NEW TECHNIQUES FOR COMPUTING THE IDEAL CLASS GROUP AND A SYSTEM OF FUNDAMENTAL UNITS IN NUMBER FIELDS

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ABSTRACT. We describe a new algorithm for computing the ideal class group, the regulator and a system of fundamental units in number fields under the generalized Riemann hypothesis. We use sieving techniques adapted from the number field sieve algorithm to derive relations between elements of the ideal class group, and $p$-adic approximations to manage the loss of precision during the computation of units. This new algorithm is particularly efficient for number fields of small degree for which a speed-up of an order of magnitude is achieved with respect to the standard methods.

1. INTRODUCTION

Let $K = \mathbb{Q}(\theta)$ be a number field of degree $n$ and discriminant $\Delta$. In this paper, we present fast methods for computing the structure of the ideal class group of the maximal order $\mathcal{O}_K$ of $K$, along with the regulator and a system of fundamental units of $\mathcal{O}_K$.

Class group and unit group computation are two of the four principal tasks for computational algebraic number theory postulated by Zassenhaus (together with the computation of the ring of integers and the Galois group). In particular, they occur in the resolution of Diophantine equations. For example, the Pell equation

$$T^2 - \Delta U^2 = 1, \quad T, U \in \mathbb{Z},$$

boils down to finding the fundamental unit in a real quadratic number field of discriminant $\Delta$ (see [13]). In addition, the Schäffer equation

$$y^2 = 1^k + 2^k + \ldots + (x - 1)^k, \quad k \geq 2,$$

can be solved using solutions to the Pell equation [17]. Unit computations are key ingredients in solving almost all Diophantine equations, for example when solving Thue equations [3]. On the other hand, the computation of the ideal class group $\text{Cl}(\mathcal{O}_K)$ of a number field $K$ allows in particular to provide numerical evidence in favor of unproven conjectures such as the heuristics of Cohen and Lenstra [9] on the ideal class group of a quadratic number field, Littlewood’s bounds [21] on $L(1, \chi)$, or Bach’s bound on the minimal bound $B$ such that ideals of norm lower than $B$ generate the ideal class group. The class group enters also into the computation of
the Mordell-Weil group of elliptic curves with the descent method, or the Brauer
group computations for representation theory [10].

In 1968, Shanks [26, 27] proposed an algorithm relying on the baby-step giant-
step method to compute the structure of the class number and the regulator of a
quadratic number field in time $O(|\Delta|^{1/4+\varepsilon})$, or $O(|\Delta|^{1/5+\varepsilon})$ under the extended
Riemann hypothesis [20]. In 1985 Pohst and Zassenhaus [23] published an algorithm
that could determine the class group of arbitrary number fields. Then, a subexponential
strategy for the computation of the group structure of the class group of an
imaginary quadratic extension was described in 1989 by Hafner and McCurley [14].

The expected running time of this method is bounded by $L_\Delta\left(\frac{1}{3}, O(1)\right)$
for certain classes of number fields where both the discriminant and
the degree tend to infinity.

Buchmann [7] generalized this result to the case of an arbitrary extension, the
heuristic complexity being valid for fixed degree $n$ and $\Delta$ tending to infinity. In a
recent work [4], Biasse described an algorithm achieving the heuristic complexity
$L_\Delta\left(\frac{1}{3}, O(1)\right)$ for certain classes of number fields where both the discriminant and
the degree tend to infinity.

In parallel of theoretical improvements, considerable efforts have been invested
to make the implementations of the subexponential methods efficient. In the qua-
dratic case, Jacobson [16] described an algorithm based on the quadratic sieve for
deriving relations between elements of $\text{Cl}(\mathcal{O}_K)$. He successfully used it for com-
puting the class group and the fundamental unit of quadratic number fields. His
implementation contained some of the practical improvements described in the con-
text of factorization such as self initialization and the single large prime variant.
This strategy was later improved by Biasse [3] who used a double large prime vari-
ant and a dedicated Gaussian elimination technique. Attempts have been made
to generalize sieving techniques to general number fields [22], but the proposed
algorithms remain impractical for the sizes of discriminant of interest.

**Our contribution.** In this paper, we present an algorithm based on sieving tech-
niques adapted from recent implementations of the number field sieve [19] for com-
puting $\text{Cl}(\mathcal{O}_K)$ under the generalized Riemann hypothesis (GRH). We also describe
a $p$-adic method for computing the regulator and a system of fundamental units.
We show that these methods allow a significant improvement for number fields of
low degree over the current state of the art.

2. **Generalities on number fields**

Let $K$ be a number field of degree $d$. It has $r_1 \leq d$ real embeddings $(\sigma_i)_{1 \leq i \leq r_1}$ and
$2r_2$ complex embeddings $(\sigma_i)_{r_1+1 \leq i \leq 2r_2}$ (coming as $r_2$ pairs of conjugates). The field
$K$ is isomorphic to $\mathcal{O}_K \otimes \mathbb{Q}$ where $\mathcal{O}_K$ denotes the ring of integers of $K$. We can
embed $K$ in $K_\mathbb{R} := K \otimes \mathbb{R} \cong \mathbb{R}^{r_1} \times \mathbb{C}^{r_2}$, and extend the $\sigma_i$’s to $K_\mathbb{R}$. Let $T_2$ be the
Hermitian form on $K_\mathbb{R}$ defined by

$$T_2(x, x') := \sum_{i} \sigma_i(x) \overline{\sigma_i(x')} ,$$

and let $\|x\| := \sqrt{T_2(x, x)}$ be the corresponding $L_2$-norm. Let $(\alpha_i)_{i \leq d}$ such that
$\mathcal{O}_K = \oplus \mathbb{Z}\alpha_i$, then the discriminant of $K$ is given by $\Delta = \text{det}^2(T_2(\alpha_i, \alpha_j))$. The
norm of an element $x \in K$ is defined by $N(x) = \prod_i |\sigma_i(x)|$. 


