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The Complexity of Coverability in ν -Petri Nets*

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Abstract

We show that the coverability problem in ν -Petri nets is complete for 'double Ackermann' time, thus closing an open complexity gap between an Ackermann lower bound and a hyper-Ackermann upper bound. The coverability problem captures the verification of safety properties in this nominal extension of Petri nets with name management and fresh name creation. Our completeness result establishes ν -Petri nets as a model of intermediate power among the formalisms of nets enriched with data, and relies on new algorithmic insights brought by the use of well-quasi-order ideals.

Categories and Subject Descriptors F.2.2 [Analysis of Algorithms and Problem Complexity]: Nonnumerical Algorithms and Problems

Keywords Well-structured transition system, formal verification, well-quasi-order, order ideal, fast-growing complexity

1. Introduction

ν-Petri nets (*ν*PN) generalise Petri nets by decorating tokens with data values taken from some infinite countable data domain \mathbb{D} . These values act as pure names: they can only be compared for equality or non-equality upon firing transitions; *ν*PNs have furthermore the ability to create *fresh* data values, never encountered before in the history of the computation. Such systems were introduced to model distributed protocols where process identities need to be taken into account (Rosa-Velardo and de Frutos-Escrig 2008, 2011), and form a restricted class of data-centric dynamic systems (Montali and Rivkin 2016). They also coincide with a restriction of the π -calculus to processes of 'depth 1' as defined by Meyer (2008), while their *polyadic* extension, which allows to manipulate tuples of tokens, is equivalent to the full π -calculus (Rosa-Velardo and Martos-Salgado 2012)—and Turing-complete.

In spite of their high expressiveness, ν PNs fit in the large family of Petri net extensions among the *well-structured* ones (Abdulla et al. 2000; Finkel and Schnoebelen 2001). As such, they still enjoy decision procedures for several verification problems, prominently safety (through the *coverability* problem) and termination. They share these properties with the other extensions of Petri nets with

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data defined by Lazić, Newcomb, Ouaknine, Roscoe, and Worrell (2008), but are something of an intermediate model. Indeed, as shown in Figure 1, they extend *unordered Petri data nets* with the ability to create fresh data values, but in turn this ability can be simulated (as far as the coverability problem is concerned) by either *ordered data Petri nets*—where $\mathbb D$ is equipped with a dense linear ordering—or *unordered data nets*—where 'whole-place operations' allow to transfer, duplicate, or destroy the entire contents of places.

The Power of Well-Structured Systems. This work is part of a general program that aims to understand the expressive power and algorithmic complexity of well-structured transition systems (WSTS), for which the complexity of the coverability problem is a natural proxy. Besides the intellectual satisfaction one might find in classifying the worst-case complexity of this problem, we hope indeed to gain new insights into the algorithmics of the systems at hand, and into their relative 'power.' A difficulty is that the generic backward coverability algorithm developed by Abdulla, Čerāns, Jonsson, and Tsay (2000) and Finkel and Schnoebelen (2001) to solve coverability in WSTS relies on well-quasi-orders (wqos), for which complexity analysis techniques are not so widely known.

Nevertheless, in a series of recent papers, the exact complexity of coverability for several classes of WSTS has been established. These complexities are expressed using ordinal-indexed fast-growing complexity classes $(\mathbf{F}_{\alpha})_{\alpha}$ (Schmitz 2016), e.g. 'Tower' complexity corresponds to the class \mathbf{F}_3 and is the first non elementary complexity class in this hierarchy, 'Ackermann' corresponds to \mathbf{F}_{ω} and is the first non primitive-recursive class, 'hyper-Ackermann' to ${\bf F}_{\omega}{}^{\omega}$ and is the first non multiply-recursive class, etc. (see Figure 4). To cite a few of these complexity results, coverability is \mathbf{F}_{ω} -complete for reset Petri nets and affine nets (Schnoebelen 2010; Figueira et al. 2011), $\mathbf{F}_{\omega^{\omega}}$ -complete for lossy channel systems (Chambart and Schnoebelen 2008; Schmitz and Schnoebelen 2011) and unordered data nets (Rosa-Velardo 2014), and even higher complexities appear for timed-arc Petri nets and ordered data Petri nets ($\mathbf{F}_{\omega^{\omega}}$ -complete, see Haddad et al. 2012) and priority channel systems and nested counter systems ($\mathbf{F}_{\varepsilon_0}$ -complete, see Haase et al. 2014; Decker and Thoma 2016); see the complexities in violet in Figure 1 for the Petri net extensions related to νPNs .

All those results rely on the same general template (see Schmitz and Schnoebelen 2013, for a gentle introduction):

- 1. for the upper bound, a *controlled bad sequence* can be extracted from any run of the backward coverability algorithm, and in turn the length of this sequence can be bounded using a *length function theorem* for the wqo at hand (e.g., Cichoń and Tahhan Bittar 1998; Figueira et al. 2011; Schmitz and Schnoebelen 2011; Rosa-Velardo 2014, for the mentioned results);
- for the lower bound, weak computers for Hardy functions and their inverses are implemented in the formalism at hand, allowing to build a working space on which a Turing or Minsky machine can be simulated.

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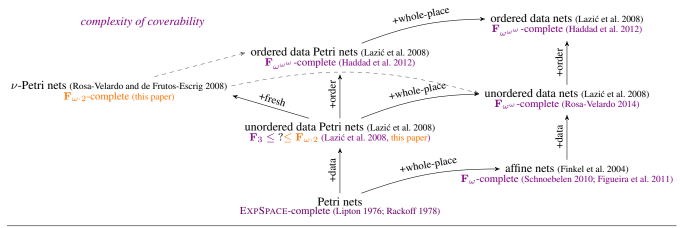


Figure 1. A short taxonomy of some data enrichements of Petri nets. Complexities in violet refer to the already known complexities for the coverability problem; the exact complexity in unordered data Petri nets is unknown at the moment. As indicated by the dashed arrows, freshness can be enforced using a dense linear order or whole-place operations.

Contributions. In this paper, we pinpoint the complexity of coverability in ν PNs by showing that it is complete for $\mathbf{F}_{\omega\cdot 2}$, i.e. for 'double Ackermann' complexity. This solves an open problem: the best known lower bound was \mathbf{F}_{ω} , from a reduction from coverability in reset Petri nets (Rosa-Velardo and de Frutos-Escrig 2011), while the best known upper bound was $\mathbf{F}_{\omega^{\omega}}$ from the more general case of unordered data nets (Rosa-Velardo 2014), leaving a considerable complexity gap.

We believe this $\mathbf{F}_{\omega\cdot 2}$ -completeness is remarkable on two counts. First, this is the first instance of a 'natural' decision problem complete for an intermediate complexity class between Ackermann and hyper-Ackermann. Second, the usual template for such complexity results, summed up in points 1 and 2 above, *fails* for νPNs , in the sense that all it could prove are the aforementioned \mathbf{F}_{ω} lower bound and $\mathbf{F}_{\omega^{\omega}}$ upper bound. As a result, we had to design new techniques, which we think are of independent interest.

These new techniques are inspired by another case where the template in 1 and 2 fails, namely that of Petri nets. Indeed, coverability in Petri nets is EXPSPACE-complete, as shown by Rackoff (1978) for the upper bound and by Lipton (1976) for the lower bound. These results however do not rely on wqos and are quite specific to Petri nets, and their generalisation to a formalism as rich as ν PNs required new insights:

For the upper bound, we analyse the complexity of the backward coverability algorithm when seen dually as computing a decreasing sequence of *downwards-closed* sets. Such sets can be represented as finite unions of *ideals* (Bonnet 1975; Finkel and Goubault-Larrecq 2009); see Section 3.

We have recently shown that, for Petri nets, this dual view allows to exhibit an invariant on the ideals appearing during the course of the execution of the backward coverability algorithm, which in turn yields a dramatic improvement on its complexity analysis from \mathbf{F}_{ω} to 2ExpTIME (Lazić and Schmitz 2015). The same bound had already been established by Bozzelli and Ganty (2011) using Rackoff's analysis, but this new viewpoint is applicable to any WSTS with effective ideal representations, and enables us to proceed along similar lines in Section 4 and to obtain the desired $\mathbf{F}_{\omega \cdot 2}$ upper bound.

 For the lower bound, we follow the pattern of Lipton's proof, in that we design an 'object-oriented' implementation of the double Ackermann function in νPNs. By this, we mean that the implementation provides an interface with increment, decrement, zero, and max operations on larger and larger *counters* up to a double Ackermannian value. This allows then the simulation of a Minsky machine working in double Ackermann space and establishes the matching $\mathbf{F}_{\omega \cdot 2}$ lower bound.

The basic building blocks of this development are Ackermannian counters reminiscent of the construction of Schnoebelen (2010) for reset Petri nets. The catch is that we need to be able to mimick this construction for non-fixed dimensions and to combine it with an iteration operator—of the kind employed recently by Lazić et al. (2016) in the context of channel systems with insertion errors to show Ackermann-hardness—, which led us to develop delicate indexing mechanisms by data values; see Section 5.

We assume the reader is already familiar with the basics of Petri nets, and start with the formal definition of ν PNs and of their semantics in the upcoming Section 2. Due to space constraints, some technical material and proofs will be found in the full version of the paper, available from https://hal.inria.fr/hal-01265302/.

