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Deep convolutional neural network with transfer learning for rectum toxicity prediction in cervical cancer radiotherapy: a feasibility study

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Abstract
Better understanding of the dose-toxicity relationship is critical for safe dose escalation to improve local control in late-stage cervical cancer radiotherapy. In this study, we introduced a convolutional neural network (CNN) model to analyze rectum dose distribution and predict rectum toxicity. Forty-two cervical cancer patients treated with combined external beam radiotherapy (EBRT) and brachytherapy (BT) were retrospectively collected, including twelve toxicity patients and thirty non-toxicity patients. We adopted a transfer learning strategy to overcome the limited patient data issue. A 16-layers CNN developed by the visual geometry group (VGG-16) of the University of Oxford was pre-trained on a large-scale natural image database, ImageNet, and fine-tuned with patient rectum surface dose maps (RSDMs), which were accumulated EBRT + BT doses on the unfolded rectum surface. We used the adaptive synthetic sampling approach and the data augmentation method to address the two challenges, data imbalance and data scarcity. The gradient-weighted class activation maps (Grad-CAM) were also generated to highlight the discriminative regions on the RSDM along with the prediction model. We compare different CNN coefficients fine-tuning strategies, and compare the predictive performance using the traditional dose volume parameters, e.g. $D_{0.1/1/2cc}$, and the texture features extracted from the RSDM. Satisfactory
1. Introduction

Cervical cancer is the third most common cause of female cancer mortality worldwide (Torre et al. 2015). Locally advanced cervical cancer, a common presentation, (Al-Mansour and Verschraegen 2010) is usually treated with external beam radiotherapy (EBRT) followed by brachytherapy (BT) with and without chemotherapy. Randomized studies have shown excellent treatment outcomes in early stage cervical cancer (Gray 2008, Haie-Meder et al. 2010). However, treatment results are inferior in advanced stages, with a five-year overall survival rate of 60–65% in stage IIB (Green et al. 2001), 25–50% in stage IIIB (Montana et al. 1986, Horiot et al. 1988), and 20–35% in stage IV disease (Rose et al. 2011).

Mounting clinical evidence correlates tumor control rate with dose. For instance, a retrospective study, RetroEMBRACE (Tanderup et al. 2016), initiated by the GEC-ESTRO group, has shown that high risk clinical target volume (CTVHR) dose $\geq 85$ Gy (EDQ210, D90) delivered in 7 weeks provides a 3-year local control rate $>94\%$ in limited-size ($20\, \text{cm}^3$), $>93\%$ in intermediate-size ($30\, \text{cm}^3$), and $>86\%$ in large-size CTVHR ($70\, \text{cm}^3$) tumors. It also concluded that increasing CTVHR volume by $10\, \text{cm}^3$ requires an additional 5 Gy for equivalent local control. However, a high dose may substantially increase toxicity risks to nearby organs at risk (OARs), such as the rectum, sigmoid, bladder, and vagina. Late rectum morbidity is associated with high rectum volume dose ($D_{2\, \text{cc}} > 75$ Gy) (Mazeron et al. 2016). Better understanding of the relationship between OAR toxicity and dose is critical for safe dose escalation to improve local control of large-size advanced-stage cervical cancer tumors.

A common limitation of current image-guided EBRT-BT techniques is that they utilize a static image during the computer tomography (CT) simulation scan and ignore anatomic variations throughout the treatment course. Even though such variations are captured by BT fractional CT scans, which are used for BT planning, the hotspots are assumed to be static. Relying on this static assumption, clinicians apply the worst-case addition method to evaluate OARs’ $D_{0.1\, \text{cc}}$, $D_{1\, \text{cc}}$, and $D_{2\, \text{cc}}$ (most exposed 0.1, 1, and $2\, \text{cm}^3$ volume) accumulative dose for toxicity prediction. Though the worst-case addition method protects OARs by overestimating OAR dose, it potentially prevents the delivery of possible high dose prescriptions to the target volume. Moreover, the reported dose volume values discard dose distribution information that may be crucial for adaptive BT planning, e.g. local characteristics of dose distribution on the rectum are related to rectum toxicity (Buettner et al. 2009, Lee et al. 2012, Wortel et al. 2015, Drean et al. 2016).

To study the correlation between received dose and induced OAR toxicity, three important problems must be addressed: (1) how to account for inter-fractional organ deformations and accurately acquire the accumulative dose received by the OARs; (2) how to utilize the OARs’ spatial dose distribution information instead of merely using one dimensional (1D)
dose volume parameters such as $D_{0.1/1/2cc}$; and (3) how to build an effective dose-toxicity prediction model with a limited patient sample size.

