Perfect class hashing and numbering for object-oriented implementation

Roland Ducournau* and Floréal Morandat

LIRMM – Université Montpellier II and CNRS, France

SUMMARY

Late binding and subtyping create run-time overhead for object-oriented languages, especially in the context of both multiple inheritance and dynamic loading, for instance for JAVA interfaces. In a previous paper, we proposed a novel approach based on perfect hashing and truly constant-time hashtables for implementing subtype testing and method invocation in a dynamic loading setting. In this first study, we based our efficiency assessment on Driesen’s abstract computational model for the time aspect, and on large-scale benchmarks for the space aspect. The conclusions were that the technique was promising but required further research in order to assess its scalability. This article presents some new results on perfect class hashing that enhance its interest. We propose and test both new hashing functions and an inverse problem which amounts to selecting the best class identifiers in order to minimize the overall hashtable size. This optimizing approach is proven to be optimal for single inheritance hierarchies. Experiments within an extended testbed with random class loading and under cautious assumptions about what should be a sensible class loading order show that perfect class hashing scales up gracefully, especially on JAVA-like multiple-subtyping hierarchies. Furthermore, perfect class hashing is implemented in the PRM compiler testbed, and compared here with the coloring technique, which amounts to maintaining the single inheritance implementation in multiple inheritance. The overall conclusion is that the approach is efficient from both time and space standpoints with the bit-wise hashing function. In contrast, the poor time efficiency of modulus hashing function on most processors is confirmed.

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1. INTRODUCTION

The implementation of object-oriented languages is an important issue in the context of both multiple inheritance and dynamic loading. Object-oriented implementation concerns a few basic mechanisms that are invoked billions of times during a 1-minute program execution. Very high efficiency is necessary to ensure this intensive use, and it is arguable that all implementations of object-oriented languages, especially in this context, should fulfill the following requirements: (i) constant-time, (ii) inlining and (iii) linear-space. Constant time is the only way to bound the worst-case behaviour. Whereas it is essential in a real-time setting, it remains highly preferable in all settings. Moreover, the basic mechanisms must be implemented by a short sequence of instructions that is inlined, thus saving a function call and reducing the time constant. Finally, space linearity ensures that the implementation will scale up gracefully with the program size; we shall see, however, that linearity must be understood with a slightly specific meaning. This general

*Correspondence to: LIRMM – Université Montpellier II and CNRS, 161 rue Ada, 34000 Montpellier, France
E-mail: Roland.Ducournau@lirmm.fr
implementation issue is exemplified by the two most used languages that support both features, namely C++ and JAVA. When the virtual keyword is used for inheritance, C++ provides a fully reusable implementation, based on subobjects, which however implies a lot of compiler-generated fields in the object layout and pointer adjustments at run-time\(^1\) [1]. Moreover, it does not meet the linear-space requirement and, until recently, there were no known efficient subtype tests available for this implementation. JAVA provides multiple inheritance of interfaces only but, even in this restricted setting, the current implementations are not constant-time (see for instance [2]). The present research was motivated by this observation—though object-oriented technology is mature, scalable implementations are urgently needed due to the ever-increasing size of object-oriented class libraries and there is still considerable doubt about the scalability of existing implementations.

In a previous work, the first author proposed a new technique, called perfect class hashing (PH), for subtyping tests and method invocation [3]. To our knowledge, this is the first and only technique that fulfills all these requirements in the multiple inheritance and dynamic loading context. However, our experiments only concluded that the technique was promising and the need for further research was stressed. Two hashing functions were actually considered, namely modulus, i.e. the remainder of integer division, and bit-wise and. These two functions involve a tradeoff between space and time efficiency, and the space efficiency of bit-wise and needs to be further improved and assessed.

In this article, we present the results of more in-depth studies and experiments that show that a variant of perfect hashing, called perfect class numbering, provides a very efficient object-oriented implementation, even with bit-wise-and and from the space standpoint.

1.1. Object-oriented implementation and perfect hashing

One typical implementation issue of object-oriented languages is late binding, which is also referred to as the message sending metaphor. The underlying principle is that the address of the actually called procedure is not statically determined at compile-time, but depends on the dynamic type of a distinguished parameter known as the receiver. An issue similar to message sending arises with attributes (aka fields, instance variables, slots, data members according to the languages), since their position in the object layout may depend on the object’s dynamic type. Subtyping introduces a last feature, i.e. run-time subtype checks, which amounts to testing whether the value of \(x\) is an instance of a class \(C\) or, equivalently, whether the dynamic type of \(x\) is a subtype of \(C\). This is the basis for so-called downcast operators. Object-oriented implementation represents the data structures and algorithms that are required for these three mechanisms, namely method invocation, attribute access and subtype testing.

When static typing is coupled with single inheritance of classes and types—we call it single subtyping—the standard implementation of object-oriented languages allows for direct access to the desired data at an invariant position. This optimizes the implementation. Otherwise, dynamic typing or multiple inheritance make it harder to retain such direct access, especially in a dynamic loading setting.

Perfect Class Hashing\(^2\) represents an alternative implementation in the context of static typing, multiple inheritance and dynamic loading. The position of all data required for implementing a class is no longer invariant with respect to the dynamic type of the object; it is instead determined by a collision-free hashing function which depends on the dynamic type. Such functions are called perfect hashing functions [4, 5].

Perfect class hashing was first applied to subtype testing. It can be formalized as follows. Let \((X, \leq)\) be a partial order that represents a class hierarchy, namely \(X\) is a set of classes and \(\leq\) the specialization relationship that supports inheritance. The subtype test amounts to checking at run-time that a class \(c\) is a superclass of a class \(d\), i.e. \(d \leq c\). An efficient implementation of this

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\(^1\) When the virtual keyword is omitted, the implementation is more efficient but no longer fully reusable because it yields repeated inheritance. The language is thus no longer compatible with both multiple inheritance and dynamic loading. In the following, we only consider C++ for the virtual semantics and implementation.

\(^2\) The technique was originally called ‘perfect hashing’. As this name generates confusion between the general hashing technique and its application to object-oriented languages, we now prefer to call it ‘perfect class hashing’.
test amounts to precomputing a data structure that allows for constant time. Dynamic loading adds a constraint, namely that the technique should preferably be incremental. Classes are loaded at run-time in a total order such that superclasses are loaded before subclasses. With an incremental technique, the data structure for the newly loaded class is computed from the data already computed for previously loaded classes, and no recomputation is needed. The perfect class hashing principle is as follows. When a class \( c \) is loaded, a unique immutable identifier \( \text{id}_c \) is associated with it and the set \( I_c = \{ \text{id}_d \mid c \preceq d \} \) of identifiers of all its superclasses is known. Thus, \( c \preceq d \) iff \( \text{id}_d \in I_c \). This set \( I_c \) is immutable, hence it can be hashed with some perfect hashing function \( h_c \), that is, a hashing function injective on \( I_c \). The previous condition becomes: \( c \preceq d \) iff \( h_c[\text{id}_c] = \text{id}_d \), whereby \( h_c \) denotes the hashtable of \( c \), i.e. a simple array. The technique is obviously incremental since all hashtables are immutable and the computation of \( h_c \) only depends upon \( I_c \). In the same article, an application to method invocation in the restricted case of JAVA interfaces was also proposed.

### 1.2. Limitations of previous work

In our previous work, we considered one-parameter collision-free hashing functions such that \( h_c(x) = \text{hash}(x, H_c) \), whereby \( H_c \) is the hashtable size. Two functions were considered for \( \text{hash} \), namely modulus (denoted \( \text{mod} \) hereafter) and bit-wise and (the exact function maps \( x \) to \( \text{and}(x, H_c - 1) \)). In both cases, \( H_c \) is defined as the least integer such that \( h_c \) is injective on the set \( I_c \). The resulting techniques are respectively denoted PH-mod and PH-and. Both functions involve a single machine instruction. However, whereas bit-wise and is a 1-cycle instruction, the latency of integer division is commonly more than 20 cycles, hence markedly higher than the total cycle count of each mechanism with the standard single-subtyping implementation. Therefore, the time efficiency of PH-mod is expected to be rather low, while that of PH-and should be high. In the aforementioned article, we computed the \( H_c \) parameters on a set of large-scale benchmarks commonly used in the object-oriented compilation community. Our requirement for space-linearity means that the memory occupation for these tables should be linear in the cardinality of the specialization relationship \( \preceq \). The results of our tests were encouraging but not perfect—namely, PH-mod required a little more than twice the cardinality of \( \preceq \), but PH-and appeared to be much more greedy, especially in the single case of a very large benchmark. Hence, the scalability of PH-and was not certain, and the main point at issue is the memory occupation of PH-and.

Our assessment of time-efficiency relied on an abstract processor model borrowed from [6]; though we believe in the model validity, it should obviously be complemented by empirical time measurement. Later, we tested perfect hashing along with a variety of implementation techniques in the PRM testbed [7]. The results mostly confirm our earlier abstract evaluation.

These first experiments left, however, a number of issues open. PH parameters depend on the class IDs which are assigned to classes as they are loaded, hence they should vary according to class loading orders. However, only a single arbitrary order was considered, and it was likely a top-down depth-first linear extension that might be far from representative of actual class loading orders. Efficient variants have also been sought. A variant based on quasi-perfect hashing (qPH) [8], which yields a 2-probe test, was proposed; it gives more compact tables at the expense of less efficient code. Another variant might be based on a 2-parameter hashing function, but we could not find a function that would reduce the table size while keeping the time overhead smaller than with modulus. We also proposed to optimize the identifier of the currently loaded class, but the technique, called perfect class numbering (PN), yielded strange results, so we did not include them in the article.

### 1.3. Contributions and plan

This article provides answers to these different issues:

- The testbed used for our previous experiments was extended in order to randomly generate class loading orders. Potentially, any order can be generated, although the complete combinatorics is intractable. However, all class loading orders are likely not sensible, and
the statistics are also restricted to concrete class loading orders, according to the assumption that only leaf classes can have instances—“make all non-leaf classes abstract” [9].

- **Perfect class numbering** is now tested in this new setting which better explains the observed odd behavior of PN-\text{and}. Several 2-parameter functions have also been proposed and tested, but only a single one is retained.
- Finally, all of these experiments are repeated with benchmarks of JAVA hierarchies that include precise information about interfaces and abstract, concrete and inner classes.

Furthermore, we present some mathematical properties of PH and PN that are the basis of simple and efficient algorithms, and also prove that PN is optimal in single-inheritance situations.

Overall, these new tests overcome all of our previous reservations about the use of perfect hashing for implementing object-oriented languages. In a dynamic loading setting, we now consider that the technique is efficient, under its PN-\text{and} form, from both space and time standpoints. In contrast, we also think that modulus is too inefficient in terms of time to be further considered. Therefore, although we developed the same formal and empirical framework for modulus, here we only present the bit-wise and results. Thus, perfect hashing should be considered by language implementers (i) for subtype testing in all languages with multiple inheritance, (ii) for implementing interfaces in all languages with multiple subtyping (e.g. JAVA, C#, ADA 2005, etc.). However, further research and tests are required before using perfect hashing for attribute access, hence complete object implementation.

The article is structured as follows. The next section presents the question of object-oriented implementation in a static typing setting. The standard single-subtyping implementation is described and several possible extensions to multiple inheritance are discussed: (i) the subobject-based implementation used by C++; (ii) the coloring approach which amounts to maintaining single-subtyping invariants in a global setting; and finally (iii) the perfect class hashing approach. Section 3 presents definitions and simple mathematical results which provide efficient algorithms and lower bounds that are proven to be the PN value in case of single inheritance. Section 4 presents experiments and statistical results in the context of full multiple inheritance, and then in the context of JAVA interfaces. All these experiments concern space-efficiency. Section 5 briefly describes experiments that have been done in the PRM compiler and summarizes this first empirical assessment of the PH run-time time-efficiency. Related works and current interface implementations are discussed in Section 6. Finally, Section 7 presents conclusions and prospects. An appendix describes algorithms for computing PH parameters.

This article is the continuation of [3]. We tried to make it as self-contained and short as possible, but we sometimes refer to the original article—hereafter it will be abbreviated PHAPST (Perfect Hashing as an Almost Perfect Subtype Test). Interested readers are referred to PHAPST for a more in-depth presentation and discussion of subtype testing and all related topics, and to [10] for a review of object-oriented implementation.

2. FROM SINGLE SUBTYPING TO MULTIPLE INHERITANCE

Single subtyping, i.e. single inheritance and static typing, allows for a straightforward constant-time implementation of the basic object-oriented mechanisms. We first describe this implementation, then explain why it does not easily generalize to multiple inheritance or dynamic typing. Finally, we present some alternatives, namely C++ subobjects, coloring and perfect hashing.

