Mobile Robot Self-Driving Through Image Classification Using Discriminative Learning of Sum-Product Networks



Undergraduate Thesis

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Abstract

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Driving has proven to be a very difficult task for machines to emulate, not only due to the inherent complexity of the problem but also because of the need for accurate real-time predictions. Nonetheless, recent advances in computer vision and machine learning have shown promising results in the real-world. Mobile robots are low-cost miniature computers with limited processing power and memory. The problem of self-driving can be similarly applied to the mobile robot domain as a down-scaled version of the same task, with an additional hardware constraint. Sum-product networks are probabilistic graphical models capable of representing tractable probability distributions containing a great number of variables. Exact inference is asymptotically linear to the number of edges in the network's graph, and its deep architecture is capable of representing a wide range of distributions. In this work, we attempt to model autonomous driving by using sum-product networks on a small mobile robot. We model this task as an imitation learning problem through image classification. We present accuracy results on an artificial self-driving dataset for different sum-product network learning algorithms, providing a comparative study not only for different network architectures, but also discriminative and generative models. Finally, we provide a real-world mobile robot implementation on a miniature computer.

Keywords: sum-product networks, probabilistic graphical models, machine learning, robotics

Abbreviations

DAG	Directed	acyclic	graph
-----	----------	---------	-------

- EM Expectation-maximization
- GD Gradient descent
- IV Indicator variable
- MAP Maximum a posteriori probability
- MPE Most probable explanation
- MPN Max-product network
- MST Minimum spanning tree
- PGM Probabilistic graphical model
 - RV Random variable
- SGD Stochastic gradient descent
- SPN Sum-product network
- SPT Sum-product tree

Symbols and notations

- μ Gaussian distribution mean
- σ Gaussian distribution standard deviation
- X Sample space of random variables
- Z Partition function
- ϕ Factor (potential)
- [X = x] Indicator function for random variable valuation X = x
 - Pa(n) Set of parent nodes of node n
 - Ch(n) Set of child nodes of node n
- Val(X) Set of possible values variable X could take

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Chapter 1

Introduction

In this chapter we first describe the motivations and objectives of this thesis. Next, we describe the structure of this document.

1.1 Motivation and objectives

Self-driving is a challenging computer vision task, mainly due to its inherent complexity and the necessity for real-time decision making. Although there have been many promising results the past few years on autonomous driving, the task still relies on the underlying problem of following a pathway through visual cues (usually road markings). A possible approach to this task is through imitation learning by means of image classification. That is, the agent tasked with driving should be able to reliably mimic human behavior by correctly classifying whether to turn, stop or go straight given an image captured in front of the car.

Mobile robots are low cost machines capable of movement. These robots are usually small, and because of their size and cost often don't have the same performance capabilities as a desktop computer. However, these domain traits make mobile robot self-driving a very similar analogue to real-world autonomous cars. Processing power and memory constraints play a big role in this case, and translate well to embedded systems present in a self-driving car.

Sum-product networks (SPNs) are probabilistic graphical models that are able to represent a wide range of tractable probability distributions of many variables. SPNs have shown impressive results in several domains, and particularly that of image classification. Their deep architecture seems to capture features and contexts well, and since inference is computed in time linear to the network's edges, SPNs are promising models for fast inference in self-driving.

In this work, we attempt to model self-driving of mobile robots through image classification. For the task of classification our objective is to use sum-product networks learned discriminatively, though we also give results for generative SPNs, comparing not only generative and discriminative learning, but also different SPN architectures.

1.2 Thesis structure

This thesis is structured as follows. In chapter 2, we first provide background on sum-product networks, where we formally define an SPN, present key properties on their structure, explain how to compute exact inference and find an approximation of the maximum a posteriori probability (MAP).

In chapter 3, we show how to compute the partial derivatives with respect to a sub-SPN and to its weights, leading on how to perform gradient descent and then on learning the weights of the network through gradient descent both generatively and discriminatively.

Chapter 4 is dedicated to algorithms for learning the structure of an SPN. We explain the two structural learning algorithms that were used in the experiments.

For ??, we first show how we model self-driving as an image classification problem. We then specify the architecture of the robot used in the experiments, giving specifications on the hardware and software used. Furthermore, we describe some concepts of control we use for navigation.

In ??, we provide classification results on many image classification datasets from various domains with different learning algorithms. We then describe the self-driving dataset used for training, and give in-dataset accuracies as well as real-world empirical results on the mobile robot itself.

Finally, in ?? we give our conclusions and provide some discussion of the results.

There is an additional section of this thesis in which we give a brief subjective insight on the work done for this thesis. We also list subjects we deemed important for the work done in this thesis.

Furthermore, ?? contains all proofs done in this thesis.

Chapter 2

Sum-product networks

In this chapter we provide some background concepts needed for defining a sumproduct network. Once this is covered, we formally define an SPN, list some interesting properties on their structure, and describe how to perform exact inference (i.e. extract the probability of evidence of some valuation) and how to find an approximation of the maximum a posteriori probability.

We leave all proofs in ??.

2.1 Background

The objective of probabilistic modelling is to compactly represent a probability distribution, be able to find a good approximation to the real function and be able to efficiently compute both the marginals and modes. Probabilistic graphical models (PGMs) attempt to solve this through the use of graphs, representing distributions as a normalized product of factors (PEARL, 1988).

$$P(X = x) = \frac{1}{Z} \prod_{k} \phi_k(x_{\{k\}})$$

Where $x \in \mathcal{X}$ is a d-dimensional vector valuation of RVs X on sample space \mathcal{X} , and factor (also called a potential) ϕ_k is a function mapping instantiations of X to a non-negative number. Z is the partition function $Z = \sum_{x \in \mathcal{X}} \prod_k \phi_k(x_{\{k\}})$ that sums out all variables and normalizes the term above it to the [0, 1] range.

A downside of this representation is that inference is exponential on the worst case, which makes learning also exponential, as it uses inference as a subroutine. To get around this problem, Darwiche proposed in Darwiche, 2003 the notion of *network polynomial*.

A network polynomial is a function over the probabilities of each instantiation. Let $\Phi(x)$ be a probability distribution. The network polynomial of $\Phi(x)$ is the function $f = \sum_{x \in \mathcal{X}} \Phi(x) \Pi(x)$, where $\Pi(x)$ is the product of the IVs of each variable on instantiation x,

where each indicator variable [Y = y] has a value of zero if $Y \neq y$ in x and a value of one otherwise (i.e. if Y = y in x or $Y \notin x$).

As an example, take the bayesian network $\mathcal{N}=A\to B$ with binary variables. Let λ_a , $\lambda_{\overline{a}}$, λ_b and $\lambda_{\overline{b}}$ be the indicator variables for when A=1, A=0, B=1 and B=0 respectively. The network polynomial of \mathcal{N} is the expression

$$f_{\mathcal{N}} = P(a)P(b|a)\lambda_a\lambda_b + P(a)P(\overline{b}|a)\lambda_a\lambda_{\overline{b}} + P(\overline{a})P(b|\overline{a})\lambda_a\lambda_{\overline{b}} + P(\overline{a})P(\overline{b}|\overline{a})\lambda_{\overline{a}}\lambda_{\overline{b}}.$$

The main advantage of this representation is to avoid recomputing terms. For instance, take an instantiation of $x = \{A = 0\}$. Then, the network polynomial will be as follows.

$$f_{\mathcal{N}}(x) = P(a)P(b|a) \cdot 0 \cdot 1 + P(a)P(\overline{b}|a) \cdot 0 \cdot 1 + P(\overline{a})P(b|\overline{a}) \cdot 1 \cdot 1 + P(\overline{a})P(\overline{b}|\overline{a}) \cdot 1 \cdot 1 =$$

$$= P(\overline{a})P(b|\overline{a}) + P(\overline{a})P(\overline{b}|\overline{a})$$

Which means we can avoid computing values from the two first terms. We can also compute the network polynomial of some unnormalized probability distribution as long as we divide by the partition function, defined as the network polynomial with all indicators set to one. Although the network polynomial has exponential size in terms of variables, computing the probability of evidence is linear in its size. By representing the network polynomial as an arithmetic circuit of sums and products, one can prove that the cost of inference is indeed polynomial.

2.2 Definitions and properties

Sum-product networks borrow many concepts from network polynomials and arithmetic circuits. There are many definitions of SPNs and in this thesis we present two. The first definition is given by the seminal article Poon and Domingos, 2011, and can be seen as a more low-level approach to defining the network. The second, based on Gens and Domingos, 2013, is a stronger definition, but one which we will use more throughout this thesis, as it lends itself better to continous data.

Let $X = \{X_1, X_2, \dots, X_n\}$ be the set of all variables. We shall call this set the root scope. Let G be a graph G. The sets of vertices and edges of G will be denoted by V(G) and E(G). We will call Ch(n) and Pa(n) the sets of children and parents of node $n \in V(G)$.

Definition 2.1 (Sum-product network; Poon and Domingos, 2011). A sum-product network (SPN) over variables X_1, X_2, \ldots, X_n is a DAG whose leaves are indicator variables $[X_1 = x_1^1], [X_2 = x_2^1], \ldots, [X_n = x_n^1], \ldots, [X_1 = x_1^d], [X_2 = x_2^d], \ldots, [X_n = x_n^d]$. Its internal nodes are weighted sums or products. Each edge coming out from a sum node n to another node j has a non-negative weight associated with it. We denote such weight by $w_{n,j}$. The value of a sum node n is $v_n = \sum_{j \in Ch(n)} w_{n,j}v_j$, where v_j is the value of node j. The value of a product node n is $v_n = \prod_{j \in Ch(n)} v_j$. The value of a leaf node is the value of the indicator variable. The value of the SPN is the value of its root node.

