Machine Learning to Build and Validate a Model for Radiation Pneumonitis Prediction in Patients with Non-Small-Cell Lung Cancer

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Running head
Machine learning build and validate RP2 prediction in NSCLC

Keywords
Machine learning; Non-small-cell lung cancer; cytokine; radiation pneumonitis;

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Statement of translational relevance
Radiation pneumonitis is a dose limiting toxicity of thoracic radiation therapy. Combining patient factors like hypertension, the lung dosimetric parameters like mean lung dose, and plasma levels of biomarkers like IL-8 and CCL2, this study built and validated internally a predictive model for radiation pneumonitis grade≥2 with AUC of 0.863 and accuracy 80% in patients with non-small-cell lung cancer who underwent radiation therapy. Should these be validated by an external database, this study will provide an opportunity to guide clinicians for personalized radiation dose prescription in future trials or clinical practice, to improve patient’s survival while limiting the risk of radiation pneumonitis.
Abstract

Purpose
Radiation pneumonitis is an important adverse event in patients with non-small-cell lung cancer (NSCLC) receiving thoracic radiation therapy (RT). However, the risk of radiation pneumonitis grade ≥ 2 (RP2) has not been well predicted. This study hypothesized that inflammatory cytokines or the dynamic changes during-RT can improve predictive accuracy for RP2.

Materials and Methods
Levels of 30 inflammatory cytokines and clinical information in patients with stages I-III NSCLC treated with RT were from our prospective studies. Statistical analysis was used to select predictive cytokine candidates and clinical covariates for adjustment. Machine learning algorithm was used to develop the generalized linear model for predicting risk RP2.

Results
A total of 131 patients were eligible, 17 (13.0%) developed RP2. IL-8 and CCL2 had significantly (Bonferroni) lower expression levels in patients with RP2 than without RP2. But none of the changes in cytokine levels during RT was significantly associated with RP2. The final predictive GLM model for RP2 was established including IL-8 and CCL2 at baseline level and two clinical variables. Nomogram was constructed based on the GLM model. The model’s predicting ability was validated in the completely independent test-set (area under curve=0.863, accuracy=80.0%, sensitivity=100%, specificity=76.5%).

Conclusion
By machine learning, this study has developed and validated a comprehensive model integrating inflammatory cytokines with clinical variables to predict RP2 before RT which provides an opportunity to guide clinicians.
Introduction

Radiation therapy (RT) plays an important role in the treatment of lung cancer, the leading cause of cancer death. Radiation induced lung toxicity (RILT) is a common and dose limiting adverse effect of thoracic RT in lung cancer patients, which may decrease quality of life, lead to pulmonary failure, and become life-threatening.\(^1,2\) Radiation pneumonitis (RP), one of the commonly reported RILT, usually occurs within 1 to 6 months after completion of RT.\(^3\) In patients treated with concurrent chemoradiation therapy,\(^4-6\) 7.0 to 32.0\% of patients have grade 2 and above (RP2) while 2.6 to 18.0\% with severe RP Grade $\geq$3.

We and others have previously demonstrated that the risk of RILT is correlated with radiation dosimetric factors, like mean lung dose (MLD)\(^2,7,8\) with AUC$<0.60$. Proteomic analysis demonstrated that molecules associated with inflammation pathways such as C4b-binding protein alpha chain (C4BPA), Complement C3 (C3) and vitronectin (VTN) had substantially higher expression levels in patients with grade $\geq$2 RILT.\(^9\) Addition of C4BPA+VTN to MLD improved the RILT predictive accuracy (AUC=0.71).\(^10\) We also found radiation-induced elevation in plasma TGF-β1 level during RT had predictive ability of RILT.\(^11\) TGF-β1 combined with MLD stratified patients for high risk of RILT.\(^12\) Additionally, we demonstrated that combining IL-8, TGF-β1, and MLD into a single model yielded a good predictive ability
In addition, baseline pulmonary function, including FEV1, FVC, and DLCO, may be related to the risk of RILT.\textsuperscript{13}

