Stereotactic Ablative Radiotherapy for the Management of Liver Metastases from Neuroendocrine Neoplasms: A Preliminary Study

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Abstract
Introduction: Liver metastases are common in patients with neuroendocrine neoplasms. The role of stereotactic ablative radiotherapy (SABR) is not well understood in this population. Objective: The objective of this study was to evaluate the safety and efficacy of SABR in treating well-differentiated neuroendocrine liver metastases (WD-NELM). Methods: A retrospective review of patients with WD-NELM treated with SABR was conducted between January 2015 and July 2019. Demographic, treatment, and clinical/radiographic follow-up data were abstracted. RECIST 1.1 criteria were applied to each individual target to evaluate the response to treatment. Local control (LC) and progression-free survival (PFS) were determined using the Kaplan-Meier methodology. Toxicity was reported according to the CTCAE v5.0. Results: Twenty-five patients with a total of 53 liver metastases treated with SABR were identified. Most patients (68%) had midgut tumors, were grade 2 (80%), and had high-volume intrahepatic and/or extrahepatic disease (76%). The median number of liver metastases treated was 2, with a median size of 2.5 cm. The median radiation dose delivered was 50 Gy/5 fractions. The median follow-up was 14 months; 24 of the 25 patients were alive at the time of analysis. The objective response rate was 32%, with improvement or stability in 96% of lesions treated. The median time to best response was 9 months. The 1-year LC and PFS were 92 and 44%, respectively. No grade 3/4 acute or late toxicity was identified. Conclusions: Liver SABR is a safe and promising means of providing LC for WD-NELM. This treatment modality should be evaluated in selected patients in concert with strategies to manage systemic disease.

Introduction

Neuroendocrine neoplasms (NENs) are diverse malignancies that vary in terms of histology and clinical aggressiveness. The 2017 World Health Organization (WHO) system classifies these tumors into grades based on the differentiation status and mitotic/Ki-67 proliferation indices [1]. Low-grade tumors may remain indolent for years, whereas poorly differentiated grade III neuroendo-
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the management of unresectable colorectal liver metasta-
ses and hepatocellular carcinoma [15]. NENs have his-
torically been considered radioresistant due to modest
responses seen with conventionally delivered radiation
treatment [16]. Current data evaluating the impact of
modern stereotactic radiation therapy for neuroen-
docrine liver metastases (NELM) are limited to case reports.
The current study outlines the experience from 2015 to
2019 at the Susan Leslie Clinic for Neuroendocrine Tu-
mors at the Odette Cancer Centre, Sunnybrook Health
Sciences Centre, in the treatment of liver metastases from
well-differentiated grade 1 or 2 NENs with SABR.

Materials and Methods

An institutional REB-approved retrospective chart review of
patients with well-differentiated NELM treated with SABR was per-
formed between January 2015 and July 2019. Demographics, treat-
ment details, and clinical/radiographic follow-up data were ab-
stracted from the patients’ clinical and radiation planning records.
Each patient was assessed in the Susan Leslie Clinic for Neuro-
endocrine Tumors, a multidisciplinary clinic dedicated to the
management of neuroendocrine tumors. In this clinic, all patients
are concomitantly assessed by physicians specialized in NET from
surgical oncology, medical oncology, radiation oncology, and en-
docrinology and when appropriate further presented at tumor
board with radiology, interventional radiology, and pathology
[17]. All patients had progressed on SSA therapy and/or everoli-
mus and were deemed not a candidate for surgical resection or
other localized therapy (based upon extent of disease, size/location
disease, progression post-local therapy, patient comorbidities,
or patient preference). Patients with ≤5 systemic metastases, as
defined by multiphasic CT and liver MRI, were classified as being
oligometastatic; those with more advanced systemic disease but
with ≤4 NELM progressing on therapy were classified as being oli-
gopressive. Representative images demonstrating this classifi-
cation are shown in Figure 1.

Patients received radiotherapy according to institutional guide-
lines using abdominal compression or active breathing control for
liver motion management and image-guided, intensity-modulated
or volumetric-modulated arc therapy. Radiation was prescribed to
ensure coverage by >95% of the prescription dose to the planning
target volume, respecting dose constraints to normal liver paren-
chyma and adjacent organs at risk. Representative treatment dose
distribution is shown in Figure 2. All patients were premedicated
with a prophylactic 5-HT3 receptor antagonist for radiation-in-
duced nausea prevention prior to each treatment delivery. Patients
were evaluated clinically during radiation therapy, at 28 days post-
therapy and then every 3 months following completion of radia-
tion therapy. All patients underwent multiphasic CT imaging on
follow-up every 3 months following treatment and evaluated with
the same criteria as RECIST 1.1, but applied to each individual tar-
get treated. Pseudoprogression was defined as initial increase in
size of lesion in the absence of clinical symptoms with subsequent
decrease in size on serial imaging [18].