2. ν -Petri Nets

We define the syntax of ν PNs exactly like Rosa-Velardo and Martos-Salgado (2012). Their semantics can be stated in terms of finitely supported partial maps from an infinite data domain $\mathbb D$ to markings in $\mathbb N^P$, telling for each data value how many tokens with that value appear in each place. However, we find it easier to work with a slightly more abstract but equivalent *multiset semantics*, which accounts for the fact that the semantics is invariant under permutations of the data domain $\mathbb D$, and eschews any explicit reference to this data domain; see Section 2.2. Following Rosa-Velardo and Martos-Salgado (2012), we also illustrate the expressive power of ν PNs in Section 2.3 by showing how they can implement reset Petri nets.

2.1 Finite Multisets

Let A be a set. Consider the *commutation* equivalence \sim over finite sequences in A^* : this is the transitive reflexive closure $\sim \stackrel{\text{def}}{=} \sim_1^*$ of the relation \sim_1 defined by $uabv \sim_1 ubav$ for all $u,v \in A^*$ and $a,b \in A$. We define (finite) multisets as \sim -equivalence classes of A^* , and write $A^{\otimes} \stackrel{\text{def}}{=} A^*/\sim$ for the set of multisets over A.

We manipulate a multiset through any of its representatives in A^* , e.g. [aab] = [aba] = [baa] are all equal to the \sim -equivalence class $\{aab, aba, baa\}$. We write accordingly '[]' for the empty

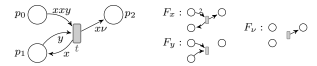


Figure 2. A ν PN and the associated flows of x, y, and ν .

multiset. Note that this viewpoint matches the definition of a finite multiset as a *finitely supported* function $m: A \to \mathbb{N}$, i.e. such that its $\begin{array}{l} \textit{support } \textit{Supp}(m) \stackrel{\text{\tiny def}}{=} \{a \in A \mid m(a) \neq 0\} \text{ is finite. For instance,} \\ \textit{Supp}([aab]) = \{a,b\} \text{ and } [aab](a) = 2 \text{ and } [aab](b) = 1. \text{ The} \end{array}$ length |m| of a multiset m is the length of any representative and satisfies $|m| = \sum_{a \in A} m(a) = \sum_{a \in Supp(m)} m(a)$ in the functional view.

Sums. Given two multisets m and m' over a set A, their sum (also called their *union*) $m \oplus m'$ is represented by the concatenation of their representatives. From the functional viewpoint, $(m \oplus m')(a) =$ m(a) + m'(a) for all $a \in A$, with length $|m \oplus m'| = |m| + |m'|$ and support $Supp(m \oplus m') = Supp(m) \cup Supp(m')$.

Embeddings. Assume (A, \leq_A) is a quasi-order (qo), i.e. that A is equipped with a reflexive transitive relation $\leq_A \subseteq A \times A$. An embedding from a multiset $m = [a_1 \cdots a_{|m|}]$ into a multiset $m' = [a'_1 \cdots a'_{|m'|}]$ is an injective function $e: \{1, \dots, |m|\} \rightarrow$ $\{1,\ldots,|m'|\}$ such that $a_i \leq_A a'_{e(i)}$ for all $1 \leq i \leq |m|$. Given such an e, we can decompose m' in a unique manner as $m'' \oplus [a'_{e(1)} \cdots a'_{e(|m|)}]$ for some m''. Note that in general $m \neq [a'_{e(1)} \cdots a'_{e(|m|)}]$, unless \leq_A is the equality relation over A. We say that m' embeds m and write $m \sqsubseteq m'$ if there exists an

embedding e from m to m'; observe that $(\overline{A}^{\otimes}, \sqsubseteq)$ is also a qo.

Markings. Let (P, =) be a finite set ordered by equality. We call a vector $m \in \mathbb{N}^P$ a marking. Markings can be added pointwise by $(\boldsymbol{m}+\boldsymbol{m}')(p)\stackrel{\text{def}}{=} \boldsymbol{m}(p)+\boldsymbol{m}'(p)$ for all $p\in P$, and compared using the *product ordering* $\boldsymbol{m}\leq \boldsymbol{m}'$, holding iff $\boldsymbol{m}(p)\leq \boldsymbol{m}'(p)$ for all $p\in P$. Note that $(\mathbb{N}^P,+,\leq)$ is isomorphic to $(P^\circledast,\oplus,\sqsubseteq)$, but we shall use the former to avoid confusion with other multisets.

2.2 Syntax and Semantics

Let \mathcal{X} and Υ be two disjoint infinite countable sets of *non-fresh variables* and *fresh variables* respectively, and let $Vars \stackrel{\text{def}}{=} \mathcal{X} \uplus \Upsilon$.

Syntax. A ν -Petri net is a tuple $N = \langle P, T, F \rangle$ where P is a finite non-empty set of places, T is a finite set of transitions disjoint from P, and $F: (P \times T) \cup (T \times P) \rightarrow Vars^{\otimes}$ is a *flow* function.

For any transition $t \in T$, let $InVars(t) \stackrel{\text{def}}{=} \bigcup_{p \in P} Supp(F(p,t))$ and $OutVars(t) \stackrel{\text{def}}{=} \bigcup_{p \in P} Supp(F(t, p))$ denote its sets of input and output variables respectively, and $Vars(t) \stackrel{\text{def}}{=} InVars(t) \cup$ OutVars(t); we require that

- 1. fresh variables are never input variables: $\Upsilon \cap InVars(t) = \emptyset$,
- 2. all the non-fresh output variables are also input variables: $OutVars(t) \cap \mathcal{X} \subseteq InVars(t).$

Writing $\mathcal{X}(t) \stackrel{\text{def}}{=} Vars(t) \cap \mathcal{X}$ and $\Upsilon(t) \stackrel{\text{def}}{=} Vars(t) \cap \Upsilon$, this entails $\mathcal{X}(t) = InVars(t) \text{ and } \Upsilon(t) = OutVars(t) \cap \Upsilon.$

For a variable $x \in Vars$, the flow of x is the function F_x : $(P \times$ $T) \cup (T \times P) \rightarrow \mathbb{N}$ defined by $F_x(p,t) \stackrel{\text{def}}{=} F(p,t)(x)$ and $F_x(t,p) \stackrel{\text{def}}{=} F(t,p)(x)$. When we fix a transition $t \in T$, we see $F_x(P,t)$ and $F_x(t,P)$ as markings in \mathbb{N}^P . Intuitively, a νPN synchronises a potentially infinite number of Petri nets acting on the same places and transitions. See Figure 2 for a depiction; as usual with Petri nets, places are represented by circles, transitions by rectangles, and non-null flows by arrows labelled with their values.

We define the size of a ν PN as $|N| \stackrel{\text{def}}{=} \max(|P|, |T|, \sum_{p,t} |F(p,t)| +$ |F(t,p)|) (this corresponds to a unary encoding of the coefficients in the multisets defined by F).

Multiset Semantics. A νPN defines an infinite transition system $\langle Confs, \rightarrow \rangle$ where $Confs \stackrel{\text{def}}{=} (\mathbb{N}^P)^{\otimes}$ is the set of *configurations* and $\rightarrow \subseteq Confs \times Confs$ is called the *step* relation.

Let us associate with any transition $t \in T$ two multisets of markings, in $(\mathbb{N}^P)^{\otimes}$, of inputs and fresh outputs respectively:

$$\operatorname{in}(t) \stackrel{\text{def}}{=} \bigoplus_{x \in \mathcal{X}(t)} [F_x(P,t)] \,, \qquad \operatorname{out}_{\Upsilon}(t) \stackrel{\text{def}}{=} \bigoplus_{\nu \in \Upsilon(t)} [F_\nu(t,P)] \,. \tag{1}$$

Given a configuration $M = [m_1 \cdots m_{|M|}]$, we say that t is *fireable* from M if there exists an embedding from in(t) into M, which here can be seen as an injective function $e: \mathcal{X}(t) \to \{1, \dots, |M|\}$ with $F_x(P,t) \leq m_{e(x)}$ for all $x \in \mathcal{X}(t)$. We call such an e a mode for t and M; given t and M there are finitely many different modes.

A mode e for t and M defines a step: it uniquely determines two configurations M' and M'' such that

$$M = M'' \oplus \bigoplus_{x \in \mathcal{X}(t)} [m_{e(x)}], \qquad (2)$$

$$M = M'' \oplus \bigoplus_{x \in \mathcal{X}(t)} [\boldsymbol{m}_{e(x)}], \qquad (2)$$

$$M' = M'' \oplus \operatorname{out}_{\Upsilon}(t) \oplus \bigoplus_{x \in \mathcal{X}(t)} [\boldsymbol{m}'_{e(x)}], \qquad (3)$$

where for all $x \in \mathcal{X}(t)$, $\boldsymbol{m}'_{e(x)} \stackrel{\text{def}}{=} \boldsymbol{m}_{e(x)} - F_x(P,t) + F_x(t,P)$. We write $M \xrightarrow{e,t} M'$ in such a case. We write as usual $M \xrightarrow{t} M'$ if there exists e for t and M such that $M \xrightarrow{e,t} M'$, and $M \to M'$ if there exists $t \in T$ such that $M \xrightarrow{t} M'$. In other words, the transition t:

- ullet applies F_x for each non-fresh variable $x \in \mathcal{X}(t)$ to a different individual marking $m_{e(x)} \ge F_x(P,t)$ of M, replacing it with the marking $m'_{e(x)}$,
- leaves the remaining markings in M'' untouched, and
- furthermore adds new markings $F_{\nu}(t, P)$ for each fresh variable $\nu \in \Upsilon(t)$ to the resulting configuration.