2.1. Single subtyping

2.1.1. Method invocation and object layout. In separate compilation of statically typed languages, late binding is generally implemented with tables that are called virtual function tables in C++ jargon. An object is laid out as an attribute table, with a header pointing at the class table which contains method addresses. Method calls are then reduced to calls to pointers to function through two extra indirections. With single inheritance, the class hierarchy is a tree and the tables implementing a class are straightforward extensions of those of its single superclass. Each class
add, to (a copy of) the tables of its direct superclass, the entries required for implementing the properties, i.e. attributes or methods, that are introduced by the considered class (a class $A$ introduces a property if $A$ defines it as a new signature that does not correspond to any property of its superclasses). Figure 1 presents a diagram of the object layout and method table in this setting, together with the corresponding code sequences in an intuitive pseudo-code borrowed from [6].

This implementation respects two essential invariants: (i) a reference to an object does not depend on the static type of the reference; (ii) the position of attributes and methods in the tables does not depend on the dynamic type of the object. Therefore all accesses to objects are straightforward, but this simplicity requires both static typing and single inheritance. From a spatial standpoint, the object layout is optimal, since there is one field per attribute, with a single extra pointer to the method table, which shares the data common to all direct instances of the considered class. The method tables are not very small, but they are also somewhat optimal. Based on the assumption that the method introduction is uniformly distributed over all classes, the total size of method tables is linear in the size of the class specialization relationship, which is assumed to be reflexive and transitive. In the worst case, this is however quadratic in the number of classes.

2.1.2. Subtype tests. The subtype test amounts to checking at run-time that a class $d$ is a superclass of a class $c$, i.e. $d \preceq c$. Usually $d$ is the dynamic type of an object, and the programmer or compiler wants to check that this object is actually an instance of $c$. In the single-inheritance context, several techniques have been proposed and are commonly used. We present only one of the simplest ones that provides a basis for further generalizations. The technique is known as Cohen’s display and was first described in [11]. It consists of assigning two integers to each class, a unique ID and a non-unique offset. Let the ID of class $C$ be $id_C$ and the offset of $C$ be $\delta(C)$. Each subclass of a class $C$ has $id_C$ stored in its method table at $\delta(C)$. Thus, an object is a direct or indirect instance of a class $C$ if and only if offset $\delta(C)$ in its method table ($tab$) contains $id_C$. This is equivalent to testing that the class, say $D$, that has instantiated the considered object is a subtype of $C$:

$$D \preceq C \Leftrightarrow tab[D, \delta(C)] = id_C$$

Originally, Cohen’s display required a set of tables separate from the method tables, and $\delta(C)$ was the depth of $C$ in the class hierarchy. However, in a statically typed language, these tables can be merged within method tables. Class offsets are ruled by the same position invariant as methods and attributes. This can be thought of as giving each class $C$ a method for checking whether an object is an instance of $C$, i.e. such that its instances can check that they are. The test fails when this pseudo-method is not found. When inlining Cohen’s display in the method table, some precautions are required to avoid bound checks and possible confusion between class IDs and method addresses without consuming unnecessary memory. This is discussed in PHAPST.
2.2. Multiple inheritance

With multiple inheritance, both invariants of reference and position cannot hold together, at least if compilation—i.e. computation of positions—is to be kept separate. For instance, in the ABCD diamond hierarchy of Figure 2, if the implementations of B and C simply extend that of A, as in single subtyping, the same offsets might be occupied by different properties in B and C, thus prohibiting a sound implementation of D. Therefore, invariants or separate computation must be dropped.

2.2.1. Multiple inheritance with static typing and dynamic linking. The ‘standard’ implementation of multiple inheritance in a static typing and separate compilation setting (i.e. that of C++) is based on subobjects. The object layout is composed of several subobjects, one for each superclass of the object’s class. Each subobject contains attributes introduced by the corresponding class, together with a pointer to a method table which contains methods known by the class. Both invariants are dropped, as both reference and position now depend on the current static type. This is the C++ implementation when the virtual keyword annotates each superclass. It is time-constant and compatible with dynamic loading, but method tables are no longer space-linear. Indeed, the number of method tables is exactly the size of the specialization relationship. Hence, the worst-case total table size of a class is now quadratic in the number of superclasses, and the total size for all classes is cubic in the class number. Furthermore, all polymorphic object manipulations—i.e. assignments and parameter passing, when the source type is a subtype of the target type—require pointer adjustments between source and target types, as they correspond to different subobjects. These adjustments can be done with offsets in method tables or explicit pointers in the object layout.

A detailed presentation of this implementation is beyond the scope of this paper. Interested readers are referred to [1] for C++ itself, [10] for a language-independent analysis of the implementation, and [12] for an analysis of multiple inheritance. Anyway, there are three main drawbacks: (i) pointer adjustments represent ubiquitous overhead throughout programs; (ii) subobjects add many compiler-generated fields in the object layout; (iii) the total size of method tables is now, in the worst case, cubic in the number of classes, instead of quadratic. Overall, this seems to be the price to be paid for multiple inheritance in a static typing and separate compilation setting. Finally, the main drawback of this implementation family is that its overhead remains even when multiple inheritance is not used. Therefore, language designers have provided alternative specification and implementation, known as non-virtual inheritance, when omitting the virtual keyword. Non-virtual inheritance gives exactly the same implementation as single subtyping in the case of single-inheritance hierarchies, but it provides degraded semantics of general multiple inheritance, hence preventing sound reusability—for instance, in the ABCD diamond example, the attributes introduced by A would be duplicated in the object layout of D.

2.2.2. Coloring for multiple inheritance with global linking. The general idea of coloring is to keep the two invariants of single inheritance, i.e. reference and position. An injective numbering of attributes, methods and classes verifies the position invariant, so this is clearly a matter of optimization for minimizing the size of all tables—or, equivalently, the number of holes, i.e. empty entries. However, this optimization must be done globally, at compile- or link-time. Thus, the technique is not incremental and not directly compatible with dynamic loading. Coloring has been independently proposed for the three mechanisms that underlie object-oriented implementation.
Perfect class hashing and numbering

Method coloring was first proposed by [13] under the name of selector coloring. Coloring was applied to attributes by [14] and to classes by [15] (under the name of pack encoding). Interested readers are referred to [16] which reviews the approach.

2.2.3. Perfect class hashing for object implementation. So far, the current situation of object-oriented implementation is as follows: (i) single-subtyping implementation meets all requirements, is compatible with dynamic loading but not with multiple inheritance; (ii) coloring, which represents its extension to multiple inheritances, is no longer compatible with dynamic loading; (iii) subobject-based implementation is compatible with both multiple inheritance and dynamic loading; however its space requirements are cubic in the number of classes and ubiquitous pointer adjustments imply marked runtime overhead.

Perfect class hashing represents an alternative approach, first applied to subtype testing. Its principle is as follows. Classes are loaded at run-time in some total order that must be a linear extension (aka topological sorting) of \((X, \preceq)\) — that is, when \(d \prec c, c\) must be loaded before \(d\). As an alternative view, when a class is loaded, its yet unloaded superclasses are recursively loaded. Anyway, when a class \(c\) is loaded, a unique and immutable identifier \(id_c\) is associated with it and the set \(I_c = \{id_d \mid c \prec d\}\) of identifiers of all its superclasses is known. Thus, \(c \preceq d\) iff \(id_d \in I_c\).

It is worth noting that the class ID is some arbitrary integer. Any injective class numbering can work, but the overall implementation may constrain class IDs to be a specific integer subset. Actually, as we shall see, the most promising scheme involves computing the ID of a class according to the IDs of its superclasses. For the moment, let us consider a simple numbering, starting from 0.

The set \(I_c\) is immutable and can be hashed with some perfect hashing function \(ht_c\), i.e. a hashing function that is injective on \(I_c\) [4, 5]. The previous condition becomes: \(c \preceq d\) iff \(ht_c(id_d) = id_d\), whereby \(ht_c\) denotes the hashtable of \(c\). The hashing function family \(ht_c\) is parametrized by the hashtable size \(H_c\), such that \(ht_c(x) = hash(x, H_c)\), where \(hash\) is some low level operation. In PHAPST, we considered modulus (mod) and bit-wise and. Both involve a single machine instruction. The technique is obviously incremental since all hashvalues are immutable and the computation of \(ht_c\) only depends upon \(I_c\). It is also time-constant and inlinable. The only point might be space-linearity, but the conclusions of PHAPST were rather positive if not definitive.

In a static typing setting, the technique can also be applied to method invocation and we proposed, in the aforementioned article, an application to JAVA interfaces. For this, as hashing method identifiers appeared to be over space-consuming, the hashtable associates, with each implemented interface, the address (or offset) of the group of methods that are introduced by the interface. Figure 3 recalls the precise implementation in this context and the corresponding code sequence. The method table is bidirectional. Positive offsets involve the method table itself, organized as with single subtyping. Negative offsets consist of the hashtable, which contains, for each implemented
interface, a two-fold entry. The object header points at its method table via the table pointer. #hashingOffset is the position of the hash parameter (h) and #htOffset is the beginning of the hashtable. At a position \(hv=\text{and}(h, \#\text{hashingOffset})\) in the hashtable, a two-fold entry is depicted that contains both the implemented interface ID, that must be compared to the target interface ID (\#interfaceId), and the address itable of the group of methods introduced by the interface that introduces the considered method. The table contains the address of the function that must be invoked, at the position \#methodOffset determined by the considered method in the method group. This method invocation technique requires static typing. However, it does not require a method to be introduced by a single class. If a method is introduced by several classes (or interfaces, as is possible in JAVA), the method entry will be replicated in each method group, with the same method address.

With rooted hierarchies, a slight optimization is possible. The position of the group of methods introduced by the root can be made invariant, and invoking such a method only requires the SST code sequence. The root can even be removed from the hashtable. Indeed, when the target is statically known, subtype testing is useless if it is the root. In contrast, when the target is not statically known (e.g. for implementing covariance), the test will just require an extra equality test against the root ID.

2.2.4. Incremental coloring (IC). Although coloring is not inherently incremental, its use for subtype testing has been proposed in the context of JAVA interfaces [17]. We called it incremental coloring (IC) in PHAPST, and extended its use to method invocation in the same way as for perfect hashing. This use of coloring with both dynamic loading and multiple inheritance yields marked overhead at load-time, since some recomputation may be required, and at run-time, since these possible recomputations increase the number of memory accesses, while degrading their locality (Figure 4). Overall, our abstract estimation was that incremental coloring should not be better than perfect hashing, or at least not sufficiently better to offset the expected load-time overhead.

2.2.5. Accessor simulation (AS). Some techniques, like perfect hashing and incremental coloring, apply only to method invocation and subtype testing. Accessor simulation is a way of applying them to attribute access. An accessor is a method that either reads or writes an attribute. Suppose that all accesses to an attribute are through an accessor. Then the attribute layout of a class does not have to be the same as the attribute layout of its superclass. A class will redefine accessors for an attribute if the attribute has a different offset in the class than it does in the superclass. True accessors require a method call for each access, which can be inefficient. However, a class can simulate accessors by
The object representation with accessor simulation coupled with class and method coloring must be compared with Fig. 1. The offset of the group of attributes introduced by a class (aGrOffset) is associated with its class ID in the method table, and an attribute position is now determined by an offset (#attrOffset) relative to this attribute group.

Figure 5. Accessor simulation with method coloring

replacing the method address in the method table with the attribute offset. This approach is called field dispatching by [18]. Another improvement is to group attributes together in the method table when they are introduced by the same class. Then one can substitute, for their different offsets, the single relative position of the attribute group, stored in the method table at an invariant position, i.e. at the class color with coloring (Fig. 5) [19, 10]. With PH and IC, the attribute-group offset is associated with the class ID and method-group address in the hash- or color-table, yielding 3-fold table entries.

Accessor simulation is a generic approach to attribute access which works with any method invocation technique; only grouping can be conditioned by static typing, since attributes must be partitioned by the classes which introduce them.

3. PERFECT CLASS HASHING AND NUMBERING

In this section, we formally define perfect class hashing (PH) and numbering (PN) and present simple theoretical results which are the basis for efficient algorithms.

3.1. Definitions

Definition 3.1 (Perfect hashing)
Let $I$ be a non-empty set of integers, and $hash: \mathbb{N} \times \mathbb{N} \to \mathbb{N}$ be a function such that $hash(x, y) < y$ and $hash(x, y) \leq x$ for all $x, y \in \mathbb{N}$. Then, the perfect hashing parameter of $I$ is the least $H \in \mathbb{N}$ such that the function $h$ that maps $x$ to $h(x) = hash(x, H)$ is injective on $I$, i.e. for all $x, y \in I$, $h(x) = h(y)$ implies $x = y$.

The definition is extended to empty sets by considering that $H = 1$ when $I = \emptyset$.

This least integer exists for mod and and (the exact function maps $x$ to and$(x, H - 1)$), but likely not for any hash function. Moreover, $H \geq n$, where $n$ is the cardinality of $I$.

