)      | 0.19            |
| Tg (ng/mL)                       | 4.2 (2.3, 9.5)  | 4.2 (2, 7.1)    | 0.266           |
| Goiter                           |                 |                 |                 |
| - Grade 0                        | 25 (24.7%)      | 64 (63.4%)      | < 0.001         |
| - Grade 1                        | 23 (22.8%)      | 36 (35.6%)      |                 |
| - Grade 2                        | 53 (52.5%)      | 1 (1.0%)        |                 |
| **Infants**                      |                 |                 |                 |
| WAZ                              | -0.34 (-0.9, 0.4) | -0.4 (-0.9, 0.4) | 0.91            |
| WLZ                              | -0.4 (-1.0, 0.5) | 0.4 (-0.4, 1.2) | < 0.001         |
| LAZ                              | -0.3 (-1.4, 0.7) | -0.9 (-1.7, -0.1) | 0.006           |
| HACZ                             | 1.3 (0.2, 1.9)  | 0.7 (0.2, 1.6)  | 0.88            |

Data are median (IQR), mean (SD) and percentages. Continuous data are analyzed using paired samples T-test or Wilcoxon, proportions (percentages) were analyzed using Chi-square test.

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significantly higher NQ (0.638) compared to infants from food insecure households and infants from literate mothers had significantly higher NQ (0.55) compared to infants from illiterate mothers. Likewise, infants whose LAZ score was above -1 SD had significantly higher NQ (0.55) compared to infants with lower LAZ scores. Moreover, infants whose mothers had BMI ≥ 120 μg/L and infants who were male had significantly higher NQ than their counterparts.

### Discussion

We examined the effects of 26 weeks of iodine supplementation on thyroid hormones, thyroglobulin and goiter of lactating women and on T4, TSH and VIP in infants. At baseline mothers and infants in the two supplemented groups (capsule and I-salt) were not significantly different in any of the biomarkers or other characteristics.

The study population had been known to have severe iodine deficiency for decades [15–18, 26]. The current study was conducted in 2013, following the rapid implementation of a salt

| Table 4. Multiple regression predicting VIP tests using stepwise criteria (n = 101). |
|----------------------------------|-----------------|-----------------|-----------------|
| **Dependent variables**          | **Predictors**  | Beta            | p               |
| Total number of shifts           | Infant UIC/Cre (μg/L) at 26 weeks | 0.24            | 0.02            |
|                                 | Maternal T3 (ng/mL) at 26 weeks | 0.21            | 0.043           |
|                                 | Maternal T4 (μg/dL) at baseline | 0.23            | 0.029           |
|                                 | Adjusted R square = 0.12          |                 |                 |
| Average look                    | Maternal Tg (ng/mL) at baseline | -0.26           | 0.008           |
|                                 | Maternal Tg (ng/mL) at baseline | -0.24           | 0.015           |
|                                 | Adjusted R square = 0.104         |                 |                 |
| Longest look                    | Maternal Tg (ng/mL) at baseline | -0.25           | 0.012           |
|                                 | Maternal Tg (ng/mL) at baseline | -0.24           | 0.016           |
|                                 | Adjusted R square = 0.10          |                 |                 |

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iodization program in 2012. Presumably as the result of the program implementation, our study population showed higher UIC and BMIC than previous studies [12].

Following 26 weeks of iodine supplementation, plasma TSH in lactating women significantly decreased by 54% in the capsule group and by 42% in the I-salt group compared to baseline. However, these improvements in TSH were not significantly different between the two groups of mothers. Santiago and colleagues reported there was no significant difference in maternal TSH between pregnant women who were supplemented with iodized salt and those who were supplemented with either 200 μg iodide per day or 300 μg iodide per day [27]. A study that compared a control group to women who received 300 μg iodine as potassium iodide during pregnancy and lactation reported that plasma T4 and T3 were significantly higher in the supplemented women; TSH was significantly higher in the control group [28].

Because a majority of a breast feeding mother’s iodine intake is secreted into the breast milk, an exclusively breastfed infant may have better iodine status than the mother [29]. Most of the infants in the current study had normal T4 and all had TSH within the normal range.

Thyroid hormones did not respond consistently to iodine supplementation in this study. Maternal TSH improved significantly following iodine supplementation, T3 and T4 showed

Table 5. Result of multiple classification analysis (MCA) for the key determinants of infants novelty quotient NQ by selected predictors and covariates (n = 101).

