Learning to control

D.A. Forsyth, UIUC
Topics

• Scamper through basic reinforcement learning ideas
• Imitation learning
  • and its variants and problems
  • as structure learning
First learned steering controller

An autonomous Land vehicle in a neural Network, Pomerleau 1989

"ALVINN:"
Markov Decision Process

- Mathematical formulation of the RL problem
- **Markov property**: Current state completely characterises the state of the world

Defined by: \((\mathcal{S}, \mathcal{A}, \mathcal{R}, \mathbb{P}, \gamma)\)

- \(\mathcal{S}\) : set of possible states
- \(\mathcal{A}\) : set of possible actions
- \(\mathcal{R}\) : distribution of reward given (state, action) pair
- \(\mathbb{P}\) : transition probability i.e. distribution over next state given (state, action) pair
- \(\gamma\) : discount factor
Markov Decision Process

- At time step $t=0$, environment samples initial state $s_0 \sim p(s_0)$
- Then, for $t=0$ until done:
  - Agent selects action $a_t$
  - Environment samples reward $r_t \sim R(. \mid s_t, a_t)$
  - Environment samples next state $s_{t+1} \sim P(. \mid s_t, a_t)$
  - Agent receives reward $r_t$ and next state $s_{t+1}$

- A policy $\pi$ is a function from $S$ to $A$ that specifies what action to take in each state
- **Objective**: find policy $\pi^*$ that maximizes cumulative discounted reward: $\sum_{t \geq 0} \gamma^t r_t$
A simple MDP: Grid World

actions = {
1. right
2. left
3. up
4. down
}

states

Set a negative “reward” for each transition (e.g. \( r = -1 \))

**Objective:** reach one of terminal states (greyed out) in least number of actions
A simple MDP: Grid World

Random Policy

Optimal Policy
The optimal policy $\pi^*$

We want to find optimal policy $\pi^*$ that maximizes the sum of rewards.

How do we handle the randomness (initial state, transition probability...)? Maximize the expected sum of rewards!

Formally: $\pi^* = \arg \max_\pi \mathbb{E} \left[ \sum_{t \geq 0} \gamma^t r_t \mid \pi \right]$ with $s_0 \sim p(s_0), a_t \sim \pi(\cdot \mid s_t), s_{t+1} \sim p(\cdot \mid s_t, a_t)$
Definitions: Value function and Q-value function

Following a policy produces sample trajectories (or paths) $s_0, a_0, r_0, s_1, a_1, r_1, \ldots$

How good is a state?
The **value function** at state $s$, is the expected cumulative reward from following the policy from state $s$:

$$V^\pi(s) = \mathbb{E} \left[ \sum_{t \geq 0} \gamma^t r_t | s_0 = s, \pi \right]$$

How good is a state-action pair?
The **Q-value function** at state $s$ and action $a$, is the expected cumulative reward from taking action $a$ in state $s$ and then following the policy:

$$Q^\pi(s, a) = \mathbb{E} \left[ \sum_{t \geq 0} \gamma^t r_t | s_0 = s, a_0 = a, \pi \right]$$
Bellman equation

The optimal Q-value function $Q^*$ is the maximum expected cumulative reward achievable from a given (state, action) pair:

$$Q^*(s, a) = \max_\pi \mathbb{E} \left[ \sum_{t \geq 0} \gamma^t r_t | s_0 = s, a_0 = a, \pi \right]$$

$Q^*$ satisfies the following Bellman equation:

$$Q^*(s, a) = \mathbb{E}_{s' \sim \mathcal{E}} \left[ r + \gamma \max_{a'} Q^*(s', a') | s, a \right]$$

**Intuition:** if the optimal state-action values for the next time-step $Q^*(s', a')$ are known, then the optimal strategy is to take the action that maximizes the expected value of $r + \gamma Q^*(s', a')$

The optimal policy $\pi^*$ corresponds to taking the best action in any state as specified by $Q^*$
Solving for the optimal policy

**Value iteration algorithm**: Use Bellman equation as an iterative update

\[ Q_{i+1}(s, a) = \mathbb{E} \left[ r + \gamma \max_{a'} Q_i(s', a') | s, a \right] \]

\( Q_i \) will converge to \( Q^* \) as \( i \rightarrow \infty \)

**What's the problem with this?**
Not scalable. Must compute \( Q(s, a) \) for every state-action pair. If state is e.g. current game state pixels, computationally infeasible to compute for entire state space!