A typical cervical cancer radiotherapy treatment course is composed of ~25 fractions EBRT treatments and ~4–6 fractions BT treatments. The substantial inter-fractional organ motion that occurs in BT treatment makes reporting the accurate cumulative dose over an entire treatment course a challenging task. It was reported that the OARs intra- and inter-fraction $D_{2cc}$ uncertainties were 20–25% (including a small fraction of 5–11% contouring uncertainties), indicating that organ motion is the major contribution to OAR dose uncertainties (Dinkla et al. 2013, Lobefalo et al. 2013, Tanderup et al. 2013). Intensity-based approaches (e.g. Demons-based) (Thirion 1998, Christensen et al. 2001, Wang et al. 2005), may not provide an optimal method for deformable image registration (DIR) in the context of cervical radiotherapy, because of the potential for large deformations in the rectum, bladder, etc, as well as the poor contrast between these OARs and their surrounding tissues. In contrast, the feature-based DIR methods (e.g. finite element model-based) (Brock et al. 2005, Xiong et al. 2006, Kaus et al. 2007, Vasquez Osorio et al. 2009, Bondar et al. 2010, Andersen et al. 2012, Wognum et al. 2013), aided by organ contours or delineated features, are more favorable for accurate anatomical mapping, especially for hollow organs with substantial deformation, such as the rectum and bladder. Our recent work (Chen et al. 2016) proposed an improved non-rigid point matching algorithm based on the ‘thin plate splines-robust point matching’ (TPS-RPM) framework (Chui and Rangarajan 2003). This novel approach was validated on a porcine bladder phantom embedded with fiducial markers as baseline, and satisfactory DIR accuracies of $3.7 \pm 1.8 \text{ mm}$ and $1.6 \pm 0.8 \text{ mm}$ were achieved for bladders with large and small deformation. It can be served as a practical tool for accurate surface matching for hollow organs.

Several groups have also investigated spatial dose distribution and rectal toxicity in prostate cancer radiotherapy (Meijer et al. 1999, Heemsbergen et al. 2005, Tucker et al. 2006, Munbodh et al. 2008, Buettner et al. 2009, 2012, Wortel et al. 2015). Most of these reported studies utilized a three-dimensional (3D) to two-dimensional (2D) mapping approach to obtain a rectum surface dose map (RSDM) for spatial dose feature extraction. For example, Munbodh et al. (2008) sought to identify dosimetric and anatomic indicators of late rectal toxicity by using the RSDM in prostate cancer patients treated with intensity modulated radiation therapy (IMRT). Buettner et al. (2012) studied the dose response of the anal sphincter region from the constructed RSDM and correlated 3D dose distributions with various side effects. Wortel et al. (2015) found substantial relationships between acute rectal toxicity and local dose distributions through RSDM in prostate cancer patients who received IMRT and 3D-conformal radiotherapy (3D-CRT). Recently, an expert system based on machine learning technic was developed to address dosimetric uncertainties caused by motion of hollow organs (e.g. rectum, bladder) for prostate cancer radiation therapy (Guidi et al. 2017). These limited but innovative studies have demonstrated the great potential of employing RSDM for rectum dose-toxicity relationship analysis.

Although most current studies focus on statistical analysis of the underlying relationship between the OARs’ toxicity and the extracted 1D dose volume parameters (ICRU 2013), or 2D/3D localized dose distribution features (Wortel et al. 2015, Drean et al. 2016), our ultimate interest lies in the possibility of employing the dose distribution information for induced OAR toxicity prediction. The recent revival of deep learning techniques highlighted by the success of deep convolutional neural networks (CNN) (LeCun et al. 2015) has brought more alternatives and opportunities for dose-toxicity prediction modeling. The CNN first gained popularity in the computer vision community and is now thriving in the medical image processing domain (Tajbakhsh et al. 2016). However, training the CNN from scratch is difficult not only because of the extensive computational requirements for the training process, but also because
of the lack of large-scale annotated medical image training datasets and the potential risk of over-fitting. In contrast with training the CNN from scratch, transfer learning fine-tunes the CNN which is pre-trained on a large labeled dataset from a different application. Recently, promising results have been reported that transfer learning can have superior, or at least, the same performance as the CNN trained from scratch for medical image classification (Shin et al 2016, Tajbakhsh et al 2016).

In this study, we introduced a CNN model to analyze rectum dose distribution and predict rectum toxicity. We adopt a transfer learning strategy to overcome the limited patient data issue by pre-training a VGG-16 CNN on a large-scale natural image database, and then fine-tuned with the patient RSDMs. A gradient-weighted class activation method that highlights the discriminative regions on the RSDM was used to correlate dose distribution and rectum toxicity when the VGG-16 CNN completes a prediction. The general algorithm workflow is depicted in figure 1.

2. Methods and materials

2.1. Patient cohort

We retrospectively collected data of 42 cervical cancer patients treated with 25 fractions (2 Gy per fraction) EBRT followed by 4 (7 Gy per fraction) or 5 fractions (6 Gy per fraction) BT. The collected data include planning images and treatment plans. The patient was scheduled for follow-up examine every 2–3 months after treatment. Patients complaining of hematochezia were further examined by colonoscopy. Patients’ rectum toxicity Grades were collected as a clinical parameter in this study, where twelve with Grade $\geq 2$ rectum toxicity were characterized as toxicity patients and thirty patients with Grade 0–1 toxicity as non-toxicity patients. To account for biologic effects of different fractionation schemes, both the BT and EBRT physical doses were converted to EQD2 doses using a linear quadratic model (Bentzen et al 2012) with an $\alpha/\beta$ ratio of 3 for rectum (Michalski et al 2010, Moulton et al 2016).

