

Breaking pairing-based cryptosystems using η_T pairing over $GF(3^{97})$

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Abstract. There are many useful cryptographic schemes, such as ID-based encryption, short signature, keyword searchable encryption, attribute-based encryption, functional encryption, that use a bilinear pairing. It is important to estimate the security of such pairing-based cryptosystems in cryptography. The most essential number-theoretic problem in pairing-based cryptosystems is the discrete logarithm problem (DLP) because pairing-based cryptosystems are no longer secure once the underlying DLP is broken. One efficient bilinear pairing is the η_T pairing defined over a supersingular elliptic curve E on the finite field $GF(3^n)$ for a positive integer n . The embedding degree of the η_T pairing is 6; thus, we can reduce the DLP over E on $GF(3^n)$ to that over the finite field $GF(3^{6n})$. In this paper, for breaking the η_T pairing over $GF(3^n)$, we discuss solving the DLP over $GF(3^{6n})$ by using the function field sieve (FFS), which is the asymptotically fastest algorithm for solving a DLP over finite fields of small characteristics. We chose the extension degree $n = 97$ because it has been intensively used in benchmarking tests for the implementation of the η_T pairing, and the order (923-bit) of $GF(3^{6 \cdot 97})$ is substantially larger than the previous world record (676-bit) of solving the DLP by using the FFS. We implemented the FFS for the medium prime case (JL06-FFS), and propose several improvements of the FFS, for example, the lattice sieve for JL06-FFS and the filtering adjusted to the Galois action. Finally, we succeeded in solving the DLP over $GF(3^{6 \cdot 97})$. The entire computational time of our improved FFS requires about 148.2 days using 252 CPU cores. Our computational results contribute to the secure use of pairing-based cryptosystems with the η_T pairing.

Keywords: pairing-based cryptosystems, η_T pairing, discrete logarithm problems, function field sieve

1 Introduction

After the advent of the tripartite Diffie-Hellman (DH) key exchange scheme [20] and ID-based encryption using pairing [11], plenty of attractive pairing-based cryptosystems have been proposed, for example, short signature [13], keyword searchable encryption [10], efficient broadcast encryption [12], attribute-based encryption [29], and functional encryption [27]. Pairing-based cryptosystems have become a major research topic in cryptography.

Pairing-based cryptosystems are constructed on the groups G_1 , G'_1 and G_2 of the same order with a bilinear pairing $G_1 \times G'_1 \rightarrow G_2$. The security of pairing-based cryptosystems is based on the difficulty in solving several number-theoretic problems such as the computational/decisional bilinear DH problem (CBDH/DBDH), strong DH problem (SDH), decisional linear problem (DLIN), and symmetric external DH problem (SXDH). However, the most important number-theoretic problem in pairing-based cryptosystems is the discrete logarithm problem (DLP) on G_1 , G'_1 , and G_2 . All the other number-theoretic problems above are no longer intractable once the DLP on G_1 , G'_1 , or G_2 is broken. Therefore, it is important to investigate the difficulty in solving the DLP.

One of the most efficient algorithms for implementing the pairing is the η_T pairing [5] defined over a supersingular elliptic curve E on the finite field $GF(3^n)$, where n is a positive integer. Since the embedding degree of E is 6, the η_T pairing can reduce a DLP over E on $GF(3^n)$, which is called an ECDLP, to a DLP over $GF(3^{6n})$. Joux proposed the (probably) first cryptographic scheme [20] that uses the pairing over E . Boneh *et al.* then applied the pairing over E to the short signature scheme [13], where a point (x, y) on E for extension degree $n = 97$ can be represented as a signature value, e.g., $x = \text{KrpIcV009CJ8iyBS8MyVknrMyE}$. At CRYPTO 2002, Barreto *et al.* presented algorithms for efficiently computing Tate pairing over E [6]. Many high-speed implementations of pairing over E have subsequently been proposed [3, 7–9, 17, 18, 24]. For many of these implementations, benchmark tests using the extension degree $n = 97$ have been conducted. Therefore, we focus on the DLP over finite field $GF(3^{6 \cdot 97})$ in this paper. The cardinality of the subgroup of the supersingular elliptic curve is 151 bits, and that of $GF(3^{6 \cdot 97})$ is 923 bits. The size of our target DLP is 247 bits larger than the

Table 1. Summary of time data for solving DLP over $GF(3^{6\cdot 97})$

phase	method	time	machine environment
collecting relations	lattice sieve	53.1 days	212 CPU cores
linear algebra	parallel Lanczos	80.1 days	252 CPU cores
individual logarithm	rationalization and special- Q descent	15.0 days	168 CPU cores
total		148.2 days	252 CPU cores

previous world record of solving the DLP over $GF(3^{6\cdot 71})$, whose cardinality is 676 bits [19]. The current world record for solving an ECDLP is the 112-bit ECDLP [14]. Pollard’s ρ method is used for solving the 112-bit ECDLP, and has not reached the ability for solving the 151-bit ECDLP over the subgroup of E .

In this paper, we analyze the difficulty in solving the DLP over $GF(3^{6\cdot 97})$ by using the function field sieve (FFS), which is known as the asymptotically fastest algorithm [1, 2]. Since the FFS proposed by Joux and Lercier (JL06-FFS) [23] is suitable for solving the DLP over a finite field whose characteristic is small, we use the JL06-FFS and propose several efficient techniques for increasing its speed. Note that the FFS generally consists of four phases: polynomial selection, collecting relations, linear algebra, and individual logarithm, and the time-consuming phases are collecting relations and linear algebra. For the collecting relations phase, we applied several techniques; lattice sieve for the JL06-FFS, lattice sieve with single instruction multiple data (SIMD), and optimization for our parameters. These techniques enable the sieving program to run about 6 times faster. In the linear algebra phase, we applied careful treatments of singleton-clique and merging [15] to the Galois action originating from extension degree 6 of $GF(3^{6\cdot 97})$, with which the size of the matrix used for the Lanczos method is reduced to approximately 30%. By implementing the JL06-FFS with our improvements, we succeeded in solving the DLP over $GF(3^{6\cdot 97})$ by using 252 CPU cores (Core2 quad, Xeon, etc) for the target problem discussed in Section 3.1. As shown in Table 1, the computations required 53.1 days for the collecting relations phase, 80.1 days for the linear algebra phase, and 15.0 days for the individual logarithm phase. Thus, a total of 148.2 days were required to solve the DLP over $GF(3^{6\cdot 97})$ by using 252 CPU cores. Our computational results contribute to the secure use of pairing-based cryptosystems with the η_T pairing.

This paper is organized as follows: In Section 2, we describe the pairing-based cryptosystems, the DLP, and an overview of the FFS. In Section 3, we present our target problem of the DLP over $GF(3^{6\cdot 97})$ and the parameter settings used for our implementation of the FFS. In Section 4, we propose our efficient implementation techniques for the lattice sieve and the parallel Lanczos method. In Section 5, we report the computational results of our implementation of solving the target problem over $GF(3^{6\cdot 97})$. Finally we give concluding remarks and a rough estimation for larger key sizes.

2 Pairing-based cryptosystems and discrete logarithm problem (DLP)

In this section, we briefly explain the security of pairing-based cryptosystems and give a general overview of the function field sieve (FFS). We also mention its parameters such as the smoothness bound B .

Before beginning the discussion, we define the discrete logarithm problem (DLP) over a finite field. Let g_F be a generator of a large multiplicative subgroup G_F of a finite field. For a given $T_F \in G_F$, a DLP over G_F is the problem of computing an integer X such that $T_F = g_F^X$. Generally, X is described as $\log_{g_F} T_F$.

The ECDLP is also defined in the same manner using the additive group operation. Let \mathcal{G}_E be a generator of a subgroup G_E of an elliptic curve E on a finite field and assume that $\mathcal{T}_E \in G_E$ is given. An ECDLP is defined as the problem of computing an integer Y satisfying $\mathcal{T}_E = [Y]\mathcal{G}_E$. In this case, we also describe Y as $\log_{\mathcal{G}_E} \mathcal{T}_E$.

2.1 Pairing-based cryptosystems and DLP

Many efficient cryptographic protocols using a bilinear pairing have been proposed (for example [10–13, 20, 27]), and high-speed implementations for the η_T pairing have been reported (for example [3, 6–9, 17, 18, 24]). We discuss the security of pairing-based cryptosystems with the η_T pairing over $GF(3^n)$ for an integer n . The security of pairing-based cryptosystems with the η_T pairing depends on the difficulty in solving the DLP over the supersingular elliptic curves. Additionally, MOV reduction [26] reduces this problem to a DLP over $GF(3^{6n})^*$ since the embedding degree of the η_T pairing is 6.

In particular, the η_T pairing is a bilinear map such that $\eta_T : G_1 \times G_1 \rightarrow G_2$, where G_1 is an additive subgroup of a supersingular elliptic curve over $GF(3^n)$, G_2 is a cyclic subgroup of $GF(3^{6n})^*$, and the cardinalities of G_1 , G_2 are the same prime number P . The security of pairing-based cryptosystems with the η_T pairing depends on the difficulty of not only an ECDLP over G_1 but also a DLP over G_2 by MOV reduction. To explain this fact, we take ID-based encryption constructed on pairing-based cryptosystems as an example. The ID-based encryption has a master key $s_{key} \in \mathbb{Z}_P$. Each user ID is deterministically transformed into a point $Q_{ID} \in G_1$, and the secret key S_{ID} is defined by $[s_{key}]Q_{ID}$. Therefore, solving the ECDLP over G_1 , namely $S_{ID} = [s_{key}]Q_{ID}$, we obtain the master key $s_{key} = \log_{Q_{ID}} S_{ID}$. Additionally, for an arbitrary point $\mathcal{R} \in G_1$, we compute $\eta_T(S_{ID}, \mathcal{R}), \eta_T(Q_{ID}, \mathcal{R}) \in G_2$, and then have $\eta_T(S_{ID}, \mathcal{R}) = \eta_T([s_{key}]Q_{ID}, \mathcal{R}) = \eta_T(Q_{ID}, \mathcal{R})^{s_{key}} \in G_2$. This implies that $s_{key} = \log_{\eta_T(Q_{ID}, \mathcal{R})} \eta_T(S_{ID}, \mathcal{R})$ is also available by solving the DLP over G_2 . In this paper, we discuss the DLP over a subgroup of $GF(3^{6n})^*$.

