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## Don't Learn What You Already Know Scheme-Aware Modeling for Profiling Side-Channel Analysis against Masking

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Abstract. Over the past few years, deep-learning-based attacks have emerged as a de facto standard, thanks to their ability to break implementations of cryptographic primitives without pre-processing, even against widely used counter-measures such as hiding and masking. However, the recent works of Bronchain and Standaert at TCHES 2020 questioned the soundness of such tools if used in an uninformed setting to evaluate implementations protected with higher-order masking. On the opposite, worst-case evaluations may be seen as possibly far from what a real-world adversary could do, thereby leading to too conservative security bounds. In this paper, we propose a new threat model that we name scheme-aware benefiting from a trade-off between uninformed and worst-case models. Our scheme-aware model is closer to a real-world adversary, in the sense that it does not need to have access to the random nonces used by masking during the profiling phase like in a worst-case model, while it does not need to learn the masking scheme as implicitly done by an uninformed adversary. We show how to combine the power of deep learning with the prior knowledge of scheme-aware modeling. As a result, we show on simulations and experiments on public datasets how it sometimes allows to reduce by an order of magnitude the profiling complexity, i.e., the number of profiling traces needed to satisfyingly train a model, compared to a fully uninformed adversary.

Keywords: Profiling Attacks  $\,\cdot\,$  Side-Channel  $\,\cdot\,$  Deep Learning  $\,\cdot\,$  Gradient Descent  $\,\cdot\,$  Masking  $\,\cdot\,$  Scheme-Aware

## 1 Introduction

**Context.** The past few years have seen the emergence of new promising lines of research in profiling Side-Chanel Analysis (SCA), which coincided with the recent advances in Machine Learning (ML) during the 2010's. Indeed, profiling attacks may be formalized as a *supervised learning* problem. As an example, the Gaussian Templates (GTs) initially proposed by Chari *et al.* in their seminal work [CRR03] are actually equivalent to a Quadratic Discriminant Analysis (QDA) in the ML terminology [HTF09]. Hence a vast investigation of relevant learning algorithms in the ML zoology, beyond those generative models [HGD<sup>+</sup>11, BL12, HZ12, LBM14, LBM15, PHG17]. In particular, following the remarkable performance of Deep Neural Networks (DNNs) in solving tasks in computer vision, the SCA community has progressively drawn its interest on such models [GHO15, MZ13, MDM16]. Nowadays, DNNs are known to be able to defeat most of the countermeasures used to protect implementations against SCA, namely de-synchronization [CDP17, KPH<sup>+</sup>19], shuffling [MDP19a, MS21] and more interestingly masking [MPP16, Tim19].

The Uninformed-vs.-Worst-Case Dichotomy. Although the supervised attack threat model introduced so far is nowadays widely adopted by the SCA community for security evaluations, one technical detail of this scenario lacks some consensus. Indeed, there exists a debate among SCA practitioners about what is known or unknown by the adversary during the profiling phase, during which one builds the attack model upon traces measured on an open clone device. In particular, whether one has access to the random nonces used by the clone device during the encryption, as part of the profiling data. This question is not trivial, since nowadays most of the counter-measures — like masking or shuffling — consist in turning a deterministic cryptographic primitive into a non-deterministic implementation. On the one hand, academia usually assumes the adversary to know the values of the random nonces, in a so-called *worst-case* threat model [ABB+20]. This model trades off some potentially conservative security levels against an easy-to-analyze evaluation approach thanks to theoretical shortcuts [DDF14, DFS15]. On the other hand, practitioners such as industrial developers and evaluators rather assume the adversary to not have access to the random nonces used by the clone device during the encryption, hence the name of uninformed threat model [MPP16, CLM20, PP20]. This scenario has the two advantages of being closer to a real-world adversary, and to be fully automatized. As a drawback, some current attacks in the uninformed settings can be much less efficient than worst-case attacks [BS20].

To what extent one threat model or another better fits the security context of the evaluation? The answer is not unique, and both threat models have their proponents and opponents. As an example, scenarios with uninformed adversary may be considered for good or bad reasons. On the one hand, it spares lots of human efforts and expertise spent in pre-processing the traces, which are taken into account in the assessment of an attack potential [SOGISS20]. On the other hand, developers may be tempted to artificially restrict the access to random nonces on the clone device given for evaluation in order to maximize the chances to pass certifications, although at the cost of a false sense of security.

Actually, both uninformed and worst-case models may be seen as the edges of a broad scope of threat models ranging from weak adversaries in the uninformed model to stronger ones in the worst-case model. Yet, realistic threat models often lie all along this spectrum. As an example, the adversary may have access to the source code of the Target of Evaluation (T.O.E.), without necessarily having the possibility to modify it. This typically covers the threat models investigated by security evaluations of many T.O.E.s, such as *native* platforms or applets [SOGISS20]. Moreover, for the security evaluation of *open* platforms, the evaluator may assume the adversary to know the source code of the cryptographic library,<sup>1</sup> but the latter one cannot be assumed to have the rights to modify the code. Even in the case where the developer is willing to collaborate with the evaluator by providing modified versions of the T.O.E. for evaluation purposes, the evaluator is then reduced to a *characterization* of the device, which differs from an *attack* as the latter one should be fully realizable without such help from the developer [SOGISS20].

Likewise, the number of different masking schemes in the literature is restricted enough so that it may be assumed to be known by the adversary. Surprisingly, to the best of our knowledge, no ML approach leveraging weaker adversaries than worst-case but still stronger than uninformed have been considered so far.<sup>2</sup> Indeed, when profiling masked implementations most of the Deep Learning (DL)-SCA literature focused on the choice of DNN architectures and hyper-parameters [KPH<sup>+</sup>19, ZBHV19]. Hence the motivations of

<sup>&</sup>lt;sup>1</sup>The code may be open-source, or at least based on an open-source implementation. See, *e.g.*, the **mbedTLS** implementations: www.mbedTLS.com, or the Agence Nationale de la Sécurité des Systèmes d'Information (ANSSI)'s protected Advanced Encryption Standard (AES): https://github.com/ANSSI-FR/SecAESSTM32.

 $<sup>^{2}</sup>$ Some previous works have proposed to inject the knowledge of additional information such as the plaintext, under the form of an extra input feature to the model, additionally to the leakage trace [HHO20]. Yet, to the best of our knowledge, no study considered how to incorporate the knowledge of the masking scheme inside the profiling model.

this work:

How can we efficiently leverage the knowledge of the masking scheme in an ML-based profiling attack, without relying on the knowledge of random nonces ?

**The Scheme-Aware Modeling.** To address this question, we investigate a new type of SCA adversary that we name *scheme-aware*. In this threat scenario, the adversary is supposed to have access to the source code of the target implementation. Concretely, this means that the adversary knows the masking scheme and order used to protect the target. Moreover, she is able to localize some Points of Interest (P.o.Is) precisely enough thanks to a careful code analysis.

We explain how these assumptions can be taken into account in a DL model. To this end, we introduce GroupRecombine, a simple neural network layer encoding the knowledge of any group-based masking scheme, under the form of a discrete convolution. Contrary to the convolutions layers used in Convolutional Neural Networks (CNNs), GroupRecombine is parameter-free, and is applied as the last layer in our model. This new layer can replace some of the upper layers of a DNN potentially carrying many learning parameters to fit, without any loss of expressiveness of the resulting architecture, thanks to the prior knowledge of the masking scheme. In addition, it can be efficiently implemented using Walsh-Hadamard (in the case of Boolean masking) or Fourier transforms (in the case of arithmetic or multiplicative masking), or a mix of both (in the case of affine masking). As a result, any model equipped with the GroupRecombine no longer requires to learn how to recombine the information gathered on each share, and may only focus on the joint learning of the leakage models of each share.

We validate our approach on simulations and on public datasets. Our experiments on the ANSSI's SCA Databases (ASCAD) emphasize some use-cases with first-order Boolean masking where very simple scheme-aware models lead to successful attacks, whereas their uninformed counter-part fails. This suggests that a significant part of the efforts spent by the SCA practitioner in an uninformed setting, *e.g.* by running huge hyper-parameter grid searches, would actually be devoted to finding a DNN architecture that efficiently captures the way to learn the masking scheme. Hence, using GroupRecombine may be seen as an efficient surrogate to this issue. As an example, we also address the challenge left by Bronchain and Standaert as a conclusion of their works at TCHES 2020, by showing on simulations how GroupRecombine could be used for profiling in presence of an affine masking scheme, without knowing the random shares to train our model.

Finally, we conclude this paper by discussing how far the scheme-aware approach could scale with an increasing masking order, by providing theoretical arguments and experimental evidence. Actually, the potential limitations of GroupRecombine that we emphasize are not restricted to our approach, and more generally cover at least any non-worst-case model trained with gradient descent, leaving open the question whether this also covers other types of profiling models in the same setting. Overall, we hope that these questions will be received as a helpful contribution to the more general debate regarding the choice of different evaluation methodologies in SCA.