To construct the ideal class group of $\mathcal{O}_K$, we rely on a generalization of the notion of ideal, namely the fractional ideals of $\mathcal{O}_K$. They can be defined as finitely generated $\mathcal{O}_K$-modules of $K$. When a fractional ideal is contained in $\mathcal{O}_K$, we refer to it as an integral ideal, which is in fact an ideal of $\mathcal{O}_K$. Otherwise, for every fractional ideal $I$ of $\mathcal{O}_K$, there exists $r \in \mathbb{Z}_{>0}$ such that $rI$ is integral. Fractional ideals are assumed to be non-zero. The sum and product of two fractional ideals of $\mathcal{O}_K$ is given by

$$IJ = \{i_1j_1 + \cdots + i_lj_l \mid l \in \mathbb{N}, i_1, \cdots, i_l \in I, j_1, \cdots, j_l \in J\}$$

$$I + J = \{i + j \mid i \in I, j \in J\}.$$  

The fractional ideals of $\mathcal{O}_K$ are invertible, that is for every fractional ideal $I$, there exists $I^{-1} := \{x \in K \mid xI \subseteq \mathcal{O}_K\}$ such that $I^{−1} = \mathcal{O}_K$. The set of fractional ideals is equipped with a norm function defined by the index $N(I) := [\mathcal{O}_K : I]$ for integral ideals, and the natural extension $N(I/J) := N(I)/N(J)$. The norm of ideals is multiplicative and for principal ideals, it agrees with the norm of the multiplicative group of fractional ideals of $\mathcal{O}_K$.

The ideal class group of $\mathcal{O}_K$ is defined by $\text{Cl}(\mathcal{O}_K) := I/\mathcal{P}$, where $I$ denotes the group of fractional ideals of $K$ and $\mathcal{P} \subseteq I$ is the subgroup of principal fractional ideals. We denote by $[a]$ the class of a fractional $a$ in $\text{Cl}(\mathcal{O}_K)$ and by $h$ the cardinality of $\text{Cl}(\mathcal{O}_K)$. Elements of $I$ admit a unique decomposition as a power product of prime ideals of $\mathcal{O}_K$ (with possibly negative exponents). An element $x \in \mathcal{O}_K$ is said to be a unit if $(x)\mathcal{O}_K = \mathcal{O}_K$, or equivalently if $N(x) = 1$. The units of $\mathcal{O}_K$ form a multiplicative group of the form

$$U = \mu \times \langle \gamma_1 \rangle \times \cdots \times \langle \gamma_r \rangle,$$

where $\mu$ is the torsion subgroup of $U$, $r := r_1 + r_2 − 1$ and the generators $\gamma_i$ of the non-torsion part are called a system of fundamental units. The regulator is an invariant of $K$ which allows us to certify the calculation of $\text{Cl}(\mathcal{O}_K)$ and $U$. It is defined as $R = \text{Vol}(\Gamma)$ where $\Gamma$ is the lattice generated by vectors of the form

$$(c_1 \log |\gamma_{i_1}|, \cdots, c_{r+1} \log |\gamma_{i_{r+1}}|),$$

with $|x|_i := |\sigma_i(x)|$ for $i \leq r + 1$, $c_1 = 1$ for $i \leq r_1$, $c_i = 2$ otherwise (for each complex embedding $\sigma_i$, if $i \leq r + 1$, then $\sigma_i = \sigma_j$ for some $j > r + 1$).

3. The subexponential strategy

The idea behind the algorithm of Buchmann [7] is to find a set of ideals $B = \{p_1, \cdots, p_N\}$ whose classes generate $\text{Cl}(\mathcal{O}_K)$, and then consider the surjective morphism

$$\mathbb{Z}^N \xrightarrow{\varphi} I \xrightarrow{\pi} \text{Cl}(\mathcal{O}_K)$$

$$(e_1, \ldots, e_N) \mapsto \prod_i p_i^{e_i} \mapsto \prod_i [p_i]^{e_i}.$$  

From the fundamental theorem of algebra, the ideal class group satisfies $\text{Cl}(\mathcal{O}_K) \simeq \mathbb{Z}^N / \ker(\pi \circ \varphi)$. Therefore, the knowledge of $\ker(\pi \circ \varphi)$, which has the structure of a $\mathbb{Z}$-lattice, enables us to derive $\text{Cl}(\mathcal{O}_K)$. In the meantime, elements of $\ker(\varphi)$ give us units as power-products of relations. From these units, we hope to derive a system of fundamental units of $\mathcal{O}_K$. The subexponential strategy can be broken down to three essential tasks: collecting relations, calculating the class group and calculating the unit group. The subexponentiality is a consequence of a careful choice of $B$.  

3.1. Relation collection. A preliminary step to the relation collection is the choice of a generating set \( B = \{ p_1, \ldots, p_N \} \) of \( \text{Cl}(O_K) \). We choose the set of prime ideals of norm bounded by an integer \( B \). The use of the Minkowski bound certifies the result unconditionally, but it causes the algorithm to take a time exponential in the size of \( \Delta \). To achieve subexponentiality, many authors chose the bound of Bach [1], who proved that under GRH, \( \text{Cl}(O_K) \) was generated by the classes of the prime ideals \( p \) satisfying \( \mathcal{N}(p) \leq 12 \log(|\Delta|)^2 \). Although asymptotically better, in practice this bound can be larger than the one described by Belabas et al. [2] who stated that under GRH, the class group was generated by the classes of the prime ideals \( p \) satisfying

\[
\sum_{(m,p): \mathcal{N}(p^m) \leq B} \frac{\log \mathcal{N}(p)}{\mathcal{N}(p^{m/2})} \left( 1 - \frac{\log \mathcal{N}(p^m)}{\log(B)} \right) > \frac{1}{2} \log |\Delta| - 1.9n - 0.785r_1 + 2.468n + 1.832r_1 \frac{\log(B)}{\log(B)},
\]

In the rest of the paper, we assume that \( B \) is constructed with the bound of Belabas et al. Indeed, Bach’s bound enlarges the dimensions of the matrices that are processed during the computation of \( \text{Cl}(O_K) \), thus inducing a slow-down that is not compensated by the fact that the relations are found more rapidly.

During the relation collection phase, we collect relations of the form

\[
(\phi_i) = p_1^{e_{i,1}} \cdots p_N^{e_{i,N}},
\]

where \( \phi_i \in K \). We progressively build the matrix \( M := (e_{i,j}) \in \mathbb{Z}^{k \times N} \) where \( k \) is the number of relations collected so far. Let \( \Lambda \subseteq \ker(\pi \circ \varphi) \) be the lattice generated by the rows of \( M \). Operations on the rows of \( M \) allow us to retrieve a basis for \( \Lambda \) and its determinant. To determine if \( \Lambda \) has rank \( N \), we perform operations modulo a random wordsize prime \( p \). In particular, the LU decomposition of \( M \) modulo \( p \) allows us to identify the prime ideals that do not contribute to the rank of \( \Lambda \). Additional relations involving these primes increase the rank of \( M \), whose rows eventually generate a finite index sublattice of \( \ker(\pi) \). To find this index, we compute the Hermite normal form (HNF) of \( M \), that is, we perform unimodular operations encoded by \( U \in \text{GL}_k(\mathbb{Z}) \) such that

\[
UM = \begin{pmatrix}
    h_{11} & 0 & \cdots & 0 \\
    \vdots & h_{22} & \ddots & \vdots \\
    \vdots & \vdots & \ddots & 0 \\
    * & * & \cdots & h_{NN} \\
0 & 0 & \cdots & 0
\end{pmatrix},
\]

with \( \forall j < i : 0 \leq h_{ij} < h_{jj} \) and \( \forall j > i : h_{ij} = 0 \). Once the HNF of \( M \) is computed, adding new rows can be done very efficiently. In the meantime, the product \( \prod_i h_{i,i} \) gives us an indication on \( [\Lambda : \ker(\pi \circ \varphi)] \), as we see in §3.2.

