

# 1 Maschine Learning

Algorithm learns class of tasks, measured by loss function, from experience.

**supervised learning** learn  $h : \Delta^* \rightarrow \Sigma^*$ ,  $h = t$ ; example:  $(x, y) \in \Delta^* \times \Sigma^*$ ,  $t(x) = y$ .

**unsupervised learning** learn  $h : \Delta^* \rightarrow \Sigma^*$ ,  $\ker(h) = \ker(t)$ ; example:  $x \in \Delta^*$ .

**reinforcement learning** learn strategy based on feedback from environment.

## 2 Supervised Learning

- model function  $t : \mathcal{M} \rightarrow \mathcal{R}$

-  $\text{supp}(t) = \{m \in \mathcal{M} \mid t(m) \neq 0\}$

-  $\bar{m} \in \text{supp}(t) \Leftrightarrow t(\bar{m}) = 1$

**Hypothesis** of A: potential result of A

**Hypothesis space**  $\mathcal{H}_A$  of A: set of all hypotheses

**h fits D** if  $h(x_i) = y_i$  for all  $(x_i, y_i) \in D$

**Version space**  $\mathcal{V}_A(D)$  of A: all hypotheses that fit D

**Inductive bias** of A: set of assumptions that A uses to predict outputs of unseen data

### 2.1 Conjunctive Clause

$\theta = (\theta_1, \dots, \theta_k), \theta_i \in M_i \cup \{\star, \perp\}$

-  $\theta_\perp = (\perp, \dots, \perp)$  most specific

-  $\theta_\star = (\star, \dots, \star)$  most general

-  $\text{supp}(h_{\theta_\perp}) = \emptyset, \text{supp}(h_{\theta_\star}) = \mathcal{M}$

-  $h_{\theta_\perp} = h_{(\theta_1, \dots, \perp, \dots, \theta_k)}$ ...

induced hypothesis  $h_\theta(m_1, \dots, m_k) = 1$  if  $\forall i : \theta_i \in \{m_i, \star\}$  else 0

$h \preceq h'$  if  $\text{supp}(h) \subseteq \text{supp}(h')$ .  $h$  is more specific (less general) than  $h'$

**Find-S Algorithm** finds most specific conjunctive clause that fits D

1. Start with  $\theta_\perp = (\perp, \dots, \perp)$
2. iterate over POSITIVE examples
3. min-generalize  $\theta$  to fit example
4.  $\perp \rightarrow a, a \rightarrow \star$

- maximal general hypothesis:

1. start at  $\theta_\star = (\star, \dots, \star)$
  2. exclude every negative example
  3.  $(\star, \dots) \rightarrow \{(b, \dots), (c, \dots)\}$
- If  $\mathcal{V}_A(D) \neq \emptyset$ , Find-S finds  $h \in \mathcal{V}_A(D)$

**disjunctive normal form**  $\Theta = \{\theta_1, \dots, \theta_m\}$

'finite set of conjunctive clauses'

induced hypothesis  $h_\Theta(\bar{m}) = 1$  if  $\exists \theta \in \Theta : h_\theta(\bar{m}) = 1$  else 0

-  $\text{supp}(\Theta) = \bigcup_{\theta \in \Theta} \text{supp}(\theta)$

- can represent all boolean functions

### Boundary sets of version space

maximally general hypotheses  $V_A^\top(D) = \{h \in V_A(D) \mid \nexists h' \in V_A(D) : h \prec h'\}$

maximally specific hypotheses  $V_A^\perp(D) = \{h \in V_A(D) \mid \nexists h' \in V_A(D) : h' \preceq h\}$

-  $h \in V_A^\perp(D)$  maximal, weil:  $\forall x \in M \setminus \text{supp}(h) : \text{supp}(h) \cup \{x\} \notin \text{supp}(V_A(D))$

Theorem:  $V_A(D) = \{h \in H_A \mid \exists h_T \in V_A^\top(D), \exists h_\perp \in V_A^\perp(D) : h_\perp \preceq h \preceq h_T\}$

->  $V_A(D)$  det. by  $V_A^\top(D)$  and  $V_A^\perp(D)$

- only 1 lower bound (in  $V_A^\perp(D)$ ), potentially multiple upper bounds (in  $V_A^\top(D)$ )

