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**Un fantastico titolo
per la mia tesi di laurea!**

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Chapter 1

Introduction

Chapter 2

State of the Art

2.1 Offline RL: Advancements and Challenges

Offline Reinforcement Learning (RL) has emerged as a promising approach for training control policies using pre-collected experiences, eliminating the need for potentially dangerous or costly online interactions. Despite significant progress in this field, several challenges persist in its practical application, particularly in control and robotics domains.

Research efforts to enhance Offline RL performance are primarily advancing along two paths.

2.1.1 Improving Robustness and Sample Efficiency

The first research direction focuses on refining existing methods to enhance their robustness and sample efficiency. [Kumar et al., 2021] propose a comprehensive workflow for offline model-free robotic reinforcement learning. Their approach addresses challenges specific to real-world robotics applications, proposing metrics and protocols to assist practitioners in selecting policy checkpoints, regularization parameters, and model architecture.

[Shin and Kim, 2023] introduce a hierarchical approach called “Guide to Control” that solves efficiently long-horizon and sparse-reward tasks from offline data. This method enhances the effectiveness of offline RL in complex environments by breaking down the overall task into more manageable sub-tasks, thereby improving learning efficiency and performance.

2.1.2 Developing Datasets and Benchmarks

The second path concentrates on creating extensive datasets and benchmarks tailored for control and robotics domains. [Gürtler et al., 2023] present a benchmark for offline RL on real-robot hardware, providing a crucial resource for evaluating algorithms in practical settings. In their work they propose a set of benchmark datasets for robotic manipulation that are intended to help improving the state-of-the-art in offline reinforcement learning.

[Liu et al., 2023] contribute datasets and benchmarks specifically designed for offline safe reinforcement learning to accelerate research in this area. Their work addresses the critical aspect of safety in robotic applications, providing resources for researchers to develop and test algorithms that not only optimize performance but also ensure safe operation in various scenarios. The suite comprises three main components: expertly crafted safe policies, D4RL-styled datasets with environment wrappers, and high-quality offline safe RL baseline implementations. The authors present a methodical data collection pipeline using advanced safe RL algorithms to generate diverse datasets across 38 popular safe RL tasks, ranging from robot control to autonomous driving. They also introduce data post-processing filters to modify dataset diversity, simulating various data collection conditions.

2.2 Data Augmentation Strategies in RL

Despite ongoing improvements in learning algorithms, the quantity of training data continues to be a critical determinant of success in machine learning processes. This is especially challenging when dealing with complex, high-dimensional inputs such as images, where the data requirements for effective training are substantial.

To address the problem of limited data availability, researchers and practitioners often turn to **data augmentation techniques**. These methods aim to expand existing datasets by generating synthetic samples that are both artificial and plausible, thereby enriching the training set without the need for additional real-world data collection.

[Mumuni and Mumuni, 2022] provide a comprehensive survey of modern data augmentation approaches, highlighting their potential in enhancing computer vision across various domains.

2.2.1 Augmentation in Online and Offline RL

Both online and offline RL approaches have incorporated data augmentation techniques to improve learning efficiency and generalization.

In the **online** RL setting, [Laskin et al., 2020] introduce Reinforcement Learning with Augmented Data (RAD), a simple module designed to enhance most RL algorithms. The authors conduct an extensive study of various data augmentation techniques for both pixel-based and state-based inputs in RL, including two new augmentations they introduce: random translate and random amplitude scale.

[Hansen and Wang, 2021] further further explore the concept of data augmentation in online RL, proposing SOft Data Augmentation (SODA), a novel approach to improve generalization in RL; unlike traditional methods that directly learn policies from augmented data, SODA decouples augmentation from policy learning. It does this by applying a soft constraint on the encoder to maximize mutual information between latent representations of augmented and non-augmented data, while using only non-augmented data for RL optimization.

In the **offline** RL setting, [Han and Kim, 2022] introduce a selective data augmentation technique using a Variational Autoencoder (VAE). The method targets sparse subspaces in the dataset, representing them in the VAE’s latent space, then sampling and decoding new data points to augment underrepresented regions. This approach aims to generate virtual data that closely aligns with the original data distribution.

[Joo et al., 2022] propose a data augmentation method for offline RL called Swapping Target Q-Value (SQV). SQV works by swapping Q-values between original and transformed images, encouraging the algorithm to treat similar states as equivalent while emphasizing differences between distinct states. This method aims to enhance pixel-based learning without auxiliary loss functions, potentially improving performance and generalization in offline RL scenarios with limited datasets.

2.2.2 Dynamics-Aware Augmentation

Traditional augmentation methods often operate without knowledge of the underlying environment dynamics, requiring careful tuning to avoid generating inconsistent samples. To mitigate this issue, dynamics-aware augmentation schemes have emerged in the robotics field.

Nonetheless, effectively implementing these techniques requires an in-depth understanding of the underlying process, which is often not available, and the development of an augmentation strategy specifically crafted for the task at hand.

Chapter 3

Prior Knowledge

3.1 Deep Learning

In the field of machine learning, the first models you are often introduced to are those for regression and classification that utilize linear combinations of fixed basis functions. According to [Bishop, 2008], these models possess useful analytical and computational properties, but their practical applicability is constrained since their capacity is limited to linear functions and they cannot understand the interaction between any two input variables. This leads to problems such as the **curse of dimensionality**, where the number of possible interactions between variables grows exponentially with the number of input variables. To address large-scale problems, it is essential to adapt the basis functions to the data, as demonstrated by support vector machines (SVMs). Alternatively, one can fix the number of basis functions in advance while allowing them to be adaptive, utilizing parametric forms for the basis functions with parameter values adjusted during training.

The most successful example of this approach in pattern recognition is the **feed-forward neural network**, also known as the multilayer perceptron (MLP).

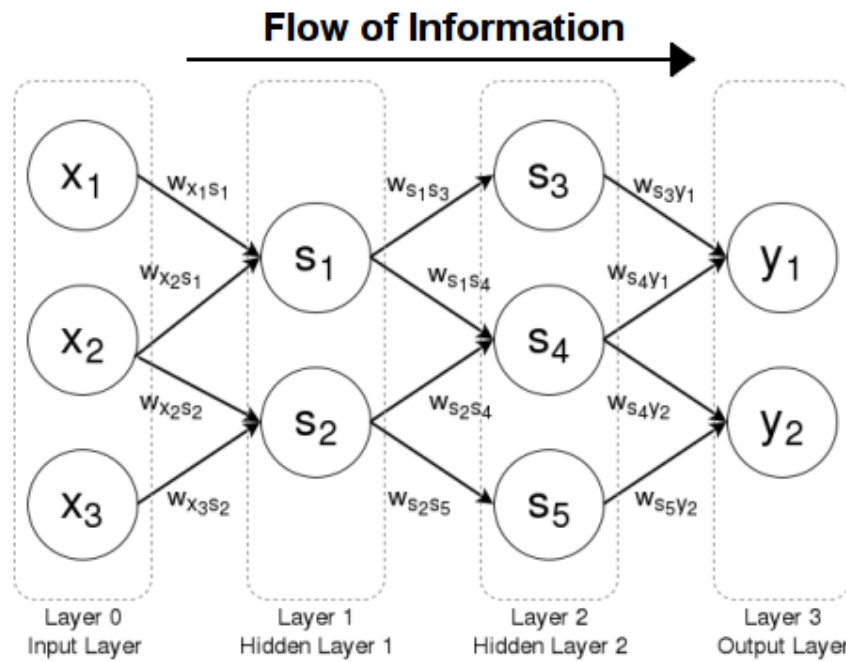


Figure 3.1: A feed-forward neural network with two hidden layers

Source: Brilliant

As explained in [Goodfellow et al., 2016], the goal of a feedforward network is to approximate a specific function f^* . For example, in a classification task, $y = f^*(x)$ maps an input x to a category y . A feedforward network defines a mapping $y = f(x; \theta)$ and learns the parameters θ that yield the best approximation of this function.

These models are called **feedforward** because information flows unidirectionally

from the input x , through intermediate computations defining f , and ultimately to the output y , without feedback loops.

The training data directly specifies the output layer's behavior but not the intermediate layers, which are called **hidden layers** because their desired outputs are not provided by the training data. The learning algorithm must determine how to best utilize these hidden layers to approximate f^* . Every layer of the network computes a non-linear transformation of the previous layer's activations, this way a complex function can be approximated by composing simpler functions, one for each layer. Layers are composed of a set of **units**, where each unit is a node that computes a non-linear function of the weighted sum of its inputs and is only connected to units in the previous and the following layer.

In order to train the network and update the weights, MLPs use the **backpropagation** technique, which computes the gradient of the loss function with respect to the weights of the network. The weights are then updated using an optimization algorithm such as **stochastic gradient descent** (SGD) or one of its adaptive variants like **Adam**, whose hyperparameters are tuned according to the task during training.

A feedforward network is called a **deep neural network** if it has more than one hidden layer, the branch of machine learning that studies deep neural networks is called **deep learning**.

They are called networks due to their structure, which involves composing multiple functions together. Typically, this composition is represented by a directed acyclic graph. For instance, functions $f^{(1)}$, $f^{(2)}$, and $f^{(3)}$ might be connected in a chain to form $f(x) = f^{(3)}(f^{(2)}(f^{(1)}(x)))$.