Perfect class hashing (PH). Let $(X, \preceq)$ be a class hierarchy equipped with some injective class identifier $id: X \to \mathbb{N}$. Then, perfect hashing applies to each class $c$ in $X$ by considering the set $I_c = \{id_d | c \preceq d\}$. The resulting parameter $H_c$ is the size of the hashtable (a simple array) that implements class $c$, and for each superclass $d$, this table contains $id_d$ at position $h_c(id_d)$. All other positions $j$ contain any integer $l$ such that $h_c(l) \neq j$. The $hash(x, y) \leq x$ constraint is not strictly necessary, but it is verified by all the tested functions. It also implies that $h_c(0) = 0$ for all $c$. With rooted class hierarchies, 0 is thus a convenient identifier for the root, and it is also a convenient value for empty entries at offset $j > 0$. Conversely, with unrooted hierarchies, any non-null integer can represent an empty entry at offset 0, but this integer must not be used as a class ID.

In the following, the notations $I$, $H$, $h$ and $n$ will be implicitly used in reference to Definition 3.1, with a subscript indicating that set $I$ actually consists of class identifiers.
Proposition 3.2 (Monotonicity)  
Perfect hashing is monotonic, that is, if \( I \subset I' \), then \( H \leq H' \).

The perfect class hashing definition holds for any injective class identifier, for instance a consecutive numbering as classes are loaded. An optimization of perfect class hashing amounts to optimizing the class identifier \( id_c \) in order to minimize the \( H_c \) parameter. Thus, instead of numbering classes as they are loaded, by incrementing a counter, the \( H_c \) parameter is first computed from the superclass identifiers, before computing the best \( id_c \) that fits the resulting hashtable.

Definition 3.3 (Perfect numbering)  
Let \( I' \) be a set of integers, and \( F \) be a set of free integers, disjoint from \( I' \). Then the perfect numbering parameter is defined as the least \( H \in \mathbb{N} \) such that: (i) there is a free integer \( id \) in \( F \), and (ii) \( H \) is the perfect hashing parameter for the set \( I = I' \cup \{ id \} \).

Perfect class numbering (PN). Then, perfect numbering applies to each class \( c \) in \( X \) by considering the set \( I'_c = \{ id_d \mid c \prec d \} \), and \( F \) is the set of integers that are not identifiers of previously loaded classes. In practice, \( F \) is large enough to ensure that \( H \) does not depend on \( F \). The resulting parameter \( H_c \) is the size of the hashtable that implements class \( c \) and \( id_c \) can be defined as the least number in \( F \) such that \( H_c \) is the perfect hashing parameter for \( I_c = I'_c \cup \{ id_c \} \).

The code generated for PN is exactly the same as that for PH (Figure 3), so they have the same time-efficiency. The only difference involves computation of the \( H_c \) and \( id_c \) parameters, and this computation involves only a slight overhead over that of PH (see Appendix A.2). Hence, PN must be preferred to PH if the expected space improvement is effective.

As loading a class triggers the loading of yet unloaded superclasses, it is interesting to extend this definition to the allocation of several identifiers.

Definition 3.4 (Perfect k-numbering)  
Let \( k > 0 \) be an integer, \( I' \) be a set of integers, and \( F \) be a set of free integers, disjoint from \( I' \). Then the perfect k-numbering parameter is defined as the least \( H \in \mathbb{N} \) such that: (i) there is a set of \( k \) free identifiers \( I'' \subset F \), and (ii) \( H \) is the perfect hashing parameter for the set \( I = I' \cup I'' \).

Perfect class k-numbering. Now, perfect numbering applies to each class \( c \) in \( X \) as follows. Let \( X' \subset X \) be the current subset of already loaded classes, \( id : X' \to \mathbb{N} \) an injective class identifier function, and \( F = \mathbb{N} \setminus id(X') \) the set of free identifiers. Let \( c \in X \setminus X' \) be some unloaded class, with \( X'_c = \{ y \in X' \mid c \prec y \} \) and \( X''_c = \{ y \in X \setminus X' \mid c \preceq y \} \). Then, \( I'_c = id(X'_c) \) and \( k \geq 1 \) is the cardinality of \( X''_c \). \( H_c \) and \( I''_c \) are successively computed, and the numbers in \( I''_c \) are finally assigned to classes in \( X''_c \).

The resulting parameter \( H_c \) is the size of the hashtable that implements \( c \), and it is uniquely determined. There are, however, many ways to define \( I''_c \) as a “minimal” set and to map \( I''_c \) to newly loaded superclasses. Moreover, only \( H_c \) is optimized, not other \( H_x \) parameters when \( c \prec x \) and \( x \) is not already loaded. It might be desired that \( c \) is a concrete class, while \( x \) is abstract and does not require a method table. However, an optimization of \( H_x \) might improve the space occupation of other concrete subclasses of \( x \). Therefore, the optimization of each \( H_x \) would be better. The previous notations are extended as follows: for each \( x \in X''_c \), \( X'_x \) and \( X''_x \) represent, respectively, the sets of previously loaded and yet unloaded superclasses of \( x \), and \( H_x \) is the perfect \( |X''_x| \)-numbering parameter of \( id(X'_x) \).

Conjecture 3.5 (Optimal perfect class k-numbering)  
There is an injective mapping \( f : X''_c \to F \) such that the PH parameter of \( id(X'_x) \cup f(X''_x) \) is \( H_x \) for each \( x \in X''_c \).

3.2. Application to bit-wise and

We now focus on the case where \( hash(x, y) = (x, y - 1) \). The effects of bit-wise hashing are not always intuitive, and we present some very simple results that might help readers. Furthermore, perfect class hashing involves a simple lower bound along with a fine optimality condition. Let \( I \)
be a set of \( n \) positive or null integers, \( m \) be an integer that serves as a bit-wise mask in the function \( h(x) = \text{and}(x, m) \), and \( H = m + 1 \) the perfect hashing parameter, i.e. the hashtable size.

**Definition 3.6 (Minimal mask)**

Given a set \( I \) of integers, a mask \( m \) is minimal for \( I \) if \( m \) provides a \( h \) function injective on \( I \) and if switching any 1-bit in \( m \) makes \( h \) non-injective.

The mask corresponding to Definition 3.1 is minimal but all minimal masks do not yield perfect hashing in the sense of this definition.

The next two propositions give lower and upper bounds to the number of 1-bits in the mask, as a function of \( n \). Of course, they yield a lower bound but no upper bound to the mask itself.

**Proposition 3.7 (PH-and lower bound)**

Hashing \( n > 0 \) integers requires a mask with at least \( \log_2 n \) 1-bits, hence \( 2^{\lceil \log_2 n \rceil} \in [n, 2n] \) is a lower bound for the \( H \) parameter.

The proof follows from the fact that a mask with \( k \) 1-bits can discriminate exactly \( 2^k \) integers.  

The point is that the hashtable capacity depends on the 1-bit count of the mask, not on the 1-bit positions, which determine its magnitude, that is, the hashtable size. Therefore, with \( \text{PH-and} \), a hashtable may have a lot of empty entries, while being full in the sense that no other number can be hashed within it. In contrast, with \( \text{mod} \), a hashtable of size \( H \) could hash \( H \) integers. In other words, with \( \text{and} \) and unlike \( \text{mod} \), \( h \) is not surjective in the \([0, H-1]\) range. This certainly accounts for the different behaviors of both hashing functions and, in practice, it prevents us from using the same algorithms for both.

**Proposition 3.8 (PH-and 1-bit count upper bound)**

A mask \( m \) minimal for a set \( I \) with cardinality \( n \) has at most \( n - 1 \) 1-bits, and this upper bound is reachable.

The proof is by induction. It is trivial for \( n = 1 \). Suppose now that it is true for any set of cardinality \( n - 1 \). Let \( I \) be a set of \( n \) integers and \( x \) be the maximum element of \( I \). Suppose that a mask \( m \) with \( n \) 1-bits is minimal for hashing \( I \). Consider now the set \( I' = I \setminus \{x\} \). According to the recurrence assumption, \( m \) is not minimal for \( I' \) and two 1-bits in \( m \) can be switched—let \( m' \) be the resulting minimal mask for \( I' \). As \( m' \) does not make a perfect hashing function on \( I' \) (otherwise we get the proof), there is a single \( y \in I' \) that agrees with \( x \) on all 1-bits of \( m' \). Hence, \( x \) and \( y \) must differ on the two bits that are 1 in \( m \) but only one is required to make a perfect hashing mask since \( y \) is unique. Hence, \( m \) is not minimal. The upper bound is reachable when each number differs from all others by exactly one 1-bit—for instance, if \( I = \{2^i \mid i = 0..n - 1\} \).

The following proposition gives an upper bound as a function of the maximum element of \( I \).

**Proposition 3.9 (PH-and magnitude upper bound)**

A minimal mask \( m \) is strictly less than \( 2^{\lceil \log_2(\max(I)+1) \rceil} \leq 2 \max(I) \) and the \( (2 \max(I)-1) \) bound is reachable.

The minimal mask is bounded by the integer formed by all the 1-bits of all integers in \( I \). Thus, an upper bound is formed by a chain of \( k \) consecutive 1-bits, where \( k - 1 \) is the rank of the leftmost 1-bit in \( \max(I) \). In the worst-case, all these 1-bits are required to form the mask—for instance, if \( k = n - 1 \) and \( I = \{0\} \cup \{2^i \mid i = 0..n - 2\} \).

**Proposition 3.10 (PH-and global upper bound)**

Let \( X \) be a class hierarchy with \( N \) classes and an injective identifier \( id : X \rightarrow [0, N - 1] \), then \( \sum_{i=1..N} 2^{\lceil \log_2(i+1) \rceil} \) is an upper bound of \( \sum H_c \).

This is a corollary of Proposition 3.9. However, as \( 2^{\lceil \log_2(i+1) \rceil} \) is strictly greater than \( i \), this upper bound is in range \([N(N+1)/2, N(N+1)]\), hence it is too high to be useful.

Given a mask \( m \), the \( m+1 \) hashtable entries can be partitioned into three sets: the entries occupied by the input numbers \( I \); unusable entries that do not fit with the mask \( m \) and can be allocated for any data; free entries that could be occupied by new numbers. The next two propositions explain it.
Proposition 3.11 (PH-and unusable entries)
Let \( m \) be a mask and \( k \) be the number of 1-bits of \( m \). Then the corresponding hashtable contains \( m + 1 - 2^k \) entries in which no number can be hashed.

Let \( m' \) be the mask formed by complementing the significant bits of \( m \), i.e. 1-bits of \( m \) are 0 in \( m' \), and 0-bits of \( m \) are 1 in \( m' \) if they are on the right of some 1-bit. Then, for any \( j \in [1..m-1] \) such that \( \text{and}(m', j) \neq 0 \), \( h^{-1}(j) = \emptyset \).

\[ \square \]

Proposition 3.12 (PH-and completion)
Let \( I \) be a set of \( n \) integers and \( m \) be the mask that forms a perfect hashing function on \( I \). Let \( k \) be the number of 1-bits of \( m \). When \( n < 2^k \), there is an infinity of sets \( J \) of cardinality \( 2^k - n \), disjoint from \( I \), such that \( m \) forms a perfect hashing function on \( I \cup J \).

This is a direct consequence of Proposition 3.7. Among the \( 2^k \) combinations of 1-bits of \( m \), only \( n \) are used by \( I \) and the remaining is free and can be used for integers in \( J \).

This means that any set can be completed to reach a \( 2^k \) cardinality while keeping the same optimal bit-wise mask. The perfect numbering algorithm follows from the next proposition.

Proposition 3.13 (Unbounded PN-and)
Let \( I' \) be a set of integers of cardinality \( n' \), and \( F = \mathbb{N} \setminus I' \) be the set of free integers. Let \( m' \) be the PH mask of the set \( I' \). Then the PN mask \( m \) is \( m' \) if the bit-wise mask \( m' \) has at least \( \log_2(n' + 1) \) 1-bits. Otherwise, \( n' = 2^k \) and \( m \) is \( m' \) with its least-weight 0-bit switched to 1. As a corollary, \( H \leq 2H' \), whereby \( H' \) is the perfect hashing parameter of the set \( I' \).

The first part is a straightforward consequence of Proposition 3.12. The corollary is implied by the fact that, in the worst-case, the least-weight 0-bit is just after the leftmost 1-bit.

This ensures that there is a subset of 1-bits in \( m \) that forms the offset of an empty entry in the new table. This proposition holds if the set \( F \) of free numbers is large enough, for instance for any finite class hierarchy with an initial set of free integers which consists of all 16- or 32-bit integers. It is worth noting that both \text{and} and \text{mod} functions present the same \( H \leq 2H' \) upper bound.

Moreover, PN-and has an interesting property—namely it is optimal, i.e. \( H_c = 2^\lceil \log_2(n_c) \rceil \), for all classes \( c \) in single inheritance, that is, such that \( c \) and all its superclasses have a single direct superclass. This is a limited converse of Proposition 3.7.

Proposition 3.14 (PN-and optimum)
Perfect numbering is optimal, i.e. \( H_c = 2^\lceil \log_2(n_c) \rceil \), for every class \( c \) in single inheritance, i.e. such that \( c \) and all its superclasses have a single direct superclass.