2.2 General overview of FFS

The FFS is the asymptotically fastest algorithm for solving a DLP over finite fields of small characteristics. Adleman proposed the first FFS in 1994 [1]. After that, several variants of the FFS have been proposed; Adleman and Huang improved the FFS [2], and Joux and Lercier proposed two more practical FFS's, JL02-FFS [22] and JL06-FFS [23]. The details of JL06-FFS are explained in Sections 3.2.

In this section, we give a general overview of an FFS that consists of four phases: polynomial selection, collecting relations, linear algebra, and individual logarithm. In the overview, we aim at computing $\log_g T$ where $T \in \langle g \rangle \subset GF(3^{6n})^*$.

Polynomial selection phase: We select κ from $\kappa = 1, 2, 3, 6$ for the coefficient field of $GF(3^\kappa)[x]$, and a bivariate polynomial $H(x, y) \in GF(3^\kappa)[x, y]$ such that H satisfies the eight conditions proposed by Adleman [1] and $\deg_y H = d_H$ for a given parameter value d_H . We compute a random polynomial $m \in GF(3^\kappa)[x]$ of degree d_m and a monic irreducible polynomial $f \in GF(3^\kappa)[x]$ such that

$$H(x, m) \equiv 0 \pmod{f}, \quad \deg f = 6n/\kappa. \quad (1)$$

We then have $GF(3^{6n}) \cong GF(3^\kappa)[x]/(f)$. Moreover, there is a surjective homomorphism

$$\xi : \begin{cases} GF(3^\kappa)[x, y]/(H) & \rightarrow & GF(3^{6n}) \cong GF(3^\kappa)[x]/(f) \\ y & \mapsto & m. \end{cases}$$

We select a positive integer B as a smoothness bound, and define a rational factor base $F_R(B)$ and an algebraic factor base $F_A(B)$ as follows.

$$F_R(B) = \{\mathfrak{p} \in GF(3^\kappa)[x] \mid \deg(\mathfrak{p}) \leq B, \mathfrak{p} \text{ is monic irreducible}\}, \quad (2)$$

$$F_A(B) = \{\langle \mathfrak{p}, y - t \rangle \in \text{Div}(GF(3^\kappa)[x, y]/(H)) \mid \mathfrak{p} \in F_R(B), H(x, t) \equiv 0 \pmod{\mathfrak{p}}\}, \quad (3)$$

where $\text{Div}(GF(3^\kappa)[x, y]/(H))$ is the divisor group of $GF(3^\kappa)[x, y]/(H)$ and $\langle \mathfrak{p}, y - t \rangle$ is a divisor generated by \mathfrak{p} and $y - t$. Note that $F_R(0) = F_A(0) = \{\emptyset\}$. We simply call the set $F_R(B) \cup F_A(B)$ a factor base and the set $F_R(k) \setminus F_R(k-1) \cup F_A(k) \setminus F_A(k-1)$ a factor base of degree k for $k = 1, 2, \dots, B$.

Collecting relations phase: We select positive integers R, S and collect a sufficient amount of pairs $(r, s) \in (GF(3^\kappa)[x])^2$ such that

$$\deg r \leq R, \deg s \leq S, \gcd(r, s) = 1, \quad (4)$$

$$rm + s = \prod_{\mathfrak{p}_i \in F_R(B)} \mathfrak{p}_i^{a_i}, \quad (5)$$

$$\langle ry + s \rangle = \sum_{\langle \mathfrak{p}_j, y - t_j \rangle \in F_A(B)} b_j \langle \mathfrak{p}_j, y - t_j \rangle, \quad (6)$$

for some non-negative integers a_i, b_j by using a sieving algorithm such as the lattice sieve discussed in Section 4.1. To efficiently compute b_j in (6), we use the following equivalent property instead of (6):

$$(-r)^{d_H} H(x, -s/r) = \prod_{\langle \mathfrak{p}_j, y - t_j \rangle \in F_A(B)} \mathfrak{p}_j^{b_j}. \quad (7)$$

The (r, s) satisfying (4), (5), and (7) is called a B -smooth pair. Let h be the class number of the quotient field of $GF(3^\kappa)(x)[y]/(H)$ and assume that h is coprime to $(3^{6n} - 1)/(3^\kappa - 1)$. Then the following congruent holds:

$$\sum_{\mathfrak{p}_i \in F_R(B)} a_i \log_g \mathfrak{p}_i \equiv \sum_{\langle \mathfrak{p}_j, y-t_j \rangle \in F_A(B)} b_j \log_g \mathfrak{s}_j \pmod{(3^{6n} - 1)/(3^\kappa - 1)}, \quad (8)$$

where $\mathfrak{s}_j = \xi(t_j)^{1/h}$, $\langle t_j \rangle = h \langle \mathfrak{p}_j, y - t_j \rangle$. We call the congruent (8) ‘‘relation’’ in this paper. Moreover, free relation [19] provides additional relations without computation with a sieving algorithm.

Linear algebra phase: We generate a system of linear equations described as a large matrix from those collected relations and reduce the rank of the matrix by filtering [15]. The reduced system of linear equations is solved using the parallel Lanczos method [4, 19] or other methods, and the discrete logarithms of elements in the factor base are obtained:

$$\log_g \mathfrak{p}_1, \dots, \log_g \mathfrak{p}_{\#F_R(B)}, \log_g \mathfrak{s}_1, \dots, \log_g \mathfrak{s}_{\#F_A(B)}.$$

Individual logarithm phase: Note that our goal is to compute $\log_g T$. Therefore, we find integers a_i, b_j such that

$$\log_g T \equiv \sum_{\mathfrak{p}_i \in F_R(B)} a_i \log_g \mathfrak{p}_i + \sum_{\langle \mathfrak{p}_j, y-t_j \rangle \in F_A(B)} b_j \log_g \mathfrak{s}_j \pmod{(3^{6n} - 1)/(3^\kappa - 1)}, \quad (9)$$

using the special- Q descent [23]. The computational time for the individual logarithm phase is smaller than those for the collecting relations and linear algebra phases.

3 Target problem for $n = 97$ and setting of parameters for FFS

We discuss solving the DLP over a subgroup of $GF(3^{6 \cdot 97})^*$, where the cardinality of the subgroup is 151 bits. To estimate the time complexity of solving such a DLP, we unintentionally set a target problem determined from the circular constant π and natural logarithm e . The details are explained in Section 3.1. To solve the target problem effectively, we select the parameter values of the FFS and estimate important numbers, e.g., the number of elements in the factor base, for it. The details are given in Section 3.2.

3.1 Target problem

For pairing-based cryptosystems, many high-speed implementations of the η_T pairing over supersingular elliptic curves on $GF(3^n)$ have been reported [3, 6–9, 17, 18, 24], and many benchmark tests using the η_T pairing have been conducted for $GF(3^{97})$. In this paper, we deal with a supersingular elliptic curve defined by

$$E := \{(x, y) \in GF(3^{97})^2 : y^2 = x^3 - x + 1\} \cup \{\mathcal{O}\},$$

where \mathcal{O} is the point at infinity. The order of the E is $3^{97} + 3^{49} + 1 = 7P_{151}$ where P_{151} is a 151-bit prime number as follows:

$$P_{151} = 2726865189058261010774960798134976187171462721.$$

Next, let G_1 be the subgroup of E of order P_{151} and let G_2 be the subgroup of $GF(3^{6 \cdot 97})^*$ of order P_{151} . Note that, since orders of G_1 and G_2 are prime numbers, every element of $G_1 \setminus \{\mathcal{O}\}$ and $G_2 \setminus \{1\}$ is a generator of G_1 and G_2 , respectively. The η_T pairing for $n = 97$ is a map from $G_1 \times G_1$ to G_2 .

Our goal is to solve the ECDLP in G_1 . To set our target problem unintentionally, we select two elements $\mathcal{Q}_\pi, \mathcal{Q}_e$ in G_1 , which correspond to the circular constant π and natural logarithm e , respectively. We explain how we select \mathcal{Q}_π and \mathcal{Q}_e as follows. First, we describe $GF(3^{97})$ as $GF(3)[x]/(x^{97} + x^{16} + 2)$, where the irreducible polynomial $x^{97} + x^{16} + 2 \in GF(3)[x]$ is well used for the fast implementation of field operations. An element in $GF(3^{97})$ is represented by $\sum_{i=0}^{96} d_i x^i$, where $d_i \in GF(3) = \{0, 1, 2\}$. To transform π and e to an element in $GF(3^{97})$ respectively, we define a bijective map $\phi : \sum_{i=0}^{96} d_i x^i \mapsto \sum_{i=0}^{96} d_i 3^i \in \mathbb{Z}$. We then transform π and e to the 3-adic integer of 97 digits as follows:

$$\begin{aligned} \lfloor \pi \cdot 3^{95} \rfloor &= (100102110122220102110021111102212222011120121212120012110010010122202221201201211121012101120022)_3, \\ \lfloor e \cdot 3^{96} \rfloor &= (220110112122110201101222210201102122220120222210212212020112112221110001202222112102102010022020)_3. \end{aligned}$$

Table 2. Number of elements \mathfrak{p} in $F_R(B)$ and $F_A(B)$ of $B = 6$

degree of \mathfrak{p}	rational	algebraic
1	27	27
2	351	338
3	6552	6552
4	132678	132496
5	2869776	2869776
6	64566684	64563408
total	67576068	67572597
$\#F_R(6) + \#F_A(6)$	135148665	

From these values, we define $\mathcal{Q}_\pi = (x_\pi, y_\pi)$ and $\mathcal{Q}_e = (x_e, y_e) \in G_1$ as follows. We first find the non-negative smallest 3-adic integers c_π and c_e such that $\phi^{-1}(\lfloor \pi \cdot 3^{95} \rfloor + c_\pi)$ and $\phi^{-1}(\lfloor e \cdot 3^{96} \rfloor + c_e)$ become x -coordinates of the elements \mathcal{Q}_π and \mathcal{Q}_e in the subgroup G_1 on the E . In fact we can set $x_\pi = \phi^{-1}(\lfloor \pi \cdot 3^{95} \rfloor + (11)_3)$ and $x_e = \phi^{-1}(\lfloor e \cdot 3^{96} \rfloor + (120)_3)$. There are two points in $G_1 \setminus \{\mathcal{O}\}$ of the same x -coordinate. We then set the corresponding y -coordinate by computing $y_\pi = (x_\pi^3 - x_\pi + 1)^{(3^{97}+1)/4}$ and $y_e = (x_e^3 - x_e + 1)^{(3^{97}+1)/4}$ in $GF(3^{97})$, respectively. These values are checkable by scripts given in Appendix B.