3.2. Class group computation. Given a matrix \( A \in \mathbb{Z}^{N \times N} \) whose rows generate \( \ker(\pi \circ \varphi) \), unimodular transformations on both rows and columns of \( A \) yield the structure of \( \text{Cl}(O_K) \). More precisely, For every non-singular matrix \( A \in \mathbb{Z}^{N \times N} \),
there exist unimodular matrices $U, V \in \mathbb{Z}^{N \times N}$ such that
\[ S := UAV = \text{diag}(d_1, \ldots, d_N) \]
where $\forall i$ such that $1 \leq i < N : d_{i+1} | d_i$. The matrix $S$ is called the Smith normal form (SNF) of $A$.

**Theorem 1.** If the rows of $A \in \mathbb{Z}^{N \times N}$ are a basis for $\ker(\pi \circ \varphi)$, and if $\text{diag}(d_1, \ldots, d_N)$ is the SNF of $A$, then
\[ \text{Cl}(\mathcal{O}_K) \simeq \mathbb{Z}/d_1 \mathbb{Z} \times \cdots \times \mathbb{Z}/d_N. \]

Once enough relations have been found, the rows of $M$ generate $\ker(\pi \circ \varphi)$, and the $N$ non-zero rows of the HNF of $M$ are a basis for $\ker(\pi \circ \varphi)$, and the SNF of $A$ gives us $\text{Cl}(\mathcal{O}_K)$. However, finding the structure of $\text{Cl}(\mathcal{O}_K)$ can also be done computing the SNF of a matrix which is in practice significantly smaller than $A$, namely the essential part of $A$. Indeed, for each matrix $H$ in HNF, there exists an index $l$ such that $\forall i > l, h_{i,i} = 1$. The upper left $l \times l$ submatrix of $H$ is called its essential part. As the classes of $p_i$ for $i > l$ are generated by those of the $p_j$, $j \leq l$, the SNF of the essential part of $A$ suffices to recover $\text{Cl}(\mathcal{O}_K)$.

### 3.3. Regulator and fundamental units computation.
Computing the regulator and a system of fundamental units of $K$ consists of finding kernel vectors of $M$. Indeed, if $X = (x_1, \ldots, x_k)$ satisfies $XM = 0$, then we have
\[ \left( \prod_i \varphi_i^{x_i} \right) \mathcal{O}_K = \mathcal{O}_K. \]
In other words, $\gamma := \prod_i \varphi_i^{x_i}$ is a unit. Every kernel vector $X$ of $M$ yields a unit, and we want to compute the group generated by all those elements as well as the regulator of this group, defined to be zero if the group is not of full rank. So far, finding of relations between units is mostly done using real linear algebra (LLL). The core problem here being the numerical instability of the matrices. This in itself is a consequence of the well-known fact that units are very large in general, writing the fundamental unit of a real quadratic field explicitly with the canonical basis needs exponentially many digits while it is always possible to find a product representation of size polynomial in $\log |\Delta|$. At the end of the procedure, we verify that the assumption we made on the completeness of the lattice of relations is true. To this end, we use an approximate of the Euler product
\[ hR = \frac{\mu(K) |\Delta|^{1/2}}{2^r(2\pi)^r} \lim_{s \to 1} \left( (s-1) \zeta_K(s) \right), \]
where $\zeta_K(s) = \sum_a \frac{1}{N(a) s}$ is the usual $\zeta$-function associated to $K$. Indeed, it allows us to derive a bound $h^*$ in polynomial time under ERH that satisfies $h^* \leq hR < 2h^*$. If the values $\det(\Gamma)$ and $\det(\Lambda)$ do not satisfy this inequality, then we need to collect more relations.

### 4. Sieving techniques
In this section, we describe sieving techniques to derive relations in $\text{Cl}(\mathcal{O}_K)$ for general number fields. This is a generalization of Jacobson’s results [16] for quadratic number fields. We provide numerical data illustrating the considerable
impact of these techniques for class group and unit group computation in the case of low degree number fields. Given a generating set \( B = \{p_1, \ldots, p_N\} \) for \( \text{Cl}(O_K) \), the usual method for deriving relations consists of computing random exponents 
\[ \vec{e} := (e_1, \ldots, e_N), \alpha \in O_K \] and a reduced ideal \( I_\vec{e} \) such that
\[ p_1^{e_1} \cdots p_N^{e_N} = (\alpha)I_\vec{e}. \]

Then, every time \( I_\vec{e} \) is \( B \)-smooth, we obtain a relation. As the arithmetic of ideals is rather expensive when \( n > 2 \), the relation search in the computer algebra software Pari [28] and versions 2.x for \( x < 18 \) of Magma [6] consists of enumerating short elements of \( I_\vec{e} \) via the Fincke-Pohst method [12].

Our method consists of deriving relations from smooth values of polynomials, thus avoiding the cost of the ideal arithmetic and of the ideal reduction. Our method for finding smooth values is based on the recent development of the number field sieve algorithm [19]. The use of trivial methods such as trial division for finding smooth values of our polynomials would yield the same theoretical complexity, but would be impractical for large discriminants. The most efficient implementation of the enumeration-based strategy for finding relations is the one of Pari. Therefore, in the following, we assess the impact of our sieving method by comparing its performance with those of Pari.

4.1. Polynomial selection. Let \( \alpha \) be a \( B \)-smooth ideal of \( O_K \). In this section, we show how to provide a polynomial \( P_\alpha \in \mathbb{Z}[X,Y] \) of degree \( n \) such that every \( (x, y) \in \mathbb{Z}^2 \) such that \( P_\alpha(x, y) \) is \( B \)-smooth yields a relation. Note that in theory, \( \alpha \) can be any ideal, however, we obtained the best results by choosing \( \alpha = O_K \).

