Phase II trial of lapatinib in adult and pediatric patients with neurofibromatosis type 2 and progressive vestibular schwannomas

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This single-institution phase II study was performed to estimate the response rate to lapatinib in neurofibromatosis type 2 (NF2) patients with progressive vestibular schwannoma (VS). Twenty-one eligible patients were enrolled. Brain and spine MRIs, including 3-dimensional volumetric tumor analysis, and audiograms were performed once at baseline and again every 12 weeks. The primary response end point was evaluable in 17 patients and defined as ≥15% decrease in VS volume. Hearing was evaluable as a secondary end point in 13 patients, with responses defined as an improvement in the pure tone average of at least 10 dB or a statistically significant increase in word recognition scores. Four of 17 evaluable patients experienced an objective volumetric response (23.5%; 95% confidence interval [CI], 10%–47%), with median time to response of 4.5 months (range, 3–12). In responders, reduction in VS volumes ranged from 15.7% to 23.9%. Four of 13 patients evaluable for hearing met hearing criteria for response (30.8%; 95% CI, 13%–58%). One sustained response exceeded 9 months in duration. Median time to overall progression (ie, volumetric progression or hearing loss) was 14 months. The estimated overall progression-free survival and volumetric progression-free survival at 12 months were 64.2% (95% CI, 36.9%–82.1%) and 70.6% (95% CI, 43.1%–86.6%), respectively. Toxicity was generally minor, and no permanent dose modifications were required. Lapatinib carries minor toxicity and has objective activity in NF2 patients with progressive VS, including volumetric and hearing responses. Future studies could explore combination therapy with other molecular targeted agents such as bevacizumab.

Keywords: acoustic neuroma, lapatinib, phase II trial, neurofibromatosis type 2, vestibular schwannoma.
first completed prospective phase II clinical trial evaluating an antitumor drug specifically in NF2 patients.

**Patients and Methods**

**Patient Eligibility and Enrollment**

Adult and pediatric patients (age ≥ 3 years) with a clinical diagnosis of NF2 according to the revised NIH criteria\(^9\) and at least one progressive VS (tumor growth or hearing progression within the past 12 months) were eligible. Histological confirmation was not required, because tumor biopsies are rarely indicated in this disease. For eligibility, tumor growth was defined as increase in tumor size of at least 2 mm in greatest diameter on conventional MRI\(^10\) or an increase in tumor volume of ≥ 15% as measured by 3-dimensional (3D) volumetric analysis. Progressive hearing progression was defined as a drop in pure tone average (PTA) of ≥ 10 dB at ≥ 2 consecutive or nonconsecutive frequencies or a drop in word recognition score (WRS) of ≥ 12% compared with the prior audiogram. Key eligibility criteria also included volumetrically measurable VS on MRI with size ≥ 0.5 cc, Karnofsky/Lansky performance score ≥ 50%, normal cardiac left ventricular ejection fraction by transthoracic echocardiogram, and adequate bone marrow, renal, and hepatic function. Exclusion criteria included any surgery within 4 weeks prior to enrollment, prior therapy with agents targeting EGFR or ErbB2, known preexisting cardiac disease, and concurrent therapy with cytochrome P450 inducers or inhibitors.

The study was conducted under a protocol approved by the institutional review board of NYU Langone Medical Center and registered at ClinicalTrials.gov (NCT00973739). Informed consent was obtained from the patients and guardians in accordance with institutional policies. All consecutive patients who met study entry criteria and who consented to participate were enrolled.

**Study Design**

This study was a prospective, open label, 2-stage phase II study. The primary and secondary end points were volumetric response and hearing response, respectively. To test the null hypothesis that the response rate is < 5% versus the alternative that the response rate is ≥ 25%, a 2-stage Simon design was used.\(^11\) Nine patients were to be enrolled in stage 1. If at least 1 patient of these 9 had a volumetric response in stage 1 at any given evaluation point, an additional 8 patients were to be enrolled in stage 2. The overall alpha level for this design was 0.05, with a power of 80%. Lapatinib was to be considered effective and of interest for further study if, after successful completion of both stages, there were at least 3 responses in the combined stages.