Again, our goal is to solve the ECDLP in G_1 , i.e., for given $\mathcal{Q}_\pi, \mathcal{Q}_e \in G_1$ we try to find integer s such that

$$\mathcal{Q}_\pi = [s]\mathcal{Q}_e. \quad (10)$$

On the other hand, the η_T pairing enables us to reduce the ECDLP in G_1 to the DLP over G_2 by the relationship $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) = \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)^s$. Therefore, we can find s by computing the discrete logarithm

$$s = \log_{\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)} \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) = \log_g \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) / \log_g \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e) \pmod{P_{151}}, \quad (11)$$

for a generator g of G_2 .

3.2 Parameter settings for FFS

In this section, we explain the parameter setting used for our implementations of the FFS. Hayashi *et al.* [19] reported that, when $n \leq 509$, the JL06-FFS [23] is more efficient for solving the DLP over $GF(3^{6n})$ than the JL02-FFS [22]. Thus, we use the JL06-FFS for our computation. In the JL06-FFS, the condition that “ r is monic” is introduced into the collecting relations phase in order to compute efficiently. For the remainder of this paper, the FFS refers to the JL06-FFS.

To solve our DLP over $GF(3^{6 \cdot 97})$, we have to select several parameter values of the FFS, such that its computational time is small enough for a fixed extension degree n . The parameter values for $n = 97$ are listed in [30, Table 3], and we use those parameter values for our computation.

We can select the parameter κ of the FFS to describe $GF(3^{6 \cdot 97})$ as $GF(3^\kappa)[x]/(f)$, where κ is in $\{1, 2, 3, 6\}$ and $f \in GF(3^\kappa)[x]$ is an irreducible polynomial of degree $6 \cdot 97/\kappa$. The appropriate value of κ is given in [30, Table 3], i.e., $\kappa = 6$. However, we select $\kappa = 3$ for the following reasons. In the linear algebra phase, filtering [15] is performed to reduce the size of the matrix. Then it is required that all elements in the factor base correspond to the memory addresses of the PC for efficient computation. The number of elements in the factor base for $\kappa = 6$ is much larger than that for $\kappa = 3$, so $\kappa = 3$ is advantageous on this point. Additionally, [30, Table 3] shows that the computational cost of the FFS for $\kappa = 3$ is only about twice as much as that for $\kappa = 6$. We conducted test runs for $\kappa = 3, 6$ in the collecting relations phase. We then noticed that our implementation for $\kappa = 3$ was much faster than for $\kappa = 6$, so we set $\kappa = 3$. For the remainder of the paper, $\kappa = 3$.

Polynomial selection phase: We select the bivariate polynomial $H(x, y)$ of the form $x + y^{d_H}$ for a given parameter d_H of the FFS in the same manner as [19]. Then we search an irreducible polynomial $f \in GF(3^\kappa)[x]$ of degree $6 \cdot 97/\kappa$ by factoring $H(x, m)$ for a random polynomial $m \in GF(3^\kappa)[x]$ whose degree is d_m . In fact, we randomly pick up m from $GF(3)[x]$, so that f is also in $GF(3)[x]$ for use of the Galois action. From [30, Table 3], we set d_H and d_m as 6 and 33, respectively. The polynomials f and m , which we used in our experiments, are provided in Appendix B.

Next, we select the smoothness bound $B = 6$ by using [30, Table 3] for (2) and (3), i.e., a rational factor base $F_R(B)$ and an algebraic factor base $F_A(B)$. The number of elements of $F_R(B)$ and $F_A(B)$ are listed in Table 2.

Collecting relations phase: In the collecting relations phase, we use the lattice sieve [28] and the free relation [19] and collect many relations (8); $(r, s) \in (GF(3^\kappa)[x])^2$ satisfying (4), (5), (7), where r is monic. The search range for the lattice sieve depends on the maximum degrees R, S of r, s . We set $R = S = 6$ based on [30, Table 3]. The lattice sieve gives a certain amount of relations for one special- Q , which is defined in Section 4.1. Therefore, we require a sufficient number of special- Q 's so that the number of relations obtained in the collecting relations phase is larger than that of all elements in the factor base. The minimum sufficient number of special- Q 's is estimated by the following process. We have to select special- Q 's from the subset $F_R(6) \setminus F_R(5)$, whose cardinality is 64566684 (See Table 2). Let θ_{min} be the minimum sufficient ratio of special- Q 's over all elements in $F_R(6) \setminus F_R(5)$. For $n = 97$ and $\kappa = 3$, we can estimate $\theta_{min} = 0.01292$ [30, Table 3]. Therefore, the number of special- Q 's must be larger than $\lceil 0.01292 \cdot 64566684 \rceil = 834202$. In our computation, we set 2500000 as the number of special- Q 's to obtain more relations than we require since we expect that these excess relations will help us reduce the size of the matrix during filtering, especially in singleton-clique.

4 Implementation

In this section, we propose the following efficient implementation techniques; the lattice sieve for JL06-FFS and optimization for our parameters in the collecting relations phase, the data structure and the parallel Lanczos method for the Galois action in the linear algebra phase, for reducing the computational cost of the FFS for solving the DLP over $GF(3^{6 \cdot 97})$. Parameters $(\kappa, d_H, d_m, B, R, S)$ are fixed as $(3, 6, 33, 6, 6, 6)$. The reasoning for this is explained in Section 3.2.

4.1 Collecting relations phase

In the collecting relations phase, we used the lattice sieve [28] in a similar fashion to factoring a large integer [25] and solving discrete logarithm problems [21, 22]. We give an overview of our implementation of the lattice sieve in the following paragraphs, and more details are described in Appendix A.

Lattice sieve for JL06-FFS: Sieving with the lattice sieve is performed for $(r, s) \in (GF(3^3)[x])^2$ such that the formula (5) given in Section 2.2 is divisible by an element Q chosen from a subset of the rational factor base $F_R(6) \setminus F_R(5)$ (this Q is called a ‘‘special- Q ’’). Recall that $\deg r$ and $\deg s$ are not greater than $R = 6$ and $S = 6$, respectively. Such (r, s) can be represented as $(r, s) = c(r_1, s_1) + d(r_2, s_2)$ for given reduced lattice bases $(r_1, s_1), (r_2, s_2) \in (GF(3^3)[x])^2$ and any $c, d \in GF(3^3)[x]$ such that $\deg(cr_1 + dr_2) \leq 6, \deg(cs_1 + ds_2) \leq 6$, then sieving is done on the bounded c - d plane. After sieving, we conduct the smoothness test [16] for ‘‘candidates’’ that are evaluated as B -smooth pairs with high probability by using the lattice sieve. Our implementation of the smoothness test is described in Appendix A.4.

A problem of applying the lattice sieve to the FFS is the condition ‘‘ r is monic’’ described in Section 3.2. Since r is represented as $cr_1 + dr_2$, it is difficult to efficiently keep r monic — it might require degree evaluations and branches. Instead of choosing monic r , we introduce the condition $r \equiv 1 \pmod{x}$. To satisfy this condition, we restrict r_1 and r_2 such that $r_1 \equiv 0 \pmod{x}$ and $r_2 \equiv 1 \pmod{x}$. Then sieving is performed on the bounded c - d plane with restriction $d \equiv 1 \pmod{x}$, whose size is reduced to $1/27$ compared with the original bounded c - d plane. This sieving procedure with the restricted condition can be implemented without extra costs such as additional degree evaluations and additional branches.

Lattice sieve with SIMD: Since operations of $GF(3)$ can be represented using logical instructions [24], operations of $GF(3^3)[x]$ can be performed using a combination of logical and shift instructions. This means SIMD implementation is appropriate for efficient computation of the lattice sieve. We represent $GF(3^3)$ as polynomial basis $GF(3)[\omega]/(\omega^3 - \omega - 1)$, and its element is represented using 6-bit $(h_1, \ell_1, h_\omega, \ell_\omega, h_{\omega^2}, \ell_{\omega^2}) \in GF(2)^6$ in our implementation. We then pack 16 elements of $GF(3^3)[x]$ of degree at most 7 into 6 registers of 128 bits, as shown in Fig. 2 in Appendix A.1, and treat 16 elements with the SIMD. Note that the upper bound of the degree of our SIMD data structure is for efficient access to each element in $GF(3^3)[x]$. On the other hand, since we choose B, R, S as all 6, the upper bound of the degrees of $c, d, r_1, s_1, r_2, s_2 \in GF(3^3)[x]$ and \mathfrak{p} in the factor base, which are treated in the lattice sieve, is also 6. Therefore, our SIMD structure can be stored elements treated in the lattice sieve.

