Relationship between electron density and effective densities of body tissues for stopping, scattering, and nuclear interactions of proton and ion beams

Nobuyuki Kanematsu† Taku Inaniwa, and Yusuke Koba
Department of Accelerator and Medical Physics, Research Center for Charged Particle Therapy, National Institute of Radiological Sciences, 4-9-1 Anagawa, Inage-ku, Chiba 263-8555, Japan
(Dated: January 11, 2012)

Purpose: In treatment planning of charged-particle radiotherapy, patient heterogeneity is conventionally modeled as variable-density water converted from CT images to best reproduce the stopping power, which may lead to inaccuracies in the handling of multiple scattering and nuclear interactions. Although similar conversions can be defined for these individual interactions, they would be valid only for specific CT systems and would require additional tasks for clinical application. This study aims to improve the practicality of the interaction-specific heterogeneity correction.

Methods: We calculated the electron densities and effective densities for stopping power, multiple scattering, and nuclear interactions of protons and ions, using the standard elemental-composition data for body tissues to construct the invariant conversion functions. We also simulated a proton beam in a lung-like geometry and a carbon-ion beam in a prostate-like geometry to demonstrate the procedure and the effects of the interaction-specific heterogeneity correction.

Results: Strong correlations were observed between the electron density and the respective effective densities, with which we formulated polyline conversion functions. Their effects amounted to 10% differences in multiple-scattering angle and nuclear-interaction mean free path for bones compared to those in the conventional heterogeneity correction. Although their realistic effect on patient dose distributions would be generally small, it could be at the level of a few percent when a carbon-ion beam traverses a large bone.

Conclusions: The present conversion functions are invariant and may be incorporated in treatment planning systems with a common function relating CT number to electron density. This will enable improved beam dose calculation while minimizing initial-setup and quality-management of the user’s specific system.

Keywords: proton beam therapy, ion beam therapy, inhomogeneity corrections, multiple scattering, nuclear interactions

I. INTRODUCTION

In the present practice of radiotherapy treatment planning, patient-specific material information is obtained from x-ray CT images. The CT number given to each image pixel represents the x-ray attenuation coefficient of the material of that pixel, from which the effective density for the treatment beam is estimated on the assumption of one-to-one correspondence. Taking water for a reference material, the effective density of a material is defined as the thickness ratio of water to the material for an equivalent dosimetric effect, which approximates to electron density for megavoltage photons or stopping-power ratio for charged particles.

The empirical relationship between CT number and effective density for a radiation of interest generally carries with it inaccuracies due to the limited tissue equivalency of sample materials. Schneider et al. developed the stoichiometric method to construct improved relations for body tissues. They modeled the physics of x-ray attenuation using measured CT numbers of well-defined materials, with which they calculated the x-ray attenuations in the standard body tissues of known elemental compositions, mass density, and electron density, in addition to their stopping-power ratios for protons, to construct the relations.

With such conversion, patients are modeled as variable-density water to assure range accuracy for charged-particle beams in conventional heterogeneity-correction algorithms. As the body tissues deviate from water in elemental compositions, these algorithms only approximately address the other beam interactions. Matsufuji et al. took the stoichiometric approach also for multiple scattering and nuclear interactions to study the validity of the conventional heterogeneity correction. They found that there would be 10%-level errors in multiple-scattering angle and nuclear-interaction mean free path of protons in bones although their effects on patient dose distributions would be small in realistic situations. Nevertheless, these definite errors may be corrected by the interaction-specific density conversions. In fact, Szymanowski and Oelfke proposed a specific CT-number conversion for multiple scattering. Palmans and Verhaegen proposed another CT-number conversion for proton nuclear interactions.

Those studies offer valid methods for improved heterogeneity correction, which however have to be applied to individual scanning conditions of individual CT systems. The resultant conversion tables must be separately set up and selectively used in treatment planning systems with periodical constancy tests or updates, which would greatly complicate the system-management tasks in the clinical environment. The purpose of this study is to establish a simpler procedure and to provide necessary data for equivalently accurate conversion. We take electron
density for a reference quantity and similarly construct conversions for the effective densities of body tissues as a function of electron density. As the conversion from CT number to electron density is commonly available in treatment planning systems, implementation of the invariant conversion functions in algorithms will offer accuracy improvement without further complicating data management.

