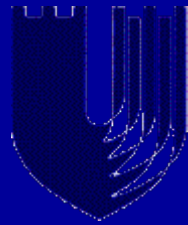


**Robust Prediction of  
Radiotherapy-induced Injury to  
Ensure Patient Safety**



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# Safety

What do planning dose constraints have to do with safety?

Everything in terms of not harming the patient!

Safety is not just about preventing mistakes that can lead to a patient being harmed. It is also about judiciously adopting guidelines (e.g., dose constraints) that can be directly linked to patient injury.

Guidelines that we usually take for granted .....

**But, it isn't that something catastrophic can happen if we use incorrect/sub-optimal guidelines for critical organs constraints, right?**

**Maybe so. This may not be the sort of disaster scenarios we associate with the word "safety". Nevertheless, adopting incorrect/sub-optimal guidelines can result in organ injury to the patient that compromises function. Also, it affects every single patient!**

**Adopting optimal guidelines can reduce injury and ensure quality of life, a broader interpretation of "safety".**

## RT-induced Pneumonitis

- The main dose-limiting toxicity for thoracic RT
- 5 - 15% of patients developed pneumonitis
- Models are needed to predict and reduce risk

## Literature

- **Identifies volumes above doses from 5 Gy to 50 Gy as being predictive of radiation pneumonitis.**
  - **Approximately 25% of volume over 20 Gy is frequently used.**
  - **Volume > 5 Gy has recently become popular.**
  - **No consensus**
- **Non-dosimetric factors largely ignored.**

## What do we need?

**A model that can be used ahead of treatment time, incorporating factors such as the dose distribution, chemotherapy, patient parameters (age, sex, etc.), which can output the probability of injury.**

**If the predicted probability of injury is high, change treatment parameters (dose, chemo).**

## What sort of model?

Complicated phenomena may be best modeled using machine-learning models (e.g., neural networks).

## Isn't this too complicated?

Do we think that modeling the weather is possible with just one or two variables? It is the same with biological injury – the simplest models have no consensus.

# Rationale for Machine Learning

**Can more easily extricate the underlying dose-effect relationship.**

Disadvantages of machine learning:

1.If not careful can overfit the model with too many features (fits the training data too well, and performs poorly on test data).

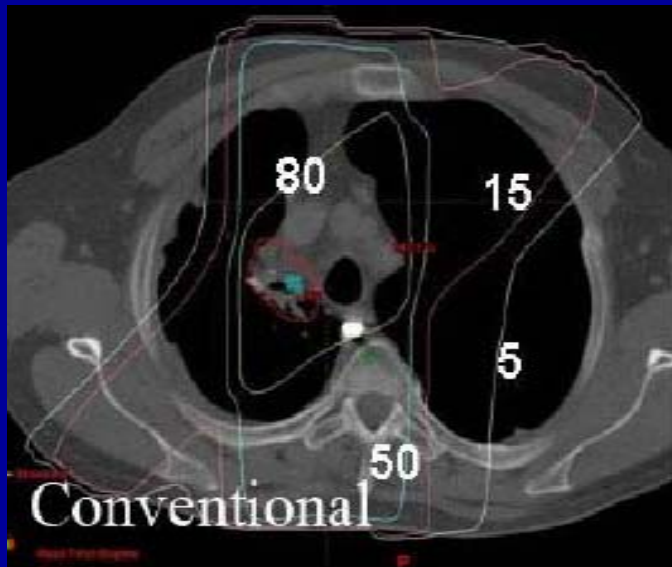
2.A particular machine learning method can have a certain bias (e.g., consistently over/underestimating injury risk in certain situations).

3.Can be hard to interpret cause-effect relationship in simple terms.



# Patient Database

- 235 lung cancer patients: 34 pneumonitis (excluding 16 “hard-to-score”)
- 3D conformal radiotherapy



- **Primary fields: AP/P**
  - 40-45 Gy
- **Boost fields:**
  - 20 Gy

# Radiation Pneumonitis

Grade	Definition
0	No increase in symptoms
1	Symptoms not requiring initiation or increase in steroids and/or oxygen
2	Symptoms requiring initiation or increase in steroids
3	Symptoms requiring oxygen
4	Symptoms requiring assisted ventilation or causing death.

# Scheme

Use both dose (lung DVH and EUD) and clinical factors to develop non-parametric models for prediction of RT-induced pneumonitis:

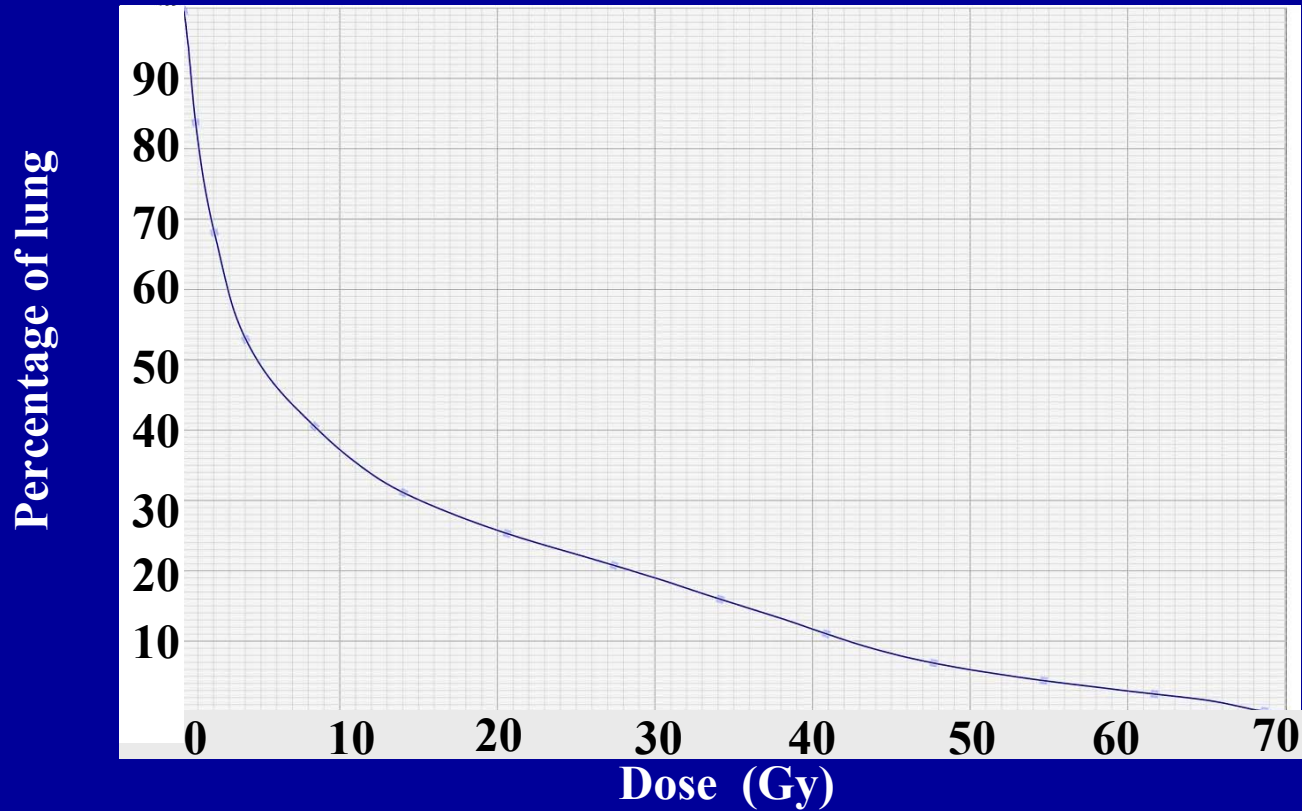
- Decision Trees
- Feed forward neural networks
- Self-organizing maps
- Support vector machines