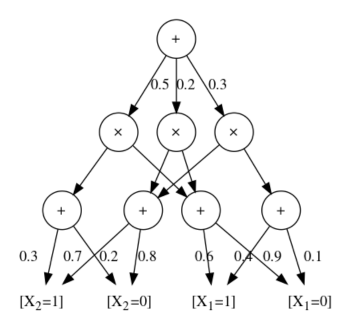


Figure 2.1: An example of an SPN.

Throughout this thesis, we denote by S(X = x) the value of an SPN S given evidence x. A sub-SPN S_n of S is the subgraph of S rooted at n. A node in an SPN is itself an SPN. When all indicator variables are set to one, the value of S is denoted by S(*). The scope of an SPN S, denoted by Sc(S), is the union set of all scopes of its children. A leaf's scope is the scope of its IV.

Definition 2.2 (Validity). An SPN is valid iff, for all evidence E = e, $S(E = e) = \Phi_S(E = e)$, where Φ_S is an unnormalized probability distribution.

Definition 2.3 (Completeness). An SPN is complete iff all children of the same sum node have the same scope.

Definition 2.4 (Consistency). An SPN is consistent iff no variable appears with a value v in one child of a product node and valued u, with $u \neq v$, in another.

Validity in an SPN means that the network correctly and efficiently computes the probability of evidence of the distribution it represents. In this document we only work with valid SPNs, as we wish to always compute exact inference. However, non-valid SPNs are an interesting field of research for approximate inference in SPNs.

A sufficient condition for validity is completeness and consistency. Yet whilst this condition is sufficient, it is not necessary, as the converse (i.e. an incomplete and inconsistent valid SPN) can hold.

Theorem 2.1 (Poon and Domingos, 2011). An SPN is valid if it is complete and consistent.

When an SPN S is valid, then S(*) is the partition function, and we can extract the probability of evidence from an SPN by computing P(X = x) = S(x)/S(*). If for every sum node all of their weights are non-negative and sum to one, then the partition function is S(*) = 1, and the SPN is the distribution itself.

Corollary 2.1 (Validity recursion; Poon and Domingos, 2011). *If an SPN S is valid, then*

all sub-SPN of S is valid.

Definition 2.5 (Decomposability). An SPN is decomposable iff no variable appears in more than one child of a product node.

In other words, an SPN is decomposable if and only if, for every product node, every child node has disjoint scopes with relation to all their other siblings. It is easy to see that decomposability implies in consistency, as there can be no inconsistency between product children since scopes are disjoint. Therefore, a complete and decomposable SPN is valid. Indeed it is much easier to produce decomposable SPNs then only consistent ones, and although this condition may seem too strong and restrictive, Peharz et al., 2015 showed that a consistent SPN is representable by a polynomially larger decomposable SPN.

So far, SPNs are restricted to the discrete domain, as we rely on IVs to define possible valuations to variables. We can generalize SPNs to the continuous by assuming an infinite number of IVs and thus replacing sum nodes whose children are IVs with integral nodes. A leaf node then becomes an integral node with infinite IVs as children. Particularly, it represents an unnormalized univariate probability distribution, such as a Gaussian. The value of this integral node n becomes the pdf $p_n(x)$. This extension brings us to a second definition of SPNs.

Definition 2.6 (Sum-product networks; Gens and Domingos, 2013). A sum-product network is defined recursively as follows.

- 1. A tractable univariate probability distribution is an SPN.
- 2. A product of SPNs with disjoint scopes is an SPN.
- 3. A weighted sum of SPNs with the same scope is an SPN, provided all weights are positive.
- 4. Nothing else is an SPN.

This second definition limits our scope to only complete and decomposable SPNs. Note that an IV is also an SPN, as we can assume that an indicator variable is a degenerate tractable univariate distribution, taking a value of one if it agrees with the given evidence and zero otherwise.

2.3 Inference

Throughout this thesis we assume that all sum nodes are normalized and sum to one, meaning the partition function is S(*) = 1 and the SPN's value is the probability itself.

Let $X = \{X_1 = x_1, X_2 = x_2, \dots, X_k = x_k\}$ be a valuation and S be an SPN. We say that X is a complete valuation if Sc(X) = Sc(S). That is, X contains a valuation for all variables in S. An incomplete valuation has some variable assignment missing.

Computing the probability of evidence is done through a bottom-up backwards pass through the SPN. To find the value of an SPN, we must know the value of the root node, which depends on all nodes below it. This is done through a topological traversal of the graph.

Finding the value of a leaf node depends on the valuation given. Let n be a leaf node, and $Sc(n) = \{X_j\}$. Let X be some valuation. Assuming the univariate probability distribution of n has fpd $p_n(x)$, then if X has a valuation $X_j = x_j$, the value of node n will be $S_n(X) = p_n(x_j)$. If X has no valuation for variable X_j , then $S_n(X)$ is the distribution's mode. Note that this holds for indicator variables, as if X has a valuation for $X_j = x_j$ and the IV matches with x_j , then $p_n(x_j) = 1$. In case it does not, $p_n(x_j) = 0$. For the incomplete case, the mode of an indicator variable is one, which holds the equivalence.

Once we compute leaf nodes, we can compute each internal node's values by following the topological order until we reach the root. For sum nodes, we compute the weighted sum $S_n(X) = \sum_{j \in Ch(n)} w_{n,j} S_j(X)$, and for products $S_n(X) = \prod_{j \in Ch(n)} S_j(X)$.

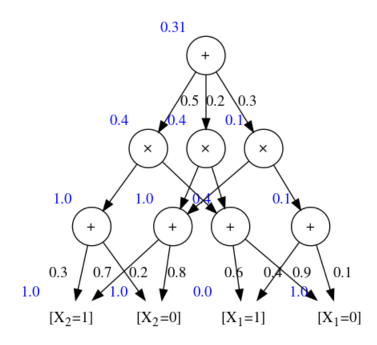


Figure 2.2: Computing the probability of evidence on a sample SPN.

Figure 2.2 shows the value of the SPN in Figure 2.1 given a valuation $X = \{X_1 = 0\}$. Values in blue are the values of each sub-SPN. Finding the probability of evidence P(X = x) is fast, as computing the value of an SPN is linear to the number of edges of the graph.

Additionally, we might want to find the probability that maximizes a certain valuation, i.e. the maximum a posteriori probability (MAP). To compute the approximate value of the MAP of some valuation X, we first transform the SPN into a max-product network (MPN) by replacing all sums with max nodes. The value of a max node is the maximum value of its weighted children. More formally, the value of an MPN's max node n is given by $M_n(X) = \max_{j \in Ch(n)} w_{n,j} M_j(X)$. Other nodes behave identically to an SPN. The computed value of an MPN is an approximation of $\max_y P(X = x, Y = y)$, where X is incomplete and Y is the set of variables that are missing. This is called the max-product algorithm.

In SPNs, computing the exact MAP was shown to be NP-hard (Peharz et al., 2015; Conaty et al., 2017; Mei et al., 2018), and better approximation algorithms were proposed as an alternative to the max-product algorithm described here. However, in this thesis,

when we talk about computing the (approximate) MAP, we are referring to the usual max-product algorithm.

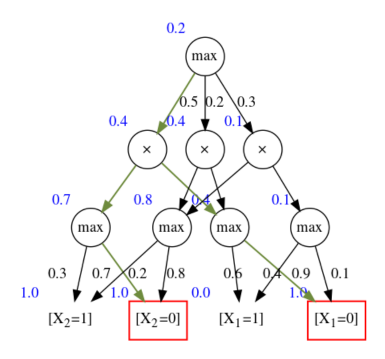


Figure 2.3: Computing the approximate MAP of an SPN through its MPN.

Once the MPN values are computed, we can find the most probable explanation (MPE) of the distribution given an evidence. This is done through a top-down forward pass, where we take a maximum sub-circuit path of the MPN by always taking the max path at a max node and taking all paths on a product node. The MPE are the maximum sub-circuit leaves' instantiations.

Figure 2.3 shows the MPN of the SPN shown in Figure 2.1 given $X = \{X_1 = 0\}$, where the numbers in blue represent the MPN values at each node, green arrows indicate the sub-circuit of maximum value and red boxes indicate the most probable valuations given evidence. The resulting MPE $\arg\max_{y\in\mathcal{Y}}P(X=\{X_1=0\},Y=y)$ is the valuation $\{X_1=0,X_2=0\}$.

Therefore, computing the probability of evidence, which is also called *soft* inference, of an SPN is done through a single bottom-up pass. Similarly, computing the MAP probability, refered to as *hard* inference, is done through a bottom-up pass on the SPN's MPN. On the other hand, finding the MPE valuations requires a bottom-up pass to first compute the MAP, and then a top-down search to find the most probable instantiations.

We provide next pseudocode for computing both soft and hard inference. We assume as input only valid, (weight) normalized SPNs. However, one could easily extend the included algorithms for unnormalized networks.

16:

Algorithm 1 SoftInference: Computes the probability of evidence in SPNs

Input A valid SPN *S* with normalized weights and a valuation *X*

```
Output The soft inference values at each node S_n
    Initialize S_n = 0
     Find topological order T of S
    for each node n \in S from T do
         if n is a leaf node then
               Let Sc(n) = \{X_k\}, p_n(x) be n's pdf and \hat{p}_n be p_n's mode
 5:
             if X_k \in X then
 6:
 7:
                   Let x_k be X_k's value in X
 8:
                   S_n \leftarrow p_n(x_k)
              else
 9:
                   S_n \leftarrow \hat{p}_n
10:
         else if n is sum node then
11:
12:
              for all j \in Ch(n) do
                   S_n \leftarrow S_n + w_{n,j}S_j
13:
         else
14:
             for all j \in Ch(n) do
15:
```

Algorithm 2 HardInference: Computes an approximation of the MAP in SPNs

Input A valid SPN *S* with normalized weights and a valuation *X*

Output The hard inference values at each node M_n

 $S_n \leftarrow S_n \cdot S_i$

17: **return** each S_n node value

```
Let M be S's MPN
     Initialize M_n = 0
     Find topological order T of M
 4: for each node n \in M from T do
        if n is a leaf node then
              Let Sc(n) = \{X_k\}, p_n(x) be n's pdf and \hat{p}_n be p_n's mode
 6:
             if X_k \in X then
 7:
                  Let x_k be X_k's value in X
 8:
 9:
                   M_n \leftarrow p_n(x_k)
             else
10:
                   M_n \leftarrow \hat{p}_n
11:
        else if n is sum node then
12:
             for all j \in Ch(n) do
13:
                   M_n \leftarrow \max(M_n, w_{n,j}M_j)
14:
        else
15:
             for all j \in Ch(n) do
16:
                   M_n \leftarrow M_n \cdot M_i
17:
18: return each M_n node value
```

Finally, we show how to algorithmically compute the MPE given some evidence.

