Star-Shaped Intense Uptake of $^{131}$I on Whole Body Scans Can Reflect Good Therapeutic Effects of Low-Dose Radioactive Iodine Treatment of 1.1 GBq

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Background: After initial radioactive iodine (RAI) treatment in differentiated thyroid cancer patients, we sometimes observe a star-shaped region of intense uptake of $^{131}$I on whole body scans (WBSs), called a ‘star artifact.’ We evaluated the clinical implications of star artifacts on the success rate of remnant ablation and long-term prognosis.

Methods: Total 636 patients who received $^{131}$I dose of 1.1 GBq for the initial RAI therapy and who did not show distant metastasis at the time of diagnosis were retrospectively evaluated. A negative second WBS was used for evaluating the ablation efficacy of the RAI therapy. Among them, 235 patients (36.9%) showed a star artifact on their first WBS.

Results: In patients with first stimulated thyroglobulin (sTg) levels $\leq$ 2 ng/mL, patients with star artifacts had a higher rate of negative second WBS compared with those without star artifacts (77.8% vs. 63.9%, $P=0.044$), and showed significantly higher recurrence-free survival ($P=0.043$) during the median 8.0 years (range, 1.0 to 10.0) of follow-up. The 5- and 10-year recurrence rates (5YRR, 10YRR) were also significantly lower in patients with star artifacts compared with those without (0% vs. 4.9%, respectively, $P=0.006$ for 5YRR; 0% vs. 6.4%, respectively, $P=0.005$ for 10YRR). However, ablation success rate or recurrence-free survival was not different among patients whose first sTg levels $>2$ ng/mL regardless of star artifacts.

Conclusion: Therefore, star artifacts at initial RAI therapy imply a good ablation efficacy or a favorable long-term prognosis in patients with sTg levels $\leq$ 2 ng/mL.

Keywords: Radioactive iodine; Remnant ablation; Star artifact; Thyroid neoplasms; Therapeutic uses
INTRODUCTION

Differentiated thyroid cancer (DTC) is known to be one of the most curable cancers, and radioactive iodine (RAI) treatment has been contributed to its excellent prognosis. Thus, still a large number of DTC has been treated with RAI therapy after total thyroidectomy, depending on the risk stratification of current guidelines [1,2].

During the initial RAI treatment in DTC, a star-shaped region of intense $^{131}$I uptake can be observed on whole body scans (WBSs) sometimes. This phenomenon is called a ‘star artifact’ (Fig. 1) and has been reported to be caused by the penetration of the collimator’s septa by the $\gamma$-rays from the high uptake of RAI in the thyroid [3,4]. Although this phenomenon is often seen in clinics, there has been no investigation into its clinical implications or outcomes.

There are a few theories regarding the high uptake of RAI that causes star artifacts. First of all, it could be caused by large amounts of remnant thyroid tissue after surgery. It was reported that patients with large amounts of remnant thyroid tissue had a higher uptake of RAI, compared with those who did not [5]. However, thyroid tissues with high expression of the sodium-iodine symporter (NIS) also could cause an excessively high uptake of RAI; active transport by the NIS leads to iodide uptake across the membranes of thyroid cancer cells [6]. Therefore, it is still questionable whether the presence of a star artifact is a poor prognostic marker representing a large amount of remnant thyroid or residual tumor that indicates the need for a more thorough evaluation, or a good prognostic marker reflecting better treatment efficacy. If it is a good prognostic marker, a relatively lower dose of second RAI ablation might be sufficient or even a diagnostic WBS might be not required in patients with a star artifact. Considering that the usefulness of the diagnostic WBS or second RAI treatment after initial low dose of RAI remnant ablation treatment is still unclear [1], determining the prognostic value of the star artifacts would be helpful for the planning of further managements.

Therefore, in this study, we investigated the clinical implications of the star artifact observed after initial RAI therapy by comparing the success rate of remnant ablation and long-term recurrence rate between patients with and without a star artifact.

METHODS

Study population

We reviewed 720 consecutive patients who underwent total thyroidectomy and RAI therapy for the treatment of DTC between January 2005 and December 2007 in the Seoul National University Hospital. Among them, 72 patients who initially received $^{131}$I dose over 1.1 GBq (30 mCi), and 12 patients showing distant metastasis at the initial WBS were excluded. The remaining 636 patients (125 men [19.7%] and 511 women [80.3%]; mean age, 59.0±11.0 years) who received 1.1 GBq of $^{131}$I were analyzed in this study. The median duration of the follow-up was 8.0 years (interquartile range [IQR], 1.0 to 10.0). The study was approved and a patient’s informed consent was waived by the Institutional Review Board (IRB) of the Seoul National University Hospital (IRB number 1104-080-359).

