Tempus Fugit: How To Plug It

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Abstract

Secret or private information may be leaked to an external attacker through the timing behaviour of the system running the untrusted code. After introducing a formalisation of this situation in terms of a confinement property, we present an algorithm which is able to transform the system into one that is computationally equivalent to the given system but free of timing leaks.

Key words: Security Analysis, Timing Leaks, Program Transformation, Probabilistic Processes

1 Introduction

This paper concerns the transformation of programs to remove covert channels. Our focus is on timing channels, which are able to signal information through the time at which an action occurs (a command is executed).

A system is vulnerable to a timing attack if there are alternative control flow paths of observably different duration; a common refinement of this definition would be just to focus on those alternative paths where the selection is guarded by confidential information. We will show that the comparison of different paths is essentially a confinement problem. We have studied confinement in [1] using probabilistic bisimulation. Specifically, we showed that two processes are probabilistically confined if and only if they share a common (probabilistic) abstract interpretation (Proposition 39 of [1]). Our approach to probabilistic abstract interpretation is based on Operator Algebra and thus

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abstractions are measurable by a norm; in particular, in *op. cit.*, we showed how to measure the difference between two processes and thus introduced the notion of approximate confinement. In *op. cit.* this measure, ϵ , was given a statistical interpretation which indicated how much work an attacker would have to perform in order to differentiate the processes.

In this paper we use our previous work to identify the need for transformation. We identify the alternative paths which fail to be bisimilar. These are then candidates for transformation. We present a transformation which establishes probabilistic bisimilarity whilst retaining input/output equivalence with the original system. This transformation involves the introduction of new "padded" control paths. This is similar to the approach of Agat [2]. However, in contrast to Agat, our work is not programming language specific; instead it is based on probabilistic transition systems which provide a very general setting. The transformed system produced by our approach is more efficient than that which would be produced by Agat's algorithm. This is because our algorithm introduces less padding steps.

In the next section, we review some background material on probabilistic transition systems and their representation as linear operators. Section 3 reviews the pertinent results from [1] and shows how to use these results to identify potential timing leaks. Section 4 and Section 6 present our algorithm for padding and its correctness. We demonstrate the algorithm via a simple example in Section 5, and present some comparisons to related work in Section 8.

2 Probabilistic Processes

Probabilistic process algebras extend the well-established classical techniques for modelling and reasoning about concurrent processes with the ability of dealing with non-functional aspects of process behaviour such as performance and reliability, or in general the quantitative counterpart of classical properties representing the functional behaviour of a system.

Several models for probabilistic processes have been proposed in the literature in recent years, which differ from each other essentially in the "amount" of nondeterminism that is allowed in the model. Replacing nondeterministic branching by a probabilistic one leads to either the *reactive* or the *generative* model introduced in [3], where the probability distribution which guides the choice may depend (reactive) or not (generative) on the action labelling the branches (or transitions). Other models allow for both nondeterministic and probabilistic branching and lead to a myriad of variants depending on the interplay between the two kinds of branching (cf. [4]). In order to describe the operational behaviour of probabilistic processes we consider *probabilistic transition systems* (PTS's), that is probabilistic extensions of the notion of labelled transition systems for classical process algebras. Probabilistic transition systems essentially consist of a set of states and a set of labels, and their behaviour can be informally described as follows: To a given action of the environment the system responds by either refusing it or making a transition to a new state according to a probability distribution. In this paper we consider only PTS's with a finite number of states and actions, which reflect the generative model of probability; they effectively correspond to a particular kind of probabilistic processes, namely finite Markov chains [5].

2.1 Probabilistic Transition Systems

Formally, we can define a probabilistic transition system in all generality as in [6]. We denote by Dist(X) the set of all probability distributions on the set X, that is of all functions $\pi: X \to [0, 1]$, such that $\sum_{x \in X} \pi(x) = 1$.

Definition 1 A probabilistic transition system is a tuple $(S, A, \rightarrow, \pi_0)$, where:

- S is a non-empty, finite set of states,
- A is a non-empty, finite set of actions,
- $\longrightarrow \subseteq S \times A \times Dist(S)$ is a transition relation, and
- $\pi_0 \in Dist(S)$ is an initial distribution on S.

For $s \in S$, $\alpha \in A$ and $\pi \in Dist(S)$ we write $s \xrightarrow{\alpha} \pi$ for $(s, \alpha, \pi) \in \longrightarrow$. By $s \xrightarrow{p:\alpha} t$ we denote the transition to individual states t with probability $p = \pi(t)$. We denote the transition probability from a state s_1 to a state s_2 with label α by $p(s_1, \alpha, s_2)$. For a set of states C we write the *accumulated* transition probability from s_1 to C as: $p(s_1, \alpha, C) = \sum_{s \in C} p(s_1, \alpha, s)$.

In the generative model the transition relation of a PTS $(S, A, \longrightarrow, \pi_0)$ is a partial function $\longrightarrow: S \hookrightarrow \text{Dist}(S \times A)$; this means that the same probability distribution is used to govern both the choice of the action and the (internal) state transition. Looking at a PTS as a Markov chain, we can identify it with the *tree* of the outcomes of the associated stochastic process; this can be seen as an experiment which takes place in stages. At each stage the probability for each possible outcome is known when the outcomes of all previous stages are known, and the number of the possible outcomes at each stage is finite. We can define the tree associated to a PTS inductively as follows.

Definition 2 Let S be a non-empty, finite set of states, and let A be a non-empty finite set of actions. Then

- $(\{s\}, A, \emptyset, \{\langle s, 1 \rangle\})$ is a tree-PTS with root $s \in S$;
- if $T_i = (S_i, A, \longrightarrow_i, \{\langle s_i, 1 \rangle\}), i = 1, \dots, m, m < \infty$ are tree-PTS's with roots $s_i \in S_i, S_i \subseteq S$ for all $i = 1, \dots, m, S_i \cap S_j = \emptyset$ for $i \neq j, s \in$ $S \setminus \bigcup_i S_i$, and $\{\langle (s_i, \alpha_i), p_i \rangle \mid s_i \in S_i, \alpha_i \in A, i = 1, \dots, m\}$ is a probability distribution, then the PTS constructed as $(\{s\} \cup \bigcup_i S_i, A, \{s \xrightarrow{p_i:\alpha_i} s_i\} \cup \bigcup \longrightarrow_i, \{\langle s, 1 \rangle\})$ is a tree-PTS with root s. We call T_i the sub-trees of T.

When the transition probabilities form a sub-probability distribution, that is a function $\pi : S \times A \to [0,1]$ with $\sum_{s \in S} \pi(s) \leq 1$, then we call the PTS a subPTS (tree-subPTS).

2.2 Linear Representation of Probabilistic Transition Systems

As in previous work - e.g. [7,1] - we recast the common relational presentation of PTS's in terms of linear maps and operators in order to provide an appropriate mathematical framework for the quantitative study of probabilistic processes.

Definition 3 Given a (sub)PTS $T = (S, A, \rightarrow, \pi_0)$ and a fixed $\alpha \in A$ then the matrix representing the probabilistic transition relation $\xrightarrow{\alpha} \subseteq S \times [0, 1] \times S$ corresponding to α is given by

$$(\mathbf{M}_{\underline{}})_{st} = \begin{cases} p \text{ iff } s \xrightarrow{p:\alpha} t \\ 0 \text{ otherwise} \end{cases}$$

where $s,t \in S$, and $(\mathbf{M})_{st}$ denotes the entry in column s and row t in the matrix \mathbf{M} . We define the matrix representation $\mathbf{M}(T)$ of the (sub)PTS P as the direct sum of the matrix representations of the transition relations $\stackrel{\alpha}{\longrightarrow}$ for each $\alpha \in A$:

$$\mathbf{M}(T) = \bigoplus_{\alpha \in A} \mathbf{M}_{\underline{}}.$$

We recall that for a set $\{\mathbf{M}_i\}_{i=1}^k$ of $n_i \times m_i$ matrices, the *direct sum* of these matrices is defined by the $(\sum_{i=1}^k n_i) \times (\sum_{i=1}^k m_i)$ "diagonal" matrix:

$$\mathbf{M} = \bigoplus_{i} \mathbf{M}_{i} = \operatorname{diag}(\mathbf{M}_{1}, \mathbf{M}_{2}, \dots, \mathbf{M}_{k})$$

Given k square, $n \times n$ matrices \mathbf{M}_i and an $m \times n$ matrix \mathbf{K} then we use the shorthand notation $\mathbf{K}(\bigoplus_{i=1}^k \mathbf{M}_i)$ for the matrix $(\bigoplus_{i=1}^k \mathbf{K})(\bigoplus_{i=1}^k \mathbf{M}_i)$, where we apply \mathbf{K} to each of the factors \mathbf{M}_i .

