Levothyroxine treatment and gastric juice pH in humans: the proof of concept

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ORIGINAL ARTICLE

Abstract

Purpose Despite the absorption of oral thyroxine (T4) occurs in the small bowel, several patients with gastric disorders show an increased need for T4. In vitro evidence suggested that medium pH variations interfere with T4 dissolution. This study was aimed at finding the proof of concept of a direct relationship between the minimal effective dose of T4 and the actual gastric juice pH.

Patients and methods Among 311 consecutively thyroxine-treated patients, 61 bearing Hashimoto’s thyroiditis (52 F/9 M; median age = 51 years) who complained persistent dyspepsia and/or upper abdominal symptoms following a noninvasive workup for gastrointestinal disorders, underwent EGDS with multiple biopsies and gastric juice pH measurement. All patients accepted to take thyroxine in fasting conditions, abstaining from eating or drinking for one hour.

Results Thyroxine requirement increased along with the rising gastric pH ($\rho = 0.4229; p = 0.0007$). A multivariate analysis revealed that gastric pH was, beside body mass index, the far more important independent variable in determining the effective dose of T4 ($\rho = 0.001$). The ROC curve revealed that the pH threshold for an increased thyroxine requirement was at 2.28, being the AUC by 78%. Subdividing patients by the histologic findings, it appeared a significant increase ($p = 0.0025$) along with the progressive damage of gastric mucosa.

Conclusion The in vivo measurement of gastric pH highlighted its key role in determining the minimal effective dose of oral T4 and may explain the interference of food, of some drugs and gut disorders on levothyroxine treatment.

Keywords Hypothyroidism · Levothyroxine dose · Levothyroxine malabsorption · Gastric juice pH · Helicobacter pylori · Atrophic gastritis

Introduction

In the era of Precision Medicine [1] the factors affecting the minimal effective dose of any drug must be considered when treating patients. Levothyroxine sodium tablet ranks among the most prescribed drugs worldwide and represents a critical-dose drug, as little variations in the blood concentration may cause treatment failure, as well as iatrogenic thyrotoxicosis [2–5]. Despite its narrow therapeutic window imposes a careful dose titration, an estimated rate of 30–60% treated patients still shows an out-of-target TSH [6], and side effects are described [7, 8]. An individualized thyroxine treatment is therefore advisable and requires the knowledge of thyroxine biochemical and pharmacologic characteristics [9–13] as well as patients’ anthropometric data and habits [14–16]. The lean body mass is a major determinant of thyroxine dose [14] but, in clinical practice, the physicians usually choose the dose based on patient weight or BMI [5]. Furthermore, to obtain the
therapeutic effect with the minimal effective dose of T4, physicians should also take into account the use of interfering foods and drugs [17, 18]. Oral thyroxine preparation is a sodium salt whose linear absorption rate in the blood lasts 60 to 90 min [19], then reaching a plateau. A seminal paper from Hays MT, demonstrated that the absorption of oral thyroxine occurs in the small bowel with different percentages [11]. Therefore, that thyroxine malabsorption may occur in patients with celiac disease, lactose intolerance, short bowel syndrome, or giardiasis is somewhat expected [17]. Nevertheless, a key role for the stomach in the oral T4 absorption process has also been suggested on a clinical ground, since several patients with gastric disorders show an increased need for thyroxine. Evidence of an increased T4 requirement has been provided in patients with Helicobacter pylori [H. pylori] infection [20, 21], chronic atrophic gastritis [21], and gastroparesis [22] as well as in patients chronically treated with proton pump inhibitors [23]. Most of these conditions feature an altered gastric juice pH since the different gastritis patterns can deeply affect gastric acid output mainly when active inflammation [i.e. H. pylori infection] or atrophy involve the acid-producing oxyntic glands [24–28]. As described for other drugs [29], gastric pH has been suggested to be a critical factor for both disaggregation and dissolution of thyroxine [30, 31]. However, the proof of concept of a relationship between the minimal effective dose of T4 and the actual gastric pH needs a direct assessment and this represented the aim of our study. Primary outcome of the study was to correlate pH values of fasting gastric acid secretion with the dose required to obtain target serum TSH in thyroxine-treated hypothyroid patients.

