Limits of End-to-End Learning

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Abstract
End-to-end learning refers to training a possibly complex learning system by applying gradient-based learning to the system as a whole. End-to-end learning systems are specifically designed so that all modules are differentiable. In effect, not only a central learning machine, but also all “peripheral” modules like representation learning and memory formation are covered by a holistic learning process. The power of end-to-end learning has been demonstrated on many tasks, like playing a whole array of Atari video games with a single architecture. While pushing for solutions to more challenging tasks, network architectures keep growing more and more complex.

In this paper we ask the question whether and to what extent end-to-end learning is a future-proof technique in the sense of scaling to complex and diverse data processing architectures. We point out potential inefficiencies, and we argue in particular that end-to-end learning does not make optimal use of the modular design of present neural networks. Our surprisingly simple experiments demonstrate these inefficiencies, up to the complete breakdown of learning.

Keywords: end-to-end machine learning

1. Introduction

We are today in the position to train rather deep and complex neural networks in an end-to-end (e2e) fashion, by gradient descent. In a nutshell, this amounts to scaling up the good old backpropagation algorithm (see Schmidhuber, 2015 and references therein) to immensely rich and complex models. However, the end-to-end learning philosophy goes one step further: carefully ensuring that all modules of a learning systems are differentiable with respect to all adjustable parameters (weights) and training this system as a whole are lifted to the status of principles.

This elegant although straightforward and somewhat brute-force technique has been popularized in the context of deep learning. It is a seemingly natural consequence of deep neural architectures blurring the classic boundaries between learning machine and other processing components by casting a possibly complex processing pipeline into the coherent and flexible modeling language of neural networks.\(^1\) The approach yields state-of-the-art results (Collobert et al., 2011; Krizhevsky et al., 2012; Mnih et al., 2015). Its appeal is a unified training scheme that makes most of the available information by taking labels (supervised learning) and rewards (reinforcement learning) into account, instead of relying only on the input distribution (unsupervised pre-training). Excellent recent examples of studies

\(^1\) In practice, some preprocessing steps like color space normalization of images are still best done outside the framework.
deeply rooted in the e2e learning philosophy—naming only a few well known ones—are the neural Turing machine (Graves et al., 2014) and the differentiable neural computer (Graves et al., 2016), value iteration networks (Tamar et al., 2016), and vision-based navigation in 3D environments (Mirowski et al., 2016). These achievements defy the well-known principal limitations of gradient descent, namely local optima and slow convergence in various circumstances depending on the exact algorithm in use, typically on badly conditioned problems.

Intuitively, both of these problems may become more severe when network architectures grow in complexity.

The current state-of-the-art is based on a long sequence of technological advancements, which took only about one decade to evolve. Major steps and factors are listed in the following.

- Interestingly, the success story of deep learning as we know it today (notwithstanding many earlier findings which, however, did not make a comparable impact at the time (LeCun et al., 2005; Schmidhuber, 2015) started with structured, layer-wise training of deep belief networks (Hinton et al., 2006) and stacked auto-encoders (Vincent et al., 2008). These techniques can be understood as the exact opposite of e2e learning. Although they mark breakthroughs of representation learning, they are widely considered to have become obsolete.
- More “classic” networks moved back into the focus due to progress on efficient GPU implementations of backpropagation for deep convolutional neural networks (CNNs) and, as a consequence, significant progress on computer vision tasks (Ciresan et al., 2011).
- GPU-based processing allowed to scale to extremely large networks and learning problems. The availability of huge training sets and the computational power to process them were key prerequisites for solving previously intractable tasks with deep learning techniques (Krizhevsky et al., 2012).
- The development went hand-in-hand with tremendous progress in the area of stochastic gradient descent (SGD). On the one hand side, linearly convergent methods for finite sums were developed (Schmidt et al., 2013; Johnson and Zhang, 2013), on the other hand, effective component-wise online adaptation techniques for learning rate parameters resulted in significant speed-ups over plain SGD. The new techniques made deep learning far more tractable and resulted in even better performance (Zeiler, 2012; Kingma and Ba, 2014).
- Various easy-to-use deep learning software libraries were developed (Jia et al., 2014; Chollet, 2015; Vedaldi and Lenc, 2015), many of which are based on powerful symbolic computation software packages like theano (Theano Development Team, 2016) and tensorflow (Abadi et al., 2016).
- The race for solving Imagenet classification (Deng et al., 2009) resulted in deeper and deeper architectures. It lead to novel, modular connectivity patterns (Szegedy et al., 2015), and also shed new light on the utility of good shortcut connections (He et al., 2016).
- The immense progress even allowed to train highly flexible networks in a (vision-based) reinforcement learning fashion (Mnih et al., 2015; Silver et al., 2016; Mirowski et al., 2016), despite the limited and sometimes delayed information contained in the reward signal.
In this paper, we take the freedom to extrapolate this development into the (foreseeable) future. We discuss the implications of applying e2e learning methods to considerably more complex systems, which we today envision to be required for solving the “holy Grale” of tasks, namely operating autonomous agents in a human environment.