Combine the four models (decision fusion) to predict injury.

- Reduce individual model bias.
- Extract features that are most important to all models.

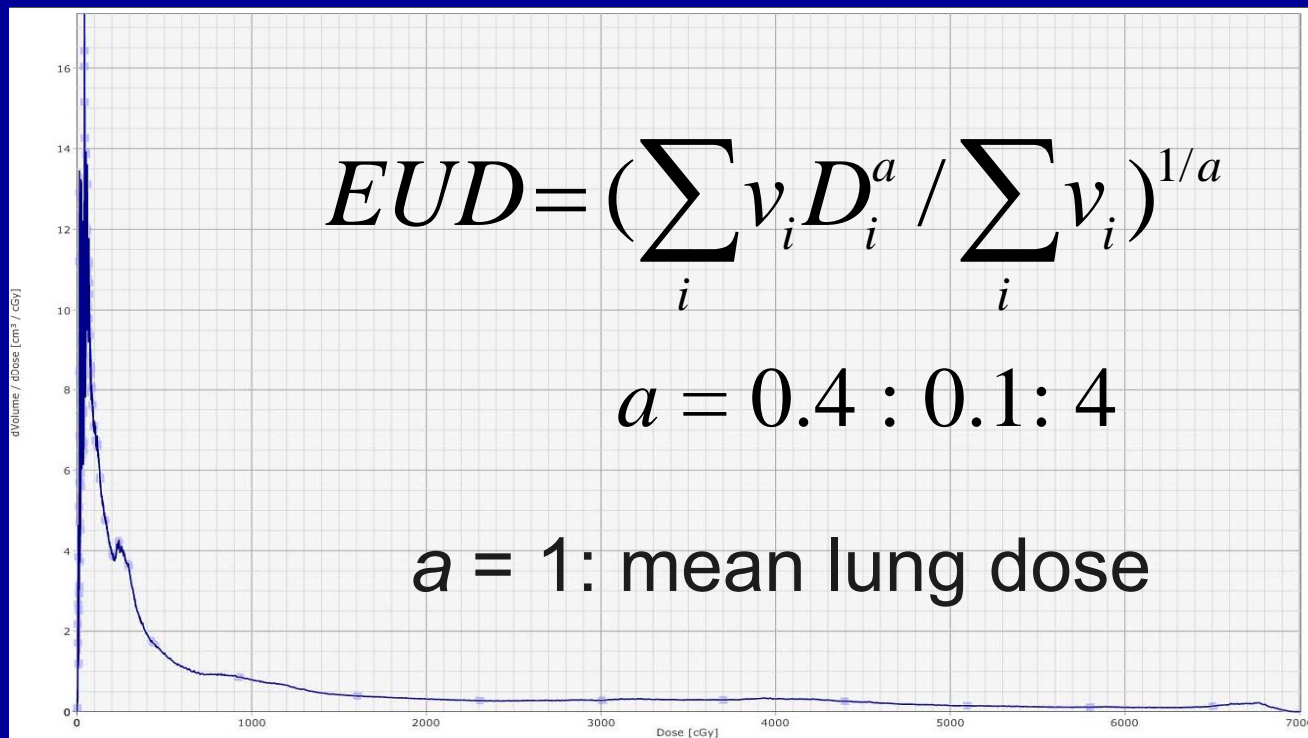
# Dose Factors

Cumulative dose-volume histogram (DVH)



# Dose Factors

Equivalent Uniform Doses (EUD) are converted from differential DVH



$$EUD = \left( \frac{\sum_i v_i D_i^a}{\sum_i v_i} \right)^{1/a}$$

$$a = 0.4 : 0.1 : 4$$

$a = 1$ : mean lung dose

$D$

# Non-dose Factors

- FEV1 (forced expiratory volume in 1 s)
- pre-RT DLCO (Carbon Monoxide diffusion capacity in lung)
- chemotherapy
- race, age, gender, tumor stage
- tumor location (right/left, up/middle/low, central/peripheral)
- histology (squamous/adenocarcinoma/non-small/small/large/other)
- once/twice daily RT
- surgery (yes or no)

- **Ten-fold cross-validation**
  - Build and test the models
- **Receiver Operating Characteristics (ROC) Curve**
  - Assess model predictive ability