Patients and methods

Patients

Patients were recruited in a tertiary Endocrinology unit between 2018 and 2020 among a sequentially examined cohort of hypothyroid outpatient, with a definite diagnosis of Hashimoto’s thyroiditis, in need for levothyroxine treatment according to the ATA guidelines (n = 311) [3] (Fig. 1). Patients with goiter and/or who underwent thyroid surgery and/or in follow-up for thyroid cancer, with non-thyroidal illnesses or severe chronic diseases, obese (BMI > 30 kg/m²), with already known diagnosis of intestinal disorders and/or previous GI surgery and pregnant or nursing were not enrolled. Furthermore, patients not compliant with levothyroxine treatment schedule, with supervening diagnosis of intestinal and/or neoplastic disorders and/or in need for not replaceable substances or drugs, known to interfere with thyroxine metabolism/absorption (see for rev ref [18]) have been progressively excluded from the study (Fig. 1).

Design of the study

Overall, 222 out of 311 patients reached the target TSH (0.8–2.5 mU/l) within the second control visit with a median thyroxine dose of 1.31 µg/kg body weight. Among these patients reaching target serum TSH, 48 complained persistent dyspepsia and/or upper abdominal symptoms, thus needing a gastroenterological evaluation (Fig. 1). Among the 89 patients not reaching target TSH, the dose of T4 has been progressively increased, until attaining the desired TSH value. However, 23 patients were excluded because of non-mendable interference issues or ascertained noncompliance (Fig. 1), while the other 66 patients underwent gastroenterological evaluation due to the persistent dyspepsia and/or the upper abdominal symptoms. Therefore, one hundred and fourteen patients underwent diagnostic workup for gastrointestinal diseases in a Gastroenterological referral center. At the end of non-invasive diagnostic workup, 44 patients were excluded from the study (18 patients with malabsorptive or inflammatory intestinal disorders, 16 refusing endoscopy, and ten patients owing symptoms resolution) (Fig. 1). The remaining 70 patients, needed to complete the diagnostic workup because of long-lasting dyspepsia or upper abdominal symptoms,
persisting despite an appropriate treatment [32]. These symptoms might have occurred isolated or concomitant with chronic unexplained micro- and macrocytic anemia and/or long lasting or refractory gastrointestinal reflux disease [32, 33]. These patients underwent upper endoscopy with multiple biopsies and gastric juice pH evaluation. The protocol was consistent with the principles of the Declaration of Helsinki and the study has been performed within the usual diagnostic workup of each single patient, upon written informed consent. The study was approved by the Sapienza University Ethical Committee. Upon endoscopic evaluation, nine additional patients were excluded (see Fig. 1). Therefore, our final study group consisted in 61 patients (52 F/9 M; median age = 51 years; IQR = 41–65 years). STROBE guidelines were followed to describe this study.

Standardization of thyroxine treatment

Following the initial clinical examination, patients have been treated with the same brand of levothyroxine sodium in tablet formulation with an age- and weight-tailored dose. As previously described [34, 35], all subjects were requested to follow a tight schedule of thyroxine ingestion, ensuring to take the tablet while in fasting condition and then abstaining from eating or drinking anything other than water for at least 1 h after T4 ingestion [16, 19]. Patient’s compliance with treatment schedule was checked by a questionnaire, accepted at the first visit and confirmed at every examination. Following a starting T4 dose of 1.0 µg/kg body weight/day, dose adjustments were made based on TSH levels in each patient and tested every three months (median follow-up = 34 months). Once reached target TSH, stability of its value had been confirmed at least in two TSH measurements.

A reference group was used to identify a threshold beyond whom an increased need for thyroxine in hypothyroid patients could be defined. This reference group consists in 140 hypothyroid with similar anthropometric characteristics, who followed exactly the same schedule of T4. These patients met all the criteria described for the study group and chiefly they did not show evidence of signs or symptoms of malabsorption nor dyspepsia, as previously described [34, 35]. The median daily dose of T4 required to obtain the target TSH (0.8–2.5 mU/l) in this reference group was 1.27 µg/kg body weight (IQR = 1.19–1.33) or a mean dose of 1.27 ± 0.12 (SD) µg/kg body weight. Therefore, the normal thyroxine requirement was calculated considering the mean daily dose of T4 in our reference group to which 1.96 standard deviations were added. Hence, in our hypothyroid patients, we considered increased a need for thyroxine overcoming 1.51 µg/kg body weight/day.