As such, and although we present experimental results later on, we consider this work primarily as a position paper. Our central claim is that e2e learning has limitations that will keep us from using it in its current form as the sole way of training networks in the future. This claim necessarily remains unproven. Also, although we point into a direction of a possible solution, we do by no means claim to have an alternative to e2e learning ready for use.

The work by Shalev-Shwartz et al. (2017) is closely related to ours. It emphasizes the reasons for failure of e2e learning on the conceptually lowest level of machine learning, namely its foundation formed by (stochastic gradient-based) optimization. This rather technical perspective is insightful, and for very specific cases it allows for an analysis in terms of well-known problems like vanishing gradients and ill-conditioned problems. The paper also emphasizes the value of problem decomposition of a complex architecture as an alternative to e2e learning. Here we take the high-level perspective of neural network architectures and the consequences of propagating information through a number of network modules as a prerequisite for e2e learning.

The remainder of this paper is organized as follows. In the next section we present and discuss the problem in an informal way and point towards a possible route to solution. We summarize this discussion by collecting merits and limitations of e2e learning for better overview. Then we underpin our arguments experimentally: we show how e2e learning can fail for the training even of rather small systems due to non-trivial couplings that originate either from the network structure itself or from the task. We close with a brief conclusion.

2. Model Engineering vs. End-to-end Learning

State-of-the-art e2e learning enjoys a high degree of automation, however, it does not work fully autonomously: the design and configuration of a learning system suited for a given task requires a certain level of experience and machine learning knowledge. Today’s deep learning systems are engineered from modules, often organized into layers and groups of layers with specific roles. Prominent examples are alternating sequences of convolution and pooling layers, fully connected layers, auto-encoders (bottleneck layers), and LSTM layers. Even dropout, which is a regularization technique and not a model component, it often thought of as a layer, which (at least in software) can be added on top of any other layer. The general proceeding is to compose a system with the capacity—in principle and by design—to exhibit a desired behavior. After the design phase, full automation takes over: weights are initialized and random, and they are subsequently trained, e2e, with variants of stochastic gradient descent.

Driving this to the extreme, one could try to design a full “brain” this way, consisting of cortical areas like visual cortex, auditory cortex, motor cortex, a hippocampus, and so forth. Of course, there is no need for the design to resemble biological brains in structure and connectivity, however, for any non-trivial agent we will surely end up with dedicated
modules for sensory data processing, natural language processing, memory, decision making, planning, motor control, and more (see Figure 1 for an illustration).

Problem decomposition is central to solving a complex problem. This divide and conquer approach is the core principle of engineering. It is underlying our network design decisions, and we are well aware that learning problems become much harder if we don’t know how to decompose them, or if we simply get it wrong.

Hence, divide-and-conquer is heavily used for the design of complex learning machines. However, training such a system in an end-to-end fashion means to explicitly ignore the problem decomposition during learning. Instead, training is done in good hope that the structural preconditioning is sufficiently strong to direct a method as simple as gradient descent from a random initial state to a highly non-trivial solution. Intuitively, this is a dangerous assumption: since the resulting optimization problem is by no means convex, learning can only work if the task is “nearly trivial” given the structure on which it operates. Data efficiency is a second concern, since unmodeled interactions between modules may require an amount of training data that grows possibly exponentially with the number of modules for the problem to be well-sampled.