II. MATERIALS AND METHODS

A. Electron density

We used 92 of the ICRU body tissues\(^2\) excluding obsolete, extreme, or artificially extracted materials such as ICRU-33 soft tissue, hydroxyapatite, calcifications, water, lipid, carbohydrate, cell nucleus, cholesterol, protein, and urinary stones. Electron density \(\rho_e\) is calculated as

\[
\rho_e = \rho_m \sum_i w_i \left( \frac{Z_i}{A_{ri}} \right)_w^{-1},
\]

where \(Z_i\) and \(A_{ri}\) are the atomic number and the atomic weight of element \(i\), \((Z/A)_w = 0.5551\) is the mean \(Z/A\) of water, and \(w_i\) and \(\rho_m\) are the elemental mass fraction and the mass density of the material. We represent \(\rho_m\), \(\rho_e\), and all other densities as non-dimensional ratios to those of water in this study.

B. Stopping power

The Bethe theory leads to the stopping-power ratio of a material to water\(^5\) or the stopping effective density, as

\[
\rho_S = \rho_e \left( \ln \frac{2v^2}{c^2 - v^2} - \ln \frac{I}{m_e c^2} \right)
\times \left( \ln \frac{2v^2}{c^2 - v^2} - \ln \frac{I_w}{m_e c^2} \right)^{-1},
\]

where \(m_e = 0.511\) MeV/c\(^2\) is the electron mass, \(v\) and \(c\) are the speeds of the projectile and light, and \(I\) and \(I_w\) are the mean excitation energies of the material and water. We calculated the \(I\) values of the body tissues with the Bragg rule

\[
\ln I = \sum_i w_i \frac{Z_i}{A_{ri}} \ln I_i \left( \sum_i w_i \frac{Z_i}{A_{ri}} \right)^{-1},
\]

where elemental \(I_i/\text{eV}\) values were H:19.2, C:81, N:82, O:106, F:112, Na:168, Mg:176, P:195, Cl:180, K:215, Ca:216, and Fe:323 for constituents of solid or liquid compounds.\(^5\) This leads to \(I_w = 75.3\) eV for water. Although recent studies suggest slightly higher \(I_w\) values,\(^2\) we used this \(I_w\) value in the denominator of Eq. (2) to cancel out the common systematic error of the compound \(I\) value in the numerator for general body tissues whose main ingredient is water. As the \(v\)-dependent variation of the \(\rho_S\) is within 1% under therapeutic conditions,\(^5\) we took the representative projectile speed \(v = 0.6\) c or the nucleon kinetic energy \(E/A = 230\) MeV to define the stopping effective density as projectile independent.

C. Multiple scattering

Scattering power \(T\) is the increase of Gaussian angular variance per transport distance\(^6\) In some formulations, it is in the form of

\[
T = \left( \frac{E_s}{p v} \right)^2 \frac{1}{X},
\]

where \(E_s = 15.0\) MeV is a constant energy, \(p\) is the particle momentum, and \(X\) is the scattering length of the material. Gottschalk derived the scattering length for heavy charged particles.\(^11\) For a composite material, it is given by

\[
\frac{1}{X} = \frac{\rho_m}{4298.4 \text{ cm}^2} \sum_i w_i \frac{Z_i^2}{A_{ri}} \left( 29.733 - \ln Z_i - \ln A_{ri} \right).
\]

This leads to \(X_w = 46.88\) cm for water. We define the scattering effective density as the scattering-power ratio of the material to water,

\[
\rho_T = \frac{T}{T_w} = \frac{X_w}{X}.
\]

D. Nuclear interactions

Silver et al. made an empirical modification to the geometric model for nucleus–nucleus collision cross section,\(^12\) resulting in

\[
\sigma_N = \pi r_0^2 \left[ A_{1/3}^{1/3} + A_{1/3}^{1/3} - b_0 \left( A_{1/3}^{1/3} + A_{1/3}^{1/3} \right) \right]^2,
\]

where we consistently use symbols \(A\) for mass number of the projectile and \(A_t\) for atomic weight of the target and introduced \(r_0 = 1.36\) fm for effective nucleon radius and \(b_0\) for overlap parameter. In their model, the energy dependence of the cross section would be almost common among target body-tissue materials in the therapeutic energy region and is thus ignored in this study.