| Variable                      | n   | Unadjusted mean | Eta (η) | Adjusted mean | Beta (β) | Sig   |
|-------------------------------|-----|-----------------|---------|---------------|----------|-------|
| Child sex                     |     |                 |         |               |          |       |
| Male                          | 50  | .548            | .214    | .546          | .205     | .027* |
| Female                        | 51  | .468            |         | .476          |          |       |
| LAZ score                     |     |                 | .207    | .240          | .015b    |       |
| < -1 SD                       | 48  | .471            |         | .465          |          |       |
| ≥ -1 SD                       | 53  | .548            |         | .554          |          |       |
| WAZ score                     |     |                 | .046    | .173          | .274     |       |
| < -1 SD                       | 24  | .523            |         | .450          |          |       |
| ≥ -1 SD                       | 77  | .503            |         | .525          |          |       |
| Infant T4 (μg/dL)             |     |                 | .15     | .06           | .326     |       |
| < 8.0                         | 9   | .56             |         | .546          |          |       |
| > 8–23                        | 92  | .51             |         | .504          |          |       |
| BMIC (μg/L)*                  |     |                 | .196    | .211          | .042b    |       |
| < 120                         | 55  | .487            |         | .485          |          |       |
| ≥ 120                         | 46  | .571            |         | .560          |          |       |
| Women’s education             |     |                 | .207    | .277          | .004*    |       |
| Illiterate                    | 38  | .458            |         | .441          |          |       |
| Literate                      | 63  | .537            |         | .548          |          |       |
| HFIAS score**                 |     |                 | .260    | .339          | .042b    |       |
| Food secure                   | 58  | .621            |         | .638          |          |       |
| Mild food insecurity          | 12  | .559            |         | .605          |          |       |
| Moderate food insecurity      | 20  | .486            |         | .480          |          |       |
| Severe food insecurity        | 11  | .472            |         | .472          |          |       |

r = 0.59; R² = 0.331; grand mean = 0.51

* Significant α .01  
  b Significant α .05

*BMIC–Breast milk iodine concentration  
  **HFIAS–Household food insecurity access scale

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decreased values after supplementation but the changes were not significant. The demands for iodine during lactation may reduce thyroid hormones if iodine intakes are marginal. However, at baseline 90% of the women had T₃ and 82% had T₄ within the reference range; values were quite similar at 26 weeks when 86% of women had T₃ and 77% had T₄ within the reference range and there was no change in Tg.

In a double-blind, randomized, controlled trial conducted in Morocco, a single capsule of 400 mg of iodine as iodized oil given to the breast-feeding mother or a single oral supplement of 100 mg of iodine given to the newborn baby did not produce a significant change in maternal or infant TSH or T₄ at 3 months, 6 months or 9 months after supplementation [30]. Such inconsistent thyroid hormone response to iodine supplement was also reported in other studies [28, 31]. For example, pregnant women showed no change or an increase in maternal TSH following iodine supplementation of 50 μg daily, 200 μg daily or 300 μg daily for six months [27, 32]. In two other randomized controlled trials, maternal free T₄ concentration decreased in both controls and treated groups and no difference was observed between groups [33].

The inconsistency of thyroid hormone response to iodine supplement could be because circulating thyroid hormones depend on iodine deficiency status, on increased needs during pregnancy and lactation and on age. A recently published review on biomarkers of iodine status suggested that T₄ and T₃ are good indicators of iodine status in areas of severe iodine deficiency and TSH was considered a sensitive marker in newborns [34]. Thyroglobulin has been recommended as a more sensitive indicator than TSH and T₄ in iodine deficient and excess areas in children because thyroglobulin is synthesized only in the thyroid [34, 35]. Another group commented that goiter rate, Tg and TSH are good markers to assess iodine status in a population but thyroid hormones are fairly imprecise to detect iodine deficiency [36]. However, unlike for children, no cutoff point for T₄ for adults has been established related to iodine status [35]. In one study, serum Tg was correlated with severity of iodine deficiency assessed by urinary iodine [34]. In the current study Tg didn’t show any correlation with urinary iodine. When thyroid markers were categorized to examine association with goiter rate, TSH at baseline was the only variable that showed significant association with goiter (X² = 5.8, p = 0.016).

In the present study maternal goiter significantly decreased in both capsule and I-salt groups despite six months of lactation. In goitrous children age 6 to 12 years in CÔte d’Ivoire, a 200 mg iodine supplement produced a significant and continuous decrease in thyroid gland volume over 30 weeks. Similar to our results there was no significant change in serum T₄ but TSH decreased significantly in the CÔte d’Ivoire study [37]. In contrast, in a severely iodine deficient area in Senegal, 480 mg of iodine given as iodized oil to adults significantly increased T₄ and decreased rate of goiter, T₃ and TSH [38].

The effect of iodine supplement on maternal thyroid disorders and in children’s neurocognitive development in severely iodine deficient areas is well established [39]. It is unclear however, the extent to which maternal iodine supplementation in areas of mild-to-moderate iodine deficiency may affect cognitive function of their offspring [10, 40]. Children of pregnant women who had received iodine supplementation from their first trimester compared with children of unsupplemented women showed no difference in mental development index measured by the Bayley Scales of Infant Development (BSID) at the age of two years [28]. However, the children of supplemented mothers had higher scores on the psychomotor development index of the BSID. On the other hand, a prospective study reported that children of mothers exposed to mild-moderate iodine deficiency showed attention deficit and hyperactivity disorders [41].

In our study, the look duration (length of the longest look) during the familiarization phase (habituation phase) as a measure of attention and processing speed was not significantly
different between groups. The mean of the longest look duration in both groups was 21.1 (18.0)s. Shorter fixation indicates better attention and faster processing speed and is a predictor of later childhood IQ [23, 42]. In the current study more than 70% of the infants in both groups were below the mean with 26% below 10s. The average look duration, which indicates the speed at which recognition of the familiar stimulus is achieved [24], likewise was not significantly different between groups.