**Solution**: use a function approximator to estimate \( Q(s, a) \). E.g. a neural network!
Solving for the optimal policy: Q-learning

Remember: want to find a Q-function that satisfies the Bellman Equation:

\[ Q^*(s, a) = \mathbb{E}_{s' \sim \mathcal{E}} \left[ r + \gamma \max_{a'} Q^*(s', a') | s, a \right] \]

**Forward Pass**

Loss function:

\[ L_i(\theta_i) = \mathbb{E}_{s, a \sim \rho(\cdot)} \left[ (y_i - Q(s, a; \theta_i))^2 \right] \]

where \( y_i = \mathbb{E}_{s' \sim \mathcal{E}} \left[ r + \gamma \max_{a'} Q(s', a'; \theta_{i-1}) | s, a \right] \)

**Backward Pass**

Gradient update (with respect to Q-function parameters \( \theta \)):

\[ \nabla_{\theta_i} L_i(\theta_i) = \mathbb{E}_{s, a \sim \rho(\cdot); s' \sim \mathcal{E}} \left[ r + \gamma \max_{a'} Q(s', a'; \theta_{i-1}) - Q(s, a; \theta_i) \right] \nabla_{\theta_i} Q(s, a; \theta_i) \]

Iteratively try to make the Q-value close to the target value \( y_i \) it should have, if Q-function corresponds to optimal Q* (and optimal policy \( \pi^* \))
Training the Q-network: Experience Replay

Learning from batches of consecutive samples is problematic:
- Samples are correlated => inefficient learning
- Current Q-network parameters determines next training samples (e.g. if maximizing action is to move left, training samples will be dominated by samples from left-hand size) => can lead to bad feedback loops

Address these problems using **experience replay**
- Continually update a **replay memory** table of transitions \((s_t, a_t, r_t, s_{t+1})\) as game (experience) episodes are played
- Train Q-network on random minibatches of transitions from the replay memory, instead of consecutive samples

Each transition can also contribute to multiple weight updates
=> greater data efficiency
Policy Gradients

What is a problem with Q-learning?
The Q-function can be very complicated!

Example: a robot grasping an object has a very high-dimensional state => hard to learn exact value of every (state, action) pair

But the policy can be much simpler: just close your hand
Can we learn a policy directly, e.g. finding the best policy from a collection of policies?
Policy Gradients

Formally, let’s define a class of parametrized policies: \( \Pi = \{\pi_\theta, \theta \in \mathbb{R}^m\} \)

For each policy, define its value:

\[
J(\theta) = \mathbb{E} \left[ \sum_{t \geq 0} \gamma^t r_t | \pi_\theta \right]
\]

We want to find the optimal policy \( \theta^* = \arg \max_\theta J(\theta) \)

How can we do this?
Gradient ascent on policy parameters!
REINFORCE algorithm

Mathematically, we can write:

\[ J(\theta) = \mathbb{E}_{\tau \sim p(\tau; \theta)} [r(\tau)] \]

\[ = \int_{\tau} r(\tau)p(\tau; \theta) d\tau \]

Where \( r(\tau) \) is the reward of a trajectory \( \tau = (s_0, a_0, r_0, s_1, \ldots) \)
**REINFORCE algorithm**

**Expected reward:**
\[ J(\theta) = \mathbb{E}_{\tau \sim p(\tau; \theta)} [r(\tau)] = \int_{\tau} r(\tau)p(\tau; \theta)d\tau \]

Now let's differentiate this:
\[ \nabla_\theta J(\theta) = \int_{\tau} r(\tau)\nabla_\theta p(\tau; \theta)d\tau \]

**Intractable! Gradient of an expectation is problematic when** \( p \) **depends on** \( \theta \)

However, we can use a nice trick:
\[ \nabla_\theta p(\tau; \theta) = p(\tau; \theta)\frac{\nabla_\theta p(\tau; \theta)}{p(\tau; \theta)} = p(\tau; \theta)\nabla_\theta \log p(\tau; \theta) \]

If we inject this back:
\[ \nabla_\theta J(\theta) = \int_{\tau} (r(\tau)\nabla_\theta \log p(\tau; \theta))p(\tau; \theta)d\tau \]

\[ = \mathbb{E}_{\tau \sim p(\tau; \theta)} [r(\tau)\nabla_\theta \log p(\tau; \theta)] \]

**Can estimate with Monte Carlo sampling**
REINFORCE algorithm

\[
\nabla_\theta J(\theta) = \int_\tau (r(\tau) \nabla_\theta \log p(\tau; \theta)) p(\tau; \theta) d\tau \\
= \mathbb{E}_{\tau \sim p(\tau; \theta)} [r(\tau) \nabla_\theta \log p(\tau; \theta)]
\]

Can we compute those quantities without knowing the transition probabilities?