## 2 Scheme-Aware Modeling and Application to Masking

In this section, we introduce the scheme-aware adversary. The idea behind this new threat model is to properly separate what can be assumed to be known by the adversary — e.g., any algorithmic and implementation aspect — from what remains unknown and therefore should be learned during the profiling phase — e.g., the device-dependent leakage model of each share. We discuss hereafter two aspects of the prior knowledge on which any scheme-aware adversary may rely, and how to leverage it.

#### 2.1 Hard-Encoding of the Discrete Convolution

Hereafter, we explain how to materialize the prior knowledge of the masking scheme into our DNN. Recall that the true model to learn may be expressed as a convolution product of elementary leakage models for each share, as stated hereafter.

**Proposition 1** ([LPR<sup>+</sup>14, Sec. 6], extended). Let  $Y_0, \ldots, Y_d \in \mathcal{Y}$  be Independent and Identically Distributed (i.i.d.) shares, uniformly drawn over the group  $(\mathcal{Y}, \star)$ . Let  $\mathbf{L} = (\mathbf{L}_0, \ldots, \mathbf{L}_d)^{\mathsf{T}}$  be a random vector denoting the leakage, and let  $\mathbf{l} = (\mathbf{l}_0, \ldots, \mathbf{l}_d)^{\mathsf{T}}$  be an observation of  $\mathbf{L}$ . Assume that any  $\mathbf{L}_i$  only depend on  $Y_i$ , i.e., any  $\mathbf{L}_i$  is independent of the  $(\mathbf{L}_j)_{j\neq i}$ . Then, the posterior Probability Mass Function (p.m.f.) of  $Y = Y_0 \star \ldots \star Y_d$ can be formulated as a discrete convolution product:

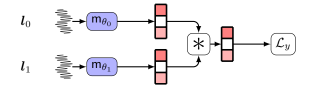
$$\mathsf{p}_{Y}(\boldsymbol{l}) = \mathsf{p}_{Y_{0}}(\boldsymbol{l}_{0}) * \dots * \mathsf{p}_{Y_{d}}(\boldsymbol{l}_{d}) \quad , \tag{1}$$

where  $\mathbf{p}_{Y_i}(\mathbf{l}_i) = \Pr(Y_i \mid \mathbf{L}_i = \mathbf{l}_i)$  denotes the conditional p.m.f. of the share  $Y_i$  given the realization  $\mathbf{l}_i$  of the leakage random vector  $\mathbf{L}_i$ .

Lomné *et al.* have given a proof of a similar result at CHES 2014, for generative models such as GTs. Proposition 1, that we prove in Appendix A, extends Lomné *et al.*'s one to discriminative models. We may leverage Proposition 1 in a scheme-aware threat model, provided that we know the inner law  $\star$  of the group  $\mathcal{Y}$ . This means that we know the discrete convolution operator in Equation 1. In other words, we no longer require to learn how to recombine the information extracted on the leakage corresponding to each share: we are reduced to jointly learn the leakage models  $l_i \mapsto \mathsf{p}_{Y_i}(l_i)$  using some corresponding estimators  $\mathsf{m}_{\theta_i}, i \in [0, d]$  respectively. Hence proposing the following model for our scheme-aware attacks:

$$\mathsf{m}_{\theta}\left(\boldsymbol{l}\right) = \mathsf{m}_{\theta_{0}}\left(\boldsymbol{l}_{0}\right) * \ldots * \mathsf{m}_{\theta_{d}}\left(\boldsymbol{l}_{d}\right) \quad . \tag{2}$$

Said more concretely, we build a model where each branch  $\mathfrak{m}_{\theta_i}$  maps its corresponding sub-leakage to a  $|\mathcal{Y}|$ -dimensional vector denoting a p.m.f. Then, all branches are combined together by computing the discrete convolution. Figure 1 depicts the idea for a first-order masking scheme: blue nodes denote branches, *i.e.* models with trainable parameters, whose goal is to modelize the conditional p.m.f.  $\mathfrak{p}_{Y_i}(l_i)$  for each share. Vectors with shades of red denote p.m.f.s over  $\mathcal{Y}$ , and the "\*" node denotes the discrete convolution with respect to the inner-law of the group  $\mathcal{Y}$ .



**Figure 1:** Scheme-aware model, with known P.o.Is for each share and a known masking scheme, but unknown random shares.

It remains to explain how the branch models can be fine-tuned so that they fit the true leakage models on each share. This can be done with Maximum Likelihood Estimation (MLE), *i.e.* by minimizing a *loss* function  $\mathcal{L}_y$  quantifying the dissimilarity between the overall output p.m.f. returned by the model  $\mathsf{m}_{\theta}(l)$  and the expected values of the target variable, that are known during profiling (for both uninformed, scheme-aware, and worst-case adversaries).

In a worst-case setting, the MLE is usually implemented by tuning of branch model separately from each other by minimizing the loss functions  $\mathcal{L}_{y_i}$  of each branch separately.

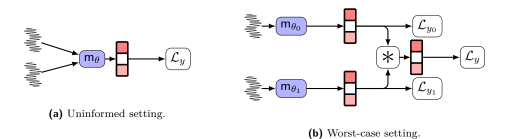


Figure 2: Other adversaries, for comparison with scheme-aware model.

Once it is done, the fine-tuned branch models are combined together with the discrete convolution. This is the approach used, *e.g.*, by Bronchain *et al.* [BS20, BDMS22]. Similarly, each branch model can compute its output p.m.f. using Gaussian Template (GT), and by converting the generative model into a discriminative one with Bayes' rule, as done somehow by Ouladj *et al.* [OGGM21]. Unfortunately, these approaches require to know the values of the shares during profiling, which we considered to be a strong assumption in real-world evaluations.

In a scheme-aware model instead, the branch models are rather *jointly* tuned by directly minimizing the overall loss function  $\mathcal{L}_y$ , averaged over a training set of traces acquired during the profiling phase. As depicted in Figure 1, computing and minimizing the overall loss function  $\mathcal{L}_y$  only requires as labels the values of the target variable during profiling, which is allowed by definition of our scheme-aware adversary.

Therefore, the scheme-aware model depicted in Figure 1 relaxes the strong assumption of random nonce knowledge during the profiling phase, but still encodes the knowledge of the masking scheme, which is no longer needed to implicitly learn from the data, contrary to the uninformed model depicted in Figure 2a. This approach sounds somewhat sub-optimal if the scheme is already known. Hence, by introducing the scheme-aware adversary that is stronger than the uninformed adversary, but weaker than the worst-case one, we expect to get a closer emulation of the actual powers of a real-world adversary.

#### 2.2 Localization of P.o.Is

In Subsection 2.1, we implicitly assumed for adversaries stronger-than-uninformed to know how to localize the P.o.Is for each share in the traces, in order to properly separate the leakages. We discuss this assumption in this section.

Usually, the P.o.I selection is done by computing some first-order statistics, such as T-tests or Signal-to-Noise Ratios (SNRs). Without knowledge of the random nonces, these tools cannot identify the right time samples, since by definition of masking, any univariate sample should be independent of the target variable Y.<sup>3</sup> This means that any non-worst-case adversary cannot identify the P.o.Is thanks to statistical tools in presence of masking.

Nevertheless, a P.o.I selection remains possible without access to the random nonces, thanks to a visual analysis of the traces, combined with a careful study of the T.O.E. source code. Indeed, a software implementation of a cryptographic primitive is typically made of (nested) loops whose number of iterations are publicly known, in virtue of the Kerckhoffs principle. This induces some sequences of (nested) patterns in the traces that can be visually identified on the raw measurements by the adversary. Moreover, this analysis can even be refined by counting the number of clock cycles for each executed instructions, and combining them with the clock and sampling frequencies in order to guess at which time sample each instruction should leak. As a consequence, it is still

<sup>&</sup>lt;sup>3</sup>Ignoring glitches or transition leakages.

possible to localize the leakage on each share, and the P.o.I selection through T-test or SNR should actually be seen as a useful but non-necessary shortcut for the evaluator to spare some time. The recent literature provides two examples of this approach. First, Masure and Strullu reported a detailed code analysis of the assembly code of the ANSSI's secure software implementation of the AES on an ARM Cortex M4, in order to extract 15,000 P.o.Is out of 1 million time samples in the raw traces, covering the leakages of the three shares used by the affine masking scheme [MS21]. Second, Egger *et al.* verified that the CPOI leakage detection method [DS16] could localize the same time windows as with the analysis of the assembly code used in the ANSSI's SCA Databases (ASCAD)-v1 dataset [EST<sup>+</sup>22, Fig. 4].