One could argue that compared to a fully connected (possibly recurrent) network modular models do indeed provide a useful problem decomposition also during learning, simply because many potential connections are not present, which is equivalent to fixing their weights to zero. However, this structure is not exploited in the optimization algorithm, which is usually blind to the structure.

Let us return to the above example of the brain-like model. Assume that each module is realized as a neural network. The roles of these modules reflect the network designer’s intention and his ideas of how to structure a possible solution to an overwhelmingly complex
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problem. However, these roles are not assigned formally to the modules. Nothing enforces that a specific module actually learns to fulfill its role and only that role. The role is only suggested by the pre-defined network structure, and the near-black-box character of complex neural networks makes even checking whether the assumptions are fulfilled or not a non-trivial task. Hence, the learning rule may come up with an alternative solution, or something that looks like a potential solution during an early training phase. More often than not, this route may lead to convergence into a poor local optimum.

Against this background it seems obvious that we should respect and even explicitly exploit the carefully chosen structure during training. There are many possible ways how to ensure that the training procedure respects our design plan. It seems that the simplest strategy is to structure the training process directly in terms of network modules and their connectivity. A simple procedure with this property is greedy, layer-wise training, however, its performance is usually inferior to e2e learning due to its inability to use labels and rewards. We consider finding a training procedure that makes the powerful of e2e learning approach aware of the network structure a grand challenge of neural network research.

The insight that existing learning methodologies may not scale to arbitrarily complex tasks and to learning systems matching this complexity is by no means new. It motivated various proposals that were popularized in the domains of control, reinforcement learning, and artificial intelligence (Ring, 1994; Abbeel and Ng, 2004; Schmidhuber, 2004; Bengio et al., 2009; Schmidhuber, 2010; Thrun and Pratt, 2012), known under different names like learning to learn, curriculum learning, continual learning, and many more. A prominent idea is to bootstrap the learning process by solving simple tasks first, in the hope that solution components of generic value evolve, which turn out to be helpful building blocks for more complex tasks later on. All of this can, but does not have to happen in an e2e fashion. The approach is very different and actually orthogonal to our vision of organizing the training process of neural networks along their structure.

3. Merits and Limitations of End-to-end Learning

Let us have a more systematic look at the strong and weak spots of the e2e learning principle. The following lists are intended to provide a hopefully useful overview of pros and cons of the method, as we see them, without claiming completeness. The purpose is to offer a basis for supporting the decision whether e2e learning may be suitable for a problem at hand or not. It is not our intent to judge the method according to the number or weight of items on either side.

Merits

- Conceptual and mathematical beauty: the system is trained in a holistic manner based on a single principle.
- Every learning step is directed at the final goal, encoded by the overall objective function. There is no need to train modules on an “auxiliary” objective,\(^2\) unrelated to the task, like contrastive divergence or reconstruction error. However, it should be noted that e2e learning can in principle incorporate auxiliary objectives, possibly

\(^2\) Here we do not refer to cases like using a differentiable loss like cross-entropy instead of the 0/1 loss, which is of final interest, but rather to using an entirely unrelated loss, often because labels and rewards have not (yet) been propagated to the module.
affecting only parts of the network, usually with the goal of avoiding slow learning and bad local optima (Mirowski et al., 2016).

- The power of e2e learning for training powerful predictors was proven many times in various domains (Collobert et al., 2011; Krizhevsky et al., 2012; Mnih et al., 2015; Silver et al., 2016).

- E2e learning is nicely consistent with the general approach of machine learning to take the human expert out of the loop and to solve problems in a purely data-driven manner.

Limitations

- The principal limitations of (stochastic) gradient descent apply, like slow convergence on ill-conditioned problems, convergence into possibly poor local optima, and vanishing gradients due to saturating non-linearities. The implications of these problems for deep e2e learning are discussed in detail by Shalev-Shwartz et al. (2017).

- In some cases, the learning signals used for e2e learning can be of inappropriate nature. For example, a network module responsible for visual representation learning can be trained together with a very different module representing a policy (Mnih et al., 2015), based on possibly sparse and delayed rewards. It seems to be far more efficient to train the vision module independently, either with unsupervised methods (exploiting the visual input is far more rich than the reward signal), or starting from one of the many available pre-trained networks.