Algorithm 3 ArgMaxSPN: Finds the MPE of a valuation on an SPN

Input A valid SPN S with normalized weights and a valuation X **Output** The arg max values of each variable according to X1: $M \leftarrow \mathsf{HardInference}(S, X)$

```
2: Let Y be a copy of X
3: Let Q be a queue
4: Push S into Q
5: for each node n \in M in Q do
        if n is a leaf node then
6:
             Let Sc(n) = \{X_k\} and p_n(x) be n's pdf
7:
             Let \hat{x} = \arg \max_{x_k} p_n(x_k) be p_n's maximum valuation
8:
            if X_k \notin X then
9:
                 Y \leftarrow Y \cup \{X_k = \hat{x}\}\
10:
        else if n is sum node then
11:
             Push maximum child M_j, j \in Ch(n) into Q
12:
        else
13:
             Push all children j \in Ch(n) into Q
14:
15: return Y
```

Chapter 3

Parameter learning

The objective of this chapter is to expose the ideas behind generative and discriminative gradient descent for parameter learning of sum-product networks. We first show how to derive the SPN with respect to its nodes and weights so that we can find the gradient of the SPN wrt its parameters (i.e. weights). This allows us to find the weight updates needed for gradient descent on SPNs. We then describe how to perform generative stochastic gradient descent, and finally discriminative gradient descent.

3.1 Derivatives

Let *S* be an SPN. We are only interested in finding the derivative of internal nodes, as leaf nodes have no weights to be updated. Our objective is to find the gradient $\partial S/\partial W$ by computing each component $\partial S/\partial w_{n,j}$, allowing us to find each weight update on the SPN.

At each weighted edge $(n \to j, w_{n,j})$, the derivative $\partial S/\partial w_{n,j}$ takes the form

$$\frac{\partial S}{\partial w_{n,j}}(X) = \frac{\partial S}{\partial S_n} \frac{\partial S_n}{\partial w_{n,j}}(X) = \frac{\partial S}{\partial S_n} \frac{\partial}{\partial w_{n,j}} \left(\sum_{i \in Ch(n)} w_{n,i} S_i(X) \right) = \frac{\partial S}{\partial S_n} S_j(X). \tag{3.1}$$

The term $\partial S/\partial S_n$ appears because of chain rule, since S_n is a function of S. This can be intuitively interpreted as taking into account the change in all nodes "above" n. So to compute the derivative wrt a weight, we need to find the derivative $\partial S/\partial S_j$ for each internal node j.

Finding $\partial S/\partial S_j$ requires analyzing two possible cases: sum and product parents of j. We now that S is a multilinear function of X, since in reality S is just a function made of sums and products. In particular, if we apply chain rule on $\partial S/\partial S_j$, we have that

$$\frac{\partial S}{\partial S_{j}}(X) = \underbrace{\sum_{\substack{n \in \operatorname{Pa}(j) \\ n: \text{ sum}}} \frac{\partial S}{\partial S_{n}} \frac{\partial S_{n}}{\partial S_{j}}(X)}_{(*)} + \underbrace{\sum_{\substack{n \in \operatorname{Pa}(j) \\ n: \text{ product}}} \frac{\partial S}{\partial S_{n}} \frac{\partial S_{n}}{\partial S_{j}}(X).$$

We expand each term at a time. Starting with the sum parents case, we can substitute the value of $S_n(X)$ with the corresponding expansion.

$$(*) = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ sum}}} \frac{\partial S}{\partial S_n} \frac{\partial}{\partial S_j} \left(\sum_{i \in \text{Ch}(n)} w_{n,i} S_i(X) \right) = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ sum}}} \frac{\partial S}{\partial S_n} w_{n,j}$$

We do the same for the product case.

$$(**) = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ product}}} \frac{\partial S}{\partial S_n} \frac{\partial}{\partial S_j} \left(\prod_{i \in \text{Ch}(n)} S_i(X) \right) = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ product}}} \frac{\partial S}{\partial S_n} \prod_{k \in \text{Ch}(n) \setminus \{j\}} S_k$$

Which brings us to the final form.

$$\frac{\partial S}{\partial S_{j}}(X) = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ sum}}} \frac{\partial S}{\partial S_{n}} w_{n,j} + \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ product}}} \frac{\partial S}{\partial S_{n}} \prod_{k \in \text{Ch}(n) \setminus \{j\}} S_{k}$$
(3.2)

Note how each $\partial S/\partial S_i$ depends on the derivative of its parents. This dependency goes all the way up to the root, where $\partial S/\partial S=1$. This derivation lends itself neatly to an algorithmic format.

Algorithm 4 Backprop: Backpropagation derivation on SPNs

Input A valid SPN S with pre-computed probabilities $S_n(X)$

Output Partial derivatives of *S* with respect to every node and weight

- 1: Înitialize $\frac{\partial S}{\partial S_n} = 0$ except $\frac{\partial S}{\partial S} = 1$ 2: **for** each node $n \in S$ in top-down order **do**
- **if** *n* is sum node **then** 3:
- 4:
- $\frac{\partial S}{\partial S_j} \leftarrow \frac{\partial S}{\partial S_j} + w_{n,j} \frac{\partial S}{\partial S_n}$ $\frac{\partial S}{\partial w_{n,i}} \leftarrow \frac{\partial S}{\partial S_n} S_j$ 5:
- 6:
- else 7:
- **for** all $j \in Ch(n)$ **do** 8:
- $\frac{\partial S}{\partial S_i} \leftarrow \frac{\partial S}{\partial S_i} + \frac{\partial S}{\partial S_n} \prod_{k \in Ch(n) \setminus \{j\}} S_k$ 9:

Computing all derivatives and forward passes is fast, as it takes linear time in the

number of edges. However, these values suffer from gradient diffusion, as their signal dwindles the deeper the network, eventually becoming zero.

A possible solution to this issue is replacing soft derivation with hard derivation. This is done by finding the derivatives of the MPN of the network instead of the SPN. This guarantees that the signal remains constant throughout the structure, at the cost of slower convergence rate. We call this hard inference derivation, as opposed to the regular soft inference derivation we covered earlier.

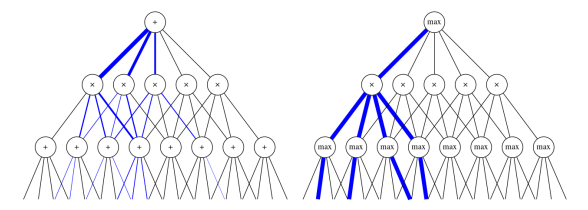


Figure 3.1: Signal difference between soft and hard derivation.

Figure 3.1 gives a visual representation of the difference between soft and hard derivation in gradient descent. MPNs preserve the signal, as the resulting gradient is constant.

To compute the hard derivatives of an SPN, we take its MPN and find its derivatives in a similar way as in soft derivation. Let M be an MPN. We shall call W the multiset of weights that a forward pass through M visits. The value of M is $M(X) = \prod_{w_i \in W} w_i^{c_i}$, where c_i is the number of times w_i appears in W. We can then take the logarithm of the MPN to end up with a friendlier expression.

$$\frac{\partial \log M}{\partial w_{n,j}} = \frac{\partial}{\partial w_{n,j}} \log \left(\prod_{w_i \in W} w_i^{c_i} \right) = \frac{1}{\prod_{w_i \in W} w_i^{c_i}} \cdot c_{n,j} w_{n,j}^{c_{n,j}-1} \cdot \prod_{w_i \in W \setminus \{w_{n,j}\}} w_i^{c_i}$$

If we assume that weights are strictly positive, the resulting expression results in the final hard derivative

$$\frac{\partial \log M}{\partial w_{n,j}} = c_{n,j} \frac{w_{n,j}^{c_{n,j}-1}}{w_{n,j}^{c_{n,j}}} = \frac{c_{n,j}}{w_{n,j}}.$$
(3.3)

Although not needed for the gradient, we can also compute the derivative in each internal node. The process is similar to soft derivation. There is no change for parent product nodes. For parent max nodes, we sum only contributions where $w_{n,j} \in W$.

$$\frac{\partial M}{\partial M_{j}} = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ max}}} \begin{cases} w_{k,n} \frac{\partial M}{\partial M_{k}} & \text{if } w_{k,n} \in W \\ 0 & \text{otherwise} \end{cases} + \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ product}}} \frac{\partial M}{\partial M_{n}} \prod_{k \in \text{Ch}(n) \setminus \{j\}} M_{k}$$
(3.4)

Computing each derivative $\partial \log M/\partial w_{n,j}$ means finding each $c_{n,j}$ count at each weight. This is done through an initial forward pass on M to find each MAP, and then finding each maximal edge in W through a backwards pass. Algorithm 5 computes the total number of occurences $c_{n,j}$ for each maximal edge $w_{n,j}$.

```
Algorithm 5 HardBackprop: Hard backpropagation derivation on SPNs
```

```
Input A valid SPN S with pre-computed MAP probabilities M_n(X) Output Counts c_{n,j} of each derivative \frac{\partial \log M}{\partial w_{n,j}}
```

```
1: Initialize c_{n,j} = 0
2: Let Q be a queue
3: Push M into Q
4: for each node n \in M in queue Q do
        if n is max node then
5:
            Let j = \arg \max_{i \in Ch(n)} w_{n,i} M_i(X) the maximum weighted child
6:
7:
            c_{n,j} \leftarrow c_{n,j} + 1
            Push M_j into Q
8:
9:
        else if n is product node then
            Push all children j \in Ch(n) into Q
10:
11: return all counts c_{n,j}
```

In summary, the derivatives of an SPN with respect to its internal nodes take values according to Table 3.1. The gradient components are shown in Table 3.2.