For probabilistic (sub-probabilistic) relations we obtain a so-called *stochastic* (*sub-stochastic*) matrix, i.e. a positive matrix where the entries in each row sum up to one (are less than or equal to one).

Matrices are not just schemes for writing down probabilities but also a way to specify *linear maps* or *operators* between/on vector spaces. In our case, PTS's are represented as operators on the vector space $\mathcal{V}(S) = \{ (v_s)_{s \in S} \mid v_x \in \mathbb{R} \}$, i.e. the set of formal linear combinations of elements in S with real coefficients. This space contains all distributions of the state space $Dist(S) \subset \mathcal{V}(S)$.

3 Process Equivalences and Confinement

The problem of preventing a system from leaking private information to unauthorised users is essentially the problem of guaranteeing the confinement [8] of the system against some specific attacks. Typically, after the introduction of the notion of noninterference [9,10], most of the work on confinement has been based on models which ultimately depend on some notion of process equivalence by identifying the absence of information flow between two processes via the indistinguishability of their behaviours [11].

In [1] it is shown how various process equivalences can be approximated, and how probabilities can be used as numerical information for quantifying such an approximation. This provides us with a quantitative measure of the indistinguishability of the process behaviour (according to the given semantics), that is in a security setting a measure of their propensity to leak information. In particular, a notion of confinement can be obtained by stating that two processes p and q are probabilistically confined iff they are probabilistic bisimilar.

The central aim of this paper is to present a transformation algorithm for PTS's which constructs bisimilar PTS's while preserving the computational effects of the original systems. This section is devoted to a formal definition of the two notions of process equivalence for PTS's at the base of our treatment, namely probabilistic bisimulation and probabilistic observables.

3.1 Probabilistic Bisimulation

Probabilistic bisimilarity is the standard generalisation of process bisimilarity to probabilistic transition systems and is due to [12].

Definition 4 A probabilistic bisimulation is an equivalence relation \sim_b on

the states of a probabilistic transition system satisfying for all $\alpha \in A$:

 $p \sim_b q \text{ and } p \xrightarrow{\alpha} \pi \Rightarrow q \xrightarrow{\alpha} \varrho \text{ and } \pi \sim_b \varrho.$

Note that in the case of generative systems considered in this paper, π and ρ are sub-probability distributions.

The notion of probabilistic bisimulation on PTS corresponds to the notion of lumped process for finite Markov chains [5]. This can be seen by noting that a probabilistic bisimulation equivalence ~ on a PTS $T = (S, A, \rightarrow, \pi_0)$ defines a probabilistic abstract interpretation of T [1]. In fact, we have shown that it is always possible to define a linear operator **K** from the space $\mathcal{V}(S)$ to the space $\mathcal{V}(S/_{\sim})$ which represents ~. This so-called classification operator is represented by the $n \times m$ -matrix:

$$(\mathbf{K})_{ij} = \begin{cases} 1 \text{ if } s_i \in C_j \\ 0 \text{ otherwise} \end{cases}$$

with $S/\sim = \{C_1, C_2, \ldots, C_m\}$ and *n* the cardinality of *S*. If $\mathbf{M}(T)$ is the operator representation of *T* then $\mathbf{K}^{\dagger}\mathbf{M}(T)\mathbf{K}$ is the abstract operator induced by \mathbf{K} where \mathbf{K}^{\dagger} is the so-called *Moore-Penrose pseudo-inverse* of \mathbf{K} . For classification operators \mathbf{K} we can construct \mathbf{K}^{\dagger} as the row-normalised transpose of \mathbf{K} . Intuitively, $\mathbf{K}^{\dagger}\mathbf{M}(T)\mathbf{K}$ is an operator which abstracts the original system *T* by encoding only the transitions between equivalence classes instead of the ones between single states. \mathbf{K} determines a partition on the state space of *T* which is exactly the lumping of the states of *T*. This can be formally deduced from the following theorem in [5].

Theorem 5 A necessary and sufficient condition for a Markov chain to be lumpable with respect to a partition $\{C_1, C_2, \ldots, C_m\}$ is that for every pair of sets C_i and C_j , the probability $p_{kC_j} = \sum_{t \in C_j} p_{it}$ of moving in one step from state s_k into set C_j have the same value for every $s_k \in C_i$. These common values $\{\tilde{p}_{ij}\}$ form the transition matrix for the lumped chain.

We can re-formulate the probabilistic bisimilarity of two processes A and B in terms of linear operators as follows, cf [1]:

Proposition 6 Given the operator representation $\mathbf{M}(A)$ and $\mathbf{M}(B)$ of two probabilistic processes A and B, then A and B are probabilistic bisimilar, that is $A \sim_b B$, iff there exist classification matrices \mathbf{K}_A and \mathbf{K}_B such that

$$\mathbf{K}_{A}^{\dagger}\mathbf{M}(A)\mathbf{K}_{A}=\mathbf{K}_{B}^{\dagger}\mathbf{M}(B)\mathbf{K}_{B}.$$

Probabilistic bisimilarity of A and B can be expressed in terms of Markov chains lumpability as follows:

Proposition 7 Given the operator representation $\mathbf{M}(A)$ and $\mathbf{M}(B)$ of two probabilistic processes A and B, then A and B are probabilistic bisimilar iff there exists a lumping \mathbf{K} of the process $\mathbf{M}(A) \oplus \mathbf{M}(B)$ of the form

$$\mathbf{K} = egin{pmatrix} \mathbf{K}_A \ \mathbf{K}_B \end{pmatrix}$$

for some classification operators \mathbf{K}_A and \mathbf{K}_B for A and B such that \mathbf{K}_A and \mathbf{K}_B are fully column-ranked.

For tree-PTS the above result essentially means that an equivalence relation on the union of the states of two processes A and B is a probabilistic bisimulation for A and B iff their roots belong to the same equivalence class.

Probabilistic bisimulation is the finest of all process equivalences as it identifies processes whose step-wise behaviour is exactly the same. From an external viewpoint other behaviours can be observed in order to distinguish two processes. In particular, a potential attacker may be able to obtain experimentally two kinds of information about a process by observing what final outcomes the process produces, and by measuring how long the process is running. Both the input/output behaviour and the timing behaviour of a process correspond to some suitable *abstractions* of the concrete behaviour of the process in question. In this paper we will concentrate on these observables and show how we can transform a process (PTS) in a way that its input/output behaviour is preserved, while its timing behaviour changes so as to become indistinguishable from another given process.

3.2 Probabilistic Observables

In our PTS model, any execution of a process leads to a single trace, i.e. a single execution path. Each time there is a probabilistic choice, one of the possible branches is chosen with some probability. The combined effect of all the choices made defines the probability of a particular path as the product of probabilities along this computational path. For finite processes we only have to consider finitely many paths of finite length and thus only need a basic probability model. This is based on a finite set of events Ω , i.e. all possible traces, such that all sets of traces have a well-defined probability attached, i.e. are measurable, and the probability function $\mathbf{P} : \mathcal{P}(\Omega) \to [0, 1]$ can be defined via a distribution, i.e. a function $\pi : \Omega \to [0, 1]$ such that $\mathbf{P}(X) = \sum_{x \in X} \pi(x)$

for all $X \subseteq \Omega$.

The notion of observables we will consider is based on a *trace semantics* for probabilistic processes whose intuition and formal definition are given below.

