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1 TITLE: POST-RADIOIODINE GRAVES' MANAGEMENT: THE PRAGMA STUDY

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74 Fairfield Hospital, Bury, UK), Thomas Grüning (Department of Nuclear Medicine, 75 University Hospitals Plymouth NHS Trust, Derriford Hospital, Plymouth, UK), Angelos 76 Kyriacou (The Christie NHS Foundation Trust, Manchester, UK) and Professor Peter 77 Trainer (The Christie Foundation Trust, Manchester, UK). The study was funded by the 78 Clinical Endocrinology Trust (UK Registered Charity Number 288679). 79 **SUMMARY** 80 81 **Objective.** Thyroid status in the months following radioiodine treatment for Graves' 82 disease can be unstable. Our objective was to quantify frequency of abnormal thyroid function post-radioiodine and compare effectiveness of common management 83 84 strategies. 85 **Design.** Retrospective, multi-centre, observational study. 86 Patients. Adult patients with Graves' disease treated with radioiodine with 12 months' 87 follow-up. 88 *Measurements.* Euthyroidism was defined as both serum thyrotropin (TSH) and free 89 thyroxine (FT4) within their reference ranges or, when only one was available, it was 90 within its reference range; hypothyroidism as TSH \geq 10 mu/L, or subnormal FT4 91 regardless of TSH; hyperthyroidism as TSH below and FT4 above their reference 92 ranges; dysthyroidism as the sum of hypo- and hyperthyroidism; subclinical 93 hypothyroidism as normal FT4 and TSH between the upper limit of normal and <10 94 mu/L; subclinical hyperthyroidism as low TSH and normal FT4 95 **Results.** Of 812 patients studied post-radioiodine, hypothyroidism occurred in 80.7% 96 and hyperthyroidism in 48.6% of patients. Three principal post-radioiodine management

97 strategies were employed: (a) anti-thyroid drugs alone, (b) levothyroxine alone and (c) 98 combination of the two. Differences among these were small. Adherence to national 99 guidelines regarding monitoring thyroid function in the first 6 months was low (21.4-100 28.7%). No negative outcomes (new-onset/exacerbation of Graves' orbitopathy, weight 101 gain, cardiovascular events), were associated with dysthyroidism. There were 102 significant differences in demographics, clinical practice, and thyroid status post-103 radioiodine between centres. 104 *Conclusions.* Dysthyroidism in the 12 months post-radioiodine was common. 105 Differences between post-radioiodine strategies were small, suggesting these 106 interventions alone are unlikely to address the high frequency of dysthyroidism. 107 CLINICAL TRIAL REGISTRATION: Clinical.trials.gov (identifier No. NCT01885533). 108 **KEY WORDS:** Graves' disease, thyroid, radioiodine, hypothyroidism, hyperthyroidism 109 **CONFLICT OF INTEREST STATEMENT:** The authors have no conflicts of interest to 110 declare. 111 **DATA AVAILABILITY STATEMENT:** The data that support the findings of this study 112 are available on request from the corresponding author.

113 **ORCiD number:** 0000-0001-7320-5574 (P. Perros)

114 **ABBREVIATIONS:** ATDs, anti-thyroid drugs; B&R, block and replace; FT3, free tri-

- iodothyronine; FT4, free thyroxine; L-T4, levothyroxine; GD, Graves' disease; GO,
- 116 Graves' orbitopathy; Q, quarter; RI, radioiodine; TFTs, thyroid function tests; TSH,
- 117 thyroid stimulating hormone.
- 118
- 119 INTRODUCTION

Radioiodine (RI) is a safe and effective treatment for Graves' disease (GD) ¹. The aim of
RI therapy is to cure the hyperthyroidism ^{2,3}. Attempts to calculate a dose of RI that
eliminate hyperthyroidism yet prevent hypothyroidism have not produced reliable
results, and have been abandoned in the UK and other countries in favour of larger,
fixed doses ^{2,4-6}. As a consequence, the majority of patients develop thyroid hormone
dependence within the first year after RI⁷.

126

127 Whether hypothyroidism should be allowed to occur post-RI before thyroid hormone 128 replacement is introduced, or be prevented, is an important question, which has 129 received little attention in recent years. The argument in favour of allowing 130 hypothyroidism to develop is to ensure that life-long treatment with levothyroxine (L-T4) 131 is necessary. The case against is based on associations between hypothyroidism with 132 impaired quality of life⁸, weight gain⁹ and Graves' orbitopathy (GO) ¹⁰⁻¹². Surveys 133 performed more than 20 years ago revealed a wide variation among clinicians agreeing 134 with the proposition that transient hypothyroidism with subsequent introduction of replacement therapy is an acceptable practice ¹³⁻¹⁵. A more recent UK-based survey ⁶ 135 and large published series ^{11,16} indicate that such variations in practice persist. 136 137 Strategies used by clinicians to bridge the transition from hyperthyroidism to euthyroidism on stable L-T4 therapy following RI include a short course of anti-thyroid 138 139 drugs (ATDs) alone, the combination of ATDs with L-T4 known as "block and replace" (B&R), or watchful monitoring with the introduction of L-T4 when needed ^{2,3,6,10}. 140 141

142 MATERIALS AND METHODS

143 **Objectives**

- 144 The primary objectives were to document the frequency of dysthyroidism in the first 12
- 145 months following RI and compare the impact of different post-RI management strategies
- 146 on thyroid status. Secondary objectives were to identify potential drivers for post-RI
- 147 dysthyroidism, relationships between post-RI dysthyroidism, and clinical outcomes and
- 148 differences between participating centres.

149 Study design

150 Retrospective, observational, multi-centre, secondary care study.

151 Inclusion Criteria

- Age \geq 18 years at the time of RI; diagnosis of GD; treatment with RI; 12 months follow-
- up after RI; most recent RI dose administered 1 or more years before enrolment.

154 **Participating centres**

- 155 Investigators were invited to participate through the *Society for Endocrinology* website
- and its newsletters (https://www.endocrinology.org/). Thirty-one NHS hospitals / centres
- 157 participated in the study
- 158 (https://web.archive.org/web/20210329111703/https://www.mapcustomizer.com/map/P
- 159 RAGMA%20centres).

