Embedding and un-embedding a categorical

Embedding the categorical output v

- \blacktriangleright we embed raw output $v \in \mathcal{V}$ into R^m as $y = \psi(v) \in \mathsf{R}^m$
- ▶ we can describe ψ by the K vectors $\psi_1 = \psi(v_1), \dots, \psi_K = \psi(v_K)$ (*i.e.*, just say what vector in \mathbb{R}^m each $v \in \mathcal{V}$ maps to)
- \blacktriangleright we call the vector ψ_i the *representative* of v_i
- we call the set $\{\psi_1, \ldots, \psi_K\}$ the *constellation*
- examples:
 - ▶ TRUE \mapsto 1, False \mapsto -1
 - ▶ TRUE \mapsto **1**, FALSE \mapsto **0**
 - ▶ Yes \mapsto 1, maybe \mapsto 0 no \mapsto -1
 - ▶ YES \mapsto (1,0), MAYBE \mapsto (0,0), NO \mapsto (0,1)
 - ▶ APPLE \mapsto (1,0,0), ORANGE \mapsto (0,1,0), BANANA \mapsto (0,0,1)
 - ▶ (Horse 3, Horse 1, Horse 2) \mapsto (3,1,2)
 - ▶ word2vec (maps 1M words to vectors in **R**³⁰⁰)

One-hot and reduced one-hot embedding

▶ one-hot embedding of K classes into \mathbf{R}^{K} : $\psi(v_i) = \psi_i = e_i$

- e.g., for Booleans: $\psi_1 = (1,0)$, $\psi_2 = (0,1)$
- ▶ reduced one-hot embedding into \mathbf{R}^{K-1} :

 \blacktriangleright choose one of the classes as the *default*, and map it to $0 \in \mathbf{R}^{K-1}$

 \blacktriangleright map the others to the unit vectors $e_1, \ldots, e_{K-1} \in \mathsf{R}^{K-1}$

▶ for Booleans:

- ▶ one-hot embedding is $\psi_1 = (1,0)$, $\psi_2 = (0,1)$
- \blacktriangleright reduced one-hot embedding is $\psi_1 = 0$, $\psi_2 = 1$

• example: $\mathcal{V} = \{$ MAYBE, YES, NO $\}$, with default MAYBE

▶ reduced one-hot embedding is ψ (MAYBE) = (0,0), ψ (YES) = (1,0), ψ (NO) = (0,1)

Classifying by un-embedding a prediction

 \blacktriangleright embed raw input to feature vector as $x = \phi(u) \in \mathsf{R}^d$

 \blacktriangleright embed raw output to representative as $y = \psi(v) \in \mathsf{R}^m$

$$\blacktriangleright$$
 create predictor $g: \mathsf{R}^d
ightarrow \mathsf{R}^m$ with $\hat{y} = g(x)$

 \blacktriangleright we hope that $\hat{y} = g(x) pprox y = \psi(v)$ (\hat{y} and y are vectors, so this means $||\hat{y} - y||_2$ small)

 \blacktriangleright to get the prediction, we *un-embed* \hat{y} to get \hat{v} : $\hat{v} = \psi^{\dagger}(\hat{y})$

- ▶ $\psi^{\dagger}: \mathbf{R}^{m} \rightarrow \mathcal{V}$ is the *un-embedding function*
- \blacktriangleright the final classifier has the form $\hat{v} = G(u) = \psi^{\dagger}(g(\phi(u)))$
- \blacktriangleright can write as $G=\psi^{\dagger}\circ g\circ \phi$
- ▶ in words: *embed*; *predict*; *un-embed*

- ▶ given prediction $\hat{y} \in \mathbf{R}^m$, we *un-embed* to get \hat{v}
- \blacktriangleright we denote our un-emdedding using the symbol $\psi^{\dagger}: \mathbf{R}^{m}
 ightarrow \mathcal{V}$
- ▶ we will use *nearest neighbor un-embedding*:

$$\psi^\dagger(\hat{y}) = rgmin_{v\in\mathcal{V}} ||\hat{y} - \psi(v)||_2$$

(we can break ties any way we like)

 \blacktriangleright i.e., we choose the raw value associated with the nearest representative to \hat{y}

Un-embedding Boolean

 \blacktriangleright embed TRUE \mapsto $1 = \psi_1$ and FALSE \mapsto $-1 = \psi_2$

un-embed via

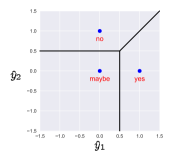
$$\psi^{\dagger}(\hat{y}) = egin{cases} ext{TRUE} & \hat{y} \geq 0 \ ext{false} & \hat{y} < 0 \end{cases}$$