Functions $f^{(1)}$ and $f^{(2)}$ must be non linear, otherwise the composition would collapse into a single linear function. These non-linear functions are called **activation functions**. The last function $f^{(3)}$ is typically a linear function that maps the output of the final hidden layer to the output layer. This is the same as applying a linear model to a transformed input $\phi(x)$, where ϕ is a nonlinear transformation. The question then becomes how to choose the mapping ϕ .

The strategy of deep learning is to learn ϕ : in this approach, we use a model

$$y = f(x; \theta, w) = \phi(x; \theta)^\top w \quad (3.1)$$

Here, we have parameters θ that are used to learn ϕ from a broad class of functions, and parameters w that map $\phi(x)$ to the desired output. This approach allows for greater flexibility: specifically, by using a broad family of functions $\phi(x; \theta)$, the human designer only needs to select the appropriate general function family rather than finding precisely an exact function.

The **universal approximation theorem** [Hornik et al., 1989] tells us that regardless of what function we are trying to learn, we know that a feedforward network with enough units will be able to *represent* this function, however we are not guaranteed that the training algorithm will be able to *learn* it.

3.1.1 Convolutional Neural Networks

Convolutional Neural Networks (CNNs) are a specialized type of neural network for processing data that has a known, grid-like topology, it is particularly effective for image recognition tasks.

The key components of CNNs are:

- **Convolutional layers:** These layers apply a convolution operation to the input, passing the result to the next layer. The input grid is convolved with a set of filters, also known as **kernels**, which are small windows that move across the input grid and apply a convolution operation between the input and the kernel weight at each position. The weight of the kernel is learned during training. Each convolutional layer applies multiple filters to the input, each producing a different **feature map**.
- **Pooling layers:** These layers downsample the feature maps produced by the convolutional layers to reduce the dimensionality of the data. This reduces the computational complexity of the network and helps to focus on the most important elements of the input.
- **Fully connected (FC) layers:** These layers connect every neuron in one layer to every neuron in another layer. It is in these layers that the features learned by the convolutional layers are used to classify the input image.

An example of a CNN classification architecture is shown in Figure 3.2.

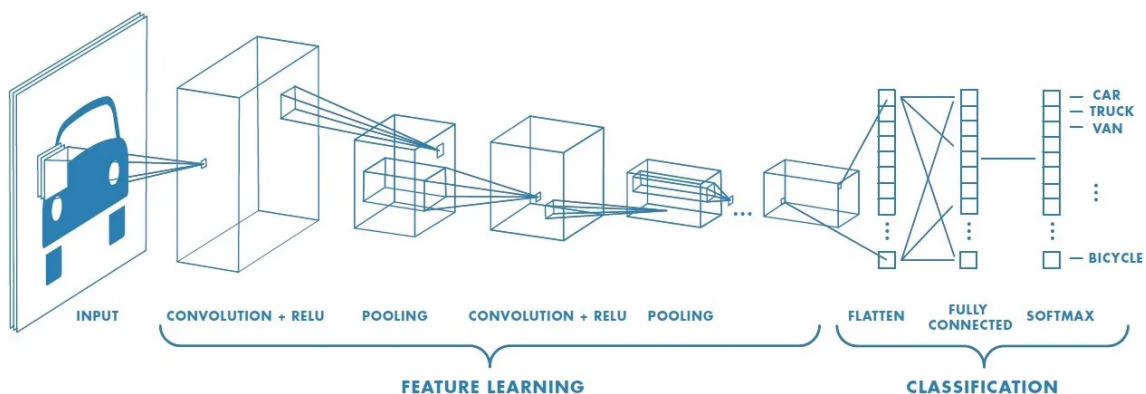


Figure 3.2: A typical CNN architecture for classification.

Source: Sumit Saha

CNNs have been shown to achieve state-of-the-art performance on a variety of computer vision tasks, including image classification, object detection and image segmentation. They have also been successfully applied to other types of data, such as speech recognition and natural language processing.

3.2 Reinforcement Learning

“**Reinforcement learning** is learning [...] how to map situations to actions so as to maximize a numerical reward signal. The learner is not told which actions to take, but instead must discover which actions yield the most reward by trying them. In the most interesting and challenging cases, actions may affect not only the immediate reward but also the next situation and, through that, all subsequent rewards. These two characteristics, **trial-and-error search** and **delayed reward**, are the two most important distinguishing features of reinforcement learning.” [Sutton and Barto, 1998]

Let’s explore the basic concepts of reinforcement learning with a simple example as described in [Zhao, 2024]. Consider a grid world scenario as shown in Figure 3.3 where a robot, referred to as the **agent**, moves between cells, occupying one cell at a time. The white cells are accessible, while the orange cells are forbidden. The goal is for the agent to reach a target cell.

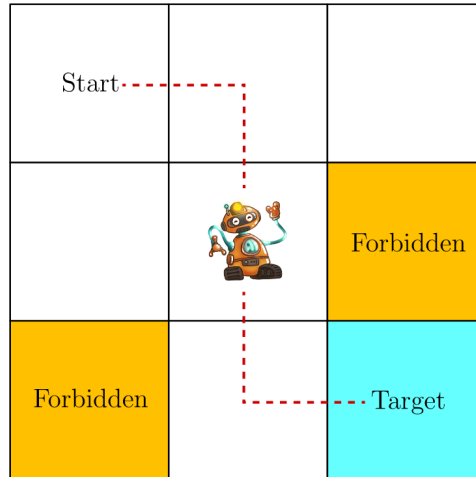


Figure 3.3: A simple reinforcement learning task

Source: [Zhao, 2024]

To achieve this, the agent needs a **policy** that guides it to the target efficiently, avoiding forbidden cells and unnecessary detours. If the grid layout is known, planning is simple. However, without prior knowledge, the agent must explore and learn through trial and error.

The agent’s status in the grid is defined by its **state** $s_i \in \mathcal{S}$, which represents its location relative to the **environment**. In the examples with nine cells, the state space is $\mathcal{S} = \{s_1, s_2, \dots, s_9\}$.

From each state, the agent can perform five **actions**: move up, right, down, left or stay put, denoted as a_1, a_2, \dots, a_5 , this set of actions is the action space $\mathcal{A} = \{a_1, \dots, a_5\}$. The available actions can vary by state, for instance, at s_1 , actions a_1 (up) and a_4 (left) would collide with the grid boundary, so the action space is $\mathcal{A}(s_1) = \{a_2, a_3, a_5\}$.

When taking an action, the agent may move from one state to another, a process known as **state transition**. For example, if the agent is at state s_1 and selects action a_2 (moving rightward), it transitions to state s_2 . This process can be represented as:

$$s_1 \xrightarrow{a_2} s_2$$

The state transition process is defined for each state and its associated actions. Mathematically, state transitions are described by conditional probabilities. For example, for s_1 and a_2 , the conditional probability distribution is:

$$\begin{aligned} p(s_1 \mid s_1, a_2) &= 0, \\ p(s_2 \mid s_1, a_2) &= 1, \\ p(s_3 \mid s_1, a_2) &= 0, \\ p(s_4 \mid s_1, a_2) &= 0, \\ p(s_5 \mid s_1, a_2) &= 0, \end{aligned}$$

indicating that taking a_2 at s_1 guarantees the agent moves to s_2 , with a probability of one, and zero probability for other states.

This is a deterministic state transition, but state transitions can also be stochastic, requiring conditional probability distributions. For instance, if random wind gusts affect the grid, taking action a_2 at s_1 might blow the agent to s_5 instead of s_2 , resulting in $p(s_5 \mid s_1, a_2) > 0$.

A **policy** is a function that maps states to actions, indicating the agent's behavior in the environment. In other words, a policy tells the agent what action to take at each state.

Mathematically, policies can be described by conditional probabilities. For example, the policy for s_1 is:

$$\begin{aligned} \pi(a_1 \mid s_1) &= 0, \\ \pi(a_2 \mid s_1) &= 1, \\ \pi(a_3 \mid s_1) &= 0, \\ \pi(a_4 \mid s_1) &= 0, \\ \pi(a_5 \mid s_1) &= 0, \end{aligned}$$

indicating that the probability of taking action a_2 at state s_1 is one, and the probabilities of taking other actions are zero.

The above policy is deterministic, but policies may generally be stochastic. For instance, let's assume that at state s_1 the agent may take actions to move either rightward or downward, each with a probability of 0.5. In this case, the policy for s_1 is:

$$\begin{aligned} \pi(a_1 \mid s_1) &= 0, \\ \pi(a_2 \mid s_1) &= 0.5, \\ \pi(a_3 \mid s_1) &= 0.5, \\ \pi(a_4 \mid s_1) &= 0, \\ \pi(a_5 \mid s_1) &= 0. \end{aligned}$$

After executing an action at a state, the agent receives a reward r as feedback from the environment. The **reward** is a function of the state and action which predicts immediate reward and is denoted as:

$$R(s_t = s, a_t = a) = \mathbb{E}[r_t | s_t = s, a_t = a] \quad (3.2)$$

and it can be positive, negative, or zero. Different rewards influence the policy the agent learns. Generally, a positive reward encourages the agent to take the corresponding action, while a negative reward discourages it.