This means that, in single inheritance, all masks are formed by a chain of \( 2^\lceil \log_2(n_c) \rceil \) rightmost consecutive 1-bits. The proof is by induction on \( n_c \), i.e. class depth. Let \( c \) be the considered class, and \( c' \) be its single direct superclass, with \( n_{c'} = n_c - 1 \). By induction, \( H_{c'} = 2^\lceil \log_2(n_{c'} - 1) \rceil \). If \( n_c = 2^k + 1 \), \( H_c = 2^k \) and the mask is full, then an extra bit is required and \( H_c = 2^{k+1} \). Otherwise, according to Proposition 3.12, the mask has some free room for an extra identifier and it can be inherited by \( c \), hence \( H_c = H_{c'} = 2^\lceil \log_2(n_c) \rceil \).

This proposition holds under the same condition as Proposition 3.13 on the set \( F \) of free integers.

The interesting point is that this perfect class 1-numbering optimum does not depend on the exact class identifiers, as any greedy choice provides a global optimum. Intuitively, all 1-bits of the mask form a rightmost prefix of the mask and are “inherited”. With multiple inheritance, when a class \( c \) has two direct superclasses \( c_1 \) and \( c_2 \), each one with a mask formed by a chain of, respectively, \( k_1 \) and \( k_2 \) 1-bits with \( k_1 \leq k_2 \), the \( k_2 \) 1-bits of the highest mask cannot always discriminate between the extra identifiers inherited from \( c_1 \), either because there are not enough 1-bits, or because there are some multiple inheritance conflicts, that is, the same 1-bits are used by each class for discriminating its proper ancestors. Hence, extra bits are required, that can be much more leftward and yield an exponential increase in the mask. This explains why PN and PH can be erratic with bit-wise \text{and} in multiple inheritance. Proposition 3.14 is important because all multiple inheritance class hierarchies are mostly in single inheritance—see, for instance, the statistics in [16]. Hence, PN-and should be optimal on a large part of each hierarchy and this should counterbalance its worst-case behaviour on the multiple inheritance core.
Finally, the optimality of PN-and with single inheritance yields formal space-linearity.

**Proposition 3.15 (Space linearity)**
For any single-inheritance hierarchy, the PN-and ratio \( \rho = \sum_c H_c / \sum_c n_c \) is in interval \([1, 2]\). The upper bound is asymptotically reachable, that is, for all \( \epsilon > 0 \), there are single-inheritance class hierarchies such that \( \rho > 2 - \epsilon \).

Consider, for instance, a chain \( A_{2^k} \prec A_{2^{k-1}} \prec \ldots \prec A_1 \) of \( 2^k \) classes, with \( A_{2^k} \) having \( x \) direct subclasses. For any such subclass \( c \), \( n_c = 2^k + 1 \), \( \log_2 n_c = k + 1 \) and \( H_c = 2^{k+1} \). So, in this framework, \( x \) can be large enough to make \( \rho > 2^{k+1}/(2^{k+1} + 1) \)

In practice, the optimal ratio \( \sum_c 2^\log_2 (n_c) / \sum_c n_c \) is always lower than 1.5 in our multiple inheritance benchmarks, and \( \sum_c H_c / \sum_c n_c \) can be very close to it in the best cases, but markedly higher in the worst cases (see Table II).

**Discussion on perfect numbering.** Its mathematical definition (Definition 3.3) corresponds to a simple algorithm (Proposition 3.13) which does not depend on the set \( F \) of free integers. In practice, class identifiers are bounded by the underlying integer implementation, e.g. 16- or 32-bit integers, and it might happen that both mathematical and algorithmic definitions differ when no free integers fit the hashtable in the algorithm. However, such a situation would be very unlikely, as the greatest allocated ID is never greater than 3 times the class number \( N \) in our tests.

Whereas the perfect class hashing definition (Definition 3.1) results in a \( \sum_c H_c \) value which is uniquely determined by the input class numbering, perfect numbering does not provide the same overall determinism, apart from the single-inheritance optimum. Indeed, the perfect numbering parameter provides only a step-by-step optimal \( H_c \), and there are several possible \( id \)s for the newly loaded class \( c \). Selecting the least \( id \) amounts to a greedy optimization that does not ensure any formal minimization of \( \sum_c H_c \). The point is even more complicated with \( k \)-numbering, since Proposition 3.13 no longer provides a step-by-step optimum. Conjecture 3.5 is quite attractive, since it relies on computation of the PH optimum for each currently loaded class.

**Proposition 3.16 (Optimal perfect class \( k \)-numbering)**
Conjecture 3.5 does not hold for bit-wise and.

Although it may work for some class loading orders of very large hierarchies (e.g. the Eclipse hierarchy in Table VII), we found simple counter-examples of the existence of a solution.

In the general case, the perfect class \( k \)-numbering problem is over-constrained and has no optimal solution, i.e. regardless of the classes that might be loaded afterwards. Only greedy optimizations can thus be considered. Moreover, while PN is markedly better than PH on average, we also observed some class loading orders where PH was better than PN. Hence, the heuristics might well be improved by further research.

### 3.3. Other hashing functions

Without loss of generality, the perfect hashing approach might be improved with small variations such as: (i) 2-probe tests, (ii) other 1-parameter 1-operation hash functions, or (iii) 2-parameter 2-operation functions. In PHAPST, we tested 2-probe quasi-perfect hashing (qPH-and); the space improvement was not marked enough to offset the expected time overhead. We could not imagine other 1-parameter operations than \texttt{mod} and \texttt{and}. Two-parameter hash functions might be an improvement from the space standpoint, but the extra instruction would likely degrade the time efficiency even if it takes a single cycle. Overall, the only efficient combination that we could imagine is bit-wise and with rightward shift, in order to truncate the mask and remove all trailing zeros. This is denoted by \texttt{PH-and+shift}.

**Definition 3.17 (PH-and+shift)**
Let \( I \) be a non-empty set of integers. Then, the PH-and+shift parameters are defined as the pair \( (H_1, H_2) \) such that \( H_1 \) is the least integer such that there is \( H_2 \) such that the function \( h \) that maps \( x \) to \( h(x) = \texttt{and(shift}(x, H_2), H_1 - 1) \) is injective on \( I \).
An algorithm is proposed in Appendix A.3. Moreover, \textit{and} and \textit{shift} can also be combined with perfect numbering. The technique, denoted PN-\textit{and+shift}, amounts to substituting PH-\textit{and+shift} for PH-\textit{and} in the algorithm for PN-\textit{and}, with the only difference that the switched 0-bit is selected between existing 1-bits, if possible, instead of being the least-weight 0-bit.

4. SPACE-EFFICIENCY TESTS

Our original testbed consists of a set of large-scale benchmarks that are commonly used in the object-oriented compilation community, together with a set of programs for computing various parameters that are either characteristics of the class hierarchies or the size requirement of various implementation techniques. This testbed thus simulates the memory occupation of these various techniques. The simulation is exact and reproducible, except for techniques that rely on heuristics like coloring or depend on some hidden run-time inputs, for instance class-loading orders. Besides PHAPST, this testbed has been used in different simulations [10, 16].

For this article, we have extended this testbed in several directions, by computing random class loading orders; by testing the new perfect hashing variants described in Section 3; by restricting class loading orders to leaf or concrete classes; finally, by extracting a set of new JAVA benchmarks that retain all class modifiers, unlike the benchmarks that are commonly used in the literature.

4.1. Class loading at random

First of all, we computed various perfect hashing parameters while loading classes at random, with the same benchmarks as in PHAPST. They represent full-multiple-inheritance hierarchies that have been extracted from real libraries of different languages. Only the class specialization relationship is considered here. With JAVA hierarchies, classes and interfaces are not distinguished from each other. Table 1 presents, for each hierarchy, its class number $N$, then the average value of $n_c$, the number of superclasses of $c$, per class (i.e. $|\preceq|/N$).

The precise testbed involves generating a random class loading order by selecting a class at random from the set of $\preceq$-\textit{maximal} yet unloaded classes until all classes are loaded. Then, for each class loading order, all PH parameters are computed. This is repeated thousands of times.

The statistics presented in Tables II and IV present minimum, average and maximum values over randomly generated class loading orders. All these numbers are normalized ratios $\rho = \sum_c H_c / \sum_c n_c$ that represent the space-linearity factor, since $\sum_c n_c = |\preceq|$. The order of magnitude

<table>
<thead>
<tr>
<th>class number</th>
<th>avg $n_c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBM-SF</td>
<td>8793</td>
</tr>
<tr>
<td>JDK1.3.1</td>
<td>7401</td>
</tr>
<tr>
<td>Java.1.6</td>
<td>5933</td>
</tr>
<tr>
<td>Orbix</td>
<td>2716</td>
</tr>
<tr>
<td>Corba</td>
<td>1699</td>
</tr>
<tr>
<td>Orbacus</td>
<td>1379</td>
</tr>
<tr>
<td>HotJava</td>
<td>736</td>
</tr>
<tr>
<td>JDK.1.0.2</td>
<td>604</td>
</tr>
<tr>
<td>Self</td>
<td>1802</td>
</tr>
<tr>
<td>Geode</td>
<td>1318</td>
</tr>
<tr>
<td>Vortex3</td>
<td>1954</td>
</tr>
<tr>
<td>Cecil</td>
<td>932</td>
</tr>
<tr>
<td>Dylan</td>
<td>925</td>
</tr>
<tr>
<td>Harlequin</td>
<td>666</td>
</tr>
<tr>
<td>Unidraw</td>
<td>614</td>
</tr>
<tr>
<td>PRMcl</td>
<td>479</td>
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<tr>
<td>Lov-obj-ed</td>
<td>436</td>
</tr>
<tr>
<td>SmartEiffel</td>
<td>397</td>
</tr>
<tr>
<td>Total</td>
<td>38784</td>
</tr>
</tbody>
</table>

Table I. Statistics on class and superclass numbers
<table>
<thead>
<tr>
<th>Benchmark</th>
<th>lower bound</th>
<th>PH-and</th>
<th>PH-and+shift</th>
</tr>
</thead>
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<tr>
<td>IBM-SF</td>
<td>1.42</td>
<td>2.0</td>
<td>3.8</td>
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<td>JDK 1.3.1</td>
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<td>1.3</td>
<td>1.6</td>
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<td>Java 1.6</td>
<td>1.27</td>
<td>1.4</td>
<td>1.6</td>
</tr>
<tr>
<td>Orbis</td>
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<td>1.3</td>
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<tr>
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<td>1.19</td>
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<td>1.8</td>
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<tr>
<td>Orbcus</td>
<td>1.20</td>
<td>1.4</td>
<td>2.0</td>
</tr>
<tr>
<td>HotJava</td>
<td>1.31</td>
<td>1.3</td>
<td>2.1</td>
</tr>
<tr>
<td>JDK 1.0.2</td>
<td>1.30</td>
<td>1.3</td>
<td>1.4</td>
</tr>
<tr>
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<td>1.9</td>
<td>3.9</td>
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<tr>
<td>Geode</td>
<td>1.48</td>
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</tr>
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<td>1.7</td>
<td>2.7</td>
</tr>
<tr>
<td>Cecil</td>
<td>1.28</td>
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<tr>
<td>Dylan</td>
<td>1.35</td>
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<td>1.6</td>
</tr>
<tr>
<td>Harlequin</td>
<td>1.32</td>
<td>1.8</td>
<td>2.9</td>
</tr>
<tr>
<td>Undrillad</td>
<td>1.27</td>
<td>1.3</td>
<td>1.0</td>
</tr>
<tr>
<td>PRMcl</td>
<td>1.31</td>
<td>1.3</td>
<td>1.5</td>
</tr>
<tr>
<td>Lov-obj-ed</td>
<td>1.38</td>
<td>2.2</td>
<td>3.8</td>
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<tr>
<td>SmartEffel</td>
<td>1.41</td>
<td>1.4</td>
<td>1.9</td>
</tr>
</tbody>
</table>

The top-left numbers represent, respectively, the total sample count (i.e., the number of class-loading orders that have been generated at random) and an estimation of the growth rate of the maximum values, expressed as the number of new maximal records per thousand. The first column presents the lower bound for bit-wise hashing, as a ratio $\sum c_2(\log_2(n_c)) / \sum n_c \in [1, 2]$. All other numbers are ratios $\rho = \sum H_s / \sum n_c$, whereby the sum is obtained for all classes, $H_s$ is the hashtable size and the denominator is the cardinality of $s$. The minimum, average ($\mu$) and maximum values of $\rho$ are presented for each technique, and the standard deviation $\sigma$ is also displayed for PH-and+shift. The ‘PH-and ref’ column recalls the tests presented in PHAPST. Italic type represents “bad” PN behaviour, namely PN max that are respectively greater than the corresponding PH min or avg. In contrast, bold type represents “good” behaviour, namely PN max that are less than PH min. Line ‘Total’ represents, for all columns, the same ratios as for each benchmark, but computed from the sum of the corresponding parameters on all benchmarks, i.e., when a column depicts some $p_b/q_b$ ratio for each benchmark $b$, the last line is $\sum p_b / \sum q_b$.