2.2. Rectum surface meshing and DIR

A rectum surface mesh was generated using rectum contours in consecutive CT slices via a particle-based surface meshing approach (Zhong et al 2013). Given the initial rectum contour
points, a high-quality isotropic triangular surface meshing was obtained by solving an inter-
particle energy function with the quasi-Newton L-BFGS optimizer.

For each patient, the rectum surface meshes in each BT fraction were generated using the
physician delineated rectum contours via the above particle-based surface meshing approach.
The obtained rectum surface mesh in the first BT fraction was used as a reference, while
those rectum surface meshes from other fractions were registered to the reference domain
via a recently developed topography-preserved point-matching deformable image registration
(TOP-DIR) algorithm (Chen et al 2016). The TOP-DIR is a local topography-preserved robust
point-matching algorithm designed for accurate dose accumulation on a hollow organ wall.
TOP-DIR can find point-to-point correspondence on an organ wall by introducing local ana-
tomic information into the iterative update of correspondence matrix computation in the ‘thin
plate splines-robust point matching’ (TPS-RPM) scheme (Chui and Rangarajan 2003). Once
the deformation vector fields (DVFs) on the rectum surface were generated by TOP-DIR, final
DVFs defined on each voxel of the dose matrix were estimated by B-spline approximation
(Zhen et al 2015).

We used the final DVFs to deform and sum all the fractional BT doses to the reference
domain (the first BT fraction) to yield a deformable cumulative BT dose. Since patients in
the study cohort received a homogenous EBRT dose in the pelvic region, the EBRT dose was
added to the BT cumulative dose without deformation. The final accumulative EBRT
+ BT rectum surface dose was employed for the subsequent dose-toxicity prediction study.

2.3. Rectum unfolding

For each evaluated patient, we generated a 2D RSDM to represent dose distribution on the
rectum surface. We used a mapping procedure proposed by Tucker et al (2006) to unfold
the rectum surface onto a rectangle in a plane. Specifically, the rectum contour centroid on each
CT slice was calculated, and $n$ rays were emitted from the centroid at evenly spaced intervals.
We recorded the intersection coordinates between the rectal contour and the emanating rays
and extracted the dose on each intersection point of the rectum wall. We then cut the rectum
at the posterior-most position on all the $z$ CT slices and unfolded it to form a flat rectangular
$n \times z$ matrix representing dose distribution on the rectum surface. Rectum unfolding is illus-
trated in figure 3.

2.4. Convolutional neural network (CNN)

We employed the flattened 2D RSDMs to train the CNN as the prediction model. The CNN is
a deep learning architecture with multiple convolutional layers responsible for detecting local
features of the inputs. A convolution layer is composed of several convolution kernels that
compute different feature maps. To enable local feature detection, each neuron of a feature
map is connected to a small neighborhood of outputs from the previous layer. The feature map
is generated by convolving the input with a learned kernel and connected to an element-wise
nonlinear activation function. A pooling layer is usually placed between two convolution lay-
ers to reduce computational complexity and achieve shift-invariance. The pooling layer takes
a small neighborhood from the convolutional layer and subsamples it to produce a single
output (e.g. average or maximum) from that neighborhood.

For image classification, the unknown network weights $W$ of a given CNN can be learned
with a labeled training image set $S = \{(x_i, y_i)\}, i = 1, \ldots, m$ by solving the optimization
problem: $\min_W \sum_{i=1}^{m} \ell(f(W; x_i), y_i)$, where $x_i$ is an input image, $y_i$ is the corresponding label,
\( f(W; x_i) \) is the prediction with network weight \( W \) for input \( x_i \), and \( \ell \) is a negative log-likelihood loss function defined in terms of the normalized soft-max probability. This optimization problem is typically solved by a stochastic gradient descent scheme. In each iteration, given a shuffled fixed size mini-batch \( I_t \in S \), the weight update is given by \( W^{(t+1)} = W^{(t)} + Z^{(t+1)} \) where \( Z^{(t+1)} = \mu Z^{(t)} - \lambda \nabla_W \left( \sum_{i \in \Gamma} \ell(f(W^{(t)} + \mu Z^{(t)}; x_i), y_i) \right) \). The free parameter \( \mu \) denotes the momentum that indicates the contribution from the previous weight update, and \( \lambda \) denotes the learning rate.

### 2.5. Transfer learning

Considering the limited number of patient data in this study, training a large CNN from scratch with random initialization would be impractical and could easily lead to over-fitting. As an alternative, we opted to utilize the learned knowledge from a pre-trained network and apply the trained parameters to a new classification task in a process called **transfer learning** (Shin et al. 2016, Tajbakhsh et al. 2016). Transfer learning begins by initializing the network with pre-trained weights from a network of the same architecture and then **fine-tunes** the parameters to accommodate the target application. Depending on the class number of the new classification task, the last fully connected layer is usually replaced with as many neurons as the new class number. In this study, we substituted the last fully connected layer with two neurons, corresponding to toxicity and non-toxicity.