However we can't find good policies by simply selecting actions with the greatest immediate rewards since they do not consider long-term outcomes. To determine a good policy, we must consider the total reward obtained in the long run and an action with the highest immediate reward may not lead to the greatest total reward.

A **trajectory** is a state-action-reward chain. For example, the agent in our example may follow this trajectory:

$$s_1 \xrightarrow{a_2, r=0} s_2 \xrightarrow{a_3, r=0} s_5 \xrightarrow{a_3, r=0} s_8 \xrightarrow{a_2, r=1} s_9.$$

The **return** of this trajectory is the sum of all rewards collected along it, in the example above:

$$\text{return} = 0 + 0 + 0 + 1 = 1$$

Returns, also called total rewards or cumulative rewards, are used to evaluate policies. Returns can also be defined for infinitely long trajectories, which may diverge. Therefore, we introduce the concept of **discounted return** for infinitely long trajectories. The discounted return is the sum of the rewards from t to T (final step):

$$G_t \doteq R_{t+1} + \gamma R_{t+2} + \gamma^2 R_{t+3} + \dots = \sum_{k=0}^{\infty} \gamma^k R_{t+k+1} \quad (3.3)$$

where $\gamma \in [0, 1]$ is the **discount factor**. If γ is close to 0, the agent emphasizes near-future rewards, resulting in a short-sighted policy. If γ is close to 1, the agent emphasizes far-future rewards.

When T is finite, we call the task episodic and each sequence up to the **terminal state** is an **episode**. Otherwise, we refer to **continuing tasks**.

The **Markov Decision Processes** (MDPs), a general framework for describing stochastic dynamical systems, allows us to formally presents these concepts. The **Markov property** refers to the **memoryless** property of a stochastic process. Mathematically, it means that

$$\begin{aligned} p(s_{t+1} \mid s_t, a_t, s_{t-1}, a_{t-1}, \dots, s_0, a_0) &= p(s_{t+1} \mid s_t, a_t), \\ p(r_{t+1} \mid s_t, a_t, s_{t-1}, a_{t-1}, \dots, s_0, a_0) &= p(r_{t+1} \mid s_t, a_t), \end{aligned} \quad (3.4)$$

where t represents the current time step and $t + 1$ represents the next time step. This indicates that the next state or reward depends only on the current state and action and is independent of the previous ones.

An MDP is defined by a tuple $\langle \mathcal{S}, \mathcal{A}, P, R, \gamma \rangle$, let us break down the components:

- \mathcal{S} : finite set of Markov states s
- \mathcal{A} : finite set of actions a
- P : state transition model for each action, a probability matrix that specifies

$$p(s_t + 1 = s' | s_t = s, a_t = a)$$

- R : reward function

$$R(s_t = s, a_t = a) = \mathbb{E}[r_t | s_t = s, a_t = a]$$

- γ : discount factor $0 \leq \gamma \leq 1$

When the policy in a MDP is fixed, it reduces to a **Markov Process** (MP), this transformation simplifies the MDP by eliminating the decision-making aspect. A Markov process is referred to as a **Markov Chain** if it operates in discrete time and the number of states is either finite or countable.

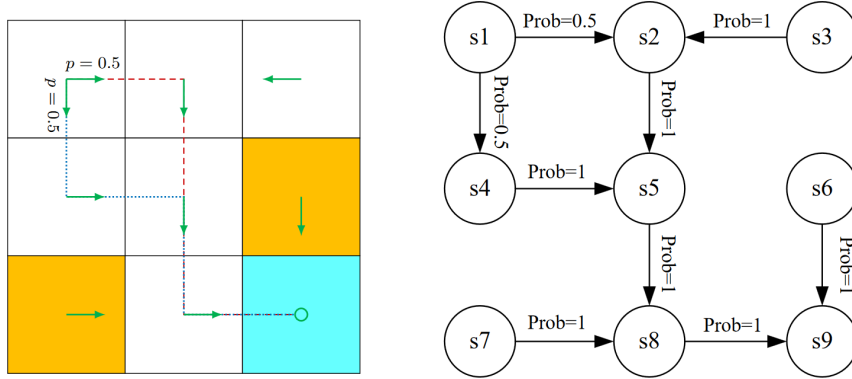


Figure 3.4: The example grid as a Markov Process graph, where the nodes represent the states and the edges represent the state transitions.

Source: [Zhao, 2024]

3.2.1 Reinforcement Learning Algorithms

In [OpenAI, 2018], the landscape of algorithms in modern reinforcement learning is explored.

The algorithms used in reinforcement learning almost always rely on **value functions**. The **value** is the expected return if the agent starts in that state or state-action pair and then follows a specific policy indefinitely.

There are four primary value functions to consider:

- **On-Policy Value Function, $V^\pi(s)$:** This function represents the expected return if the agent starts in state s and acts according to the policy π :

$$V^\pi(s) = \mathbb{E}_{\tau \sim \pi} [R(\tau) \mid s_0 = s] \quad (3.5)$$

- **On-Policy Action-Value Function, $Q^\pi(s, a)$:** This function gives the expected return if the agent starts in state s , takes an arbitrary action a , and then always follows the policy π :

$$Q^\pi(s, a) = \mathbb{E}_{\tau \sim \pi} [R(\tau) \mid s_0 = s, a_0 = a] \quad (3.6)$$

- **Optimal Value Function, $V^*(s)$:** This function provides the expected return if the agent starts in state s and follows the optimal policy for the environment:

$$V^*(s) = \max_{\pi} \mathbb{E}_{\tau \sim \pi} [R(\tau) \mid s_0 = s] \quad (3.7)$$

- **Optimal Action-Value Function, $Q^*(s, a)$:** This function represents the expected return if the agent starts in state s , takes an arbitrary action a , and then follows the optimal policy for the environment:

$$Q^*(s, a) = \max_{\pi} \mathbb{E}_{\tau \sim \pi} [R(\tau) \mid s_0 = s, a_0 = a] \quad (3.8)$$

Figure 3.5 presents a partial taxonomy of the modern reinforcement learning algorithms. One key distinction is whether the agent utilizes (or learns) a model of the environment. A model, in this context, refers to a function that predicts state transitions and rewards.

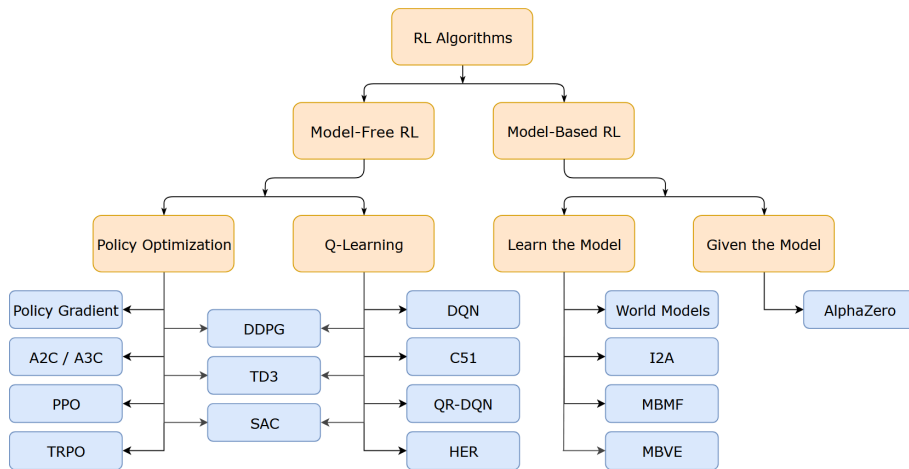


Figure 3.5: Partial taxonomy of algorithms in modern RL.

Source: [OpenAI, 2018]

The primary advantage of having a model is that it enables the agent to plan by predicting future states and rewards. However, the major drawback is that an accurate model of the environment is often unavailable to the agent. Algorithms which use a model are known as **model-based** methods, while those that do not are referred to as **model-free** methods. Our focus will be on the latter.

Model-free methods can be categorized into two main approaches for representing and training agents:

- **Policy Optimization:** explicitly represents a policy as $\pi_\theta(a | s)$ and optimizes the parameters θ either directly via gradient ascent on the performance objective $J(\pi_\theta)$ or indirectly by maximizing local approximations of $J(\pi_\theta)$. Typically, this optimization is performed **on-policy**, meaning that each update uses data collected while the agent is acting according to the most recent version of the policy.
- **Q-Learning:** involves learning an approximator $Q_\theta(s, a)$ for the optimal action-value function $Q^*(s, a)$. Unlike policy optimization, Q-learning is generally performed **off-policy**, allowing updates to use data collected at any time during training, irrespective of the agent's policy at the time of data collection. The corresponding policy is derived from the relationship between Q^* and π^* , where the actions taken by the Q-learning agent are determined by

$$a(s) = \arg \max_a Q_\theta(s, a) \quad (3.9)$$

An example of a Q-learning algorithm is Deep Q-Network (**DQN**), which uses a deep neural network to approximate the optimal action-value function in environments with large state spaces, and its variants such as Double DQN (**DDQN**) and Dueling Double DQN (**D3QN**).