### Candidate Elimination Algorithm

Output: DNF for  $V_A^\top(D)$  and  $V_A^\perp(D)$

1.  $S_\perp = \{\theta_\perp\}, S_\star = \{\theta_\star\}$

2. for  $1 \leq i \leq n : y_i = 1$  (pos. xmpls)

1. keep only fitting h from  $S_\star$

2.  $\forall \theta \in S_\perp : h_\theta(x_i) = 0$

- remove  $\theta$ , add all min generalizations  $\theta'$  of  $\theta$  that fit  $x_i$  to  $S_\perp$

3. keep only most specific h in  $S_\perp$

3. for  $1 \leq i \leq n : y_i = 0$  (neg. xmpls)

1. keep only fitting h from  $S_\perp$

2.  $\forall \theta \in S_\star : h_\theta(x_i) = 1$

- remove  $\theta$ , add all min specializations  $\theta'$  of  $\theta$  that fit  $x_i$  to  $S_\star$ , for which a more specific  $\theta_\perp \in S_\perp$  exists!

3. keep only most general h in  $S_\star$

-  $V_A^\top = \{h_\theta \mid \theta \in S_\star\}, V_A^\perp = \{h_\theta \mid \theta \in S_\perp\}$

- Concept indentified if:  $S_\perp = S_\star$  and  $|S_\star| = 1$ .  $V_A(D) = \emptyset$  if  $S_\perp = \emptyset \vee S_\star = \emptyset$

### 2.2 Decision Trees

**Splitting**  $\Pi = \{M_1, \dots, M_p\}$  is finite partition of (sub)feature Space  $\mathcal{M}'$

- induces splitting of  $\{1, \dots, n\}$  into  $I_{D'}(M_1), \dots, I_{D'}(M_p)$  (sets of indices)

- monotonic splits: based on 1 feature

- simple split: monotonic, into all realizations  $M = \{\bar{m} \in M \mid m_1 = a, \dots\}$

- binary split: monotonic, into 2 sets  $M = \{\bar{m} \in M \mid m_1 \in A\} \cup \{\bar{m} \in M \mid m_1 \notin A\}$

- induced hypothesis  $h_T(\bar{m}) = T(v)$ , where  $v$  is unique leaf s.t.  $\bar{m} \in M_v$

- simple decision trees can represent all hypotheses

### Decision Tree Quality Measures

- Number of leaves
- Height (max number of constraints to check)
- External path length (sum of all path lengths from root to leaf)

- Weighted external path length (sum of all path lengths from root to leaf, weighted by number of examples classified in that leaf)

Theorem: Given D and bound b, its NP hard to decide existence of decision tree T s.t.  $h_T$  fits D and T has ext. p.l.  $\leq b$

- Majority Class  $\text{Maj}_D(M')$  maj.  $r \in R$

- Number of Misclassifications:  $Err_D(M', r)$  in feature subspace  $M'$  with majority class  $r$

-  $Err_D(T)$ : sum up all  $Err_D(M_v, T(v))$

**Pure Node** v if  $Err_D(M_v, T(v)) = 0$

- class distribution  $p_D^{M'}(r) : p(r)$  in  $M'$

- **Impurity Function**  $\iota : [0, 1]^R \rightarrow \mathbb{R}$  if

- $\iota(p)$  is minimal  $\forall p : p(r) = 1$
- $\iota$  symmetric in classes
- $\iota$  is maximal for uniform distr.

gets probability distribution as input

1.  $\iota(p) = 1 - \max_{r \in R} p(r)$

2. Entropy  $H(p) = -\sum_{r \in R} p(r) \log_2 p(r)$

3. Gini Impurity  $G(p) = 1 - \sum_{r \in R} p(r)^2$

- Impurity of  $M'$  is  $\iota_D(M') = \iota(p_D^{M'})$

### Impurity Reduction of splitting

$\Pi = \{M'_1, \dots, M'_p\}$  of  $M'$  is:

$\iota_D(\Pi) = \iota_D(M') - \sum_{i=1}^p \frac{|\iota_D(M'_i)|}{|\iota_D(M')|} \iota_D(M'_i)$

### Tree Construction for $M' \subseteq M$

1. if no elements in  $M'$ : new leave  $v$ :  $T(v) = \text{Maj}_D(M)$

2. if  $\iota(M') \leq \epsilon$ : new leaf  $v$ :  $T(v) = \text{Maj}_D(M')$

3. else: select split  $\Pi$  of  $M'$  with maximal impurity reduction

- strict imp. fct.: concave at every point

-  $\iota_D(\Pi) \geq 0 \forall \Pi$  and strict imp. fct.  $\iota$

**ID3** simple D.T., monothetic simple splits, impurity function: entropy.

inductive bias: local optimization (greedy)

**CART** D.T., binary splits, impurity function: Gini impurity

- true loss of  $h \in H_A$ : misclassifications:  $l^*(h) = \sum_{\bar{m} \in M} (1 - \delta_{h(\bar{m}), t(\bar{m})})$

-  $h$  **overfits** D if  $\exists h' \in H_A :$

$l(h, D) < l(h', D)$  and  $l^*(h) > l^*(h')$

when: training data: noisy, small, biased

- Training Data: optimize loss here

- Validation: optimize hyperparameters

- Test Data: final estimation (true loss)

-  $h$  **overfits**  $(D, D_V)$  if  $\exists h' \in H_A :$

$l(h, D) < l(h', D) \wedge l(h', D_V) < l(h, D_V)$

true loss of  $h$  estimated by  $l(h, D_V)$

Countermeasures to overfitting:

- increase data quality/quantity
- early stopping (no more splits)
- threshold large -> omits useful splits
- threshold small -> Large Tree
- regularization (penalize model complexity in training process)

**Pruning**: turn inner node  $v$  into leave with label  $\text{Maj}(M_v)$

### D.T. pruning Algorithm

1. given fully trained D.T.

2. prune every inner node als long as pruning doesnt increase validation loss:  $l(h'_T, D_V) \leq l(h_T, D_V)$

### 2.3 Linear Regression

$t : M \rightarrow R, M \subseteq \mathbb{R}^k, R \subseteq \mathbb{R}$

- find  $w = (w_0, \dots, w_k) \in \mathbb{R}^{k+1}$  s.t.:

$h_w(x_1, \dots, x_k) = w_0 x_0 + \sum_{i=1}^k w_i x_i$

approximates  $t \Leftrightarrow w$  minimizes  $l(h_w, D)$

- note:  $w_0 = 1$  always! ( $w_0$  is bias)

### Analytical Solution

1. Partial derivatives:  $\frac{\partial l(h_w, D)}{\partial w_i}$

2. Set to 0, put in values from  $D$

3. Solve LGS with Gauss for  $w_i$

### SGD (Iterative Solution)

1. initialize  $w$  randomly

2. choose random  $1 \leq i \leq n, T++$

3.  $\delta = y_i - h_w(x_i)$  (residual)

4.  $\Delta w = \delta \cdot x_i$  (derivatives)

5.  $w = w + \eta \cdot \Delta w$  (parameters)

6. If  $\neg$ converged  $\rightarrow 2$ , else return  $w$

- Pros: simple, robust to noisy data, representation independent

- Cons: stability, convergence problems, sensitive to learning rate  $\eta$

### BGD (accumulate derivatives $\forall i$ )

1. initialize  $w$  randomly

2. For each  $1 \leq i \leq n, T++$ :

2.1.  $\delta = y_i - h_w(x_i)$

2.2.  $\Delta w = \Delta w + \delta \cdot x_i$

3.  $w = w + \eta \cdot \Delta w$

4. If  $\neg$ converged  $\rightarrow 2$ , else return  $w$

- sequence of examples (batch) are processed together, before updating  $w$

### IGD

1. initialize  $w$  randomly

2. For each  $1 \leq i \leq n, T++$ :

2.1.  $\delta = y_i - h_w(x_i)$

2.2.  $\Delta w = \delta \cdot x_i$

2.3.  $w = w + \eta \cdot \Delta w$

3. If  $\neg$ converged  $\rightarrow 2$ , else return  $w$

Property	SGD stochastic	IGD iterative	BGD batch	MBGD mini-batch
Batch size	1	1	n	varies
Batch selection	random	sequential	sequential	sequential
Parallelization	difficult	difficult	trivial	trivial
Space requirement	low	low	high	varies
Stuck local minimum	no	no	yes	varies
Convergence speed	slow	slow	fast	varies