We used the state-of-the-art VGG-16 (Simonyan and Zisserman 2014) as our network architecture. This CNN achieved substantially improved performance over the other networks in the ImageNet Large Scale Visual Recognition Challenge (ILSVRC) 2012 (Russakovsky et al. 2015). The VGG-16 network (figure 1) is a deep CNN that consists of 16 layers, including 13 convolution layers and 3 fully-connected layers (termed fc6-fc8). All the convolution layers are built with fixed filters with a size of \( 3 \times 3 \), and the stride and padding are fixed at 1. There are 5 groups of convolution layers (termed conv1-conv5), each of which is followed by a max pooling layer with a window size of \( 2 \times 2 \) that down-samples the images to reduce computational burden and control over-fitting. The filter numbers of conv1-conv5 are 64, 128, 256, 512, and 512. We pre-trained the VGG-16 CNN with the ImageNet (Deng et al. 2009), which is currently the largest image dataset, with more than 1.2 million natural images of 1000 object categories. These 1000 object categories represent objects that we come across in our day-to-day lives, such as species of dogs, cats, various household objects, vehicle types etc. Given an input test image, the ImageNet-trained VGG-16 CNN generates the test image’s probabilities (or label) belonging to each of the 1000 object class. Options to utilize the pre-trained parameters in VGG-16 include: (1) ‘shallow tuning’: fine-tune only the last few fully connected layers; and (2) ‘deep tuning’: fine-tune all the network layers. In this study, considering the substantial difference between the source application (natural image) and target application (dose image), we opted to fine-tune all layers in VGG-16 and compare its gains over the ‘shallow tuning’.

### 2.6. Gradient-weighted class activation map (Grad-CAM)

In addition to predicting rectum toxicity using VGG-16 CNN, we would like to understand what features have been learned and where these features are located. To identify and locate the learned features that distinguish toxicity from non-toxicity, we used the gradient-weighted class activation mapping (Grad-CAM) (Ramprasaath et al. 2016) to highlight
discriminative regions on the RSDM when VGG-16 completes prediction. Denote the last convolutional layer produce \( K \) feature maps \( A_k^l \in \mathbb{R}^{u \times v} \) and a Grad-CAM map \( L_{Grad-CAM}^{k} \in \mathbb{R}^{u \times v} \). To obtain the Grad-CAM map, the gradient of score \( y \) with respect to feature maps \( A_k^l \) of a convolutional layer, i.e. \( \frac{\partial y}{\partial A_k^l} \), is flowed back and global-average-pooled to obtain the weights \( a_k^l \) by \( a_k^l = 1/Z \sum \sum \frac{\partial y}{\partial A_k^l} \). The Grad-CAM map \( L_{Grad-CAM}^{k} \) is calculated as \( L_{Grad-CAM}^{k} = ReLU\left( \sum_k a_k^l A_k^l \right) \), where \( ReLU\left( \cdot \right) \) is the rectified linear unit. The final Grad-CAM map is obtained by up-sampling \( L_{Grad-CAM}^{k} \) to the size of the input image.

2.7. Strategies to avoid over-fitting

Though training the large VGG-16 from scratch is resolved by utilizing transfer learning, we still need to fine-tune the million-parameter network, which is challenging. To reduce the chance of over-fitting, two issues, including data imbalance and data scarcity, need to be addressed. The training patient cohort in this study is limited and not balanced, where the toxicity cases constitute only a small minority of the data. Learning from imbalanced data is risky because the network may tend to predict more often on the majority non-toxicity class to score an overall high accuracy, and scarify the sensitivity of identifying the minority toxicity cases.

As for data imbalance, we employed the adaptive synthetic sampling approach (ADASYN) (He et al 2008) to generate synthetic minority toxicity data to balance the training data set. The ADASYN determines a weighted distribution for different minority class samples according to their level of difficulty in learning. For each minority class sample, the more adjacent majority samples it has in a certain neighboring range, the more synthetic examples will be created. Specifically, given the labeled training set \( S = \{ (x_i, y_i) \}, i = 1, \ldots, m \) with a minority class \( S_{\text{min}} \) and a majority class \( S_{\text{maj}} \), the number of data needed to synthesize is determined by \( G = (m_t - m_s) \times \beta \), where \( m_t \) and \( m_s \) represent the numbers of minority and majority class samples, respectively, and \( \beta \in [0, 1] \) is a parameter used to specify the desired balance level after generating the synthetic data. For each \( x_i \in S_{\text{min}} \), we can find \( K \) nearest neighbors based on the Euclidean distance and calculate the density distribution \( r_i = \Delta_i/K, i = 1, \ldots, m_s \), where \( \Delta_i \) is the number of examples in the \( K \) nearest neighbors of \( x_i \) that belongs to the majority class, and \( Z \) is a normalized constant so that \( \sum r_i = 1 \). The number of synthetic samples needed for each minority sample \( x_i \in S_{\text{min}} \) is given by \( g_i = r_i \times G \). For each of the \( g_i \) synthetic data of \( x_i \in S_{\text{min}} \), one minority example \( x_{ij} \) is chosen at random from the \( K \) nearest neighbors of \( x_i \), and the synthetic data can be generated by \( s_i = x_i + (x_{ij} - x_i) \times \lambda \), where \( \lambda \) is a random number \( \lambda \in [0, 1] \).

As for data scarcity, data augmentation was employed to artificially increase the size of the training dataset. Random geometric image transformation, including translation, rotation, scaling, flipping, as well as augmenting image intensity values, such as blurring and noise addition, were applied to the ADASYN balanced data prior to each fine-tuning iteration to increase the training image dataset to 10 times of its original size. This data augmentation strategy has been shown to be helpful for avoiding over-fitting and successful generalization (Ronneberger et al 2015, Milletari et al 2016, Kayalibay et al 2017).

