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**Course website** 

KYOTO UNIVERSITY

Statistical Machine Learning Theory

#### Semi-supervised, Active, and Transfer Learning

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#### Topics:

#### Semi-supervised, active, and transfer learning

- Semi-supervised learning
  - Weighted maximum likelihood estimation
  - Graph-based methods (e.g. label propagation)
  - Self-training
- Active learning
  - Uncertainty sampling
  - Estimated model change
- Transfer learning
  - Covariate shift using with weighted ML estimation
  - Shared parameters and domain specific parameters

#### Semi-supervised learning and active learning: Learning with labeled and unlabeled data

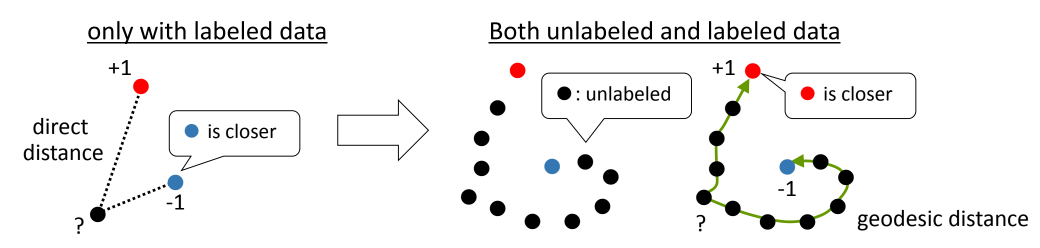
We have both labeled and unlabeled instances

-Labeled data: 
$$\{(x^{(1)}, y^{(1)}), ..., (x^{(N)}, y^{(N)})\}$$

- -Unlabeled data:  $\{x^{(N+1)}, ..., x^{(N+M)}\}$
- -Usually,  $N \ll M$
- Semi-supervised learning uses unlabeled data as well as labeled data
- Active learning
  - -has accesses to an oracle to give labels to unlabeled data
  - has to choose which unlabeled data to query next

# Role of unlabeled data in supervised learning: Information of the input data distribution

- Data generation process
  - -Input  $oldsymbol{x}$  is generated by input data distribution  $\mathcal{D}_{oldsymbol{x}}$
  - -Output y for x is generated by conditional distribution  $\mathcal{D}_{y|x}$
- lacktriangle Unlabeled data can be used for capturing  $\mathcal{D}_{oldsymbol{x}}$ 
  - Input data distribution, input space metric, or better representations



### **Semi-supervised Learning**

#### Semi-supervised learning problem: Learning with labeled and unlabeled data

We have both labeled and unlabeled instances

-Labeled data 
$$L = \{ (x^{(1)}, y^{(1)}), ..., (x^{(N)}, y^{(N)}) \}$$

- -Unlabeled data  $U = \{x^{(N+1)}, \dots, x^{(N+M)}\}$
- Estimate a deterministic mapping  $f: \mathcal{X} \to \mathcal{Y}$  (often with a confidence value) or a conditional probability P(y|x)

#### Typical approaches of semi-supervised learning: Learning with labeled and unlabeled data

- Weighted maximum likelihood estimation
- Graph-based learning
- Self-training
- Clustering
- Generative models

### Weighted maximum likelihood: Estimate input distribution to weight labeled instances

The original goal of ML estimation is to maximize

$$E_{x,y}[\log P(y|x)] = \int \log P(y|x) dp(x) dp(y|x) \approx \frac{1}{N} \sum_{i=1}^{N} \log P(y^{(i)}|x^{(i)})$$

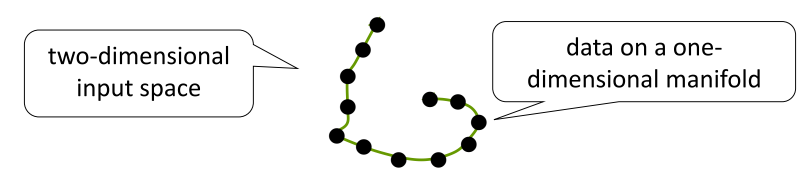
- -Each training data instance is equally weighted
- Weighted maximum likelihood: Each training data instance is weighted according to p(x)

maximize 
$$\sum_{i=1}^{N} p(\mathbf{x}^{(i)}) \log P(\mathbf{y}^{(i)}|\mathbf{x}^{(i)})$$

-p(x) is estimated using unlabeled data (but not practical)