There are also hybrid approaches that combine elements of both policy optimization and Q-learning. In this spectrum of algorithms we can find **actor-critic methods**, which consist of two components: an **actor** that makes actions and a **critic** that evaluates them. An example of an actor-critic algorithm is Twin-Delayed DDPG (**TD3**).

If samples are collected during training the reinforcement learning is considered **online**, otherwise, if the training set is fixed, it is **offline**.

3.3 Generative Adversarial Networks

Generative Adversarial Networks (GANs) is a framework for estimating generative models via an adversarial process, initially introduced by [Goodfellow et al., 2014]. This framework involves training two models: a **generative** model G , which captures the data distribution, and a **discriminative** model D , which predicts samples as either originating from the true data distribution or the generative model. The

training objective for G is to maximize the likelihood of D making incorrect classifications. This adversarial process can be conceptualized as a minimax two-player game. In the theoretical space of arbitrary functions G and D , a unique solution exists where G accurately recovers the training data distribution and D outputs $\frac{1}{2}$ for all inputs (in other words, it reaches maximum uncertainty). When G and D are defined by multilayer perceptrons, the entire system can be trained using backpropagation.

To learn the generator's distribution p_G over data x , we define a prior $p_z(z)$ on input noise variables $z \in \Omega_Z$, called **latent distribution**, and represent a mapping to the data space as $G(z; \theta_g)$, where $G : \Omega_Z \rightarrow \Omega_X$ is a differentiable function parameterized by θ_g . Additionally, we define a second network $D(x; \theta_d)$ that outputs a single scalar, representing the probability that x originated from the true data distribution p_x rather than from p_G . The discriminative model D is trained to maximize the probability of correctly labeling both the training examples and the samples generated by G . Simultaneously, the generative model G is trained to minimize $\log(1 - D(G(z)))$.

Formally, D and G engage in the following two-player minimax game with the value function $V(G, D)$:

$$\min_G \max_D V(D, G) = \mathbb{E}_{x \sim p_x(x)} [\log D(x)] + \mathbb{E}_{z \sim p_z(z)} [\log(1 - D(G(z)))]. \quad (3.10)$$

We can see a graphical representation of the GAN framework in Figure 3.6.

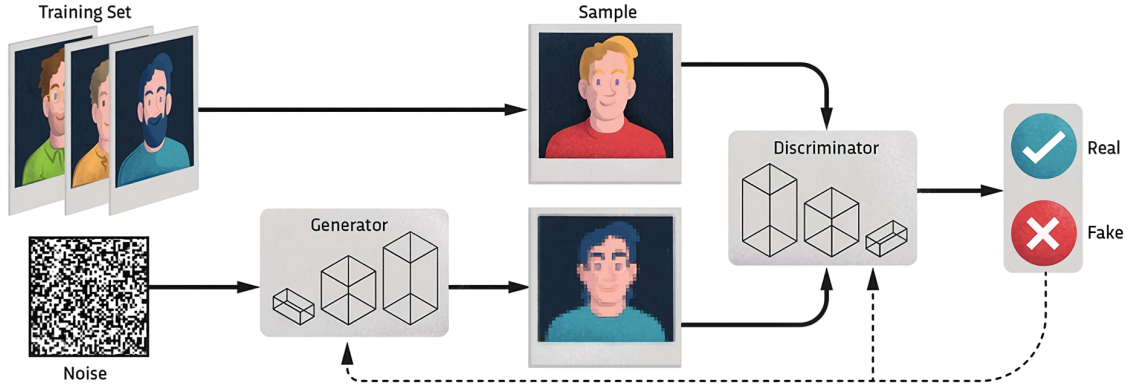


Figure 3.6: GAN framework applied to human faces generation task

Source: Yashwant Singh Kaurav

3.3.1 Bidirectional GANs

Generative Adversarial Networks (GANs) have the potential to be used for unsupervised learning of rich feature representations across various data distributions. However, an obvious issue arises because the GAN framework inherently lacks an

inverse mapping from generated data back to the latent representation. This limitation means that while the generator can map latent samples to generated data, there is no mechanism to map data back to the latent space.

To address this issue, [Donahue et al., 2017] introduced a novel framework called Bidirectional Generative Adversarial Networks (BiGAN). The BiGAN framework, whose architecture is illustrated in Figure 3.7, enhances the standard GAN model (as introduced by [Goodfellow et al., 2014]) by incorporating an **encoder**. This encoder, denoted as $E : \Omega_X \rightarrow \Omega_Z$, maps data x to latent representations z , complementing the generator G .

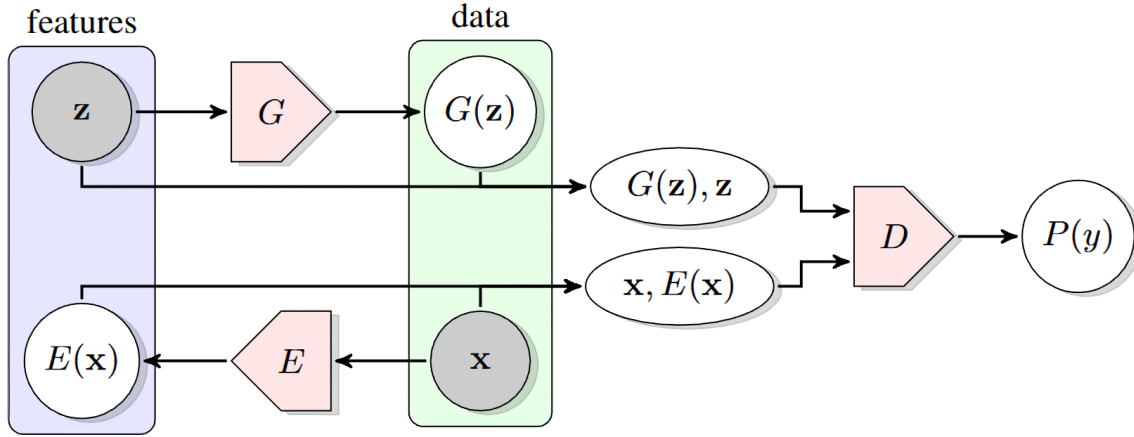


Figure 3.7: BiGAN architecture with generator, discriminator, and encoder.

Source: [Donahue et al., 2017]

In BiGAN, the discriminator D operates not only in the data space (distinguishing between x and $G(z)$) but also jointly in the data and latent spaces. Specifically, the discriminator evaluates pairs of data and their corresponding latent representations, distinguishing between real $(x, E(x))$ and generated $(G(z), z)$. Here, the latent component is either an encoder output $E(x)$ or a generator input z .

An essential aspect of BiGAN is that the encoder E is designed to learn an inverse mapping of the generator G . Despite the encoder and generator not directly interacting, since $E(G(z))$ is not explicitly computed and the generator does not use $E(x)$, the framework ensures that the encoder effectively inverts the generator's mapping.

The training objective for BiGANs is formulated as a minimax problem involving the generator G , the encoder E , and the discriminator D . This objective can be written as:

$$\min_{G, E} \max_D V(D, E, G)$$

where the value function $V(D, E, G)$ is defined as:

$$V(D, E, G) := \mathbb{E}_{x \sim p_X} \underbrace{\left[\mathbb{E}_{z \sim p_E(\cdot|x)} [\log D(x, z)] \right]}_{\log D(x, E(x))} + \mathbb{E}_{z \sim p_Z} \underbrace{\left[\mathbb{E}_{x \sim p_G(\cdot|z)} [\log(1 - D(x, z))] \right]}_{\log(1 - D(G(z), z))}. \quad (3.11)$$

In simpler terms, this objective consists of two components:

1. The expectation over the data distribution p_X and the encoder's latent space distribution $p_E(\cdot|x)$, which aims to maximize $\log D(x, E(x))$.
2. The expectation over the prior distribution p_Z and the generator's data distribution $p_G(\cdot|z)$, which aims to maximize $\log(1 - D(G(z), z))$.

The optimization of this minimax objective is performed using an alternating gradient-based approach, similar to the method introduced by [Goodfellow et al., 2014] for GANs.

3.3.2 Bidirectional Conditional GANs

Conditional GAN (cGAN) ([Mirza and Osindero, 2014]) is a variant of standard GANs designed to enable the conditional generation of data samples based on both latent variables (intrinsic factors) and known auxiliary information (extrinsic factors) such as class labels or associated data from other modalities. However, cGANs fails to achieve several key properties:

1. The ability to disentangle intrinsic and extrinsic factors during the generation process.
2. The ability to separate the components of extrinsic factors from each other, ensuring that the inclusion of one factor minimally impacts the others.

[Jaiswal et al., 2018] introduced the Bidirectional Conditional GAN (BiCoGAN). BiCoGAN enhances the cGAN framework by simultaneously training an encoder along with the generator and discriminator. This encoder learns inverse mappings of data samples to both intrinsic and extrinsic factors, thereby overcoming deficiencies in prior approaches. In Figure 3.8, we present the architecture of BiCoGAN.

