Liothyronine and levothyroxine prescribing in England: A comprehensive survey and evaluation

Mike Stedman¹ | Peter Taylor² | Lakdasa Premawardhana² | Onyebuchi Okosieame² | Colin Dayan² | Adrian H. Heald³,⁴

¹Res Consortium, Andover, UK
²Thyroid Research Group, Systems Immunity Research Institute, Cardiff University School of Medicine, Cardiff, UK
³The School of Medicine, Manchester Academic Health Sciences Centre, University of Manchester, Manchester, UK
⁴Department of Endocrinology and Diabetes, Salford Royal Hospital, Salford, UK

Abstract

Introduction: The approach to thyroid hormone replacement varies across centres, but the extent and determinants of variation is unclear. We evaluated geographical variation in levothyroxine (LT4) and liothyronine (LT3) prescribing across General Practices in England and analysed the relationship of prescribing patterns to clinical and socioeconomic factors.

Methods: Data was downloaded from the NHS monthly General Practice Prescribing Data in England for the period 2011-2020.

Results: The study covered a population of 19.4 million women over 30 years of age, attending 6,660 GP practices and being provided with 33.7 million prescriptions of LT4 and LT3 at a total cost of £90 million/year. Overall, 0.5% of levothyroxine treated patients continue to receive liothyronine. All Clinical Commission Groups (CCGs) in England continue to have at least one liothyronine prescribing practice and 48.5% of English general practices prescribed liothyronine in 2019-2020. Factors strongly influencing more levothyroxine prescribing (model accounted for 62% of variance) were the CCG to which the practice belonged and the proportion of people with diabetes registered on the practice list plus antidepressant prescribing, with socioeconomic disadvantage associated with less levothyroxine prescribing. Whereas factors that were associated with increased levels of liothyronine prescribing (model accounted for 17% of variance), were antidepressant prescribing and % of type 2 diabetes mellitus individuals achieving HbA1c control of 58 mmol/mol or less. Factors that were associated with reduced levels of liothyronine prescribing included smoking and higher obesity rates.

Conclusion: In spite of strenuous attempts to limit prescribing of liothyronine in general practice a significant number of patients continue to receive this therapy, although there is significant geographical variation in the prescribing of this as for levothyroxine, with specific general practice and CCG-related factors influencing prescribing of both levothyroxine and liothyronine across England.
Primary hypothyroidism affects 2%-5% of the general population and is predominantly managed in the community. The majority of individuals with hypothyroidism are effectively treated with levothyroxine (LT4), but a proportion of patients suffer persistent symptoms, despite achieving biochemical control with levels of free thyroxine (FT4) and thyroid-stimulating hormone (TSH) within the laboratory reference ranges. Some patients who remain dissatisfied with LT4 therapy report improved well-being when prescribed combination therapy with liothyronine (LT3) and LT4. LT3 represents the synthetic form of the metabolically active thyroid hormone and was originally widely prescribed when synthetic thyroid hormones first replaced animal thyroid extracts in the 1950s. However, LT4 monotherapy has since prevailed as the treatment of choice for hypothyroidism because of its more favourable pharmacokinetic profile and once daily dosing schedule, coupled with the failure of randomised controlled trials to show superiority of combination therapy over LT4 alone. Furthermore, early clinical studies used unduly high doses of LT3 that sometimes resulted in unpleasant adverse effects from iatrogenic thyrotoxicosis.

Accordingly, existing clinical guidelines do not support the routine use of LT3 in the management of hypothyroidism. The 2016 British Thyroid Association (BTA) position statement recommends that a trial of combination therapy can be considered in patients who unambiguously do not derive symptomatic benefit from LT4 alone, a position that is broadly consistent with the guidelines of the European and American Thyroid Associations. In the UK, LT3 prescribing has additionally been restricted by serial price increases following transition from the proprietary to the generic product in 2007. Cost considerations have in recent years prompted a series of local prescribing policies aimed at curtailing LT3 prescribing. In the wake of these policies, an analysis by Taylor et al noted a substantial fall in LT3 use in England, a trend that varied geographically and was more pronounced in economically deprived areas. However, the drivers of thyroid hormone prescribing at practice level and the extent of prescribing variation for both LT4 and LT3 are unclear. In the present study, we evaluated geographical variation in LT4 and LT3 prescribing at general practice level in England and examined the factors associated with prescribing variation.