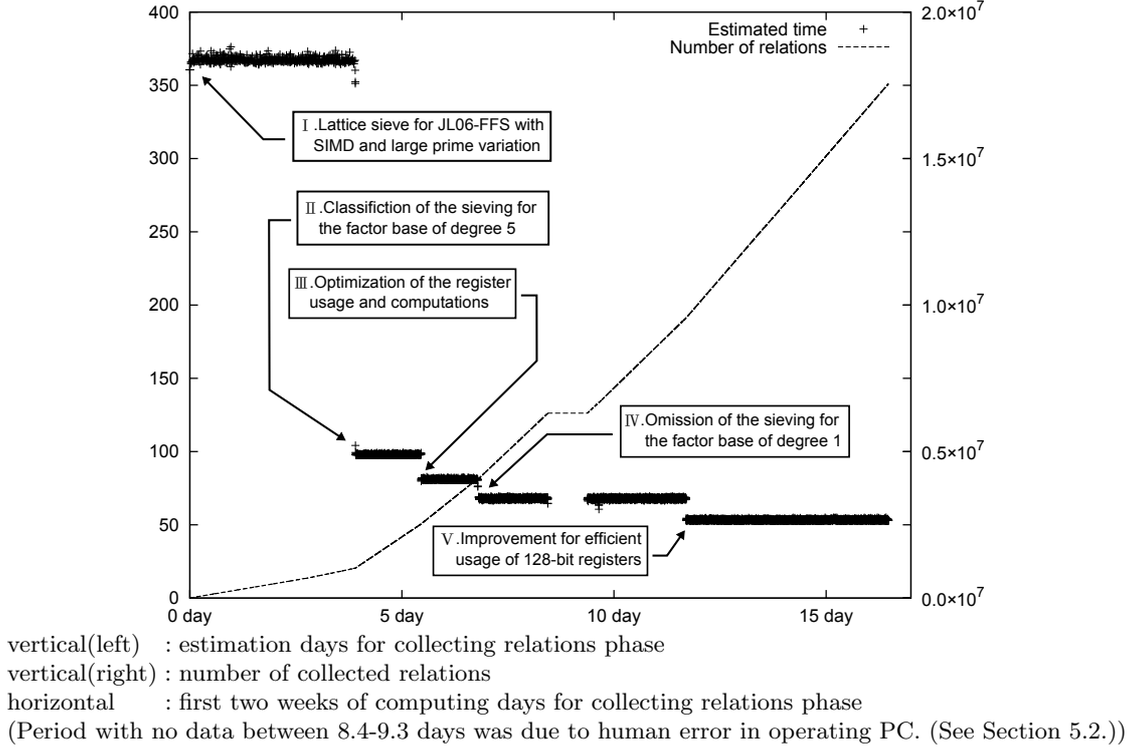


Fig. 1. Our improvement in collecting relations phase for first two weeks

History of our optimizations: Figure 1 shows the process of our improvements in the collecting relations phase for the first two weeks. We improved our implementation of the lattice sieve four times during this period. We first used large prime variation to omit sieving for the factor base of degree 6 and implemented the lattice sieve for the FFS with the SIMD implementation. We then ran the program for the first four days (stage I in Fig. 1). At that point, the estimated total number of days for the collecting relations phase was about 360 days. While the sieving program was running, we found that sieving for the factor base of degree 5 requires heavier computation than sieving for the factor bases of degree 1, 2, 3 and 4. Therefore, we improved sieving for the factor base of degree 5; thus, our sieving program became over 3 times faster than before (stage II in Fig. 1). Next, we optimized register usage for input values and omitted wasteful computations (stage III in Fig. 1). Additionally, we omitted sieving for the factor base of degree 1 (stage IV in Fig. 1), since that computational time was larger than the computational time for the factor bases of degree 2, 3, 4, and 5. Moreover, we improved our sieving program to use 128-bit registers more efficiently (stage V in Fig. 1). Finally, our sieving program became about 6 times faster than the first one (stage I in Fig. 1) and the estimated total number of days for the collecting relations phase became about 53.1 days. In the next paragraph, we explain the details of the improvement in stage II, which is the most effective and important improvement in our implementation of the lattice sieve.

Details of stage II: In the lattice sieve, the main computation of sieving for given lattice bases $(r_1, s_1), (r_2, s_2) \in (GF(3^3)[x])^2$ is as follows. For fixed $d \in GF(3^3)[x]$, whose degree is upper-bounded by a degree bound D , we compute $c_0 \equiv -d(r_1t + s_1)^{-1}(r_2t + s_2) \pmod{\mathfrak{p}}$ for all pairs $(\mathfrak{p}, t) \in \{(\mathfrak{p}, t) \mid \mathfrak{p} \in F_R(B), t \equiv m \pmod{\mathfrak{p}}\} \cup \{(\mathfrak{p}, t) \mid \langle \mathfrak{p}, y-t \rangle \in F_A(B)\}$, and compute $c \in GF(3^3)[x]$, whose degree is upper-bounded by a degree bound C , such that $c = c_0 + k\mathfrak{p}$ where $k \in GF(3^3)[x]$. We call the computation “sieving at d ” in this section. For given lattice bases, sieving at d is performed for all d of degree not larger than D . Note that c_0 does not need to be computed when $(r_1t + s_1) \equiv 0 \pmod{\mathfrak{p}}$; therefore we assume $(r_1t + s_1) \not\equiv 0 \pmod{\mathfrak{p}}$ in the following description.

In stage I of our implementation, we found that the time of sieving at d for $\deg \mathfrak{p} = 5$ takes over 100 msec, but each sieving time at d for $\deg \mathfrak{p} = 1, 2, 3$ and 4 takes about 10 msec or less. Therefore, we tried to improve the sieving of degree 5. When we compute c_0 for \mathfrak{p} of degree 5, the degree of c_0 becomes 4 with probability about $26/27$. On the other hand, the degree of the lattice bases r_1, s_1, r_2, s_2 is 3 in most cases because the degree of

special- Q is 6. On such bases, degree bounds C and D can be chosen as 3 to satisfy condition (4), i.e., $\deg r \leq 6$ and $\deg s \leq 6$. These facts show that about 26/27 of the computation of sieving for \mathfrak{p} of degree 5 are waste computations. Therefore, we discuss how to sieve only with the polynomial c_0 , whose degree is not larger than 3, as follows.

Let $\alpha \in GF(3^3)[x]$ be $-(r_1t + s_1)^{-1}(r_2t + s_2) \bmod \mathfrak{p}$, then we have $c_0 = d\alpha \bmod \mathfrak{p}$. Let $\alpha_i \in GF(3^3)$ be the coefficient of the fourth-order term of $x^i\alpha \bmod \mathfrak{p}$ for $i = 0, 1, 2, 3$. Since $\deg d \leq 3$, d is represented as $d_3x^3 + d_2x^2 + d_1x + 1$ for $d_3, d_2, d_1 \in GF(3^3)$. Recall that we restricted $d \equiv 1 \bmod x$ in our implementation of the lattice sieve. Here we know that the degree of c_0 is not larger than 3 if $d_3\alpha_3 + d_2\alpha_2 + d_1\alpha_1 + \alpha_0 = 0$. Therefore, it is sufficient to perform sieving at d for \mathfrak{p} in the factor base of degree 5 for only d satisfying the following property:

$$d_1 = \begin{cases} -K\alpha_1^{-1} & \text{if } \alpha_1 \neq 0 \\ \text{any element in } GF(3^3) & \text{if } \alpha_1 = 0 \text{ and } K = 0 \end{cases} \quad (12)$$

where $K = d_3\alpha_3 + d_2\alpha_2 + \alpha_0$. When $\alpha_1 = 0$ and $K = 0$, we should compute c_0 for d whose d_1 is any element in $GF(3^3)$, and we cannot cut off any d_1 ; therefore, we assume that $\alpha_1 \neq 0$ in the following description. Suppose that we now fix lattice bases $(r_1, s_1), (r_2, s_2)$ and a pair (\mathfrak{p}, t) where $\deg \mathfrak{p} = 5$, then each α_i for $i = 0, 1, 2, 3$ is also fixed. Therefore, since K depends on d_2 and d_3 , the d_1 satisfying (12) is given by d_2 and d_3 and uniquely determined for given d_2 and d_3 . This implies that, since d_1 is in $GF(3^3)$ whose cardinality is 27, we can ignore 26 d_1 's not satisfying (12) for given d_2 and d_3 . In fact, the time of sieving at d for all pairs (\mathfrak{p}, t) where $\deg \mathfrak{p} = 5$ is reduced to about 1.5 msec by ignoring d_1 not satisfying (12). Note that we need to compute K for given d_2 and d_3 for all pairs (\mathfrak{p}, t) . The time of computing K for all (\mathfrak{p}, t) takes about 150 msec in our implementation. Therefore, for all pairs (\mathfrak{p}, t) where $\deg \mathfrak{p} = 5$, the computations of K and sieving at d require about 7.1 msec at stage II, which is over 10 times faster than the computation of sieving at d at stage I. As a result, our implementation of the lattice sieve at stage II becomes over 3 times faster than that at stage I.

4.2 Linear algebra phase

After the collecting relations phase, we obtain a system of linear equations modulo P_{151} , which is described in Section 2.1. The Galois action [19, 23] can reduce the number of variables of the system of linear equations to one-third. Additionally, after the Galois action, the numbers of equations and variables of the system of linear equations can be further reduced using filtering [15], i.e., singleton-clique and merging. To solve the system of linear equations defined by this reduced matrix, we use the parallel Lanczos method [4, 19].

Galois action: The Galois action to $GF(3^{6 \cdot 97})/GF(3^{3 \cdot 97})$ enables us to reduce the number of variables of the system of linear equations to one-third (details of the Galois action are discussed in [19, 23]). However, when we use the Galois action, 151-bit large integers such as $e_0 + e_1\tau + e_2\tau^2$, where $\tau = 3^{97^2} \bmod P_{151}$ and e_i is a small integer of a few bits, are added to elements of the system of linear equations. This unfortunate fact eventually increases the data size of the reduced matrix; therefore, high-capacity memory is required. To allay the increase in the representation size of the elements, we store only a triplet (e_1, e_2, e_3) in the PC memory, not a large 151-bit integer. Since e_i is small enough to be represented by 8 bits, the size of the elements is reduced from 151 to 24 bits on average. We call this representation the “ τ -adic structure”. Note that the τ -adic structure is used for the Galois action and singleton-clique.

Singleton-clique: Singleton-clique [15] deletes unnecessary rows and columns to reduce the size of the matrix. In our implementation, singleton-clique is performed by maintaining 20000 more rows than columns to prevent accidentally decreasing the rank of the matrix.

Merging: Merging [15] is a weight-controlled Gaussian elimination, where the weight is the number of non-zero elements of a matrix. For some small integer k , the column with a weight smaller than or equal to k is deleted by row eliminations with controlling the pivot selection so that the weight of the matrix is as small as possible. This operation is called k -way merging. During the merging computation, elements of the matrix generally become large since row eliminations are computed in merging. On the τ -adic structure, we must conduct merging under the restriction that e_i is not larger than 8 bits; therefore, results of row elimination, which cannot be stored on the τ -adic structure, often appear. On the other hand, there is no such restriction on a large 151-bit integer structure because such a structure can represent all integers in $\mathbb{Z}_{P_{151}}$. This means that the size of the merged matrix on a large 151-bit integer structure is smaller than that on the τ -adic structure. Thus, a large 151-bit

integer structure is better for merging than the τ -adic structure if the matrix represented by a large 151-bit integer structure can be stored on the PC memory. Fortunately, the size of the matrix reduced using singleton-clique is small enough to store the matrix represented by a large 151-bit integer structure on the PC memory; therefore, we convert the data representation of the matrix from the τ -adic structure to a large 151-bit integer structure.