Inference	Partial derivatives wrt internal node j
Soft	$\frac{\partial S}{\partial S_j} = \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ sum}}} w_{n,j} \frac{\partial S}{\partial S_n} + \sum_{\substack{n \in \text{Pa}(j) \\ n: \text{ product}}} \frac{\partial S}{\partial S_n} \prod_{k \in \text{Ch}(n) \setminus \{j\}} S_k$
Hard	$\frac{\partial M}{\partial M_{j}} = \sum_{\substack{n \in \operatorname{Pa}(j) \\ n: \max}} \begin{cases} w_{k,n} \frac{\partial M}{\partial M_{k}} & \text{if } w_{k,n} \in W, \\ 0 & \text{otherwise.} \end{cases} + \sum_{\substack{n \in \operatorname{Pa}(j) \\ n: \text{ product}}} \frac{\partial M}{\partial M_{n}} \prod_{k \in \operatorname{Ch}(n) \setminus \{j\}} M_{k}$

Table 3.1: Partial derivatives for the SPN wrt internal nodes.

Inference	Partial derivatives wrt weight $w_{n,j}$
Soft	$\frac{\partial S}{\partial w_{n,j}} = S_j \frac{\partial S}{\partial S_n}$
Hard	$\frac{\partial M}{\partial w_{n,j}} = M_j \frac{\partial M}{\partial M_n}$

Table 3.2: *Partial derivatives for the SPN wrt weights.*

3.2 Generative gradient descent

Once computed all derivatives, we update each node with the resulting gradient component. For generative gradient descent, where we are learning a joint probability distribution P(X, Y), our objective is to find the gradient of the log-likelihood

$$\frac{\partial}{\partial W}\log P(X,Y) = \frac{\partial}{\partial W}\log S(X,Y) = \frac{1}{S(X,Y)}\frac{\partial S}{\partial W}(X,Y) \propto \frac{\partial S}{\partial W}(X,Y).$$

Since the gradient is proportional to the derivative of the weights, our weight update becomes

$$\Delta w_{n,j} = \eta \frac{\partial S}{\partial w_{n,j}}(X, Y),$$

where η is the learning rate. An L2 regularization factor can be added to the expression above, leaving us with the final generative gradient descent weight update

$$\Delta w_{n,j} = \eta \frac{\partial S}{\partial w_{n,j}}(X, Y) - 2\lambda w_{n,j}, \tag{3.5}$$

where λ is the regularization constant. We call this soft generative gradient descent. It is now easy to visualize why gradient diffusion occurs with soft derivation. Component $\partial S/\partial w_{n,j}$ depends on partial derivative $\partial S/\partial S_n$. Assuming normalized weights, the root node derivative $\partial S/\partial S=1$ and each subsequent descendant node becomes smaller and smaller.

Weight update for hard derivation comes directly from Equation 3.3. Since we are interested in the log-likelihood of the joint distribution

$$\frac{\partial}{\partial W}\log P(X,Y) = \frac{\partial}{\partial W}\log M(X,Y),$$

we get, for each component $w_{n,j}$, the weight update

$$\Delta w_{n,j} = \eta \frac{c_{n,j}}{w_{n,j}}.$$

In a similar fashion to soft generative gradient descent, we can apply L2 regularization to each weight update.

$$\Delta w_{n,j} = \eta \frac{c_{n,j}}{w_{n,j}} - 2\lambda w_{n,j} \tag{3.6}$$

So for generative gradient descent we get the following weight updates.

Inference	Weight updates
Soft	$\Delta w_{n,j} = \eta \frac{\partial S}{\partial w_{n,j}}(X, Y) - 2\lambda w_{n,j}$
Hard	$\Delta w_{n,j} = \eta \frac{c_{n,j}}{w_{n,j}} - 2\lambda w_{n,j}$

Table 3.3: *Generative gradient descent weight updates with L2 regularization.*

Algorithm 6 and Algorithm 7 show pseudocode for both soft and hard generative stochastic gradient descent, though it is easy to extend both to mini-batch versions. From now on we denote soft generative gradient descent and hard generative gradient descent as SGGD and HGGD for short.

Algorithm 6 SoftGenGD: Soft generative stochastic gradient descent for SPNs

Input A valid SPN S, learning rate η , regularization constant λ and a dataset D **Output** S with learned weights

```
1: repeat
2: for each instance I \in D do
3: Compute SoftInference(S, I)
4: Compute Backprop(S)
5: for each sum node n \in S do
6: w_{n,j} \leftarrow \eta \frac{\partial S}{\partial w_{n,j}} - 2\lambda w_{n,j}
7: Normalize weights
8: until convergence
```

Algorithm 7 HardGenGD: Hard generative stochastic gradient descent for SPNs

Input A valid SPN S, learning rate η , regularization constant λ and a dataset D **Output** S with learned weights

```
1: repeat
2: for each instance I \in D do
3: Compute HardInference(S, I)
4: Compute HardBackprop(S)
5: for each sum node n \in S do
6: w_{n,j} \leftarrow \eta \frac{c_{n,j}}{w_{n,j}} - 2\lambda w_{n,j}
7: Normalize weights
```

8: until convergence

3.3 Discriminative gradient descent

The goal of discriminative learning is optimizing the conditional probability distribution P(Y|X), where Y and X are query and evidence variables. To compute the gradient of this distribution we maximize the conditional log-likelihood (Gens and Domingos, 2012).

$$\frac{\partial}{\partial W} \log P(Y|X) = \frac{\partial}{\partial W} \log \left(\frac{P(Y,X)}{P(X)} \right) = \frac{\partial}{\partial W} \log P(Y,X) - \frac{\partial}{\partial W} \log P(X)$$

Through chain rule, we get the form

$$\begin{split} \frac{\partial}{\partial W} \log P(Y,X) - \frac{\partial}{\partial W} \log P(X) &= \frac{1}{P(Y,X)} \frac{\partial}{\partial W} P(Y,X) - \frac{1}{P(X)} \frac{\partial}{\partial W} P(X) \\ &= \frac{1}{S(Y,X)} \frac{\partial}{\partial W} S(Y,X) - \frac{1}{S(X)} \frac{\partial}{\partial W} S(X). \end{split}$$

We can update our weights discriminatively by taking each gradient component

$$\Delta w_{n,j} = \eta \left(\frac{1}{S(Y,X)} \frac{\partial S(Y,X)}{\partial w_{n,j}} - \frac{1}{S(X)} \frac{\partial S(X)}{\partial w_{n,j}} \right).$$

With L2 regularization, soft discriminative gradient descent has the following form.

$$\Delta w_{n,j} = \eta \left(\frac{1}{S(Y,X)} \frac{\partial S(Y,X)}{\partial w_{n,j}} - \frac{1}{S(X)} \frac{\partial S(X)}{\partial w_{n,j}} \right) - 2\lambda w_{n,j}$$
(3.7)

For hard inference we want to optimize the following expression.

$$\frac{\partial}{\partial W} \log \tilde{P}(Y|X) = \frac{\partial}{\partial W} \log \left(\frac{\tilde{P}(Y,X)}{\tilde{P}(X)} \right) = \frac{\partial}{\partial W} \log \left(\frac{M(Y,X)}{M(X)} \right)$$

Where \tilde{P} is the MAP probability of the distribution. As usual, we apply chain rule, yielding

$$\frac{\partial}{\partial W}\log\left(\frac{M(Y,X)}{M(X)}\right) = \frac{\partial}{\partial W}\log M(Y,X) - \frac{\partial}{\partial W}\log M(X).$$

But we know from Equation 3.3 that the derivatives of the logs have a particular expression based on the counts of visited weights. We substitute the equation above with the earlier results from hard derivation, giving us the following equation for each gradient component.

$$\frac{\partial}{\partial w_{n,j}} \log \left(\frac{M(Y,X)}{M(X)} \right) = \frac{\partial}{\partial w_{n,j}} \log M(Y,X) - \frac{\partial}{\partial w_{n,j}} \log M(X) = \frac{\Delta c_{n,j}}{w_{n,j}}$$

Where $\Delta c_{n,j}$ is the difference between the first counting, restricted to (Y, X), and the second restricted to only X.

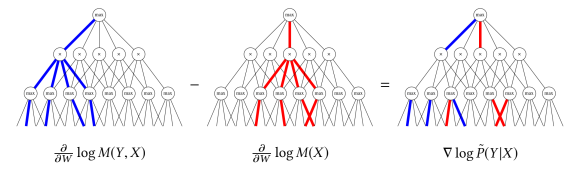


Figure 3.2: Hard discriminative gradient descent counts visualization.

Figure 3.2 shows the hard discriminative gradient descent difference derived from $\frac{\partial}{\partial W} \log \tilde{P}(Y|X)$. The first pass, shown in the image with blue edges, counts the maximum edges given the Y, X valuation. The second pass, in red, is the evidence pass on X. The gradient is then computed by finding the difference between the two countings. On the right-hand side of the expression portrayed in Figure 3.2, blue edges mean a positive count $c_{n,j}$ and red edges represent a negative count. Edges coming out from product nodes are not colored, as they are not weighted.