**Treatment**

Lapatinib was supplied by GlaxoSmithKline and administered in continuous 4-week courses. Pediatric patients < 18 years of age received 900 mg/m\(^2\) twice daily, up to a maximum dose of 750 mg twice daily, according to published phase I data.\(^12\) Patients ≥ 18 years of age received the standard recommended adult dose of 1500 mg once daily. For drug-induced diarrhea, a weight-based dose of loperamide was administered as needed. For treatment-related acneiform rash, clindamycin and benzoyl peroxide topical gel were prescribed as needed. Clinical evaluations, including a complete physical and neurological exam, complete blood count with differential, comprehensive metabolic panel, and serum pregnancy test (for females of child-bearing potential), were performed at baseline and every 4 weeks thereafter. To monitor for potential cardiotoxicity, echocardiograms and electrocardiograms were obtained at baseline and every 3 courses thereafter. Patients were allowed to remain on study unless volumetric progression, objective hearing deterioration, or unacceptable toxicity occurred. Adverse events were graded using version 3.0 of the NCI Common Toxicity Criteria (CTCAE). For treatment interruptions due to adverse events, therapy had to be held until toxicity was sufficiently improved, to grade ≤ 2 or ≤ baseline.

**Response Evaluation**

Baseline MRIs were required within 30 days and baseline audiograms within 14 days prior to starting lapatinib. A target tumor was defined as any volumetrically measurable VS. Volumetric response (primary end point) and hearing response (secondary end point) were assessed at the end of every third 4-week course and compared with baseline. On-study imaging consisted of MRIs of the brain and entire spine, and 3D tumor volumetrics were obtained on postcontrast, T1-weighted magnetization-prepared rapid acquisition with gradient echo sequences, a 1-mm slice thickness, and no gap, using semi-automated segmentation software (Vitrea platform). As volumetric measurements are superior to traditional tumor measurements in regard to sensitivity, reliability, and reproducibility,\(^13\) volumetrics have become the modality of choice for defining and assessing imaging response in NF2 clinical trials.\(^14,15\) We defined volumetric response or progression as a ≥ 15% decrease or increase, respectively, in VS volume compared with baseline. The ≥ 15% threshold was determined after intraobserver variability was found to be negligible compared with the 15% change of interest for tumor volumes > 0.5 cc in a pilot study (data not shown). Interobserver variability was also found to be low but was eliminated by assigning each patient to a specific radiology technician for volumetric measurement. Other NF2-related tumors, such as additional cranial nerve schwannomas and intracranial and intraspinal meningiomas and ependymomas, were also monitored radiologically and clinically.
Serial audiological evaluations were used to assess hearing response, including determination of pure tone thresholds and WRS. WRS was tested using the 50-item recorded CID (Central Institute for the Deaf)-W22 monosyllable word list. Hearing response or progression was defined as a clinically significant increase or decrease, respectively, in the WRS. WRS represents the most clinically relevant objective measure of hearing quality in NF2 patients and has therefore been suggested as a trial end point. PTAs were calculated by the mean of the individual threshold frequencies at 500, 1000, 2000, and 4000 Hz and was recorded for each ear. An increase of ≥10 dB in the PTA between any follow-up assessment and the baseline value was considered hearing deterioration, while an improvement of ≥10 dB indicated a clinically significant improvement, as previously suggested.

Pharmacokinetics
Blood samples for measurement of lapatinib plasma concentration were collected from pediatric patients immediately prior to a scheduled lapatinib dose, as well as at 2, 4, and 6 hours (optional) after, as previously described.

Statistical Analysis
Progression-free survival (PFS) was measured from date of enrollment to date of volumetric or hearing progression. PFS was analyzed using the Kaplan–Meier method in terms of overall PFS (volumetric or hearing progression), volumetric progression, and hearing progression. Point estimates for PFS with 95% confidence intervals (CIs) were calculated from Kaplan–Meier curves.