RAI ablation and definition of a ‘star artifact’ on WBS images

For $^{131}$I RAI therapy, 608 patients discontinued thyroxine replacement at least 4 weeks prior to $^{131}$I administration and were switched to triiodothyronine, which was discontinued 2 weeks before $^{131}$I administration. Iodine restriction diet was started 2 weeks before $^{131}$I administration. Serum samples for thyroglobulin (Tg) and anti-thyroglobulin antibody (TgAb) measurement were collected 1 hour before $^{131}$I administration. The other 28 patients underwent recombinant human thyroid stimulating hormone (rhTSH; Thyrogen, Genzyme Corp., Cambridge, MA, USA).

Fig. 1. Representative $^{131}$I whole body scan images (A-C) with and (D) without star artifact after 1st 1.1 GBq of radioactive iodine treatment.
USA) therapy; after adhering to iodine restriction diet for 2 weeks, rhTSH was administered (0.9 mg intramuscular) for 2 consecutive days. Tg and TgAb levels at the time of ablation were measured on the 5th day after the first rhTSH injection, which corresponded to the 3rd day after \(^{131}\)I administration [7]. Thyroxine was stopped the day before the 1st administration of rhTSH, and started again the day after WBS. For both stimulation methods, serum levels of thyroid stimulating hormone (TSH) over 30 IU/mL were considered sufficient.

A post-therapy WBS was acquired 2 days after \(^{131}\)I administration using a large field-of-view gamma camera (ON 410, Ohio Nuclear, Solon, OH, USA) with a high-energy general purpose collimator. A 20% symmetric window was centered at 364 keV. A total of 100,000 counts were accumulated in each case, and anterior images of the neck, chest, and abdomen were obtained.

The presence of star artifacts was assessed according to the pattern of \(^{131}\)I uptake in the remnant thyroid on WBS after the initial RAI administration. As shown in Fig. 1, the star artifact in our study was defined as a star-shaped region of intense uptake of \(^{131}\)I, reviewed by two independent physicians (S.H.K. and J.A.L.). The final consensus for the discrepant cases was made after discussion with another physician (Y.I.P.).

Follow-up protocols and the definition of recurrence
In some patients (n=50) who showed no or faint RAI uptake only in the thyroid bed with stimulated thyroglobulin (sTg) \(<0.2\) ng/mL at the time of initial ablation, neck ultrasonography was repeated and the Tg level on TSH suppression was measured 6 to 12 months after initial ablation. Otherwise, 2nd WBS using 0.15 to 1.1 GBq (4 to 30 mCi) of \(^{131}\)I was performed (n=586). Regular follow-up included a physical examination, measurement of TSH, free thyroxine, total triiodothyronine, serum Tg and TgAb levels every 6 to 12 months, and neck ultrasonography every 1 to 3 years. Levothyroxine was administered to achieve serum TSH levels during the first 5 to 10 years after initial treatment [8].

Clinical recurrence was defined as a reappearance of pathologically proven malignant tissue and/or an appearance of definitive metastatic lesions in other organs by WBS, positron emission tomography computed tomography (PET-CT), or CT, assessed by radiologic specialists [8].

Serum Tg and TgAb measurement
Serum Tg was measured with an immunoradiometric assay (RIA Tgplus, BRAHMS GmbH, Hennigsdorf, Germany). The analytical sensitivity read at the optimal curve was 0.08 ng/mL, and the functional assay sensitivity (20% interassay coefficient of variation) was 0.2 ng/mL. TSH level was measured with an immunoradiometric assay (TSH-CTK-3, DiaSorin, Saluggia, Italy). The analytical sensitivity was 0.04 mU/L, and the functional assay sensitivity was 0.07 mU/L. TgAb was measured with a radioimmunometric assay (anti-Tg, BRAHMS GmbH). The analytical sensitivity was 5.5 U/mL, and the functional assay sensitivity was \(<20\) U/mL. TgAb was defined as positive when the levels were >60 U/mL.

Statistical analysis
Categorical variables are presented as numbers and percentages, and continuous variables are expressed as mean±standard deviation (SD) for normal distribution, or medians with IQR for nonnormal distribution. Pearson chi-square test was used to compare categorical variables. Continuous variables were compared using Student t test for normally distributed variables and the Mann-Whitney U test for nonnormal distributions. Recurrence-free survival (RFS) curves were constructed using the Kaplan-Meier method and compared using the log-rank test. All P values were two-sided with P<0.05 considered statistically significant. All statistical analyses were performed with SPSS software version 22.0 (IBM Co., Armonk, NY, USA).