The actual weight update has a similar form to hard gradient descent.

$$\Delta w_{n,j} = \eta \frac{\Delta c_{n,j}}{w_{n,j}}$$

With L2 regularization we get

$$\Delta w_{n,j} = \eta \frac{\Delta c_{n,j}}{w_{n,j}} - 2\lambda w_{n,j}. \tag{3.8}$$

In a similar fashion to generative gradient descent, we denote by HDGD and SDGD hard discriminative gradient descent and soft discriminative gradient descent respectively.

We now build a discriminative gradient descent table for each inference type. Just like in generative gradient descent, we add an L2 term to it.

Inference	Weight updates
Soft	$\Delta w_{n,j} = \eta \left(\frac{1}{S(Y,X)} \frac{\partial S(Y,X)}{\partial w_{n,j}} - \frac{1}{S(X)} \frac{\partial S(X)}{\partial w_{n,j}} \right) - 2\lambda w_{n,j}$
Hard	$\Delta w_{n,j} = \eta rac{\Delta c_{n,j}}{w_{n,j}} - 2\lambda w_{n,j}$

Table 3.4: *Discriminative gradient descent weight updates with L2 regularization.*

We now finally show an algorithmic form to HDGD and SDGD. Note how in discriminative gradient descent we have two passes through the network. We can avoid recomputing node values by memoizing nodes that have no query variables in descendant's scopes (Gens and Domingos, 2012).

Algorithm 8 SoftDiscGD: Soft discriminative stochastic gradient descent for SPNs

Input A valid SPN S, query variables Y, learning rate η , regularization constant λ and a dataset D

Output *S* with learned weights

```
1: repeat
```

- **for** each instance $I \in D$ **do**
- Compute SoftInference(S, I) and store them in S_n^+ for each node n3:
- Compute Backprop(S^+) and store them in $\frac{\partial S_n^+}{\partial w_{n,j}}$ 4:
- Compute SoftInference($S, I \setminus Y$) and store them in S_n^- for each node n
- Compute Backprop(S^-) and store them in $\frac{\partial S_n^-}{\partial w_n}$
- 7:

7: **for** each sum node
$$n \in S^- \cup S^+$$
 do
8: $w_{n,j} \leftarrow \eta \left(\frac{1}{S^+} \frac{\partial S^+}{\partial w_{n,j}} - \frac{1}{S^-} \frac{\partial S^-}{\partial w_{n,j}} \right) - 2\lambda w_{n,j}$

- Normalize weights
- 10: **until** convergence

Algorithm 9 HardDiscGD: Hard discriminative stochastic gradient descent for SPNs

Input A valid SPN S, query variables Y, learning rate η , regularization constant λ and a dataset D

Output *S* with learned weights

```
1: repeat
2: for each instance I \in D do
3: Compute HardInference(S, I) and store them in M_n^+ for each node n
4: Compute HardBackprop(M^+) and store them in c_{n,j}^+
5: Compute HardInference(S, I \setminus Y) and store them in M_n^- for each node n
6: Compute HardBackprop(M^-) and store them in c_{n,j}^-
7: for each sum node n \in S do
8: w_{n,j} \leftarrow \eta\left(\frac{c_{n,j}^+ - c_{n,j}^-}{w_{n,j}}\right) - 2\lambda w_{n,j}
9: Normalize weights
```

10: until convergence

Chapter 4

Structure learning

In this chapter we cover two structure learning algorithms we use for image classification. The first is based on Dennis and Ventura, 2012. The second is a variation of Gens and Domingos, 2013's structure learning schema. Once we cover both algorithms, we explain how we add a slight modification to the first architecture. We have empirically found that this change increased image classification accuracy significantly. We call this the "classification architecture".

4.1 The Dennis-Ventura architecture

Let us first formalize the notion of dataset. We call a dataset a set of *instances*, where each instance is a set we call *valuation* or *instantiation*. As we have mentioned before, a valuation may be incomplete, meaning that an instantiation of some random variable may be missing from the instance. In this thesis we assume complete data, as both structure learning algorithms do not admit incomplete datasets.

Having said that, let D be a complete dataset. Since D is complete, for each instance I we can map each random variable X from I to a number, yielding an ordered vector (X_1, X_2, \ldots, X_m) equivalent to I. We do the same for each instance I. The vector (I_1, I_2, \ldots, I_n) is a representation of D. This way, D could be seen as a $m \times n$ matrix. We denote by D^T the transpose of the matrix representation of the dataset D. Let T be a subscope, that is, a subset of the set of all variables in the SPN. We use the notation D_T to represent the matrix of all instances from D but restricted only to elements from random variables in T.

Since we are restricted to the image classification domain, we give some semantic meaning to datasets. If D is a dataset, then each instance $I \in D$ can be seen as a vector containing all the pixel values of an image plus a classification label. Each variable is a pixel from the image, and each variable value is the pixel's color intensity. If the dataset D is a vector of images and their labels, the transpose D^{T} is a vector of variables and their values in each image.

Just like in Poon and Domingos, 2011, the Dennis-Ventura algorithm uses the notion

of similarity between local variables. This local neighborhood is called a Region. A Region represents a cluster of pixels that has some semantic value when grouped together. Contrastingly, a Decomposition represents independence between variables. In an SPN, a Region is graphically represented by a set of sum nodes, whilst a Decomposition is a set of products.

To learn an SPN structure from data, Dennis and Ventura, 2012 uses a *region graph*, which is a simplified representation of an SPN made out of Region nodes and Decomposition nodes.

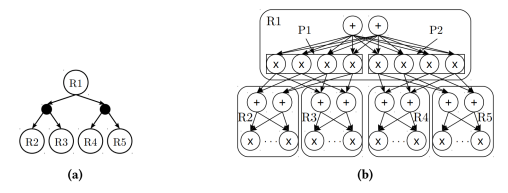


Figure 4.1: Dennis-Ventura region graph and translated SPN as shown in Dennis and Ventura, 2012.

The region graph is generated by recursively finding two subregions from a parent region through the use of k-means clustering. Let R be a region, and D_R^{T} the transposed dataset restricted to R's scope. We partition R into two subregions R_1 and R_2 by k-means clustering D_R^{T} , yielding two subclusters $D_{R_1}^{\mathsf{T}}$ and $D_{R_2}^{\mathsf{T}}$. We then apply recursion on the two subregions. At each clustering step, we connect regions R to a decomposition node P, which is then connected to each R_i node. Our stop criteria is when $Sc(R_i) = 1$. The root node is a special case. We run k-means cluster on D, and for each D_i cluster, we construct a sub-SPN for each root child with D_i .

Once created, the region graph is then translated to a valid SPN. Each region node R is translated to a set of SPN nodes. If Sc(R) = 1, then these nodes are g univariate gaussian distributions, where each gaussian is a different quantile of the distribution of the pixel. Else, m sum nodes are created. Partition nodes are translated to product nodes. Edges are added such that every product child node of a region is connected to all sum nodes in the region. Each of these product nodes are then connected to a distinct pair of sum nodes from both region's children subregions, meaning that each decomposition node contains 2^m products.

With the architecture done, we apply parameter learning on the SPN to learn weights. This is done either through gradient descent or EM. In this thesis we apply generative and discriminative gradient descent to the architecture.

4.2 The Gens-Domingos schema

In Gens and Domingos, 2013, Gens and Domingos describe a flexible schema for structure learning of SPNs. The schema is based on the idea that the scope in a sum node's child mean that these variables are similar (and by consequence dissimilar to the variables in other sibling nodes' scopes), and that variables in a product's child mean that variables are dependent of each other (and analogally to sums, independent of other siblings).

This interpretation of SPNs yields an adaptable and open schema of learning. Sum nodes are created through clustering, as each cluster has some similarity aspect given some metric. Meanwhile, product nodes are created through statistical variable independence algorithms. When the scope of this partitioning of variables is one, we create a univariate distribution over the partitioned dataset.

Algorithm 10 GensArch: Gens-Domingos structure learning schema

```
Input Set of instances D and scope X
Output SPN structure learned from D and X
 1: if |X| = 1 then
 2:
         return univariate distribution over D_X
 3: else
         Partition X into P_1, P_2, \ldots, P_m such that every P_i is independent of P_j, i \neq j
 4:
         if m > 1 then
 5:
              Let \pi be a new product node
 6:
              for i \leftarrow 1, \ldots, m do
 7:
                  p_i \leftarrow GensArch(D_{P_i}, P_i)
 8:
                  \pi.AddChild(p_i)
 9:
              return \pi
10:
         else
11:
              Cluster D such that Q_1, Q_2, \ldots, Q_n are D's clusters
12:
              Let \sigma be a new sum node
13:
              for i \leftarrow 1, \ldots, n do
14:
15:
                  s_i \leftarrow GensArch(Q_i, X)
                  w \leftarrow |Q_i|/|D|
16:
                                                                           ▶ w is edge \sigma \rightarrow s_i's weight
                  \sigma.AddChild(s_i, w)
17:
18:
              return \sigma
```

Our implementation was done by using DBSCAN, a density based clustering algorithm that automatically decides the number of clusters to generate (ESTER et al., 1996), for clustering and the traditional G-test for variable independence. We also implemented k-means, k-mode and k-median for clustering and Pearson's χ^2 -square for independence testing. We found that DBSCAN yielded the best accuracy results on clustering, but took a long time for training. Using k-means clustering with k=2 yielded worse, but comparable results, but with the upside of being fast to train. The G-test provided the best variable independence results.

Finding independent variable subsets can be done by iteratively comparing variables

pairwise. The connected components of the resulting disconnected graph are independent subsets. This brute force approach is intractable, as the G-test already takes O(|X||Y|) time, where X and Y are random variables and $|\cdot|$ indicates the number of categories of the RV, and testing each pair of variables exhaustively is exponential.