Results
Patients
Twenty-one patients were enrolled between October 2009 and March 2011. There were 13 males (61.9%) and 8 females (38.1%), and participants were a median age of 28 years at enrollment (range, 10–51), including 4 pediatric patients <18 years. Three patients (patients 2, 3, and 5) had familial NF2; the remainder were sporadic NF2 patients. All patients or their legal representatives provided written informed consent for treatment. Four patients were nonevaluable (NE). Two patients were nonevaluable due to coming off study prior to the first scheduled response evaluation: one was diagnosed with sarcoidosis (patient NE1) and one withdrew from the study for personal reasons (patient NE2). One patient (NE3) elected to remain on study medication despite being found ineligible due to small tumor size <0.5 cc. Stage 2 of the study was opened for enrollment after the first response was observed on stage 1. Accrual to this study was closed after the planned enrollment of 17 evaluable subjects was reached, who had a total of 22 measurable (ie, target) VS tumors. Characteristics of all evaluable patients are summarized in Table 1.

Treatment
The 17 evaluable patients received a total of 190 four-week courses of lapatinib. The median number of courses received was 12 (range, 3–21). Five patients did not complete the planned 12 months of therapy, all due to volumetric progression. One of these patients (patient 4) came off study for a tumor volume increase of >15%, although this initial measurement was revised down to +10.4% after a secondary review performed on all patients prior to publication disclosed a technical error in the volumetric measurement for this patient only. He therefore would have been eligible to continue on study in retrospect but remained in the progressive disease category for study analysis. Two patients progressed after receiving the 12 months of therapy. The estimated volumetric PFS at 12 months was 70.6% (95% CI, 43.1%–86.6%). Of note, 2 patients on lapatinib (patients 6 and 10) with stable VS and hearing suffered from progressively growing meningioma requiring surgical resection. Both patients continued on study with a brief interruption of lapatinib perisurgically.

The initial study design limited lapatinib to 12 cycles, but the protocol was subsequently amended to allow further treatment for patients with evidence of continued clinical benefit. Seven patients chose to continue on lapatinib beyond the 12th course, but all were eventually taken off study due to either MRI progression (n = 1), hearing deterioration (n = 1), increased size of the tumor’s cystic component (n = 1), prolonged adverse event (delayed postsurgical wound healing; n = 1), or patient and/or family preference (n = 3).

Toxicity
All 21 enrolled participants were available for toxicity monitoring. Observed toxicity was generally minor (CTCAE 3.0 grades 1 and 2) and most commonly included rash (53%) and less commonly diarrhea, fatigue, nail changes, headache, and elevation of alanine aminotransferase and aspartate aminotransferase. A single patient (4.8%) experienced a grade 3 toxicity (ie, delayed wound healing possibly related to lapatinib after surgery for progressive meningioma). No cardiotoxicity and no grade 4 toxicity were observed.

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Volumetric and Hearing Responses

Baseline patient characteristics and responses to treatment are summarized in Tables 1 and 2, respectively, with additional details included in the online supplementary Table S1. Considering all target tumors, best volumetric change ranged from −23.9% (shrinkage) to +17.57% (progression) compared with the baseline measurement (see Fig. 3). We observed a volumetric response in 4 subjects (patients 6, 9, 10, and 13), with a median time to response of 4.5 courses (range, 3–12). The total volumetric response rate in evaluable patients was therefore 23.5% (95% CI, 10.0%–47.4%). In responders, reduction in VS volumes ranged from −15.74% to −23.9%. In addition, one subject (patient 9) had a significant volume reduction in both target VS tumors. Three of the responders had follow-up neuroimaging studies that showed a small increase in their tumor volumes, although none reached baseline tumor size.