Descriptive statistics were performed, and variables were pre-
sented categorically as proportions or continuously using medians.
One-year LC and systemic control were calculated. Toxicity was evaluated retrospectively and reported as per the CTCAE v5.0 criteria. R Studio version 1.1.463 was used for all statistical analyses.

Results

Twenty-five patients with a total of 53 individual liver metastases treated with SABR were identified. The median follow-up was 14 months (range 2–54 months). Twenty-four of the 25 patients were still alive at the time of data analysis. Patient demographics are reported in Table 1. Most patients (68%) had midgut tumors (small bowel and pancreas), were grade 2 (80%), and were treated for oligoprogressive disease (76%). All patients had progressed on SSA, of which 2 had also progressed on everolimus. Fifteen patients (64%) had prior local therapy to the liver: surgery (9), liver-directed therapy (6), or both (2). Almost all patients (96%) remained on SSA despite having radiographic disease progression.

Radiation treatment details are outlined in Table 2. The median number of liver metastases treated per patient was 2 (range: 1–4), with a median size of 2.5 cm (range 0.7–9.7 cm). The median radiation dose delivered was 50 Gy/5 fractions (range 25–60 Gy/3 fractions), with a median biologically effective dose (BED10) of 100 (range 39–180).

Of the 53 treated metastases, 3 patients with 6 total metastases were defined as stable disease due to radiation treatment effects that either obscured accurate size definition or led to pseudoprogression (defined as an initial increase in size of the lesion but subsequent decrease on serial imaging). The evaluated overall response was complete response = 2, partial response = 13, stable disease = 30, and progressive disease = 2. The best treatment response according to changes in axial diameter was −100% (median −16%, range −100 to 47%), graphically represented in Figure 3. The objective response rate was 32%, seen in 12 patients. Initial disease improvement or stability was seen in 96% of all lesions treated. The median time to best response was 9 months (range 3–16 months). LC at 1 year was 92% (CI 0.847–0.999%). Systemic progression-free survival (PFS) at 1 year was 44% (CI 0.28–
0.70%). Local progression was only observed in patients with grade 2 histology.

Table 3 outlines identified acute and late toxicities. Three patients (12%) experienced acute grade 2 fatigue, and 4% experienced grade 2 nausea. One patient (4%) continued to have late grade 2 fatigue. No patients experienced any grade 3 or 4 acute or late toxicity.

**Discussion/Conclusion**

The potentially indolent course of well-differentiated NEN necessitates a treatment that maximizes disease control, maintains quality of life, and minimizes acute and chronic side effects of therapy. As there are no large comparative clinical trials that have evaluated systemic and liver-directed therapies, treatment selection is best determined by a multidisciplinary team that weighs the benefits and toxicity of each therapy in the context of the patient’s goals, the burden of functional symptoms, and the pattern of metastatic disease [19].

Surgical resection is a first-line option for functional tumors to reduce hormonal burden and should also be considered for cytoreduction of low-volume disease in properly selected patients [8, 19, 20]. However, patient or technical factors must be considered, given an acute surgical morbidity rate of 18% [8]. TAE is an alternative treatment for diffuse disease and/or symptomatic hormone production that has early tumor response and control rates of 50–66% and ∼80%, respectively. It is associated with a serious acute adverse event rate upward of 8% and grade 3 toxicity ranging from 0 to 25% [21–23]. Similarly, treatment with selective internal radiation therapy (SIRT) is also effective with a reported tumor response of ∼55% and control at 85–90%, with acute side effects seen in 30% of patients and grade 3 toxicity in 10% of patients [24, 25]. RFA is another well-tolerated liver-directed therapy effective in decreasing hormonal burden with reported tumor control rates of 60–80% and a modest complication rate of ∼10% [9, 26]. Peptide receptor radionuclide therapy may also be utilized in patients who have progressed on SSA [12]; however, it remains unclear whether patients with liver-only or low-volume oligometastatic dis-

| Variable                  | N = 25 |
|---------------------------|--------|
| Sex                       |        |
| Female                    | 14     |
| Male                      | 11     |
| Median age (range)        | 70 (46–88) |
| Histology                 |        |
| Small bowel               | 8      |
| Duodenum                  | 2      |
| Pancreas                  | 7      |
| Colorectal                | 4      |
| Lung                      | 3      |
| Renal                     | 1      |
| Grade                     |        |
| 1                         | 5      |
| 2*                        | 20     |
| Metastases treated per patient, n | |
| 1                         | 10     |
| 2                         | 6      |
| 3                         | 5      |
| 4                         | 4      |
| All sites of metastatic disease treated | |
| Yes (oligometastatic disease) | 6     |
| No (oligoprogressive disease) | 19   |
| Concurrent SSA use        |        |
| Yes                       | 24     |
| No                        | 1      |
| Prior liver surgery       |        |
| Yes                       | 9      |
| No                        | 16     |
| Prior liver-directed therapy |        |
| Yes                       | 6      |
| No                        | 19     |

SSA, somatostatin analogs. * Atypical lung carcinoid classified as grade 2.