Instead of testing every variable pairwise, we constructed a dependency graph. Each vertex from the dependency graph represents a variable. An edge between two vertices means the two variables are dependent. To find independent partitions in a dataset it suffices to find a spanning tree of the dependency graph. We do this through Kruskal's MST union-find algorithm. The resulting connected components of the spanning tree are the partitions we wish to find. This significantly reduces complexity. However, we have found that it still accounts for approximately 90% of the algorithm's runtime.

We speculate that a better approach to variable indepedence would be finding approximate spanning trees on the graph. Many independence tests resulted in a completely dependent graph, but with cuts that could possibly yield better accuracy and runtime performance.

Our Gens-Domingos implementation generated very deep and expressive SPNs, resulting in good accuracy results. However, the algorithm only generates SPTs, as once each step (either clustering or variable independence) is concluded, the function never returns to the same node.

4.3 The classification architecture

The Dennis-Ventura structure learning algorithm is able to model classification problems by partitioning data into l clusters, where l is the total number of labels, and assigning a sub-SPN for each cluster. One can interpret each sub-SPN as a model of each label. However, clustering may not select the right classification instances for each label, as we have no control over which labels fit each cluster. This effect is intensified on datasets containing a large number of data.

We try to solve this problem by simplifying the model. Instead of generating sub-SPNs through clustering, we restrict each label to its own SPN. In our architecture, we create a single sum node as root, representing the image and its classification label. For each label l, we construct a sub-SPN S_l such that the SPN is still valid. This is done by assigning a product node as S_l 's root. Let Y be the classification variable. Each of these products are then connected to an indicator variable $[Y = y_l]$ and a sub-SPN restricted to only data where $Y = y_l$.

Figure 4.2 shows the graphical representation of the classification architecture. The SPN is still valid, as the product node guarantees decomposability and the root sum node is complete. Each sub-SPN $S|Y=y_i$ is then constructed with the Dennis-Ventura algorithm, but restricted to data with the $Y=y_i$ valuation.

In practice, this architecture yielded much better results than the original clustering method. Furthermore, it is possible to easily parallelize each $S|Y = y_i$ learning procedures for faster learning runtime. Similarly, since each $S|Y = y_i$ is independent of other restricted

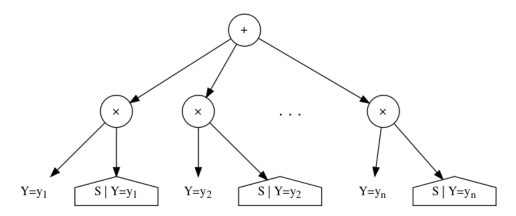


Figure 4.2: The classification architecture for the Dennis-Ventura structure.

SPNs, we can compute SPN values concurrently, allowing for faster inference. We cover this more thoroughly in chapter 5.

Interestingly, when this architecture model was applied to the Gens-Domingos algorithm, accuracy decreased. This is possibly due to the limitations of trees in SPNs. Another possible reason is that the Gens-Domingos better captures interactions between labels and pixels than only between pixels.

Modelling the problem

In this chapter we formally define a possible model for the mobile robots self-driving problem. We first present this problem as an imitation learning through image classification problem. We then describe the dataset intended as training set for image classification. Finally, we discuss the problems of inference speed we should encounter when dealing with a mobile robot, and provide a concurrent programming solution for our particular problem.

5.1 The problem

One of the main problems of self-driving is to follow a certain pathway. In real life, a self-driving car should be able to maintain itself centered on a lane, more specifically inbetween lane markings. We address this particular subproblem of self-driving. This is achieved by considering this situation as an imitation learning application.

Imitation learning consists of an agent accurately mimicking human behavior. In our case, we wish for such an agent to simulate the behavior of keeping a car centered on a single lane. We model this particular situation by use of image classification. The agent, in this case the self-driving car, should reliably identify when to turn and when to go straight my solely "looking" at the road. This can be achieved through the use of image classification, as a turn tends to have different visual features then a straight lane.

Whilst this comes naturally to humans, machines have trouble identifying these features by themselves. Noise and object occlusion play a big role in how reliably the agent behaves. A possible obstruction of the agent's view could turn fatal in a real-life scenario. However, with the advent of more complex models in machine learning, modelling this problem through image classification has become a feasible solution.

Our approach to self-driving in mobile robots consists of a very simplified and purely reactive image classification problem. The mobile robot should follow a lane and turn accordingly based on its image input of its front view.

We define a classification variable, which we will denote by *Y*, as an indicator of what

the robot should do. The function Val(Y) defines the set of all possible values of Y. In our case, $Val(Y) = \{L, R, U\}$, each meaning that the robot should "go left", "go right" and "go straight" respectively.

Let $X = \{X_1, X_2, \dots, X_n\}$ be the set of variables that compose an image, where each X_i represents a pixel of a flattened image. Our entire scope is defined by the set $W = X \cup Y$. Our objective is to reliably guess Y's value based solely on the values of X. That is, we wish to find

$$\underset{y \in \text{Val}(Y)}{\text{arg max}} P(Y = y | X) = \underset{y \in \text{Val}(Y)}{\text{arg max}} \frac{P(Y = y, X)}{P(X)} \propto \underset{y \in \text{Val}(Y)}{\text{arg max}} P(Y = y, X). \tag{5.1}$$

Where we assume the existence of an underlying probability distribution that correctly models the classification problem. Ultimately, our goal is to find this distribution by "learning" from data through the learning algorithms described in chapter 3 and chapter 4.

Once learned, the SPN is able to find the MAP probabilities and states that correctly predict the most probable values of Y given an image. Note how the LHS of Equation 5.1 can be computed with a single pass with the approximate max-product algorithm. Alternatively, we can also compute the exact max values by computing each possible $y \in Val(Y)$ with a single forward pass through the SPN.

5.2 The dataset

For training, we used Moraes and Salvatore's self-driving dataset (Moraes and Salvatore, 2018). Every image has dimensions 80×45 , with three additional channels for RGB. The dataset is split into three sets, corresponding to training, test and validation data. Each image contains a label indicating whether the robot should go straight, turn left or turn right. These actions are labeled as 0, 1 and 2.



Figure 5.1: Sample images from training dataset.

Figure 5.1 showcases sample images from the training dataset. The leftmost image has label L, the one on the middle is U and the one on the right R. It is possible to observe that images do not have uniform lightning and lane markings are irregular. This adds a noise effect to the images.

The original dataset is already reduced in size. However, the presence of color is not so important to identify the correct values of *Y*. If we compare Figure 5.1 and Figure 5.2, lane markings are still very much visible.



Figure 5.2: Grayscale sample images from training dataset.

We can try to further reduce the complexity of the dataset at the same time preserving its most informational features by attempting to reduce the number of possible values of each pixel through image quantization. This transformation turned out to be very meaningful in terms of both training speed and accuracy, as we detail in a later chapter. However, noise is still very much present in the images, as Figure 5.3 shows.



Figure 5.3: *Quantized sample images from training dataset.*

Another possible transformation we can apply on the dataset is binarization. However, traditional "hard" binarization with a fixed k threshold on the image could potentially cause completely black or white images due to a poor choice of k. This can countered with two possible solutions. Either through adaptive threshold where we choose k either by a mean measurement or through a gaussian, or by use of Otsu's binarization (OTSU, 1979). We found that Otsu's method, coupled with a prior gaussian blur transformation on each image, proved the most capable of correctly applying binarization in our dataset.



Figure 5.4: Binarized sample images from training dataset.

Figure 5.4 shows the final result of applying a combination of gaussian blur and Otsu's binarization.

One last transformation we tested our models on was histogram equalization. Equalization was done in order to reshape the pixel value histogram to an approximately uniform distribution. This was done in an attempt to increase the accuracy of the Gens-Domingos algorithm. As mentioned in chapter 4, the Gens-Domingos schema attempts to find partitions of independent variables. We use an implementation of the G-test, which works best when there are sufficient samples for every variable category. The original

dataset has a skewed histogram that contains almost no pixel values in the extreme ranges (either too white or too black). This transformation added a lot of noise to the dataset. In spite of this, we empirically found that with the use of equalization accuracy was increased by more or less 5%.

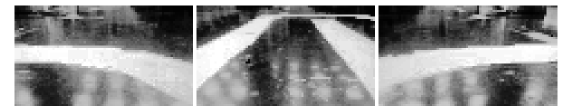


Figure 5.5: Equalized sample images from training dataset.

5.3 The model

Our classification model should be able to accurately infer the most probable action to be taken given the front camera feed's image. Although a simple decision model could be used for such a task, in many cases we wish to maintain an uncertainty measurement (e.g. a probability distribution) as a means to quantify error. Error and noise is often a problem in robotics, one which can be mitigated through probabilistic localization algorithms. For our particular case, we discard these problematic issues and focus solely on the problem of computing a valid probability distribution that accurately models our classification problem. SPNs have full probabilistic semantics and are able to represent local and sparse variable interactions due to their deep architecture. This allows for a reliable probabilistic model for modelling our classification problem.

We use two SPN architectures for our problem. We model the first using the Dennis-Ventura algorithm described in section 4.1. This model in particular uses the classification architecture also cited in chapter 4. The second model uses the Gens-Domingos structural algorithm (section 4.2).

As we have previously mentioned, we wish to compute the MPE

$$\underset{y \in \text{Val}(Y)}{\text{arg max}} P(Y = y, X) = \underset{y \in \text{Val}(Y)}{\text{arg max}} S(Y = y, X) \approx y|_{M(Y = y, X)}, y \in \text{Val}(Y). \tag{5.2}$$

Where $y|_{M(Y=y,X)}$ signals an MPN forward and backward pass to compute the most probable explanation of variable *Y* given *X* as evidence on the underlying SPN.