The distribution of the PTA and WRS for each ear is shown in the online supplementary Table S2. For all ears, the mean baseline PTA and WRS were 66.93 dB (SD ± 36.41) and 36.14% (SD ± 46.12), respectively. Four patients were deaf bilaterally at study enrollment and were therefore not evaluable for hearing response. Four patients (patients 3, 5, 9, and 11) experienced an improvement in their WRS sufficient to meet the definition of a clinical response, as established a priori. The responses were observed after a median of 3 courses (range, 3–9). Considering 4 responders of 13 evaluable patients, excluding deaf patients, the hearing response rate was 30.8% (95% CI, 12.7%–57.6%). Improvement in WRS was sustained in patient 3 only and exceeded 9 months in duration. The serial audiological measurements for each individual patient are available in the online supplementary Table S3. Of note, a combined volumetric and WRS hearing response of a tumor was observed in patient 9 only.

Regarding PTA, no patient had an improvement of ≥10 dB, indicative of a response. However, 2 patients reached the threshold of ≥10 dB deterioration in their PTAs, patient 3 (who experienced a concomitant deterioration in WRS) and patient 6 (who later suffered from a drop in WRS); the data are summarized in the online supplementary Table S4.

Progression-free Survival and Median Time to Progression

Seven patients experienced volumetric progression, and 2 patients experienced hearing progression. At 12 months, estimated PFS was 70.6% (95% CI, 43.1%–86.6%) for volumetrics, 88.9% (95% CI, 43.3%–98.4%) for hearing, and 64.2% (95% CI, 36.9%–82.1%) for overall progression (ie, volumetric or hearing progression). The median time to progression (hearing or volumetric) was 14 months (see Fig. 1).

Pharmacokinetics

Informed consent/assent was required for pharmacokinetic blood sampling and was provided by 2 of 3 evaluable pediatric patients. The maximum plasma concentrations observed were 3.24 and 3.36 μg/mL (see Fig. 2), similar to prior pharmacokinetic data from a pediatric phase I trial, which showed a median peak plasma concentration at steady state of 6.2 μg/mL (range, 3.1–10.3) at the same dose level used in our study (ie, 900 mg/m² twice daily).12
Table 2. Summary of evaluable patient treatment outcomes on study

<table>
<thead>
<tr>
<th>Patient</th>
<th>Treatment Duration (courses)</th>
<th>Best Reduction in Target Left/Right Tumor Volume (% baseline volume)</th>
<th>Progression of Target Left/Right Tumor Volume (% baseline volume)</th>
<th>Best Change in WRS in Target Left/Right ear (absolute score in %; difference from baseline)</th>
<th>Best Change in PTA (dB) for Target Tumor in Left/Right (absolute PTA; difference from baseline)</th>
<th>Reason for Treatment Discontinuation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>21</td>
<td>−13.73</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Patient preference</td>
</tr>
<tr>
<td>2</td>
<td>3</td>
<td>No reduction</td>
<td>+17.57</td>
<td>0 (−4)</td>
<td>56.25 (+10.00)</td>
<td>R VS growth</td>
</tr>
<tr>
<td>3</td>
<td>12</td>
<td>−3.90/−11.48                                                      +20.78/+19.67</td>
<td>100/NA (+16)</td>
<td>28.75/NA (0)</td>
<td>R and L VS growth</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>NA</td>
<td>+10.41</td>
<td>NA</td>
<td>NA</td>
<td>R VS growth</td>
</tr>
<tr>
<td>5</td>
<td>15</td>
<td>−6.25</td>
<td>NA</td>
<td>44 (+40)</td>
<td>51.25 (−3.75)</td>
<td>Patient preference</td>
</tr>
<tr>
<td>6</td>
<td>21</td>
<td>−16.57</td>
<td>NA</td>
<td>80 (+8)</td>
<td>48.75 (+10.00)</td>
<td>Progressive hearing loss</td>
</tr>
<tr>
<td>7</td>
<td>6</td>
<td>No reduction</td>
<td>+10.89/+16.89</td>
<td>NA/NA</td>
<td>NA/NA</td>
<td>R VS growth</td>
</tr>
<tr>
<td>8</td>
<td>6</td>
<td>No reduction</td>
<td>+21.14</td>
<td>100 (+4)</td>
<td>21.25 (−5.00)</td>
<td>L VS growth</td>
</tr>
<tr>
<td>9</td>
<td>12</td>
<td>−23.90/−20.25</td>
<td>NA/NA</td>
<td>NA/100 (+8)</td>
<td>NA/23.75 (−3.75)</td>
<td>Growth of R VS cystic component</td>
</tr>
<tr>
<td>10</td>
<td>15</td>
<td>−19.79</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Delay in wound healing</td>
</tr>
<tr>
<td>11</td>
<td>9</td>
<td>−3.33</td>
<td>NA</td>
<td>96 (+12)</td>
<td>31.25 (−2.50)</td>
<td>Progressive hearing loss</td>
</tr>
<tr>
<td>12</td>
<td>12</td>
<td>−14.15</td>
<td>NA</td>
<td>100 (0)</td>
<td>3.75 (+1.25)</td>
<td>Completed 12 cycles</td>
</tr>
<tr>
<td>13</td>
<td>12</td>
<td>−15.74</td>
<td>NA</td>
<td>100 (0)</td>
<td>10.00 (−2.50)</td>
<td>Completed 12 cycles</td>
</tr>
<tr>
<td>14</td>
<td>16</td>
<td>−7.31</td>
<td>NA</td>
<td>100 (+4)</td>
<td>21.25 (0)</td>
<td>Patient preference</td>
</tr>
<tr>
<td>15</td>
<td>12</td>
<td>−1.77</td>
<td>+22.12</td>
<td>NA</td>
<td>NA</td>
<td>R VS growth</td>
</tr>
<tr>
<td>16</td>
<td>3</td>
<td>−3.64/NA</td>
<td>NA/+18.64</td>
<td>32/NA (−4)</td>
<td>63.75/NA (−2.50)</td>
<td>R VS growth</td>
</tr>
<tr>
<td>17</td>
<td>12</td>
<td>−5.73/−3.91</td>
<td>NA/NA</td>
<td>100/96 (0/+4)</td>
<td>10.00/40.00 (−1.25/−7.50)</td>
<td>Completed 12 cycles</td>
</tr>
</tbody>
</table>