Table II. Statistics on random class loading orders

The top-left numbers represent, respectively, the total sample count (i.e., the number of class-loading orders that have been generated at random) and an estimation of the growth rate of the maximum values, expressed as the number of new maximal records per thousand. The first column presents the lower bound for bit-wise hashing, as a ratio $\sum c_2(\log_2(n_c)) / \sum n_c \in [1, 2]$. All other numbers are ratios $\rho = \sum H_s / \sum n_c$, whereby the sum is obtained for all classes, $H_s$ is the hashtable size and the denominator is the cardinality of $s$. The minimum, average ($\mu$) and maximum values of $\rho$ are presented for each technique, and the standard deviation $\sigma$ is also displayed for PH-and+shift. The ‘PH-and ref’ column recalls the tests presented in PHAPST. Italic type represents “bad” PN behaviour, namely PN max that are respectively greater than the corresponding PH min or avg. In contrast, bold type represents “good” behaviour, namely PN max that are less than PH min. Line ‘Total’ represents, for all columns, the same ratios as for each benchmark, but computed from the sum of the corresponding parameters on all benchmarks, i.e., when a column depicts some $p_b/q_b$ ratio for each benchmark $b$, the last line is $\sum p_b / \sum q_b$.

The first conclusion is that simple perfect hashing with consecutive numbering is really bad. In most benchmarks, the results presented in PHAPST (column ‘PH-and ref’) are close to the PH-and minimum values, hence rather optimistic. This is disappointing, since we expected some average behavior when taking an arbitrary order. Moreover, the PH-and variations are marked, as the max/min ratio is greater than 20 for the five largest benchmarks. In the worst cases, PH-and can be dramatically inefficient—$\rho$ can be greater than 200—and it only depends on the class loading order which is a problem input.

In contrast, perfect numbering produces markedly better results. The PH-and average is always lower than the PH-and minimum, except again in a few cases; however, the PH-and maximum...
Table III. Statistics on class, leaf and superclass numbers

<table>
<thead>
<tr>
<th>Class</th>
<th>Leaf</th>
<th>Superclass</th>
<th>Avg n&lt;sub&gt;c&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBM-SF</td>
<td>8793</td>
<td>6001</td>
<td>8.8</td>
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<tr>
<td>JDK1.3.1</td>
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<td>Java.1.6</td>
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<td>311</td>
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<td>38784</td>
<td>27633</td>
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is generally higher than the PH-and average, though markedly lower than its maximum. PN-and and PH-and+shift yield similar results but PN-and+shift improves on both, especially in the worst cases (maximum values); however, for many benchmarks, the worst-case size remains more than twice that of the PH-and estimation in PHAPST.

These results are rather negative. Even when considering the most compact variant, i.e. PN-and+shift, worst cases (column 'max') remain over space-consuming for many benchmarks (especially IBM-SF and Geode). As PN-and and PN-and+shift are certainly the most promising techniques, we produced more statistics for them. The positive point is that their standard deviations σ is rather low. For all benchmarks but IBM-SF and Geode, σ is indeed less than 2 for PN-and, and than 1 for PN-and+shift. As an empirical verification of the Bienaymé-Chebyshev inequality, we also observed that, for each benchmark, the proportion of class loading orders that exceed μ + 2σ is less than 5%, for both PN-and and PN-and+shift.

In PHAPST, we did not conclude on PN because our first results seemed erratic. Actually PH-and itself is erratic, and PN is always a marked improvement over PH. However, the technique remains space-inefficient with bit-wise and in the worst-case class loading orders, even though these worst cases are not frequent. But now, the PN-and average is always better than the PH-and number in PHAPST. Therefore, restricting class loading orders might make PN-and acceptable in all cases. Indeed, all linear extensions are not sensible class loading orders. Thus, modelling class loading orders and testing perfect hashing on “sensible” class loading orders are still open issues.

4.2. Random leaf-class loading

The main issue with these first tests is that the bit-wise and hashing function appears to be highly dependent on the class loading order. Class loading depends on precise implementation of runtime systems like virtual machines, and the class loading order depends on the considered program behavior. However, without loss of generality, one may assume that class loading is always triggered by the need to instantiate a yet unloaded class, as all other uses of yet unloaded classes could be made lazy. Hence, only concrete, i.e. non-abstract, classes must be ordered and abstract classes are only inserted when needed in concrete-class orders, in such a way that superclasses are always loaded before subclasses.

4.2.1. Principle. As our benchmarks do not record whether a class is abstract or concrete, we consider an assumption that is often advocated, namely “make all non-leaf classes abstract” [9]. This is indeed a common methodological recommendation; see for instance [23]. Table III presents
the leaf number and the average $n_c$ per leaf for all benchmarks. On average, according to this assumption, there would be a little more than one abstract class to four classes. This is an upper bound of the actual ratio, since it is meaningless for a leaf class to be abstract, and a little more than the ratio of one to six advocated by [24] (however, at a time when class hierarchies were markedly smaller).

An informal algorithm is as follows. An unloaded leaf $c$ is selected at random, and the set $X_c = \{ x \mid c \subseteq x \}$ is partitioned into two subsets $X'_c$ and $X''_c$, that contain, respectively, already loaded and yet unloaded superclasses, including $c$. $X''_c$ is then ordered in a top-down linear extension. For each class $x \in X''_c$, in this order, $id_x$ and $H_x$ are computed according to the considered algorithm (PH or PN). In the PN case, instead of considering each class in $X''_c$ separately, an alternative is a global optimization of $X''_c$ numbering, with perfect $k$-numbering (Definition 3.4). There are actually pros and cons for both approaches, since PN-and is optimal for single-inheritance classes (Proposition 3.14) and the simple optimal problem (Conjecture 3.5) has no solution in the general case. However, our experiments show that global optimization gives slightly better results.

There is however a huge number of linear extensions, namely factorial of the number of leaves, which may be as many as 6000 in the largest benchmarks (Table IV). The number of orders is thus above $10^{20000}$, obviously, the point cannot be to compute exact statistics. It is possible to somewhat reduce this combinatorial explosion by partitioning the set of leaves according to their parents; indeed, the way leaves with the same parents are ordered with each other is not significant. Therefore, a two-step selection avoids equivalent orders. Sets are represented as lists, and a set of equivalent leaves is first selected at random, then the first element in this set is picked. The number of orders is now $\text{fact}(\sum_{i=1}^k p_i) \times \prod_{i=1}^k \text{fact}(p_i)$, instead of $\text{fact}(\sum_{i=1}^k p_i)$, whereby $k$ is the number of equivalence classes and $p_i$ the cardinality of each of them. In this way, the combinatorics reduces to about $10^{12000}$ orders; this is better but still exhaustively intractable ($10^{30}$ nanoseconds represent a round upper bound for the solar system life). Though we were unable to prove that there were no frequent worse cases, our experiments tended to quickly converge, that is, after some thousands of tests, the rate of new maximal records was rather low, i.e. less than one to a thousand, the average remained stable and the growth of the maximum extremely slow. Leaf partition is also used for the statistics in Table II.

<table>
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<tr>
<th>PH-and+shift</th>
<th>PH-and</th>
<th>PN-and</th>
<th>useful PN-and</th>
<th>lower bound</th>
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<td>max</td>
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<td>1.4</td>
<td>1.7</td>
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</table>

All numbers have the same meaning as in Table II, but \( \sum_i \) sums are now restricted to leaf classes, and the useful part of PN-and is presented instead of PN-and+shift. In the lower bound and PN ‘min’ columns, bold type also indicates that the optimum value is reached.
and shift method tables. As the resulting ratios do not markedly differ between the two forms, we only present \(k\) numbering a set of classes as a whole. We tested 1- and the best orders. Regarding PN, it is not clear whether the optimization takes advantage of loading and explore all possible orders (the combinatorics is too large), whereas leaf-class ordering focuses on original class order was a depth-first top-down linear extension, hence a possible leaf-class ordering.

In this setting, all the PH-and values are markedly improved (Table IV), and the PHAPST ratios in Table II are now close to the average PH ratios in Table IV. This is explained by the fact that our original class order was a depth-first top-down linear extension, hence a possible leaf-class ordering. The fact that the minimum values are greatly improved confirms that our random testbed cannot explore all possible orders (the combinatorics is too large), whereas leaf-class ordering focuses on the best orders. Regarding PN, it is not clear whether the optimization takes advantage of loading and numbering a set of classes as a whole. We tested 1- and \(k\)-numbering; on average, the resulting \(H_c\) parameter does not significantly differ, regardless of whether all yet unloaded classes are numbered as a whole (\(k\)-numbering), or whether they are numbered one by one by successive applications of PN in a top-down ordering (1-numbering). Hence, the improvement over PH is mostly due to the selection of specific class loading orders.

The most interesting observation is that the erratic behavior of PH-and has been markedly reduced (the max/min ratio is now always less than 10, instead of 35) and most worst-case orders have been ruled out (the worst-case ratio \(\rho\) hardly exceeds 50, instead of 200). On most benchmarks, the maximum PN-and values in Table IV are now lower than the corresponding PH-and minimal values in Table II. The standard deviation is now less than 1 on all benchmarks; the proportion of leaf-class orders that exceed \(\mu + 2\sigma\) is still less than 5%; and \(\mu + 2\sigma\) is always less than 5. Moreover, for the useful PN-and ratio, \(\mu + 2\sigma\) is always less than 3, and the maximal values are markedly lower than that of PN-and. Hence, a large part of the extra memory occupation is not wasted but can be reused for specific static allocations. Overall, PN-and produces excellent results. On average, its space-occupation ratio is close to 2 for most benchmarks, and perfect hashing is thus obtained at a cost similar to that of efficient linear probing. The ratio is often very close to its \(2^\lceil \log_2(\text{nc}) \rceil\) lower bound, though it does not exclude a risk of a bad-case class loading order. However this risk is low, as bad cases are infrequent; and it is not fatal, as the resulting memory occupation would be large but not unreasonable. Actually, it would seem that there is no need for any other hashing function, and PN-and+shift is not sufficiently better to offset its expected time-overhead.

Of course, it would be interesting to confirm that leaf-class orders are representative of actual class loading orders. The reality is likely midway between both models. On the one hand, programmers
Perfect class hashing and numbering

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<th>number</th>
<th>total</th>
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From top to bottom, three benchmark groups: complete libraries, SPECjvm2008 and DaCapo benchmarks. The first columns represent the class and interface numbers. The next section represents the average and maximum numbers of superclasses and implemented interfaces, per concrete class. For instance, on average on the whole Java-6-sun library, a concrete class extends 3.1 classes and implements 1.4 interfaces, and the maximum values are, respectively, 10 and 20. Moreover, the maximum value of the union of both is 29. \( n_c \) is now the number of interfaces implemented by concrete class \( c \). 'Total' lines include all benchmarks of each group.

Table VI. Statistics on classes and interfaces for JAVA benchmarks

and class hierarchies only partly comply with Meyers’ directive. On the other hand, concrete classes would represent only a possibility, since a concrete class can be instantiated in some programs, but the class may also remain abstract in many others. This concerns only non-leaf classes, since a non-instantiated leaf would never be loaded.

4.2.3. Loading a subset of all classes. However, apart from some small benchmarks like PRMcl and SmartEiffel, our benchmarks do not represent actual programs, but whole class libraries. Therefore, a real program would only load a subset of these libraries, and it was reported in [17] that hardly one thousand classes were actually loaded from similar several-thousand class hierarchies. Therefore, we also tested PH-and variants when only a subset of all classes was loaded. The principle is the same as random class ordering, except that class loading ends when a fraction of all classes are loaded. In practice, we tested it with fractions running from 20% to 60%. The class number limitation improves all PH parameters, but the bad features of PH-and remain, though at a smaller scale. Table V presents only the statistics of PN-and in this new setting, for the restriction to leaf classes. The table shows marked improvement as the class number decreases.

4.3. Application to JAVA-like multiple subtyping

Perfect hashing can also be applied to JAVA-like interfaces in a restricted form. In JAVA, the extends relationship between classes is in single inheritance, thus Cohen’s display and usual single-subtyping implementation works, and it is more efficient than PH for classes (Section 2.1). Therefore, only the implements relationship between classes and interfaces needs to be hashed.
All numbers are ratios $\rho = \sum_c n_c / \sum_c H_c$, whereby the sum is done on concrete classes only (the denominator is the cardinality of the implements relationship, restricted to concrete classes). As $n_c$ can be zero, the lower bound is now $\rho = \sum_c o_c / \sum_c n_c$, where $o_c = 1$ if $n_c = 0$ and $2^{\lceil \log_2(n_c) \rceil}$ otherwise.

Table VII. Statistics of PN- and for JAVA benchmarks over random concrete-class loading orders.

The exact relation is not the explicit implements relationship, but rather its closure with class specialization, i.e. if $B$ extends $A$ and $A$ implements $I$, then $B$ implements $I$. Overall, only classes require a hashtable and only interfaces are numbered and hashed. In this setting, the hashtable is used for subtype tests whose target is an interface, and for method invocations when the receiver is typed by an interface and the method is not introduced by the $\text{Object}$ root.