Recalling chapter 2, we can compute this probability in two different ways. Either by computing each Y value by means of a forward pass on the SPN S(Y = y, X), or through the approximate MAP M(Y = y, X). These two options both carry disadvantages. On one hand, computing each Y value through a forward pass on each possible valuation can cause inference to become very slow, especially when hardware is as limited as in a mobile robot. On the other hand, the max-product algorithm is a fast inference method, though

approximate. Furthermore, the approximate method tends to favor shallower paths due to its Viterbi-style features.

We compromise with exact inference, but with an addendum. We take advantage of our hardware's multi-cored CPU by running exact inference in parallel. Since the Gens-Domingos and Dennis-Ventura architectures are very distinct structure-wise, we apply a different implementation for each.

As mentioned in chapter 4, the Dennis-Ventura structure we implemented follows a particular classification architecture that models each set of images of a certain label as a separate, independent sub-SPN. This allows for an easy parallel programming implementation, as each CPU core can be assigned to a single sub-SPN, and thus to a label. Each core will then be linked to a particular robot command. Once all cores finish inference, we then compare which sub-SPN returned the highest probability. This is only advantageous if the number of labels is low. In our case, |Val(Y)| = 3, meaning this method is feasible for our problem.

This method does not work as well on the Gens-Domingos structure, as we cannot guarantee if an independency or clustering step on the root node has yielded a sufficient number of nodes for each core. However, a similar method is used, where a concurrency queue of n most processes stores each child of the root node, allowing for them to be run in parallel. Since the Gens-Domingos structural algorithm produces SPTs, every child of a node is guaranteed to be independent of its siblings.

Hardware

In this chapter we dive into the hardware specifications of our mobile robot. Our robot is composed of two main units: the Berry and the Brick. The former is composed of a Raspberry Pi and a webcam, and serve as the brain for our robot. The latter is a Lego Mindstorms robot, which contains two differential motors for each wheel and a small processing unit, called the Brick, used for issuing low-level commands to the wheels. We first explore the Berry, and then describe the Brick.

6.1 The Berry

The main processing unit of our robot consists of a Raspberry Pi 3 Model B. With a Broadcom Quad-Core BCM2837 with 64-bit at 1.4 GHz, it is a small, yet powerful microcomputer. Its architecture is based on ARMv7, and has four processing cores.

Through four USB 2.0 ports, we are able to connect the Raspberry Pi with the motor part of our robot and a small portable webcam that will be used for input.

A MicroSD with 16GB provides both the Raspbian operating system, which is based on Debian, as well as an additional 1 GB swap memory space, as the Berry contains only 1GB SDRAM. The remaining amount is used as storage.

Although the Berry has reasonable processing power for its size, training is done offline in a desktop computer, with only inference done on the micro-computer. With its connected webcam, the Berry receives each image frame, applies some pre-processing to the image (which will be detailed in chapter 7), feeds it to the SPN as evidence, computes the most probable classification label, and finally sends this to the external unit, i.e. the Brick, responsible for dealing with the robot's motion. This is all done concurrently, as we can dedicate three cores to classification, and the remaining unit to camera capture, image pre-processing and message passing.

6.2 The Brick

The Lego Mindstorms robot is composed of three main parts: the brick, which is the Lego Mindstorm's processing unit that handles the motors, and two differential motors that are able to give motion to the robot in a somewhat precise manner through the use of tachometers. In this document, when we say the Brick, we are referring to the whole set of brick and motors.

In our experiments we used the Lego Mindstorms NXT. Its main processor is an Atmel AT91SAM7S256 with a 48 MHz clock and 32-bit ARMv4 architecture. It has 64 KB of RAM and 256 KB of flash storage. A USB port allows for local input and output from and to the Brick.

Low-level handling is done on the Brick. Once it receives a command to be executed (i.e. the classification label passed by the Berry), the Brick needs to interpret it and execute the desired movement. We use the leJOS NXJ API¹ for low-level motor programming. The Brick's cycle is as follows:

Algorithm 11 Brick: The Brick's cycle

```
1: Connect and power up motors
2: Let UP \leftarrow 0x00
3: Let LEFT \leftarrow 0x01
4: Let RIGHT \leftarrow 0x02
5: Let OUIT \leftarrow 0x03
6: Let NOOP ← 0 \times 04
7: Let M_L and M_R be the left and right side motors respectively
8: Let k be some speed constant
   while true do
        if input size > 0 then
10:
            Read input byte and store in variable c
11:
            if c is QUIT then
12:
13:
                Disconnect and power down
            else if c is UP then
14:
                Set M_L's power to k
15:
                Set M_R's power to k
16:
            else if c is LEFT then
17:
                Set M_L's power to 2k
18:
                Set M_R's power to 3k
19:
            else if c is RIGHT then
20:
                Set M_L's power to 3k
21:
                Set M_R's power to 2k
22:
```

No complex control is done on the Brick. A standard "do something until told otherwise" is implemented. This is reliant on the Berry's inference speed, but we found that inference was fast enough for this to work well on slow speeds.

¹Available at http://lejos.org/nxj.php

Setting the motors' power is straightforward with leJOS. Two function calls are enough for our case. The problem is in choosing k. If k is too big, not only the Berry might not have enough time to compute the labels, but also turns may not be as steep as the lane requires. In the case of a too small k, the challenge of autonomous driving becomes null. We experimented on values of k and chose k = 150 for our Lego Mindstorms. We recognize that a better control solution to this would be preferred, but for our case this suffices.

6.3 Bridging the two

Communication between the two modules, Berry and Brick, is done through a USB cable. We use a Go codebase on the Berry, opting to use Google's GoUSB² as a low-level interface to handle the Raspberry Pi's data output, sending each predicted label as a byte value.

At each cycle, the Brick checks for new input by reading from an input stream. This is done through Java's java.io.DataInputStream, which translates the incoming USB data into Java bytes.

This hierarchization of Berry to Brick allows for the unit with most processing power, i.e. the Berry, to take on the heavy load, leaving only the necessary dedicated low-level motor handling to the Brick, which is very limited in terms of CPU and memory.

Every USB device contains a pair of IDs that are essential for identification. The vendor ID is used to identify which company or organization created the device. Whilst the product ID is used to identify specific products created by the company, in our case the Lego Mindstorms NXT version 2. The vendor and product IDs of the Brick are 0x0694 and 0x0002. These IDs allow us to cycle through each active USB device and select exactly the one we need.

USB devices may contain multiple functionalities, such as acting as a power source or as an input/output stream. These are called configurations, and are usually indexed by a number. In this thesis we are interested in the read and write configuration of the Brick, indexed by the number 1.

Each configuration contains different interfaces that can be seen as virtual devices to the physical USB. We selected interface number 0, with alternative interface 0.

When writing and reading from and to a USB device, we must define an output and input endpoint. There are a total of 30 endpoints, where input endpoints are indexed from 0×81 to $0\times8f$, and output endpoints from 0×01 to $0\times0f$. Two additional in/out endpoints 0×80 and 0×00 are control endpoints used internally by the USB device. For our Brick we used the input 0×82 and output 0×01 endpoints.

²Available at https://github.com/google/gousb

Experiments and benchmarks

In this chapter we apply our model on an artificial self-driving dataset, running several experiments and measuring how good our models perform with each setup. We first describe the different configurations we use for training and testing. We then show results on accuracy for each of the models and pre-processing transformations. Finally we show how fast each model is.

7.1 Pre-processing

Before training and inference, we apply different image transformations to the dataset. As mentioned in chapter 5, we use three main transformations: binarization, quantization and equalization. In all cases we first convert the original RGB colored dataset to grayscale.

7.1.1 Binarization

The binarization process was done by first converting the original dataset to grayscale, followed by applying a gaussian blur filter with a (1, 1) kernel and (1, 1) standard deviation, and finally using Otsu's binarization. We chose this particular process since standard hard threshold binarization was unable to produce clear images of the track lines.



Figure 7.1: Binarization using hard threshold.

Figure 7.1 shows how hard thresholding can produce noisy images, as the applied threshold does not depend on local neighborhoods. Images are labeled as left, up and right

respectively.



Figure 7.2: Binarization using Otsu's threshold.

Figure 7.2 shows a sample of the dataset after applying Otsu's binarization.

7.1.2 Quantization

The Gens-Domingos algorithm, as mentioned in chapter 4, has two main steps: a clustering phase and an independency test part. Our independency test implementation in specific uses the standard G-test statistical independence test based on contingency tables, where each frequency of the categories of each two variables are laid out on a matrix and their likelihood ratios are computed. This test takes only O(nm), where n and m are the number of categories of each variable, but each two variables must be tested pairwise with all others, and although we use a spanning-tree heuristic, it still grows fast with n and m.

We empirically found that $\max\{n, m\}$ and training set size are directly correlated to the model's accuracy and speed. If n and m are big and the training set size is small, then accuracy will fall. On the other hand, accuracy increases if n and m are small and the training set is small. However, when both n, m and training set are big, we achieve much better accuracy results, but inference speed increases as well.

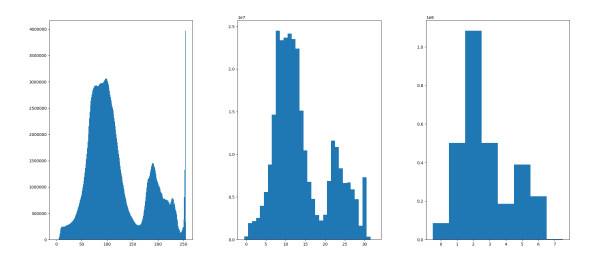


Figure 7.3: Histogram for dataset pixel values on 8-bit, 5-bit and 3-bit image resolutions.