3.2.1 Trace Semantics

Formally, the trace semantics of a process T is defined via the notion of probabilistic result of a computational path.

Definition 8 For a computational path $\pi = s_0 \xrightarrow{p_1:\alpha_1} s_1 \xrightarrow{p_2:\alpha_2} \dots \xrightarrow{p_n:\alpha_n} s_n$, we define its probabilistic result $\mathcal{R}(\pi)$ as the pair $\langle \alpha_1 \alpha_2 \dots \alpha_n, prob(\pi) \rangle$, where $prob(\pi) = \prod_i p_i$. The trace associated to π is $t(\pi) = \alpha_1 \alpha_2 \dots \alpha_n$.

Since different computational paths may have the same associated trace (cf. Example 9), we need to sum up the probabilities associated to each such computational path in order to appropriately define a probabilistic result. Formally, this can be done by defining for a process T and trace t the set

 $[\pi]_{T,t} = \{\pi \mid \pi \text{ a computational path for } T \text{ and } t(\pi) = t\},\$

which collects all the computational paths with trace t. Then we define for each trace t the computational result $\mathcal{R}([\pi]_{T,t})$ as the pair $\langle t, \sum_{\pi \in [\pi]_{T,t}} prob(\pi) \rangle$.

We can now define the trace semantics of T as the set

 $\mathcal{S}(T) = \{\mathcal{R}([\pi]_{T,t}) \mid t = t(\pi) \text{ and } \pi \text{ is a computational path for } T\}.$

Example 9 Consider the following simple processes with action $A = \{a, b, e\}$:



It is easy to see that the possible traces for the first two processes are given by the sequence ab while for process T_3 we have the traces abe and aeb and for process T_4 we have aeb and ab.

We can thus describe the trace semantics of these four processes by:

$$\mathcal{S}(T_1) = \{ \langle ab, 1 \rangle \}$$

$$\mathcal{S}(T_2) = \{ \langle ab, 1 \rangle \}$$

$$\mathcal{S}(T_3) = \{ \langle abe, 0.5 \rangle, \langle aeb, 0.5 \rangle \}$$

$$\mathcal{S}(T_4) = \{ \langle aeb, 0.5 \rangle, \langle ab, 0.5 \rangle \}$$

3.2.2 I/O Observables

The notion of input/output observables, which we will refer to as I/O observables from now on, correspond to the most abstract observation of a process behaviour as it just considers the final result of an execution. In our setting, these final results are identified with *probability distributions* over the set of all possible *abstract traces*. Abstract traces are defined in terms of a function [[.]] which allows us to avoid specifying a concrete semantics for the labels, thus keeping our approach as general as possible. In fact, in a PTS we can think of labels as some kind of "instructions" which are performed by the process, e.g. $a = \mathbf{x} := \mathbf{x}-\mathbf{1}^n$, $b = \mathbf{x} := 2*\mathbf{x}^n$, etc. The final result of an execution is thus the accumulated effect of executing these instructions. Depending on the specific application, the function [[.]] will be defined so as to abstract from the appropriate computational details of the concrete traces.

For our purposes, we can leave the actual nature of [.] unspecified and only assume that there is a neutral instruction e, i.e. a label which represents something like a **skip** statement, such that for any trace $t_1t_2 \ldots t_n$ we have:

$$\llbracket t_1 t_2 \dots t_n \rrbracket \equiv \llbracket t_1 t_2 \dots t_i e t_{i+1} t_n \rrbracket$$

i.e. putting e anywhere in the execution (including at the end or the beginning) does not change the computational result. We will identify traces if the only difference between them are some occurrences of this neutral label e; this introduces an equivalence relation \equiv on the set of all traces.

Clearly, different concrete traces might be identified by $[\![.]\!]$; for example, if a = "x := x+1" and b = "x := x-1" one would usually have $[\![ab]\!] = [\![e]\!] = [\![ba]\!]$.

Definition 10 Given a tree-PTS T, we define its I/O observables as

$$\mathcal{O}(T) = \{ \mathcal{R}([\pi]_{T, \llbracket t \rrbracket}) \mid \pi \text{ is a computational path for } T \}.$$

Proposition 11 For any tree-PTST, its I/O observables $\mathcal{O}(T)$ define a probability distribution on the set of states S.

PROOF. By induction on the tree structure of T.

- If $T = (\{s\}, A, \emptyset, \{\langle s, 1 \rangle\})$ then $\mathcal{O}(T)$ is obviously a probability distribution on S.
- If $T = (\{s\} \cup \bigcup_i S_i, A, \{s \xrightarrow{p_i:\alpha_i} s_i\} \cup \bigcup \longrightarrow_i, \{\langle s, 1 \rangle\})$, then by the inductive hypothesis we have for all $i = 1, \ldots, m$ that $\mathcal{O}(T_i)$ is a probability distribution on S_i . Therefore, $\mathcal{O}(T) = \sum_{i=1}^m p_i \cdot \mathcal{O}(T_i)$ is a probability distribution on $\bigcup_i S_i$. \Box

Based on this notion of I/O observables we can now define when two processes are I/O equivalent.

Definition 12 Two tree-PTS's A and B are I/O equivalent, denoted by $A \sim_{io} B$, if they define the same probability distribution on the set of the equivalence classes of abstract traces, i.e. iff $\mathcal{O}(A) = \mathcal{O}(B)$.

Example 13 If we consider again the four processes from Example 9, we get the following I/O observables:

 $\mathcal{O}(T_1) = \{ \langle \llbracket ab \rrbracket, 1 \rangle \}$ $\mathcal{O}(T_2) = \{ \langle \llbracket ab \rrbracket, 1 \rangle \}$ $\mathcal{O}(T_3) = \{ \langle \llbracket abe \rrbracket, 0.5 \rangle, \langle \llbracket aeb \rrbracket, 0.5 \rangle \} = \{ \langle \llbracket ab \rrbracket, 1 \rangle \}$ $\mathcal{O}(T_4) = \{ \langle \llbracket aeb \rrbracket, 0.5 \rangle, \langle \llbracket ab \rrbracket, 0.5 \rangle \} = \{ \langle \llbracket ab \rrbracket, 1 \rangle \}$

i.e. all four processes are I/O equivalent.

3.3 Timing Behaviour

An external observer may have the ability to measure the time a process needs to perform a given task. In order to do so she just needs to observe "execution times", while the concrete nature of the performed instruction is irrelevant for her purposes. In our model, the timing behaviour of a process can therefore be defined by abstracting from labels (instructions) and assuming that there is a function |.| from labels to real numbers which measures the time it takes to execute the instruction represented by the label.

One can think of various choices for the timing function. For example, one can assume (as in [13]) that there is only one special label t with |t| = 1 which indicates time progress, while all other labels take no execution time; in this case the time flow is measured by the number of these special labels occurring in a trace. Alternatively, one can assign a different duration to each label.

In our model we assume that all labels need a strictly positive time to be executed, i.e. $|\alpha| > 0$ for all labels, and that the execution time of all labels is the same. In particular, while in general one could have various neutral labels e with different executions time, e.g. $|e_0| = 0$, $|e_1| = 1, \ldots$, we will assume that there is only one neutral element, denoted by $\sqrt{}$, which consumes exactly one time unit.

Formally we define the time semantics of our processes in terms of the notion of *time bisimulation*, that is a bisimulation relation on PTS's where transitions are labelled only by action $\sqrt{}$.

Definition 14 Given a tree-PTS $T = (S, A, \rightarrow, \pi_0)$, we define its depleted or ticked version \hat{T} , as the PTS $(S', A', \rightarrow', \pi'_0)$ with S' = S, $A' = \{\sqrt\}$, $\pi'_0 = \pi_0$, and \longrightarrow' defined by

$$(s,\sqrt{\pi}) \in \mathcal{A}$$
 iff $(s,\alpha,\pi) \in \mathcal{A}$ for some $\alpha \in A$.

Time bisimilarity is defined as a probabilistic bisimilarity (cf. Definition 4) for depleted versions of PTS.