We have: \( p(\tau; \theta) = \prod_{t \geq 0} p(s_{t+1}|s_t, a_t) \pi_\theta(a_t|s_t) \)

Thus: \( \log p(\tau; \theta) = \sum_{t \geq 0} \log p(s_{t+1}|s_t, a_t) + \log \pi_\theta(a_t|s_t) \)

And when differentiating: \( \nabla_\theta \log p(\tau; \theta) = \sum_{t \geq 0} \nabla_\theta \log \pi_\theta(a_t|s_t) \)

Doesn't depend on transition probabilities!

Therefore when sampling a trajectory \( \tau \), we can estimate \( J(\theta) \) with

\[
\nabla_\theta J(\theta) \approx \sum_{t \geq 0} r(\tau) \nabla_\theta \log \pi_\theta(a_t|s_t)
\]
Intuition

Gradient estimator: \( \nabla_\theta J(\theta) \approx \sum_{t \geq 0} r(\tau) \nabla_\theta \log \pi_\theta(a_t|s_t) \)

Interpretation:
- If \( r(\tau) \) is high, push up the probabilities of the actions seen
- If \( r(\tau) \) is low, push down the probabilities of the actions seen

Might seem simplistic to say that if a trajectory is good then all its actions were good. But in expectation, it averages out!

However, this also suffers from high variance because credit assignment is really hard. Can we help the estimator?
Variance reduction

Gradient estimator: \[ \nabla_{\theta} J(\theta) \approx \sum_{t \geq 0} r(\tau) \nabla_{\theta} \log \pi_{\theta}(a_t | s_t) \]

First idea: Push up probabilities of an action seen, only by the cumulative future reward from that state

\[ \nabla_{\theta} J(\theta) \approx \sum_{t \geq 0} \left( \sum_{t' \geq t} r_{t'} \right) \nabla_{\theta} \log \pi_{\theta}(a_t | s_t) \]

Second idea: Use discount factor \( \gamma \) to ignore delayed effects

\[ \nabla_{\theta} J(\theta) \approx \sum_{t \geq 0} \left( \sum_{t' \geq t} \gamma^{t'-t} r_{t'} \right) \nabla_{\theta} \log \pi_{\theta}(a_t | s_t) \]
Variance reduction: Baseline

**Problem:** The raw value of a trajectory isn’t necessarily meaningful. For example, if rewards are all positive, you keep pushing up probabilities of actions.

**What is important then?** Whether a reward is better or worse than what you expect to get

**Idea:** Introduce a baseline function dependent on the state. Concretely, estimator is now:

\[
\nabla_\theta J(\theta) \approx \sum_{t \geq 0} \left( \sum_{t' \geq t} \gamma^{t'-t} r_{t'} - b(s_t) \right) \nabla_\theta \log \pi_\theta(a_t | s_t)
\]
How to choose the baseline?

$$\nabla_\theta J(\theta) \approx \sum_{t \geq 0} \left( \sum_{t' \geq t} \gamma^{t' - t} r_{t'} - b(s_t) \right) \nabla_\theta \log \pi_\theta(a_t | s_t)$$

A simple baseline: constant moving average of rewards experienced so far from all trajectories.

Variance reduction techniques seen so far are typically used in “Vanilla REINFORCE”.
How to choose the baseline?

A better baseline: Want to push up the probability of an action from a state, if this action was better than the expected value of what we should get from that state.

Q: What does this remind you of?

A: Q-function and value function!

Intuitively, we are happy with an action $a_t$ in a state $s_t$ if $Q^\pi(s_t, a_t) - V^\pi(s_t)$ is large. On the contrary, we are unhappy with an action if it’s small.

Using this, we get the estimator: $\nabla_\theta J(\theta) \approx \sum_{t \geq 0} (Q^{\pi^\theta}(s_t, a_t) - V^{\pi^\theta}(s_t)) \nabla_\theta \log \pi^\theta(a_t | s_t)$
Actor-Critic Algorithm

**Problem:** we don’t know Q and V. Can we learn them?

*Yes*, using Q-learning! We can combine Policy Gradients and Q-learning by training both an **actor** (the policy) and a **critic** (the Q-function).