In the BiCoGAN framework, the generator $G(\tilde{z}; \theta_G)$ learns a mapping from the distribution $p_{\tilde{Z}}$, where $\tilde{z} = [z, c]$, to p_G , with the goal of making p_G approximate p_X . Concurrently, the encoder $E(x; \theta_E)$ learns a mapping from p_X to p_E , aiming to make p_E approximate $p_{\tilde{Z}}$. The discriminator D evaluates real or fake decisions using pairs $(\tilde{z}, G(\tilde{z}); \theta_D)$ and $(E(x), x; \theta_D)$.

The encoder in BiCoGAN must effectively learn the inverse mapping from data x to latent variables z and conditions c , just as the generator must incorporate both to produce data samples that can deceive the discriminator. This requirement follows from the invertibility under the optimality theorem of BiGANs. However, achieving this optimality is challenging in practice, especially when the prior vector contains structured or complex information. While intrinsic factors z are sampled from a simple latent distribution, extrinsic factors c , such as class labels or object attributes, have specialized and complex distributions that are more difficult to model.

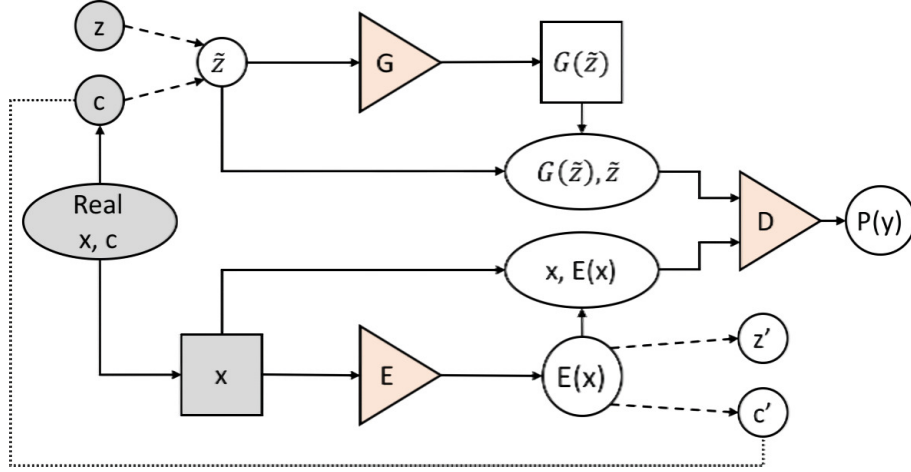


Figure 3.8: BiCoGAN architecture with generator, discriminator, and encoder.

Source: [Jaiswal et al., 2018]

To address this challenge, we introduce the **extrinsic factor loss** (EFL) as a mechanism to guide BiCoGANs in better encoding extrinsic factors. During training, the condition c associated with each real data sample is known and can be used to improve the learning of inverse mappings from x to c . The specific form of EFL depends on the nature of c and the dataset or domain in question.

The BiCoGAN value function can be expressed as:

$$\begin{aligned}
 V(D, G, E) := & \mathbb{E}_{x \sim p_X(x)} [\log D(E(x), x)] \\
 & + \gamma \mathbb{E}_{(x, c) \sim p_X(x, c)} [\text{EFL}(c, E_c(x))] \\
 & + \mathbb{E}_{z \sim p_{\tilde{z}}(\tilde{z})} [\log(1 - D(\tilde{z}, G(\tilde{z})))]
 \end{aligned} \tag{3.12}$$

where γ is the **extrinsic factor loss weight** (EFLW), defined as:

$$\gamma = \min(\alpha e^{\rho t}, \phi)$$

Here, α is the initial value of γ , ϕ is its maximum value, ρ controls the rate of exponential increase, and t indicates the number of epochs the model has been trained.

3.3.3 Wasserstein GANs

In a Generative Adversarial Network (GAN), the interaction between the generator G and the discriminator D is formalized as a minimax optimization problem:

$$\min_G \max_D \mathbb{E}_{x \sim p_X} [\log(D(x))] + \mathbb{E}_{\tilde{x} \sim p_G} [\log(1 - D(\tilde{x}))], \tag{3.13}$$

where p_X represents the real data distribution, and p_G represents the model distribution implicitly defined by $\tilde{x} = G(z)$, with $z \sim p(z)$. If the discriminator is optimally trained before each update of the generator's parameters, minimizing this

objective corresponds to minimizing the Jensen-Shannon divergence between p_X and p_G . However, this often leads to vanishing gradients as the discriminator saturates.

An alternative approach involves using the **Earth-Mover distance** (also known as the Wasserstein-1 distance), $W(q, p)$, which measures the minimum cost of transporting mass to transform distribution q into distribution p . Under mild assumptions, $W(q, p)$ is continuous and differentiable almost everywhere.

The **Wasserstein GAN** (WGAN) ([Gulrajani et al., 2017]) modifies the value function using the Kantorovich-Rubinstein duality:

$$\min_G \max_{D \in \mathcal{D}} \mathbb{E}_{x \sim p_X} [D(x)] - \mathbb{E}_{\tilde{x} \sim p_G} [D(\tilde{x})],$$

where \mathcal{D} is the set of 1-Lipschitz functions, and p_G is again the model distribution defined by $\tilde{x} = G(z)$, with $z \sim p(z)$. When the discriminator (referred to as the **critic** in this context) is optimal, minimizing the value function with respect to the generator's parameters minimizes $W(p_X, p_G)$.

The WGAN value function leads to a critic function with more well-behaved gradients with respect to its input, facilitating easier optimization of the generator. Empirically, the WGAN value function has been observed to correlate better with sample quality compared to the traditional GAN value function.

To enforce the Lipschitz constraint on the critic, a **gradient penalty** is used. A differentiable function is 1-Lipschitz if its gradients have a norm of at most 1 everywhere. Therefore, the gradient norm of the critic's output with respect to its input is directly constrained. The new objective is:

$$L = \mathbb{E}_{\tilde{x} \sim p_G} [D(\tilde{x})] - \mathbb{E}_{x \sim p_X} [D(x)] + \lambda \mathbb{E}_{\hat{x} \sim p_{\hat{X}}} [(\|\nabla_{\hat{x}} D(\hat{x})\|_2 - 1)^2],$$

where the first two terms represent the original critic loss and the third term is the gradient penalty.

3.4 Causal Inference

The **causal inference** is a discipline that studies the relationships of cause-and-effect between variables. Specifically, it is concerned with inferring the causes of an observed phenomenon, distinguishing between **correlation** and **causality**. We are going to explore the basic concepts of causal inference as described in [Neal, 2020].

3.4.1 Correlation does not imply Causation

Consider the following scenario: you come across some data showing a correlation between wearing shoes to bed and waking up with a headache. It appears that most people who wear shoes to bed tend to wake up with a headache, while those who don't wear shoes to bed typically do not wake up with a headache. It is common for people to interpret such data, where there is an **association**, as indicating that

wearing shoes to bed causes headaches. This is especially true if they are seeking a reason to avoid wearing shoes to bed.

We can explain the association between wearing shoes to bed and waking up with a headache without implying that one causes the other. Both events are actually caused by a **common factor**: drinking the night before. This relationship is depicted in Figure 3.9.

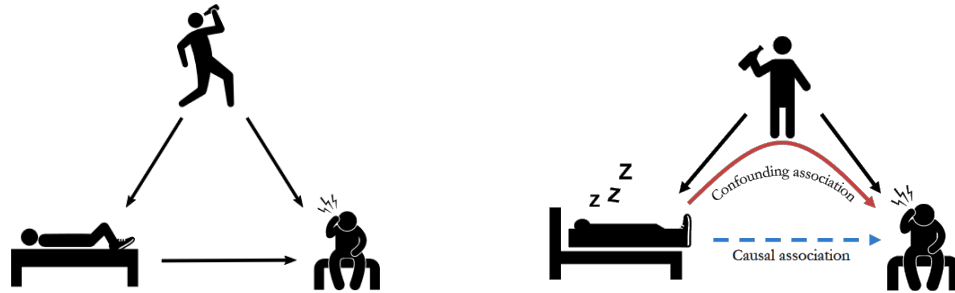


Figure 3.9: A confounding variable causes both wearing shoes to bed and waking up with a headache.

Source: [Neal, 2020]

This kind of variable is often referred to as a **confounder** and we refer to this type of association as a **confounding association** since the observed relationship is influenced by a confounder.

3.4.2 The Flow of Association and Causation in Graphs

In the context of modeling associations between variables, we utilize Directed Acyclic Graphs (DAGs). The following assumptions are pertinent to this modeling approach:

- **Minimality Assumption**: composed of:
 - **Local Markov Assumption** (LMA): Each node is conditionally independent of its non-descendants given its parents.
 - **Dependency of Adjacent Nodes**: Nodes that are adjacent in the DAG are dependent.
- **Bayesian Network Factorization** (BNF): The joint probability distribution can be factorized as

$$P(x_1, \dots, x_n) = \prod_i P(x_i \mid \text{pa}_i) \quad (3.14)$$

where pa_i denotes the parents of node x_i . If P factorizes G , then P is Markovian with respect to G , in fact the Local Markov Assumption is valid if and only if the Bayesian Network Factorization is also valid.

- **Causal Edges Assumption**: In a DAG, every parent is a direct cause of all its children.