Definition 15 Given two PTS's A and B, we say that A and B are time bisimilar, denoted by $A \sim_{tb} B$, iff there is a probabilistic bisimulation on the states of the ticked versions of A and B.

Example 16 The following two processes P_1 and P_2 are not probabilistic bisimilar; they are nevertheless time bisimilar as one can easily see by considering their ticked versions \hat{P}_1 and \hat{P}_2 :



On the other hand it is easy to see that two deterministic processes, i.e. processes which correspond to a single trace, are time bisimilar if and only if they have the same length, as the following example illustrates:



Our model for timing leaks essentially follows the noninterference based approach to confinement recalled at the beginning of this section. In particular, we define timing attacks in terms of the ability of distinguishing the timing behaviour of two processes expressed via the notion of time bisimilarity.

In [1] we have used the same idea to model probabilistic covert channels. The notion of process equivalence used in that paper is an approximate version of probabilistic bisimilarity as the purpose there is to provide an an estimate of the confinement of the system, rather than just checking whether it is confined or not. We aim to extending this current work toward a similar direction (cf. Section 8). To this purpose, the linear representation of PTS, i.e. their correspondence with processes which are Markov chains, is essential; in fact, as shown in [1], it provides the necessary quantitative framework for defining numerical estimates.

Definition 17 We say that two processes T_1 and T_2 are confined against timing attacks or time confined *iff their associated PTS are time bisimilar*.

We can re-formulate this definition of timing leaks in terms of lumping as done in Proposition 6 for probabilistic bisimilarity. This will be the base for the definition of the transformation technique we will present in Section 4.

Definition 18 Given the operator representation $\mathbf{M}(T) = \bigoplus_{\alpha \in A} \mathbf{M}_{\alpha}(T)$ of a probabilistic process T then the linear representation of its ticked version is given by:

$$\widehat{\mathbf{M}}(T) = \sum_{\alpha \in A} \mathbf{M}_{\alpha}(T)$$

Definition 19 We say that A and B are time bisimilar, $A \sim_{tb} B$, iff there exist classification matrices \mathbf{K}_A and \mathbf{K}_B , such that

$$\mathbf{K}_{A}^{\dagger}\widehat{\mathbf{M}}(A)\mathbf{K}_{A} = \mathbf{K}_{B}^{\dagger}\widehat{\mathbf{M}}(B)\mathbf{K}_{B}.$$

In the following we present a technique for closing possible covert timing channels by transforming the system into an equivalent one which is free from timing leaks. The transformation algorithm we present is applicable to finite tree-PTS's. It uses ideas from the optimal lumping algorithm in [14] and the padding techniques from [2].

Intuitively, given a system formed by processes A and B, the algorithm is based on the construction of a lumping for the Markov chain formed by the direct sum of the two Markov chains representing A and B. By Proposition 7, the resulting lumped process will be a probabilistic bisimulation for A and B iff the root nodes of A and B belong to the same class (or they are essentially the same state in the lumped process). In this case no transformation is needed as the two processes are confined against timing attacks; otherwise the algorithm produces two new processes A' and B' which exhibit the same I/O behaviour as A and B but are also time confined. The following diagram shows the effect of the transformation.



4 Transforming Probabilistic Transition Systems

Checking whether two systems are bisimilar or not, aka the bisimulation problem in the literature, is important not only in concurrency theory but also in many other fields such as verification and model checking, where bisimulation is used to minimise the states space, but also in computer security where many non-interference properties can be specified in terms of bisimulation equivalence [15,1]. Various algorithms have been proposed which solve this problem in linear time, the most important one being the algorithm proposed by Paige and Tarjan in [16]. This algorithm has been adapted in [14] for probabilistic processes; in particular, the Derisavi et al. algorithm constructs the optimal lumping quotient of a finite Markov chain in linear time, and can then be used to check the probabilistic bisimilarity of two PTS's.

In this paper we take a transformational approach: we not only check whether two probabilistic processes are bisimilar or not, but in the case where the processes are not bisimilar then we transform them into two bisimilar ones. The transformation we propose is inspired by the Derisavi et al. algorithm but does not aim to construct an "optimal" partition in the sense of [16]. It rather aims to add missing states and adjust the transition probabilities when it is needed to make two given processes bisimilar. It also preserves the probabilistic observable of the original processes.

The algorithm we present consists in essentially two intertwined procedures: one constructs a lumping partition for the PTS union of two tree-PTS's; the other "fixes" the partition constructed by the lumping procedure if some properties are violated which guarantee the partition to be a probabilistic bisimulation for the given processes. Fixing corresponds to adding some missing states (padding) and adjusting the transition probabilities in order to achieve bisimilarity. Moreover, the transformation is performed in a way that the original I/O observables of the processes is preserved. The overall procedure is repeated until a partition is constructed which effectively makes the two processes probabilistically bisimilar.

Before embarking on a more detailed description of our algorithm, we introduce some necessary notions and definitions. In the rest of this paper we focus on finite tree-structured PTS.

For a PTS $(S, A, \longrightarrow, \pi_0)$ we will consider partitions $P = \{B_1, \ldots, B_n\}$ of S; we will refer to the B_i 's as the *blocks* of the partition P. Given a state $s \in S$ and a partition P we define the cumulative probability of s with respect to $B \in P$ as $p(s, B) = \sum \{q \mid s \xrightarrow{q} t \text{ with } t \in B\}$. We further define the backward image for a block B and a probability q as $pre_{q, \longrightarrow}(B) = \{s \mid \exists B' \subseteq B : p(s, B') = q\}$; this corresponds to the set of all states from which B (i.e a state in B) can be reached with probability q. We will often omit the index \longrightarrow when it is clear from the context.

For a tree-PTS T we will refer to HEIGHT(T) as the maximal length of a path in T from the root to a terminal state. The n layer cut-off of a tree-PTS Tis defined inductively as (i) the set of terminal nodes for n = 0, and (ii) for n = i + 1 the set of nodes with only direct links (paths of length one) to nodes in the i layer cut-off. The procedure CUTOFF(T, n) returns the n layer cut-off of T which we also denote by $T|_n$. The complement of the n layer cut-off is sometimes called the n layer top and the top nodes in the n layer cut-off form the n layer.

Auxiliary Procedures. An important feature of the transformation of PTS's we present in the following is the creation of new states. These will be copies of existing states or of a distinguished additional "dummy state" d. We will denote copies of a given state s by s', s'', s''', \ldots . For copies of the dummy state we usually will omit the primes. For a tree-PTS $T = (S, A, \longrightarrow, \{\langle r, 1 \rangle\})$ we define its copy as the tree-PTS $T' = (S', A, \longrightarrow', \{\langle r', 1 \rangle\})$ with $S' = \{s' \mid s \in S\}$ and $s' \xrightarrow{p:\alpha} t'$ iff $s \xrightarrow{p:\alpha} t$.

In the main algorithm we assume a routine COPYTREE(T,s) which returns a copy of the sub-tree of a tree-PTS T which is rooted in state s (where s is a state in T). We will also use a routine DEPLETETREE(T, s) which takes a tree-PTS T and returns \check{T} , i.e. a copy of T where all states are replaced by copies of the dummy state d and all transitions are labelled by $\sqrt{}$.

Another auxiliary procedure is LINKING (T_1, s, α, p, T_2) whose effect is to introduce a (copy) of a tree-PTS T_1 into an existing tree-PTS T_2 at state s; the linking transition will be labelled by α and the associated transition proba-

```
1: procedure LUMPING(T)

2: Assume: T is a tree-PTS

3: P \leftarrow \{S\}

4: while \negSTABLE(P, T) do

5: choose B \in P

6: P \leftarrowSPLITTING(B, P)

7: end while

8: end procedure
```

Algorithm 1. Lumping Algorithm

bility will be p. A related procedure called JOINING (T_1, s, p, T_2) introduces a (copy) of a tree-PTS T_2 into an existing tree-PTS T_1 at state s by identifying the root of T_1 with s and by multiplying initial transitions with probability p.