Let \( \alpha \) and \( \beta \) be two independent elements of \( \mathfrak{a} \). Then, we create by interpolation a \( P_\alpha \) such that
\[ \forall x, y \in \mathbb{Z}_2^2, P_\alpha(x, y) = N(x\alpha + y\beta). \]

Every time \( \phi_{x,y} := x\alpha + y\beta \) has a smooth norm, we add the relation corresponding to the principal ideal \( (\phi_{x,y}) \) to the relation matrix. Before applying sieving algorithms to \( P_\alpha \) to derive relations, we need to ensure that it is likely to yield enough smooth values. Polynomial selection is an important part of the number field sieve algorithm, and so it is in our algorithm. However, the specificities of our context prevent us from directly adapting the methods of NFS for selecting the sieving polynomial. First of all, we can afford to find relations with many different choices of \( \alpha \) and \( \beta \), whereas the choice of a sieving polynomial in the NFS algorithm is fixed.

We require that our choices of \( \alpha \) and \( \beta \) yield polynomials with small coefficients, and that we have a sufficient randomization at the infinite places to avoid drawing \( \phi_{x,y} \) spanning the same subgroup of the unit group of \( O_K \).

To randomize the choice \( \alpha, \beta \), we consider random coefficients \( a_1, \ldots, a_n \in \mathbb{R}^n \) such that \( \sum_{i \leq n} a_i = 0 \). For every such \( n \)-tuple \( \vec{a} \), we define the embedding
\[ \psi_{\vec{a}}: \alpha \mapsto (a_1 \log |\alpha|_1, \ldots, a_n \log |\alpha|_n). \]

For every choice of \( \vec{a} \), the set of elements of the form \( \psi_{\vec{a}}(\alpha) \) for \( \alpha \in \mathfrak{a} \) is a lattice \( \Lambda_{\vec{a}} \) of \( \mathbb{R}^n \) for which we can find an LLL reduced basis for the norm
\[ T_{\vec{a}}^2 : (x_1, \ldots, x_n) \mapsto e^{2a_1 x_1^2 + \ldots + 2a_n x_n^2}. \]

For every choice of \( \vec{a} \), the first two vectors \( \alpha, \beta \) of an LLL reduced basis of \( \Lambda_{\vec{a}} \) are potential candidates for the creation of a polynomial yielding smooth values.
Every time we draw such a couple of elements of \( a \), we need to make sure that they do not generate the same \( \mathbb{Z} \)-module as another pair previously used. To prevent this from happening, every time we draw a couple \( \alpha, \beta \) by the previous method, we express them in terms of the canonical \( \mathbb{Z} \)-basis of \( \mathcal{O}_K \). Thus, to every couple \( \alpha, \beta \) corresponds the matrix \( M_{\alpha, \beta} \in \mathbb{Z}^{2 \times n} \) of their coordinates. The HNF of \( M_{\alpha, \beta} \) uniquely represents the \( \mathbb{Z} \)-module spanned by \( (\alpha, \beta) \). Thus, to avoid duplicates, we store a hash of the HNF of \( \mathbb{Z}_p \), derive all the others by translation by \( (0,0) \). From (1), we know that once we have one root \( \alpha, \beta \), we express them in terms of the canonical \( \mathbb{Z} \)-basis of \( \Lambda_\mathbb{Z} \) for \( \tilde{a} = (a_1, \ldots, a_n) \).

Thus, to every couple \( (\alpha, \beta) \) to draw relations. We summarize the procedure of the selection of a sieving polynomial in Algorithm 1.

Algorithm 1 Polynomial selection

| Input: | \( a, (A_1, \ldots, A_n) \), HashTable |
|---|---|
| Output: | Sieving polynomial \( P_{\alpha, \beta} \) corresponding to \( \alpha, \beta \in a \) |
| 1: while | a new \( \alpha, \beta \) has not been found do |
| 2: Draw | \( a_1 \leq A_1, \ldots, a_n \leq A_n \) at random such that \( a_1 + \ldots + a_n = 0 \) |
| 3: Let | \( \alpha, \beta \) be the first two elements of a LLL-reduced basis of \( \Lambda_\mathbb{Z} \) for \( \tilde{a} = (a_1, \ldots, a_n) \) |
| 4: Compute the | hash \( h_{\alpha, \beta} \) of the HNF of \( M_{\alpha, \beta} \) |
| 5: if | \( h_{\alpha, \beta} \notin \text{HashTable} \) then |
| 6: Compute by | interpolation \( P_{\alpha, \beta} \in \mathbb{Z}[X,Y] \) such that \( P_{\alpha, \beta}(x, y) = N(x\alpha + y\beta) \) |
| 7: end if |
| 8: end while |
| 9: return | \( \alpha, \beta, P_{\alpha, \beta} \) |

4.2. Line sieving. The quadratic sieve algorithm [25] used to derive smooth values of a binary quadratic form generalizes to the case of polynomials of arbitrary degree. Its design follows from the observation that if \( P \in \mathbb{Z}[X,Y] \) is a polynomial of degree \( n \), then

\[
\forall y_0 \in \mathbb{Z}, \ p \mid P(r_p, y_0) \Rightarrow \forall i \in \mathbb{Z}, \ p \mid P(r_p + ip, y_0).
\]

Given \( y_0 \in \mathbb{Z} \), we wish to find the \( x \in [-I/2, I/2] \) such that \( P(x, y_0) \) is \( B \)-smooth, where \( B \) is the bound on the norm of the prime ideals in the factor base. Instead of trying them all, we prefer to isolate a short list of good candidates that we test by trial division. If \( p \mid P(x, y_0) \) for many \( p \leq B \), then \( P(x, y_0) \) is likely to be \( B \)-smooth. From (1), we know that once we have one root \( r_p \) of \( P(X, y_0) \) mod \( p \), then we can derive all the others by translation by \( (p, 0) \). Line sieving consists of initiating to zero an array \( S \) of length \( I \) whose cells represent the \( x \in [-I/2, I/2] \). Then, for each \( p \leq B \), we compute the smallest roots \( x_p \in [-I/2, I/2] \) of \( P(X, y_0) \) mod \( p \) and repeat

\[
S[x_p] \leftarrow S[x_p] + \log(p), \quad x_p \leftarrow x_p + p.
\]

Then, whenever \( S[x] \approx \log(P(x, y_0)) \) for \( x \in [-I/2, I/2] \), the value \( P(x, y_0) \) is likely to be \( B \)-smooth. We summarize this procedure in Algorithm 2.