Parallel Lanczos method: By using the parallel Lanczos method [4, 19], we solve the system of linear equations defined by the matrix reduced via the Galois action, singleton-clique, and merging. For parallel computing, the matrix should be split into sub-matrices, i.e., split into $N = N_1 \times N_2$ sub-matrices for N nodes, and nodes communicate among N_1 nodes or N_2 nodes. To reduce the synchronization time before communicating among N_1 nodes or N_2 nodes, the matrix is split so that each sub-matrix has almost the same weight. Our machine environment for the parallel Lanczos method consisted of 22 nodes, and each node had 12 CPU cores and 2 NICs ((h) in Table 9 in Appendix C). The 2 NICs were connected to a 48-port Gbit HUB, i.e., 44 ports were used for connecting 22 nodes. All 22 nodes could be used, so we had a choice for machine environment; $20 = 5 \times 4$, $21 = 7 \times 3$ or $22 = 11 \times 2$. Using 20 nodes requires the least communication costs but the most computational costs, and using 22 nodes requires the most communication costs but the least computational costs. Using 21 nodes was the best for our implementation; therefore, we used 21 nodes for the computation of the parallel Lanczos method.

For computation in the parallel Lanczos method, many modular multiplications of 151-bit integers \times 151-bit integers modulo P_{151} are required due to the Galois action. We implemented Montgomery multiplication optimized to 151-bit integers using assembly language. Our program then becomes several times faster than straightforward modular multiplication using GMP (<http://gmplib.org/>) for multiple precision arithmetic.

After the computation of the parallel Lanczos method started, we improved our codes of the parallel Lanczos method (for example, efficient register usage, overlapping communications and computations). These improvements are about 15% faster than our initial implementation.

4.3 Individual logarithm phase

As mentioned in Section 3.1, $\log_g \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi)$ and $\log_g \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)$ are required to solve our target problem. To compute them, rationalization and special- Q descent [23] were used. For simplicity, let T be $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi)$, or $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)$ in the following paragraphs.

Rationalization: To reduce the computational costs of the special- Q descent, which is described in the next paragraph, we randomize T such that the randomized element is M -smooth for a small enough integer $M > B$ by the following process. First, we randomize T by $z \equiv g^\gamma T \pmod{f}$ for a random integer $\gamma \in \mathbb{Z}_{P_{151}}$. We then rationalize z as $z \equiv z_1/z_2 \pmod{f}$ where degrees of z_1 and z_2 are about $\deg f/2$. Note that for an integer M , the probability that both z_1 and z_2 are M -smooth is usually higher than the probability that z is M -smooth, and it is better to rationalize to obtain M -smooth elements. We gather many such pairs (z_1, z_2) and calculate upper bounds M_1, M_2 of the degrees of the irreducible factors of z_1, z_2 for each pair, respectively. Note that if both M_1 and M_2 are small, the computational time of special- Q descent decreases. Therefore, we search a pair (z_1, z_2) such that those upper bounds M_1, M_2 are small enough. Since the logarithm of the target element T is described as

$$\log_g T \equiv \log_g z_1 - \log_g z_2 - \gamma \pmod{P_{151}},$$

we perform special- Q descent for all irreducible factors of such z_1, z_2 to compute those logarithms $\log_g z_1, \log_g z_2$.

Special- Q descent: M_i smooth elements z_i obtained by the rationalization, where $M_i > B$ for $i = 1, 2$, contain some irreducible factors of degree larger than B whose logarithms are not computed in the linear algebra phase. To compute these logarithms, the special- Q descent [23] is usually used. First, we perform special- Q descent for irreducible factors \mathfrak{p} of z_i , i.e., the lattice sieve is conducted with \mathfrak{p} as special- Q , and search $(r, s) \in (GF(3^3)[x])^2$ such that the degrees of the irreducible factors of polynomials (5) and (7), except \mathfrak{p} , are less than $\deg \mathfrak{p}$. When a relation generated by such (r, s) has variables corresponding to irreducible polynomials \mathfrak{p}' or prime divisors $\langle \mathfrak{p}', y - t \rangle$ such that $\deg \mathfrak{p}' > B = 6$, the lattice sieve is continued with \mathfrak{p}' or $\langle \mathfrak{p}', y - t \rangle$ as special- Q . Note that the degree of special- Q is descended by doing such a procedure recursively, finally reaching $B = 6$. After all computations of special- Q descent, we can find the logarithm of \mathfrak{p} by backtracking all obtained relations.

Table 3. Number of collected relations in collecting relations phase

lattice sieve	159032292 relations obtained from 2500000 special- Q 's (64.91 relations/special- Q , 389 sec/special- Q)
	153815493 unique (non-duplicated) relations obtained from 2449991 unique special- Q 's in the above 2500000 special- Q 's
free relation	33786299 relations
total	187602242 relations (consist of 134697663 elements in the factor base)

5 Experimental results

We succeeded in solving a DLP over $GF(3^{6 \cdot 97})$ by using the FFS with our efficient implementation techniques discussed in Section 4. In this section, we report our computation results, such as the computational time of each phase of the FFS and the number of relations.

5.1 Polynomial Selection:

The FFS has six parameters $\kappa, d_H, d_m, B, R,$ and S , as defined in Section 2.2, and we set $(\kappa, d_H, d_m, B, R, S) = (3, 6, 33, 6, 6, 6)$ for our target problem, based on the reason given in Section 3.2. In the polynomial selection phase, we can extract appropriate polynomials such as the definition polynomial $H(x, y)$ of a function field described in Section 3.2 in one minute, so the computational cost of the polynomial selection phase is negligibly small.

5.2 Collecting relations phase

In the collecting relations phase, we search many relations that are equations of the form (8) to generate a system of linear equations by using the lattice sieve and the free relation. We explain our computational results of the collecting relations phase, e.g., the number of relations obtained in this phase, the computational time of the lattice sieve for one special- Q .

Lattice sieve: Each special- Q has to be chosen from $F_R(6) \setminus F_R(5)$. The number of elements of $F_R(6) \setminus F_R(5)$ is 64563408, and the size of the table of those elements is about 500 MB. Since our program of the lattice sieve is computed using many nodes, it is not convenient to pick up the element from that 500-MB table as a special- Q . Therefore, we selected a special- Q by randomly generating an irreducible polynomial in $GF(3^3)[x]$ of degree 6, which is in $F_R(6) \setminus F_R(5)$, and iterated the computation of the lattice sieve for the special- Q .

We prepared 47 PCs (in total 212 CPU cores) of (a)-(c) and (e)-(g) in Table 9 in Appendix C for the lattice sieve. The computation of the lattice sieve began on May 14, 2011, and we continued optimizing our program of the collecting relations phase. Figure 1 shows the process of our improvements in the collecting relations phase for the first two weeks. The total time for the collection of relations shortened due to our improvements. In the center of Fig. 1, there is a period in which no relations were obtained, from 8.5 to 9.4 days. This was due to human error; therefore, we wasted computer power during one day since a set of “seeds” to compute the program for each PC had been exhausted in the server PC during this period. As discussed in Section 4.1, we applied several improvements to our program of the collecting relations phase; lattice sieve for JL06-FFS, the lattice sieve with SIMD, and optimization for our parameters. Finally, as Table 8 in Appendix C shows, the computation finished on September 9, 2011 and required 118 days including the loss-time of some programming errors, updating our codes, and power outages. The real computational time of the lattice sieve was equivalent to 53.1 days using 212 CPU cores such as Xeon E5440.

Table 3 summarizes the process of generating relations in the collecting relations phase. The first to third rows in the right column in Table 3 mean that we randomly selected 2500000 special- Q 's from 64563408 elements in $F_R(B)$ of degree 6 and obtained 159032292 relations by using the lattice sieve. For one special- Q , our program generated 64.91 relations on average, and this computation required 389 sec. Since the special- Q 's were randomly selected from the 64563408 elements, there were several duplicate special- Q 's. Therefore, there also existed duplicate relations given by those duplicate special- Q 's. The fourth to fifth rows in the right column in Table 3 imply that, after removing such duplicate relations from all the obtained relations, 153815493 unique (non-duplicated) relations remained. It might seem that the number of duplicate relations is very small compared to the integer factorization case using the number field sieve. This arises from the fact that the size of the sieving space in our parameters is so large compared to that case.

Table 4. Compressing matrix using Galois action, singleton-clique and merging

method	size of matrix
before compressing	187602242 equations \times 134697663 variables
Galois action	159394665 equations \times 45049572 variables
singleton-clique	14060794 equations \times 14040791 variables
6-way merging	6141443 equations \times 6121440 variables

Table 5. Matrices after merging and computational time of parallel Lanczos method for matrices

k -way merging	#row	#column	weight of matrix	weight/#row	estimated time (parallel Lanczos method)
-	14060794	14040791	261454353	18.595	236.3 days
2	11156100	11136095	253671023	22.738	158.1 days
3	8573837	8553832	301766399	35.196	108.9 days
4	7324638	7304633	365468344	49.896	92.5 days
5	6622387	6602382	409017570	61.763	84.2 days
6	6141443	6121440	455224330	74.123	80.07 days
7	5754191	5734186	506738984	88.064	77.79 days
8	5479424	5459419	556209807	101.509	77.07 days
9	5251949	5231944	607254883	115.625	77.08 days
10	5076238	5056233	656586248	129.345	77.70 days

Free relation: The free relation gives us additional relations not generated by a sieving algorithm such as the lattice sieve. The details of the free relation is given in [19]. As shown in the sixth row in the right column in Table 3, the free relation gave us 33786299 relations. The seventh row in the right column in Table 3 means that we obtained 187602242 relations in total by adding relations given by the free relation to those generated using the lattice sieve. Eventually, we obtained a system of linear equations consisting of 187602242 equations and 134697663 variables. Note that there are 451002 elements in the factor base, which does not appear in the 187602242 relations

5.3 Linear algebra phase

Galois action : As mentioned in Section 4.2, the Galois action reduced the size of the matrix generated in the collecting relations phase to one-third since $\kappa = 3$. In fact, as the third row in the right column in Table 4 shows, the number of variables was reduced from 134697663 to 45049572, and the number of equations decreased from 187602242 to 159394665. This implies that the Galois action is appropriate for reducing variables of the matrix. To allay the fact that the size of each element of the matrix increases from a few bits to 151 bits due to the Galois action, we used the τ -adic structure mentioned in Section 4.2.

Singleton-clique : After using the Galois action, we additionally reduce the variables and equations of the matrix by singleton-clique [15]. To positively obtain the solutions of the system of linear equations in the linear algebra phase, we performed singleton-clique while keeping the number of variables larger than that of the equations by 20000. Therefore, as the fourth row in the right column of Table 4 shows, we obtained the matrix consisting of 14060794 rows and 14040791 columns. This computation takes 3 hours with a PC of (e) in Table 9 in Appendix C.