160 Enrolment and data collection

- 161 Patients were identified from registries of RI administration and endocrine departmental
- 162 databases at each institution. Following enrolment, the medical records were used to
- 163 extract relevant information. All paper records were pseudo-anonymized and entered in
- a central electronic database. Recruitment commenced in March 2013 and ended in
- 165 February 2015.

166	Definitions
167	Patients were considered to have GD when there was biochemical evidence of
168	thyrotoxicosis (low serum TSH with elevated serum FT3 and / or FT4 levels) and one or
169	more of the following: (a) Diffuse uptake on thyroid isotope scan, (b) elevated serum
170	TSH receptor antibody levels, (c) clinical evidence of GO, (d) diffuse goitre by palpation
171	and positive thyroid peroxidase antibodies. Patients were considered to have GO if they
172	had eye features class II or greater according to the NOSPECS classification ¹⁷ .
173	Exacerbation of GO was defined as recorded evidence of worsening symptoms and / or
174	eye signs.
175	Based on the results of TFTs performed in local laboratories, patients were classified
176	as:
177	• Hypothyroid: serum TSH above and FT4 below the reference range, or serum
178	TSH \geq 10 mU/l associated with a normal serum FT4, or TSH \geq 10 mU/l without an
179	available FT4 level, or serum FT4 less than the reference range regardless of the
180	serum TSH concentration
181	Hyperthyroid: serum TSH below and serum FT4 above the reference ranges
182	• Subclinical hypothyroid: serum TSH above the reference range but <10 mU/L
183	and serum FT4 within the reference range
184	Subclinical hyperthyroidism: serum FT4 within and serum TSH below the
185	reference range
186	• Euthyroid: both serum TSH and FT4 within the reference range or, when only
187	one was available, within the reference range.

188 Data on weight gain were extracted from medical records.

210	Missing data
209	(dysthyroid was the sum of hypo- and hyperthyroid episodes).
208	number of hypo-, hyper- and dysthyroid episodes for each patient were calculated
207	and hyperthyroidism in the same Q, they were both counted. From the above, the total
206	only the most abnormal result was included. When there were episodes of both hypo-
205	sets of TFTs were available in the same Q showing either hypo- or hyperthyroidism,
204	classified thyroid profile within 3 months of the previous was not counted. When multiple
203	abnormal biochemistry would be correctable within 3 months, therefore a similarly
202	An assumption was made that, following detection of hypo- or hyperthyroid TFTs, the
201	hyperthyroid biochemistry
200	"Subclinical hyperthyroid" when one or more TFTs showed subclinical
199	hypothyroid biochemistry
198	"Subclinical hypothyroid" when one or more TFTs showed subclinical
197	"Euthyroid" when TFTs showed euthyroid biochemistry
196	"Hypothyroid" when one or more TFTs showed hypothyroid biochemistry
195	"Hyperthyroid" when one or more TFTs showed hyperthyroid biochemistry
194	For further analyses each Q for each patient was coded as:
193	Q1=0-91 days, Q2=92-182, Q3=183-274 days, Q4=275-365 after RI treatment.
192	The 12-month follow-up period after RI was split into 3-month blocks or quarters (Qs):
191	Data handling
190	classifications were based.
189	Each centre used local laboratories and reference ranges upon which the above

- For the analyses pertaining to thyroid status during the 12 months following RI, 27
- 212 patients were excluded because they had no TFTs available. For other analyses (not
- 213 involving TFTs) data from all 812 patients were included.
- 214 Biochemical assays

215 Biochemical data on thyroid function were derived from the local laboratories. The 216 assay platforms used were: Siemens Advia Centaur XP, Siemens Vista, Roche Cobas 217 800, Roche Modular, Centaur, Abbott Architect, Beckman Coulter DxI. The most 218 commonly used reference ranges for TSH (0.35-4.5 mU/L) and FT4 (10-22 pmol/L) 219 were used for normalisation of all TSH and FT4 data, so as to make them comparable for statistical analyses. Validated formulas for normalisation were used¹⁸. To calculate 220 221 values from laboratory x to laboratory y according to the following formula (y represents 222 a normalized value from laboratory x to laboratory y, x is a measured concentration at 223 laboratory x, Uy is the upper reference level for laboratory y and Ux is the upper 224 reference level for laboratory x):

 $y = x \frac{U_y}{U_x}$

It has been reported that patients on L-T4 for primary hypothyroidism have a higher mean FT4 serum concentration than healthy euthyroid people¹⁹ and this fact may need to be taken into account when interpreting TFTs of such patients. However, for the purposes of this study the same normalised reference range for FT4 was used for all patients.

231 Statistical analyses

232 Statistical analyses were undertaken using STATA (STATA Corp LLC, College Station,

233 TX, USA. Version 16) for primary and secondary outcomes. Parametric and non-

234 parametric tests, linear and logistic regression analyses were used. All p values are 235 two-sided and a value of 0.05 considered to indicate statistical significance. The effects 236 of post-RI treatment strategies were examined using a linear mixed model. The model 237 included age, gender, smoking habit, dose of RI, and centre ID. Logistic regression was 238 used to examine: (a) associations between new-onset and exacerbation of GO and age. 239 gender, smoking, time from diagnosis of GD, dose of RI, prophylactic steroid use and 240 thyroid status post-RI; (b) associations between changes in body weight and age, 241 gender, smoking, thyroid status post-RI, use of prophylactic steroids, centre ID; (c) 242 associations between cardiovascular events and age, gender, smoking, dose of RI, thyroid status post-RI. A linear mixed model was used to explore differences between 243 244 centres using age, gender, smoking, proportion of patients who had previously been 245 treated with ATDs with curative intent, GO prior to RI, use of prophylactic steroids, dose 246 of RI, weight change and thyroid status in the model.

247 **Regulatory approvals**

Ethical approval was granted from the National Research Ethics Service (IRAS
reference 110269). The study was adopted by the National Institute of Health Research
Clinical Research Network, and received Research and Development and Caldicott
Guardian approval from each of the sites.