- The valuable information contained in the problem decomposition that resulted in a specific network design is ignored during e2e training. Intuitively, this can be dangerous, in particular if modules interact in a non-trivial way, since they can hamper each other’s learning progress. Examples in section 4 demonstrate that such interactions can slow down learning significantly, to the point of a complete breakdown. It seems that, at least in principle, the decomposition into modules could be used to device structured training schemes with the potential to overcome these problems.

Trade-Offs

- In a general analysis, the convergence speed of gradient-based methods is independent of the number of parameters (Polyak, 1963). Therefore it can be expected that training all modules of a network at once takes fewer gradients (data samples) than training them independently, e.g., in a greedy layer-wise or otherwise structured manner. However, this argument can be disputed since the learning speed depends on the conditioning of the problem, which can well be significantly better for sub-problems involving only few modules.

This summary does not yield a general conclusion. We leave it as such and instead turn to concrete learning systems designed to stress the limits of e2e learning.

4. Experiments

In previous sections we have made seemingly speculative claims about possible issues resulting from applications of e2e learning to networks consisting of interacting modules. In this section we will demonstrate that such problems indeed exist. We aim to validate the following claims empirically:
- As the network complexity or the task difficulty grows, e2e learning can become inefficient.
- For too complex tasks or networks, e2e learning can fail completely.
- Training modules one at a time can solve this issue.\textsuperscript{3}

We demonstrate these claims based on experiments in the following.\textsuperscript{4}

### 4.1. Scalable Stacking

This first series of experiments relies on a network structure with freely scalable complexity, i.e., a class of networks with parameterizable number of modules. It is not the purpose to design a learning system that solves a realistic, complex task, but to demonstrate certain effects in as clean as possible system. Therefore we aim to keep things plain and simple.

We start with a rather trivial supervised classification problem with $B \in \mathbb{N}$ classes. Inputs are one-hot-encoded numbers in the range $1, \ldots, B$, and so are the target outputs, which agree with the inputs. We apply a standard setup, which is mini-batch gradient descent training with the ada-delta method (Zeiler, 2012), minimizing the cross-entropy loss.

While in a real application network modules may have widely varying structure and connectivity, we define a single module to be applied in a sequence of arbitrary length (or depth).

**Network Module.** The module takes one-hot encoded numbers in the range $1, \ldots, B$ as input, sends them through a bottleneck layer of size $b \ll B$ with sigmoidal (logistic) activation function, and propagates the result to the output layer of size $B$ with softmax activation. The goal of learning is to represent the identity mapping restricted to one-hot encoded vectors; for chained modules, the overall chain shall represent the identity mapping.

For $B = 2^b$ the problem can be solved by a demultiplexer/multiplexer logic circuit, which translates one-hot-encoded numbers to a binary representation and back. We verified empirically that networks can reliably learn solutions even for values of $B$ significantly larger than $2^b$, e.g., $B = 64$ and $b = 4$. In principle, even as few as $b = 2$ hidden units suffice, for any number $B$ of inputs/outputs. In the sequel we resort to the rather simple problem with parameters $B = 10$ and $b = 4$. Including the biases, the total number of weights per network module is as small as 94. This module fulfills our requirements of performing a non-trivial computation, and being stackable without limit.

**Stacked Modules.** In a first experiment we train $N$ network modules stacked on top of each other. The data set consists of 10 points, with input and target corresponding to the 10 possible one-hot encodings. We train the network until it has learned the identity mapping, i.e., the classification error on the training data drops to zero. Note that we are not interested in the question whether the network generalizes, which is trivial in this case since the training data cover all possible one-hot encodings.

\textsuperscript{3} Caveat: we do not claim to offer a constructive alternative to e2e learning. We do not know a way how to train modules in isolation that works generically across a large number of tasks and could act as a plug-in replacement for e2e learning.