To even further reduce the chance of getting over-fitting, we opted to only ‘slightly’ fine-tune the VGG-16 CNN with the augmented training dataset, i.e. a minimal number of training epoch of one was used for fine-tuning, intending to maintain the classification capability that mostly learned from the large natural image dataset ImageNet, and also adapt the network with the flexibility to the new application on the dose image.
2.8. Implementation details

For all patients, the rectum was extracted between the level of the ischial tuberosity and the rectosigmoid junction, with rectum length ranging from 6 cm to 9 cm in the patient cohort. For rectum surface meshing, a region of interest (ROI) with a size of $200 \times 200 \times 100$ (234 mm $\times$ 234 mm $\times$ 200 mm) encompassing the rectum was extracted, and the segmented rectums in each BT fraction were converted to surface meshes with 1500 vertices for the subsequent TOP-DIR surface point matching. For rectum surface unfolding, $n = 30$ rays were emitted from the centroid at a spaced interval of $12^\circ$. The flattened RSDM had a size of $30 \times z$, where $z$ is a sampling number along the superior-inferior direction. We set $z = 35$, which approximates the resolution of $0.5$ cm in the SI direction. To accommodate the VGG-16 architecture designed for color image inputs, the flattened grey-scale 2D RSDMs were coded to RGB images with a red-blue colormap, where the two extrema, red and blue, correspond to the maximum 90 Gy and the minimum 50 Gy in EQD2 doses (biologic equivalent dose in 2 Gy fractions). The colorized RSDMs were resampled to $224 \times 224$ before being fed to VGG-16. For all of the evaluations, a learning rate $\lambda = 0.0001$ were empirically chosen with a training mini-batch size of 5.

The rectum surface meshing was programmed under the Microsoft Visual C++ 2010 platform. The TOP-DIR was implemented on the compute unified device architecture (CUDA) programming environment, and the rectum unfolding and the ADASYN algorithm were coded in Matlab R2015b. The VGG-16 was built on Python 2.7 equipped with two machine learning libraries: Lasagne (Dieleman et al. 2015) and Theano (Theano Development Team 2016).

2.9. Prediction quantification and comparisons with predictions via $D_{0.1/1/2cc}$ and texture features

To quantitatively evaluate the registration accuracy of the rectum surface, we employed four similarity metrics (Chen et al. 2015b, 2016): the Dice’s coefficient (DC), the percent error (PE), the mean vertex-to-vertex distance (VVD), and the Hausdorff distance (HD). Higher DC or lower PE, VVD, and HD indicate better results.

The prediction performance was quantified by the mean accuracy ($\text{ACC} = (\text{TP} + \text{TN})/(\text{TP} + \text{FP} + \text{FN} + \text{TN})$), sensitivity ($\text{SEN} = \text{TP}/(\text{TP} + \text{FN})$), specificity ($\text{SPE} = \text{TN}/(\text{TN} + \text{FP})$), and the area under the receiver operating characteristic curve (AUC), where TP is true positive, TN is true negative, FP is false positive, and FN is false negative. The repeated stratified 10-fold cross validation (CV) and the leave-one-out cross validation (LOOCV) method were used to assess the prediction performance.

For comparison, we evaluated the advantage of the proposed model over using the dose volume parameters, i.e. $D_{0.1cc}$, $D_{1cc}$, and $D_{2cc}$ (most exposed 0.1, 1, and 2 cm$^3$ volume), for toxicity prediction. The $D_{0.1/1/2cc}$ were first computed from the EBRT + BT EQD2 dose by the ‘worst-case scenario’ (WS) addition method (ICRU 2013), and a logistic regression was performed on the extracted $D_{0.1/1/2cc}$ to calculate the toxicity probability.

We also extract 43 texture features from the RSDM, including 3 first-order gray level statistical features, 9 gray level co-occurence matrix (GLCM) texture features, 13 gray level run-length matrix (GLRLM) texture features, 13 gray level size zone matrix (GLSZM) texture features, and 5 neighborhood gray-tone difference matrix (NGTDM) texture features (Vallieres et al. 2015). Statistical analysis was first performed to screen out those features with statistical significance, on which logistic regression was performed to estimate toxicity.

We used the Mann–Whitney test to perform the statistical analyses in this study. Results were considered to be statistically significant if $p < 0.05$. 

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3. Results

3.1. Rectum DIR

Though substantial deformation usually occurs between two fractional rectums, the TOP-DIR can still produce satisfactory registration results (as example cases shown in figure 2). For all 42 evaluated patients, there are 198 BT fractions in total, and 156 DIRs have been performed. The average DC, PE, VVD, and HD before and after TOP-DIR registration over the patient group are summarized in table 1. The mean DC increases from 0.71 to 0.86, and the mean PE, VVD, and HD decrease from 0.62, 1.5 mm, and 7.0 mm to 0.26, 0.7 mm, and 3.9 mm, respectively. These quantitative results indicate that the TOP-DIR based rectum surface registration achieved high DIR accuracy.