Lumping. As already mentioned, our algorithm is inspired by the Paige-Tarjan algorithm for constructing bisimulation equivalences [16], i.e. *stable* partitions, for a given transition system. The same ideas can be used for probabilistic transition relations \xrightarrow{p} ; in this case we say that a partition $P = \{B_1, \ldots, B_n\}$ of a set of states is stable with respect to \xrightarrow{p} iff for all blocks B_i and B_j in P and probability q we have that $pre_q(B_i) \cap B_j = \emptyset$ or $B_j \subseteq pre_q(B_i)$.

The construction of the lumping partition is done via the refinement of an initial partition which identifies all states in one class and proceeds as in the algorithm LUMPING (cf Algorithm 1). Starting with the initial partition $P = \{S\}$ the algorithm incrementally refines it by a splitting procedure. For a block B_i the splitting procedure SPLITTING (B_i, P) refines the current partition P by replacing each block $B_j \in P$ with $pre_p(B_i) \cap B_j \neq \emptyset$ for some probability p by the new blocks $pre_p(B_i) \cap B_j$ and $B_j \setminus pre_p(B_i)$. The block B_i is called a *splitter*. We will also assume the definition of a procedure STABLE(P, T) which returns *true* iff partition P is stable for PTS T.

In order to construct a time bisimulation \sim_{tb} for two tree-PTS's T_1 and T_2 we apply a slightly modified lumping procedure (cf Algorithm 2) to the ticked version \hat{T} of the union PTS $T = T_1 \oplus T_2$. The procedure MAXLUMPING (T_1, T_2) takes two tree-PTS's $T_1 = (S_1, A, \longrightarrow, \{\langle r_1, 1 \rangle\})$ and $T_2 = (S_2, A, \longrightarrow, \{\langle r_2, 1 \rangle\})$ and tries to construct a time bisimulation for T_1 and T_2 by lumping T.

At each step the procedure constructs a partition $P = \{B_1, B_2, \ldots, B_m\}$ of the set $S_1 \cup S_2$ and identifies a set of *critical blocks* $C = \{C_1, \ldots, C_k\} \subseteq P$. A critical block C_j is an equivalence class which has representatives in only one of the two PTS's, i.e. such that $C_j \cap S_1 = \emptyset$ or $C_j \cap S_2 = \emptyset$. The procedure works on "layers", i.e. at each iteration MAXLUMPING (T_1, T_2) partitions the nodes in layer n by choosing a splitter among the blocks in the n layer cut-off;

1: procedure MAXLUMPING (T_1, T_2) Assume: $T_1 = (S_1, A, \longrightarrow, \{\langle r_1, 1 \rangle\})$ is a tree-PTS 2: Assume: $T_2 = (S_2, A, \longrightarrow, \{\langle r_2, 1 \rangle\})$ is a tree-PTS 3: 4: $n \leftarrow 0$ \triangleright Base Case $P \leftarrow \{S_1 \cup S_2\}$ \triangleright Partition 5: $S \leftarrow \{S_1 \cup S_2\}$ \triangleright Splitters 6: 7: while $n \leq \text{HEIGHT}(T_1 \oplus T_2)$ do $TT \leftarrow \text{CUTOFF}(T_1 \oplus T_2, n)$ 8: $\triangleright n$ layer cut off 9: $P \leftarrow \{B \cap TT \mid B \in P\}$ \triangleright Partition below current layer $S \leftarrow P \cup \{(S_1 \cup S_2) \setminus TT\}$ 10: \triangleright Splitters (below and top) while $S \neq \emptyset$ do 11: choose $B \in S, S \leftarrow S \setminus B$ 12: \triangleright Choose a splitter $P \leftarrow \text{Splitting}(B, P)$ \triangleright Split (current layer) 13: $C \leftarrow \{B \in P \mid B \cap S_1 = \emptyset \lor B \cap S_2 = \emptyset\}$ 14: \triangleright Critical class(es) 15:if $C \neq \emptyset$ then 16:return C \triangleright Found critical class(es) 17:end if end while 18:19: $n \leftarrow n+1$ \triangleright Go to next level 20: end while return C 21: 22: end procedure

Algorithm 2. Maximal Lumping Algorithm

it moves on to the next layer only when all such blocks have been considered.

Padding. The padding procedure identifies the critical blocks by calling the procedure MAXLUMPING (T_1, T_2) and then transforms the two processes T_1 and T_2 in order to "fix" the transitions to the critical blocks. The transformation is performed by the procedure FIXIT which we will describe later. In the special case where the critical blocks do not contain any representative of either of the two processes, say T_1 , the fixing consists in padding T_1 with a depleted sub-tree formed by its root only which is a dummy state. This is linked to the root of T_1 via a ticked transition with probability 1 by the LINKING procedure. In the general (more symmetric) case where some of the critical blocks contain only states from one process and some other critical blocks only states from the other process the transformation is made by the FIXIT procedure.

When among the critical blocks of the current partition there is at least one block C_k which contains only representatives of T_1 and one block C_j which contains only representatives of T_2 , the fixing is more elaborate and involves both processes. In this case there must exist states $s_1 \in C_k$ and $s_2 \in C_j$ with different probabilities of reaching some other blocks B_i of the partition.

```
1: procedure PADDING(T_1, T_2)
 2:
           C \leftarrow \text{MAXLUMPING}(T_1, T_2)
                                                                                   \triangleright identify critical classes
 3:
           repeat
                 if C = \{C_i\}_i with C_i \cap S_1 = \emptyset, \forall C_i then
 4:
                      D \leftarrow (\{d\}, \{\sqrt{\}}, \emptyset, \{\langle d, 1\rangle\})
 5:
                      T_1 \leftarrow \text{LINKING}(D, d, \sqrt{1}, T_1)
 6:
 7:
                 else if C = \{C_i\}_i with \forall C_i : C_i \cap S_2 = \emptyset then
                      D \leftarrow (\{d\}, \{\sqrt\}, \emptyset, \{\langle d, 1\rangle\})
 8:
 9:
                      T_2 \leftarrow \text{Linking}(D, d, \sqrt{1, T_2})
                else if C = \{C_i\}_i with C_j \subseteq S_1 \neq \emptyset \neq C_k \subseteq S_2 then
10:
11:
                      choose s_1 \in C_k
12:
                      choose s_2 \in C_i
                      FIXIT(T_1, s_1, T_2, s_2)
13:
14:
                 end if
15:
                 C \leftarrow \text{MAXLUMPING}(T_1, T_2)
                                                                                                  \triangleright i.e. T_1 \sim_{tb} T_2
16:
           until C = \emptyset
           return T_1, T_2
17:
18: end procedure
```

Algorithm 3. Padding Algorithm

The FIXIT procedure selects two blocks B_k and B_l in previous layer such that the probabilities p_{1k}, p_{1l} of the transitions from s_1 to B_k and B_l and the probabilities p_{2k}, p_{2l} of the transitions from s_2 to B_k and B_l are such that $p_{1k} > p_{2k}$ and $p_{1l} < p_{2l}$. This choice is always possible. In fact, the sum of all probabilities from s_1 and s_2 to other classes sum up to one, i.e. $\sum_i p_{1i} = 1 =$ $\sum_i p_{2i}$ where *i* covers all classes B_i reachable from s_1 or s_2 . As s_1 and s_2 are in different (critical) blocks the probabilities must differ for at least one class B_i , i.e. there exists at least one index k with $p_{1k} \neq p_{2k}$; thus we have that either $p_{1k} > p_{2k}$ or $p_{1k} < p_{2k}$. Without loss of generality, let $p_{1k} > p_{2k}$, i.e. we found the class B_k as required. We now have $\sum_{i\neq k} p_{1i} + p_{1k} = 1 = \sum_{i\neq k} p_{2i} + p_{2k}$ and thus $\sum_{i\neq k} p_{1i} < \sum_{i\neq k} p_{2i}$. As the summation is over the same number of probabilities, i.e. positive numbers, there must be at least one summand in $\sum_{i\neq k} p_{2i}$ such that $p_{2l} > p_{1l}$.