\textsuperscript{4} Our code is based on the Keras library (Chollet, 2015). It is available from the first author’s homepage: \url{https://www.ini.rub.de/PEOPLE/glasmtbl/code/limits-of-e2e-learning/code-supplement.zip}
In figure 2 we provide the number of epochs until training is completed. Note that each epoch corresponds to only 10 gradients, therefore the numbers are large. We see a roughly exponential increase of the training effort. For \( N = 5 \) modules (not shown), 8 out of 10 training runs hit the limit of \( 10^9 \) gradients computations (\( 10^8 \) epochs) without solving the task.

In a control experiment, the weights between the different modules were shared. Hence the module received gradients from all layers at once. The scaling was similar, but the numbers were even worse.

This is an interesting and insightful result. All networks are rather small, featuring only a few hundred weights. Training a single module is easy. However, training multiple modules at once is much harder, with effort growing exponentially, by roughly one order of magnitude per module. This implies that training of multiple modules interferes in a non-trivial and highly disruptive way.

It can be argued that the problem is easily avoided by a residual learning approach (He et al., 2016). This is certainly true for our setup, however, recall that the simplistic architecture is supposed to act as a drastically simplified model of a complex network where different modules have different roles. For example, feeding raw vision input directly into a high-level decision making module (e.g., behavior planning) usually does not make a whole lot of sense, since the decision making module relies on high-level cues that are output of the vision module, possibly further processed by other modules into a stable scene representation.

**Handwritten Digit Classification.** We extend the above experiment to test how the difficulty to learn with multiple modules affects the training of proven components. For this purpose we created a fairly straightforward network for classifying handwritten digits from the well-known MNIST data set. The network consists of the following layers: \( 5 \times 5 \) convolution (relu), \( 2 \times 2 \) max-pooling, \( 5 \times 5 \) convolution (relu), \( 2 \times 2 \) max-pooling, fully connected layer with 200 nodes (relu), and finally a fully connected output layer with 10 nodes (softmax). We refer to this rather plain architecture as the “basic MNIST module”. It achieves non-trivial results (\( \approx 99.3\% \) correct after about 10 epochs). Then we stack \( N \geq 0 \) of the above described modules on to of the basic MNIST module. The network is trained in an e2e manner until the validation error stalls for at least 20 epochs in a row.
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Figure 3: Number of epochs (left) and resulting classification accuracy (right) for the handwritten digit recognition task, with $N \in \{0, 1, 2, 3, 4\}$ modules stacked on top of the basic MNIST module. The solid lines are averaged over 10 runs, individual runs are represented as black circles. Note that the performance values tend to cluster roughly at multiple of 10%, corresponding to the overwhelming majority of a classes being classified correctly.

Validation accuracies and numbers of epochs for this experiment are provided in figure 3. It is understood that the results are no better than in the first experiment; actually, they are worse. With a single module on top of the basic MNIST module, the system achieves an accuracy of nearly 99% correct, however, with significantly increased training effort compared to the basic MNIST module and slightly worse accuracy (and with frequent failures if the early stopping criterion is set to 10 instead of 20 epochs without improvement of the validation error). For $N = 2$ the performance drops significantly, in only 3 of 10 runs does the network manage to classify all digit classes correctly. For $N = 3$, this rate drops to 1 out of 10 runs, and with $N = 4$ modules, learning fails entirely. The weights remain very close to their initial values.

Figure 4 illustrates the learned filters of the first network layer at the end of the epoch with best validation accuracy, i.e., the network selected by early stopping. Various orientations are clearly recognizable for the first three networks. All networks are initialized exactly the same, hence they evolve similar filters. The networks with $N \geq 3$ modules fail to evolve meaningful filters; indeed, the filter weights remain close to their initial values.

As a control experiment, we tested the same networks with the basic MNIST module initialized with well-tuned filters, and all further modules initialized at random as before. The results were overall comparable. With $N = 4$ modules on top the networks did not learn, however, training also did not destroy the existing filters.

We find that stacking non-trivial modules on top of the basic MNIST module results in significant slow-downs, and quickly to a complete breakdown of end-to-end training. In this case, none of the layers is able to learn, not even the first convolution layer, which is
Figure 4: Visualization of the convolution filters in the first layer before and after training, and the corresponding validation accuracy, for the first out of 10 runs.

closest to the data. This is easily explained by the fact that the gradient at each node is affected by all weights during the forward pass, and by all subsequent weights during the backwards pass. Hence, as long as none of the components is reasonably well-trained, no component can expect to receive a meaningful update direction, unless there is a rather broad and hopefully straight path from random initialization to the goal state. This is in contrast to greedy, layer-wise training, which is entirely unaffected by later layers, and by their detrimental effect on the gradient.