252

253 **RESULTS**

A total of 812 patients were included from 31 UK centres. The baseline characteristicsare shown in Table 1.

256 Thyroid function tests

257 A total of 3,951 sets of TFTs (3,616 paired TSH and FT4, 328 TSH only, 7 FT4 only) 258 were recorded over 12 months following RI treatment in 785 patients. The TSH and FT4 259 values across time are shown in Figure 1. Categorisation of the data shown in Figure 1 260 into thyroid status were, euthyroid 23.9% (945/3951), hypothyroid 31.9% (1262/3951), 261 hyperthyroid 12.0% (475/3951), subclinical hypothyroid 8.3% (328/3951), subclinical 262 hyperthyroid 20.7% (816/3951), The median number of tests per patient per Q was 1 263 (range 0-7) for Q1, 2 (range 0-5) for Q2, 1 (range 0-6) for Q3, and 1 (range 0-7) for Q4. 264 Hypothyroidism peaked in Q2 (61.8%) and declined to 17.8% in Q4. Hyperthyroidism 265 was highest in Q1 (26.3%) and reached a trough in Q4 (13.4%). Hypothyroidism was 266 most prevalent in Q2 (60.2%) and lowest in Q4 (18.6%), while euthyroidism was lowest 267 in Q2 (11.3%) and highest in Q4 (33.0%). Subclinical hyper- and hypothyroidism varied 268 between 9.1-23% and 3.9-11.4% respectively (Figure 2). The overall risk of patients 269 experiencing at least one episode of hypo- or hyperthyroidism in the 12 months 270 following RI (calculated from a subgroup of 358 patients who had at least one set of 271 TFTs for every Q), was 80.7% and 48.6% respectively (Table 2). Conversely, only 9.2% 272 of patients avoided dysthyroidism during the 12 months post-RI. TSH values peaked in 273 Q2 and were lowest in Q4. There were no differences in serum FT4 levels across Qs 274 (Table 3). It may be argued that hyperthyroidism in Q1 is to be expected and that a 275 single episode of hypothyroidism is acceptable in order to confirm the need for life-long 276 thyroid hormone replacement, however, 26.8% of patients experienced more than one 277 episode of hypothyroidism and 54.8% of the hyperthyroid episodes occurred after Q1 278 (Figures 1 and 2).

- 279
- Ultimate and penultimate TFTs before commencement of L-T4 treatment

280 In a subset of patients (61.7%, 484/785), dates were available for starting L-T4 281 treatment. For this group of patients, it was possible to explore: (a) thyroid status before 282 starting L-T4, (b) how promptly L-T4 was started after the blood test, (c) whether 283 dysthyroidism in the last (ultimate) set of TFTs before commencing L-T4 could have 284 been predicted by the previous (penultimate) set of TFTs. At the time of the ultimate 285 TFTs before starting L-T4, 77% (373/484) of patients were hypothyroid. Hypothyroid 286 patients were commenced on L-T4 treatment within a median of 7.8 days (range 0-161) 287 from the date of the ultimate hypothyroid TFTs. In 67.8% (328/484) of patients 288 penultimate TFTs were available. Penultimate TFTs were taken a median of 48 days 289 (range 2-203) before the ultimate TFTs, and a median of 60 days (range 1-342) since 290 RI. Of these 328 patients the penultimate TFTs showed: subclinical hyperthyroidism in 291 37.5% (123/328), euthyroidism in 23.8% (78/328), hypothyroidism in 18.3% (60/328), 292 hyperthyroidism in 18.0% (59/328) and subclinical hypothyroidism in 2.4% (8/328). The 293 probability of hypothyroidism in the ultimate TFT was highest (90%) if the penultimate 294 TFT was also hypothyroid, and lowest (75.6%) if the penultimate TFT was euthyroid. 295 Post-RI management strategies and thyroid status outcomes 296 Of the 785 patients who had follow-up TFTs after RI, the post-RI treatment strategy was 297 recorded in 91.6% of cases (719/785): 35.5% (255/719) received ATDs alone, 15.2% 298 (109/719) B&R, and 49.4% (355/719) L-T4 alone. There were some differences in 299 baseline characteristics between the three management strategy categories 300 (Supplementary Table 1). Table 3B shows the frequencies in thyroid status for the entire 301 cohort and by treatment strategy for each Q. Using a liner mixed model that included 302 age, gender, smoking habit, dose of RI, and centre ID, and considering thyroid status as

- 303 a categorical variable (hypothyroid, hyperthyroid or dysthyroid), the only difference
- 304 between the management strategy groups was a lower risk of hyperthyroidism
- 305 associated with the use of L-T4 alone compared to other treatment strategies (p<0.02,
- 306 Figure 3).

307 Efficacy of RI treatment

- 308 Hyperthyroid thyroid function tests in Q4 were used as a surrogate measure of failure of
- 309 RI treatment. Using this criterion (and data from 516/785 patients with available TFTs in
- 310 Q4), RI failed in 13.4% (69/516) of patients.

311 Changes in body weight

- 312 Data on body weight were available in 74.0% (601/812) of patients. The majority
- 313 (73.9%) gained weight within a year of being treated with RI by a mean of 3.0 kg (SD
- 4.3). This amount of weight seems modest compared to that reported by other studies ⁹,
- 315 however most patients in PRAGMA had relapsed thyrotoxicosis and were probably not
- 316 as thyrotoxic. Multiple linear regression showed no association with demographic
- 317 variables, smoking status, post-RI thyroid status, use of prophylactic steroids for GO, or
- 318 post-RI treatment strategy, after adjusting for centre ID.

