

# [The potential of heavy-ion therapy to improve outcomes for locally advanced non-s...](https://assignbuster.com/the-potential-of-heavy-ion-therapy-to-improve-outcomes-for-locally-advanced-non-small-cell-lung-cancer/)

[Health & Medicine](https://assignbuster.com/essay-subjects/health-n-medicine/)

The treatment of unresectable locally advanced non-small cell lung cancer (LA-NSCLC) remains a daunting challenge. To date, the best median survival achieved in a randomized prospective bi-modality clinical trial for LA-NSCLC is 28. 7 months for patients who received standard chemoradiotherapy on NRG Oncology RTOG 0617 ( [1](#B1) ). Emerging data from RTOG 0617 and other institutions have also implicated radiation dose to the heart as a driver of cardiovascular events, survival, and patient-reported quality of life ( [2](#B2) – [5](#B5) ). To improve survival and quality of life for LA-NSCLC, it is critical to explore emerging therapeutic technologies. One such technology is radiation therapy delivered with heavy ions, such as carbon ions. Heavy-ion therapy has both unique biological and physical advantages that may improve local control while also reducing radiation exposure to non-target organs at risk such as the heart. With the technology implemented at several Asian and European centers in conjunction with the planned development of several therapeutic heavy-ion centers in the United States, there will be real opportunity to exploit this technology to gain ground and improve the therapeutic ratio in LA-NSCLC.

LA-NSCLC is highly resistant to conventionally fractionated radiation therapy with cooperative group studies showing locoregional failure rates greater than 60% after definitive chemoradiation ( [6](#B6) ). From a radiation biology standpoint, there is considerable rationale to support use of heavy-ion therapy to improve local control for these patients. Upon encountering target tissue, interaction of charged heavy ions with matter yields the highest linear energy transfer (LET) of any currently available form of clinical radiation ( [7](#B7) ). This in turn results in unique clustered DNA lesions, resulting in a lower oxygen enhancement ratio and higher relative biological effectiveness (RBE) that is on the order of threefold greater than photon radiotherapy ( [7](#B7) ). Early preclinical data have also suggested tumors with EGFR mutation may be more susceptible to heavy-ion therapy than photon irradiation, suggesting that heavy-ion therapy may provide opportunities to further tailor radiation therapy in the heterogeneous genetic landscape of LA-NSCLC ( [8](#B8) ). Taken together, these properties suggest that heavy ions could improve local control by overcoming DNA repair pathways that confer radiation resistance and provide patient-tailored options for LA-NSCLC.

The physical properties of heavy-ion therapy also provide dosimetric advantages for LA-NSCLC ( [9](#B9) ). With particle therapy, the energy deposited in tissue increases with depth eventually coming to an abrupt stop, known as the Bragg peak. Using particles, practitioners have the ability to stop the dose deposition at a specified point, reducing or eliminating exposure to tissues distal to the target volumes. These advantages are particularly relevant to recent findings in RTOG 0617 where the importance of radiation conformity to prevent pneumonitis and reduce cardiac doses has been established ( [10](#B10) , [11](#B11) ). Just as proton therapy can reduce cardiac doses in comparison to photon therapy ( [12](#B12) ), heavy ions also have the potential to further reduce cardiac and pulmonary doses ( [9](#B9) , [13](#B13) ). First, the Bragg Peak can reduce radiation doses distal to the target to a far greater degree than photon irradiation. Second, heavy ions exhibit less scattering than protons because of their mass, resulting in a sharper lateral penumbra than proton or photon radiotherapy. Third, pencil beam scanning and arc technologies are expected to provide unprecedented geometric avoidance of non-target organs at risk. The combination of these physical advantages with motion management and Monte Carlo algorithms for plan optimization has potential to produce dramatic improvements in the conformity of the high, intermediate, and low dose regions. While proton therapy exhibits some of these characteristics such as the Bragg Peak, proton therapy has more lateral scattering and lacks the LET of heavy-ion therapy.