The pathogenesis of RILT is described as multiple inter-reacting cellular activities such as hypoxia, fibrogenesis, inflammation, and angiogenesis.\textsuperscript{14} It is known that RILT combined the events of RP and radiation induced lung fibrosis (RILF) together, though RP and RILF have different biopathophysiological mechanisms. RP is associated with inflammatory reaction, while the latter is direct results of fibrosis and scar formation. The biomarkers of RP have been studied more extensively for clinical prediction. Kim JY, et al.\textsuperscript{15} found that TGF-\(\beta\)1 level became significantly higher at 4 weeks after RT (\(p = 0.007\)). Variations of circulating IL-1A, IL-6, and IL-10 were also significant with RP (\(p < 0.05\)).\textsuperscript{16,17} Serum superoxide dismutase (SOD) has the predictive ability of RP with a sensitivity of about 0.8, and a specificity of about 0.7.\textsuperscript{18,19} Additionally, some groups revealed a number of single-nucleotide polymorphism markers (SNPs)\textsuperscript{20-23} were significantly correlated with the incidence of RP, including TGF-\(\beta\)1 rs1982073 with RP\textsuperscript{2} (hazard ratio = 0.489);\textsuperscript{20} TGF-\(\beta\)1 rs11466345 with RP3 (hazard ratio = 2.295).\textsuperscript{21} Genetic variation in the pro-inflammatory genes IL-1A, IL-8, TNF, TNFRSF1B, and MIF also significantly increased the risk of RP;\textsuperscript{22} MTHFR rs1801131 with RP2 (hazard ratio = 0.37).\textsuperscript{23} These studies suggested that individual patient's genetic makeup and cytokine milieu may play critical roles.
in an individual’s response to RP2 development. However, no study to date has reported good validated models to predict the risk of RP.

In this study, we hypothesized that cytokines may play a vital role in predicting RP. We measured the plasma levels of representative cytokines of having immunomodulating and inflammatory effects, including interleukin, colony stimulating factor, interferon, tumor necrosis factor, transforming growth factor, growth factor and chemokine families, and even their changing dynamic during the course of radiation. This study aimed to build and validate a model to predict RP2 by using plasma cytokine in patients with NSCLC who underwent radiation therapy.

Materials and Methods

Study population

Eligible subjects included patients with stages I-III NSCLC undergoing radiation alone or combined radiation with chemotherapy (UMCC 2003.073, UMCC 2003.076, NCT00603057, NCT01190527). Patients with a life expectancy of less than 6 months were excluded as they might not benefit from local radiation and might not be assessable for late lung toxicity. No restrictions were placed on either the degree of weight loss or pulmonary compromise, or oxygen dependency. All clinical data, including clinical parameters, grading of RP, and blood samples, were prospectively collected.
Radiation therapy

All patients received daily fractionated 3D conformal external beam radiotherapy technique with or without sequential or concurrent chemotherapy. No patients treated with stereotactic body RT were included in the analysis. In general, the radiation dose prescription is limited by MLD of 20 Gy or normal lung tissue complication probability of 15%-17.5%. The lung dosimetric factors were computed with subtraction of gross tumor volume (GTV) overlapping with normal lung.

Cytokines measurement

A total 30 inflammation modulating cytokines were measured, including: EGF, VEGF, CCL2, CCL3, CCL4, CCL11, CX3CL1, CXCL10, G-CSF, GM-CSF, IFN-γ, TNF-α, IL-1a, IL-1b, IL-1r, IL-2, IL-4, IL-5, IL-6, IL-7, IL-8, IL-10, IL-12p40, IL-12p70, IL-13, IL-15, IL-17, sCD40l, TGF-α, and TGF-β1. Cytokine measurements were performed in plasma samples at 3 time-points: at baseline (within 2 weeks before the start of RT) and at 2 and 4 weeks during RT. Cytokines were measured in pg/mL as previously described.