RESULTS
Baseline characteristics
Among the 636 participants, 235 patients (36.9%) exhibited the star artifacts (group with star artifacts) (Fig. 1A-C) and 401 patients (63.1%) did not (group without star artifacts) (Fig. 1D) after the first RAI therapy. The baseline characteristics of the two groups are shown in Table 1. In the group with star artifacts, the patients were younger (P=0.002), and had a lower proportion of papillary thyroid cancer (P=0.001), compared to the other group. Among the patients without TgAb, sTg levels were significantly higher in the group with star artifacts than in the group without (P<0.001). In addition, there was a higher proportion of patients with sTg \(\leq 2\) ng/mL in the group without star artifacts than in the group with star artifacts (P=0.005). There were no significant differences in gender, follow-up duration, tumor size, multiplicity, extrathyroidal extension, nodal metastasis status, proportions of patients with TgAb, and total dose of RAI between the two groups.
Success rate of ablation after initial RAI treatment in TgAb negative patients

There was no significant difference in the success rate after initial RAI treatment between patients with and without star artifacts, as defined by a negative result in the second WBS. In the subgroup analysis depending on the initial sTg levels, the ablation success rate was higher in the group with star artifacts compared with those without star artifact among patients whose initial sTg levels were <0.2 ng/mL ($P=0.027$), ≤1 ng/mL ($P=0.025$), or ≤2 ng/mL ($P=0.044$). However, the ablation success rate defined by both a negative WBS and sTg <0.2 ng/mL was not different between groups even after the same subgroup analysis (Table 2).

Next, we analyzed odds ratios (OR) regarding ablation success rate, which defined by a negative WBS only, according to the presence of star artifacts in each initial sTg cut-off level (<0.2, ≤1, and ≤2 ng/mL). We defined ‘lower sTg’ as initial sTg levels same or lower than a cut-off value, and ‘higher sTg’
as initial sTg levels higher than a cut-off value. Compared to the patients with lower sTg without star artifacts, ablation success rates were significantly higher in patients with lower sTg levels with star artifact (OR, 2.579; 95% confidence interval [CI], 1.141 to 5.818; \( P = 0.023 \)) for sTg cut-off of 0.2 ng/mL (OR, 2.431; 95% CI, 1.150 to 5.141; \( P = 0.020 \)) for 1 ng/mL (OR, 1.973; 95% CI, 1.031 to 3.779; \( P = 0.040 \)). In patients with higher sTg with or without star artifact, ablation success rates were not different from those with lower sTg without star artifact (Table 3).

However, as expectedly, ablation success rates relative to patients with a star artifact but with higher sTg levels were not significantly increased nor decreased in all patients without star artifacts and higher sTg levels in all cut-off values (OR, 1.692; 95% CI, 0.692 to 4.135; \( P = 0.249 \)) for sTg cut-off of 0.2 ng/mL (OR, 1.788; 95% CI, 0.724 to 4.417; \( P = 0.208 \)) for 1 ng/mL (OR, 1.519; 95% CI, 0.592 to 3.893; \( P = 0.384 \)) for 2 ng/mL).

### Comparison of recurrence rate between groups with and without star artifacts

Next, we compared the recurrence rate according to the presence of star artifacts at the 1st WBS. During a median follow-up of 8.0 years (IQR, 1.0 to 10.0), recurrence was detected in 46 patients (7.2%); those who had sTg \( \leq 2 \) ng/mL showed a lower rate of clinical recurrence than those who had sTg > 2 ng/mL (3.1% vs. 19.4%, log rank \( P < 0.001 \)).

Among them, 16 patients (6.8%) with star artifacts and 30 patients (7.5%) without star artifacts developed clinical recurrence (\( P = 0.874 \)). There were no significant differences in RFS between the groups with and without star artifact; median RFS were 22.1 and 20.5 years, respectively (\( P = 0.636 \)). 5YRR and 10YRR were also not different (Fig. 2A).

We performed a subgroup analysis according to the presence of sTg Ab. Among the 311 patients with positive TgAb, there were no significant differences in RFS between the groups with and without star artifacts (median RFS, 19.1 and 21.2 years, respectively; \( P = 0.466 \)). There also was no difference in 5YRR or 10YRR between the groups (Fig. 2B). In a total 325 patients with negative TgAb, similar results were observed; the presence of a star artifact did not influence the recurrence at all (data not shown).