In the era of anti-PD-1-/PDL-1-targeted therapies, another important biological advantage of heavy-ion therapy is its potential immunostimulatory effects. While radiation therapy is known to have complex reactions with the immune system and tumor microenvironment, heavy ions have may have unique immune effects that are distinct from photons or proton therapy. Because of their high LET and RBE, preclinical evidence suggests that heavy ions induce non-apoptotic cell death that is independent of the typical p53, bax/bcl, and p21 signal transduction cascades ( [14](#B14) , [15](#B15) ). These alternative forms of cell death could provide more diverse tumor epitopes for cytotoxic T-cells to prime immune responses. Another immunological advantage of heavy-ion therapy stems from the unique physical properties discussed earlier. Reducing low and intermediate dose exposure has the potential to reduce integral dose that can cause lymphopenia and hematologic toxicity ( [16](#B16) – [18](#B18) ). In RTOG 0617, Grade 3+ hematological toxicity was observed in 56% of patients with only one-third of patients completing consolidative chemotherapy ( [1](#B1) ), highlighting the need to reduce the myelosuppressive effects of definitive chemoradiotherapy. Thus, by reducing lymphopenia from exposure of circulating lymphocytes and also stimulating the local production of immunological epitopes, heavy-ion therapy might be a potent weapon to illicit *in situ* vaccine responses.

Conventionally fractionated radiation therapy as a single modality has dismal cure rates ( [19](#B19) ), and the primary benefit of concurrent chemotherapy in historic combined modality trials has been to improve local control ( [20](#B20) – [22](#B22) ), which comes at the cost of myelosuppression, esophagitis, and peripheral neuropathy in LA-NSCLC. However, heavy-ion therapy can deliver doses to tumor targets with greater potency than concurrent chemoradiation without the systemic side effects of chemotherapy. This raises the question of whether heavy-ion therapy can obviate the need for concurrent cytotoxic therapy. Concurrent chemotherapy has significant downsides including high rates of severe esophagitis and immunosuppression. A Phase I–II of carbon ion therapy for patients with LA-NSCLC who were medically unfit to receive concurrent chemotherapy from the Research Institute for Charged Particle Therapy in Chiba, Japan showed promising results ( [23](#B23) ). In this study, dose was escalated from 68 to 72 cobalt Gy equivalents without dose limiting toxicity. Oncological outcomes were also favorable with a 2-year local control rate of 93. 1% and overall survival of 51. 9%. There has also been success using carbon ion therapy without cytotoxic chemotherapy for medically inoperable early-stage NSCLC ( [13](#B13) , [24](#B24) – [26](#B26) ). Exploration of heavy-ion technology as a strategy to avoid concurrent cytotoxic chemotherapy should be encouraged as a way to reduce toxicities and health-care costs for patients without compromising oncological efficacy.

Currently used fractionation schedules for carbon therapy differ greatly from standard photon techniques. Given differential biological effects on tumors normal tissues as a function of dose per fraction, hypofractionated approaches are commonplace in the practice of heavy-ion therapy. The use of hypofractionated treatment schedules may allow for a greater number of patients to be treated at select centers at a reduced cost. Coupled with the potential for reduced toxicity and improved outcomes, this could make heavy-ion therapy a cost effective treatment, despite the high upfront costs of building such facilities likely on the order of $200–300 million (USD).

As the number of heavy-ion therapy centers is expected to increase in the United States in the coming years, there will be opportunity to explore its role for LA-NSCLC. We have outlined rationale for robust exploration of heavy-ion therapy in LA-NSCLC for the purpose of improving what are presently suboptimal outcomes in this population.

## Author Contributions

All authors have contributed to conceptualization and writing of this manuscript.

## Conflict of Interest Statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

## References

1. Bradley JD, Paulus R, Komaki R, Masters G, Blumenschein G, Schild S, et al. Standard-dose versus high-dose conformal radiotherapy with concurrent and consolidation carboplatin plus paclitaxel with or without cetuximab for patients with stage IIIA or IIIB non-small-cell lung cancer (RTOG 0617): a randomised, two-by-two factorial phase 3 study. *Lancet Oncol* (2015) 16(2): 187–99. doi: 10. 1016/S1470-2045(14)71207-0

2. Movsas B, Hu C, Sloan J, Bradley J, Komaki R, Masters G, et al. Quality of life analysis of a radiation dose-escalation study of patients with non-small-cell lung cancer: a secondary analysis of the radiation therapy oncology group 0617 randomized clinical trial. *JAMA Oncol* (2016) 2(3): 359–67. doi: 10. 1001/jamaoncol. 2015. 3969