Evaluation of radiation pneumonitis

The primary endpoint of radiation therapy was RP2, defined as RP grade ≥2. RP was diagnosed and graded based on a modified criteria combining...
RTOG/SWOG/CTCAE. The detailed grading definitions were previously described in our previous publications,\textsuperscript{2,12} consistent with a recent update from the expert panel of an AAPM task (Supplementary Table S1).\textsuperscript{26}

**Statistical analysis**

Patients with detectable levels of all 30 cytokines were eligible. Chi-square test, Fisher’s exact test, and logistic regression were applied for univariate clinical variables’ analyses, in order to select covariates available for model development. GLMM (generalized linear mixed model) with Bonferroni multiplicity correction was used to assess the potential importance of each cytokine as well as its dynamics (at baseline, 2 and 4 weeks during-RT), after adjusting for potential prognostic covariates as identified in the univariate clinical variables’ analyses. Machine learning algorithm was used for developing the final prediction GLM (generalized linear model) model of risk RP2 based on selected cytokines and clinical covariates, as described in the next section. The nomogram was built based on the GLM model for risk RP2. ROC curves and its corresponding AUC value, accuracy, sensitivity, specificity, positive predictive value (PPV) and negative predictive value (NPV) were calculated to show the performance of the GLM model. All analyses were performed after data normalization: Log-transformed all cytokine levels and GTV. All statistical tests were 2-sided, and the overall adjusted p-value threshold was 0.05. All analyses were performed using the "R" version
Machine learning

Given the concern about generalization performance, machine learning was adopted here to build the generalized linear model (GLM). The methodology of the process can be divided into the following steps. 1) Divide the dataset into a training cross-validation set (trainCV-set) and a test-set (80% and 20%). 2) Use ten times 10-fold cross-validation (CV) in the trainCV-set to avoid over-fitting. In the progress of CV, the trainCV-set was divided into training-set and CV-set (9 folds and 1 fold); the most fitted GLM model was generated on training-set by model performance criteria AIC; and the mean standard error (MSE) of the GLM model was calculated on the CV-set. 3) Select the model, which had an MSE value closest to the mean MSE value, as the final model for predicting RP2. 4) Tested the final GLM model in test-set, which is unused and is completely independent with the trainCV-set. Given the imbalanced data (17 cases with RP2), we random resample those datasets carefully in keeping the ratio of RP2 not to zero.

Results

Patients Characteristics and RP2

A total of 131 consecutive patients with NSCLC met the study criteria. Median age was 65.7 years (range 59.4-73.9), 22.9% were female. 84.7% were
treated with a combination of chemotherapy and RT. Seventeen of 131 patients (13.0%) developed RP2 at a minimum follow-up of 12 months.

Table 1 shows the clinical variables included in this study, including gender, age, Karnofsky performance status (KPS), smoking history, Chronic obstructive pulmonary disease (COPD), Cardiovascular disease (CVD), Hypertension, tumor location, tumor clinical stage, gross tumor volume (GTV), whether received chemotherapy, mean lung dose (MLD), mean heart dose (MHD), and equivalent dose in 2Gy fraction (EQD2). Under univariate analysis, MLD was significantly correlated with the risk of RP2 (p-value=0.029). Three candidate clinical variables with p-value<0.1, including MLD, Hypertension, and GTV were selected as covariates for further multivariable analysis.

**Analysis of single cytokine with RP2**

To identify the most influential inflammatory cytokines in predicting RP2, the associations between the risk of RP2 and cytokine expression level and rate of level change were analyzed using GLMM models. Time points (baseline, 2 and 4 weeks during RT) were defined as random effects in GLMM model. The suitable GLMM model for every single cytokine, adjusted by the three selected clinical variables (Hypertension, GTV, and MLD), was determined by the minimum AIC score. Analysis showed that two cytokines (IL-8 and CCL2)
had significantly (p-value < 0.0017, Bonferroni) lower expression levels in patients with RP2 than those without RP2 (Figure 1 and Table 2). However, none of the temporal rate of cytokine’s level during RT was statistically significant for risk RP2 after multiplicity adjustment. The full data was supplied in Supplementary Table S2.

**Multivariable model for predicting RP2**

Based on the above analysis, the expression levels of IL-8 and CCL2 at baseline as possible early predictors of RP2, along with the three clinical variables (Hypertension, GTV, and MLD), might be the candidates for the final multivariable GLM models for predicting the risk of RP2. To guarantee the predictors following a normal distribution and orthogonality, their distributions (IL-8, CCL2, and GTV in log transform) and correlations were shown in Figure 2. It can be seen that there were no strong correlations among them, while a straightforward relationship between MLD and GTV was found. According to their p-values (Table 1) and further clinical usage of the model, MLD was retained as one predictive candidate in the final multivariable model.