However, when we re-analyzed the subgroups without TgAb according to the initial sTg levels, among the 219 patients whose sTg level was \( \leq 2 \) ng/mL, patients with star artifacts showed significantly higher RFS than those without star artifacts (log rank \( P = 0.043 \)). The median RFS could not be calculated because there was no case of recurrence in the group with star artifacts, but 5YRR and 10YRR were significantly lower in patients with star artifact than those without star artifact (0% vs. 4.9%, \( P = 0.006 \) for 5YRR; 0% vs. 6.4%, \( P = 0.005 \) for 10YRR) (Fig. 2C). However, in the 112 patients whose sTg level was > 2 ng/mL, there were no significant differences in RFS between groups with and without star artifacts (median RFS, 9.4 and 8.8 years, respectively; \( P = 0.462 \)). Both 5YRR and 10YRR were not different either (Fig. 2D).

### DISCUSSION

In this study, we found that star artifacts resulting from the strong uptake of \( ^{131}I \) could be a good prognostic factor reflecting higher ablation success rates after the initial RAI therapy or long-term prognosis in patients with sTg level \( \leq 2 \) ng/dL. The sTg level has been thought a well-known predictive marker for recurrence in DTC because it could reflect the amount of residual mass of thyroid, including cancer, and patients with higher sTg levels have shown a higher probability of recurrence when...
they are negative for TgAb [9]. In cases with high sTg levels, star artifacts might indicate a large amount of residual thyroid mass; if that were the case, the presence of a star artifact would be a poor prognostic factor. On the other hand, in cases with low sTg levels, star artifacts might instead reflect an increased RAI uptake by the residual thyroid tissue. If the expression of NIS in the residual tissue is upregulated, the RAI uptake would be greater even if the residual mass was lower. In such cases, the star artifact could be a good prognostic factor to predict a successful ablation after RAI therapy.

In the study, the recurrence rate was significantly higher in patients with sTg levels >2 ng/mL than in those with sTg levels ≤2 ng/mL, as previously reported [10]. Within the subgroup of patients with sTg levels >2 ng/mL, star artifacts was not associated with the rate of recurrence, and it was same when the analysis was performed in patients with a sTg value of 2 to 10 ng/mL. In other words, in patients with higher Tg levels, which may represent a larger residual mass, it is difficult to conclude whether the star artifact indicates a high RAI uptake capability or large amount of residual thyroid. It is suspected that the prognostic power of high sTg levels might be strong enough to mask the effects of the star artifact. On the other hand, in patients with sTg levels ≤2 ng/mL, RFS and the proportion of negative subsequent WBS was significantly higher in patients with star artifacts than in those without. This implicates that the presence of a star artifact in patients with low sTg levels, which likely represent a small residual thyroid mass, reflects the high RAI uptake of the residual thyroid, resulting in good therapeutic effects.
A putative explanation of the good prognostic value of star artifacts could be a different degree of differentiation of remnant thyroid tissue, mainly because of NIS expression. NIS is a well-known main carrier of iodide transport in thyrocytes [6]. In a previous study, patients with positive NIS immunostaining responded to RAI treatment better than those with negative immunostaining, although the authors did not provide recurrence or mortality data [11]. Therefore, we could hypothesize that when the remnant tumor size is similar, a higher uptake of RAI due to a higher expression of NIS could increase the efficacy of RAI treatment, and that might lead to a better prognosis. Additionally, it has been proposed that measuring NIS expression in postoperative tissue might predict the effectiveness of RAI treatment, and could be helpful to determine individualized doses of RAI [11]. Unfortunately, we did not measure the expression of NIS in this study, and further study is required to prove the hypothesis.

On the other hand, there still exists a possibility that the star artifact might reflect the high burden of remnant thyroid tissue and lead to a poor prognosis. Schneider et al. [5] reported that recurred patients had a 10-fold higher uptake-to-dose ratio of RAI in remnant thyroid compared with those who did not. They suggested that the high uptake of RAI in remnant thyroid could be a risk factor for a recurrent disease. However, the previous study did not measure sTg levels, which might provide information about the residual amount of thyroid tissue, unlike the present study, in which the analysis was stratified according to sTg levels. The subgroup of patients with higher sTg levels had a higher recurrence rate and no significant difference in recurrence rates between patients with and without star artifacts. Therefore, sTg levels could be a better marker for reflecting the burden of remnant thyroid cancer or poor prognosis than star artifacts in those with high sTg levels.