319 Graves' orbitopathy

- A minority of patients (18.2%, 148/812) had GO prior to treatment with RI. The median
- time from diagnosis of GD to RI for patients with GO was 31.9 months (range 0.9-226.5)
- 322 and not statistically different to patients without GO. Current smoking was associated
- with a greater risk of GO prior to RI (28.3%) compared to non-smokers (14.5%)
- 324 (p<0.001). New-onset GO after RI developed in 3.5% (23/664), while exacerbation of
- 325 pre-existing GO in 41.9% (62/148) of patients. Logistic regression showed that current

326 smoking status and a lower dose of most recent RI were the only two factors that were 327 predictive of new-onset of GO (p=0.029 and p=0.027 respectively). Prophylactic 328 steroids were administered in 47.3% (70/148) of patients with pre-existing GO, and in 329 0.3% (2/664) patients without GO. The rate of exacerbation of GO after RI in patients 330 with pre-existing GO who received prophylactic steroids (24.3%, 17/70) was no different 331 to those who did not receive steroids (17.9%, 14/78, p=NS). The rates of referral to 332 Ophthalmology were 82.6% (19/23) for new-onset and 41.9% (26/62) for exacerbation 333 of pre-existing GO. Specific treatments for GO were administered in 13.4% (23/172) of 334 patients after RI and all took place after referral to Ophthalmology. The commonest 335 treatment was steroids (47.8%, 11/23) followed by surgical orbital decompression (26.1%, 6/23), lid surgery (17.4%, 4/23), radiotherapy (4.3%, 1/23) and squint surgery 336 337 (4.3%, 1/23).

338 Cardiovascular events post-RI

Data on cardiovascular events were available in 97% (788/812) of patients and
occurred in 1.2% (10/788) after RI (atrial fibrillation 1.0%, atrial fibrillation associated
with acute coronary syndrome 0.1%, stroke 0.1%). Logistic regression showed no
associations between age, gender, smoking, dose of RI, thyroid status post-RI or
treatment strategy post-RI.

344 Adherence to guidelines

Adherence to the 2007 national guidelines²⁰ was high in relation to dose of RI (93.1%), timing of initiation of ATDs after RI when indicated (93.8%), measurement of both FT4 and TSH (91.7%), and measurement of TFTs at 7-9 months (75.0%) and 9-12 months

- 348 (84%). Adherence was low to the recommendations that TFTs should be measured at
- about 6 weeks post-RI (21.4%), 12-14 weeks (28.7%) and 24-26 weeks (21.4%).

350 Differences between centres

- Differences between centres were noted in patient age (p<0.001), gender (p<0.05),
- 352 current smoking status (p<0.05), previous treatment with ATDs with curative intent
- 353 (p<0.001), prevalence of GO prior to RI (p<0.001), use of prophylactic steroids for
- 354 prevention of exacerbation or new-onset of GO (p<0.001), dose of RI administered
- 355 (p<0.001), weight change (p<0.001), and number of hypothyroid (p<0.05), hyperthyroid
- 356 (p<0.05) and dysthyroid episodes (p<0.05) (Supplementary Figure 1).
- 357

358 **DISCUSSION**

359 One of the main findings of PRAGMA was the high frequency of dysthyroidism in the 360 first 12 months post-RI. Only 9.2% of patients avoided dysthyroidism, while 80.7% and 361 48.6% experienced at least one episode of biochemical hypo- or hyperthyroidism 362 respectively. Hypothyroidism was most likely to occur in Q2, while hyperthyroidism was 363 commonest in Q1; thus, the first 6 months after RI define the time window of the highest 364 risk of dysthyroidism. More than a guarter (26.8%) of patients suffered two or more 365 hypothyroid episodes. These findings suggest that management of many patients may 366 be suboptimal. Paradoxically, one of the contributors to the high frequency of 367 hypothyroidism may be misinterpretation of professional guidelines. The American Thyroid Association guidelines ² state "The goal of radioiodine therapy in Graves" 368 disease is to control hyperthyroidism by rendering the patient hypothyroid". This 369 370 statement was probably intended to emphasise the futility of striving to achieve

371 euthyroidism without thyroid hormone substitution by using small doses of RI, and the 372 inevitability of thyroid failure, rather than encourage clinicians to allow patients to 373 become hypothyroid. The UK national guidelines available at the time of the study state 374 "hypothyroidism in the first six months after treatment may be transient in over half of 375 the patients, and long-term thyroxine replacement should not be given unless it is clear *that hypothyroidism is permanent*"²⁰. This recommendation is based on a cited study by 376 377 Aizawa et al (1997)²¹, whereby relatively small calculated doses of RI were used 378 (ranging from 171-219 MBg), in contrast to current practices in the UK, the rest of Europe and North America, which range between 400 and 800 MBg^{2,4,5}. PRAGMA 379 380 shows that when 400-800 MBg of RI is used, the probability of a hypothyroid episode in 381 the first 6 months being persistent, if not treated, is 90%. An important question is 382 whether dysthyroidism can be prevented in the year following RI. Some studies have 383 shown that it is possible to achieve lower rates of dysthyroidism than PRAGMA in the 384 first year after RI ^{10,12,22} (incidence of hypothyroidism and subclinical hypothyroidism 385 less than 5.5% and 14% respectively), though it is unclear which are the important 386 components that determine success and how much different strategies (use of ATDs 387 alone, B&R or L-T4 alone) contribute. There were no major differences between the 388 three main post-RI strategies (although a non-significant trend of an association 389 between the use of B&R and greater rates of euthyroidism achieved was noted (Table 390 3B)), suggesting that these interventions alone are unlikely to address the high 391 frequency of dysthyroidism. Probable contributors to dysthyroidism post-RI include: (a) 392 suboptimal level of biochemical monitoring, especially in the first 6 months; (b) non-

adherence by patients with treatment; (c) and reluctance by physicians to introduce full
 replacement doses of L-T4; (d) rapid change in biochemical status post-RI.