3. Guberina M, Eberhardt W, Stuschke M, Gauler T, Heinzelmann F, Cheufou D, et al. Heart dose exposure as prognostic marker after radiotherapy for resectable stage IIIA/B non-small-cell lung cancer: secondary analysis of a randomized trial. *Ann Oncol* (2017) 28(5): 1084–9. doi: 10. 1093/annonc/mdx069

4. Wang K, Eblan MJ, Deal AM, Lipner M, Zagar TM, Wang Y, et al. Cardiac toxicity after radiotherapy for stage III non-small-cell lung cancer: pooled analysis of dose-escalation trials delivering 70 to 90 Gy. *J Clin Oncol* (2017) 35(13): 1387–94. doi: 10. 1200/JCO. 2016. 70. 0229

5. Dess RT, Sun Y, Matuszak MM, Sun G, Soni PD, Bazzi L, et al. Cardiac events after radiation therapy: combined analysis of prospective multicenter trials for locally advanced non-small-cell lung cancer. *J Clin Oncol* (2017) 35(13): 1395–402. doi: 10. 1200/JCO. 2016. 71. 6142

6. Machtay M, Paulus R, Moughan J, Komaki R, Bradley JE, Choy H, et al. Defining local-regional control and its importance in locally advanced non-small cell lung carcinoma. *J Thorac Oncol* (2012) 7(4): 716–22. doi: 10. 1097/JTO. 0b013e3182429682

7. Durante M, Loeffler JS. Charged particles in radiation oncology. *Nat Rev Clin Oncol* (2010) 7(1): 37–43. doi: 10. 1038/nrclinonc. 2009. 183

8. Amornwichet N, Oike T, Shibata A, Nirodi CS, Ogiwara H, Makino H, et al. The EGFR mutation status affects the relative biological effectiveness of carbon-ion beams in non-small cell lung carcinoma cells. *Sci Rep* (2015) 5: 11305. doi: 10. 1038/srep11305

9. Kubo N, Saitoh JI, Shimada H, Shirai K, Kawamura H, Ohno T, et al. Dosimetric comparison of carbon ion and X-ray radiotherapy for stage IIIA non-small cell lung cancer. *J Radiat Res* (2016) 57(5): 548–54. doi: 10. 1093/jrr/rrw041

10. Chun SG, Hu C, Choy H, Komaki RU, Timmerman RD, Schild SE, et al. Impact of intensity-modulated radiation therapy technique for locally advanced non-small-cell lung cancer: a secondary analysis of the NRG oncology RTOG 0617 randomized clinical trial. *J Clin Oncol* (2017) 35(1): 56–62. doi: 10. 1200/JCO. 2016. 69. 1378

11. Chun SG, Hu C, Bradley JD. Reply to D. Ball et al. *J Clin Oncol* (2017) 35(13): 1493–4. doi: 10. 1200/JCO. 2016. 71. 5755

12. Tucker SL, Liu A, Gomez D, Tang LL, Allen P, Yang J, et al. Impact of heart and lung dose on early survival in patients with non-small cell lung cancer treated with chemoradiation. *Radiother Oncol* (2016) 119(3): 495–500. doi: 10. 1016/j. radonc. 2016. 04. 025

13. Shirai K, Kawashima M, Saitoh JI, Abe T, Fukata K, Shigeta Y, et al. Clinical outcomes using carbon-ion radiotherapy and dose-volume histogram comparison between carbon-ion radiotherapy and photon therapy for T2b-4N0M0 non-small cell lung cancer-A pilot study. *PLoS One* (2017) 12(4): e0175589. doi: 10. 1371/journal. pone. 0175589

14. Durante M, Brenner DJ, Formenti SC. Does heavy ion therapy work through the immune system? *Int J Radiat Onco Biol Phys* (2016) 96(5): 934–6. doi: 10. 1016/j. ijrobp. 2016. 08. 037

15. Loeffler JS, Durante M. Charged particle therapy – optimization, challenges and future directions. *Nat Rev Clin Oncol* (2013) 10(7): 411–24. doi: 10. 1038/nrclinonc. 2013. 79