4.2. Sequence Forecasting

We consider a classic sequence learning problem introduced by Hochreiter and Schmidhuber (1997) for demonstrating the power of LSTM cells. We are given two sequences \((x, a_1, a_2, \ldots, a_n, x)\) and \((y, a_1, a_2, \ldots, a_n, y)\), where the symbols are one-hot encoded. The task is to predict the next input \((x\) or \(y)\) for the duration of the whole sequence. The authors apply a straightforward network architecture with a hidden LSTM layer and skip connections. They train the network in two phases: first only the skip connections (a non-recursive and essentially linear model) are trained until convergence, then a single LSTM cell is added. This is to avoid what they call the “abuse problem”: this problem decomposition avoid the “abuse” of the LSTM cell, e.g., as a bias units.

We investigate this phenomenon more closely by training networks with the above problem decomposition, and alternatively in an e2e fashion with the LSTM unit added from the
start. We suspect that due to modern techniques like Glorot initialization (Glorot and Bengio, 2010) and adaptive learning rates (Zeiler, 2012) the problem is less severe compared to the 1990es, however, figure 5 shows that the effect is still pronounced.

Already in this 20 years old experiment it was found that e2e learning can be inefficient. However, the problem was not recognized as an instance of a more general phenomenon.

4.3. Planning

In the board game of RoboRally, in each turn a simplistic mobile robot is pre-programmed for five movements steps in a row, in a race to reach a goal location. Most of the fun of the game comes from the impossibility of reliable planning due to unforeseen interactions with other robots, programmed by other players, resulting in robots being pre-programmed with the best intentions, but moving straight into fatal disaster. For our planning task, we ignore these interactions and consider a simplified setup that is more alike to a classic grid world navigation task, however, we stick to the game element of pre-programming five moves in a row.

The robot’s state is given by its position (2D integer coordinates) and orientation (north, south, east, west) in a grid world. It plays only a single turn, consisting of five elementary actions chosen from the set move_forward, turn_left, turn_right, and wait. A grid cell is either a normal (empty) cell, the goal cell, a wall (obstacle), a bottomless pit, a cell with a laser, or a conveyor belt. Moving into a wall is a legal and available action, but does not change the robot’s state. Entering a pit cell kills the robot instantly, entering the goal cell results in immediate success. In these cases the robot stops moving (ignoring further actions), so the states are terminal. However, each episode is always run for five steps and the robot keeps collecting rewards. Hence reaching the goal quickly is beneficial. For simplicity, in our setup all conveyor belts transport the robot one field to the north (after its own movement), and lasers (usually damaging the robot) have no effect other than giving negative reward. Rewards are structured as follows: goal +10, pit -10, laser -1, and in order to encourage exploration, the wait action is penalized with an “intrinsic” reward of -1. This is a quite simple and

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5. Designer: Richard Garfield; Publisher (as of 2017): Hasbro, Inc.
6. The wait action can be useful when interacting with certain game elements like conveyor belts. The original game dynamics are more complex, adding multi-step and backwards movements, and in particular by executing the steps of multiple robots in a specific order.
actually deterministic Markov decision process (MDP). We used the map shown in figure 6, which allows the robot to reach the goal in only four steps (optimally: turn_right followed three times by move_forward).

Playing this game successfully and even optimally is not hard at all, since there are only \(4^5 = 1024\) possible action sequences, 25 of which reach the goal, and the environment is deterministic. Our agent proceeds by planning its actions based on a forward model, which by itself is a neural network and must be learned from interactions. Given a state-action pair \((s, a)\) with \(s\) and \(a\) each one-hot-encoded (representing explicit probabilities, the outer product representing the joint distribution acts as the actual input), the forward model maps to a probability distribution over successor states. It is implemented by a fully connected network layer with linear activation and probability simplex constraints. Expected immediate rewards are modeled analogously with a linear layer without any constraints. This model is extremely simple in nature; it is essentially linear. The model can capture the true transition and reward structure of the MDP exactly, and it is suitable for propagating distributions over states and actions arbitrarily far into the future. This property makes the model suitable for planning, e.g., by executing mental trials. Of course, the predictions will be off as long as the model is not yet well trained. The actions are encoded as probability distributions, realized as a softmax layer receiving five one-hot encoded “time step” inputs.