395

396 New-onset of GO after RI was uncommon in the PRAGMA cohort and similar to one of 397 the largest published series ²³. The negative association between the dose of the most 398 recent RI and new-onset of GO is an interesting observation and may relate to the 399 hypothesis that "total thyroid ablation" is beneficial in GO²⁴. Prophylactic steroids did 400 not seem to prevent exacerbations of pre-existing GO, which has been noted in other studies ^{25,26}, and may be related to the dose and route of administration ²⁷. In the 401 402 majority of patients, new-onset and exacerbation of pre-existing GO triggered referrals 403 to Ophthalmology (79.2% and 86.7% respectively), and subsequently most referred 404 patients received treatment. This contrasts to a European survey conducted in 2006 405 which showed a reluctance among endocrinologists to refer patients to Ophthalmology 406 ²⁸ and suggests that the management of GO in the UK may be improving, possibly in 407 response to the efforts of TEAMeD (Thyroid Eye Disease Amsterdam Declaration 408 Implementation Group, UK) ^{29,30}. Cardiovascular events after RI were reassuringly 409 uncommon after RI and similar to that reported for the background population in 410 England ³¹. Significant differences in patient outcomes were noted between centres, 411 which may be explained partly by differences in patient demographics and therefore 412 case-mix, and requires further attention. Despite the high frequency of dysthyroidism in 413 the first 12 months post-RI, there were no discernible negative effects on patient 414 outcomes, such as increased risk of GO, or cardiovascular events.

415

416 The principal strength of PRAGMA is the large number of patients and multi-centre participation. Based on available UK data ^{32, 33, 34}, it is estimated that the PRAGMA 417 418 cohort represented about 10% of the UK population of patients with GD treated with RI 419 per year. Given the participation of 31 centres and their wide geographical distribution 420 across the UK, it can be inferred that the findings of PRAGMA are likely to be 421 representative of UK patients and practises. The fact that most of the PRAGMA cohort 422 had previously been treated with ATDs with curative intent concurs with current 423 practices in Europe ^{6,35}, and the USA ³⁶. In view of the above, and the fact that the 424 number of patients included in PRAGMA is one of the largest in the literature, this 425 suggests that the findings generated by PRAGMA are also likely to be of relevance to 426 other European and North American populations of adult patients with GD treated with 427 RI. The study is also subject to weaknesses. The data are retrospective, there are likely 428 to be selection biases, there were missing data, and it was not possible to validate the 429 data independently due to limited resources.

430

431 Based on the findings of PRAGMA, some simple measures may reduce the frequency 432 of post-RI dysthyroidism: (a) adherence to the recent NICE guidelines which 433 recommend monitoring of TSH, FT4 and FT3 6 weekly during the first 6 months following RI until TSH is in the reference range³⁷; (b) patient engagement (patients 434 435 being informed of the high risk of dysthyroidism, the importance of adherence to 436 medication, the importance of frequent monitoring and need to modify their medication 437 following results of blood tests); (c) initiation of L-T4 treatment when thyroid 438 biochemistry shows subclinical hypothyroidism or hypothyroidism; (d) use of full

replacement doses of L-T4 from the outset as recommended by NICE ³⁷. Additional
prospective studies are required to define the efficacy and cost effectiveness of other
strategies for the post-RI management of patients with GD.

442

In conclusion, dysthyroidism in the first 12 months after RI, especially hypothyroidism, is
very common and often recurrent, suggesting suboptimal management. The findings of
PRAGMA indicate that guidance from professional organizations on whether avoidance
of dysthyroidism after RI should be pursued rigorously by clinicians for all patients with
GD post-RI, would be valuable.

448

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453 contributed to the acquisition, analysis, and interpretation of the data. A.B. and M.P.Z.

454 performed the statistical analyses. All authors contributed to the discussion, edited and

455 critically reviewed the manuscript. P.P. is the guarantor of this work and, as such, had

456 full access to all the data in the study and takes responsibility for the integrity of the data

457 and the accuracy of the data analysis. All authors read and approved the final

458 manuscript.

459

460 **ADDITIONAL INFORMATION**

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- 464 *Data availability:* The datasets generated during and/or analysed during the current
- study are not publicly available but are available from the corresponding author on
- 466 reasonable request.

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- 565

566 LEGENDS FOR FIGURES AND TABLES

567

568 **FIGURE 1**

- 569 The distribution of serum TSH (left panel) and FT4 (right panel) across time is shown for
- all patients with recorded thyroid function tests (n=785). The y axes show the serum
- 571 TSH and FT4 concentrations. The x axis shows time. The reference ranges for
- 572 normalized TSH and FT4 were 0.30-45 mU/L and 9-22 pmol/L respectively.
- 573

574 **FIGURE 2**

- 575 Thyroid status across time for all 785 patients with available thyroid function tests after
- 576 radioiodine. The y axis shows frequency of euthyroidism, hypothyroidism
- 577 hyperthyroidism , subclinical hypothyroidism and subclinical hyperthyroidism. The x
- 578 axes show time across 3-month blocks (quarters Q1-Q4). The horizontal brackets
- 579 denote statistically significant pairs (p<0.05, chi-squared tests).
- 580

581 **FIGURE 3**

- 582 Thyroid status for patients treated with anti-thyroid drugs alone post-radioiodine (dark
- 583 grey columns), block and replace (white columns) and levothyroxine alone (light grey
- columns). The x axes show time across 3-month blocks (quarters Q1-Q4). Use of L-T4

- alone was associated with a lower risk of hyperthyroidism compared to other treatment
- 586 strategies (p<0.02, linear mixed model). ATDs: anti-thyroid drugs; B&R: block and
- 587 replace. L-T4: levothyroxine
- 588
- 589

590 SUPPLEMENTARY FIGURE 1

591 Differences between centres. The vertical axes indicate the parameters of interest

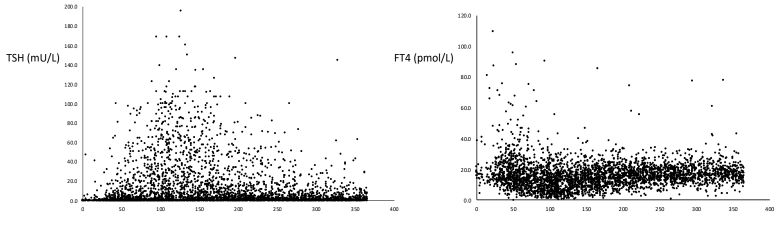
592 (mean and 95% CI). The horizontal axes denote the centre identification numbers. All