Example 1. Consider the transition t in Figure 2 acting on P = $\{p_0, p_1, p_2\}$ and a configuration $M = [\boldsymbol{m}_1 \boldsymbol{m}_2 \boldsymbol{m}_3]$ where $\boldsymbol{m}_1 =$ (2,1,1), $m_2 = (2,0,0)$, and $m_3 = (1,1,0)$.

We have in(t) = [(2,0,0)(1,1,0)] and $out_{\gamma}(t) = [m_4]$ with $m_4 = (0,0,1)$, and three possible modes. We can have $e_1(x) = m_1$, resulting in $m'_1 = (0, 2, 2)$, and $e_1(y) = m_3$, resulting in $m_3' = (0,0,0)$, hence $M \xrightarrow{e_1,t} [m_1'm_2m_3'm_4]$. Another possibility is to have $e_2(x) = m_2$ yielding $m_2' = (0,1,1)$ and $e_2(y) = m_1$ yielding $m_1'' = (1,0,1)$, showing that $M \xrightarrow{e_2,t} [m_1''m_2'm_3m_4]$, and a last possibility is to have $e_3(x) = m_2$ and $e_3(y) = m_3$, resulting in a step $M \xrightarrow{e_3,t} [m_1m_2'm_3'm_4]$.

2.3 Example: Reset Petri Nets

Rosa-Velardo and Martos-Salgado (2012) show that νPNs are able to simulate reset Petri nets, an extension of Petri nets with special arcs that empty a place upon firing a transition. A remarkable aspect of the construction we are going to sketch here is that three places and a simple addressing mechanism are enough to simulate reset Petri nets with an arbitrary number of places—recall that the latter have an Ackermannian-hard coverability problem (Schnoebelen 2010). This explains why we will be able to push the lower bound beyond Ackermann-hardness in Section 5, where we design more involved addressing mechanisms.

Given any reset Petri net with places $P = \{p_0, \dots, p_{n-1}\}$, we build a ν PN with three places a, \bar{a} , and v. The intuition is for a

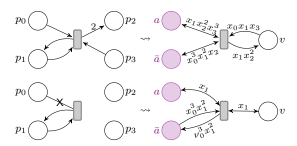


Figure 3. Examples of simulations of reset Petri net transitions (left) by a ν PN (right).

and \bar{a} to maintain an *addressing* mechanism for the original places in P, while v maintains the actual token counts of the original net. The places a and \bar{a} use n different data values, each with distinct counts of tokens; more precisely, all the reachable configurations M are of the form $[m_0 \cdots m_{n-1}] \oplus M'$ where $m_i(a) = i$ and $m_i(\bar{a}) = n - 1 - i$ for all $0 \le i < n$, and all the markings m in M' are *inactive*, i.e. with $m(a) + m(\bar{a}) < n - 1$. Each active marking m_i simulates the place p_i of the original net by holding in $m_i(v)$ the number of tokens in place p_i .

For instance, the top of Figure 3 shows how a transition of a Petri net with 4 places (on the left) can be simulated with this construction (on the right). The flows of each variable x_0, x_1, x_2, x_3 with places a and \bar{a} identify uniquely the places p_0, p_1, p_2, p_3 of the original net, while the flows with place v update the token counts accordingly.

The interest of this addressing mechanism is that it allows to simulate reset transitions, like the one on the bottom left of Figure 3 that empties place p_0 upon firing. This is performed by creating a fresh marking \mathbf{m}'_0 with $\mathbf{m}'_0(a) = 0$, $\mathbf{m}'_0(\bar{a}) = 3$, and $\mathbf{m}'_0(v) = 0$; after the transition step, we will have $\mathbf{m}_0(a) = \mathbf{m}_0(\bar{a}) = 0$ and \mathbf{m}_0 might still have some leftover tokens in v, but it is inactive and will be ignored in the remainder of the computation.

3. Backward Coverability

The decision problem we are interested in is *coverability*:

input: a ν PN, and two configurations M_0, M_1

question: does there exist $M \supseteq M_1$ such that $M_0 \to^* M$?

We instantiate in this section the backward coverability algorithm from (Lazić and Schmitz 2015) for νPNs . This algorithm is a dual of the classical algorithm of Abdulla et al. (2000) and Finkel and Schnoebelen (2001): instead of building an increasing chain $U_0 \subsetneq U_1 \subsetneq \cdots$ of upwards-closed sets U_k of configurations that can cover the target M_1 in at most k steps, it constructs instead a decreasing chain $D_0 \supsetneq D_1 \supsetneq \cdots$ of downwards-closed sets D_k of configurations that cannot cover the target in k or fewer steps (see Section 3.3). Like the usual backward algorithm, the termination and correctness of this dual version hinges on the fact that $\langle Confs, \rightarrow, \sqsubseteq \rangle$ is a WSTS (see Section 3.1). We need however an additional ingredient, which is a means of effectively representing and computing our downwards-closed sets D_k of configurations. We rely for this on *ideals* of $(Confs, \sqsubseteq)$, which play the same role as finite bases in the classical algorithm; see Section 3.2.

3.1 ν -Petri Nets are Well-Structured

Well-Quasi-Orders. Let (A, \leq_A) be a qo. Given a set $S \subseteq A$, its *downward-closure* is $\downarrow S \stackrel{\text{def}}{=} \{a \in A \mid \exists s \in S : a \leq_A s\};$ when S is a singleton $\{s\}$ we write more simply $\downarrow s$. A set $D \subseteq A$ is *downwards-closed* (also called *initial*) if $\downarrow D = D$. Upward-closures $\uparrow S$ and upwards-closed subsets $\uparrow U = U$ are defined similarly.

A well-quasi-order (wqo) is a qo (A, \leq_A) where every bad sequence a_0, a_1, \ldots of elements over A, i.e. with $a_i \not\leq_A a_j$ for all i < j, is finite (Higman 1952). Equivalently, it is a qo with the descending chain property: all the chains $D_0 \supsetneq D_1 \supsetneq \cdots$ of downwards-closed subsets $D_j \subseteq A$ are finite. Equivalently, it has the finite basis property: any non-empty subset $S \subseteq A$ has a finite number of minimal elements (and at least one minimal element) up to equivalence.

For instance, any finite set P equipped with equality forms a wqo (P,=): its downwards-closed subsets are singletons $\{p\}$ for $p \in P$, and its chains of downwards-closed sets are of length at most one. Assuming (A, \leq_A) is a wqo, then finite multisets over A provide another instance: $(A^{\circledast}, \sqsubseteq)$ is also a wqo as a consequence of Higman's Lemma. Hence both the sets of markings (\mathbb{N}^P, \leq) and of configurations $(Confs, \sqsubseteq)$ of a νPN are wqos.

Compatibility. The transition system $\langle Confs, \rightarrow \rangle$ defined by a νPN further satisfies a *compatibility* condition with the embedding relation: if $M_1 \sqsubseteq M_1'$ and $M_1 \to M_2$, then there exists $M_2' \supseteq M_2$ with $M_1' \to M_2'$. In other words, \sqsubseteq is a simulation relation on the transition system $\langle Confs, \rightarrow \rangle$. Since $(Confs, \sqsubseteq)$ is a wqo, this transition system is therefore *well-structured* (Abdulla et al. 2000; Finkel and Schnoebelen 2001).

3.2 Effective Ideal Representations

Ideals. Let (A, \leq_A) be a wqo. An *ideal* I of A is a non-empty, downwards-closed, and (up-) *directed* subset of A; this last condition enforces that, if a, a' are in I, then there exists $b \in I$ that dominates both: $a \leq_A b$ and $a' \leq_A b$. For example, looking again at the case of finite sets (P, =), we can see that singletons $\{p\}$ are ideals. In fact, more generally $\downarrow a$ for $a \in A$ is always an ideal of A. But there can be other ideals, e.g. I^{\circledast} is an ideal of A^{\circledast} if I is an ideal of A.

The key property of wqo ideals is that any downwards-closed set D over a wqo has a unique decomposition as a finite union $D=I_1\cup\dots\cup I_n$, where the I_j 's are incomparable for inclusion—this was shown e.g. by Bonnet (1975), and by Finkel and Goubault-Larrecq (2009) in the context of complete WSTS (and generalised to Noetherian topologies). Ideals are also irreducible: if $I\subseteq D_1\cup D_2$ for two downwards-closed sets D_1 and D_2 , then $I\subseteq D_1$ or $I\subseteq D_2$.

Effective Representations. Although ideals provide finite decompositions for downwards-closed sets, they are themselves usually infinite, and some additional effectiveness assumptions are necessary to employ them in algorithms. In this paper, we will say that a wqo (A, \leq_A) has effective ideal representations (see Finkel and Goubault-Larrecq 2009; Goubault-Larrecq et al. 2016, for more stringent requisites) if every ideal can be represented, and there are algorithms on those representations:

- (CI) to check $I \subseteq I'$ for two ideals I and I',
- (II) to compute the ideal decomposition of $I \cap I'$ for two ideals I and I',
- (CU') to compute the ideal decomposition of the residual $A \setminus \uparrow a = \{a' \in A \mid a \not\leq_A a'\}$ for any a in A.