Un-embedding one-hot

- \blacktriangleright take $\mathcal{V} = \{1, \dots, K\}$
- ▶ one-hot embedding: $\psi_i = e_i$, i = 1, ..., K
- \blacktriangleright un-embed via $\psi^{\dagger}(\hat{y}) = \operatorname{argmin}_i ||y e_i||_2$
- \blacktriangleright can be expressed as $\psi^{\dagger}(\hat{y}) = \operatorname{argmax}_{i} \hat{y}_{i}$
- \blacktriangleright *i.e.*, we guess class associated with the largest entry in \hat{y}
- reason:
 - $\blacktriangleright \ ||\hat{y} e_i||_2^2 = ||\hat{y}||_2^2 + 1 2\hat{y}^{\mathsf{T}} e_i = ||\hat{y}||_2^2 + 1 2\hat{y}_i$
 - \blacktriangleright first two terms don't depend on i, so we just choose i to maximize \hat{y}_i

Un-embedding yes, maybe, no



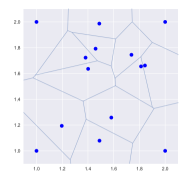
▶ embed YES \mapsto (1,0), MAYBE \mapsto (0,0), NO \mapsto (0,1) (reduced one-hot)

un-embed via

$$\psi^{\dagger}(\hat{y}) = egin{cases} ext{YES} & \hat{y}_1 > 1/2, \; \hat{y}_1 > \hat{y}_2 \ ext{MAYBE} & \hat{y}_1 < 1/2, \; \hat{y}_2 < 1/2 \ ext{NO} & \hat{y}_2 > 1/2, \; \hat{y}_1 < \hat{y}_2 \end{cases}$$

(can choose any value on boundaries)

Voronoi diagram



- ▶ ψ^{\dagger} partitions R^m into the K regions $\{y \mid \psi^{\dagger}(y) = v_i\}$, for i = 1, ..., K
- ▶ regions are *polyhedra* (of points closer to one representative than all others)
- called Voronoi diagram
- **b** boundaries between regions are perpendicular bisectors between pairs of representatives ψ_i, ψ_j

Loss function and empirical risk

Parametrized predictor

- \blacktriangleright we use parametrized predictor $g_{\theta}: \mathsf{R}^d \to \mathsf{R}^m$
- \triangleright θ is a parameter that we can choose
- predictor $g_{ heta}$ gives classifier $v = G(u) = \psi^{\dagger}(g_{ heta}(\psi(u)))$

- \blacktriangleright we'll choose θ using ERM and a training data set
- > we validate the predictor by performance metric on a test data set

Examples of parametrized predictors for classification

tree-based predictor (called a *classification tree*)

 $\blacktriangleright~\theta$ encodes tree, feature to split at each node, threshold, leaf values

 \blacktriangleright each leaf has a value of \hat{y}

neural network

- \triangleright θ gives offset and weights in the different layers
- \blacktriangleright \hat{y} is output of last layer
- ▶ linear predictor
 - \triangleright θ is a $d \times m$ parameter matrix

$$\triangleright \ \hat{y} = g_{\theta}(x) = \theta^{\mathsf{T}} x$$

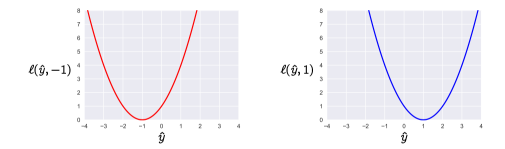
Loss function for classifiers

- \blacktriangleright we use a loss function $\ell: {\mathsf R}^m \times {\mathcal V} \to {\mathsf R}$
- \blacktriangleright $\ell(\hat{y}, y)$ is how much prediction $\hat{y} \in \mathsf{R}^m$ bothers us when observed value is $y \in \{\psi_1, \dots, \psi_K\}$
- \blacktriangleright the only possible values of y are ψ_1, \ldots, ψ_K , so we can simply give the K functions of \hat{y}

$$\ell(\hat{y},\psi_j), \quad j=1,\ldots,K$$

- ▶ $\ell(\hat{y}, \psi_j)$ is how much we dislike predicting \hat{y} when $y = \psi_j$
- ▶ typically $\ell(\hat{y}, \psi_j)$ is nonnegative, and small when $\hat{y} \approx \psi_j$
- ▶ square loss: $\ell(\hat{y}, \psi_j) = ||\hat{y} \psi_j||_2^2$
- ▶ we'll see far better loss functions for classifiers later

Square loss for Boolean classification



ERM and RERM

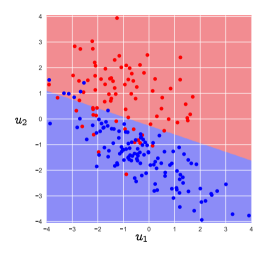
 \blacktriangleright we are given a training data set $x^1, \ldots, x^n, y^1, \ldots, y^n$, and a parametrized predictor $g_ heta$