- 593 parameters shown in the figure were statistically different when tested by centre to the
- level of p<0.05. The upper panel shows differences in the primary outcomes. A: number
- of hypothyroid episodes RI; B: number of hyperthyroid episodes post-RI; C: number of
- 596 episodes post-RI. The lower panel shows differences between centres in baseline
- 597 characteristics. RI: radioiodine; GO: Graves' orbitopathy; centre ID: centre identification
- 598 number.
- 599
- 600 **TABLE 1**
- 601 Baseline characteristics of patients.
- 602
- 603 **TABLE 2**
- 604 Cumulative rates of euthyroidism and dysthyroidism progressing through quarters.
- 605
- 606 **TABLE 3**

- 607 Thyroid function tests by quarter (3-month blocks, or Qs) (A) and thyroid status by post-
- 608 radioiodine treatment strategy (B).
- 609

610 SUPPLEMENTARY TABLE 1

- 611
- 612 Baseline characteristics of all patients and in patients and in patients treated with anti-
- 613 thyroid drugs, block and replace and L-T4 alone post-radioiodine (RI).
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Time since radioiodine (days)

Figure 1

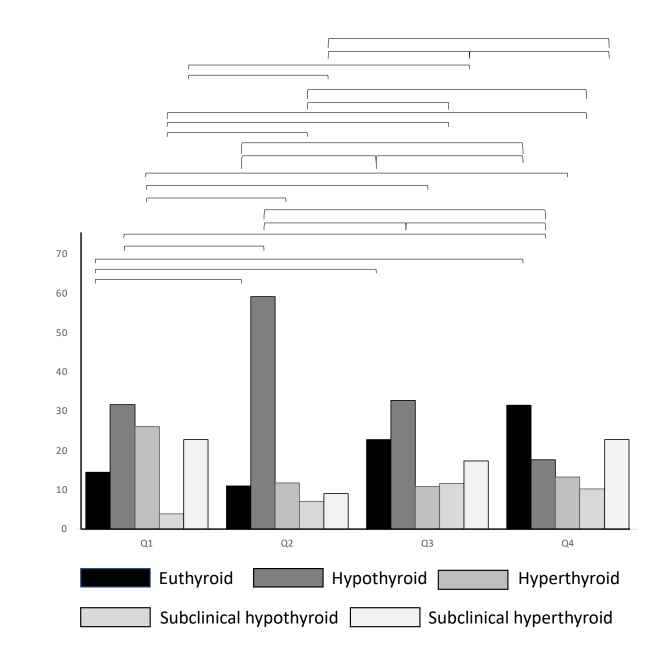


FIGURE 2

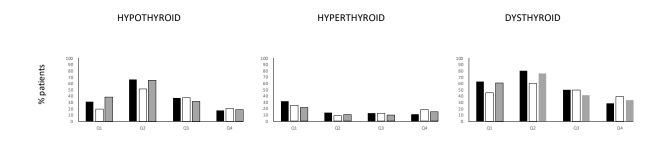


FIGURE 3

627 **TABLE 1**

628 Baseline characteristics of patients.

629 Patients enrolled

630	Total	812
631	Per centre	
632	Mean (SD)	26.1 (16.0)
633	Median (range)	21 (5-66)
634	95% CI	20.3-32.0
635	Age at time of most recent RI tr	eatment
636	Mean (SD) (years)	49.8 (14.2)
637	Median (range)	50 (18-89)
638	95% CI	48.8-50.8
639	Missing data (%, n)	2.0 (16/812)
640	Female	
641	% (n)	75.6 (615/812)
642	Missing data (%, n)	0 (0/812)

643 Activity of most recent RI

644	Mean (SD) (MBq)	481.8 MBq (101.8)
645	Median (range)	416 (330-809)
646	95% CI	474.7-488.9
647	Missing data (%, n)	2.0 (17/812)
648	Cumulative activity of RI	
649	Mean (SD) (MBq)	527.0 (196.0)
650	Median (range)	424.0 (330-1750)
651	95% CI	513.3-540.7
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661 **TABLE 2**

Rates of euthyroidism and dysthyroidism progressing through quarters (Qs) 1-4 in a subgroup of 358 patients who had at

least one set of thyroid function tests in each Q. Each Q represents a 3-month block from the date of radioiodine (RI)

664 treatment.665

666		CUMULATIVE RATES			
667		Q1*	Q2*	Q3*	Q4*
668					
669	Euthyroidism (%, n)	18.4 (66/358)	5.0 (18/358)	5.0 (18/358)	5.0 (18/358)
670					
671					
672	Hypothyroidism (%, n)	26.5 (95/358)	72.9 (261/358)	79.3 (284/358)	80.7 (289/358)
673					
674	Hyperthyroidism (%, n)	31.3 (112/358)	39.1 (140/358)	45.3 (162/358)	48.6 (174/358)
675					
676	Subclinical hypothyroid (%, n)	22.3 (8/358)	0.3 (1/358)	2.0 (7/358)	2.0 (7/358)
677					
678	Subclinical hyperthyroID (%, n)	26.0 (93/358)	7.0 (25/358)	2.2 (8/358)	2.2 (8/358)
679					
680	Missing data (%, n)	0 (0/358)	0 (0/358)	0 (0/358)	0 (0/358)
681					

682 *The sum of the rows in all Qs is greater than 100% because some patients had both hypo- and hyperthyroid episodes in

683 TABLE 3

684

685 Thyroid function tests by quarter (3-month blocks, or Qs) (A) and thyroid status by post-radioiodine treatment strategy (B).