## Polynomial Regression

Approach: 1. prepare nonlinear combinations of features as features (curse of dimensionality: max  $k$  features  $5k \leq n$ )

2. then perform linear regression on expanded feature space with SGD, BGD, IGD or analytical approach.

3. Project solution back to input (feature) space

- keep original features

- for  $m_i^3$  also include  $m_i^2$

- increase complexity -> increase risk of overfitting

### Regularization

'penalize model complexity in training process', optimize for  $l'(w, D)$ :

$l'(w, D) = l(h_w, D) + \frac{\lambda}{k} \cdot r(w)$

- Lasso Regression:  $r(w) = \sum_{i=1}^k |w_i|$

- Ridge Regression:  $r(w) = \sum_{i=1}^k w_i^2$

-  $\lambda$  big -> more regularization -> less complex model

-  $k$ : num of features (excluding bias  $w_0$ )

### Develop (S,B,I)GD for specific loss function $l(x)$ :

1. get 1st derivative  $l'(x)$  of loss:

for 1 Data example:  $n = 1$ , leave out  $\sum$

2. find  $-\delta = h_w(x_i) - y_i$  in 1st derivative

3. replace line 4 (derivatives) with:

$\Delta w = l'(x)$  but substitute  $\delta$  (! - !)

### Logistic Regression (Classification)

'find optimal hyperplane  $w'$  by optimization, to get  $h_w$ , to get classifier:

$h_w^c(z) = 1$  if  $h_w(z) \geq \frac{1}{2}$ , 0 otherwise

- 'discriminative' classifier

- only reasonable for binary  $R = \{0, 1\}$  ( $|R| > 2$  induces order bias)

- Training: optimizes  $w$  for  $l(h_w, D)$

- Prediction: performed by  $h_w^c$  (different)

- Discriminating Hyperplane 1 Dimension

> logistic function:  $h_w^\sigma(z) = \sigma(h_w(z))$

> logistic classifier:  $h_w^{c,\sigma}(z) = 1$  if  $h_w^\sigma(z) \geq \frac{1}{2}$ , 0 otherwise

- 'generative' classifier

-  $h_w^\sigma(z)$  gives prop, that  $z$  is class 1

> MLE Maximum Likelihood Estimator given  $H_A$  for  $D$  is:  $\hat{h} = \arg \max_{h \in H_A} P[D; h]$

## 2.4 Support Vector Machines

'learn optimal discriminating Hyperplane with maximal margin directly'

$$H(w) = \{(z_1, \dots, z_k) \in \mathbb{R}^k | w_0 + w_1x_1 + \dots + w_kx_k = 0\}$$

-  $H_1$  closest  $x_i$  with  $y_i = 1$  ( $wz^T = 1$ )

-  $H_0$  closest  $x_i$  with  $y_i = 0$  ( $wz^T = -1$ )

- Hyperplanes  $H_1, H_0$  parallel to  $H(w)$

$$\text{Margin distance}(H_1, H_0) = \frac{2}{\|w\|}$$

- Hinge Loss: only falsely classified data causes loss

**Hard Margin SVM:** no misclassifications/boundary violations

$$\hat{w} = \arg \min_w \frac{1}{2} \hat{w}^T \hat{w}, l_h(h_w, D) = 0$$

**Soft Margin SVM:**  $\lambda$  trades margin size against boundary violations

$\lambda$  small: larger margin, more violations

$\lambda$  big: smaller margin, less violations

$$\hat{w} = \arg \min_w \frac{1}{2} \hat{w}^T \hat{w} + \lambda l_h(h_w, D)$$

**Kernel Trick:** Kernels permits nonlinear separation in input space  $\mathbb{R}^k$  (through linear separation in  $\mathbb{R}^{k+d}$ )