## 3 Scheme-Aware Modeling with DNNs

Now we have introduced the scheme-aware adversary in the case of masking and explained the intuition behind its advantages, we discuss in this section how to concretely implement it with Deep Neural Networks (DNNs). First, we discuss in Subsection 3.1 how to concretely minimize the overall loss function with scheme-aware models, by introducing a new DNN layer called GroupRecombine. Then, we explain in Subsection 3.2 how our approach can be extended to many types of masking schemes. Finally, we argue in Subsection 3.3 why we implemented our own version of GroupRecombine, instead of relying on native building blocks of most DL frameworks.

#### 3.1 Implementing the Backward Propagation

To optimize a function based on DNNs, the most widely used approach is to use Gradient Descent (GD)-based optimizers. To this end, we need to compute the derivatives of our model when using a recombination layer. These derivatives are computed with the backward propagation algorithm [BPRS17], who leverages the chaining rule to reduce the computation of the derivatives for a composed function to the computation of derivatives for each elementary function. Our models being made of regular DNN layers for which the backward propagation is already hard-coded, we are then reduced to specify how to back-propagate the gradient through the discrete convolution only. We do this by implementing GroupRecombine, a parameter-free DNN layer consisting in the discrete convolution, augmented with backward propagation.

We briefly explain hereafter how the backward propagation can be computed in GroupRecombine. Thanks to the convolution theorem, the discrete convolution layer can itself be seen as a composition of a fast transform (and its inverse) and an element-wise product of d + 1 vectors. Using the chaining rule, we are again reduced to compute the backward propagation for each mapping in the composition. The (inverse) fast transform is a linear mapping, so its differential coincides with the mapping itself. In other words, the backward pass of the fast transform is the same as its forward pass. Regarding the element-wise product, it is a multi-linear mapping whose backward pass is already hard-coded in the DL frameworks such as Tensorflow or Pytorch. By putting things together, we obtain the backward pass through our GroupRecombine.

*Remark* 1. The backward pass of the GroupRecombine layer can also be directly hard-coded without decomposing with fast transforms. Interestingly, this approach coincides with implementing the update rule "from functions to variables" in the Belief - Propagation (BP) algorithm [BS21, Eq. (4)].

#### 3.2 Handling other Types of Masking

GroupRecombine works for any type of group-based masking scheme, *e.g.* Boolean [CJRR99, GP99], arithmetical [CG00], or multiplicative masking [von01]. In the latter case, one should recall that for any finite field  $(\mathcal{Y}, \oplus, \times)$ , the group  $(\mathcal{Y}, \times)$  is in bijection with  $(\mathbb{Z}_{|\mathcal{Y}|-1}, +)$ . In other words, the GroupRecombine for multiplicative masking can be implemented by using the GroupRecombine for arithmetical masking, and to permuting the entries of input and output vectors using discrete log / alog tables. As a result, it becomes also straightforward to handle less common types of masking, such as affine [FMPR11], by combining several types of GroupRecombine for different masking schemes. We will apply GroupRecombine to affine masking in Subsection 4.2.

Although we did not test it yet, extending GroupRecombine to inner-product masking schemes [BFG<sup>+</sup>17, BFG15] should be feasible as well. Indeed, inner-product masking derives from Boolean masking by applying a public linear mapping, that could be handled by hard-coding the corresponding permutation of the entries in the input and output p.m.f.s, similarly to the transformation from arithmetical to multiplicative masking.

#### 3.3 Using Native Convolutions in DL Frameworks?

We implemented our own version of the discrete convolution used in GroupRecombine. Since discrete convolutions are widely used in DL, one might wonder why not using such layers natively implemented in the main frameworks such as Tensorflow or Pytorch. There are two main reasons for that.

First, for Boolean masking, the discrete convolution is not natively implemented in DL frameworks like Pytorch [PGM<sup>+</sup>19], or Tensorflow [AAB<sup>+</sup>15]. Even for arithmetical masking the discrete convolution must be circular, whereas the convolution layers used in DL frameworks are usually not circular and use zero-padding to deal with side effects.

Second, even if circular padding were used, the convolution layers proposed in the DL frameworks rely on a naive computation of the convolution product — *i.e.* not based on fast transforms. The reason is that in computer vision-based DL, the filter size W is often too small for the computation with fast transform — with complexity  $\mathcal{O}(W \cdot \log_2(W))$  — to be significantly more efficient that the naive approach of complexity  $\mathcal{O}(W^2)$  [VJM<sup>+</sup>15].<sup>4</sup> In our context where the convolutions are often computed over  $W = 2^n$  classes, where n is the bit-size of the target, using a non-naive approach becomes more efficient than the naive one.

That is why we implement GroupRecombine using fast transforms. For Boolean masking, we use the Walsh-Hadamard (WH) transform, whereas for arithmetical masking, we use the Fast Fourier Transform (FFT). Both WH and FFT can be implemented with Pytorch on (General Purpose) Graphic Processing Unit (GPU) with a CUDA backend: the latter one is natively implemented in the framework, while for the former one, we leverage the implementation developed by Thomas *et al.* [TGD<sup>+</sup>18]. Overall, our GroupRecombine layer results in a parameter-free layer that can be easily integrated into the Pytorch framework.<sup>5</sup>

## 4 Analyzing Performances of GroupRecombine

Now we introduced our GroupRecombine, we would like to compare its performances with uninformed and worst-case settings. In this section, we show the advantages of using

 $<sup>^{4}</sup>$ Computer vision applications of DL usually prefer filter sizes lower than 4, as it maximizes the ratio between the size of receptive fields of the convolution layers and the number of parameters to train [SZ15]. This rational only holds for 2D (images) or 3D (videos) data, and no longer holds for 1D time series like SCA traces.

 $<sup>{}^{5}</sup>$ We illustrate this claim with some code snippet in the supplementary material of this submission. Although we did not test, we expect the implementation to be as straightforward in Tensorflow or Keras as it is in Pytorch.

GroupRecombine, both on simulated experiments and on real experimental data. We first describe the settings of our experiments in Subsection 4.1. Then, we report and discuss results on simulations in Subsection 4.2, and on experiments in Subsection 4.3.

#### 4.1 Settings for Comparison

For a fair comparison, we would like to show that all other things being equal, using GroupRecombine leads to better performance. This requires to properly define the types of adversaries against which we test GroupRecombine, and how to assess the comparison between each model.

#### 4.1.1 Spectrum of Adversaries under Test

To this end, we describe hereafter the different adversaries that we consider.

- Worst-case. This model is depicted in Figure 2b. Each branch model  $\mathfrak{m}_{\theta_i}$  is learned independently from each other, based on a restricted amount of corresponding P.o.Is, and by minimizing the loss function  $\mathcal{L}_{y_i}$  based on the corresponding share  $y_i$ . Concretely, each branch model is instantiated with a one-hidden-layer Multi-Layer Perceptron (MLP) with N = 1,000 neurons, a Rectified Linear Unit (ReLU) activation function on the hidden layer, and a softmax activation function on the output layer. Following the standard practice in DL, Batch Normalization (BN) is also applied before ReLU. Once all branch models are trained, they are passed through GroupRecombine to infer on the validation traces.
- Scheme-aware. It is the same as the worst-case setting, but the trainings of the branch models are done *jointly*, using the loss function  $\mathcal{L}_y$  computed from the labels of the target y, as the random nonces are no longer known. To better evidence the advantages of our scheme-aware model, we decline three different versions:
  - Known P.o.Is and scheme (SA). This corresponds to the model depicted in Figure 1. Each branch model is only fed with the appropriate P.o.Is, and the recombination is done with GroupRecombine.
  - Known masking scheme only (SA \ P.o.Is). This corresponds to the model depicted in Figure 3b. This is the same model as the previous one, except that each branch model is fed with raw traces, instead of separate P.o.Is.
  - Known P.o.Is only (SA  $\setminus$  Enc). This corresponds to the model depicted in Figure 3a. The model is the same as with the one with known P.o.Is and masking scheme, except that the GroupRecombine layer is replaced by another one-hidden-layer MLP with N' = 100 neurons.

The two latter versions are downgraded compared to the former one. Therefore, if our scheme-aware model is sound, it is expected to work better than its downgraded versions.

• Uninformed setting. This model is depicted in Figure 2a. Since the adversary is not assume to know neither the P.o.I location, nor the underlying masking scheme, the uninformed setting is a one-hidden-layer MLP that is fed with raw traces. For consistency with worst-case and scheme-aware models with known P.o.Is and scheme, we keep the number of hidden neurons constant. Therefore, our MLP in the uninformed setting has  $(d + 1) \cdot N$  neurons.