### Table 2. Liver metastasis treatment characteristics

| Variable                                      | Total liver metastases treated (n = 53) |
|-----------------------------------------------|----------------------------------------|
| Median size of metastases (range)             | 2.5 cm (0.7–9.7 cm)                    |
| Size grouping of metastases, cm               |                                        |
| >1.0                                          | 3                                      |
| 1.0–1.9                                       | 14                                     |
| 2–2.9                                         | 19                                     |
| 3–3.9                                         | 5                                      |
| >4                                            | 12                                     |
| Median biological equivalent dose (range)     | 100 Gy<sub>10</sub> (39–180)           |
| Radiation dose/fractionation                  |                                        |
| 30 Gy/5                                       | 2                                      |
| 35 Gy/5                                       | 3                                      |
| 40 Gy/5                                       | 10                                     |
| 45 Gy/5                                       | 6                                      |
| 50 Gy/5                                       | 20                                     |
| 60 Gy/5                                       | 3                                      |
| 60 Gy/3                                       | 2                                      |
| Other                                         | 7                                      |
ease would benefit from a more targeted ablative approach.

The ENETS Consensus Guidelines for managing liver metastases recommend ablation (RFA, TAE, or SIRT) for low-volume liver metastases [19]. Historically, radiation has not been considered, likely due to initial experience using outdated techniques and dose/fractionation tailored for widespread metastatic disease [27]. Contemporary studies utilizing modern radiation techniques have demonstrated radiographic and clinical response to primary and metastatic disease [16, 28, 29]. Radiobiological models utilize the alpha/beta ratio to determine the response of tumors to varied dose/fraction radiation schemes; most rapidly growing histologies will have an alpha/beta value of ~10, while slower growing tumors will have a lower value and thus be more responsive to high-dose per fraction treatment [30]. Given the overall low proliferative rate of neuroendocrine tumors, compared to conventional radiation treatment, the use of SABR is predicted to be more effective. This premise is strengthened by a multigene expression index for radiosensitivity correlated with response to SABR for liver metastases [31], in which small bowel neuroendocrine malignancies were predicted to be highly radiosensitive, whereas pancreatic and large bowel

**Table 3. Radiation-related toxicity**

| Acute toxicity   | Incidence (n = 25) | Late toxicity (n = 25) | Incidence (n = 25) |
|------------------|--------------------|------------------------|--------------------|
| **Fatigue**      |                    |                        |                    |
| Grade 0          | 18 (72%)           | Grade 0                | 23 (92%)           |
| Grade 1          | 4 (16%)            | Grade 1                | 1 (4%)             |
| Grade 2          | 3 (12%)            | Grade 2                | 1 (4%)             |
| Grade 3+         | 0 (0%)             | Grade 3+               | 0 (0%)             |
| **Nausea**       |                    |                        |                    |
| Grade 0          | 17 (68%)           | Grade 0                | 24 (96%)           |
| Grade 1          | 7 (28%)            | Grade 1                | 1 (4%)             |
| Grade 2          | 1 (4%)             | Grade 2                | 0 (0%)             |
| Grade 3+         | 0 (0%)             | Grade 3+               | 0 (0%)             |
| **Abdominal pain** |                  |                        |                    |
| Grade 0          | 23 (92%)           | Grade 0                | 25 (100%)          |
| Grade 1          | 2 (8%)             | Grade 1                | 0 (0%)             |
| Grade 2          | 0 (0%)             | Grade 2                | 0 (0%)             |
| Grade 3+         | 0 (0%)             | Grade 3+               | 0 (0%)             |
| **Other**        |                    |                        |                    |
| Grade 0          | 24 (96%)           | Grade 0                | 25 (100%)          |
| Grade 1          | 1 (4%)             | Grade 1                | 0 (0%)             |
| Grade 2          | 0 (0%)             | Grade 2                | 0 (0%)             |
| Grade 3+         | 0 (0%)             | Grade 3+               | 0 (0%)             |
emerging LC strategy. Specific to liver metastases, an abstract by Bignardi et al. [33] described 2 patients with liver metastases from nonfunctional low-grade small bowel and pancreatic neuroendocrine tumors treated to 45–60 Gy, with reported LC greater than 40 months. A recent report by Ohri et al. [15] evaluating liver SABR for metastases of all histologies demonstrated improved control rates with BED_{10} ≥100 Gy. There was a nonsignificant trend toward improved LC with higher BED radiation in our series (data not shown), supporting that BED_{10} ≥100 Gy should be considered when treating NELM.