3.2. Rectum unfolding

An example case of a rectum surface dose rendered in 3D and the corresponding unfolded 2D RSDM may be seen in figures 3(a) and (b), respectively. The cut is positioned along the posterior-most end of the rectum, and therefore, the horizontal axis of the flattened surface dose map denotes the position along the contour circumference, starting clockwise (view from rectum top to bottom) in the directional order of ‘posterior’, ‘left’, ‘anterior’, ‘right’, and ‘posterior’. The vertical axis lies along the superior and inferior direction.

The averaged RSDMs for patients with and without toxicity are shown in figures 6(b) and (c), respectively. These figures show that the high dose region is at the top part of the rectum in patients who developed toxicity. This observation is analogous to those reported by Heemsbergen et al (2005) and Munbodh et al (2008), who studied rectum toxicity in prostate cancer patients with EBRT and also observed similar upward shift of the high dose region. We
can also see from figures 6(b) and (c) that the average dose is higher in the high dose region of the toxicity group and covers a larger rectum region. However, as pointed out by Munbodh et al (2008), the high dose region in the average RSDM should be interpreted with caution: the larger high dose region found in the average toxicity group RSDM does not necessarily mean that individual RSDMs of each toxicity patient will show a larger high dose region. One possible explanation for this phenomenon is that the high dose regions in the non-toxicity group are more dispersed and therefore appear smaller in the averaged RSDM.

A pixel-wise Mann–Whitney test performed on the RSDMs between the toxicity and non-toxicity groups generated a pixel-wise $p$-value map; only small $p$ values ($< 0.05$) are shown in figure 6(a). The region that shows an evident statistical difference ($p < 0.01$, arrow in figure 6(a)) is located on the upper high dose region of the rectum.

3.3. Prediction performance comparisons

Table 2 lists the prediction results by logistic regression on the cumulative $D_{0.1/1/2cc}$ EQD2 doses extracted from the rectum surface via the WS addition method. Though the mean $D_{0.1/1/2cc}$ on the toxicity group are generally higher than the non-toxicity group, the prediction capability of the logistic regression on the $D_{0.1/1/2cc}$ is only moderate, with SEN: 56.7% and 52.8%, SPE: 50% and 60.0%, and AUC: 0.47 and 0.58 in LOOCV and 10-fold CV, respectively. These results suggest that the 1D dose volume parameters $D_{0.1/1/2cc}$, which exclude spatial dose information, are not powerful rectum toxicity predictors.

Forty-three texture features were extracted from the RSDM, and only four features were found to be statistical significant ($p < 0.05$), including two GLCM (contrast, correlation) and two NGTDM (coarseness, complexity), which were further used for toxicity prediction by logistic regression. As summarized in table 3, texture features from the RSDM can generate better predictive performance than the $D_{0.1/1/2cc}$, with SEN: 72.1% and 70.8%, SPE: 59.0% and 58.0%, and AUC: 0.7 and 0.7 for 10-fold CV and LOOCV, respectively. This implied that features from the 2D RSDM might offer more useful dosimetric information to correlate rectal toxicity and dose distribution.
For the proposed model, we have compared different fine-tuning strategies and validated the gains of transfer learning over training the VGG-16 from scratch. The comparison results are summarized in Table 4 and Figures 4 and 5. The VGG-16 network was either ‘shallowly’ fine-tuned on the last few fully connected layers (e.g. only fc8, or fc7/fc6–fc8), or fine-tuned ‘deeper’ on all the layers (e.g. conv1–fc8). The VGG-16 performs moderately when trained from scratch or with only the last few fully connected layers fine-tuned. Incremental performance may be observed when more convolutional layers are included in fine-tuning. The ROC analysis for LOOCV (Figure 4) also favors the inclusion of more layers for fine-tuning, and similar trends were also observed in 10-fold CV. Furthermore, decreased prediction performances were seen when more training epochs were used (Figure 5), implying a tendency of getting over-fitting. Therefore, we opted to fine-tune all layers of VGG-16 with a small training epoch number of one for the rest of the evaluations in this study. Fully fine-tuning all layers of VGG-16 achieves satisfactory prediction results, with an SEN: 61.1% and 75%, SPE: 70% and 83.3%, and AUC: 0.70 and 0.89 respectively for 10-fold CV and LOOCV.

Better performance was seen in the LOOCV than the 10-fold CV, this might attribute to the smaller size of training dataset used in each fold of validation in the 10-fold CV, where such performance discrepancy can be evident in a small sample scenario. Though 10-fold CV is a well-accepted validation procedure in the data mining field with large sample size, the LOOCV can also produce unbiased validation when the available data are rare, especially in the medical domain where only limited data samples are available (Refaeilzadeh et al 2009).

### 3.4. Grad-CAM map

In LOOCV, for each testing RSDM input into the final trained VGG-16, a Grad-CAM map is generated along with the output prediction. The Grad-CAM map is resampled to the size of the input image and normalized to [0, 1], with higher values representing greater importance in making the prediction decision. The Grad-CAM map can help understand how the network scores the input RSDM with learned features (e.g. dose distribution pattern) from different regions of the input RSDM. Generally, regions of the Grad-CAM map with larger values

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**Table 2.** The accumulative $D_{0.1/2cc}$ EQD2 doses (Gy, mean ($\mu$) ± SD ($\sigma$)) obtained by the ‘worst-case scenario’ addition method and the corresponding prediction by logistic regression.