- with suitable Kernel: no actual computations in  $\mathbb{R}^{k+d}$

-> no additional effort for non linear classification (linear classifier for free)

**D Linearly Separable if:**

$$\exists w_0, w_1, \dots, w_k : y'_i \cdot (w_0 + w_1x_1 + \dots + w_kx_k) > 0$$

where:  $y'_i = 1$  if  $(y_i = 1)$ ,  $-1$  if  $(y_i = 0)$

1.  $\forall i$  where  $y_i = 1 : h(x_i) > 0$

2.  $\forall i$  where  $y_i = 0 : h(x_i) < 0$

Proof by finding  $w$ , then transform to discriminating hyperplane (eg point  $x$ ):

## loss functions (and derivatives)

- **(0-1 loss)**  $l_{0/1}(h, D) = \sum_{i=1}^n (1 - \delta_{y_i, h(x_i)})$   
->  $\delta_{ij} = 1$  if  $i = j, 0$  otherwise.
- **(Mean Squared Error)**  $l_2(h, D) = \frac{1}{2n} \sum_{i=1}^n (h(x_i) - y_i)^2$   
 $\frac{\partial l(h, D)}{\partial w_p} = \frac{1}{n} \sum_{i=1}^n (h(x_i) - y_i)x_p = \frac{1}{n} \sum_{i=1}^n (w_0 + w_1x_{i1} + \dots + w_px_{ip} - y_i)x_p$
- **(Ridge Regression)**  $l'(w, D) = \frac{1}{2n} \sum_{i=1}^n (h(x_i) - y_i)^2 + \frac{\lambda}{k} \sum_{i=1}^k w_i^2$   
 $\frac{\partial l'(w, D)}{\partial w_p} = \frac{1}{n} \sum_{i=1}^n (h(x_i) - y_i)x_p + 2\lambda w_p, w_p \in \{0, w_1, \dots, w_k\}$
- **(Logistic Loss)**  $\ell_\sigma(h, D) = -\frac{1}{n} \sum_{i=1}^n (y_i \cdot \log(h(x_i)) + (1 - y_i) \cdot \log(1 - h(x_i)))$   
 $\frac{\partial \ell_\sigma(h_w, D)}{\partial w_p} = -\frac{1}{n} \sum_{i=1}^n (y_i - h_w^\sigma(x_i)) \cdot x_ip$
- **(Hinge Loss)**  $l_h(h, D) = \frac{1}{n} \sum_{i=1}^n \max(0, 1 - (2y_i - 1)h(x_i))$
- **(Cross Entropy Loss)**  $l_H(h, D) = \frac{1}{n} \sum_{i=1}^n H(1_{y_i}, h(x_i))$   
->  $H(1_y, p) = -\sum_{r \in R} 1_y(r) \log p(r) = -\log p(y)$

$$- w_0 + w_1x_1 = 0 \Rightarrow x = -\frac{w_0}{w_1}$$

Disprove by finding contradiction in System of inequations.

## 2.5 Neural Networks

### Perception Hypothesis

$$h_w^H(z) = H(\sum_{i=0}^k w_i z_i) = 1 \text{ if } wz^T \geq 0$$

### Perceptron Training

1. initialize  $w$  randomly,  $T = 0$
2. Select random  $1 \leq i \leq n$
3.  $\delta = y_i - h_w^H(x_i)$
4.  $\Delta w = \delta \cdot x_i$
5.  $w = w + \eta \cdot \Delta w$
6. If  $l(h_w^H, D) \neq 0$  to 2, else return  $w$

Rosenblatt: If D linrly seprble: PT terminates after finitely many corrections

Property	Gradient descent	PT algorithm
Loss function Discriminator	$\ell_2$ or $\ell_\sigma$ hyperplane	0-1 loss $\ell_{0/1}$ hyperplane
Data inseparable Perfect separation	robust potentially	no termination guaranteed
Parameter updates on Correction size	surrogate error scaled by steepness	class error fixed

Nonlinear seperation possible through network of perceptrons

$$\text{Network } N = (V, E, wt), wt : E \rightarrow \mathbb{R}$$

- State:  $s : V \rightarrow \mathbb{R}$

$$s'(v') = a(\sum_{v \in V} s(v) \cdot wt(v, v'))$$

$a$  (weighted sum of previous states)

- OR:  $N = W \in \mathbb{R}^{(V \cup I) \times V}$  (adj matrix)

- FeedForward N: no loops, no cycles,

every node reachable from 1 input node  $w_0 * i_0 = w_0$  is bias, noted above node

Characteristics: 1. input, 2. internal nodes, 3. layers, 4. type (FFN/RNN?)