Although there is no established standard for number of looks a child would make during the habituation phase or trials to criterion, infants were not significantly different between groups. However, the longer it takes an infant to reach the habituation criterion (i.e., the greater the number of looks before criterion is reached), the longer it is taking the infant to learn the stimulus. In this regard 43% of the infants had number of looks above the median and 35% had number of looks below the median of 6 (5, 7.7); those with few looks learned the habituation face more quickly [43]. In total (both groups combined) the mean NQ was 0.51 (0.19) which is similar with the mean novelty preference reported by Oakes and Kovack-Lesh [44] and is higher than the mean NQ reported by Kennedy and colleagues eight years earlier (before introduction of iodized salt) in infants 6–8 months age in a similar area in Ethiopia [21].

The VIP paradigm has been used to assess infants’ and children’s information processing speed, recognition memory and attention in relation to various factors. Infants whose mothers had high blood DHA concentration showed faster information processing speed over the first year as well as less distractibility in the second year than those whose mothers had low blood DHA [45]. A study conducted in Ethiopian infants showed that growth was associated with increased performance in the VIP [21] paradigm. In another study, increased prenatal alcohol exposure was correlated with longer looks, indicating slower information processing [46]. Similarly, exposure to toxic substances during pregnancy resulted in poorer recognition memory during the first year of the child’s life [47].

In the present study we have examined possible associations and predictor variables of VIP of infants. Accordingly, NQ was affected by infants’ TSH, T4, anthropometry indices, maternal BMI, and education as well as household food security state. Moreover, male infants remembered the familiar face and learned faster than female infants. Not much evidence is available regarding iodine intake on infant cognition performance by sex. One study done in 1982 in school children from an area of endemic goiter reported that girls showed better cognition improvement as a result of iodine supplement than boys [48]. Another study from iodine sufficient and mildly iodine deficient areas reported similar results [49]. However in this study, because male and female infants didn’t differ in either UIC, T4 or TSH, the difference observed in NQ could be independent of iodine intake. For instance, it could be related to amount of breast milk consumed or infant stimulation by caretakers [50, 51].

Based on the multiple regression analyses maternal thyroid hormones at baseline were significant predictors of some of the VIP variables. Concentrations one week after delivery may reflect the maternal thyroid hormone status during pregnancy which might have affected the child’s brain development. However, none of the VIP variables were significantly different by maternal goiter grade at baseline. Moreover, comparison of thyroid biomarkers and VIP data for mothers with UIC above and below 100 ug/L were not significantly different.

We performed reliability checks for longest look and novelty preference. The correlation for longest look was 0.78. However, correlations for novelty preference were lower. This poor reliability was likely due to the quality of the recorded video and the fact that the stimuli were presented on a single laptop screen, which made it difficult to tell on the recorded video whether the child was looking at one of the pictures or farther to the side and thus away from the pictures. For the future this problem could be solved by having a better video recording and/or by using two laptop screens placed farther apart.
A limitation of this study was the relatively small sample size which was calculated based on outcomes for the original study [12]. Based on the post-hoc power calculation, the null association between the intervention and T₃ and T₄ may be due to insufficient power.

In conclusion, novelty quotient of infants was predicted in the VIP paradigm by higher maternal education, BMIC and LAZ, as well as by better household food security, and male sex. Attention to iodine status and other modifiable factors has the potential to impact optimal child development.

Although the study population had a long history of iodine deficiency, salt iodization had been implemented during the previous year. The present study found a median UIC at baseline that indicated community level iodine sufficiency based on WHO standards with median UIC less than 100 μg/L classified as iodine deficient. The median UIC at baseline for the mothers was 107 (71, 161) μg/L and for the infants was 218 (108, 356) μg/L. Nonetheless, on the basis of UIC, 46% of the women and 21% of the infants would be classified as at least mildly iodine deficient at baseline [4]. But effects of maternal iodine supplementation on thyroid function may have been muted because nearly half of the women were iodine sufficient. Biomarkers measured in this study would not be expected to be as responsive to iodine supplementation in iodine sufficiency. In the present study, only TSH and goiter showed improvement with supplementation in women and the only biomarker that showed association with goiter was TSH. Further investigation of TSH as a biomarker for assessment of iodine nutrition even in adults may be warranted. Moreover, because several minerals including iron and selenium play essential roles in thyroid hormone metabolism, it is important to assess the impact of these nutrients in both mothers and their infants [52].

This study confirmed that iodized salt showed similar effects on iodine status and thyroid hormones as a daily iodine capsule supplement if the salt is adequately iodized and closely monitored. Data for the present study were collected within a year after the salt iodization program was launched in Ethiopia, so it is too early to comment on the long term effects of an adequate supply of iodine to the population or on the quality control of the iodization process. Large scale and repeated monitoring studies likely are warranted in the interest of optimizing iodine status of the Ethiopian population.

**Supporting information**

S1 File. Study protocol.  
(PDF)

S2 File. CONSORT checklist.  
(DOC)

S1 Dataset. Data.  
(PDF)

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