 \blacktriangleright empirical risk associated with loss function ℓ is

$$\mathcal{L}(heta) = rac{1}{n}\sum_{i=1}^n \ell(\hat{y}^i,y^i) = rac{1}{n}\sum_{i=1}^n \ell(g_ heta(x^i),y^i)$$

- **ERM**: choose θ to minimize $\mathcal{L}(\theta)$
- \blacktriangleright in most cases, we need to resort to numerical optimization to find θ
- ▶ regularized ERM: choose θ to minimize $\mathcal{L}(\theta) + \lambda r(\theta)$
- ▶ r is the regularizer and $\lambda > 0$ is the regularization hyper-parameter

- \blacktriangleright linear predictor $\hat{y} = \theta^{\mathsf{T}} x$
- ▶ square loss $\ell(\hat{y}, \psi_j) = ||\hat{y} \psi_j||_2^2$
- ▶ square regularizer $r(\theta) = ||\theta||_F^2$
- called *least squares classifier*
- can solve RERM problem exactly using least squares
- ▶ we'll see better losses for classifiers later



▶ $u \in \mathsf{R}^2$, embedded as $x = (1, u_1, u_2)$; $v \in \{-1, 1\}$, embedded as y = v

▶ square loss and regularizer

ERM for Neyman-Pearson metric

Neyman-Pearson meetric

- ▶ suppose we care about the Neyman-Pearson metric, $\sum_{j=1}^{K} \kappa_j E_j$
- ▶ E_j is rate of mistaking v_j for some other class; κ is a weight vector
- \triangleright κ_j is how much we care about mistaking v_j , relative to others
- \blacktriangleright to reflect different costs for different errors, we scale the losses by κ_i

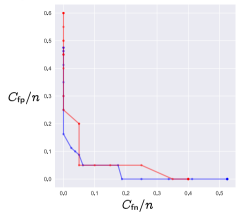
 \blacktriangleright if $\widetilde{\ell}(\widehat{y},\psi_j)$, $j=1,\ldots K$ are the unweighted losses, we use

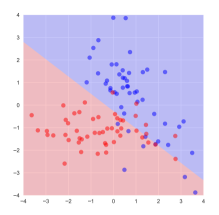
$$\ell(\hat{y},\psi_j)=\kappa_j \widetilde{\ell}(\hat{y},\psi_j), \hspace{1em} j=1,\ldots,K$$

- ▶ Boolean classifier, with $\psi_1 = -1$, $\psi_2 = 1$
- \blacktriangleright we care about Neyman-Pearson metric, $\kappa E_{\rm fn} + E_{\rm fp}$
- \blacktriangleright $\kappa > 0$ is how much we care about false negatives relative to false positive
- ▶ we use loss function

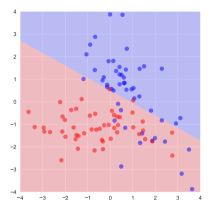
$$\ell(\hat{y},y) = egin{cases} (\hat{y}-y)^2 & ext{if } y = -1 \ \kappa(\hat{y}-y)^2 & ext{otherwise} \end{cases}$$

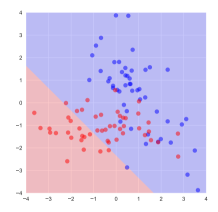
which gives more weight to deviating from the positive representative ψ_2





- ▶ square loss, sum squares regularizer
- ▶ left hand plot shows training errors in blue, test errors in red
- ▶ right hand plot shows minimum-error classifier (*i.e.*, $\kappa = 1$)





- $\blacktriangleright\,$ left hand plot shows predictor when $\kappa=0.4$
- \blacktriangleright right hand plot shows predictor when $\kappa = 4$

Summary

Summary

a *classifier* is a predictor, when the raw output is categorical $v \in \mathcal{V} = \{v_1, \dots, v_K\}$

- ▶ called a *Boolean classifier* when $|\mathcal{V}| = K = 2$, *multi-class classifier* when K > 2
- ▶ judged by various *error rates*, summarized in a *confusion matrix*, on test data

fitting a classifier to a training data set via ERM or RERM

- ▶ we embed the raw output v into R^m using ψ , with $\psi_i = \psi(v_i)$ the *representative* of class i
- \blacktriangleright we build a predictor for y, given x
- we *un-embed* a prediction $\hat{y} \in \mathbf{R}^m$ to a class prediction $\hat{v} = \psi^{\dagger}(\hat{y})$, using nearest neighbor
- ▶ there are special loss functions for fitting classifiers, that we'll see later