4.3. Lattice sieving. Let \( P_{\alpha, \beta}(X,Y) \in \mathbb{Z}[X,Y] \) be the sieving polynomial described in § 4.1 B the bound on the norm of the ideals in the factor base, and \( I,J \in \mathbb{Z}_{>0} \). Every couple \( (x,y) \in [-I/2, I/2] \times [1, J] \) such that \( P_{\alpha, \beta}(x,y) \) is \( B \)-smooth yields a relations. Therefore, one can repeat the line sieving operation on
Algorithm 2 Line sieving

Input: $P \in \mathbb{Z}[X,Y]$, $I, B, y_0 \in \mathbb{Z}$

Output: Smooth values of $P(X, y_0)$ in $[-I/2, I/2]$  
1: $L \leftarrow \emptyset$, $\forall x \in [-I/2, I/2]$, $S[x] \leftarrow 0$.
2: for $p \leq B$ do
3: Let $x_p$ be the smallest root of $P(X, y_0) \mod [-I/2, I/2]$.
4: while $r_p \leq I/2$ do
5: $S[x_p] \leftarrow S[x_p] + \log(p)$, $x_p \leftarrow x_p + p$.
6: end while
7: end for
8: for $x \in [-I/2, I/2]$ do
9: if $S[x] \approx P(x, y_0)$ then
10: If $P(x, y_0)$ is $B$-smooth, $L \leftarrow L \cup \{x\}$.
11: end if
12: end for
13: return $L$

$P_{\alpha, \beta}(X, y_0)$ for every $y_0 \in [1, J]$. This method is efficient when sieving with primes $p < I$, but when the primes are significantly larger than $I$, the root computation at Step 3 of Algorithm 2 is often performed for nothing since there is a good chance that none of the $x \in [-I/2, I/2]$ will be a root of $P_{\alpha, \beta}(X, y_0) \mod p$. A way around that is to have an array $S$ of length $IJ$ representing $[-I/2, I/2]$ and to fill it by line sieving methods for the primes $p < I$ and by lattice sieving for the other primes.

The lattice sieve was first described by Pollard [24]. Since then, it has been extensively studied and improved in the past 15 years, and the most recent developments of this method yield the factorization of RSA768 (see [19]). This strategy relies on a one-time enumeration of roots of $P_{\alpha, \beta}(X, Y) \mod p$ in $[-I/2, I/2] \times [1, J]$. The entry $x \leq IJ$ of the array $S$ that we use to store the logarithmic contributions corresponds to the couple $(i, j) \in [-I/2, I/2] \times [1, J]$ where

$$i = (x - I/2) \mod I$$

$$j = (x - i - I/2)/I.$$

As in the line sieving case, every entry of $S$ is initialized to zero, and for every $p \leq B$ and every $(i, j) \in [-I/2, I/2] \times [1, J]$ such that $p \mid P_{\alpha, \beta}(i, j)$, we want to perform the operation $S[x] \leftarrow S[x] + \log p$. Line sieving repeated on every line $j \leq J$ allows us to efficiently do this for $p < I$. For the others, we followed the approach of [13], as it is done in [19] for the factorization of RSA768. By [13 Prop. 1], we know that for every $p$ such that we have a root $r_p$ of $P_{\alpha, \beta}(X, 1)$ modulo $p$, there exists a basis $\{(a, b), (c, d)\}$ of the couples $(i, j)$ such that $p \mid P_{\alpha, \beta}(i, j)$ that satisfies

- $b > 0$ and $d > 0$
- $-I < a \leq 0 \leq c < I$
- $c - a \geq I$.

This basis is computed via an algorithm described in [13] that relies on the continued fraction expansion of $r_p$. To fill the array $S$, we start from $(i, j) = (0, 0)$ which is a common root modulo all primes. Then, by induction, we construct the next pair $(i', j')$ from $(i, j)$ by choosing

- $(i, j) + (a, b)$ if $i \geq -a$
Then we find additional relations with enumeration.

To do this, we used a modified version of the function `bnfinit` its contribution to the performances of the class group and unit group computation. 

compute the roots of $P \mod q$ occur in a relation, one can use the special-

strategy consists of sieving with a polynomial $P_q$ derived from the original sieving polynomial $P$ such that

$$\forall (i, j) \in [-I/2, I/2] \times [1, J], \exists (x, y) \in \mathbb{Z}^2, P_q(i, j) = P(x, y)$$

This strategy was used by Pollard in his original paper [24] to sieve on the rational side, but most current implementations use it in the rational side as well [19].

To create $P_q$ for a given $q$, we need a root $r_q$ of $P$ modulo $q$. Then, we find a reduced basis $(a_0, b_0), (a_1, b_1)$ of the lattice spanned by the vectors $(r_q, 0), (r_q, 1)$. The polynomial $P_q$ is then simply given by

$$P_q(i, j) = P(ia_0 + ja_1, ib_0 + jb_1).$$

The reduced basis is given by successive Gaussian reductions, as explained in [13].

Then, to sieve with a given polynomial $P$, we repeat the procedure described in §4.3 for many different polynomials of the form $P_q$. Fortunately, once the roots of $P \mod p$ for all $p \leq B$ have been computed, it is possible to use these values to compute the roots of $P_q \mod p$ for $p \leq B$. Indeed,

$$P(ia_0 + ja_1, ib_0 + jb_1) \equiv 0 \mod p$$

means that there is some root $r_p$ of $P(X, 1) \mod p$ such that $r_p = \frac{ia_0 + ja_1}{ia_0 + ja_1} \mod p$.

This implies that we have $P_q(r_p^q, 1) \equiv 0 \mod p$ for

$$r_p^q = \frac{i}{j} = -\frac{a_1 - r_pb_1}{a_0 - r_pb_0} \mod p,$$

which gives us a root of $P_q(X, 1) \mod p$ from $(a_0, b_0), (a_1, b_1)$ and a root of $P(X, 1) \mod p$. We summarize our procedure to derive relations from an ideal $\mathfrak{a} \subseteq \mathcal{O}_K$ in Algorithm [3]

4.5. Overall relation collection phase. A necessary condition to compute the class group and the unit group is to produce a full-rank relation matrix $M$. Our sieving methods allow us to derive relations in $\text{Cl}(\mathcal{O}_K)$ very rapidly, but it is hard to force a given prime to occur in a relation. The best performance is obtained by sieving with the trivial ideal $\mathcal{O}_K$. If we want to see a given prime ideal $p | (p)$ occur in a relation, one can use the special-$q$ with $q = p$, or sieve with the ideal $p$. 