### Induced Regression Hypothesis

$$h_W^v(z) : \mathbb{R}' \rightarrow \mathbb{R}, h_W^v(z) = s_v^{(m)}$$

Forward Pass with fixed output node

## Neural Network Training

compute gradients gradients

$$> \text{Gradients of Nodes } \frac{\partial L}{\partial r_v} =$$

$$- (s_v - y) \cdot H(r_v) \text{ if } v \text{ designated o node}$$

- 0 if  $v$  other o node

$$- H(r_v) \cdot \sum_{v' \in V} W_{vv'} \cdot \frac{\partial L}{\partial r_{v'}} \text{ otherwise}$$

$$> \text{Gradients of Edges } \frac{\partial L}{\partial W_{vv'}} : \text{'State of predecessor} \cdot \text{gradient of successor'}$$

### FF NN Training with SGD

1. initialize  $W$  randomly
2. choose random  $1 \leq i \leq n, T++$
3. compute states  $s_v$  (forward pass)
4. compute gradients  $\Delta L$  for  $(x_i, y_i)$  (backward pass)
5.  $W = W - \eta \cdot \Delta L$  (update weights)
6. If -converged  $\rightarrow 2$ , else return  $W$

- RNNs good for sequential data, gives 'sequence regression hypothesis'

**m-unroll of RNN:** FNN with m layers. backpropagation through time: unroll RNN, compute gradients for each layer, then average gradients for each parameter over all layers

### Transformer

'multiclass classification with attention'

- softmax  $\sigma : \mathbb{R}^V \rightarrow \mathbb{R}^V$  outputs distribution over inputs

- input -> (word) embedding -> encoder

->attention layer -> decoder -> output

- FF attention layer indicates rellevant source parts

### 3 Unsupervised Learning

$h : M \rightarrow \{1, \dots, c\}$  assigns each point a

**cluster:** names  $\{1, \dots, c\}$  arbitrary

- loss is avg of squared euclidian distance to cluster mean (centroid)

### c-means (Lloyds) Algorithm

1. init c centroids  $m_1, \dots, m_c$  random
2. Add each  $i$  of  $x_i$  to closest centroids  $(m_j)$  Set  $I_j$
3. Update centroids to mean of assigned points.  $m_j = \frac{1}{n} \sum_{i \in I_j} x_i$

$$- \text{induced hypothesis: } h_{m_1, \dots, m_c}(x) = \arg \min_{1 \leq j \leq c} \|x - m_j\|^2$$

-  $c$  needs to eb known in advance: elbow method: plot loss for  $1 \leq c \leq c_{\max}$ , look for 'elbow' where loss reduction flattens

### DBSCAN (Density Based Spatial Clustering of Applications with Noise)

- Hyperparameters:  $\epsilon$  (radius),  $c_{\min}$  (min points in  $\epsilon$ -neighborhood to be core )

**DBSCAN algorithm** [Ester, Kriegel, Sander, Xu 1996]
 

- Compute  $C \leftarrow \{x_i | 1 \leq i \leq n, x_i \text{ is core}\}$  (initialize core points)
- $\alpha \leftarrow 1$  and  $B \leftarrow C$  (first cluster and need to handle all core points)
- While  $B \neq \emptyset$ 
  - Select  $x \in B, C_\alpha \leftarrow \emptyset$  and  $C_\alpha \leftarrow \{x\}$  (select core point and setup cluster)
  - While  $C_\alpha \neq C_\alpha'$ 
    - Select  $x \in B, C_\alpha' \leftarrow \emptyset$  and  $C_\alpha' \leftarrow \{x\}$  (as long as cluster grows)
    - $C_\alpha \leftarrow C_\alpha \cup C_\alpha'$  (expand cluster)
    - $C_\alpha \leftarrow \{x_i | 1 \leq i \leq n, \exists x' \in C_\alpha: \|x - x'\| \leq \epsilon\}$  (determine neighbors)
  - $B \leftarrow B \setminus C_\alpha$  and  $\alpha \leftarrow \alpha + 1$  (cluster handled and prepare next cluster)
- Output clusters  $C_1, \dots, C_{\alpha-1}$  (remaining points are outliers)