The current study is the first study regarding the clinical outcomes of star artifacts, which found in almost 40% of patients after the initial RAI treatment. There has only been a case report [4] and a review that mentioned star artifacts [3]. Because most hospitals perform measurements of dose-to-response ratio during WBS only in research settings [5], it is more practical to base the assessment on the presence of star artifacts, rather than measuring dose-to-response ratio. In addition, we performed a stratified analysis according to the patients’ sTg levels, allowing us to distinguish patients with a high burden of remnant thyroid cancer. Lastly, the duration of follow-up in this study was quite long (median, 8 years), which enables a comparison of long-term recurrence, but not mortality data.

The study also has a few limitations. As previously described, we could not evaluate NIS expression in postoperative thyroid tissue to confirm our hypothesis. Also, because the incidence of star artifacts could be variable by doses of RAI therapy and the acquisition methods such as camera and collimators, the incidence of star artifact in this study cannot be generalized in other settings. In addition, the presence of star artifacts was determined subjectively, and we were unable to assess the correlation between dose-to-response ratio and star artifacts in this retrospective study. Dose-to-response ratio can be measured during the WBS, but this is not a routine process in the hospital. However, the star artifact itself could be useful in clinical settings, because the measurement of dose-to-response ratio is rarely done in most hospitals.

In conclusion, in patients with low sTg, a star-shaped intense uptake on WBS, a star artifact, may reflect the positive therapeutic efficacy of initial RAI therapy in patients with DTC. It might be applicable as a prognostic factor in patients with sTg levels \(\leq 2 \text{ ng/mL}\).

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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AUTHOR CONTRIBUTIONS

Conception or design: Y.J.P., J.A.L., S.H.K. Acquisition, analysis, or interpretation of data: J.A.L., S.H.K. Drafting the work or revising: S.H.K., J.A.L. Final approval of the manuscript: S.H.K., J.A.L., Y.S.S., S.M., Y.A.K., M.J.K., S.W.C., J.H.M., K.H.Y., D.J.P., B.Y.C., Y.J.P.

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REFERENCES

1. Haugen BR, Alexander EK, Bible KC, Doherty GM, Mandel SJ, Nikiforov YE, et al. 2015 American Thyroid Association management guidelines for adult patients with thyroid nodules and differentiated thyroid cancer: the American Thyroid Association guidelines task force on thyroid nodules and differentiated thyroid cancer. Thyroid 2016;26:1-133.

2. Chung JK, Cheon GJ. Radioiodine therapy in differentiated thyroid cancer: the first targeted therapy in oncology. Endocrinol Metab (Seoul) 2014;29:233-9.

3. Wartofsky L, Van Nostrand D. Thyroid cancer. Totowa: Humana Press Inc.; 2006. Chapter 36 part A, Stunning: untoward effect of 131I thyroid imaging prior to radioablation therapy; p. 337-45.

4. Anastasios G, Foteini C, Dimitrios T, Philippos K. Functioning metastasis of papillary thyroid carcinoma originating as a double star effect on the post-therapeutic radioiodine whole body scan. Intern Med 2008;47:1155-6.

5. Schneider DF, Ojomo KA, Chen H, Sippel RS. Remnant uptake as a postoperative oncologic quality indicator. Thyroid 2013;23:1269-76.

6. Filetti S, Bidart JM, Arturi F, Caillou B, Russo D, Schlumberger M. Sodium/iodide symporter: a key transport system in thyroid cancer cell metabolism. Eur J Endocrinol 1999;141:443-57.

7. Mazzaferri EL, Kloos RT. Using recombinant human TSH in the management of well-differentiated thyroid cancer: current strategies and future directions. Thyroid 2000;10:767-78.

8. Choi H, Lim JA, Ahn HY, Cho SW, Lee KE, Kim KW, et al. Secular trends in the prognostic factors for papillary thyroid cancer. Eur J Endocrinol 2014;171:667-75.

9. Gibelli B, Tredici P, De Cicco C, Bodei L, Sandri MT, Renne G, et al. Preoperative determination of serum thyroglobulin to identify patients with differentiated thyroid cancer who may present recurrence without increased thyroglobulin. Acta Otorhinolaryngol Ital 2005;25:94-9.

10. Kim TY, Kim WB, Kim ES, Ryu JS, Yeo JS, Kim SC, et al. Serum thyroglobulin levels at the time of 131I remnant ablation just after thyroidectomy are useful for early prediction of clinical recurrence in low-risk patients with differentiated thyroid carcinoma. J Clin Endocrinol Metab 2005;90:1440-5.

11. Min JJ, Chung JK, Lee YJ, Jeong JM, Lee DS, Jang JJ, et al. Relationship between expression of the sodium/iodide symporter and 131I uptake in recurrent lesions of differentiated thyroid carcinoma. Eur J Nucl Med 2001;28:639-45.