We have adapted all perfect hashing and perfect numbering functions to JAVA interfaces and tested them on new class hierarchies that have been extracted from DaCapo [25] and SPECjvm2008 (http://www.spec.org/jvm2008/) benchmarks, or full Java 1.6 libraries§. Table VI presents the number of classes and interfaces for each benchmark, along with the average numbers of extended classes and implemented interfaces. Each benchmark represents an approximation of the set of classes that could be loaded when the corresponding archive (.jar) file is loaded. These sets are generated by meta-programming, and are closed under two relationships: specialization (i.e. implements and extends) and method signature. The only classes that might defy this analysis are those that are instantiated within a method while being only a strict subclass of the collected classes. These new benchmarks provide extra information about classes, namely if they are abstract, along with their nesting and visibility. Therefore, the algorithm described in Section 4.2 has been improved in order to take the new information into account. Concrete classes are considered, instead of leaf-classes, and complex loading rules are taken into account for inner classes. Indeed, JAVA inner classes are archived in separate .class files, and they can be loaded independently of their enclosing class. However, non-static inner classes are only instantiable from their enclosing class or a subclass of it. In contrast, static inner classes can be instantiated from the enclosing class, the package or the whole hierarchy according to their visibility. Finally, inner classes can be instantiated.

§These new benchmarks are available on http://www.lirmm.fr/~morandat/benchmarks.
through invocation of a virtual or static method (both of a class), or by initializing a static variable (of a class or even an interface). Overall, the enclosing class or some class of the same package must often be loaded before the considered inner class can be loaded.

The application of Definition 3.1 is now slightly different, and resembles an application to unrooted hierarchies, since Object is considered as a class. Moreover, a class does not encode itself in its hashtable. Thus, for a class $c$ that does not implement any interface other than Object, $I_c = \emptyset$, $n_c = 0$ and $H_c = 1$, and the single table entry must then contain any non-null integer that is not an interface ID. Proposition 3.14 optimum now occurs when interfaces are in single inheritance, and with the difference that $H_c = 1$ instead of $2^{\lceil \log_2(n_c) \rceil}$ when $n_c = 0$. Therefore, the global optimum is unbounded. Finally, the PH algorithms are applied in the same manner as described in Section 4.2.1, except that $X'_c$ and $X''_c$ are now sets of, respectively, already loaded and still unloaded interfaces implemented by concrete class $c$.

Table VII presents the statistics of PN-and over these random concrete-class loading orders. The results are clearly satisfactory. Apart from Java-6-sun and Eclipse which represent complete libraries that will never be loaded as a whole in a single program, all other benchmarks represent set of classes, a large part of which could be loaded in a single run. Their PN-and parameter is quite good, namely lower than 2 on average, and $\mu + 2\sigma$ rarely exceeds 2. These data will be further discussed in Section 6. The overall conclusion is that PN-and would also provide a scalable implementation for JAVA-like interfaces.

5. TIME-EFFICIENCY TESTS

This section presents the results of experiments that provide empirical assessment of the PH runtime efficiency and complement the PHAPST abstract assessment.

5.1. Abstract assessment

In Driesen’s [2001] computational model, the cycle count of a code sequence is a function of the processor latencies for different machine instructions like load ($L$ latency) or branch ($B$ latency). We add to these two parameters the integer division latency ($D$) for PH-mod. Besides these latencies, processors are characterized by their instruction-level parallelism. Overall, each code sequence can be associated with a cycle count which represents the time necessary to execute the sequence when all data are cached. Driesen’s model is purely static. Therefore, cache misses are not considered, whereas each one would represent a latency far greater than the cycle count of the sequences considered here. Moreover, well-predicted branchings cost nothing, whereas a misprediction costs $B$ cycles. These two dynamic features, i.e. cache misses and mispredictions, can however be considered as a risk. For instance, incremental coloring introduces extra cache-miss risks, since three memory areas are concerned, instead of one in all other techniques. Moreover, it also introduces extra misprediction risks for subtype testing, since it requires an extra bound check. There are thus two different ways of failing (instead of one). Table VIII presents the results of these analyses for the considered techniques.

5.2. Empirical runtime assessment in the PRM testbed

We also implemented PH in the compiler of the PRM language, and tested its time efficiency on a real program, namely the PRM compiler itself. In this setting, perfect hashing is compared to different techniques on a variety of processors. These experiments are original, as they compare different implementation techniques, in a common framework that allows a fair comparison, with all other things being equal. In this section, we only summarize the results of these tests. Interested readers are referred to [7] for a more complete report.

Tested techniques. The PRM testbed has been used for comparing a variety of techniques from which we extract four families:
Link-time coloring (Section 2.2.2), as a generalization of single-subtyping implementation, represents a reference, i.e. the zero point of our scale; this is the best known technique that does not involve global optimization, but it requires global linking; it is also the technique used in JAVA for implementing classes, hence for method invocation and subtype testing when the receiver’s type or the target type is a class, and for all attribute accesses.

- Compile-time coloring, with type analysis for detecting monomorphic call sites, represents another reference, close to the optimum, and the difference between link-time and compile-time coloring can be considered as the unit of our scale.
- All PH variants have been tested. As the tests concern only time-efficiency, PH and PN do not differ apart from method table sizes, but this slight difference should not affect cache misses.
- The incremental version of coloring (IC) (Section 2.2.4) represents an alternative to PH in spite of its expected load-time overhead.

All techniques are considered for application to the three basic mechanisms required by object-oriented programming: (i) subtype tests, (ii) method invocation and (iii) attribute access. For attribute access, two variants are considered, with attribute coloring (AC) or accessor simulation (AS). The tests with accessor simulation provide an assessment of the use of the considered technique (IC or PH) for implementing full multiple inheritance, for instance as an alternative to the subobject-based implementation of C++. Attribute coloring is the most efficient implementation, that of JAVA for instance, so combining PH or IC with attribute coloring provides an assessment of the considered techniques when applied to JAVA interfaces. Of course, this assessment requires some extrapolation, as PRM classes are not JAVA interfaces. However, this experiment is much more demanding than actual JAVA tests, since the tested technique here represents all method invocations instead of being reserved to interface-typed receivers. Therefore, if the overhead is low in the PRM testbed, it should be even lower with actual JAVA programs.

Tested program. We implemented all these techniques in the PRM compiler, which is dedicated to exhaustive assessment of various implementation techniques and compilation schemes [26, 7]. The test involves meta-programming, as the benchmark program is the compiler itself, which is written in PRM and compiles the PRM source code into C code. The PRM compiler is actually not compatible with dynamic loading, and the code for PH or IC has been generated at link-time exactly as if it were generated at load-time, and the usual optimizations of the PRM compiler (see [26]) are deactivated. The class load ordering does not matter since here we only consider time measurement, and the class hierarchy is small enough (compared to the largest benchmarks in Section 4) for making size variations insignificant.

The tested program makes intensive usage of the compared object-oriented mechanisms. The respective numbers of method invocations and attribute accesses are about 1.8 and 2.6 billion. The program size is also significant; it is the PRMcl benchmark in Tables I to IV. Load-time
PERFECT CLASS HASHING AND NUMBERING

Each sub-table presents the results for a cluster of processors, and each number is the unweighted mean of the corresponding measures for all processors in the considered cluster. Each row describes a method invocation and subtype testing technique, and the two columns represent the overhead vs. pure coloring (link-time MC-AC), respectively with attribute coloring (AC) and accessor simulation (AS).

Table IX. Comparison of execution time according to implementation techniques and processors

<table>
<thead>
<tr>
<th>Technique</th>
<th>AC</th>
<th>AS</th>
<th>AC</th>
<th>AS</th>
<th>AC</th>
<th>AS</th>
</tr>
</thead>
<tbody>
<tr>
<td>compile-time MC</td>
<td>-1.0</td>
<td>-0.3</td>
<td>-1.0</td>
<td>0.5</td>
<td>-1.0</td>
<td>0.2</td>
</tr>
<tr>
<td>link-time MC</td>
<td>0.0</td>
<td>0.5</td>
<td>0.0</td>
<td>1.2</td>
<td>0.0</td>
<td>1.1</td>
</tr>
<tr>
<td>IC</td>
<td>0.5</td>
<td>1.2</td>
<td>1.1</td>
<td>3.1</td>
<td>0.7</td>
<td>2.9</td>
</tr>
<tr>
<td>PH-and</td>
<td>0.4</td>
<td>1.3</td>
<td>1.3</td>
<td>4.3</td>
<td>0.5</td>
<td>2.9</td>
</tr>
<tr>
<td>PH-and+shift</td>
<td>0.6</td>
<td>1.8</td>
<td>1.5</td>
<td>4.9</td>
<td>0.8</td>
<td>4.2</td>
</tr>
<tr>
<td>PH-mod</td>
<td>3.0</td>
<td>9.2</td>
<td>6.0</td>
<td>17.9</td>
<td>1.9</td>
<td>9.3</td>
</tr>
<tr>
<td>Unit #processor</td>
<td>19.0%</td>
<td>12.8%</td>
<td>10.7%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster</td>
<td>4</td>
<td>2</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Time-measurement conditions. Tested variants differ only by the underlying implementation technique, with all other things being equal. The compilation testbed is deterministic, and two compilations of the same program by the same compiler executable produce exactly the same executable. Moreover, when applied to a program, two compiler variants produce exactly the same code, and it has been verified with the `diff` command on both C and binary files. Overall, the effect of memory locality should be roughly constant, apart from specific effects due to the considered techniques. The tests were performed on several processors from different families and manufacturers: Intel® Pentium™, AMD® Athlon™, SUN® UltraSPARC™ or PowerPC™. All processors run under various 32-bit versions of Unix (Linux, Solaris, BSD).

5.3. Results and discussion

The results presented in [7] are here abstracted in two ways (Table IX). First, the scale is changed and the unit is now the difference in percentage between link-time and compile-time colorings. Second, a few processors are added, and the whole processor set is partitioned into 3 clusters by a \( k \)-mean algorithm, with each processor being represented by a 10-dimension point in this new scale.

- Cluster 1 gathers processors of different families (UltraSPARC, PowerPC, and Pentium) that are generally older than other processors (from 2001 to 2006). The unit is high and PH-and overhead is about half a unit with AC and slightly more than a unit with AS.
- Cluster 2 consists of AMD processors (from 2003 and 2006). The unit is average and all overheads are exaggerated.
- Cluster 3 consists of recent Pentiums (from 2006 to 2009). The unit is low, PH-and overhead is slightly higher than in cluster 1 with AC, but markedly higher with AS. Actually, cluster 3 resembles cluster 1 for AC, and cluster 2 for AS.
- PH-and is generally slightly better than IC with AC, not with AS, but the differences are not marked. Moreover, the extra instructions of PH-and+shift entails low extra overhead.
- In contrast, the overhead of PH-mod is markedly higher than that of PH-and. Coupled with the abstract cycle count, this is the reason why we decided to give up PH-mod and we do not present the theoretical and experimental analyses that we have undertaken.

These empirical results can be partly explained by comparing them to the theoretical predictions in Table VIII.

- For method invocation and subtype testing, PH-and adds a few loads from a memory area that is already used by the reference technique, hence without extra cache misses, plus a
few 1-cycle instructions; these extra cycles represent real overhead that is, however, slight in comparison with the overall method call cost; PH-and+shift adds a load and a shift that are partly or even totally done in parallel, hence with slight overhead.

- PH-mod adds high integer division latency, commonly estimated at about 20-25 cycles [27], hence much more expensive than extra loads; $D$ latency can be estimated at $D \approx 1 + \frac{F \Delta}{1.8}$, whereby $\Delta$ is the difference between the PH-mod and PH-and measures (in seconds), $F$ is the processor frequency (in GHz), and 1.8 is the number of invocations (in billions). This estimation is about 25-35 cycles (as expected), or even more, for cluster 1 and 2, but less than 10 for processors in cluster 3. This is confirmed by the fact that most processors in cluster 3 use the Radix–16 algorithm, rather than Radix–4 which was used in older Pentiums [28].

- Accessor-simulation replaces the single coloring load by a sequence that adds several loads from a memory area (i.e. the method table) that was not already used—hence it increases the cache miss risks with marked overhead. This overhead increases with the cost of the underlying method invocation technique, from method coloring to PH-and and PH-mod. The measures provide the same kind of estimation of $D$ at $1 + \frac{F \Delta}{4.3}$, whereby 4.3 is the total number of mechanism invocations, in billions. On all processors, the estimation is slightly higher than the previous one by a few cycles.

- In Table VIII, the difference between IC and PH-and is not significant, and the tests confirm it, as extra indirects in IC and extra computations in PH-and cause similar overheads. However, IC implies access to extra memory areas, a first one for the color and a second one for the recomputable color table, hence extra cache misses are expected, that are likely underestimated in our tests. However, we have no explanation for the fact that IC is slightly better than PH-and with AS, whereas the relation is inverted with AC (eg in cluster 1). Overall, as the PH load-time cost is also far lower than that of IC, PH-and must be preferred to IC.