- The actor decides which action to take, and the critic tells the actor how good its action was and how it should adjust.
- Also alleviates the task of the critic as it only has to learn the values of (state, action) pairs generated by the policy.
- Can also incorporate Q-learning tricks e.g. experience replay.
- **Remark:** we can define by the **advantage function** how much an action was better than expected

\[ A^\pi(s, a) = Q^\pi(s, a) - V^\pi(s) \]
Why so many RL algorithms?

• Different tradeoffs
  • Sample efficiency
  • Stability & ease of use

• Different assumptions
  • Stochastic or deterministic?
  • Continuous or discrete?
  • Episodic or infinite horizon?

• Different things are easy or hard in different settings
  • Easier to represent the policy?
  • Easier to represent the model?
Reinforcement Learning: Learning policies guided by **sparse** rewards, e.g., win or not the game.

- Good: simplest, cheapest form of supervision
- Bad: High sample complexity

Where is it successful so far?

- in simulation, where we can afford a lot of trials, easy to parallelize
- not in robotic systems:
  1. action execution takes long
  2. we cannot afford to fail
  3. safety concerns

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Ideally we want **dense in time** rewards to closely guide the agent closely along the way.

Who will supply those shaped rewards?

1. **We will manually design them**: “cost function design by hand remains one of the ‘black arts’ of mobile robotics, and has been applied to untold numbers of robotic systems”

2. **We will learn them from demonstrations**: “rather than having a human expert tune a system to achieve desired behavior, the expert can demonstrate desired behavior and the robot can tune itself to match the demonstration”

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Learning from demonstrations a.k.a. Imitation Learning:
Supervision through an expert (teacher) that provides a set of demonstration trajectories: sequences of states and actions.

Imitation learning is useful when is easier for the expert to demonstrate the desired behavior rather than:

a) coming up with a reward that would generate such behavior,
b) coding up the desired policy directly.
The Imitation Learning problem

The agent (learner) needs to come up with a policy whose resulting state, action trajectory \textit{distribution matches} the expert trajectory \textit{distribution}.

Does this remind us of something…?
GANs! Generative Adversarial Networks (on state-action trajectories)

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Generative Adversarial Networks, Goodfellow et al. 2014
The Imitation Learning problem: Challenge

Actions along the trajectories are interdependent, as actions determine state transitions and thus states and actions down the road.

interdependent labels -> structure prediction

Action interdependence in time:

Algorithms developed in Robotics for imitation learning found applications in structured predictions problems, such as, sequence generation/labelling e.g. parsing.
Imitation Learning

For taking this structure into account, numerous formulations have been proposed:

- Direct: Supervised learning for policy (mapping states to actions) using the demonstration trajectories as ground-truth (a.k.a. behavior cloning) + ways to handle the neglect of action interdependence.

- Indirect: Learning the latent rewards/goals of the teacher and planning under those rewards to get the policy, a.k.a. Inverse Reinforcement Learning (next lecture)

Experts can be:

- Humans

- Optimal or near Optimal Planners/Controllers

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Imitation Learning as Supervised Learning

Driving policy: a mapping from (history of) observations to steering wheel angles

Behavior Cloning=Imitation Learning as Supervised learning

- Assume actions in the expert trajectories are i.i.d.
- Train a classifier or regressor to map observations to actions at each time step of the trajectory.
Classifier or regressor?

Because multiple actions $u$ may be plausible at any given observation $o$, policy network $p_{\pi_\theta}(u_t|o_t)$ usually is not a regressor but rather:

- A classifier (e.g., softmax output and cross-entropy loss, after discretizing the action space)

$$J(\theta) = -\sum_{i=1}^{m} \sum_{k=1}^{K} 1_{y(i)=k} \log[P(y(i) = k|x(i); \theta)]$$

- A GMM (mixture components weights, means and variances are parametrized at the output of a neural net, minimize GMM loss, (e.g., Hand writing generation Graves 2013)

- A stochastic network (previous lecture)
Independent in time errors

error at time $t$ with probability $\varepsilon$

$E[\text{Total errors}] \approx \varepsilon T$
As you get further off the path, the probability of making an error grows, cause the classifier thinks this state is rare.