All these effectiveness assumptions are true of the representations for (\mathbb{N}^P,\leq) and $(\mathit{Confs},\sqsubseteq)$ described by Goubault-Larrecq et al. (2016), which we recall next.

Extended Markings. Let $\mathbb{N}_{\omega} \stackrel{\text{def}}{=} \mathbb{N} \uplus \{\omega\}$, where ' ω ' denotes a new top element with $\omega + n = \omega - n = \omega > n$ for all $n \in \mathbb{N}$. An *extended marking* is a vector $e \in \mathbb{N}_{\omega}^{P}$. The product ordering and pointwise sum operations are lifted accordingly. Then the ideals of (\mathbb{N}^{P}, \leq) are exactly the sets

$$\llbracket \boldsymbol{e} \rrbracket \stackrel{\text{def}}{=} \{ \boldsymbol{m} \in \mathbb{N}^P \mid \boldsymbol{m} \le \boldsymbol{e} \}$$
 (4)

defined by extended markings $e \in \mathbb{N}^P_\omega$. Note that \mathbb{N}^P_ω contains \mathbb{N}^P as a substructure. Regarding effectiveness assumptions, let us just mention that, for (CI), $[\![e]\!] \subseteq [\![e']\!]$ iff $e \leq e'$. See (Goubault-Larrecq et al. 2016) for more details.

Extended Configurations. Note that, since $(\mathbb{N}^P_\omega, \leq)$ is a qo, $((\mathbb{N}^P_\omega)^\circledast, \sqsubseteq)$ is also a qo—they are in fact both wqos. An *extended configuration* is a pair (B,S) comprising a finite *base* multiset $B \in (\mathbb{N}^P_\omega)^\circledast$ and a finite *star* set $S \subseteq \mathbb{N}^P_\omega$. Then the ideals of $(Confs, \sqsubseteq)$ are exactly the sets

$$[\![B,S]\!] \stackrel{\text{def}}{=} \{M \in Confs \mid \exists E \in S^{\circledast} : M \sqsubseteq B \oplus E\}$$
 (5)

defined by extended configurations.

This representation is however not canonical, in the sense that there can be $(B,S) \neq (B',S')$ with $[\![B,S]\!] = [\![B',S']\!]$. For instance, if $e \geq e'$, then for all extended configurations (B,S),

$$[B, S \cup \{e, e'\}] = [B, S \cup \{e\}],$$
 (6)

$$[\![B \oplus [e'], S \cup \{e\}]\!] = [\![B, S \cup \{e\}]\!].$$
 (7)

In fact, those are the only two situations, and reading equations (6) and (7) left-to-right as *reduction* rules—which are furthermore confluent—we can associate to any extended configuration (B,S) a unique *reduced* extended configuration. Such an extended configuration (B,S) is such that S is an antichain and, for all extended markings $e \in S$ and $e' \in Supp(B)$, $e \not\geq e'$. Reduced extended configurations provide canonical representatives for the ideals of $(Confs, \sqsubseteq)$; we write XConfs for the set of all reduced extended configurations. In the following, for an ideal I of $(Confs, \sqsubseteq)$ we write (B(I), S(I)) for its canonical representative in XConfs.

Observe that XConfs also embeds Confs as an isomorphic substructure: any configuration M can be associated to the extended configuration (M,\emptyset) .

Regarding effectiveness assumptions, we shall only comment on (CI) and refer the reader to (Goubault-Larrecq et al. 2016) for details. Given two reduced extended configurations (B,S) and (B',S') in XConfs, $[\![B,S]\!]\subseteq [\![B',S']\!]$ iff $\exists E\in S'^\circledast$ such that $B\sqsubseteq B'\oplus E$, and $S\subseteq_H S'$, where ' \subseteq_H ' denotes the *Hoare* ordering: $S\subseteq_H S'$ iff for all $e\in S$ there exists $e'\in S'$ such that $e'\subseteq e'$.

3.3 Backward Coverability Algorithm

Consider a ν PN and a target configuration M_1 . Define

$$D_* \stackrel{\text{def}}{=} \{ M \in Confs \mid \forall M' \supseteq M_1 . M \not \sim M' \}$$
 (8)

as the set of configurations that do not cover M_1 . The purpose of the backward coverability algorithm is to compute D_* ; solving a coverability instance with source configuration M_0 then amounts to checking whether M_0 belongs to D_* .

Let us define the reachability relation in at most $k \in \mathbb{N}$ steps by $\rightarrow^{\leq 0} \stackrel{\text{def}}{=} \{(M,M) \mid M \in Confs\}$ and $\rightarrow^{\leq k+1} \stackrel{\text{def}}{=} \rightarrow^{\leq k} \cup \{(M,M'') \mid \exists M' \in Confs : M \rightarrow M' \rightarrow^{\leq k} M''\}$. The idea of the algorithm is to compute successively for every k the set D_k of configurations that do *not* cover M_1 in k or fewer steps:

$$D_k \stackrel{\text{def}}{=} \{ M \in Confs \mid \forall M' \supseteq M_1 . M \not\stackrel{\text{def}}{=} M' \} . \tag{9}$$

As shown in (Lazić and Schmitz 2015, Claim 3.2) these over-approximations D_k can be computed inductively on k:

$$D_0 = Confs \setminus \uparrow M_1$$
, $D_{k+1} = D_k \cap \operatorname{Pre}_{\forall}(D_k)$, (10)

where for any set $S \subseteq \mathit{Confs}$ its set of $\mathit{universal\ predecessors}$ is

$$\operatorname{Pre}_{\forall}(S) \stackrel{\text{def}}{=} \{ M \in \operatorname{Confs} \mid \forall M'. (M \to M' \Rightarrow M' \in S) \}.$$
 (11)

This set is downwards-closed if S is downwards-closed (Lazić and Schmitz 2015, Claim 3.3). We need here to check an additional effectiveness assumption for νPNs (which holds, see the full paper):

(**Pre**) the ideal decomposition of $Pre_{\forall}(D)$ is computable for all downwards-closed D,

where D is given as a finite set of ideal representations. Then D_0 is computed using (CU'), and at each iteration the intersection of $\operatorname{Pre}_{\forall}(D_k)$ with D_k is also computable by (Pre) and (II).

The algorithm terminates as soon as $D_k \subseteq D_{k+1}$, and then $D_{k+j} = D_k = D_*$ for all j. This is guaranteed to arise eventually by the descending chain condition, since otherwise we would have an infinite descending chain of downwards-closed sets $D_0 \supsetneq D_1 \supsetneq D_2 \supsetneq \cdots$. The termination check $D_k \subseteq D_{k+1}$ is effective by (CI): by ideal irreducibility, $D_k = I_1 \cup \cdots \cup I_n \subseteq J_1 \cup \cdots \cup J_r = D_{k+1}$ for ideals I_1, \ldots, I_n and ideals J_1, \ldots, J_m if and only if for all $1 \le i \le n$ there exists $1 \le j \le r$ such that $I_i \subseteq J_j$.

4. Complexity Upper Bounds

We establish in this section a double Ackermann upper bound on the complexity of coverability in ν PNs. The main ingredient to that end is a combinatorial statement on the length of controlled descending chains of downwards-closed sets, and we define in Section 4.2 control functions and exhibit a control on the descending chain $D_0 \supset D_1 \supset \cdots$ built by the backward coverability algorithm for a ν PN. One can extract a controlled bad sequence from such a controlled descending chain (see Section 4.3), from which the length function theorem of Rosa-Velardo (2014) yields in turn an hyper-Ackermann upper bound. In order to obtain the desired double Ackermann upper bound, we need to refine this analysis. We observe in Section 4.4 that the descending chains for ν PNs enjoy an additional star monotonicity property. This in turn allows to prove the upper bound by extracting Ackermann-controlled bad sequences of extended markings; see Theorem 9. The final step is to put this upper bound in the complexity class $\mathbf{F}_{\omega \cdot 2}$.

4.1 Fast-Growing Complexity Classes

In order to express the non-elementary functions required for our complexity statements, we employ a family of subrecursive functions $(h_{\alpha})_{\alpha}$ indexed by ordinals α known as the *Cichoń* hierarchy (Cichoń and Tahhan Bittar 1998).

Ordinal Terms. We use ordinal terms α in Cantor Normal Form (CNF), which can be written as terms $\alpha = \omega^{\alpha_1} + \cdots + \omega^{\alpha_n}$ where $\alpha_1 \geq \cdots \geq \alpha_n$ are themselves written in CNF. Using such notations, we can express any ordinal below ε_0 , the minimal fixpoint of $x = \omega^x$. The ordinal 0 is obtained when n = 0; otherwise if $\alpha_n = 0$ the ordinal α is a successor ordinal $\omega^{\alpha_1} + \cdots + \omega^{\alpha_{n-1}} + 1$, and if $\alpha_n > 0$ the ordinal α is a limit ordinal. We usually write ' λ ' to denote limit ordinals; any such limit ordinal can be written uniquely as $\gamma + \omega^{\beta}$ with $\beta > 0$.