Note that in GroupRecombine, the approximation error is null. This means that provided that each branch model  $\mathbf{m}_{\theta_i}$  can exactly computes the true leakage model  $\mathbf{p}_{\mathbf{Y}_i}(\boldsymbol{l}_i)$ , then the whole scheme-aware model using GroupRecombine exactly implements the true conditional

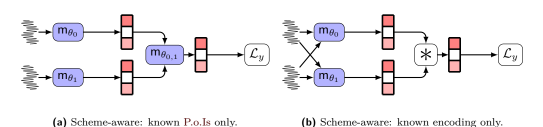


Figure 3: "Downgraded" scheme-aware models. In blue: blocks with trainable parameters.

p.m.f. of the target variable Y. On the contrary, since the discrete convolution is a non-linear mapping, it cannot be exactly instantiated by a MLP with ReLU activation function [Yar17, Thm. 6]. Nevertheless, the simulations of Masure *et al.* at CHES 2020 suggest that the approximation error can be made negligible with an architecture identical to the one considered here for models in the uninformed setting [MDP19a]. Hence, the comparison between uninformed models and the other ones remains fair. Following the empirical study of Perin and Picek [PP20], we train all our models by minimizing the Negative Log Likelihood (NLL) loss function, by using the Adaptive Moment Estimation (Adam) optimizer [KB15], with a learning rate of  $10^{-4}$ .

#### 4.1.2 Performance Metrics and Quantifying Complexity

To assess the quality of a model, we measure the Perceived Information (PI), as it has been shown to be strongly related to the minimum number of traces required to succeed a key recovery with a Maximum Likelihood Distinguisher (MLD) [MDP19a].<sup>6</sup>

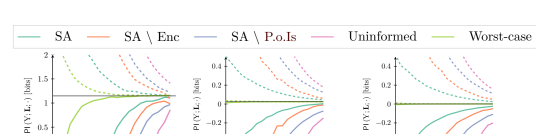
Based on this, we can assess the *profiling* complexity. It is measured in terms of the amount of profiling traces needed to reach the optimal value of PI. To this end, we plot the *learning curves* depicting the evolution of the PI with respect to the number  $N_p$  of profiling traces used to train the model. More precisely, for each value of  $N_p$  and for each trained model we keep the model at the epoch when the training loss is minimal. The dashed curve corresponds to the PI computed over the training set, whereas the plain curve corresponds to the PI computed on a validation set. This is equivalent to assessing the model's quality when it is not combined with any other regularization technique, in order to assess the sole effect of GroupRecombine. Since an adversary is likely to combine the use of GroupRecombine with the use of a validation loss, we also plot a modified type of learning curve, where we keep the trained model at the epoch when the validation loss — instead of the training loss — is minimal.

Remark 2. For consistency when the number  $N_p$  increases, we use optimizers with full-batch, therefore one optimization step always equals one epoch.

#### 4.2 Results on Simulation

In this subsection, we present the results of our simulations. We first describe hereafter the simulation framework. For each trace, a (d + 1)-sharing  $(Y_0, \ldots, Y_d)$  is drawn uniformly from  $\mathcal{Y}^{d+1}$ , where  $\mathcal{Y} = \llbracket 0, 255 \rrbracket$ . Then, each sub-leakage is drawn as  $\boldsymbol{l}_i = \operatorname{hw}(Y_i) + B$ , where hw stands for the function mapping a binary variable to its Hamming weight, and where B is a Gaussian noise with standard deviation of  $\sigma$ . Meanwhile, the label Y is computed as  $Y_0 \star \ldots \star Y_d$ . Unless in Subsubsection 4.2.3, we consider Boolean masking, *i.e.*  $\star$  is the bit-wise addition  $\oplus$ .

<sup>&</sup>lt;sup>6</sup>We also have the following relationship with the NLL loss function  $\mathcal{L}(\mathbf{m}_{\theta})$  used for training: PI (Y; **L**;  $\mathbf{m}_{\theta}$ ) =  $n - \mathcal{L}(\mathbf{m}_{\theta})$ , where *n* stands for the bit-size of the target variable Y.



105

 $N_{i}$ 

-0.4

10

10

 $10^{6}$ 

10

 $N_{t}$ 

-0.4

10

 $N_p$ (b) W/o validation, SNR = 1. (c) W/o validation, SNR = 0.1. (a) W/o validation, SNR = 200. ·10<sup>-</sup> 10-1.53  $\mathsf{Pl}\left(\mathbf{Y}; \mathbf{L}; \cdot\right)$  [bits] 2  $PI(Y; L; \cdot)$  [bits]  $\mathsf{Pl}\left(Y;\mathbf{L};\cdot\right)\left[\mathrm{bits}\right]$ ſ 0.50 0  $10^{3}$  $10^{5}$  $10^{4}$  $10^{4}$ 10 10  $10^{4}$ 105 10  $N_r$  $N_{\tau}$  $N_r$ 

(d) With validation, SNR = 200. (e) With validation, SNR = 1. (f) With validation, SNR = 0.1.

Figure 4: Learning curves for models against a first-order Boolean masking.

Since the leakage model is known in our simulated framework, we can also compute a Monte-Carlo (MC) estimation of the Mutual Information (MI), using the true Probability Density Function (p.d.f.) used to sample the simulated traces. To this end,  $N_v = 200,000$  validation traces are simulated, leading to an unbiased estimation error of roughly  $\frac{1}{\sqrt{N}}$ .

#### 4.2.1 First-Order Boolean Masking

0

 $10^{3}$ 

 $10^{4}$ 

 $10^{5}$ 

The results for two 8-bit Boolean shares are depicted on Figure 4. We can see on Figure 4a, that the light green curves (depicting the worst-case model) enjoy the fastest convergence towards the black horizontal line denoting the MI, whereas the pink curves (depicting the model in the uninformed setting) suffer from the slowest convergence. Between those curves, the dark green, orange and blue curves denoting the different variants of scheme-aware models enjoy convergence at an intermediate speed. The same sketch can also be observed on Figure 4b and Figure 4c. Moreover, the same observation can be made when assuming that the adversary has a validation set of traces in order to measure the PI in a non-biased way, according to Figures 4d, 4e, 4f. This can be interpreted as the fact that, regardless of the noise level, scheme-aware models enjoy a higher profiling complexity than that of worst-case models, but still lower than that of models in the uninformed setting.

#### 4.2.2 Second-Order Boolean Masking

We push our simulated experiments one step forward, adding a third Boolean share into the leakage. The results are shown in Figure 5. When analyzing the learning curve in Figure 5a, we may notice two main differences compared to the 2 share simulation. First, the pink, blue and orange curves have been shifted to the right, meaning that their corresponding profiling complexity has been increased by an order of magnitude. Interestingly, the green curve denoting the best scheme-aware model has merely been shifted, meaning that

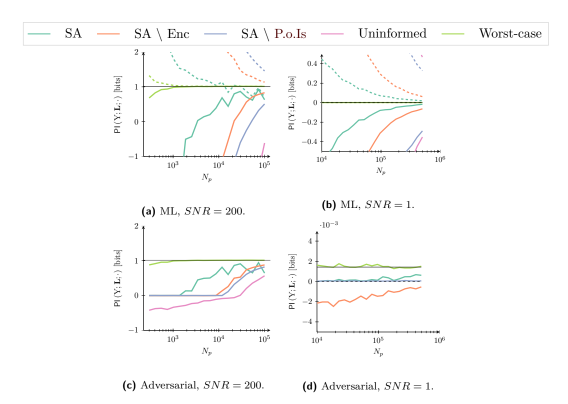


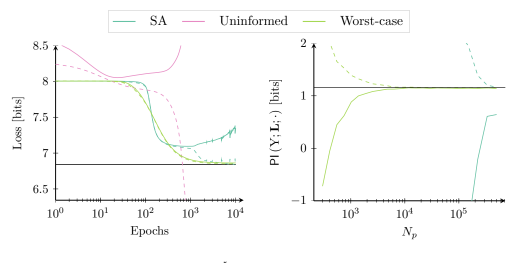
Figure 5: Learning curves for models against a second-order Boolean masking.

its profiling complexity did not change much compared when adding a third share into the experiment. Nevertheless, as a second observation, it is noticeable that the green learning curve seems less *smooth* than in the previous experiment. This may testify an increasing *optimization* complexity, *i.e.* the fact that the minimizer encounters difficulties to decrease the training loss. We will thoroughly discuss this phenomenon in Section 5. But more generally, all our simulations so far show that the downgraded scheme-aware models (depicted by blue and orange curves) perform less than the full scheme-aware (denoted by dark green curves). This confirms the soundness of our approach, as the good performance does not come from a side-effect. Hence, in the remaining of the paper, we only focus on non-downgraded models for the scheme-aware adversary.