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A Reduction of Imitation Learning and Structured Prediction to No-Regret Online Learning, Ross et al. 2011

error at time $t$ with probability $\varepsilon$

$$E[\text{Total errors}] \approx \varepsilon(T + (T-1) + (T-2) + \ldots + 1) \propto \varepsilon T^2$$
“In addition, the network must not solely be shown examples of accurate driving, but also how to recover (i.e. return to the road center) once a mistake has been made. Partial initial training on a variety of simulated road images should help eliminate these difficulties and facilitate better performance.” ALVINN: An autonomous Land vehicle in a neural Network, Pomerleau 1989
Data Distribution Mismatch!

\[ p_{\pi^*}(o_t) \neq p_{\pi_\theta}(o_t) \]
SL succeeds when training and test data distributions match, that is a fundamental assumption.
Demonstration Augmentation: ALVINN 1989

- Using graphics simulator for road images and corresponding steering angle ground-truth
- Online adaptation to human driver steering angle control
- 3 layers, fully connected layers, very low resolution input from camera and lidar.

“In addition, the network must not solely be shown examples of accurate driving, but also how to recover (i.e. return to the road center) once a mistake has been made. Partial initial training on a variety of simulated road images should help eliminate these difficulties and facilitate better performance.” ALVINN: An autonomous Land vehicle in a neural Network, Pomerleau 1989

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Additional, left and right cameras with automatic grant-truth labels to recover from mistakes.

“DAVE-2 was inspired by the pioneering work of Pomerleau [6] who in 1989 built the Autonomous Land Vehicle in a Neural Network (ALVINN) system. Training with data from only the human driver is not sufficient. The network must learn how to recover from mistakes. ...”

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Data Augmentation (3): Trails 2015

A Machine Learning Approach to Visual Perception of Forest Trails for Mobile Robots Giusti et al.
DAGGER (in simulation)

Dataset AGGregation: bring learner’s and expert’s trajectory distributions closer by labelling additional data points resulting from applying the current policy

1. train $\pi_\theta(u_t|o_t)$ from human data $\mathcal{D}_{\pi^*} = \{o_1, u_1, \ldots, o_N, u_N\}$
2. run $\pi_\theta(u_t|o_t)$ to get dataset $\mathcal{D}_{\pi} = \{o_1, \ldots, o_M\}$
3. Ask human to label $\mathcal{D}_{\pi}$ with actions $u_t$
4. Aggregate: $\mathcal{D}_{\pi^*} \leftarrow \mathcal{D}_{\pi^*} \cup \mathcal{D}_{\pi}$
5. GOTO step 1.

Problems:
- execute an unsafe/partially trained policy
- repeatedly query the expert

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* A Reduction of Imitation Learning and Structured Prediction to No-Regret Online Learning, Ross et al. 2011
DAGGER (in simulation)

Dataset AGGregation: bring learner’s and expert’s trajectory distributions closer by labelling additional data points resulting from applying the current policy

1. train $\pi_\theta(u_t|o_t)$ from human data $D_{\pi^*} = \{o_1, u_1, ..., o_N, u_N\}$
2. run $\pi_\theta(u_t|o_t)$ to get dataset $D_\pi = \{o_1, ..., o_M\}$
3. Ask human to label $D_\pi$ with actions $u_t$
4. Aggregate: $D_{\pi^*} \leftarrow D_{\pi^*} \cup D_\pi$
5. GOTO step 1.

Notice you might not actually need a human here - if your states are discretized, and you have enough data, you might get this by matching

Problems:

- execute an unsafe/partially trained policy
- repeatedly query the expert

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Initialize $\mathcal{D} \leftarrow \emptyset$.
Initialize $\hat{\pi}_1$ to any policy in $\Pi$.

for $i = 1$ to $N$ do

Let $\pi_i = \beta_i \pi^* + (1 - \beta_i) \hat{\pi}_i$.
Sample $T$-step trajectories using $\pi_i$.
Get dataset $\mathcal{D}_i = \{(s, \pi^*(s))\}$ of visited states by $\pi_i$ and actions given by expert.
Aggregate datasets: $\mathcal{D} \leftarrow \mathcal{D} \cup \mathcal{D}_i$.
Train classifier $\hat{\pi}_{i+1}$ on $\mathcal{D}$.

end for

Return best $\hat{\pi}_i$ on validation.