For a compound material of mass density \(\rho_m\), the number of element-i nuclei per volume is proportional to \(\rho_m w_i / A_{ri}\). The total nuclear cross section per volume is thus proportional to \(\sum_i \sigma_N \rho_m w_i / A_{ri}\), which is proportional to the incidence of nuclear interactions. We define the nuclear effective density as the ratio of the incidence of nuclear interactions in the material to that in
water, resulting in
\[ \rho_N = \sum_i \frac{\sigma_{Ni}}{A_i} w_i \rho_m \left( \frac{\sigma_N}{A_r} \right)^{-1}, \]
\[ \left( \frac{\sigma_N}{A_r} \right)_w = \frac{2 \sigma_{NH} + \sigma_{NO}}{M_w}, \]
where \( \sigma_{NH} \) and \( \sigma_{NO} \) are the cross sections of hydrogen and oxygen nuclei and \( M_w = 18.015 \) is the molecular weight of water. Unlike the other effective densities, we formulated the nuclear effective density as projectile dependent.

E. Example of corrective conversions

To demonstrate the procedure and to evaluate the significance of the interaction-specific corrective conversion, we carried out simplistic proton and carbon-ion beam-transport calculations in one-dimensional geometries, each with and without the corrective conversions. We applied a proton beam of range 9.8 cm in water to a lung-like geometry consisting of staked slabs of 1-cm soft tissue, 2-cm bone, 1-cm soft tissue, 8-cm lung, and 3-cm tumor. We applied a carbon-ion beam of range 22.4 cm in water to a prostate-like geometry consisting of stacked slabs of 6-cm soft tissue, 6-cm bone, 4-cm soft tissue, and 4-cm tumor. We assigned electron densities 1.0 for soft tissue and tumor, 1.4 for bone, and 0.25 for lung and set the beam ranges to coincide with the tumor depths without the correction. These beams were point-like mono-directional at the phantom entrance and were transported through the slabs at 1-mm intervals with quantities of interest,
\[ d(x) = \int_0^x \rho(x') \, dx' \]
\[ \sigma_y(x) = \left( \int_0^x (x - x')^2 \rho(x') T_w(x') \, dx' \right)^{1/2}, \]
evaluated as a function of transport distance \( x \), where generic depth, beam size, and density, \( (d, \sigma_y, \rho) \), represent the effective depths and the beam sizes calculated with the respective effective densities. For the integration, the Fermi-Eyges theory in the numerical form \(^{14}\) was implemented with a semi-empirical scattering power for water, \( T_w \). We are interested in the beam size corrected for multiple scattering, \( \sigma_yT \), and the effective depth corrected for nuclear interactions, \( d_N \), to compare to the effective depth and beam size in the conventional heterogeneity correction, \( (d_S, \sigma_yS) \). The effective depths are geometric parameters that will not depend on beam properties except for the \( d_N \) that is slightly different between protons and ions. We specifically refer to the effective depths of the distal end of the target as the effective ranges.