After that, FIXIT selects four states $t_{1k} \in B_k \cap S_1$, $t_{2k} \in B_k \cap S_2$, $t_{1l} \in B_l \cap S_1$, and $t_{2l} \in B_l \cap S_2$ with $s_1 \xrightarrow{q_{1k}} t_{1k}$, $s_1 \xrightarrow{q_{1l}} t_{1l}$, $s_2 \xrightarrow{q_{2k}} t_{2k}$, and $s_2 \xrightarrow{q_{2l}} t_{2l}$ and the corresponding sub-trees T_{1k}, T_{1l}, T_{2k} and T_{2l} which are rooted in t_{1k}, t_{1l}, t_{2k} and t_{2l} . Note that since B_k and B_l are in the CUTOFF(n-1) they are not critical blocks, thus it is always possible to find such state. The probabilities q_{1l} and q_{2k} stay unchanged but q_{1k} and q_{2l} will be decreased by the maximal possible amount. Ideally, this decrement should be r_k and r_l , but if q_{1k} and q_{2l} are smaller than r_k and r_l we only reduce q_{1k} and q_{2l} to zero, i.e. effectively we remove the corresponding sub-trees.

The states s_1 and s_2 now have the same probabilities to reach the classes B_l and

1: procedure FIXIT (T_1, s_1, T_2, s_2) 2: for all B_i blocks do \triangleright determine all possible changes Assume: $s_1 \xrightarrow{p_{1i}} B_i$ and $s_2 \xrightarrow{p_{2i}} B_i$ 3: 4: $p'_i \leftarrow \min(p_{1i}, p_{2i})$ end for 5:Assume: B_k is such that $p'_k = p_{2k}$ 6: 7: Assume: B_l is such that $p'_l = p_{1l}$ $r_k \leftarrow p_{1k} - p'_k$ 8: 9: $r_l \leftarrow p_{2l} - p'_l$ choose $t_{1k} \in B_k \cap S_1$ and T_{1k} sub-tree starting with $s_1 \xrightarrow{1} t_{1k}$ 10:choose $t_{2k} \in B_k \cap S_2$ and T_{2k} sub-tree starting with $s_1 \xrightarrow{1} t_{2k}$ 11:choose $t_{1l} \in B_l \cap S_1$ and T_{1l} sub-tree starting with $s_2 \xrightarrow{1} t_{1l}$ 12:choose $t_{2l} \in B_l \cap S_2$ and T_{2l} sub-tree starting with $s_2 \xrightarrow{1} t_{2l}$ 13: \triangleright make maximal possible changes of probabilities 14: Assume: $s_1 \xrightarrow{q_{1k}} t_{1k}, s_1 \xrightarrow{q_{1l}} t_{1l}, s_2 \xrightarrow{q_{2k}} t_{2k}, \text{ and } s_2 \xrightarrow{q_{2l}} t_{2l}$ 15: $r_{1k} \leftarrow \min(q_{1k}, r_k)$ 16: $q_{1k} \leftarrow (q_{1k} - r_{1k})$ 17: $r_{2l} \leftarrow \min(q_{2l}, r_l)$ 18: $q_{2l} \leftarrow (q_{2l} - r_{2l})$ 19:20: \triangleright construct and link up "compensating" subtrees $X_{kl}^1, Y_{kl}^1 \leftarrow \text{Padding}(T_{1k}, \hat{T}_{1l})$ 21: $X_{kl}^2, Y_{kl}^2 \leftarrow \text{PADDING}(\hat{T}_{2k}, T_{2l})$ 22: 23: JOINING $(T_1, s_1, r_{1k}, X_{kl}^1)$ JOINING $(T_2, s_2, r_{2l}, Y_{kl}^2)$ 24: 25: end procedure

Algorithm 4. Fixing Algorithm

 B_k , i.e. have become bisimilar. However, this transformations has two flaws: (i) the system is no longer a PTS as the probabilities from s_1 and s_2 no longer add up to one, (ii) the distribution on the terminal states reachable in T_{1k} and T_{2l} are reduced. In order to fix this we have to construct "compensating trees" which are bisimilar and which reproduce the missing probabilities for the outputs. These two trees are constructed via calling PADDING on a copy of T_{1k} and a depleted version of T_{1l} , and a copy of T_{2l} and a depleted version of T_{2k} respectively. Finally, these "compensating trees" are linked to s_1 and s_2 with a \sqrt{label} and the missing probabilities r_{1k} and r_{2l} .

This way we obtain again a proper tree-PTS with normalised probability distributions for s_1 and s_2 . Furthermore, the "compensating trees" produce exactly the same output as T_{1k} and T_{2l} , thus the resulting PTS's have the same overall output as the original ones. Finally, by construction the compensating trees are bisimilar as T_{1k} and T_{2k} are bisimilar; the same holds for T_{1l} and T_{2l} . Consider two processes A and B (with $p \neq q$ and $p, q \notin \{0, 1\}$) defined as:



The aim is to construct two processes A' and B' such that $A' \sim_{tb} B'$ and $A \sim_{io} A'$ and $B \sim_{io} B'$. Note that A and B do not have the same I/O observables, i.e. $A \not\sim_{io} B$, and that we also do not require this for the transformed processes.

If we call MAXLUMPING(A, B) we have as first splitter the set of all states, i.e. $S_1 = \{1, 2, ..., 8\}$. The backward image of S_1 , i.e. calling SPLITTING($S_1, \{S_1\}$), allows us to distinguish the terminal nodes from 'internal' nodes, i.e. we get the partition $K_1 = \{\{2, 4, 6, 8\}, \{1, 3, 5, 7\}\}$. Obviously, both processes have representatives in each of the two blocks of this partition.

If we continue lumping using the splitter $S_2 = \{2, 4, 6, 8\}$, i.e. the terminal nodes, we get a refined partition $K_2 = \{\{2, 4, 6, 8\}, \{3, 7\}, \{1\}, \{5\}\}$. The probability of reaching S_2 from nodes 3 and 7 is the same (namely 1) while 1 and 5 reach the terminal nodes with different probabilities (namely p and q) and thus form two separate blocks.

The splitting K_2 now is not properly balanced, i.e. the class $\{1\}$ has no representatives in B and $\{5\}$ has no representatives in A. MAXLUMPING(A, B) will thus report $\{1\}$ and $\{5\}$ as critical classes.

In order to fix the 'imbalance' between A and B we have to do two things: (i) minimising/unifying the probabilities of transitions emitting from 1 and 5, and (ii) introducing subtrees X and Y to compensate for the missing outputs and to re-establish normalised probabilities.

• The first step thus is to transform A and B into the following form:



These two transformed versions of A and B are now probabilistically tick bisimilar but unfortunately they have different outputs, i.e. they are not I/O equivalent; moreover they fail to fulfill the normalisation condition for PTS and Markov chains.

• To compensate for the "missing" probability $r = 1 - \min(p, q) - \min(1 - p, 1-q)$ we have to construct subtrees X and Y which reproduce the missing output states and which are bisimilar. As we have a critical class, namely $C_k = \{1\} \in A$ and $C_j = \{5\} \in B$, in each process we have to consider the third case (line 10) in PADDING. As the critical classes are singletons we have to take $s_1 = 1$ and $s_2 = 5$ in when we call FIXIT(A, 1, B, 5).

There are now for each of the states $s_1 = 1$ and $s_2 = 5$ two blocks they can make a move to, namely $B_k = \{3, 7\}$ and $B_l = \{2, 4, 6, 8\}$, but with different probabilities. We have now to chose in FIXIT (lines 10 to 13) representative states for these two blocks in each of the two processes A and B. In our example these states are uniquely determined, $t_{1k} = 3$, $t_{1l} = 2$, $t_{2k} = 7$, and $t_{2l} = 6$. Furthermore, we have to consider the following sub-trees of A and B which are rooted in these representative states:



We assume w.l.o.g. p > q, i.e. $\min(p,q) = q$ and $\min(1-p, 1-q) = 1-p$ (the other case is symmetric) and thus $r_{1k} = r_{2l} = p-q$. The compensation tree X_{kl}^1 for A thus must fulfill $X_{kl}^1 \sim_{io} T_{1l}$ and for Y_{kl}^2 in B we have $Y_{kl}^2 \sim_{io} T_{2k}$ and furthermore $X_{kl}^1 \sim_{tb} Y_{kl}^2$. In order to construct X_{kl}^1 and Y_{kl}^2 we thus just have to call (recursively) PADDING (T_{1k}, \hat{T}_{1l}) and PADDING (\hat{T}_{2k}, T_{2l}) .