A possible explanation for poor results with small training sets is the low number of pixel intensity values for too extreme values, as there are fewer samples where pixels are either too bright or too dark, as Figure 7.3 shows.

Quantizing the dataset resulted in a significant improvement in accuracy to the model when training with a small set of images (\leq 300). We found that when we increased the number of training images, accuracy depended less on quantization, but inference time increased, as the model grew in depth. We thus needed to find a balance between network complexity and accuracy.

We faced two possible solutions to this problem. Either implement an exact independence test, such as the Fisher exact test (FISHER, 1922), or perform equalization on the dataset. The former was unfortunately not an option, as we found that there were no libraries in Go or C that provided a general case implementation of the Fisher exact test, and implementing our own within our time constraints was out of question. We chose the latter, applying histogram equalization on the entire dataset.

7.1.3 Equalization

Equalization was done using OpenCV. We attempted two methods of histogram equalization: traditional equalization through brightness and contrast normalization, and Contrast Limited Adaptive Histogram Equalization (CLAHE) (Zuidervald, 1994). We found that the CLAHE method resulted in images that were very similar to the output of the traditional method. Since these transformations are also expected to be applied on-the-fly during inference, we chose the standard equalization for its speed. Figure 7.4 shows how the dataset pixel histogram looks like after equalization.

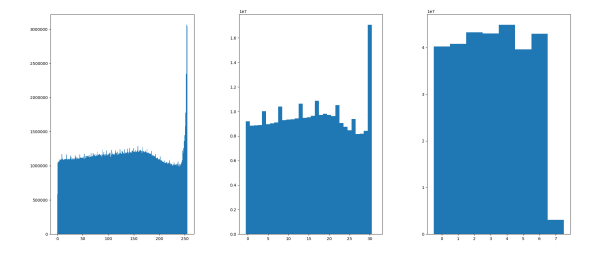


Figure 7.4: Histogram for equalized dataset with 8-bit, 5-bit and 3-bit image resolutions.

When coupling quantization and equalization, we were able to achieve \approx 81% accuracy. Interestingly, these transformations proved to be harmful for the Dennis-Ventura architecture. In fact, the structure yielded better results with higher resolutions compared to

lower. Equalization also had little to no effect on this architecture, increasing accuracy in 1% or 2% when training with a small dataset, and having no effect when training with many samples. We attribute this phenomenom to the classification architecture we discussed in chapter 4. Since each sub-SPN is essentially modelling each class as a separate, independent image model to the other classes, the more details in the image, the easier the model can distinguish from other label images. Furthermore, since the Dennis-Ventura algorithm does not need to run an independence test, it does not suffer from its drawbacks and thus does not depend on an equalized histogram.

7.2 Setups

For our experiments, we tested two structure learning algorithms, the Dennis-Ventura and Gens-Domingos architectures, and for each of these models we evaluated accuracy when applying either generative or discriminative weight learning. We additionally ran tests without applying weight learning to serve as reference.

For the Dennis-Ventura algorithm, we discarded pre-clustering, opting to use the classification architecture mentioned in chapter 4, as it resulted in much better accuracy. We also fixed the number of sums per region and gaussians per pixel to four, and the similarity threshold to 0.975.

For the Gens-Domingos algorithm, we tested two implementations. The first uses k-means for the clustering step with k=2. The second uses DBSCAN, with parameters $\epsilon=4$ the maximum radius of a point neighborhood, and m=4 the minimum number of points to describe a dense region. Both were set with a p-value of 0.01 for the independence step. We refer to the k-means implementation as k-GD, and the DBSCAN variation as DBSCAN-GD.

We found that when using DBSCAN-GD, the resulting structure was too complex, causing inference to take too long. For instance, when running inference on k-GD, average prediction took about 0.4 second. When using DBSCAN, the average time of prediction was 12.33 seconds. Size-wise, the DBSCAN variation had a structure 18 times bigger. However, in terms of accuracy, DBSCAN-GD showed impressive results, with accuracies ranging from 98% to 100%. Despite these results, a model that takes too long for inference is not adequate for a self-driving application. For this reason, we show only comparisons with the k-GD model.

In both training and testing, we first apply image transformations (e.g. quantizing, binarization or equalization) to the dataset and then train or perform prediction with a particular model. The applied image transformation is always identical in training and testing. For example, a valid training pipeline would be choosing to apply 3-bit quantization and equalization to a training dataset, train an SPN structure using k-GD, perform discriminative gradient descent on the resulting structure to learn its weights, and finally save the model for testing. The equivalent testing pipeline for this training pipeline would be applying the same image transformation, in our case 3-bit quantization and equalization, and for each image find the arg $\max_{u \in Y} P(Y = y|X)$.

Both Gens-Domingos and Dennis-Ventura algorithms generate a structure as deep as the number of training samples. The deeper the structure, the more expressive it is. We found that accuracy with models trained with 1000 samples had much better accuracies than with 500. However, a more complex network means inference will take longer. We attempted to keep inference at less than a second per prediction. This was done by restricting the training dataset to 500 images.

We use a particular set of notations for our experiments. For image transformations, we denote by Q_n as applying an n-bit quantization of the dataset. An E means the dataset was equalized, and a B means it was binarized. Any combination of image transformation is signalled with a + sign. A \emptyset means there were no image transformations done to the dataset apart from grayscale conversion.

For learning algorithms, a GD means we are using k-means Gens-Domingos and DV Dennis-Ventura. This is then followed by the weight learning algorithm used. The letters g, d and s mean we either applied generative gradient descent, discriminative gradient descent or no weight learning.

7.3 Accuracy

In this section we show accuracy results in each setup. All values are in percentage of hits.

Accuracy (%)	DV+g	DV+d	DV+s	GD+g	GD+d	GD+s
В	76.0	00.0	76.6	00.0	78.2	78.8
Q_2	75.6	00.0	76.6	0.00	00.0	77.6
$Q_2 + E$	75.8	00.0	76.6	0.00	00.0	78.0
Q_3	76.0	00.0	76.6	0.00	71.8	79.2
$Q_3 + E$	75.2	76.6	76.6	0.00	80.2	80.2
Q_4	75.8	00.0	76.6	0.00	00.0	79.8
$Q_4 + E$	75.6	00.0	76.6	0.00	00.0	82.0
Q_5	76.0	76.6	76.6	0.00	73.8	78.0
$Q_5 + E$	75.6	76.6	76.6	0.00	81.2	78.8
Q_6	76.0	00.0	76.6	0.00	00.0	80.8
$Q_6 + E$	75.6	00.0	76.6	0.00	00.0	77.6
Q_7	76.6	0.00	76.6	0.00	00.0	79.2
$Q_7 + E$	76.6	00.0	76.6	0.00	00.0	79.6
Ø	75.6	76.6	76.6	0.00	69.6	80.4
E	75.6	76.6	76.6	0.00	78.6	79.0

Table 7.1: Accuracy values for each possible model permutation.

Table 7.1 shows some interesting results. The first is that generative gradient descent on the Dennis-Ventura architecture had a small to no negative impact on the performance of the network.

The Gens-Domingos architecture, on the other hand, achieved better results, but showed a more unpredictable behavior.

7.4 Speed

Training (mins)	DV+g	DV+d	DV+s	GD+g	GD+d	GD+s
В	54m54s	48m42s	00m19s	00.0	0.47	05m50s
Q_2	53m46s	0.00	00m17s	00.0	0.42	04m44s
$Q_2 + E$	53m00s	0.00	00m15s	00.0	0.33	04m55s
Q_3	60m19s	00.0	00m15s	00.0	0.40	05m23s
$Q_3 + E$	62m06s	76.6	00m22s	00.0	0.34	04m28s
Q_4	50m14s	0.71	00m18s	00.0	0.31	04m39s
$Q_4 + E$	50m19s	0.00	00m16s	00.0	0.37	05m43s
Q_5	54m22s	0.61	00m15s	00.0	0.34	04m55s
$Q_5 + E$	50m32s	0.66	00m24s	00.0	0.33	04m06s
Q_6	50m54s	0.00	00m14s	00.0	0.37	04m19s
$Q_6 + E$	56m32s	0.00	00m15s	00.0	0.34	05m23s
Q_7	56m58s	00.0	00m15s	00.0	0.40	05m52s
$Q_7 + E$	60m13s	0.00	00m15s	00.0	0.42	05m52s
Ø	62m14s	0.57	00m15s	00.0	0.52	06m45s
E	57m23s	0.57	00m15s	00.0	0.37	04m58s

Table 7.2: Average time in minutes for training each model.

Inference (secs)	DV+g	DV+d	DV+s	GD+g	GD+d	GD+s
В	0.58	00.0	0.79	00.0	78.8	0.47
Q_2	0.56	00.0	0.63	00.0	77.6	0.42
$Q_2 + E$	0.62	00.0	0.61	00.0	78.0	0.33
Q_3	0.65	00.0	0.59	00.0	79.2	0.40
$Q_3 + E$	0.57	76.6	0.69	00.0	80.2	0.34
Q_4	0.56	0.71	0.65	00.0	79.8	0.31
$Q_4 + E$	0.57	00.0	0.61	00.0	82.0	0.37
Q_5	0.56	0.61	0.66	00.0	78.0	0.34
$Q_5 + E$	0.57	0.66	0.62	00.0	78.8	0.33
Q_6	0.65	00.0	0.62	00.0	80.8	0.37
$Q_6 + E$	0.57	00.0	0.60	00.0	77.6	0.34
Q_7	0.56	0.00	0.59	00.0	79.2	0.40
$Q_7 + E$	0.56	0.00	0.58	00.0	79.6	0.42
Ø	0.64	0.57	0.58	00.0	80.4	0.52
E	0.62	0.57	0.59	0.00	79.0	0.37

Table 7.3: Average time in seconds to predict a single image.

Real world

In this chapter we finally apply our model to a real-world implementation of mobile robot lane following.

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