Methods

Thyroid diagnostic workup

The diagnosis of Hashimoto’s thyroiditis was based on the presence of high titer anti-thyroperoxidase antibodies [anti-TPO Abs], a characteristic ultrasonographic pattern and hypothyroidism [serum TSH > 10 mU/l]. Serum TSH and FT4 and anti-TPO Abs levels were assayed by commercial assays [Thermo Scientific, BRAHMS TSH RIA, Germany].

Gastrointestinal diagnostic workup

As a policy of our Center, all patients undergo an accurate anamnestic interview and those with clinical suspicion and gastrointestinal signs or symptoms are referred to the abovementioned Gastroenterological center to enter a diagnostic workup, as previously described [17]. As shown in Fig. 1, 114 patients underwent a noninvasive gastrointestinal screening. Briefly, the presence of lactose intolerance was evaluated by lactose H2 breath test; the first line of celiac disease investigation was based on the Elisa measurement of serum anti-transglutaminase autoantibodies (AXA Diagnostics Srl, Pomezia, Italy; cut-off title <10 U/ml). Serum gastrin 17 levels were assayed by chemiluminescence assay (Biokit Healthcare, Helsinki, Finland normal range, 13–115 pg/ml) and serum anti-parietal cells antibodies were measured by Autoantibody test system (A. Menarini Diagnostics s.r.l.—Firenze, Italy; cut-off title< 40).

Upper endoscopy

All patients underwent upper endoscopy in the morning, fasting from at least eight hours. All endoscopy procedures were performed between 8:00 and 10:00 a.m. All subjects were sedated using intravenous midazolam [range to 2–5 mg]. Following exclusion of macroscopic gastro-duodenal lesions, 5 biopsies were collected in the stomach using a standard biopsy forceps, two from antrum, one from the incisura angularis, two from mid-body and two from fundus during retroversion, in accordance with the updated Sydney System [36].

Gastric juice aspiration and gastric pH value assessment

Immediately after the insertion of the endoscope into the stomach, 5 mL of gastric juice were aspirated in fundus and mid-body [greater curve] by means of a sterile teflon catheter and collected into a sterile trap connected with the suction line of the endoscope. After collection, the pH of gastric juice was measured using a glass electrode pH–meter and afterwards titrated with a 1 N solution of NaOH to evaluate the actual H+ concentration in each sample.
Histologic assessment

All gastric and duodenal biopsies were fixed and stained with haematoxylin and eosin for histopathologic examination. Modified Giemsa staining was used to evaluate *H. pylori* infection. Assessment of gastritis patterns was done according to the updated Sydney scoring system [36] [0 = absence, 1 = mild, 2 = moderate, 3 = severe] evaluating the acute inflammation (neutrophil infiltrate), chronic inflammation (mononuclear cell infiltrate), glandular atrophy, intestinal metaplasia, *Helicobacter pylori* infection [36]. Atrophy of the fundic mucosa was defined as focal or complete replacement of oxyntic glands by metaplastic pyloric or intestinal glands. An expert pathologist blinded for patient’s clinical diagnosis performed the histopathologic examination. Based on histological reports, patients were divided in 5 groups. Group 0 when no alterations in the whole gastric mucosa were found; Group 1, when only antral superficial gastritis was present; Group 2, when a superficial chronic pangastritis was described; Group 3, in the presence of active superficial pangastritis; Group 4, when atrophic chronic gastritis was found.

Statistical analysis

A descriptive analysis was performed calculating median and interquartile range (IQR) for continue variables and absolute and relative frequencies for qualitative variables. Chi-squared and Fisher’s tests were used to compare proportions of different groups. After testing for normality with Shapiro-Wilk test, Student’s *t* test and Pearson *r* test or analogous nonparametric Mann–Whitney and Spearman *ρ* tests were used to compare different groups or assess correlation between variables, respectively. To assess the role of pH in determining thyroxine dose, a multiple regression analysis was built, including, beside pH, also age, sex, BMI, TSH, FT4, histologic pattern of gastritis and histologic detection of *H. Pylori*. For purpose of analysis, given the skewed distribution of the thyroxine dose, generalized linear regression analysis was carried out. Coefficient, standard error (SE) and *p* values were calculated for each of the independent variables. Statistical significance was defined as a two-sided *p* value < 0.05 for all analyses, which were carried out using STATA statistical software, version 15 (Stata Corp., College Station, TX, USA). Prof. Corrado De Vito, coauthor of the manuscript, analyzed the data.