Boldface values signify that the value met the clinical definition for response.
Abbreviations: R, right; L, left; VS, vestibular schwannoma; WRS, word recognition score; PTA, pure tone average; NA, not applicable.
Discussion

NF2 is an autosomal-dominant genetic disease with an incidence of approximately 1 per 30,000. The majority of NF2 patients develop progressive hearing loss in adolescence or young adulthood due to unilateral or bilateral VS. Additional intracranial and spinal tumors, including meningiomas, schwannomas, and ependymomas, are also highly prevalent. NF2-related tumors, although mostly slow growing, cause considerable morbidity and mortality, particularly when diagnosed at a young age, and result in a significantly reduced life expectancy. The available treatment options for these neoplasms, which often occur at multiple sites simultaneously, are noncurative and mostly limited to surgery and/or radiation therapy. Surgical resection of VS often leads to complete deafness and facial nerve injury, while radiation therapy may increase the risk for developing secondary tumors. Therefore, effective medical treatment options are urgently needed for this disease.

Although chemotherapy is effective for low-grade brain tumors, such as astrocytomas, no such therapy has been validated in NF2. However, many conventional chemotherapeutic agents are unsuitable for NF2 patients due to neuro- and/or ototoxicity, and clinicians are reluctant to use mutagenic chemotherapy in patients with loss of tumor suppressor function, such as NF2.
patients. Presently, no known effective chemotherapy option exists for the treatment of schwannomas. However, recent retrospective studies have shown that bevacizumab, a monoclonal anti–vascular endothelial growth factor (VEGF) antibody, may result in tumor shrinkage and hearing improvement in patients with NF2 and VS,\textsuperscript{21,22} although “rebound” tumor growth has been observed after discontinuation of the drug.\textsuperscript{23} Based on encouraging preclinical data\textsuperscript{6} and a favorable safety profile, we conducted a phase II clinical trial using lapatinib in NF2 patients with progressive VS.

Lapatinib was generally well tolerated in this patient population, with toxicities that were usually mild and manageable.