Algorithm 3.1: DAGGER Algorithm.
Structured prediction

Structured prediction: a learner makes predictions over a set of interdependent output variables and observes a joint loss.

Example: part of speech tagging

```
x = the monster ate the sandwich
y = Dt Nn Vb Dt Nn
```

A structured prediction problem consists of an input space $\mathcal{X}$, an output space $\mathcal{Y}$, a fixed but unknown distribution $\mathcal{D}$ over $\mathcal{X} \times \mathcal{Y}$, and a non-negative loss function $l(y^*, \hat{y}) \rightarrow \mathbb{R}^{\geq 0}$ which measures the distance between the true $y^*$ and predicted $\hat{y}$ outputs. The goal of structured learning is to use $N$ samples $(x_i, y_i)_{i=1}^N$ to learn a mapping $f : \mathcal{X} \rightarrow \mathcal{Y}$ that minimizes the expected structured loss under $\mathcal{D}$.  

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Traditional strategy

- Construct a parametric cost function $H(\mathcal{X}, \mathcal{Y}; \theta)$

- So that, for training $X^*$

$$\arg\max_{\mathcal{Y}} H(\mathcal{X}^*, \mathcal{Y}; \theta)$$

- is close to correct $Y^*$

  - (see movies for some details on construction)
Sequence labelling:
Part of speech tagging

\[
x = \text{the monster ate the sandwich}
\]
\[
y = \text{Dt Nn Vb Dt Nn}
\]
HMM: Making scribal Latin searchable

• **Goal:** make the ink in a handwritten text searchable

• **Issue:** not a good idea to transcribe

• **Strategy:**
  • compute $\log P(\text{ink} | \text{known sequence})$
  • for a line
  • known sequence can be a regular expression
    • eg (character)^* mihi (character)^*
    • ex: check you can do this w/ DP
  • rank lines by this, report
HMM: Making scribal Latin searchable

- **Goal**: make the ink in a handwritten text searchable

- **Issue**: few examples of glyphs
  - hard to label

- **Strategy**:
  - doesn’t really matter
    - like a substitution cypher - letter frequencies are what’s important
  - AND you can grow the pool of examples:
    - when you see “interrogave?unt” you know it’s “interrogaverunt”
    - so you can get another glyph
Figure 1: **Left**, a full page of our manuscript, a 12’th century manuscript of Terence’s Comedies obtained from [1]. **Top right**, a set of lines from a page from that document and **bottom right**, some words in higher resolution. Note: (a) the richness of page layout; (b) the clear spacing of the lines; (c) the relatively regular handwriting.
Figure 7: The handwritten text does not fully correspond to the transcribed version; for example, scribes commonly write “michi” for the standard “mihi”. Our search process reflects the ink fairly faithfully, however. Left the first four lines returned for a search on the string “michi”; right the first four lines returned for a search on the string “mihi”, which does not appear in the document. Note that our search process can offer scholars access to the ink in a particular document, useful for studying variations in transcription, etc.
Figure 8: Searches on short strings produce substrings of words as well as words (we show the first two lines returned from a search for “tu”).
Optimizing Graphical Models for Structured prediction

Graph labelling

- Encode output labels as a MRF
- Learn parameters of that model to:
  - maximize $p(\text{true labels} | \text{input})$
  - minimize $\text{loss}(\text{true labels, predicted labels})$

Let $G = (V, E)$ be a graph such that

$Y = (Y_v)_{v \in V}$, so that $Y$ is indexed by the vertices of $G$. Then $(X, Y)$ is a conditional random field when the random variables $Y_v$, conditioned on $X$, obey the Markov property with respect to the graph:

$p(Y_v | X, Y_w, w \neq v) = p(Y_v | X, Y_w, w \sim v)$, where $w \sim v$ means that $w$ and $v$ are neighbors in $G$. 

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Optimizing Graphical Models for Structured prediction

- Encode output labels as a MRF
- Learn parameters of that model to:
  - maximize $p(\text{true labels} \mid \text{input})$
  - minimize $\text{loss(true labels, predicted labels)}$

- Assumed Independence assumptions may not hold

- Computationally intractable with too many “edges” or non-decomposable loss functions (that involve many $y$s)
Instead: Decomposition of label

Sequence generation/labelling:

We can define an ordering and generate labels one at a time, where each output generated depends on all previous ones. E.g., sequential data admits the natural sequential ordering.