But what is a **cause**? A variable X is said to be a cause of a variable Y if X can change in response to changes in Y .

In Figure 3.10, we can see how implementing each one of these assumptions leads to having different relationships between the variables in the graph.

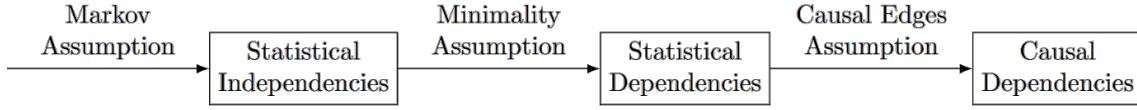


Figure 3.10: Different relationships between variables in a graph based on the assumptions.

Source: [Neal, 2020]

This brings us to the core of this section: understanding the flow of association and causation in DAGs. We can comprehend this flow in general DAGs by analyzing the minimal building blocks of graphs.

By **flow of association**, we mean whether any two nodes in a graph are associated or not associated. In other words, whether two nodes are (statistically) dependent or (statistically) independent. Additionally, we will explore whether two nodes are conditionally independent or not.

Consider a graph consisting of just two unconnected nodes, as depicted in Figure 3.11.



Figure 3.11: Graph with two unassociated nodes.

These nodes are not associated simply because there is no edge between them. This can be demonstrated by considering the factorization of $P(x_1, x_2)$ given by the Bayesian Network Factorization:

$$P(x_1, x_2) = P(x_1)P(x_2)$$

This factorization immediately proves that the two nodes X_1 and X_2 are unassociated (independent) in this basic structure. The key assumption here is that P is Markov with respect to the graph in Figure 3.11.

In contrast, if there is an edge between the two nodes (as in Figure 3.12), then the nodes are associated. The assumption used here is the causal edges assumption,

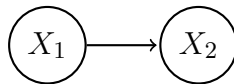


Figure 3.12: Graph with two nodes X_1 and X_2 with an arrow from X_1 to X_2 , indicating that X_1 is a cause of X_2 .

which indicates that X_1 is a cause of X_2 . Since X_1 is a cause of X_2 , X_2 must be able to change in response to changes in X_1 , establishing their association. Generally, any time two nodes are adjacent in a causal graph, they are associated.

Let's now consider more the building blocks of DAGs as shown in Figure 3.13, where we have a **chain**, a **fork** and an **immorality**.

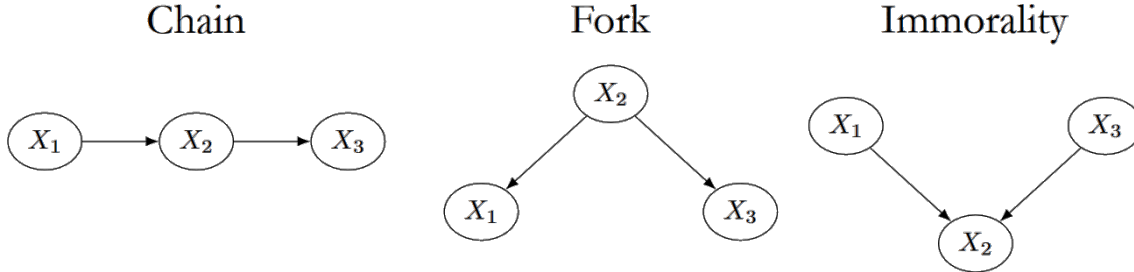


Figure 3.13: Building blocks of DAGs: chain, fork, and immorality.

Source: [Neal, 2020]

Chains and forks share the same set of dependencies. In both structures, X_1 and X_2 are dependent, and X_2 and X_3 are dependent for the same reason discussed for the graph in Figure 3.12. Adjacent nodes are always dependent when we make the causal edges assumption. But does association flow from X_1 to X_3 through X_2 in chains and forks?

In chain graphs, X_1 and X_3 are usually dependent simply because X_1 causes changes in X_2 , which then causes changes in X_3 . In a fork graph, X_1 and X_3 are also usually dependent because the value that X_2 takes on determines both the value that X_1 takes on and the value that X_3 takes on. In other words, X_1 and X_3 are associated through their shared common cause. However, while in chains the association between X_1 and X_3 is a causal one, in forks it is not.

Chains and forks also share the same set of independencies. When we condition on X_2 in both graphs, it blocks the flow of association from X_1 to X_3 . This is due to the local Markov assumption; each variable can locally depend only on its parents. Therefore, when we condition on X_2 (the parent of X_3 in both graphs), X_3 becomes independent of X_1 (and vice versa).

We refer to this independence as an instance of a blocked path. These blocked paths are illustrated in Figure 3.14.

In contrast to chains and forks, in an immorality, $X_1 \perp X_3$. We observe that conditioning on a collider can turn a blocked path into an unblocked path. The parents X_1 and X_3 are not associated in the general population, but when we condition on their shared child X_2 taking on a specific value, they become associated. Conditioning on the collider X_2 allows association to flow along the path $X_1 \rightarrow X_2 \leftarrow X_3$, despite the fact that it does not when we do not condition on X_2 . In Figure 3.15, we can see the immorality with the blocked path and the unblocked path.

Conditioning on **descendants of a collider** also induces association between the parents of the collider. The intuition is that if we learn something about a collider's

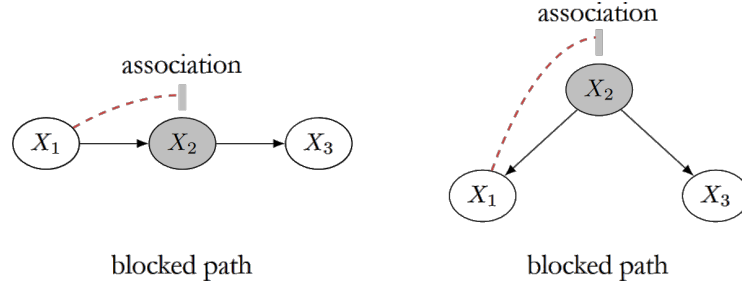


Figure 3.14: Blocked paths in chain and fork graphs.

Source: [Neal, 2020]

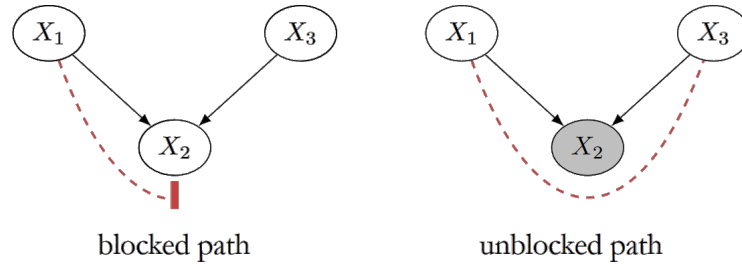


Figure 3.15: Immortality with a blocked path and an unblocked path.

Source: [Neal, 2020]

descendant, we usually also learn something about the collider itself because there is a direct causal path from the collider to its descendants. In Figure 3.16, we can see a conditioning on a descendant of a collider.

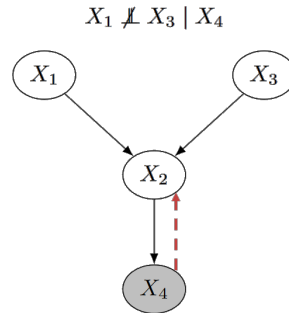


Figure 3.16: Conditioning on a descendant of a collider.

Source: [Neal, 2020]

Now let's formally codify the concept of a **blocked path**: a path between nodes X and Y is **blocked** by a (potentially empty) conditioning set Z if either of the following is true:

1. Along the path, there is a chain $\cdots \rightarrow W \rightarrow \cdots$ or a fork $\cdots \leftarrow W \rightarrow \cdots$, where W is conditioned on ($W \in Z$).
2. There is a **collider** W on the path that is not conditioned on ($W \notin Z$) and none of its descendants are conditioned on ($de(W) \not\subseteq Z$).

An **unblocked path** is simply a path that is not blocked. The graphical intuition to have in mind is that **association** flows along unblocked paths and does not flow along blocked paths.

Now, we are ready to introduce **d-separation**: two (sets of) nodes X and Y are **d-separated** by a set of nodes Z if all of the paths between (any node in) X and (any node in) Y are blocked by Z ([Pearl, 1988]).

Otherwise, they are **d-connected**.

3.4.3 Potential Outcomes and the Fundamental Problem of Causal Inference

The central issue driving the need for causal inference is that correlation does not imply causation. If these two concepts were equivalent, causal inference would be easy. This raises the critical question: how can we determine when one event causes another and, therefore, infer causality?

To address this, we introduce the concept of **potential outcomes**. Suppose, as shown in Figure 3.17, you have a headache, and you know that taking a pill would alleviate the headache, whereas not taking the pill would result in the headache persisting. In this scenario, you can infer a causal effect of the pill on the headache. However, if not taking the pill also resulted in the headache disappearing, you would conclude that there is no causal effect. This is the intuition behind potential outcomes.