- 686
- 687

689

688 Α

690 Q1 Q2 Q3 Q4 691 Serum TSH (mu/L) 692 Mean (SD) 8.9 (18.5)* 29.7 (36.0)* 11.3 (19.1)* 5.3 (11.4) 693 Median (range) 0.3 (0.0-122.9) 13.1 (0.0-195.9) 2.6 (0.0-146.8) 1.2 (0.0-144.7) 694 95% CI 7.5-10.2 27.0-32.3 9.9-12.8 5.3-6.4 695 724 713 647 455 n 696 697 Serum FT4 (pmol/L) Mean (SD) 18.0 (13.2) 12.5 (9.1) 16.6 (7.1) 18.1 (7.5) 698 699 Median (range) 14.4 (1.2-109.7) 10.7 (0.4-90.2) 15.8 (0.8-74.4) 17.4 (2.5-77.9) 16.0-17.2 11.8-13.2 700 95% CI 17.0-19.0 17.5-18.8 689 689 597 456 701 n 702

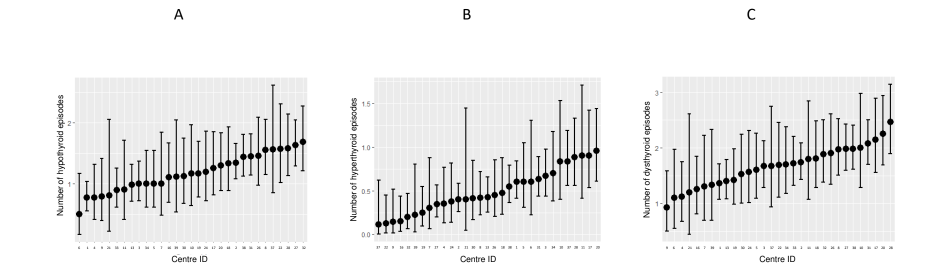
- 703 **B****
- 704

705					
706		Q1	Q2	Q3	Q4
707	All patients (%, n)				
708	Euthyroid	14.5 (105/723)	11.0 (79/715)	22.7 (147/648)	31.5 (165/524)
709	Hypothyroid	31.7 (229/723)	59.2 (423/715)	32.7 (212/648)	17.7 (93/524)
710	Hyperthyroid	26.1 (189/723)	11.7 (84/715)	10.8 (70/648)	13.2 (69/524)
711	Subclinical hypothyroid	3.9 (28/723)	7.0 (50/715)	11.6 (75/648)	10.3 (54/524)
712	Subclinical hyperthyroid	22.8 (165/723)	9.0 (64/715)	17.3 (112/648)	22.7 (119/524)
713	Other§	1.0 (7/723)	2.1 (15/715)	4.9 (32/648)	4.6 (24/524)
714	Missing data	0.1 (1/724)	0 (0/715)	0 (0/648)	0 (0/524)
715					
716	Anti-thyroid drugs alone (%, n)				
717	Euthyroid	15.5 (37/239)	6.6 (16/242)	19.3 (43/223)	34.1 (63/185)
718	Hypothyroid	29.7 (71/239)	63.2 (153/242)	36.3 (81/223)	17.8 (33/185)
719	Hyperthyroid	30.1 (72/239)	13.2 (32/242)	12.6 (28/223)	10.8 (20/185)
720	Subclinical hypothyroid	2.9 (7/239)	7.9 (19/242)	12.1 (27/223)	11.4 (21/185)
721	Subclinical hyperthyroid	20.1 (48/239)	8.3 (20/242)	17.5 (39/223)	22.7 (42/185)
722	Other§	1.7 (4/239)	0.8 (2/242)	2.2 (5/223)	3.2 (6/185)
723	Missing data	0 (0/239)	0 (0/242)	0 (0/223)	0 (0/185)
724					
725	Block and replace (%, n)				
726	Euthyroid	20.5 (17/83)	24.2 (22/91)	21.9 (21/96)	32.5 (26/80)

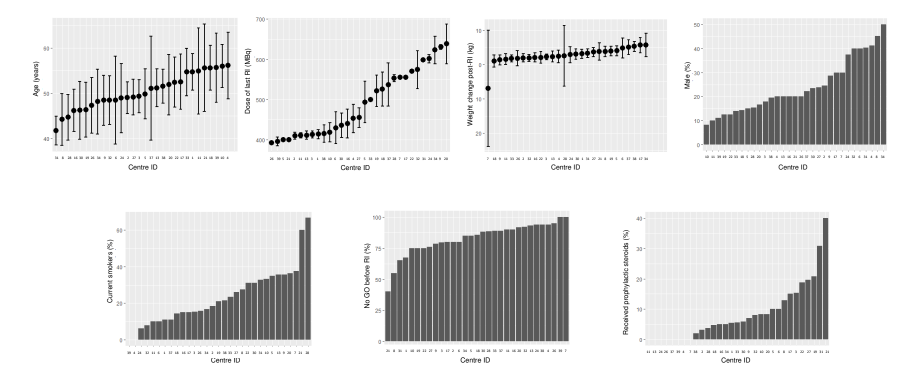
	Hypothyroid	19.3 (16/83)	49.4 (45/91)	36.4 (35/96)	20.0 (16/80)
8	Hyperthyroid	26.5 (22/83)	8.9 (8/91)	11.5 (11/96)	17.5 (14/80)
)	Subclinical hypothyroid	3.6 (3/83)	6.6 (6/91)	11.5 (11/96)	8.7 (7/80)
)	Subclinical hyperthyroid	26.5 (22/83)	6.6 (6/91)	13.5 (13/96)	16.3 (13/80)
l	Other§	3.6 (3/83)	4.3 (4/91)	5.2 (5/96)	5.0 (4/80)
2	Missing data	0 (0/83)	0 (0/91)	0 (0/96)	0 (0/80)
3					
4 L	L-T4 alone (%, n)				
5	Euthyroid	11.8 (44/373)	10.7 (38/356)	24.5 (74/302)	28.0 (67/239)
6	Hypothyroid	37.8 (141/373)	62.4 (222/356)	31.5 (95/302)	18.5 (44/239)
7	Hyperthyroid	22.5 (84/373)	11.0 (39/356)	9.9 (30/302)	14.6 (35/239)
3	Subclinical hypothyroid	4.6 (17/373)	6.2 (22/356)	11.6 (35/302)	10.1 (24/239)
)	Subclinical hyperthyroid	23.3 (87/373)	7.3 (26/356)	15.2 (46/302)	23.4 (56/239)
)	Other§	(0/373)	2.4 (9/356)	7.3 (22/302)	5.4 (13/239)
l	Missing data	0 (0/373)	0 (0/356)	0 (0/302)	0 (0/239)
2					
3					

TSH concentration; hyperthyroid: serum TSH below and serum FT4 above the reference ranges; subclinical hypothyroid: serum TSH above the reference range but <10 mU/L and serum FT4 within the reference range; subclinical hyperthyroidism: serum FT4 within and serum TSH below
 the reference range; euthyroid: both serum TSH and FT4 within the reference range or, when only one was available, within the reference range.