Results

The anthropometric and thyroid biochemical features in the whole study group has been summarized in Tables 1a and 1b. Median pH in gastric juice in the whole sample was 2.28 (IQR = 1.47–6.4) and median H⁺ concentration was 52 mEq/l (IQR = 5–76). In the whole sample the median dose of T4 required to obtain the desired TSH was 100 μg/day (IQR = 86–125) that following normalization by individual weights was equal to 1.47 μg/kg body weight/day (IQR = 1.32–1.72).

A direct correlation between the required T4 dose and an increased pH has been observed in the whole sample (*ρ* = 0.4229; *p* = 0.0007) (Fig. 2a). This finding was confirmed.
by the inverse correlation between the dose of T4 and the actual H⁺ juice concentration ($\rho = -0.3987; \ p = 0.0018$).

When analyzing the whole sample of the patients, the expected minimal effective dose ($1.27 \pm 0.24 \ \mu g/kg/day$), as in the reference group, was sufficient to obtain the target serum TSH in 36 patients. The remaining 25 patients required a higher T4 dose ranging from 1.55 to 2.42 $\mu g/kg/day$. We further analyzed the dose of thyroxine by subdividing our whole sample based on gastric pH in two groups: group A ($n = 29$) with pH $\leq 2.0$, representing the normal fasting pH in healthy human stomach [37–39], and group B ($n = 32$) with pH $> 2.0$. As shown in Table 1b, the anthropometric characteristics and thyroid function tests were similar in these two groups.

In Group A, the median required T4 dose was $1.39 \ \mu g/kg/day$ (IQR = 1.26–1.47 $\mu g/kg/day$), being 23% lower than the dose of $1.71 \ \mu g/kg/day$ (IQR = 1.46–1.94 $\mu g/kg/day$) required by patients of group B ($p < 0.0001$) (Fig. 3). The distribution of patients requiring an increased dose was uneven inside the two pH-related groups. In fact, two patients out of 29 (7%) in the group A (normal pH) needed an increased T4 dose as compared with 23/32 patients (72%) in the group 2 (increased pH). The ROC curve (using $1.51 \ \mu g/kg/day$ as cut-off) revealed that the pH threshold that discriminated patients with increased thyroxine requirement was at 2.28, being the area under the curve by 78% (Fig. 4).

### The dose of thyroxine in relation with gastritis patterns

The need for thyroxine was then analyzed in relation with the histologic grading of mucosal gastric alterations. Overall, the histologic characterization of gastric mucosa revealed that: (a) 4/61 (6.6%) patients had minimal or no alterations (Group 0); (b) 8/61 (13.1%) patients had mild superficial chronic gastritis of the antrum (Group 1); (c) 20/61 (32.7%) patients had mild superficial chronic pangastritis (Group 2); (d) 7/61 (11.5%) had H. Pylori-related moderate/severe active superficial chronic pangastritis, (Group 3); (e) the remaining 22/61 patients (36.1%) showed gastric atrophy (Group 4). In this latter group, 18 out of 22 (82%) patients had an H. Pylori-related atrophic pangastritis while the remaining four patients (17%) had an autoimmune fundus/corpus-restricted atrophic gastritis. Table 2a describes the features of gastric function and immunology of the four groups. As expected from the literature [24–27, 36, 37], gastric pH was similar in the Groups 0, 1 and 2 (median pH 1.70 vs 1.39 vs 1.61, respectively; $p = 0.1291$). In contrast, Group 3 and 4 had significant higher gastric pH than 0 + 1 + 2 groups, again without differences between their median values ($p = 0.1347$). Therefore, statistical analysis
Table 2 a Gastric features of patients subdivided by histologic grading of gastric damage. b Characteristics of patients subdivided by positivity or negativity of H. pylori infection