We therefore have to compare twice two trees in order to obtain a padded version (calling PADDING recursively):



A dotted transition denotes a ticked or depleted transition: it means that no computation takes place during this transition (i.e. the state does not change) only time passes. Empty nodes \circ denote copies of the dummy state: they can be seen as place-holders that inherit the state from the previous node. Obviously these two sub-trees X_{kl}^1 and Y_{kl}^2 fulfill the needed requirements regarding bisimilarity and I/O equivalence.

• Finally we have to introduce these compensatory sub-trees into the transformed versions of A and B to obtain (with $r = 1 - \min(p, q) - \min(1 - p, 1 - q)$):



6 Correctness of Padding

In order to show that PADDING (T_1, T_2) indeed produces PTS's T'_1 and T'_2 which are (i) I/O-equivalent to T_1 and T_2 respectively, and (ii) time bisimilar, we first describe the effect of various operations performed in PADDING on the observables and (non)bisimilar states.

Given two states s_1 and s_2 in two tree-subPTS's T_1 and T_2 respectively and a bisimulation relation ~ on the union $T = T_1 \oplus T_2$ then we define new subPTS's $T_{s_1}^{\min}$ and $T_{s_2}^{\min}$ as a common minimisation of the transition probabilities of s_1 and s_2 by changing the transition probabilities from each of the two states to any class C_k of the bisimulation relation ~ and all labels α as follows:

$$p'(s_1, \alpha, C_k) = \min(p(s_1, \alpha, C_k), p(s_2, \alpha, C_k))$$

$$p'(s_2, \alpha, C_k) = \min(p(s_1, \alpha, C_k), p(s_2, \alpha, C_k))$$

$$p'(s_i, \alpha, s_j) = p(s_i, \alpha, s_j) \text{ for } i \notin \{1, 2\}$$

where $p(\cdot)$ denotes the probabilities in T and $p'(\cdot)$ the probabilities in $T_{s_j}^{\min}$, j = 1, 2.

Note that, since it is possible to change the accumulated probabilities $p(s, \alpha, C)$ from states to classes in various ways by adjusting the state-to-state probabilities $p(s, \alpha, s')$, the common minimisation is not unique.

Lemma 20 Given a state s_1 in T_1 and a state s_2 in T_2 and a bisimulation relation \sim on $T = T_1 \oplus T_2$, then the common minimisation $T_{s_1}^{\min}$ and $T_{s_2}^{\min}$ are such that $s_1 \sim s_2$.

PROOF. This follows directly from the definition of $p'(\cdot)$. \Box

Consider two tree-subPTS's, T and S, and a state t in T. The insertion of S into T at t with probability q and label λ is the tree-subPTS $T \triangleleft_{a;\lambda}^t S$ with

$$p'(t_i, \alpha, t_j) = p(t_i, \alpha, t_j)$$
 for $t_i, t_j \in T$ and all labels α
 $p'(s_k, \alpha, s_l) = p(s_k, \alpha, s_l)$ for $s_i, s_j \in S$ and all labels α
 $p'(t, \lambda, s_0) = q$

where s_0 is the root of S, $p(\cdot)$ denotes the probabilities in T and S, and $p'(\cdot)$ the probabilities in $T \triangleleft_{a:\alpha}^t S$.

A special case of insertion is the extension of a tree with a dummy state r, which is used in the padding algorithm to deal with the case of a critical block whose intersection with one of the two tree-PTS's is empty. In this case, T_2 is the degenerated tree with the new state r as the only state, and the label on the transition from r to S_2 is $\lambda = e$. The resulting tree-PTS, $r \triangleleft_{1:e}^{t_0} S$, where t_0 is the root of S, is in this case time bisimilar (but in general not bisimilar) to $T_1 \triangleleft_{a;\lambda}^{t_1} S_1$.

The insertion of sub-trees changes the observables as follows.

Lemma 21 Given two tree-subPTS's, T and S, and a state t in T, let $\mathcal{O}(T)$ and $\mathcal{O}(S)$ be the observables of T and S and π_0 be the (unique) path from the root of T to t. Then the observables of the insertion of S into T with probability p on action α has the following observables:

$$\mathcal{O}(T \triangleleft_{p:\alpha}^{t} S) = \mathcal{O}(T) + p \ (\pi_{0}\alpha \cdot \mathcal{O}(S)),$$

i.e. the observables $\mathcal{O}(T)$ of T plus the "p-weighted" paths in $\pi_0 \alpha \cdot \mathcal{O}(S)$.

As a special case, for the extension of a tree with a dummy state $\mathcal{O}(r \triangleleft_{1:e}^{t_0} S)$, we have:

$$\mathcal{O}(r \triangleleft_{1:e}^{t_0} S) = \mathcal{O}(r) + 1 \ (e \cdot \mathcal{O}(S)) = \mathcal{O}(S),$$

i.e. the original observables are not changed.

Let T be a tree-subPTS and let S be a sub-tree-subPTS of T. Then the partial deletion of S in T, denoted by $T \not\triangleleft_{p'}^p S$, is the tree-subPTS on the states of T obtained by changing the entry probability into S from T, i.e. the probability with which the root of S can be reached from $T \setminus S$, from p to p'. For p' = 0, this operation corresponds to the deletion of S in T.

The (partial) deletion of a sub-tree changes the observables as follows.

Lemma 22 Given a tree-subPTS T and a sub-tree-subPTS S of T with observables $\mathcal{O}(T)$ and $\mathcal{O}(S)$ respectively, and a (unique) path π_0 from the root of T to the root of S, then the observables of $T \not\triangleleft_{p'}^p S$ are given by:

$$\mathcal{O}(T \not\triangleleft_{p'}^p S) = \mathcal{O}(T) - (p - p')(\pi_0 \cdot \mathcal{O}(S)).$$

PROOF. Each path reaching a leaf through S is not completely removed but only its impact on the observables is reduced. In T the probability is a product of the form $\prod_k p_k \cdot p \cdot \prod_l p_l$ where the p_k indicate the computational steps in π_0 , the probability p is the original entry probability into S, and p_l are the probabilities in S. In $T \not\triangleleft_{p'}^p S$ the probability of this path is changed to: $\prod_k p_k \cdot p' \cdot \prod_l p_l$. If we therefore remove the contribution of these paths in the original observables and add the contribution in the new subPTS we obtain the desired result: $\mathcal{O}(T \not\triangleleft_{p'}^p S) = \mathcal{O}(T) - p\pi_0 \cdot \mathcal{O}(S) + p'\pi_0 \cdot \mathcal{O}(S) =$ $\mathcal{O}(T) - (p-p')\pi_0 \cdot \mathcal{O}(S)$. If p' = 0 then the subtree S is effectively eliminated and so are all the paths through S to the leaves. As the observables are distributions over paths we simply have to eliminate the contribution of these paths through S from the observables of T. The contribution of these paths is not just their "S-stage" but has to be prefixed by the contribution until S, i.e. its root, is reached, this is given by the unique path π_0 . \Box

We can now show the main theorem establishing that the padding is correct.

Theorem 23 Given two tree-PTS's T_1 and T_2 then PADDING (T_1, T_2) returns the tree-PTS's T'_1 and T'_2 such that $T'_1 \sim_{tb} T'_2$, $T'_1 \sim_{io} T_1$, and $T'_2 \sim_{io} T_2$.

PROOF.