Fundamental Sequences. For all x in \mathbb{N} and limit ordinals λ , we use a standard assignment of *fundamental sequences* $\lambda(0) < \lambda(1) < \cdots < \lambda(x) < \cdots < \lambda$ with supremum λ . Fundamental sequences are defined by transfinite induction by:

$$(\gamma+\omega^{\beta+1})(x)\stackrel{\mathrm{def}}{=} \gamma+\omega^{\beta}\cdot(x+1),\ (\gamma+\omega^{\lambda})(x)\stackrel{\mathrm{def}}{=} \gamma+\omega^{\lambda(x)}.$$

For instance, $\omega(x) = x + 1$, $\omega^2(x) = \omega \cdot (x + 1)$, $\omega^{\omega}(x) = \omega^{x+1}$, etc.

The Cichoń Hierarchy. Let $h: \mathbb{N} \to \mathbb{N}$ be a strictly increasing function. The *Cichoń* functions $(h_{\alpha}: \mathbb{N} \to \mathbb{N})_{\alpha}$ are defined by

$$h_0(x) \stackrel{\text{def}}{=} 0$$
, $h_{\alpha+1}(x) \stackrel{\text{def}}{=} 1 + h_{\alpha}(h(x))$, $h_{\lambda}(x) \stackrel{\text{def}}{=} h_{\lambda(x)}(x)$.

For instance, $h_k(x)=k$ for all finite k (thus $h_1\neq h$), but for limit ordinals λ , $h_\lambda(x)$ performs a form of diagonalisation: for instance, setting $H(x)\stackrel{\text{def}}{=} x+1$ the successor function, $H_\omega(x)=H_{x+1}(x)=x+1$, $H_{\omega^2}(x)=(2^{x+1}-1)(x+1)$ is a function

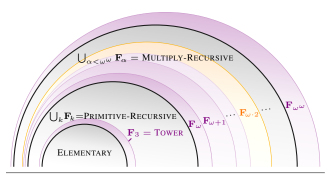


Figure 4. Pinpointing $\mathbf{F}_{\omega \cdot 2}$ among the complexity classes beyond ELEMENTARY.

of exponential growth, while H_{ω^3} is a non elementary function akin to a tower of exponentials of height x, H_{ω^ω} is a non primitive-recursive function with growth similar to the Ackermann function, and $H_{\omega^{\omega^\omega}}$ is a non multiply-recursive function characteristic of hyper-Ackermannian complexity.

The Cichoń functions are weakly increasing. If $g(x) \leq h(x)$ for all x, then also $g_{\alpha}(x) \leq h_{\alpha}(x)$ for all x. Finally, if $\alpha < \beta$, then h_{α} is eventually bounded by h_{β} : there exists x_0 such that for all $x \geq x_0$, $h_{\alpha}(x) \leq h_{\beta}(x)$.

Complexity Classes. Following (Schmitz 2016), we can define complexity classes for computations with time or space resources bounded by Cichoń functions of the size of the input. We concentrate in this paper on the double Ackermann complexity class. For $\alpha>2$, let $\mathscr{F}_{<\alpha}$ denote the set of number-theoretic functions computable in deterministic time bounded by H_{β} for $\beta<\omega^{\alpha}$, which we can write as:

$$\mathscr{F}_{<\alpha} = \bigcup_{\beta < \omega^{\alpha}} \text{FDTIME}(H_{\beta}(n)) .$$
 (12)

This class coincides with $\bigcup_{\beta<\alpha} \mathscr{F}_{\beta}$ in the *extended Grzegorczyk hierarchy* $(\mathscr{F}_{\alpha})_{\alpha}$ (Wainer 1970).

Let h be any primitive-recursive function, i.e. any function in $\mathscr{F}_{<\omega}$. Then we can define $\mathbf{F}_{\omega\cdot 2}$ by (see Schmitz 2016, Theorem 4.2):

$$\mathbf{F}_{\omega \cdot 2} = \bigcup_{p \in \mathscr{F}_{<\omega \cdot 2}} \mathsf{DTIME}(h_{\omega^{\omega \cdot 2}}(p(n))) \ . \tag{13}$$

This is the set of decision problems solvable with resources bounded by a doubly Ackermann function $h_{\omega^{\omega\cdot 2}}$ applied to some 'slower' function p of the size of the input. The definition is tailored to define completeness for $\mathbf{F}_{\omega\cdot 2}$ through many-one reductions in $\mathscr{F}_{<\omega\cdot 2}$. Although we know many examples of problems complete for the related classes \mathbf{F}_{ω} and $\mathbf{F}_{\omega^{\omega}}$ (see Figure 4 for a depiction), this is the first time we encounter the class $\mathbf{F}_{\omega\cdot 2}$.

4.2 Controlled Descending Sequences

Consider some set A with a norm $\|.\|: A \to \mathbb{N}$. Given a strictly increasing control function $g: \mathbb{N} \to \mathbb{N}$ and an initial norm $n \in \mathbb{N}$, we say that a sequence a_0, a_1, \ldots of elements from A is strongly (g, n)-controlled if $\|a_0\| \le n$ and $\|a_{i+1}\| \le g(\|a_i\|)$ for all i. A less stringent, amortised requisite is to ask $\|a_i\| \le g^i(n)$ for all i, where g^i is the ith iterate of g; we say in that case that the sequence is (g, n)-controlled.

These notions can be applied to sequences D_0, D_1, \ldots of downwards-closed subsets of $(Confs, \sqsubseteq)$ seen as finite sets of reduced extended configurations in XConfs. Let us therefore equip extended configurations $(B,S) \in XConfs$ and extended markings $e \in \mathbb{N}^P_\omega$ with the following norm: $\|B,S\| \stackrel{\mathrm{def}}{=} \max(\|B\|,\|S\|)$, $\|B\| \stackrel{\mathrm{def}}{=} \max_{e \in Supp(B)}(|B|,\|e\|)$, $\|S\| \stackrel{\mathrm{def}}{=} \max_{e \in S(B)}(\|e\|)$, and

 $\|e\| \stackrel{\text{def}}{=} \max_{p \in P \mid e(p) < \omega} e(p)$. For a finite set D of extended configurations, we then set $\|D\| \stackrel{\text{def}}{=} \max_{(B,S) \in D} \|B,S\|$.

By controlling how big the extended configurations of $\operatorname{Pre}_{\forall}(D)$ can grow as a function of $\|D\|$, we show that the descending chain $D_0 \supseteq D_1 \supseteq \cdots$ computed by the backward coverability algorithm for νPNs is strongly controlled (see the full paper):

Lemma 2 (Strong Control for ν PNs). The descending chain computed by the backward coverability algorithm for a ν PN N and target configuration M is strongly (g,n)-controlled for $g(x) \stackrel{\text{def}}{=} x + |N|$ and $n \stackrel{\text{def}}{=} ||M||$.

4.3 Length Functions Theorems

Length function theorems are combinatorial statements that provide upper bounds on the lengths ℓ of (g, n)-controlled sequences $a_0, a_1, \ldots, a_{\ell}$.

Bad Sequences of Extended Markings. A first example of a length function theorem is the following Ackermannian upper bound for bad sequences e_0, e_1, \ldots of extended markings in \mathbb{N}^{P}_{ω} : combining Corollary 2.25 and Theorem 2.34 from (Schmitz and Schnoebelen 2012) (see also Appendix A of Lazić and Schmitz 2015):

Fact 3 (Length Function Theorem for Bad Sequences in \mathbb{N}^P_ω). Let n>0. Any (g,n)-controlled bad sequence e_0,e_1,\ldots,e_ℓ of extended markings in $(\mathbb{N}^P_\omega,\leq)$ has length bounded by $h_{\omega^{|P|+1}}(n\cdot|P|!)$, where $h(x)\stackrel{\text{def}}{=}|P|\cdot q(x)$.

Proper Ideals in Descending Chains. When considering a descending chain $D_0 \supseteq D_1 \supseteq \cdots \supseteq D_\ell$ of downwards-closed subsets of some wqo (A, \leq_A) , where each set D_k is represented as a finite set of ideals, observe that we can extract at each step $0 \le k < \ell$ an ideal I_k from the decomposition of D_k that disappears in the next decomposition D_{k+1} . We call such an ideal *proper*; it satisfies $I_k \subseteq D_k$ but $I_k \not\subseteq D_{k+1}$, and as a consequence $I_k \not\subseteq I_{k'}$ for all k' > k since $D_{k'} \subseteq D_{k+1}$ in this case. Hence we can extract a sequence $I_0, I_1, \ldots, I_{\ell-1}$ of ideals, which is a bad sequence for the inclusion ordering.

