Patterns of thyroid hormone levels in pediatric medullary thyroid carcinoma patients on vandetanib therapy

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Abstract

Background: Tyrosine kinase inhibitors (TKIs) have been associated with elevated TSH as a drug class effect. Prior studies of vandetanib in adults with medullary thyroid carcinoma (MTC) described an increase in levothyroxine (LT) requirement. We studied TSH, free T4, and LT dosing in children and adolescents enrolled in the phase I/II trial of vandetanib for medullary thyroid cancer (MTC).

Methods: Data from 13 patients with multiple endocrine neoplasia type 2B (MEN 2B) and MTC were analyzed [6 M, 7 F, median age 13.0 y (9.1-17.3)] Eleven patients (85%) had undergone prior thyroidectomy and all received single-drug therapy with vandetanib for > 6 months. Confirmed compliance with vandetanib (67–150 mg/m²/day) and LT was a necessary inclusion criterion.

Results: While on vandetanib treatment, all 11 athyreotic patients exhibited significantly increased TSH levels. The baseline TSH level was 4.37 mIU/ml (0.08 - 23.30); in comparison, the first peak TSH concentration on vandetanib was 15.70 mIU/ml (12.50 - 137.00, p = 0.0010). The median time to reach the initial peak of elevated TSH was 1.8 months (0.3 - 9.3). Free T4 levels remained within the normal reference range. An increase from a baseline LT dose of 91 mcg/m²/day (±24) to 116 mcg/m²/day (±24) was required in order to resume normative TSH levels (p = 0.00005), equal to an increase of 36.6% (±16.56) in the dosage of LT in mcg/day. For the 2 patients with intact thyroid glands, free T4 and TSH remained normal over a combined 6 patient years of follow up.

Conclusions: In our cohort of pediatric MTC patients, athyreotic patients with preexisting hypothyroidism developed increased TSH and reduced free T4 during the first few months of treatment with vandetanib, necessitating an increase in LT dosage. Additional patients with normal thyroid function before treatment and intact glands (n = 2) maintained normal thyroid function tests during treatment. Elevated TSH in athyreotic patients may be due to an indirect effect of vandetanib on the metabolism of thyroid hormone, or to altered TSH sensitivity at the pituitary. Proper recognition and management of abnormal thyroid hormone levels is critical in growing children on TKIs.

Trial registration: ClinicalTrials.gov Identifier: NCT00514046

Keywords: Multiple Endocrine Neoplasia type 2 B (MEN2B), Medullary thyroid carcinoma (MTC), Tyrosine kinase inhibitor (TKIs), Vandetanib
Background
Hereditary medullary thyroid carcinoma (MTC) is a manifestation of multiple endocrine neoplasia (MEN) type 2A and MEN 2B syndromes caused by germline, activating mutations in the RET (REarranged during Transfection) proto-oncogene (10q11.2). MTC accounts for 5-10% of all pediatric thyroid malignancies and its annual incidence is 0.3-2 cases per 10^6 children (1 M: 1 F) [1-3].

Only 25% of MTC cases are hereditary (Familial MTC (FMTC), MEN2A, MEN2B) while the majority, 75% of MTC cases, are sporadic. Hereditary MTC is multifocal, bilateral, indolent and usually presents with metastasis at the time of diagnosis. If diagnosed while the tumor is confined to the thyroid, MTC has a favorable prognosis (10 year survival rate 70-80%) [4]. Common sites of metastatic disease include cervical and mediastinal lymph nodes, as well as lungs, liver, and bone; in the case of metastatic disease the 10 year survival rate drops to approximately 40% [4-7]. Metastatic or locally advanced MTC is unresponsive to cytotoxic chemotherapy or radiation [4,8]. Targeted therapy with Tyrosine Kinase inhibitors (TKIs) are now approved to treat progressive and advanced MTC. TKIs compete with the ATP-binding domains of the TK catalytic unit inhibiting the activation of oncogenic intracellular signaling pathways [9]. Vandetanib (Caprelsa, Astra Zeneca Pharmaceuticals) is an orally bioavailable multi-RTK (receptor Tyrosine kinase) inhibitor that blocks the mutant RET gene product and has antitumor activity in adults with hereditary MTC [10,11]. At the National Cancer Institute, a phase I/II trial of vandetanib for children and adolescents with MTC was conducted to define a recommended dose and assess antitumor activity, as described by Fox et al. [12]. A common non-dose limiting toxicity was TSH elevation necessitating an increase in levothyroxine (LT) dosage – however, this was only noted in athyreotic patients who were previously on a stable dose. TKIs have been associated with elevated TSH as a drug class effect [13]; two prior studies of vandetanib in adults with MTC described an increase in LT requirement [14,15]. Little is known about the effect of vandetanib for children and adolescents with MTC, NCT00514046.