#### 4.2.3 Affine Masking

We then move our simulation from a second order Boolean masking scheme to an affine scheme. Hereupon, Bronchain and Standaert argued that learning an affine scheme with an uninformed model turns out to be hard, as emphasized by their simulations [BS20]. There, the authors considered a slightly different leakage model for the multiplicative share  $\alpha$  of the affine sharing.<sup>7</sup> Indeed, from their experimental measurements, they were able to recover the multiplicative share with almost 100% accuracy with their worst-case attack, meaning that the leakage model of the multiplicative share  $\alpha$  was injective. This can be explained by the fact that the concrete implementation of all affine schemes known in the literature are table-based [FMPR11, MS21], meaning that the values  $x \times \alpha$  are sequentially processed for  $x \in [1, 2^n - 1]$  during the pre-computation phase, which may leak a lot [TWO14].

 $<sup>^{7}</sup>$ For completeness, we also provide in supplementary material the results of our simulation without changing the leakage model. The conclusions remain the same.



(a) Optimization curves for  $N_p = 5 \cdot 10^5$ . (b) Learning curves.

Figure 6: Comparison between models for an affine masking.

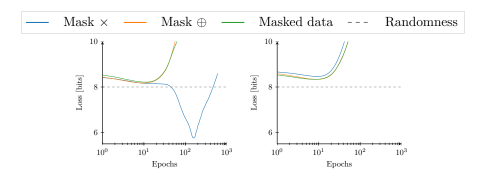
To take this into account in their subsequent simulations, Bronchain and Standaert emulated the leakage of the multiplicative share  $\alpha$  with an identity model. Nevertheless, Cristiani et al. experimentally showed that learning such a leakage model with DNNs could be hard [CLM20, Fig. 3], whereas replacing the identity by a one-hot encoding could make the problem much easier. A similar experiment on images, conducted under the *coordinate* transform problem terminology, led to similar conclusions [LLM<sup>+</sup>18]. This suggests that beside not being physically realistic, the identity leakage model could make the problem artificially much harder. That is why we revisit Bronchain and Standaert's experiment by changing the leakage model regarding the multiplicative share. Hereafter, additionally to the leakages on the other shares, we consider that the adversary has access to the values of  $hw(x \times \alpha)$ , for  $x \in [1, 2^n - 1]$ , where hw denotes the Hamming weight leakage model.<sup>8</sup> The simulation results are presented in Figure 6. While our model in the uninformed setting is not able to get a positive PI, as depicted by the pink curve on Figure 6a, we can see that our scheme-aware model leveraging the knowledge of both P.o.Is and masking scheme is able to get a validation loss below the 8-bit random threshold, denoting an effective model. Even though it does not contradict the previous conclusions of Bronchain and Standaert regarding the efficiency of models in the uninformed setting, the outcomes of our scheme-aware model show that not having access to the random nonces does not necessarily lead to an unsuccessful attack.

#### 4.2.4 Does Each Branch Actually Learn True Leakage Distributions?

In view of the good results obtained by our GroupRecombine in our simulations, we may wonder whether the intermediate output p.m.f.s returned by each branch in a scheme-aware attack — in the known-P.o.Is setting — were also good estimates of the true p.m.f. of each share. Can we compare the performance of our scheme-aware attacks against a worst-case adversary on each share separately?

At first sight, there is no reason why it would be possible, since the mapping  $(\mathbf{p}, \mathbf{p}') \mapsto \mathbf{p} * \mathbf{p}'$  is not invertible. As an example, if  $\tau_h$  denotes the translation operator, *i.e.*,  $\tau_h(\mathbf{p}) =$ 

<sup>&</sup>lt;sup>8</sup>Note that this leakage model seems to behave similarly to a one-hot encoding, as it is possible to guess the value of  $\alpha$  by looking at the entry x of the leakage for which the leakage equal n. This would denote the value x such that  $x \times \alpha = 2^n - 1$ . Therefore, this leakage model somehow acts like a one-hot encoding, up to a permutation of the encoding entries.



(a) With injective leakage model (b) With Hamming weight leakage for  $\alpha$ .

Figure 7: Branch losses.

 $p[\cdot \star h]$ , then the convolution product is known to be *co-variant* with translation, *i.e.*,  $\tau_h(\mathbf{p}) * \mathbf{p}' = \tau_h(\mathbf{p} * \mathbf{p}')$ . As a corollary, we have  $\tau_h(\mathbf{p}) * \tau_{h^{-1}}(\mathbf{p}') = \tau_{h\star h^{-1}}(\mathbf{p} * \mathbf{p}') = \mathbf{p} * \mathbf{p}'$ . In other words, even if our scheme-aware model could reach the optimal performance, the leakage models on each share could at best be learned up to a shift of the probabilities.

But what about with affine masking? Here the additive and multiplicative shares do not play the exact same role, so the argument about translation covariance no longer holds. Does this mean that some branches could actually learn the true leakage model on their respective share? To clarify this question, we also monitored the loss function on each branch output against the labels of each share, during training. These branch losses were computed over the validation set. We plot on Figure 7 such metrics, monitored during the simulation in affine masking of Subsubsection 4.2.3. We notice in Figure 7a that the loss for the branches of additive shares in orange and green diverge, as expected earlier. But surprisingly, the loss on the multiplicative branch in blue goes below the 8-bit randomness threshold before starting over-fitting. This means that the multiplicative branch may be used in this case to infer on the value of the multiplicative share, despite not having known at all the values of the multiplicative share during training. Interestingly, we also note that the branch loss in blue escapes its plateau after 50 epochs, whereas at the same epoch in Figure 6a, the overall training and validation losses for the scheme-aware model are still stuck on the plateau. This denotes that at this epoch, the learning has somehow started, even if this does not reflect in the value of the overall loss.

Actually, our observations should be slightly mitigated, as they seem to be leakagemodel-dependent. Indeed, we replicated our simulation, by replacing our injective leakage model for the multiplicative share by a simpler (non-injective) Hamming weight leakage model. The corresponding results shown on Figure 7b indicates this time that the multiplicative branch is not able to reach a positive PI, so it cannot be used to infer on the multiplicative share in this case. Still, we argued in Subsubsection 4.2.3 that the latter leakage model is less representative than the former one.

### 4.3 Application on Experimental Data

Now we established the interest of GroupRecombine on simulations, we would like to test it under experimental traces. To this end, we replicate the same experiments as with the simulation described in Subsection 4.2 on some public datasets using masking.

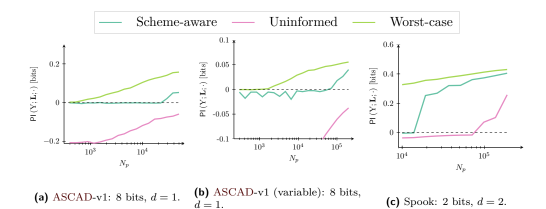


Figure 8: Learning curves (best validation loss).

#### 4.3.1 Experiments on ASCAD-v1

We start with the ASCAD-v1 dataset, published in 2018 by Benadjila *et al.* [BPS<sup>+</sup>20]. It deals with a first-order masked implementation of AES, with a Boolean scheme based on table re-computation. The cryptographic primitive is implemented on an 8-bit ATMega8515 device on which the Electro-Magnetic (EM) field emanations are measured. Two versions of the dataset are proposed: one so-called *fixed* with measurements acquired on a 700 time samples window using a fixed encryption key, and a *variable* dataset with measurements on a 1,400 time samples window using a variable encryption key for profiling traces. Both windows cover the look-up of the re-computed Sbox when applying the SubBytes operation on the third byte of the AES state during the first round. Since on both datasets, the data dimensionality is much higher than in our simulations whereas the number of profiling traces remains of same order of magnitude as in our simulations, the DNNs used in our simulations are more likely to over-fit. That is why we reduce the number of neurons in the hidden layer of the branches from 1,000 to 100.

**Results on Fixed Dataset.** We report hereafter the outcomes of our trainings on the fixed dataset. When the threat model assumes to know the P.o.I location, the P.o.I selection has been done by splitting the 700 time samples into two halves, the first 350 time samples containing some leakages about the masked share while the second 350 time samples containing leakages about the mask. This pre-processing is suboptimal compared to a P.o.I selection with SNR, but reflects more what an adversary can do with a visual trace analysis with the help of the source code, and can even be further refined with a thorough code analysis, as argued in Subsection 2.2. The results are depicted on Figure 8a. First, it can be seen that when using the full profiling trace set — *i.e.*,  $N_p = 50,000$  — the validation loss eventually diverges, meaning that when using only shallow MLPs like in our experiments, none of the different threat models would lead to an effective attack without further pre-processing. Nevertheless, we can see that the validation losses in light green, green and blue have their minimum value below the 8-bit threshold, meaning that selecting the best model based on a validation loss would eventually lead to successful attacks.