The proof is by induction on the layers in the union $T_1 \oplus T_2$. We assume that k is the number of layers, i.e. $k = max\{\text{HEIGHT}(T_1), \text{HEIGHT}(T_2)\}$. We will use the notation T_i^n , i = 1, 2, to indicate the *n*-stage transformation of T_i : this is the tree-PTS formed by the *n* layer top of T_i and the *n* layer cut-off of T'_i , that is an intermediate version of T_i and T'_i where the *n* bottom layers are already padded but the top still needs to be fixed. Thus, $T'_1 = T_1^k$ and $T'_2 = T_2^k$. We will also use the notation $s \bigtriangleup T$ to indicate that the state *s* has a transition to some classes of the partition T on some action with some probability.

The algorithm PADDING progresses through the trees T_1 and T_2 layer by layer starting from the terminal states or leaves (base case). The sub-routine MAXLUMPING tries to identify non-bisimilar states in T_1 and T_2 . When the procedure enters the layer n + 1 then the *n*-cutoffs are time bisimilar, i.e. $T_1|_n \sim_{tb} T_2|_n$ (induction hypothesis). Moreover, we will show that at each layer $n, T_1^n \sim_{io} T_1$, and $T_2^n \sim_{io} T_2$, which implies the second part of the theorem's claim.

Calling the sub-routine MAXLUMPING has three possible outcomes: (i) no mismatches (i.e. critical blocks) are found, (ii) there is only one mismatch state on level n + 1, i.e. one of the two trees is higher, or (iii) there are two states s_1 and s_2 in T_1 and T_2 respectively which are not bisimilar. If case (i) holds, then T_1 and T_2 are already bisimilar and the algorithm terminates. So we prove the correctness of the algorithm in the other two cases.

k = 0 Since all states at layer 0 are leaves (there are no outgoing transitions), clearly, we have that $T_1|_0 \sim_{tb} T_2|_0$, and $T_1^0 \sim_{io} T_1$, and $T_2^0 \sim_{io} T_2$.

 $k \rightarrow k+1$ In the general case (iii) the procedure FIXIT() is called which proceeds in two phases: In the first phase the procedure constructs a common minimisation $T_{s_1}^{min}$ and $T_{s_2}^{min}$ of $s_1 \triangle T_1|_n$ and $s_2 \triangle T_2|_n$ with respect to two non bisimilar states s_1 and s_2 at level n + 1. In the bisimulation relation constructed by calling MAXLUMPING, the partition of the states in $s_1 \Delta T_1|_n$ and $s_2 \triangle T_2|_n$ is such that $s_1 \not\sim_{tb} s_2$. However, by Lemma 20 we have that after the common minimisation s_1 and s_2 are now bisimilar. As a result, the number of non-bisimilar states at layer n+1 is strictly reduced. We observe that case (ii) is the special case of (iii) in which either s_1 or s_2 is a dummy state d and the transition to the n layer cut-off is labelled by e. In phase two FIXIT() inserts subtrees S_1 and S_2 in $T_{s_1}^{min}$ and $T_{s_2}^{min}$ respectively. Since these subtrees are constructed via the padding procedure on copies and depleted versions of subtrees S'_1 and S'_2 of T_1 and T_2 of height smaller than n, by the induction hypothesis they are time bisimilar. This guarantees that we still have $T'_1|_n \sim_{tb} T'_2|_n$, where $T'_i|_n$, j = 1, 2, is the n layer cut-off after the insertion of subtrees S_1 and S_2 . Moreover, by Lemma 22 and Lemma 21 we have

$$\mathcal{O}(T_1^{n+1}) = \mathcal{O}(T_{s_1}^{\min} \triangleleft_{p_1 - \min(p_1, p_2)}^{s_1} S_1)$$

= $\mathcal{O}(T_1) - (p_1 - \min(p_1, p_2)) \cdot \mathcal{O}(S_1) + (p_1 - \min(p_1, p_2) \cdot \mathcal{O}(S_1))$
= $\mathcal{O}(T_1).$

and similarly for T_2^{n+1} .

As we have only finitely many states in a layer we will eventually reach the situation where there are no non-bisimilar states in layer n+1 or more generally in the n+1 cut-off. The algorithm therefore terminates in each layer and as we also have only finitely many layers the algorithm terminates with $T'_1 \sim_{tb} T'_2$, $T'_1 \sim_{io} T_1$, and $T'_2 \sim_{io} T_2$. \Box

7 Comparisons and Related Work

7.1 Comparison with Agat's transformation

In [2] Agat proposes a transformation technique in the context of a type system with security types. The type system, besides detecting illegal information flow, transforms conditional statements that branch on high-security data into new statements where both conditional branches have been made bisimilar, thus making the new statement immune to timing attacks and the associated high-security data safe. The conditional statement is transformed by adding a 'dummy copy' of each branch to its counterpart such that, when the new branches are compared against each other, the original code is matched against its dummy copy. If we adapt this technique to PTS's and apply it to the processes A and B considered before, we get the transformed processes shown below:



According to Agat's transformation, a 'dummy execution' of B should follow A and a 'dummy execution' of A should precede B. Given that A has two final states, two dummy copies of B are appended to process A. Similarly, a dummy copy of A is put before B. However, in order to preserve the tree structure of the process, a new copy of B is made so it can be joined to one of the final states of the dummy copy of A.

Clearly the two transformed trees are time bisimilar and the new processes are I/O-equivalent to their original versions. It is also clear that this transformation not only creates a greater number of states than our proposed solution but, more importantly, generates processes with longer execution time. The work by Agat [2] is certainly the most closely related to ours; in fact we are not aware of other approaches to closing timing leaks which exploit program transformation techniques. Apart from the basic differences between our transformation algorithm and the Agat transformation type system explained in the previous section, our approach is in some sense more abstract than [2] in that it does not address a specific language; rather it is in principle adaptable to any probabilistic language whose semantics can be expressed in terms of our PTS model.

Timing attacks are prominently studied in security protocol analysis, and in general in the analysis of distributed programs specifically designed to achieve secure communication over inherently insecure shared media such as the Internet. In this setting, timing channels represent a serious threat as shown for example in [17] where a timing attack is mounted on RSA encryption, and in [18] where it is shown how by measuring the time required by certain operation a web site can learn information about the user's past activities; or in [19] where an attack is described over Abadi's private authentication protocol [20].

The various approaches which have been proposed in the literature for the time analysis of security and cryptographic protocols mainly exploit languages and tools based on formal methods (timed automata, model checkers and process algebra). For example, [21] develop formal models and a timed automata based analysis for the Felten and Schneider's web privacy attack mentioned above. A process algebraic approach is adopted in [13] and [22] where the problem of timing information leakage is modelled as non-interference property capturing some time-dependent information flows. A different approach is the one adopted in [23] where static analysis techniques are used to verify communication protocols against timing attacks.

8 Conclusion and Further Work

In this paper we have investigated possible countermeasures against timing attacks in a process algebraic setting. The aim is to transform systems such that their timing behaviour becomes indistinguishable – i.e. they become bisimilar – while preserving their computational content – i.e. I/O behaviour. Our particular focus in this work was on finite, tree-like, probabilistic systems and probabilistic bisimulation.

Simple obfuscation methods have been considered before, e.g. by Agat [2],

which may lead to a substantial increase of the (average) running time of the transformed processes. Our approach attempts to minimise this additional running time overhead introduced by the transformation. The padding algorithm achieves this aim by patching time leaks 'on demand' whenever the lumping algorithm which aims to establish the bisimilarity of two processes encounters a critical situation. Further work will be directed towards improving the padding algorithm further. At various stages of the algorithm we make non-deterministic choices (of certain states or classes); it will be important to investigate strategies to achieve optimal choices.

We also plan to investigate probabilistic and conditional padding. The current algorithm introduces time leak fixes whenever they appear to be necessary. The resulting processes are therefore perfectly bisimilar. However, one could also think of applying patches randomly or only if the expected time leak exceeds a certain threshold. One would then obtain approximate or ε -bisimilar systems – in the sense of our previous work [7,1] – but could expect a smaller increase of the (average) running time and/or the size of the system or program considered. Optimising the trade-off between vulnerability against timing attacks – measured quantitatively by ε – and the additional costs – running time, system size etc. – requires a better understanding of the relation between these different factors. To this aim we believe we can exploit well established techniques and results from the fields of decision theory and non-linear optimisation.

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