Methods
Data from 13 patients with MEN 2B and MTC were analyzed [6 M, 7 F, median age 13.0 y (9.1-17.3)] (Table 1). Patients were enrolled in the phase I/II trial of vandetanib for Multiple Endocrine Neoplasia Type 2B (MEN 2B) and MTC, NCT00514046. Informed consent from the patients’ parents (and assent from older children) was obtained for all patients. Eleven patients (85%) had undergone prior thyroidectomy and all received single-drug therapy with vandetanib for > 6 months. Confirmed compliance with vandetanib (67–150 mg/m^2/day) and LT was a necessary inclusion criterion (Table 1). Patients were instructed to take LT on an empty stomach and not with other medications or calcium/soy containing products. One patient was excluded due to non-compliance with his replacement with LT. Data were analyzed using paired t-tests for normally distributed data and the Wilcoxon signed rank test for non-parametric data, and are reported as mean (±SD) or median (range).

Results
While on vandetanib treatment, all 11 athyreotic patients exhibited significantly increased TSH levels. The baseline TSH level was 4.37 mclU/ml (0.08 - 23.30). In comparison, the first peak TSH concentration on vandetanib was 15.70 mclU/ml (12.50 - 137.00, p = 0.0010) (Figure 1). The median time it took to reach the first peak of elevated TSH was 1.8 months (0.3 - 9.3). Free T4 levels remained within the normal reference range, yet significantly decreased from baseline levels of 1.47 ng/dL (±0.21) to 1.27 ng/dL (±0.30) when measured at the time of maximum TSH (p = 0.039) (Figure 2). TSH levels normalized after subsequent increases in LT dose. An increase from a baseline LT dose of 91 mcg/m^2/day (±24) to 116 mcg/m^2/day (±24) was required in order to resume normative TSH levels (p = 0.00005), (Figure 3), equal to an increase of 36.6% (±16.56) in the dosage of LT in mcg/day. There were no clinical sequelae as a result of the elevations in TSH. Thyroid hormone levels increased, either due to a change in LT dose, or in one patient, due to vandetanib being held for oral surgery without adjustment of the LT dose to account for the effect of the TKI. Linear growth was closely monitored for the duration of the study. The median percentile of height for age at baseline was 30 (<3-96)% and increased to 55 (<3-96)% at the last evaluation (P = 0.03). The median percentile of weight for age at baseline was 9 (<3-96)% and increased to 20 (<3-91)% at last evaluation (P = 0.48). For the 2 patients with thyroid glands still intact, free T4 averaged 1.17 ± 0.15, (normal range 0.8-1.5 ng/dL) and TSH levels averaged 3.48 ± 2.18, (normal range 0.4-4 mclU/ml) over a combined 6 patient years of follow up.