The general slow growth rate of neuroendocrine disease must be considered when evaluating response on serial scans. In this series, patients were progressing on SSA therapy and the majority of patients had grade 2 disease, suggesting that observed LC is reflective of treatment effect as opposed to indolent behavior of disease. Our data demonstrating an objective response rate of 32% and 1-year LC rate of 92% suggest that SABR is an effective treatment option to maximize LC with inherent patient advantages: minimally invasive, done as an outpatient, flexible with respect to tumor location, favorable side effect profile, and in the context of larger metastases, potentially superior and more cost-effective than RFA [34, 35]. Patients may have a prolonged disease course with NET; it is important to further develop the armamentarium for management of metastatic disease. Surgery should be the treatment of choice when it is appropriate and recommended. Given the excellent results of local therapy provided by RFA, TAE, and SIRT, future evaluation needs to be done to assess where SABR may be applied as an emerging LC strategy.

One important finding is the median time to maximal response of 9 months. This highlights a limitation of response evaluation criteria that rely on tumor dimensions, particularly when applied to therapies whose activity is not immediately reflected by change in tumor size. Traditional RECIST criteria have been criticized for their application in post-SABR response assessment [17]. The ablative nature of SABR therapy and its effect on the tumor microvasculature provide an opportunity to develop alternative response evaluation criteria sensitive to the tumor microenvironment. Dynamic contrast-enhanced and other functional imaging techniques are promising modalities under investigation for early response monitoring in a variety of solid tumor sites [36, 37]; we are prospectively evaluating alternative imaging response criteria in a subset of the patients currently being reported.

The role of SABR within the oligometastatic paradigm is intriguing. Emerging data in other histologies suggest that PFS and OS can be improved by treating all sites of metastatic disease with SABR in concert with standard of care systemic therapy [38, 39]. Our results in liver metastases and recent case studies in primary pulmonary carcinoids [40, 41] demonstrate that SABR is an effective treatment modality for neuroendocrine cancers. Given the tolerability of SABR and SSA therapies, it can be hypothesized that clinical outcomes may be improved in patients with a low metastatic disease burden by combining SSAs and upfront SABR to all sites of disease, potentially delaying the transition to more toxic second-line therapy.

Limitations of this study include retrospective data collection, small sample size, and that it pre-dates the availability of Gallium-68 PET to evaluate the extent of disease. Given the increased sensitivity/specificity of Ga-68 PET over CT and MRI [42, 43], there may have been occult disease within the liver that was not treated with ablative therapy, which may have impacted the overall PFS rates. However, this would be true for any local therapeutic option. Any future studies evaluating the role of SABR in the upfront setting would require Ga-68 PET to establish true oligometastatic disease presentations. However, given the nature of the Susan Leslie Clinic for Neuroendocrine Tumors at the Odette Cancer Centre, there are significant strengths: this is the largest series to date specific to treatment of liver metastases; all patients were reviewed by multidisciplinary team before being offered SABR, ensuring appropriateness of patient selection; all treatment plans were generated using a standardized departmental guidelines and underwent expert peer QA to ensure high-quality therapy; and all patients had clinical and radiographic follow-up at 1 treatment center to ensure consistent image acquisition/interpretation and patient evaluation.

This is the first series to specifically evaluate SABR as a treatment modality for well-differentiated NELM. SABR is a safe and promising treatment option for patients with limited volume disease; it may complement or provide an additional strategy for NELM alongside more established liver-directed therapies such as TAE, RFA, or SIRT. Further prospective studies are required to determine the optimal treatment modality and sequencing for patients with low-volume metastatic disease.
Statement of Ethics

This retrospective study was approved and overseen by the Sunnybrook Research Institute Research Ethics Board (REB #:192-2019).

Conflict of Interest Statement

J.M.H. has no conflicts of interest to declare. H.T.-K.C. has no conflicts of interest to declare. W.C. has no conflicts of interest to declare. A.T. has no conflicts of interest to declare. L.E.D. has no conflicts of interest to declare. W.C. has no conflicts of interest to declare. All authors provided substantial contributions. J.H.: analysis/interpretation of data, manuscript creation, and final approval. L.D.: analysis/interpretation of data, manuscript creation, and final approval. H.T.-K.C.: analysis/interpretation of data, manuscript creation, and final approval. S.M.: conception of work, data acquisition/interpretation, manuscript creation, and final approval.

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