# Ten-Fold Cross-Validation

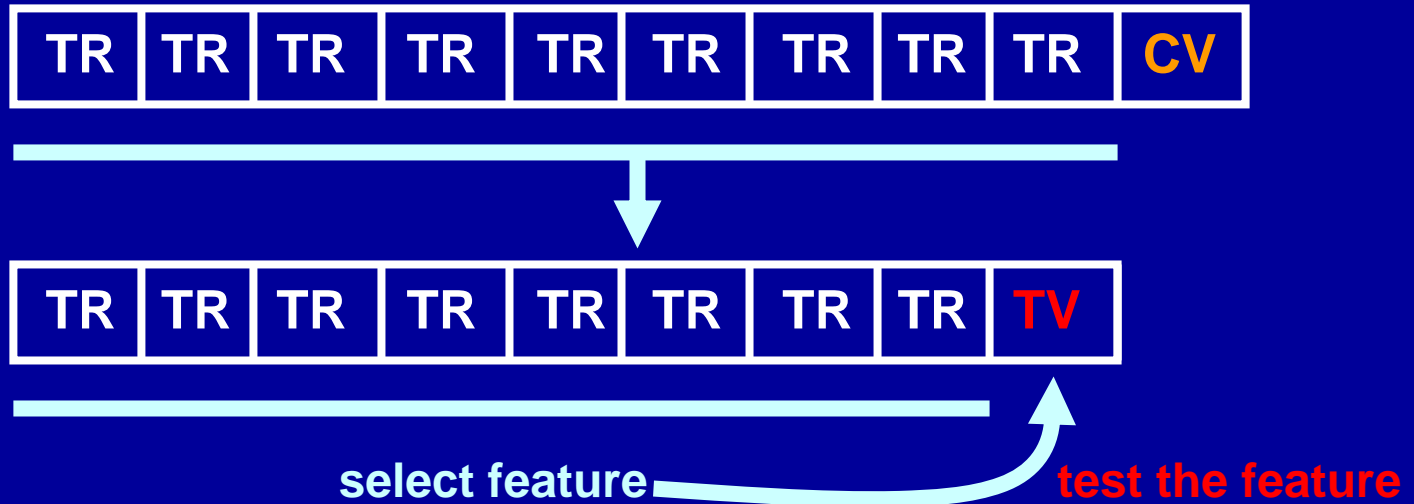


TR: training data

CV: cross-validation



# Feature selection



TR: training data, TV: training validation, CV: cross-validation

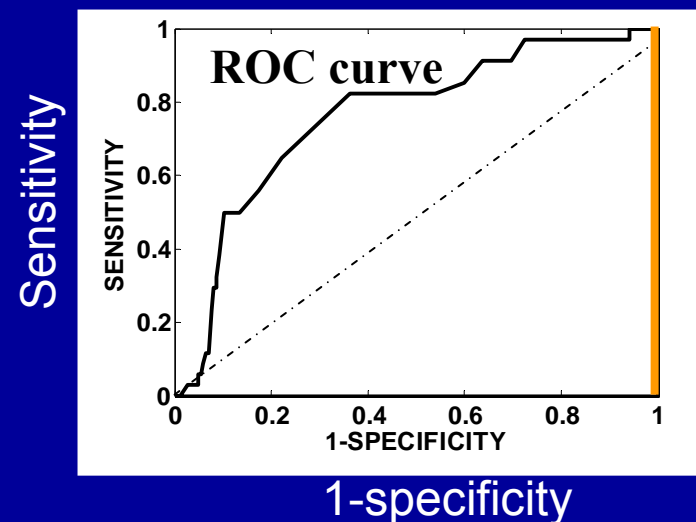
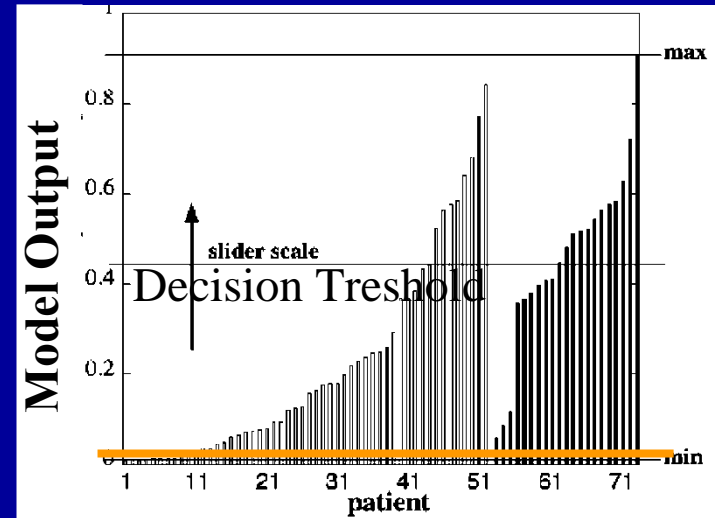
In general, the model was sequentially built by adding, substituting, deleting features.

# Receiver Operating Characteristics (ROC) Curve

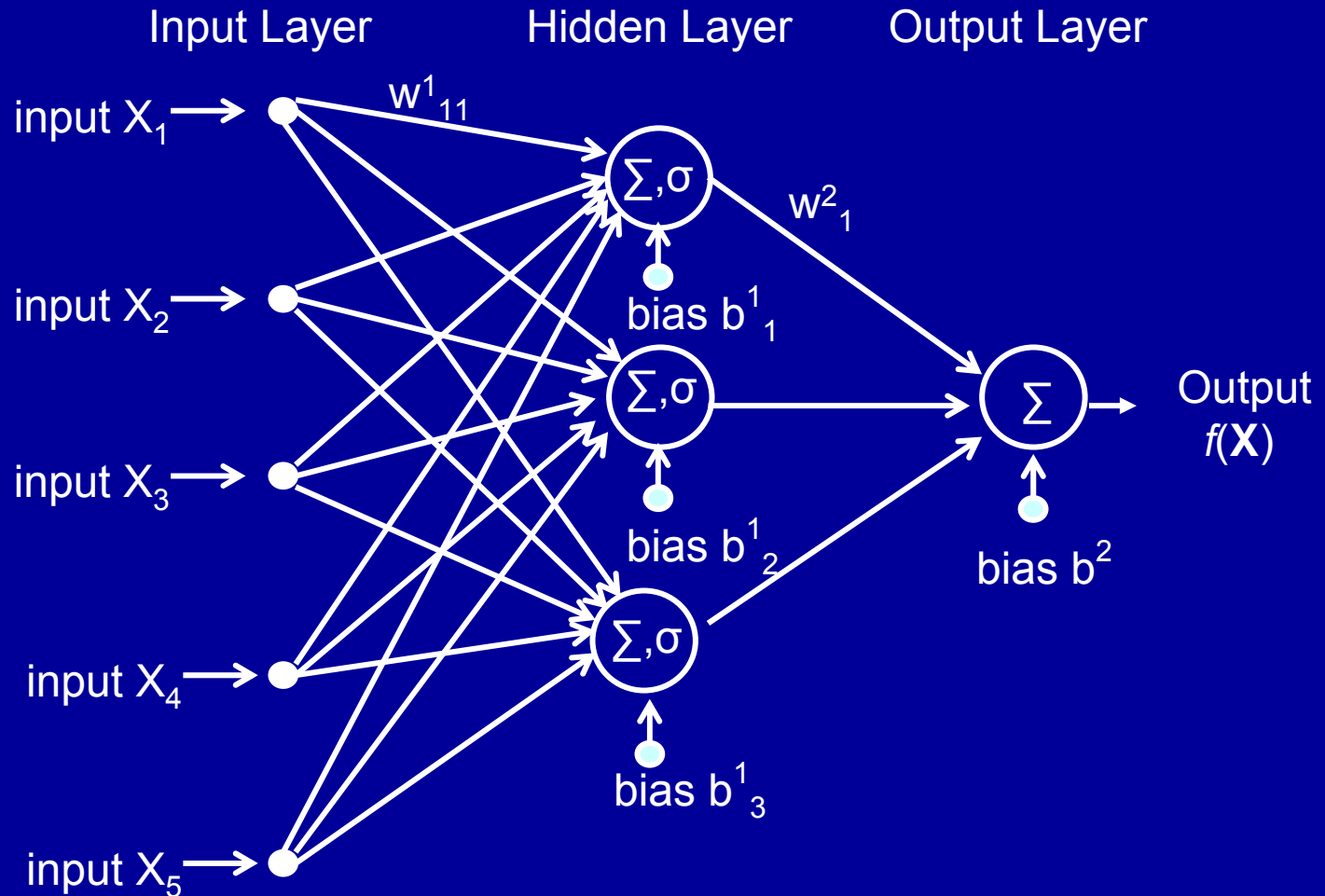
$$\text{Sensitivity} = \frac{\text{Model-identified True Positives}}{\text{True Positives}}$$