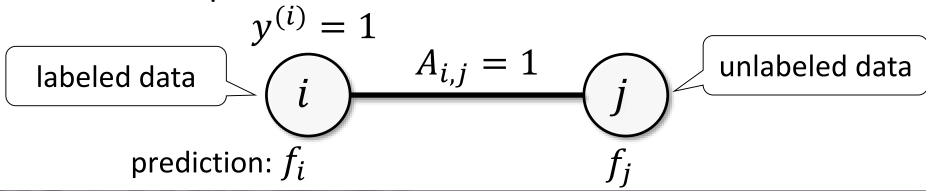
#### Graph-based method: Capture intrinsic shape of input space

- Basic idea: construct a graph capturing the intrinsic shape of the input space, and make predictions on the graph
- Assumption: Data lie on a manifold in the feature space
- The graph represent adjacency relationships among data
  - -K-nearest neighbor graph (e.g.  $A_{i,j} = \{0, 1\}$ )
  - -Edge-weighted graph with e.g.  $A_{i,j} = \exp \left(-\| \ \pmb{x}^{(i)} \pmb{x}^{(j)} \|_2^2\right)$



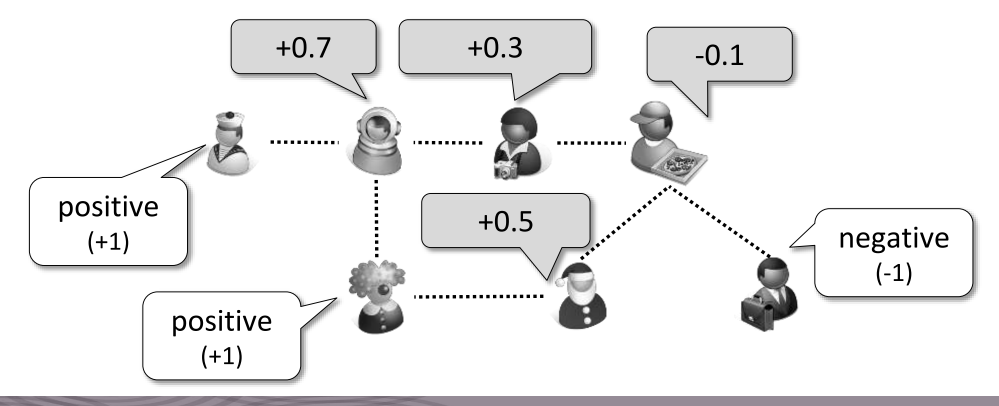
#### Label propagation: Simple graph-based method

- Basic idea: Adjacent instances tend to have the same label
  - Note that we have test instances (i.e. transductive setting)
- minimize<sub>f</sub>  $\sum_{i=1}^{N} (f_i y^{(i)})^2 + \gamma \sum_{i,j} A_{i,j} (f_i f_j)^2$ 
  - -1st term: (squared) loss function to fit to labeled data
  - 2nd term: regularization function to make adjacent nodes to have similar predictions



# Illustrative example of label propagation: Infection prediction on social network

- Predict if people are infected by some disease
  - Test results are known for some people
  - Infections spread over social networks



#### Self-training: Believe what you believe

- Procedure:
- 1. Initialization: train a classifier using labeled dataset L
- 2. Use the classifier to assign temporary labels to unlabeled dataset  $\it U$
- 3. Train a classifier using L and U (with the temporary labels)
- 4. Return to Step 2
- For probabilistic classifier, use the weighted ML estimation:

$$\text{maximize} \sum_{i \in L} \log p(y^{(i)}|\boldsymbol{x}^{(i)}) + \sum_{i \in U} \sum_{\hat{y}} p(\hat{y}|\boldsymbol{x}^{(i)}) \log p(\hat{y}|\boldsymbol{x}^{(i)})$$

#### **Active Learning**

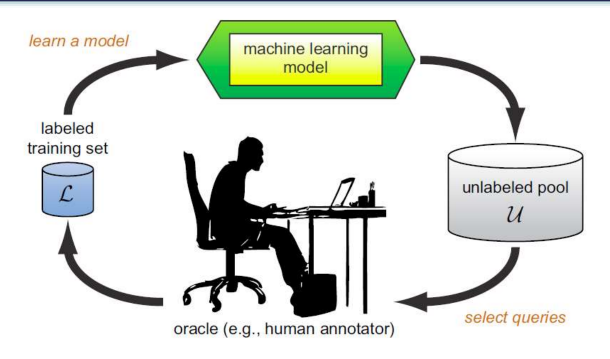


Figure 1: The pool-based active learning cycle.