However, even after using those methods, some prime ideals still do not contribute to the rank of $M$. Rather than sieving in random power-products involving missing primes, one might prefer to switch to enumeration-based methods to complete the relation search. To identify the primes that need to appear in a relation, we perform an $LU$ decomposition of the relation matrix modulo a random wordsize prime. We try to produce enough relations with sieving so that the rank of $M$ is $97\%$ of $\#B$.

Then we find additional relations with enumeration.

To assess the advantage of sieving over enumeration techniques, we need to isolate its contribution to the performances of the class group and unit group computation. To do this, we used a modified version of the function `bnfinit` of the computer.
Algorithm 3 Sieving procedure

Input:  \( a \subseteq O_K, B = \{ p \mid N(p) \leq B \}, I, J \in \mathbb{Z}_{>0} \).
1: Select \( \alpha, \beta \in C_K \) and a sieving polynomial \( P_{\alpha, \beta} \) with Algorithm 1.
2: \( \forall p \leq B \), compute the roots of \( P_{\alpha, \beta}(X, 1) \mod p \).
3: for \( q \leq B \) do
4: Compute \( P_q \) and its roots modulo the \( p \leq B \) as in §4.4.
5: Let \( S \) be an array of size \( IJ \) initialized to 0.
6: for \( p \leq I \) do
7: Do \( S[x] \leftarrow S[x] + \log(p) \) for each \( x \) representing \( (i, j) \in [-I/2, I/2] \times [1, J] \) such that \( p \mid P_q(i, j) \) by repeating Algorithm 2 for each line \( j \leq J \).
8: end for
9: for \( p > I \) do
10: Calculate a basis \( \{ (a, b), (c, d) \} \) of the lattice of points in \( [-I/2, I/2] \times [1, J] \) such that \( p \mid P_q(i, j) \) by using the method of §4.3.
11: Do \( S[x] \leftarrow S[x] + \log(p) \) for each \( x \) representing \( (i, j) \in [-I/2, I/2] \times [1, J] \) such that \( p \mid P_q(i, j) \) by using the method of §4.3.
12: end for
13: end for
14: for \( x \leq IJ \) do
15: if \( S[x] \approx \log(P_q(i, j)) \) where \( x \) represent \( (i, j) \in [-I/2, I/2] \times [1, J] \) then
16: If \( \log(P_q(i, j)) \) is \( B \)-smooth, store the corresponding relation.
17: end if
18: end for

Algorithm 4 Full rank relation matrix computation

Input:  \( K, B \)
Output: A full-rank relation matrix for the primes of norm bounded by \( B \)
1: \( B \leftarrow \{ p \mid N(p) \leq B \} = \{ p_1, \ldots, p_N \} \).
2: Derive \( N \) relations by repeating Algorithm 3 with \( a = (1) \). Let \( M \) be the relation matrix.
3: Perform an LU decomposition of \( M \) and let EmptyList be the list of zero columns.
4: for \( p \in \text{EmptyList} \) do
5: Sieve with \( p \), update \( M \).
6: end for
7: Update EmptyList by updating the LU decomposition of \( M \).
8: for \( p \in \text{EmptyList} \) do
9: Find a relation involving \( p \) by enumerating short elements in random power-products.
10: end for
11: return \( M \)

algebra software Pari that accepts in input a list of precomputed relations. We interfaced via SAGE this version of Pari with a developing version of Magma containing a function creating relations with the sieving algorithm. The Magma function tries to create enough relations so that the rank of \( M \) be 97% of \( \#B \) and passes it to Pari which adds new relations with enumeration methods and
calculates the class group and the unit group. We compared the performance of this approach to the traditional `bnfinit` function of Pari. There are two main reasons for using a hybrid version. The first one is that Pari’s implementation of enumeration techniques is the most efficient. As these are necessary to finish the creation of the relation matrix after calling the sieving algorithm, it is interesting to see how the two perform together. Another reason for this choice is the fact that many different algorithms contribute to the computation of the class group and the unit group. In particular, we use time-consuming linear algebra methods such as the HNF computation. Our methodology avoids the risk of seeing the influence of the quality of the implementation of other algorithms occurring in the class group and unit group computation.

We performed our computations on a 2.6 GHz Opteron with 4GB of memory. We used a branch of the developping version 2.6.0 of Pari provided by Loïc Grenier and the developping version of Magma, interfaced via Sage 4.7.2. We allocated 3GB of memory to the computation made with Pari. For each size $d$, we drew at random 10 number fields with discriminant satisfying $\log_2 |\Delta| = d$. For each discriminant, we computed the class group and the unit group with `bnfinit`, which we refer to as the “Pari” method, and with the hybrid version which we refer to as the “sieving+Pari” method. The average timings, in CPU sec (rounded to the nearest integer), are presented in Table 1. They illustrate the impact of sieving methods for small degree number fields. It is very strong for degree 3,4 and 5 number field, for which we often witness a speed-up of a factor at least 10 while it is rather moderate for degree 6 number fields, and negligible for number fields of degree 7,8. Finding smooth values of a polynomial gets more difficult when we increase its degree, but it is not the only reason why the impact of sieving decreases with the degree. Indeed, for degree 6 number field, our sieving algorithm still derives relations at a competitive pace, but there are many linear dependencies whereas enumeration allows a more targeted search, thus avoiding linear dependencies. To put

### Table 1. Impact of sieving on class group and unit group computation of small degree number fields

| $n$ | $\log_2 |\Delta|$ | Pari | Pari+Sieving |
|-----|-----------------|------|-------------|
| 3   | 120             | 76   | 11          |
| 3   | 140             | 694  | 66          |
| 3   | 160             | 6828 | 333         |
| 3   | 180             | 29807| 2453        |
| 4   | 120             | 38   | 7           |
| 4   | 140             | 366  | 24          |
| 4   | 160             | 4266 | 175         |
| 4   | 180             | 31661| 1201        |
| 5   | 120             | 33   | 18          |
| 5   | 140             | 295  | 64          |
| 5   | 160             | 3402 | 378         |
| 5   | 180             | 16048| 2342        |
| 6   | 120             | 40   | 111         |
| 6   | 140             | 294  | 161         |
| 6   | 160             | 1709 | 1012        |
| 6   | 180             | 14549| 8413        |
Table 2. Impact of the quadratic sieve on fields generated by $X^2 + 4(10^n + 1)$

| $n$  | 20 | 30 | 40 | 45 | 50 | 54 |
|------|----|----|----|----|----|----|
| Magma 2.18 | 0.7 | 6  | 22 | 128| 170| 1453|
| Pari + Sieving | 0.5 | 5 | 44 | 271| 593| 1085|
| Pari    | 0.2 | 3 | 66 | 556| 1562| 9251|

these improvements into perspective, we show in Table 2 the impact of Jacobson’s self initializing quadratic sieve [16] which is implemented in Magma 2.18. The timings for “Pari” and “Pari + Sieving” are derived under the same setting as for Table 1. In addition, we added the performances of of Magma 2.18 which uses different methods for linear algebra. Timings for the same serie of number fields were reported by Jacobson in [16, Table A.3] on a 296Mhz SUN processor (for a fair comparison one has to take into account the verification time since the timings of Table 1 and Table 2 correspond to a certification under GRH).