$$\text{Specificity} = \frac{\text{Model-identified True Negatives}}{\text{True Negatives}}$$

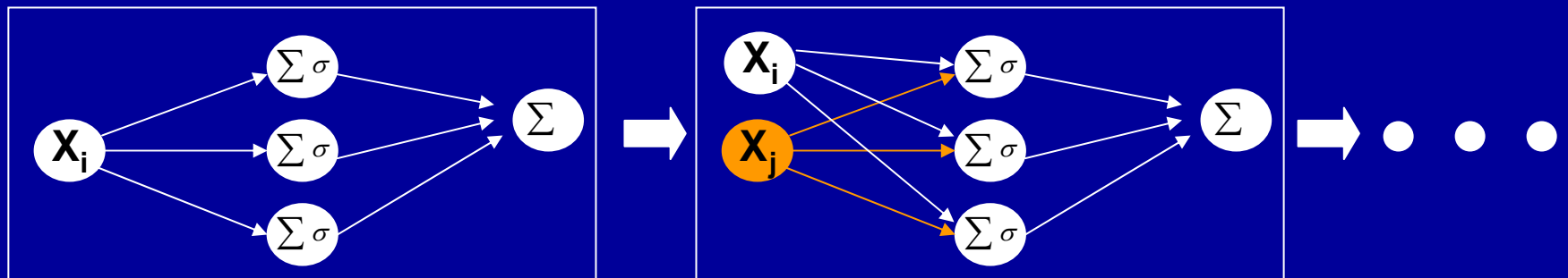
- Area under ROC curve
- Perfect model: area = 1
- Model as good as chance: area = 0.5