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#### Active learning: Learning with a label oracle

- lacksquare Start with only unlabeled data  $U = \{x^{(1)}, ..., x^{(N)}\}$
- At each round, an active learner can query an unlabeled instance to be labeled by an oracle
  - then update the predictor using current labeled (and unlabeled) data
- An active learning algorithm determines the query strategy specifying which unlabeled instance should be queried next

### Active learning query strategies: Choose the most "informative" instance

- Basic idea: Query the instance whose label is the most informative
- Several basic strategies to choose "informative" instance
  - -Query the instance with the most uncertain label
  - Query the instance which will gives the largest expected model change

**—** . . .

#### Uncertainty sampling: Query the instance with the most uncertain label

- In a linear classifier  $f(x) = \text{sign}(w^T x)$ ,  $|w^T x|$  indicates "confidence level" of the prediction
  - -For multi-class classification,
    - use  $\max_k \mathbf{w}^{(k)\intercal} \mathbf{x}$
    - or, margin  $\max_k \mathbf{w}^{(k)\intercal} \mathbf{x}$  secondbest<sub>k</sub>  $\mathbf{w}^{(k)\intercal} \mathbf{x}$
  - -For probabilistic classifiers, the entropy  $\sum_{y} -P(y|x) \log P(y|x)$  is used as an uncertainty measure
- Query  $x^{(i)}$  with the lowest confidence/highest uncertainty

### Differences among confidence level, margin, and entropy [Settles, 2010. page 14]

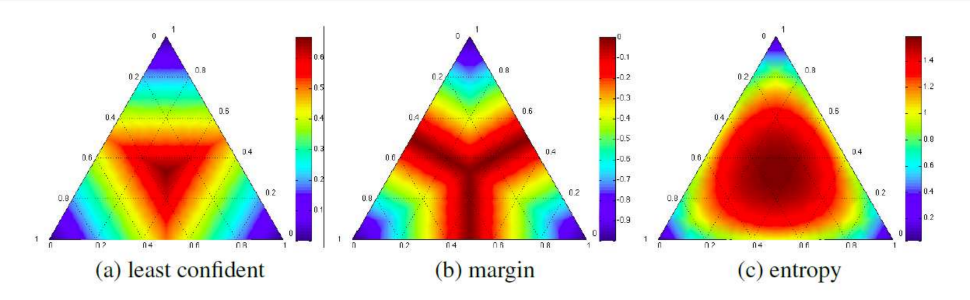


Figure 5: Heatmaps illustrating the query behavior of common uncertainty measures in a three-label classification problem. Simplex corners indicate where one label has very high probability, with the opposite edge showing the probability range for the *other* two classes when that label has very low probability. Simplex centers represent a uniform posterior distribution. The most informative query region for each strategy is shown in dark red, radiating from the centers.

## limitation of uncertainty sampling: Uncertainty sampling is based on local information

- Querying the least confident instance cares only about the local information
- Obtaining one labeled instance can make an impact on the whole model
- We should take the amount of the "impact" of a label into account

#### Expected model change:

#### Query the instance which gives the largest model change

- Choose instance x which gives the largest (expected) gradient of the objective function:  $\sum_{y} -P(y|x) \parallel \nabla_{w} J(L \cup (x,y)) \parallel$ 
  - Assume gradient-based learning methods are used
    - e.g. gradient descent  $\mathbf{w}^{\text{new}} \leftarrow \mathbf{w}^{\text{old}} \gamma \nabla_{\mathbf{w}} J(L \cup (\mathbf{x}, \mathbf{y}))$  when new labeled instance  $(\mathbf{x}, \mathbf{y})$  is added to L
- Choose an instance which gives the largest information gain

$$\sum_{y} -P(y|\mathbf{x}) \sum_{i=N+1}^{N+M} \sum_{y'} P_{\mathbf{w}^{\text{new}}}(y'|\mathbf{x}^{(i)}) \log P_{\mathbf{w}^{\text{new}}}(y'|\mathbf{x}^{(i)})$$

 $-P_{w^{\text{new}}}$ : model after update with new labeled data (x, y)

### **Transfer Learning**

#### Transfer learning:

#### Training and test data come from different distributions

- Training dataset and test dataset are sampled from different distributions
- In the standard settings, an input x is sampled from  $\mathcal{D}_x$ , and an output y is sampled from  $\mathcal{D}_{y|x}$  (in both training and test)
- In transfer learning,
  - -Training data come from  $\mathcal{D}_x^{\mathrm{train}}$  and  $\mathcal{D}_{y|x}^{\mathrm{train}}$
  - -Test data come from  $\mathcal{D}_x^{\mathrm{test}}$  and  $\mathcal{D}_{y|x}^{\mathrm{test}}$
- Example: Domain adaptation
  - -Classification of general text documents and medical texts