Merging : Before using the parallel Lanczos method to solve the system of linear equations reduced by singleton-clique, we can additionally reduce the matrix by merging [15]. We performed 2- to 10-way merging for the matrix reduced by singleton-clique, and estimated the computational time of our implementation of the parallel Lanczos method from the size of the reduced matrix in Table 5. The table shows that the 8-way merged matrix is better than the others in terms of the computational time of the parallel Lanczos method. However, we used the 6-way merged matrix in our computation, not the 8-way merged matrix, for the following reason. We began with the computation of 2-way merging for the matrix reduced by singleton-clique. After finishing this computation, we started 3-way merging for the 2-way merged matrix. In the same manner, we also performed 2- to 10-way merging in turn. Until 6-way merging, computation was done using a PC of (e) in Table 9 in Appendix C, which had 16-GB RAM. In 7-way merging, the computation was running out of the 16-GB RAM, so 7-way merging was not able to be computed on the PC. Therefore, we started the computation of the parallel

Table 6. Computational time of parallel Lanczos method for matrix reduced by 6-way merging

calculation time/loop	626.3 msec
synchronization time/loop	46.5 msec
communication time/loop	457.3 msec
total time/loop	1130.1 msec
number of loops	6121438
total time of parallel Lanczos method	80.1 days

Lanczos method for the 6-way merged matrix. After that, we had an opportunity to use a workstation that had 128-GB RAM and performed 7- to 10-way merging on the workstation. Eventually, we noticed that the computational cost of the Lanczos method for the 8-way merged matrix was three days less than that for 6-way merging. However, by this time, the computation of the parallel Lanczos method for the 6-way merged matrix had already been running for about seven days, so we continued computing the parallel Lanczos method for the 6-way merged matrix. The fifth row in the right column of Table 4 means that the 6-way merged matrix consisted of 6121440 variables and 6141443 equations. The entire computation for our merging took about 10 hours.

Parallel Lanczos method : We used the parallel Lanczos method [4, 19] to solve the system of linear equations defined by the 6-way merged matrix. Note that this matrix is sparse and defined over $\mathbb{Z}_{P_{151}}$, where P_{151} is the 151-bit prime number given in Section 3.1. The computation of the parallel Lanczos method started on January 16, 2012, and was conducted on 21 PCs (in total 252 CPU cores) of (h) in Table 9 in Appendix C, which were connected via a 48-port Gbit HUB. As mentioned in Section 4.2, we continued improving our codes of the parallel Lanczos method after computation began. The computational times of our improved codes of the parallel Lanczos method are listed in Table 6. The number of loops of this method is equal to the rank of the 6-way merged matrix, and was at most 6121440. The computation for one loop required 626.3 msec on average. Additionally, for one loop, the synchronization time for communication among nodes was 46.5 msec on average, and the communication time for exchanging data among nodes was 457.3 msec on average. Therefore, the average time for one loop was 1130.1 msec. Finally, computation finished on April 14, 2012. The number of loops was 6121438 and the computation for the parallel Lanczos method took 90 days including time losses similar to our implementation of the lattice sieve. The real computational time is equivalent to 80.1 days using 252 CPU cores such as Xeon X5650.

5.4 Individual logarithm phase

Our target is to compute the discrete logarithm $\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e))$ and $\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi))$ for some $g \in G_2$, as mentioned in Section 3.1.

Rationalization: Let g be a polynomial $(x + \omega)^{(3^{6 \cdot 97} - 1)/P_{151}} \in G_2$, where ω is a polynomial basis of $GF(3^3) \cong GF(3)[\omega]/(\omega^3 - \omega - 1)$. Note that g is a generator of $G_2 \subset GF(3^{6 \cdot 97})^*$ and $x + \omega$ is a monic irreducible polynomial in $F_R(B)$ of degree 1. We search a pair (z_1, z_2) (and (z'_1, z'_2)) $\in (GF(3^3)[x])^2$ such that $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e) \cdot g^{\gamma_1} = z_1/z_2$ (and $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) \cdot g^{\gamma_2} = z'_1/z'_2$), where z_i (and z'_i) are M_i -smooth for some $\gamma_1, \gamma_2 \in \mathbb{Z}_{P_{151}}$ and $i = 1, 2$. To reduce the computational time of the following special- Q descent, we tried to find as small an M_i as possible within our computational resources for $i = 1, 2$. In fact, we found z_1 and z_2 , which are 13- and 15-smooth (and z'_1 and z'_2 which are 15- and 14-smooth), respectively. These computations were conducted on 168 CPU cores (PCs of (a)-(d) in Table 9 in Appendix C) and required 7 days for each computation.

$$\begin{aligned} \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e) \cdot g^{\gamma_1} &= (13\text{-smooth})/(15\text{-smooth}) \\ \gamma_1 &= 2514037766787322013334785428291787565870435706 \\ \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) \cdot g^{\gamma_2} &= (15\text{-smooth})/(14\text{-smooth}) \\ \gamma_2 &= 2657516740789758289434702436228062607247517136 \end{aligned}$$

Special- Q descent: We performed special- Q descent for each irreducible factor of smooth elements obtained by the rationalization. First, we discuss the detailed results of the computation for $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)$. Table 7 shows

Table 7. Computational time for special- Q descent in individual logarithm phase

degree of special- Q	#special- Q	time for one special- Q
15	1	159 sec
14	3	71 sec
13	4	39 sec
12	6	62 sec
11	21	41 sec
10	12	53 sec
9	22	594 sec
8	40	747 sec
7	194	3287 sec

the number of irreducible factors in the computations of special- Q descent and the time for computing them for each special- Q . These computations were conducted on 168 CPU cores (PCs of (a)-(d) in Table 9 in Appendix C) and took about 0.5 days in total. The computation for $\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi)$ also took about 0.5 days in total on the same machine environment. Thus, as shown in Table 1, the computation of the individual logarithm phase took 15 days; 7 days (for rationalization) \times 2 elements + 0.5 days (for special- Q descent) \times 2 elements.

By using the logarithms of the corresponding elements in the factor base obtained from the linear algebra phase, we could compute $\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e))$ and $\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi))$. The logarithm of each element is as follows:

$$\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)) = 1540966625957007958347823268423957036469656370,$$

$$\log_g(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi)) = 1630281950635507295663809171217833096970449894.$$

Finally, we obtained the logarithm of the target element $\log_{\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)}(\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi))$ as follows:

$$\log_{\eta_T(\mathcal{Q}_\pi, \mathcal{Q}_e)} \eta_T(\mathcal{Q}_\pi, \mathcal{Q}_\pi) = 1752799584850668137730207306198131424550967300. \quad (13)$$

This is the solution of the ECDLP of equation $\mathcal{Q}_\pi = [s]\mathcal{Q}_e$ in (10). Scripts for checking this solution are provided in Appendix B.

6 Concluding remarks

We evaluated the security of pairing-based cryptosystems using the η_T pairing over supersingular elliptic curves on finite field $GF(3^n)$. We focused on the case of $n = 97$ since many implementers have reported practically relevant high-speed implementations of the η_T pairing with $n = 97$ in both software and hardware. In particular, we examined the difficulty in solving the discrete logarithm problem (DLP) over $GF(3^{6 \cdot 97})$ by our implementation of the function field sieve (FFS).

To reduce the computational cost of the FFS for solving the DLP over $GF(3^{6 \cdot 97})$, we proposed several efficient implementation techniques. In the collecting relations phase, we implemented the lattice sieve for the JL06-FFS with SIMD and introduced improvements by optimizing for factor bases of each degree; therefore, our lattice sieve for the JL06-FFS became about 6 times faster than the first one. The main difference from the number field sieves for integer factorization is the linear algebra phase, namely, we have to deal with a large modulus of 151-bit prime for the computation of the FFS. We thus performed filtering (singleton-clique and merging) by carefully considering the data structure of large integers developing from the Galois action, so that we can efficiently conduct the parallel Lanczos method. From the above improvements, we succeeded in solving the DLP over $GF(3^{6 \cdot 97})$ in 148.2 days by using PCs with 252 CPU cores. Our computational results contribute to the security estimation of pairing-based cryptosystems using the η_T pairing. In particular, they show that the security parameter of such pairing-based cryptosystems must be chosen with $n > 97$.

Finally, we show a very rough estimation of required computational power for solving the DLP over $GF(3^{6n})$ with $n > 97$. Our experiment on the DLP over $GF(3^{6n})$ with $n = 97$ used 252 CPU cores of mainly 2.67 GHz Xeon for 148.2 days, which are equivalent to $2^{62.9}$ clock cycles. From the analysis of [30], the computational complexities of breaking the DLP over $GF(3^{6n})$ with $n = 163$ and 193 become $2^{15.4}$ and $2^{19.1}$ times larger than that with $n = 97$, respectively. Therefore, we could estimate that about $2^{78.3}$ and $2^{82.0}$ clock cycles are required for breaking the DLP over $GF(3^{6n})$ with $n = 163$ and 193, respectively. On the other hand, the currently fastest supercomputer K has a throughput of about 10.5 petaflop/s from <http://www.top500.org/>, and it performs about $2^{78.1}$ floating-point operations for one year. If one floating-point operation on the CPU of the K is equivalent to one clock cycle of logical operation on the Xeon core, we might be able to break the DLP over $GF(3^{6 \cdot 163})$ using our implementation on supercomputer K for one year.

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A Details of our implementation of collecting relations phase

In this appendix, we describe the details of our implementations of the collecting relations phase discussed in Section 4.1. As mentioned in Section 3.2, parameters $(\kappa, d_H, d_m, B, R, S)$ are fixed as $(3, 6, 33, 6, 6, 6)$.

Naive lattice sieve: First, we begin with the explanation of the naive lattice sieve for introducing notations. We define $\bar{F}_R(B) = \{(\mathfrak{p}, t) \mid \mathfrak{p} \in F_R(B), t \equiv m \pmod{\mathfrak{p}}\}$ and $\bar{F}_A(B) = \{(\mathfrak{p}, t) \mid \langle \mathfrak{p}, y - t \rangle \in F_A(B)\}$. In the lattice sieve, sieving is performed on $(r, s) \in (GF(3^\kappa)[x])^2$ such that $rm + s$ (resp. $r^6x + s^6$ due to $H(x, y) = x + y^6$) is divisible by q , where $(q, u) \in \bar{F}_R(B)$ (resp. $\bar{F}_A(B)$). This $Q = (q, u)$ is called ‘‘special- Q ’’⁴. The (r, s) in the lattice sieve is represented as $c(r_1, s_1) + d(r_2, s_2)$ for $c, d \in GF(3^3)[x]$ of degree upper-bounded by C and D , respectively, where $(r_1, s_1), (r_2, s_2)$ are reduced lattice bases of $(0, q), (1, -t)$, and sieving is performed on the bounded c - d plane.