(a)

| Characteristics of patients subdivided by positivity or negativity of H. pylori | n | pH  | [H+] (mEq/l) | H. pylori infection | APCAb positivity | Gastrin (pg/ml) |
|---|---|---|---|---|---|---|
| Minimal or no alteration (GROUP 0) | 4 | 1.70 (1.64-1.87) | 64 (50-78) | 0 | 0 | 22 (11-30) |
| Chronic superficial mild antal gastritis (GROUP 1) | 8 | 1.39 (1.20-1.42) | 83 (74-93) | 0 | 1 | 40 (39-40) |
| Chronic superficial mild pangastritis (GROUP 2) | 20 | 1.61 (1.30-2.09) | 74 (54-83) | 0 | 4 | 21 (12-38) |
| Chronic active superficial moderate/severe pangastritis (GROUP 3) | 7 | 2.44 (2.18-6.87) | 51 (1-63) | 6 | 4 | 253 (144-320) |
| Gastric atrophy (GROUP 4) | 22 | 6.57 (5.96-7.09) | 5 (0-13) | 17 | 18 | 460 (279-561) |

(b)

| Patients (n) | pH  | Age (years) | Weight (kg) | TSH (mU/l) | FT4 (ng/dl) | APCAb positivity | Gastrin (pg/ml) |
|---|---|---|---|---|---|---|---|
| H. pylori – | 37 | 1.62 (1.37-2.18) | 73 (52-84) | 1.23 (0.82-2.03) | 1.25 (1.13-1.32) | 8/29 | 39 (19-105) |
| H. pylori + | 24 | 6.5 (3.80-6.98) | 5 (1-28) | 1.36 (1.05-1.80) | 1.23 (1.15-1.34) | 20/4 | 357 (243-525) |

\[ p < 0.0001 \]

Discussion

Our data represent the first direct evidence of the role of gastric pH in reaching the pharmacologic thyroid homeostasis. In fact, this is the first study to correlate that the effective dose of T4 (p = 0.001) (Table 3).

Multivariate analysis

Beside the expected and recognized role of BMI, the results of multivariate analysis revealed that gastric pH was the far more important independent variable in determining the effective dose of T4 (p = 0.001) (Table 3).

Multivariate evidence of H. pylori infection and levothyroxine requirement

The possible role of H. pylori infection in the subsequent changes of gastric pH, prompted us to analyze whether its distribution may affect the daily dose of oral T4. In the whole sample the prevalence of H. pylori infection was 24 out of 29 patients (83%) from groups 3 and 4 had histologic evidence of H. pylori infection. Amongst the 37 H. pylori-positive patients, only 9 patients showed a pH >2 while, in the group H. pylori-negative, only 2/24 had pH ≤ 2. In the first direct evidence of the role of H. pylori infection was 39.5% (24/61 patients). Interestingly, no infected patients were recognized in the whole sample. The prevalence of H. pylori infection was 66% of those in groups 3 compared with 60% of those in groups 4 (p = 0.0003).

The overall mechanism that impairs intestinal absorption of thyroxine in hypo-chlorhydric patients is matter of further studies.

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debate. Up to now, some evidence indirectly suggested the involvement of pH in shaping the minimal effective oral dose of T4 in vivo [20, 21, 28]. A growing body of in vitro evidence indicates that, similarly to other drugs, the thyroxine molecule is very sensitive to environmental pH variations [9, 10, 29]. In murine models, changes in blood pH may affect the partition between the fast exchanging pool of thyroxine between tissues and the intravascular milieu [40]. From a pharmacologic standpoint, the role of gastric pH was highlighted by Seng et al., who suggested that a bioequivalence in healthy volunteers might differ from the one in patients bearing altered gastric pH [50]. In a clinical study, a more efficient absorption of thyroxine has been observed when vitamin C was simultaneously taken in patients bearing both hypothyroidism and gastritis [51]. In these patients with impaired acid secretion, ascorbic acid may lower gastric pH thus facilitating the T4 absorption process [51]. More recently, a pharmacokinetic study in healthy euthyroid subjects by Chon et al. [52] showed that simultaneous milk ingestion decreases oral levothyroxine absorption. Interestingly, the pH of cow’s milk is ~6.6 and it contains about 1 g of calcium per liter. Both these factors may impair the optimal absorption of LT4, leading to an increased need for therapeutic dose of the hormone [54].