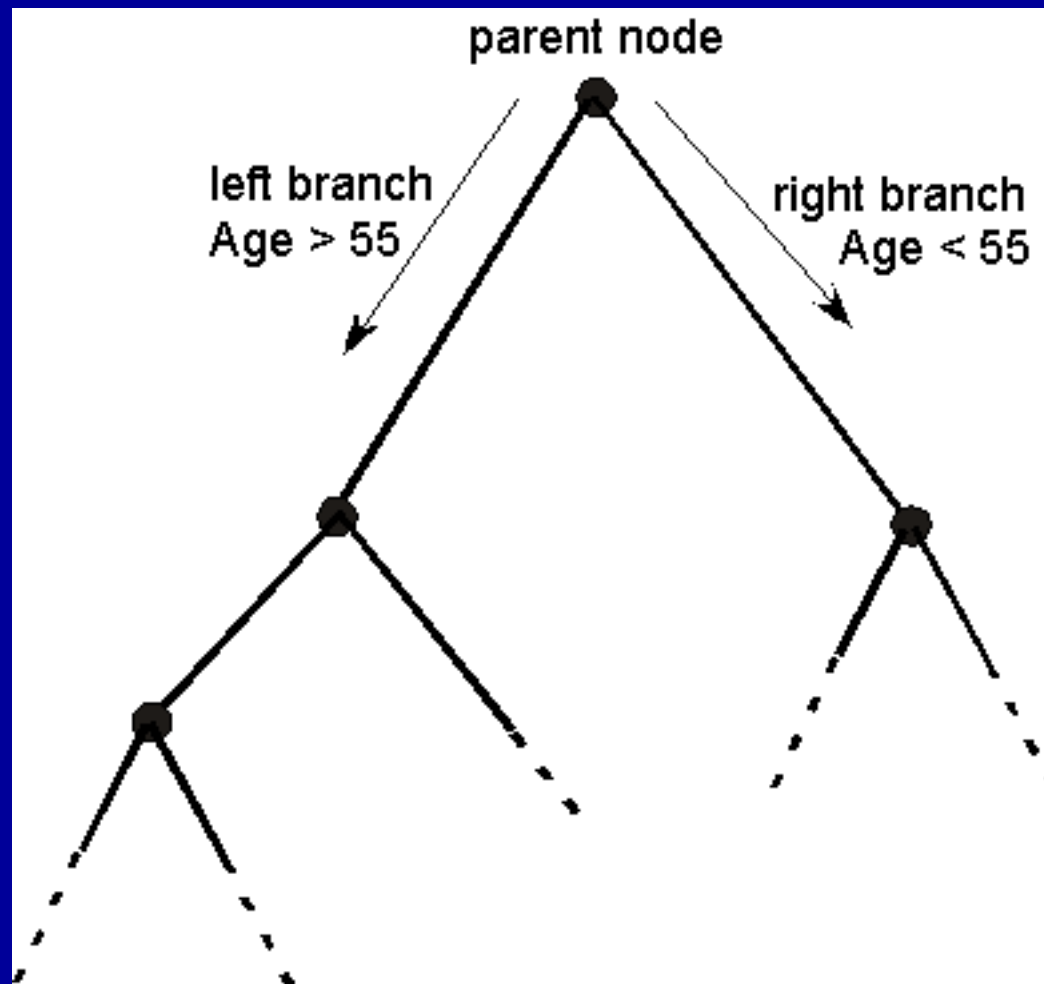
# Feed-Forward Neural Network



# Growing the Network: Construction Algorithm

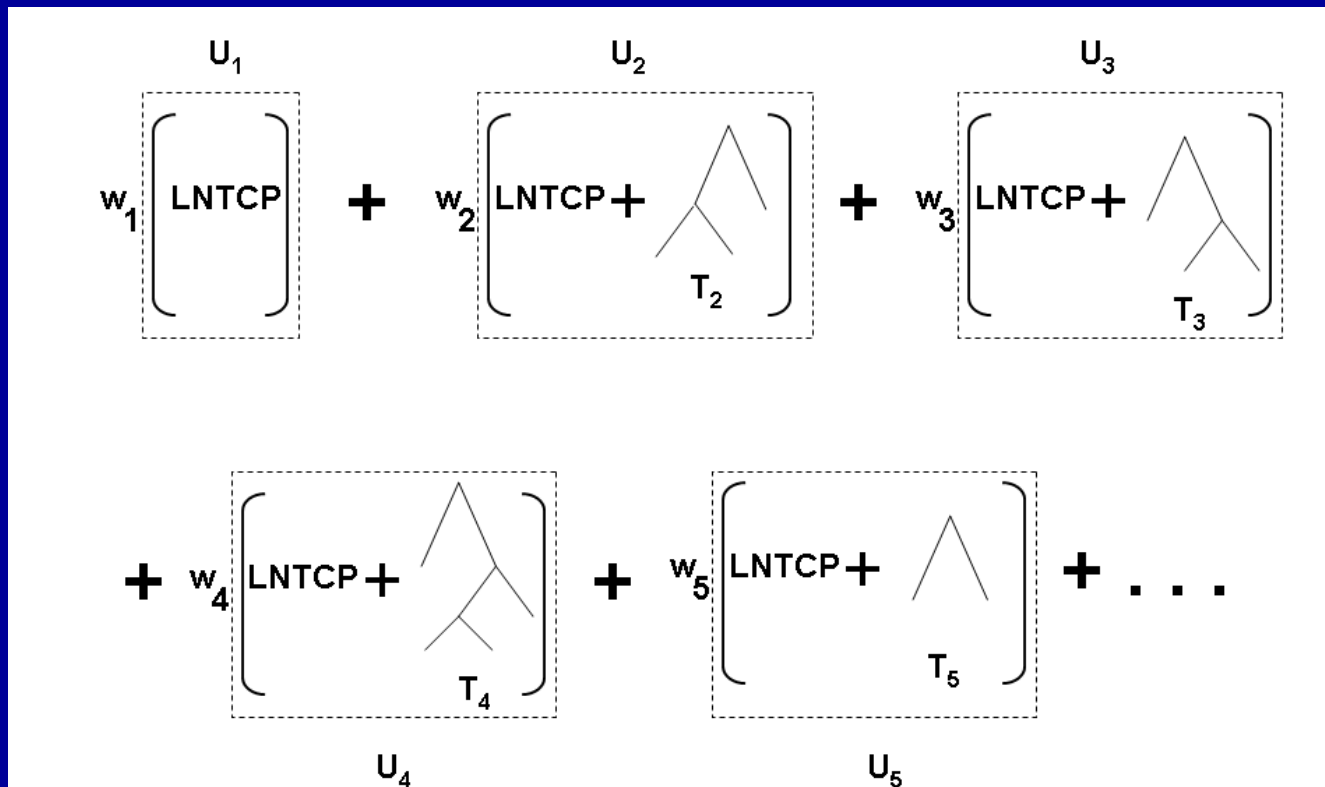


# Decision Trees

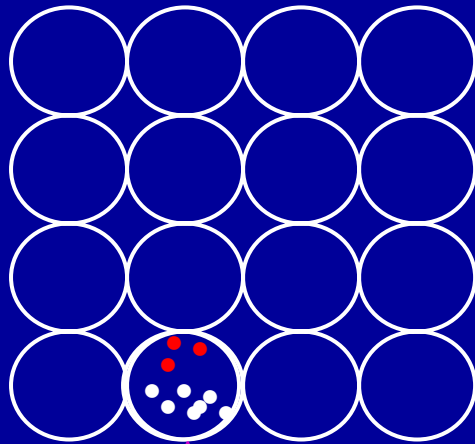


# Model Building

Model is built by combining weighted predictive units. Each predictive unit is composed of the Lyman NTCP added to a decision tree. Weighting is achieved by a statistical methodology termed AdaBoost.



# Self-Organizing Map (SOM)



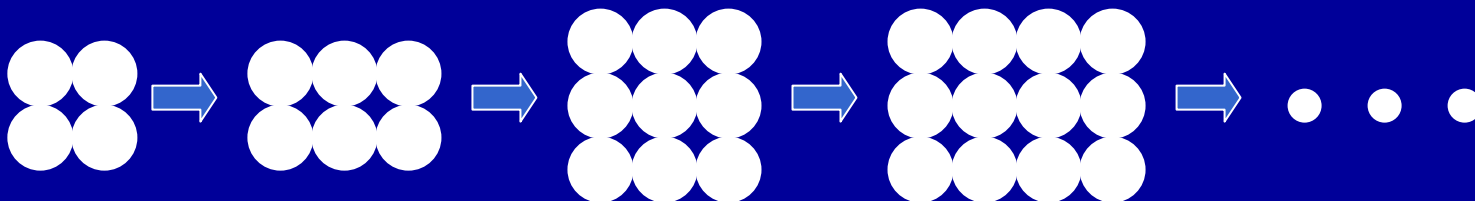
Mean lung dose ~ 15 Gy  
Chemotherapy  
Female

Patients with similar features are clustered in the same region.