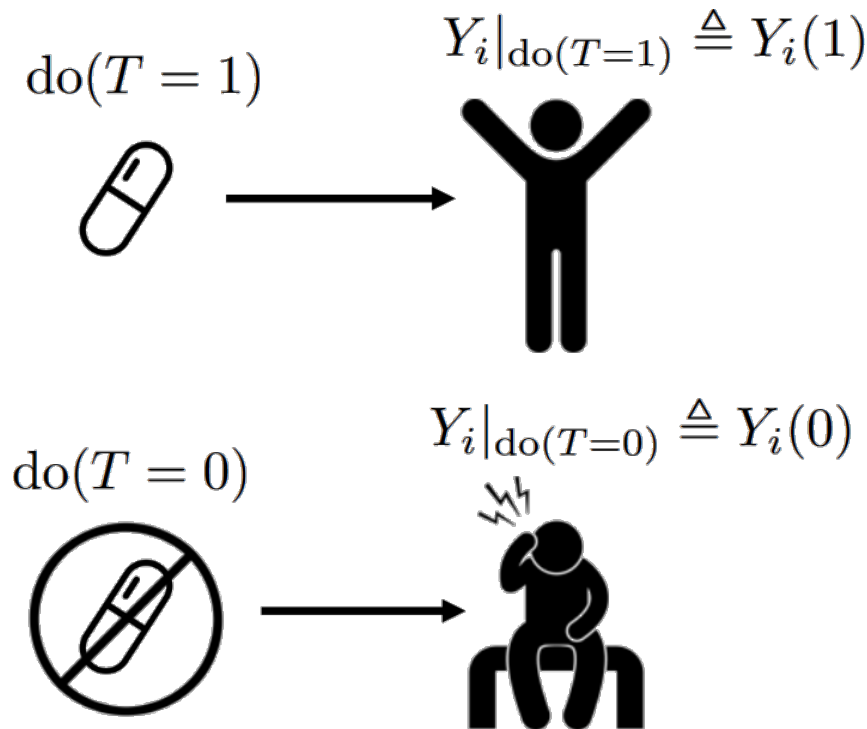


Figure 3.17: Potential outcomes for a causal effect.

Source: [Neal, 2020]

Let's define some notation:

- T : Observed treatment
- Y : Observed outcome
- i : Denotes a specific unit or individual
- $Y_i|_{\text{do}(T=1)} \triangleq Y_i(1)$: Potential outcome under treatment
- $Y_i|_{\text{do}(T=0)} \triangleq Y_i(0)$: Potential outcome under no treatment
- **Causal Effect** = $Y_i(1) - Y_i(0)$

To account for individual differences, we measure the **Average Treatment Effect** (ATE), given by:

$$\mathbb{E}[Y(1)] - \mathbb{E}[Y(0)]. \quad (3.15)$$

A fundamental problem in causal inference is the challenge of distinguishing between the observational distribution $P(Y|T = t)$ and the interventional distribution $P(Y|\text{do}(T = t))$. These distributions are generally not equal; if they were, correlation would indeed imply causation. The intervention $T = t$ on a population is denoted by the do-operator, which allows us to measure the causal effect from the interventional distribution $P(Y|\text{do}(T = t))$.

However, we often cannot intervene on the entire population and can only access the observational distribution. The distinction between these distributions is illustrated in Figure 3.18. Intervening with $\text{do}(T = 0)$ (**factual**) usually precludes access to $\text{do}(T = 1)$ (**counterfactual**).

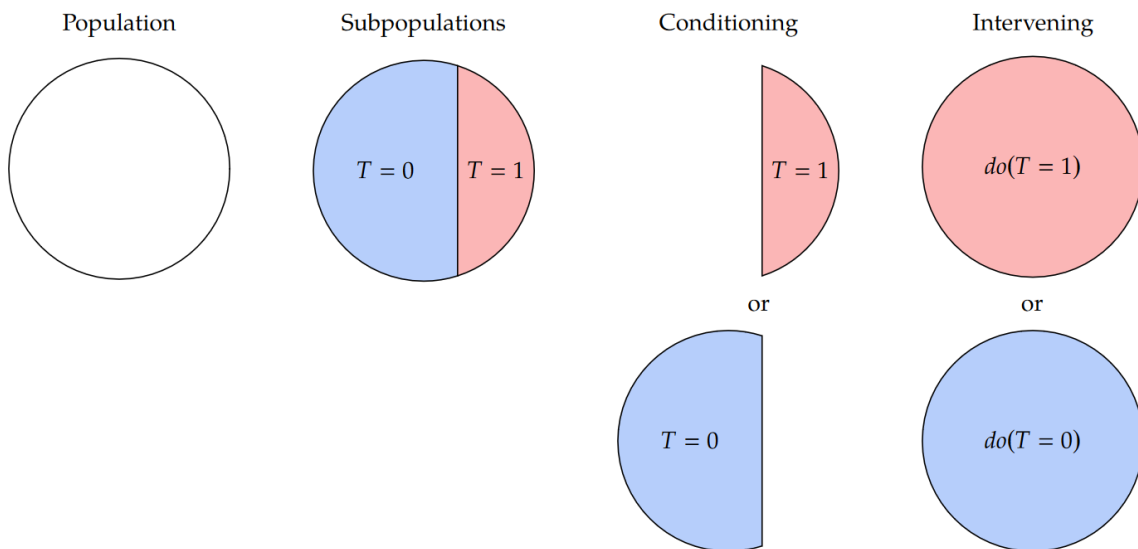


Figure 3.18: The difference between observational and interventional distributions.

Source: [Neal, 2020]

One approach to address this issue is through a **Randomized Controlled Trial** (RCT). In an RCT, participants are randomly assigned to treatment or control groups, ensuring that $(Y(1), Y(0)) \perp\!\!\!\perp T$, under the assumption of exchangeability. **Exchangeability** means that the treatment groups are comparable in the sense that if they were swapped, the new treatment group would exhibit the same outcomes as the original treatment group, and the new control group would exhibit the same outcomes as the original control group. The randomization process ensures that, as we can see in Figure 3.19, the causal association between the treatment and the outcome is identifiable since there is no confounding due to a missing connection between the treatment and the cause.

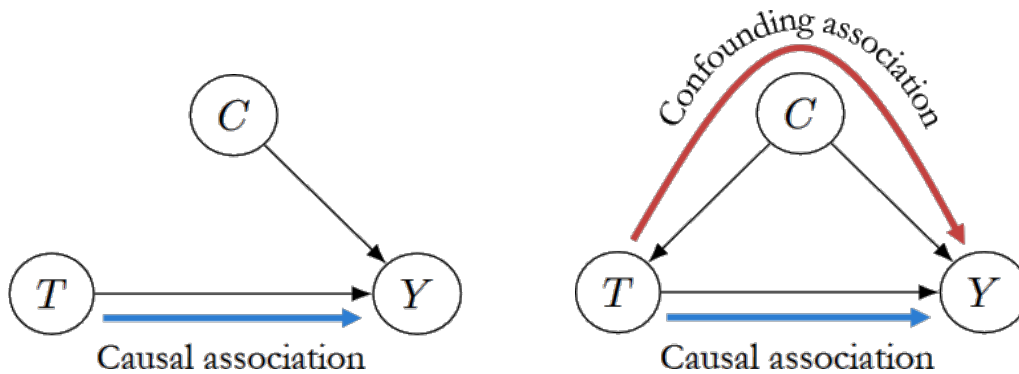


Figure 3.19: Randomized Controlled Trial (RCT) design vs. Observational Study.

Source: [Neal, 2020]

However, it is not always feasible to randomize treatment due to several reasons:

- **Ethical reasons:** For instance, it would be unethical to randomize people to smoke in order to measure the effect on lung cancer.
- **Infeasibility:** It is impractical to randomize countries into different economic systems (e.g. communist vs. capitalist) to measure the effect on GDP.
- **Impossibility:** Certain changes, such as altering a person's DNA at birth to measure the effect on breast cancer, are simply not possible.

We must then find a way to estimate causal effects from observational data.

3.4.4 Causal Models and Identification

Identification is the process of moving from a **causal estimand** to a **statistical estimand**. As we have seen in 3.4.3, conditioning on $T = t$ means that we are restricting our focus to the subset of the population who received treatment t . In contrast, an **intervention** (denoted with the **do-operator**) involves taking the entire population and giving everyone treatment t .

Interventional distributions such as $P(Y \mid \text{do}(T = t))$ are conceptually quite different from the **observational distribution** $P(Y)$ since the latter do not include

the do-operator. Because of this, we can observe data from them without conducting any experiments. This is why we call data from $P(Y, T, X)$ **observational data**. If we can reduce an expression Q with do in it (an interventional expression) to one without do in it (an observational expression), then Q is said to be **identifiable**. We will refer to an estimand as a **causal estimand** when it contains a do-operator, and as a **statistical estimand** when it does not contain a do-operator.

Before describing a very important assumption, we must specify what a **causal mechanism** is, we will refer to the causal mechanism that generates X_i as the conditional distribution of X_i given all of its causes: $P(x_i \mid \text{pa}_i)$.

To achieve many causal identification results, the main assumption we will make is that interventions are local: intervening on a variable X_i only changes the causal mechanism for X_i . In this sense, the causal mechanisms are **modular**.