## Covariate shift: Input distributions are different

Covariate shift: only the input distributions are different

$$-\mathcal{D}_{x}^{\text{train}} \neq \mathcal{D}_{x}^{\text{test}}$$

- $-\mathcal{D}_{y|x}^{\text{train}} = \mathcal{D}_{y|x}^{\text{test}}$ : conditional distributions are the same
- -Training dataset is labeled and test dataset is unlabeled
- Occurs when sampling of labeled data is constrained
  - Labels are obtained only from the targets to which some actions are taken (e.g. responses to direct mails)
  - Labels can only be taken in controlled environments (e.g., in-vitro experiments)
  - Active learning controls the training distribution

# Maximum likelihood learning under covariate shift: Maximize likelihood for test input distribution

- The distribution on which we want to work well is the test input distribution  $p^{\text{test}}(\mathbf{x})$
- In maximum likelihood estimation, we want to maximize

$$E_X^{\text{test}}[\log P(y|\mathbf{x})] = \int p^{\text{test}}(\mathbf{x}) \log P(y|\mathbf{x}) d\mathbf{x}$$

- -Note that the expectation is taken over  $p^{\text{test}}(x)$
- However, we can not directly evaluate the objective function
  - We do not have label information for test dataset

#### Covariate shift learning only with training labels: Weighted maximum likelihood with density ratio

Use the importance sampling

$$E_X^{\text{test}}[\log P(y|\mathbf{x})] = \int \frac{p^{\text{test}}(\mathbf{x})}{p^{\text{train}}(\mathbf{x})} p^{\text{train}}(\mathbf{x}) \log P(y|\mathbf{x}) d\mathbf{x}$$

$$\approx \frac{1}{N} \sum_{i=1}^{N} \frac{p^{\text{test}}(\mathbf{x}^{(i)})}{p^{\text{train}}(\mathbf{x}^{(i)})} \log P(y^{(i)}|\mathbf{x}^{(i)})$$

$$= \frac{1}{N} \sum_{i=1}^{N} \omega(\mathbf{x}^{(i)}) \log P(\mathbf{y}^{(i)} | \mathbf{x}^{(i)})$$
 training data  $(\mathbf{x}^{(i)}, \mathbf{y}^{(i)})$  is weighted with  $\omega(\mathbf{x}^{(i)})$ 

weighted with  $\omega(x^{(i)})$ 

-Weighted ML estimation with weight  $\omega(x^{(i)}) = \frac{p^{\operatorname{test}}(x^{(i)})}{\operatorname{ntrain}(x^{(i)})}$ 

#### Practical considerations:

#### Density ratio estimation and adaptive importance

- Estimation of the density ratio  $\omega(x) = \frac{p^{\operatorname{test}(x)}}{p^{\operatorname{train}(x)}}$  is required
  - -Density estimation of  $p^{\mathrm{test}}$  and  $p^{\mathrm{train}}$
  - –Some approaches directly estimate  $\omega$
- Adaptive importance weighted ML estimation:

-Practically 
$$\omega^{\lambda}(\mathbf{x}^{(i)}) = \left(\frac{p^{\operatorname{test}}(\mathbf{x}^{(i)})}{p^{\operatorname{train}}(\mathbf{x}^{(i)})}\right)^{\lambda}$$
  $(0 \le \lambda \le 1)$  works better

# Transfer learning of different conditional distributions: Adaptation to model changes

Transfer learning of different conditional distributions

$$-\mathcal{D}_{y|x}^{\text{train}} \neq \mathcal{D}_{y|x}^{\text{test}}$$

- $-\mathcal{D}_x^{\mathrm{train}} = \mathcal{D}_x^{\mathrm{test}}$ : Input distributions are the same
- -Labels are available in both training and test datasets
- Adaptation to changes of predictive models
  - Transfer knowledge from a general task to a specific task (and vice versa)
  - Model changes over time

#### A simple approach to model change adaptation: Shared parameters and domain specific parameters

- Assume linear models (e.g.  $f(x) = sign(w^T x)$ )
  - –The source domain model has  $w^{(s)}$ , while the target domain model has  $w^{(t)}$
- The models have shared parts and domain specific parts
  - –Source domain model  $\mathbf{w}^{(\mathrm{s})} = \mathbf{v}^{(0)} + \mathbf{v}^{(\mathrm{s})}$
  - -Target domain model  $\boldsymbol{w}^{(t)} = \boldsymbol{v}^{(0)} + \boldsymbol{v}^{(t)}$
  - -Equivalent to setting  $\mathbf{w}=(\mathbf{v}^{(0)},\mathbf{v}^{(s)},\mathbf{v}^{(t)})$  and  $\widetilde{\mathbf{x}}=(\mathbf{x},\mathbf{x},\mathbf{0})$  for the source domain and  $\widetilde{\mathbf{x}}=(\mathbf{x},\mathbf{0},\mathbf{x})$  for the target domain
- Standard classification methods are applicable