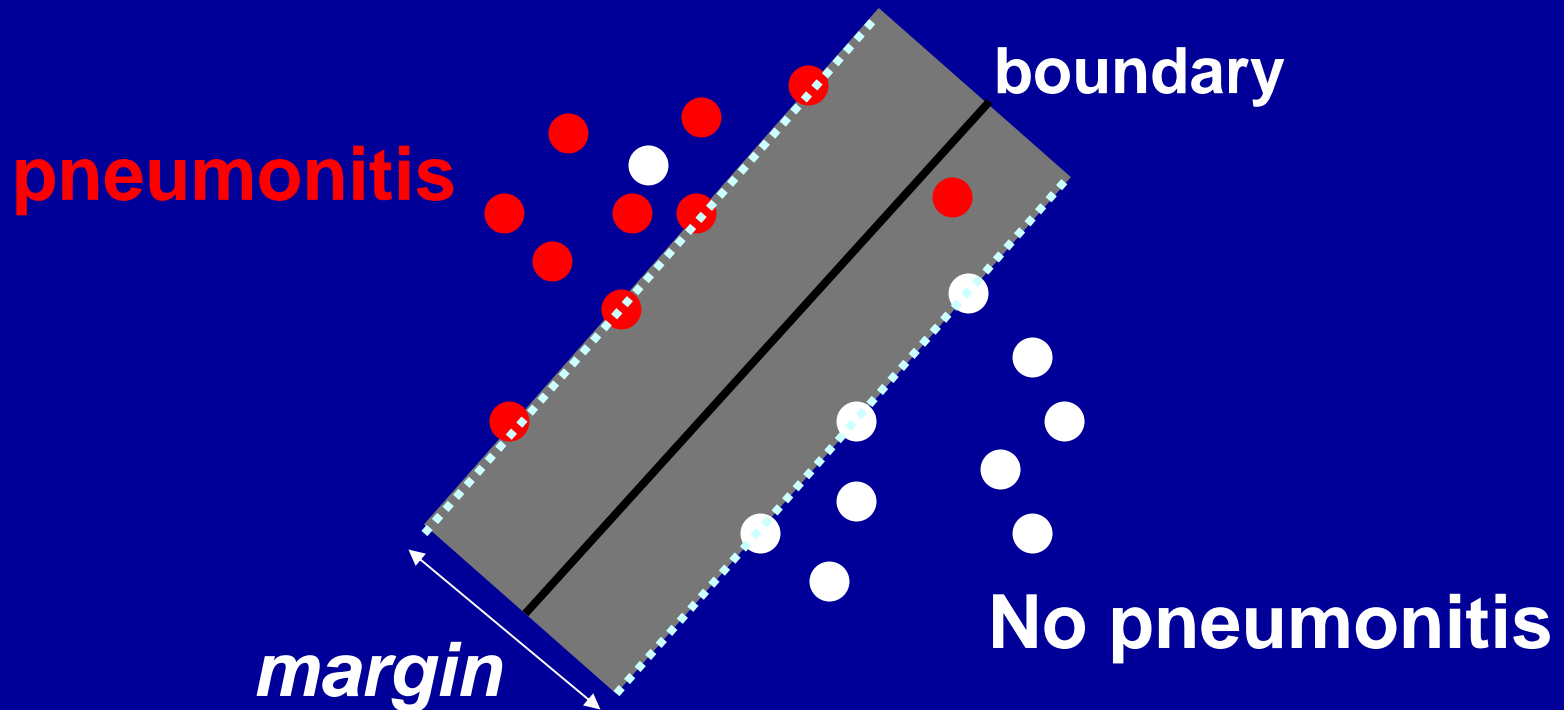
$$risk = \frac{N_p}{N_n + N_p}$$

3: pneumonitis •  
7: no pneumonitis •

$$Risk = \frac{3}{3 + 7} = 30\%$$



# Support Vector Machine (SVM)



Boundary is optimized to maximize the width of margin.  
SVM (unlike LDA) is not as affected by outliers.