- results in set of cluster and outliers (noise), not part of any cluster

### Hierarchical Clustering

1. Start with singleton clusters
2. greedily merge minimal scoring clusters to new clusters
3. Stop combination once cluster-cluster score  $>$  threshold  $\theta$

### 4 Reinforcement Learning

- Agent -> Environment

- Env -> (state, reward) -> Agent

- Environment is propabilistic NFA; has p-distribution for each action in state  $q$  (not fully known, completely observable)

- Trace: starts in initial state, respects  $\Delta$ , ends in terminal state:  $q_0 \rightarrow^{a_1} q_1 \dots$

- **(average within-cluster variance)**  $l_V(h, D) = \frac{1}{n} \sum_{i=1}^n \|x_i - \mu_i\|^2$  where  $\mu_i$  is centroid of cluster that  $x_i$  belongs to (euclidean distance to cluster mean)

### Ableitungsregeln

- Produktregel:  $(f \cdot g)(x)' = f'(x) \cdot g(x) + f(x) \cdot g'(x)$
- Quotientenregel:  $(\frac{f}{g})'(x) = \frac{f'(x)g(x) - f(x)g'(x)}{g^2(x)}$
- Kettenregel:  $(f \circ g)'(x) = (f' \circ g)(x) \cdot g'(x) = f'(g(x)) \cdot g'(x)$   
 $(f \circ g \circ h)'(x) = f'(g(h(x))) \cdot g'(h(x)) \cdot h'(x)$
- $(\frac{1}{g})'(x) = -\frac{g'(x)}{g^2(x)}$
- $(\frac{1}{x^n})' = -nx^{-n-1}$
- $\log_a'(x) = \frac{1}{x \ln(a)}$
- $\ln_e'(x) = \frac{1}{x}$
- $|x'| = \frac{x}{|x|} = \text{sgn}(x)$  for  $x \neq 0$

- Rewards:  $R : \Delta \rightarrow \mathbb{R}$ , reward for taking transition (mostly only for terminal transitions: win/lose)

- Policy: assigns propability to each action in each state:  $P : Q \times \Sigma \rightarrow [0, 1]$   
 $P(a|q)$  determines Agents action  
 $P$  detm. if  $\forall q \in Q : \exists a \in \Sigma : P(a|q) = 1$

### Bellman equation

- estimate value of States: to steer towards high-value states

$$V(q) = \max_{a \in \Sigma} \sum_{q' \in Q} P(q'|q, a) \cdot (R(q, a, q') + \gamma V(q'))$$

'likelyhood of transisiton  $\delta$ . direct reward + discounted future reward' (Q-Value  $Q_V(q, a)$  is the same  $V(q)$  but for specific action  $a$ )

-  $\gamma$  discount factor: future rewards discounted by  $\gamma^n$  for n steps in future

**Value iteration:** for every state compute  $V(q)$  until convergence (finds optimal Q-values  $V^*$ )

given  $V^*$ : optimal policy selects action with highest expected reward:

### Policy iteration

- Q-Value with deterministic policy

$$P: Q_V^P(q) = \sum_{q' \in Q} p(q'|q, P(q)) \cdot (R(q, P(q), q') + \gamma V(q'))$$

1. for every state compute  $Q_V^P(q)$  until convergence
2. update policy to take optimal action
3. If -converged  $\rightarrow 1$

- both methods are model based: they have perfect information (exploitation)

### No perfect information: model free

-  $\epsilon$ -greedy policy: with prop  $\epsilon$  select random action (exploration), else optimal action (exploitation)

- Q-Learning: extension of Val Iteration

- SARSA: extension of Policy Iteration