TABLE I: Polyline conversion functions from electron density \( \rho_e \) to stopping effective density \( \rho_s \), scattering effective density \( \rho_T \), and nuclear effective density \( \rho_N \) for protons and ions of \( 4 \leq A \leq 16 \).

| \( \rho_e \)  | 0   | 0.9 | 0.9 | 1.035 | 1.4 | 2.0  |
|-------------|-----|-----|-----|-------|-----|------|
| \( \rho_s/\rho_e \) | 1.000 | 1.000 | 1.028 | 1.000 | 0.973 | 0.942 |
| \( \rho_T/\rho_e \) | 0.995 | 0.955 | 0.77 | 0.995 | 1.32 | 1.66 |
| \( \rho_N/\rho_e \) for protons | 0.992 | 0.992 | 1.07 | 0.992 | 0.89 | 0.78 |
| \( \rho_N/\rho_e \) for ions | 0.987 | 0.987 | 1.12 | 0.987 | 0.81 | 0.64 |

III. RESULTS

A. Corrective conversion functions

Figure \( \text{I} \) shows correspondences between the electron density and the effective densities of body tissues for proton and ion beams. There was a concentration of tissues around \( (\rho_e, \rho_s/\rho_e, \rho_T/\rho_e, \rho_N/\rho_e) = (1.035, 1.000, 0.995, 0.99) \). The low density \( \rho_e = 0.258 \) for the lung tissue was attributed to the air content. Generally in the \( \rho_e \) region, a weak negative correlation between \( \rho_s/\rho_e \) and \( \rho_e \) was observed. That was attributed to the low \( I \) values of carbon-rich adipose tissues in the low \( \rho_e \) region and the high \( I \) values of calcium-rich bone tissues in the high \( \rho_e \) region. Simplistically for a single-element material, Coulomb scattering depends on \( \rho_m Z^2/A_r \) whereas the electron density depends on \( \rho_m Z/A_r \). As there is approximate linear relationship among \( \rho_m, A_r, Z \), and \( \rho_e \) for common materials, the ratio \( \rho_T/\rho_e \propto Z \) showed a strong positive correlation with \( \rho_e \). Similarly, because the nuclear effective density depends approximately on \( \rho_m Z^2/A_r \), the ratio \( \rho_N/\rho_e \propto A_r^{2/3}/Z \) showed a negative correlation with \( \rho_e \). The \( \rho_N/\rho_e \) ratios for protons largely deviated from those for the ions due to the distinctive formulation of the overlap parameter \( b_0 \) in Eq. \( \text{I} \).

For the conversion functions, we set a discontinuity point at \( \rho_e = 0.9 \), where none of these tissues existed, an inflection point at the center of the tissue concentration, and another inflection point at \( \rho_e = 1.4 \) in the bone-tissue region. Similarly to the lung tissues, tissue-air mixing may occur on the surface of skin and epithelial tissues due to finite spatial resolution of the CT image, for which we assigned constant \( \rho/\rho_e \) ratios in the \( \rho_e < 0.9 \) region. Although material mixing may generally confuse any conversion schemes, its influence is limited as there normally are only a few tissue boundaries along the beam path. Because the variation of \( \rho_T/\rho_e \) among the ions was small, we defined one conversion function for the ions of \( 4 \leq A \leq 16 \). Table \( \text{I} \) shows the resultant conversion factors as polyline functions, which are also shown in Fig. \( \text{I} \).
FIG. 1: Ratios of (a) stopping effective density $\rho_s$, (b) scattering effective density $\rho_T$, (c) nuclear effective density $\rho_N$ for protons, and (d) nuclear effective density $\rho_N$ for helium ions ($\times$), carbon ions ($\triangle$), and oxygen ions (+) to electron density $\rho_e$ with the polyline conversion functions (solid lines).

B. Example of corrective conversions

Figures 2(a) and 2(c) show the development of the depth corrections for stopping and nuclear interactions, $(d_s - d_e)$ and $(d_N - d_e)$, for the proton beam in the lung-like geometry and for the carbon-ion beam in the prostate-like geometry. The corrections of the stopping effective depth, which would assure the range accuracy, were within about ±0.1 cm in both cases. The bone thickness mainly determined the corrections of the effective ranges. The nuclear effective range ($R_N = 21.6$ cm) deviated from the stopping effective range ($R_S = 22.3$ cm) by −0.7 cm in the prostate-like geometry.

Similarly, figures 2(b) and 2(d) show the development of the beam sizes. The uncorrected beam size $\sigma_{yS}$ and the conventional beam size $\sigma_{yE}$ were almost indistinguishable in these cases, while the corrected beam size $\sigma_{yT}$ deviated from them by about 5% for the proton beam in the lung-like geometry. The bone thickness to scatter and the distance to travel determined the amount of the correction.