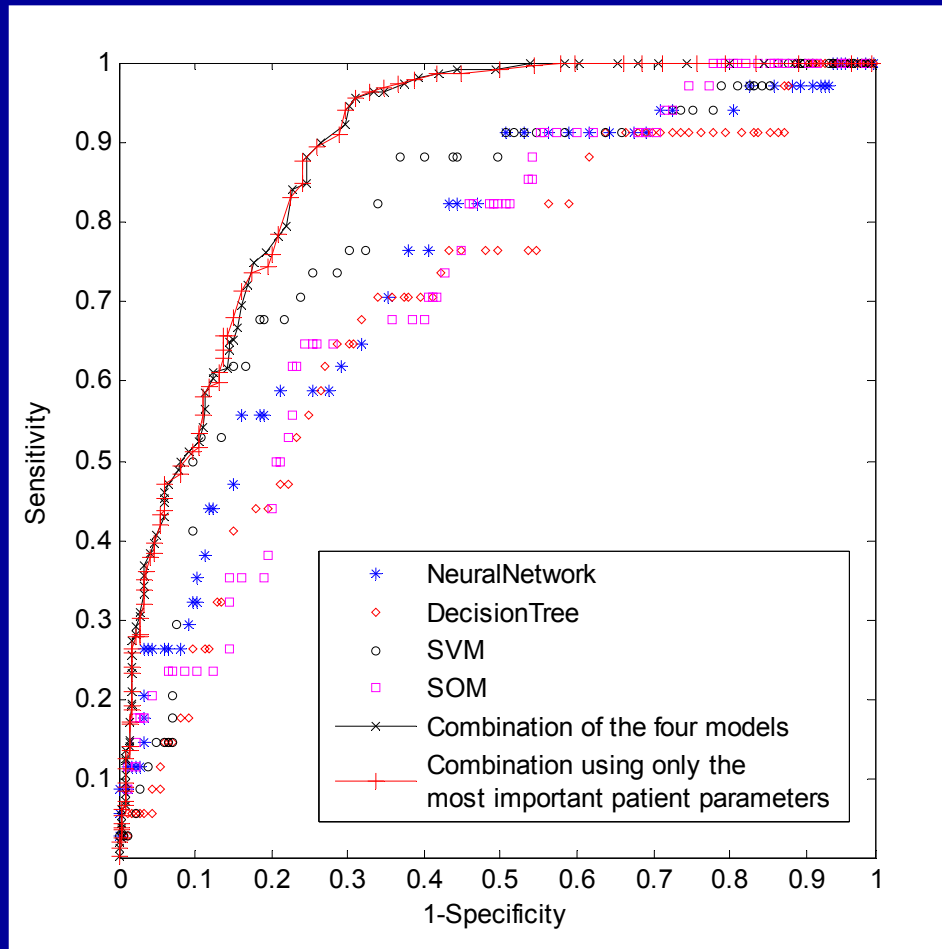
# Decision Fusion of Four Models

**Initially:**

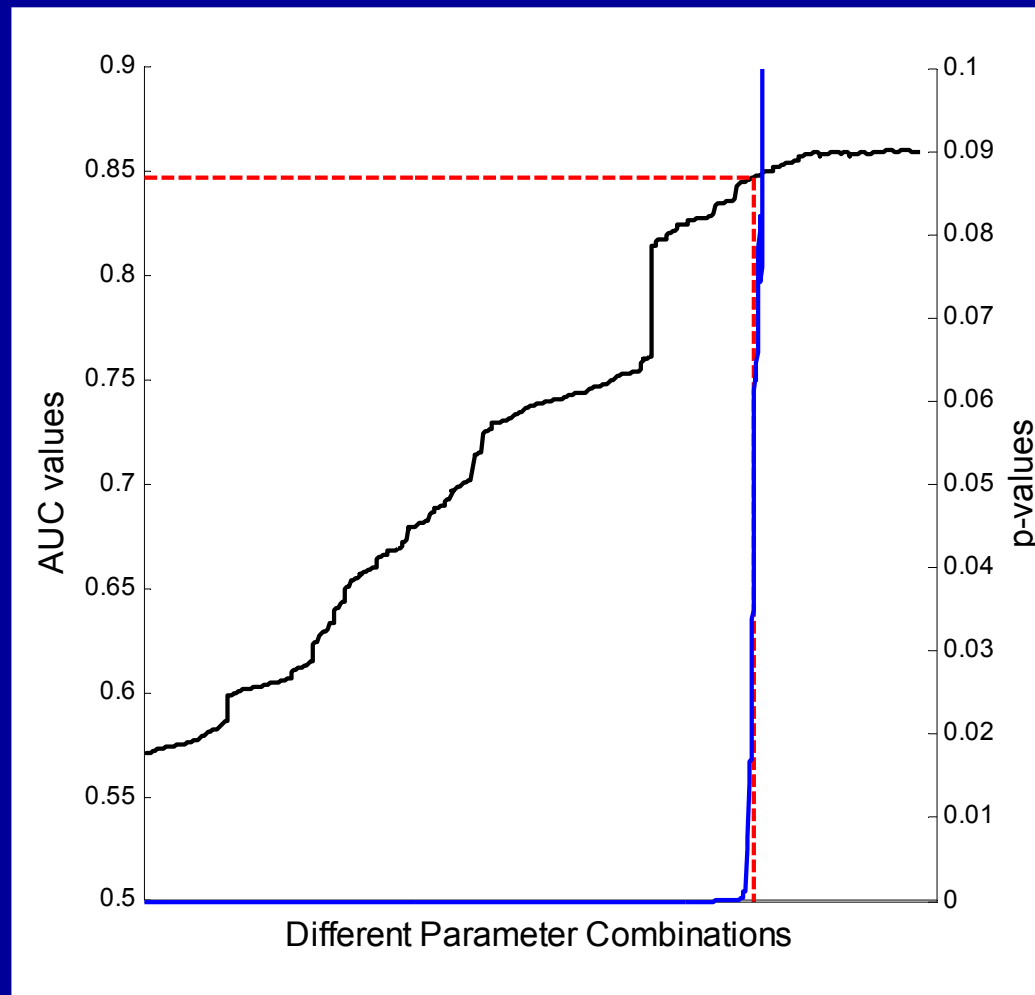
$$\text{Injury risk} = \frac{\text{NNET} + \text{DT} + \text{SOM} + \text{SVM}}{4}$$

**Refined Later:**

**Injury risk = Bayesian**



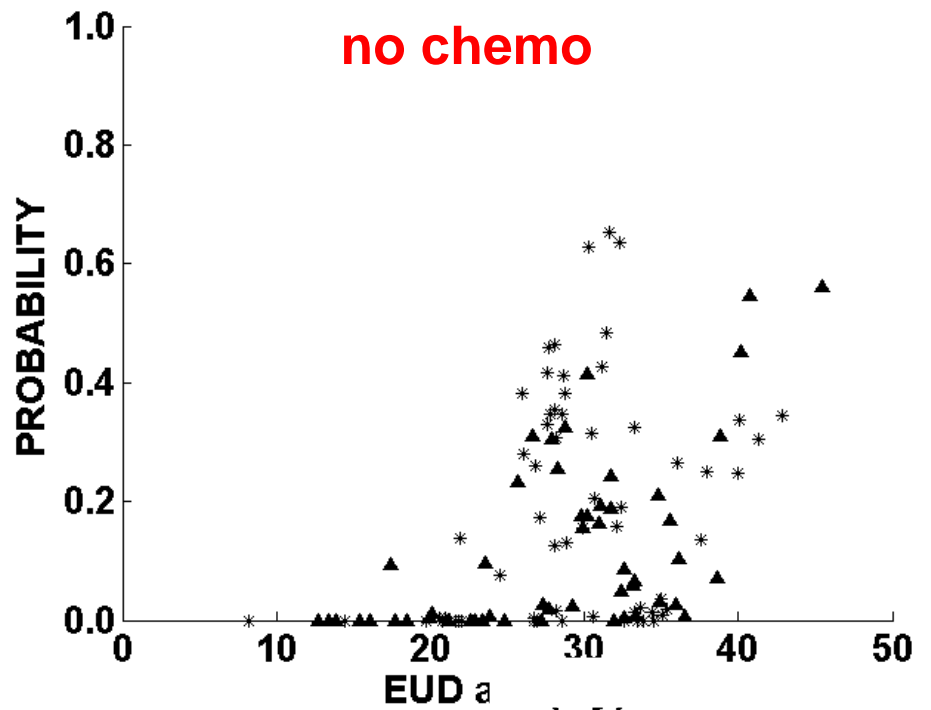
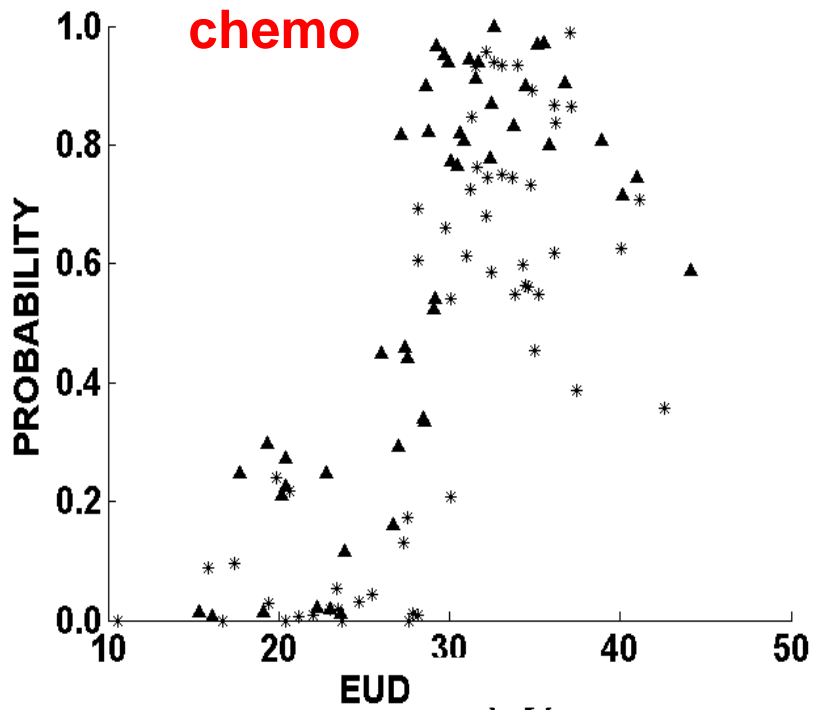
**Fused Decision Fusion AUC = 0.85**