Table 1 Patient Characteristics

| Number of patients | 13 |
|-------------------|----|
| Gender (male/female) | 6/7 |
| Age at start of enrollment on clinical trial (median, range) | 13.0 years (9.1-17.3) |
| RET mutation | M918T (n = 13) |
| Baseline LT dose | 100 mcg/day (57–200) |
Discussion
Innovative therapeutic agents that target genetic alterations (selective cancer therapy) associated with the development of MTC seem promising to treat progressive and advanced MTC [16]. Even if TKIs combine a high therapeutic window with less toxicity than conventional chemotherapy, vandetanib's adverse effects varied in respect to the administered dose and included rash, nausea, hypertension, headache, ECG QTc prolongation and endocrine effects [17]. In a randomized controlled trial of vandetanib for adults with MTC in which 90% of patients enrolled had prior thyroidectomy, increased dosing of LT was required in 49.3% of vandetanib treated patients compared to 17.2% of placebo-treated patients [14]. In our cohort of pediatric MTC patients, patients who were rendered hypothyroid after thyroidectomy developed increased TSH and reduced free T4 during the first few months of treatment with vandetanib, necessitating an increase in LT dosage. Over the duration of this ongoing study, we continue to adjust LT dosing individualized to each patient as dosages of TKIs are altered, and patients grow and progress through puberty. Interestingly, while on vandetanib, 2 patients with normal pretreatment thyroid functions and intact glands continued to have TFTs within the normal range without requiring thyroid hormone treatment. Several studies report that TKIs either induce recurrence of hypothyroidism in hypothyroid patients, increasing their needs in LT replacement, or induce hypothyroidism in patients with previously normal thyroid function [14,18-20]. In a study looking at endocrine function in 35 adults with thyroid cancer on vandetanib, LT dose had to be increased for 26 patients, 5 with differentiated thyroid cancer and 21 with MTC [15]. Similar to our findings with children, none of the three adult nonthyroidectomized MTC patients needed LT treatment during the study according to normal serial TSH levels on vandetanib [15]. Elevated TSH in athyreotic patients may be due to an indirect effect of vandetanib on the metabolism of thyroid hormone, or with thyroid hormone action at the pituitary level. Impaired type-2 iodothyronine deiodinase and reduction of T3 generation from T4 during the first few months of treatment with vandetanib, necessitating an increase in LT dosage.

TKIs are altered, and patients grow and progress through puberty. Interestingly, while on vandetanib, 2 patients with normal pretreatment thyroid functions and intact glands continue to have TFTs within the normal range without requiring thyroid hormone treatment. Several studies report that TKIs either induce recurrence of hypothyroidism in hypothyroid patients, increasing their needs in LT replacement, or induce hypothyroidism in patients with previously normal thyroid function [14,18-20]. In a study looking at endocrine function in 35 adults with thyroid cancer on vandetanib, LT dose had to be increased for 26 patients, 5 with differentiated thyroid cancer and 21 with MTC [15]. Similar to our findings with children, none of the three adult nonthyroidectomized MTC patients needed LT treatment during the study according to normal serial TSH levels on vandetanib [15]. Elevated TSH in athyreotic patients may be due to an indirect effect of vandetanib on the metabolism of thyroid hormone, or with thyroid hormone action at the pituitary level. Impaired type-2 iodothyronine deiodinase and reduction of T3 generation from T4 during the first few months of treatment with vandetanib, necessitating an increase in LT dosage.

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T4, however, in hypothyroid patients relying on LT substitution alone, increased TSH results.

Diarrhea (the primary dose limiting toxicity in the pediatric clinical trial) may have contributed to decreased thyroid hormone absorption [12,20]. At baseline nine children had diarrhea while 80% had grade 1–2 diarrhea [12] during the initial 2 cycles and fewer children presented diarrhea in advanced stages of the study. Patients with MEN2B may have intestinal ganglioneuromatosis, which may play a role in the altered thyroid hormone absorption. One proxy to screen for malabsorption is linear growth, however the overall improvement in linear growth while on vandetanib suggests that malabsorption of levothyroxine is not the cause of lowered TSH levels.

Conclusions
Clinicians need to be aware of the potential downside of both over and undertreatment with thyroid hormone to ensure clinical stability. Compared to previously reported clinical trial in adults in which increased dosing of LT was required in 49.3% of athyreotic patients on vandetanib, 91% of athyreotic children in the present study required increased dosing of LT in order to maintain normal TSH values. Proper recognition and management of abnormal thyroid hormone levels is critical in growing children on TKIs.

Abbreviations
TKIs: Tyrosine kinase inhibitors; LT: Levothyroxine; MTC: Medullary thyroid carcinoma; MEN 2B: Multiple Endocrine Neoplasia Type 2B; TTFs: Thyroid function tests; RET: Rearranged during Transfection; CEA: Carcinoembryonic antigen; FMTC: Familial MTC.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
ML: Conceived of the study and drafted the manuscript. AG and EB: Performed data collection and analysis, and helped to draft the manuscript. NS: Participated in the design of the study and performed the statistical analysis. EP, MC, LM, and SA: Participated in the design of the study and provided ongoing oncological care to the patients, and directed the clinical trial. FB, BW and CS conceived of the study, and participated in its design and helped to draft the manuscript. All authors read and approved the final manuscript.

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