Moreover, we can still observe the same hierarchy between the threat models as in the simulations conducted in Subsubsection 4.2.1. The model in the worst-case setting leads to a PI close to 0.2 bit, which is the highest lower bound of MI reported in the literature on ASCAD-v1 [CLM20]. Then, the scheme-aware model leveraging both P.o.I location and knowledge of the masking scheme reaches a PI of 0.05 bits, whereas the scheme-aware model exploiting the knowledge of the masking scheme only obtains a PI of 0.02 bits. Finally, the scheme-aware model leveraging the P.o.Is location only and the model in the

uninformed setting cannot get a positive PI during the whole training.

We validate this observations by reproducing the trainings on a lower number of profiling traces. For each of these trainings, the best PI on the validation loss is reported on Figure 8a. As can be noticed, the previous observations regarding the hierarchy of the threat models remains true.

**Results on Variable Dataset.** We repeated the same experiments on the variable dataset. We first tried by splitting the traces into two contiguous parts of 700 points each, as with the fixed dataset. Unfortunately, none of the non-worst-case models could succeed in getting a positive PI, which suggests that our P.o.I selection method for our scheme-aware attacks was not sufficient, at least for the amount of traces available in this public dataset. Therefore, we refined the P.o.I selection, by narrowing the windows for the two shares. For the share  $r_{out}$ , we selected the range [0, 300], whereas for the share  $Y \oplus r_{out}$ , we selected the range  $[v_{0,300}]$ , whereas for the share  $Y \oplus r_{out}$ , we selected the range  $[v_{0,300}]$ . This refined P.o.I selection is possible even without knowledge of  $r_{out}$  during profiling, thanks to a careful assembly code analysis, similar to the one recently conducted by Egger *et al.* [EST<sup>+</sup>22, Fig. 4]. For consistency in our comparisons, we also feed the model in the uninformed setting with the two restricted P.o.I windows, stacked together.

We then report on Figure 8b the results obtained for this second attempt on the variable dataset. Like with the fixed dataset, we can see that the model uninformed is unable to get a positive PI with the 600 P.o.Is given as an input. On the contrary, it took 70,000 profiling traces to get a positive PI for our best scheme-aware model (green curve in Figure 8b). This is much more than for the worst-case model that only required 2,000 profiling traces (light green curve). Still, the results obtained on both ASCAD-v1 datasets confirm that it is possible to succeed an attack by using small MLPs, provided that the recombination is cleverly done, *e.g.* with GroupRecombine.

#### 4.3.2 Experiment for Second-Order Masking

Provided with the promising results presented in Figure 4 and Figure 6, and on the good experimental verifications on the ASCAD-v1 datasets, we pushed our experiments one step further by trying to extend our attacks to higher-order masking. To this end, we report positive results on a second-order Boolean masking, and negative results on an affine masking.

**Results on Clyde.** We first report our results obtained on a second-order Boolean masking. To this end, we considered the CHES 2020 CTF dataset.<sup>9</sup> More precisely, we considered the traces depicting the software implementation of Clyde protected with a 3-sharing. Clyde is a bit-slice SPN cipher, whose state is made of four 32-bit words, with a 4-bit Sbox. The authors of the CTF provided the traces with a baseline attack recovering an 8-bit secret chunk, by targeting two bits in each of the secret words. We propose hereafter to replicate one the 2-bit key recovery with GroupRecombine. To this end, we target the  $15^{th}$  and  $16^{th}$  bits of the first word. The dataset is made of 200,000 traces from which we use 190,000 of them for training and the remaining 10,000 for validation. Using the training traces, we first compute the SNR of each share. Then, we keep some contiguous windows around the three main peaks of SNR for each share. As argued in Subsection 2.2, we assume that an adversary without access to the random shares during profiling could have been able to select the same P.o.Is, thanks to a joint analysis of the traces and the source code of the implementation.<sup>11</sup> Then, these P.o.Is are fed to GroupRecombine, using the

<sup>&</sup>lt;sup>9</sup>https://ctf.spook.dev/.

 $<sup>^{10}</sup>$ The highest peak of SNR is around 0.3.

<sup>&</sup>lt;sup>11</sup>The source code is available at https://git-crypto.elen.ucl.ac.be/spook/masked\_spook\_sw.

same architecture for the branches as the ones used in our experiments on ASCAD-v1. The results are reported in Figure 8c. It shows that the scheme-aware model is able to get a positive PI with less than 20,000 profiling traces, whereas the uninformed adversary requires around 70,000 profiling traces. In other words, a scheme-aware adversary can spare some profiling complexity.

Attempt on ASCAD-v2. Provided with the promising results presented in Figure 6 on the affine masking, and on the good experimental verifications on the ASCAD-v1 datasets, we pushed our experiments one step further by trying to replicate the attack of affine masking on an actual implementation of the affine masking. To this end, we used the ASCAD-v2 dataset [MS21]. It is made of 500,000 traces, each having 15,000 time samples coming from two contiguous parts of the raw power consumption traces acquired on an STM32 Cortex-M3 device. There, the authors explain that the first window covers P.o.Is of the multiplicative share only, whereas the second window covers P.o.Is of the additive share and the masked data. Since the implementation also uses shuffling to protect the sensitive data, we artificially deactivate the latter counter-measure, by relabeling the masked data thanks to the knowledge of the random seeds used for permutation.

Unfortunately, we could not get effective attacks using scheme-aware models with simple MLPs as branch models, whereas the same model trained in a worst-case scenario could get a positive PI. This negative result should nevertheless been mitigated. Indeed, the authors of [MS21] reported some successful worst-case attacks on the dataset, leveraging the knowledge of at least one share during profiling, but did not succeed in attacking the dataset with a model in the uninformed setting. To the best of our knowledge, no successful attack has ever been reported using non-worst-case models, since the release of the dataset in early 2021. We will discuss in Section 5 the potential reasons behind this difficulty.

## 5 Discussion

We have seen that using scheme-aware adversaries could mitigate some drawbacks of uninformed adversaries. In this section, we discuss some parts of our results. Subsection 5.1 argues that changing the type of DNN architecture in the branches of a scheme-aware model should not affect the comparative advantage of GroupRecombine with respect to the uninformed approach. Finally, Subsection 5.2 questions to what extent non-worst-case approaches could efficiently work against higher-order masking schemes.

#### 5.1 On the Choice of Architecture for the Branches

In our experiments involving scheme-aware attacks, we used the same architecture for the branches of our GroupRecombine model, namely a one-hidden-layer MLP with 100 or 1,000 neurons. Naturally, better performances could have been obtained by investigating other types of DL architectures. As a consequence, our models in the uninformed setting are not necessarily the best ones. Actually, our results report unsuccessful uninformed attacks with one-hidden-layer MLPs on ASCAD, whereas the literature reports much better results on this dataset, by using deeper MLPs — up to 6 layers according to Benadjila *et al.* [BPS<sup>+</sup>20] — or CNNs [KPH<sup>+</sup>19, MDP19b].

Therefore, one might wonder whether our comparison is biased towards GroupRecombine. Hereupon, we stress that all trainable models depicted in Figure 1 and in Figure 3 have been instantiated with the simplest DNN architecture one may use. The fact that the worst-case models instantiated with such simple branches reached the optimal performances on our simulations, and reached levels of PI close to the state of the art on the ASCAD-v1 dataset. Naturally, it may be possible to use other branch models, such as CNNs, with **GroupRecombine**. But our experiments suggest that using shallow MLPs is often sufficient for optimal leakage modeling — at least provided that other types of counter-measures are ignored. In other words, this suggests that the main efforts spent by the DL practitioner in designing more complex architectures in an uninformed setting would actually serve at learning how to recombine the information gathered by the first layers of the DNN on each share, according to the masking scheme. Hence, by hard-encoding the masking scheme with **GroupRecombine**, we expect the DL practitioner to spare a significant amount of time spent, *e.g.*, in running exhaustive/random search of hyper-parameters, which is acknowledged to be the bottleneck task in DL-based SCA [BPS<sup>+</sup>20, RWPP21, WPP20].

Likewise, no regularization technique — e.g., weight decay, dropout — have been considered in this study, so adding them could have naturally improved the results. However, we argue that the effect of regularization techniques is *orthogonal* to the effect of using **GroupRecombine**. Indeed, beside being hyper-parameter-free contrary to other types of regularization, our recombination layer does not act on the bias-variance tradeoff, as most of regularizers do [SB14, Chap. 5]. This means that, provided that the assumed masking scheme is the right one, the regularization effect of the recombination layer never degrades the approximation capacity of our model as argued at the end of Subsubsection 4.1.1, contrary to what all other types of regularization are likely to do.