Image generation/labelling:

Here again we can define an ordering:

Pixel Recurrent Neural Networks, van den Oord et al

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Structured prediction as sequential decision making

When y decomposes in an ordered manner, a sequential decision making process emerges.
Structured prediction as sequential decision making

When $y$ decomposes in an ordered manner, a sequential decision making process emerges.
Structured prediction as sequential decision making

\[ x = \text{the monster ate the sandwich} \]

\[ y = \text{Dt Nn Vb Dt Nn} \]

- Example: Sequence labelling
- State: captures input sequence \( x \) and whatever labels (here part of speech tags) we have produced so far
- Actions: Next label to output
- Policy: a mapping of the input \( x \) and labels generated so far to the next label
- Reward: agreement of the predicted \( \hat{y} \) with ground-truth \( y^* \): \( \ell(e) = \ell(y^*, y_e) \)

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Recurrent Neural Networks

- RNNs tie the weights at each time step
- Condition the neural network on all previous inputs
- In principle, any interdependencies can be modeled between inputs and outputs, as well as between output labels.
- In practice, limitations from SGD training, capacity, initialization etc.

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Recurrent Neural Networks

For sequence labelling problems, actions of the labelling policies are $y_t$, e.g., part of speech tags

For sequence generation, actions of the labelling policies are $y_t = x_{t+1}$, e.g., word in answer generation  

$P(x_{t+1} = v_j | x_t, ..., x_1) = \hat{y}_{t,j}$

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Recurrent Neural Networks

The network is typically trained to maximize the log-likelihood of the output sequences given the input sequences of a training set $\mathcal{D} = \{(x^{(i)}, y^{(i)})\}$:

$$
\theta^* = \arg \max_{\theta} \log \sum_{(x^{(i)}, y^{(i)}) \in \mathcal{D}} P_{\theta}(y^{(i)}, x^{(i)})
$$

If the likelihood of an example decomposes over individual time steps:

$$
\log P_{\theta}(y|x) = \sum_{t} \log P_{\theta}(y_t|h_t)
$$

Else loss is computed at the end of the sequence and is back propagated through time.

A learned policy is the inference function of the model:

$$
\hat{\theta}(h_t) = \arg \max_{y} P(y_t = y|h_t; \theta)
$$

The reference policy is the policy that always outputs the true labels:

$$
\theta^*(h_t) = y_t
$$

Fragkiadaki, ND
Recurrent Neural Networks

The regular training procedure of RNNs treat true labels $y_t$ as actions while making forward passes. Hence, the learning agent follows trajectories generated by the reference policy rather than the learned policy. In other words, it learns:

$$\hat{\theta}^{sup} = \arg \min_{\theta} \mathbb{E}_{h \sim d_{\pi^*}} [l_{\theta}(h)]$$

However, our true goal is to learn a policy that minimizes error under its own induced state distribution:

$$\hat{\theta} = \arg \min_{\theta} \mathbb{E}_{h \sim d_{\theta}} [l_{\theta}(h)]$$
DAGGER for sequence labelling/generation with RNNs

function TRAIN(N, α)
  Initialize α = 1.
  Initialize model parameters θ.
  for i = 1..N do
    Set α = α · p.
    Randomize a batch of labeled examples.
    for each example (x, y) in the batch do
      Initialize h₀ = Φ(X).
      Initialize \( \mathcal{D} = \{(h₀, y₀)\} \).
      for t = 1..|Y| do
        Uniformly randomize a floating-number \( \beta \in [0, 1) \).
        if \( \alpha < \beta \) then
          Use true label: \( \tilde{y}_{t-1} = y_{t-1} \)
        else
          Use predicted label: \( \tilde{y}_{t-1} = \arg\max_y P(y \mid h_{t-1}; \theta) \).
        end if
        Compute the next state: \( h_t = f_\theta(h_{t-1}, \tilde{y}_{t-1}) \).
        Add example: \( \mathcal{D} = \mathcal{D} \cup \{(h_t, y_t)\} \).
      end for
    end for
  end for
  Online update θ by \( \mathcal{D} \) (mini-batch back-propagation).
end function
Imitation Learning

Two broad approaches:

- **Direct**: Supervised training of policy (mapping states to actions) using the demonstration trajectories as ground-truth (a.k.a. behavior cloning)

- **Indirect**: Learn the unknown reward function/goal of the teacher, and derive the policy from these, a.k.a. **Inverse Reinforcement Learning**