2 | METHODS

2.1 | Data sources

We used a series of NHS general practice datasets to analyse thyroid hormone prescriptions at general practice level in England for the period 2019-2020. The study was conducted on data from the National Health Service (NHS) operational year April 2019–Mar 2020, around which data is normally collated.

What’s known
- Recent prescribing policies across the United Kingdom have imposed significant restrictions on liothyronine prescribing in general practice driven by the prohibitive costs and uncertain benefits of liothyronine in the management of hypothyroidism.
- However, the impact of these policies on liothyronine usage and costs are still unclear.

What’s new
- In spite of strenuous attempts to limit prescribing of liothyronine in general practice, a significant number of patients continue to receive this therapy, although there is significant geographical variation in the prescribing of this, as for levothyroxine with general practice and clinical commissioning group (CCG) level factors a significant determinant.

Information on practice populations and patient distribution by age and sex was obtained from the General Practice registration dataset. This dataset contains a record of all registered patients in GP practices in England broken down by age-bands and sex within Clinical Commissioning Group (CCG) areas. In addition, we collated data from the NHS general practice workforce statistics dataset comprising information on staff numbers including GPs, nurses and other clinical and non-clinical staff working in general practices in England. The Quality Outcome Frameworks dataset was used to document chronic disease prevalence, care quality indicators, as well as social deprivation measures and urban/rural location of the general practice. We also extracted practice level data from the NHS General practice patient survey on rates of patient satisfaction and confidence in chronic disease management as well as ethnicity data for each practice.

Medication use (LT4, LT3 and antidepressants) was obtained from published practice-level monthly prescriptions issued by the NHS Business Service Authority. Prescriptions were extracted by dose and British National Formulary (BNF) code and quantified according to the Defined Daily Dose (DDD). DDD is a standard method of comparing doses of a given medication and is taken as the average maintenance dose per day of a drug administered for its main indication in adults. The amount of active agent was converted to defined daily doses by applying the levels given in the World Health Organisation and Anatomical Therapeutic Chemical (WHO/ATC) classification. For LT3 which is most often used in combination with levothyroxine, an adjusted dose of 20 mcg/day was applied and for LT4 100 mcg/day was used. All the data used in the analyses presented are publicly available and can be made available on request from MS, co-author.
2.2 | Data analysis

As the prevalence of hypothyroidism is higher in women, we calibrated prescribing of thyroid hormone replacement against the number of women in each general practice older than 30 years. In other words, the number of prescriptions as DDD was normalised for comparison between general practices by the number of women aged over 30 years old\textsuperscript{12} as this 38% of the population contains 85% of the patients with hypothyroidism.\textsuperscript{13} The gender and age mix for each practice were taken from the population data at general practice level. A multivariate regression model was used to identify the possible drivers of LT4 and LT3 prescribing. The outcome variables were (a) the amount of LT4 prescribed as a percentage of women aged >30 years, and (b) the amount of LT3 prescribed as a percentage of LT4 prescriptions.

The factors included in the model were as follows: (1) the local population characteristics (age, gender, ethnicity, social deprivation, location, urban vs rural, north vs south, east vs west); (2) the chronic disease burden of the population (rates of hypertension, diabetes, chronic obstructive pulmonary disease (COPD), asthma, cancer, depression and antidepressant prescription); (3) the general practitioner characteristics (age, gender, country of qualification); (4) general practice service performance based on levels of glycosylated haemoglobin HbA1c control reported in the national diabetes audit (NDA); and (5) the patient survey feedback regarding service satisfaction and patient confidence in managing their own long-term conditions.

The stepwise regression was first run with the all the local GP practice factors and then to establish the effect of the CCG. The local average CCG value for each prescribing variable was added as an additional factor for each practice into the stepwise regression to determine how much of the variation in local practices could be explained by their local CCG average for LT4 and LT3 prescribing. The difference in $r^2$ and standardised beta value for the CCG average was an indicator of the size of impact of the CCG on the model.