#### 5.2 The Initial Plateau: An Effect of Masking

Although not noticeable from the learning curves, it turned out that in all our experiments and simulations presented so far, the optimization curves -i.e. denoting the evolution of the loss function through the epochs — depicted an initial *plateau* for both training and validation loss. An example of such a plateau is shown in Figure 6a. Namely, when targeting some leakage induced by masking, the Gradient Descent (GD)-based optimization algorithm starts its procedure being stuck on a plateau whose level coincides with full randomness. This plateau is not a simulation artifact, as it can be observed in many other studies investigating uninformed adversaries against masking. See, e.g., the works of Timon [Tim19, Fig. 5], Perin and Picek [PP20, Figs. 5b, 6b, 7, 8, 13], Cristiani et al. [CLM20, Fig. 8], [CLM21, Fig. 3], and Lu et al. [LZC<sup>+</sup>21, Fig. 10 – 11, left]. Moreover, some of these figures suggest that the higher the masking order, the longer the initial plateau. As a more recent example, Gohr et al. also emphasized, in a similar context, some impact of the masking order on the performance of trained models [GLS22, Fig. 6]. Interestingly, the initial plateau barely happens when targeting unprotected implementations, or even leakages protected by shuffling [MDP19a, Fig. 2, right] or de-synchronization [CDP17, Fig. 8], [KPH<sup>+</sup>19, Fig. 10], [MBC<sup>+</sup>20, Fig. 6], suggesting this plateau is closely linked to the use of masking. These intriguing observations call for further explanation and verification: is this plateau really due to masking, and if so to what extent it affects the optimization?

#### 5.2.1 Empirical Verification on Exhaustive Datasets

To address these questions, we repeat our simulations for a Boolean masking, by using here a noise-free, *exhaustive* dataset, *i.e.* for which the training and validation loss are equal — which is made possible thanks to the discrete nature of our noise-free leakage model. Thus, the profiling complexity is nullified, allowing to focus only on the optimization complexity.<sup>12</sup> The setting of this simulation is voluntarily much simpler rather than realistic, so that the optimization complexity may be seen as a lower bound of what a real-world adversary could expect.

 $<sup>^{12}</sup>$ This setting therefore emulates the common assumption of an adversary with *unlimited* profiling power. The interested reader may find in Appendix B a discussion about how to efficiently emulate this case.

In order to quantitatively measure the optimization complexity, we define the *weak learning* threshold, set to  $\mathcal{L}(\theta) = n - \epsilon$ . The weak learning threshold corresponds to an adversary with an *effective* model, *i.e.* a model wit strictly positive PI, up to an  $\epsilon$ -margin. More concretely, the weak learning threshold can be used to measure the length of the plateau in the optimization curve. We also define the *strong learning* threshold, set to  $\mathcal{L}(\theta) = \mathsf{MI}(Y; \mathbf{L}) + \epsilon$ . The strong learning threshold corresponds to an optimal adversary from an information theoretic point of view, up to an  $\epsilon$ -margin. Hence, the optimization complexity can be measured in terms of the number of epochs required to reach the weak and strong learning thresholds.

The results of our simulations with an exhaustive dataset are showed on Figure 9, for  $\epsilon = 0.05$ , and averaged over 5 different seeds. Note that here we fixed the number of neurons in the uninformed setting, so the curves Figure 9a and Figure 9b are not directly comparable with each other. For the scheme-aware model, we can observe that the green

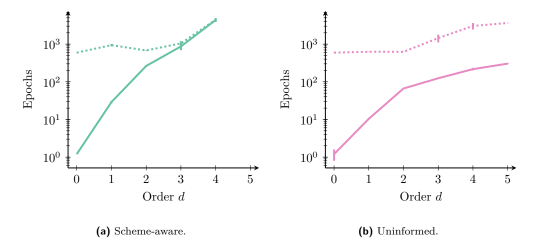


Figure 9: Number of epochs to get weak (plain curves) and strong (dotted curves) learning.

plain curve goes from 1 epoch for d = 0 to 4,000 epochs for d = 4. This denotes an exponential increase of the optimization complexity in weak learning with the masking order. Since the optimization complexity in strong learning (denoted by the dotted curves) is strictly higher than the one in weak learning, we can also deduce that the optimization complexity in strong learning will follow an exponential trend.

For the uninformed model, we may also notice a dramatic increase of the optimization complexity in weak learning, from one epoch without masking to 200 epochs with 6 shares. Unfortunately, Figure 9 does not provide enough evidence to conclude in a sharp way on the same exponential increase of the optimization complexity as observed with scheme-aware models. Indeed, as the size of the exhaustive dataset also increases exponentially, it no longer fits into our 48 GB Nvidia RTX A6000 GPU when  $d \geq 6$ . Does this suggest that uninformed models could efficiently scale with the masking order in terms of optimization complexity, whereas scheme-aware models do not?

#### 5.2.2 A Theoretical Argument that holds for Every Non-Worst-Case Model

Actually, we argue in this section that uninformed models should also face the exponential increase of optimization complexity with respect to the masking order. Our point relies on a theorem proved by Shalev-Shwartz *et al.*, in an almost similar problem [SSS17]. There, the authors investigated to what extent some tasks may be learned in an "end-to-end" manner — *i.e.* uninformed in our terminology —, or by "decomposition" in more elementary learning problems — *i.e.* worst-case in our terminology — would be more

efficient. They emphasized that some problems could be hard to learn with gradient descent in the uninformed setting, as stated hereafter.

**Theorem 1** ([SSS17, Thm. 3], informal). Let **L** denote a d-tuple ( $\mathbf{L}_1, \ldots, \mathbf{L}_d$ ) of input instances, and assume that each  $\mathbf{L}_i$  is i.i.d. standard Gaussian in  $\mathbb{R}^p$ . Define the target function  $h_{\mathbf{u}}(\mathbf{l}) = \prod_{i=1}^d \operatorname{sign}(\mathbf{u}^{\mathsf{T}} \mathbf{l}_i)$ , for some normalized hyperplane  $\mathbf{u} \in \mathbb{R}^p$ . Let  $\mathsf{m}_{\theta}$  be a predictor differentiable with respect to its parameter  $\theta$ , such that  $\mathop{\mathbb{E}}_{\mathbf{L}} \left[ \| \nabla_{\theta} \mathbf{m}_{\theta} \|^2 \right] \leq G(\theta)^2$ , and let  $\mathcal{L}(\theta) = \mathop{\mathbb{E}}_{\mathbf{L}} \left[ \ell \left( \mathsf{m}_{\theta}(\mathbf{L}), h_{\mathbf{u}}(\mathbf{L}) \right) \right]$  be the loss function to minimize, for some smooth function  $\ell(\cdot)$ . Then,

$$\mathbb{E}_{\boldsymbol{u}}\left[\left\|\nabla_{\boldsymbol{\theta}}\mathcal{L}\left(\boldsymbol{\theta}\right) - \mathbb{E}_{\boldsymbol{u}}\left[\nabla_{\boldsymbol{\theta}}\mathcal{L}\left(\boldsymbol{\theta}\right)\right]\right\|^{2}\right] \leq G(\boldsymbol{\theta})^{2} \cdot \mathcal{O}\left(\sqrt{\frac{d\log(p)}{p}}\right)^{a} \quad . \tag{3}$$

Let us interpret the meaning of this theorem from our SCA point of view. Consider one bit, masked with d shares, and assume that the leakage distribution conditionally to each share is noise-free and the same for each bit. Therefore, the leakage model to be learned is denoted by the decision surface materialized by the hyperplane  $\boldsymbol{u}$ , and the masked target bit can be expressed as  $h_{\boldsymbol{u}}(\boldsymbol{l})$ .<sup>13</sup> Note that in a worst-case scenario where the adversary has unlimited profiling powers, the true decision surface  $\boldsymbol{u}$  is known, so the masked bit would be successfully recovered in one trace in the attack phase.

Yet, what Theorem 1 tells us is that the success rate could be much worse for a real-world adversary with limited computational powers using gradient descent. Indeed, the authors of [SSS17] interpret the left hand-side of Equation 3 as a measure of the feedback *signal* returned by the labels through the gradient of the loss function. Then, this result tells us that this feedback signal decreases exponentially fast with the masking order, provided that the dimensionality p of each sub-leakage is high enough.

As a result, the trajectory taken by the parameter  $\theta$  during the gradient descent depends less and less on the features of the target function to learn, denoted by the vector  $\boldsymbol{u}$ , as the masking order increases. Abbe *et al.* showed at NEURIPS'20 that this exponential decrease in the feedback signal would result in an exponential growth in the number of steps in the gradient descent needed to escape the initial plateau [AS20, Thm. 3, Cor. 2]. , [AKem<sup>+</sup>21, ACHM22]. This suggests that the non-exponential trend observed in Figure 9 can be regarded as a simulation artifact since we had p = 1, whereas in most non-worst-case attacks the dimensionality p of each leakage is typically much higher, such that we have  $\frac{d\log(p)}{p} \ll 1$ . Overall, we conclude that the hardness when tackling profiling attacks in non-worst-case settings is mainly due to the optimization procedure, *i.e.* based on gradient descent in this paper, than on the choice of models.