From among such pairs (r, s) , we search a pair (c, d) such that $rm + s$ (resp. $r^6x + s^6$) is divisible by \mathfrak{p} , where $(\mathfrak{p}, t) \in S_R \subseteq \bar{F}_R(B)$ (resp. $S_A \subseteq \bar{F}_A(B)$) is not equal to (q, u) . We call sets S_R and S_A the rational sieving factor base and algebraic sieving factor base, respectively. The pair (c, d) is represented as $(c_0 + k\mathfrak{p}, d)$ for $k \in GF(3^3)[x]$ and a fixed d , where

$$c_0 \equiv \begin{cases} -d(r_1t + s_1)^{-1}(r_2t + s_2) \pmod{\mathfrak{p}} & \text{if } (r_1t + s_1) \not\equiv 0 \pmod{\mathfrak{p}} \\ \text{any polynomials} & \text{if } (r_1t + s_1) \equiv 0 \text{ and } d \equiv 0 \pmod{\mathfrak{p}} \end{cases}$$

This fact enables us to know which polynomial is the factor of $rm + s$ (resp. $r^6x + s^6$) without factoring these polynomials. Thus, by computing $c_0 + k\mathfrak{p}$ for a fixed d and each $\mathfrak{p} \in S_R$ (resp. S_A), we can obtain (c, d) such that $rm + s$ (resp. $r^6x + s^6$) is divisible by \mathfrak{p} . For one (r, s) and a certain number of factors \mathfrak{p} of $rm + s$ (resp. $r^6x + s^6$), if the sum of the degrees of all the \mathfrak{p} reaches a threshold, the $rm + s$ (resp. $r^6x + s^6$) is B -smooth with high probability. Such (r, s) is called a ‘‘candidate’’. For each candidate, we determine whether it is a B -smooth pair by using the smoothness test [16].

In practice, a threshold is given by $\deg(rm + s) - \epsilon$ (resp. $\deg(r^6x + s^6) - \epsilon$), where ϵ is a non-negative integer, called a ‘‘margin’’. For a larger margin, we may obtain more relations but have to conduct additional smoothness tests. Conversely, for a smaller margin, we may lose some relations; however, the number of computations of the smoothness test decreases. Therefore, we should optimize the margin for efficient sieving.

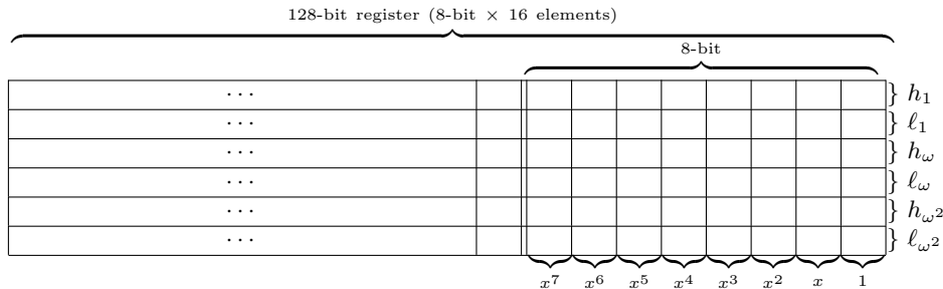
A.1 SIMD implementation

In the lattice sieve, we treat elements in $GF(3^3)[x]$ of the degree at most 6 since parameters B, R , and S are selected as 6, as mentioned in Section 3.2. As described in Section 4.1, we represent $GF(3^3)$ as polynomial basis $GF(3)[\omega]/(\omega^3 - \omega - 1)$ using 6-bit $(h_1, \ell_1, h_\omega, \ell_\omega, h_{\omega^2}, \ell_{\omega^2}) \in GF(2)^6$. Then we can compute operations of $GF(3)$ for coefficients of ω^i for $i = 0, 1, 2$ by logical instructions, as described in [24], so we can also compute operations of $GF(3^3)$ by logical instructions. In our implementation, we represent 16 elements in $GF(3^3)[x]$ of degree at most 7 using 6 registers of 128 bits, as shown in Fig. 2. Then we can compute an addition and a subtraction of $GF(3^3)[x]$ by logical instructions, and a multiplication by x and a division by x by shift instructions. These operations are computed for 16 elements in $GF(3^3)[x]$ with the SIMD.

A.2 Large prime variation

As shown in Table 2, the number of elements in the factor base of degree 6 is over 20 times larger than that of elements in the factor base of degree not larger than 5. Therefore, to compute the lattice sieve efficiently,

⁴ In Section 4.1, we defined special- Q as $Q \in F_R(B)$, not $Q \in \bar{F}_R(B)$ for simplicity.



Note: an element in $GF(3^3) \cong GF(3)[\omega]/(\omega^3 - \omega - 1)$ is represented using 6-bit $(h_1, \ell_1, h_\omega, \ell_\omega, h_{\omega^2}, \ell_{\omega^2}) \in GF(2)^6$.

Fig. 2. 128-bit data structure of 16 elements in $GF(3^3)[x]$ for SIMD implementation

we should omit sieving for the factor base of degree 6, i.e., define the rational sieving factor base $S_R = \bar{F}_R(5)$ and the algebraic sieving factor base $S_A = \bar{F}_A(5)$. This means we deal with the factor base of degree 6 as large primes, which are usually used for efficient computation of the number field sieve. After sieving, $rm + s$ and $r^6x + s^6$ from a candidate (r, s) can be described as a product $(\prod_{\deg \mathfrak{p} \leq 5} \mathfrak{p})L$, where $L \in GF(3^3)[x]$ has no prime factors of degree less than 6. Note that, since \mathfrak{p} can be found by sieving, the degree of L is easily computable. To use large prime variation efficiently, the margin ϵ in the lattice sieve should be large enough. We determined experimentally that $\epsilon = 20$ is an optimized value in our first implementation (stage I at Fig. 1 in Section 4.1).

A.3 Omitting sieving for factor base of degree 1

In our implementation of stage III, we found that the sieving for the factor base of degree 1 is over 1.5 times slower than that for factor bases of other degrees. On the other hand, the sieving for the factor base of degree 1 brings in less information for finding candidates; therefore, we omit the sieving for the factor base of degree 1, i.e., we define the rational sieving factor base $S_R = \bar{F}_R(5) \setminus \bar{F}_R(1)$ and algebraic sieving factor base $S_A = \bar{F}_A(5) \setminus \bar{F}_A(1)$. Through our experiment, we found that an optimized value of the margin ϵ changed to $\epsilon = 21$ from $\epsilon = 20$. This improvement makes our program about 25% faster than that of stage III.

A.4 Smoothness test

To determine that a candidate is truly a B -smooth pair, the smoothness test [16] is frequently conducted. Therefore, an efficient implementation of the smoothness test is necessary to reduce the total computational cost of the collecting relations phase.

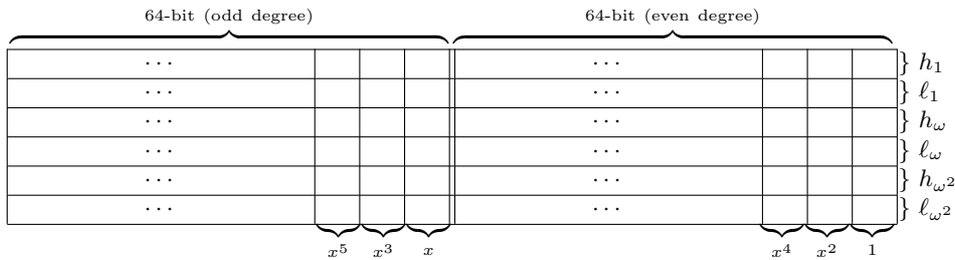
Let U be $rm + s$ or $r^6x + s^6$, then U is tested for B -smoothness by computing

$$U' \prod_{i=\lceil B/2 \rceil}^B (x^{3^i} - x) \bmod U, \quad (14)$$

where U' is a formal derivative of U . The (14) is equal to 0 when U is B -smooth.

The main part of the smoothness test is the computation of cubic residue modulo U , i.e., $A^3 \bmod U$ for $A = \sum A_i x^i \in GF(3^3)[x]$ and $A_i \in GF(3^3)$. In fact, $A^3 \bmod U$ is equal to $\sum A_i^3 (x^{3^i} \bmod U)$; therefore, a cubic residue modulo U can be computed without a reduction by table lookup for computing $x^{3^i} \bmod U$. Moreover, since $x^{3^i} \bmod U = (x^{3(\deg U - 1)} \bmod (Ux^{3(\deg U - 1 - i)})) / x^{3(\deg U - 1 - i)}$ for $i = 0, \dots, \deg U - 1$, the table of $x^{3^i} \bmod U$ can be computed by computing a reduction $x^{3(\deg U - 1)} \bmod U$. This technique makes our implementation of the smoothness test several times faster than that of our initial implementation.

Since input elements of the smoothness test are $rm + s$ and $r^6x + s^6$, whose degree is much larger than 7, they cannot be stored on the SIMD structure for the lattice sieve described in Appendix A.1. Therefore, we introduce a new structure for the smoothness test described in Fig. 3, which uses 6 registers of 128 bits for efficient computing. Note that an element in $GF(3^3)$ in the new structure is represented similar to the SIMD structure for the lattice sieve, i.e., using 6-bit $(h_1, \ell_1, h_\omega, \ell_\omega, h_{\omega^2}, \ell_{\omega^2}) \in GF(2)^6$, for converting efficiently from the SIMD structure to the new structure. As shown in Fig. 3, coefficients of odd degrees of an element in $GF(3^3)[x]$ are stored in the left 64 bits of 128-bit registers, and coefficients of even degrees are stored in the right 64 bits of 128-bit registers. Thus, this data structure can represent an element of $GF(3^3)[x]$, whose degree is at most 127, by using 6 registers of 128 bits. Note that we separate coefficients of odd and even degrees so that it is not necessary to use bit-shift instructions for all 128 bits, which cannot be computed in one instruction.



Note: an element in $GF(3^3) \cong GF(3)[\omega]/(\omega^3 - \omega - 1)$ is represented using 6-bit $(h_1, \ell_1, h_\omega, \ell_\omega, h_{\omega^2}, \ell_{\omega^2}) \in GF(2)^6$.