All analyses were conducted on 64 bit excel with power pivot and Analyse-it add in (Microsoft EXCEL). $\chi^2$ test was used to compare proportions. A $P$ value < .05 was considered significant.

3 | RESULTS

The study covered a population of 19.4 million women over 30 years of age, attending 6660 GP practices and being provided with 33.7 million prescriptions of LT4 and LT3 at a total cost of £90 million/year.

3.1 | Variation in LT4 and LT3 prescribing (Figure 1)

In England, there are 135 local clinical commissioning groups of different population sizes. There was some variation in the identification and treatment of hypothyroidism with LT4 across different CCGs: the median was 7.0% of the population of women age >30 years. The lowest decile region was 5.5% while the highest decile was 8.3% of women >30 years old (factorial variation of 1.5). Variation in use of LT3 was higher. The median was 0.4% of those being treated with Levothyroxine. The lowest decile was 0.1% while the highest decile was 1.4% of those treated with LT4 (factorial variation of 14.0) (Figure 1). All CCGs had at least one LT3 prescribing practice, with 51.5% of general practices not prescribing any LT3.

3.2 | Geographical variation by CCG (Figure 2)

There was a significant geographical variation across CCGs in England in rates of both LT4 and LT3 prescribing—adjusted for the proportion of women over the age of 30 in each CCG. In some areas such as the South-West, Herefordshire and Lincolnshire, higher prescribing of LT4 mapped to higher LT4 prescribing. In other areas such
as North Cumbria and County Durham, there was relatively high prescribing of LT4 but not LT3. Conversely in some areas such as Cheshire and Kent and Medway there was relatively high LT3 prescribing and lower prescribing of LT4.

Overall there are more CCGs in the North of England (defined as a latitude more northerly than 52.6 degrees north) in the lowest tertile of LT3 prescribing, $X^2$ 3.4, $P = .008$ (Table 1).

### 3.3 | Regression modelling

#### 3.3.1 | Levothyroxine prescribing (Figure 3)

For local levels of levothyroxine excluding the effects of CCG guidance, the stepwise regression analysis could explain 54% of the variation between practices. When CCG average for ADDD levothyroxine as percentage of population women age >30 years was included as a factor, the regression model could explain 62% of the variation between general practices, the CCG component having a standardised beta of 0.38 and so the strongest impact. Other factors related to more LT4 prescribing were the proportion of older women in the general practice, the proportion of people registered with diabetes mellitus, the proportion of people with significant social disadvantage, higher smoking rates and a higher proportion of people of black and ethnic minority (BME) ethnicity. General practices relatively more northerly in location tended to prescribe less levothyroxine.

#### 3.3.2 | Liothyronine prescribing (Figure 4)

For local levels of LT3 prescribing, we considered all practices including those that prescribed none. The base analysis of local factors could only explain 6% of the variation between practices, however, by including the CCG average value effect, the stepwise regression analysis could explain 17% of the variation between practices, with the CCG component having a standardised beta of 0.34. Other factors that were associated with increased levels of LT3 prescribing were rates of antidepressant prescribing and percentage of type 2 diabetes mellitus individuals achieving HbA1c control of 58 mmol/mol or less. Factors that were associated with reduced levels of LT3 prescribing included smoking and obesity rates and diabetes prevalence on the practice list.

#### 3.3.3 | Comparison of factors related to levothyroxine and liothyronine prescribing

The main discretionary drivers of local thyroid prescribing for both levothyroxine and liothyronine were local CCG guidance, and practice use of antidepressant. However the impact of CCG guidance on liothyronine prescribing was much greater than for levothyroxine with tripling of the $r^2$. Specifically, this was much higher than for levothyroxine where $r^2$ increased by 20% when the CCG factor was included.

#### 3.3.4 | Year on year prescribing

The $R^2$ value for the factors relating to change in year on year prescribing for levothyroxine (6%) and liothyronine (2%) were low, indicating that the factors that we have access to (including which CCG they belong to) do not relate to difference in change in prescribing year on year, between practices (data not shown).