IV. DISCUSSION

A distinctive feature of this study is the generality of the conversion functions. For example, Szymanowski and Oellke constructed a conversion function for the angular scaling factor or $\sqrt{\rho_T/\rho_S}$ in our notation. While their function was only valid for a specific CT system, the $\rho_e/\rho_T$ function in Table I is generally valid. As the biological data and the physical models are common or equivalent, the direct CT-number conversion and the corrective conversion through $\rho_e$ are essentially equivalent in accuracy, assuming that the corrective conversion is used with a stoichiometrically constructed CT-number–$\rho_e$ table.

Intrinsic error of CT number, such as that from x-ray hardening effect (typically 1% for soft tissues and 2% for bones) influences both conversion schemes in a similar way.

Matsufuji et al. found that, for a typical bone of $\rho_e \approx 1.35$, the errors of the conventional heterogeneity correction would be −10% in multiple-scattering angle and −10% in mean free path of proton–nuclear interaction. For a tissue of $\rho_e = 1.35$, the polyline functions in Table I give $\rho_T/\rho_e = 1.21$ and $\rho_N/\rho_e = 0.904$ for protons. Because the multiple-scattering angle is proportional to $\sqrt{\rho_T}$ and the mean free path is proportional to $1/\rho_N$, the $\rho_T$ and $\rho_N$ conversions would make +10% and +11% corrections to multiple-scattering angle and mean free path, respectively. The +10% and +11% corrections correspond to 9% and 10% effects in the corrected values, which agree with their original estimations. They also simulated realistic situations, in which bone was not dominant in the beam path, and concluded that the effects on patient dose distributions would be small, which we confirmed in our examples.

The nuclear interactions cause attenuation of primary...
particles. Lee et al. proposed an approximation formula for proton fluence \( \Phi(R) = \Phi(0) (1 + 0.012R/cm) \) as a linear function of residual range \( R \) in water\(^{12} \) to which the nuclear effective range \( R_N \) should be applied in the interaction-specific heterogeneity correction. For the proton-like geometry ignoring the small \( R_N \) difference between protons and ions, the error of the nuclear effective range with the conventional heterogeneity correction \( (\Delta R = R_S - R_N = 0.7 \text{ cm}) \) would cause a relative error of \(-0.7\%\) to the proton survival \( \Phi(0)/\Phi(R) \) and consequently to the dose to the tumor. Palmans and Verhaegen reported up to 0.5\% dosimetric effects for their examples that contained less bone than ours\(^2 \). For carbon-ion beams, the dosimetric effect would be a few or more times larger for their large nuclear cross section, i.e. a few percent, which could be clinically influential.

The fragmentation processes in ion beams are further complex. In the participant–spectator model, the target nucleus will not directly influence how the projectile nucleus may break up.\(^{14} \) There are some experimental data for the relative yields of projectile fragments,\(^{19} \) which may be thus reasonably invariant to target materials. Due to experimental difficulties in precise measurement of a treatment-beam spectrum, Monte Carlo simulation will be useful for building a detailed beam model in numerical or analytical form.\(^{20} \)

V. CONCLUSIONS

For general body tissues, we found strong correlations between electron density and the effective densities that characterize the strengths of stopping power, multiple scattering, and nuclear interactions of the projectile protons and ions. The stopping effective density deviated from the electron density by up to a few percent and the scattering and nuclear effective densities deviated by up to a few tens percent, which are consistent with other studies. To correct those errors, invariant corrective conversion functions from the electron density to the individual effective densities were constructed. Their effect on patient dose may be negligibly small in general, although it could be at the level of a few percent when an ion beam traverses a large bone.

The electron density should be derived from the CT number with a conversion table that has been constructed with the best accuracy, i.e. stoichiometrically, for treatment planning of proton and ion-beam therapy. The proposed extension of heterogeneity correction will enable improved beam dose calculation without seriously sacrificing simplicity or efficiency of the patient model and the algorithms, while quality-management tasks for CT and treatment panning systems will be kept as they are in the current clinical practice.