16. Yovino S, Kleinberg L, Grossman SA, Narayanan M, Ford E. The etiology of treatment-related lymphopenia in patients with malignant gliomas: modeling radiation dose to circulating lymphocytes explains clinical observations and suggests methods of modifying the impact of radiation on immune cells. *Cancer Invest* (2013) 31(2): 140–4. doi: 10. 3109/07357907. 2012. 762780

17. Pignalosa D, Lee R, Hartel C, Sommer S, Nikoghosyan A, Debus J, et al. Chromosome inversions in lymphocytes of prostate cancer patients treated with X-rays and carbon ions. *Radiother Oncol* (2013) 109(2): 256–61. doi: 10. 1016/j. radonc. 2013. 09. 021

18. Tang C, Liao Z, Gomez D, Levy L, Zhuang Y, Gebremichael RA, et al. Lymphopenia association with gross tumor volume and lung V5 and its effects on non-small cell lung cancer patient outcomes. *Int J Radiat Oncol Biol Phys* (2014) 89(5): 1084–91. doi: 10. 1016/j. ijrobp. 2014. 04. 025

19. Perez CA, Pajak TF, Rubin P, Simpson JR, Mohiuddin M, Brady LW, et al. Long-term observations of the patterns of failure in patients with unresectable non-oat cell carcinoma of the lung treated with definitive radiotherapy. Report by the Radiation Therapy Oncology Group. *Cancer* (1987) 59(11): 1874–81. doi: 10. 1002/1097-0142(19870601)59: 11 <1874:: AID-CNCR2820591106> 3. 0. CO; 2-Z

20. Curran WJ Jr, Paulus R, Langer CJ, Komaki R, Lee JS, Hauser S, et al. Sequential vs. concurrent chemoradiation for stage III non-small cell lung cancer: randomized phase III trial RTOG 9410. *J Natl Cancer Inst* (2011) 103(19): 1452–60. doi: 10. 1093/jnci/djr325

21. Belani CP, Choy H, Bonomi P, Scott C, Travis P, Haluschak J, et al. Combined chemoradiotherapy regimens of paclitaxel and carboplatin for locally advanced non-small-cell lung cancer: a randomized phase II locally advanced multi-modality protocol. *J Clin Oncol* (2005) 23(25): 5883–91. doi: 10. 1200/JCO. 2005. 55. 405

22. Furuse K, Fukuoka M, Kawahara M, Nishikawa H, Takada Y, Kudoh S, et al. Phase III study of concurrent versus sequential thoracic radiotherapy in combination with mitomycin, vindesine, and cisplatin in unresectable stage III non-small-cell lung cancer. *J Clin Oncol* (1999) 17(9): 2692–9. doi: 10. 1200/JCO. 1999. 17. 9. 2692

23. Takahashi W, Nakajima M, Yamamoto N, Yamashita H, Nakagawa K, Miyamoto T, et al. A prospective nonrandomized phase I/II study of carbon ion radiotherapy in a favorable subset of locally advanced non-small cell lung cancer (NSCLC). *Cancer* (2015) 121(8): 1321–7. doi: 10. 1002/cncr. 29195

24. Miyamoto T, Yamamoto N, Nishimura H, Koto M, Tsujii H, Mizoe JE, et al. Carbon ion radiotherapy for stage I non-small cell lung cancer. *Radiother Oncol* (2003) 66(2): 127–40. doi: 10. 1016/S0167-8140(02)00367-5

25. Miyamoto T, Baba M, Sugane T, Nakajima M, Yashiro T, Kagei K, et al. Carbon ion radiotherapy for stage I non-small cell lung cancer using a regimen of four fractions during 1 week. *J Thorac Oncol* (2007) 2(10): 916–26. doi: 10. 1097/JTO. 0b013e3181560a68

26. Iwata H, Demizu Y, Fujii O, Terashima K, Mima M, Niwa Y, et al. Long-term outcome of proton therapy and carbon-ion therapy for large (T2a-T2bN0M0) non-small-cell lung cancer. *J Thorac Oncol* (2013) 8(6): 726–35. doi: 10. 1097/JTO. 0b013e318288ab02