### 4 | DISCUSSION

Undetected hypothyroidism causes significant morbidity and may be complicated by cardiovascular disease, lipid disorders and neurocognitive impairment. In pregnancy, uncorrected maternal hypothyroidism increases risk of fetal loss, while neglected disease in the elderly may culminate in life-threatening metabolic decompensation. Furthermore, there remains a subset of individuals who report reduced quality of life with health needs that are not met with LT4. Despite a spate of recent guidelines and prescribing policies, the real-world approach to thyroid hormone replacement remains inconsistent and driven by factors that are still unclear. Here, we have evaluated variation in LT4 and LT3 prescribing across general practices in England and determined factors that influence prescribing.

We found significant variation in the use of LT3 and LT4 with a higher degree of variation seen with LT3 prescribing. Although all CCGs had at least one LT3 prescribing practice, about half of practices did not prescribe any LT3. The regression analysis indicated that the CCG that a general practice belongs to has the greatest influence on LT4 and LT3 prescribing. Other factors related to increased LT4 prescribing were the proportion of people registered with diabetes and chronic obstructive pulmonary disease at a general practice and rates of antidepressant prescribing. Interestingly, older general practitioners tended to prescribe more LT4. Factors associated with...
less LT4 prescribing included the proportion of people with obesity and of people with significant socioeconomic deprivation. For LT3, factors that were associated with increased prescribing were antidepressant use and the percentage of individuals with type 2 diabetes achieving HbA1c control of 58 mmol/mol or less, whereas obesity, diabetes and smoking were associated with reduced prescribing.

In spite of strenuous attempts to limit LT3 prescribing in general practice, our findings show that a significant number of doctors continue to prescribe LT3. In England priorities for primary care are set by the local CCGs of which there are 135 of different sizes. Notably, all CCGs had at least one LT3 prescribing practice, suggesting a continued need for LT3 whether driven by patients or

FIGURE 2  (A) Map of Variation in levothyroxine and liothyronine prescribing in England by CCG. (B) The key to the England CCG map with each numbered CCG described
their clinicians. Although cost considerations have led to prescribing policies designed to reduce LT3 prescription, the continued use of LT3 may have been encouraged by various sets of guidance published in the last decade that now allow LT3 prescriptions in carefully selected individuals. Rates of LT4 prescribing, on the other hand, are influenced by differing views on the laboratory TSH thresholds for LT4 initiation. Although these thresholds have progressively reduced over the years, our results suggest that there is a significant variation with respect to screening and treatment initiation in patients with hypothyroidism. The influence of

FIGURE 2 Continued
of CCGs through local medicine management committees appear to play an over-riding role in the approach to both LT3 and LT4 prescribing.

The increased LT4 prescribing seen in association with comorbidities like diabetes, chronic obstructive pulmonary disease and depression may reflect frequent testing and diagnosis in these groups of patients due to their more frequent contact with general practice. The association of both LT4 and LT3 prescribing to rates of antidepressant prescribing may as alluded be a measure of how an individual general practice may be attuned to their practice population in relation to the realities of living with long-term conditions and their consequences.15 In addition, patients with depression are more likely to have frequent contacts with their practices and therefore more likely to be tested for unresolved symptoms.15 Reduced LT4 prescription was also seen with social deprivation, practices in more northern regions of England, and black and ethnic minority individuals suggesting inequalities in care access. Similar variations were reported by Taylor et al.1 Thus 50 years after the seminal paper by Taylor et al in 1970,16 the combination of LT4 and LT3 still finds favour with a good number of practitioners and many patients.

4.1 | Strengths/Limitations

A strength of this study is in the use of real-world general practice data collated at a national level, better to understand the factors influencing thyroid hormone prescribing across all CCGs in England. Although we have not looked at data from the other nations that make up the UK, our findings are likely to be applicable to the other parts of the UK and other parts of the world where there is a large differential between the costs of LT4 and LT3. Also we have not specifically evaluated the prescribing of NDT which is used by a small proportion of patients with hypothyroidism.
The matter of longitudinal trends in prescribing LT4 and LT3 in England is the subject of a separate paper.17

4.2 Conclusion

There is significant geographical variation in the prescribing of LT4 and LT3 in general practice. The CCG where any general practice is located appears to be the over-riding influence on thyroid hormone prescribing with the influence much greater for LT3 than for LT4 prescribing.