As an application, consider a (g,n)-controlled descending chain $S_0 \supsetneq_H S_1 \supsetneq_H \cdots \supsetneq_H S_\ell$ of antichains $S_k \subseteq \mathbb{N}^P_\omega$ for the Hoare ordering. Each antichain S_k is in fact an ideal representation for the downwards-closed set of markings $D_k = \bigcup_{e \in S} \llbracket e \rrbracket \subseteq \mathbb{N}^P$, i.e. this defines a descending chain $D_0 \supsetneq D_1 \supsetneq \cdots \supsetneq D_\ell$ (the reader can check that $S \supsetneq_H S'$ iff the associated downwards-closed sets are strictly included: $\bigcup_{e \in S} \llbracket e \rrbracket \supsetneq \bigcup_{e' \in S'} \llbracket e' \rrbracket$). As pointed out just before, we can extract a bad sequence $e_0, e_1, \ldots, e_{\ell-1}$ of extended markings in \mathbb{N}^P_ω representing proper ideals. Furthermore, this sequence is also (g,n)-controlled, thus Fact 3 can be applied:

Corollary 4 (Length Function Theorem for Hoare-Descending Chains over \mathbb{N}^P_ω). Let n>0. Any (g,n)-controlled descending chain $S_0 \supsetneq_H S_1 \supsetneq_H \cdots \supsetneq_H S_\ell$ of antichains of $(\mathbb{N}^P_\omega, \leq)$ has length at most $h_{\omega^{|P|+1}}(n\cdot |P|!)+1$, where $h(x)\stackrel{\text{def}}{=} |P|\cdot g(x)$.

4.4 Star-Monotone Descending Chains

Let us lift the step relation \to to work over ideals. Define for any ideal $S\subseteq Confs$

$$\operatorname{Post}_{\exists}(S) \stackrel{\text{def}}{=} \{ M' \in \operatorname{Confs} \mid \exists M \in S . M \to M' \} . \tag{14}$$

Then for any ideal I of $(Confs, \sqsubseteq)$, $\downarrow Post_{\exists}(I)$ is downwards-closed with a unique decomposition into maximal ideals. We follow Blondin et al. (2014) and write ' $I \to J$ ' if J is an ideal from the decomposition of $\downarrow Post_{\exists}(I)$. We will use the following fact proven in (Lazić and Schmitz 2015, Claim 4.2):

Fact 5 (Proper Transition Sequences). If I_{k+1} is a proper ideal of D_{k+1} , then there exist an ideal J and a proper ideal I_k of D_k such that $I_{k+1} \to J \subseteq I_k$.

We can check that this step relation lifted to ideals is monotone in the star set (see the full paper):

Lemma 6 (Ideal Steps are Star-Monotone). *If* I *and* J *are two ideals and* $I \rightarrow J$, *then* $S(I) \subseteq_H S(J)$.

We say that a descending chain $D_0 \supseteq D_1 \supseteq \cdots \supseteq D_\ell$ of downwards-closed subsets of Confs is star-monotone if for all $0 \le k < \ell - 1$ and all proper ideals I_{k+1} in the decomposition of D_{k+1} , there exists a proper ideal I_k in the decomposition of D_k such that $S(I_{k+1}) \subseteq_H S(I_k)$.

Lemma 7 (ν PN Descending Chains are Star-Monotone). *The descending chains computed by the backward coverability algorithm for \nuPNs are star monotone.*

Proof. Let $D_0 \supsetneq D_1 \supsetneq \cdots \supsetneq D_\ell$ be the descending chain computed for our νPN . Suppose $0 \le k < \ell - 1$ and I_{k+1} is a proper ideal in the decomposition of D_{k+1} . By Fact 5, there exists a proper ideal I_k in the decomposition of D_k and an ideal J such that $I_{k+1} \to J$ and $J \subseteq I_k$. By Lemma 6, $S(I_{k+1}) \subseteq_H S(J)$, and by (CI), $S(J) \subseteq_H S(I_k)$.

The crux of our proof is the following theorem:

Theorem 8 (Length Function Theorem for Star-Monotone Descending Chains over $(\mathbb{N}^P_\omega)^{\circledast}$). Let n>0. Any strongly (g,n)-controlled star-monotone descending chain $D_0\supseteq D_1\supseteq\cdots\supseteq D_\ell$ of configurations in $(\mathbb{N}^P_\omega)^{\circledast}$ has length at most $h_{\omega^{\omega\cdot 2}}(|P|+n)$ for some h primitive-recursive in g.

Proof idea. We prove the theorem in the full paper, but provide here a quick overview of its proof.

Since the sequence $D_0 \supseteq D_1 \supseteq \cdots \supseteq D_\ell$ is star-monotone, starting from some proper ideal $I_{\ell-1}$ in the decomposition of $D_{\ell-1}$, we can find a sequence of proper ideals $I_0,\ldots,I_{\ell-1}$ such that $S(I_k) \supseteq_H S(I_{k+1})$ for all $0 \le k < \ell-1$. Let $S_j \stackrel{\text{def}}{=} S(I_j)$. We then extract a subsequence with $S_{i_0} \supseteq_H S_{i_1} \supseteq_H \cdots \supseteq_H S_{i_r}$ such that $i_0 \stackrel{\text{def}}{=} 0$ and $S_0 \equiv_H S_1 \equiv_H \cdots \equiv_H S_{i_1-1} \supseteq_H S_{i_1} \equiv_H \cdots \equiv_H S_{i_2-1} \supseteq_H S_{i_2} \cdots S_{i_r-1} \supseteq_H S_{i_r} \equiv_H \cdots \equiv_H S_{\ell-1}$, where two star sets are *Hoare-equivalent*, noted $S \equiv_H S'$, iff $S \subseteq_H S'$ and $S \supseteq_H S'$. Without loss of generality, we can also assume that $S(I) \equiv_H S_{i_j+1-1}$ for all proper ideals I in a segment $D_{i_j,\ldots,D_{i_j+1}-1}$ of the computation.

We analyse independently the length of a 'Hoare-equivalent' segment where $S_{i_j} \equiv_H S_{i_j+1} \equiv_H \cdots \equiv_H S_{i_j+1-1}$ and the length of the 'Hoare-descending' chain where $S_{i_0} \supsetneq_H S_{i_1} \supsetneq_H \cdots \supsetneq_H S_{i_r}$. For the former, we show that the associated sequence of bases $B_{i_j}, B_{i_j+1}, \ldots, B_{i_{j+1}-1}$ is a bad sequence controlled by $(g, \|D_{i_j}\|)$, where all the B_k can be treated as $(|P| \cdot \|D_{i_j}\|)$ -dimensional vectors in $\mathbb{N}_{\omega}^{|P| \cdot \|D_{i_j}\|}$. We can therefore apply Fact 3 to this sequence and obtain an Ackermannian control (a, n) on the sequence $S_{i_0} \supsetneq_H S_{i_1} \supsetneq_H \cdots \supsetneq_H S_{i_r}$. In turn, this sequence is bounded thanks to Corollary 4 by $a_{\omega^\omega}(n \cdot |P|!)$, a function that nests an Ackermannian blowup at each of its Ackermannian-many steps. An analysis of this last function yields the result.

Together with the primitive-recursive control (g,n) exhibited in Lemma 2 and the star-monotonicity of the descending chains computed by the backward algorithm shown in Lemma 7, Theorem 8 provides an upper bound in $\mathbf{F}_{\omega \cdot 2}$ as defined in Equation 13:

Theorem 9. The coverability problem for νPNs is in $\mathbf{F}_{\omega \cdot 2}$.

5. Complexity Lower Bounds

5.1 Ackermann Functions

When it comes to lower bounds, we find it more convenient to work with a variant of the functions from Section 4.1 called the *Ackermann hierarchy*. Here we shall only need the functions $(A_{\alpha})_{\alpha<\omega\cdot 2}$ from this hierarchy, which can be defined as follows for all k and x in \mathbb{N} :

$$\begin{aligned} A_1(x) &\stackrel{\text{def}}{=} 2x \;, & A_{k+2}(x) &\stackrel{\text{def}}{=} A_{k+1}^x(1) \;, \\ A_{\omega}(x) &\stackrel{\text{def}}{=} A_{x+1}(x) \;, & A_{\omega+k+1}(x) &\stackrel{\text{def}}{=} A_{\omega+k}^x(1) \;. \end{aligned}$$

The double Ackermann function is then defined as

$$A_{\omega \cdot 2}(x) \stackrel{\text{def}}{=} A_{\omega + x + 1}(x) ; \qquad (15)$$

note that this is considerably larger than $A_{\omega}(A_{\omega}(x))$. We can employ the function $A_{\omega \cdot 2}$ instead of $H_{\omega^{\omega \cdot 2}}$ since, by (Schmitz 2016, Theorem 4.1),

$$\mathbf{F}_{\omega \cdot 2} = \bigcup_{p \in \mathscr{F}_{<\omega \cdot 2}} \mathsf{DTIME}(A_{\omega \cdot 2}(p(n))) \ . \tag{16}$$

5.2 Routines, Libraries, and Programs

To present our lower bound construction, we shall develop some simple and limited but convenient mechanisms for programming with νPNs .

Syntax of Routines and Libraries. Let a library mean a sequence of named routines

$$\ell_1: R_1, \ldots, \ell_K: R_K,$$

where ℓ_1, \ldots, ℓ_K are pairwise distinct labels. In turn, a *routine* is a sequence of commands $c_1, \ldots, c_{K'}$, where each c_i for i < K' is one of the following:

- a νPN transition,
- a nondeterministic jump goto G for a nonempty subset G of $\{1, \ldots, K'\}$, or
- a subroutine invocation call ℓ' ;

and $c_{K'}$ is return.

The call ℓ' commands should be thought of as invoking subroutines from another, lower level, library which remains to be provided and composed with this library.