Assumption of Modularity / Independent Mechanisms / Invariance:

If we intervene on a set of nodes $S \subseteq [n]^1$, setting them to constants, then for all i , we have the following:

1. If $i \notin S$, then $P(x_i \mid \text{pa}_i)$ remains unchanged.
2. If $i \in S$, then $P(x_i \mid \text{pa}_i) = 1$ if x_i is the value that X_i was set to by the intervention; otherwise, $P(x_i \mid \text{pa}_i) = 0$.

The modularity assumption allows us to encode many different interventional distributions in a single graph. For example, it could be the case that $P(Y)$, $P(Y \mid \text{do}(T = t))$, $P(Y \mid \text{do}(T = t'))$, and $P(Y \mid \text{do}(T_2 = t_2))$ are all completely different distributions that share almost nothing. If this were the case, each of these distributions would need its own graph. However, by assuming modularity, we can encode them all with the same graph that we use to encode the joint $P(Y, T, T_2, \dots)$, and we can know that all of the factors (except the ones that are intervened on) are shared across these graphs.

The **causal graph** for interventional distributions, as shown in an example in Figure 3.20, is simply the same graph used for the observational joint distribution, but with all of the edges to the intervened node(s) removed. This is because the probability for the intervened factor has been set to 1, allowing us to ignore that factor.

Recall from subsection 3.4.2, that **causal association** flows from T to Y along directed paths, while **non-causal association** can flow along other paths from T to Y unless they are blocked by either a non-collider that is conditioned on or a collider that is not conditioned on. These non-directed unblocked paths from T to Y are known as **backdoor paths** because they have an edge that goes in the “backdoor” of the T node. By conditioning on certain variables, we can block these paths and identify causal quantities like $P(Y \mid \text{do}(T = t))$.

Our goal is to transform the causal estimand $P(y \mid \text{do}(T = t))$ into a statistical estimand that relies only on the observational distribution. We start by assuming

¹We use $[n]$ to refer to the set $\{1, 2, \dots, n\}$.

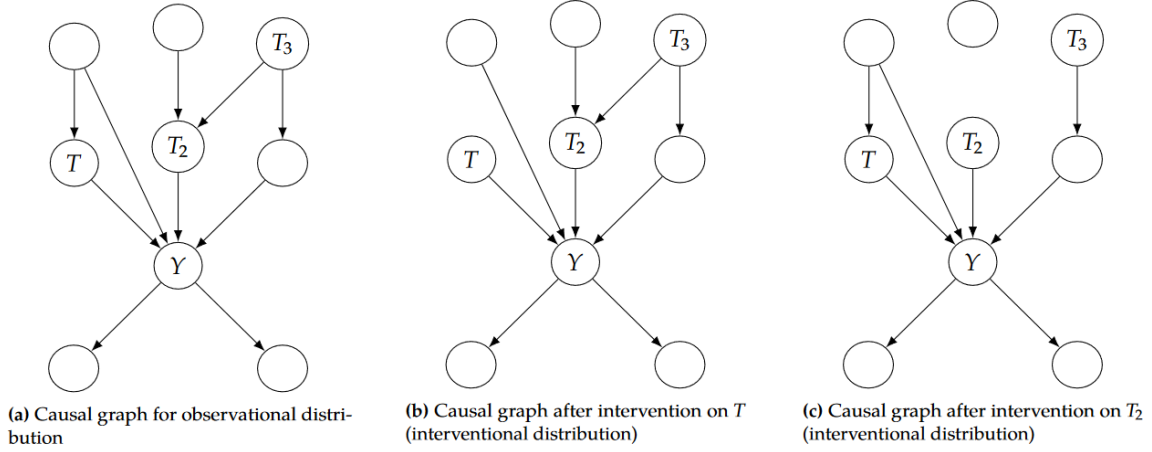


Figure 3.20: Intervention as edge deletion in causal graphs.

Source: [Neal, 2020]

we have a set of variables W that satisfy the **backdoor criterion**:

A set of variables W satisfies the backdoor criterion relative to T and Y if the following are true:

1. W blocks all backdoor paths from T to Y .
2. W does not contain any descendants of T .

When W satisfies the backdoor criterion, it becomes a **sufficient adjustment set**. Additionally, we must ensure **positivity**, which means that all subgroups of the data with different covariates have some probability of receiving any value of treatment. Formally, we define **positivity** for binary treatment as follows:

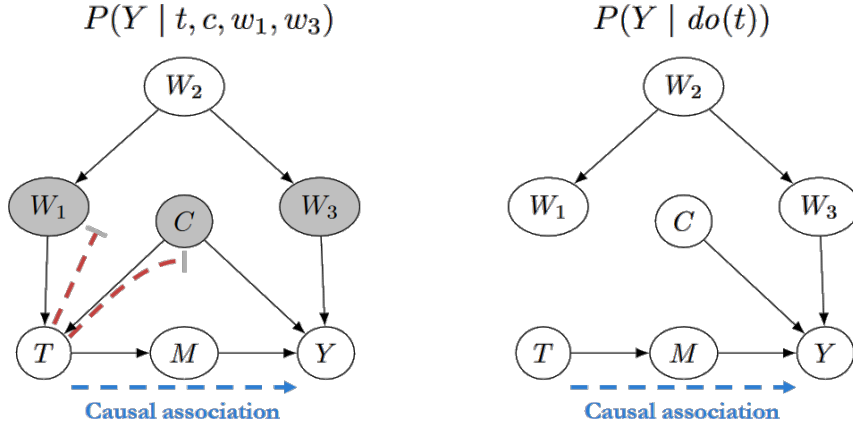
For all values of covariates x present in the population of interest (i.e., x such that $P(X = x) > 0$),

$$0 < P(T = 1 \mid X = x) < 1$$

Backdoor Adjustment: Given the modularity assumption, that W satisfies the backdoor criterion, and positivity, we can identify the causal effect of T on Y :

$$P(y \mid \text{do}(T = t)) = \sum_w P(y \mid T = t, W = w)P(w) \quad (3.16)$$

This theorem connects to **d-separation**. We can use the backdoor adjustment if W **d-separates** T from Y in the manipulated graph. Recall that isolating the causal association means identifying it. We achieve this if T is d-separated from Y in the manipulated graph or if Y is d-separated from T in the manipulated graph, conditional on W . In Figure 3.21, we can see how the backdoor paths that are blocked by conditioning on W allow us to identify the causal effect of T on Y .

Figure 3.21: Backdoor paths blocked by conditioning on W .

Source: [Neal, 2020]

3.4.5 Structural Causal Models

We need to be able to state that A is a **cause** of B , meaning that changing A results in changes in B , but changing B does not result in changes in A . This relationship is represented by the following **structural equation**:

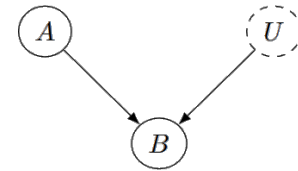
$$B := f(A) \quad (3.17)$$

where f is some function that maps A to B .

However, the mapping between A and B in Equation 3.17 is **deterministic**. Ideally, we'd like to allow it to be **probabilistic**, which allows room for some unknown causes of B that factor into this mapping. We can then write the following:

$$B := f(A, U) \quad (3.18)$$

where U is some unobserved random variable. The graph for this simple structural equation is shown in Figure 3.22.

Figure 3.22: Graph of 3.18. The dashed node U means that U is unobserved.

Source: [Neal, 2020]

While we have shown a single **structural equation** in Equation 3.18, there can be a large collection of structural equations in a single model, commonly labeled M . For example, we write structural equations for the causal model in Figure 3.23

below:

$$M : \begin{cases} B := f_B(A, U_B) \\ C := f_C(A, B, U_C) \\ D := f_D(A, C, U_D) \end{cases} \quad (3.19)$$

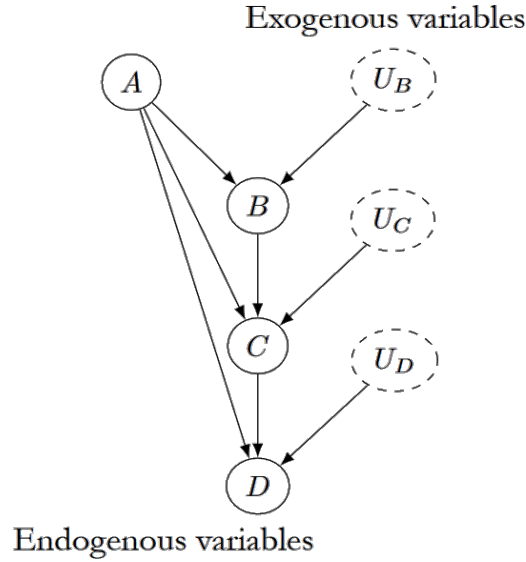


Figure 3.23: Graph of 3.19.

Source: [Neal, 2020]

The variables for which we write structural equations are known as **endogenous variables**. These are the variables whose causal mechanisms we are modeling. In contrast, **exogenous variables** are variables that do not have any parents in the causal graph.

Structural Causal Model (SCM): a structural causal model is a tuple consisting of the following sets:

1. A set of endogenous variables V
2. A set of exogenous variables U
3. A set of functions f , one to generate each endogenous variable as a function of other variables

Chapter 4

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