750 §TSH and FT4 did not conform to any of the listed categories (both elevated or both reduced).



SUPPLEMENTARY FIGURE 1 (UPPER PANEL)



- 55 SUPPLEMENTARY FIGURE 1 (LOWER PANEL)

759 SUPPLEMENTARY TABLE 1

760 Baseline characteristics of all patients and in patients and in patients treated with anti-thyroid drugs, block and replace

and L-T4 alone post-radioiodine (RI).

763 764	ALL ENROLLED PATIENTS STRATEGY		PATIENT GROUPS BY POST-RI MANAGEMENT		
764 765			Anti-thyroid	Block and	L-T4
766 767	alone		drugs alone	replace	
768 769	n	812	255	109	355
770					
771 772 773	Age at time of most recent RI treatment				
774	Mean (SD) (years) 49.8 (14.2)	48.3 (13.8)*	52.1 (13.6)*	50.0 (14.6)	
775	Median (range)	50 (18-89)	49.0 (19-83)	50.0 (22-86)	50.0
776		(18-89)			
777	95% CI	48.8-50.8	46.5-50.0	49.5-54.6	48.4-
778		51.5			

779	Missing data (%, n)	2.0 (16/812)	2.0 (5/255)	0 (0/109)	2.2
780		(8/355)			
781	Female				
782	% (n)	75.6 (615/812)	72.2 (184/255)	74.3 (81/109)	78.3
783	(278/355)				
784	Missing data (%, n)	0 (0/812)	0 (0/255)	0 (0/109)	0
785	(0/355)				
786	Activity of most recent RI				
787	Mean (SD) (MBq)	481.8 MBq (101.8)	508.0 (111.0)**	459.9 (91.4)***	472.0
788		(97.5)			
789	Median (range)	416 (330-809)	531 (364-809)	400.0 (390-800)	410.5
790		(330-800)			
791	95% CI	474.7-488.9	494.2-521.8	442.5-477.3	461.7-
792		482.2			
793	Missing data (%, n)	2.0 (17/812)	2.0 (5/255)	0.9 (1/109)	2.0
794		(7/355)			

795 Time from diagnosis of Graves'

796 707						
797 798	Mean (SD) (months)	38.3 (43.7)	36.4 (35.8)	37.4 (31.4)	41.3	
799	(49.3)					
800	Median (range)	26.9 (-8.9-458.0)§	27.3 (-8.9-283.1)	27.2 (1.0-186.0)	28.3	
801	(0-464.1)					
802	95% CI	35.3-41.3	32.0-40.8	31.4-43.3	36.1-	
803	46.5					
804	Missing data (%, n)	1.2 (10/812)	0 (0/255)	0 (0/109)	1.1	
805	(4/355)					
806	Smoking status					
807	Never smoked (%, n)	52.3 (401/767)	47.8 (122/243)	59.6 (62/104)	50.6	
808	(174/336)					
809	Ex-smoker (%, n)	26.1 (200/767)	26.3 (64/243)	22.1 (23/104)	26.8	
810	(90/336)					
811	Current smoker (%, n)	21.6 (166/767)	23.5 (57/243)	18.3 (19/104)	21.4	
812	(72/336)					

813	Missing data (%, n)	5.5 (45/812)	2.4 (12/255)	4.6 (5/109)	5.4
814	(19/355)				
815 816 817 818	Treatment with curative intent for hyperthyroidism prior to most recent radioiodine treatment				
819	No treatment (%, n)	14.9 (121/812)	1.1 (28/255)	10.1 (11/109)	20.3
820	(72/355) ^a				
821	Course of anti-thyroid				
822	drugs (%, n)	75.1 (610/812)	78.4 (200/255)	79.8 (87/109)	69.6
823	(247/355) ^a				
824	Thyroidectomy (%, n)	0.6 (5/812)	0.8 (2/255)	1.8 (2/109)	0.3
825	(1/355)				
826	Radioiodine (%, n)	9.4 (76/812)	9.8 (25/255)	8.2 (9/109)	9.9
827	(35/355)				
828	Missing data (%, n)	0 (0/812)	0 (0/255)	0 (0/109)	0
829	(0/355)				
830 831	Graves' orbitopathy before RI (%, n)	18.2 (148/812)	20.8 (53/255)	28.4 (31/109) ^b	13.2
832	(47/355) ^c				
833	Missing data (%, n)	0 (0/812)	0 (0/255)	0 (0/109)	0
834	(0/355)				

- *The mean age of patients treated with block and replace was significantly greater than patients treated with anti-thyroid drugs alone (unpaired t-
- test p=0.017); all other comparisons of age between groups were statistically insignificant.
- **Patients treated with anti-thyroid drugs had received a higher dose of RI than the entire group (p=0.000), than the groups treated with block and replace (p=0.000), and the group treated with L-T4 (p=0.000).
- 840 ***Patients treated with block and replace had received a lower dose of RI than the entire group (p=0.028).
- 841 §In a single case, the diagnosis of Graves' disease as the cause of thyrotoxicosis was made after the RI.
- ^aPatients treated with L-T4 were less likely to have received definitive treatment before RI compared to patients treated with anti-thyroid drugs
 alone (p=0.02, Chi-squared test) and patients treated with block ad replace (p=0.02, Chi-squared test).
- ^bPatients treated with block and replace had a higher frequency of Graves' orbitopathy before RI treatment than the rest of the cohort (p=0.014)
- 845 and patients treated with L-T4 (p=0.000, Chi-squared test)..
- 846 Patients treated with L-T4 had a lower frequency of Graves' orbitopathy before RI treatment than the rest of the cohort (p=0.04, Chi-squared test),
- 847 than patients treated with anti-thyroid drugs alone (p=0.015, Chi-squared test), and lower than patients treated with block and replace (p=0.000,
- 848 *Chi-squared test).*
- 849