The final multivariable logistic prediction model was generated by machine learning as described above. This included two cytokines (IL-8 and CCL2) and two clinical variables (Hypertension and MLD), as shown in Table 2. The nomogram for predicting risk RP2 was constructed using the GLM model, as
shown in Figure 3. To evaluate the final model’s generalization performance, this model was validated in test-set which was completely independent with trainCV-set. The predictive performances on test-set were the following:

AUC=0.863 (95%CI=0.676~1, p-value=0.027), accuracy=80.0%, sensitivity=100%, specificity=76.5%, PPV=42.9%, NPV=100%. On the full dataset, the final GLM model had classified performances as the following:

AUC=0.881 (95%CI=0.799~0.963, p-value=1.299e-6), accuracy=82.0%, sensitivity=86.7%, specificity=82.0%, PPV=44.8%, NPV=97.3%. The ROC curve of the full dataset was compared with those based on univariate models and the model from our previous study for RILT, as shown in Figure 4.

**Discussion**

In this study, we studied 30 cytokines from 131 NSCLC patients enrolled in prospective clinical trials. Although our initial hypothesis was the temporal change of cytokines during RT can predict RP2 better than baseline measurements alone, the results of this study demonstrated that the expression levels of two cytokines (IL-8 and CCL2) were statistically significant for RP2, while none of the rates in the change of cytokines was significant for RP2. This has an important meaning that the individual’s intrinsic micro-environment of patients, especially inflammation cytokine levels before RT, plays an important role in the following progress of RP2 in the patient with NSCLC underwent RT.
As described above, previous groups have reported that biologic factors for RP prediction. In smaller datasets, TGF-β1, IL-6, IL-1A, SOD, GPX have shown significant p-values with RP. Specifically, IL-6 and IL-1A had been studied for their prediction ability as reported sensitivities of 50% and 53% respectively. Rs1982073 in TGF-β1, rs11466345 in TGF-β1 and rs10898880 in ATG16L2 had hazard ratios (0.489, 2.2 and 1.8 respectively). Other research groups used simple clinical variables to predicting the RP progression, such as MLD (odds ratio=2.02) and lung receiving 20 Gy of radiotherapy (odds ratio=1.41). Notably, Valdes G et al. used machine learning in predicting RP. They found radiation dosimetric parameters and patients' race were important features in RUSBoost algorithm, however the accuracy of their classification is limited and the algorithm is difficult for a clinical usage. We compared them in details as shown in Supplementary Table S3. While in our study, we developed and validated a predictive model for RP2 with AUC=0.863, using stringent statistical method and machine learning approach. This GLM model included IL-8 and CCL2 at baseline level and two clinical variables (MLD and Hypertension) as early predictors of RP2. We also validated in the completely independent test-set, with the model predictive values of over 80% (AUC=0.863, Sensitivity=100%, Specificity=0.765%), numerically better than those previous reports for RP.
Furthermore, since it predicts RP2 based on cytokines at the baseline, our model may provide an opportunity to personalize radiation treatment guidance before RT start. To our knowledge, this is the first study to validate that IL-8 and CCL2 as the early predictors for RP2, particularly to predict RP2 before RT start. Using the nomogram in Figure 3, the risk of RP2 can be calculated based on the patient's IL-8 and CCL2 levels, hypertension, and MLD values before RT start. Even the MLD value can be modulated to control the RP2 risk, which may contribute to patients' overall survival. Therefore, predicting the risk of RP2 before RT may provide guidance for the aggressiveness of the RT treatment or prescribing anti-inflammatory treatment.