Overall, the results of these tests are quite satisfactory, and PH-and is even better than expected for method invocation and subtype testing on many processors. It should thus provide very high efficiency for implementing interfaces in JAVA and .NET. When also used for attribute access, the overhead becomes less reasonable, and the results of PH-and with AS can actually be considered as bad. However, preliminary tests with the C++-like subobject-based implementation (Section 2.2.1) show that PH-and with AS is not less efficient than subobjects [29].

Of course, the validity of such experiments that rely on a single program may be questioned. This single-benchmark limitation is inherent to our experimentation. The PRM compiler is the only one that proposes such versatility in the basic implementation of object-oriented programs. The compensation is that the language has been developed with this single goal in mind, and its compiler is the only large-scale program written in PRM. This is however a large, fully object-oriented program, which intensively uses the basic mechanisms that are tested. Moreover, the experiments compare two implementations with all other things being equal and, apart from IC, these implementations present similar processor cache-miss risks. With IC, the effect of cache misses is likely underestimated, since all color tables are here allocated in the same memory area, whereas load-time recomputations should scatter them in the heap. Therefore, when the differences are marked, their sign should hold for all programs and only the order of magnitude should vary. In contrast, the comparison between IC and PH cannot conclude, because differences are rather low on most processors, and increased cache-miss risks might penalize IC. However, we prefer PH because of their respective load-time behaviours.

6. RELATED WORKS

Perfect class hashing is proposed here as a potential alternative to the implementation approaches that are currently undertaken in C++ and JAVA-like languages. The key features of these languages are static typing, multiple inheritance (possibly reduced to multiple subtyping) and dynamic loading or linking. For these two language families, there is some evidence that their implementations
do not meet our constant-time, linear-space and inlining criteria. The original subobject-based implementation of C++ has been briefly reviewed in Section 2.2.1. It yields marked overhead, with many compiler-generated pointers in the object layout and pointer adjustments at runtime. Moreover, before perfect hashing, there were no known constant-time subtype tests for this implementation. From the memory occupation standpoint, subobjects present a cubic worst case, and large scale experiments show that the spatial cost can be markedly higher than for all other techniques [10]. Moreover, the aforementioned comparison between PH- and subobjects show that they provide similar time-efficiency [29]. The rest of this section will now consider the case of JAVA and .NET languages.

6.1. JAVA-like runtime systems

Modern runtime systems are dedicated to languages like JAVA and C# that use only a limited form of multiple inheritance, namely multiple subtyping of interfaces. In this setting, the implementation of classes is that of single-subtyping, and interfaces are usually implemented in rather ad hoc ways. Currently, almost all known JAVA interface implementations do not meet one of two requirements for constant-time or incrementality. For instance, the proposal of [17] is not inherently incremental, so it yields potentially high load-time overhead, together with extra run-time indirections. All other techniques reported in [2, 30, 31, 17] are not time-constant. The only exception might be the proposal of using large direct access tables in SableVM [32], but it does not meet the linear-space requirement. However, the empty slots of these huge tables are used for allocating other data, and we do not know the extent to which it offsets the nonlinear size. The same trick is possible for PH- and according to Proposition 3.11. However, in SableVM, it cannot be used for subtype testing, for which empty entries represent failure, i.e. useful information. Therefore, the total occupied size is linear in the product of class and interface numbers.

Non-constant-time techniques generally rely on searching and caching. For instance, the interface that introduces the invoked method is cached in the method table. Then, for each method invocation, the interface ID is compared with the cache. A cache hit thus provides the right interface-table address, whereas a cache miss requires the interface ID to be searched in a data structure, before filling the cache. However, even with cache hits, the cycle count for method invocation is $3L + B + 2$ (Figure 6), which is midway between PH- and $(4L + B + 3)$ and single subtyping $(2L + B)$. This cache can also be used for subtyping tests, as in [31] where the underlying search is naively sequential. The cycle count is now $2L + 2$ with cache hits, but not lower than PH- and with cache misses. As for all cache-based techniques, the resulting efficiency depends on the actual cache-hit rate. For instance, empirical cache-hit rates between 50% and 99% are reported in [17] for subtype testing. This rate can be improved with multiple caches, whereby each interface is statically
assigned to a specific cache. We tested this possibly original approach in the PRM testbed with quadruple caches and an overall cache-hit rate about 80%. The experiments show that this improves only on very inefficient underlying techniques, that is PH-mod, on all processors but cluster 3 [7]. Moreover, caches markedly increase memory occupation in both method tables and code sequences.

An alternative involves polymorphic inline caches (PIC) [33] which consist of testing the receiver’s dynamic type against a few expected types. It yields a series of conditional branchings. Under the CWA, all types can be exhausted, and this is called binary tree dispatch (BTD) [34, 35]. Experiments in the PRM testbed show that the best techniques involve coloring for highly polymorphic sites and BTD for other sites [7]. However, under the OWA, all types cannot be exhausted, and the code sequence must include a case for cache misses, eg PH. The resulting efficiency still depends on the cache-miss rate, hence on the way the expected types are computed.

Overall, the reasons for the apparent efficiency of these runtime systems must be searched elsewhere than in the used implementation technique. A first reason is that, in JAVA-like languages, the implementation question concerns only interfaces, i.e. a subset of all classes, instead of all classes as in PRM. Hence, cached entities are fewer and cache-hit rates higher. This is explicit in Tables II and VII, where the average number of implemented interfaces is seldom greater than 2, while the average number of superclasses is about 7.

However, the main reason for this apparent efficiency likely follows from the implied compilation scheme, namely just-in-time (JIT) compilers, rather than from the used implementation techniques. Indeed, JIT compilers rely on a provisional closed-world assumption that allows the compiler to perform many optimizations that are currently valid. For instance, a method call can be handled as a static call, or as a class invocation (instead of an interface invocation), because this optimization is possible in the current state of the system. Later on the compilation of some new code may force the compiler to recompile the optimized code because the underlying optimizations are no longer valid. With this approach, the runtime system must first warm up before reaching its cruising speed whereby the code is optimized and no further recompilation is required.

6.2. Comparison with perfect hashing

In the Java benchmarks in Table VII, abstract classes are known and the statistics represent the real memory occupation of a program that would load exactly those classes and interfaces. Of course, there is no program that would load the complete Java-6-sun or Eclipse libraries, but these benchmarks show insight into possible material limitations. For instance, in the Eclipse benchmark, the maximum number of superclasses, i.e. the longest inheritance chain, is 16, the maximum number of implemented interfaces is 28, and the maximum number of extended classes or implemented interfaces is 36. This means that there are programs that require an efficient implementation for these specific cases.

With a midsize benchmark like Derby, the PN-and ρ ratio (i.e. μ + 2σ) is not greater than 2.1 for 93% of concrete-class-loading orders. As the average number of implemented interfaces is 2.1, this means that the entire interface implementation requires about 4.4 (i.e. 2.1 × 2.1) entries per class plus a word for the $H_e$ parameter. Each entry is two-fold, and this amounts to 10 words per class. Moreover, the average number of superclasses is 3.1. This is the average size of Cohen’s display. Overall, about 13 words per class allow for both subtype tests and interfaces. When considering only subtype testing, 9 words are enough.

It is worthwhile comparing these concrete cases with the technique used for subtyping tests in the HotSpot virtual machine [31]. The proposed technique involves a data structure with 3 words for the cache and 8 words for fixed-size Cohen’s display, whereas the tested benchmarks do not require more than 5 words for Cohen’s display (8 words are proposed for the sake of “robustness”). Moreover, a so-called “secondary list” is used for implementing interface cache misses and extra superclasses that exceed Cohen’s display fixed size (this is what we call the “underlying implementation”), but it is neither precisely described, nor counted in these 11 words. Overall, the proposed technique is not better than PH on midsize benchmarks like Derby, and it cannot take actual Java hierarchies into account, as the class inheritance chains can be markedly longer than 5 or even 8, and cache misses would make the technique quite inefficient in the case of
Eclipse classes with 36 superclasses and implemented interfaces. The length of the “secondary list” would be 28, and sequential search would be painful. Actually, it would seem that the technique is only tailored to the SPECjvm98 (http://www.spec.org/jvm98/) benchmarks, which are markedly smaller than SPECjvm2008 and DaCapo, without considering the need to scale up gracefully on larger programs.

Both techniques could be compacted by using 16-bit integers, but our experiments with the PrM testbed show that the gain in method table size is offset by a slight increase in code size. Therefore, it should be considered only for processors that are equipped with specific half-word instructions. Overall, we consider that perfect hashing ensures several advantages over the technique proposed in [31]. PN-and is indeed constant-time, inlinable, scalable, and it addresses both subtype testing and method invocation mechanisms.

Another comparison can be made with the technique proposed for method invocation in JikesRVM (formerly Jalapeño) [2]. Method identifiers are hashed in a fixed-size hashtable whose entries contain addresses of stub functions that handle collisions. This paper presents benchmarks with two values, namely 5 and 40, for the hashtable size. In JikesRVM 3.1.0, this value is 29. The first value is very low, and certainly too low to be time-efficient. In contrast, 29 and 40 are markedly higher than PH average.

Finally, it is quite hard to compare PH and PIC. Indeed, in the best cases (no cache misses) PIC is far better, but in the worst cases (no cache hits), the winner is PH. There are actually several points against PIC: (i) before JIT compilation, profiling is needed to identify the frequent types that must be expected; (ii) this profiling must be continued in the compiled code if the code is intended to remain optimized in case of a change of frequent types; (iii) the code sequences are markedly longer. It is likely that (ii) significantly degrades the efficiency. Overall, PIC makes the compilation highly complicated in contrast with the simplicity of PH.

7. CONCLUSIONS AND PROSPECTS

Our previous works on perfect class hashing [3] concluded that the technique was promising and deserved further consideration. However, a more in-depth assessment was also required. The tests presented here now allow us to draw some new and much more definitive conclusions about the time and space efficiency of perfect class hashing.

To our knowledge, PH is, together with C++ subobject-based implementation, the only constant-time technique for method invocation that allows for both multiple inheritance and dynamic loading at reasonable spatial cost. Furthermore, its space requirements are roughly linear in the cardinality of the specialization relation and far lower than that of subobjects. PH is also the only constant-time and linear-space technique for subtyping tests that allows for both multiple inheritance and dynamic loading. These assertions hold when PH is used under the perfect class numbering variant, with both mod and and hashing functions. However, the time constant of mod is too high, even on recent processors which use the Radix-16 division algorithm. Therefore, PH-mod should be reserved to processors with very efficient integer division, that is if any exists. In contrast, the time efficiency of PH-and is better than expected when used for method invocation and subtyping tests. Its overhead vs. coloring is not insignificant, but it seems to be the price to be payed for implementing interfaces when they are intensively used, and the tested alternatives like caching are actually markedly less scalable. PN-and is thus certainly a very good solution for implementing JAVA interfaces—this is actually the best solution that we are aware of. PN-and+shift could also be envisaged from the time standpoint, but it does not represent a neat spatial improvement on PN-and when leaves are assumed to be the only concrete classes.

The main contribution of this article is, however, from the space standpoint. We have proposed and tested, here, (i) an optimized approach, namely perfect class numbering; (ii) a more systematic testbed based on random class loading; and (iii) a more realistic model based on concrete leaf classes. It follows from these new tests that PH-and is inherently erratic and over space-consuming in the worst-case class loading orders. Perfect class numbering provides a marked improvement over plain perfect hashing, with both mod and and hashing functions. However, in spite of its
optimality in single inheritance, it is still over space-consuming in the worst cases, but combining it with shift provides an improvement that is slight on average but marked in the worst cases. Finally, the assumption that all non-leaf classes are abstract makes the behaviour of PH and PN markedly more regular; in practice, the PN-and worst cases are now infrequent and they yield reasonable memory overhead. Furthermore, a large part of the extra memory occupation could be used for allocating static data.

Moreover, when it is also used for attribute access, the perfect hashing overhead becomes unreasonable on many processors—hence, it might not be a direct alternative to the C++ subobject-based implementation.

The prospects of this work are manifold:

- The first experiments in PRM led us expect that PH-and should be very efficient for implementing JAVA interfaces—thus experiments in a production virtual machine represent the next step to confirm it. Regarding runtime systems and JIT compilers, the point at issue cannot be to substitute perfect hashing for both caching and JIT compilers. The proposal is rather to use perfect hashing, i.e. PN-and, as the single underlying technique for implementing interfaces, and to adapt current JIT compilers to this technique. The observed efficiency should not be drastically changed, but our claim is that the resulting implementation should be both simple and scalable. Constant time is also an essential argument for real-time systems.

This simple goal is, however, not so easy to meet. Indeed, to be efficient, perfect hashing requires a very simple but uncommon method-table layout, namely with bidirectionality and per interface grouping. Moreover, an efficient implementation of Cohen’s display, without bound checks, requires allocation of method tables in dedicated areas [3]. For instance, we tried to implement PH in JikesRVM [2, 30], but we gave up because the VM overall implementation does not meet these requirements, and the expected improvement would be offset by unnecessary indirections and bound checks.