DISCLOSURES

None of the co-authors has any conflict of interest in relation to the work presented here.

DATA AVAILABILITY STATEMENT

All the data used in the analyses presented is publically available and can be also made available from MS, co-author on application.

ORCID

Mike Stedman https://orcid.org/0000-0002-0491-7823
Adrian H. Heald https://orcid.org/0000-0002-9537-4050

REFERENCES

1. Taylor PN, Albrecht D, Scholz A, et al. Global epidemiology of hyperthyroidism and hypothyroidism. Nat Rev Endocrinol. 2018;14:301-316.
2. Saravanan P, Chau WF, Roberts N, Vedhara K, Greenwood R, Dayan CM. Psychological well-being in patients on “adequate” doses of L-thyroxine: results of a large, controlled community-based questionnaire study. Clin Endocrinol (Oxf). 2002;57:577-578.
3. Smith RN, Taylor SA, Massey JC. Controlled clinical trial of combined triiodothyronine and thyroxine in the treatment of hypothyroidism. Br Med J. 1970;1:145-148.
4. Wiersinga WM, Duntas L, Fadeyev V, Nygaard B, Vanderpump MP. 2012 ETA guidelines: the use of L-T4 + L-T3 in the treatment of hypothyroidism. Eur Thyroid J. 2012;1:55-71.
5. Jonklaas J, Bianco AC, Bauer AJ, et al. American thyroid association task force on thyroid hormone replacement. Guidelines for the treatment of hypothyroidism: prepared by the American thyroid association task force on thyroid hormone replacement. Thyroid. 2014;24:1670-1751.
6. Okosie O, Gilbert J, Abraham P, et al. Management of primary hypothyroidism: statement by the British Thyroid Association Executive Committee. Clin Endocrinol (Oxf). 2016;84:799-808.
7. https://digital.nhs.uk/data-and-information/publications/statistics/lpractice-level-prescribing-data. Accessed October 1, 2020.
8. https://digital.nhs.uk/data-and-information/publications/statistics/lgeneral-and-personal-medical-services. Accessed October 2, 2020.
9. https://qof.digital.nhs.uk/. Accessed October 1, 2020.
10. https://www.gp-patient.co.uk/. Accessed October 1, 2020.
11. WHO Collaborating Centre for Drug Statistics Methodology. Defined Daily Dose (DDD). ATC/DDD index. http://www.whocc.no/atc_ddd_index. Accessed October 3, 2020.
12. https://www.ons.gov.uk/peoplepopulationandcommunity/populationandmigration/demography.
13. Giorda CB, Carnà P, Romeo F, Costa G, Tartaglino B, Gnavi R. Prevalence, incidence and associated comorbidities of treated hypothyroidism: an update from a European population. Eur J Endocrinol. 2017;176:533-542.
14. Taylor PN, Iqbal A, Minassian C, et al. Falling threshold for treatment of borderline elevated thyrotropin levels-balancing benefits and risks: evidence from a large community-based study. JAMA Intern Med. 2014;174:32-39.
15. Heald AH, Stedman M, Davies M, Farman S, Taylor D, Bailey S, Gadsby R. Quantifying the impact of patient-practice relationship quality on the levels of the average annual antidepressant practice prescribing rate in primary care in England. Prim Care Companion CNS Disord. 2020;22.
16. Taylor S, Kapur M, Adie R. Combined thyroxine and triiodothyronine for thyroid replacement therapy. BMJ. 1970;2:270-271.
17. Stedman M, Taylor P, Premawardhana L, Okosie O, Dayan C, Heald AH. Trends in costs and prescribing for liothyronine and levothyroxine in England and Wales 2011–2020. Clinical Endocrinology. 2021;94:980–989. https://doi.org/10.1111/cen.14414.

How to cite this article: Stedman M, Taylor P, Premawardhana L, Okosie O, Dayan C, Heald AH. Liothyronine and levothyroxine prescribing in England: A comprehensive survey and evaluation. Int J Clin Pract. 2021;75:e14228. https://doi.org/10.1111/ijcp.14228