---

1 U. Schneider, E. Pedroni, and A. Lomax, “The calibration of CT Hounsfield units for radiotherapy treatment planning,” Phys. Med. Biol. 41 111–124 (1996).
2 ICRU, Tissue substitutes in radiation dosimetry and measurement, ICRU Report 44 (1989).
3 N. Matsufuji, H. Tomura, Y. Futami, H. Yamashita, A. Higashi, S. Minohara, M. Endo, and T. Kanai, “Relationship between CT number and electron density, scatter angle and nuclear reaction for hadron-therapy treatment planning,” Phys. Med. Biol. 43 3261–3275 (1998).
4 H. Szymanowski and U. Oelike, “CT calibration for two-dimensional scaling of proton pencil beams,” Phys. Med. Biol. 48 861–874 (2003).
5 H. Palmans and F. Verhaegen, “Assigning nonelastic nuclear interaction cross sections to Hounsfield units for Monte Carlo treatment planning of proton beams,” Phys. Med. Biol. 50 991–1000 (2005).
6 ICRU, Photon, electron, proton and neutron interaction data for body tissues, ICRU Report 46 (1992).
7 ICRU, Stopping powers and ranges for protons and alpha particles, ICRU Report 49 (1993).
8 ICRU, Stopping powers for electrons and positrons, ICRU Report 37 (1984).
9 P. Sigmund, A. Schinner, and H. Paul, “Errata and addenda for ICRU Report 73, stopping of ions heavier than helium, Journal of the ICRU vol. 5 no. 1 (2005),” Journal of the ICRU (2009).
10 N. Kanematsu, N. Matsufuji, R. Kohno, S. Minohara, and T. Kanai, “A CT calibration method based on the poly-binary tissue model for radiotherapy treatment planning,” Phys. Med. Biol. 48 1053–1064 (2003).
11 B. Gottschalk, “On the scattering power of radiotherapy protons,” Med. Phys. 37 352–367 (2010).
12 L. Sihver, C. H. Tsao, R. Silberberg, T. Kanai, and A. F. Barghouty, “Total reaction and partial cross section calculations in proton–nucleus \((Z_t \leq 26)\) and nucleus–nucleus reactions \((Z_p \leq Z_t \leq 6)\),” Phys. Rev. C 47 1225–1236 (1993).
13 L. Sihver and D. Mancusi, “Present status and validation of HIBRAC,” Radiation Measurements 44 38–46 (2009).
14 N. Kanematsu, “Alternative scattering power for Gaussian beam model of heavy charged particles,” Nucl. Instrum. Methods B 266 5056–5062 (2008).
15 N. Kanematsu, “Semi-empirical formulation of multiple scattering for the Gaussian beam model of heavy charged particles stopping in tissue-like matter,” Phys. Med. Biol. 54 N67–73 (2009).
B. Schaffner and E. Pedroni, “The precision of proton range calculations in proton radiotherapy treatment planning: experimental verification of the relation between CTHU and proton stopping power,” Phys. Med. Biol. 43 1579–92 (1998).

M. Lee, A. E. Nahum, and S. Webb, “An empirical method to build up a model of proton dose distribution for a radiotherapy treatment-planning package,” Phys. Med. Biol. 38 989–998 (1993).

G. Baur, F. Rössel, D. Trautmann, and R. Shyam, “Fragmentation processes in nuclear reactions,” Phys. Rep. 111 333–371 (1984).

N. Matsufuji, A. Fukumura, M. Komori, T. Kanai, and T. Kohno, “Influence of fragment reaction of relativistic heavy charged particles on heavy-ion radiotherapy,” Phys. Med. Biol. 48 1605–1623 (2003).

J. Kempe and A. Brahme, “Analytical theory for the fluence, planar fluence, energy fluence, planar energy fluence and absorbed dose of primary particles and their fragments in broad therapeutic light ion beams,” Physica Medica 26 6–16 (2010).