- Experiments with the PRM testbed must be generalized to other processor families, manufacturers or architectures—especially for testing more efficient integer divisions and 64-bit implementations; other programs than the PRM compiler would also be of great interest.

- Full-multiple-inheritance languages do not provide such large and rich benchmarks as JAVA-like languages, and information about concrete classes is generally not available. Therefore, our experiments in full multiple inheritance rely on the assumption that only the leaf ordering matters. This assumption should be confirmed either empirically or theoretically. However, empirical experiments are usually at a smaller scale than simulations. For instance, in [17], a report on experiments on JAVA programs based on almost ten-thousand class libraries shows that each run hardly loads one thousand classes and interfaces. Bit-wise and perfect class hashing and numbering are always very good on one-thousand class benchmarks. It would thus seem that the scalability of these techniques can be proven only by simulations or theoretical means. This is, however, somewhat unavoidable, since scalability is the ability to gracefully scale up over future programs that do not yet exist. Conversely, empirical experiments, in spite of their smaller scale, could provide models of class load orderings that would be less theoretical than the slogan “make all non-leaf classes abstract”.

As PN-and provides a technique for implementing all three basic object-oriented mechanisms—namely method invocation, subtype testing and attribute access—that is efficient at least for the first two, it is essential to compare PH with the C++ subobject-based technique. Both techniques are indeed the only ones that ensure constant-time method invocation in a multiple inheritance and dynamic loading framework, while being inherently incremental. We are currently conducting these tests in the PRM testbed, and our aforementioned preliminary results show that perfect hashing coupled with attribute coloring is much more efficient than subobjects [29]. However, this combination of PH and coloring is not compatible with dynamic loading, and the compatible combination, with accessor simulation instead of coloring, is not better than subobjects. The tests must be completed for precise comparisons and firm conclusions, but the overall conclusion is that

both implementations are not that efficient. In practice, the efficiency of the C++ language comes from the fact that virtual inheritance is only marginal in C++ programs.

Despite this apparent inefficiency, an efficient solution to full multiple inheritance implementation in a dynamic loading setting might be based on perfect hashing, accessor simulation and JIT optimizations, and there is some evidence that such a solution could be markedly more efficient than current subobject-based implementations. We now envisage designing a virtual machine for the PRM language, with a JIT compiler adapted to full multiple inheritance and perfect hashing. In this context a basic JIT optimization would be to use the single-subtyping implementation instead of perfect hashing until it becomes impossible, then to recompile the concerned methods. Preliminary simulations show that the single-subtyping implementation could be used on most attribute accesses, and on many method invocations, thus making a full multiple-inheritance virtual machine as efficient as a JAVA-like program making intensive use of interfaces [36]. Moreover, the recompilation cost would remain markedly low.

An extra contribution of this article concerns the methodology used for simulating load-time effects at random on very-large-scale hierarchies (Section 4). This approach is currently used to simulate the effects of alternative decisions in the design of the PRM virtual machine. It provides a preliminary assessment of recompilation costs and overall expected efficiency in the worst case. Such random simulations could also be used for assessing the scalability of many optimizations currently used in virtual machines and JIT compilers.

APPENDIX A. ALGORITHMS

The computation of perfect hashing parameters is presented in a simple COMMON LISP code [37], as in PHAPST. In the following algorithms, simplicity takes priority over efficiency.

A.1. Perfect class hashing with bit-wise and

PH-and could be based on a function that takes successive numbers until it finds a perfect hashing function. This works well with modulus but, with bit-wise and, the PH parameter closely depends on the significant bits of the input numbers. Indeed, with bit-wise and, a hashtable can have empty entries that cannot be filled by any number, and the useful entries depend only on the 1-bits of the mask (Proposition 3.7). Therefore, an algorithm working at the bit level is certainly preferable. By the way, it is also more efficient as it yields a logarithmic complexity.

The basic version of PH-and first computes a mask composed of all discriminant bits, i.e. bits which are not 0 or 1 in all numbers. The resulting integer gives a perfect hashing function since all integer pairs in the input differ by at least one bit in the mask. Then the function checks each 1-value bit by decreasing weight and switches the bit when it is not required for injectivity.

```lisp
defun ph-and (ln) ;; LN is an integer list
  (if (null (cdr ln))
    1
    (let ((mask (logxor (apply #’logior ln) (apply #’logand ln))))
      ;; MASK consists of all discriminant bits
      (let ((mask (logxor mask (ash 1 b))) ;; NEW = MASK with switched bit
        (when (ph-and-p ln new) (setf mask new))))
      finally return (1+ mask))))
```

In this code, logxor, logior and logand are COMMON LISP integer functions for bit-wise operations: exclusive and inclusive or, and and. (highest-bit n) gives the position of the leftmost 1-bit of a positive integer n. (logbitp n b) tests if the b-th bit of n is 1. (ash n b) shifts n left by b positions (when b is positive).

```lisp
defun ph-and-p (ln mask)
  (loop for i in ln
```
for \( hv \) = (logand i mask) do
    (if (eql (aref *ht* hv) mask)
        (return nil)
    (setf (aref *ht* hv) mask))
finally return t)

The \textit{ph-and-p} function checks that its \textit{mask} parameter forms a perfect hashing function for the \textit{ln} identifier list. \textit{*ht*} is a global variable bound to an array that is presumed to be large enough and is initialized only once in \textit{ph-and} (this is a slight improvement over the PHAPST version). A successful call to \textit{ph-and-p} leaves the hashtable filled by numbers in \textit{ln} on length \( hc \). However, \textit{ph-and} often finishes with an unsuccessful call to \textit{ph-and-p}. Therefore a subsequent call to \textit{ph-and-p} may be required in the following functions, in order to reset the hashtable.

With consecutive numbering of class IDs, the \textit{ph-and} complexity is \( O(id_c + n_c \log id_c) \). Indeed, \( id_c \) is the maximum of \textit{ln} input, and \textit{mask} is bounded by \( 2id_c \) (Proposition 3.9). The \textit{*ht*} initialization is thus in \( O(id_c) \), and the main loop involves \( O(\log id_c) \) calls to \textit{ph-and-p}, which itself is in \( O(n_c) \).

\section*{A.2. Perfect class numbering}

\textit{PN-and} is slightly more complicated than \textit{PH-and}. The algorithm is based on Proposition 3.13 as follows. It first computes the \textit{PH-and} parameter for the input list of numbers. If the bit-wise \textit{mask} does not contain enough 1-bits (\texttt{logcount} is the function that returns the 1-bit count of its argument), the rightmost 0-bit is switched. Then \textit{*ht*} is reset by \textit{ph-and-p} and the function ends by selecting free identifiers. It returns two values\footnote{The function here uses the \texttt{COMMON LISP} feature called \textit{multiple values}, with the \texttt{values} special form. Another special form that is used hereafter, i.e. \texttt{multiple-value-bind}, binds a list of variables to such multiple values.}, namely the set \( idc \) of new identifiers and the hashtable size \( hc \).

\begin{verbatim}
defun pn-and* (ln n)
    (let ((mask (1- (ph-and ln))))
        (loop for b from 0 by 1
            while (> (+ (length ln) n) (ash 1 (logcount mask))) ;; when there are not enough 1-bits
                unless (logbitp b mask) ;; rightmost 0-bit
                do (setf mask (logxor mask (ash 1 b)))) ;; switch
        (ph-and-p ln mask);
        (values (compute-least-free-ids-and n mask) (1+ mask))))

defun compute-least-free-ids-and (n mask)
    (loop for i in *free* until (= n 0)
        for hv = (logand i mask)
        unless (eql (aref *ht* hv) mask)
        collect (progn (setf (aref *ht* hv) mask)
                        (decf n)
                        i)))
\end{verbatim}

*\textit{free*} is a data structure that represents the set of free identifiers. The loop on \textit{*free*} is actually slightly more complicated, as \textit{*free*} is a more efficient data structure than a simple list, e.g. a union of intervals. It might also be optimized to take advantage of the bit-wise structure of free numbers. However, it is certainly not straightforward, and this naive loop is likely a good tradeoff between simplicity and efficiency.

Finally, the newly allocated class identifiers must be assigned to the set \( X''_c \) of newly loaded classes, and there are many ways to do this. In this respect, (\textit{pn-and* ln n}) is certainly not exactly equivalent to iterating (\textit{pn-and* ln 1}) \( n \) times. Their respective results must depend, however, on the structure of the class hierarchy, and only the latter form ensures the single-inheritance optimum. Consider, for instance, the \textit{ABCD} diamond of Figure 2. Suppose that only \( A \) is already numbered, so all other classes must be numbered as a whole (i.e. \( n=3 \)). Proposition 3.14 states that (\textit{pn-and ln}) is optimal in the presence of single inheritance. Hence, it should be preferred for \( B \) and \( C \), especially if they have many subclasses. On the contrary, (\textit{pn-and* ln ...)}
n) should likely be preferred for $D$, especially if $D$ has many subclasses. So the overall efficiency depends on the whole class hierarchy, which is unpredictable when the considered classes are loaded. Our experiments are actually not conclusive, as the differences are not significant.

Without considering the free-number computation, $pn$-and* adds $O(\log n'_{id'} \log id'_c)$ to the $O(n_{id'} \log id'_c)$ $ph$-and complexity, where $id'_c$ is now the maximum inherited class ID, and $n_{id'}'$ and $n_{id''}$ are the respective numbers of already loaded and yet unloaded classes (i.e. $n_{id''}$ is $n$). The free-number computation is in $O(id'_c)$, where $id'_c$ is the maximum returned free number, which will be assigned to the current class $c$. Of course, this analysis does not provide formal complexity, since it is a function of the output, instead of the input. However, as we empirically observed that the maximum allocated identifier does not exceed $3N$, where $N$ is the total class number, this provides rough complexity on average.

A.3. Bit-wise shift

The combination of $and$ and $shift$ (Definition 3.17) relies on a similar algorithm, but it is now linear instead of logarithmic. It is indeed exponential in the 1-bit count. The function is like $ph$-and except that the 1-bits of the mask are not scanned in the same order. When both leftmost and rightmost 1-bits are switchable, both solutions are computed and the best one is returned. Otherwise, the left- or right-most 1-bits are switched while it is possible, and finally the remaining inner 1-bits are switched in decreasing weight. We present a simplified version. The $best$-and+s function returns a single bit-wise mask whose rightmost 1-bit gives the parameter $H_2$, and $H_1$ is obtained by shifting the mask by $H_2$ positions.

```lisp
defun ph-and+s (ln)  
(if (null (cdr ln))  
  (values 1 0) ;; returns $H_1$ and $H_2$  
  (let ((mask (logxor (apply #'logior ln) (apply #'logand ln))))  
    (lbit))  
  (fill *ht* nil :start 0 :end (1+ mask)) ;; resets *HT*  
  (setf mask (best-and+s mask)  
    lbit (lowest-bit mask))  
  (values (1+ (ash mask (- lbit))) lbit))))

defun best-and+s (ln mask)  
(let* ((hbit (highest-bit mask))  
   (lbit (lowest-bit mask))  
   (newh (logxor mask (ash 1 hbit)))  
   (newl (logxor mask (ash 1 lbit))))  
(if (ph-and-p ln newh)  
  (if (ph-and-p ln newl)  
    (let ((mh (best-and+s ln newh))) ;; exponential  
      (ml (best-and+s ln newl))) ;; part  
      (if (<= (ash mh (- (lowest-bit mh)))  
        (ash ml (- (lowest-bit ml))))  
        (mh ml))))  
  (best-and+s ln newh)  
  (best-and+s ln newl)  
  ;; switches inner 1-bits by decreasing weight  
  (loop for b from (1- hbit) by 1 downto (1+ lbit)  
    when (logbitp b mask) do  
    (let ((new (logxor mask (ash 1 b))))  
      (when (ph-and-p ln new)  
        (setf mask new))))  
  (finally return mask))))))
```

The combination with perfect numbering gives the following algorithm. The difference with respect to $pn$-and* is that the switched bit is first searched in the range between leftmost and rightmost 1-bits, if any, and otherwise the mask is extended, preferably on the right.

```lisp
defun pn-and+s* (ln n)  
(multiple-value-bind  
  (hc mask lbit hbit) (ph-and+s ln)  
  (loop while (> (+ (length ln) n) (ash 1 (logcount mask))) do
```
(loop for b from lbit by 1 ;; rightmost
  unless (logbitp b mask) do ;; inner 0-bit
    (when (and (> b hbit) (> lbit 0))
      (setf b (1- lbit))
    )
    (setf mask (logxor mask (ash l b))) ;; switch
    (hc (1+ (ash mask (- lbit))))
  (return)))
(ph-and-p ln mask) ;; ends like PN-AND*
(values (compute-least-free-ids-and n mask) hc))

REFERENCES


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