5. Computing the unit group

Assume that we have created a relation matrix $(e_{i,j})$ corresponding to the relations

$$(\phi_i) = p_1^{e_{1,1}} \cdots p_N^{e_{N,N}}.$$ 

Every kernel vector allows us to derive a unit of $O_K$. Let $\beta_1, \cdots, \beta_k$ be a generating system of the unit created so far. We compute a new unit $\beta'$, and we wish to find a new minimal generating set for $(\beta_1, \cdots, \beta_k, \beta)$. Usually this is done by computing (real) logarithms of the units followed by some approximate linear algebra to find a (tentative) relation as well as the (tentative) new basis. This then is followed by an exact verification of the relation to guarantee correctness. The difficulty comes from the fact that the entries in the real matrix differ vastly in size - by several orders of magnitude - thus making it necessary to work with a huge precision, in fact the precision is also subexponential in the discriminant for guaranteed results.

Here, we propose to use $p$-adic logarithms instead. The key advantage comes from the much better control of error propagation in the linear algebra: unless division by non-units happens, linear algebra does not increase errors. However, while the correctness is based on the unproven Leopold conjecture about the non-vanishing of the $p$-adic regulator, this is not a problem in practice: any relation found by the $p$-adic method can easily be verified unconditionally, thus a failure of the algorithm would provide a counter example to Leopold’s conjecture.

We start by choosing a prime $p$ such that the $p$-adic splitting field $K_p$ has moderate degree, here we allow at most degree 2. Then we have $n$ embeddings $\phi_i$ of $K \to K_p$, and we define a map $L_p : K^* \to K_p^n : x \mapsto (\log \phi_i(x))_i$ where $\phi_i$ is the usual $p$-adic logarithm extended to $K_p$. In order to estimate the necessary $p$-adic precision, we also need the usual real logarithmic embedding, denoted by $L : K^* \to \mathbb{R}^{r+1}$. We are looking for a (rational) solution $(x_i)$ to $\sum x_i L_p(\beta_i) = L_p(\beta')$. Using $p$-adic linear algebra we will instead get a $p$-adic solution (or a proof that $\beta'$ is independent). Using standard rational reconstruction techniques, we derive the rational solution from the $p$-adic one and then the integral relation between the units. In order to estimate the $p$-adic precision, we bound numerator and denominator using Cramers rule and universal lower bounds on the
logarithms of units. The rational solution then also satisfies $\sum x_i L(\beta_i) = L(\beta')$. Let $(\alpha_i)$ be a basis for $\langle \beta_1, \ldots, \beta_n, \beta' \rangle$. By Cramer’s rule, we write

$$x_i = \frac{\det(L_p(\beta_1), \ldots, L_p(\beta'), \ldots, L_p(\beta_s))/\det(L_p(\beta_1), \ldots, L_p(\beta_s))}{\det(L_p(\beta_1), \ldots, L_p(\beta_s))/\det(L_p(\alpha_1), \ldots, L_p(\alpha_s))}$$

Since the (unknown) $(\alpha_i)$ form a basis, we see that

$$\det(L_p(\beta_1), \ldots, L_p(\beta'), \ldots, L_p(\beta_s))/\det(L_p(\alpha_1), \ldots, L_p(\alpha_s))$$

is an integer and the same is true for $L$ instead of $L_p$, thus we can write $x_i$ as a quotient of integers. In either case, to make sense of the determinants, we will have to select an appropriate number of rows to make the matrices square. To bound the integers, we make use of the Hadamard bound for $\det(L(\beta_1), \ldots, L(\beta'), \ldots, L(\beta_s))$ and some universal lower bound for $\det(L(\alpha_1))$. For the lower bound we use lower bounds of logarithms of non-torsion units ([11 3.5]): $\|L(\alpha_i)\|_2 \geq \frac{21 \log d}{128}$, or, if the unit group has full rank, $s = r = r_1 + r_2 - 1$, we use lower regulator bounds, possibly coming from the Euler product. Having obtained bounds from the real logarithm ($L$) with low precision, we calculate the $p$-adic precision required to find $x_i$ using $p$-adic linear algebra and rational reconstruction. In the course of the computation it can happen that the $p$-adic determinants ($p$-adic regulators) have non-trivial valuation. In this case we have to restart the computation with a correspondingly higher precision to account for the loss. Since the Leopold-conjecture is non-proven as of now, we also need to verify the solution by computing a low-precision estimate for $\| \sum x_i L(\beta_i) - L(\beta') \|$ to compare it to the lower bound used above.

From the relation $x_i$, we can easily obtain a presentation of the new basis $\alpha_i$ in terms of the $\beta_i$, $\beta'$. For optimization, we then proceed to compute a new basis $\hat{\alpha}_i$ such that the real logarithms are (roughly) $LLL$-reduced. We note that we do not rely on any $LLL$ estimates here, so any heuristic algorithm that aims at reducing the apparent size will do. Since we do not have any $LLL$ algorithm that will accept real input (as opposed to rational), it is important that this does not influence the correctness.

5.1. Advantages of the $p$-adic method. There are two core advantages of the $p$-adic logarithms over the ordinary, complex, ones: first, the linear algebra problems we need to solve in order to find dependencies or relations between units have a much simpler error analysis. In fact, contrary to the complex case, it is possible through the use of ring based operations to solve linear equations without any additional loss of precision. This is very important in the context of unit computation since the matrices representing the image of $L(\alpha)$ are very badly conditioned for classical numerical methods. The other advantage of the $p$-adic logarithms is more subtle: if we assume Leopold’s conjecture to hold for the field(s) we are interested in, then instead of doing linear algebra over $\mathbb{R}$ with a precision of say $q$ to find dependencies, it is sufficient to work with a real precision of $q/2$ and a $p$-adic precision of $q/2$ as well. Thus, assuming classical multiplication, we gain a factor of about 4 through the use of lower precision. Using fast multiplication (in high precision), the gain is smaller but still noticeable. But the most important advantage is the much easier precision control: instead of complicated and very delicate estimates for linear algebra problems, all we need are upper bounds on linear combinations with integral coefficients - which are trivial to obtain.