Semantics of Programs. When a library contains no subroutine calls, we say it is a *program*. The denotation of a program L as above is a $\nu PN \mathcal{N}(L)$ constructed so that:

- The places of $\mathcal{N}(L)$ are all the places that occur in L, and four special places $p, \overline{p}, p', \overline{p}'$. Places $\langle p, \overline{p} \rangle$ are used to store the pair of numbers $\langle i, K-i \rangle$ where $\ell_i: R_i$ is the routine being executed, and then places $\langle p', \overline{p}' \rangle$ to store the pair of numbers $\langle i', K'-i' \rangle$ where i' is the current line number in routine R_i and K' is the maximum number of lines in any R_1, \ldots, R_K .
- Each transition of $\mathcal{N}(L)$ either executes a transition command $c_{i'}$ inside some R_i ensuring that $\langle p, \overline{p} \rangle$ contains $\langle i, K i \rangle$ and modifying the contents of $\langle p', \overline{p}' \rangle$ from $\langle i', K' i' \rangle$ to $\langle i' + 1, K' (i' + 1) \rangle$, or similarly executes a nondeterministic jump command.

Initial and Final Tape Contents. We shall refer to the special p, \overline{p} , p', \overline{p}' as *control places*, to the rest as *tape places*, and to markings of the latter places as *tape contents*. For two tape contents M and M', we say that a routine $\ell_i: R_i$ can *compute* M' from M if and only if $\mathcal{N}(L)$ can reach in finitely many steps M' with the control at the last line of R_i from M with the control at the first line of R_i ; when no M' is computable from M by $\ell_i: R_i$, we say that the routine *cannot terminate* from M.

Interfaces and Compositions of Libraries. For a library L, let us write $\Lambda_{\rm in}(L)$ (resp., $\Lambda_{\rm out}(L)$) for the set of all routine labels that are invoked (resp., provided) in L. We say that libraries L_0 and L_1 are compatible if and only if $\Lambda_{\rm in}(L_0)$ is contained in $\Lambda_{\rm out}(L_1)$. In that case, we can compose them to produce a library $L_0 \circ L_1$ in which tape contents of L_1 persist between successive invocations of its routines, as follows:

- $\Lambda_{\rm in}(L_0 \circ L_1) = \Lambda_{\rm in}(L_1)$ and $\Lambda_{\rm out}(L_0 \circ L_1) = \Lambda_{\rm out}(L_0)$.
- $L_0 \circ L_1$ has an additional place w used to store the name space of L_0 (i.e., for each name manipulated by L_0 , one token labelled by it) and an additional place \overline{w} for the same purpose for L_1 .
- For each routine $\ell: R$ of L_0 , the corresponding routine $\ell: R \circ L_1$ of $L_0 \circ L_1$ is obtained by ensuring that the transition commands in R (resp., L_1) maintain the name space stored on place w (resp., \overline{w}), and then inlining the subroutine calls in R.

5.3 Counter Libraries

Our main technical objective, after which it will be easy to arrive at the claimed lower bound for νPN coverability, is to construct libraries that implement increments, decrements and zero tests on a pair of counters whose values range up to a bound which is doubly-Ackermannian in the sizes of the libraries.

To begin, we define the general notion of libraries that provide the operations we need on a pair of bounded counters, as well as what it means for a stand-alone such library to be correct up to a specific bound. A key step is then to consider counter libraries that may not be programs, i.e. may invoke operations on another pair of counters (which we call *auxiliary*). We define correctness of such libraries also, where the bounds of the provided counters may depend on the bounds of the auxiliary counters.

As illustrations of both notions of correctness, we provide examples that will moreover be used in the sequel.

Interfaces of Counter Libraries. Letting Γ denote the set of labels of operations on pairs of bounded counters

$$\Gamma \stackrel{\text{def}}{=} \{init, eq, i.inc, i.dec, i.iszero, i.ismax : i \in \{1, 2\}\},\$$

we regard L to be a *counter* library if and only if $\Lambda_{\rm out}(L)=\Gamma$ and $\Lambda_{\rm in}(L)\subseteq\Gamma$.

Correct Counter Programs. When L is also a program, and N is a positive integer, we say that L is N-correct if and only if, after initialisation, the routines behave as expected with respect to the bound N. Namely, for every tape contents M which can be computed from the empty tape contents by a sequence σ of operations from Γ , provided init occurs only as the first element of σ and letting n_i be the difference between the numbers of occurrences in σ of i.inc and i.dec, we must have for both $i \in \{1, 2\}$:

- eq can terminate from M if and only if $n_1 = n_2$;
- i.inc can terminate if and only if $n_i < N-1$;
- i.dec can terminate if and only if $n_i > 0$;
- *i.iszero* can terminate if and only if $n_i = 0$;
- *i.ismax* can terminate if and only if $n_i = N 1$.

Example: Enumerated Counter Program. For any positive integer N, it is trivial to implement a pair of N-bounded counters by manipulating the values and their complements directly. Let Enum(N) be a counter program which uses four places e_1 , \overline{e}_1 , e_2 , \overline{e}_2 and such that for both $i \in \{1, 2\}$:

• routine *init* puts N-1 tokens onto \overline{e}_1 and N-1 tokens onto \overline{e}_2 , all carrying a fresh name d;

- routine eq guesses $n \in \{0, \ldots, N-1\}$, takes $\langle n, N-1-n, n, N-1-n \rangle$ tokens from places $\langle e_1, \overline{e}_1, e_2, \overline{e}_2 \rangle$, and then puts them back;
- routine *i.inc* moves a token from \overline{e}_i to e_i ;
- routine *i.dec* moves a token from e_i to \overline{e}_i ;
- routine i.iszero takes N-1 tokens from place \overline{e}_i and then puts them back;
- routine i.ismax takes N-1 tokens from place e_i and then puts them back.

It is simple to verify that Enum(N) is computable in space logarithmic in N, and that:

Lemma 10. For every N, the counter program Enum(N) is N-correct.

Note that the size of Enum(N) is at least polynomial in N, whereas our technical aim is to build correct counter programs whose bounds are doubly-Ackermannly larger than their sizes.

Correct Counter Libraries. Given a counter library L, and given a function $F: \mathbb{N}^+ \to \mathbb{N}^+$, we say that L is F-correct if and only if, for all N-correct counter programs C, $L \circ C$ is F(N)-correct.

We employ Acker, a counter library such that the bound of the provided counters equals the Ackermann function $A_{\omega}(N)$ applied to the bound N of the auxiliary counters. This is an adaptation of the construction by Schnoebelen (2010) of Ackermannian values in reset Petri nets, using the addressing mechanism described in Section 2.3 to simulate an N-dimensional reset Petri net; see the full paper for details.

Lemma 11. The counter library Acker is A_{ω} -correct.

5.4 An Iteration Operator

The most complex part of our construction is an operator $-^*$ whose input is any counter library L. Its output L^* is also a counter library, which essentially consists of an arbitrary number of copies of L composed in sequence. Namely, for any N-correct counter program C, the counter operations provided by $L^* \circ C$ behave in the same way as those provided by

$$\overbrace{L \circ \cdots \circ L}^{N} \circ Enum(1).$$

Hence, when L is F-correct, we have that L^* is F'-correct, where $F'(x) = F^x(1)$. Recall that Enum(1) provides trivial counters, i.e. with only one possible value, so its testing operations are essentially no-ops whereas its increments and decrements cannot terminate successfully.

The main idea for the definition of L^* is to combine a distinguishing of name spaces as in the composition of libraries with an arbitrarily wide indexing mechanism like the one employed in Section 2.3. The key insight here is that a whole collection of 'addressing places' $\langle a_i, \bar{a}_i \rangle_i$ as used in Section 2.3 can be simulated by adding one layer of addressing.

More precisely, numbering the copies of L by $0,\ldots,N-1$, writing $\ell_1:R_1,\ldots,\ell_K:R_K$ for the routines of L where $\ell_1=init$ (since L is a counter library, it has $K=|\Gamma|=10$ routines) and writing K' for the maximum number of lines in any R_1,\ldots,R_K,L^* can maintain the control and the tape of each copy of L in the implicit composition as follows:

• To record that the program counter of the *i*th copy of L is currently in routine $\ell_j: R_j$ at line $j', \langle i, N-1-i, j, K-j, j', K'-j', 1 \rangle$ tokens carrying a separate name d_i are kept on special places $\langle w, \overline{w}, p, \overline{p}, p', \overline{p}', \overline{t} \rangle$.

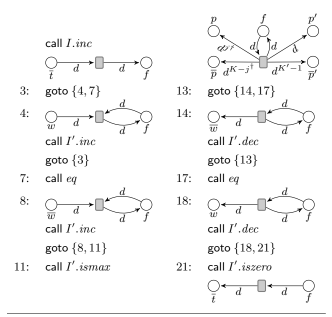


Figure 5. Performing a call $\ell_{j^{\dagger}}$ provided I < N - 1. At the beginning, I' is assumed to be zero, and the same is guaranteed at the end.