Besides these nutritional interferences, it should be noted that the presence of H. pylori infection may also act as a confounding factor since:

a. gastric juice pH may vary according to the phase of infection. In fact, chronic infection with H. pylori may feature either decreased or increased acid secretion, along with the distribution and/or the severity of gastritis [55].

b. while autoimmune atrophic gastritis is substantially irreversible, H. pylori-related atrophic gastritis is reversible in about 20% of cases [56].

It follows that, when an H. pylori infection occurs in patients treated with levothyroxine, an accurate follow-up of thyroid homeostasis is recommended until eradication has been confirmed.

From a therapeutic standpoint, recent evidence suggested that novel pharmacologic preparations of thyroxine might be more effective than levothyroxine tablet formulation, in patients with impaired gastric acidic output [57]. Indeed, pilot studies have provided evidence that switching from tablet formulation to liquid [58] or softgel T4 preparations

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Table 3 Results of the multiple linear regression to identify predictors of thyroxine dose

| Variable                          | Coefficient | SE a  | pV   |
|----------------------------------|-------------|-------|------|
| pH (continue)                    | 0.067       | 0.020 | 0.001|
| Age (years, continue)            | -0.003      | 0.002 | 0.187|
| Sex (0 = male; 1 = female)       | -0.154      | 0.085 | 0.071|
| BMI (kg/m²)                      | 0.030       | 0.005 | 0.001|
| Gastritis pattern                |             |       |      |
| No alteration Ref.               |             |       |      |
| Antral superficial gastritis     | -0.108      | 0.141 | 0.433|
| Superficial chronic pangastritis | 0.007       | 0.109 | 0.950|
| Active superficial pangastritis  | -0.091      | 0.169 | 0.592|
| Atrophic chronic pangastritis    | -0.195      | 0.166 | 0.240|
| TSH (mU/L, continue)             | 0.016       | 0.035 | 0.651|
| Helicobacter pylori (0 = absent; 1 = present) | 0.045 | 0.094 | 0.632|

a Standard error
[59], was associated to superior pharmacokinetic parameters in patients with impaired gastric acidity [50] and to a lower therapeutic dose in patients bearing gastric disorders such as *H. pylori* infection [60]. These novel preparations, as compared to tablet, require no disaggregation process and showed a better dissolution profile (softgel T4) [42] or even no need for dissolution (liquid T4) [57].

The clinical relevance of the present findings stems from the following:

- the high frequency of gastric disorders in general population and in thyroid autoimmune disorders [61–63],
- the prescription rate of proton pump inhibitors [2],
- the impact of food co-ingestion with oral thyroxine in term of absorption efficiency [12, 52].

Last, but not least, the use of healthcare resources is increased in patients experiencing multiple levothyroxine dose adjustments and an increased costs for the health systems has been proven [64].

A major limitation of this study comes from the number of patients which however is hardly possible to overcome. In fact, the invasive procedure required to obtain gastric juice pH and the limited number of patients with stably increased pH prevent from designing studies in large populations.

In conclusion, our data highlight a key role of gastric pH in vivo in shaping the minimal effective dose of thyroxine in hypothyroid patients. Therefore, when prescribing oral thyroxine, physicians should keep in mind all the physiologic, nutritional, pharmacologic, and pathologic conditions that may impair the gastric acid secretion.

**Author contributions** C.V., C.S., and M.C. conceived of, designed the study, and wrote the final version of the manuscript. G.B. and B.P. performed endoscopies, collected gastroenterological data, and revised the manuscript. C.D.V. performed the statistical analysis. M.G.S., L.G., and I.S. organized the data and wrote the firstdraft of the article. A.C. and G.S. performed the literature search and contribute to the analysis of gastroenterological data. All authors contributed significantly to the article and approved the submitted version.

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**Compliance with ethical standards**

**Conflict of interest** M.C. was invited lecturer in international symposia and received honorarium from Institut Biochimique SA (IBSA), Pambio Noranco, CH. The other authors have no conflicts to declare.

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