We should also mention that one disadvantage of the $p$-adic method lies in the total lack of control over the real size of the units, thus it needs to be paired
with a crude (and uncritical for correctness) size reduction algorithm. Also, it is (currently) not possible to forego completely the use of complex (or real) logarithms, as the $p$-adic method is not capable to proving a unit to be torsion - without the knowledge of bounds on the real size.

### 5.2. Lower bound from Euler Product

Suppose that, as in the class group algorithm, we are given an approximation of the Euler product, ie. we have a real number $E$ such that $1/\sqrt{2} \leq hR/E \leq \sqrt{2}$. After the relation matrix has full rank, and assuming the factorbasis is large enough for correctness, we have an upper bound for the class number, thus a lower bound for $R$. This lower bound will be several orders of magnitude larger than the universal bounds available otherwise.

### 5.3. Saturation

After the initial steps of the algorithm, when the relation matrix has full rank, we have a tentative class number $h$ and a tentative regulator $R$. Experimentally, at this point, $hR$ does not approximate the Euler product very well - the product will be off by several orders of magnitude. However, after we found one or two more relations, the product has the same size than the Euler product, it frequently even looks like only a factor of 2 is missing in either $h$ or $R$.

To find the last missing relation can easily take more time than the entire previous run therefore we suggest using saturation methods instead. At this point in the algorithm the relations define a subgroup $U$ of the $S$-unit group $U_S$ where $S$ is the factor basis. From the Euler product we know that the index $(U_S : U) =: b$ is small, lets say $b < B$. For any prime $p|b$ there is some $u \in U_S \setminus U$ such that $u^p \in U$. Let us fix the prime $p$. For any prime ideal $Q \notin S$ such that $p|N(Q) - 1$ we can define the map $\phi_Q : U \to \mathbb{F}^\times_Q/(\mathbb{F}_Q^\times)^p$ mapping $S$-units into the multiplicative group of the residue class field modulo $p$-th powers.

The Cebotaref theorem [29] guarantees that if $u \in U$ is not a $p$-th power, there will be some $Q$ such that $\phi_Q(u)$ is non-trivial, ie. $u$ is not a $p$-th power modulo $Q$. We now simply intersect $\ker \phi_Q$ for several $Q$ until either the intersection is $U^p$ or it is does not change for five consecutive $Q$. We expect that any $u \in U/\bigcap \ker \phi_Q$ will have a $p$-th root in $U_S$ but not in $S$.

Therefore $u^p = u$ is a new relation that will change $hR$ by $p$. Repeating this for all $p < B$ until we cannot enlarge $U$ any more we find the missing relations.

### 5.4. Representation

During the execution of the algorithm, all ($S$-)units are naturally represented as power products of the relations coming from the sieving (or the saturation). It is well known that the explicit representation of the units with respect to a fixed basis for the field can require exponentially large coefficients, so it is important to operate on the power products as much as possible. However, even the exponent vectors constructed for the basis of the unit group, or the saturation, will become huge, so we need to “size reduce” the power products. In particular, this happens even if the resulting element is not too large. Using ideas of [8] for compact representations and [15] for reduced divisors in function fields, we can find a representation for those elements that depends only on the logarithmic size (and the number field) rather than the execution path. For any prime $p$ we can write any unit

$$ u = \prod r_i^{e_i} = \prod a_i^{p^{e_i}} $$

with elements such that the size of $a_i$ depends on the discriminant and $p$ only. The length of the product comes from $L(u)$. Furthermore, in this presentation it is easy to test for $p$-th powers as only $a_0$ needs to be tested and this is a small element.
5.5. Example. To illustrate the power of the p-adic method, we look at a totally real quartic field generated by a root of

\[ x^4 + 17211x^3 + 5213x^2 - 176910463x - 4958. \]

The discriminant \( \Delta \) of the maximal order has 38 digits. In the course of the computation, we found 534 relations involving prime ideals of norm up to 3000 = \( 0.4 \log^2 |\Delta| \) describing a trivial class group. We then searched for 5 further relations to obtain units \( u_i \) (1 \( \leq \) i \( \leq \) 5). As power products of the relations, the units are given via exponent vectors \( e_i \) with \( ||e_i||_\infty \) ranging between \( 10^{80} \) and \( 10^{100} \) and \( 20 < ||e_i||_1/||e_i||_\infty < 92 \). So, while not uniformly large, the exponents are non-sparse, involving huge integers. Using a decimal precision of 170 digits, we establish that the logarithms of the units are roughly \( ||L(u_i)||_\infty \approx 10^{160} \). The first three units are indeed independent, giving a basis for a subgroup of full rank, the fourth is then dependent. Choosing the prime \( p = 10337 \) we get \( \mathbb{Q}_p \) as a splitting field. Using a p-adic precision of 245 digits (ie. working in \( \mathbb{Z}_p \) mod \( p^{245} \)), we compute the dependency for the fourth unit, involving exponents of around \( 10^{360} \). The new unit group is then tentatively LLL reduced, producing a new basis where the \( ||L(\tilde{u}_i)||_\infty \) are bounded by \( 10^7 \) only. The last unit then involves a much smaller dependency, here the exponents are only around \( 10^{60} \).

Unfortunately, looking at the Euler product, the unit group is not complete. However, the saturation technique outlined above takes 1sec to determine that the product of the three basis elements is (proabably) a square. Finding a better representation where the exponents are all powers of 2 takes less than 1 sec and then we can enlarge the unit group easily.

Due to the implementation, the p-adic precision used was actually higher: changing (increasing) precision is very computationally expensive, so we try to avoid this and simply double the precision. We used a precision of 320 for the p-adics and a maximal precision of 1000 for the real precision. The computation of the log is the dominating part, we spent 50sec or 90% of the total processing time here.

6. Conclusion

We introduced new techniques to enhance the performances of the subexponential methods for computing the class group and the unit group of a number field. In particular, sieving allows a speed-up of an order of magnitude for number fields of small degree. These techniques could be developed even further. Indeed, we have not taken into account all the improvements to sieving techniques describes in the context of the number fields sieve algorithm such as large prime variations or cache-friendly methods. It is also notable that fast techniques for deriving relations in the class group of a small degree number field have applications in evaluating isogenies between small genus curves via complex multiplication methods. Indeed, in that case, evaluating isogenies between genus \( g \) curves involves relations in the class group of a degree \( 2g \) number field.

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