## 6 Is Profiling in a Non-Worst-Case Hard Anyway?

In this paper, we have shown how a real-world adversary could leverage some prior knowledge from the source code of a target implementation, by substituting uninformed attacks with scheme-aware adversaries. As a result, we evidenced how scheme-aware modeling could dramatically improve the efficiency of a side-channel attack in the context of masking, from a profiling complexity point of view. We also showed that the main difficulty for the adversary is due to the drawbacks of GD-based optimization procedures, rather than the selection procedure of the appropriate hyper-parameters of a model. Interestingly, this difficulty is expected to increase with the masking order, which opens a new fundamental question for the SCA developers:

 $<sup>^{13}</sup>h_u$  can be seen as the xor function, where the inputs are remapped over  $\{-1, +1\}$ .

Is the conditional p.m.f. of an intermediate computation that is protected against masking efficiently learnable in a non-worst-case setting?

On the one hand, our problem is somehow close to the well-known Learning Parity with Noise (LPN) problem. This problem is cryptographically hard, as it is the root for some lattice-based cryptographic primitives [GRS08, Pie12]. Yet, without noise this problem becomes efficiently learnable, as it can be solved with Gaussian elimination. Nevertheless, it has been shown that any GD-based approach would result in an exponential optimization complexity [Tho96], [AS20, Thm. 6], [SSS17, Thm. 1]. This provides an example of easy learning problem where GD-based learning can fail. In this respect, it might be interesting to study to what extent the scattershot encoding introduced by Gohr et al. could be an efficient solution [GLS22]. On the other hand, some recent results in learning theory suggest that profiling masked implementations in non-worst-case settings could be hard regardless of the nature of the learning algorithm used by the adversary. Indeed, under the assumption that the leakage model has an additive Gaussian noise, the leakage distribution can be expressed as a Gaussian mixture, with a number of modes increasing exponentially with the masking order. Some recent works showed that there are some Gaussian mixtures for which — under cryptographic assumptions — there is no learning algorithm able to scale polynomially with the number of modes both in terms of computational and profiling complexity [BRST21, GVV22]. In other words, any generative model used for profiling in a non-worst-case setting, may be prone to fail when facing higher-order masking, regardless of the profiling method used by the adversary. Whether this limitation also translates to discriminative models like MLPs or CNNs is a great open question.

This open question naturally has strong impact on the understanding of side-channel evaluation contexts [ABB<sup>+</sup>20]. If the problem is hard, then there is a gap between the profiling complexity in the worst-case and uninformed contexts, increasing with the number of shares. At the extreme, one could imagine that implementation security could also rely on this complexity (i.e., claim an implementation secure if no model can be learned in an uninformed context). On the one hand, this would give some theoretical background to current evaluation approaches that consider attacks using implementation knowledge as more critical. On the other hand, profiling remains a one-time effort and is highly dependent on even mild assumptions that adversaries could make about the implementations. So such an extreme view seems very risky and a more conservative approach would then just be to consider that the possible gap between worst-case and uninformed profiling offers some welcome security margin against very powerful attacks which may help preserving implementation security in the longer term. If the problem is not hard, then the worst-case approach becomes even more unavoidable, as it provides a shortcut to the security level that will be reached even by practical adversaries. We hope that the scheme-aware context can help illuminating this fundamental question in the future. In this respect, the investigation of combined countermeasures (e.g., masking + shuffling or desynchronization) appears as a natural target, and it would be interesting to study how to adapt scheme-aware modeling to these more challenging contexts.

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### A Duality between Masking and Convolutions

Proof of Proposition 1. Let  $\mathbf{p}_{\mathbf{Y}}(\boldsymbol{l})[y]$  denote the probability  $\Pr(\mathbf{Y} = y \mid \mathbf{L} = \boldsymbol{l})$ , and let  $\overline{\mathbf{Y}}$  denote the d + 1 sharing of  $\mathbf{Y}$ , namely  $\overline{\mathbf{Y}} = (\mathbf{Y}_0, \dots, \mathbf{Y}_d)$ . By applying Bayes' theorem we get:

$$\mathbf{p}_{\mathbf{Y}}(\boldsymbol{l})[y] = \frac{\Pr\left(\mathbf{Y}=\boldsymbol{y}\right)}{\Pr\left(\mathbf{L}=\boldsymbol{l}\right)} \Pr\left(\mathbf{L}=\boldsymbol{l} \mid \mathbf{Y}=\boldsymbol{y}\right)$$
(4)

Denoting  $\Pr(\mathbf{L} = \mathbf{l} \mid \mathbf{Y} = y)$  by  $\mathsf{p}_{\mathbf{L}}(y)[\mathbf{l}]$ , and using the total probabilities formula d times, we expand the latter term as follows:

$$\mathsf{p}_{\mathbf{L}}(y)[\boldsymbol{l}] = \sum_{y_1,\dots,y_d \in \mathcal{Y}} \Pr\left(\mathbf{L} = \boldsymbol{l} \mid (\mathbf{Y}, \mathbf{Y}_1, \dots, \mathbf{Y}_d) = (y, y_1, \dots, y_d)\right) \cdot \prod_{i=1}^d \Pr\left(\mathbf{Y}_i = y_i\right) \quad .$$
(5)

By noting  $y_0 \triangleq y \star (y_1 \star \ldots \star y_d)^{-1}$ , and since the mapping  $(y, y_1, \ldots y_d) \mapsto (y_0, y_1, \ldots y_d)$  is invertible we may reformulate the conditional probability as follows:

$$\Pr\left(\mathbf{L} = \boldsymbol{l} \mid (\mathbf{Y}, \mathbf{Y}_1, \dots, \mathbf{Y}_d) = (y, y_1, \dots, y_d)\right) = \Pr\left(\mathbf{L} = \boldsymbol{l} \mid \overline{\mathbf{Y}} = (y_0, \dots, y_d)\right) .(6)$$

Moreover, according to assumption (b), we have:

$$\Pr\left(\mathbf{L} = \boldsymbol{l}\right) = \prod_{i=0}^{d} \Pr\left(\mathbf{L}_{i} = l_{i}\right) , \qquad (7)$$

$$\Pr\left(\mathbf{L} = \boldsymbol{l} \mid \overline{\mathbf{Y}} = (y_0, \dots, y_d)\right) = \prod_{i=0}^{a} \Pr\left(\mathbf{L}_i = l_i \mid \mathbf{Y}_i = y_i\right) .$$
(8)

Finally, we may use the uniform assumption to remark that:  $\Pr(Y = y) = \Pr(Y_0 = y_0)$ . We may now combine Equations (4), (5), (6), (7), (8) and the latter fact:

$$\mathsf{p}_{\mathbf{Y}}(\boldsymbol{l})[y] = \sum_{y_1,\ldots,y_d \in \mathcal{Y}} \mathsf{p}_{\mathbf{Y}_0}(\boldsymbol{l}_0) \left[ y \star (y_1 \star \ldots \star y_d)^{-1} \right] \mathsf{p}_{\mathbf{Y}_1}(\boldsymbol{l}_1)[y_1] \cdot \ldots \cdot \mathsf{p}_{\mathbf{Y}_d}(\boldsymbol{l}_d)[y_d] .$$

## **B** Optimizing Simulations in An Exhaustive Dataset

Even for a noise-free leakage, computing the loss function to minimize in a naive way would become quickly intractable, as it would result in a sum over all possible sharings of Y, *i.e.*,  $2^{n \cdot (d+1)}$  terms.

Hopefully, we can do much better in our simulated framework, as the conditional probability distribution  $\Pr(Y \mid \mathbf{L})$  and the marginal distribution of leakages  $\Pr(\mathbf{L})$  can be used to rephrase the terms in the loss function as follows:

$$\mathcal{L}(\theta) = \sum_{l_0} \dots \sum_{l_d} \mathbb{P}_{\mathrm{r}(\mathrm{Y} \mid \mathbf{L} = (\boldsymbol{l}_0, \dots, \boldsymbol{l}_d))} [\mathsf{m}_{\theta}(\boldsymbol{l}_0, \dots, \boldsymbol{l}_d)] \cdot \prod_{i=0}^d \Pr(\mathrm{L}_i = l_i) \quad .$$
(9)

The sum to compute in Equation 9 contains  $|\mathcal{L}|^{d+1}$  terms, where  $|\mathcal{L}|$  denotes the leakage space of one share. In the case where the leakage model is highly non-injective such as with Hamming weights — *i.e.*  $|\mathcal{L}| = n + 1$  —, computing the latter sum turns out to be much more efficient. For this model, and assuming that the shares are uniformly distributed, the marginal distribution  $\Pr(\mathbf{L})$  is a joint distribution of d binomial laws  $\mathcal{B}(n, 1/2)$ .

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