**The different machine learning models constituting the fusion pick slightly different variables. However, the most common and important variables are sufficient to almost fully characterize system.**

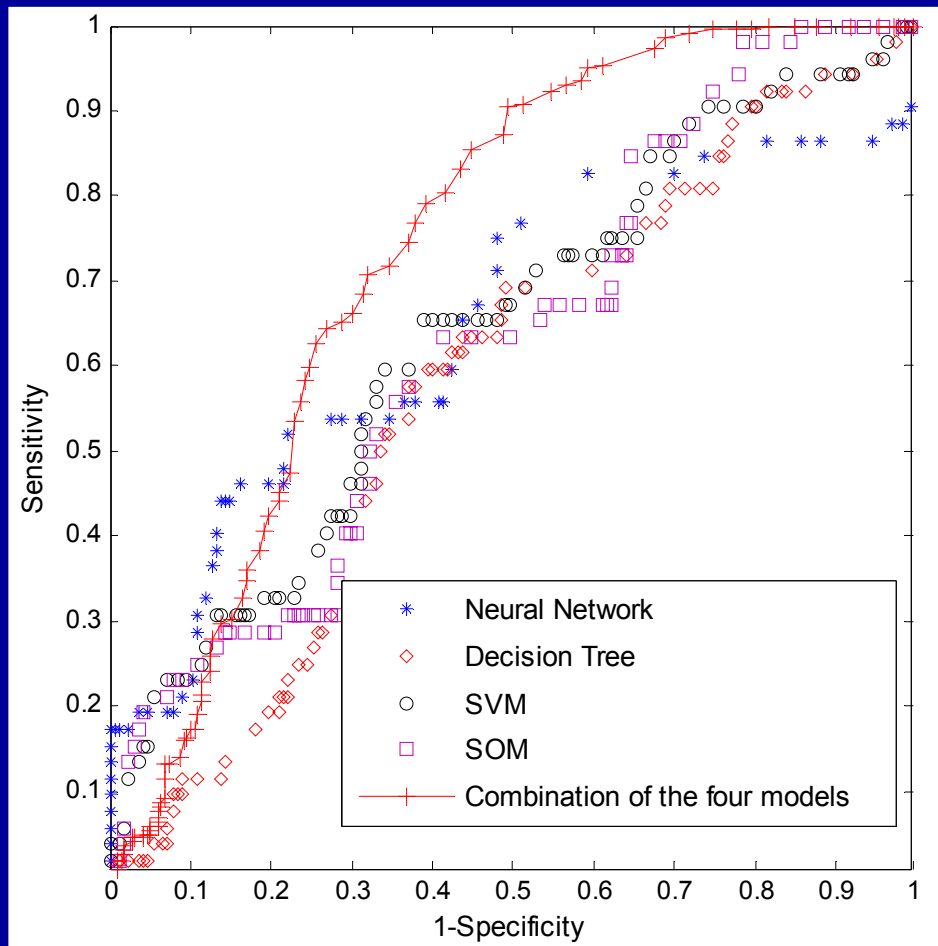
Parameters	% Frequency	Parameters	% Frequency
EUD <sub>1,3</sub>	50%	V <sub>30</sub>	49%
Chemotherapy prior to Radiotherapy	52%	Female gender	44%
Non-small Cell	34%	Squamous Cell	38%
Adenocarcinoma	41%	Small Cell Histology	39%
Central Tumor Location	42%	Inferior Tumor location	37%
Number of fractions (BID)	27%	FEV <sub>1</sub>	13%

**Fusion Identified Features: EUD<sub>1,3</sub>, V<sub>30</sub>, Tumor location, Histology, Female gender, and Chemotherapy schedule**

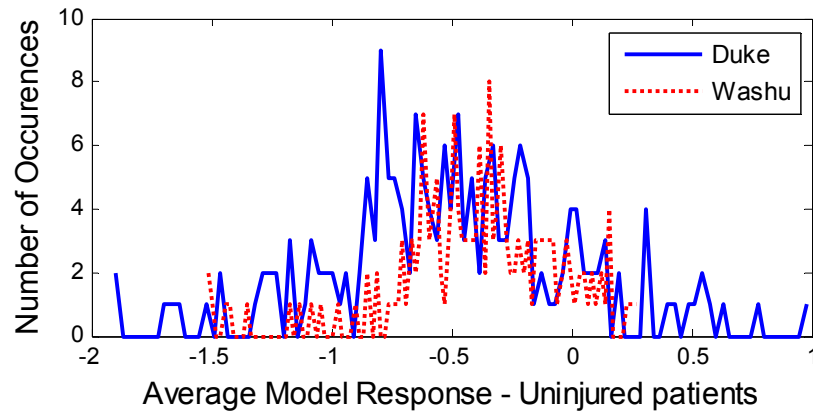
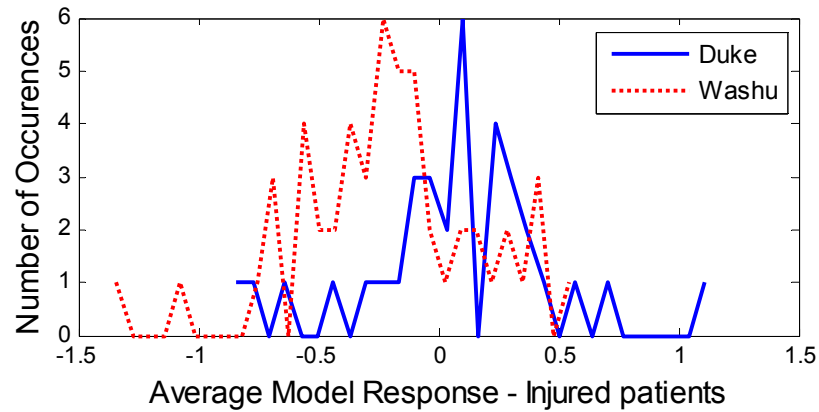


# Testing on External Dataset

Washington University dataset:  
219 patients (52 developed RP post-radiotherapy)



**AUC = 0.73**



**There is a rough equivalence between the Duke and WashU responses from the fusion model.**

# Conclusions

- Fusing machine learning models can reduce individual model bias, i.e., function like an almost bias-free model.
- Different models can predict different features. Fusion allows us to extract common “consensus” features.
- Testing on an external dataset demonstrates robustness.

**Ensuring reduced patient complications should be a primary safety goal!**