- The current height *i* of the stack of subroutine calls is kept in one of the auxiliary counters, and we have that:
 - for all i' < i, the program counter of the i'th copy of L is at some subroutine invocation call ℓ' such that the program counter of the (i'+1)th copy of L is in the routine named ℓ' ;
 - for all i'>i, there are $\langle i',N-1-i',0,0,0,0,1\rangle$ tokens carrying $d_{i'}$ on places $\langle w,\overline{w},p,\overline{p},p',\overline{p}',\overline{t}\rangle$.
- For every name manipulated by the *i*th copy of L, $\langle i, N-1-i, 1 \rangle$ tokens carrying it are kept on special places $\langle w, \overline{w}, t \rangle$. Thus, places t and \overline{t} are used to distinguish these names from the artificial names that record the control.

To define L^* , its places are all the places that occur in L, plus nine special places $w, \overline{w}, p, \overline{p}, p', \overline{p}', \overline{t}, t$ and f. Writing N for the bound of the auxiliary counters and I, I' for the two auxiliary counters, routine $\ell_j: R_j^*$ of L^* is defined to execute:

- If $\ell_j = init$, initialise the auxiliary counters (by calling their init routine), and then using the auxiliary counters and place f, for each $i \in \{0, \ldots, N-1\}$, put $\langle i, N-1-i, 1 \rangle$ tokens carrying a fresh name d_i onto places $\langle w, \overline{w}, \overline{t} \rangle$.
- Put $\langle j,K-j,1,K'-1\rangle$ tokens carrying d_0 onto places $\langle p,\overline{p},p',\overline{p}'\rangle$. (I will always be 0 at this point.)
- Repeatedly, using I' and place f, identify j' and j'' such that there are $\langle I, N-1-I, j', K-j', j'', K'-j'', 1 \rangle$ tokens carrying d_I on places $\langle w, \overline{w}, p, \overline{p}, p', \overline{p}', \overline{t} \rangle$, and advance the Ith copy of L by performing the command c at line j'' in routine $\ell_{j'}: R_{j'}$ of L as follows:
 - If c is a ν PN transition, use I' and place f to maintain the Ith name space, i.e. to ensure that all names manipulated by c have $\langle I, N-1-I, 1 \rangle$ tokens on places $\langle w, \overline{w}, t \rangle$.
 - If c is a nondeterministic jump goto G, choose $j^{\ddagger} \in G$ and ensure that there are $\langle j^{\ddagger}, K' j^{\ddagger} \rangle$ tokens carrying d_I on places $\langle p', \overline{p}' \rangle$.

- If c is a subroutine invocation call $\ell_{j^{\dagger}}$ and I < N 1, put $\langle j^{\dagger}, K j^{\dagger}, 1, K' 1 \rangle$ tokens carrying d_{I+1} on places $\langle p, \overline{p}, p', \overline{p}' \rangle$, and increment I. Example code that implements this can be found in Figure 5.
- If c is a subroutine invocation call ℓ' , I=N-1 and ℓ' is not an increment or a decrement (of the trivial counter program Enum(1)), simply increment the program counter by moving a token carrying d_I from place \overline{p}' to place p'. When ℓ' is an increment or a decrement, L^* blocks at this point.
- In the remaining case, c is return. Remove the tokens carrying d_I from places $\langle p, \overline{p}, p', \overline{p}' \rangle$. If I > 0, move a token carrying d_{I-1} from \overline{p}' to place p' and decrement I. Otherwise, exit the loop.

We observe that L^* is computable from L in logarithmic space.

Lemma 12. For every F-correct counter library L, we have that L^* is $\lambda x. F^x(1)$ -correct.

Proof. We argue by induction on N that, for every N-correct counter program $C, L^* \circ C$ is $F^N(1)$ -correct.

The base case N=1 is straightforward. Suppose C is 1-correct, i.e. provides counters with only one possible value. By the definition of L^* , when the bound of the auxiliary counters is 1, only one copy of L is simulated. Hence $L^* \circ C$ as a counter program is indistinguishable from $L \circ Enum(1)$. The latter is $F^N(1)$ -correct, i.e. F(1)-correct, because L is assumed F-correct and Enum(1) is 1-correct by Lemma 10.

For the inductive step, suppose C is (N+1)-correct. For any tape contents M of $L^* \circ C$ and $i \in \{0,\ldots,N\}$, let M_i denote the subcontents belonging to the ith copy of L, i.e. the restriction of M to the names that label $\langle i,N-i,1\rangle$ tokens on places $\langle w,\overline{w},t\rangle$ and to the places of L.

By the inductive hypothesis and Lemma 10, $L^* \circ Enum(N)$ is $F^N(1)$ -correct, and so $L \circ (L^* \circ Enum(N))$ is $F^{N+1}(1)$ -correct. For any tape contents M' of the latter counter program and $i \in \{0,\ldots,N\}$, let M_i' denote: if i=0, the subcontents of the left-hand L; otherwise, the subcontents belonging to the (i-1)th copy of L in $L^* \circ Enum(N)$.

The required conclusion that $L^* \circ C$ is $F^{N+1}(1)$ -correct is implied by the next claim, proven in the full paper:

Claim 12.1. For every tape contents M and M' which $L^* \circ C$ and $L \circ (L^* \circ Enum(N))$ (respectively) can compute from the empty tape contents by a sequence σ of counter operations where init occurs only as the first element, we have that:

- 1. $M_i = M'_i$ for all $i \in \{0, ..., N\}$;
- 2. for every counter operation $op \neq init$, $L^* \circ C$ can complete op from M if and only if $L \circ (L^* \circ Enum(N))$ can complete op from M'.

5.5 Doubly-Ackermannian Minsky Machines

We are now equipped to reduce from the following $\mathbf{F}_{\omega \cdot 2}$ -complete problem (cf. Schmitz 2016, Section 2.3.2):

Given a deterministic Minsky machine \mathcal{M} , does it halt while the sum of counters is less than $A_{\omega \cdot 2}(|\mathcal{M}|)$?

and thereby establish our lower bound. The idea here is classical: simulate ${\cal M}$ by a reset Petri net on a doubly-Ackermannian budget, and if it halts then check that the simulation was accurate.

Theorem 13. The coverability problem for νPNs is $\mathbf{F}_{\omega \cdot 2}$ -hard.

Proof. Suppose \mathcal{M} a deterministic Minsky machine. Let $L_{|\mathcal{M}|}$ be the counter library $(\cdots (Acker^*)^* \cdots)^*$ with $|\mathcal{M}| + 1$ nested

iteration operators. By lemmata 10, 11 and 12, we have that $L_{|\mathcal{M}|}$ is $A_{\omega+|\mathcal{M}|+1}$ -correct and that $L_{|\mathcal{M}|} \circ Enum(|\mathcal{M}|)$ is $A_{\omega\cdot 2}(|\mathcal{M}|)$ -correct. Finally, let $Sim(\mathcal{M})$ be a one-routine library that uses one of the pair of counters provided by the counter program $L_{|\mathcal{M}|} \circ Enum(|\mathcal{M}|)$ as follows:

- Initialise $L_{|\mathcal{M}|} \circ Enum(|\mathcal{M}|)$.
- Simulate \mathcal{M} where zero tests are performed as resets (cf. Section 2.3) and where the difference between the total number of increments and the total number of decrements is maintained in a counter T of $L_{|\mathcal{M}|} \circ Enum(|\mathcal{M}|)$. Any attempt to increment T beyond its maximum value blocks the simulation.
- If M halts, check that the sum of its counters is at least T, i.e. decrease T to zero while at each step decrementing some counter of M.

Observe that the latter check succeeds if and only if there was no reset of a non-zero counter, i.e. all zero tests in the simulation were correct. Hence, \mathcal{M} halts while the sum of its counters is less than $A_{\omega \cdot 2}(|\mathcal{M}|)$ if and only if the one-routine program

$$Test(\mathcal{M}) = Sim(\mathcal{M}) \circ (L_{|\mathcal{M}|} \circ Enum(|\mathcal{M}|))$$

can terminate, i.e. the $\nu PN \mathcal{N}(Test(\mathcal{M}))$ can cover the marking in which the control places point to the last line of $Test(\mathcal{M})$.

Since the iteration operator is computable in logarithmic space and increases the number of places by adding a constant, we have that the counter library $L_{|\mathcal{M}|}$ and thus also the νPN $\mathcal{N}(Test(\mathcal{M}))$ are computable in time elementary in $|\mathcal{M}|$, and that their numbers of places are linear in $|\mathcal{M}|$. We conclude the $\mathbf{F}_{\omega\cdot 2}$ -hardness by the closure under any sub-doubly-Ackermannian reduction (i.e. in $\mathcal{F}_{<\omega\cdot 2}$), and therefore certainly any elementary one (Schmitz 2016, Section 2.3.1).

6. Concluding Remarks

In this paper, we have shown that coverability in ν -Petri nets is complete for double Ackermann time, i.e. $\mathbf{F}_{\omega\cdot 2}$ -complete. In order to solve this open problem, we have applied a new technique to analyse the complexity of the backward coverability algorithm using ideal representations—thereby demonstrating the versatility of this technique designed in (Lazić and Schmitz 2015)—, and pushed for the first time the 'object oriented' construction of Lipton (1976) beyond Ackermann-hardness. This is also the first known instance of a natural decision problem for double Ackermann time.

Our $\mathbf{F}_{\omega \cdot 2}$ upper bound furthermore improves the best known upper bound for coverability in unordered data Petri nets. In this case however, the currently best known lower bound is hardness for \mathbf{F}_3 , which was proven by Lazić et al. already in 2008, leaving quite a large complexity gap.

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