The overall agent consists of two trainable modules, the world model mapping state \(s\) and action \(a\) to successor state \(s'\) and reward \(r\), and the programming module (action selector module), mapping time \(t \in \{1, 2, 3, 4, 5\}\) to an action \(a\) from the action set defined above. The full system is displayed in figure 6.

The system is trained in an end-to-end manner for 10,000 episodes. In each step, an episode is planned. The plan is refined by a gradient step on the programming module, based on the total reward predicted by the forward model. I.e., the agent improves its policy based on its current world model. Then an actual action sequence is sampled from the updated action distribution (the actual action in the RoboRally game), and the episode is executed in the environment. Note that this setup differs from standard reinforcement learning, since the agent essentially performs only a single action with a “reactive” policy, consisting of five sub-actions, which must be pre-programmed and hence planned upfront. Then the world model module is updated based on the observed successor states and rewards.\(^7\)

This setup is obviously not the most efficient possible for the task at hand, which can of course be solved easily, e.g., by tabular reinforcement learning, or simply by brute force enumeration of all action sequences, or assuming possible stochasticity, by multi-armed bandit algorithms. Therefore it must be emphasized that this is a toy experiment, the only purpose of which is to demonstrate the effects of unmodeled dependencies between learning modules. Our learning system indeed has the property of interest, namely a non-trivial mutual dependency of the two learning modules: the action selector depends on the quality of the world model for correct forward planning, and the world model depends on the action selector to perform sufficient exploration so that it gets to see the relevant parts of the state space.

Despite the simplicity of the task, learning both modules in the above described intertwined manner works only in 47 out of 100 runs. In the other 53 runs the system quickly

\(^7\) Our learning system is related to value iteration networks (Tamar et al., 2016), however, in our case the term planning actually refers to performing a full “mental trial” based on a learned forward model.
converges to a local optimum, consisting of turning movements, since they avoid negative rewards. This task is on the edge of the manageable difficulty; even slightly more complex tasks result in near certain failure. This is despite the facts that both modules can represent the optimal solution exactly (the solution is realizable), the time horizon is rather small, the environment consists of only 24 states with four actions, and learning any of the modules in isolation works flawlessly: learning the MDP from random actions works fine, and so does learning the optimal policy from the exact MDP or a pre-trained forward model. This result suggests a simple solution: first train the world model till convergence based on random actions, then train the action selector. This decomposition of the training

8. Care must be taken not to insert too many pits, since otherwise the chance of dying during exploration exceeds that of reaching the goal. That turns “not moving” into a local optimum, which is easily reachable from the initial random policy. An explicit exploration strategy would be needed to overcome this problem.
process is exactly in line with the problem decomposition underlying the network design. Of course, in its simple form this solution is dissatisfactory since random exploration is often inefficient, but it is a simplistic demonstration showing that a learning process organized along the network structure can indeed be a viable solution.

5. Conclusion

We have demonstrated that end-to-end learning can be very inefficient for training neural network models composed of multiple non-trivial modules. End-to-end learning can even break down entirely; in the worst case none of the modules manages to learn. In contrast, each module is able to learn if the other modules are already trained and their weights frozen. This suggests that training of complex learning machines should proceed in a structured manner, training simple modules first and independent of the rest of the network.

Our example problems are necessarily somewhat contrived. Considering neural networks designed for solving real tasks, whether the limits of end-to-end will show up in the foreseeable future or not remains to be seen. At this point we simply want to raise awareness for the existence of limitations. To overcome these problems in a principled manner, we believe that new structured learning paradigms are needed, which should be in line with the network structure, and which may or may not contain greedy learning and end-to-end learning as techniques. We are convinced that such structured learning paradigms would allow us to push the boundaries of training complex learning systems beyond the current state-of-the-art.

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