Our model is promising with a predictive accuracy of 0.86. The relatively high accuracy of our final GLM model may partially contribute to consideration of clinically important variables. MLD's contribution in RP2 is consistent with previous literatures studied in RILT.7,8,10-12 Interestingly, hypertension was found that have the contribution to RP2. The biologic mechanism is unclear regarding the relationship between hypertension and RP2. It is possible that the progression of hypertension is associated with inflammation and fibrosis. However, whether inflammation is the cause or effect of hypertension is not well understood.33 Of additional note, the effect of Tumor Location on RP2 was also tested here as it was previously reported in both patient34 and...
animal studies.\textsuperscript{35,36} Although we considered it as a candidate predictor, tumor location parameter was not significant so as to be included in multivariate consideration, but not included in the final predictive GLM model. This result does not necessarily imply that tumor location is insignificant, since our sample included 20\% tumor with unknown location. More so, this may be due to the fact that the majority of our patients were stage III with some component of central diseases.

It is known that cytokines play an important role in RP2. In our study, lower IL-8 level was found statistically significant with higher risk of RP2, which is consistent with previous literature on RILT. Both of our previous studies\textsuperscript{8,37} and Hart et al.\textsuperscript{38} had reported that low IL-8 was correlated with an increased risk of RILT. IL-8 has chemotactic activity for leukocytes and induces collagen synthesis and cell proliferation\textsuperscript{39} in animal studies, but it has been consistently found to have an anti-inflammatory effect in humans.\textsuperscript{12,37,38,40} Furthermore, it has been shown that neutrophils penetrate the injury site and perform the critical tasks of dismantling injured vessels and creating channels for new vascular regrowth, which is important for full repair of the sterile injury.\textsuperscript{41} This discovery strongly supported our results that higher level of IL-8 before RT was correlated with lower risk of RP. We believe that higher level of IL-8 can chemotaxis more neutrophils, to migrate toward the site of injury caused by radiation. As enough neutrophils be recruited and activated in the repair
process, the progress of RP will not be happened because the radiation injury was almost being repaired.

This study is the first to demonstrate that low CCL2 level was associated with the increased risk of RP2. CCL2 was also known as involved in attracting neutrophils in animal studies. These factors may work together with IL-8 in RP process, as they are recognized in other conditions like inflammation of vascular disease. It may be reasonable to hypothesize that long-term overexpression of CCL2 in humans may play the same anti-inflammation role as IL-8. This needs to be validated and be a focus of future research.

It is interesting to note that neither the rate of change in TGF-β1 during treatment nor the baseline level of TGF-β1 was significantly associated with RP2; and its’ AUC for RP2 on the dataset was 0.507, as shown in Figure 3. This was different from some previous studies that investigated plasma TGF-β1 as a predictor for RILT, including our own studies.\textsuperscript{8,11-13,15,20,22,37,42} This controversial result may be multifactorial. Firstly, definitions of RP and RILT were not consistent as described in the introduction section, which may have confounded the results. The role of TGF-β1 may be more prominent for fibrosis.\textsuperscript{37,43} Secondly, TGF-β1 can be produced by both tumor and normal tissues, which seriously confound its role on RP2. The insignificant results of TGF-β1 on RP2 does not override its effect on RILF or RILT. Studies with
larger numbers of events or stratified analysis with consideration of TGF-β1
effect on tumor are needed.

While this study may be somewhat limited in the number of RP2 events and
number of cytokines tested, this has been corrected by Bonferroni test, a
stringent methodology for correction as well as by the use of machine
learning algorithms, a stringent methodology for ensuring model’s
generalization. Our study continues to show that inflammatory cytokines play
an important role in the evolution of radiation pneumonitis and further clinical
studies leveraging these relationships are warranted.

In summary, this study demonstrated that two inflammatory cytokines (IL-8
and CCL2) have strong correlation with RP2, while the temporal rate of
cytokine levels had no statistical significance with RP2. According to machine
learning algorithms, we established a predictive GLM model which included
mean lung dose, hypertension and both IL-8 and CCL2 at baseline levels as
early predictors of radiation pneumonitis. The predictive performance of the
model was validated in the independent test-set with an AUC=0.863,
Sensitivity=100%, and Specificity= 76.5%. This model and its nomogram, if
further validated externally, can provide an opportunity of guiding
personalized lung cancer treatment plan according to individual’s
inflammatory cytokines.
Reference