Our volumetric and audiological response rates of 23.5% and 30.8%, respectively, appear superior to a recently published retrospective series of 10 patients treated with erlotinib, an EGFR inhibitor.\textsuperscript{24} In that series, no objective volumetric responses in the primary target tumors were observed, although 3 patients experienced stable disease volumetrically, with maximum tumor shrinkage of −14% in 1 patient. Prolonged stable disease was observed in 3 patients, and only 1 transient hearing response by WRS was observed. The median time to progression for either tumor growth or hearing loss in our study was 14 months, which is preferred over the 7.1 months in the erlotinib series.

Suggested response criteria for NF2 clinical trials\textsuperscript{14} published after the conception of this study have proposed to define a response as a reduction of ≥20% in tumor volume and to consider a reduction between 5% and <20% as a minor response. According to these response definitions, we would have obtained major and minor response rates of 5.9% (1 of 17 evaluable patients) and 32.9% (9 of 17 evaluable patients), respectively.

We did not observe any imaging responses in meningiomas, and these tumors continued to progress in many of our patients during the study period. These observations are consistent with recently published results of a phase II trial using the EGFR inhibitors erlotinib and gefitinib in patients with progressive meningiomas, where no response was seen.\textsuperscript{25}

The only other antitumor drug reported in the literature leading to tumor shrinkage and hearing improvement in NF2 patients is bevacizumab. The first and, to date, largest retrospective series of NF2 patients treated with bevacizumab showed a reduction in VS size in 9 of 10 patients (90%), with a range of −5% to −44% in tumor volume.\textsuperscript{21} Applying our volumetric response criterion of ≥15% in volume reduction, 7 of 10 patients (70%) were responders in that study, which appears superior to the 23.5% observed in our prospective trial. In the bevacizumab series, 4 of 7 patients (57.1%) had significant hearing improvement, which also appears superior to our results with lapatinib.

One possible limitation of lapatinib’s activity in our study might be suboptimal drug delivery to tumor tissue. Although responses to lapatinib have been observed in some HER2-positive breast cancer patients with brain metastases,\textsuperscript{26,27} it is not known whether VS is protected by the blood–brain barrier, the blood–nerve barrier, or the blood–cerebrospinal fluid barrier. The achievable tissue concentration of lapatinib in VS may be lower than that in tumors outside the CNS, limiting efficacy. Clinical data on brain metastasis show tissue lapatinib concentrations averaging approximately 7-fold higher than plasma concentrations.\textsuperscript{28} To determine the achievable intratumoral lapatinib concentration in VS and the effects on EGFR/ErbB2 signaling, we are currently conducting a lapatinib “phase 0” or pharmacodynamic trial in patients undergoing VS surgery (ClinicalTrials.gov identifier NCT00863122).\textsuperscript{15}

In summary, our study indicates that lapatinib is well tolerated and promotes antitumor activity, including hearing responses, in a subset of NF2 patients with progressive VS. Our study results serve as a valuable benchmark for comparison with future efficacy trials in this patient population. Further studies will be needed to better define the role of lapatinib in the treatment of VS in NF2 patients, and possible combination therapies with other molecular targeted agents should be explored. In a variety of preclinical models, the EGFR/ErbB2 signaling pathway and VEGF-dependent angiogenesis are functionally linked, and VEGF may play a role in the acquired resistance to ErbB receptor antagonists.\textsuperscript{29} Combination therapy with bevacizumab and lapatinib showed activity in a recent phase II study for HER2-overexpressing metastatic breast cancer in a heavily pretreated patient population\textsuperscript{30} and should be investigated in NF2 patients as well.

**Supplementary Material**

Supplementary material is available online at Neuro-Oncology (http://neuro-oncology.oxfordjournals.org).

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**Conflict of interest statement.** M. A. K. received funding for this study, as well as for a separate pharmacokinetic/pharmacodynamic study with lapatinib from GlaxoSmithKline under institutional clinical trial agreements. K. M. K. is an employee of GlaxoSmithKline and has stock ownership in the company. All other authors declare that they have no relevant conflicts of interest.

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