<table>
<thead>
<tr>
<th>Patient Cohort</th>
<th>Predictive model evaluation method</th>
<th>Logistic regression</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Toxity</td>
<td>Non-toxicity</td>
</tr>
<tr>
<td>$\mu$</td>
<td>$D_{0.1cc}$</td>
<td></td>
</tr>
<tr>
<td>($\sigma$)</td>
<td>$D_{1cc}$</td>
<td></td>
</tr>
<tr>
<td>$D_{2cc}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>10-fold CV</td>
<td>LOOCV</td>
</tr>
<tr>
<td>$D_{0.1cc}$</td>
<td>85.6 ± 11.9</td>
<td>81.8 ± 8.1</td>
</tr>
<tr>
<td>$D_{1cc}$</td>
<td>72.8 ± 8.1</td>
<td>70.2 ± 4.7</td>
</tr>
<tr>
<td>$D_{2cc}$</td>
<td>66.8 ± 6.9</td>
<td>64.4 ± 3.8</td>
</tr>
</tbody>
</table>

**Table 3.** Prediction on statistical significant texture features by logistic regression.

<table>
<thead>
<tr>
<th>Texture features with $p &lt; 0.05$</th>
<th>Predictive model evaluation method</th>
<th>Logistic Regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>GLCM(contrast, correlation);</td>
<td>10-fold CV</td>
<td></td>
</tr>
<tr>
<td>NGTDM (coarseness, complexity)</td>
<td>LOOCV</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SEN 72.1%</td>
<td>SPE 59.0%</td>
</tr>
<tr>
<td></td>
<td>SEN 70.8%</td>
<td>SPE 58.0%</td>
</tr>
</tbody>
</table>
correspond to regions in the RSDM whose learned features are more important for toxicity prediction.

Figures 6(d) and (e), respectively, show the average Grad-CAM maps of the toxicity and non-toxicity patients. For the toxicity group, the salient regions of the Grad-CAM are located on the upper rectum, which corresponds to the high dose regions in the RSDM. In contrast, the salient regions in the non-toxicity group are mostly located in the low dose regions in RSDM. The mean dose comparisons of different salient regions between the two groups are shown in figure 6(f). The Grad-CAM maps suggest that the VGG-16 network utilizes learned features from the high and low dose regions in the input RSDM to yield a final prediction score. In addition, the salient regions of the average Grad-CAM of the toxicity group (figure 6(d)) have a similar distribution with the p-value map (figure 6(a)), especially in the regions with small $p < 0.01$ (arrow in figures 6(a) and (d)), suggesting that dose features from these regions in the RSDM are critical for discriminating toxicity from non-toxicity. When comparing the average toxicity Grad-CAM with the average non-toxicity Grad-CAM, the salient regions of the toxicity group may be found mainly in the high dose regions. In contrast, the salient

Table 4. Prediction performance by training from scratch or fine-tuning different layers of VGG-16.

<table>
<thead>
<tr>
<th>Fine-tuning strategy</th>
<th>10-fold CV</th>
<th>LOOCV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SEN</td>
<td>SPE</td>
</tr>
<tr>
<td>Train from scratch</td>
<td>55.6%</td>
<td>67.7%</td>
</tr>
<tr>
<td>Fine-tune only fc8</td>
<td>41.7%</td>
<td>77.4%</td>
</tr>
<tr>
<td>Fine-tune fc7–fc8</td>
<td>41.7%</td>
<td>80.4%</td>
</tr>
<tr>
<td>Fine-tune fc6–fc8</td>
<td>43.5%</td>
<td>82.2%</td>
</tr>
<tr>
<td>Fine-tune conv1–fc8</td>
<td>61.1%</td>
<td>70.0%</td>
</tr>
</tbody>
</table>

Figure 4. ROC analysis of VGG-16 trained from scratch or fine-tuned with different layers using LOOCV.
regions of the non-toxicity group spread out across the low dose regions. This implies that the VGG-16 CNN used dose features from the high and low dose regions to make prediction decisions, and such features could be geometric features, such as shape, location, area, perimeter (or perhaps their ratios), etc, from the high and low dose regions of the rectum.

4. Discussion and conclusions

In this paper, we have successfully developed a rectum toxicity prediction model by applying a pre-trained CNN network to unfolded RSDMs from cervical cancer patients treated with combined BT and EBRT. A state-of-the-art VGG-16 CNN pre-trained on a large-scale natural image dataset, ImageNet, was used as the network structure and was further fine-tuned on the training RSDMs cohort to adapt to our medical application. Satisfactory prediction results achieved in this work have demonstrated the feasibility of transferring the learned CNN knowledge from natural image to medical image, even though the substantial difference between the two applications suggests that such transfer may be impossible. To the best of our knowledge, this is the first attempt to apply the transfer learning of CNN to radiation dose distribution analysis.