43. Tsoutsou PG, Koukourakis MI. Radiation pneumonitis and fibrosis: Mechanisms underlying
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<tr>
<td>1$^{st}$ – 3$^{rd}$ Qu</td>
<td>(4.5 – 19.1)</td>
<td>(4.3 – 18.6)</td>
<td>(9.6 – 19.8)</td>
<td></td>
</tr>
<tr>
<td>EQD2 (Gy) #</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>70.0</td>
<td>70.0</td>
<td>76.2</td>
<td>0.308</td>
</tr>
<tr>
<td>1$^{st}$ – 3$^{rd}$ Qu</td>
<td>(65.0 – 78.0)</td>
<td>(65.0 – 77.9)</td>
<td>(65.0 – 81.9)</td>
<td></td>
</tr>
<tr>
<td>Chemotherapy $^$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>20 (15.3%)</td>
<td>17 (14.9%)</td>
<td>3 (17.6%)</td>
<td>0.724</td>
</tr>
<tr>
<td>Yes</td>
<td>111 (84.7%)</td>
<td>97 (85.1%)</td>
<td>14 (82.4%)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviation: KPS = Karnofsky Performance Status; COPD = Chronic obstructive pulmonary disease; CVD = Cardiovascular disease; GTV = gross tumor volume; MLD = mean lung dose; MHD = mean heart dose; EQD2 = equivalent dose in 2 Gy fractions; $^\$ Fisher’s exact test; & Chi-square test; # logistic regression; ~QU=quartile * p-value < 0.05; • < 0.1
Table 2. Single cytokines’ level and the final GLM model for predicting risk RP2

<table>
<thead>
<tr>
<th>variables</th>
<th>Estimate coefficient</th>
<th>Standard error</th>
<th>Odds ratio (^9)</th>
<th>confidence interval 95%</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single Cytokines</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IL-8</td>
<td>-1.177</td>
<td>0.281</td>
<td>0.308</td>
<td>0.178 – 0.535</td>
<td>2.8e-5 **</td>
</tr>
<tr>
<td>CCL2</td>
<td>-0.979</td>
<td>0.295</td>
<td>0.376</td>
<td>0.211 – 0.669</td>
<td>8.9e-4 **</td>
</tr>
<tr>
<td>(Intercept)</td>
<td>-0.569</td>
<td>3.038</td>
<td>0.566</td>
<td>0.001 – 226.743</td>
<td>0.851</td>
</tr>
<tr>
<td>The final GLM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IL-8</td>
<td>-0.887</td>
<td>0.371</td>
<td>0.412</td>
<td>0.183 – 0.810</td>
<td>0.017 *</td>
</tr>
<tr>
<td>CCL2</td>
<td>-1.190</td>
<td>0.471</td>
<td>0.304</td>
<td>0.107 – 0.734</td>
<td>0.011 *</td>
</tr>
<tr>
<td>Hypertension</td>
<td>1.993</td>
<td>1.086</td>
<td>7.337</td>
<td>1.128 – 90.945</td>
<td>0.066</td>
</tr>
<tr>
<td>MLD</td>
<td>0.321</td>
<td>0.161</td>
<td>1.378</td>
<td>1.058 – 2.017</td>
<td>0.046 *</td>
</tr>
</tbody>
</table>

\(^9\) Ratio of levels in patient with RP2 / without RP2
*p-value < 0.05; ** < 0.0017 (Bonferroni)

Figure Legends

Figure 1. Temporal data of cytokine expression levels: (left) IL-8, (right) CCL2 (means ± 95% confidence interval)

Figure 2. Visualization of the continuous predicting candidates’ distribution and correlation. Corr= Pearson Correlation Coefficient. (x- and y-coordinate are predictors’ level).

Figure 3. The nomogram for risk RP2, constructed based on the final GLM model. It based on two cytokines (IL-8 and CCL2) and two clinical variables (Hypertension, MLD).

Figure 4. ROC curves for risk RP2 on the whole dataset, comparing the final GLM model with single cytokines (IL-8, CCL2, TGF-β1); and comparing with IL-8+TGF-β1+MLD, which were referenced from our previous study for predicting RILT.\(^8\)
Figure 1
Figure 3

Points

IL8

CCL2

Hypertension

MLD

Total Points

Risk

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