Fig. 3. 128-bit data structure of element in $GF(3^3)[x]$ for smoothness test

B Scripts for checking solution

We provide the following PARI/GP (<http://pari.math.u-bordeaux.fr/>) scripts for making sure that the equation (13) is true. Note that the target problem defined by the left side of (13) is equivalent to the ECDLP on G_1 described in Section 3.1. Therefore, the following scripts use the additive group operation over supersingular elliptic curve E . The variables `Ell`, `Qpi`, `Qe`, and `Sol` correspond to E , the points Q_π , Q_e , and the solution (13), respectively. The function `ellpow(Ell,Qe,Sol)` means the point $[Sol]Q_e \in E$.

```
Ell=ellinit([0,0,0,-1,1])
pi3c=[1,0,0,1,0,2,1,1,0,1,2,2,2,2,0,1,0,2,1,1,0,0,2,1,1,1,1,1,0,2,2,1,2,2,2,0,1,1,1,2,0,1,2,1,2,1,2,1,2,0,0,1,2,1,1,0,0,1,0,\
0,1,0,1,2,2,2,0,2,2,2,1,2,0,1,2,0,1,2,1,1,1,2,1,0,1,2,1,0,1,2,1,0,1,1,2,0,1,1,0]
e3c=[2,2,0,1,1,0,1,1,2,1,2,2,1,1,0,2,0,1,1,0,1,2,2,2,1,0,2,0,1,1,0,2,1,2,2,2,2,0,1,2,0,2,2,2,2,1,0,2,1,2,2,1,2,0,2,0,1,1,2,1,\
1,2,2,2,1,1,1,0,0,0,1,2,0,2,2,2,1,1,2,1,0,2,1,0,2,0,1,0,0,2,2,2,1,0]
xpi=Mod(Pol(Mod(pi3c,3)),x^97+x^16+2)
xe=Mod(Pol(Mod(e3c,3)),x^97+x^16+2)
ypi=(xpi^3-xpi+1)^((3^97+1)/4)
ye=(xe^3-xe+1)^((3^97+1)/4)
Qpi=[xpi,ypi]
Qe=[xe,ye]
Sol=1752799584850668137730207306198131424550967300
ellpow(Ell,Qe,Sol)==Qpi
```

By using the following commands, we can confirm that the target problem is defined by constants π and e in the manner explained in Section 3.1. Both the following second and third commands must return the output value 1.

```
default(realprecision,50)
subst(Pol(pi3c),x,3)==ffloor(Pi*3^95)+(1*3^1+1*3^0)
subst(Pol(e3c),x,3)==ffloor(exp(1)*3^96)+(1*3^2+2*3^1+0*3^0)
```

We also provide the following scripts corresponding to the polynomials f and m , which were explained in Section 3.2 and selected in the polynomial selection phase of the FFS for our experiments. In our experiments, the finite field $GF(3^{6 \cdot 97})$ is described as $GF(3^3)[x]/(f)$, where $f \in GF(3^3)[x]$ is irreducible of degree 194 and every coefficient of f is in $GF(3)$. Additionally, we describe the finite field $GF(3^3)$ as $GF[\omega]/(\omega^3 - \omega + 1)$. The script `f_vec` means the coefficients of f . For $m \in GF(3)[x]$ satisfying that $H(x, m) \equiv 0 \pmod{f}$ of (1) given in Section 2.2, the script `m_vec` corresponds to the list of the coefficients of m . We can check $H(x, m) \equiv 0 \pmod{f}$ by the script `Mod(m^6+x, f)`. The scripts `etaT_pi_pi` and `etaT_pi_e` mean $\eta_T(Q_\pi, Q_\pi)$ and $\eta_T(Q_\pi, Q_e)$, respectively. We can make sure that our solution is true by checking that the final command returns the output value 1.

```
f_vec=[1,0,1,2,1,1,2,0,1,0,1,2,2,1,0,0,2,0,1,1,1,0,0,2,0,2,1,1,2,0,1,1,1,1,0,0,0,0,2,0,2,1,2,2,2,0,0,1,0,1,1,1,0,0,2,0,2,1,2,2,\
1,0,1,2,1,0,2,2,2,0,0,0,0,2,0,2,1,2,2,1,0,2,1,2,2,0,0,1,0,1,2,2,1,0,2,2,2,1,0,2,2,2,0,0,1,0,0,2,0,1,1,1,0,0,2,1,2,2,2,0,\
0,2,0,2,2,2,0,0,1,0,1,2,1,2,2,1,0,2,2,0,0,1,0,1,2,0,1,1,0,0,2,2,2,0,1,1,1,2,0,1,0,1,2,2,1,0,0,2,0,1,1,1,\
1,0,0,2,0,2,2,2,0,1,1,1]
m_vec=[1,0,0,1,2,2,2,1,2,1,2,0,2,0,1,1,2,1,2,0,2,1,2,1,2,2,2,1,0,2,1,2,0]
f = Pol(Mod(f_vec,3))
m = Pol(Mod(m_vec,3))
Mod(m^6+x,f)
etaT_pi_pi_vec=[\
222,002,100,122,100,010,221,111,001,110,001,222,212,101,112,100,021,101,200,122,221,000,012,000,221,100,120,120,022,001,\
112,011,112,000,121,212,120,121,011,211,111,021,110,100,210,220,101,021,002,211,210,011,022,220,122,112,221,021,011,102,\
202,111,222,101,212,210,121,210,112,120,020,202,200,122,112,211,201,021,021,222,010,200,220,211,012,112,102,010,202,211,\
021,002,122,210,011,011,211,222,000,020,120,002,011,120,011,111,002,022,102,122,221,011,010,000,201,211,022,011,211,212,\
200,221,100,202,102,212,001,002,202,000,111,022,122,120,212,210,021,002,100,120,022,010,022,010,212,112,100,201,210,121,\
010,001,012,222,100,021,010,100,001,021,102,021,222,110,210,211,202,012,112,000,111,000,020,121,001,100,112,011,222,221,\
000,102,000,220,100,210,000,202,012,102,022,200,011,210]
etaT_pi_e_vec=[\
210,122,210,002,221,110,210,120,102,122,020,121,101,222,001,101,121,200,000,212,122,212,012,022,222,111,202,222,210,012,\
201,201,010,010,111,220,100,001,212,001,012,200,020,002,111,020,020,000,011,120,002,001,010,200,101,200,100,220,011,020,\
222,200,211,111,000,221,212,200,222,112,022,100,221,022,200,022,212,111,110,011,120,221,022,020,021,112,020,022,022,201,\
022,121,222,201,101,221,000,001,220,222,111,012,211,201,010,011,221,211,010,220,102,211,000,100,120,122,210,122,000,222,\
222,101,212,111,100,011,111,210,010,110,100,100,011,212,102,122,201,011,102,211,102,012,010,210,010,022,200,202,022,221,\
021,201,011,220,221,100,202,002,112,210,122,110,222,201,211,100,121,112,021,020,010,212,001,022,011,110,101,220,200,121,\
102,222,221,210,201,000,002,200,212,212,002,111,220,122]
```

```
vec2poly(list) ={\
local(i,j,c,tmp,poly);\
poly=0;for(i=1,length(list),c=0;tmp=list[i];\
for(j=0,2,c+=Mod(tmp%10,3)*w^j;tmp\=10);\
poly=poly*x+Mod(c,w^3-w-1));\
return(Mod(poly,f));}
etaT_pi_pi=vec2poly(etaT_pi_pi_vec)
etaT_pi_e=vec2poly(etaT_pi_e_vec)
p151=(3^97+3^49+1)/7

Sol=1752799584850668137730207306198131424550967300
lift(etaT_pi_e^Sol) == etaT_pi_pi
```

C Time-line of experiments and machine environment

Here we provide two tables, Table 8 and 9, which present the time-line of our experiments and machine environment, respectively.

We begin with Table 8. The second to fourth rows correspond to the computation of the lattice sieve in the collecting relations phase, the fifth to sixth rows correspond to the computation of the parallel Lanczos method in the linear algebra phase, and the seventh to tenth rows correspond to the computation of the rationalization and that of the special- Q descent in the individual logarithm step. The column “days” means our working time including the loss-time of some programming errors, updating our codes, and power outages. As shown in Table 8, we required 118 days for computing the lattice sieve, 90 days for computing the parallel Lanczos method, and 15 days for computing the rationalization and the special- Q descent.

Next, specifications of our machine environment are listed in Table 9. We used PCs (a)-(c) and (e)-(g) for the collecting relations phase, (e) for computing the filtering, (h) for computing the parallel Lanczos method, and (a)-(d) for the individual logarithm phase. Note that #cores means the number of CPU cores in a node, not a CPU, and #nodes means the number of PCs we used. For example, (h) in Table 9 means that we used 22 nodes, each of which had 12 CPU cores of Xeon X5650 and 12-GB RAM. Since a Xeon X5650 has 6 CPU cores, a node of (h) has 2 CPUs; therefore, each node had 12 CPU cores in total, as shown in the table.

Table 8. Time-line of our experiments

date	days	remarks
2011.5.14	-	The computation of the lattice sieve started. (47 nodes, 212 CPU cores)
2011.6.15	32	Minimal required number of relations was collected. (834202 special- Q 's)
2011.9.9	118	The computation of the lattice sieve finished. (2500000 special- Q 's)
2012.1.16	-	The computation of the parallel Lanczos method started. (21 nodes, 252 CPU cores)
2012.4.14	90	The computation of the parallel Lanczos method finished.
2012.2.3	-	The computation of the rationalization started. (20 nodes, 168 CPU cores)
2012.2.17	14	The computation of the rationalization finished.
2012.2.27	14	The computation of the special- Q descent started. (20 nodes, 168 CPU cores)
2012.2.28	15	The computation of the special- Q descent finished.
2012.4.24		The obtained logarithm was verified and our experiments completely finished.

Table 9. Machine environment for our experiments

	CPU	memory	#cores	#nodes
(a)	Core2 quad Q9450 2.66GHz	4GB	4	10
(b)	Core2 quad Q9650 3.0GHz	4GB	4	10
(c)	Xeon X3460 2.8GHz	8GB	4	10
(d)	Xeon E3-1275 3.4GHz	16GB	4	12
(e)	Xeon E5440 2.83GHz	16GB	8	5
(f)	Xeon X5355 2.66GHz	16GB	8	1
(g)	Xeon L5420 2.50GHz	4GB	4	11
(h)	Xeon X5650 2.67GHz	12GB	12	22