The first step toward estimating an accumulative EBRT + BT dose is to accurately align the underlying anatomy. Therefore, the total EBRT + BT dose is ideally obtained by DIR, which provides means for accumulating dose at tissue voxel level and theoretically can predict
accurate dose absorption and patient’s response to a course of treatment. In our EBRT treatment plan regimen, a homogenous dose distribution (hot spot < 107%) often covers the entire pelvic region. Under this scenario, even if the rectum has a large motion, it still receives almost the same dose as planned. On the other hand, DIR between EBRT and BT CTs is a non-trivial task because of the clinical use of the intracavitary applicator in BT. Registering the BT CT image with applicator to the EBRT CT image without applicator (or vice versa) is difficult, if not impossible, since the point-to-point correspondence assumption is usually violated in most DIR algorithms. There are several reported attempts to address this issue (Berendsen et al 2013, Vásquez Osorio et al 2015), for example, Berendsen et al (2013) proposed a DIR with penalty term that minimizes the volume of the missing structure for cervical MR images with and without applicator. Vásquez Osorio et al (2015) validated a structure-wise registration with vector field integration to map the largely deformed anatomies between EBRT and BT. However, these novel approaches need comprehensive validations before they can be confidently applied in a clinical setting. Thus adding EBRT to BT without deformation is a good approximation without knowing the uncertainties brought by the EBRT-BT DIR.

The hollow anatomic structure of the rectum makes it possible to take advantage of the CNN, which takes 2D images as input. The flattened 2D RSDM contains the spatial dose distribution information and is intrinsically superior to the 1D dose volume parameters, e.g. $D_{0.1/1/2cc}$, in revealing the dose-toxicity relationship, as shown by the superior prediction performance achieved via the proposed model compared to the prediction using logistic regression on $D_{0.1/1/2cc}$. Furthermore, the proposed model can easily extend its application to other critical organs with hollow structures, such as the bladder, vagina, etc.

Figure 6. (a): Pixel-wise $p$-value map (Mann–Whitney test) with small $p < 0.05$ between (b) and (c): the average rectum RSDM of the toxicity and non-toxicity patients. The arrow in (a) indicates regions with $p < 0.01$. (d) and (e): Average Grad-CAM map of the toxicity and non-toxicity groups. The arrow indicates the salient region. (f): Box plot of the mean dose in different salient regions extracted from the Grad-CAM map. The boxes run from the 25th to 75th percentile; the two ends of the whiskers represent the 5% and 95% of the data, and the horizontal line in the box represents the median value. The circles represent outliers.
In this study, we used VGG-16 as the CNN architecture mainly because of its outstanding performance in the large scale natural images classification challenge (Simonyan and Zisserman 2014). Other state-of-the-art CNN structures, such as CifarNet (Krizhevsky 2009), AlexNet (Krizhevsky et al 2012), GoogLeNet (Szegedy et al 2014), etc, may yield even better prediction results. Nevertheless, we did not validate these networks, since the purpose of this study was to demonstrate the possibility of transferring learned knowledge between two distinct applications, instead of comparing prediction performances among the currently available CNNs.

Though recent studies have demonstrated the successful application of transfer learning in medical fields with different image modalities, such as x-ray, CT, ultrasound, colonoscopy, etc, (Bar et al 2015, Chen et al 2015a, Ginneken et al 2015, Shin et al 2016, Tajbakhsh et al 2016), the rationale for its effectiveness is still under investigation. The ‘transferability’ of knowledge imbedded in a pre-trained network depends on the difference between the two databases: the one on which the CNN was trained and the one to which the learned knowledge will be transferred (Tajbakhsh et al 2016). The difference between natural images and textured medical images (e.g. CT, MR) is considerable, and the difference between natural images and radiation dose images is even greater. As a result, fine-tuning all of the CNN layers might be necessary (Tajbakhsh et al 2016). The experimental results in this study support the claim that the CNN tends to behave more stably when lower convolutional layers are also fine-tuned.

This work also demonstrates the gain of transfer learning over training from scratch. Given the relatively small patient sample size, training a CNN from scratch is difficult, if not impossible, because a small sample size might not be sufficient to train a large number of network parameters in a deep CNN. For transfer learning, in contrast, the network parameters have been initialized and only need further fine-tuning. Previous studies have shown that transferring features followed by further fine-tuning can result in networks that generalize better than those trained directly on the target dataset (Yosinski et al 2014). One thing we should emphasize is that the dose-toxicity prediction model established in this study is preliminary, since limited patient data was used. A large dataset is a necessity to obtain a more powerful and accurate prediction model.

In this study, we also try the Grad-CAM to reveal the possible location of rectal toxicity. We found that the small $p$-value regions on the RSDM (figure 6(a)), which indicate the difference between the toxicity and non-toxicity RSDMs, coincide with the salient regions of the average Grad-CAM of the toxicity group. Each Grad-CAM map was generated by the prediction model along with every input testing RSDM, so the salient region of the averaged Grad-CAM map informs us where those important dose features come from when the prediction model makes classification decisions. The small $p$-value regions on RSDM and the salient region of the toxicity group’s Grad-CAM both point to the superior part of the rectum, suggesting that the superior rectum might imply a possible positional explanation for rectum toxicity. This observation accords with those reported by Heemsbergen et al (2005) and Munbodh et al (2008), who found that the high dose was shifted upward to the top part of the rectum in their studies of rectum toxicity in prostate cancer patients. However, further in-depth investigations are still needed to provide more solid clinical evidence to support the claim that the superior rectum is mainly responsible for rectum toxicity.

Acknowledgments

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